

Coronavirus Disease 2019 (COVID-19) Treatment Guidelines

Credit NIAID-RML

Downloaded from <u>https://www.covid19treatmentguidelines.nih.gov/</u> on 2/5/2024 Visit <u>https://www.covid19treatmentguidelines.nih.gov/</u> to access the most up-to-date guideline.



How to Cite the COVID-19 Treatment Guidelines:

COVID-19 Treatment Guidelines Panel. Coronavirus Disease 2019 (COVID-19) Treatment Guidelines. National Institutes of Health. Available at <u>https://www.covid19treatmentguidelines.nih.gov/</u>. Accessed [insert date].

The COVID-19 Treatment Guidelines Panel regularly updates the recommendations in these guidelines as new information on the management of COVID-19 becomes available. The most recent version of the guidelines can be found on the COVID-19 Treatment Guidelines website (<u>https://www.covid19treatmentguidelines.nih.gov/</u>).

Table of Contents

What's New in the Guidelines
Guidelines Development
Overview
Overview of COVID-19.9Testing for SARS-CoV-2 Infection.14Prevention of SARS-CoV-2 Infection18Clinical Spectrum of SARS-CoV-2 Infection.22Prioritization of Anti-SARS-CoV-2 Therapies for the Treatment of COVID-19 in Nonhospitalized Patients When There Are Logistical Constraints.31
Clinical Management of Adults
Clinical Management of Adults Summary
Clinical Management of Children
Clinical Management of Children Summary
Critical Care for Adults
Care of Critically III Adults With COVID-19 (Summary Recommendations)
Critical Care for Children
Introduction to Critical Care Management of Children With COVID-19
Antiviral Agents, Including Antibody Products
Antiviral Agents, Including Antibody Products (Summary Recommendations)
COVID-19 Treatment Guidelines 2

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Remdesivir	170
Table 4a. Remdesivir: Selected Clinical Trial Data	174
Ritonavir-Boosted Nirmatrelvir (Paxlovid)	182
Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications	189
Molnupiravir	196
Anti-SARS-CoV-2 Monoclonal Antibodies	202
Table 4b. Anti-SARS-CoV-2 Monoclonal Antibodies: Selected Clinical Trial Data	204
COVID-19 Convalescent Plasma	208
Table 4c. COVID-19 Convalescent Plasma: Selected Clinical Trial Data	214
Interferons	223
Table 4d. Interferons: Selected Clinical Trial Data	226
Table 4e. Characteristics of Antiviral Agents, Including Antibody Products	233
Immunomodulators	
Immunomodulators (Summary Recommendations)	238
Systemic Corticosteroids	239
Table 5a. Systemic Corticosteroids: Selected Clinical Trial Data	245
Inhaled Corticosteroids	255
Table 5b. Inhaled Corticosteroids: Selected Clinical Trial Data	258
Interleukin-6 Inhibitors	265
Table 5c. Interleukin-6 Inhibitors: Selected Clinical Trial Data	270
Janus Kinase Inhibitors	278
Table 5d: Janus Kinase Inhibitors: Selected Clinical Trial Data	282
Abatacept	289
Inflixamab	292
Interleukin-1 Inhibitors	295
Vilobelimab	
Table 5e. Characteristics of Immunomodulators	305
Antithrombotic Therapy in Patients With COVID-19	315
Table 6a. Anticoagulant Therapy: Selected Clinical Trial Data	330
Table 6b. Antiplatelet Therapy: Selected Clinical Trial Data	344
Miscellaneous Drugs	
Miscellaneous Drugs (Summary Recommendations)	349
Fluvoxamine	
Table 7a. Fluvoxamine: Selected Clinical Trial Data	353
Intravenous Immunoglobin	361
Ivermectin	
Table 7b. Ivermectin: Selected Clinical Trial Data	367
Metformin	382

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Table 7c. Metformin Selected Clinical Trial Data
Table 7d. Characteristics of Miscellaneous Drugs
Supplements
Supplements (Summary Recommendations)
Vitamin C
Vitamin D
Zinc
Considerations for Using Concomitant Medications in Patients With COVID-19
Special Populations
Special Considerations in People Who Are Immunocompromised
Special Considerations in Adults and Children With Cancer
Special Considerations in Solid Organ Transplant, Hematopoietic Cell Transplant,
and Cellular Immunotherapy Candidates, Donors, and Recipients
Special Considerations During Pregnancy and After Delivery
Pregnancy, Lactation, and COVID-19 Therapeutics
Influenza and COVID-19 450
Special Considerations in People With HIV 456
Appendix A, Table 1. COVID-19 Treatment Guidelines Panel Members
Appendix A, Table 2. Panel on COVID-19 Treatment Guidelines Financial Disclosure for Companies Related to COVID-19 Treatment or Diagnostics

What's New in the Guidelines

Last Updated: December 20, 2023

The *Coronavirus Disease 2019 (COVID-19) Treatment Guidelines* is published in an electronic format that can be updated in step with the rapid pace and growing volume of information regarding the treatment of COVID-19.

The COVID-19 Treatment Guidelines Panel (the Panel) is committed to updating this document to ensure that health care providers, patients, and policy experts have the most recent information regarding the optimal management of COVID-19 (see the <u>Panel Roster</u> for a list of Panel members).

New Guidelines sections and recommendations and updates to existing Guidelines sections are developed by working groups of Panel members. All recommendations included in the Guidelines are endorsed by a majority of Panel members (see <u>Guidelines Development</u> for additional details on the development process).

Major revisions to the Guidelines within the past month are as follows:

December 20, 2023

Future of the Guidelines

In response to the rapidly evolving COVID-19 pandemic, the National Institutes of Health assembled a panel of experts to provide practical recommendations for health care providers and issued the first version of the COVID-19 Treatment Guidelines on April 21, 2020. For close to 4 years, the Panel has critically reviewed the growing body of research data on COVID-19 and used that information to develop and revise their recommendations for treating patients with this disease.

The federal COVID-19 Public Health Emergency ended in May 2023. The last update of the Guidelines will be published in early 2024. The Guidelines website will remain available for several months and will provide a downloadable version of the final publication of the Guidelines.

Key Updates to the Guidelines

Fluvoxamine

Six randomized, placebo-controlled trials evaluated the use of fluvoxamine in nonhospitalized adults with COVID-19. Most of these studies showed that, compared to placebo, fluvoxamine did not improve clinical outcomes in these patients. Therefore, the Panel **recommends against** the use of **fluvoxamine** for the treatment of COVID-19 in nonhospitalized patients (**AIIa**).

Vitamin C

Two large, harmonized, randomized, multinational trials evaluated the use of intravenous vitamin C in hospitalized patients with COVID-19. The trials included patients who were critically ill and patients who were not critically ill. Enrollment in both studies was terminated because of futility and a potential for harm. After reviewing the results of these studies, the Panel **recommends against** the use of vitamin C for the treatment of COVID-19 in hospitalized patients (**AIIa**).

Minor Updates to the Guidelines

Several other sections of the Guidelines were updated to remove outdated information and add new references.

COVID-19 Treatment Guidelines

Guidelines Development

Last Updated: October 10, 2023

The COVID-19 Treatment Guidelines were developed to provide clinicians with guidance on caring for patients with COVID-19. Because clinical information about the optimal management of COVID-19 is evolving quickly, these Guidelines are updated frequently to reflect newly published data and other authoritative information.

Panel Composition

Members of the COVID-19 Treatment Guidelines Panel (the Panel) are appointed by the Panel co-chairs based on their clinical experience and expertise in patient management, translational and clinical science, and/or the development of treatment guidelines. Panel members include representatives from federal agencies, health care organizations, academic institutions, and professional societies. Federal agencies and professional societies represented on the Panel include:

- American Association of Critical-Care Nurses
- American Association for Respiratory Care
- American College of Chest Physicians
- American College of Emergency Physicians
- American College of Obstetricians and Gynecologists
- American Society of Hematology
- American Thoracic Society
- Biomedical Advanced Research and Development Authority
- Centers for Disease Control and Prevention
- Department of Defense
- Department of Veterans Affairs
- Food and Drug Administration
- Infectious Diseases Society of America
- National Institutes of Health
- Pediatric Infectious Diseases Society
- Society of Critical Care Medicine
- Society of Infectious Diseases Pharmacists

The inclusion of representatives from professional societies does not imply that these societies have endorsed all elements of the Guidelines.

The names and affiliations of the Panel members, ex officio members, consultants, and members of the Guidelines support team are provided in <u>Appendix A, Table 1</u>. Financial disclosures for the Panel members can be found in <u>Appendix A, Table 2</u>.

Development of the Guidelines

Each section of the Guidelines is developed by a working group of Panel members with expertise in the

area addressed in the section. Each working group is responsible for identifying relevant information and published scientific literature and for conducting a systematic, comprehensive review of that information and literature. The working groups propose updates to the Guidelines based on the latest published research findings and clinical information.

New Guidelines sections and recommendations are reviewed and voted on by the voting members of the Panel. To be included in the Guidelines, a recommendation statement must be endorsed by a majority of voting members; this applies to recommendations for and against treatments and cases where there is insufficient evidence to recommend either for or against treatments. Updates to existing sections that do not affect the rated recommendations are approved by Panel co-chairs without a Panel vote. Panel members are required to keep all Panel deliberations and unpublished data that are evaluated during the development of the Guidelines confidential.

Method of Synthesizing Data and Formulating Recommendations

The working groups critically review and synthesize the available data to develop recommendations. During this process, the Panel evaluates the data, including the source of the data, the type of study (e.g., randomized controlled trial, prospective or retrospective cohort study, case series, in vitro study), the quality and suitability of the methods, the number of participants, and the effect sizes observed.

The recommendations in these Guidelines are based on scientific evidence and expert opinion. Each recommendation includes 2 ratings: an uppercase letter (**A**, **B**, or **C**) that indicates the strength of the recommendation and a Roman numeral with or without a lowercase letter (**I**, **IIa**, **IIb**, or **III**) that indicates the quality of the evidence that supports the recommendation (see Table 1).

The ratings for the quality of the evidence reflect both the likelihood of bias in the treatment effect estimate and the precision of the estimate. A rating of **I** corresponds to a low likelihood of bias and a high precision, a rating of **IIa** (for randomized trials) or **IIb** (for observational studies) corresponds to a moderate likelihood of bias and a moderate or high precision, and a rating of **III** corresponds to a high likelihood of bias (for any type of study).

Strength of Recommendation	Evidence for Recommendation			
 A: Strong recommendation for the statement B: Moderate recommendation for the statement C: Weak recommendation for the statement 	I: <i>High quality of evidence:</i> 1 or more randomized trials without major limitations, ^a well-powered subgroup analyses of such trials, or meta-analyses without major limitations			
	IIa: Moderate quality of evidence: Randomized trials and subgroup analyses of randomized trials that do not meet the criteria for a I rating			
	IIb: <i>Moderate quality of evidence:</i> Observational studies without major limitations ^b			
	III: Expert opinion			
^a The rating may be lower than I in cases where trials have produced conflicting results.				
^b This category also includes meta-analyses of observational studies.				

Table 1. Recommendation Rating Scheme

To develop the recommendations in these Guidelines, the Panel uses data from the rapidly growing body of research on COVID-19. The Panel also relies heavily on experience with other diseases, supplemented with the members' clinical experience with COVID-19.

In general, the recommendations in these Guidelines fall into the following categories:

- The Panel recommends using [blank] for the treatment of COVID-19 (rating). Recommendations in this category are based on evidence that the potential benefits of using this intervention outweigh the potential risks.
- There is insufficient evidence for the Panel to recommend either for or against the use of [blank] for the treatment of COVID-19 (no rating). This statement is used when there are currently not enough data to support a recommendation, or when the available data are conflicting.
- The Panel recommends against the use of [blank] for the treatment of COVID-19, except in a clinical trial (rating). This recommendation is used in cases where the available data have shown no benefit of using this intervention for the treatment of COVID-19 and/or the intervention has demonstrated safety concerns. More results from clinical trials are needed to further define the role of these interventions in treating COVID-19.
- The Panel recommends against the use of [blank] for the treatment of COVID-19 (rating). This recommendation is used in cases where the available data show that there is no benefit of using this intervention to treat COVID-19 and/or the safety concerns for the intervention outweigh any potential benefits.

Evolving Knowledge on Treatments for COVID-19

Several agents (i.e., baricitinib, ritonavir-boosted nirmatrelvir [Paxlovid], remdesivir, tocilizumab) are currently approved by the Food and Drug Administration for the treatment of COVID-19, and a number of other agents have received Emergency Use Authorizations. An array of drugs that are approved for other indications and multiple investigational agents are being studied for the treatment of COVID-19 in clinical trials around the globe. Information about these trials can be found at <u>ClinicalTrials.gov</u>. In addition, providers can access and prescribe investigational drugs or agents that are approved or licensed for other indications through various mechanisms, including Emergency Investigational New Drug applications, compassionate use or expanded access programs with drug manufacturers, and/or off-label use.

Whenever possible, the Panel recommends that promising, unapproved, or unlicensed treatments for COVID-19 be studied in well-designed, controlled clinical trials. This recommendation also applies to drugs that have been approved or licensed for indications other than the treatment of COVID-19. The Panel recognizes the critical importance of clinical research in generating evidence to address unanswered questions regarding the safety and efficacy of potential treatments for COVID-19. However, the Panel also realizes that many patients and providers who cannot access these potential treatments via clinical trials still seek guidance about whether to use them.

New data on the treatment of COVID-19 are emerging rapidly. Some of these data are being published in peer-reviewed journals, and some can be found in manuscripts that have not yet been peer reviewed or in press releases. The Panel continuously reviews the available data and assesses their scientific rigor and validity. These sources of data and the clinical experiences of the Panel members are used to determine whether new recommendations or changes to the current recommendations are warranted.

Finally, it is important to stress that the rated treatment recommendations in these Guidelines should not be considered mandates. What to do or not to do for an individual patient is ultimately decided by the patient and their provider.

Overview of COVID-19

Last Updated: December 20, 2023

Epidemiology

Individuals of all ages are at risk of SARS-CoV-2 infection. However, the probability of severe COVID-19 is higher in people aged \geq 65 years, those living in nursing homes or long-term care facilities, those who are not vaccinated against COVID-19 or who have poor responses to COVID-19 vaccines, and those with certain chronic medical conditions. Data on comorbid health conditions among patients with COVID-19 indicate that patients with cardiovascular disease, chronic kidney disease, chronic obstructive pulmonary disease, diabetes with complications, neurocognitive disorders, and obesity are at increased risk of severe COVID-19. The risk appears to be higher in patients with multiple comorbid conditions. Other conditions that may lead to a high risk of severe COVID-19 include cancer, cystic fibrosis, immunocompromising conditions, liver disease (especially in patients with cirrhosis), pregnancy, and sickle cell disease. Transplant recipients and people who are taking immunosuppressive medications are also at high risk of severe COVID-19.¹ See <u>Clinical Spectrum of SARS-CoV-2 Infection</u> for a description of the clinical manifestations of SARS-CoV-2 infection and a discussion of the spectrum of disease.

Although COVID-19 vaccination does not eliminate the risk of SARS-CoV-2 infection, vaccination does significantly reduce the risk of COVID-19–related morbidity and mortality, particularly in individuals who are at high risk of progressing to severe disease.^{2,3}

Racial and Ethnic Minorities and Other Marginalized Groups

Communities that have been historically marginalized or made socially vulnerable due to a lack of access to health care or an inability to socially isolate are at increased risk of SARS-CoV-2 acquisition, COVID-19–related hospitalization, and death. These communities include racial and ethnic minorities, essential non-health care workers, and some people with disabilities.

Key Considerations

- The COVID-19 Treatment Guidelines Panel recommends that health care providers, health care systems, and payers ensure equitable access to high-quality care and treatment for all patients, regardless of race, ethnic identity, or other minoritized identity or social status (AIII). "Minoritized" refers to social groups that have been deprived of power and status by the dominant culture in society and encompasses not just racial identities but other identities as well, including gender identity and sexual orientation.⁴
- Promoting equitable care for these groups must include considering the full range of medical, demographic, and social factors that may negatively impact health outcomes.
- Clinicians should be aware that pulse oximeters may not accurately detect hypoxemia in people with darker skin pigmentation.^{5,6} This may delay treatment and lead to worse clinical outcomes in patients with COVID-19.⁷ See <u>Clinical Spectrum of SARS-CoV-2 Infection</u> for more information.
- Supporting equitable access to high-quality care and treatment for all patients is now an imperative for all health care organizations accredited by the Joint Commission, as well as a priority for the Centers for Disease Control and Prevention (CDC) and other public health agencies.

COVID-19-Related Health Outcomes

Historical structural inequities significantly contribute to the health disparities experienced by racial and ethnic minority groups (e.g., Black/African American people, Hispanic people, American Indian/ Alaska Native people).⁸ Some data have highlighted that select racial and ethnic minority groups experience higher rates of COVID-19, subsequent hospitalization, and death in relation to their share of the total U.S. population. Black/African American people, Hispanic people, and American Indian/Alaska Native people also experience rates of hospitalization that are more than 2 times higher and rates of COVID-19–related death that are approximately 2 times higher than those experienced by White people. The largest disparities were observed among American Indian/Alaska Native people, who experienced a rate of hospitalization almost 3 times higher and a rate of death 2.1 times higher than White people.⁹

The increased risk of severe COVID-19 among racial and ethnic minority groups may be partly attributed to higher rates of comorbid conditions in these populations (e.g., cardiovascular disease, diabetes, chronic kidney disease, hypertension, obesity, pulmonary disease).⁹

Disparities in Access to Care

Members of racial and ethnic minority groups have an increased risk of exposure to COVID-19 and decreased access to care. Large-scale mobility data reveals that people living in lower-income communities were less able to physically isolate during COVID-19 emergency declarations,¹⁰ as members of these communities were frequently unable to work from home.¹¹ A 2020 study evaluating access to health care resources in New York City found that in areas of the city where the majority of the population was Black/African American and Hispanic, there were higher COVID-19 positivity rates and fewer licensed hospital beds and intensive care unit beds than in areas where the majority of the population was White.¹²

Disparities in Access to COVID-19 Treatments

Data from 41 U.S. health care systems reveal racial and ethnic disparities in the use of anti-SARS-CoV-2 monoclonal antibodies (mAbs) for the treatment of COVID-19.¹³ Black/African American patients, Asian patients, and patients of other races were, respectively, 22.4%, 48.3%, and 46.5% less likely to receive anti-SARS-CoV-2 mAbs for the treatment of COVID-19 than White patients.^{13,14} Disparities have also been observed in the dispensing rates for ritonavir-boosted nirmatrelvir (Paxlovid) and molnupiravir. One study reported that between April and July 2022, Black/African American patients were prescribed ritonavir-boosted nirmatrelvir 35.8% less often than White patients, and Hispanic patients were prescribed this drug 29.9% less often than White patients.¹⁵ Despite a greater number of dispensing sites in neighborhoods with a higher social vulnerability, oral antivirals were prescribed at a lower rate for patients with COVID-19 who were living in these areas than in those with a lesser degree of social vulnerability.¹⁶ These disparities are not limited to outpatient settings. One retrospective cohort study of veterans hospitalized with COVID-19 reported that Black veterans had lower odds of receiving COVID-19–specific treatments, including systemic steroids, remdesivir, and immunomodulators, than White veterans.¹⁷

Other Marginalized Groups

Other marginalized groups also experience worse outcomes for COVID-19. Hospitalization rates for COVID-19 among Medicare beneficiaries who were eligible for disability were approximately 50% higher than those among people who were eligible for Medicare based on age alone, and this discrepancy disproportionately affected Black/African American people, Hispanic people, and American Indian/Alaska Native people.¹⁸

Migrants, refugees, and essential non-health care workers (e.g., food supply, food service, public transportation, and agricultural workers) also have disproportionately high rates of COVID-19 cases and deaths. These high rates can be attributed to overcrowding, an inability to physically isolate, and inadequate access to health care.¹⁹⁻²¹

Given the pervasiveness of disparities in access to care and provision of treatment, it is imperative for clinicians, working with others on the patient care team, to assess the social factors that contribute to access and quality gaps and to strive to provide equitable treatment to all patients. These issues have been identified as a strategic priority by the <u>Joint Commission</u> and the <u>CDC</u>.

SARS-CoV-2 Variants

Like other RNA viruses, SARS-CoV-2 is constantly evolving through random mutations. New mutations can potentially increase or decrease infectiousness and virulence. In addition, mutations can increase the virus' ability to evade adaptive immune responses from previous SARS-CoV-2 infections or vaccination. This viral evolution may increase the risk of reinfection or decrease the efficacy of vaccines.²² There is evidence that some SARS-CoV-2 variants have reduced susceptibility to plasma from people who were previously infected or immunized, as well as to anti-SARS-CoV-2 mAbs.²³⁻²⁵

Since December 2020, the World Health Organization has assigned Greek letter designations to several identified variants. A SARS-CoV-2 variant designated as a variant of concern displays certain characteristics, such as increased transmissibility or virulence. In addition, vaccines and therapeutics may have decreased effectiveness against variants of concern, and the mutations found in these variants may interfere with the targets of diagnostic tests. The variant of interest designation has been used for important variants that are not fully characterized; however, organizations do not use the same variant designations, and they may define their variant designations differently.^{26,27}

In September 2021, the CDC added a new designation for variants: <u>variants being monitored</u>. The data on these variants indicate a potential or clear impact on approved or authorized medical countermeasures, or these variants are associated with cases of more severe disease or increased transmission rates. However, these variants are either no longer detected or are circulating at very low levels in the United States; therefore, they do not pose a significant and imminent risk to public health in the United States.

The Omicron variant was designated as a variant of concern in November 2021 and rapidly became the dominant variant across the globe. The Omicron subvariants BA.1, BA.1.1, and BA.2 emerged in early to mid-2022, followed by the subvariants BA.4, BA.5, BQ.1, BQ.1.1, XBB, EG.5, HV.1, and FL.1.5.1. The newer Omicron subvariants are generally more transmissible than previous variants and are not susceptible to any of the anti-SARS-CoV-2 mAbs that were previously authorized for the treatment and prevention of COVID-19.^{24,25,28,29}

Data on the emergence, transmission, and clinical relevance of these new variants are rapidly evolving; this is especially true for research on how variants might affect transmission rates, disease progression, vaccine development, and the efficacy of current therapeutics. Because the research on variants is moving quickly and the classification of the different variants may change over time, websites such as the CDC's <u>COVID Data Tracker</u>, <u>CoVariants.org</u>, and the World Health Organization's <u>Tracking</u> <u>SARS-CoV-2 Variants</u> provide regular updates on data for SARS-CoV-2 variants.

References

1. Centers for Disease Control and Prevention. Underlying medical conditions associated with higher risk for severe COVID-19: information for healthcare professionals. 2023. Available at: <u>https://www.cdc.gov/</u>

coronavirus/2019-ncov/hcp/clinical-care/underlyingconditions.html. Accessed November 13, 2023.

- 2. Link-Gelles R, Weber ZA, Reese SE, et al. Estimates of bivalent mRNA vaccine durability in preventing COVID-19-associated hospitalization and critical illness among adults with and without immunocompromising conditions—VISION Network, September 2022–April 2023. *MMWR Morb Mortal Wkly Rep.* 2023;72(21):579-588. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37227984</u>.
- 3. Our World in Data. United States: COVID-19 weekly death rate by vaccination status. 2022. Available at: <u>https://ourworldindata.org/grapher/united-states-rates-of-covid-19-deaths-by-vaccination-status-by-vaccine?country=12-17~12~12</u>. Accessed November 13, 2023.
- 4. American Medical Association. Advancing health equity: a guide to language, narrative and concepts. 2021. Available at: <u>https://ama-assn.org/equity-guide</u>.
- Valbuena VSM, Seelye S, Sjoding MW, et al. Racial bias and reproducibility in pulse oximetry among medical and surgical inpatients in general care in the Veterans Health Administration 2013–19: multicenter, retrospective cohort study. *BMJ*. 2022;378:e069775. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35793817</u>.
- Chesley CF, Lane-Fall MB, Panchanadam V, et al. Racial disparities in occult hypoxemia and clinically based mitigation strategies to apply in advance of technological advancements. *Respir Care*. 2022;67(12):1499-1507. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35679133</u>.
- Fawzy A, Wu TD, Wang K, et al. Racial and ethnic discrepancy in pulse oximetry and delayed identification of treatment eligibility among patients with COVID-19. *JAMA Intern Med.* 2022;182(7):730-738. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35639368</u>.
- 8. National Academies of Sciences, Engineering, and Medicine. *Communities in Action: Pathways to Health Equity*. 2017. National Academies Press; 2017. Available at: https://www.ncbi.nlm.nih.gov/pubmed/28418632.
- 9. Mackey K, Ayers CK, Kondo KK, et al. Racial and ethnic disparities in COVID-19-related infections, hospitalizations, and deaths: a systematic review. *Ann Intern Med.* 2021;174(3):362-373. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33253040.
- Weill JA, Stigler M, Deschenes O, Springborn MR. Social distancing responses to COVID-19 emergency declarations strongly differentiated by income. *Proc Natl Acad Sci U S A*. 2020;117(33):19658-19660. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32727905</u>.
- 11. Economic Policy Institute. Not everybody can work from home: Black and Hispanic workers are much less likely to be able to telework. 2020. Available at: <u>https://www.epi.org/blog/black-and-hispanic-workers-are-much-less-likely-to-be-able-to-work-from-home</u>. Accessed November 13, 2023.
- 12. Douglas JA, Subica AM. COVID-19 treatment resource disparities and social disadvantage in New York City. *Prev Med.* 2020;141:106282. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33035550</u>.
- Wiltz JL, Feehan AK, Molinari NM, et al. Racial and ethnic disparities in receipt of medications for treatment of COVID-19—United States, March 2020–August 2021. *MMWR Morb Mortal Wkly Rep.* 2022;71(3):96-102. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35051133</u>.
- Wu EL, Kumar RN, Moore WJ, et al. Disparities in COVID-19 monoclonal antibody delivery: a retrospective cohort study. *J Gen Intern Med*. 2022;37(10):2505-2513. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35469360</u>.
- Boehmer TK, Koumans EH, Skillen EL, et al. Racial and ethnic disparities in outpatient treatment of COVID-19—United States, January–July 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(43):1359-1365. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36301738</u>.
- 16. Gold JAW, Kelleher J, Magid J, et al. Dispensing of oral antiviral drugs for treatment of COVID-19 by ZIP code-level social vulnerability—United States, December 23, 2021–May 21, 2022. MMWR Morb Mortal Wkly Rep. 2022;71(25):825-829. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35737571.
- 17. Castro AD, Mayr FB, Talisa VB, et al. Variation in clinical treatment and outcomes by race among US veterans hospitalized with COVID-19. *JAMA Netw Open*. 2022;5(10):e2238507. Available at: <u>https://www.</u>

ncbi.nlm.nih.gov/pubmed/36282499.

- Yuan Y, Thierry JM, Bull-Otterson L, et al. COVID-19 cases and hospitalizations among Medicare beneficiaries with and without disabilities—United States, January 1, 2020–November 20, 2021. MMWR Morb Mortal Wkly Rep. 2022;71(24):791-796. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35709015.
- 19. Hayward SE, Deal A, Cheng C, et al. Clinical outcomes and risk factors for COVID-19 among migrant populations in high-income countries: a systematic review. *J Migr Health*. 2021;3:100041. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33903857.
- Lewnard JA, Mora AM, Nkwocha O, et al. Prevalence and clinical profile of severe acute respiratory syndrome coronavirus 2 infection among farmworkers, California, USA, June–November 2020. *Emerg Infect Dis.* 2021;27(5):1330-1342. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33657340</u>.
- Heinzerling A, Vergara XP, Gebreegziabher E, et al. COVID-19 outbreaks and mortality among public transportation workers—California, January 2020–May 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(33):1052-1056. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35980867</u>.
- 22. Walensky RP, Walke HT, Fauci AS. SARS-CoV-2 variants of concern in the United States—challenges and opportunities. *JAMA*. 2021;325(11):1037-1038. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33595644.
- 23. Wang P, Nair MS, Liu L, et al. Antibody resistance of SARS-CoV-2 variants B.1.351 and B.1.1.7. *Nature*. 2021;593(7857):130-135. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33684923</u>.
- 24. Cameroni E, Bowen JE, Rosen LE, et al. Broadly neutralizing antibodies overcome SARS-CoV-2 Omicron antigenic shift. *Nature*. 2022;602(7898):664-670. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35016195.
- 25. Liu L, Iketani S, Guo Y, et al. Striking antibody evasion manifested by the Omicron variant of SARS-CoV-2. *Nature*. 2022;602(7898):676-681. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35016198</u>.
- 26. World Health Organization. Tracking SARS-CoV-2 variants. 2023. Available at: <u>https://www.who.int/en/activities/tracking-SARS-CoV-2-variants</u>. Accessed November 13, 2023.
- 27. Centers for Disease Control and Prevention. SARS-CoV-2 variant classifications and definitions. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/variants/variant-classifications.html</u>. Accessed November 13, 2023.
- 28. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Evusheld (tixagevimab co-packaged with cilgavimab). 2023. Available at: <u>https://www.fda.gov/media/154701/download</u>.
- 29. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for bebtelovimab. 2022. Available at: <u>https://www.fda.gov/media/156152/download</u>.

Testing for SARS-CoV-2 Infection

Last Updated: December 20, 2023

Summary of Testing for SARS-CoV-2 Infection

The COVID-19 Treatment Guidelines Panel (the Panel) defers to the Centers for Disease Control and Prevention (CDC) for recommendations on diagnostic testing for SARS-CoV-2 infection. The Panel also defers to the CDC for recommendations on the use of testing for screening purposes, such as for screening among people who are asymptomatic but have had recent known or suspected exposure to SARS-CoV-2. Some key CDC recommendations include:

- For diagnosing current SARS-CoV-2 infection, the CDC recommends using either a nucleic acid amplification test (NAAT) or an antigen test and using a specimen from the upper respiratory tract (e.g., nasal, nasopharyngeal).
- There may be a window period of up to 5 days after exposure before viral antigens or nucleic acids can be detected.
- NAATs are the most sensitive tests for detecting current SARS-CoV-2 infection. Because antigen tests are less sensitive than NAATs, the Food and Drug Administration recommends repeating antigen tests that produce negative results in certain circumstances, such as when clinical suspicion of COVID-19 is high in people who are symptomatic or when people who are asymptomatic have had known or suspected exposure to SARS-CoV-2.
- Antibody tests should not be used to diagnose current SARS-CoV-2 infection.
- Currently, antibody tests are not recommended for assessing SARS-CoV-2 immunity following COVID-19 vaccination or for assessing the need for vaccination in a person who is unvaccinated.

Diagnostic Testing for SARS-CoV-2 Infection

For diagnosing current SARS-CoV-2 infection, the Centers for Disease Control and Prevention (CDC) recommends using either a nucleic acid amplification test (NAAT) or an antigen test.¹ Testing may also be used for screening and to determine the length of a patient's isolation period.² There may be a window period of up to 5 days after exposure before viral antigens or nucleic acids can be detected.

A number of diagnostic tests for SARS-CoV-2 infection (e.g., NAATs, antigen tests) have received Food and Drug Administration (FDA) Emergency Use Authorizations (EUAs) for use in laboratories and points of care (e.g., physician offices, pharmacies, long-term care facilities, school clinics) and for self-administered testing.³ An influenza and SARS-CoV-2 multiplex NAAT that can simultaneously detect and differentiate between influenza A, influenza B, and SARS-CoV-2 also received an EUA from the FDA.⁴ The FDA also granted authorization to market the first over-the-counter, at-home, molecular NAAT (i.e., Cue COVID-19) and antigen test (i.e., Flowflex COVID-19) for use in people with symptomatic COVID-19.

For diagnosing current SARS-CoV-2 infection, the CDC recommends using a specimen from the upper respiratory tract (e.g., nasal, nasopharyngeal).⁵ Testing lower respiratory tract specimens is also an option in certain circumstances (e.g., in those receiving mechanical ventilation). For details about collecting and handling specimens for COVID-19 testing, please refer to the CDC's recommendations.

Antigen Testing for SARS-CoV-2 Infection

Antigen-based diagnostic tests are widely used at home, at the point of care, and in the laboratory because of their low cost, rapid turnaround time, and availability. Antigen tests and laboratory-based NAATs have similar high specificity. False positive test results can occur with antigen tests, although they are unlikely when the tests are used correctly.⁶ The likelihood of a false positive antigen test result is higher when the expected probability of SARS-CoV-2 infection is low. Because antigen tests are less sensitive than NAATs, the FDA recommends repeating antigen tests that produce negative results in certain

circumstances, such as when clinical suspicion of COVID-19 is high in people who are symptomatic or when people who are asymptomatic have had known or suspected exposure to SARS-CoV-2.

Nucleic Acid Amplification Testing for SARS-CoV-2 Infection

NAATs, such as reverse transcription polymerase chain reaction–based diagnostic tests, which detect viral nucleic acids,⁷ are the most sensitive tests for detecting current SARS-CoV-2 infection.

Diagnostically, some NAATs may produce false negative results if a mutation occurs in the part of the virus's genome that is assessed by that test.⁸ The FDA monitors the potential effects of SARS-CoV-2 genetic variations on NAAT results and issues updates when specific variations could affect the performance of NAATs that have received EUAs.⁹ A single negative test result does not exclude the possibility of SARS-CoV-2 infection in people who have a high likelihood of infection based on their exposure history or clinical presentation.¹⁰

Reinfection

Reinfection has been reported in people after an initial diagnosis of SARS-CoV-2 infection. Because reinfection can be difficult to distinguish from persistent shedding (i.e., positive NAAT results persisting for weeks or months), the CDC recommends using an antigen test instead of a NAAT in patients who have symptoms compatible with SARS-CoV-2 infection who are within 90 days of recovering from a previous SARS-CoV-2 infection. Because intermittent detection of viral RNA can occur, a negative result on an initial NAAT followed by a positive result on a subsequent test does not necessarily mean a person has been reinfected.¹¹ When the results for an initial and subsequent test are positive, comparative viral sequence data from both tests are needed to distinguish between the persistent presence of viral fragments and reinfection. In the absence of viral sequence data, the cycle threshold (Ct) value from a positive NAAT result may provide information about whether a newly detected infection is related to the persistence of viral fragments or to reinfection. The Ct value is the number of PCR cycles at which the nucleic acid target in the sample becomes detectable. In general, the Ct value is inversely related to the SARS-CoV-2 viral load. Because the clinical utility of Ct values is unclear, an expert should be consulted if these values are used to guide clinical decisions.

Serologic or Antibody Testing for Diagnosis of SARS-CoV-2 Infection

Unlike NAATs and antigen tests, which detect the presence of SARS-CoV-2, serologic or antibody tests can detect recent or prior SARS-CoV-2 infection or vaccination. The CDC recommends that antibody tests should not be used to diagnose current SARS-CoV-2 infection.¹² It may take 21 days or longer after symptom onset for seroconversion to occur (i.e., the development of detectable immunoglobulin M or immunoglobulin G antibodies to SARS-CoV-2).¹³⁻¹⁸

No serologic tests for SARS-CoV-2 have been approved by the FDA. Some, but not all, commercially available serologic tests for SARS-CoV-2 have received EUAs from the FDA.¹⁹ Several professional societies and federal agencies, including the <u>Infectious Diseases Society of America</u>, the <u>CDC</u>, and the <u>FDA</u>, provide guidance on the use of serologic testing for SARS-CoV-2.

Serologic Testing and Immunity to SARS-CoV-2 Infection

Currently, antibody tests are not recommended for assessing SARS-CoV-2 immunity following COVID-19 vaccination or for assessing the need for vaccination in a person who is unvaccinated.

The FDA has issued EUAs for more than 80 SARS-CoV-2 serologic tests since the beginning of the pandemic. However, these tests are not currently authorized for routine use in making individual medical decisions.¹⁹ SARS-CoV-2 serologic tests are authorized for detecting antibodies, but their

ability to predict protective immunity has not been validated. Most of these tests are not standardized. Furthermore, as SARS-CoV-2 is not a well-conserved virus, mutations in the receptor binding domain of the virus could lead to decreased binding affinity between antibodies and SARS-CoV-2–specific antigens.

If a serologic test is performed, the result should be interpreted with caution. First, it remains unclear how long SARS-CoV-2 antibodies persist following infection or vaccination. A negative serologic test result also does not preclude prior SARS-CoV-2 infection or vaccination against COVID-19. Second, some people who are infected with SARS-CoV-2 or who are vaccinated against COVID-19 (e.g., those who are immunocompromised) may not develop measurable levels of antibodies. It is presumed that those who do not have measurable antibodies after vaccination are at higher risk of SARS-CoV-2 infection than those who have measurable antibodies. Third, because nucleocapsid proteins are not a constituent of the vaccines that are currently approved by the FDA, available through EUAs, or in late-stage clinical trials, serologic tests that detect antibodies by recognizing nucleocapsid proteins should be used to distinguish between antibody responses to natural infection and vaccine-induced antibody responses to the SARS-CoV-2 spike protein antigen.

Assuming that the test is reliable, serologic tests that identify recent or prior SARS-CoV-2 infection may be used to determine who may be eligible to donate COVID-19 convalescent plasma and may aid in diagnosing multisystem inflammatory syndrome in children (MIS-C) and multisystem inflammatory syndrome in adults (MIS-A).

References

- 1. Centers for Disease Control and Prevention. Overview of testing for SARS-CoV-2, the virus that causes COVID-19. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/testing-overview.html</u>. Accessed December 7, 2023.
- Centers for Disease Control and Prevention. Isolation and precautions for people with COVID-19. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/your-health/isolation.html</u>. Accessed December 5, 2023.
- 3. Food and Drug Administration. In vitro diagnostics EUAs. 2023. Available at: <u>https://www.fda.gov/medical-devices/covid-19-emergency-use-authorizations-medical-devices/in-vitro-diagnostics-euas</u>. Accessed December 4, 2023.
- 4. Centers for Disease Control and Prevention. CDC's influenza SARS-CoV-2 multiplex assay. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/lab/multiplex.html</u>. Accessed December 12, 2023.
- Centers for Disease Control and Prevention. Interim guidelines for collecting and handling of clinical specimens for COVID-19 testing. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/lab/</u> guidelines-clinical-specimens.html. Accessed December 7, 2023.
- 6. Centers for Disease Control and Prevention. Considerations for SARS-CoV-2 antigen testing for healthcare providers testing individuals in the community. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/lab/resources/antigen-tests-guidelines.html</u>. Accessed December 11, 2023.
- 7. Centers for Disease Control and Prevention. Nucleic acid amplification tests (NAATs). 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/lab/naats.html</u>. Accessed December 11, 2023.
- 8. Food and Drug Administration. Genetic variants of SARS-CoV-2 may lead to false negative results with molecular tests for detection of SARS-CoV-2—letter to clinical laboratory staff and health care providers. 2021. Available at: <u>https://www.fda.gov/medical-devices/letters-health-care-providers/genetic-variants-sars-cov-2-may-lead-false-negative-results-molecular-tests-detection-sars-cov-2</u>. Accessed December 7, 2023.
- 9. Food and Drug Administration. SARS-CoV-2 viral mutations: impact on COVID-19 tests. 2023. Available at: https://www.fda.gov/medical-devices/coronavirus-covid-19-and-medical-devices/sars-cov-2-viral-mutations-

impact-covid-19-tests. Accessed December 7, 2023.

- Kucirka LM, Lauer SA, Laeyendecker O, Boon D, Lessler J. Variation in false-negative rate of reverse transcriptase polymerase chain reaction-based SARS-CoV-2 tests by time since exposure. *Ann Intern Med*. 2020;173(4):262-267. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32422057</u>.
- Xiao AT, Tong YX, Zhang S. Profile of RT-PCR for SARS-CoV-2: a preliminary study from 56 COVID-19 patients. *Clin Infect Dis.* 2020;71(16):2249-2251. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32306036.
- 12. Centers for Disease Control and Prevention. Interim guidelines for COVID-19 antibody testing. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/testing/antibody-tests-guidelines.html</u>. Accessed December 7, 2023.
- Guo L, Ren L, Yang S, et al. Profiling early humoral response to diagnose novel coronavirus disease (COVID-19). *Clin Infect Dis*. 2020;71(15):778-785. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32198501</u>.
- Haveri A, Smura T, Kuivanen S, et al. Serological and molecular findings during SARS-CoV-2 infection: the first case study in Finland, January to February 2020. *Euro Surveill*. 2020;25(11):2000266. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32209163</u>.
- 15. Long QX, Liu BZ, Deng HJ, et al. Antibody responses to SARS-CoV-2 in patients with COVID-19. *Nat Med.* 2020;26(6):845-848. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32350462</u>.
- Okba NMA, Müller MA, Li W, et al. Severe acute respiratory syndrome coronavirus 2–specific antibody responses in coronavirus disease patients. *Emerg Infect Dis*. 2020;26(7):1478-1488. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32267220</u>.
- Xiang F, Wang X, He X, et al. Antibody detection and dynamic characteristics in patients with coronavirus disease 2019. *Clin Infect Dis*. 2020;71(8):1930-1934. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32306047</u>.
- 18. Zhao J, Yuan Q, Wang H, et al. Antibody responses to SARS-CoV-2 in patients with novel coronavirus disease 2019. *Clin Infect Dis*. 2020;71(16):2027-2034. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32221519</u>.
- 19. Food and Drug Administration. EUA authorized serology test performance. 2022. Available at: <u>https://www.fda.gov/medical-devices/coronavirus-disease-2019-covid-19-emergency-use-authorizations-medical-devices/eua-authorized-serology-test-performance</u>. Accessed December 7, 2023.

Prevention of SARS-CoV-2 Infection

Last Updated: December 20, 2023

Summary Recommendation

• The COVID-19 Treatment Guidelines Panel (the Panel) recommends COVID-19 vaccination for everyone who is eligible according to the Centers for Disease Control and Prevention's Advisory Committee on Immunization Practices (AI).

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

General Prevention Measures

Transmission of SARS-CoV-2 occurs primarily through exposure to respiratory droplets.¹ Exposure can occur when individuals inhale droplets or particles that contain the virus or touch mucous membranes with hands that have been contaminated with the virus. Exhaled droplets or particles can also deposit the virus onto exposed mucous membranes.

The risk of SARS-CoV-2 transmission can be reduced by covering coughs and sneezes, wearing a well-fitted mask around others, and isolating when experiencing symptoms. Frequent handwashing also effectively reduces the risk of infection.² Health care providers should follow the Centers for Disease Control and Prevention (CDC) recommendations for infection control and the appropriate use of personal protective equipment.³

COVID-19 Vaccines

Recommendation

• The Panel recommends COVID-19 vaccination for everyone who is eligible according to the CDC's Advisory Committee on Immunization Practices (AI).

Rationale

Vaccination is the most effective way to prevent COVID-19. Two 2023–2024 mRNA vaccines, BNT162b2 (Pfizer-BioNTech) and mRNA-1273 (Moderna), and the 2023–2024 recombinant spike protein with adjuvant vaccine NVX-CoV2373 (Novavax)⁴ are currently available in the United States. The adenovirus vector vaccine Ad26.COV2.S (Johnson & Johnson/Janssen) is no longer available in the United States.

COVID-19 vaccination is recommended for everyone aged ≥ 6 months in the United States. The Food and Drug Administration (FDA) Emergency Use Authorization fact sheet and the product label for each vaccine provide detailed information on the vaccination schedule and the doses that are approved or authorized for that vaccine. The type and dose of vaccine and the timing of the doses depend on the recipient's age and underlying medical conditions. The CDC regularly updates the clinical considerations for the COVID-19 vaccines currently approved by the FDA or authorized for use in the United States.⁵

Adverse Events

COVID-19 vaccines are safe and effective. Local and systemic adverse events are relatively common with these vaccines. Most of the adverse events that occurred during vaccine trials were mild or

moderate in severity (i.e., they did not prevent vaccinated people from engaging in daily activities) and resolved after 1 or 2 days. There have been a few reports of severe allergic reactions following COVID-19 vaccination, including rare reports of patients who experienced anaphylaxis after receiving an mRNA vaccine.^{6,7}

Thrombosis with thrombocytopenia syndrome is a serious condition characterized by blood clots in large blood vessels and low platelet levels. The prevalence of the syndrome was approximately 4 per million among people who received the Johnson & Johnson/Janssen vaccine.^{8,9} That vaccine is no longer available in the United States. If a patient experiences thrombosis and thrombocytopenia syndrome after receiving a COVID-19 vaccine outside of the United States, a hematologist should be consulted about evaluation and management.

Myocarditis and pericarditis after COVID-19 vaccination are rare, and most of the reported cases were very mild and self-limiting.¹⁰ These conditions have occurred most often in male adolescents, young adults, and people who have received mRNA vaccines.

The results of recent studies suggest that adults aged ≥ 18 years who received the Johnson & Johnson/Janssen vaccine have an increased risk of Guillain-Barré syndrome.¹¹ In contrast, people who received mRNA vaccines do not have an increased risk of Guillain-Barré syndrome.¹²

The CDC monitors severe adverse events, such as strokes, and provides regular updates on <u>selected</u> <u>adverse events of COVID-19 vaccines</u>.

Vaccination in Pregnant and Lactating People

Pregnant and lactating individuals were not included in the initial COVID-19 vaccine trials. However, the CDC and the American College of Obstetricians and Gynecologists recommend vaccination for pregnant and lactating people. This recommendation is based on the accumulated safety and efficacy data on the use of these vaccines in pregnant people, as well on as the increased risk of severe disease in pregnant individuals with COVID-19.¹³⁻¹⁷ These organizations also recommend vaccination for people who are trying to become pregnant or who may become pregnant in the future. The American College of Obstetricians and Gynecologists provides guidance for clinicians on counseling pregnant patients about COVID-19 vaccination.¹⁸

Pre-Exposure Prophylaxis

As of January 2024, no biomedical intervention other than vaccines prevents COVID-19 disease. Previously, the FDA authorized the use of the anti-SARS-CoV-2 monoclonal antibodies tixagevimab plus cilgavimab (Evusheld) as pre-exposure prophylaxis (PrEP) of COVID-19 in people who were not expected to mount an adequate immune response to COVID-19 vaccination and in people with COVID-19 vaccine contraindications.¹⁹ Due to the increased prevalence of Omicron subvariants that are not susceptible to tixagevimab plus cilgavimab, this combination is not currently authorized by the FDA for use as PrEP of COVID-19.²⁰ It remains critical that these individuals:

- Keep up to date with COVID-19 vaccination unless a contraindication exists.
- Take precautions to avoid infection. The CDC provides <u>information on the prevention of</u> <u>COVID-19</u> in people who are immunocompromised.
- Be tested for SARS-CoV-2 infection if they experience signs and symptoms consistent with COVID-19 and, if infected, promptly seek medical attention.

Post-Exposure Prophylaxis

As of January 2024, no biomedical intervention other than vaccines prevents disease after exposure to SARS-CoV-2. Previously, the FDA authorized the use of the anti-SARS-CoV-2 monoclonal antibody products bamlanivimab plus etesevimab and casirivimab plus imdevimab as post-exposure prophylaxis (PEP) in certain people at high risk of progression to severe COVID-19. However, the Omicron subvariants are not susceptible to these products; therefore, their use as SARS-CoV-2 PEP is not recommended.

References

- 1. Centers for Disease Control and Prevention. COVID-19 overview and infection prevention and control priorities in non-U.S. healthcare settings. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/non-us-settings/overview/index.html</u>. Accessed December 1, 2023.
- 2. Centers for Disease Control and Prevention. How to protect yourself and others. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/prevent-getting-sick/prevention.html</u>. Accessed December 1, 2023.
- Centers for Disease Control and Prevention. Interim infection prevention and control recommendations for healthcare personnel during the coronavirus disease 2019 (COVID-19) pandemic. 2023. Available at: <u>https:// www.cdc.gov/coronavirus/2019-ncov/hcp/infection-control-recommendations.html</u>. Accessed December 1, 2023.
- 4. Food and Drug Administration. FDA authorizes updated Novavax COVID-19 vaccine formulated to better protect against currently circulating variants. 2023. Available at: <u>https://www.fda.gov/news-events/press-announcements/fda-authorizes-updated-novavax-covid-19-vaccine-formulated-better-protect-against-currently</u>. Accessed December 1, 2023.
- Centers for Disease Control and Prevention. Use of COVID-19 vaccines in the United States. 2023. Available at: <u>https://www.cdc.gov/vaccines/covid-19/clinical-considerations/covid-19-vaccines-us.html</u>. Accessed December 1, 2023.
- 6. Centers for Disease Control and Prevention. Interim considerations: preparing for the potential management of anaphylaxis after COVID-19 vaccination. 2022. Available at: <u>https://www.cdc.gov/vaccines/covid-19/clinical-considerations/managing-anaphylaxis.html</u>. Accessed December 1, 2023.
- Food and Drug Administration. Fact sheet for healthcare providers administering vaccine (vaccination providers): Emergency Use Authorization (EUA) of the Janssen COVID-19 vaccine to prevent coronavirus disease 2019 (COVID-19). 2022. Available at: <u>https://www.fda.gov/media/146304/download</u>.
- See I, Su JR, Lale A, et al. US case reports of cerebral venous sinus thrombosis with thrombocytopenia after Ad26.COV2.S vaccination, March 2 to April 21, 2021. *JAMA*. 2021;325(24):2448-2456. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/33929487</u>.
- 9. See I, Lale A, Marquez P, et al. Case series of thrombosis with thrombocytopenia syndrome after COVID-19 vaccination—United States, December 2020 to August 2021. *Ann Intern Med.* 2022;175(4):513-522. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35038274.
- Centers for Disease Control and Prevention. Selected adverse events reported after COVID-19 vaccination. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/vaccines/safety/adverse-events.html</u>. Accessed December 1, 2023.
- Hanson KE, Goddard K, Lewis N, et al. Incidence of Guillain-Barré syndrome after COVID-19 vaccination in the Vaccine Safety Datalink. *JAMA Netw Open*. 2022;5(4):e228879. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35471572</u>.
- 12. Abara WE, Gee J, Marquez P, et al. Reports of Guillain-Barré syndrome after COVID-19 vaccination in the United States. *JAMA Netw Open*. 2023;6(2):e2253845. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36723942</u>.
- 13. Centers for Disease Control and Prevention. COVID-19 vaccines while pregnant or breastfeeding. 2023.

COVID-19 Treatment Guidelines

Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/vaccines/recommendations/pregnancy.html</u>. Accessed December 1, 2023.

- Shimabukuro TT, Kim SY, Myers TR, et al. Preliminary findings of mRNA COVID-19 vaccine safety in pregnant persons. *N Engl J Med*. 2021;384(24):2273-2282. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33882218</u>.
- Zauche LH, Wallace B, Smoots AN, et al. Receipt of mRNA COVID-19 vaccines and risk of spontaneous abortion. *N Engl J Med.* 2021;385(16):1533-1535. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34496196.
- 16. Goldshtein I, Nevo D, Steinberg DM, et al. Association between BNT162b2 vaccination and incidence of SARS-CoV-2 infection in pregnant women. JAMA. 2021;326(8):728-735. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34251417</u>.
- 17. Collier AY, McMahan K, Yu J, et al. Immunogenicity of COVID-19 mRNA vaccines in pregnant and lactating women. *JAMA*. 2021;325(23):2370-2380. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33983379</u>.
- American College of Obstetricians and Gynecologists. COVID-19 vaccination considerations for obstetricgynecologic care. 2023. Available at: <u>https://www.acog.org/clinical/clinical-guidance/practice-advisory/</u> <u>articles/2020/12/covid-19-vaccination-considerations-for-obstetric-gynecologic-care</u>. Accessed December 1, 2023.
- Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Evusheld (tixagevimab co-packaged with cilgavimab). 2023. Available at: <u>https://www.fda.gov/media/154701/download</u>.
- 20. Food and Drug Administration. FDA announces Evusheld is not currently authorized for emergency use in the U.S. 2023. Available at: <u>https://www.fda.gov/drugs/drug-safety-and-availability/fda-announces-evusheld-not-currently-authorized-emergency-use-us</u>. Accessed December 1, 2023.

Clinical Spectrum of SARS-CoV-2 Infection

Last Updated: March 6, 2023

Patients with SARS-CoV-2 infection can experience a range of clinical manifestations, from no symptoms to critical illness. In general, adults with SARS-CoV-2 infection can be grouped into the following severity of illness categories; however, the criteria for each category may overlap or vary across clinical guidelines and clinical trials, and a patient's clinical status may change over time.

- *Asymptomatic or presymptomatic infection:* Individuals who test positive for SARS-CoV-2 using a virologic test (i.e., a nucleic acid amplification test [NAAT] or an antigen test) but have no symptoms consistent with COVID-19.
- *Mild illness:* Individuals who have any of the various signs and symptoms of COVID-19 (e.g., fever, cough, sore throat, malaise, headache, muscle pain, nausea, vomiting, diarrhea, loss of taste and smell) but do not have shortness of breath, dyspnea, or abnormal chest imaging.
- *Moderate illness:* Individuals who show evidence of lower respiratory disease during clinical assessment or imaging and who have an oxygen saturation measured by pulse oximetry $(SpO_2) \ge 94\%$ on room air at sea level.
- *Severe illness:* Individuals who have $\text{SpO}_2 < 94\%$ on room air at sea level, a ratio of arterial partial pressure of oxygen to fraction of inspired oxygen ($\text{PaO}_2/\text{FiO}_2$) <300 mm Hg, a respiratory rate >30 breaths/min, or lung infiltrates >50%.
- *Critical illness:* Individuals who have respiratory failure, septic shock, and/or multiple organ dysfunction.

 SpO_2 is a key parameter for defining the illness categories listed above. However, pulse oximetry has important limitations (discussed in more detail below). Clinicians who use SpO_2 when assessing a patient must be aware of those limitations and conduct the assessment in the context of that patient's clinical status.

Patients aged \geq 50 years are at a higher risk of progressing to severe COVID-19. Other underlying conditions associated with a higher risk of severe COVID-19 include asthma, cancer, cardiovascular disease, chronic kidney disease, chronic liver disease, chronic lung disease, diabetes, advanced or untreated HIV infection, obesity, pregnancy, cigarette smoking, and being a recipient of immunosuppressive therapy or a transplant.¹ Health care providers should closely monitor patients with these conditions until they achieve clinical recovery.

The initial evaluation for patients may include chest imaging (e.g., X-ray, ultrasound or computed tomography scan) and an electrocardiogram. Laboratory testing should include a complete blood count with differential and a metabolic profile, including liver and renal function tests. Although inflammatory markers such as C-reactive protein (CRP), D-dimer, and ferritin are not routinely measured as part of standard care, results from such measurements may have prognostic value.²⁻⁴

The definitions for the severity of illness categories also apply to pregnant patients. However, the threshold for certain interventions is different for pregnant and nonpregnant patients. For example, oxygen supplementation for pregnant patients is generally used when SpO_2 falls below 95% on room air at sea level to accommodate the physiologic changes in oxygen demand during pregnancy and to ensure adequate oxygen delivery to the fetus.⁵

If laboratory parameters are used for monitoring pregnant patients and making decisions about interventions, clinicians should be aware that normal physiologic changes during pregnancy can alter several laboratory values. In general, leukocyte cell count increases throughout gestation and delivery and peaks during the immediate postpartum period. This increase is mainly due to neutrophilia.⁶ D-dimer and CRP levels also increase during pregnancy and are often higher in pregnant patients than in nonpregnant patients.⁷ Detailed information on treating COVID-19 in pregnant patients can be found in <u>Special Considerations During Pregnancy and After Delivery</u> and in the pregnancy considerations subsections in the Guidelines.

In children with COVID-19, radiographic abnormalities are common and, for the most part, should not be the only criteria used to determine the severity of illness. The normal values for respiratory rate also vary with age in children; therefore, hypoxemia should be the primary criterion used to define severe COVID-19, especially in younger children. In a small subset of children and young adults, SARS-CoV-2 infection may be followed by the severe inflammatory condition multisystem inflammatory syndrome in children (MIS-C).^{8,9} This syndrome is discussed in detail in <u>Special Considerations in Children</u>.

Clinical Considerations for the Use of Pulse Oximetry

During the COVID-19 pandemic, the use of pulse oximetry to assess and monitor patients' oxygenation status increased in hospital, outpatient health care facility, and home settings. Although pulse oximetry is useful for estimating blood oxygen levels, pulse oximeters may not accurately detect hypoxemia under certain circumstances. To avoid delays in recognizing hypoxemia, clinicians who use pulse oximetry to assist with clinical decisions should keep these limitations in mind.

Pulse oximetry results can be affected by skin pigmentation, thickness, or temperature. Poor blood circulation or the use of tobacco or fingernail polish also may affect results. The Food and Drug Administration (FDA) advises clinicians to refer to the label or manufacturer website of a pulse oximeter or sensor to ascertain its accuracy.¹⁰ The FDA also advises using pulse oximetry only as an estimate of blood oxygen saturation, because an SpO₂ reading represents a range of arterial oxygen saturation (SaO₂). For example, an SpO₂ reading of 90% may represent a range of SaO₂ from 86% to 94%.

Several published reports have compared measurements of SpO_2 and SaO_2 in patients with and without COVID-19. The studies demonstrated that occult hypoxemia (defined as $\text{SaO}_2 < 88\%$ despite $\text{SpO}_2 > 92\%$) was more common in patients with darker skin pigmentation, which may result in adverse consequences.¹¹⁻¹³ The likelihood of error was greater in the lower ranges of SpO_2 . In 2 studies, greater incidences of occult hypoxemia were observed in patients who were Black, Hispanic, or Asian than in patients who were White.^{11,12} In 1 of these studies, occult hypoxemia was associated with more organ dysfunction and hospital mortality.¹³

A 5-hospital registry study of patients evaluated in the emergency department or hospitalized for COVID-19 found that 24% were not identified as eligible for treatment due to overestimation of SaO_2 . The majority of patients (55%) who were not identified as eligible were Black. The study also examined the amount of time delay patients experienced before their treatment eligibility was identified. The median delay for patients who were Black was 1 hour longer than the delay for patients who were White.¹⁴

In pulse oximetry, skin tone is an important variable, but accurately measuring oxygen saturation is a complex process. One observational study in adults was unable to identify a consistently predictable difference between SaO_2 and SpO_2 over time for individual patients.¹¹ Factors other than skin pigmentation (e.g., peripheral perfusion, pulse oximeter sensor placement) are likely involved.

Despite the limitations of pulse oximetry, an FDA-cleared pulse oximeter for home use can contribute

to an assessment of a patient's overall clinical status. Practitioners should advise patients to follow the manufacturer's instructions for use, place the oximeter on the index or ring finger, and ensure the hand is warm, relaxed, and held below the level of the heart. Fingernail polish should be removed before testing. Patients should be at rest, indoors, and breathing quietly without talking for several minutes before testing. Rather than accepting the first reading, patients or caretakers should observe the readings on the pulse oximeter for \geq 30 seconds until a steady number is displayed and inform their health care provider if the reading is repeatedly below a previously specified value (generally 95% on room air at sea level).^{10,15} Pulse oximetry has been widely adopted as a remote patient monitoring tool, but when the use of pulse oximeters is compared with close monitoring of clinical progress via video consultation, telephone calls, text messaging, or home visits, there is insufficient evidence that it improves clinical outcomes.^{16,17}

Not all commercially available pulse oximeters have been cleared by the FDA. SpO₂ readings obtained through non-FDA-cleared devices, such as over-the-counter sports oximeters or mobile phone applications, lack sufficient accuracy for clinical use. Abnormal readings on these devices should be confirmed with an FDA-cleared device or an arterial blood gas analysis.^{18,19}

Regardless of the setting, SpO_2 should always be interpreted within the context of a patient's entire clinical presentation. A patient's signs and symptoms (e.g., dyspnea, tachypnea, chest pain, changes in cognition or attentional state, cyanosis) should be given greater weight than a pulse oximetry result.

Asymptomatic or Presymptomatic Infection

Asymptomatic SARS-CoV-2 infection can occur, although the percentage of patients who remain truly asymptomatic throughout the course of infection is variable and incompletely defined. The percentage of individuals who present with asymptomatic infection and progress to clinical disease is unclear. Some asymptomatic individuals have been reported to have objective radiographic findings consistent with COVID-19 pneumonia.^{20,21}

Mild Illness

Patients with mild illness may exhibit a variety of signs and symptoms (e.g., fever, cough, sore throat, malaise, headache, muscle pain, nausea, vomiting, diarrhea, loss of taste and smell). They do not have shortness of breath, dyspnea on exertion, or abnormal imaging. Most patients who are mildly ill can be managed in an ambulatory setting or at home through telemedicine or telephone visits. No imaging or specific laboratory evaluations are routinely indicated in otherwise healthy patients with mild COVID-19. Patients aged \geq 50 years and those with underlying comorbidities are at higher risk of disease progression and are candidates for antiviral therapy. See Therapeutic Management of Nonhospitalized Adults With COVID-19 for recommendations regarding anti-SARS-CoV-2 therapies.

Moderate Illness

Moderate illness is defined as evidence of lower respiratory disease during clinical assessment or imaging, with $\text{SpO}_2 \ge 94\%$ on room air at sea level. Given that pulmonary disease can progress rapidly in patients with COVID-19, patients with moderate disease should be closely monitored. See <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for recommendations regarding anti-SARS-CoV-2 therapies in patients at high risk of progression to severe disease.

Severe Illness

Patients with COVID-19 are considered to have severe illness if they have $SpO_2 < 94\%$ on room air at sea level, $PaO_2/FiO_2 < 300$ mm Hg, a respiratory rate >30 breaths/min, or lung infiltrates >50%.

These patients may experience rapid clinical deterioration and should be given oxygen therapy and be hospitalized. See <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> for treatment recommendations.

Critical Illness

SARS-CoV-2 infection can cause acute respiratory distress syndrome, virus-induced distributive (septic) shock, cardiac shock, an exaggerated inflammatory response, thrombotic disease, and exacerbation of underlying comorbidities.

The clinical management of patients with COVID-19 who are in the intensive care unit should include treatment with immunomodulators, and, in some cases, the addition of remdesivir. These patients should also receive treatment for any comorbid conditions and nosocomial complications. For more information, see <u>Critical Care for Adults</u> and <u>Therapeutic Management of Hospitalized Adults With</u> <u>COVID-19</u>.

Infectious Complications in Patients With COVID-19

Some patients with COVID-19 may have additional infections when they present for care or that develop during the course of treatment. These coinfections may complicate treatment and recovery. Older patients or those with certain comorbidities or immunocompromising conditions may be at higher risk for these infections. The use of immunomodulators such as dexamethasone, interleukin-6 inhibitors (e.g., tocilizumab, sarilumab), or Janus kinase inhibitors (e.g., baricitinib, tofacitinib) to treat COVID-19 may also be a risk factor for infectious complications. However, when these therapies are used appropriately, the benefits outweigh the risks.

Infectious complications in patients with COVID-19 can be categorized as follows:

- Coinfections at presentation: Although most individuals present with only SARS-CoV-2 infection, concomitant viral infections, including influenza and other respiratory viruses, have been reported.²² Community-acquired bacterial pneumonia has also been reported, but it is uncommon, with a prevalence that ranges from 0% to 6% of people with SARS-CoV-2 infection.^{22,23} Antibacterial therapy is generally not recommended unless additional evidence for bacterial pneumonia is present (e.g., leukocytosis, the presence of a focal infiltrate on imaging).
- *Reactivation of latent infections:* There are case reports of underlying chronic hepatitis B virus and latent tuberculosis infections reactivating in patients with COVID-19 who receive immunomodulators as treatment, although the data are currently limited.²⁴⁻²⁶ Reactivation of herpes simplex virus and varicella zoster virus infections have also been reported.²⁷ Cases of severe and disseminated strongyloidiasis have been reported in patients with COVID-19 during treatment with tocilizumab and corticosteroids.^{28,29} Many clinicians would initiate empiric treatment (e.g., with the antiparasitic drug ivermectin), with or without serologic testing, in patients who require immunomodulators for the treatment of COVID-19 and have come from areas where *Strongyloides* is endemic (i.e., tropical, subtropical, or warm temperate areas).³⁰
- *Nosocomial infections:* Hospitalized patients with COVID-19 may acquire common nosocomial infections, such as hospital-acquired pneumonia (including ventilator-associated pneumonia), line-related bacteremia or fungemia, catheter-associated urinary tract infection, and *Clostridioides difficile*-associated diarrhea. Early diagnosis and treatment of these infections are important for improving outcomes in these patients.
- *Opportunistic fungal infections:* Invasive fungal infections, including aspergillosis and mucormycosis, have been reported in hospitalized patients with COVID-19.³¹⁻³⁴ Although these

infections are relatively rare, they can be fatal, and they may be seen more commonly in patients who are immunocompromised or receiving mechanical ventilation. The majority of mucormycosis cases have been reported in India and are associated with diabetes mellitus or the use of corticosteroids.^{35,36} The approach for managing these fungal infections should be the same as the approach for managing invasive fungal infections in other settings.

SARS-CoV-2 Reinfection and Breakthrough Infection

As seen with other respiratory viral infections, reinfection after recovery from prior infection has been reported for SARS-CoV-2.³⁷ Reinfection may occur as initial immune responses to the primary infection wane over time. Data regarding the prevalence, risk factors, timing, and severity of reinfection are evolving and likely vary depending on the circulating variants. Breakthrough SARS-CoV-2 infections (i.e., infection in people who completed the primary vaccine series with or without booster doses) also occurs.³⁸ When compared with infection in people who are unvaccinated, breakthrough infection appears less likely to lead to severe illness or symptoms that persist ≥ 28 days.³⁸⁻⁴³ The time to breakthrough infection (i.e., solid organ or bone marrow transplant recipients or people with HIV) than for those with no immunocompromising conditions.³⁸ For information on diagnostic testing in the setting of suspected reinfection, see Testing for SARS-CoV-2 Infection. In addition, information about the epidemiology, diagnosis, and evaluation of suspected SARS-CoV-2 reinfection or breakthrough infection is provided by the Centers for Disease Control and Prevention (CDC).

Although data are limited, no evidence suggests that the treatment of suspected or documented SARS-CoV-2 reinfection or breakthrough infection should be different from the treatment used during the initial infection, as outlined in <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> and <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>.

Prolonged Viral Shedding, Persistent Symptoms, and Other Conditions After SARS-CoV-2 Infection

Symptomatic SARS-CoV-2 infection is typically associated with a decline in viral shedding and resolution of COVID-19 symptoms over days to weeks. However, in some cases, reduced viral shedding and symptom resolution are followed by viral or symptom rebound. People who are immunocompromised may experience viral shedding for many weeks. Some people experience symptoms that develop or persist for more than 4 weeks after the initial COVID-19 diagnosis.

Viral or Symptom Rebound Soon After COVID-19

Observational studies and results from clinical trials of therapeutic agents have described SARS-CoV-2 viral or COVID-19 symptom rebound in patients who have completed treatment for COVID-19.⁴⁴⁻⁴⁶ Viral and symptom rebounds have also occurred when anti-SARS-CoV-2 therapies were not used.^{46,47} Typically, this phenomenon has not been associated with progression to severe COVID-19.

Prolonged Viral Shedding in Patients Who Are Immunocompromised

Patients who are immunocompromised may experience prolonged shedding of SARS-CoV-2 with or without COVID-19 symptoms.^{48,49} Prolonged viral shedding may affect SARS-CoV-2 testing strategies and isolation duration for these patients. In some cases, the prolonged shedding may be associated with persistent COVID-19 symptoms. Currently, the <u>evidence is insufficient</u> to guide any clinical recommendations for managing the treatment of these individuals. Limited data support using combination antiviral therapy or extending the duration of COVID-19 therapies beyond the durations authorized or approved by the FDA. See <u>Special Considerations in People Who</u>

<u>Are Immunocompromised</u> for more information on the clinical management of people who are immunocompromised.

Persistent, New, or Recurrent Symptoms More Than 4 Weeks After SARS-CoV-2 Infection

Some patients report persistent, new, or recurrent symptoms and conditions (e.g., cardiopulmonary injury, neurocognitive impairment, new-onset diabetes) more than 4 weeks after the initial COVID-19 diagnosis.⁵⁰ The nomenclature for this phenomenon is evolving; no clinical terminology has been established. The terminology used includes long-COVID, post-COVID-19 condition, post-COVID-19 syndrome, and post-acute sequelae of SARS-CoV-2. Patients who have these symptoms or conditions have been called "long haulers."

Data on the incidence, natural history, and etiology of these symptoms are emerging. However, reports on these syndromes have several limitations, such as differing case definitions, a lack of comparator groups, and overlap between the reported symptoms and the symptoms of post-intensive care syndrome that have been described for patients without COVID-19. In addition, many reports only included patients who attended post-COVID clinics. Details on the pathogenesis, clinical presentation, and treatment for these conditions are beyond the scope of these Guidelines. The <u>CDC</u> provides information about the timeframes, presentation of symptoms, and management strategies for post-COVID conditions. Research on the prevalence, characteristics, and pathophysiology of persistent symptoms and conditions after COVID-19 is ongoing, including research through the National Institutes of Health's <u>RECOVER Initiative</u>, which aims to inform potential therapeutic strategies.

MIS-C and multisystem inflammatory syndrome in adults (MIS-A) are serious postinfectious complications of SARS-CoV-2 infection. For more information on these syndromes, see <u>Therapeutic</u> <u>Management of Hospitalized Children With MIS-C, Plus a Discussion on MIS-A</u>.

References

- 1. Centers for Disease Control and Prevention. Underlying medical conditions associated with higher risk for severe COVID-19: information for healthcare professionals. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/clinical-care/underlyingconditions.html</u>. Accessed February 15, 2023.
- Tan C, Huang Y, Shi F, et al. C-reactive protein correlates with computed tomographic findings and predicts severe COVID-19 early. *J Med Virol*. 2020;92(7):856-862. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32281668.
- 3. Berger JS, Kunichoff D, Adhikari S, et al. Prevalence and outcomes of D-dimer elevation in hospitalized patients with COVID-19. *Arterioscler Thromb Vasc Biol.* 2020;40(10):2539-2547. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32840379</u>.
- 4. Casas-Rojo JM, Antón-Santos JM, Millán-Núñez-Cortés J, et al. Clinical characteristics of patients hospitalized with COVID-19 in Spain: results from the SEMI-COVID-19 registry. *Rev Clin Esp (Barc)*. 2020;220(8):480-494. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32762922</u>.
- 5. Society for Maternal-Fetal Medicine. Management considerations for pregnant patients with COVID-19. 2021. Available at: <u>https://s3.amazonaws.com/cdn.smfm.org/media/2734/SMFM_COVID_Management_of_COVID_pos_preg_patients_2-2-21_(final).pdf</u>.
- Abbassi-Ghanavati M, Greer LG, Cunningham FG. Pregnancy and laboratory studies: a reference table for clinicians. *Obstet Gynecol*. 2009;114(6):1326-1331. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/19935037</u>.
- Anderson BL, Mendez-Figueroa H, Dahlke JD, et al. Pregnancy-induced changes in immune protection of the genital tract: defining normal. *Am J Obstet Gynecol*. 2013;208(4):321.e1-321.e9. Available at: <u>https://www. ncbi.nlm.nih.gov/pubmed/23313311</u>.

- Riphagen S, Gomez X, Gonzalez-Martinez C, Wilkinson N, Theocharis P. Hyperinflammatory shock in children during COVID-19 pandemic. *Lancet*. 2020;395(10237):1607-1608. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32386565</u>.
- 9. Verdoni L, Mazza A, Gervasoni A, et al. An outbreak of severe Kawasaki-like disease at the Italian epicentre of the SARS-CoV-2 epidemic: an observational cohort study. *Lancet*. 2020;395(10239):1771-1778. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32410760.
- Food and Drug Administration. Pulse oximeter accuracy and limitations: FDA safety communication. 2022. Available at: <u>https://www.fda.gov/medical-devices/safety-communications/pulse-oximeter-accuracy-and-limitations-fda-safety-communication</u>. Accessed February 17, 2023.
- 11. Chesley CF, Lane-Fall MB, Panchanadam V, et al. Racial disparities in occult hypoxemia and clinically based mitigation strategies to apply in advance of technological advancements. *Respir Care*. 2022;67(12):1499-1507. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35679133.
- Valbuena VSM, Seelye S, Sjoding MW, et al. Racial bias and reproducibility in pulse oximetry among medical and surgical inpatients in general care in the Veterans Health Administration 2013–19: multicenter, retrospective cohort study. *BMJ*. 2022;378:e069775. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35793817.
- 13. Wong AI, Charpignon M, Kim H, et al. Analysis of discrepancies between pulse oximetry and arterial oxygen saturation measurements by race and ethnicity and association with organ dysfunction and mortality. *JAMA Netw Open*. 2021;4(11):e2131674. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34730820</u>.
- Fawzy A, Wu TD, Wang K, et al. Racial and ethnic discrepancy in pulse oximetry and delayed identification of treatment eligibility among patients with COVID-19. *JAMA Intern Med.* 2022;182(7):730-738. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35639368</u>.
- Luks AM, Swenson ER. Pulse oximetry for monitoring patients with COVID-19 at home. Potential pitfalls and practical guidance. *Ann Am Thorac Soc.* 2020;17(9):1040-1046. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32521167</u>.
- 16. Alboksmaty A, Beaney T, Elkin S, et al. Effectiveness and safety of pulse oximetry in remote patient monitoring of patients with COVID-19: a systematic review. *Lancet Digit Health*. 2022;4(4):e279-e289. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35337644</u>.
- Lee KC, Morgan AU, Chaiyachati KH, et al. Pulse oximetry for monitoring patients with COVID-19 at home—a pragmatic, randomized trial. *N Engl J Med.* 2022;386(19):1857-1859. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35385625</u>.
- Harskamp RE, Bekker L, Himmelreich JCL, et al. Performance of popular pulse oximeters compared with simultaneous arterial oxygen saturation or clinical-grade pulse oximetry: a cross-sectional validation study in intensive care patients. *BMJ Open Respir Res.* 2021;8(1):e000939. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34489238</u>.
- Lipnick MS, Feiner JR, Au P, Bernstein M, Bickler PE. The accuracy of 6 inexpensive pulse oximeters not cleared by the Food and Drug Administration: the possible global public health implications. *Anesth Analg.* 2016;123(2):338-345. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27089002</u>.
- 20. Zhang R, Ouyang H, Fu L, et al. CT features of SARS-CoV-2 pneumonia according to clinical presentation: a retrospective analysis of 120 consecutive patients from Wuhan city. *Eur Radiol*. 2020;30(8):4417-4426. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32279115</u>.
- 21. Inui S, Fujikawa A, Jitsu M, et al. Chest CT findings in cases from the cruise ship Diamond Princess with coronavirus disease 2019 (COVID-19). *Radiol Cardiothorac Imaging*. 2020;2(2):e200110. Available at: https://pubmed.ncbi.nlm.nih.gov/33778566.
- Kim D, Quinn J, Pinsky B, Shah NH, Brown I. Rates of co-infection between SARS-CoV-2 and other respiratory pathogens. *JAMA*. 2020;323(20):2085-2086. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/32293646</u>.

- 23. Kubin CJ, McConville TH, Dietz D, et al. Characterization of bacterial and fungal infections in hospitalized patients with coronavirus disease 2019 and factors associated with health care-associated infections. *Open Forum Infect Dis.* 2021;8(6):ofab201. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34099978.
- 24. Garg N, Lee YI. Reactivation TB with severe COVID-19. *Chest*. 2020;158(4):A777. Available at: <u>https://journal.chestnet.org/article/S0012-3692(20)32910-X/fulltext</u>.
- 25. Rodríguez-Tajes S, Miralpeix A, Costa J, et al. Low risk of hepatitis B reactivation in patients with severe COVID-19 who receive immunosuppressive therapy. *J Viral Hepat*. 2021;28(1):89-94. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32969557</u>.
- 26. Aldhaleei WA, Alnuaimi A, Bhagavathula AS. COVID-19 induced hepatitis B virus reactivation: a novel case from the United Arab Emirates. *Cureus*. 2020;12(6):e8645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32550096</u>.
- 27. Xu R, Zhou Y, Cai L, et al. Co-reactivation of the human herpesvirus alpha subfamily (herpes simplex virus-1 and varicella zoster virus) in a critically ill patient with COVID-19. *Br J Dermatol*. 2020;183(6):1145-1147. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32790074</u>.
- 28. Lier AJ, Tuan JJ, Davis MW, et al. Case report: disseminated strongyloidiasis in a patient with COVID-19. *Am J Trop Med Hyg.* 2020;103(4):1590-1592. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32830642</u>.
- 29. Marchese V, Crosato V, Gulletta M, et al. Strongyloides infection manifested during immunosuppressive therapy for SARS-CoV-2 pneumonia. *Infection*. 2021;49(3):539-542. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32910321.
- Stauffer WM, Alpern JD, Walker PF. COVID-19 and dexamethasone: a potential strategy to avoid steroidrelated Strongyloides hyperinfection. *JAMA*. 2020;324(7):623-624. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32761166</u>.
- Salmanton-García J, Sprute R, Stemler J, et al. COVID-19-associated pulmonary aspergillosis, March–August 2020. Emerg Infect Dis. 2021;27(4):1077-1086. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33539721.
- Chong WH, Neu KP. Incidence, diagnosis and outcomes of COVID-19-associated pulmonary aspergillosis (CAPA): a systematic review. *J Hosp Infect*. 2021;113:115-129. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33891985</u>.
- 33. Machado M, Valerio M, Álvarez-Uría A, et al. Invasive pulmonary aspergillosis in the COVID-19 era: an expected new entity. *Mycoses*. 2021;64(2):132-143. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33210776</u>.
- 34. Yusuf E, Seghers L, Hoek RAS, et al. Aspergillus in critically ill COVID-19 patients: a scoping review. *J Clin Med.* 2021;10(11):2469. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34199528.
- 35. Singh AK, Singh R, Joshi SR, Misra A. Mucormycosis in COVID-19: a systematic review of cases reported worldwide and in India. *Diabetes Metab Syndr*. 2021;15(4):102146. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34192610</u>.
- 36. Pal R, Singh B, Bhadada SK, et al. COVID-19-associated mucormycosis: an updated systematic review of literature. *Mycoses*. 2021;64(12):1452-1459. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133798</u>.
- 37. Cohen JI, Burbelo PD. Reinfection with SARS-CoV-2: implications for vaccines. *Clin Infect Dis*. 2021;73(11):e4223-e4228. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33338197</u>.
- 38. Sun J, Zheng Q, Madhira V, et al. Association between immune dysfunction and COVID-19 breakthrough infection after SARS-CoV-2 vaccination in the US. *JAMA Intern Med.* 2022;182(2):153-162. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34962505.
- 39. Link-Gelles R, Ciesla AA, Roper LE, et al. Early estimates of bivalent mRNA booster dose vaccine effectiveness in preventing symptomatic SARS-CoV-2 infection attibutable to Omicron BA.5- and XBB/ XBB.1.5-related sublineages among immunocompetent adults—increasing community access to testing program, United States, December 2022–January 2023. MMWR Morb Mortal Wkly Rep. 2023;72(5):119-124.

Available at: https://pubmed.ncbi.nlm.nih.gov/36730051.

- 40. Antonelli M, Penfold RS, Merino J, et al. Risk factors and disease profile of post-vaccination SARS-CoV-2 infection in UK users of the COVID Symptom Study app: a prospective, community-based, nested, case-control study. *Lancet Infect Dis*. 2022;22(1):43-55. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34480857.
- Kuodi P, Gorelik Y, Zayyad H, et al. Association between BNT162b2 vaccination and reported incidence of post-COVID-19 symptoms: cross-sectional study 2020–21, Israel. NPJ Vaccines. 2022;7(1):101. Available at: https://pubmed.ncbi.nlm.nih.gov/36028498.
- 42. Centers for Disease Control and Prevention. Rates of laboratory-confirmed COVID-19 hospitalizations by vaccination status. 2022. Available at: <u>https://covid.cdc.gov/covid-data-tracker/#covidnet-hospitalizations-vaccination</u>. Accessed February 16, 2023.
- 43. Centers for Disease Control and Prevention. Rates of COVID-19 cases and deaths by vaccination status. 2023. Available at: <u>https://covid.cdc.gov/covid-data-tracker/#rates-by-vaccine-status</u>. Accessed February 20, 2023.
- 44. Charness ME, Gupta K, Stack G, et al. Rebound of SARS-CoV-2 infection after nirmatrelvir-ritonavir treatment. N Engl J Med. 2022;387(11):1045-1047. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36069968.</u>
- 45. Boucau J, Uddin R, Marino C, et al. Characterization of virologic rebound following nirmatrelvir-ritonavir treatment for COVID-19. *Clin Infect Dis.* 2023;76(3):e526-e529. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35737946.
- 46. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Lagevrio (molnupiravir) capsules. 2023. Available at: <u>https://www.fda.gov/media/155054/download</u>.
- 47. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Paxlovid. 2023. Available at: <u>https://www.fda.gov/media/155050/download</u>.
- 48. Qutub M, Aldabbagh Y, Mehdawi F, et al. Duration of viable SARS-CoV-2 shedding from respiratory tract in different human hosts and its impact on isolation discontinuation polices revision: a narrative review. *Clin Infect Pract.* 2022;13:100140. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35190799</u>.
- 49. Tarhini H, Recoing A, Bridier-Nahmias A, et al. Long-term severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infectiousness among three immunocompromised patients: from prolonged viral shedding to SARS-CoV-2 superinfection. *J Infect Dis.* 2021;223(9):1522-1527. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33556961.
- Centers for Disease Control and Prevention. Post-COVID conditions: information for healthcare providers. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/clinical-care/post-covid-conditions.html</u>. Accessed February 17, 2023.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Prioritization of Anti-SARS-CoV-2 Therapies for the Treatment of COVID-19 in Nonhospitalized Patients When There Are Logistical Constraints

Last Updated: December 20, 2023

The prioritization guidance in this section should be used **only** when logistical constraints limit the availability of therapies. When there are no logistical constraints, clinicians can prescribe therapies to treat SARS-CoV-2 infection to any eligible individual following the recommendations in these Guidelines.

If it is necessary to triage patients for receipt of anti-SARS-CoV-2 therapies, the COVID-19 Treatment Guidelines Panel (the Panel) suggests prioritizing patients based on their clinical risk factors for severe disease, their vaccination status, and their ability to mount an adequate immune response to COVID-19 vaccination or SARS-CoV-2 infection.

Prioritization schemes should include a plan for equitable distribution of scarce resources to individuals who may have less knowledge of or access to these therapies. The availability and distribution of recommended therapies should be monitored to ensure that access to products is equitable.

Patient Prioritization for Treatment

The Panel recommends using **ritonavir-boosted nirmatrelvir** (**Paxlovid**) to treat nonhospitalized adults (**AIIa**) and adolescents (**BIII**) who have mild to moderate COVID-19 and are at high risk of progressing to severe disease.

Remdesivir is a recommended option if ritonavir-boosted nirmatrelvir cannot be used. However, some treatment facilities may not have the ability to provide a 3-day course of remdesivir intravenous infusions to all eligible patients. In these situations, prioritizing patients who will benefit the most from the therapy becomes necessary. If administration of remdesivir is not feasible, clinicians should review the Panel's recommendations in <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for alternative treatment options.

The prioritization scheme below is based on 4 key elements: age, vaccination status, immune status, and clinical risk factors. For a list of risk factors, see the Centers for Disease Control and Prevention (CDC) webpage <u>Underlying Medical Conditions Associated With Higher Risk for Severe COVID-19</u>. The groups are listed by tier in descending order of priority.

Tier	Risk Group		
1	• People who are immunocompromised and are not expected to mount an adequate immune response to COVID-19 vaccination or SARS-CoV-2 infection due to their underlying conditions, regardless of vaccine status (see Immunocompromising Conditions below); <i>or</i>		
	• Unvaccinated individuals at the highest risk of severe disease (anyone aged ≥75 years or anyone aged ≥65 years with additional risk factors)		
2	• Unvaccinated individuals not included in Tier 1 who are at risk of severe disease (anyone aged ≥65 years or anyone aged <65 years with clinical risk factors)		
3	• Vaccinated individuals at risk of severe disease (anyone aged ≥65 years or anyone aged <65 years with clinical risk factors)		
	• Vaccinated individuals who are not up to date with their immunizations are likely at higher risk for severe disease; patients within this tier who are in this situation should be prioritized for treatment.		

COVID-19 Treatment Guidelines

Immunocompromising Conditions

The CDC website <u>COVID-19 Vaccines for People Who Are Moderately or Severely</u> <u>Immunocompromised</u> provides a list of moderate or severe immunocompromising conditions.

If there are logistical constraints to providing the Panel's recommended therapies to all individuals who are moderately to severely immunocompromised, the Panel suggests prioritizing patients who are least likely to mount an adequate response to COVID-19 vaccination or SARS-CoV-2 infection and are at risk for severe outcomes. This includes, but is not limited to, patients who:

- Are receiving active treatment for solid tumor and hematologic malignancies.
- Have hematologic malignancies (e.g., chronic lymphocytic lymphoma, non-Hodgkin lymphoma, multiple myeloma, acute leukemia) and are known to have poor responses to COVID-19 vaccines, regardless of the treatment status for the hematologic malignancy.
- Received a solid organ or islet transplant and are receiving immunosuppressive therapy.
- Received chimeric antigen receptor T cell therapy or a hematopoietic cell transplant and are within 2 years of transplantation or are receiving immunosuppressive therapy.
- Have a moderate or severe primary immunodeficiency (e.g., severe combined immunodeficiency, DiGeorge syndrome, Wiskott-Aldrich syndrome, common variable immunodeficiency disease).
- Have advanced or untreated HIV infection (defined as people with HIV and CD4 T lymphocyte cell counts <200 cells/mm³, a history of an AIDS-defining illness without immune reconstitution, or clinical manifestations of symptomatic HIV).
- Are receiving active treatment with high-dose corticosteroids (i.e., ≥20 mg prednisone or equivalent per day for ≥2 weeks), alkylating agents, antimetabolites, transplant-related immunosuppressive drugs, cancer chemotherapeutic agents classified as severely immunosuppressive, or immunosuppressive or immunomodulatory biologic agents (e.g., B cell-depleting agents).

If logistical constraints preclude administration of remdesivir to all prioritized patients, the Panel suggests further prioritizing patients who are more severely immunocompromised and have additional risk factors for severe disease.

Clinical Risk Factors

Some of the most important risk factors for severe COVID-19 include age (risk increases with each decade after age 50),¹ receipt of cancer treatment, immunocompromising conditions or receipt of immunosuppressive medications, cardiovascular disease, chronic kidney disease, chronic lung disease, diabetes, obesity (i.e., body mass index \geq 30), and pregnancy. For a complete list of risk factors, including information on the relative risk of severe disease, see the CDC webpage <u>Underlying</u> <u>Medical Conditions Associated With Higher Risk for Severe COVID-19</u>. Of note, the likelihood of developing severe COVID-19 increases when a person has multiple comorbidities.² For people who are not immunocompromised, vaccination with a primary COVID-19 vaccine series and booster doses dramatically reduces the risk of progressing to severe disease.

Although children with COVID-19 have substantially lower mortality than adults with COVID-19, severe disease can occur, especially in those with risk factors. See <u>Table 3b</u> for the Panel's framework for assessing the risk of progression to severe COVID-19 in children.

References

- 1. Centers for Disease Control and Prevention. Underlying medical conditions associated with higher risk for severe COVID-19: information for healthcare professionals. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/clinical-care/underlyingconditions.html</u>. Accessed October 17, 2023.
- 2. Rosenthal N, Cao Z, Gundrum J, Sianis J, Safo S. Risk factors associated with in-hospital mortality in a U.S. national sample of patients with COVID-19. *JAMA Netw Open*. 2020;3(12):e2029058. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33301018</u>.

Clinical Management of Adults Summary

Last Updated: October 10, 2023

Two main processes are thought to drive the pathogenesis of COVID-19. Early in the clinical course, the disease is primarily driven by the replication of SARS-CoV-2. Later in the clinical course, the disease appears to be driven by a dysregulated immune/inflammatory response to SARS-CoV-2 that leads to tissue damage. Based on this understanding, therapies that directly target SARS-CoV-2 are anticipated to have the greatest effect early in the course of the disease, while immunosuppressive/anti-inflammatory therapies are likely to be more beneficial in the later stages of COVID-19.

The clinical spectrum of SARS-CoV-2 infection includes asymptomatic or presymptomatic infection and mild, moderate, severe, and critical illness. Table 2a provides guidance for clinicians on the therapeutic management of nonhospitalized adult patients. This includes patients who do not require hospitalization or supplemental oxygen and those who have been discharged from an emergency department or a hospital. Table 2b provides guidance on the therapeutic management of hospitalized adult patients according to their disease severity and oxygen requirements.

Table 2a. Therapeutic Management of Nonhospitalized Adults With Mild to Moderate COVID-19Who Do Not Require Supplemental Oxygen

Patient Disposition	Panel's Recommendations		
All Patients	 Symptom management should be initiated for all patients (AIII). The Panel recommends against the use of dexamethasone^a or other systemic corticosteroids in the absence of another indication (AIIb). 		
Patients Who Are at High Risk of Progressing to Severe COVID-19 ^{b,c}	 Preferred therapies. Listed in order of preference: Ritonavir-boosted nirmatrelvir (Paxlovid)^d (Alla); see footnote on drug interactions^e Remdesivir^{d,f} (Blla) 		
	 Alternative therapy. For use when the preferred therapies are not available, feasible to use, or clinically appropriate: Molnupiravir^{d,g,h} (Clla) 		
Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.			

^a There is currently a lack of safety and efficacy data on the use of dexamethasone in outpatients with COVID-19. Using systemic glucocorticoids in outpatients with COVID-19 may cause harm.

- ^b For a list of risk factors, see the CDC webpage <u>Underlying Medical Conditions Associated With Higher Risk for Severe</u> <u>COVID-19</u>. When deciding whether to prescribe antiviral treatment to a patient who has been vaccinated, clinicians should be aware of the conditions associated with a high risk of disease progression. These conditions include older age, a prolonged amount of time since the most recent vaccine dose (e.g., >6 months), and a decreased likelihood of an adequate immune response to vaccination due to a moderate to severe immunocompromising condition or the receipt of immunosuppressive medications. The number and severity of risk factors also affects the level of risk.
- ^c For a discussion of potential treatment options for patients who are immunocompromised and have prolonged COVID-19 symptoms and evidence of ongoing viral replication, see below and <u>Special Considerations in People Who Are</u> <u>Immunocompromised</u>.
- ^d If a patient requires hospitalization after starting treatment, the full treatment course can be completed at the health care provider's discretion.
- ^e Ritonavir-boosted nirmatrelvir has significant drug-drug interactions. Clinicians should carefully review a patient's concomitant medications and evaluate potential drug-drug interactions. See <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications</u> for more information.
- ^f Administration of remdesivir requires an IV infusion once daily for 3 days.
- ⁹ Molnupiravir appears to have lower efficacy than the other options recommended by the Panel. Therefore, it should be considered when the other options are not available, feasible to use, or clinically appropriate.
- ^h The Panel **recommends against** the use of **molnupiravir** for the treatment of COVID-19 in pregnant patients unless there are no other options and therapy is clearly indicated **(AIII)**.

Key: CDC = Centers for Disease Control and Prevention; IV = intravenous; the Panel = the COVID-19 Treatment Guidelines Panel

COVID-19 Treatment Guidelines

Table 2b. Therapeutic Management of Hospitalized Adults With COVID-19

Disease Severity	Recommendations for Antiviral or Immunomodulator Therapy		Recommendations for
Disease Sevenity	Clinical Scenario	Recommendation	Anticoagulant Therapy
Hospitalized for Reasons Other Than COVID-19	Patients with mild to moderate COVID-19 who are at high risk of progressing to severe COVID-19 ^{a,b}	See <u>Therapeutic Management of Nonhospitalized Adults With</u> <u>COVID-19</u> .	For patients without an indication for therapeutic anticoagulation: • Prophylactic dose of heparin, unless
Hospitalized but Does Not Require Supplemental Oxygen	All patients	The Panel recommends against the use of dexamethasone (Alla) or other systemic corticosteroids (AllI) for the treatment of COVID-19.°	contraindicated (AI); (BIII) for pregnant patients
	Patients who are at high risk of progressing to severe COVID-19 ^{a,b}	Remdesivir ^d (BIIb) for patients who are immunocompromised; (BIII) for other high-risk patients	
Hospitalized and Requires Conventional Oxygen ^e	Patients who require minimal conventional oxygen	Remdesivir ^{d,f} (Blla)	For nonpregnant patients with D-dimer levels above the ULN who do not have an
	Most patients	Use dexamethasone plus remdesivir ^f (Blla). If remdesivir cannot be obtained, use dexamethasone (Bl) .	increased bleeding risk: • Therapeutic dose of heparin ^h (Clla)
	Patients who are receiving dexamethasone and who have rapidly increasing oxygen needs and systemic inflammation	Add 1 of the following immunomodulators: ⁹ Preferred • PO baricitinib (Blla) • IV tocilizumab (Blla) Alternatives • IV abatacept (Clla) • IV infliximab (Clla)	 For other patients: Prophylactic dose of heparin, unless contraindicated (AI); (BIII) for pregnant patients
Hospitalized and Requires HFNC Oxygen or NIV	All patients	 Dexamethasone should be administered to all patients (AI). If not already initiated, promptly add 1 of the following immunomodulators: <i>Preferred</i> PO baricitinib^{g,i} (AI) <i>Preferred Alternative</i> IV tocilizumab^{g,i} (BIIa) Additional Alternatives (Listed in Alphabetical Order) IV abatacept^{g,i} (CIIa) IV infliximab^{g,i} (CIIa) Add remdesivir to 1 of the options above in certain patients (for examples, see footnote).¹ 	 For patients without an indication for therapeutic anticoagulation: Prophylactic dose of heparin, unless contraindicated (AI); (BIII) for pregnant patients For patients who are started on a therapeutic dose of heparin in a non-ICU setting and then transferred to the ICU, the Panel recommends switching to a prophylactic dose of heparin, unless there is another indication for therapeutic anticoagulation (BIII).
Hospitalized and Requires MV or ECMO	All patients	 Dexamethasone should be administered to all patients (AI). If the patient has not already received a second immunomodulator, promptly add 1 of the following (listed in alphabetical order): PO baricitinib^{i,k} (BIIa) IV tocilizumab^{i,k} (BIIa) 	anucuayulaliuli (Dili) .

COVID-19 Treatment Guidelines

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

- ^a For a list of risk factors, see the CDC webpage <u>Underlying Medical Conditions Associated With Higher Risk for Severe</u> <u>COVID-19</u>.
- ^b Ritonavir-boosted nirmatrelvir (Paxlovid) has not been studied in hospitalized patients. The FDA product label for ritonavirboosted nirmatrelvir allows for its use in hospitalized patients with mild to moderate COVID-19 (i.e., those who do not require supplemental oxygen) who are at high risk of progressing to severe COVID-19 and who are within 5 days of symptom onset.
- ^c Corticosteroids that are prescribed for an underlying condition should be continued.

^d Evidence suggests that the benefit of remdesivir is greatest when the drug is given early in the course of COVID-19 (e.g., within 10 days of symptom onset).

- ^e Conventional oxygen refers to oxygen supplementation that is not HFNC oxygen, NIV, MV, or ECMO.
- ^f If these patients progress to requiring HFNC oxygen, NIV, MV, or ECMO, the full course of remdesivir should still be completed.
- ⁹ If none of the preferred or alternative options are available or feasible to use, the JAK inhibitor **PO tofacitinib (Clla)** or the IL-6 inhibitor **IV sarilumab (Clla)** can be used in combination with dexamethasone. Sarilumab is only commercially available as a SUBQ injection; see <u>Table 5e</u> for information regarding the preparation of an IV infusion using the SUBQ product.
- ^h Contraindications for the use of therapeutic anticoagulation in patients with COVID-19 include a PLT <50 x 10⁹/L, Hgb <8 g/dL, the need for dual antiplatelet therapy, bleeding within the past 30 days that required an ED visit or hospitalization, a history of a bleeding disorder, or an inherited or active acquired bleeding disorder.
- ⁱ Dexamethasone should be initiated immediately, without waiting until the second immunomodulator can be acquired. If other immunomodulators cannot be obtained or are contraindicated, use dexamethasone alone **(AI)**.
- ^j Examples of patients who may benefit most from adding remdesivir include patients who are immunocompromised **(BIIb)**; patients with evidence of ongoing viral replication (e.g., those with a low Ct value, as measured by an RT-PCR result or with a positive rapid antigen test result) **(BIII)**; or patients who are ≤ 10 days from symptom onset **(CIIa)**. For more information on immunocompromising conditions, see <u>Special Considerations in People Who Are Immunocompromised</u>.
- ^k If PO baricitinib and IV tocilizumab are not available or feasible to use, **PO tofacitinib** can be used instead of PO baricitinib (**Clla**), and **IV sarilumab** can be used instead of IV tocilizumab (**Clla**).

Key: CDC = Centers for Disease Control and Prevention; Ct = cycle threshold; ECMO = extracorporeal membrane oxygenation; ED = emergency department; FDA = Food and Drug Administration; HFNC = high-flow nasal cannula; Hgb = hemoglobin; ICU = intensive care unit; IL = interleukin; IV = intravenous; JAK = Janus kinase; MV = mechanical ventilation; NIV = noninvasive ventilation; the Panel = the COVID-19 Treatment Guidelines Panel; PLT = platelet count; PO = oral; RT-PCR = reverse transcription polymerase chain reaction; SUBQ = subcutaneous; ULN = upper limit of normal

General Management of Nonhospitalized Adults With Acute COVID-19

Last Updated: September 26, 2022

Summary Recommendations
• Management of nonhospitalized patients with acute COVID-19 should include providing supportive care, taking steps to reduce the risk of SARS-CoV-2 transmission (including isolating the patient), and advising patients on when to contact a health care provider and seek an in-person evaluation (AIII).
• When possible, patients with symptoms of COVID-19 should be triaged via telehealth visits to determine whether they require COVID-19-specific therapy and in-person care (AIII).
 Patients who are at high risk of progression to severe COVID-19 may be eligible for pharmacologic therapy. See <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for specific recommendations.
• At a minimum, health care providers should use telehealth to closely follow patients with dyspnea, and in-person monitoring of these patients should be considered (AIII).
• Patients with persistent or progressive dyspnea, especially those who have an oxygen saturation measured by pulse oximetry (SpO ₂) ≤94% on room air at sea level or have symptoms that suggest high acuity (e.g., chest pain or tightness, dizziness, confusion, other mental status changes), should be referred to a health care provider for an in-person evaluation (AIII).
 Clinicians should be aware that using pulse oximeters to measure oxygen saturation has important limitations. Therefore, SpO₂ results should be considered in the context of the patient's clinical condition. See <u>Clinical Spectrum of SARS-CoV-2 Infection</u> for more information.
Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Introduction

This section of the Guidelines is intended to provide general information to health care providers who are caring for nonhospitalized adults with COVID-19. The COVID-19 Treatment Guidelines Panel's (the Panel) recommendations for pharmacologic management can be found in <u>Therapeutic Management</u> of Nonhospitalized Adults With COVID-19. The Panel recognizes that there are times during the COVID-19 pandemic when the distinction between outpatient and inpatient care may be less clear. Patients with COVID-19 may receive care outside traditional ambulatory care or hospital settings if there is a shortage of hospital beds, staff, or resources. In addition, asymptomatic SARS-CoV-2 infection or mild disease may be diagnosed during a patient's hospital admission for a non-COVID-19 condition. Health care providers should use their judgment when deciding whether the guidance offered in this section applies to individual patients.

This section focuses on the evaluation and management of:

- Adults with COVID-19 in an ambulatory care setting
- Adults with COVID-19 following discharge from the emergency department (ED)
- Adults with COVID-19 following inpatient discharge

Outpatient evaluation and management in each of these settings may include some or all of the following: telemedicine, remote monitoring, in-person visits, and home visits by nurses or other health care providers.

Data from the United States show that racial and ethnic minorities experience higher rates of COVID-19, hospitalization, and death.¹⁻⁵ In addition, inequitable receipt of COVID-19 treatments by race, ethnicity,

and socioeconomic status has been reported.⁶⁻⁸ The underlying causes of these observed disparities may include barriers to telehealth visits, transportation challenges, inadequate insurance coverage, a lack of primary care providers, and hesitancy about receiving treatment. To reduce COVID-19 treatment disparities, providers must be aware of the problem and provide patient-centered care. All patient groups must have equal access to treatments, regardless of race, ethnicity, or other minoritized status.

Managing Patients With COVID-19 in an Ambulatory Care Setting

Approximately 80% of patients with COVID-19 who are unvaccinated have mild illness that does not require medical intervention or hospitalization,⁹ and the proportion is likely higher in patients who are vaccinated. Most patients with mild COVID-19 (defined as the absence of viral pneumonia and hypoxemia) can be managed in an ambulatory care setting or at home. Patients with moderate COVID-19 (those with viral pneumonia but without hypoxemia) or severe COVID-19 (those with dyspnea, hypoxemia, or lung infiltrates >50%) need in-person evaluation and close monitoring, as pulmonary disease can progress rapidly and require hospitalization.

When managing outpatients with COVID-19, clinicians should provide supportive care, take steps to reduce the risk of SARS-CoV-2 transmission as recommended by the Centers for Disease Control and Prevention (CDC),^{10,11} and advise patients on when to seek an in-person evaluation.¹² Supportive care includes managing symptoms (as described below), ensuring that patients are receiving the proper nutrition, and being cognizant of the risks of social isolation, particularly for older adults.¹³ Health care providers should identify patients who are at high risk of progression to severe COVID-19. These patients may be candidates for antiviral therapy, including treatment with an anti-SARS-CoV-2 monoclonal antibody (mAb). See <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for more information.

Older patients and those with chronic medical conditions have a higher risk of hospitalization and death. However, SARS-CoV-2 infection may cause severe disease and death in patients of any age, even in the absence of risk factors. In the care of older adults with COVID-19, factors such as cognitive impairment, frailty, the risk of falls, and polypharmacy should be considered. The decision to monitor a patient in the outpatient setting should be made on a case-by-case basis.

Assessing the Need for In-Person Evaluation

When possible, patients with symptoms of COVID-19 should be triaged via telehealth visits to determine whether they require COVID-19-specific therapy and in-person care (**AIII**). Outpatient management may include the use of patient self-assessment tools. During the initial triage, clinic staff should determine which patients are eligible to receive supportive care at home and which patients warrant an in-person evaluation.¹⁴ Local emergency medical services, if called by the patient, may also be helpful when deciding whether an in-person evaluation is indicated.

At a minimum, health care providers should use telehealth to closely follow patients with dyspnea, and in-person monitoring of these patients should be considered (**AIII**). Patients with persistent or progressive dyspnea, especially those who have an oxygen saturation measured by pulse oximetry $(SpO_2) \leq 94\%$ on room air at sea level or have symptoms that suggest high acuity (e.g., chest pain or tightness, dizziness, confusion, other mental status changes), should be referred to a health care provider for an in-person evaluation (**AIII**).

Clinicians who use SpO_2 results when assessing patients must be aware of the important limitations of pulse oximeters and conduct assessments in the context of a patient's clinical condition. See <u>Clinical</u> <u>Spectrum of SARS-CoV-2 Infection</u> for more information.

The criteria used to determine the appropriate clinical setting for an in-person evaluation may vary by location and institution. It may also change over time as new data and treatment options emerge. There should be a low threshold for in-person evaluation of older people and those with medical conditions associated with an increased risk of progression to severe COVID-19. Individuals who perform the initial triage should use their clinical judgment to determine whether patients require ambulance transport.

In some settings where clinical evaluation is challenged by geography, health care provider home visits may be used to evaluate patients.¹⁵ Patients who are homeless should be provided with housing where they can adequately self-isolate. Providers should be aware of the potential adverse effects of prolonged social isolation, including depression and anxiety.¹³ All outpatients should receive instructions regarding self-care, isolation, and follow-up, and they should be advised to contact a health care provider or a local ED for any worsening symptoms.¹¹

Clinical Considerations When Managing Patients in an Ambulatory Care Setting

Patients with SARS-CoV-2 infection may be asymptomatic or experience symptoms that are indistinguishable from other acute viral or bacterial infections (e.g., fever, cough, sore throat, malaise, muscle pain, headache, gastrointestinal symptoms). People who have symptoms compatible with COVID-19 should undergo diagnostic SARS-CoV-2 testing (see <u>Testing for SARS-CoV-2 Infection</u>). Considering other possible etiologies of symptoms is important, including other respiratory viral infections (e.g., influenza), community-acquired pneumonia, congestive heart failure, asthma or chronic obstructive pulmonary disease exacerbations, and streptococcal pharyngitis.

Although mild dyspnea is common, worsening dyspnea and severe chest pain or tightness suggest the development or progression of pulmonary involvement. In studies of patients who developed acute respiratory distress syndrome, progression occurred a median of 2.5 days after the onset of dyspnea.¹⁶⁻¹⁸ At a minimum, health care providers should use telehealth to closely follow patients with dyspnea, and in-person monitoring of these patients should be considered (**AIII**). Patients with persistent or progressive dyspnea, especially those who have an SpO₂ \leq 94% on room air at sea level or have symptoms that suggest high acuity (e.g., chest pain or tightness, dizziness, confusion, other mental status changes), should be referred to a health care provider for an in-person evaluation (**AIII**).

If an adult patient has access to a pulse oximeter at home, SpO_2 measurements can be used to help assess overall clinical status. Patients should be advised to use pulse oximeters on warm fingers rather than cold fingers for better accuracy. Patients should inform their health care providers if the value is repeatedly below 95% on room air at sea level. Pulse oximetry may not accurately detect hypoxemia, especially in patients who have dark skin pigmentation.^{19,20}

Not all commercially available pulse oximeters have been cleared by the Food and Drug Administration (FDA). SpO₂ readings obtained through non-FDA-cleared devices, such as over-the-counter sports oximeters or mobile phone applications, lack sufficient accuracy for clinical use.^{21,22} Abnormal readings on these devices should be confirmed with an FDA-cleared device or an arterial blood gas analysis. Importantly, SpO₂ readings should only be interpreted within the context of a patient's entire clinical presentation (i.e., results should be disregarded if a patient is complaining of increasing dyspnea). See <u>Clinical Spectrum of SARS-CoV-2 Infection</u> for more information regarding the limitations of pulse oximetry.

Counseling Regarding the Need for Follow-Up

Health care providers should identify patients who are at high risk of disease progression. These patients may be candidates for antiviral therapy, including treatment with an anti-SARS-CoV-2 mAb (see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>). Clinicians should ensure that

COVID-19 Treatment Guidelines

these patients receive adequate medical follow-up. The frequency and duration of follow-up will depend on the risk for severe disease, the severity of symptoms, and the patient's ability to self-report worsening symptoms. Health care providers should determine whether a patient has access to a phone, computer, or tablet for telehealth; whether they have adequate transportation for clinic visits; and whether they have regular access to food. The clinician should also confirm that the patient has a caregiver who can assist with daily activities if needed.

All patients and/or their family members or caregivers should be counseled about the warning symptoms that should prompt re-evaluation through a telehealth visit or an in-person evaluation in an ambulatory care setting or ED. These symptoms include new onset of dyspnea; worsening dyspnea (particularly if dyspnea occurs while resting or if it interferes with daily activities); dizziness; and mental status changes, such as confusion.

Managing Adults With COVID-19 Following Discharge From the Emergency Department

There are no fixed criteria for admitting patients with COVID-19 to the hospital; criteria may vary by region and hospital facility. Patients with severe disease are typically admitted to the hospital. Rarely, a patient with severe disease may not be admitted due to a high prevalence of infection and limited hospital resources. In addition, patients who could receive appropriate care at home but are unable to be adequately managed in their usual residential setting are candidates for temporary shelter in supervised facilities, such as a COVID-19 alternative care facility.²³ For example, patients who are living in multigenerational households or are homeless may not be able to self-isolate and should be provided resources such as dedicated housing units or hotel rooms, when available. Unfortunately, dedicated residential care facilities for patients with COVID-19 are not widely available, and community-based solutions for self-care and isolation should be explored.

Treatment with an antiviral agent or anti-SARS-CoV-2 mAb is recommended for patients with mild to moderate COVID-19 who are not on supplemental oxygen and are at high risk of clinical progression (see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>).

In rare cases where institutional resources (e.g., inpatient beds, staff members) are scarce, it may be necessary to discharge an adult patient and provide an advanced level of home care, including supplemental oxygen (if indicated), pulse oximetry, and close follow-up. Although early discharge of patients with severe disease is not generally recommended by the Panel, it is recognized that these management strategies are sometimes necessary. In these situations, some institutions have provided frequent telemedicine follow-up visits for these patients or a hotline that allows patients to speak with a clinician when necessary. Home resources should be assessed before a patient is discharged from the ED. Outpatients should have a caregiver and access to a device suitable for telehealth. Patients and/ or their family members or caregivers should be counseled about the warning symptoms that should prompt re-evaluation by a health care provider.

If a patient is not being admitted to the hospital, the Panel **recommends against** the use of anticoagulants and antiplatelet therapy in the ED for the prevention of venous thromboembolism (VTE) or arterial thrombosis, except in a clinical trial (**AIIa**). This recommendation does not apply to patients with other indications for antithrombotic therapy. For more information, see <u>Antithrombotic Therapy in</u> <u>Patients With COVID-19</u>. Patients should be encouraged to ambulate, and activity should be increased according to the patient's tolerance.

Managing Adults With COVID-19 Following Hospital Discharge

Most patients who are discharged from the hospital setting should have a follow-up visit with a health *COVID-19 Treatment Guidelines*

care provider soon after discharge. Whether an in-person or telehealth visit is most appropriate depends on the clinical and social situation. In some cases, adult patients are deemed to be stable for discharge from the inpatient setting, although they still require supplemental oxygen. Special consideration may be given to continuing certain therapeutics (e.g., dexamethasone) in this setting. For more information, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>. When possible, these individuals should receive oximetry monitoring and close follow-up through telehealth visits, visiting nurse services, or in-person clinic visits.

The Panel **recommends against** routinely continuing VTE prophylaxis after hospital discharge for patients with COVID-19 unless they have another indication or are participating in a clinical trial (**AIII**). For more information, see <u>Antithrombotic Therapy in Patients With COVID-19</u>. Patients should be encouraged to ambulate, and activity should be increased according to the patient's tolerance.

Considerations in Pregnancy

Managing pregnant outpatients with COVID-19 is similar to managing nonpregnant patients (see <u>Special</u> <u>Considerations in Pregnancy</u>). Clinicians should offer supportive care and therapeutics as indicated, take steps to reduce the risk of SARS-CoV-2 transmission, and provide guidance on when to seek an in-person evaluation. The American College of Obstetricians and Gynecologists (ACOG) has published recommendations on how to use telehealth for prenatal care and how to modify routine prenatal care when necessary to decrease the risk of SARS-CoV-2 transmission to patients, caregivers, and staff.²⁴

In pregnant patients, SpO₂ should be maintained at 95% or above on room air at sea level; therefore, the threshold for monitoring pregnant patients in an inpatient setting may be lower than in nonpregnant patients. At this time, there are no changes to fetal monitoring recommendations in the outpatient setting, and fetal surveillance and management should be similar to the fetal surveillance and management used for pregnant patients with medical illness.^{25,26} However, these monitoring strategies can be discussed on a case-by-case basis with an obstetrician. Pregnant and lactating patients should be given the opportunity to participate in clinical trials of outpatients with COVID-19 to help inform decision-making in this population.

References

- 1. Azar KMJ, Shen Z, Romanelli RJ, et al. Disparities in outcomes among COVID-19 patients in a large health care system in California. *Health Aff (Millwood)*. 2020;39(7):1253-1262. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32437224</u>.
- 2. Gold JAW, Wong KK, Szablewski CM, et al. Characteristics and clinical outcomes of adult patients hospitalized with COVID-19—Georgia, March 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(18):545-550. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32379729</u>.
- 3. Gross CP, Essien UR, Pasha S, et al. Racial and ethnic disparities in population-level COVID-19 mortality. *J Gen Intern Med.* 2020;35(10):3097-3099. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32754782</u>.
- 4. Islam SJ, Nayak A, Hu Y, et al. Temporal trends in the association of social vulnerability and race/ethnicity with county-level COVID-19 incidence and outcomes in the USA: an ecological analysis. *BMJ Open*. 2021;11(7):e048086. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34301657.
- Price-Haywood EG, Burton J, Fort D, Seoane L. Hospitalization and mortality among Black patients and White patients with COVID-19. *N Engl J Med*. 2020;382(26):2534-2543. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32459916</u>.
- 6. Azam TU, Berlin H, Anderson E, et al. Differences in inflammation, treatment, and outcomes between Black and non-Black patients hospitalized for COVID-19: a prospective cohort study. *Am J Med.* 2022;135(3):360-368. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34793753</u>.
- 7. Wiltz JL, Feehan AK, Molinari NM, et al. Racial and ethnic disparities in receipt of medications for treatment

of COVID-19—United States, March 2020–August 2021. *MMWR Morb Mortal Wkly Rep.* 2022;71(3):96-102. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35051133</u>.

- Wu EL, Kumar RN, Moore WJ, et al. Disparities in COVID-19 monoclonal antibody delivery: a retrospective cohort study. *J Gen Intern Med*. 2022;37(10):2505-2513. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35469360</u>.
- Stokes EK, Zambrano LD, Anderson KN, et al. Coronavirus disease 2019 case surveillance—United States, January 22–May 30, 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(24):759-765. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32555134</u>.
- 10. Centers for Disease Control and Prevention. How to protect yourself and others. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/prevent-getting-sick/prevention.html</u>. Accessed November 7, 2022.
- 11. Centers for Disease Control and Prevention. If you are sick or caring for someone. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/if-you-are-sick</u>. Accessed September 15, 2022.
- 12. Cheng A, Caruso D, McDougall C. Outpatient management of COVID-19: rapid evidence review. *Am Fam Physician*. 2020;102(8):478-486. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33064422</u>.
- Morrow-Howell N, Galucia N, Swinford E. Recovering from the COVID-19 pandemic: a focus on older adults. J Aging Soc Policy. 2020;32(4-5):526-535. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32336225</u>.
- 14. Centers for Disease Control and Prevention. Symptoms of COVID-19. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/symptoms-testing/symptoms.html</u>. Accessed August 23, 2023.
- 15. Close RM, Stone MJ. Contact tracing for Native Americans in rural Arizona. *N Engl J Med.* 2020;383(3):e15. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32672426</u>.
- Wang D, Hu B, Hu C, et al. Clinical characteristics of 138 hospitalized patients with 2019 novel coronavirusinfected pneumonia in Wuhan, China. *JAMA*. 2020;323(11):1061-1069. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32031570</u>.
- Arentz M, Yim E, Klaff L, et al. Characteristics and outcomes of 21 critically ill patients with COVID-19 in Washington state. *JAMA*. 2020;323(16):1612-1614. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32191259</u>.
- Yang X, Yu Y, Xu J, et al. Clinical course and outcomes of critically ill patients with SARS-CoV-2 pneumonia in Wuhan, China: a single-centered, retrospective, observational study. *Lancet Respir Med.* 2020;8(5):475-481. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32105632</u>.
- Luks AM, Swenson ER. Pulse oximetry for monitoring patients with COVID-19 at home. Potential pitfalls and practical guidance. *Ann Am Thorac Soc.* 2020;17(9):1040-1046. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32521167</u>.
- 20. Sjoding MW, Dickson RP, Iwashyna TJ, Gay SE, Valley TS. Racial bias in pulse oximetry measurement. *N Engl J Med.* 2020;383(25):2477-2478. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33326721</u>.
- 21. Harskamp RE, Bekker L, Himmelreich JCL, et al. Performance of popular pulse oximeters compared with simultaneous arterial oxygen saturation or clinical-grade pulse oximetry: a cross-sectional validation study in intensive care patients. *BMJ Open Respir Res.* 2021;8(1):e000939. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34489238</u>.
- 22. Lipnick MS, Feiner JR, Au P, Bernstein M, Bickler PE. The accuracy of 6 inexpensive pulse oximeters not cleared by the Food and Drug Administration: the possible global public health implications. *Anesth Analg.* 2016;123(2):338-45. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27089002</u>.
- 23. Meyer GS, Blanchfield BB, Bohmer RMJ, Mountford MB, Venderwagen WC. Alternative care sites for the COVID-19 pandemic: the early U.S. and U.K. experience. *NEJM Catalyst*. 2020. Available at: <u>https://catalyst.nejm.org/doi/full/10.1056/CAT.20.0224</u>.
- 24. American College of Obstetricians and Gynecologists. COVID-19 FAQs for obstetrician-gynecologists, telehealth. 2020. Available at: <u>https://www.acog.org/clinical-information/physician-faqs/covid-19-faqs-for-ob-</u>

gyns-telehealth. Accessed September 20, 2022.

- 25. Society for Maternal-Fetal Medicine. Management considerations for pregnant patients with COVID-19. 2021. Available at: <u>https://s3.amazonaws.com/cdn.smfm.org/media/2734/SMFM_COVID_Management_of_COVID_pos_preg_patients_2-2-21_(final).pdf</u>.
- 26. American College of Obstetricians and Gynecologists. COVID-19 FAQs for obstetrician-gynecologists, obstetrics. 2022. Available at: <u>https://www.acog.org/clinical-information/physician-faqs/covid-19-faqs-for-ob-gyns-obstetrics</u>. Accessed November 25, 2022.

Therapeutic Management of Nonhospitalized Adults With COVID-19

Last Updated: November 2, 2023

Symptom management should be initiated for all nonhospitalized adults with mild to moderate COVID-19. For adults who are at high risk of progression to severe disease, several antiviral therapeutic options are available to reduce the risk of hospitalization or death. The COVID-19 Treatment Guidelines Panel's (the Panel) recommendations on the use of these drugs for the treatment of COVID-19 are outlined in this section.

The main goal of therapeutic management for nonhospitalized patients is to prevent progression to severe disease, hospitalization, or death. Other goals may include accelerating symptom recovery, viral clearance, and prevention of long-term sequelae. Current data on the impact of therapy on these secondary goals are limited.

Several factors affect the selection of the best treatment option for a specific patient. These factors include the clinical efficacy and availability of the treatment option, the feasibility of administering parenteral medications, the potential for significant drug-drug interactions, the patient's pregnancy status, the time from symptom onset, and the in vitro activities of the available products against the currently circulating SARS-CoV-2 variants and subvariants.

Most of the data that support the use of the recommended treatment options come from clinical trials that enrolled individuals who were at high risk of disease progression and who had no pre-existing immunity from COVID-19 vaccination or prior SARS-CoV-2 infection. Accordingly, the proportion of hospitalizations and deaths in the placebo arms of these trials was high compared to what has been seen more recently in populations where most people are vaccinated or have had prior SARS-CoV-2 infection. Although these trials demonstrated the efficacy of using antiviral drugs in high-risk populations, it is difficult to know their precise effectiveness in the current setting because of the low rates of hospitalization and death among those who have been vaccinated.

Nevertheless, some patients continue to have an increased risk of disease progression, and it is in those people that therapies are most likely to be beneficial. Patients who are at the highest risk are older patients (i.e., those aged >50 years and especially those aged \geq 65 years) and patients who are unlikely to have an adequate immune response to COVID-19 vaccines due to a moderate to severe immunocompromising condition or the receipt of immunosuppressive medications. Other risk factors include lack of vaccination or incomplete vaccination; a prolonged amount of time since the most recent vaccine dose (e.g., >6 months); and conditions such as obesity, diabetes, and chronic respiratory, cardiac, or kidney disease.¹

People who are members of racial and ethnic minority groups have higher rates of hospitalization and death from COVID-19 than people who are White.² Disparities in the use of antiviral treatments in patients who are not White have been reported; therefore, attention to equitable access is critical.^{3,4}

The Panel's recommendations reflect the available data on the benefits of using antiviral therapies to prevent progression to severe COVID-19. The Panel will consider the potential benefits of available therapies for other outcomes, such as symptom recovery, as those data emerge.

Table 2a outlines the Panel's recommendations for the therapeutic management of nonhospitalized adults with COVID-19. For recommended doses of the agents listed in Table 2a, see <u>Table 4e</u>.

Table 2a. Therapeutic Management of Nonhospitalized Adults With Mild to Moderate COVID-19Who Do Not Require Supplemental Oxygen

Patient Disposition	Panel's Recommendations	
All Patients	 Symptom management should be initiated for all patients (AIII). The Panel recommends against the use of dexamethasone^a or other systemic corticosteroids in the absence of another indication (AIIb). 	
Patients Who Are at High Risk of Progressing to Severe COVID-19 ^{6,c}	 Preferred therapies. Listed in order of preference: Ritonavir-boosted nirmatrelvir (Paxlovid)^d (Alla); see footnote on drug interactions^e Remdesivir^{d,f} (Blla) 	
	Alternative therapy. For use when the preferred therapies are not available, feasible to use, or clinically appropriate: • Molnupiravir ^{d,g,h} (Clla)	
Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.		

^a There is currently a lack of safety and efficacy data on the use of dexamethasone in outpatients with COVID-19. Using systemic glucocorticoids in outpatients with COVID-19 may cause harm.

- ^b For a list of risk factors, see the CDC webpage <u>Underlying Medical Conditions Associated With Higher Risk for Severe</u> <u>COVID-19</u>. When deciding whether to prescribe antiviral treatment to a patient who has been vaccinated, clinicians should be aware of the conditions associated with a high risk of disease progression. These conditions include older age, a prolonged amount of time since the most recent vaccine dose (e.g., >6 months), and a decreased likelihood of an adequate immune response to vaccination due to a moderate to severe immunocompromising condition or the receipt of immunosuppressive medications. The number and severity of risk factors also affects the level of risk.
- ^c For a discussion of potential treatment options for patients who are immunocompromised and have prolonged COVID-19 symptoms and evidence of ongoing viral replication, see below and <u>Special Considerations in People Who Are</u> <u>Immunocompromised</u>.
- ^d If a patient requires hospitalization after starting treatment, the full treatment course can be completed at the health care provider's discretion.
- ^e Ritonavir-boosted nirmatrelvir has significant drug-drug interactions. Clinicians should carefully review a patient's concomitant medications and evaluate potential drug-drug interactions. See <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications</u> for more information.
- ^f Administration of remdesivir requires an IV infusion once daily for 3 days.
- ⁹ Molnupiravir appears to have lower efficacy than the other options recommended by the Panel. Therefore, it should be considered when the other options are not available, feasible to use, or clinically appropriate.
- ^h The Panel **recommends against** the use of **molnupiravir** for the treatment of COVID-19 in pregnant patients unless there are no other options and therapy is clearly indicated **(AIII)**.

Key: CDC = Centers for Disease Control and Prevention; IV = intravenous; the Panel = the COVID-19 Treatment Guidelines Panel

Symptom Management

Treatment of symptoms includes using over-the-counter antipyretics, analgesics, or antitussives for fever, headache, myalgias, and cough. Patients should be advised to drink fluids regularly to avoid dehydration. Rest is recommended as needed during the acute phase of COVID-19, and ambulation and other forms of activity should be increased according to the patient's tolerance. Patients should be educated about the variability in time to symptom resolution and complete recovery. When possible, patients with symptoms of COVID-19 should be triaged via telehealth visits to determine whether they require COVID-19–specific therapy and in-person care (**AIII**).

At a minimum, health care providers should use telehealth to closely follow patients with dyspnea, and in-person monitoring of these patients should be considered (**AIII**). Patients with persistent or progressive dyspnea, especially those who have an oxygen saturation measured by pulse oximetry \leq 94% on room air at sea level or have symptoms that suggest high acuity (e.g., chest pain or tightness, dizziness, confusion, other mental status changes), should be referred to a health care provider for an in-person evaluation (**AIII**).

Rationale for the Panel's Recommendations

The Panel's recommendations for the antiviral agents that are used to treat nonhospitalized patients with mild to moderate COVID-19 who are at high risk of clinical progression are based on the results of clinical trials. The Panel **recommends against** using anti-SARS-CoV-2 monoclonal antibodies for the treatment of COVID-19 (**AIII**) because the dominant Omicron subvariants in the United States are not expected to be susceptible to these products. See <u>Anti-SARS-CoV-2 Monoclonal Antibodies</u> for more information.

The Panel favors the use of ritonavir-boosted nirmatrelvir (Paxlovid) in most high-risk, nonhospitalized patients with mild to moderate COVID-19. When ritonavir-boosted nirmatrelvir is not clinically appropriate (e.g., because of significant drug-drug interactions), the Panel recommends using remdesivir. Ritonavir-boosted nirmatrelvir has high efficacy; has been shown to reduce hospitalization and death when administered to high-risk, unvaccinated, nonhospitalized patients within 5 days of symptom onset;⁵ and is an oral medication, whereas remdesivir requires intravenous (IV) administration.

The Panel's recommendation for remdesivir is based on a Phase 3, randomized, placebo-controlled trial that reported high clinical efficacy in high-risk, nonhospitalized patients with COVID-19 who were unvaccinated.⁶ However, in some settings, daily IV administration of remdesivir for 3 days may be a logistical challenge.

The Panel recommends **molnupiravir** as a therapeutic option when the other recommended antiviral treatment options are not available, feasible to use, or clinically appropriate (**CIIa**). Molnupiravir appears to have lower clinical efficacy than the other treatment options, although no randomized studies have compared these therapies directly. The rationale for each of the Panel's recommendations is discussed below.

Currently, data on the use of combinations of antiviral agents for the treatment of COVID-19 are limited. Clinical trials are needed to determine whether combination therapy has a role in the treatment of COVID-19.

Strategies for the Use of Ritonavir-Boosted Nirmatrelvir

Because ritonavir is a strong cytochrome P450 3A4 inhibitor and a P-glycoprotein inhibitor, it may increase blood concentrations of certain concomitant medications and increase the potential for serious drug toxicities. Therefore, the Food and Drug Administration (FDA) prescribing information includes a boxed warning for significant drug-drug interactions with ritonavir-boosted nirmatrelvir.⁷ Clinicians should consider both the potential benefits of treatment with ritonavir-boosted nirmatrelvir and the potential risks related to drug-drug interactions.

Many drug-drug interactions between ritonavir-boosted nirmatrelvir and concomitant medications **can be safely managed** (e.g., with certain statins, calcium channel blockers, or direct oral anticoagulants). If a significant drug-drug interaction is identified, prescribers should consider consulting with a pharmacist.

The following resources are available to assist in identifying and managing drug-drug interactions:

• Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant

COVID-19 Treatment Guidelines

Medications

- The Liverpool COVID-19 Drug Interactions website
- The <u>University of Waterloo/University of Toronto drug interaction guide</u>
- The FDA prescribing information for ritonavir-boosted nirmatrelvir

The use of ritonavir-boosted nirmatrelvir may be challenging in patients with severe renal impairment and in patients receiving certain transplant-related immunosuppressants or chemotherapy. The FDA prescribing information states that until more data are available, ritonavir-boosted nirmatrelvir is not recommended in patients with an estimated glomerular filtration rate (eGFR) of <30 mL/min.⁷ Although data on dose adjustments are limited, some groups have proposed dosing adjustments in patients with an eGFR of <30 mL/min or for patients receiving hemodialysis.⁸⁻¹¹

The decision to prescribe ritonavir-boosted nirmatrelvir to patients receiving calcineurin and mammalian target of rapamycin inhibitors should always be made in consultation with the patient's specialist providers. Among reports submitted to the FDA Adverse Events Reporting System, the most commonly reported concomitant medications resulting in serious adverse reactions, including fatal events, were calcineurin inhibitors (e.g., tacrolimus).¹² Ritonavir-boosted nirmatrelvir may be prescribed to select patients if an expert in managing the interaction is available and close therapeutic drug monitoring is logistically feasible. Otherwise, an alternative therapy for COVID-19 should be considered. See the American Society of Transplantation statement for additional information.

Interactions between ritonavir-boosted nirmatrelvir and chemotherapeutic agents should also be managed in consultation with the patient's specialist providers. For guidance on managing these interactions, refer to the FDA prescribing information for ritonavir-boosted nirmatrelvir and the prescribing information for the chemotherapeutic agent.⁷ The <u>University Health Network/Kingston</u> <u>Health Sciences Centre</u> provides an additional resource for evaluating drug-drug interactions between ritonavir-boosted nirmatrelvir and chemotherapeutic agents.

Strategies for the Use of Remdesivir

Advanced planning (e.g., reserving infusion slots, identifying alternative infusion sites) may be needed to increase access to IV remdesivir. IV remdesivir can be administered in skilled nursing facilities, home health care settings, and outpatient facilities such as infusion centers. If treatment facilities cannot provide a 3-day course of remdesivir IV infusions to all eligible patients, prioritizing patients who will benefit the most from the therapy becomes necessary. The prioritization scheme below is based on 4 key elements: age, vaccination status, immune status, and clinical risk factors. For a list of risk factors, see the Centers for Disease Control and Prevention (CDC) webpage <u>Underlying Medical Conditions Associated With Higher Risk for Severe COVID-19</u>. The groups are listed by tier in descending order of priority.

Tier	Risk Group
-	 Immunocompromised individuals not expected to mount an adequate immune response to COVID-19 vaccination or SARS-CoV-2 infection due to their underlying conditions,^a regardless of vaccine status; or
	 Unvaccinated individuals at the highest risk of severe disease (anyone aged ≥75 years or anyone aged ≥65 years with additional risk factors).
2	 Unvaccinated individuals not included in Tier 1 who are at risk of severe disease (anyone aged ≥65 years or anyone aged <65 years with clinical risk factors)
3	 Vaccinated individuals at risk of severe disease (anyone aged ≥65 years or anyone aged <65 years with clinical risk factors)^b

^a See the CDC website <u>COVID-19 Vaccines for People Who Are Moderately or Severely Immunocompromised</u> for a discussion

COVID-19 Treatment Guidelines

of immunocompromising conditions.

^b Vaccinated individuals who are not up to date with their immunizations are likely at higher risk for severe disease; patients within this tier who are in this situation should be prioritized for treatment. See the CDC webpage <u>Stay Up to Date with</u> <u>COVID-19 Vaccines</u> for more information.

See <u>Prioritization of Anti-SARS-CoV-2 Therapies for the Treatment of COVID-19 in Nonhospitalized</u> <u>Patients When There Are Logistical Constraints</u> for more information.

Patients Who Are Immunocompromised and Have Prolonged Symptoms and Evidence of Ongoing Viral Replication

For patients who are immunocompromised and have prolonged COVID-19 symptoms and evidence of ongoing viral replication (e.g., those with a low cycle threshold value, as measured by a reverse transcription polymerase chain reaction result or with a positive rapid antigen test result) despite receiving a course of antiviral therapy, the optimal management is unknown. Case reports and case series have documented the treatment of these patients with additional antiviral treatments, prolonged courses of antiviral treatments, high-titer COVID-19 convalescent plasma (CCP), or combination therapy.¹³⁻¹⁷ The data for these approaches are not definitive, but some Panel members would use 1 or more of the following treatment options:

- Longer and/or additional courses of ritonavir-boosted nirmatrelvir
- Longer and/or additional courses of remdesivir
- High-titer CCP from a vaccinated donor who recently recovered from COVID-19 likely caused by a SARS-CoV-2 variant similar to the variant causing the patient's illness

For further discussion of these potential treatment options, see <u>Special Considerations in People Who</u> <u>Are Immunocompromised</u>.

Additional Information on Ritonavir-Boosted Nirmatrelvir

Nirmatrelvir is an orally bioavailable protease inhibitor that is active against M^{PRO}, a viral protease that plays an essential role in viral replication.¹⁸ The FDA has approved ritonavir-boosted nirmatrelvir for the treatment of mild to moderate COVID-19 in nonhospitalized adults who are at high risk of progressing to severe COVID-19.⁷

Patients should complete the 5-day treatment course of ritonavir-boosted nirmatrelvir, which was shown to be efficacious in the EPIC-HR trial.⁵ If a patient requires hospitalization after starting treatment, the full 5-day treatment course of ritonavir-boosted nirmatrelvir should be completed unless there are drug-drug interactions that preclude its use.

In the EPIC-HR trial, ritonavir-boosted nirmatrelvir reduced the risk of hospitalization or death by 89% compared to placebo in unvaccinated, nonhospitalized adults with laboratory-confirmed SARS-CoV-2 infection.⁵ This efficacy is comparable to the efficacies reported in similar patient populations for remdesivir (87% relative reduction)^{5,7} and greater than the efficacy reported for molnupiravir in this setting (31% relative reduction).⁶

Because ritonavir-boosted nirmatrelvir has the potential for significant drug-drug interactions with concomitant medications, this regimen may not be the optimal choice for all patients (see <u>Drug-Drug</u><u>Interactions Between Ritonavir-Boosted Nirmatrelvir [Paxlovid] and Concomitant Medications</u>). However, because ritonavir-boosted nirmatrelvir is the only highly effective oral antiviral available for the treatment of COVID-19, drug-drug interactions that can be safely managed should not preclude the use of this medication.

For more information on the use of ritonavir-boosted nirmatrelvir, see <u>Ritonavir-Boosted Nirmatrelvir</u> (<u>Paxlovid</u>). See Viral Rebound and Symptom Recurrence below for information regarding SARS-CoV-2 viral rebound in patients who have completed treatment with ritonavir-boosted nirmatrelvir.

Additional Information on Remdesivir

Remdesivir is a nucleotide prodrug of an adenosine analog that inhibits SARS-CoV-2 replication. It is approved by the FDA for the treatment of COVID-19 in adults and children aged \geq 28 days and weighing \geq 3 kg who are hospitalized with COVID-19 and for those with mild to moderate COVID-19 who are not hospitalized and are at high risk of progressing to severe disease. In the PINETREE trial, nonhospitalized patients with mild to moderate COVID-19 who were unvaccinated and at high risk of progressing to severe disease received 3 days of IV remdesivir or placebo. Use of remdesivir resulted in an 87% relative reduction in the risk of hospitalization or death.¹⁹⁻²¹

Remdesivir should be administered in a setting where severe hypersensitivity reactions, such as anaphylaxis, can be managed. Patients should be monitored during the infusion and observed for at least 1 hour after the infusion as clinically appropriate.

For more information, see Remdesivir.

Additional Information on Molnupiravir

Molnupiravir is the oral prodrug of beta-D-N4-hydroxycytidine, a ribonucleoside that has shown antiviral activity against SARS-CoV-2 in vitro and in clinical trials.²²⁻²⁴ The FDA issued an Emergency Use Authorization for molnupiravir for the treatment of mild to moderate COVID-19 in nonhospitalized patients aged ≥ 18 years who are at high risk of disease progression and for whom alternative treatment options are not accessible or clinically appropriate.

The MOVe-OUT trial enrolled nonhospitalized adults who were unvaccinated and at high risk of progression to severe disease in the pre-Omicron era. The study found that molnupiravir reduced the rate of hospitalization or death by 31% compared to placebo.²⁵ A secondary analysis of MOVe-OUT trial data revealed that patients who received molnupiravir and progressed to hospitalization were less likely to need respiratory interventions than patients who received placebo and progressed to hospitalization.²⁶

The PANORAMIC trial enrolled participants during a period when the Omicron variant was circulating.²⁷ The participants were nonhospitalized adults with COVID-19 who were at high risk of progressing to severe disease, and 94% had received at least 3 doses of a COVID-19 vaccine. The study found that the use of molnupiravir plus usual care did not reduce the primary composite outcome of hospitalization or death compared to usual care alone. The rates of this composite outcome were low (1%) in both arms. Molnupiravir plus usual care was superior to usual care alone for several secondary clinical endpoints. For example, patients who received molnupiravir plus usual care reported recovering from COVID-19 an estimated 4 days earlier than those who received usual care alone. However, because the PANORAMIC trial was an open-label study and the patients knew whether they were receiving molnupiravir or not, this may have affected their reported symptoms. As a result, these findings are less reliable than those from a placebo-controlled trial.

Although the different COVID-19 treatment options have not been directly compared in clinical trials, the Panel recommends using molnupiravir as an alternative therapy when ritonavir-boosted nirmatrelvir and remdesivir are not available, feasible to use, or clinically appropriate, because molnupiravir appears to have lower clinical efficacy than these other options.

Molnupiravir is a mutagenic ribonucleoside antiviral agent, and there is a theoretical risk that the drug will be metabolized by the human host cell and incorporated into the host DNA, leading to mutations.

The available genotoxicity data and the 5-day duration of treatment led the FDA to conclude that molnupiravir has a low risk for genotoxicity.²⁸

The Panel **recommends against** the use of **molnupiravir** for the treatment of COVID-19 in pregnant patients unless there are no other options and therapy is clearly indicated (**AIII**). People who engage in sexual activity that may result in conception should use effective contraception during and following treatment with molnupiravir.

Fetal toxicity has been reported in animal studies of molnupiravir.²⁸ However, when other therapies are not available, pregnant patients with COVID-19 who are at high risk of progressing to severe disease may reasonably choose molnupiravir after being fully informed of the risks, particularly if they are beyond the time of embryogenesis (i.e., >10 weeks' gestation). See <u>Pregnancy, Lactation, and COVID-19 Therapeutics</u> for more information.

For more information, see Molnupiravir.

Viral Rebound and Symptom Recurrence

Observational studies and the EPIC-HR and MOVe-OUT trials have described SARS-CoV-2 viral rebound and the recurrence of COVID-19 symptoms in some patients who have completed treatment with ritonavir-boosted nirmatrelvir or molnupiravir.^{7,28-31} The frequency, mechanism, and clinical implications of these events are unclear. Viral rebound and the recurrence of COVID-19 symptoms can also occur in the absence of treatment.^{7,29-32}

To date, the recurrence of COVID-19 symptoms and virus detection following the use of antiviral therapies has not been associated with progression to severe COVID-19. Therefore, concerns about the recurrence of symptoms or viral rebound **should not** be a reason to avoid using antiviral therapies.^{28,33-35} There are insufficient data on whether a longer course of ritonavir-boosted nirmatrelvir or molnupiravir will prevent viral rebound or symptom recurrence. There also are insufficient data on the efficacy of administering a second course of antiviral therapy to treat viral rebound or symptom recurrence.

Immunomodulators

The Panel **recommends against** the use of **dexamethasone** or other systemic corticosteroids to treat outpatients with mild to moderate COVID-19 who do not require hospitalization or supplemental oxygen (**AIIb**). Patients with COVID-19 who are receiving **dexamethasone** or another corticosteroid for an underlying condition should continue this therapy as directed by their health care provider (**AIII**).

Medicare and FDA data show a significant increase in the number of prescriptions for systemic corticosteroids among nonhospitalized patients with COVID-19³⁶ despite a lack of safety and efficacy data on the use of systemic corticosteroids in this setting. Systemic glucocorticoids may cause harm in nonhospitalized patients with COVID-19. Results from 1 randomized controlled trial and 1 observational cohort study did not demonstrate a clinical benefit of dexamethasone among hospitalized patients who did not require supplemental oxygen,³⁷ and dexamethasone may potentially cause harm in these patients.³⁸

In the RECOVERY trial, the use of dexamethasone had no effect on mortality among hospitalized patients with COVID-19 who did not require supplemental oxygen (rate ratio 1.19; 95% CI, 0.91– 1.55).³⁷ A large observational study of patients at Veterans Affairs hospitals reported no survival benefit for dexamethasone among patients with COVID-19 who did not require supplemental oxygen. Instead, these patients had an increased risk of 90-day mortality (HR 1.76; 95% CI, 1.47–2.12).³⁷ However, hospitalized patients with COVID-19 are likely to have an increased risk of mortality compared to nonhospitalized patients, which is a limitation of observational trial data.

COVID-19 Treatment Guidelines

Concomitant Medication Management

In general, a patient's usual medication and/or supplement regimen should be continued after the diagnosis of COVID-19 (see <u>Considerations for Using Concomitant Medications in Patients With</u> <u>COVID-19</u>). Angiotensin-converting enzyme (ACE) inhibitors; angiotensin receptor blockers (ARBs); statin therapy; nonsteroidal anti-inflammatory drugs; and oral, inhaled, and intranasal corticosteroids that are prescribed for comorbid conditions should be continued as directed (**AIIa** for ACE inhibitors and ARBs; **AIII** for other medications). Patients should be advised to avoid the use of nebulized medications in the presence of others to avoid potential aerosolization of SARS-CoV-2.³⁹ In patients with HIV, antiretroviral therapy should not be switched or adjusted for the theoretical purpose of preventing or treating SARS-CoV-2 infection (**AIII**). For more information, see <u>Special Considerations</u> in <u>People With HIV</u>.

When a patient is receiving an immunomodulating medication, the prescribing clinician or an expert in the subspecialty should be consulted about the risks and benefits associated with a temporary dose reduction or discontinuation. These risks and benefits will depend on the medication's indication and the severity of the underlying condition (see <u>Special Considerations in People Who Are</u> <u>Immunocompromised</u>).

Before prescribing ritonavir-boosted nirmatrelvir, clinicians should carefully review the patient's concomitant medications, including over-the-counter medications and herbal supplements, to evaluate potential drug-drug interactions.

References

- 1. Centers for Disease Control and Prevention. Underlying medical conditions associated with higher risk for severe COVID-19: information for healthcare professionals. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/clinical-care/underlyingconditions.html</u>. Accessed October 17, 2023.
- 2. Mackey K, Ayers CK, Kondo KK, et al. Racial and ethnic disparities in COVID-19-related infections, hospitalizations, and deaths: a systematic review. *Ann Intern Med.* 2021;174(3):362-373. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33253040.
- 3. Wu EL, Kumar RN, Moore WJ, et al. Disparities in COVID-19 monoclonal antibody delivery: a retrospective cohort study. *J Gen Intern Med*. 2022;37(10):2505-2513. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35469360.
- 4. Gold JAW, Kelleher J, Magid J, et al. Dispensing of oral antiviral drugs for treatment of COVID-19 by ZIP code-level social vulnerability—United States, December 23, 2021–May 21, 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(25):825-829. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35737571.
- Hammond J, Leister-Tebbe H, Gardner A, et al. Oral nirmatrelvir for high-risk, nonhospitalized adults with COVID-19. N Engl J Med. 2022;386(15):1397-1408. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35172054</u>.
- Gottlieb RL, Vaca CE, Paredes R, et al. Early remdesivir to prevent progression to severe COVID-19 in outpatients. *N Engl J Med.* 2022;386(4):305-315. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34937145</u>.
- 7. Ritonavir-boosed nirmatrelvir (Paxlovid) [package insert]. Food and Drug Administration. 2023. Available at: <u>https://www.accessdata.fda.gov/drugsatfda_docs/label/2023/217188s000lbl.pdf</u>.
- Hiremath S, Blake PG, Yeung A, et al. Early experience with modified dose nirmatrelvir/ritonavir in dialysis patients with coronavirus disease 2019. *Clin J Am Soc Nephrol*. 2023;18(4):485-490. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36723285</u>.
- Toussi SS, Neutel JM, Navarro J, et al. Pharmacokinetics of oral nirmatrelvir/ritonavir, a protease inhibitor for treatment of COVID-19, in subjects with renal impairment. *Clin Pharmacol Ther*. 2022;112(4):892-900. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35712797</u>.

- 10. University of Liverpool. Prescribing resources. 2022. Available at: <u>https://covid19-druginteractions.org/</u> prescribing_resources. Accessed July 13, 2023.
- 11. Ontario Health. COVID-19 supplemental clinical guidance #4: nirmatrelvir/ritonavir (Paxlovid) use in patients with advanced chronic kidney disease and patients on dialysis with COVID-19. 2022. Available at: https://www.ontariohealth.ca/sites/ontariohealth/files/2022-04/PaxlovidClinicalGuide.pdf.
- 12. Food and Drug Administration Center for Drug Evaluation and Research. Antimicrobial drugs advisory committee meeting. 2023. Available at: <u>https://www.fda.gov/media/168508/download</u>.
- Huygens S, Gharbharan A, Serroukh Y, et al. High-titer convalescent plasma plus nirmatrelvir/ritonavir treatment for non-resolving COVID-19 in six immunocompromised patients. *J Antimicrob Chemother*. 2023;78(7):1644-1648. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37248664</u>.
- 14. Brosh-Nissimov T, Ma'aravi N, Leshin-Carmel D, et al. Combination treatment of persistent COVID-19 in immunocompromised patients with remdesivir, nirmatrelvir/ritonavir and tixagevimab/cilgavimab. J Microbiol Immunol Infect. 2023;Published online ahead of print. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/37805361/</u>.
- 15. Mikulska M, Sepulcri C, Dentone C, et al. Triple combination therapy with two antivirals and monoclonal antibodies for persistent or relapsed SARS-CoV-2 infection in immunocompromised patients. *Clin Infect Dis.* 2023;77(2):280-286. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36976301.
- Graziani L, Gori L, Manciulli T, et al. Successful use of nirmatrelvir/ritonavir in immunocompromised patients with persistent and/or relapsing COVID-19. *J Antimicrob Chemother*. 2023;78(2):555-558. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36544352</u>.
- 17. Trottier CA, Wong B, Kohli R, et al. Dual antiviral therapy for persistent coronavirus disease 2019 and associated organizing pneumonia in an immunocompromised host. *Clin Infect Dis.* 2023;76(5):923-925. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36281907</u>.
- Pillaiyar T, Manickam M, Namasivayam V, Hayashi Y, Jung SH. An overview of severe acute respiratory syndrome-coronavirus (SARS-CoV) 3CL protease inhibitors: peptidomimetics and small molecule chemotherapy. *J Med Chem.* 2016;59(14):6595-6628. Available at: https://www.ncbi.nlm.nih.gov/pubmed/26878082.
- Fiaschi L, Dragoni F, Schiaroli E, et al. Efficacy of licensed monoclonal antibodies and antiviral agents against the SARS-CoV-2 Omicron sublineages BA.1 and BA.2. *Viruses*. 2022;14(7):1374. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35891355</u>.
- Takashita E, Kinoshita N, Yamayoshi S, et al. Efficacy of antiviral agents against the SARS-CoV-2 Omicron subvariant BA.2. *N Engl J Med*. 2022;386(15):1475-1477. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35263535</u>.
- 21. Takashita E, Yamayoshi S, Simon V, et al. Efficacy of antibodies and antiviral drugs against Omicron BA.2.12.1, BA.4, and BA.5 subvariants. *N Engl J Med.* 2022;387(5):468-470. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35857646</u>.
- 22. Fischer WA II, Eron JJ Jr, Holman W, et al. A Phase 2a clinical trial of molnupiravir in patients with COVID-19 shows accelerated SARS-CoV-2 RNA clearance and elimination of infectious virus. *Sci Transl Med.* 2022;14(628):eabl7430. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34941423.
- 23. Zhou S, Hill CS, Sarkar S, et al. Beta-d-N4-hydroxycytidine inhibits SARS-CoV-2 through lethal mutagenesis but is also mutagenic to mammalian cells. *J Infect Dis*. 2021;224(3):415-419. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33961695</u>.
- 24. Zou R, Peng L, Shu D, et al. Antiviral efficacy and safety of molnupiravir against Omicron variant infection: a randomized controlled clinical trial. *Front Pharmacol*. 2022;13:939573. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35784723.
- 25. Jayk Bernal A, Gomes da Silva MM, Musungaie DB, et al. Molnupiravir for oral treatment of COVID-19 in nonhospitalized patients. *N Engl J Med*. 2022;386(6):509-520. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34914868</u>.

COVID-19 Treatment Guidelines

- 26. Johnson MG, Puenpatom A, Moncada PA, et al. Effect of molnupiravir on biomarkers, respiratory interventions, and medical services in COVID-19: a randomized, placebo-controlled trial. *Ann Intern Med.* 2022;175(8):1126-1134. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35667065</u>.
- 27. Butler CC, Hobbs FDR, Gbinigie OA, et al. Molnupiravir plus usual care versus usual care alone as early treatment for adults with COVID-19 at increased risk of adverse outcomes (PANORAMIC): an open-label, platform-adaptive randomised controlled trial. *Lancet*. 2023;401(10373):281-293. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36566761.
- 28. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Lagevrio (molnupiravir) capsules. 2023. Available at: <u>https://www.fda.gov/media/155054/download</u>.
- Charness ME, Gupta K, Stack G, et al. Rebound of SARS-CoV-2 infection after nirmatrelvir-ritonavir treatment. N Engl J Med. 2022;387(11):1045-1047. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36069968</u>.
- Anderson AS, Caubel P, Rusnak JM, Investigators E-HT. Nirmatrelvir-ritonavir and viral load rebound in COVID-19. N Engl J Med. 2022;387(11):1047-1049. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36069818</u>.
- Boucau J, Uddin R, Marino C, et al. Characterization of virologic rebound following nirmatrelvir-ritonavir treatment for coronavirus disease 2019 (COVID-19). *Clin Infect Dis*. 2023;76(3):e526-e529. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35737946</u>.
- 32. Deo R, Choudhary MC, Moser C, et al. Symptom and viral rebound in untreated SARS-CoV-2 infection. *Ann Intern Med.* 2023;176(3):348-354. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36802755</u>.
- 33. Soares H, Baniecki ML, Cardin R, et al. Viral load rebound in placebo and nirmatrelvir-ritonavir treated COVID-19 patients is not associated with recurrence of severe disease or mutations. *Res Sq.* 2022;Preprint. Available at: <u>https://www.researchsquare.com/article/rs-1720472/v1</u>.
- Ranganath N, O'Horo JC, Challener DW, et al. Rebound phenomenon after nirmatrelvir/ritonavir treatment of coronavirus disease 2019 (COVID-19) in high-risk persons. *Clin Infect Dis*. 2023;76(3):e537-e539. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35698452</u>.
- 35. Centers for Disease Control and Prevention. COVID-19 rebound after Paxlovid treatment. 2022. Available at: <u>https://emergency.cdc.gov/han/2022/han00467.asp</u>. Accessed July 13, 2023.
- 36. Bradley MC, Perez-Vilar S, Chillarige Y, et al. Systemic corticosteroid use for COVID-19 in US outpatient settings from April 2020 to August 2021. JAMA. 2022;327(20):2015-2018. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35394487</u>.
- Crothers K, DeFaccio R, Tate J, et al. Dexamethasone in hospitalised COVID-19 patients not on intensive respiratory support. *Eur Respir J*. 2022;60(1):2102532. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34824060</u>.
- 38. RECOVERY Collaborative Group. Dexamethasone in hospitalized patients with COVID-19. *N Engl J Med.* 2021;384(8):693-704. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32678530</u>.
- 39. Cazzola M, Ora J, Bianco A, Rogliani P, Matera MG. Guidance on nebulization during the current COVID-19 pandemic. *Respir Med.* 2021;176:106236. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33248363.

Therapeutic Management of Hospitalized Adults With COVID-19

Last Updated: October 10, 2023

Table 2b. Therapeutic Management of Hospitalized Adults With COVID-19

Disease Severity	Recommendati	Recommendations for		
DISEASE SEVENLY	Clinical Scenario	Recommendation	Anticoagulant Therapy	
Hospitalized for Reasons Other Than COVID-19	Patients with mild to moderate COVID-19 who are at high risk of progressing to severe COVID-19 ^{a,b}	See <u>Therapeutic Management of Nonhospitalized Adults With</u> <u>COVID-19</u> .	For patients without an indication for therapeutic anticoagulation: • Prophylactic dose of heparin, unless	
Hospitalized but Does Not Require Supplemental	All patients	The Panel recommends against the use of dexamethasone (Alla) or other systemic corticosteroids (AllI) for the treatment of COVID- $19.^{\circ}$	contraindicated (AI); (BIII) for pregnant patients	
Oxygen	Patients who are at high risk of progressing to severe COVID-19 ^{a,b}	Remdesivir ^d (BIIb) for patients who are immunocompromised; (BIII) for other high-risk patients		
Hospitalized and Requires Conventional Oxygen ^e	Patients who require minimal conventional oxygen	Remdesivir ^{d,f} (Blla)	For nonpregnant patients with D-dimer levels above the ULN who do not have an	
	Most patients	Use dexamethasone plus remdesivir ^f (Blla). If remdesivir cannot be obtained, use dexamethasone (Bl) .	increased bleeding risk: • Therapeutic dose of heparin ^h (Clla)	
	Patients who are receiving dexamethasone and who have rapidly increasing oxygen needs and systemic inflammation	Add 1 of the following immunomodulators: ⁹ <i>Preferred</i> • PO baricitinib (Blla) • IV tocilizumab (Blla) <i>Alternatives</i>	 For other patients: Prophylactic dose of heparin, unless contraindicated (AI); (BIII) for pregnant patients 	
		 IV abatacept (CIIa) IV infliximab (CIIa) 		

Disease Severity	Recommendations for Antiviral or Immunomodulator Therapy		Recommendations for	
Disease Seventy	Clinical Scenario	Recommendation	Anticoagulant Therapy	
Hospitalized and Requires HFNC Oxygen or NIV	All patients	 Dexamethasone should be administered to all patients (AI). If not already initiated, promptly add 1 of the following immunomodulators: <i>Preferred</i> PO baricitinib^{9,i} (AI) <i>Preferred Alternative</i> IV tocilizumab^{9,i} (BIIa) <i>Additional Alternatives (Listed in Alphabetical Order)</i> IV abatacept^{9,i} (CIIa) IV infliximab^{9,i} (CIIa) Add remdesivir to 1 of the options above in certain patients (for examples, see footnote).ⁱ 	 For patients without an indication for therapeutic anticoagulation: Prophylactic dose of heparin, unless contraindicated (AI); (BIII) for pregnant patients For patients who are started on a therapeutic dose of heparin in a non-ICU setting and then transferred to the ICU, the Panel recommends switching to a prophylactic dose of heparin, unless there is another indication for therapeutic anticoagulation (BIII). 	
Hospitalized and Requires MV or ECMO	All patients	 Dexamethasone should be administered to all patients (AI). If the patient has not already received a second immunomodulator, promptly add 1 of the following (listed in alphabetical order): PO baricitinib^{i,k} (BIIa) IV tocilizumab^{i,k} (BIIa) 		

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

^a For a list of risk factors, see the CDC webpage Underlying Medical Conditions Associated With Higher Risk for Severe COVID-19.

^b Ritonavir-boosted nirmatrelvir (Paxlovid) has not been studied in hospitalized patients. The FDA product label for ritonavir-boosted nirmatrelvir allows for its use in hospitalized patients with mild to moderate COVID-19 (i.e., those who do not require supplemental oxygen) who are at high risk of progressing to severe COVID-19 and who are within 5 days of symptom onset.

° Corticosteroids that are prescribed for an underlying condition should be continued.

^d Evidence suggests that the benefit of remdesivir is greatest when the drug is given early in the course of COVID-19 (e.g., within 10 days of symptom onset).

^e Conventional oxygen refers to oxygen supplementation that is not HFNC oxygen, NIV, MV, or ECMO.

^f If these patients progress to requiring HFNC oxygen, NIV, MV, or ECMO, the full course of remdesivir should still be completed.

⁹ If none of the preferred or alternative options are available or feasible to use, the JAK inhibitor **PO tofacitinib (Clla)** or the IL-6 inhibitor **IV sarilumab (Clla)** can be used in combination with dexamethasone. Sarilumab is only commercially available as a SUBQ injection; see <u>Table 5e</u> for information regarding the preparation of an IV infusion using the SUBQ product.

^h Contraindications for the use of therapeutic anticoagulation in patients with COVID-19 include a PLT <50 x 10⁹/L, Hgb <8 g/dL, the need for dual antiplatelet therapy, bleeding within the past 30 days that required an ED visit or hospitalization, a history of a bleeding disorder, or an inherited or active acquired bleeding disorder.

ⁱ Dexamethasone should be initiated immediately, without waiting until the second immunomodulator can be acquired. If other immunomodulators cannot be obtained or are contraindicated, use dexamethasone alone **(AI)**.

^j Examples of patients who may benefit most from adding remdesivir include patients who are immunocompromised (BIIb); patients with evidence of ongoing viral

COVID-19 Treatment Guidelines

replication (e.g., those with a low Ct value, as measured by an RT-PCR result or with a positive rapid antigen test result) (**BIII**); or patients who are ≤ 10 days from symptom onset (**CIIa**). For more information on immunocompromising conditions, see <u>Special Considerations in People Who Are Immunocompromised</u>.

^k If PO baricitinib and IV tocilizumab are not available or feasible to use, **PO tofacitinib** can be used instead of PO baricitinib (**Clla**), and **IV sarilumab** can be used instead of IV tocilizumab (**Clla**).

Key: CDC = Centers for Disease Control and Prevention; Ct = cycle threshold; ECMO = extracorporeal membrane oxygenation; ED = emergency department; FDA = Food and Drug Administration; HFNC = high-flow nasal cannula; Hgb = hemoglobin; ICU = intensive care unit; IL = interleukin; IV = intravenous; JAK = Janus kinase; MV = mechanical ventilation; NIV = noninvasive ventilation; the Panel = the COVID-19 Treatment Guidelines Panel; PLT = platelet count; PO = oral; RT-PCR = reverse transcription polymerase chain reaction; SUBQ = subcutaneous; ULN = upper limit of normal

Two main processes are thought to drive the pathogenesis of COVID-19. Early in the clinical course, the disease is primarily driven by the replication of SARS-CoV-2. Later in the clinical course, the disease is driven by a dysregulated immune/inflammatory response to SARS-CoV-2 infection that may lead to further tissue damage and thrombosis. Based on this understanding, therapies that directly target SARS-CoV-2 are anticipated to have the greatest effect early in the course of the disease, whereas immunosuppressive, anti-inflammatory, and antithrombotic therapies are likely to be more beneficial after COVID-19 has progressed to stages characterized by hypoxemia and endothelial dysfunction.

Currently, most people in the United States have some degree of immunity to SARS-CoV-2 due to COVID-19 vaccination or SARS-CoV-2 infection. The increase in population immunity and the change in variants have led to a decrease in the rate of severe disease caused by COVID-19. Because other co-existing disease processes can cause hypoxemia in patients who test positive for SARS-CoV-2 infection, clinicians should perform the appropriate evaluations to rule out alternative diagnoses.

Below is a summary of the rationale for the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations on the therapeutic management of hospitalized patients with COVID-19. For dosing information for each of the recommended drugs, please refer to Table 2c below. For more information about these therapies and the evidence that supports the Panel's recommendations, please refer to the specific drug pages and clinical data tables.

Patients Who Are Hospitalized for Reasons Other Than COVID-19 and Who Do Not Require Supplemental Oxygen

Hospitalized patients with COVID-19 who do not require supplemental oxygen are a heterogeneous population. Some patients may be hospitalized for reasons other than COVID-19 but may also have mild to moderate COVID-19 (see <u>Clinical Spectrum of SARS-CoV-2 Infection</u>). In these cases, patients who are at high risk of progressing to severe COVID-19 may benefit from antiviral therapy.

Remdesivir is approved by the Food and Drug Administration (FDA) for the treatment of COVID-19 in adult and pediatric patients aged ≥ 12 years and weighing ≥ 40 kg who:

- Are hospitalized; or
- Are not hospitalized, have mild to moderate COVID-19, and are at high risk of progressing to severe COVID-19.

Ritonavir-boosted nirmatrelvir (Paxlovid) is approved by the FDA and molnupiravir has an Emergency Use Authorization from the FDA for use in patients with mild to moderate COVID-19 who are at high risk of progressing to severe disease. These therapies can be used in hospitalized patients who qualify for therapy if they were admitted to the hospital for a diagnosis other than COVID-19. The Panel's recommendations for these patients are the same as those for nonhospitalized patients (see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>).

Patients Who Are Hospitalized for COVID-19 and Who Do Not Require Supplemental Oxygen

Recommendations

- The Panel recommends using **remdesivir** for the treatment of COVID-19 in patients who do not require supplemental oxygen and who are immunocompromised (**BIIb**) and for other patients who are at high risk of progressing to severe disease (**BIII**).
- Remdesivir should be administered for 5 days or until hospital discharge, whichever comes first.

The rationale for using remdesivir in high-risk patients is based on several lines of evidence. In a trial conducted predominantly among hospitalized patients with COVID-19 who were not receiving supplemental oxygen at enrollment, a 5-day course of remdesivir was associated with greater clinical improvement when compared with standard of care.¹ Evidence from the PINETREE trial also suggests that early therapy reduces the risk of progression, although that study was performed in high-risk, unvaccinated, nonhospitalized patients with \leq 7 days of symptoms who received a 3-day course of remdesivir.

Other studies have not shown a clinical benefit of remdesivir in this group of hospitalized patients with COVID-19. In the ACTT-1 trial, remdesivir showed no significant benefit in hospitalized patients with mild to moderate disease; however, only 13% of the study population did not require supplemental oxygen.² In the large Solidarity trial, the use of remdesivir was not associated with a survival benefit among the subset of hospitalized patients who did not require supplemental oxygen.³ See <u>Table 4a</u> for more information.

The aggregate data on using remdesivir to treat all high-risk patients show a faster time to recovery in patients who received remdesivir but no clear evidence of a survival benefit. Therefore, the Panel recommends using **remdesivir** in hospitalized patients with COVID-19 who are at high risk of progressing to severe disease (**BIII**).

In a large, retrospective cohort study of hospitalized patients with COVID-19 who were immunocompromised (n = 28,338), patients who received remdesivir had a lower risk of mortality than those who did not receive remdesivir.⁴ Forty percent of patients in this cohort were not receiving supplemental oxygen at baseline; mortality was reduced in this subset of patients. Therefore, the Panel recommends using **remdesivir** in hospitalized patients with COVID-19 who are immunocompromised (**BIIb**).

For information on medical conditions that confer high risk, see the Centers for Disease Control and Prevention webpage <u>People With Certain Medical Conditions</u>.

Recommendation

• The Panel **recommends against** the use of **dexamethasone** (**AIIa**) or other systemic corticosteroids (**AIII**) for the treatment of COVID-19 in patients who do not require supplemental oxygen.

In the RECOVERY trial, no survival benefit was observed for dexamethasone among the subset of patients with COVID-19 who did not require supplemental oxygen at enrollment.⁵ In an observational cohort study of U.S. veterans, the use of dexamethasone was associated with higher mortality in hospitalized patients with COVID-19 who did not require supplemental oxygen.⁶

There are insufficient data to inform the use of other systemic corticosteroids in hospitalized patients with COVID-19. Patients who are receiving corticosteroid treatment for an underlying condition should continue to receive corticosteroids. See <u>Table 5a</u> for more information.

COVID-19 Treatment Guidelines

Patients Who Require Conventional Oxygen

Patients with COVID-19 who require conventional oxygen (i.e., those who do not require high-flow nasal cannula [HFNC] oxygen, noninvasive ventilation [NIV], or mechanical ventilation) are a heterogeneous population. Although the oxygen requirement qualifies all these patients as having severe disease, some of these patients will improve after a short period with or without treatment; others will develop progressive disease. There is no consensus on which clinical or laboratory parameters should be used to determine a patient's risk of progression and guide therapy.

Recommendation

• For patients with COVID-19 who only require minimal conventional oxygen, the Panel recommends using **remdesivir** without dexamethasone (**BIIa**).

In these patients, the hyperinflammatory state for which corticosteroids might be most beneficial may not yet be present or fully developed. In a subgroup analysis during the ACTT-1 trial, remdesivir significantly reduced the time to clinical recovery and significantly reduced mortality among the subset of patients who were receiving conventional oxygen at enrollment.² Evidence from ACTT-1 and a pooled analysis of individual data from 9 randomized controlled trials⁷ suggest that remdesivir will have its greatest benefit when administered early in the clinical course of COVID-19 (e.g., within 10 days of symptom onset). See <u>Table 4a</u> for more information.

Recommendations

- For most patients with COVID-19 who require conventional oxygen, the Panel recommends using **dexamethasone plus remdesivir (BIIa)**.
- If dexamethasone is not available, an equivalent dose of another corticosteroid (e.g., **prednisone**, **methylprednisolone**, **hydrocortisone**) may be used (**BIII**).

The results of several studies suggest that the use of remdesivir plus dexamethasone improves clinical outcomes among hospitalized patients with COVID-19. In the CATCO trial, in which 87% of patients received corticosteroids and 54% were on conventional oxygen, remdesivir significantly reduced the need for mechanical ventilation among the subset of patients who did not require mechanical ventilation at enrollment when compared with standard of care.⁸ In the Solidarity trial, in which approximately two-thirds of the patients received corticosteroids, remdesivir significantly reduced mortality among the large subset of patients (n > 7,000) who were receiving conventional or HFNC oxygen at enrollment.³ See <u>Table 4a</u> for more information.

An individual patient-level meta-analysis of 8 clinical trials examined the efficacy of using remdesivir in hospitalized patients with COVID-19.⁷ This meta-analysis found that remdesivir significantly reduced the number of patients who required mechanical ventilation or who died by Day 28 in the combined subgroups of patients who did not require oxygen or who were receiving conventional oxygen at baseline. However, the effect of remdesivir was not evaluated separately in the subgroup of patients who were receiving conventional oxygen at enrollment.

Recommendation

• If remdesivir is not available, the Panel recommends using **dexamethasone alone** in patients with COVID-19 who require conventional oxygen (**BI**).

In the RECOVERY trial, the use of dexamethasone 6 mg once daily for 10 days or until hospital discharge significantly reduced mortality among the subset of patients who were receiving oxygen (defined as receiving oxygen supplementation but not mechanical ventilation or extracorporeal membrane oxygenation [ECMO]) at enrollment.⁵ Remdesivir was administered to <1% of the study *COVID-19 Treatment Guidelines*

participants. Results for patients who were only receiving conventional oxygen at enrollment were not available. See <u>Table 5a</u> for more information.

Recommendation

- For patients with COVID-19 who have rapidly increasing oxygen needs and systemic inflammation, the Panel recommends adding 1 of the following immunomodulators to dexamethasone:
 - Preferred Second Immunomodulators
 - Oral (PO) baricitinib (BIIa)
 - Intravenous (IV) tocilizumab (BIIa)
 - Alternative Second Immunomodulators
 - IV abatacept (CIIa)
 - IV infliximab (CIIa)

If none of these options are available or feasible to use, the Janus kinase (JAK) inhibitor **PO tofacitinib** (**CIIa**) or the interleukin (IL)-6 inhibitor **IV sarilumab** (**CIIa**) can be used in combination with dexamethasone. Sarilumab is only commercially available as a subcutaneous (SUBQ) injection; see <u>Table 5e</u> for information regarding the preparation of an IV infusion using the SUBQ product.

Several large randomized controlled trials have evaluated the use of dexamethasone in combination with a second immunomodulator, including:

- Abatacept, a cytotoxic T-lymphocyte-associated antigen 4 agonist⁹
- Baricitinib, a JAK inhibitor¹⁰⁻¹³
- Infliximab, a tumor necrosis factor inhibitor⁹
- Tocilizumab, an IL-6 inhibitor^{12,14,15}

These studies included patients who required conventional oxygen only, as well as those with increasing oxygen needs and/or elevated levels of inflammatory markers. Subgroup analyses in these trials have not clearly defined which patients in this heterogeneous group are most likely to benefit from adding a second immunomodulator to corticosteroid therapy. The study endpoints for these trials included progression to more severe disease, the need for mechanical ventilation, and death. Nonetheless, some trials suggest that adding a second immunomodulator provides benefits to patients who require conventional oxygen, especially those with rapidly increasing oxygen requirements and systemic inflammation.

The Panel recommends either baricitinib or tocilizumab as the preferred second immunomodulator because both are approved by the FDA for the treatment of COVID-19, and data from multiple clinical trials have demonstrated that these agents provide a clinical benefit in patients with COVID-19 who require conventional oxygen.^{5,10-16} There is also more clinical experience with the use of these 2 agents in this setting than other potential treatment options.

The ACTIV-1 immune modulator trial was a double-blind, multi-arm, placebo-controlled, randomized trial in moderately to severely ill adults hospitalized with COVID-19.⁹ The trial separately evaluated treatment with the immunomodulators abatacept, cenicriviroc, and infliximab versus placebo. All arms received standard care, and the separate analyses included data from a shared placebo arm. The primary endpoint was time to recovery by Day 28. Key secondary endpoints included clinical status at Day 14 and mortality through Days 28 and 60. The majority of patients received corticosteroids (>89%) and remdesivir (>93%).

COVID-19 Treatment Guidelines

None of the study drugs had a significant effect on the time to recovery. Mortality by Day 28 was lower among patients in the abatacept and infliximab arms than among those in the shared placebo arm. Based on the results of this trial, abatacept or infliximab may be considered alternatives to baricitinib or tocilizumab. There are no studies that directly compare the use of abatacept or infliximab to the use of baricitinib or tocilizumab in people with COVID-19.

When baricitinib, tocilizumab, abatacept, or infliximab are not available or feasible to use, the JAK inhibitor tofacitinib or the IL-6 inhibitor sarilumab may be used as alternative agents. Tofacitinib decreased the risk for respiratory failure or death among hospitalized patients with COVID-19 in the STOP-COVID trial,¹⁷ and sarilumab reduced mortality and the duration of organ support to the same degree as tocilizumab in the REMAP-CAP trial.^{12,14}

Use of Anticoagulants

- The Panel recommends using a **therapeutic dose of heparin** for nonpregnant patients with D-dimer levels above the upper limit of normal who require conventional oxygen and who do not have an increased bleeding risk (**CIIa**).
- Patients who do not meet the criteria for therapeutic heparin noted above, including pregnant individuals, should receive a **prophylactic dose of heparin** unless this drug is contraindicated **(AI)**; **(BIII)** for pregnant patients.

The Panel's recommendations for the use of heparin are based on data from 3 open-label randomized controlled trials that compared the use of therapeutic doses of heparin to prophylactic or intermediate doses of heparin in hospitalized patients who did not require intensive care unit (ICU)-level care. Pooled data from the ATTACC/ACTIV-4a/REMAP-CAP trials reported more organ support-free (i.e. alive and free of ICU-based organ support) days for patients in the therapeutic heparin arm than in the usual care arm, but there was no difference between the arms in mortality or length of hospitalization.¹⁸ The RAPID trial compared a therapeutic dose of heparin to a prophylactic dose in hospitalized patients with moderate COVID-19. There was no statistically significant difference between the arms in the occurrence of the primary endpoint (which was a composite endpoint of ICU admission and initiation of NIV or mechanical ventilation), but the therapeutic dose of heparin reduced 28-day mortality.¹⁹ In the HEP-COVID trial, venous thromboembolism (VTE), arterial thromboembolism, and death by Day 30 occurred significantly less frequently in patients who received a therapeutic dose of heparin than in those who received a prophylactic dose of heparin, but there was no difference in mortality by Day 30 between the arms.²⁰

Patients Who Require High-Flow Nasal Cannula Oxygen or Noninvasive Ventilation

In these patients, systemic inflammation contributes to hypoxemia, and thus these patients may benefit from receiving a second immunomodulator in addition to dexamethasone. There is no consensus on which clinical or laboratory parameters reliably predict the risk of progression to mechanical ventilation or death.

The available evidence suggests that the benefits of adding baricitinib or tocilizumab to dexamethasone treatment outweigh the potential risks in patients with COVID-19 who require HFNC oxygen or NIV. Although the combination of dexamethasone and secondary immunomodulating medications may increase the risk of opportunistic infections or the risk of reactivating latent infections, there are insufficient data to make recommendations about initiating prophylaxis against these infections.

Recommendations

• Dexamethasone should be administered to all patients with COVID-19 who require HFNC

COVID-19 Treatment Guidelines

oxygen or NIV (AI).

- If not already initiated, promptly add 1 of the following immunomodulators to dexamethasone:
 - Preferred Second Immunomodulator
 - PO baricitinib (AI)
 - Preferred Alternative Second Immunomodulator
 - IV tocilizumab (BIIa)
 - Additional Alternative Second Immunomodulators (Listed in Alphabetical Order)
 - IV abatacept (CIIa)
 - IV infliximab (CIIa)

If none of these options are available or feasible to use, **PO tofacitinib** (**CIIa**) or **IV sarilumab** (**CIIa**) can be used in combination with dexamethasone. Sarilumab is only commercially available as a SUBQ injection; see <u>Table 5e</u> for information regarding the preparation of an IV infusion using the SUBQ product.

Clinicians should make a significant effort to obtain and administer 1 of the recommended second immunomodulating medications. However, dexamethasone should be initiated immediately, without waiting until the second immunomodulator can be acquired. Dexamethasone was used as a single-agent immunomodulatory strategy in the RECOVERY trial and demonstrated a survival benefit among patients who required supplemental oxygen.⁵ In this trial, the treatment effect for dexamethasone was not evaluated separately for those who required conventional oxygen and those who required HFNC oxygen or NIV (see <u>Systemic Corticosteroids</u>).

Several large randomized controlled trials have demonstrated that patients with COVID-19 who require HFNC oxygen or NIV benefit from combining dexamethasone with an additional immunomodulator, such as a JAK inhibitor or an IL-6 inhibitor. The quality of the evidence and the totality of the data support a stronger recommendation for baricitinib than for tocilizumab. See <u>Table 5c</u> and <u>Table 5d</u> for more information. Every effort should be made to obtain baricitinib or tocilizumab. Other immunomodulators, including abatacept and infliximab, have shown a clinical benefit in people with COVID-19 in a randomized controlled trial.

Two large randomized controlled trials (the RECOVERY and COV-BARRIER trials) both reported a survival benefit among hospitalized patients with COVID-19 who required HFNC oxygen or NIV and who received baricitinib plus dexamethasone.¹³ Data from the ACTT-2¹⁰ and ACTT-4²¹ trials support the overall safety of using baricitinib in combination with remdesivir and the potential for a clinical benefit of this combination, but neither trial studied baricitinib in combination with dexamethasone as the standard of care. A retrospective analysis of data from 11 U.S. health systems suggests that the use of baricitinib may be associated with fewer adverse effects than tocilizumab, including fewer secondary infections, thrombotic events, and cases of acute liver injury.²²

The use of tocilizumab in combination with corticosteroids reduced in-hospital mortality in patients with rapid respiratory decompensation who were admitted to the ICU in the REMAP-CAP trial.¹⁴ Similar results were reported during the RECOVERY trial, although patients were only randomized into the tocilizumab arm if they had oxygen saturation <92% on room air and C-reactive protein levels \geq 75 mg/L.¹⁵ Both REMAP-CAP and RECOVERY evaluated the efficacy of adding tocilizumab to standard care; in both cases, standard care included dexamethasone therapy. Other randomized trials that have evaluated the use of tocilizumab have demonstrated mixed results, including a lack of benefit when tocilizumab was administered without dexamethasone as part of standard care.²³⁻²⁶

In the ACTIV-1 trial, which evaluated the use of abatacept, cenicriviroc, and infliximab in hospitalized patients with COVID-19, neither abatacept nor infliximab demonstrated a significant effect on the primary endpoint of time to recovery. In the subgroup of patients who received HFNC oxygen or NIV, mortality at Day 28 (a secondary outcome) was lower in both the abatacept and the infliximab arms than in the shared placebo arm.

Combinations of 3 immunomodulators (e.g., dexamethasone plus baricitinib plus tocilizumab) have not been studied in clinical trials. Although some patients in the baricitinib arm of the RECOVERY trial also received tocilizumab, data from the study are insufficient to issue a recommendation. When both agents are used, there is a potential for greater risk of secondary infections.¹³

The clinical trial data cited above informed the Panel's recommendations for adding a second immunomodulator to dexamethasone in hospitalized patients who require HFNC oxygen or NIV. Based on these clinical trial results, the Panel recommends baricitinib over tocilizumab as the second immunomodulator. See <u>Table 5c</u> and <u>Table 5d</u> for more information. The evidence for the use of either abatacept or infliximab in people with COVID-19 is derived from a single study, while multiple trials have demonstrated a beneficial effect of using baricitinib or tocilizumab.

Recommendations

- For certain hospitalized patients who require HFNC oxygen or NIV, the Panel recommends adding **remdesivir** to 1 of the recommended immunomodulator combinations. Examples of patients who may benefit most from adding remdesivir include:
 - Patients who are immunocompromised (BIIb)
 - Patients with evidence of ongoing viral replication (e.g., those with a low cycle threshold [Ct] value, as measured by a reverse transcription polymerase chain reaction [RT-PCR] result or with a positive rapid antigen test result) (**BIII**)
 - Patients who are ≤ 10 days from symptom onset (CIIa)

Clinical trial data have not clearly established that remdesivir reduces the time to recovery or improves survival in patients who require HFNC oxygen or NIV. However, because clinical trials have found that remdesivir prevents clinical progression in patients who are not on mechanical ventilation, some patients receiving HFNC oxygen or NIV might benefit from receiving remdesivir. In the Solidarity trial, remdesivir had a modest but statistically significant effect on reducing the risk of death or progression to ventilation in patients who were receiving oxygen but who were not ventilated at baseline.³ However, these effects could not be evaluated separately for patients who required conventional oxygen supplementation and those who required HFNC oxygen or NIV.³ In the CATCO trial, among the patients who were not receiving mechanical ventilation at baseline, 8% of patients who received remdesivir required mechanical ventilation compared to 15% of those who received standard of care (relative risk 0.53; 95% CI, 0.38–0.75).⁸ See <u>Table 4a</u> for more information.

The Panel's rationale for recommending remdesivir for certain patients who require HFNC oxygen or NIV is discussed below. This discussion includes examples of patients who may benefit most from receiving remdesivir. In addition, clinicians may extend the course of remdesivir beyond 5 days in this population based on clinical response.

Patients Who Are Immunocompromised

People who are immunocompromised already have difficulty achieving viral clearance. The use of immunomodulators to treat COVID-19 may further impair this process. Because SARS-CoV-2 replication may be prolonged in these patients, remdesivir may help enhance viral clearance and improve outcomes. In a large, retrospective study of a cohort of patients who were immunocompromised, patients

who received remdesivir had a lower risk of mortality than those who did not receive remdesivir; however, only 19% of the patients in the study were receiving HFNC oxygen or NIV.⁴ For more information, see <u>Special Considerations in People Who Are Immunocompromised</u>.

Patients With Suspected Ongoing Viral Replication

Hospitalized patients who require HFNC oxygen or NIV are routinely treated with 2 immunomodulators to prevent or mitigate inflammatory-mediated injury. These treatments may impair the patient's ability to achieve viral clearance; thus, directly treating the virus with remdesivir may theoretically help improve outcomes. Substantial evidence from studies of other viral diseases supports the benefits of reducing the viral burden. Ct values can be obtained from some SARS-CoV-2 RT-PCR assays, and these values may be used as a proxy for the level of ongoing viral replication (low Ct values correspond to higher viral loads). While this information is not available on all RT-PCR platforms, Ct values may be helpful in informing decisions regarding the use of remdesivir. Positive rapid antigen test results are also consistent with higher viral loads.²⁷

Patients Who Are Within 10 Days of Symptom Onset

Active viral replication occurs early in the course of SARS-CoV-2 infection. Evidence from the ACTT-1 and PINETREE trials suggests that remdesivir will have the greatest benefit when administered early in the clinical course of COVID-19. In the ACTT-1 trial, remdesivir demonstrated a greater benefit in patients who were enrolled within 10 days of symptom onset than in those who were enrolled later in the disease course.^{2,28}

Use of Anticoagulants

- The Panel recommends using a **prophylactic dose of heparin** as VTE prophylaxis, unless a contraindication exists (**AI**); (**BIII**) for pregnant patients.
- For patients who start on a therapeutic dose of heparin in a non-ICU setting due to COVID-19 and then transfer to the ICU, the Panel recommends switching from the therapeutic dose to a **prophylactic dose of heparin**, unless VTE is confirmed (**BIII**).
- The Panel **recommends against** the use of a therapeutic dose of anticoagulation for VTE prophylaxis, except in a clinical trial (**BI**).

The multiplatform randomized controlled trial REMAP-CAP/ACTIV-4a/ATTACC compared the effectiveness of a therapeutic dose of heparin to standard care in critically ill patients with COVID-19. The study did not show an increase in the number of organ support-free days or the probability of survival to hospital discharge among patients who received therapeutic doses of anticoagulation.¹⁶ See Antithrombotic Therapy in Patients With COVID-19 for more information.

Patients Who Require Mechanical Ventilation or Extracorporeal Membrane Oxygenation

Recommendations

- **Dexamethasone** should be administered to all patients with COVID-19 who require mechanical ventilation or ECMO (**AI**).
- If the patient has not already received a second immunomodulator in addition to dexamethasone, promptly add 1 of the following (listed in alphabetical order):
 - PO baricitinib (BIIa)
 - IV tocilizumab (BIIa)

Dexamethasone was shown to reduce mortality in critically ill patients with COVID-19 in a metaanalysis that aggregated 7 randomized trials and included data on 1,703 critically ill patients. The largest trial included in the meta-analysis was the RECOVERY trial, which had a subgroup of patients who were receiving mechanical ventilation (see <u>Systemic Corticosteroids</u> and <u>Table 5a</u>).⁵ Subsequent studies of immunomodulator therapy suggest that using a second immunomodulator in combination with dexamethasone is more effective in patients with COVID-19 who require mechanical ventilation or ECMO.

Clinical trials that have evaluated combining IL-6 inhibitors or JAK inhibitors with corticosteroids for the treatment of patients with COVID-19 provide the most robust evidence for the Panel's recommendations.

Clinical trials of tocilizumab have reported an overall survival benefit in patients with hypoxemia and signs of systemic inflammation (RECOVERY)¹⁵ and in patients who are critically ill and require organ support (REMAP-CAP).¹⁴ Although these studies included patients who were receiving mechanical ventilation at randomization, the studies were not specifically powered to assess the effectiveness of IL-6 inhibitors in these patients. Other studies of tocilizumab in critically ill patients did not find a survival benefit, although the time between initiation of organ support in the ICU and study enrollment differed across the studies (see <u>Table 5c</u>).^{23,26} The use of corticosteroids also varied across the studies.

An extension of the COV-BARRIER trial compared the efficacy of baricitinib to placebo in 101 critically ill patients with COVID-19. The study reported significant reductions in mortality (relative reduction of 46% at 28 days and 44% at 60 days) and no major adverse events among patients who received baricitinib.²⁹ Systematic reviews of JAK inhibitors confirm the efficacy of using baricitinib in hospitalized patients with COVID-19 who require oxygen support. There is a lower certainty of evidence for patients who were receiving mechanical ventilation or ECMO, and baricitinib may have modestly attenuated efficacy in this group.³⁰ Baricitinib or tocilizumab should only be administered in combination with dexamethasone or another corticosteroid.

In the ACTIV-1 trial, the use of abatacept, cenicriviroc, or infliximab did not reduce the time to recovery or mortality in patients with COVID-19 who required mechanical ventilation or ECMO. Therefore, these immunomodulators are not recommended for these patients.

Considerations for the Use of Remdesivir

Remdesivir is most effective against COVID-19 in patients who are earlier in the course of the disease and who do not require mechanical ventilation or ECMO. Among patients who were receiving mechanical ventilation or ECMO during the Solidarity trial, there was a trend toward an increase in mortality for patients treated with remdesivir.³ For patients who progress to requiring mechanical ventilation or ECMO after they initiate remdesivir, the Panel suggests continuing remdesivir until the treatment course is completed.

Subgroup analyses from 2 randomized trials suggest there is no clinical benefit to using a combination of remdesivir and dexamethasone in patients who are receiving mechanical ventilation or ECMO.^{2,3} The data are inconclusive on whether corticosteroid therapy may delay viral clearance in patients with COVID-19.³¹⁻³⁵

Use of Anticoagulants

- The Panel recommends using a **prophylactic dose of heparin** as VTE prophylaxis, unless a contraindication exists (**AI**); (**BIII**) for pregnant patients.
- For patients who start on a therapeutic dose of heparin in a non-ICU setting due to COVID-19 and then transfer to the ICU, the Panel recommends switching from the therapeutic dose to a **prophylactic dose of heparin**, unless there is another indication for therapeutic anticoagulation (**BIII**).

• The Panel **recommends against** the use of a therapeutic dose of anticoagulation for VTE prophylaxis, except in a clinical trial (**BI**).

Patients who required mechanical ventilation or ECMO were included in the multiplatform REMAP-CAP/ACTIV-4a/ATTACC trial that studied therapeutic dose of heparin.¹⁶ Because these studies reported no benefits of using therapeutic doses of heparin, the recommendations for using prophylactic doses of heparin in hospitalized patients who require mechanical ventilation or ECMO are the same as those for patients who require HFNC oxygen or NIV.

Table 2c. Dosing Regimens for the Drugs Recommended in Table 2b

The drugs in this table are listed in alphabetical order.

Drug Name	Dosing Regimen	Comments
Abatacept	Abatacept 10 mg/kg actual body weight (up to 1,000 mg) administered as a single IV dose	No adjustment based on eGFR
Baricitinib	BAR dose is dependent on eGFR; duration of therapy is up to 14 days or until hospital discharge (whichever comes first).	• eGFR \geq 60 mL/min/1.73 m ² : BAR 4 mg PO once daily • eGFR 30 to <60 mL/min/1.73 m ² : BAR 2 mg PO once daily • eGFR 15 to <30 mL/min/1.73 m ² : BAR 1 mg PO once daily • eGFR <15 mL/min/1.73 m ² : BAR is not recommended .
Dexamethasone	DEX 6 mg IV or PO once daily for up to 10 days or until hospital discharge (whichever comes first)	 If DEX is not available, an equivalent dose of another corticosteroid may be used. For more information, see <u>Systemic Corticosteroids</u>.
Infliximab	Infliximab 5 mg/kg actual body weight administered as a single IV dose	No adjustment based on eGFR
Heparin	Therapeutic dose of SUBQ LMWH or IV UFH	• Administer for 14 days or until hospital discharge (whichever comes first) unless there is a diagnosis of VTE or another indication for therapeutic anticoagulation.
	Prophylactic dose of SUBQ LMWH or SUBQ UFH	Administer for the duration of the hospital stay.
Remdesivir	RDV 200 mg IV once, then RDV 100 mg IV once daily for 4 days or until hospital discharge (whichever comes first)	 If the patient is hospitalized for reasons other than COVID-19, the treatment duration is 3 days. For more information, see <u>Therapeutic Management of</u> <u>Nonhospitalized Adults With COVID-19</u>. If the patient progresses to more severe illness, complete
		 For a discussion on using RDV in patients with renal insufficiency, see <u>Remdesivir</u>.
Sarilumab	Use the single-dose, prefilled syringe (not the prefilled pen) for SUBQ injection. Reconstitute sarilumab 400 mg in 100 cc 0.9% NaCl and administer as an IV infusion over 1 hour.	 In the United States, the currently approved route of administration for sarilumab is SUBQ injection. In the REMAP-CAP trial, the SUBQ formulation was used to prepare the IV infusion.
Tocilizumab	Tocilizumab 8 mg/kg actual body weight (up to 800 mg) administered as a single IV dose	 In clinical trials, a third of the participants received a second dose of tocilizumab 8 hours after the first dose if no clinical improvement was observed.
Tofacitinib	Tofacitinib 10 mg P0 twice daily for up to 14 days or until hospital discharge (whichever comes first)	• eGFR <60 mL/min/1.73 m ² : tofacitinib 5 mg PO twice daily

Key: BAR = baricitinib; DEX = dexamethasone; eGFR = estimated glomerular filtration rate; IV = intravenous; LMWH = low-molecular-weight heparin; NaCl = sodium chloride; PO = oral; RDV = remdesivir; SUBQ = subcutaneous; UFH = unfractionated heparin; VTE = venous thromboembolism

COVID-19 Treatment Guidelines

References

- 1. Spinner CD, Gottlieb RL, Criner GJ, et al. Effect of remdesivir vs standard care on clinical status at 11 days in patients with moderate COVID-19: a randomized clinical trial. *JAMA*. 2020;324(11):1048-1057. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32821939.
- Beigel JH, Tomashek KM, Dodd LE, et al. Remdesivir for the treatment of COVID-19—final report. *N Engl J Med.* 2020;383(19):1813-1826. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32445440</u>.
- 3. WHO Solidarity Trial Consortium. Remdesivir and three other drugs for hospitalised patients with COVID-19: final results of the WHO Solidarity randomised trial and updated meta-analyses. *Lancet*. 2022;399(10339):1941-1953. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35512728.
- 4. Mozaffari E, Chandak A, Gottlieb RL, et al. Remdesivir reduced mortality in immunocompromised patients hospitalized for COVID-19 across variant waves: findings from routine clinical practice. *Clin Infect Dis.* 2023;Published online ahead of print. Available at: https://pubmed.ncbi.nlm.nih.gov/37556727.
- 5. RECOVERY Collaborative Group. Dexamethasone in hospitalized patients with COVID-19. *N Engl J Med.* 2021;384(8):693-704. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32678530</u>.
- Crothers K, DeFaccio R, Tate J, et al. Dexamethasone in hospitalised COVID-19 patients not on intensive respiratory support. *Eur Respir J*. 2022;60(1):2102532. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34824060</u>.
- Amstutz A, Speich B, Mentré F, et al. Effects of remdesivir in patients hospitalized with COVID-19: a systematic review and individual patient data meta-analysis of randomised controlled trials. *Lancet Respir Med.* 2023;11(5):453-464. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36828006</u>.
- 8. Ali K, Azher T, Baqi M, et al. Remdesivir for the treatment of patients in hospital with COVID-19 in Canada: a randomized controlled trial. *CMAJ*. 2022;194(7):E242-E251. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35045989.
- 9. O'Halloran JA, Ko ER, Anstrom KJ, et al. Abatacept, cenicriviroc, or infliximab for treatment of adults hospitalized with COVID-19 pneumonia: a randomized clinical trial. *JAMA*. 2023;330(4):328-339. Available at: https://pubmed.ncbi.nlm.nih.gov/37428480.
- Kalil AC, Patterson TF, Mehta AK, et al. Baricitinib plus remdesivir for hospitalized adults with COVID-19. N Engl J Med. 2021;384(9):795-807. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33306283</u>.
- Marconi VC, Ramanan AV, de Bono S, et al. Efficacy and safety of baricitinib for the treatment of hospitalised adults with COVID-19 (COV-BARRIER): a randomised, double-blind, parallel-group, placebo-controlled Phase 3 trial. *Lancet Respir Med*. 2021;9(12):1407-1418. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34480861</u>.
- 12. REMAP-CAP Investigators. Effectiveness of tocilizumab, sarilumab, and anakinra for critically ill patients with COVID-19: the REMAP-CAP COVID-19 immune modulation therapy domain randomized clinical trial. *medRxiv*. 2021;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2021.06.18.21259133v2</u>.
- RECOVERY Collaborative Group. Baricitinib in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial and updated meta-analysis. *Lancet*. 2022;400(10349):359-368. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35908569</u>.
- 14. REMAP-CAP Investigators. Interleukin-6 receptor antagonists in critically ill patients with COVID-19. *N Engl J Med.* 2021;384(16):1491-1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631065</u>.
- 15. RECOVERY Collaborative Group. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2021;397(10285):1637-1645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33933206</u>.
- REMAP-CAP, ACTIV-4a, and ATTACC Investigators. Therapeutic anticoagulation with heparin in critically ill patients with COVID-19. *N Engl J Med*. 2021;385(9):777-789. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34351722</u>.

17. Guimarães PO, Quirk D, Furtado RH, et al. Tofacitinib in patients hospitalized with COVID-19 pneumonia. *N* COVID-19 Treatment Guidelines 67

Engl J Med. 2021;385(5):406-415. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34133856.

- ATTACC, ACTIV-4a, and REMAP-CAP Investigators. Therapeutic anticoagulation with heparin in noncritically ill patients with COVID-19. *N Engl J Med.* 2021;385(9):790-802. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34351721</u>.
- 19. Sholzberg M, Tang GH, Rahhal H, et al. Effectiveness of therapeutic heparin versus prophylactic heparin on death, mechanical ventilation, or intensive care unit admission in moderately ill patients with COVID-19 admitted to hospital: RAPID randomised clinical trial. *BMJ*. 2021;375:n2400. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34649864.
- 20. Spyropoulos AC, Goldin M, Giannis D, et al. Efficacy and safety of therapeutic-dose heparin vs standard prophylactic or intermediate-dose heparins for thromboprophylaxis in high-risk hospitalized patients with COVID-19: the HEP-COVID randomized clinical trial. *JAMA Intern Med.* 2021;181(12):1612-1620. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34617959.
- 21. Wolfe CR, Tomashek KM, Patterson TF, et al. Baricitinib versus dexamethasone for adults hospitalised with COVID-19 (ACTT-4): a randomised, double-blind, double placebo-controlled trial. *Lancet Respir Med.* 2022;10(9):888-899. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35617986</u>.
- Peterson JH, Paranjape NS, Grundlingh N, Priestley JL. Outcomes and adverse effects of baricitinib versus tocilizumab in the management of severe COVID-19. *Crit Care Med.* 2023;51(3):337-346. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36729439</u>.
- 23. Rosas IO, Bräu N, Waters M, et al. Tocilizumab in hospitalized patients with severe COVID-19 pneumonia. *N Engl J Med.* 2021;384(16):1503-1516. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631066</u>.
- 24. Salama C, Han J, Yau L, et al. Tocilizumab in patients hospitalized with COVID-19 pneumonia. *N Engl J Med*. 2021;384(1):20-30. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33332779</u>.
- 25. Stone JH, Frigault MJ, Serling-Boyd NJ, et al. Efficacy of tocilizumab in patients hospitalized with COVID-19. *N Engl J Med*. 2020;383(24):2333-2344. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33085857</u>.
- 26. Rosas IO, Diaz G, Gottlieb RL, et al. Tocilizumab and remdesivir in hospitalized patients with severe COVID-19 pneumonia: a randomized clinical trial. *Intensive Care Med*. 2021;47(11):1258-1270. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34609549.
- 27. Regan J, Flynn JP, Choudhary MC, et al. Detection of the Omicron variant virus with the Abbott BinaxNow SARS-CoV-2 rapid antigen assay. *Open Forum Infect Dis.* 2022;9(3):ofac022. Available at: https://pubmed.ncbi.nlm.nih.gov/35169591.
- Gottlieb RL, Vaca CE, Paredes R, et al. Early remdesivir to prevent progression to severe COVID-19 in outpatients. *N Engl J Med.* 2022;386(4):305-315. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34937145</u>.
- 29. Ely EW, Ramanan AV, Kartman CE, et al. Efficacy and safety of baricitinib plus standard of care for the treatment of critically ill hospitalised adults with COVID-19 on invasive mechanical ventilation or extracorporeal membrane oxygenation: an exploratory, randomised, placebo-controlled trial. *Lancet Respir Med.* 2022;10(4):327-336. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35123660.
- 30. Zhang X, Shang L, Fan G, et al. The efficacy and safety of Janus kinase inhibitors for patients with COVID-19: a living systematic review and meta-analysis. *Front Med (Lausanne)*. 2021;8:800492. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35155477.
- 31. Chen Y, Li L. Influence of corticosteroid dose on viral shedding duration in patients with COVID-19. *Clin Infect Dis.* 2021;72(7):1298-1300. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32588884</u>.
- 32. Li S, Hu Z, Song X. High-dose but not low-dose corticosteroids potentially delay viral shedding of patients with COVID-19. *Clin Infect Dis*. 2021;72(7):1297-1298. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32588877.
- 33. Ding C, Feng X, Chen Y, et al. Effect of corticosteroid therapy on the duration of SARS-CoV-2 clearance in

patients with mild COVID-19: a retrospective cohort study. *Infect Dis Ther*. 2020;9(4):943-952. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32986226</u>.

- 34. Liu J, Zhang S, Dong X, et al. Corticosteroid treatment in severe COVID-19 patients with acute respiratory distress syndrome. J Clin Invest. 2020;130(12):6417-6428. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33141117</u>.
- 35. Spagnuolo V, Guffanti M, Galli L, et al. Viral clearance after early corticosteroid treatment in patients with moderate or severe COVID-19. *Sci Rep.* 2020;10(1):21291. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33277573.

Clinical Management of Children Summary

Last Updated: August 22, 2023

Data from the Centers for Disease Control and Prevention demonstrate that SARS-CoV-2 infection and severe disease and death due to COVID-19 occur less often in children than in adults.¹⁻⁴ Although only a small percentage of children with COVID-19 will require medical attention, the percentage of intensive care unit admissions among hospitalized children is comparable to that for hospitalized adults with COVID-19.⁵⁻¹⁶

Observational studies and meta-analyses have found that children with certain comorbidities have a higher risk of severe COVID-19. These comorbidities include cardiac disease, neurologic disorders, prematurity (in young infants), diabetes, obesity (particularly severe obesity), chronic lung disease, feeding tube dependence, and immunocompromised status.¹⁷⁻²⁰ Demographic factors, such as age (<1 year and 10–14 years)²¹ and non-White race/ethnicity,^{12,22-24} have also been associated with severe disease. However, many studies did not assess the relative severity of underlying medical conditions in children with severe COVID-19.

In general, COVID-19 has similar clinical manifestations and disease stages in children and adults, including an early phase driven by viral replication and a late phase that appears to be driven by a dysregulated immune/inflammatory response to SARS-CoV-2 that leads to tissue damage. Respiratory complications in young children that can occur during the early clinical phase include croup and bronchiolitis. In addition, a small number of children who have recovered from acute SARS-CoV-2 infection develop multisystem inflammatory syndrome in children (MIS-C) 2 to 6 weeks after infection. MIS-C is a postinfectious inflammatory condition that can lead to severe organ dysfunction.

The published guidance on treating COVID-19 in children has been mostly extrapolated from recommendations for adults with COVID-19, recommendations for children with other viral infections, and expert opinion.²⁵⁻²⁷ Applying adult data from COVID-19 trials to children is a unique challenge because most children experience a mild course of illness with COVID-19. Relative to adults, children with COVID-19 have substantially lower mortality and less need for hospitalization. Because of these differences in epidemiology and disease severity, the effect sizes for children are likely to be smaller than those observed in adults; therefore, to produce a beneficial outcome, the number needed to treat is higher. Collectively, these factors influence the risk versus benefit balance for pharmacologic therapies in children.

In the absence of sufficient clinical trial data on the treatment of children with COVID-19, the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations for the therapeutic management of children are based largely on safety and efficacy data from clinical trials in adults, the child's risk of disease progression, and expert opinion. In general, the data from clinical trials in adults are most applicable when treating older children with severe COVID-19 and predominantly lower respiratory tract disease. It is challenging to develop recommendations for children with SARS-CoV-2 infection who present with clinical syndromes that are also associated with other respiratory viruses (e.g., bronchiolitis, croup, asthma) using data from clinical trials in adults. There is no evidence to suggest that these syndromes should be managed differently when caused by SARS-CoV-2 infection. Clinical judgment is needed when applying recommendations for treating adults with these clinical syndromes to children.

The Panel's recommendations for the management of children with COVID-19 or MIS-C are summarized in the tables below. Table 3a provides recommendations for the therapeutic management of nonhospitalized children with COVID-19. The Panel's recommendations are stratified by age (per the Food and Drug Administration Emergency Use Authorizations) and risk level. See <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u> for more information. Table 3b includes a framework to help clinicians evaluate the risk for severe COVID-19 based on patient conditions and COVID-19 vaccination status.

The recommendations for hospitalized children in Table 3c are stratified by disease severity. See <u>Therapeutic Management of Hospitalized Children With COVID-19</u> for more information. Table 3d summarizes the recommendations for the therapeutic management of MIS-C. For the rationale behind these recommendations and supporting data, see <u>Therapeutic Management of Hospitalized Children</u> <u>With MIS-C</u>, <u>Plus a Discussion on MIS-A</u>.

Disk of Course OOUID 40	Panel's Recommendations		
Risk of Severe COVID-19	Aged 12–17 Years	Aged <12 Years	
Symptomatic, Regardless of Risk Factors	• Provide supportive care (AIII).	• Provide supportive care (AIII).	
High Risk ^{a,b}	 Use 1 of the following options (listed in order of preference):^c Ritonavir-boosted nirmatrelvir (Paxlovid) within 5 days of symptom onset (BIII) Remdesivir within 7 days of symptom onset (CIII) 	 Ritonavir-boosted nirmatrelvir is not authorized by the FDA for use in children aged <12 years. There is insufficient evidence to recommend either for or against the routine use of remdesivir. Consider treatment based on age and other risk factors. 	
Intermediate Risk ^{b,d}	• There is insufficient evidence to recommend either for or against the use of any antiviral therapy. Consider treatment based on age and other risk factors.	• There is insufficient evidence to recommend either for or against the routine use of remdesivir.	
Low Risk ^{b,e}	 Manage with supportive care alone (BIII). 	Manage with supportive care alone (BIII).	
Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.			

^a Molnupiravir is not authorized by the FDA for use in children aged <18 years and should not be used.

- ^b See Table 3b for the Panel's framework for assessing the risk of progression to severe COVID-19 based on patient conditions and COVID-19 vaccination status.
- $^{\circ}$ Initiate treatment as soon as possible after symptom onset.
- ^d The relative risk of severe COVID-19 for intermediate-risk patients is lower than the risk for high-risk patients but higher than the risk for low-risk patients.
- ^e Low-risk patients include those with comorbid conditions that have a weak or unknown association with severe COVID-19. Patients with no comorbidities are included in this group.

Key: FDA = Food and Drug Administration; the Panel = the COVID-19 Treatment Guidelines Panel

Table 3b. The Panel's Framework for Assessing the Risk of Progression to Severe COVID-19Based on Patient Conditions and COVID-19 Vaccination Status

Conditions	Risk Level by Vaccination Status ^a		
Conditions	Unvaccinated	Primary Series	Up to Date
Strong or Consistent Association With Progre	ssion to Severe COVID-	19	•
Moderately or severely immunocompromised (see <u>Special Considerations in People Who Are</u> <u>Immunocompromised</u>)	High		
 Obesity (BMI ≥95th percentile for age), especially severe obesity (BMI ≥120% of 95th percentile for age)^b 			
 Medical complexity with dependence on respiratory technology^c 			
 Severe neurologic, genetic, metabolic, or other disability that results in impaired airway clearance or limitations in self care or activities of daily living 	High	Intermediate	
 Severe asthma or other severe chronic lung disease requiring ≥2 inhaled or ≥1 systemic medications daily 			
Severe congenital or acquired cardiac disease			
Multiple moderate to severe chronic diseases			
Moderate or Inconsistent Association With Prog	ression to Severe COVI)-19	
• Aged <1 year			
 Prematurity in children aged ≤2 years 			
Sickle cell disease	Intermediate		
 Diabetes mellitus (poorly controlled) 			
 Nonsevere cardiac, neurologic, or metabolic disease^d 			
Weak or Unknown Association With Progression to Severe COVID-19			
Mild asthma			
Overweight	Low		
Diabetes mellitus (well controlled)			

^a **Unvaccinated** = individuals who are not eligible for COVID-19 vaccination or are <2 weeks from the final dose of the primary series. **Vaccinated with primary series** = individuals who completed the primary series of 2 or 3 doses (the current CDC term is "fully vaccinated") and are >2 weeks after the final dose of the primary series but have not received a booster, if they are eligible for a booster. Children aged <5 years are not currently eligible for booster doses. **Vaccinated and up to date** = individuals who received the recommended booster dose(s) if eligible or have completed the primary series but are not yet eligible for a booster. See the <u>COVID-19 vaccination schedule</u> from the CDC for more information.

^b The degree of risk conferred by obesity in younger children is less clear than it is in older adolescents.

° This includes patients with a tracheostomy and those who require NIV.

^d The data for this group are particularly limited.

Key: BMI = body mass index; CDC = Centers for Disease Control and Prevention; NIV = noninvasive ventilation; the Panel = the COVID-19 Treatment Guidelines Panel

Table 3c. Therapeutic Management of Hospitalized Chi	ildren With COVID-19

Disease Severity	Panel's Recommendations
Hospitalized for COVID-19	For children aged ≥12 years admitted for COVID-19, use prophylactic anticoagulation unless contraindicated (BIII) . ^a
Does Not Require Supplemental Oxygen	For children admitted for COVID-19 who are at the highest risk of progression to severe COVID-19, ^b consider using remdesivir ^c for children aged 12–17 years (CIII) . There is insufficient evidence for using remdesivir in children aged 28 days to <12 years.
	For children admitted for reasons other than COVID-19 who have mild to moderate COVID-19 and are at the highest risk of progression, ^b refer to <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u> .
Requires Conventional Oxygen ^d	Use 1 of the following options:
	• Remdesivir ^c (BIII)
	 Dexamethasone plus remdesivir^c for children with increasing oxygen needs, particularly adolescents (BIII)
	Use 1 of the following options:
	Dexamethasone (BIII)
Requires Oxygen Through High-Flow	 Dexamethasone plus remdesivir^c (BIII)
Device or NIV ^e	For children who do not have rapid (e.g., within 24 hours) improvement in oxygenation after initiation of dexamethasone, baricitinib ^f or tocilizumab can be considered for children aged 12–17 years (BIII) and for children aged 2–11 years (CIII).
Requires MV or ECMO [®]	Dexamethasone ⁹ (AIII)
	For children who do not have rapid (e.g., within 24 hours) improvement in oxygenation after initiation of dexamethasone, baricitinib ^f or tocilizumab may be considered for children aged 12–17 years (BIII) and for children aged 2–11 years (CIII) .
	ves a rating for the strength of the recommendation (A, B, or C) and a rating II). See <u>Guidelines Development</u> for more information.

- ^a Weighing the risk factors for thrombosis and bleeding, some Panel members would use prophylactic anticoagulation for children aged <12 years who are hospitalized for COVID-19.
- ^b For example, for children who are severely immunocompromised regardless of COVID-19 vaccination status and those who are unvaccinated and have additional risk factors for progression (see <u>Therapeutic Management of Nonhospitalized</u> <u>Children With COVID-19</u>).
- ^c The clinical benefit of remdesivir is greatest if it is initiated within 10 days of symptom onset. Remdesivir should be given for 5 days or until hospital discharge, whichever comes first.
- ^d Conventional oxygen refers to oxygen supplementation that is not high-flow oxygen, NIV, MV, or ECMO.
- ^e Patients who are receiving NIV or MV at baseline and require a substantial increase in baseline support should be treated per the recommendations for patients requiring new NIV or MV.
- ^f **Tofacitinib** is an alternative if baricitinib is not available (**BIII**).
- ⁹ For children who started receiving remdesivir before admission to the ICU, the remdesivir should be continued to complete the treatment course.

Key: ECMO = extracorporeal membrane oxygenation; ICU = intensive care unit; MV = mechanical ventilation; NIV = noninvasive ventilation; the Panel = the COVID-19 Treatment Guidelines Panel

Table 3d. Therapeutic Management of Hospitalized Pediatric Patients With MIS-C

	Panel's Recommendations
	Initial treatment for MIS-C includes both immunomodulatory and antithrombotic therapy.
	Initial Immunomodulatory Therapy
	 IVIG 2 g/kg IBW (up to a maximum total dose of 100 g) IV plus low to moderate dose methylprednisolone (1–2 mg/kg/day) IV^a or another glucocorticoid at an equivalent dose^a (AIIb).
	• Glucocorticoid monotherapy, only if IVIG is unavailable or contraindicated (Blla).
	• IVIG monotherapy, only if glucocorticoids are contraindicated (BIIb).
	Intensification Immunomodulatory Therapy
	 Intensification therapy is recommended for children with refractory MIS-C who do not improve within 24 hours of receiving initial immunomodulatory therapy (AIII). One of the following can be used (listed in alphabetical order):
	 High-dose anakinra 5–10 mg/kg IV or SUBQ once daily (BIIb)
MIS-C	 Higher-dose glucocorticoid (e.g., methylprednisolone 10–30 mg/kg/day IV or equivalent glucocorticoid) (BIIb)^b
	 Infliximab^c 5–10 mg/kg IV for 1 dose (BIIb)
	Antithrombotic Therapy
	 Low-dose aspirin (3–5 mg/kg/day, up to maximum dose of 81 mg/day) PO for all patients without risk factors for bleeding (AIII), <u>AND</u>
	 Anticoagulation for patients who fall under 1 of the following clinical scenarios:
	 Therapeutic anticoagulation for patients with large CAAs according to the American Heart Association guidelines for Kawasaki disease (AIII).
	 Therapeutic anticoagulation for patients with moderate to severe LV dysfunction who have no risk factors for bleeding (AIII).
	 For patients with MIS-C who do not have large CAAs or moderate to severe LV dysfunction, consider prophylactic or therapeutic anticoagulation on an individual basis, taking into consideration risk factors for thrombosis and bleeding. See Table 3e for additional information.
	ne Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating orts it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

^a Duration of therapy may vary. For more information, see <u>Therapeutic Management of Hospitalized Children With MIS-C.</u> Plus a Discussion on MIS-A.

^b In certain patients with severe illness, intensification therapy may include dual therapy with higher-dose glucocorticoids and infliximab or anakinra. Anakinra and infliximab **should not be given** in combination.

^c Infliximab **should not be used** in patients with macrophage activation syndrome.

Key: CAA = coronary artery aneurysm; IBW = ideal body weight; IV = intravenous; IVIG = intravenous immunoglobulin; LV = left ventricular; MIS-C = multisystem inflammatory syndrome in children; PO = oral; SUBQ = subcutaneously

References

- 1. Centers for Disease Control and Prevention. COVID-19 weekly cases and deaths per 100,000 population by age, race/ethnicity, and sex. 2023. Available at: <u>https://covid.cdc.gov/covid-data-tracker/#demographicsovertime</u>. Accessed June 23, 2023.
- Centers for Disease Control and Prevention. Provisional COVID-19 deaths: focus on ages 0–18 years. 2023. Available at: <u>https://data.cdc.gov/NCHS/Provisional-COVID-19-Deaths-Focus-on-Ages-0-18-Yea/nr4s-juj3</u>.

Accessed June 23, 2023.

- Centers for Disease Control and Prevention. Demographic trends of COVID-19 cases and deaths in the US reported to CDC. 2023. Available at: <u>https://covid.cdc.gov/covid-data-tracker/#demographics</u>. Accessed June 23, 2023.
- 4. Centers for Disease Control and Prevention. COVID-NET laboratory-confirmed COVID-19 hospitalizations. 2023. Available at: <u>https://covid.cdc.gov/covid-data-tracker/#covidnet-hospitalization-network</u>. Accessed June 23, 2023.
- 5. Dong Y, Mo X, Hu Y, et al. Epidemiology of COVID-19 among children in China. *Pediatrics*. 2020;145(6):e20200702. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32179660</u>.
- CDC COVID-19 Response Team. Coronavirus disease 2019 in children—United States, February 12–April 2, 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(14):422-426. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32271728.
- Cui X, Zhang T, Zheng J, et al. Children with coronavirus disease 2019: a review of demographic, clinical, laboratory, and imaging features in pediatric patients. *J Med Virol*. 2020;92(9):1501-1510. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32418216</u>.
- 8. Livingston E, Bucher K. Coronavirus disease 2019 (COVID-19) in Italy. *JAMA*. 2020;323(14):1335. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32181795</u>.
- Tagarro A, Epalza C, Santos M, et al. Screening and severity of coronavirus disease 2019 (COVID-19) in children in Madrid, Spain. *JAMA Pediatr*. 2020;Published online ahead of print. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32267485</u>.
- DeBiasi RL, Song X, Delaney M, et al. Severe coronavirus disease-2019 in children and young adults in the Washington, DC, metropolitan region. *J Pediatr*. 2020;223:199-203.e1. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32405091</u>.
- 11. Chao JY, Derespina KR, Herold BC, et al. Clinical characteristics and outcomes of hospitalized and critically ill children and adolescents with coronavirus disease 2019 at a tertiary care medical center in New York City. *J Pediatr.* 2020;223:14-19.e2. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32407719.
- Swann OV, Holden KA, Turtle L, et al. Clinical characteristics of children and young people admitted to hospital with COVID-19 in United Kingdom: prospective multicentre observational cohort study. *BMJ*. 2020;370:m3249. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32960186</u>.
- Götzinger F, Santiago-García B, Noguera-Julián A, et al. COVID-19 in children and adolescents in Europe: a multinational, multicentre cohort study. *Lancet Child Adolesc Health*. 2020;4(9):653-661. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32593339</u>.
- Shi DS, Whitaker M, Marks KJ, et al. Hospitalizations of children aged 5–11 years with laboratory-confirmed COVID-19—COVID-NET, 14 states, March 2020–February 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(16):574-581. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35446827</u>.
- 15. Siegel DA, Reses HE, Cool AJ, et al. Trends in COVID-19 cases, emergency department visits, and hospital admissions among children and adolescents aged 0–17 years—United States, August 2020–August 2021. MMWR Morb Mortal Wkly Rep. 2021;70(36):1249-1254. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34499628</u>.
- Marks KJ, Whitaker M, Anglin O, et al. Hospitalizations of children and adolescents with laboratoryconfirmed COVID-19—COVID-NET, 14 states, July 2021–January 2022. *MMWR Morb Mortal Wkly Rep*. 2022;71(7):271-278. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35176003</u>.
- Wanga V, Gerdes ME, Shi DS, et al. Characteristics and clinical outcomes of children and adolescents aged <18 years hospitalized with COVID-19—six hospitals, United States, July–August 2021. *MMWR Morb Mortal Wkly Rep.* 2021;70(5152):1766-1772. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34968374</u>.
- 18. Choi JH, Choi SH, Yun KW. Risk factors for severe COVID-19 in children: a systematic review and metaanalysis. *J Korean Med Sci.* 2022;37(5):e35. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35132841</u>.

- 19. Shi Q, Wang Z, Liu J, et al. Risk factors for poor prognosis in children and adolescents with COVID-19: a systematic review and meta-analysis. *EClinicalMedicine*. 2021;41:101155. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34693233</u>.
- 20. Kompaniyets L, Agathis NT, Nelson JM, et al. Underlying medical conditions associated with severe COVID-19 illness among children. *JAMA Netw Open*. 2021;4(6):e2111182. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34097050.
- 21. Harwood R, Yan H, Talawila Da Camara N, et al. Which children and young people are at higher risk of severe disease and death after hospitalisation with SARS-CoV-2 infection in children and young people: a systematic review and individual patient meta-analysis. *EClinicalMedicine*. 2022;44:101287. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35169689.
- 22. Holmes L Jr, Wu C, Hinson R, et al. Black-White risk differentials in pediatric COVID-19 hospitalization and intensive care unit admissions in the USA. *J Racial Ethn Health Disparities*. 2023;10(3):1187-1193. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35604543.
- Antoon JW, Grijalva CG, Thurm C, et al. Factors associated with COVID-19 disease severity in US children and adolescents. *J Hosp Med.* 2021;16(10):603-610. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34613896.
- 24. Saatci D, Ranger TA, Garriga C, et al. Association between race and COVID-19 outcomes among 2.6 million children in England. *JAMA Pediatr.* 2021;175(9):928-938. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34152371.
- 25. Chiotos K, Hayes M, Kimberlin DW, et al. Multicenter initial guidance on use of antivirals for children with coronavirus disease 2019/severe acute respiratory syndrome coronavirus 2. *J Pediatric Infect Dis Soc*. 2020;9(6):701-715. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32318706</u>.
- 26. Dulek DE, Fuhlbrigge RC, Tribble AC, et al. Multidisciplinary guidance regarding the use of immunomodulatory therapies for acute coronavirus disease 2019 in pediatric patients. *J Pediatric Infect Dis Soc.* 2020;9(6):716-737. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32808988</u>.
- 27. Wolf J, Abzug MJ, Anosike BI, et al. Updated guidance on use and prioritization of monoclonal antibody therapy for treatment of COVID-19 in adolescents. *J Pediatric Infect Dis Soc*. 2022;11(5):177-185. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35107571.

Special Considerations in Children

Last Updated: July 21, 2023

Key Considerations

- SARS-CoV-2 infection is generally milder in children than in adults, and a substantial proportion of children with the infection are asymptomatic.
- Most nonhospitalized children with COVID-19 will not require any specific therapy.
- Children with ≥1 of the following comorbidities are at risk of severe COVID-19: cardiac disease, neurologic disorders, prematurity (in young infants), diabetes, obesity (particularly severe obesity), chronic lung disease, feeding tube dependence, and immunocompromised status. Age (<1 year and 10–14 years) and non-White race/ethnicity are also associated with severe disease.
- The data on the pathogenesis and clinical spectrum of SARS-CoV-2 infection are more limited for children than for adults.
- Vertical transmission of SARS-CoV-2 appears to be rare, but suspected or probable cases of vertical transmission have been described.
- A small subset of children and young adults with SARS-CoV-2 infection may develop multisystem inflammatory syndrome in children (MIS-C). Many patients with MIS-C require intensive care management. The majority of children with MIS-C do not have underlying comorbidities.
- Data on the prevalence of post-COVID conditions in children are limited but suggest that younger children may have fewer persistent symptoms than older children and adults.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

This section provides an overview of the epidemiology and clinical spectrum of disease, including COVID-19, multisystem inflammatory syndrome in children (MIS-C), and post-COVID conditions. This section also includes information on risk factors for severe COVID-19, vertical transmission, and infants born to a birth parent with SARS-CoV-2 infection. Throughout this section, COVID-19 refers to the acute, primarily respiratory illness due to infection with SARS-CoV-2. MIS-C refers to the postinfectious inflammatory condition.

Epidemiology

Data from the Centers for Disease Control and Prevention (CDC) demonstrate that SARS-CoV-2 infection and severe disease and death due to COVID-19 occur less often in children than in adults.¹⁻⁴ According to a report from the CDC, by February 2022, approximately 75% of children and adolescents had serologic evidence of prior SARS-CoV-2 infection.⁵

The data on the pathogenesis and clinical spectrum of SARS-CoV-2 infection in children are still limited compared to the data for adults. Although only a small percentage of children with COVID-19 will require medical attention, the percentage of intensive care unit (ICU) admissions among hospitalized children is comparable to that for hospitalized adults with COVID-19.⁶⁻¹⁷

Children from some racial and ethnic groups experience disproportionate rates of COVID-19-related hospitalization, which may be a result of barriers to accessing health care and economic and structural inequities. From 2020 to 2021, Black/African American children with COVID-19 in the United States were 2 times more likely to be hospitalized and 5 times more likely to be admitted to the ICU than White children.¹⁸

A U.S. study of children with COVID-19 who were hospitalized between April and September 2020 reported an association between race/ethnicity and disease severity.¹⁹ In a large United Kingdom study, admission to critical care was independently associated with hospitalized children who self-reported as being of Black ethnicity.¹³ A study in England reported that children who identified as Asian were more likely than children who identified as White to be hospitalized for COVID-19 and to be admitted to an ICU.²⁰ The study also found that children who identified as Black or as mixed or other races/ethnicities had significantly more hospitalizations than children who identified as White.

Clinical Manifestations of COVID-19

The signs and symptoms of SARS-CoV-2 infection in symptomatic children may be similar to those in adults; however, a greater proportion of children may be asymptomatic or have only mild illness when compared with adults. Although the true incidence of asymptomatic SARS-CoV-2 infection is unknown, a small study reported that 45% of children who underwent surveillance testing at the time of hospitalization for a non-COVID-19 indication had asymptomatic infection.²¹ The most common signs and symptoms of COVID-19 in hospitalized children are fever, nausea/vomiting, cough, shortness of breath, and upper respiratory symptoms.^{13,22} The signs and symptoms of COVID-19 may overlap significantly with those of influenza and other respiratory and enteric viral infections. Critical disease, including respiratory failure, acute respiratory distress syndrome, and, less commonly, shock, may occur in children with COVID-19.^{23,24} The overall incidence of SARS-CoV-2 infection and, by extension, COVID-19-related hospitalizations among children has increased substantially with the emergence of recent variants, particularly the Omicron variant.^{17,25} For more information, see <u>Therapeutic Management of Hospitalized Children With COVID-19</u> and <u>Introduction to Critical Care Management of Children With COVID-19</u>.

Risk Factors for Severe COVID-19

Observational studies and meta-analyses have found that children with certain comorbidities have a higher risk of severe COVID-19. These comorbidities include cardiac disease, neurologic disorders, prematurity (in young infants), diabetes, obesity (particularly severe obesity), chronic lung disease, feeding tube dependence, and immunocompromised status.²⁶⁻²⁹ Demographic factors, such as age (<1 year and 10–14 years)³⁰ and non-White race/ethnicity,^{13,18-20} have also been associated with severe disease. However, many studies did not assess the relative severity of underlying medical conditions.

Many published studies reported an increased relative risk of severe disease in children with comorbidities, but the absolute risk of severe COVID-19 among children remains low. However, protocolized admissions for certain populations (e.g., febrile young infants) may confound the association between comorbidities and severe COVID-19. Most children who have been hospitalized for severe COVID-19 have not been fully vaccinated, as many were not eligible for COVID-19 vaccination because of their age at the time these studies were conducted. The CDC has additional information on the underlying conditions that are <u>risk factors for severe COVID-19</u>.

The children who are most likely to benefit from treatment are nonhospitalized children with mild to moderate COVID-19 who are at the highest risk of severe COVID-19 (e.g., those with severe comorbidities). For a description of children who are considered to be at high risk of severe COVID-19 and the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations for their treatment, see <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u>.

Age

Among all children, infants and adolescents have the highest risk of COVID-19-related ICU admission or death. From March 2020 to mid-August 2021, U.S. children aged <5 years had the highest cumulative

COVID-19-related hospitalization rates, followed closely by adolescents.³¹ Children aged 5 to 11 years had the lowest hospitalization rates. From July to August 2021, when the Delta variant was the dominant variant, 25% of 713 children admitted to 6 U.S. hospitals were aged <1 year, 17% were aged 1 to 4 years, 20% were aged 5 to 11 years, and 38% were aged 12 to 17 years.²⁶ From March 2020 to mid-June 2021, 26.5% of 3,116 U.S. children hospitalized for COVID-19 were admitted to an ICU.³¹

An individual patient data meta-analysis reported that patients aged <1 year and those aged 10 to 14 years had the highest risks of ICU admission and death among hospitalized children with COVID-19.³⁰ Another meta-analysis reported that neonates, but not infants aged 1 to 3 months, had an increased risk of severe COVID-19 compared with other pediatric age groups.²⁷ When Omicron was the dominant circulating variant, hospitalization rates among children and adolescents were higher than when the Delta variant was dominant, and they were highest for children aged <5 years.^{25,32} However, the proportion of hospitalized children who required ICU admission was significantly lower when the Omicron variant was dominant.

Comorbidities

Several chronic conditions are prevalent in hospitalized children with COVID-19. When the Delta variant was the dominant variant in the United States, 68% of hospitalized children had \geq 1 underlying medical conditions, such as obesity (32%), asthma or reactive airway disease (16%), or feeding tube dependence (8%). Obesity was present in approximately a third of hospitalized children aged 5 to 11 years, 60% of whom had a body mass index (BMI) \geq 120% of the 95th percentile. For adolescents, 61% had obesity; of those patients, 61% had a BMI \geq 120% of the 95th percentile.²⁶

Meta-analyses and observational studies identified risk factors for ICU admission, mechanical ventilation, or death among hospitalized children with COVID-19.²⁷⁻²⁹ These risk factors included prematurity in young infants, obesity, diabetes, chronic lung disease, cardiac disease, neurologic disease, and immunocompromising conditions. Another study found that having a complex chronic condition that affected ≥ 2 body systems or having a progressive chronic condition or continuous dependence on technology for ≥ 6 months (e.g., dialysis, tracheostomy with ventilator assistance) was significantly associated with an increased risk of moderate or severe COVID-19.³³ The study also found that children with hospitalization within the previous 12 months) had a higher risk of critical COVID-19 or death than those with less severe conditions. The CDC has additional information on the underlying conditions that are risk factors for severe COVID-19.

Having multiple comorbidities increases the risk of severe COVID-19 in children. A meta-analysis of data from children hospitalized with COVID-19 found that the risk of ICU admission was greater for children with 1 chronic condition than for those with no comorbidities, and the risk increased substantially as the number of comorbidities increased.³⁰

COVID-19 Vaccination

Staying up to date with COVID-19 vaccinations remains the most effective way to prevent severe COVID-19. See the CDC webpages <u>Stay Up to Date With COVID-19 Vaccines</u> and <u>Use of COVID-19</u> <u>Vaccines in the United States</u> for more information on COVID-19 vaccination schedules.

The estimates for vaccine effectiveness against severe COVID-19 in adolescents aged 12 to 18 years exceeded 90% while the Delta variant was the dominant variant in the United States.^{34,35} When Omicron was the dominant variant, vaccine effectiveness against hospitalization for noncritical COVID-19 was 20% in adolescents; vaccine effectiveness against critical illness was 79% in these patients.³⁵

79

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

In children aged 5 to 11 years, vaccine effectiveness against hospitalization was more variable, with an estimated effectiveness of 68% after Omicron became the dominant variant in the United States. An Italian study estimated that vaccine effectiveness was 38% in this group of children during this period.^{35,36}

See Prevention of SARS-CoV-2 Infection for more information about COVID-19 vaccines.

Mortality

Death from COVID-19 is uncommon in children. Risk factors for death include having chronic conditions, such as neurologic or cardiac disease, and having multiple comorbidities. Among children aged <21 years in the United States, the number of deaths associated with COVID-19 has been higher for children aged 10 to 20 years, especially for young adults aged 18 to 20 years, and for those who identify as Hispanic, Black, or American Indian/Alaskan Native.^{37,38}

A systematic review and meta-analysis reported that neurologic or cardiac comorbidities were associated with the greatest increase in risk of death among hospitalized children with COVID-19.³⁰ In the same study, an individual patient data meta-analysis found that the risk of COVID-19-related death was greater for children with 1 chronic condition than for those with no comorbidities, and the risk increased substantially as the number of comorbidities increased.

Vertical Transmission and Infants Born to People With SARS-CoV-2 Infection

Systematic reviews and meta-analyses have reported that confirmed vertical transmission of SARS-CoV-2 appears to be rare, and severe maternal COVID-19 has been associated with SARS-CoV-2 infection in babies.³⁹ In 2 large, combined cohorts of pregnant individuals from the United States and United Kingdom, SARS-CoV-2 infection was reported in 1.8% and 2% of the babies born to people with SARS-CoV-2 infection.⁴⁰ A systematic review and meta-analysis of prospective observational studies from high-income countries estimated that the frequency of SARS-CoV-2 infection in infants born to people with SARS-CoV-2 infection is 2.3%.⁴¹

Case reports have described intrauterine fetal demise during the third trimester of pregnancy in individuals with mild COVID-19 due to infection with the Delta variant.^{42,43} These individuals had evidence of placental SARS-CoV-2 infection, placental malperfusion, and placental inflammation. One case report described a person with asymptomatic SARS-CoV-2 infection and severe preeclampsia who gave birth at 25 weeks of gestation by emergency cesarean delivery. The neonate died on Day 4, and evidence of SARS-CoV-2 infection was found in placental tissues and in the infant's lungs and vascular endothelium at autopsy.⁴⁴ Evidence of placental SARS-CoV-2 infection was reported in 5 stillbirths and for 1 live-born neonate in Sweden.⁴⁵

A systematic review of neonatal SARS-CoV-2 infections reported that 70% were due to postpartum transmission, and 30% were due to vertical transmission from an infected birth parent.⁴⁶ Two systematic reviews reported that newborn infants rooming-in with the birth parent did not have an increased risk of SARS-CoV-2 transmission when compared with newborns who were isolated from the birth parent.^{41,47}

Detection of SARS-CoV-2 RNA in the breast milk of individuals with confirmed cases of COVID-19 is very uncommon.⁴⁸ Currently, there is no evidence of SARS-CoV-2 transmission through breast milk.⁴⁹ Breast milk from people with SARS-CoV-2 infection can contain antibodies to SARS-CoV-2.^{50,51} For information regarding the safety of feeding infants breast milk from individuals who are receiving treatment for COVID-19, see <u>Pregnancy, Lactation, and COVID-19 Therapeutics</u>.

Multisystem Inflammatory Syndrome in Children

A small subset of children and young adults with SARS-CoV-2 infection, including those with *COVID-19 Treatment Guidelines*

asymptomatic infection, may develop MIS-C. This syndrome is also called pediatric inflammatory multisystem syndrome—temporally associated with SARS-CoV-2 (PIMS-TS). Although the case definitions for these syndromes differ slightly, they are likely the same disease. The syndrome was first described in Europe, where previously healthy children with severe inflammation and Kawasaki disease-like features were identified as having current or recent infection with SARS-CoV-2.

The clinical spectrum of MIS-C has been described in the United States and is similar to that described for PIMS-TS. MIS-C is consistent with a postinfectious inflammatory syndrome related to SARS-CoV-2.^{52,53} Most patients with MIS-C have serologic evidence of previous SARS-CoV-2 infection, but only a minority have had a positive reverse transcription polymerase chain reaction (RT-PCR) result for SARS-CoV-2 at presentation.^{54,55}

The peak population-based incidence of MIS-C lags about 4 weeks behind the peak of acute pediatric COVID-19-related hospitalizations. Adults may develop a similar syndrome called multisystem inflammatory syndrome in adults (MIS-A), although it is not clear if this postinfectious complication is similar to MIS-C.⁵⁴⁻⁵⁶ Published data that characterize the condition are limited.

Although risk factors for the development of MIS-C have not been established, an analysis of MIS-C cases in the United States found that ICU admission was more likely for patients aged 6 to 12 years than for younger children, and it was more likely for children who identified as non-Hispanic Black than for those who identified as non-Hispanic White.⁵⁷ Unlike most children who present with severe COVID-19, the majority of children who present with MIS-C do not seem to have common underlying comorbidities other than obesity.⁵⁷ In addition, children whose deaths were related to MIS-C were less likely to have underlying medical conditions than children who died of COVID-19.³⁸

Several studies have suggested that COVID-19 vaccination protects against the development of MIS-C.^{58,59} The development of MIS-C after COVID-19 vaccination is very rare.^{58,60} Following the emergence of the Omicron variant, the incidence of MIS-C and the clinical severity of MIS-C have declined.^{61,62} This decline may be a result of several factors; for example, more children have now received COVID-19 vaccines and had prior exposure to SARS-CoV-2, both of which may provide some protection against MIS-C. In addition, the Omicron viral genome is less likely to trigger hyperinflammation than the viral genomes of other SARS-CoV-2 variants.

Clinical Manifestations of Multisystem Inflammatory Syndrome in Children

The CDC and the Council of State and Territorial Epidemiologists (CSTE) issued an updated case definition for MIS-C on January 1, 2023.⁶³ The 2023 CSTE/CDC Surveillance Case Definition for MIS-C is an individual aged <21 years who:

- Presents with fever,^a laboratory evidence of inflammation,^b and illness with a clinical severity that requires hospitalization or results in death, with new-onset clinical manifestations in ≥2 categories (i.e., cardiac, shock, hematologic, gastrointestinal, dermatologic)^c; and
- Does not have a more likely alternative diagnosis; and
- Has a positive viral test result from:
 - Either a molecular test that detects SARS-CoV-2 RNA or a SARS-CoV-2 antigen test up to 60 days prior to or during hospitalization or in a post-mortem specimen; *or*
 - A test that detects SARS-CoV-2-specific antibodies associated with current illness; or
- Has a close contact with a confirmed or probable case of COVID-19 in the 60 days prior to hospitalization; *or*

• Has a death certificate that lists MIS-C as an underlying cause of death or a significant condition COVID-19 Treatment Guidelines

contributing to death.

^a Subjective or documented fever $\geq 38.0^{\circ}$ C.

^b C-reactive protein level \geq 3.0 mg/dL (30 mg/L).

^c See Table A for a list of categories for these organ manifestations.

Table A. Clinical Manifestation Criteria for the 2023 CSTE/CDC MIS-C Surveillance Case Definition

Clinical Manifestation	Criteria
Cardiac Involvement	Left ventricular ejection fraction <55%
	 Coronary artery dilatation, aneurysm, or ectasia
	• Troponin levels elevated above laboratory normal range or indicated as elevated in a clinical note
Shock	Clinician diagnosis, as documented in clinical note
Hematologic Involvement	 Thrombocytopenia (i.e., platelet count <150,000 cells/µL)
	 Lymphopenia (i.e., absolute lymphocyte count <1,000 cells/µL)
Gastrointestinal Involvement	Abdominal pain
	Vomiting
	• Diarrhea
Dermatologic/Mucocutaneous Involvement	• Rash
	Inflammation of the oral mucosa
	Conjunctivitis or conjunctival injection
	Extremity findings (e.g., erythema, edema)

Key: CDC = Centers for Disease Control and Prevention; CSTE = Council of State and Territorial Epidemiologists; MIS-C = multisystem inflammatory syndrome in children

Distinguishing MIS-C from other febrile illnesses in the community setting remains challenging, but the presence of persistent fever, multisystem manifestations, and laboratory abnormalities could help early recognition.⁶⁴ The clinical spectrum of hospitalized cases has included younger children with mucocutaneous manifestations that overlap with Kawasaki disease, older children with more multiorgan involvement and shock, and patients with respiratory manifestations that overlap with COVID-19.

Patients with MIS-C are often critically ill, and up to 80% of children require ICU admission; however, data collected while Omicron was the dominant variant in the United States suggest that the cases of MIS-C reported during this period were less severe than those reported when other variants were dominant.^{61,65} Most patients with MIS-C have markers of cardiac injury or dysfunction, including elevated levels of troponin and brain natriuretic protein; higher levels of these markers are associated with ICU admission, myocardial dysfunction, and shock.⁵⁷ In these cases, echocardiographic findings may include impaired left ventricular function, coronary artery dilations, and, rarely, coronary artery aneurysms. During the period when Omicron was the dominant variant in the United States, the clinical phenotype of MIS-C appeared to be more consistent with classic Kawasaki disease.^{61,65} The reported mortality in the United States for hospitalized children with MIS-C is 1% to 2%. Longitudinal studies to examine the long-term sequelae of MIS-C are currently ongoing.

The pathogenesis of MIS-C is still being elucidated and may include distinct humoral immune responses, innate immune activation, or a superantigen effect. Differences between MIS-C and typical Kawasaki disease have been demonstrated in terms of epidemiology, cytopenias, cytokine expression, and elevation

COVID-19 Treatment Guidelines

of inflammatory markers. Immunologic profiling has also shown differences in cytokine expression (tumor necrosis factor-alpha and interleukin-10) between MIS-C and COVID-19 in children.⁶⁶⁻⁶⁸

For the Panel's recommendations on the treatment of MIS-C, see <u>Therapeutic Management of</u> <u>Hospitalized Children With MIS-C, Plus a Discussion on MIS-A</u>.

Post-COVID Conditions

The persistent symptoms after COVID-19 that have been described in children are similar to those seen in adults. The terminology for these collective symptoms is evolving and includes long COVID, post-COVID-19 condition, and post-acute sequelae of SARS-CoV-2 infection (PASC). The data on the incidence of post-COVID conditions in children are limited and somewhat conflicting, but the overall incidence appears to be lower in children than in adults (see <u>Clinical Spectrum of SARS-CoV-2</u> Infection).⁶⁹⁻⁷³ However, given the high overall rate of SARS-CoV-2 infection in children, the burden of post-COVID conditions in children may be quite large.

Case definitions for post-COVID conditions vary between studies, which makes determining the true incidence of these conditions challenging. The incidence of post-COVID symptoms in children appears to increase with age. The most common symptoms reported include persistent fatigue, headache, shortness of breath, sleep disturbances, gastrointestinal symptoms, and an altered sense of smell.⁷⁴ Cardiopulmonary injury, neurocognitive impairment, and new-onset diabetes may occur. However, some studies did not include control groups of people who did not have SARS-CoV-2 infection, and this makes it challenging to assess the relative risk of these symptoms.

Details on the pathogenesis, clinical presentation, and treatment for post-COVID conditions in children are beyond the scope of these Guidelines. The CDC provides <u>additional information</u> about the incidence, presentation, and management strategies for post-COVID conditions in children as well as adults. Additional research is needed to define the incidence, pathophysiology, spectrum, and severity of post-COVID conditions in children and to identify the optimal strategies for the prevention, diagnosis, and treatment of these conditions.

References

- 1. Centers for Disease Control and Prevention. COVID-19 weekly cases and deaths per 100,000 population by age, race/ethnicity, and sex. 2022. Available at: <u>https://covid.cdc.gov/covid-data-tracker/#demographicsovertime</u>. Accessed June 23, 2023.
- Centers for Disease Control and Prevention. Provisional COVID-19 deaths: focus on ages 0–18 years. 2022. Available at: <u>https://data.cdc.gov/NCHS/Provisional-COVID-19-Deaths-Focus-on-Ages-0-18-Yea/nr4s-juj3</u>. Accessed June 23, 2023.
- Centers for Disease Control and Prevention. Demographic trends of COVID-19 cases and deaths in the US reported to CDC. 2022. Available at: <u>https://covid.cdc.gov/covid-data-tracker/#demographics</u>. Accessed June 23, 2023.
- 4. Centers for Disease Control and Prevention. COVID-NET laboratory-confirmed COVID-19 hospitalizations. 2022. Available at: <u>https://covid.cdc.gov/covid-data-tracker/#covidnet-hospitalization-network</u>. Accessed June 23, 2023.
- Clarke KEN, Jones JM, Deng Y, et al. Seroprevalence of infection-induced SARS-CoV-2 antibodies—United States, September 2021–February 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(17):606-608. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35482574</u>.
- 6. Dong Y, Mo X, Hu Y, et al. Epidemiology of COVID-19 among children in China. *Pediatrics*. 2020;145(6):e20200702. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32179660</u>.

- CDC COVID-19 Response Team. Coronavirus disease 2019 in children—United States, February 12–April 2, 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(14):422-426. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/32271728</u>.
- 8. Cui X, Zhang T, Zheng J, et al. Children with coronavirus disease 2019: a review of demographic, clinical, laboratory, and imaging features in pediatric patients. *J Med Virol*. 2020;92(9):1501-1510. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32418216.
- 9. Livingston E, Bucher K. Coronavirus disease 2019 (COVID-19) in Italy. *JAMA*. 2020;323(14):1335. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32181795.
- Tagarro A, Epalza C, Santos M, et al. Screening and severity of coronavirus disease 2019 (COVID-19) in children in Madrid, Spain. *JAMA Pediatr*. 2020;Published online ahead of print. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32267485</u>.
- DeBiasi RL, Song X, Delaney M, et al. Severe coronavirus disease-2019 in children and young adults in the Washington, DC, metropolitan region. *J Pediatr*. 2020;223:199-203.e1. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32405091</u>.
- 12. Chao JY, Derespina KR, Herold BC, et al. Clinical characteristics and outcomes of hospitalized and critically ill children and adolescents with coronavirus disease 2019 at a tertiary care medical center in New York City. J Pediatr. 2020;223:14-19.e2. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32407719</u>.
- Swann OV, Holden KA, Turtle L, et al. Clinical characteristics of children and young people admitted to hospital with COVID-19 in United Kingdom: prospective multicentre observational cohort study. *BMJ*. 2020;370:m3249. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32960186</u>.
- Götzinger F, Santiago-García B, Noguera-Julián A, et al. COVID-19 in children and adolescents in Europe: a multinational, multicentre cohort study. *Lancet Child Adolesc Health*. 2020;4(9):653-661. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32593339</u>.
- Shi DS, Whitaker M, Marks KJ, et al. Hospitalizations of children aged 5–11 years with laboratory-confirmed COVID-19—COVID-NET, 14 states, March 2020–February 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(16):574-581. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35446827</u>.
- 16. Siegel DA, Reses HE, Cool AJ, et al. Trends in COVID-19 cases, emergency department visits, and hospital admissions among children and adolescents aged 0–17 years—United States, August 2020–August 2021. MMWR Morb Mortal Wkly Rep. 2021;70(36):1249-1254. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34499628</u>.
- Marks KJ, Whitaker M, Anglin O, et al. Hospitalizations of children and adolescents with laboratoryconfirmed COVID-19—COVID-NET, 14 states, July 2021–January 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(7):271-278. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35176003</u>.
- Holmes L Jr, Wu C, Hinson R, et al. Black-White risk differentials in pediatric COVID-19 hospitalization and intensive care unit admissions in the USA. *J Racial Ethn Health Disparities*. 2023;10(3):1187-1193. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35604543</u>.
- Antoon JW, Grijalva CG, Thurm C, et al. Factors associated with COVID-19 disease severity in US children and adolescents. *J Hosp Med*. 2021;16(10):603-610. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34613896.
- 20. Saatci D, Ranger TA, Garriga C, et al. Association between race and COVID-19 outcomes among 2.6 million children in England. *JAMA Pediatr.* 2021;175(9):928-938. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34152371</u>.
- Poline J, Gaschignard J, Leblanc C, et al. Systematic severe acute respiratory syndrome coronavirus 2 screening at hospital admission in children: a French prospective multicenter study. *Clin Infect Dis*. 2021;72(12):2215-2217. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32710743</u>.
- 22. Kim L, Whitaker M, O'Halloran A, et al. Hospitalization rates and characteristics of children aged <18 years hospitalized with laboratory-confirmed COVID-19—COVID-NET, 14 states, March 1–July 25, 2020.

MMWR Morb Mortal Wkly Rep. 2020;69(32):1081-1088. Available at: <u>https://www.ncbi.nlm.nih.gov/</u>pubmed/32790664.

- 23. Feldstein LR, Tenforde MW, Friedman KG, et al. Characteristics and outcomes of US children and adolescents with multisystem inflammatory syndrome in children (MIS-C) compared with severe acute COVID-19. *JAMA*. 2021;325(11):1074-1087. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33625505</u>.
- 24. Bhalala US, Gist KM, Tripathi S, et al. Characterization and outcomes of hospitalized children with coronavirus disease 2019: a report from a multicenter, viral infection and respiratory illness universal study (coronavirus disease 2019) registry. *Crit Care Med*. 2022;50(1):e40-e51. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34387240.
- 25. Marks KJ, Whitaker M, Agathis NT, et al. Hospitalization of infants and children aged 0–4 years with laboratory-confirmed COVID-19—COVID-NET, 14 states, March 2020–February 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(11):429-436. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35298458.
- Wanga V, Gerdes ME, Shi DS, et al. Characteristics and clinical outcomes of children and adolescents aged <18 years hospitalized with COVID-19—six hospitals, United States, July–August 2021. *MMWR Morb Mortal Wkly Rep.* 2021;70(5152):1766-1772. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34968374</u>.
- 27. Choi JH, Choi SH, Yun KW. Risk factors for severe COVID-19 in children: a systematic review and metaanalysis. *J Korean Med Sci.* 2022;37(5):e35. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35132841</u>.
- 28. Shi Q, Wang Z, Liu J, et al. Risk factors for poor prognosis in children and adolescents with COVID-19: a systematic review and meta-analysis. *EClinicalMedicine*. 2021;41:101155. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34693233.
- 29. Kompaniyets L, Agathis NT, Nelson JM, et al. Underlying medical conditions associated with severe COVID-19 illness among children. *JAMA Netw Open*. 2021;4(6):e2111182. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34097050.
- 30. Harwood R, Yan H, Talawila Da Camara N, et al. Which children and young people are at higher risk of severe disease and death after hospitalisation with SARS-CoV-2 infection in children and young people: a systematic review and individual patient meta-analysis. *EClinicalMedicine*. 2022;44:101287. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35169689</u>.
- Delahoy MJ, Ujamaa D, Whitaker M, et al. Hospitalizations associated with COVID-19 among children and adolescents—COVID-NET, 14 states, March 1, 2020–August 14, 2021. *MMWR Morb Mortal Wkly Rep.* 2021;70(36):1255-1260. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34499627</u>.
- 32. Dorabawila V, Hoefer D, Bauer UE, et al. Risk of infection and hospitalization among vaccinated and unvaccinated children and adolescents in New York after the emergence of the Omicron variant. *JAMA*. 2022;327(22):2242-2244. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35559959.
- 33. Forrest CB, Burrows EK, Mejias A, et al. Severity of acute COVID-19 in children <18 years old March 2020 to December 2021. *Pediatrics*. 2022;149(4):e2021055765. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35322270.
- 34. Olson SM, Newhams MM, Halasa NB, et al. Effectiveness of BNT162b2 vaccine against critical COVID-19 in adolescents. *N Engl J Med.* 2022;386(8):713-723. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35021004.
- Price AM, Olson SM, Newhams MM, et al. BNT162b2 protection against the Omicron variant in children and adolescents. *N Engl J Med*. 2022;386(20):1899-1909. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35353976/</u>.
- 36. Sacco C, Del Manso M, Mateo-Urdiales A, et al. Effectiveness of BNT162b2 vaccine against SARS-CoV-2 infection and severe COVID-19 in children aged 5-11 years in Italy: a retrospective analysis of January-April, 2022. *Lancet*. 2022;400(10346):97-103. Available at: https://pubmed.ncbi.nlm.nih.gov/35780801/.
- 37. Bixler D, Miller AD, Mattison CP, et al. SARS-CoV-2-associated deaths among persons aged <21 years— United States, February 12–July 31, 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(37):1324-1329. Available

at: https://www.ncbi.nlm.nih.gov/pubmed/32941417.

- 38. McCormick DW, Richardson LC, Young PR, et al. Deaths in children and adolescents associated with COVID-19 and MIS-C in the United States. *Pediatrics*. 2021;148(5):e2021052273. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34385349.
- 39. Allotey J, Chatterjee S, Kew T, et al. SARS-CoV-2 positivity in offspring and timing of mother-to-child transmission: living systematic review and meta-analysis. *BMJ*. 2022;376:e067696. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35296519.
- 40. Mullins E, Hudak ML, Banerjee J, et al. Pregnancy and neonatal outcomes of COVID-19: coreporting of common outcomes from PAN-COVID and AAP-SONPM registries. *Ultrasound Obstet Gynecol.* 2021;57(4):573-581. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33620113</u>.
- 41. Morniroli D, Vizzari G, Tosi M, et al. Mother-to-child transmission of SARS-CoV-2 infection in high-income countries: a systematic review and meta-analysis of prospective observational studies. *Sci Rep.* 2023;13(1):8813. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/37258854/</u>.
- 42. Shook LL, Brigida S, Regan J, et al. SARS-CoV-2 placentitis associated with B.1.617.2 (Delta) variant and fetal distress or demise. *J Infect Dis.* 2022;225(5):754-758. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35024844.
- 43. Guan M, Johannesen E, Tang CY, et al. Intrauterine fetal demise in the third trimester of pregnancy associated with mild infection with the SARS-CoV-2 Delta variant without protection from vaccination. *J Infect Dis*. 2022;225(5):748-753. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35024853</u>.
- 44. Reagan-Steiner S, Bhatnagar J, Martines RB, et al. Detection of SARS-CoV-2 in neonatal autopsy tissues and placenta. *Emerg Infect Dis.* 2022;28(3):510-517. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35138244.
- 45. Zaigham M, Gisselsson D, Sand A, et al. Clinical-pathological features in placentas of pregnancies with SARS-CoV-2 infection and adverse outcome: case series with and without congenital transmission. *BJOG*. 2022;129(8):1361-1374. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35243759</u>.
- 46. Raschetti R, Vivanti AJ, Vauloup-Fellous C, et al. Synthesis and systematic review of reported neonatal SARS-CoV-2 infections. *Nat Commun.* 2020;11(1):5164. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33060565</u>.
- Walker KF, O'Donoghue K, Grace N, et al. Maternal transmission of SARS-CoV-2 to the neonate, and possible routes for such transmission: a systematic review and critical analysis. *BJOG*. 2020;127(11):1324-1336. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32531146</u>.
- 48. Kumar J, Meena J, Yadav A, Kumar P. SARS-CoV-2 detection in human milk: a systematic review. *J Matern Fetal Neonatal Med.* 2021;35(25):5456-5463. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33550866</u>.
- Centeno-Tablante E, Medina-Rivera M, Finkelstein JL, et al. Transmission of SARS-CoV-2 through breast milk and breastfeeding: a living systematic review. *Ann N Y Acad Sci.* 2021;1484(1):32-54. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32860259</u>.
- 50. Bäuerl C, Randazzo W, Sánchez G, et al. SARS-CoV-2 RNA and antibody detection in breast milk from a prospective multicentre study in Spain. *Arch Dis Child Fetal Neonatal Ed.* 2022;107(2):216-221. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34417223</u>.
- Duncombe CJ, McCulloch DJ, Shuey KD, et al. Dynamics of breast milk antibody titer in the six months following SARS-CoV-2 infection. *J Clin Virol*. 2021;142:104916. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34315010</u>.
- 52. Riphagen S, Gomez X, Gonzalez-Martinez C, Wilkinson N, Theocharis P. Hyperinflammatory shock in children during COVID-19 pandemic. *Lancet*. 2020;395(10237):1607-1608. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32386565.
- 53. Whittaker E, Bamford A, Kenny J, et al. Clinical characteristics of 58 children with a pediatric inflammatory multisystem syndrome temporally associated with SARS-CoV-2. *JAMA*. 2020;324(3):259-269. Available at:

https://www.ncbi.nlm.nih.gov/pubmed/32511692.

- 54. Dufort EM, Koumans EH, Chow EJ, et al. Multisystem inflammatory syndrome in children in New York State. *N Engl J Med*. 2020;383(4):347-358. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32598830</u>.
- 55. Feldstein LR, Rose EB, Horwitz SM, et al. Multisystem inflammatory syndrome in U.S. children and adolescents. *N Engl J Med*. 2020;383(4):334-346. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32598831.
- 56. Morris SB, Schwartz NG, Patel P, et al. Case series of multisystem inflammatory syndrome in adults associated with SARS-CoV-2 infection—United Kingdom and United States, March–August 2020. MMWR Morb Mortal Wkly Rep. 2020;69(40):1450-1456. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33031361.
- 57. Abrams JY, Oster ME, Godfred-Cato SE, et al. Factors linked to severe outcomes in multisystem inflammatory syndrome in children (MIS-C) in the USA: a retrospective surveillance study. *Lancet Child Adolesc Health*. 2021;5(5):323-331. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33711293.
- 58. Zambrano LD, Newhams MM, Olson SM, et al. Effectiveness of BNT162b2 (Pfizer-BioNTech) mRNA vaccination against multisystem inflammatory syndrome in children among persons aged 12–18 years— United States, July–December 2021. MMWR Morb Mortal Wkly Rep. 2022;71(2):52-58. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35025852.
- 59. Levy M, Recher M, Hubert H, et al. Multisystem inflammatory syndrome in children by COVID-19 vaccination status of adolescents in France. *JAMA*. 2022;327(3):281-283. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34928295</u>.
- 60. Yousaf AR, Cortese MM, Taylor AW, et al. Reported cases of multisystem inflammatory syndrome in children aged 12–20 years in the USA who received a COVID-19 vaccine, December, 2020, through August, 2021: a surveillance investigation. *Lancet Child Adolesc Health*. 2022;6(5):303-312. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35216660.
- 61. McCrindle BW, Harahsheh AS, Handoko R, et al. SARS-CoV-2 variants and multisystem inflammatory syndrome in children. *N Engl J Med*. 2023;388(17):1624-1626. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36947454/</u>.
- 62. Cohen JM, Carter MJ, Cheung CR, Ladhani S. Lower risk of multisystem inflammatory syndrome in children with the Delta and Omicron variants of severe acute respiratory syndrome coronavirus 2. *Clin Infect Dis*. 2023;76(3):e518-e521. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35788276/</u>.
- 63. Centers for Disease Control and Prevention. Information for healthcare providers about multisystem inflammatory syndrome in children (MIS-C). 2023. Available at: <u>https://www.cdc.gov/mis/mis-c/hcp_cstecdc/index.html</u>. Accessed June 23, 2023.
- 64. Carlin RF, Fischer AM, Pitkowsky Z, et al. Discriminating multisystem inflammatory syndrome in children requiring treatment from common febrile conditions in outpatient settings. *J Pediatr*. 2021;229:26-32.e2. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33065115</u>.
- 65. Godfred-Cato S, Bryant B, Leung J, et al. COVID-19-associated multisystem inflammatory syndrome in children—United States, March–July 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(32):1074-1080. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32790663</u>.
- 66. Lee PY, Day-Lewis M, Henderson LA, et al. Distinct clinical and immunological features of SARS-CoV-2induced multisystem inflammatory syndrome in children. *J Clin Invest*. 2020;130(11):5942-5950. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32701511</u>.
- 67. Rowley AH, Shulman ST, Arditi M. Immune pathogenesis of COVID-19-related multisystem inflammatory syndrome in children. *J Clin Invest*. 2020;130(11):5619-5621. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32870815</u>.
- 68. Diorio C, Henrickson SE, Vella LA, et al. Multisystem inflammatory syndrome in children and COVID-19 are distinct presentations of SARS-CoV-2. *J Clin Invest*. 2020;130(11):5967-5975. Available at: <u>https://www.ncbi.</u>

nlm.nih.gov/pubmed/32730233.

- 69. Zimmermann P, Pittet LF, Curtis N. How common is long COVID in children and adolescents? *Pediatr Infect Dis J*. 2021;40(12):e482-e487. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34870392</u>.
- 70. Zimmermann P, Pittet LF, Curtis N. Long COVID in children and adolescents. *BMJ*. 2022;376:o143. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35058281</u>.
- 71. Molteni E, Sudre CH, Canas LS, et al. Illness duration and symptom profile in symptomatic UK school-aged children tested for SARS-CoV-2. *Lancet Child Adolesc Health*. 2021;5(10):708-718. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34358472</u>.
- 72. Zheng YB, Zeng N, Yuan K, et al. Prevalence and risk factor for long COVID in children and adolescents: a meta-analysis and systematic review. J Infect Public Health. 2023;16(5):660-672. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36931142/</u>.
- 73. Pinto Pereira SM, Mensah A, Nugawela MD, et al. Long COVID in children and youth after injection or reinfection with the Omicron variant: a prospective observational study. *J Pediatr*. 2023;Published online ahead of print. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/37172813/</u>.
- 74. Fainardi V, Meoli A, Chiopris G, et al. Long COVID in children and adolescents. *Life (Basel)*. 2022;12(2):285. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35207572</u>.

Therapeutic Management of Nonhospitalized Children With COVID-19

Last Updated: December 28, 2022

This section outlines the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations for the therapeutic management of nonhospitalized children (i.e., pediatric patients aged <18 years) with mild to moderate COVID-19. These recommendations are also for children who have mild to moderate COVID-19 and are hospitalized for reasons other than COVID-19. For patients aged \geq 18 years, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>. Throughout this section, the term "COVID-19" refers to the acute, primarily respiratory illness caused by infection with SARS-CoV-2. For the Panel's recommendations for managing multisystem inflammatory syndrome in children (MIS-C), see <u>Therapeutic Management of Hospitalized Children With MIS-C, Plus a Discussion on MIS-A</u>.

Treatment Considerations for Children With COVID-19

Currently, no results from pediatric clinical trials that evaluated the treatment of COVID-19 have been published. Data evaluating the use of pharmacologic therapy in children with COVID-19 are limited largely to descriptive reports.^{1,2} Therefore, more high-quality randomized trials, observational studies, and pharmacokinetic studies are urgently needed. Whenever possible, clinical trials of therapeutics and multicenter observational cohorts should enroll children with COVID-19.

The current recommendations for treating COVID-19 in children have been mostly extrapolated from recommendations for adults with COVID-19, recommendations for children with other viral infections, and expert opinion.³⁻⁵ Applying adult data from COVID-19 trials to children is a unique challenge, because most children experience a mild course of illness with COVID-19. Relative to adults, children with COVID-19 have substantially lower mortality and less need for hospitalization. Because of these differences in epidemiology and disease severity, the effect sizes for children are likely to be smaller than those observed in adults; therefore, to produce a beneficial outcome, the number needed to treat is higher. Collectively, these factors influence the risk versus benefit balance for pharmacologic therapies in children.

Other challenges are the uncertainty about which comorbid conditions place children at the highest risk of severe COVID-19 and the uncertainty about the absolute magnitude of the increased risk from those comorbid conditions. Clinicians need to consider the number and severity of a child's comorbid conditions when making decisions about pharmacologic treatments for COVID-19. For more information on risk factors for children with COVID-19, see <u>Special Considerations in Children</u>.

Recommendations

In the absence of sufficient clinical trial data on the treatment of children with COVID-19, the Panel's recommendations for the therapeutic management of nonhospitalized children are based largely on adult safety and efficacy data from clinical trials (see Table 3a). No pediatric comparative studies have been published; therefore, all the quality of evidence ratings for the Panel's recommendations in this section are based on expert opinion (i.e., a **III** rating).

The majority of children with mild to moderate COVID-19 will not progress to more severe illness; therefore, the Panel recommends managing these patients with supportive care alone (**AIII**). The risks and benefits of therapy should be assessed based on COVID-19 disease severity, age, vaccination status,

and the presence of underlying medical conditions that may place the patient at high risk of severe COVID-19.

Disk of Course OOUID 40	Panel's Recommendations	
Risk of Severe COVID-19	Aged 12–17 Years	Aged <12 Years
Symptomatic, Regardless of Risk Factors	• Provide supportive care (AIII).	• Provide supportive care (AIII).
High Risk ^{a,b}	 Use 1 of the following options (listed in order of preference):^c Ritonavir-boosted nirmatrelvir (Paxlovid) within 5 days of symptom onset (BIII) Remdesivir within 7 days of symptom onset (CIII) 	 Ritonavir-boosted nirmatrelvir is not authorized by the FDA for use in children aged <12 years. There is insufficient evidence to recommend either for or against the routine use of remdesivir. Consider treatment based on age and other risk factors.
Intermediate Risk ^{b,d}	• There is insufficient evidence to recommend either for or against the use of any antiviral therapy. Consider treatment based on age and other risk factors.	• There is insufficient evidence to recommend either for or against the routine use of remdesivir.
Low Risk ^{b,e}	 Manage with supportive care alone (BIII). 	 Manage with supportive care alone (BIII).
Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating		

Table 3a. Therapeutic Management	of Nonhospitalized Childr	en With COVID-19
Table Sa. Therapeutic Management	of Nonnospitalized Children	

^a Molnupiravir is not authorized by the FDA for use in children aged <18 years and should not be used.

for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

^b See Table 3b for the Panel's framework for assessing the risk of progression to severe COVID-19 based on patient conditions and COVID-19 vaccination status.

° Initiate treatment as soon as possible after symptom onset.

^d The relative risk of severe COVID-19 for intermediate-risk patients is lower than the risk for high-risk patients but higher than the risk for low-risk patients.

^e Low-risk patients include those with comorbid conditions that have a weak or unknown association with severe COVID-19. Patients with no comorbidities are included in this group.

Key: FDA = Food and Drug Administration; the Panel = the COVID-19 Treatment Guidelines Panel

Table 3b. The Panel's Framework for Assessing the Risk of Progression to Severe COVID-19Based on Patient Conditions and COVID-19 Vaccination Status

Conditions	Risk Le	vel by Vaccination Sta	itus ^a
Conditions	Unvaccinated	Primary Series	Up to Date
Strong or Consistent Association With Progres	ssion to Severe COVID- [.]	19	
Moderately or severely immunocompromised (see <u>Special Considerations in People Who Are</u> <u>Immunocompromised</u>)	High		
 Obesity (BMI ≥95th percentile for age), especially severe obesity (BMI ≥120% of 95th percentile for age)^b 			
 Medical complexity with dependence on respiratory technology^c 		Intermediate	
 Severe neurologic, genetic, metabolic, or other disability that results in impaired airway clearance or limitations in self care or activities of daily living 	High		
 Severe asthma or other severe chronic lung disease requiring ≥2 inhaled or ≥1 systemic medications daily 			
Severe congenital or acquired cardiac disease			
Multiple moderate to severe chronic diseases			
Moderate or Inconsistent Association With Prog	ression to Severe COVIE)-19	
 Aged <1 year Prematurity in children aged ≤2 years 			
Sickle cell disease	Intermediate		
Diabetes mellitus (poorly controlled)			
 Nonsevere cardiac, neurologic, or metabolic disease^d 			
Weak or Unknown Association With Progression to Severe COVID-19			
Mild asthma			
Overweight	Low		
Diabetes mellitus (well controlled)			

^a **Unvaccinated** = individuals who are not eligible for COVID-19 vaccination or are <2 weeks from the final dose of the primary series. **Vaccinated with primary series** = individuals who completed the primary series of 2 or 3 doses (the current CDC term is "fully vaccinated") and are >2 weeks after the final dose of the primary series but have not received a booster, if they are eligible for a booster. Children aged <5 years are not currently eligible for booster doses. **Vaccinated and up to date** = individuals who received the recommended booster dose(s) if eligible or have completed the primary series but are not yet eligible for a booster. See the current <u>COVID-19 vaccination schedule</u> from the CDC for more information.

^b The degree of risk conferred by obesity in younger children is less clear than it is in older adolescents.

 $^{\circ}$ This includes patients with a tracheostomy and those who require NIV.

^d The data for this group are particularly limited.

Key: BMI = body mass index; CDC = Centers for Disease Control and Prevention; NIV = noninvasive ventilation; the Panel = the COVID-19 Treatment Guidelines Panel

Rationale for the Panel's Framework for Assessing the Risk of Progression to Severe COVID-19

Although mortality associated with COVID-19 in children is low overall, severe disease can occur, especially in those with risk factors.⁶ Risk stratification for severe disease in children remains challenging. Imprecise definitions of comorbid conditions, insufficient granularity for differentiating the severity of comorbidities (e.g., mild vs. severe lung disease, poorly controlled vs. well-controlled diabetes), and small sample sizes limit the conclusions that can be drawn from individual studies and make comparing findings across studies difficult.

Furthermore, asymptomatic SARS-CoV-2 infection detected during admission screening for children who are hospitalized for reasons other than COVID-19 may affect the estimated risk of severe COVID-19, particularly for patient groups that may have protocolized admissions (e.g., children with febrile neutropenia, infants aged <90 days with fever). In addition, the published studies that have evaluated these associations in children are limited largely to case series without control groups and observational studies with methodologic limitations.

Despite these challenges, a risk-stratification framework needs to be developed that will allow clinicians to identify the patients who are most likely to benefit from receiving treatment. These patients can be prioritized in situations where supply or logistical constraints make it impossible to offer therapy to all eligible patients. Both the Pediatric Infectious Diseases Society and the American Academy of Pediatrics advocate for a risk-stratified approach to identifying the patients who are at the highest risk of progression to severe COVID-19 among those eligible for therapies under Food and Drug Administration (FDA) Emergency Use Authorizations (EUAs).^{5,7}

The Panel's approach to risk stratification and prioritization considers COVID-19 vaccination status, immune function, clinical risk factors, the strength of the evidence that demonstrates an association between each clinical risk factor and severe disease, and expert opinion.^{6,8-21} See <u>Special Considerations</u> in <u>Children</u> for more information on clinical risk factors. The Panel suggests that decisions regarding treatment be individualized, particularly for patients in the intermediate risk category. Clinicians should consider the number and severity of comorbid conditions, the child's vaccination status, and the time since vaccination.

Comorbid conditions associated with severe COVID-19 are separated into the following categories in Table 3b:

- *Strong or Consistent Association With Progression to Severe COVID-19:* Comorbid conditions for which the published literature most consistently supports an increased risk of severe COVID-19. Patients in this category are moderately or severely immunocompromised, at risk of severe COVID-19, and not expected to develop an adequate immune response to COVID-19 vaccination.
- *Moderate or Inconsistent Association With Progression to Severe COVID-19:* Comorbid conditions and ages for which the published literature supports an association with severe COVID-19, but the association may be moderate or inconsistent across studies. In addition, the absolute risk of progression to severe disease or death is likely modest for any of the patients in this category.
- Weak or Unknown Association With Progression to Severe COVID-19: Comorbid conditions for which the data suggesting an association with severe COVID-19 are weak or for which an association is unknown. Patients with no comorbidities are included in this category.

Vaccination Status

Because COVID-19 vaccines are highly effective in preventing severe disease, individuals who are not immunocompromised and are up to date on their vaccines (i.e., those who have received the recommended booster dose[s], if eligible, or who have completed the primary series but are not yet eligible for a booster)²² are likely to have a low absolute risk of severe disease. Therefore, the potential benefit from antiviral treatment is less clear for these patients. Patients who have had the primary series of vaccinations (i.e., those who are fully vaccinated but not up to date) may have a lower level of protection against severe disease than patients who are up to date, but the data comparing these groups are limited. However, evidence suggests that vaccine protection against severe COVID-19 wanes over time, particularly protection against the Omicron variant of concern (VOC) and its subvariants.²³⁻²⁶ Clinicians should consider the time since a child's vaccination when making treatment decisions.

Health Disparities

COVID-19–related outcomes are worse among medically underserved populations, although this factor is not strictly a comorbid condition. Some racial and ethnic groups experience disproportionate rates of COVID-19 hospitalization and are less likely to receive specific therapies.²⁷⁻³⁰ These factors may be relevant when making clinical decisions about treatment.^{31,32} See <u>Special Considerations in Children</u> for more information.

Rationale for the Panel's Recommendations

Ritonavir-Boosted Nirmatrelvir (Paxlovid)

Ritonavir-boosted nirmatrelvir has received an FDA EUA for the treatment of mild to moderate COVID-19 in nonhospitalized adolescents aged \geq 12 years and weighing \geq 40 kg who are at high risk of progression to severe COVID-19.³³

The EPIC-HR trial enrolled adults aged ≥ 18 years who were at high risk of severe COVID-19; they were randomized to receive ritonavir-boosted nirmatrelvir or placebo. The primary outcome of COVID-19– related hospitalization or all-cause mortality occurred in 8 of 1,039 patients (0.8%) who received ritonavir-boosted nirmatrelvir and in 66 of 1,046 patients (6.3%) who received placebo, an 89% relative risk reduction.³⁴ No pediatric patients were included in the trial, and no pediatric safety data were made available.

Ritonavir has been used extensively in pediatric patients as a pharmacokinetic booster for the treatment of HIV and hepatitis C virus infection, and it has a known and tolerable side effect profile. In the FDA EUA, the dose of ritonavir-boosted nirmatrelvir authorized for adolescents aged ≥ 12 years and weighing ≥ 40 kg is expected to result in a drug exposure similar to that observed in adults.³³

Given the high efficacy of ritonavir-boosted nirmatrelvir in adults, its overall manageable side effect profile, the pediatric clinical experience with ritonavir, and the convenience of an oral medication, the Panel recommends the use of **ritonavir-boosted nirmatrelvir (Paxlovid)** for nonhospitalized adolescents aged \geq 12 years and weighing \geq 40 kg who have mild to moderate COVID-19 and are at the highest risk of progression to severe COVID-19 (**BIII**). Ritonavir-boosted nirmatrelvir is expected to be active against all Omicron subvariants, although clinical efficacy data are currently limited.³⁵⁻³⁷

Because of the potential for significant drug-drug interactions with some concomitant medications, ritonavir-boosted nirmatrelvir may not be the safest choice for some patients. See <u>Ritonavir-Boosted</u> <u>Nirmatrelvir (Paxlovid)</u> and <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid)</u> and <u>Concomitant Medications</u> for more information.

Remdesivir

Remdesivir is approved by the FDA for use in hospitalized and nonhospitalized pediatric patients aged \geq 28 days and weighing \geq 3.0 kg.³⁸ Remdesivir is expected to be active against the Omicron VOC and its subvariants, although clinical efficacy data are currently limited.^{37,39-41}

In a study that included nonhospitalized adults with mild to moderate COVID-19 who were at high risk of progression to severe disease, administering an intravenous (IV) infusion of remdesivir once daily for 3 days resulted in an 87% relative reduction in the risk of hospitalization or death when compared with placebo.⁴² Although adolescents aged \geq 12 years were eligible for inclusion, the trial included only 8 patients aged <18 years; therefore, no conclusions regarding the efficacy of remdesivir in children can be made from this trial. In addition, clinical experience data from hospitalized children with COVID-19 who received remdesivir through a compassionate use program have been reported.^{2,43} Given the demonstrated efficacy of remdesivir in the overall study population, its overall favorable side effect profile, and clinical experience with remdesivir in hospitalized children, **remdesivir**, as an alternative to ritonavir-boosted nirmatrelvir, can be considered for children aged \geq 12 years who are at the highest risk of progression to severe COVID-19 (**CIII**).

There is insufficient evidence to recommend either for or against the routine use of remdesivir for the treatment of COVID-19 in nonhospitalized children aged <12 years who are at the highest risk of progression to severe disease or who are at intermediate risk of severe disease. Administering remdesivir requires performing an IV infusion once daily for 3 days, so logistical constraints may preclude the use of remdesivir in many settings.

Pharmacologic Therapies Not Recommended for Nonhospitalized Children With COVID-19

Molnupiravir

The FDA EUA for molnupiravir is limited to people aged ≥ 18 years, and there are no data on the safety of using molnupiravir in children.⁴⁴ The mechanism of action of molnupiravir has raised concerns about potential mutagenesis in mammalian cells. See <u>Molnupiravir</u> and <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for additional information.

Corticosteroids

Corticosteroids are not indicated for the treatment of COVID-19 in nonhospitalized children. However, corticosteroids should be used per usual standards of care in children with asthma and croup triggered by SARS-CoV-2 infection. Children with COVID-19 who are receiving corticosteroids for an underlying condition should continue this therapy as directed by their health care providers.

Other Therapeutic Agents

For other therapies that have been studied or are under investigation for the treatment of COVID-19, see <u>Therapies</u>.

References

- 1. Schuster JE, Halasa NB, Nakamura M, et al. A description of COVID-19-directed therapy in children admitted to US intensive care units 2020. *J Pediatric Infect Dis Soc.* 2022;11(5):191-198. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35022779</u>.
- 2. Goldman DL, Aldrich ML, Hagmann SHF, et al. Compassionate use of remdesivir in children with severe COVID-19. *Pediatrics*. 2021;147(5):e2020047803. Available at: <u>https://www.ncbi.nlm.nih.gov/</u>

pubmed/33883243.

- 3. Chiotos K, Hayes M, Kimberlin DW, et al. Multicenter initial guidance on use of antivirals for children with coronavirus disease 2019/severe acute respiratory syndrome coronavirus 2. *J Pediatric Infect Dis Soc*. 2020;9(6):701-715. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32318706</u>.
- 4. Dulek DE, Fuhlbrigge RC, Tribble AC, et al. Multidisciplinary guidance regarding the use of immunomodulatory therapies for acute coronavirus disease 2019 in pediatric patients. *J Pediatric Infect Dis Soc.* 2020;9(6):716-737. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32808988.
- 5. Wolf J, Abzug MJ, Anosike BI, et al. Updated guidance on use and prioritization of monoclonal antibody therapy for treatment of COVID-19 in adolescents. *J Pediatric Infect Dis Soc*. 2022;11(5):177-185. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35107571.
- 6. Havers FP, Whitaker M, Self JL, et al. Hospitalization of adolescents aged 12–17 years with laboratoryconfirmed COVID-19—COVID-NET, 14 states, March 1, 2020–April 24, 2021. *MMWR Morb Mortal Wkly Rep.* 2021;70(23):851-857. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34111061.
- 7. Wolf J, Abzug MJ, Wattier RL, et al. Initial guidance on use of monoclonal antibody therapy for treatment of coronavirus disease 2019 in children and adolescents. *J Pediatric Infect Dis Soc*. 2021;10(5):629-634. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33388760</u>.
- 8. Graff K, Smith C, Silveira L, et al. Risk factors for severe COVID-19 in children. *Pediatr Infect Dis J*. 2021;40(4):e137-e145. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33538539</u>.
- Fernandes DM, Oliveira CR, Guerguis S, et al. Severe acute respiratory syndrome coronavirus 2 clinical syndromes and predictors of disease severity in hospitalized children and youth. *J Pediatr*. 2021;230:23-31. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33197493</u>.
- Götzinger F, Santiago-García B, Noguera-Julián A, et al. COVID-19 in children and adolescents in Europe: a multinational, multicentre cohort study. *Lancet Child Adolesc Health*. 2020;4(9):653-661. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32593339</u>.
- Kompaniyets L, Agathis NT, Nelson JM, et al. Underlying medical conditions associated with severe COVID-19 illness among children. *JAMA Netw Open*. 2021;4(6):e2111182. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34097050</u>.
- Feldstein LR, Tenforde MW, Friedman KG, et al. Characteristics and outcomes of US children and adolescents with multisystem inflammatory syndrome in children (MIS-C) compared with severe acute COVID-19. *JAMA*. 2021;325(11):1074-1087. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33625505</u>.
- Mukkada S, Bhakta N, Chantada GL, et al. Global characteristics and outcomes of SARS-CoV-2 infection in children and adolescents with cancer (GRCCC): a cohort study. *Lancet Oncol.* 2021;22(10):1416-1426. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34454651</u>.
- CDC COVID-19 Response Team. Coronavirus disease 2019 in children—United States, February 12–April 2, 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(14):422-426. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32271728.
- 15. Chao JY, Derespina KR, Herold BC, et al. Clinical characteristics and outcomes of hospitalized and critically ill children and adolescents with coronavirus disease 2019 at a tertiary care medical center in New York City. J Pediatr. 2020;223:14-19. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32407719</u>.
- DeBiasi RL, Song X, Delaney M, et al. Severe coronavirus disease-2019 in children and young adults in the Washington, DC, metropolitan region. *J Pediatr*. 2020;223:199-203. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32405091</u>.
- 17. Dong Y, Mo X, Hu Y, et al. Epidemiology of COVID-19 among children in China. *Pediatrics*. 2020;145(6):e20200702. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32179660</u>.
- Sun D, Li H, Lu XX, et al. Clinical features of severe pediatric patients with coronavirus disease 2019 in Wuhan: a single center's observational study. *World J Pediatr*. 2020;16(3):251-259. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32193831</u>.

- 19. Zachariah P, Johnson CL, Halabi KC, et al. Epidemiology, clinical features, and disease severity in patients with coronavirus disease 2019 (COVID-19) in a children's hospital in New York City, New York. *JAMA* Pediatr. 2020;174(10):e202430. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32492092.
- Delahoy MJ, Ujamaa D, Whitaker M, et al. Hospitalizations associated with COVID-19 among children and adolescents—COVID-NET, 14 states, March 1, 2020–August 14, 2021. MMWR Morb Mortal Wkly Rep. 2021;70(36):1255-1260. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34499627</u>.
- Wanga V, Gerdes ME, Shi DS, et al. Characteristics and clinical outcomes of children and adolescents aged <18 years hospitalized with COVID-19—six hospitals, United States, July–August 2021. *MMWR Morb Mortal Wkly Rep.* 2021;70(5152):1766-1772. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34968374</u>.
- 22. Centers for Disease Control and Prevention. Stay up to date with COVID-19 vaccines including boosters. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/vaccines/stay-up-to-date.html</u>. Accessed October 27, 2022.
- 23. Dorabawila V, Hoefer D, Bauer UE, et al. Risk of infection and hospitalization among vaccinated and unvaccinated children and adolescents in New York after the emergence of the Omicron variant. *JAMA*. 2022;327(22):2242-2244. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35559959.
- 24. Klein NP, Stockwell MS, Demarco M, et al. Effectiveness of COVID-19 Pfizer-BioNTech BNT162b2 mRNA vaccination in preventing COVID-19-associated emergency department and urgent care encounters and hospitalizations among nonimmunocompromised children and adolescents aged 5–17 years—VISION Network, 10 states, April 2021–January 2022. MMWR Morb Mortal Wkly Rep. 2022;71(9):352-358. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35239634.
- 25. Castelli JM, Rearte A, Olszevicki S, et al. Effectiveness of mRNA-1273, BNT162b2, and BBIBP-CorV vaccines against infection and mortality in children in Argentina, during predominance of Delta and Omicron COVID-19 variants: test negative, case-control study. *BMJ*. 2022;379:e073070. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36450402.
- 26. Chemaitelly H, AlMukdad S, Ayoub HH, et al. COVID-19 vaccine protection among children and adolescents in Qatar. N Engl J Med. 2022;387(20):1865-1876. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36322837.
- Antoon JW, Grijalva CG, Thurm C, et al. Factors associated with COVID-19 disease severity in US children and adolescents. *J Hosp Med.* 2021;16(10):603-610. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34613896.
- 28. Parcha V, Booker KS, Kalra R, et al. A retrospective cohort study of 12,306 pediatric COVID-19 patients in the United States. *Sci Rep.* 2021;11(1):10231. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33986390</u>.
- 29. Holmes L, Jr, Wu C, Hinson R, et al. Black-White risk differentials in pediatric COVID-19 hospitalization and intensive care unit admissions in the USA. *J Racial Ethn Health Disparities*. 2022;Published online ahead of print. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35604543</u>.
- 30. Woodruff RC, Campbell AP, Taylor CA, et al. Risk factors for severe COVID-19 in children. *Pediatrics*. 2022;149(1):e2021053418. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/34935038/</u>.
- Wiltz JL, Feehan AK, Molinari NM, et al. Racial and ethnic disparities in receipt of medications for treatment of COVID-19—United States, March 2020–August 2021. *MMWR Morb Mortal Wkly Rep.* 2022;71(3):96-102. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35051133</u>.
- 32. Gold JAW, Kelleher J, Magid J, et al. Dispensing of oral antiviral drugs for treatment of COVID-19 by ZIP code-level social vulnerability—United States, December 23, 2021–May 21, 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(25):825-829. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35737571.
- 33. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Paxlovid. 2022. Available at: <u>https://www.fda.gov/media/155050/download</u>.
- 34. Hammond J, Leister-Tebbe H, Gardner A, et al. Oral nirmatrelvir for high-risk, nonhospitalized adults with COVID-19. *N Engl J Med*. 2022;386(15):1397-1408. Available at: <u>https://www.ncbi.nlm.nih.gov/</u>

pubmed/35172054.

- 35. Greasley SE, Noell S, Plotnikova O, et al. Structural basis for the in vitro efficacy of nirmatrelvir against SARS-CoV-2 variants. *J Biol Chem.* 2022;298(6):101972. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/35461811</u>.
- 36. Rai DK, Yurgelonis I, McMonagle P, et al. Nirmatrelvir, an orally active M^{PRO} inhibitor, is a potent inhibitor of SARS-CoV-2 variants of concern. *bioRxiv*. 2022;Preprint. Available at: https://www.biorxiv.org/content/10.1101/2022.01.17.476644v1.
- 37. Vangeel L, Chiu W, De Jonghe S, et al. Remdesivir, molnupiravir and nirmatrelvir remain active against SARS-CoV-2 Omicron and other variants of concern. *Antiviral Res.* 2022;198:105252. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35085683</u>.
- 38. Remdesivir (Veklury) [package insert]. Food and Drug Administration. 2022. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2022/214787Orig1s015lbl.pdf.
- Fiaschi L, Dragoni F, Schiaroli E, et al. Efficacy of licensed monoclonal antibodies and antiviral agents against the SARS-CoV-2 Omicron sublineages BA.1 and BA.2. *Viruses*. 2022;14(7):1374. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35891355</u>.
- Takashita E, Kinoshita N, Yamayoshi S, et al. Efficacy of antiviral agents against the SARS-CoV-2 Omicron subvariant BA.2. *N Engl J Med*. 2022;386(15):1475-1477. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35263535</u>.
- 41. Takashita E, Yamayoshi S, Simon V, et al. Efficacy of antibodies and antiviral drugs against Omicron BA.2.12.1, BA.4, and BA.5 subvariants. *N Engl J Med.* 2022;387(5):468-470. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35857646</u>.
- 42. Gottlieb RL, Vaca CE, Paredes R, et al. Early remdesivir to prevent progression to severe COVID-19 in outpatients. *N Engl J Med.* 2022;386(4):305-315. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34937145</u>.
- 43. Méndez-Echevarría A, Pérez-Martínez A, Gonzalez Del Valle L, et al. Compassionate use of remdesivir in children with COVID-19. *Eur J Pediatr*. 2021;180(4):1317-1322. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33200304</u>.
- 44. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Lagevrio (molnupiravir) capsules. 2022. Available at: <u>https://www.fda.gov/media/155054/download</u>.

Therapeutic Management of Hospitalized Children With COVID-19

Last Updated: July 21, 2023

This section outlines the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations for the therapeutic management of children (i.e., pediatric patients aged <18 years) who are hospitalized for COVID-19. Throughout this section, the term "COVID-19" refers to the acute, primarily respiratory illness due to infection with SARS-CoV-2. Multisystem inflammatory syndrome in children (MIS-C) refers to the postinfectious inflammatory condition.

Treatment Considerations for Children With COVID-19

Currently, no pediatric clinical trial results evaluating the treatment of COVID-19 have been published. Data evaluating pharmacologic therapy in children with COVID-19 are limited largely to descriptive reports.^{1,2} Therefore, more high-quality randomized trials, observational studies, and pharmacokinetic studies are urgently needed. Whenever possible, clinical trials of therapeutics and multicenter observational cohorts should enroll children with COVID-19.

Published guidance documents on the treatment of COVID-19 in children have been mostly extrapolated from recommendations for adults with COVID-19, recommendations for children with other viral infections, and expert opinion.³⁻⁵ Applying adult data from COVID-19 trials to children is a unique challenge because most children experience a mild course of illness with COVID-19. Relative to adults, children with COVID-19 have substantially lower mortality and less need for hospitalization. Because of these differences in epidemiology and disease severity, the effect sizes for children are likely to be smaller than those observed in adults; therefore, to produce a beneficial outcome, the number needed to treat is higher. Collectively, these factors influence the risk versus benefit balance for pharmacologic therapies in children.

Other challenges are the uncertainty about which comorbid conditions place children at the highest risk of severe COVID-19 and the uncertainty about the absolute magnitude of the increased risk from those comorbid conditions. For children with COVID-19, the number and severity of their comorbid conditions influence decisions about pharmacologic treatment. For more information on risk factors for children with COVID-19, see <u>Special Considerations in Children</u>.

Recommendations

In the absence of sufficient clinical trial data on the treatment of children with COVID-19, the Panel's recommendations for the therapeutic management of hospitalized children are based largely on adult safety and efficacy data from clinical trials, the child's risk of disease progression, and expert opinion (see Table 3c). For the Panel's recommendations for adults, see <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>.

In general, adult data are most applicable to older children with severe COVID-19 and predominantly lower respiratory tract disease. Extrapolation of adult data to children with SARS-CoV-2 infection who present with clinical syndromes common to other respiratory viruses (e.g., bronchiolitis, croup, asthma) is challenging. No evidence indicates that these syndromes should be managed differently when caused by SARS-CoV-2 infection. Clinical judgment is needed when applying these recommendations to patients, particularly young children.

Disease Severity	Panel's Recommendations
Hospitalized for COVID-19	For children aged \geq 12 years admitted for COVID-19, use prophylactic anticoagulation unless contraindicated (BIII) . ^a
Does Not Require Supplemental Oxygen	For children admitted for COVID-19 who are at the highest risk of progression to severe COVID-19, ^b consider using remdesivir ^c for children aged 12–17 years (CIII) . There is insufficient evidence for using remdesivir in children aged 28 days to <12 years.
	For children admitted for reasons other than COVID-19 who have mild to moderate COVID-19 and are at the highest risk of progression, ^b refer to <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u> .
	Use 1 of the following options:
Requires Conventional Oxygen ^d	• Remdesivir ^c (BIII)
	 Dexamethasone plus remdesivir^c for children with increasing oxygen needs, particularly adolescents (BIII)
	Use 1 of the following options:
	Dexamethasone (BIII)
Requires Oxygen Through High-Flow	 Dexamethasone plus remdesivir^c (BIII)
Device or NIV ^e	For children who do not have rapid (e.g., within 24 hours) improvement in oxygenation after initiation of dexamethasone, baricitinib ^f or tocilizumab can be considered for children aged 12–17 years (BIII) and for children aged 2–11 years (CIII).
	Dexamethasone ⁹ (AIII)
Requires MV or ECMO ⁹	For children who do not have rapid (e.g., within 24 hours) improvement in oxygenation after initiation of dexamethasone, baricitinib ^f or tocilizumab may be considered for children aged 12–17 years (BIII) and for children aged 2–11 years (CIII).
Each recommendation in the Guidelines recei	ves a rating for the strength of the recommendation (A, B, or C) and a rating

Table 3c. Therapeutic Management of Hospitalized Children With COVID-19

^a Weighing the risk factors for thrombosis and bleeding, some Panel members would use prophylactic anticoagulation for children aged <12 years who are hospitalized for COVID-19.

- ^b For example, for children who are severely immunocompromised regardless of COVID-19 vaccination status and those who are unvaccinated and have additional risk factors for progression (see <u>Therapeutic Management of Nonhospitalized</u> <u>Children With COVID-19</u>).
- ^c The clinical benefit of remdesivir is greatest if it is initiated within 10 days of symptom onset. Remdesivir should be given for 5 days or until hospital discharge, whichever comes first.
- ^d Conventional oxygen refers to oxygen supplementation that is not high-flow oxygen, NIV, MV, or ECMO.

for the evidence that supports it (I, IIa, IIb, or III). See Guidelines Development for more information.

- ^e Patients who are receiving NIV or MV at baseline and require a substantial increase in baseline support should be treated per the recommendations for patients requiring new NIV or MV.
- ^f Tofacitinib is an alternative if baricitinib is not available (BIII).
- ⁹ For children who started receiving remdesivir before admission to the ICU, the remdesivir should be continued to complete the treatment course.

Key: ECMO = extracorporeal membrane oxygenation; ICU = intensive care unit; MV = mechanical ventilation; NIV = noninvasive ventilation; the Panel = the COVID-19 Treatment Guidelines Panel

COVID-19 Treatment Guidelines

Rationale for the Panel's Recommendations for Drug Therapies

Remdesivir

Remdesivir is approved by the Food and Drug Administration (FDA) for hospitalized and nonhospitalized pediatric patients aged ≥ 28 days and weighing ≥ 3 kg.⁶ Remdesivir is expected to be active against the Omicron variant of concern, although in vitro and in vivo data are currently limited (see <u>Remdesivir</u>).⁷ For most hospitalized patients, remdesivir should be administered for 5 days or until the patient is ready for discharge, whichever comes first. Treatment may be extended to 10 days for severely ill patients who have not clinically improved or for patients who are severely immunocompromised.

In a trial conducted predominantly among hospitalized patients with COVID-19 who did not receive supplemental oxygen at enrollment, a 5-day course of remdesivir was associated with greater clinical improvement when compared with the standard of care.⁸ Remdesivir was also studied in ACTT-1, a double-blind, placebo-controlled, randomized trial for hospitalized adults with COVID-19 who received remdesivir for 10 days (or until hospital discharge) or placebo.⁹ The study reported that the remdesivir arm had a shorter time to clinical recovery than the placebo arm (10 days vs. 15 days; P < 0.001). A subgroup analysis demonstrated that patients who received conventional oxygen therapy had the greatest benefit. No benefit was detected for patients who did not receive supplemental oxygen or for those who received noninvasive ventilation (NIV) or mechanical ventilation. No statistically significant differences in mortality or in the need for new mechanical ventilation were detected, and the benefit of remdesivir in this study was limited to patients with symptoms for <10 days.

Three open-label trials in adults compared remdesivir to a local standard of care.^{10,11} The World Health Organization's Solidarity trial enrolled hospitalized adult patients with COVID-19 in 35 countries. In the overall cohort, no difference in hospital mortality was demonstrated (14.5% in the remdesivir arm vs. 15.6% in the usual care arm; rate ratio 0.91; 95% CI, 0.82–1.02; P = 0.12). However, in the subset of patients receiving supplemental oxygen but not NIV or mechanical ventilation, remdesivir significantly reduced the risk of in-hospital mortality by 13% (14.6% vs. 16.3%; rate ratio 0.87; 95% CI, 0.76–0.99; P = 0.03).¹⁰

The CATCO study demonstrated similar findings. Treatment with remdesivir, when compared with standard care, reduced the need for mechanical ventilation in hospitalized adults with COVID-19 (8% vs. 15%; relative risk 0.53; 95% CI, 0.38–0.75). In this study, 87% of patients in both the remdesivir arm and standard of care arm received dexamethasone.¹² In contrast to these 2 studies, the DisCoVeRy trial demonstrated no difference for any clinical outcome when the use of remdesivir plus usual care was compared to usual care alone.¹¹

The efficacy of remdesivir has not been evaluated in clinical trials of hospitalized children with COVID-19. A Phase 2/3, single-arm, open-label study evaluated the safety, tolerability, and pharmacokinetics of remdesivir in 53 hospitalized children with COVID-19.¹³ Children weighing 3 kg to <40 kg received remdesivir 5 mg/kg on Day 1, followed by remdesivir 2.5 mg/kg daily. Adverse events included acute kidney injury (11%) and an increase in alanine transaminase levels (8%). However, this study did not have a placebo group, limiting the ability to draw conclusions regarding the significance of these adverse events. Published observational data are limited to descriptive case series.^{1,2}

Findings from the adult trials and the pediatric pharmacokinetic study led the Panel to recommend **remdesivir** for hospitalized children who have a new or increasing need for conventional oxygen (**BIII**) and to recommend **dexamethasone plus remdesivir** for children who require oxygen through a high-flow device or NIV (**BIII**). It is not known if remdesivir offers an additional clinical benefit to standard care in younger children with SARS-CoV-2 infection who are receiving respiratory support for bronchiolitis, asthma, or croup.

COVID-19 Treatment Guidelines

For children hospitalized for COVID-19 who do not require supplemental oxygen, the Panel recommends **remdesivir** for children aged 12 to 17 years who are at the highest risk for progression to severe disease (CIII). This recommendation was extrapolated from the findings of the PINETREE study, which demonstrated a reduction in hospitalization among high-risk, unvaccinated adults treated in the outpatient setting.¹⁴ However, there is insufficient evidence for or against the use of remdesivir in children aged 28 days to <12 years and weighing ≥ 3 kg who do not require supplemental oxygen. Given the reported clinical experience with the use of remdesivir among younger patients,¹³ the use of remdesivir in high-risk, younger children who do not require supplemental oxygen may be considered on a case-by-case basis.

Dexamethasone

Dexamethasone was evaluated in the RECOVERY trial, which was an open-label, randomized trial conducted in the United Kingdom.¹⁵ The trial compared the use of up to 10 days of dexamethasone 6 mg, administered by intravenous injection or orally, with usual care among hospitalized adults with COVID-19. The primary outcome was all-cause mortality at 28 days, which occurred in 22.9% of patients randomized to receive dexamethasone versus 25.7% of patients randomized to receive usual care (age-adjusted rate ratio 0.83; 95% CI, 0.75–0.93; P < 0.001). Patients who required mechanical ventilation or extracorporeal membrane oxygenation (ECMO) had the greatest effect size (29.3% vs. 41.4%; rate ratio 0.64; 95% CI, 0.51–0.81). No difference in outcomes was observed for patients who did not require supplemental oxygen (17.8% vs. 14.0%; rate ratio 1.19; 95% CI, 0.92–1.55). For the 28-day mortality outcome, a difference between arms was observed for patients who required supplemental oxygen (23.3% vs. 26.2%; rate ratio 0.82; 95% CI, 0.72–0.94). However, it should be noted that these patients were a heterogeneous group, including those who received either conventional oxygen or NIV. See Systemic Corticosteroids for detailed information.

The safety and efficacy of using dexamethasone or other corticosteroids for the treatment of COVID-19 have not been evaluated in pediatric patients. Given that the mortality for adults in the placebo arm in the RECOVERY trial was substantially greater than the mortality generally reported for children with COVID-19, caution is warranted when extrapolating from recommendations for adults and applying them to patients aged <18 years.

However, because of the effect size observed in the RECOVERY trial, the Panel recommends the use of dexamethasone for children who require mechanical ventilation or ECMO (AIII). The Panel also recommends the use of **dexamethasone**, with or without concurrent **remdesivir**, for children who require oxygen through a high-flow device or NIV (BIII). The Panel does not recommend routine use of corticosteroids for children who require only conventional oxygen, but corticosteroids can be considered in combination with remdesivir for patients with increasing oxygen needs, particularly adolescents (BIII).

There is evidence demonstrating that the use of corticosteroids does not benefit infants with viral bronchiolitis not related to COVID-19, and current American Academy of Pediatrics guidelines recommend against the use of corticosteroids in this population.¹⁶ There are no COVID-19-specific data to support the use of corticosteroids in children with bronchiolitis due to SARS-CoV-2 infection. Corticosteroids should be used per the usual standards of care in children with asthma and croup triggered by SARS-CoV-2.

The use of dexamethasone for the treatment of severe COVID-19 in children who are profoundly immunocompromised has not been evaluated, and there is a potential risk of harm. Therefore, the use of corticosteroids should be considered on a case-by-case basis in consultation with relevant specialists, and the benefits and risks of the therapy should be weighed. If dexamethasone is not available, alternative glucocorticoids such as prednisone, methylprednisolone, or hydrocortisone can be considered. The dexamethasone dose for pediatric patients is 0.15 mg/kg (with a maximum dose of 6 COVID-19 Treatment Guidelines

mg) once daily for ≤ 10 days.

Baricitinib

The Janus kinase inhibitor baricitinib was approved by the FDA for the treatment of COVID-19 in hospitalized adults. An FDA Emergency Use Authorization (EUA) for baricitinib remains active for the treatment of COVID-19 in hospitalized children aged 2 to 17 years who require supplemental oxygen, NIV, mechanical ventilation, or ECMO.¹⁷

In the COV-BARRIER trial, adults with COVID-19 pneumonia were randomized to receive baricitinib or standard care. Patients treated with baricitinib showed a reduction in mortality when compared with those who received standard care; the reduction was greatest in patients who received high-flow oxygen or NIV. Similarly, the ACTT-2 trial in adults showed that patients who received baricitinib plus remdesivir had improved time to recovery when compared with patients who received remdesivir alone. This effect was most pronounced in patients who received high-flow oxygen or NIV.¹⁸ In the ACTT-4 trial, 1,010 patients were randomized 1:1 to receive baricitinib plus remdesivir or dexamethasone plus remdesivir. The study reported no difference between the arms for the outcome of mechanical ventilation-free survival.¹⁹

In the RECOVERY trial, 8,156 patients, including 33 children aged 2 to 17 years, were randomized to receive baricitinib or usual care (95% received corticosteroids).²⁰ Treatment with baricitinib was associated with a 13% proportional reduction in mortality, with the greatest effect size occurring in patients who received NIV. The RECOVERY investigators included these patients in a meta-analysis and found that treatment with baricitinib was associated with a 20% proportional reduction in mortality (rate ratio 0.80; 95% CI, 0.72–0.89; P < 0.0001). See Janus Kinase Inhibitors and Therapeutic Management of Hospitalized Adults With COVID-19 for additional information. These data in adults indicate that baricitinib is likely to be most beneficial for patients receiving noninvasive forms of respiratory support.

Several open-label trials and cohort studies have evaluated baricitinib in children with autoinflammatory and rheumatic diseases, including many children aged <5 years, and found the treatment was well tolerated; however, the pharmacokinetics of baricitinib in younger children are not well studied.²¹⁻²⁴ Information on the safety and effectiveness of the use of baricitinib in children with COVID-19 is limited to case reports.

In contrast to the strong recommendation for its use for adults, baricitinib is not considered the standard of care for all children who require high-flow oxygen or NIV because of the low mortality in children with COVID-19 (especially in young children) and the limited data on the use of baricitinib in these children.

Extrapolating from clinical trials among adults with COVID-19, the Panel recommends that:

- For children who require oxygen through a high-flow device or NIV *and* do not have rapid (e.g., within 24 hours) improvement in oxygenation after initiation of dexamethasone, **baricitinib** can be considered for children aged 12 to 17 years (**BIII**) and for children aged 2 to 11 years (**CIII**).
- For children who require mechanical ventilation or ECMO *and* do not have rapid (e.g., within 24 hours) improvement in oxygenation after initiation of dexamethasone, **baricitinib** may be considered for children aged 12 to 17 years (**BIII**) and for children aged 2 to 11 years (**CIII**).

Clinicians should consult with specialists experienced in treating children with immunosuppression (e.g., with pediatric infectious disease, pediatric rheumatology) when considering administering baricitinib to hospitalized children with COVID-19. Data from adults indicate that baricitinib should be initiated

promptly; ideally, it should be initiated at the onset of clinical deterioration or respiratory failure.

Tofacitinib

There are no data on the efficacy of tofacitinib in pediatric patients with COVID-19; the Panel's recommendation is extrapolated from data in adults. The STOP-COVID trial compared tofacitinib to the standard of care in adults hospitalized for COVID-19 pneumonia.²⁵ The standard of care included glucocorticoids for most patients. The study demonstrated a reduction in mortality and respiratory failure at Day 28 for the tofacitinib arm when compared with the placebo arm. Tofacitinib has been studied less extensively than baricitinib for the treatment of COVID-19. Thus, tofacitinib, as an alternative to baricitinib, is recommended to be used in combination with dexamethasone in adults with COVID-19 who require high-flow oxygen or NIV. See Janus Kinase Inhibitors and Therapeutic Management of Hospitalized Adults With COVID-19 for additional information.

No trials have evaluated the safety of using tofacitinib in children with COVID-19. Overall, there has been more clinical experience with the use of tofacitinib than baricitinib in children, particularly when used in children with juvenile idiopathic arthritis (JIA) as young as 2 years of age. A Phase 1 study was conducted to define the pharmacokinetics and safety of using tofacitinib in children,²⁶ and a Phase 3, double-blind, randomized, placebo-controlled trial investigated the efficacy of using tofacitinib in children with JIA.²⁷ Tofacitinib is available as a liquid formulation for children.

Given the established safety of tofacitinib in the pediatric population, **tofacitinib** can be considered an alternative for children hospitalized for COVID-19 if baricitinib is not available (**BIII**). The dose of tofacitinib that should be used to treat hospitalized children with COVID-19 has not been established. As with baricitinib, the dose of tofacitinib for hospitalized children with COVID-19 likely needs to be higher than the dose typically used to treat pediatric rheumatologic diseases. Therefore, clinicians should consult with specialists experienced in treating children with immunosuppression (e.g., with pediatric infectious disease, pediatric rheumatology) when considering administering tofacitinib to hospitalized children with COVID-19.

Tocilizumab

Tocilizumab is an interleukin (IL)-6 inhibitor that has received an FDA EUA for the treatment of hospitalized adults and children with COVID-19 who are aged ≥ 2 years, receiving systemic corticosteroids, and require supplemental oxygen, NIV, mechanical ventilation, or ECMO.²⁸ Two large randomized controlled trials (REMAP-CAP and RECOVERY) conducted among hospitalized adults with COVID-19 have demonstrated reductions in mortality with the use of tocilizumab. See Interleukin-6 Inhibitors and Therapeutic Management of Hospitalized Adults With COVID-19 for additional information.

The RECOVERY trial was an open-label study that included hospitalized adults who had an oxygen saturation of <92% on room air or were receiving supplemental oxygen therapy; patients also had C-reactive protein levels \geq 75 mg/L.²⁹ Patients were randomized to receive tocilizumab plus usual care or usual care alone. Mortality at 28 days was significantly lower in the tocilizumab arm compared to the usual care arm. The REMAP-CAP trial included adults with suspected or confirmed COVID-19 who were admitted to an intensive care unit and received either respiratory (i.e., NIV or mechanical ventilation) or cardiovascular organ (i.e., vasopressor/inotrope) support.³⁰ Patients were randomized within 24 hours of organ failure to receive either tocilizumab or sarilumab (the majority received tocilizumab) or to receive standard care. The median number of organ support-free days was higher for those who received tocilizumab than for those who received standard care, and in-hospital mortality was lower in the combined tocilizumab or sarilumab arm than in the standard care arm. In both

studies, the majority of patients received dexamethasone (82% in the RECOVERY trial and 93% in the REMAP-CAP trial).

Studies have evaluated the use of tocilizumab for the treatment of non-COVID-19 conditions in children, including JIA³¹⁻³⁵ and chimeric antigen receptor T cell-related cytokine release syndrome.³⁶ The FDA approved tocilizumab for use in children aged ≥ 2 years for these indications.³¹⁻³⁵ The use of tocilizumab for children with severe cases of COVID-19 has been described only in case series.³⁷⁻³⁹

Extrapolating from clinical trials among adults with COVID-19, the Panel recommends that:

- For children who require oxygen through a high-flow device or NIV *and* who do not have rapid (e.g., within 24 hours) improvement in oxygenation after initiation of dexamethasone, **tocilizumab** can be considered for children aged 12 to 17 years (**BIII**) and for children aged 2 to 11 years (**CIII**).
- For children who require mechanical ventilation or ECMO *and* who do not have rapid (e.g., within 24 hours) improvement in oxygenation after initiation of dexamethasone, if tocilizumab has not been started, addition of **tocilizumab** may be considered for children aged 12 to 17 years (**BIII**) and for children aged 2 to 11 years (**CIII**).

Data from REMAP-CAP and RECOVERY are most likely to be applicable to high-risk adolescent patients. Clinicians should consult with specialists experienced in treating children with immunosuppression (e.g., with pediatric infectious disease, pediatric rheumatology) when considering the use of tocilizumab in younger children with COVID-19.

Sarilumab

Sarilumab, a monoclonal antibody that blocks IL-6 receptors, is not authorized by the FDA for the treatment of COVID-19. Data evaluating the efficacy of sarilumab for the treatment of COVID-19 hyperinflammation are limited, and there is a lack of pediatric dosing information. Therefore, the Panel **recommends against** the use of **sarilumab** in hospitalized children with COVID-19, except in a clinical trial (**AIII**).

Anticoagulation in Children With COVID-19

Recommendations

- The Panel recommends prophylactic anticoagulation for children aged ≥12 years who are hospitalized for COVID-19, unless there are contraindications (**BIII**).
- Weighing the risk factors for thrombosis and bleeding, some Panel members would use prophylactic anticoagulation for children aged <12 years who are hospitalized for COVID-19. Institutional standards for anticoagulation should be followed.
- There is insufficient evidence for the Panel to recommend either for or against the use of therapeutic anticoagulation in children of any age with COVID-19.

Limited data characterize the risk of thromboembolic disease in children with COVID-19. Among children who do not have COVID-19, most thromboembolic events occur in neonates and adolescents.^{40,41} In a multicenter, retrospective cohort study that included 814 pediatric patients with COVID-19 or MIS-C,⁴² thromboembolic events were detected in 2.1% of patients with COVID-19 and in 6.5% of patients with MIS-C.

Limited data inform the clinical use of anticoagulation among children with COVID-19. Only the COVAC-TP trial has evaluated the dose, safety, and efficacy of anticoagulant prophylaxis in children

with COVID-19 or MIS-C.⁴³ In this multicenter, Phase 2 clinical trial of children hospitalized with COVID-19-related illness (including MIS-C) in the United States, a starting dose of enoxaparin 0.5 mg/kg achieved targeted anticoagulant activity (as measured by antifactor Xa level) in the majority of patients with few dose changes, and no patients experienced clinically relevant bleeding as defined by the International Society on Thrombosis and Haemostasis.⁴⁴ In this trial, thromboembolic events occurred in 2 patients (5.3%; 90% CI, 1.0%–15.7%); both events were related to central venous catheters.⁴³ These results raise the question of whether prophylactic doses of anticoagulants sufficiently reduce thromboembolism risk in children hospitalized with COVID-19 or MIS-C.

To date, no clinical trial has evaluated the safety and efficacy of therapeutic anticoagulation in hospitalized children with COVID-19. Therefore, the Panel has determined that there is insufficient evidence to recommend either for or against the use of therapeutic anticoagulation in children of any age with COVID-19.

References

- 1. Schuster JE, Halasa NB, Nakamura M, et al. A description of COVID-19-directed therapy in children admitted to US intensive care units 2020. J Pediatric Infect Dis Soc. 2022;11(5):191-198. Available at: https://www. ncbi.nlm.nih.gov/pubmed/35022779.
- 2. Goldman DL, Aldrich ML, Hagmann SHF, et al. Compassionate use of remdesivir in children with severe COVID-19. Pediatrics. 2021;147(5):e2020047803. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33883243.
- 3. Chiotos K, Hayes M, Kimberlin DW, et al. Multicenter initial guidance on use of antivirals for children with coronavirus disease 2019/severe acute respiratory syndrome coronavirus 2. J Pediatric Infect Dis Soc. 2020;9(6):701-715. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32318706.
- 4. Dulek DE, Fuhlbrigge RC, Tribble AC, et al. Multidisciplinary guidance regarding the use of immunomodulatory therapies for acute coronavirus disease 2019 in pediatric patients. J Pediatric Infect Dis Soc. 2020;9(6):716-737. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32808988.
- 5. Wolf J, Abzug MJ, Anosike BI, et al. Updated guidance on use and prioritization of monoclonal antibody therapy for treatment of COVID-19 in adolescents. J Pediatric Infect Dis Soc. 2022;11(5):177-185. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35107571.
- 6. Remdesivir (Veklury) [package insert]. Food and Drug Administration. 2022. Available at: https://www. accessdata.fda.gov/drugsatfda_docs/label/2022/214787Orig1s015lbl.pdf.
- 7. Vangeel L, Chiu W, De Jonghe S, et al. Remdesivir, molnupiravir and nirmatrelvir remain active against SARS-CoV-2 Omicron and other variants of concern. Antiviral Res. 2022;198:105252. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35085683.
- 8. Spinner CD, Gottlieb RL, Criner GJ, et al. Effect of remdesivir vs standard care on clinical status at 11 days in patients with moderate COVID-19: a randomized clinical trial. JAMA. 2020;324(11):1048-1057. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32821939.
- 9. Beigel JH, Tomashek KM, Dodd LE, et al. Remdesivir for the treatment of COVID-19—final report. N Engl J Med. 2020;383(19):1813-1826. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32445440.
- 10. WHO Solidarity Trial Consortium. Remdesivir and three other drugs for hospitalised patients with COVID-19: final results of the WHO Solidarity randomised trial and updated meta-analyses. Lancet. 2022;399(10339):1941-1953. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35512728.
- 11. Ader F, Bouscambert-Duchamp M, Hites M, et al. Remdesivir plus standard of care versus standard of care alone for the treatment of patients admitted to hospital with COVID-19 (DisCoVeRy): a Phase 3, randomised, controlled, open-label trial. Lancet Infect Dis. 2022;22(2):209-221. Available at: https://www.ncbi.nlm.nih. gov/pubmed/34534511.
- 12. Ali K, Azher T, Baqi M, et al. Remdesivir for the treatment of patients in hospital with COVID-19 in Canada: COVID-19 Treatment Guidelines 105

a randomized controlled trial. *CMAJ*. 2022;194(7):E242-E251. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35045989</u>.

- 13. Ahmed A, Rojo P, Agwu A, et al. Remdesivir treatment for COVID-19 in hospitalized children: CARAVAN interim results. 2022. Available at: <u>https://www.croiconference.org/abstract/remdesivir-treatment-for-covid-19-in-hospitalized-children-caravan-interim-results</u>. Accessed June 28, 2023.
- Gottlieb RL, Vaca CE, Paredes R, et al. Early remdesivir to prevent progression to severe COVID-19 in outpatients. *N Engl J Med.* 2022;386(4):305-315. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34937145.
- 15. RECOVERY Collaborative Group. Dexamethasone in hospitalized patients with COVID-19. *N Engl J Med*. 2021;384(8):693-704. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32678530</u>.
- Ralston SL, Lieberthal AS, Meissner HC, et al. Clinical practice guideline: the diagnosis, management, and prevention of bronchiolitis. *Pediatrics*. 2014;134(5):e1474-e1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/25349312</u>.
- 17. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization (EUA) of baricitinib. 2022. Available at: <u>https://www.fda.gov/media/143823/download</u>.
- Kalil AC, Patterson TF, Mehta AK, et al. Baricitinib plus remdesivir for hospitalized adults with COVID-19. N Engl J Med. 2021;384(9):795-807. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33306283</u>.
- Wolfe CR, Tomashek KM, Patterson TF, et al. Baricitinib versus dexamethasone for adults hospitalised with COVID-19 (ACTT-4): a randomised, double-blind, double placebo-controlled trial. *Lancet Respir Med*. 2022;10(9):888-899. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35617986</u>.
- 20. RECOVERY Collaborative Group. Baricitinib in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial and updated meta-analysis. *Lancet*. 2022;400(10349):359-368. Available at: https://pubmed.ncbi.nlm.nih.gov/35908569.
- Sanchez GAM, Reinhardt A, Ramsey S, et al. JAK1/2 inhibition with baricitinib in the treatment of autoinflammatory interferonopathies. *J Clin Invest*. 2018;128(7):3041-3052. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29649002</u>.
- 22. Kim H, Dill S, O'Brien M, et al. Janus kinase (JAK) inhibition with baricitinib in refractory juvenile dermatomyositis. *Ann Rheum Dis*. 2021;80(3):406-408. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32843325.
- 23. Vanderver A, Adang L, Gavazzi F, et al. Janus kinase inhibition in the Aicardi-Goutieres syndrome. *N Engl J Med.* 2020;383(10):986-989. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32877590</u>.
- 24. Kim H, Brooks KM, Tang CC, et al. Pharmacokinetics, pharmacodynamics, and proposed dosing of the oral JAK1 and JAK2 inhibitor baricitinib in pediatric and young adult CANDLE and SAVI patients. *Clin Pharmacol Ther.* 2018;104(2):364-373. Available at: https://www.ncbi.nlm.nih.gov/pubmed/29134648.
- 25. Guimarães PO, Quirk D, Furtado RH, et al. Tofacitinib in patients hospitalized with COVID-19 pneumonia. *N Engl J Med.* 2021;385(5):406-415. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133856</u>.
- 26. Ruperto N, Brunner HI, Zuber Z, et al. Pharmacokinetic and safety profile of tofacitinib in children with polyarticular course juvenile idiopathic arthritis: results of a Phase 1, open-label, multicenter study. *Pediatr Rheumatol Online J*. 2017;15(1):86. Available at: https://www.ncbi.nlm.nih.gov/pubmed/29282090.
- 27. Ruperto N, Brunner HI, Synoverska O, et al. Tofacitinib in juvenile idiopathic arthritis: a double-blind, placebo-controlled, withdrawal Phase 3 randomised trial. *Lancet*. 2021;398(10315):1984-1996. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34767764.
- 28. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Actemra (tocilizumab). 2021. Available at: <u>https://www.fda.gov/media/150321/download</u>.
- 29. RECOVERY Collaborative Group. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2021;397(10285):1637-1645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33933206</u>.

- 30. REMAP-CAP Investigators. Interleukin-6 receptor antagonists in critically ill patients with COVID-19. *N Engl J Med.* 2021;384(16):1491-1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631065</u>.
- 31. Brunner HI, Ruperto N, Zuber Z, et al. Efficacy and safety of tocilizumab in patients with polyarticular-course juvenile idiopathic arthritis: results from a Phase 3, randomised, double-blind withdrawal trial. Ann Rheum Dis. 2015;74(6):1110-1117. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/24834925</u>.
- Brunner HI, Ruperto N, Zuber Z, et al. Efficacy and safety of tocilizumab for polyarticular-course juvenile idiopathic arthritis in the open-label two-year extension of a Phase III trial. *Arthritis Rheumatol*. 2021;73(3):530-541. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32951358</u>.
- 33. De Benedetti F, Brunner HI, Ruperto N, et al. Randomized trial of tocilizumab in systemic juvenile idiopathic arthritis. N Engl J Med. 2012;367(25):2385-2395. Available at: https://www.ncbi.nlm.nih.gov/pubmed/23252525.
- 34. DeWitt EM, Kimura Y, Beukelman T, et al. Consensus treatment plans for new-onset systemic juvenile idiopathic arthritis. *Arthritis Care Res (Hoboken)*. 2012;64(7):1001-1010. Available at: https://www.ncbi.nlm.nih.gov/pubmed/22290637.
- 35. Horneff G, Schulz AC, Klotsche J, et al. Experience with etanercept, tocilizumab and interleukin-1 inhibitors in systemic onset juvenile idiopathic arthritis patients from the BIKER registry. *Arthritis Res Ther*. 2017;19(1):256. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29166924</u>.
- 36. Kotch C, Barrett D, Teachey DT. Tocilizumab for the treatment of chimeric antigen receptor T cell-induced cytokine release syndrome. *Expert Rev Clin Immunol*. 2019;15(8):813-822. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31219357</u>.
- 37. Derespina KR, Kaushik S, Plichta A, et al. Clinical manifestations and outcomes of critically ill children and adolescents with coronavirus disease 2019 in New York City. *J Pediatr*. 2020;226:55-63.e2. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32681989</u>.
- 38. Shekerdemian LS, Mahmood NR, Wolfe KK, et al. Characteristics and outcomes of children with coronavirus disease 2019 (COVID-19) infection admitted to US and Canadian pediatric intensive care units. JAMA Pediatr. 2020;174(9):868-873. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32392288</u>.
- 39. Götzinger F, Santiago-García B, Noguera-Julián A, et al. COVID-19 in children and adolescents in Europe: a multinational, multicentre cohort study. *Lancet Child Adolesc Health*. 2020;4(9):653-661. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32593339.
- 40. Stein PD, Kayali F, Olson RE. Incidence of venous thromboembolism in infants and children: data from the National Hospital Discharge Survey. *J Pediatr*. 2004;145(4):563-556. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/15480387</u>.
- 41. O'Brien SH, Stanek JR, Witmer CM, Raffini L. The continued rise of venous thromboembolism across US children's hospitals. *Pediatrics*. 2022;149(3):e2021054649. Available at: https://pubmed.ncbi.nlm.nih.gov/35156127.
- 42. Whitworth H, Sartain SE, Kumar R, et al. Rate of thrombosis in children and adolescents hospitalized with COVID-19 or MIS-C. *Blood*. 2021;138(2):190-198. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33895804.
- 43. Sochet AA, Morrison JM, Jaffray J, et al. Enoxaparin thromboprophylaxis in children hospitalized for COVID-19: a Phase 2 trial. *Pediatrics*. 2022;150(1):e2022056726. Available at: https://pubmed.ncbi.nlm.nih.gov/35484817.
- 44. Mitchell LG, Goldenberg NA, Male C, et al. Definition of clinical efficacy and safety outcomes for clinical trials in deep venous thrombosis and pulmonary embolism in children. *J Thromb Haemost*. 2011;9(9):1856-1858. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21884565</u>.

COVID-19 Treatment Guidelines

Therapeutic Management of Hospitalized Children With MIS-C, Plus a Discussion on MIS-A

Last Updated: July 21, 2023

This section outlines the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations for the therapeutic management of pediatric patients with multisystem inflammatory syndrome in children (MIS-C). The case definition for MIS-C from the Council of State and Territorial Epidemiologists (CSTE) and the Centers for Disease Control and Prevention (CDC) includes individuals aged <21 years.¹ The recommendations in this section encompass this age group. No randomized controlled trials have compared different treatment approaches for MIS-C. However, data from descriptive and observational comparative effectiveness studies are available to guide treatment for MIS-C. For information on the clinical manifestations of MIS-C, see <u>Special Considerations in Children</u>.

Multisystem Inflammatory Syndrome in Adults

It should be noted that adults can present with a syndrome similar to MIS-C, termed multisystem inflammatory syndrome in adults (MIS-A).² The published literature on MIS-A is restricted to small case series and a single observational epidemiological study that provide little data to guide treatment decisions for patients with MIS-A.³⁻⁵ Although the therapeutic management of MIS-A has not been studied, it is reasonable to extrapolate from data on treating patients with MIS-C to aid in the management of individuals with MIS-A.

Table 3d. Therapeutic Management of Hospitalized Pediatric Patients With MIS-C

	Panel's Recommendations		
	Initial treatment for MIS-C includes both immunomodulatory and antithrombotic therapy.		
	Initial Immunomodulatory Therapy		
	 IVIG 2 g/kg IBW (up to a maximum total dose of 100 g) IV plus low to moderate dose methylprednisolone (1–2 mg/kg/day) IV^a or another glucocorticoid at an equivalent dose^a (AIIb). 		
	• Glucocorticoid monotherapy, only if IVIG is unavailable or contraindicated (Blla).		
	• IVIG monotherapy, only if glucocorticoids are contraindicated (BIIb).		
	Intensification Immunomodulatory Therapy		
	 Intensification therapy is recommended for children with refractory MIS-C who do not improve within 24 hours of receiving initial immunomodulatory therapy (AIII). One of the following can be used (listed in alphabetical order): 		
	 High-dose anakinra 5–10 mg/kg IV or SUBQ once daily (BIIb) 		
MIS-C	 Higher-dose glucocorticoid (e.g., methylprednisolone 10–30 mg/kg/day IV or equivalent glucocorticoid) (BIIb)^b 		
	 Infliximab^c 5–10 mg/kg IV for 1 dose (BIIb) 		
	Antithrombotic Therapy		
	 Low-dose aspirin (3–5 mg/kg/day, up to maximum dose of 81 mg/day) PO for all patients without risk factors for bleeding (AIII), <u>AND</u> 		
	 Anticoagulation for patients who fall under 1 of the following clinical scenarios: 		
	 Therapeutic anticoagulation for patients with large CAAs according to the American Heart Association guidelines for Kawasaki disease (AIII). 		
	 Therapeutic anticoagulation for patients with moderate to severe LV dysfunction who have no risk factors for bleeding (AIII). 		
	 For patients with MIS-C who do not have large CAAs or moderate to severe LV dysfunction, consider prophylactic or therapeutic anticoagulation on an individual basis, taking into consideration risk factors for thrombosis and bleeding. See Table 3e for additional information. 		
	ne Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating orts it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.		

^a Duration of therapy may vary. See Table 3e and text below.

^b In certain patients with severe illness, intensification therapy may include dual therapy with higher-dose glucocorticoids and infliximab or anakinra. Anakinra and infliximab **should not be given** in combination.

^c Infliximab **should not be used** in patients with macrophage activation syndrome.

Key: CAA = coronary artery aneurysm; IBW = ideal body weight; IV = intravenous; IVIG = intravenous immunoglobulin; LV = left ventricular; MIS-C = multisystem inflammatory syndrome in children; PO = oral; SUBQ = subcutaneously

	Dosing Regimens		
	For infants, children, and adolescents unless otherwise specified. The doses listed are for FDA-approved indications for other diseases or from reported experiences or clinical trials.	Adverse Events	Monitoring Parameters
Intravenous Immunoglobulin	IVIG 2 g/kg IBW (up to a maximum total dose of 100 g) IV In the event of cardiac dysfunction or fluid overload, consider administering IVIG in divided doses (1 g/kg IBW per dose IV every 24 hours for 2 doses).	 Hypersensitivity Fever Chills Flushing Hemolytic anemia 	 Renal function Urine output CBC with differential Infusion or injection-related AEs Anaphylaxis Signs and symptoms of hemolysis
Methyl- prednisolone	Methylprednisolone 1–2 mg/kg IV every 12 hours If the patient does not respond to 1–2 mg/kg IV every 12 hours, increase the dose to 10–30 mg/kg/day (up to maximum of 1,000 mg/day) IV for 1–3 days.	 Adrenal suppression Hyperglycemia Sodium retention Fluid retention Leukocytosis Immune suppression 	Blood pressureCBC with differentialBMP
Anakinra	Anakinra 5–10 mg/kg/day IV (preferred) or SUBQ in 1 to 4 divided doses	 Headache Fever Hypersensitivity Immune suppression Transaminitis 	 CBC with differential LFTs SCr
Infliximab	Infliximab 5–10 mg/kg IV for 1 dose	 Infusion-related reaction Headache Immune suppression 	 Monitor vital signs every 2–10 minutes during infusion. CBC with differential
Aspirin	Aspirin 3–5 mg/kg (up to maximum of 81 mg) PO once daily	Gastrointestinal ulcersHypersensitivityRenal dysfunction	Signs or symptoms of bleedingRenal function
Enoxaparin	 Enoxaparin Prophylaxis Aged >2 Months to <18 Years 0.5 mg/kg (up to maximum of 30 mg) SUBQ every 12 hours Enoxaparin Treatment Aged >2 Months to <18 Years 1 mg/kg SUBQ every 12 hours Monitor antifactor Xa activity (treatment goal: 0.5 to 1). 	 Increased risk of bleeding Thrombocytopenia 	CBC with differentialRenal function

Table 3e. Dosing Regimens for the Drugs Recommended for the Treatment of MIS-C

Key: AE = adverse effect; BMP = blood mineral panel; CBC = complete blood count; FDA = Food and Drug Administration; IBW = ideal body weight; IV = intravenous; IVIG = intravenous immunoglobulin; LFT = liver function test; MIS-C = multisystem inflammatory syndrome in children; PO = oral; SCr = serum creatinine; SUBQ = subcutaneous

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Treatment Considerations for Children With MIS-C

Initial Immunomodulatory Therapy for MIS-C

The Panel recommends consulting with a multidisciplinary team when managing immunomodulatory therapy for children with MIS-C (AIII). The multidisciplinary team may include experts in cardiology, hematology, infectious disease, intensive care, and rheumatology. MIS-C is defined by multiorgan dysfunction, and input from other pediatric subspecialists may be needed depending on the presentation of the individual patient. Thus, children with MIS-C should be cared for at centers with access to these pediatric specialists.

Intravenous immunoglobulin (IVIG) and glucocorticoids are the most commonly used immunomodulatory medications in reported cohorts of children with MIS-C.⁶⁻¹⁴ The American College of Rheumatology has outlined initial diagnostic and treatment considerations for patients with MIS-C and recommends using IVIG in combination with glucocorticoids as first-tier therapy for most hospitalized children with MIS-C.¹⁵ Several nonrandomized studies suggest that the use of IVIG plus glucocorticoids is associated with less treatment failure, faster recovery of cardiac function, shorter intensive care unit (ICU) stays, and less need for treatment escalation than IVIG monotherapy.^{7,16-20} Based on these data, the Panel recommends using **IVIG** in combination with low to moderate doses of glucocorticoids for children hospitalized with MIS-C (AIIb).

IVIG should be given at a dose of 2 g/kg of ideal body weight, with a maximum total dose of 100 g. The patient's cardiac function and fluid status should be monitored carefully during the IVIG infusion. IVIG can be given in divided doses of 1 g/kg of ideal body weight over 2 days if there is a concern about the patient's fluid status. Methylprednisolone 1 to 2 mg/kg/day, or another glucocorticoid at an equivalent dose, is considered low to moderate glucocorticoid dosing. Once there is clinical improvement (i.e., the child is afebrile, end organ dysfunction resolves, and inflammatory markers are trending downward), a steroid taper should be initiated. Typically, the taper lasts for several weeks to avoid rebound inflammation and is guided by the clinical status of the patient.

Glucocorticoid monotherapy is an alternative initial treatment for MIS-C. Some studies have shown that patients treated with this approach had similar outcomes to patients treated with IVIG monotherapy and IVIG plus glucocorticoids.^{18,20,21} However, secondary analyses indicate that patients who were initially treated with IVIG plus glucocorticoids had faster time to improvement, less need for treatment escalation, and faster time to defervescence than patients who received glucocorticoid monotherapy.²⁰ Thus, the combination of IVIG and glucocorticoids appears to provide additional benefits that are not provided by glucocorticoid monotherapy.

Initial treatment that includes IVIG is also beneficial because it reduces the frequency of coronary artery aneurysms (CAAs) in patients with Kawasaki disease.^{14,22} Kawasaki disease is increasingly difficult to differentiate from MIS-C, and more recent SARS-CoV-2 variants have resulted in MIS-C presentations that are similar to Kawasaki disease.²³ Distinguishing MIS-C from Kawasaki disease is further complicated by the fact that seropositivity for SARS-CoV-2 is now widespread, making it difficult to establish the epidemiological link required for the MIS-C diagnosis. For these reasons, the Panel recommends using **IVIG** plus glucocorticoids as the initial therapy for patients with MIS-C (**AIIb**). **Glucocorticoid monotherapy** is recommended **only** if IVIG is unavailable or contraindicated (**BIIa**). **IVIG monotherapy** is recommended **only** if glucocorticoids are contraindicated (**BIIb**).

Clinical Data on Initial Immunomodulatory Therapy for MIS-C

Intravenous Immunoglobulin in Combination With Glucocorticoids

No randomized clinical trials evaluating the use of IVIG plus glucocorticoids for the treatment of MIS-C

COVID-19 Treatment Guidelines

have been completed. The comparative benefit of adding steroids to IVIG for MIS-C treatment has been estimated in observational cohort studies that used statistical techniques to adjust for confounders. The first of these studies employed observational data from a national surveillance system cohort in France and used propensity matching to compare short-term outcomes in children with MIS-C who were treated initially with IVIG 2 gm/kg alone or IVIG plus methylprednisolone (most patients received 1.6–2 mg/ kg/day for 5 days).¹⁶ The study team observed a lower risk of treatment failure (defined as a fever that persisted for 2 days after treatment or recurrent fever within 7 days), less need for hemodynamic support, less severe left ventricular dysfunction, and shorter ICU stays among the children who were initially treated with the combination therapy.¹⁶ This was a small study, and only 32 patients treated with IVIG plus methylprednisolone and 64 patients treated with IVIG alone could be matched based on propensity score.

A larger study in the United States analyzed data from the Overcoming COVID-19 surveillance registry to evaluate immunomodulatory therapy for MIS-C. The study included 103 patients who received initial treatment with IVIG plus glucocorticoids and an equal number of propensity score-matched patients who received IVIG alone. The risk of cardiovascular dysfunction on or after Day 2 was measured among these patients using a composite outcome of left ventricular ejection fraction of <55% or vasopressor use. The composite outcome occurred in 17% of patients in the IVIG plus glucocorticoids arm and in 31% of patients in the IVIG alone arm (risk ratio 0.56; 95% CI, 0.34–0.94).¹⁷ In addition, patients treated with the combination of IVIG and glucocorticoids were less likely to require adjunctive immunomodulatory therapy than those treated with IVIG alone. Methylprednisolone, the glucocorticoid that was prescribed most often, was administered to 353 patients (68% of patients, including nonpropensity score-matched patients, in the entire cohort). Among these patients, the dosing of methylprednisolone ranged from 2 mg/kg/day in 284 patients (80%) to 10 to 30 mg/kg/day in 69 patients (20%).

A third study, the international, observational BATS study, compared patients with MIS-C who received IVIG alone (n = 246) to those who received IVIG plus glucocorticoids (n = 208). This study found similar rates for the composite outcome of inotropic support or mechanical ventilation by Day 2 or later or death in both treatment arms. The composite outcome occurred in 44 of 221 patients (21%) in the IVIG alone arm and in 56 of 180 patients (31%) in the IVIG plus glucocorticoids arm (OR 0.77; 95% CI, 0.33–1.82). However, escalation of immunomodulatory treatment was less common among the patients who received IVIG plus glucocorticoids than among those who received IVIG alone (OR 0.18; 95% CI, 0.10–0.33). It is notable that the study also allowed for the inclusion of patients who had any inflammatory illness after acute COVID-19 but who did not meet the CDC or World Health Organization (WHO) criteria for MIS-C. This multicenter study included sites from 34 counties, which introduced the potential for more variability in supportive care. In addition, the overall percentage of patients with abnormal cardiac findings (12% of the 538 patients) was lower than in other cohorts.¹⁸

Intravenous Immunoglobulin Monotherapy

The use of IVIG is long established for patients with Kawasaki disease, a syndrome that has overlapping manifestations with MIS-C, and thus the product's safety profile is well understood. In patients with Kawasaki disease, IVIG prevents the development of CAAs,^{22,24} a complication also observed in some patients with MIS-C. IVIG is the most frequently used therapy for MIS-C. In a national survey of U.S. institutional protocols for managing MIS-C, IVIG was the first-line therapy in 98% of 40 participating centers.²⁵

Data on the efficacy of IVIG in patients with MIS-C is extrapolated from case series that show mostly favorable outcomes. In a series of 539 MIS-C cases, 77% of the children received IVIG. A sizeable proportion of these children had reduced left ventricular ejection fraction at admission (172 of 503 evaluable patients [34.2%]); the symptom resolved by Day 30 in 156 of the children (90.7%). Although

these studies have not described the occurrence of specific adverse events related to IVIG use, the dosing used (IVIG 2 g/kg) has a well-established safety profile when used for Kawasaki disease.¹⁴

A limitation of all published studies on IVIG use for MIS-C is the frequent and often rapid sequential addition of other immunomodulatory therapies, such as corticosteroids. In addition, there is accumulating evidence that glucocorticoids given in combination with IVIG are more effective as treatment for MIS-C. However, IVIG monotherapy may be a reasonable treatment option for a small subset of patients with MIS-C who are stable (i.e., not in shock or with organ-threatening disease) and have contraindications for glucocorticoid therapy. Such contraindications may include concern about the impact of corticosteroids on the diagnostic evaluation or an underlying medical condition.

Glucocorticoid Monotherapy

The observational BATS study also compared initial treatment with IVIG (n = 246) to treatment with glucocorticoids (n = 99) and found no differences in primary or secondary outcomes between these 2 cohorts.¹⁸ However, in a subgroup analysis of patients who met the WHO criteria for MIS-C, the glucocorticoid arm (n = 78) had significantly fewer patients who required respiratory support by Day 2 or later or who died than the IVIG arm (n = 192).

In a subsequent publication, the BATS consortium reported on additional patients with MIS-C who were enrolled in the study (over 2,000 patients in total).²⁰ The study had 2 primary outcomes. The first was a composite of the need for inotropic or ventilator support on or after Day 2 or death. The second was time to improvement by 1 level on an ordinal severity scale. In this larger study, there was once again no difference in the primary outcomes among the arms in a propensity-weighted analysis (combination therapy with IVIG plus glucocorticoids was compared to IVIG alone, and glucocorticoid monotherapy was compared to IVIG alone).

In secondary analyses, there were lower rates of treatment escalation among patients who received combination therapy than among those who received IVIG alone, and lower rates of treatment escalation among patients who received glucocorticoid monotherapy than among those who received IVIG alone. There was faster time to improvement, less need for treatment escalation, and lower rates of persistent fever on Day 2 in the combination therapy arm compared to the glucocorticoid monotherapy arm. The frequency of CAAs measured at hospital discharge and the severity of CAAs were similar in these treatment arms. Of the 236 patients with documented CAAs during the initial hospitalization, 196 had follow-up echocardiograms. Over 90% of the CAAs resolved, with similar rates of resolution across the treatment groups.

As in the initial publication for the observational BATS study, the inclusion criteria are broad and the patients did not need to meet the full WHO case definition for MIS-C. Compared to the other treatment arms, a greater proportion of the patients in the IVIG plus glucocorticoid arm met the WHO case definition for MIS-C, were ventilated and/or treated with inotropes at Day 0, and had CAAs (even before the initiation of immunomodulators). Many patients received additional immunomodulatory agents after Day 1, including 230 of 487 patients in the initial glucocorticoids alone group who also received IVIG. Finally, COVID-19 vaccination has been associated with reduced incidence and severity of MIS-C, but this was not evaluated in the study.^{26,27}

To date, the only randomized trial that evaluated treatments in patients with MIS-C was conducted in Switzerland.²¹ This open-label, multicenter study compared methylprednisolone 10 mg/kg per day for 3 days (n = 37) to a single dose of IVIG 2 gm/kg (n = 38). In this study, patients met the criteria for the case definition of pediatric multisystem inflammatory syndrome—temporally associated with SARS-CoV-2 (PMIS-TS). There was no difference in the primary outcome of length of hospital stay or death between the 2 arms. The length of hospital stay from admission to discharge was 6 days for both arms (estimated effect size -0.037 of the \log_{10} transformed times; 95% CI, -0.13 to 0.065; P = 0.42). No deaths were reported in either arm. In a secondary analysis, 27% of patients in the glucocorticoid arm required respiratory support compared to 55% of those treated with IVIG, which was a significant difference. There was no difference in the occurrence of coronary artery enlargement between the 2 arms. The small sample size in this study limited the power for treatment comparisons, and many patients received additional therapies for MIS-C after randomization.

Intensification Immunomodulatory Therapy for MIS-C

Children with MIS-C typically respond briskly to immunomodulatory therapy and show clinical improvements within the first 24 hours of treatment. Treatment response is characterized by resolution of fever, improvement of organ function, and reduced levels of inflammatory markers, particularly C-reactive protein. In contrast, refractory disease is often accompanied by persistent fever, worsening organ dysfunction, and increasing levels of inflammatory markers. Intensification therapy is recommended for children with refractory MIS-C who do not improve within 24 hours of receiving initial immunomodulatory therapy (AIII). Children with uncontrolled MIS-C despite treatment with IVIG and low to moderate doses of glucocorticoids will often continue to deteriorate without further intervention, and this decline in clinical status can be quite rapid.

No comparative studies have evaluated intensification therapies for MIS-C. The data on this topic are limited to results from cohort studies in patients with MIS-C, expert opinion, and experience in treating other hyperinflammatory syndromes in children, such as Kawasaki disease and macrophage activation syndrome. For children with refractory MIS-C, the Panel recommends providing additional immunomodulatory therapy (in alphabetical order) with **anakinra (BIIb)**, **higher-dose glucocorticoids** (**BIIb**), or **infliximab (BIIb**). Currently, there is insufficient evidence to determine which of these agents is most effective for intensification therapy in patients with refractory MIS-C. In patients with refractory severe disease, some Panel members would use dual therapy with **higher-dose glucocorticoids** and **anakinra (BIII)** or **higher-dose glucocorticoids** and **infliximab (BIII)** or **intensification therapy**. Anakinra and infliximab **should not be used** in combination. A second dose of IVIG is not commonly reported in the literature as a strategy for intensification therapy in patients with MIS-C. This may be due to the high rates of IVIG resistance, the rapid pace of disease escalation, and the risk for fluid overload in patients with MIS-C.¹⁰ Therefore, the Panel **recommends against** a second dose of **IVIG** for intensification therapy in patients with refractory MIS-C (**BIII**).

Patients with MIS-C who receive multiple immunomodulatory agents are at risk for infection and need to be monitored carefully. Most children with MIS-C were previously healthy. In patients who have an immune disorder or are taking immunosuppression therapy, the risk of infection is greater. The risks and benefits of using immunomodulatory agents in patients with MIS-C who are immunocompromised need to be evaluated on a case-by-case basis.

Clinical Data on Intensification Immunomodulatory Therapy for MIS-C

High-Dose Glucocorticoids

High-dose glucocorticoid therapy is defined as methylprednisolone (or an equivalent corticosteroid) dosed at 10 to 30 mg/kg/day and given intravenously (IV). Often, this higher dose of glucocorticoids is given for 1 to 3 days before returning to low to moderate doses (1–2 mg/kg/day). Multiple observational studies have evaluated the use of high-dose glucocorticoids (methylprednisolone 10–30 mg/kg/day) in children with MIS-C.^{17,28-30} In addition, single-center treatment protocols for MIS-C that incorporate high-dose glucocorticoids into the treatment algorithm have been published. Implementation of the protocols has resulted in positive clinical outcomes in patients with MIS-C.¹⁹ There is substantial experience with using high-dose glucocorticoids in pediatric patients with other inflammatory

conditions, such as Kawasaki disease and macrophage activation syndrome.

Anakinra

Anakinra is the most commonly used biologic medication for the treatment of MIS-C in the United States.²⁵ Multiple noncomparative, observational cohorts have reported on the use of anakinra in patients with MIS-C.^{10,11,13,31-33} This medication has been used extensively and has a good safety record in pediatric patients with other hyperinflammatory syndromes (e.g., systemic juvenile idiopathic arthritis, macrophage activation syndrome).³⁴⁻³⁶ Anakinra has also been used successfully to treat IVIG-resistant Kawasaki disease. Anakinra has a short half-life (4–6 hours), and the medication can be stopped quickly, which many providers regard as a benefit relative to longer-acting immunomodulators. High-dose anakinra (5–10 mg/kg/day) is recommended for patients with MIS-C based on the demonstrated efficacy of high-dose anakinra in patients with macrophage activation syndrome. The duration of anakinra therapy varies in the literature and is used by some patients for long periods (e.g., up to 2 weeks) as a steroid-sparing agent.

Infliximab

The Panel recommends a single dose of infliximab 5 to 10 mg/kg IV as an option for intensification therapy. Infliximab has been studied for the treatment of MIS-C in a single-center retrospective study that compared patients treated with IVIG alone (n = 20) to those treated with IVIG and a single dose of infliximab 10 mg/kg IV (n = 52).³⁷ Of note, infliximab was used as the first-line therapy in this study, and the patients were not treated with glucocorticoids. The patients who received IVIG and infliximab were more likely to be admitted to the ICU and had more severe illness than those who received IVIG alone. However, the patients who received the combination therapy were less likely to require additional therapy after 24 hours (the primary outcome). In addition, patients who received IVIG and infliximab had shorter stays in the ICU and improved cardiac outcomes. These results show that infliximab has a therapeutic effect in patients with MIS-C.

Infliximab is approved by the Food and Drug Administration for use in children with inflammatory bowel disease and is used widely to treat juvenile idiopathic arthritis. Infliximab has been employed in IVIG-resistant Kawasaki disease.^{38,39} Although the half-life of infliximab in patients with MIS-C is unknown, it likely has effects that persist for several weeks. This extended period of drug activity can allow for a steroid-sparing effect in patients with MIS-C.

Antithrombotic Therapy for MIS-C

There is general agreement that patients with MIS-C who do not have risk factors for bleeding should receive low-dose aspirin (AIII). This recommendation is largely due to experience in treating children with Kawasaki disease and the likelihood of analogous platelet activation and endothelial dysfunction in children with MIS-C.⁴⁰ Children treated with aspirin and steroids should also receive prophylactic H2 blockers or proton pump inhibitors. Patients with MIS-C who have large CAAs (Z-score ≥ 10) should receive therapeutic anticoagulation according to the <u>American Heart Association guidelines</u> for Kawasaki disease (AIII). Children with left ventricular dysfunction are at risk for intracardiac thrombosis. Patients with MIS-C and moderate-to-severe left ventricular dysfunction should receive therapeutic anticoagulation, unless it is contraindicated due to bleeding risk factors (AIII).

There is less consensus on the use of either prophylactic or therapeutic anticoagulation in patients with MIS-C who do not have large CAAs and/or moderate to severe left ventricular dysfunction. Children with MIS-C have marked elevations in D-dimer levels and other abnormalities of coagulation, which suggests that they may be at increased risk for thrombosis.⁴¹ In a multicenter retrospective study of children with acute COVID-19 and MIS-C, the independent risk factors for thrombosis included indwelling catheters, older age (>12 years), malignancy, admission to the ICU, and elevated D-dimer

levels.⁴² In a multicenter, Phase 2 trial of enoxaparin thromboprophylaxis in children hospitalized for COVID-19 and MIS-C (COVAC-TP), children with MIS-C frequently exhibited hyperfibrinogenemia and had significantly elevated D-dimer levels compared to children with primary SARS-CoV-2 infection.⁴³ There are limited published data on the risk of bleeding in children with MIS-C who are managed with anticoagulant thromboprophylaxis. Major bleeding events (as defined by the International Society on Thrombosis and Haemostasis) were observed in patients with MIS-C who were treated with anticoagulation in the aforementioned retrospective study⁴² but not in the COVAC-TP trial, which employed prophylactic dosing of enoxaparin and permitted the use of aspirin at a dose of up to 5 mg/kg/day.⁴³ However, 5% of patients developed catheter-related thromboembolic events despite the use of enoxaparin thromboprophylaxis in the COVAC-TP trial.

Given the uncertainty regarding the benefit of anticoagulation for MIS-C, prophylactic or therapeutic anticoagulation for children with MIS-C who do not have large CAAs or moderate to severe left ventricular dysfunction should be considered on a case-by-case basis, taking into account the risk factors for thrombosis and bleeding.

Antiviral Therapy for MIS-C

The role of SARS-CoV-2 antiviral therapy in treating MIS-C has not been systematically studied; however, it is not expected to be beneficial because MIS-C is considered an immune-mediated phenomenon that occurs weeks after primary SARS-CoV-2 infection. Therefore, the Panel **recommends against** the use of SARS-CoV-2 antiviral therapy for patients with MIS-C (AIII).

Critical Care Management

Shock occurs in approximately 50% of patients with MIS-C and may include elements of distributive, cardiogenic, or hypovolemic shock.^{14,44,45} In general, clinicians should manage shock in patients with MIS-C per the usual critical care standards as outlined in the Pediatric Surviving Sepsis Campaign Guidelines.⁴⁶

References

- Centers for Disease Control and Prevention. Information for healthcare providers about multisystem inflammatory syndrome in children (MIS-C). 2023. Available at: <u>https://www.cdc.gov/mis/mis-c/hcp_cstecdc/ index.html</u>. Accessed June 23, 2023.
- Centers for Disease Control and Prevention. Multisystem inflammatory syndrome in adults (MIS-A) case definition information for healthcare providers. 2021. Available at: <u>https://www.cdc.gov/mis/mis-a/hcp.html</u>. Accessed July 6, 2023.
- Morris SB, Schwartz NG, Patel P, et al. Case series of multisystem inflammatory syndrome in adults associated with SARS-CoV-2 infection—United Kingdom and United States, March–August 2020. MMWR Morb Mortal Wkly Rep. 2020;69(40):1450-1456. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33031361.
- Kunal S, Ish P, Sakthivel P, Malhotra N, Gupta K. The emerging threat of multisystem inflammatory syndrome in adults (MIS-A) in COVID-19: a systematic review. *Heart Lung*. 2022;54:7-18. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35306376</u>.
- 5. Melgar M, Abrams JY, Godfred-Cato S, et al. A multicenter retrospective cohort study to characterize patients hospitalized with MIS-A and COVID-19 in the United States, 2020-2021. *Clin Infect Dis.* 2023;Published online ahead of print. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37384794</u>.
- 6. Verdoni L, Mazza A, Gervasoni A, et al. An outbreak of severe Kawasaki-like disease at the Italian epicentre of the SARS-CoV-2 epidemic: an observational cohort study. *Lancet*. 2020;395(10239):1771-1778. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32410760.

- 7. Belhadjer Z, Auriau J, Méot M, et al. Addition of corticosteroids to immunoglobulins is associated with recovery of cardiac function in multi-inflammatory syndrome in children. *Circulation*. 2020;142(23):2282-2284. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33112651.
- 8. Toubiana J, Poirault C, Corsia A, et al. Kawasaki-like multisystem inflammatory syndrome in children during the COVID-19 pandemic in Paris, France: prospective observational study. *BMJ*. 2020;369:m2094. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32493739.
- Whittaker E, Bamford A, Kenny J, et al. Clinical characteristics of 58 children with a pediatric inflammatory multisystem syndrome temporally associated with SARS-CoV-2. *JAMA*. 2020;324(3):259-269. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32511692</u>.
- Pouletty M, Borocco C, Ouldali N, et al. Paediatric multisystem inflammatory syndrome temporally associated with SARS-CoV-2 mimicking Kawasaki disease (Kawa-COVID-19): a multicentre cohort. *Ann Rheum Dis*. 2020;79(8):999-1006. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32527868</u>.
- Feldstein LR, Rose EB, Horwitz SM, et al. Multisystem inflammatory syndrome in U.S. children and adolescents. *N Engl J Med*. 2020;383(4):334-346. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32598831</u>.
- 12. Dufort EM, Koumans EH, Chow EJ, et al. Multisystem inflammatory syndrome in children in New York State. *N Engl J Med.* 2020;383(4):347-358. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32598830</u>.
- 13. Lee PY, Day-Lewis M, Henderson LA, et al. Distinct clinical and immunological features of SARS-CoV-2induced multisystem inflammatory syndrome in children. *J Clin Invest*. 2020;130(11):5942-5950. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32701511.
- Feldstein LR, Tenforde MW, Friedman KG, et al. Characteristics and outcomes of US children and adolescents with multisystem inflammatory syndrome in children (MIS-C) compared with severe acute COVID-19. *JAMA*. 2021;325(11):1074-1087. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33625505</u>.
- Henderson LA, Canna SW, Friedman KG, et al. American College of Rheumatology clinical guidance for multisystem inflammatory syndrome in children associated with SARS-CoV-2 and hyperinflammation in pediatric COVID-19: version 3. *Arthritis Rheumatol*. 2022;74(4):e1-e20. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35118829</u>.
- 16. Ouldali N, Toubiana J, Antona D, et al. Association of intravenous immunoglobulins plus methylprednisolone vs immunoglobulins alone with course of fever in multisystem inflammatory syndrome in children. *JAMA*. 2021;325(9):855-864. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33523115</u>.
- 17. Son MBF, Murray N, Friedman K, et al. Multisystem inflammatory syndrome in children—initial therapy and outcomes. *N Engl J Med*. 2021;385(1):23-34. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133855</u>.
- McArdle AJ, Vito O, Patel H, et al. Treatment of multisystem inflammatory syndrome in children. N Engl J Med. 2021;385(1):11-22. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133854</u>.
- 19. Jonat B, Gorelik M, Boneparth A, et al. Multisystem inflammatory syndrome in children associated with coronavirus disease 2019 in a children's hospital in New York City: patient characteristics and an institutional protocol for evaluation, management, and follow-up. *Pediatr Crit Care Med.* 2021;22(3):e178-e191. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33003176.
- Channon-Wells S, Vito O, McArdle AJ, et al. Immunoglobulin, glucocorticoid, or combination therapy for multisystem inflammatory syndrome in children: a propensity-weighted cohort study. *Lancet Rheumatol*. 2023;5(4):e184-e199. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36855438</u>.
- 21. Welzel T, Atkinson A, Schöbi N, et al. Methylprednisolone versus intravenous immunoglobulins in children with paediatric inflammatory multisystem syndrome temporally associated with SARS-CoV-2 (PIMS-TS): an open-label, multicentre, randomised trial. *Lancet Child Adolesc Health*. 2023;7(4):238-248. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36746174</u>.
- 22. Newburger JW, Takahashi M, Burns JC, et al. The treatment of Kawasaki syndrome with intravenous gamma globulin. *N Engl J Med.* 1986;315(6):341-347. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/2426590</u>.

- 23. McCrindle BW, Harahsheh AS, Handoko R, et al. SARS-CoV-2 variants and multisystem inflammatory syndrome in children. *N Engl J Med.* 2023;388(17):1624-1626. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36947454.</u>
- 24. Furusho K, Kamiya T, Nakano H, et al. High-dose intravenous gammaglobulin for Kawasaki disease. *Lancet*. 1984;2(8411):1055-1058. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/6209513</u>.
- 25. Dove ML, Jaggi P, Kelleman M, et al. Multisystem inflammatory syndrome in children: survey of protocols for early hospital evaluation and management. *J Pediatr*. 2021;229:33-40. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33075369.
- 26. Levy M, Recher M, Hubert H, et al. Multisystem inflammatory syndrome in children by COVID-19 vaccination status of adolescents in France. *JAMA*. 2022;327(3):281-283. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34928295</u>.
- 27. Zambrano LD, Newhams MM, Olson SM, et al. BNT162b2 mRNA vaccination against coronavirus disease 2019 is associated with a decreased likelihood of multisystem inflammatory syndrome in children aged 5–18 years—United States, July 2021–April 2022. *Clin Infect Dis.* 2023;76(3):e90-e100. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35924406</u>.
- 28. Chiotos K, Bassiri H, Behrens EM, et al. Multisystem inflammatory syndrome in children during the coronavirus 2019 pandemic: a case series. *J Pediatric Infect Dis Soc*. 2020;9(3):393-398. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32463092.
- 29. Newburger JW, Sleeper LA, McCrindle BW, et al. Randomized trial of pulsed corticosteroid therapy for primary treatment of Kawasaki disease. *N Engl J Med*. 2007;356(7):663-675. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/17301297</u>.
- 30. Inoue Y, Okada Y, Shinohara M, et al. A multicenter prospective randomized trial of corticosteroids in primary therapy for Kawasaki disease: clinical course and coronary artery outcome. *J Pediatr.* 2006;149(3):336-341. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/16939743</u>.
- Çağlayan Ş, Sönmez HE, Yener GO, et al. Anakinra treatment in multisystemic inflammatory syndrome in children (MIS-C) associated with COVID-19. *Front Pediatr*. 2022;10:942455. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36061383</u>.
- 32. Brisca G, Consolaro A, Caorsi R, et al. Timely recognition and early multi-step antinflammatory therapy may prevent ICU admission in patients with MIS-C: proposal for a severity score. *Front Pediatr.* 2021;9:783745. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/34988039</u>.
- 33. Chang JC, Young CC, Muscal E, et al. Variation in early anakinra use and short-term outcomes in multisystem inflammatory syndrome in children. *Arthritis Rheumatol*. 2023;75(8):1466-1476. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36908050</u>.
- 34. Eloseily EM, Weiser P, Crayne CB, et al. Benefit of anakinra in treating pediatric secondary hemophagocytic lymphohistiocytosis. *Arthritis Rheumatol*. 2020;72(2):326-334. Available at: https://www.ncbi.nlm.nih.gov/pubmed/31513353.
- 35. Quartier P, Allantaz F, Cimaz R, et al. A multicentre, randomised, double-blind, placebo-controlled trial with the interleukin-1 receptor antagonist anakinra in patients with systemic-onset juvenile idiopathic arthritis (ANAJIS trial). *Ann Rheum Dis.* 2011;70(5):747-754. Available at: https://www.ncbi.nlm.nih.gov/pubmed/21173013.
- 36. Ter Haar NM, van Dijkhuizen EHP, Swart JF, et al. Treatment to target using recombinant interleukin-1 receptor antagonist as first-line monotherapy in new-onset systemic juvenile idiopathic arthritis: results from a five-year follow-up study. *Arthritis Rheumatol*. 2019;71(7):1163-1173. Available at: https://www.ncbi.nlm.nih.gov/pubmed/30848528.
- 37. Cole LD, Osborne CM, Silveira LJ, et al. IVIG compared to IVIG plus infliximab in multisystem inflammatory syndrome in children. *Pediatrics*. 2021;148(6):e2021053702. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34548377</u>.

- 38. Mori M, Hara T, Kikuchi M, et al. Infliximab versus intravenous immunoglobulin for refractory Kawasaki disease: a Phase 3, randomized, open-label, active-controlled, parallel-group, multicenter trial. *Sci Rep.* 2018;8(1):1994. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29386515</u>.
- 39. Yamaji N, da Silva Lopes K, Shoda T, et al. TNF-alpha blockers for the treatment of Kawasaki disease in children. *Cochrane Database Syst Rev.* 2019;8(8):CD012448. Available at: https://www.ncbi.nlm.nih.gov/pubmed/31425625.
- 40. McCrindle BW, Rowley AH, Newburger JW, et al. Diagnosis, treatment, and long-term management of Kawasaki disease: a scientific statement for health professionals from the American Heart Association. *Circulation*. 2017;135(17):e927-e999. Available at: https://www.ncbi.nlm.nih.gov/pubmed/28356445.
- 41. Ankola AA, Bradford VR, Newburger JW, et al. Coagulation profiles and viscoelastic testing in multisystem inflammatory syndrome in children. *Pediatr Blood Cancer*. 2021;68(12):e29355. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34532964.
- 42. Whitworth H, Sartain SE, Kumar R, et al. Rate of thrombosis in children and adolescents hospitalized with COVID-19 or MIS-C. *Blood*. 2021;138(2):190-198. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33895804.
- 43. Sochet AA, Morrison JM, Jaffray J, et al. Enoxaparin thromboprophylaxis in children hospitalized for COVID-19: a Phase 2 trial. *Pediatrics*. 2022;150(1):e2022056726. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35484817</u>.
- 44. Abrams JY, Oster ME, Godfred-Cato SE, et al. Factors linked to severe outcomes in multisystem inflammatory syndrome in children (MIS-C) in the USA: a retrospective surveillance study. *Lancet Child Adolesc Health*. 2021;5(5):323-331. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33711293</u>.
- 45. Godfred-Cato S, Bryant B, Leung J, et al. COVID-19-associated multisystem inflammatory syndrome in children—United States, March–July 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(32):1074-1080. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32790663</u>.
- 46. Weiss SL, Peters MJ, Agus MSD, et al. Perspective of the Surviving Sepsis Campaign on the management of pediatric sepsis in the era of coronavirus disease 2019. *Pediatr Crit Care Med.* 2020;21(11):e1031-e1037. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32886460</u>.

Care of Critically III Adults With COVID-19

Last Updated: December 20, 2023

Summary Recommendations

Hemodynamics

- For adults with COVID-19 and shock, the COVID-19 Treatment Guidelines Panel (the Panel) recommends using dynamic parameters, skin temperature, capillary refilling time, and/or lactate levels over static parameters to assess fluid responsiveness (**BIIa**).
- For the acute resuscitation of adults with COVID-19 and shock, the Panel recommends using buffered/balanced crystalloids (BIIa).
- For the acute resuscitation of adults with COVID-19 and shock, the Panel **recommends against** the initial use of **albumin** for resuscitation **(BI)**.
- For adults with COVID-19 and shock, the Panel recommends **norepinephrine** as the first-choice vasopressor (AI).
- For adults with COVID-19 and shock, the Panel recommends titrating vasoactive agents to target a mean arterial pressure (MAP) of 60 to 65 mm Hg, over higher MAP targets (BI).
- The Panel **recommends against** using hydroxyethyl starches for intravascular volume replacement in adult patients with COVID-19 and sepsis or septic shock (AI).
- When norepinephrine is available, the Panel **recommends against** using **dopamine** for adult patients with COVID-19 and shock **(AI)**.
- As a second-line vasopressor, the Panel recommends adding either vasopressin (up to 0.03 units/min) (Blla) or epinephrine (Bllb) to norepinephrine to raise MAP to target or adding vasopressin (up to 0.03 units/min) (Blla) to decrease norepinephrine dosage.
- The Panel recommends against using low-dose dopamine for renal protection (AI).
- The Panel recommends using **dobutamine** in adult patients with COVID-19 who show evidence of cardiac dysfunction and persistent hypoperfusion despite adequate fluid loading and the use of vasopressor agents **(BIII)**.
- The Panel recommends that all adult patients with COVID-19 who require vasopressors have an arterial catheter placed as soon as practical, if resources are available (**BIII**).
- For adult patients with refractory septic shock who have completed a course of corticosteroids to treat COVID-19, the Panel recommends using low-dose corticosteroid therapy ("shock-reversal") over no corticosteroid therapy (Blla).

Oxygenation and Ventilation

- For adults with COVID-19 and acute hypoxemic respiratory failure despite conventional oxygen therapy, the Panel recommends starting therapy with high-flow nasal cannula (HFNC) oxygen; if patients fail to respond, noninvasive ventilation or intubation and mechanical ventilation should be initiated (**Blla**).
- For adults with COVID-19 and acute hypoxemic respiratory failure despite conventional oxygen therapy who do not have an indication for endotracheal intubation and for whom HFNC oxygen is not available, the Panel recommends performing a closely monitored trial of noninvasive ventilation (**BIIa**).
- For adults with persistent hypoxemia who require HFNC oxygen and for whom endotracheal intubation is not indicated, the Panel recommends a trial of awake prone positioning (**Blla**).
- The Panel **recommends against** the use of awake prone positioning as a rescue therapy for refractory hypoxemia to avoid intubation in patients who otherwise meet the indications for intubation and mechanical ventilation (AIII).
- If intubation becomes necessary, the procedure should be performed by an experienced practitioner in a controlled setting due to the enhanced risk of exposing health care practitioners to SARS-CoV-2 during intubation (AIII).
- For mechanically ventilated adults with COVID-19 and acute respiratory distress syndrome (ARDS):
 - The Panel recommends using low tidal volume (VT) ventilation (VT 4–8 mL/kg of predicted body weight) over higher VT ventilation (VT >8 mL/kg) (AI).
 - The Panel recommends targeting plateau pressures of <30 cm H₂0 (Alla).

Summary Recommendations, continued

- The Panel recommends using a conservative fluid strategy over a liberal fluid strategy (Blla).
- The Panel recommends against the routine use of inhaled nitric oxide (Alla).
- For mechanically ventilated adults with COVID-19 and moderate to severe ARDS:
 - The Panel recommends using a higher positive end-expiratory pressure (PEEP) strategy over a lower PEEP strategy (Blla).
 - For mechanically ventilated adults with COVID-19 and refractory hypoxemia despite optimized ventilation, the Panel recommends prone ventilation for 12 to 16 hours per day over no prone ventilation (**Blla**).
 - The Panel recommends using, as needed, intermittent boluses of neuromuscular blocking agents or a continuous neuromuscular blocking agent infusion to facilitate protective lung ventilation (Blla).
- For mechanically ventilated adults with COVID-19, severe ARDS, and hypoxemia despite optimized ventilation and other rescue strategies:
 - The Panel recommends using an inhaled pulmonary vasodilator as a rescue therapy; if rapid improvement in oxygenation is not observed, the treatment should be tapered (CIII).
 - The Panel recommends using recruitment maneuvers rather than not using recruitment maneuvers (Clla).
 - If recruitment maneuvers are used, the Panel **recommends against** the use of staircase (incremental PEEP) recruitment maneuvers (Alla).

Pharmacologic Interventions

- In the absence of a proven or suspected secondary infection, the Panel **recommends against** the use of empiric broad-spectrum antimicrobials in patients with severe or critical COVID-19 (BIII).
- As with any hospitalized patient, patients with COVID-19 who receive antimicrobials should be reassessed daily to minimize the adverse consequences of unnecessary antimicrobial therapy (AIII).

Extracorporeal Membrane Oxygenation

• There is insufficient evidence for the Panel to recommend either for or against the use of extracorporeal membrane oxygenation in adults with COVID-19–associated ARDS and refractory hypoxemia.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Introduction to Critical Care Management of Adults With COVID-19

Last Updated: May 31, 2022

COVID-19 can progress to critical illness, including hypoxemic respiratory failure, acute respiratory distress syndrome (ARDS), septic shock, cardiac dysfunction, thromboembolic disease, hepatic and/ or renal dysfunction, central nervous system disease, and exacerbation of underlying comorbidities in both adults and children. In addition, multisystem inflammatory syndrome in adults (MIS-A) can occur several weeks or months after SARS-CoV-2 infection, which can lead to critical illness.

Many of the initial recommendations for the management of critically ill adults with COVID-19 in these Guidelines were extrapolated from experience with other causes of sepsis and respiratory failure.¹ However, there is now a rapidly growing body of evidence regarding the management of critically ill patients with COVID-19.

Treating patients with COVID-19 in the intensive care unit (ICU) often requires managing underlying illnesses or COVID-19-related morbidities. As with any patient who is admitted to the ICU, clinicians also need to focus on preventing ICU-related complications.

Selected Clinical Manifestations of COVID-19 Critical Illness

Inflammatory Response Due to COVID-19 in Adults

Patients with COVID-19 may express increased levels of pro-inflammatory cytokines and antiinflammatory cytokines, which has previously been referred to as "cytokine release syndrome" or "cytokine storm." However, these terms are both imprecise and misnomers, because the magnitude of cytokine elevation in many patients with COVID-19 is modest compared to that in patients with many other critical illnesses, such as sepsis and ARDS.^{2,3} In addition, some patients with elevated cytokine levels have no specific pathology that can be attributed to the elevated levels.

Patients with COVID-19 and severe pulmonary involvement often manifest extrapulmonary disease and exhibit laboratory markers of acute inflammation. Patients with these manifestations of severe pulmonary disease typically progress to critical illness 10 to 12 days after the onset of COVID-19 symptoms.

Multisystem Inflammatory Syndrome in Adults

There are case reports describing patients who had evidence of acute or recent SARS-CoV-2 infection (confirmed by a nucleic acid amplification test [NAAT] or an antigen or antibody test) with minimal respiratory symptoms but with laboratory markers of severe inflammation (e.g., elevated levels of C-reactive protein [CRP], ferritin, D-dimer, cardiac enzymes, liver enzymes, and creatinine) and various other symptoms, including fever and shock. These patients also had signs of cardiovascular, gastrointestinal, dermatologic, and neurologic disease. This constellation of signs and symptoms has been designated MIS-A.⁴ To date, most adults with MIS-A have survived. This syndrome is similar to multisystem inflammatory syndrome in children (MIS-C), which is much more well described.

The current case definition for MIS-A from the <u>Centers for Disease Control and Prevention</u> states that patients must be aged ≥ 21 years, be hospitalized for ≥ 24 hours or have an illness that results in death, and meet the clinical and laboratory criteria outlined below. The patient should not have a more likely alternative diagnosis for the illness (e.g., bacterial sepsis, exacerbation of a chronic medical condition).

Clinical Criteria

Patients must have a subjective or documented fever (\geq 38.0°C) for \geq 24 hours prior to hospitalization or within the first 3 days of hospitalization and at least 3 of the following clinical criteria, which must have occurred prior to hospitalization or within the first 3 days of hospitalization. At least 1 must be a primary clinical criterion.

- Primary clinical criteria:
 - Severe cardiac illness. This includes myocarditis; pericarditis; coronary artery dilatation/ aneurysm; or new-onset right or left ventricular dysfunction (left ventricular ejection fraction <50%), second- or third-degree atrioventricular block, or ventricular tachycardia. Cardiac arrest alone does not meet this criterion.
 - Rash AND nonpurulent conjunctivitis
- Secondary clinical criteria:
 - New-onset neurologic signs and symptoms. These include encephalopathy in a patient without prior cognitive impairment, seizures, meningeal signs, or peripheral neuropathy (including Guillain-Barré syndrome).
 - Shock or hypotension that are not attributable to medical therapy (e.g., sedation, renal replacement therapy)
 - Abdominal pain, vomiting, or diarrhea
 - Thrombocytopenia (platelet count <150,000 cells/µL)

Laboratory Criteria

- The presence of laboratory evidence of inflammation AND SARS-CoV-2 infection
- Elevated levels of at least 2 of the following:
 - CRP
 - Ferritin
 - Interleukin (IL)-6
 - Erythrocyte sedimentation rate
 - Procalcitonin
- A positive SARS-CoV-2 test result for current or recent infection using a reverse transcription polymerase chain reaction, serology, or antigen test

These criteria must be met by the end of Day 3 of hospitalization, where the date of hospital admission is Day 0.

Because there is no specific diagnostic test for MIS-A, diagnosis of this inflammatory syndrome is one of exclusion after other causes (e.g., bacterial sepsis) have been eliminated. Although there are currently no controlled clinical trial data in patients with MIS-A to guide treatment of the syndrome, case reports have described the use of intravenous immunoglobulin, corticosteroids, or anti-IL-1 receptor antagonist therapy.⁵⁻⁷

COVID-19-Induced Cardiac Dysfunction, Including Myocarditis

The published literature describes cardiac injury or dysfunction in up to 24% of adults who are hospitalized with COVID-19.⁸ COVID-19 may be associated with an array of cardiovascular complications, including acute coronary syndrome, myocarditis, stress (Takotsubo) cardiomyopathy,

COVID-19 Treatment Guidelines

arrythmias, and thromboembolic disease.9

Thromboembolic Events and COVID-19

Critically ill adults with COVID-19 have been observed to have a prothrombotic state and higher rates of venous thromboembolic disease. In some studies, thromboemboli have been diagnosed even in patients who received chemical prophylaxis with heparinoids.¹⁰⁻¹² Autopsy studies provide additional evidence of both thromboembolic disease and microvascular thrombosis in patients with COVID-19.¹³ Some authors have called for routine surveillance of ICU patients for venous thromboembolism.¹⁴ See Antithrombotic Therapy in Patients With COVID-19 for a more detailed discussion.

Renal and Hepatic Dysfunction Due to COVID-19

Although SARS-CoV-2 is primarily a pulmonary pathogen, renal and hepatic dysfunction are consistently described in adults with severe COVID-19.¹⁵ In a 2020 multicenter cohort study of critically ill adults in the United States, 20.6% of patients developed acute kidney injury (AKI) that was treated with renal replacement therapy (RRT).¹⁶ In a cohort of critically ill adults in Brazil, the development of an AKI that required RRT was associated with poor prognosis.¹⁷

Other Intensive Care Unit-Related Complications

When treating patients with COVID-19, clinicians also need to minimize the risk of conventional ICU complications. Patients who are critically ill with COVID-19 are at risk for nosocomial infections, such as ventilator-associated pneumonia, hospital-acquired pneumonia, catheter-related bloodstream infections, and other complications of critical illness care.

Critically ill patients with COVID-19 may also experience prolonged delirium and/or encephalopathy. The risk factors that are associated with delirium include the use of mechanical ventilation, restraints, benzodiazepines, opioids, vasopressors, and antipsychotics.^{18,19} Neurological manifestations of COVID-19 have been described in a significant proportion of hospitalized patients and are more frequent in patients with severe disease.²⁰ Autopsy studies have reported both macrovascular and microvascular thrombosis with evidence of hypoxic ischemia.²¹ Adequate management of critically ill patients with COVID-19 includes paying careful attention to best sedation practices and monitoring for stroke.

Important Considerations in the Care of Critically III Patients With COVID-19

Interactions Between Drugs Used to Treat COVID-19 and Drugs Used to Treat Comorbidities

All ICU patients should be routinely monitored for drug-drug interactions. The potential for drug-drug interactions between investigational medications or medications that are used off-label to treat COVID-19 and concurrent drugs should be considered.

Sedation Management in Adults With COVID-19

International guidelines provide recommendations on the prevention, detection, and treatment of pain, sedation, and delirium in ICU patients.^{22,23} Sedation management strategies, such as maintaining a light level of sedation (when appropriate) and minimizing sedative exposure, have shortened the duration of mechanical ventilation and the length of stay in the ICU for patients without COVID-19.^{24,25}

The Society of Critical Care Medicine's (SCCM) ICU Liberation Campaign promotes the ICU Liberation Bundle (A-F) to improve post-ICU patient outcomes. The A-F Bundle includes the following elements:

A. Assess, prevent, and manage pain;

COVID-19 Treatment Guidelines

- B. Both spontaneous awakening and breathing trials;
- C. Choice of analgesia and sedation;
- D. Delirium: assess, prevent, and manage;
- E. Early mobility and exercise; and
- F. Family engagement and empowerment.

The A-F Bundle also provides frontline staff with practical application strategies for each element.²⁶ The A-F Bundle should be incorporated using an interprofessional team model. This approach helps standardize communication among team members, improves survival, and reduces long-term cognitive dysfunction of patients.²⁷ Despite the known benefits of the A-F Bundle, its impact has not been directly assessed in patients with COVID-19; however, use of the Bundle should be encouraged, when appropriate, to improve ICU patient outcomes. Prolonged mechanical ventilation of COVID-19 patients, coupled with deep sedation and potentially neuromuscular blockade, increases the workload of ICU staff. Additionally, significant drug shortages may force clinicians to use older sedatives with prolonged durations of action and active metabolites, impeding routine implementation of SCCM's <u>PADIS</u> <u>guidelines</u>. This puts patients at additional risk for ICU and post-ICU complications.

Post-Intensive Care Syndrome

Post-intensive care syndrome (PICS) is a spectrum of cognitive, psychiatric, and/or physical disability that affects survivors of critical illness and persists after a patient leaves the ICU.²⁸ Patients with PICS may present with varying levels of impairment, including profound muscle weakness (ICU-acquired weakness); problems with thinking and judgment (cognitive dysfunction); and mental health problems, such as problems sleeping, post-traumatic stress disorder (PTSD), depression, and anxiety. ICU-acquired weakness affects 33% of all patients who receive mechanical ventilation, 50% of patients with sepsis, and \leq 50% of patients who remain in the ICU for \geq 1 week.²⁹⁻³¹ Cognitive dysfunction affects 30% to 80% of patients discharged from the ICU.³²⁻³⁴ About 50% of ICU survivors do not return to work within 1 year after discharge.³⁵ Although no single risk factor has been associated with PICS, there are opportunities to minimize the risk of PICS through medication management (using the A-F Bundle), physical rehabilitation, follow-up clinics, family support, and improved education about the syndrome. PICS also affects family members who participate in the care of their loved ones. In 1 study, a third of family members who had major decision-making roles experienced mental health problems, such as depression, anxiety, and PTSD.³⁶

Some patients with COVID-19 who have been treated in the ICU express manifestations of PICS.³⁷ Although specific therapies for COVID-19-induced PICS are not yet available, physicians should maintain a high index of suspicion for cognitive impairment and other related problems in survivors of severe or critical COVID-19 illness.

Advance Care Planning and Goals of Care

The advance care plans and the goals of care for all critically ill patients must be assessed at hospital admission and regularly thereafter. This is an essential element of care for all patients. Information on palliative care for patients with COVID-19 can be found on the <u>National Coalition for Hospice and</u> <u>Palliative Care website</u>.

To guide shared decision making in cases of serious illness, advance care planning should include identifying existing advance directives that outline a patient's preferences and values. Values and care preferences should be discussed, documented, and revisited regularly for patients with or without prior directives. Specialty palliative care teams can facilitate communication between clinicians and surrogate

decision makers, support frontline clinicians, and provide direct patient care services when needed.

Surrogate decision makers should be identified for all critically ill patients with COVID-19 at hospital admission. Infection-control policies for COVID-19 often create communication barriers for surrogate decision makers, and most surrogates will not be physically present when discussing treatment options with clinicians. Many decision-making discussions will occur via telecommunication.

Acknowledgments

The Surviving Sepsis Campaign (SSC), an initiative supported by SCCM and the European Society of Intensive Care Medicine, issued *Guidelines on the Management of Critically Ill Adults with Coronavirus Disease 2019* (COVID-19) in March 2020, and a revised version was published in March 2021.¹ The COVID-19 Treatment Guidelines Panel (the Panel) has based the recommendations in this section on the SSC COVID-19 guidelines with permission, and the Panel gratefully acknowledges the work of the SSC COVID-19 Guidelines Panel. The Panel also acknowledges the contributions and expertise of Andrew Rhodes, MBBS, MD, of St. George's University Hospitals in London, England, and Waleed Alhazzani, MBBS, MSc, of McMaster University in Hamilton, Canada.

References

- 1. Alhazzani W, Evans L, Alshamsi F, et al. Surviving Sepsis Campaign guidelines on the management of adults with coronavirus disease 2019 (COVID-19) in the ICU: first update. *Crit Care Med.* 2021;49(3):e219-e234. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33555780</u>.
- 2. Leisman DE, Ronner L, Pinotti R, et al. Cytokine elevation in severe and critical COVID-19: a rapid systematic review, meta-analysis, and comparison with other inflammatory syndromes. *Lancet Respir Med.* 2020;8(12):1233-1244. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33075298</u>.
- 3. Sinha P, Matthay MA, Calfee CS. Is a "cytokine storm" relevant to COVID-19? *JAMA Intern Med.* 2020;180(9):1152-1154. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32602883</u>.
- Morris SB, Schwartz NG, Patel P, et al. Case series of multisystem inflammatory syndrome in adults associated with SARS-CoV-2 infection—United Kingdom and United States, March-August 2020. MMWR Morb Mortal Wkly Rep. 2020;69(40):1450-1456. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33031361.
- Sansone M, Studahl M, Berg S, Gisslen M, Sundell N. Severe multisystem inflammatory syndrome (MIS-C/A) after confirmed SARS-CoV-2 infection: a report of four adult cases. *Infect Dis (Lond)*. 2022:1-6. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35034549</u>.
- Martins A, Policarpo S, Silva-Pinto A, et al. SARS-CoV-2-related multisystem inflammatory syndrome in adults. *Eur J Case Rep Intern Med*. 2021;8(11):003025. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34912751</u>.
- Belay ED, Godfred Cato S, Rao AK, et al. Multisystem inflammatory syndrome in adults after SARS-CoV-2 infection and COVID-19 vaccination. *Clin Infect Dis*. 2021. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34849680</u>.
- 8. Zou F, Qian Z, Wang Y, Zhao Y, Bai J. Cardiac injury and COVID-19: a systematic review and meta-analysis. *CJC Open*. 2020;2(5):386-394. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32838255</u>.
- Nishiga M, Wang DW, Han Y, Lewis DB, Wu JC. COVID-19 and cardiovascular disease: from basic mechanisms to clinical perspectives. *Nat Rev Cardiol*. 2020;17(9):543-558. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32690910</u>.
- Llitjos JF, Leclerc M, Chochois C, et al. High incidence of venous thromboembolic events in anticoagulated severe COVID-19 patients. *J Thromb Haemost*. 2020;18(7):1743-1746. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32320517</u>.

- 11. Helms J, Tacquard C, Severac F, et al. High risk of thrombosis in patients with severe SARS-CoV-2 infection: a multicenter prospective cohort study. *Intensive Care Med.* 2020;46(6):1089-1098. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32367170.
- 12. Klok FA, Kruip M, van der Meer NJM, et al. Incidence of thrombotic complications in critically ill ICU patients with COVID-19. *Thromb Res.* 2020;191:145-147. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32291094.
- Menter T, Haslbauer JD, Nienhold R, et al. Postmortem examination of COVID-19 patients reveals diffuse alveolar damage with severe capillary congestion and variegated findings in lungs and other organs suggesting vascular dysfunction. *Histopathology*. 2020;77(2):198-209. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32364264.
- 14. Tavazzi G, Civardi L, Caneva L, Mongodi S, Mojoli F. Thrombotic events in SARS-CoV-2 patients: an urgent call for ultrasound screening. *Intensive Care Med.* 2020;46(6):1121-1123. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32322918.
- Arentz M, Yim E, Klaff L, et al. Characteristics and outcomes of 21 critically ill patients with COVID-19 in Washington state. *JAMA*. 2020;323(16):1612-1614. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32191259</u>.
- Gupta S, Coca SG, Chan L, et al. AKI treated with renal replacement therapy in critically ill patients with COVID-19. *J Am Soc Nephrol.* 2021;32(1):161-176. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33067383</u>.
- Samaan F, Carneiro de Paula E, de Lima Souza FBG, et al. COVID-19-associated acute kidney injury patients treated with renal replacement therapy in the intensive care unit: a multicenter study in Sao Paulo, Brazil. *PLoS One*. 2022;17(1):e0261958. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35030179</u>.
- Helms J, Kremer S, Merdji H, et al. Neurologic features in severe SARS-CoV-2 infection. N Engl J Med. 2020;382(23):2268-2270. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32294339</u>.
- 19. Pun BT, Badenes R, Heras La Calle G, et al. Prevalence and risk factors for delirium in critically ill patients with COVID-19 (COVID-D): a multicentre cohort study. *Lancet Respir Med*. 2021;9(3):239-250. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33428871.
- 20. Mao L, Jin H, Wang M, et al. Neurologic manifestations of hospitalized patients with coronavirus disease 2019 in Wuhan, China. *JAMA Neurol*. 2020;77(6):683-690. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32275288</u>.
- 21. Solomon IH, Normandin E, Bhattacharyya S, et al. Neuropathological features of COVID-19. *N Engl J Med.* 2020;383(10):989-992. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32530583</u>.
- 22. Barr J, Fraser GL, Puntillo K, et al. Clinical practice guidelines for the management of pain, agitation, and delirium in adult patients in the intensive care unit. *Crit Care Med.* 2013;41(1):263-306. Available at: https://www.ncbi.nlm.nih.gov/pubmed/23269131.
- 23. Devlin JW, Skrobik Y, Gelinas C, et al. Clinical practice guidelines for the prevention and management of pain, agitation/sedation, delirium, immobility, and sleep disruption in adult patients in the ICU. *Crit Care Med.* 2018;46(9):e825-e873. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30113379</u>.
- Kress JP, Vinayak AG, Levitt J, et al. Daily sedative interruption in mechanically ventilated patients at risk for coronary artery disease. *Crit Care Med.* 2007;35(2):365-371. Available at: https://www.ncbi.nlm.nih.gov/pubmed/17205005.
- 25. Girard TD, Kress JP, Fuchs BD, et al. Efficacy and safety of a paired sedation and ventilator weaning protocol for mechanically ventilated patients in intensive care (Awakening and Breathing Controlled trial): a randomised controlled trial. *Lancet*. 2008;371(9607):126-134. Available at: https://www.ncbi.nlm.nih.gov/pubmed/18191684.
- 26. Society of Critical Care Medicine. ICU liberation bundle (A-F). Available at: <u>https://www.sccm.org/ICULiberation/ABCDEF-Bundles</u>. Accessed March 28, 2022.

- 27. Barnes-Daly MA, Phillips G, Ely EW. Improving hospital survival and reducing brain dysfunction at seven California community Hospitals: implementing PAD guidelines via the ABCDEF bundle in 6,064 patients. *Crit Care Med.* 2017;45(2):171-178. Available at: https://www.ncbi.nlm.nih.gov/pubmed/27861180.
- 28. Society of Critical Care Medicine. Post-intensive care syndrome. 2013. Available at: <u>https://www.sccm.org/MyICUCare/THRIVE/Post-intensive-Care-Syndrome</u>.
- 29. Fan E, Dowdy DW, Colantuoni E, et al. Physical complications in acute lung injury survivors: a two-year longitudinal prospective study. *Crit Care Med*. 2014;42(4):849-859. Available at: https://www.ncbi.nlm.nih.gov/pubmed/24247473.
- 30. De Jonghe B, Sharshar T, Lefaucheur JP, et al. Paresis acquired in the intensive care unit: a prospective multicenter study. *JAMA*. 2002;288(22):2859-2867. Available at: https://www.ncbi.nlm.nih.gov/pubmed/12472328.
- 31. Ali NA, O'Brien JM, Jr., Hoffmann SP, et al. Acquired weakness, handgrip strength, and mortality in critically ill patients. Am J Respir Crit Care Med. 2008;178(3):261-268. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/18511703</u>.
- 32. Pandharipande PP, Girard TD, Jackson JC, et al. Long-term cognitive impairment after critical illness. *N Engl J Med.* 2013;369(14):1306-1316. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/24088092</u>.
- 33. Iwashyna TJ, Ely EW, Smith DM, Langa KM. Long-term cognitive impairment and functional disability among survivors of severe sepsis. *JAMA*. 2010;304(16):1787-1794. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/20978258</u>.
- 34. Mikkelsen ME, Christie JD, Lanken PN, et al. The adult respiratory distress syndrome cognitive outcomes study: long-term neuropsychological function in survivors of acute lung injury. *Am J Respir Crit Care Med*. 2012;185(12):1307-1315. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/22492988</u>.
- 35. Kamdar BB, Sepulveda KA, Chong A, et al. Return to work and lost earnings after acute respiratory distress syndrome: a 5-year prospective, longitudinal study of long-term survivors. *Thorax*. 2018;73(2):125-133. Available at: https://www.ncbi.nlm.nih.gov/pubmed/28918401.
- 36. Azoulay E, Pochard F, Kentish-Barnes N, et al. Risk of post-traumatic stress symptoms in family members of intensive care unit patients. *Am J Respir Crit Care Med.* 2005;171(9):987-994. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/15665319</u>.
- 37. Carfi A, Bernabei R, Landi F, Gemelli Against C-P-ACSG. Persistent symptoms in patients after acute COVID-19. *JAMA*. 2020;324(6):603-605. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32644129</u>.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Hemodynamics for Adults

Last Updated: July 8, 2021

Most of the hemodynamic recommendations below are similar to those previously published in the *Surviving Sepsis Campaign: International Guidelines for Management of Sepsis and Septic Shock: 2016.* Ultimately, adult patients with COVID-19 who require fluid resuscitation or hemodynamic management of shock should be treated and managed identically to adult patients with septic shock.¹

Recommendation

• For adults with COVID-19 and shock, the COVID-19 Treatment Guidelines Panel (the Panel) recommends using dynamic parameters, skin temperature, capillary refilling time, and/or lactate levels over static parameters to assess fluid responsiveness (**BIIa**).

Rationale

In a systematic review and meta-analysis of 13 randomized clinical trials in intensive care unit (ICU) patients without COVID-19 (n = 1,652),² dynamic assessment to guide fluid therapy reduced mortality (risk ratio 0.59; 95% CI, 0.42–0.83), ICU length of stay (weighted mean difference -1.16 days; 95% CI, -1.97 to -0.36), and duration of mechanical ventilation (weighted mean difference -2.98 hours; 95% CI, -5.08 to -0.89). Dynamic parameters used in these trials included stroke volume variation (SVV), pulse pressure variation (PPV), and stroke volume change with passive leg raise or fluid challenge. Passive leg raising, followed by PPV and SVV, appears to predict fluid responsiveness with the greatest accuracy.³ The static parameters included components of early goal-directed therapy (e.g., central venous pressure, mean arterial pressure [MAP]).

Resuscitation of patients with shock who do not have COVID-19 based on serum lactate levels has been summarized in a systematic review and meta-analysis of 7 randomized clinical trials (n = 1,301). Compared with central venous oxygen saturation-guided therapy, early lactate clearance-directed therapy was associated with a reduction in mortality (relative ratio 0.68; 95% CI, 0.56–0.82), shorter ICU stay (mean difference -1.64 days; 95% CI, -3.23 to -0.05), and shorter duration of mechanical ventilation (mean difference -10.22 hours; 95% CI, -15.94 to -4.50).⁴

Recommendation

• For the acute resuscitation of adults with COVID-19 and shock, the Panel recommends using buffered/balanced crystalloids over unbalanced crystalloids (**BIIa**).

Rationale

A pragmatic randomized trial compared the use of balanced and unbalanced crystalloids for intravenous (IV) fluid administration in critically ill adults without COVID-19 (n = 15,802). The rate of the composite outcome of death, new renal-replacement therapy, or persistent renal dysfunction was lower in the balanced crystalloids group than in the unbalanced crystalloids group (OR 0.90; 95% CI, 0.82-0.99; P = 0.04).⁵ A secondary analysis compared outcomes in a subset of patients with sepsis (n = 1,641). Compared to treatment with unbalanced crystalloids, treatment with balanced crystalloids resulted in fewer deaths (aOR 0.74; 95% CI, 0.59-0.93; P = 0.01) and more vasopressor-free and renal replacement-free days.⁶ A subsequent meta-analysis of 21 non-COVID-19 randomized controlled trials (n = 20,213) that included the pragmatic trial cited above compared balanced crystalloids to 0.9% saline

for resuscitation of critically ill adults and children. The trial reported nonsignificant differences between the treatment groups in hospital mortality (OR 0.91; 95% CI, 0.83–1.01) and acute kidney injury (OR 0.92; 95% CI, 0.84–1.00).⁷

Recommendation

• For the acute resuscitation of adults with COVID-19 and shock, the Panel **recommends against** the initial use of **albumin** for resuscitation (**BI**).

Rationale

A meta-analysis of 20 non-COVID-19 randomized controlled trials (n = 13,047) that compared the use of albumin or fresh-frozen plasma to crystalloids in critically ill patients found no difference in all-cause mortality between the treatment groups.⁸ In contrast, a meta-analysis of 17 non-COVID-19 randomized controlled trials (n = 1,977) that compared the use of albumin to crystalloids specifically in patients with sepsis observed a reduction in mortality among the patients who received albumin (OR 0.82; 95% CI, 0.67–1.0; P = 0.047).⁹ Given the higher cost of albumin and the lack of a definitive clinical benefit, the Panel **recommends against** the routine use of **albumin** for initial acute resuscitation of patients with COVID-19 and shock (**BI**).

Recommendation

• For adults with COVID-19 and shock, the Panel recommends **norepinephrine** as the first-choice vasopressor (**AI**).

Rationale

Norepinephrine increases MAP due to its vasoconstrictive effects, with little change in heart rate and less increase in stroke volume compared to dopamine. Dopamine increases MAP and cardiac output, primarily due to an increase in stroke volume and heart rate. Norepinephrine is more potent than dopamine and may be more effective at reversing hypotension in patients with septic shock. Dopamine may be particularly useful in patients with compromised systolic function, but it causes more tachycardia and may be more arrhythmogenic than norepinephrine.¹⁰ It may also influence the endocrine response via the hypothalamic pituitary axis and have immunosuppressive effects.¹¹ A systematic review and meta-analysis of 11, non-COVID-19 randomized controlled trials that compared vasopressors used to treat patients with septic shock found that norepinephrine use resulted in lower all-cause mortality (risk ratio 0.89; 95% CI, 0.81–0.98) and a lower risk of arrhythmias (risk ratio 0.48; 95% CI, 0.40–0.58) than dopamine use.¹² Although the beta-1 activity of dopamine would be useful in patients with myocardial dysfunction, the greater risk of arrhythmias limits its use.^{13,14}

Recommendation

• For adults with COVID-19 and shock, the Panel recommends titrating vasoactive agents to target a MAP of 60 to 65 mm Hg, over higher MAP targets (**BI**).

Rationale

A recent individual patient-data meta-analysis of 2, non-COVID-19 randomized controlled trials (n = 894) comparing higher versus lower blood pressure targets for vasopressor therapy in adult patients with shock reported no significant difference between the patients in the higher and lower target groups in 28-day mortality (OR 1.15; 95% CI, 0.87–1.52), 90-day mortality (OR 1.08; 95% CI, 0.84–1.44), myocardial injury (OR 1.47; 95% CI, 0.64–3.56), or limb ischemia (OR 0.92; 95% CI, 0.36–2.10).¹⁵ The risk of arrhythmias was increased in patients allocated to the higher target group (OR 2.50; 95%

CI, 1.35–4.77). Similarly, the recently published "65 Trial," a randomized clinical trial in patients without COVID-19 (n = 2,463), reported no significant difference in mortality between patients with vasopressor therapy guided by a MAP target of 60 to 65 mm Hg and those with treatment guided by a higher, standard of care MAP target (41% vs. 43.8%; RR 0.93; 95% CI, 0.85–1.03).¹⁶ With an indication of improved outcome with lower MAP targets (and no firm indication of harm), the Panel recommends titrating vasoactive agents to a MAP target of 60 to 65 mm Hg (**BI**).

Additional Recommendations for Adults With COVID-19 and Shock Based on General Principles of Critical Care

- The Panel **recommends against** using hydroxyethyl starches for intravascular volume replacement in adult patients with COVID-19 and sepsis or septic shock (**AI**).
- When norepinephrine is available, the Panel **recommends against** using **dopamine** for adult patients with COVID-19 and shock (AI).
- As a second line vasopressor, the Panel recommends adding either **vasopressin** (up to 0.03 units/ min) (**BIIa**) or **epinephrine** (**BIIb**) to norepinephrine to raise MAP to target or adding vasopressin (up to 0.03 units/min) (**BIIa**) to decrease norepinephrine dosage.
- The Panel recommends against using low-dose dopamine for renal protection (AI).
- The Panel recommends using **dobutamine** in adult patients with COVID-19 who show evidence of cardiac dysfunction and persistent hypoperfusion despite adequate fluid loading and the use of vasopressor agents (**BIII**).
- The Panel recommends that all adult patients with COVID-19 who require vasopressors have an arterial catheter placed as soon as practical, if resources are available (**BIII**).
- For adult patients with refractory septic shock who have completed a course of corticosteroids to treat COVID-19, the Panel recommends using low-dose corticosteroid therapy ("shock-reversal") over no corticosteroid therapy (**BIIa**).
 - A typical corticosteroid regimen in septic shock is hydrocortisone 200 mg IV per day administered either as an infusion or in intermittent doses. The duration of hydrocortisone therapy is usually a clinical decision.
 - Adult patients who are receiving corticosteroids for COVID-19 are receiving sufficient replacement therapy such that they do not require additional hydrocortisone.

References

- Rhodes A, Evans LE, Alhazzani W, et al. Surviving Sepsis Campaign: international guidelines for management of sepsis and septic shock: 2016. *Crit Care Med.* 2017;45(3):486-552. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28098591</u>.
- 2. Bednarczyk JM, Fridfinnson JA, Kumar A, et al. Incorporating dynamic assessment of fluid responsiveness into goal-directed therapy: a systematic review and meta-analysis. *Crit Care Med.* 2017;45(9):1538-1545. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28817481</u>.
- 3. Bentzer P, Griesdale DE, Boyd J, MacLean K, Sirounis D, Ayas NT. Will this hemodynamically unstable patient respond to a bolus of intravenous fluids? *JAMA*. 2016;316(12):1298-1309. Available at: https://www.ncbi.nlm.nih.gov/pubmed/27673307.
- 4. Pan J, Peng M, Liao C, Hu X, Wang A, Li X. Relative efficacy and safety of early lactate clearance-guided therapy resuscitation in patients with sepsis: a meta-analysis. *Medicine (Baltimore)*. 2019;98(8):e14453. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30813144</u>.
- Semler MW, Self WH, Wanderer JP, et al. Balanced crystalloids versus saline in critically ill adults. *N Engl J Med*. 2018;378(9):829-839. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29485925</u>.

- Brown RM, Wang L, Coston TD, et al. Balanced crystalloids versus saline in sepsis. A secondary analysis of the SMART clinical trial. *Am J Respir Crit Care Med.* 2019;200(12):1487-1495. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31454263</u>.
- 7. Antequera Martin AM, Barea Mendoza JA, Muriel A, et al. Buffered solutions versus 0.9% saline for resuscitation in critically ill adults and children. *Cochrane Database Syst Rev.* 2019;7:CD012247. Available at: https://www.ncbi.nlm.nih.gov/pubmed/31334842.
- Lewis SR, Pritchard MW, Evans DJ, et al. Colloids versus crystalloids for fluid resuscitation in critically ill people. *Cochrane Database Syst Rev.* 2018;8:CD000567. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30073665</u>.
- 9. Delaney AP, Dan A, McCaffrey J, Finfer S. The role of albumin as a resuscitation fluid for patients with sepsis: a systematic review and meta-analysis. *Crit Care Med.* 2011;39(2):386-391. Available at: https://www.ncbi.nlm.nih.gov/pubmed/21248514.
- 10. Regnier B, Rapin M, Gory G, Lemaire F, Teisseire B, Harari A. Haemodynamic effects of dopamine in septic shock. *Intensive Care Med.* 1977;3(2):47-53. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/893773</u>.
- 11. Beck G, Brinkkoetter P, Hanusch C, et al. Clinical review: immunomodulatory effects of dopamine in general inflammation. *Crit Care*. 2004;8(6):485-491. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/15566620</u>.
- Avni T, Lador A, Lev S, Leibovici L, Paul M, Grossman A. Vasopressors for the treatment of septic shock: systematic review and meta-analysis. *PLoS One*. 2015;10(8):e0129305. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26237037</u>.
- Regnier B, Safran D, Carlet J, Teisseire B. Comparative haemodynamic effects of dopamine and dobutamine in septic shock. *Intensive Care Med.* 1979;5(3):115-120. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/500939</u>.
- De Backer D, Creteur J, Silva E, Vincent JL. Effects of dopamine, norepinephrine, and epinephrine on the splanchnic circulation in septic shock: which is best? *Crit Care Med.* 2003;31(6):1659-1667. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/12794401</u>.
- 15. Lamontagne F, Day AG, Meade MO, et al. Pooled analysis of higher versus lower blood pressure targets for vasopressor therapy septic and vasodilatory shock. *Intensive Care Med.* 2018;44(1):12-21. Available at: https://www.ncbi.nlm.nih.gov/pubmed/29260272.
- 16. Lamontagne F, Richards-Belle A, Thomas K, et al. Effect of reduced exposure to vasopressors on 90-day mortality in older critically ill patients with vasodilatory hypotension: a randomized clinical trial. *JAMA*. 2020;323(10):938-949. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32049269</u>.

Oxygenation and Ventilation for Adults

Last Updated: December 20, 2023

The COVID-19 Treatment Guidelines Panel's (the Panel) recommendations in this section were informed by the Surviving Sepsis Campaign <u>guidelines for managing sepsis</u> and <u>guidelines for managing</u> <u>COVID-19</u> in adults.

Severe illness in people with COVID-19 typically occurs approximately 1 week after the onset of symptoms. The most common symptom is dyspnea, which is often accompanied by hypoxemia. Patients with severe disease typically require supplemental oxygen and should be monitored closely for worsening respiratory status, because some patients may progress to acute respiratory distress syndrome (ARDS).

Goal of Oxygenation

The optimal oxygen saturation measured by pulse oximetry (SpO_2) in adults with COVID-19 who are receiving supplemental oxygen is unknown. However, a target SpO_2 of 92% to 96% seems logical, considering that indirect evidence from patients without COVID-19 suggests that an $SpO_2 < 92\%$ or >96% may be harmful.^{1,2} Special care should be taken when assessing SpO_2 in patients with darker skin pigmentation, as recent reports indicate that occult hypoxemia (defined as arterial oxygen saturation $[SaO_2] < 88\%$ despite an $SpO_2 > 92\%$) is more common in these patients.^{3,4} See <u>Clinical Spectrum of SARS-CoV-2 Infection</u> for more information.

The potential harm of maintaining an SpO₂ <92% was demonstrated during a trial that randomly assigned patients with ARDS who did not have COVID-19 to either a conservative oxygen strategy (target SpO₂ of 88% to 92%) or a liberal oxygen strategy (target SpO₂ \geq 96%).¹ The trial was stopped early due to futility after enrolling 205 patients, but increased mortality was observed at Day 90 in the conservative oxygen strategy arm (between-group risk difference 14%; 95% CI, 0.7% to 27%), and a trend toward increased mortality was observed at Day 28 (between-group risk difference 8%; 95% CI, -5% to 21%).

The results of a meta-analysis of 25 randomized trials that involved patients without COVID-19 demonstrated the potential harm of maintaining an SpO₂ >96%.² This study found that a liberal oxygen supplementation strategy (a median fraction of inspired oxygen [FiO₂] of 0.52) was associated with an increased risk of in-hospital mortality (relative risk 1.21; 95% CI, 1.03–1.43) when compared with a more conservative SpO₂ supplementation strategy (a median FiO₂ of 0.21).

Acute Hypoxemic Respiratory Failure

In adults with COVID-19 and acute hypoxemic respiratory failure, conventional oxygen therapy may be insufficient to meet the oxygen needs of the patient. Options for providing enhanced respiratory support include using high-flow nasal canula (HFNC) oxygen, noninvasive ventilation (NIV), intubation and mechanical ventilation, or extracorporeal membrane oxygenation. In this section, mechanical ventilation refers to the delivery of positive pressure ventilation through an endotracheal or tracheostomy tube. NIV refers to the delivery of either continuous positive airway pressure (CPAP) or bilevel positive airway pressure (e.g., BiPAP) through a noninvasive interface, such as a face mask or nasal mask.

Nonmechanically Ventilated Adults With Acute Hypoxemic Respiratory Failure

High-Flow Nasal Cannula Oxygen and Noninvasive Ventilation

Recommendations

• For adults with COVID-19 and acute hypoxemic respiratory failure despite conventional oxygen

COVID-19 Treatment Guidelines

therapy, the Panel recommends starting therapy with HFNC oxygen; if patients fail to respond, NIV or intubation and mechanical ventilation should be initiated (**BIIa**).

• For adults with COVID-19 and acute hypoxemic respiratory failure despite conventional oxygen therapy who do not have an indication for endotracheal intubation and for whom HFNC oxygen is not available, the Panel recommends performing a closely monitored trial of NIV (**BIIa**).

Rationale

Several studies have informed clinical practice on the optimal oxygen delivery system for patients with COVID-19 and acute hypoxemic respiratory failure. A randomized study of 711 patients with COVID-19 in 34 intensive care units (ICUs) in France compared HFNC oxygen delivery to oxygen delivery through a nonrebreather mask.⁵ The patients had acute respiratory failure with a ratio of arterial partial pressure of oxygen to fraction of inspired oxygen (PaO₂/FiO₂) \leq 200 mm Hg. The mean FiO₂ was 0.58 in both arms. Although the difference between arms for the primary endpoint of 28-day mortality was not statistically significant (10% in the HFNC oxygen arm vs. 11% in the conventional oxygen arm; absolute difference -1.2%; 95% CI, -5.8% to 3.4%; *P* = 0.60), the intubation rate was significantly lower in the HFNC oxygen arm than in the conventional oxygen arm. Unless a contraindication exists, most Panel members would switch to HFNC oxygen delivery for patients with respiratory failure who do not require mechanical ventilation but have worsening hypoxemia or increased work of breathing despite receiving conventional oxygen at flow rates up to 10 L/min.

For patients with COVID-19 and acute hypoxemic respiratory failure who do not respond to conventional oxygen therapy, HFNC oxygen is preferred over NIV. No studies directly compare HFNC oxygen with mask-delivered NIV in patients with COVID-19; therefore, this guidance is based on an unblinded clinical trial in patients without COVID-19 who had acute hypoxemic respiratory failure.⁶ Study participants were randomized to receive HFNC oxygen, conventional oxygen therapy, or NIV. The patients in the HFNC oxygen arm had more ventilator-free days (mean 24 days) than those in the conventional oxygen therapy arm (mean 22 days) or the NIV arm (mean 19 days; P = 0.02). In addition, the conventional oxygen therapy arm (HR 2.01; 95% CI, 1.01–3.99) and the NIV arm (HR 2.50; 95% CI, 1.31–4.78) had higher 90-day mortality than the HFNC oxygen arm. In the subgroup of patients with severe hypoxemia (those with PaO₂/FiO₂ ≤200 mm Hg), the HFNC oxygen arm had a lower intubation rate than the conventional oxygen therapy arm (HR 2.07) and the NIV arm (HR 2.57).

The trial's findings were corroborated by a meta-analysis of 8 trials with 1,084 participants that assessed the effectiveness of oxygenation strategies.⁷ Compared to NIV, HFNC oxygen reduced the rate of intubation (OR 0.48; 95% CI, 0.31–0.73) and ICU mortality (OR 0.36; 95% CI, 0.20–0.63).

One small study compared the use of NIV delivered by a helmet device to HFNC oxygen in patients with COVID-19. The HENIVOT trial randomized 109 patients with moderate to severe COVID-19 (defined as those who had $PaO_2/FiO_2 \leq 200 \text{ mm Hg}$) to receive either NIV via a helmet device or HFNC oxygen.⁸ The study found no difference between the arms for the primary outcome of respiratory support–free days. However, only 30% of patients in the NIV arm required endotracheal intubation compared to 51% of patients in the HFNC oxygen arm (P = 0.03).

Two larger studies compared the use of NIV with conventional oxygen therapy in patients with COVID-19. The RECOVERY-RS trial was an adaptive randomized controlled trial that was essentially conducted as 2 separate trials that compared NIV and HFNC oxygen to the same conventional oxygen therapy control group.⁹ The trial was stopped early and enrolled fewer than a third of the planned sample size of 4,002 participants. Between April 2020 and May 2021, 1,273 adults with acute hypoxemic respiratory failure related to COVID-19 were randomized to receive NIV (n = 380), HFNC oxygen (n = 418), or conventional oxygen therapy (n = 475). The primary endpoint was a composite of endotracheal intubation or death within 30 days. The proportion of patients who met the primary endpoint was

significantly lower in the NIV arm than in the conventional oxygen therapy arm (36.3% vs. 44.4%; P = 0.03). This difference was not due to mortality but was entirely due to a reduction in the number of patients who required intubation. There was no significant difference between the HFNC oxygen arm and the conventional oxygen therapy arm in the occurrence of the primary endpoint (44.3% vs. 45.1%; P = 0.83).

There was substantial crossover between the arms, but an inverse probability weighting analysis that corrected for the bias this may have introduced did not change the results.⁹ Adverse events were more common in the NIV arm. Initially, a comparison between NIV and HFNC oxygen was not planned, but a post hoc analysis found that the proportion of patients who required endotracheal intubation or who died was lower in the NIV arm than in the HFNC oxygen arm (34.6% vs. 44.3%; P = 0.02).

In contrast to the RECOVERY-RS trial, the HiFlo-COVID trial randomized 220 patients with COVID-19 to receive HFNC oxygen or conventional oxygen therapy and found that a smaller proportion of patients in the HFNC oxygen arm required intubation (34.3% vs. 51.0%; P = 0.03).¹⁰ Patients in the HFNC arm also had a shorter median time to recovery (11 vs. 14 days; P = 0.047).

The conflicting results of these studies make drawing inferences from the data difficult. Additionally, the RECOVERY-RS trial was stopped long before it reached its planned sample size for reasons not related to futility, efficacy, or harm; inferring benefit in this context is questionable. The Panel recognizes that for patients who need more oxygen support than a conventional nasal cannula can provide, most clinicians will administer oxygen via HFNC and subsequently progress to NIV if needed. Therefore, the pertinent clinical question is whether HFNC oxygen or NIV should be used when a patient does not respond to conventional oxygen therapy. Other than the post hoc analysis in the RECOVERY-RS trial, no study has specifically investigated this question.

NIV is an aerosol-generating procedure, and studies of SARS-CoV show that it may increase the risk of nosocomial transmission.^{11,12} For patients with SARS-CoV-2, it remains unclear whether the use of HFNC oxygen results in a lower risk of nosocomial transmission than the use of NIV.

Awake Prone Positioning in Nonmechanically Ventilated Adults

Recommendations

- For adults with persistent hypoxemia who require HFNC oxygen and for whom endotracheal intubation is not indicated, the Panel recommends a trial of awake prone positioning (**BIIa**).
- The Panel **recommends against** the use of awake prone positioning as a rescue therapy for refractory hypoxemia to avoid intubation in patients who otherwise meet the indications for intubation and mechanical ventilation (**AIII**).

Additional Considerations

- Patients who can adjust their position independently and tolerate lying prone can be considered for awake prone positioning.
- Awake prone positioning is acceptable and feasible for pregnant patients and can be performed in the left lateral decubitus position or the fully prone position.¹³
- Some patients do not tolerate awake prone positioning. Failure rates as high as 63% have been reported in the literature.¹⁴
- Awake prone positioning **should not be used** as a substitute for intubation and mechanical ventilation in patients with refractory hypoxemia who otherwise meet the indications for these interventions.
- Awake prone positioning may be infeasible or impractical in patients with:

- Spinal instability
- Facial or pelvic fractures
- An open chest or unstable chest wall
- Awake prone positioning should be used with caution in patients with confusion, delirium, or hemodynamic instability; patients who cannot independently change position; or patients who have had recent abdominal surgery, nausea, or vomiting.

Rationale

Awake prone positioning, or having a nonintubated patient lie on the stomach, may improve oxygenation and prevent the patient from progressing to requiring intubation and mechanical ventilation. Although prone positioning has been shown to improve oxygenation and outcomes in patients with moderate to severe ARDS who are receiving mechanical ventilation,^{15,16} there is less evidence regarding the benefit of prone positioning in awake patients who require supplemental oxygen without mechanical ventilation. Several case series of patients with COVID-19 who required oxygen or NIV have reported that awake prone positioning improved oxygenation,¹⁷⁻²⁰ and some series have also reported low intubation rates after awake prone positioning.^{19,20}

The Awake Prone Positioning Meta-Trial Group has conducted the largest trial on awake prone positioning.²¹ This study was a prospective, multinational meta-trial of 6 open-label, randomized, controlled, superiority trials that compared awake prone positioning to standard care in adults who required HFNC oxygen for acute hypoxemic respiratory failure due to COVID-19.

The study enrolled 1,126 patients between April 2, 2020, and January 26, 2021, and the intention-to-treat analysis included 1,121 patients.²¹ Of the 564 patients who underwent awake prone positioning, 223 (40%) met the composite primary endpoint of intubation or death within 28 days of enrollment. Among the 557 patients who received standard care, 257 (46%) met the primary endpoint (relative risk 0.86; 95% CI, 0.75–0.98). The incidence of intubation by Day 28 was lower in the awake prone positioning arm than in the standard care arm (HR intubation 0.75; 95% CI, 0.62–0.91). There was no difference in 28-day mortality between the awake prone positioning arm and the standard care arm (HR mortality 0.87; 95% CI, 0.68–1.11).

During the first 14 days of the study, the median daily duration of awake prone positioning was 5.0 hours (IQR 1.6–8.8 hours).²¹ However, the median daily duration varied from 1.6 hours to 8.6 hours across the individual trials. Longer daily durations for awake prone positioning were associated with treatment success by Day 28. This study evaluated the incidences of certain adverse events, including skin breakdown, vomiting, and central or arterial line dislodgment. These events occurred infrequently during the study, and the incidences were similar in each arm. No cardiac arrests occurred during awake prone positioning.

The optimal daily duration of awake prone positioning is unclear. In the meta-trial of awake prone positioning, only 25 of 151 patients (17%) who had an average of \geq 8 hours of awake prone positioning per day met the primary endpoint of intubation or death when compared with 198 of 413 patients (48%) who remained in awake prone positioning for <8 hours per day.²¹ This result is consistent with past clinical trials of prone positioning in mechanically ventilated patients with ARDS, in which clinical benefits were observed after longer durations of prone positioning.^{15,16}

Intubation for Mechanical Ventilation

Recommendation

• If intubation becomes necessary, the procedure should be performed by an experienced

COVID-19 Treatment Guidelines

practitioner in a controlled setting due to the enhanced risk of exposing health care practitioners to SARS-CoV-2 during intubation (AIII).

Rationale

It is essential to closely monitor hypoxemic patients with COVID-19 for signs of respiratory decompensation. To ensure the safety of both patients and health care workers, intubation should be performed in a controlled setting by an experienced practitioner.

Mechanically Ventilated Adults

General Considerations

Recommendations

For mechanically ventilated adults with COVID-19 and ARDS:

- The Panel recommends using low tidal volume (VT) ventilation (VT 4–8 mL/kg of predicted body weight) over higher VT ventilation (VT >8 mL/kg) (AI).
- The Panel recommends targeting plateau pressures of <30 cm H₂O (AIIa).
- The Panel recommends using a conservative fluid strategy over a liberal fluid strategy (BIIa).
- The Panel recommends against the routine use of inhaled nitric oxide (AIIa).

Rationale

There is no evidence that the ventilator management of patients with hypoxemic respiratory failure due to COVID-19 should differ from the ventilator management of patients with hypoxemic respiratory failure due to other causes.

Positive End-Expiratory Pressure and Prone Positioning in Mechanically Ventilated Adults With Moderate to Severe ARDS

Recommendations

For mechanically ventilated adults with COVID-19 and moderate to severe ARDS:

- The Panel recommends using a higher positive end-expiratory pressure (PEEP) strategy over a lower PEEP strategy (**BIIa**).
- For mechanically ventilated adults with COVID-19 and refractory hypoxemia despite optimized ventilation, the Panel recommends prone ventilation for 12 to 16 hours per day over no prone ventilation (**BIIa**).

Rationale

PEEP is beneficial in patients with ARDS because it prevents alveolar collapse, improves oxygenation, and minimizes atelectrauma, a source of ventilator-induced lung injury. A meta-analysis of individual patient data from the 3 largest trials that compared lower and higher levels of PEEP in patients without COVID-19 found that less ICU mortality and in-hospital mortality was associated with higher levels of PEEP in those with moderate (PaO_2/FiO_2 100–200 mm Hg) and severe ($PaO_2/FiO_2 < 100$ mm Hg) ARDS.²²

Although there is no clear standard for a high level of PEEP, a conventional threshold is >10 cm H_2O .²³ Recent reports have suggested that, in contrast to patients with ARDS not caused by COVID-19, some patients with moderate or severe ARDS due to COVID-19 have normal static lung compliance. In these patients, high levels of PEEP may cause harm by compromising hemodynamics and cardiovascular performance.^{24,25} Other studies have reported that patients with moderate to severe ARDS due to

COVID-19 had low lung compliance, similar to the lung compliance seen in patients with conventional ARDS.²⁶⁻²⁹ These seemingly contradictory observations suggest that patients with COVID-19 and ARDS are a heterogeneous population, and assessments for responsiveness to high levels of PEEP should be individualized based on oxygenation and lung compliance. Clinicians should monitor patients for known side effects of high levels of PEEP, such as barotrauma and hypotension.

In the prepandemic PROSEVA study of patients with moderate to severe early ARDS ($PaO_2/FiO_2 < 150$ mm Hg) who required mechanical ventilation, the patients who were randomized to undergo prone positioning for ≥ 16 hours per day had improved survival compared to those who remained in the supine position throughout the course of mechanical ventilation.¹⁵ A meta-analysis evaluated the results of the PROSEVA study and 7 other randomized controlled trials that investigated the use of prone positioning in people with ARDS.³⁰ A subgroup analysis revealed that mortality was reduced among patients who remained prone for ≥ 12 hours per day when compared with patients who remained in the supine position (risk ratio 0.74; 95% CI, 0.56–0.99). Prone positioning improved oxygenation in all the trials. Patients in the prone positioning arms had higher PaO_2/FiO_2 on Day 4 than those in the supine positioning arms (mean difference 23.5 mm Hg; 95% CI, 12.4–34.5).

The use of prone positioning may be associated with serious adverse events, including unplanned extubation or central catheter removal. However, the meta-analysis found no differences between the prone positioning and supine positioning arms in the frequency of these events.³⁰ The use of prone positioning was associated with an increased risk of pressure sores (risk ratio 1.22; 95% CI, 1.06–1.41) and endotracheal tube obstruction (risk ratio 1.76; 95% CI, 1.24–2.50) in the 3 studies that evaluated these complications.

Neuromuscular Blockade in Mechanically Ventilated Adults With Moderate to Severe ARDS Recommendation

For mechanically ventilated adults with COVID-19 and moderate to severe ARDS:

• The Panel recommends using, as needed, intermittent boluses of neuromuscular blocking agents (NMBAs) or a continuous NMBA infusion to facilitate protective lung ventilation (**BIIa**).

Rationale

Although the use of NMBAs in patients with ARDS reduces ventilator dyssynchrony, a large multicenter trial across several ICUs reported no significant difference in mortality between patients who received deep sedation and continuous NMBA infusion and patients who received a usual-care approach of lighter sedation without routine NMBAs.³¹

Rescue Therapies for Mechanically Ventilated Adults With ARDS

Recommendations

For mechanically ventilated adults with COVID-19, severe ARDS, and hypoxemia despite optimized ventilation and other rescue strategies:

- The Panel recommends using an inhaled pulmonary vasodilator as a rescue therapy; if rapid improvement in oxygenation is not observed, the treatment should be tapered (**CIII**).
- The Panel recommends using recruitment maneuvers rather than not using recruitment maneuvers (CIIa).
- If recruitment maneuvers are used, the Panel **recommends against** the use of staircase (incremental PEEP) recruitment maneuvers (**AIIa**).

Rationale

A recruitment maneuver refers to a temporary increase in airway pressure during mechanical ventilation to open collapsed alveoli and improve oxygenation. No studies have assessed the effect of recruitment maneuvers on oxygenation in patients with severe ARDS due to COVID-19. However, a systematic review and meta-analysis of 6 trials of recruitment maneuvers in patients with ARDS who did not have COVID-19 found that recruitment maneuvers reduced mortality, improved oxygenation 24 hours after the maneuver, and decreased the need for rescue therapy.³² Because recruitment maneuvers can cause barotrauma or hypotension, patients should be closely monitored during the maneuvers. If a patient decompensates during recruitment maneuvers, the maneuvers should be stopped immediately.

The importance of properly performing recruitment maneuvers was illustrated by an analysis of 8 randomized controlled trials in patients without COVID-19 (n = 2,544) that found that recruitment maneuvers did not reduce in-hospital mortality (risk ratio 0.90; 95% CI, 0.78–1.04).²³ However, a subgroup analysis found that traditional recruitment maneuvers significantly reduced in-hospital mortality (risk ratio 0.85; 95% CI, 0.75–0.97). Mortality was higher among patients treated with incremental PEEP titration recruitment maneuvers than among those treated with traditional recruitment maneuvers, but this difference was not statistically significant (risk ratio 1.06; 95% CI, 0.97–1.17).

There are no prospective trials of pulmonary vasodilators in people with COVID-19. However, a metaanalysis of mostly small, retrospective trials did not show improved outcomes.³³ A Cochrane review of 13 trials evaluated the use of inhaled nitric oxide in patients with ARDS who did not have COVID-19 and found no reduction in mortality.³⁴ Because the review showed a transient benefit for oxygenation, it is reasonable to attempt using inhaled nitric oxide as a rescue therapy in patients with COVID-19 and severe ARDS after other options have failed. However, if the use of nitric oxide does not improve a patient's oxygenation, it should be tapered quickly to avoid rebound pulmonary vasoconstriction, which may occur when nitric oxide is discontinued after prolonged use.

References

- Barrot L, Asfar P, Mauny F, et al. Liberal or conservative oxygen therapy for acute respiratory distress syndrome. *N Engl J Med*. 2020;382(11):999-1008. Available at: <u>https://www.ncbi.nlm.nih.gov/</u> pubmed/32160661.
- Chu DK, Kim LH, Young PJ, et al. Mortality and morbidity in acutely ill adults treated with liberal versus conservative oxygen therapy (IOTA): a systematic review and meta-analysis. *Lancet.* 2018;391(10131):1693-1705. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29726345</u>.
- Chesley CF, Lane-Fall MB, Panchanadam V, et al. Racial disparities in occult hypoxemia and clinically based mitigation strategies to apply in advance of technological advancements. *Respir Care*. 2022;67(12):1499-1507. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35679133</u>.
- Valbuena VSM, Seelye S, Sjoding MW, et al. Racial bias and reproducibility in pulse oximetry among medical and surgical inpatients in general care in the Veterans Health Administration 2013–19: multicenter, retrospective cohort study. *BMJ*. 2022;378:e069775. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35793817</u>.
- 5. Frat JP, Quenot JP, Badie J, et al. Effect of high-flow nasal cannula oxygen vs standard oxygen therapy on mortality in patients with respiratory failure due to COVID-19: the SOHO-COVID randomized clinical trial. *JAMA*. 2022;328(12):1212-1222. Available at: https://pubmed.ncbi.nlm.nih.gov/36166027.
- Frat JP, Thille AW, Mercat A, et al. High-flow oxygen through nasal cannula in acute hypoxemic respiratory failure. *N Engl J Med*. 2015;372(23):2185-2196. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/25981908</u>.
- 7. Ni YN, Luo J, Yu H, et al. The effect of high-flow nasal cannula in reducing the mortality and the rate of endotracheal intubation when used before mechanical ventilation compared with conventional oxygen therapy

and noninvasive positive pressure ventilation: a systematic review and meta-analysis. *Am J Emerg Med.* 2018;36(2):226-233. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28780231</u>.

- 8. Grieco DL, Menga LS, Cesarano M, et al. Effect of helmet noninvasive ventilation vs high-flow nasal oxygen on days free of respiratory support in patients with COVID-19 and moderate to severe hypoxemic respiratory failure: the HENIVOT randomized clinical trial. *JAMA*. 2021;325(17):1731-1743. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33764378.
- Perkins GD, Ji C, Connolly BA, et al. Effect of noninvasive respiratory strategies on intubation or mortality among patients with acute hypoxemic respiratory failure and COVID-19: the RECOVERY-RS randomized clinical trial. *JAMA*. 2022;327(6):546-558. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35072713</u>.
- Ospina-Tascón GA, Calderón-Tapia LE, García AF, et al. Effect of high-flow oxygen therapy vs conventional oxygen therapy on invasive mechanical ventilation and clinical recovery in patients with severe COVID-19: a randomized clinical trial. *JAMA*. 2021;326(21):2161-2171. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34874419.
- 11. Yu IT, Xie ZH, Tsoi KK, et al. Why did outbreaks of severe acute respiratory syndrome occur in some hospital wards but not in others? *Clin Infect Dis.* 2007;44(8):1017-1025. Available at: https://www.ncbi.nlm.nih.gov/pubmed/17366443.
- Tran K, Cimon K, Severn M, Pessoa-Silva CL, Conly J. Aerosol generating procedures and risk of transmission of acute respiratory infections to healthcare workers: a systematic review. *PLoS One*. 2012;7(4):e35797. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/22563403</u>.
- 13. Society for Maternal-Fetal Medicine. Management considerations for pregnant patients with COVID-19. 2021. Available at: <u>https://s3.amazonaws.com/cdn.smfm.org/media/2734/SMFM_COVID_Management_of_COVID_pos_preg_patients_2-2-21_(final).pdf</u>.
- Hallifax RJ, Porter BM, Elder PJ, et al. Successful awake proning is associated with improved clinical outcomes in patients with COVID-19: single-centre high-dependency unit experience. *BMJ Open Respir Res.* 2020;7(1):e000678. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32928787</u>.
- 15. Guérin C, Reignier J, Richard JC, et al. Prone positioning in severe acute respiratory distress syndrome. *N Engl J Med.* 2013;368(23):2159-2168. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/23688302</u>.
- 16. Fan E, Del Sorbo L, Goligher EC, et al. An official American Thoracic Society/European Society of Intensive Care Medicine/Society of Critical Care Medicine clinical practice guideline: mechanical ventilation in adult patients with acute respiratory distress syndrome. *Am J Respir Crit Care Med.* 2017;195(9):1253-1263. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28459336</u>.
- Caputo ND, Strayer RJ, Levitan R. Early self-proning in awake, non-intubated patients in the emergency department: a single ED's experience during the COVID-19 pandemic. *Acad Emerg Med.* 2020;27(5):375-378. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32320506</u>.
- 18. Elharrar X, Trigui Y, Dols AM, et al. Use of prone positioning in nonintubated patients with COVID-19 and hypoxemic acute respiratory failure. *JAMA*. 2020;323(22):2336-2338. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32412581.
- Sartini C, Tresoldi M, Scarpellini P, et al. Respiratory parameters in patients with COVID-19 after using noninvasive ventilation in the prone position outside the intensive care unit. *JAMA*. 2020;323(22):2338-2340. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32412606</u>.
- 20. Sun Q, Qiu H, Huang M, Yang Y. Lower mortality of COVID-19 by early recognition and intervention: experience from Jiangsu province. *Ann Intensive Care*. 2020;10(1):33. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32189136</u>.
- 21. Ehrmann S, Li J, Ibarra-Estrada M, et al. Awake prone positioning for COVID-19 acute hypoxaemic respiratory failure: a randomised, controlled, multinational, open-label meta-trial. *Lancet Respir Med.* 2021;9(12):1387-1395. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34425070</u>.
- 22. Briel M, Meade M, Mercat A, et al. Higher vs. lower positive end-expiratory pressure in patients with

acute lung injury and acute respiratory distress syndrome: systematic review and meta-analysis. *JAMA*. 2010;303(9):865-873. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/20197533</u>.

- 23. Alhazzani W, Møller MH, Arabi YM, et al. Surviving Sepsis Campaign: guidelines on the management of critically ill adults with coronavirus disease 2019 (COVID-19). *Intensive Care Med.* 2020;46(5):854-887. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32222812</u>.
- 24. Marini JJ, Gattinoni L. Management of COVID-19 respiratory distress. *JAMA*. 2020;323(22):2329-2330. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32329799</u>.
- 25. Tsolaki V, Siempos I, Magira E, et al. PEEP levels in COVID-19 pneumonia. *Crit Care*. 2020;24(1):303. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32505186</u>.
- Bhatraju PK, Ghassemieh BJ, Nichols M, et al. COVID-19 in critically ill patients in the Seattle region—case series. N Engl J Med. 2020;382(21):2012-2022. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32227758</u>.
- 27. Cummings MJ, Baldwin MR, Abrams D, et al. Epidemiology, clinical course, and outcomes of critically ill adults with COVID-19 in New York City: a prospective cohort study. *Lancet*. 2020;395(10239):1763-1770. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32442528</u>.
- Schenck EJ, Hoffman K, Goyal P, et al. Respiratory mechanics and gas exchange in COVID-19-associated respiratory failure. *Ann Am Thorac Soc.* 2020;17(9):1158-1161. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32432896.
- 29. Ziehr DR, Alladina J, Petri CR, et al. Respiratory pathophysiology of mechanically ventilated patients with COVID-19: a cohort study. *Am J Respir Crit Care Med.* 2020;201(12):1560-1564. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32348678.
- Munshi L, Del Sorbo L, Adhikari NKJ, et al. Prone position for acute respiratory distress syndrome. A systematic review and meta-analysis. *Ann Am Thorac Soc.* 2017;14(suppl 4):S280-S288. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29068269</u>.
- National Heart, Lung, and Blood Institute PETAL Clinical Trials Network. Early neuromuscular blockade in the acute respiratory distress syndrome. *N Engl J Med.* 2019;380(21):1997-2008. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31112383</u>.
- Goligher EC, Hodgson CL, Adhikari NKJ, et al. Lung recruitment maneuvers for adult patients with acute respiratory distress syndrome. A systematic review and meta-analysis. *Ann Am Thorac Soc.* 2017;14(suppl 4):S304-S311. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29043837</u>.
- 33. Khokher W, Malhas S, Beran A, et al. Inhaled pulmonary vasodilators in COVID-19 infection: a systematic review and meta-analysis. *J Intensive Care Med*. 2022;37(10):1370-1382. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35915994</u>.
- 34. Gebistorf F, Karam O, Wetterslev J, Afshari A. Inhaled nitric oxide for acute respiratory distress syndrome (ARDS) in children and adults. *Cochrane Database Syst Rev.* 2016(6):CD002787. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27347773</u>.

Pharmacologic Interventions for Critically III Patients

Last Updated: December 20, 2023

Empiric Broad-Spectrum Antimicrobial Therapy

Recommendations

- In the absence of a proven or suspected secondary infection, the COVID-19 Treatment Guidelines Panel (the Panel) **recommends against** the use of empiric broad-spectrum antimicrobials in patients with severe or critical COVID-19 (**BIII**).
- As with any hospitalized patient, patients with COVID-19 who receive antimicrobials should be reassessed daily to minimize the adverse consequences of unnecessary antimicrobial therapy (AIII).

Rationale

Variable rates of community- and hospital-acquired infections have been reported in adult patients with COVID-19. Bacterial coinfection at the time of hospitalization has been reported in 1% to 3.5% of patients with COVID-19.^{1,2} Secondary infections have been reported in 14% to 37% of patients in intensive care units, but the reported rates have been influenced by differences in the severity of illness, duration of hospitalization, method of diagnosis, and time period studied.^{3,4}

No clinical trials have evaluated the use of empiric broad-spectrum antimicrobials in patients with severe or critical COVID-19 or other coronavirus infections. Routine, empiric use of antimicrobials in patients with severe or critical COVID-19 **is not recommended (BIII)**. This recommendation is intended to mitigate the unintended consequences of antimicrobial side effects and resistance. The use of antimicrobials may be considered in specific situations, such as the presence of a lobar infiltrate on a chest X-ray, leukocytosis, an elevated serum lactate level, microbiologic data, or shock.

The use of antimicrobials in patients with severe or critical COVID-19 should follow guidelines established for other hospitalized patients (i.e., for hospital-acquired pneumonia, ventilator-associated pneumonia, or bloodstream infections associated with central lines). It is unclear whether using corticosteroids or other immunomodulatory agents recommended in the Guidelines should alter such approaches.

Therapeutic Management of Hospitalized Adults With COVID-19

For the Panel's recommendations on the use of abatacept, baricitinib, dexamethasone, infliximab, remdesivir, and tocilizumab, see <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>.

Immune-Based Therapy

For recommendations on the use of immunomodulators in patients with COVID-19, see <u>Immunomodulators</u>.

Antithrombotic Therapy

For the Panel's recommendations regarding the use of antithrombotic therapy in critical care settings, see <u>Antithrombotic Therapy in Patients With COVID-19</u> and <u>Therapeutic Management of Hospitalized</u> COVID-19 Treatment Guidelines

Adults With COVID-19.

References

- 1. Langford BJ, So M, Raybardhan S, et al. Bacterial co-infection and secondary infection in patients with COVID-19: a living rapid review and meta-analysis. *Clin Microbiol Infect*. 2020;26(12):1622-1629. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32711058.
- 2. Garcia-Vidal C, Sanjuan G, Moreno-García E, et al. Incidence of co-infections and superinfections in hospitalized patients with COVID-19: a retrospective cohort study. *Clin Microbiol Infect*. 2021;27(1):83-88. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32745596</u>.
- Zangrillo A, Beretta L, Scandroglio AM, et al. Characteristics, treatment, outcomes and cause of death of invasively ventilated patients with COVID-19 ARDS in Milan, Italy. *Crit Care Resusc.* 2020;22(3):200-211. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32900326</u>.
- Yang X, Yu Y, Xu J, et al. Clinical course and outcomes of critically ill patients with SARS-CoV-2 pneumonia in Wuhan, China: a single-centered, retrospective, observational study. *Lancet Respir Med.* 2020;8(5):475-481. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32105632</u>.

Extracorporeal Membrane Oxygenation for Adults

Last Updated: December 20, 2023

Recommendation

• There is insufficient evidence for the COVID-19 Treatment Guidelines Panel to recommend either for or against the use of extracorporeal membrane oxygenation (ECMO) in adults with COVID-19–associated acute respiratory distress syndrome (ARDS) and refractory hypoxemia.

Rationale

ECMO has been used as a rescue therapy in patients with ARDS caused by COVID-19 and refractory hypoxemia. However, there is no conclusive evidence that ECMO is responsible for better clinical outcomes, regardless of the cause of hypoxemic respiratory failure.¹⁻⁴

The clinical outcomes for patients with ARDS who are treated with ECMO are variable and depend on multiple factors, including the etiology of hypoxemic respiratory failure, the severity of pulmonary and extrapulmonary illness, the presence of comorbidities, and the ECMO experience of the individual center.⁵⁻⁷ Several multicenter, observational cohort studies from the first half of 2020⁸⁻¹⁰ reported that patients who required ECMO for COVID-19 had similar mortality to patients in a 2018 randomized study who did not have COVID-19 but had ARDS and received ECMO.³

However, subsequent observational studies reported that in patients who required ECMO for COVID-19, outcomes in late 2020 and early 2021 were worse than outcomes in spring 2020.^{11,12} The largest analysis used data from 4,812 patients in the international Extracorporeal Life Support Organization Registry who had COVID-19 and received ECMO in 2020.¹¹ At centers that provided ECMO throughout 2020, patients who started ECMO before May 1, 2020, had a 90-day mortality of 36.9% after ECMO initiation (95% CI, 34.1% to 39.7%). At the same centers, patients who initiated ECMO between May 2 and December 31, 2020, had a 90-day mortality of 51.9% (95% CI, 50.0% to 53.8%). Furthermore, at centers that started using ECMO for patients with COVID-19 after May 1, 2020, the 90-day mortality after ECMO initiation was 58.9% (95% CI, 55.4% to 62.3%). These observational data should be interpreted with caution, as they may reflect a changing case mix of patients with COVID-19 who were referred for ECMO.

Three target emulation trials compared the efficacy of ECMO and conventional mechanical ventilation in patients with severe COVID-19–associated ARDS.^{10,13,14} The largest of these trials included 844 patients with COVID-19 who had hypoxemic respiratory failure and were receiving ECMO.¹⁴ The study reported that the patients who received ECMO had lower 60-day mortality than the patients who received only conventional mechanical ventilation (26% vs. 33.2%; risk difference –7.1%; 95% CI, –8.2% to –6.1%; risk ratio 0.78; 95% CI, 0.75–0.82). Favorable ECMO outcomes were associated with the following factors: aged <65 years, a ratio of arterial partial pressure of oxygen to fraction of inspired oxygen <80 mm Hg, \leq 10-day duration of mechanical ventilation, and >15 cm H₂O driving pressure.

Ultimately, the benefits of ECMO cannot be clearly defined for patients with COVID-19 and severe ARDS because no randomized controlled trials have evaluated the use of ECMO in this population.

Clinicians interested in pursuing ECMO for patients with COVID-19 and severe ARDS should consider transferring care to high-volume ECMO centers. These patients should be entered into clinical trials or

registries so more informative data can be obtained. More information on the use of ECMO in patients with COVID-19 can be found on the <u>Extracorporeal Life Support Organization website</u>.

References

- Peek GJ, Mugford M, Tiruvoipati R, et al. Efficacy and economic assessment of conventional ventilatory support versus extracorporeal membrane oxygenation for severe adult respiratory failure (CESAR): a multicentre randomised controlled trial. *Lancet*. 2009;374(9698):1351-1363. Available at: <u>https://www.ncbi.</u> <u>nlm.nih.gov/pubmed/19762075</u>.
- 2. Pham T, Combes A, Rozé H, et al. Extracorporeal membrane oxygenation for pandemic influenza A(H1N1)induced acute respiratory distress syndrome: a cohort study and propensity-matched analysis. *Am J Respir Crit Care Med.* 2013;187(3):276-285. Available at: https://www.ncbi.nlm.nih.gov/pubmed/23155145.
- Combes A, Hajage D, Capellier G, et al. Extracorporeal membrane oxygenation for severe acute respiratory distress syndrome. *N Engl J Med*. 2018;378(21):1965-1975. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29791822</u>.
- 4. Munshi L, Walkey A, Goligher E, et al. Venovenous extracorporeal membrane oxygenation for acute respiratory distress syndrome: a systematic review and meta-analysis. *Lancet Respir Med.* 2019;7(2):163-172. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30642776</u>.
- Bullen EC, Teijeiro-Paradis R, Fan E. How I select which patients with ARDS should be treated with venovenous extracorporeal membrane oxygenation. *Chest.* 2020;158(3):1036-1045. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32330459</u>.
- Henry BM, Lippi G. Poor survival with extracorporeal membrane oxygenation in acute respiratory distress syndrome (ARDS) due to coronavirus disease 2019 (COVID-19): pooled analysis of early reports. *J Crit Care*. 2020;58:27-28. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32279018</u>.
- Mustafa AK, Alexander PJ, Joshi DJ, et al. Extracorporeal membrane oxygenation for patients with COVID-19 in severe respiratory failure. *JAMA Surg.* 2020;155(10):990-992. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32780089</u>.
- Barbaro RP, MacLaren G, Boonstra PS, et al. Extracorporeal membrane oxygenation support in COVID-19: an international cohort study of the Extracorporeal Life Support Organization Registry. *Lancet*. 2020;396(10257):1071-1078. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32987008</u>.
- Schmidt M, Hajage D, Lebreton G, et al. Extracorporeal membrane oxygenation for severe acute respiratory distress syndrome associated with COVID-19: a retrospective cohort study. *Lancet Respir Med*. 2020;8(11):1121-1131. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32798468</u>.
- 10. Shaefi S, Brenner SK, Gupta S, et al. Extracorporeal membrane oxygenation in patients with severe respiratory failure from COVID-19. *Intensive Care Med.* 2021;47(2):208-221. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33528595.
- Barbaro RP, MacLaren G, Boonstra PS, et al. Extracorporeal membrane oxygenation for COVID-19: evolving outcomes from the international Extracorporeal Life Support Organization Registry. *Lancet*. 2021;398(10307):1230-1238. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34599878</u>.
- Broman LM, Eksborg S, Lo Coco V, et al. Extracorporeal membrane oxygenation for COVID-19 during first and second waves. *Lancet Respir Med.* 2021;9(8):e80-e81. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34146489</u>.
- 13. Hajage D, Combes A, Guervilly C, et al. Extracorporeal membrane oxygenation for severe acute respiratory distress syndrome associated with COVID-19: an emulated target trial analysis. *Am J Respir Crit Care Med*. 2022;206(3):281-294. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35533052</u>.
- Urner M, Barnett AG, Bassi GL, et al. Venovenous extracorporeal membrane oxygenation in patients with acute COVID-19 associated respiratory failure: comparative effectiveness study. *BMJ*. 2022;377:e068723. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35508314</u>.

Introduction to Critical Care Management of Children With COVID-19

Last Updated: May 31, 2022

COVID-19 may lead to critical illness in children, including hypoxemic respiratory failure, acute respiratory distress syndrome (ARDS), septic shock, cardiac dysfunction, thromboembolic disease, hepatic or renal dysfunction, central nervous system disease, and exacerbation of underlying comorbidities. In addition, multisystem inflammatory syndrome in children (MIS-C) is a rare, postinfectious complication of SARS-CoV-2 and is frequently associated with critical illness.

Data informing the optimal management of children with acute COVID-19 or MIS-C are limited. In general, management should follow the principles of pediatric critical care usually applied to non-COVID-19-related illness, such as the <u>Pediatric Acute Lung Injury Consensus Conference (PALICC)</u> recommendations and the <u>Surviving Sepsis Campaign International Guidelines for the Management</u> of <u>Septic Shock and Sepsis-Associated Organ Dysfunction in Children</u>. For patients with COVID-19 in the intensive care unit (ICU), treatment often requires managing underlying illnesses other than COVID-19 that may have contributed to the need for ICU admission, as well as managing COVID-19 complications. Finally, prevention of ICU-related complications is critical to achieving optimal clinical outcomes for any patient admitted to the ICU.

Selected Clinical Manifestations of COVID-19 Critical Illness

Inflammatory Response

Patients with COVID-19 may develop a hyperinflammatory state, which appears to be distinct from classic "cytokine storm" syndromes (e.g., macrophage activation syndrome in juvenile idiopathic arthritis, familial hemophagocytic lymphohistiocytosis). This phenomenon is less well-described in children than in adults.

Multisystem Inflammatory Syndrome in Children

MIS-C is a rare, postinfectious complication of SARS-CoV-2 that is characterized by persistent fever, systemic inflammation, and multisystem organ dysfunction. The majority of children with MIS-C require ICU-level care, primarily for shock and for vasopressor and inotropic support.¹⁻³ For details on the definition of MIS-C, clinical features, and recommended treatments, see <u>Special Considerations in</u> <u>Children and Therapeutic Management of Hospitalized Children With MIS-C</u>, Plus a Discussion on <u>MIS-A</u>.

Cardiac Dysfunction, Including Myocarditis

Although cardiac involvement is common in patients with MIS-C,^{2,4} cardiac manifestations have rarely been described in children with acute COVID-19. Myocarditis, cardiac conduction abnormalities, and coronary artery aneurysms have been reported in patients with MIS-C. Myocarditis may also occur after SARS-CoV-2 vaccination, particularly in adolescent males, although the clinical course generally is relatively mild.⁵

Thromboembolic Events

Limited data characterize the prevalence of thromboembolic disease in children with COVID-19 or MIS-C. In a multicenter, retrospective cohort study including 814 hospitalized patients with COVID-19 or MIS-C, thromboembolic events were detected in 2.1% of patients with COVID-19 and 6.5% of

patients with MIS-C.⁶ The same study conducted a multivariable analysis and found that the following variables were associated with increased risk of thromboembolic events: children aged \geq 12 years, MIS-C, central venous catheters, and underlying malignancies. See <u>Antithrombotic Therapy in Patients</u> <u>With COVID-19</u> for additional recommendations.

Acute Kidney Injury

Acute kidney injury is estimated to occur in 12% to 44% of hospitalized children with COVID-19 or MIS-C, but the need for renal replacement therapy is extremely rare.⁷⁻¹⁰

Neurologic Involvement

Neurologic involvement is common in children with COVID-19 or MIS-C and is estimated to occur in approximately 30% to 40% of children hospitalized with these conditions.^{2,11} Severe neurologic manifestations, including severe encephalopathy, stroke, demyelinating conditions, cerebral edema, and Guillain-Barré syndrome, have also been described.¹¹

Important Considerations in the Care of Critically III Patients With COVID-19

Considerations for the care of children with COVID-19 or MIS-C should generally follow the usual principles of pediatric critical care. Sedation management and considerations related to post-intensive care syndrome–pediatric (PICS-p) are discussed below. See <u>Oxygenation and Ventilation for Children</u>, <u>Hemodynamic Considerations for Children</u>, and <u>Extracorporeal Membrane Oxygenation for Children</u> for more information on pediatric critical care.

Sedation Management

Guidelines for the management of pain, agitation, neuromuscular blockade, delirium, and early mobility (PANDEM) in infants and children admitted to the pediatric ICU have recently been published.¹² In general, children with COVID-19 or MIS-C who require mechanical ventilation should be managed per the usual critical care for patients with respiratory failure who require mechanical ventilation. The usual care includes sedation with the minimal effective dose required to tolerate mechanical ventilation, optimize gas exchange, and minimize the risk of ventilator-induced lung injury. Using validated pain and sedation scales, the critical care team should set a sedation/pain target based on the phase of ventilation.

Two large randomized controlled trials examined the use of protocols to manage sedation titration in children receiving mechanical ventilation.^{13,14} In both studies, participants received usual care or protocol-driven care implemented by nurses. The studies found that the use of the protocols did not demonstrate a significant benefit on outcomes, such as the duration of ventilation. However, a patient's risk of harm from protocolized sedation is generally low, which led the Society of Critical Care Medicine to issue a conditional recommendation, based on low-level evidence, in its PANDEM clinical practice guidelines suggesting the use of protocolized sedation in children who are critically ill and receiving mechanical ventilation.¹²

Studies evaluating data on the effect of early mobility protocols on critically ill children are limited. One trial evaluated the safety and feasibility of early mobilization in 58 patients who were randomized to receive usual care or early physical therapy, occupational therapy, and speech therapy consultation within 72 hours of admission to the pediatric ICU.¹⁵ Although no differences between the arms were demonstrated for clinical, functional, or quality of life outcomes, the study found that the early rehabilitation consultations were safe and feasible.

Ongoing trials are measuring the effect of early mobilization on patient-centered outcomes in children receiving mechanical ventilation. The PANDEM guideline statement issued by the Society of Critical

Care Medicine conditionally recommends, based on a low quality of evidence, implementing early mobilization strategies in children when feasible, which likely would apply to children with COVID-19 or MIS-C.¹²

Post-Intensive Care Syndrome

In recent years, there has been a growing awareness that PICS can occur in pediatric patients. PICS-p has been demonstrated to have a multifaceted effect on the physical, cognitive, emotional, and social health of child survivors of critical illness and their families.¹⁶ Furthermore, many pediatric survivors of sepsis or ARDS manifest significant impairments in physical, cognitive, and emotional health.¹⁷⁻¹⁹ Although no clear data characterize the prevalence of PICS-p or long-term morbidity in children with COVID-19 or MIS-C, the prevalence is expected to be similar to that observed in other populations with similar illness severities.

Acknowledgments

For these pediatric recommendations, the COVID-19 Treatment Guidelines Panel integrated the recommendations from pediatric-specific guidelines, including the European Society of Paediatric and Neonatal Intensive Care's recommendations²⁰ for the care of critically ill children with COVID-19 and the Surviving Sepsis Campaign's perspective on managing sepsis in children with COVID-19.²¹ In addition, recommendations from several non-COVID-19-specific treatment guidelines, such as the *Surviving Sepsis Campaign International Guidelines for the Management of Septic Shock and Sepsis-Associated Organ Dysfunction in Children*,²² the PALICC recommendations,²³ and the Society of Critical Care Medicine's PANDEM guidelines,¹² were integrated.

References

- 1. Abrams JY, Oster ME, Godfred-Cato SE, et al. Factors linked to severe outcomes in multisystem inflammatory syndrome in children (MIS-C) in the USA: a retrospective surveillance study. *Lancet Child Adolesc Health*. 2021;5(5):323-331. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33711293</u>.
- Feldstein LR, Tenforde MW, Friedman KG, et al. Characteristics and outcomes of US children and adolescents with multisystem inflammatory syndrome in children (MIS-C) compared with severe acute COVID-19. *JAMA*. 2021;325(11):1074-1087. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33625505</u>.
- 3. Alsaied T, Tremoulet AH, Burns JC, et al. Review of cardiac involvement in multisystem inflammatory syndrome in children. *Circulation*. 2021;143(1):78-88. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33166178.
- 4. Valverde I, Singh Y, Sanchez-de-Toledo J, et al. Acute cardiovascular manifestations in 286 children with multisystem inflammatory syndrome associated with COVID-19 infection in Europe. *Circulation*. 2021;143(1):21-32. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33166189.
- 5. Jain SS, Steele JM, Fonseca B, et al. COVID-19 vaccination-associated myocarditis in adolescents. *Pediatrics*. 2021;148(5):e2021053427. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34389692</u>.
- Whitworth H, Sartain SE, Kumar R, et al. Rate of thrombosis in children and adolescents hospitalized with COVID-19 or MIS-C. *Blood*. 2021;138(2):190-198. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33895804.
- 7. Raina R, Chakraborty R, Mawby I, et al. Critical analysis of acute kidney injury in pediatric COVID-19 patients in the intensive care unit. *Pediatr Nephrol*. 2021;36(9):2627-2638. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33928439.
- 8. Kari JA, Shalaby MA, Albanna AS, et al. Acute kidney injury in children with COVID-19: a retrospective study. *BMC Nephrol.* 2021;22(1):202. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34059010</u>.
- 9. Basalely A, Gurusinghe S, Schneider J, et al. Acute kidney injury in pediatric patients hospitalized with acute

COVID-19 Treatment Guidelines

COVID-19 and multisystem inflammatory syndrome in children associated with COVID-19. *Kidney Int.* 2021;100(1):138-145. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33675848</u>.

- Bjornstad EC, Krallman KA, Askenazi D, et al. Preliminary assessment of acute kidney injury in critically ill children associated with SARS-CoV-2 infection: a multicenter cross-sectional analysis. *Clin J Am Soc Nephrol.* 2021;16(3):446-448. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33144276</u>.
- LaRovere KL, Riggs BJ, Poussaint TY, et al. Neurologic involvement in children and adolescents hospitalized in the United States for COVID-19 or multisystem inflammatory syndrome. *JAMA Neurol*. 2021;78(5):536-547. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33666649</u>.
- Smith HAB, Besunder JB, Betters KA, et al. 2022 Society of Critical Care Medicine clinical practice guidelines on prevention and management of pain, agitation, neuromuscular blockade, and delirium in critically ill pediatric patients with consideration of the ICU environment and early mobility. *Pediatr Crit Care Med.* 2022;23(2):e74-e110. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/3511943</u>8.
- Curley MA, Wypij D, Watson RS, et al. Protocolized sedation vs usual care in pediatric patients mechanically ventilated for acute respiratory failure: a randomized clinical trial. *JAMA*. 2015;313(4):379-389. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/25602358</u>.
- Blackwood B, Tume LN, Morris KP, et al. Effect of a sedation and ventilator liberation protocol vs usual care on duration of invasive mechanical ventilation in pediatric intensive care units: a randomized clinical trial. *JAMA*. 2021;326(5):401-410. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34342620</u>.
- 15. Fink EL, Beers SR, Houtrow AJ, et al. Early protocolized versus usual care rehabilitation for pediatric neurocritical care patients: a randomized controlled trial. *Pediatr Crit Care Med.* 2019;20(6):540-550. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30707210</u>.
- Manning JC, Pinto NP, Rennick JE, Colville G, Curley MAQ. Conceptualizing post intensive care syndrome in children—the PICS-p framework. *Pediatr Crit Care Med.* 2018;19(4):298-300. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29406379</u>.
- Herrup EA, Wieczorek B, Kudchadkar SR. Characteristics of postintensive care syndrome in survivors of pediatric critical illness: a systematic review. *World J Crit Care Med.* 2017;6(2):124-134. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28529914</u>.
- Keim G, Watson RS, Thomas NJ, Yehya N. New morbidity and discharge disposition of pediatric acute respiratory distress syndrome survivors. *Crit Care Med.* 2018;46(11):1731-1738. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30024428</u>.
- Watson RS, Asaro LA, Hutchins L, et al. Risk factors for functional decline and impaired quality of life after pediatric respiratory failure. *Am J Respir Crit Care Med.* 2019;200(7):900-909. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31034245</u>.
- 20. Rimensberger PC, Kneyber MCJ, Deep A, et al. Caring for critically ill children with suspected or proven coronavirus disease 2019 infection: recommendations by the Scientific Sections' Collaborative of the European Society of Pediatric and Neonatal Intensive Care. *Pediatr Crit Care Med.* 2021;22(1):56-67. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33003177</u>.
- 21. Weiss SL, Peters MJ, Agus MSD, et al. Perspective of the Surviving Sepsis Campaign on the management of pediatric sepsis in the era of coronavirus disease 2019. *Pediatr Crit Care Med.* 2020;21(11):e1031-e1037. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32886460</u>.
- 22. Weiss SL, Peters MJ, Alhazzani W, et al. Surviving Sepsis Campaign international guidelines for the management of septic shock and sepsis-associated organ dysfunction in children. *Pediatr Crit Care Med.* 2020;21(2):e52-e106. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32032273.
- Pediatric Acute Lung Injury Consensus Conference Group. Pediatric acute respiratory distress syndrome: consensus recommendations from the Pediatric Acute Lung Injury Consensus Conference. Pediatr Crit Care Med. 2015;16(5):428-439. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/25647235</u>.

Hemodynamic Considerations for Children

Last Updated: May 31, 2022

Children with acute COVID-19 infrequently experience shock requiring hemodynamic support. However, similar to children with sepsis or septic shock from other causes, children with COVID-19 and shock should be evaluated and managed per the *Surviving Sepsis Campaign International Guidelines for the Management of Septic Shock and Sepsis-Associated Organ Dysfunction in Children*.^{1,2}

Shock occurs in approximately half of the patients with multisystem inflammatory syndrome in children (MIS-C); reported prevalence ranges from 35% to 80%.³⁻⁵ Limited data inform optimal hemodynamic management for MIS-C. Given that the physiology observed in patients with MIS-C results from a combination of distributive, cardiogenic, and, occasionally, hypovolemic shock, the COVID-19 Treatment Guidelines Panel (the Panel) suggests that clinicians use the management principles outlined in the Surviving Sepsis Campaign's guidelines for children, as well as the principles for clinical management of heart failure and general critical care, as appropriate. The Panel's recommendations apply to the care of children and infants >37 weeks gestational age.

Recommendation

• For children with COVID-19 or MIS-C and shock, the COVID-19 Treatment Guidelines Panel (the Panel) recommends a target mean arterial pressure (MAP) between the fifth and fiftieth, or greater than the fiftieth, percentiles for age (AIII).

Rationale

There are no clinical trials that support specific hemodynamic targets for children with septic shock due to COVID-19, MIS-C, or any other etiology. The panel members for the pediatric Surviving Sepsis Campaign guidelines were divided on the most appropriate MAP target and made no specific recommendation for a target MAP. Therefore, for children with COVID-19 or MIS-C, clinicians should use the same approach used for children without COVID-19 and target a MAP between the fifth and fiftieth, or greater than the fiftieth, percentiles for age. When MAP cannot be reliably measured, systolic blood pressure is a reasonable alternative.²

Recommendation

• The Panel recommends that, when available, a combination of serial clinical assessments; cardiac ultrasound or echocardiography; and/or laboratory markers, including lactate levels, should be used to monitor the response to resuscitation in children with COVID-19 or MIS-C and shock (**BIII**).

Rationale

Observational data from children with non-COVID-19-related sepsis suggest that using clinical assessment alone limits the ability to classify patients with sepsis as having "warm" (i.e., likely to require fluid or vasopressors) or "cold" (i.e., likely to require inotropes) shock, when compared with assessments that include objective measures of cardiac output/index or systemic vascular resistance.^{6,7} Cardiac ultrasonography can be performed at the bedside and serially, and it may provide additional clinical data on volume responsiveness and cardiac function.⁸ Data from studies evaluating use of cardiac ultrasound in children with COVID-19 and MIS-C are limited to reports from case series.⁹

However, given the spectrum of hemodynamic perturbations observed and because approximately a third of children with MIS-C exhibit left ventricular dysfunction, cardiac ultrasonography may have particular value in MIS-C.⁴

Elevated lactate level is associated with worse outcomes in children with non-COVID-19-related sepsis, although the specific threshold is unknown and has varied from 2 mmol/L to 4 mmol/L across studies.^{10,11} Data on serial lactate measures are limited to a single observational study demonstrating an association between normalization in lactate and a decreased risk of persistent organ dysfunction in children with non-COVID-19-related sepsis (adjusted relative risk 0.47; 95% CI, 0.29–0.78).¹² The role of serial lactate measures has not been systematically evaluated for COVID-19 or MIS-C. An observational study of 1,080 children with MIS-C demonstrated an association between elevated markers of inflammation (e.g., C-reactive protein, procalcitonin), brain natriuretic peptide (BNP) or N-terminal proBNP (NT-proBNP), and troponin and the presence of cardiac dysfunction, shock, and the need for intensive care unit admission. However, the timing of the laboratory values in the study was not available, so the elevated markers may reflect, rather than predict, severe illness.³

Recommendation

• The Panel recommends administration of balanced **crystalloids** rather than 0.9% saline for the initial resuscitation of children with shock due to COVID-19 or MIS-C (**CIIb**).

Rationale

No published clinical trials directly compare balanced/buffered crystalloids with 0.9% saline administered to children with sepsis of any etiology, although an international randomized trial is underway (ClinicalTrials.gov Identifier <u>NCT04102371</u>). Two observational studies using administrative data compared the use of balanced/buffered crystalloids to 0.9% saline in propensity-matched cohorts of children with non-COVID-19-related severe sepsis or septic shock. One of the studies compared patients who received any or only Ringer's lactate solution in the first 3 days of admission with patients who received only normal saline. The study demonstrated no differences between the arms for 30-day mortality or frequency of acute kidney injury.¹³

The other study compared patients receiving only balanced fluids with those receiving only 0.9% saline. The study demonstrated that the balanced-fluid arm had lower mortality (12.5% vs. 15.9%; OR 0.76; 95% CI, 0.62–0.93; P = 0.007), reduced acute kidney injury (16.0% vs. 19.2%; OR 0.82; 95% CI, 0.68–0.98; P = 0.028), and fewer days on vasoactive infusions (3.0 days vs. 3.3 days; P < 0.001) than the saline arm.¹⁴ No published studies focused on patients with COVID-19 or MIS-C, although hyponatremia is common in patients with MIS-C, and decisions about the type of fluid therapy used should be individualized for this population.

Recommendations

- The Panel recommends the use of **epinephrine** or **norepinephrine** rather than dopamine in children with COVID-19 or MIS-C and shock (**BIIa**).
- There is insufficient evidence to differentiate between norepinephrine or epinephrine as a first-line vasoactive drug in children with COVID-19 or MIS-C. The choice of vasoactive agent should be individualized and based on clinical examination, laboratory data, and data from cardiac ultrasound or echocardiography.

Rationale

Use of vasoactive infusions should be considered for children with shock due to COVID-19 if signs of

COVID-19 Treatment Guidelines

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

shock persist after resuscitation with 40 mL/kg to 60 mL/kg of fluid, or sooner if there is evidence of cardiac dysfunction or signs of fluid overload (e.g., tachypnea, hepatomegaly). Similar principles may be applied to patients with MIS-C, particularly because their clinical presentation overlaps significantly with the clinical presentation of children with septic shock due to other causes. However, given the high prevalence of cardiac dysfunction in patients with MIS-C, clinicians should consider performing echocardiography or cardiac ultrasound early in the initial resuscitation if MIS-C is suspected and consider initiating a vasoactive infusion if cardiac dysfunction is identified.

Data from pediatric studies comparing vasopressors are limited, and there are no data specific to patients with COVID-19 or MIS-C. Two small pediatric trials compared epinephrine with dopamine in patients with non-COVID-19-related fluid-refractory septic shock.^{15,16} One study randomized 63 children to receive dopamine 5 µg/kg/min to 10 µg/kg/min and 57 children to receive epinephrine 0.1 µg/kg/min to 0.3 µg/kg/min. Mortality by Day 28 was 14.2% in the dopamine arm and 7% in the epinephrine arm (OR 6.5; 95% CI, 1.1–37.8; P = 0.03). In the other study, 31 children were randomized to receive incremental doses of dopamine 10 µg/kg/min to 20 µg/kg/min, and 29 children were randomized to receive incremental doses of epinephrine 0.1 to 0.3 µg/kg/min. The primary outcome of shock resolution within 1 hour occurred in 4 children (13%) receiving dopamine and 12 children (41%) receiving epinephrine (OR 4.8; 95% CI, 1.3–17.2; P = 0.019).

No pediatric trials have compared norepinephrine to other vasoactive agents in patients with sepsis, but based on data from studies of adults, the pharmacologic properties of norepinephrine and dopamine (see <u>Hemodynamics for Adults</u>), and the 2020 Surviving Sepsis Campaign guidelines for children, norepinephrine is suggested over dopamine.²

Collectively, this evidence is insufficient to recommend norepinephrine versus epinephrine as a first-line vasoactive agent in children with COVID-19 or MIS-C. Further, given the varied physiology observed with MIS-C in particular, decisions about which vasopressor to use should be individualized based on clinical and laboratory data and findings from bedside cardiac ultrasound or echocardiography.

Recommendation

• There is insufficient evidence for the Panel to recommend either for or against the use of inodilators (including dobutamine or milrinone) in children with COVID-19 or MIS-C who show evidence of cardiac dysfunction and persistent hypoperfusion despite adequate fluid loading and the use of vasopressor agents.

Rationale

Data from studies evaluating use of inodilators in children with COVID-19, MIS-C, and non-COVID-19-related sepsis are limited to reports from case series. However, the majority of the pediatric Surviving Sepsis Campaign guidelines panel (77%) would use an inodilator at least some of the time for patients with non-COVID-19-related sepsis, cardiac dysfunction, and persistent hypoperfusion despite adequate fluid loading and the use of vasopressor agents.² Expert consultation from specialists in pediatric cardiology and critical care medicine is recommended in this scenario.

Additional Recommendations

- For the acute resuscitation of children with COVID-19 or MIS-C and shock, the Panel recommends the use of **crystalloids** rather than albumin (**AIIb**).
- The Panel **recommends against** using **hydroxyethyl starches** for intravascular volume replacement in children with COVID-19 or MIS-C and sepsis or septic shock (**AIII**).

- For children with refractory shock who have recently completed a course of corticosteroids to treat COVID-19, the Panel recommends using low-dose corticosteroid therapy ("shock-reversal") over no corticosteroid therapy (**CIII**).
 - Children who are currently receiving corticosteroids for COVID-19 or MIS-C are generally receiving sufficient glucocorticoid replacement therapy and do not require additional hydrocortisone for refractory shock.

References

- 1. Weiss SL, Peters MJ, Agus MSD, et al. Perspective of the Surviving Sepsis Campaign on the management of pediatric sepsis in the era of coronavirus disease 2019. *Pediatr Crit Care Med*. 2020;21(11):e1031-e1037. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32886460</u>.
- 2. Weiss SL, Peters MJ, Alhazzani W, et al. Surviving Sepsis Campaign international guidelines for the management of septic shock and sepsis-associated organ dysfunction in children. *Pediatr Crit Care Med.* 2020;21(2):e52-e106. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32032273.
- 3. Abrams JY, Oster ME, Godfred-Cato SE, et al. Factors linked to severe outcomes in multisystem inflammatory syndrome in children (MIS-C) in the USA: a retrospective surveillance study. *Lancet Child Adolesc Health*. 2021;5(5):323-331. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33711293.
- Feldstein LR, Tenforde MW, Friedman KG, et al. Characteristics and outcomes of US children and adolescents with multisystem inflammatory syndrome in children (MIS-C) compared with severe acute COVID-19. *JAMA*. 2021;325(11):1074-1087. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33625505</u>.
- Godfred-Cato S, Bryant B, Leung J, et al. COVID-19-associated multisystem inflammatory syndrome in children—United States, March–July 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(32):1074-1080. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32790663</u>.
- Egan JR, Festa M, Cole AD, et al. Clinical assessment of cardiac performance in infants and children following cardiac surgery. *Intensive Care Med.* 2005;31(4):568-573. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/15711976</u>.
- Razavi A, Newth CJL, Khemani RG, Beltramo F, Ross PA. Cardiac output and systemic vascular resistance: clinical assessment compared with a noninvasive objective measurement in children with shock. *J Crit Care*. 2017;39:6-10. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28088009</u>.
- 8. Ranjit S, Aram G, Kissoon N, et al. Multimodal monitoring for hemodynamic categorization and management of pediatric septic shock: a pilot observational study. *Pediatr Crit Care Med.* 2014;15(1):e17-26. Available at: https://www.ncbi.nlm.nih.gov/pubmed/24196006.
- 9. Kennedy TM, Dessie A, Kessler DO, et al. Point-of-care ultrasound findings in multisystem inflammatory syndrome in children: a cross-sectional study. *Pediatr Emerg Care*. 2021;37(6):334-339. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33871226</u>.
- Scott HF, Brou L, Deakyne SJ, et al. Association between early lactate levels and 30-day mortality in clinically suspected sepsis in children. *JAMA Pediatr.* 2017;171(3):249-255. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28068437</u>.
- 11. Bai Z, Zhu X, Li M, et al. Effectiveness of predicting in-hospital mortality in critically ill children by assessing blood lactate levels at admission. *BMC Pediatr*. 2014;14:83. Available at: https://www.ncbi.nlm.nih.gov/pubmed/24673817.
- Scott HF, Brou L, Deakyne SJ, et al. Lactate clearance and normalization and prolonged organ dysfunction in pediatric sepsis. *J Pediatr*. 2016;170:149-155 e141-144. Available at: https://www.ncbi.nlm.nih.gov/pubmed/26711848.
- Weiss SL, Keele L, Balamuth F, et al. Crystalloid fluid choice and clinical outcomes in pediatric sepsis: a matched retrospective cohort study. *J Pediatr*. 2017;182:304-310 e310. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28063688</u>.

- Emrath ET, Fortenberry JD, Travers C, McCracken CE, Hebbar KB. Resuscitation with balanced fluids is associated with improved survival in pediatric severe sepsis. *Crit Care Med.* 2017;45(7):1177-1183. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28437373</u>.
- 15. Ventura AM, Shieh HH, Bousso A, et al. Double-blind prospective randomized controlled trial of dopamine versus epinephrine as first-line vasoactive drugs in pediatric septic shock. *Crit Care Med.* 2015;43(11):2292-2302. Available at: https://www.ncbi.nlm.nih.gov/pubmed/26323041.
- 16. Ramaswamy KN, Singhi S, Jayashree M, Bansal A, Nallasamy K. Double-blind randomized clinical trial comparing dopamine and epinephrine in pediatric fluid-refractory hypotensive septic shock. *Pediatr Crit Care Med.* 2016;17(11):e502-e512. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27673385</u>.

Oxygenation and Ventilation for Children

Last Updated: September 26, 2022

The COVID-19 Treatment Guidelines Panel's (the Panel) recommendations in this section were informed by recommendations from the Surviving Sepsis Campaign's guidelines for managing <u>adult</u> <u>sepsis</u>, <u>pediatric sepsis</u>, and <u>COVID-19</u>, as well as by recommendations from the 2015 Pediatric Acute Lung Injury Consensus Conference (<u>PALICC</u>).

Goal of Oxygenation

Recommendations

- A target oxygen saturation measured by pulse oximetry (SpO₂) of 92% to 97% is recommended for most children with COVID-19 who require supplemental oxygen (**AIIb**).
- For children with severe pediatric acute respiratory distress syndrome (PARDS; i.e., with an oxygenation index ≥16 or SpO₂ index ≥12.3), an SpO₂ <92% can be considered to minimize exposure to a high fraction of inspired oxygen (FiO₂), but prolonged periods of SpO₂ <88% should be avoided (CIII).

Rationale

The optimal SpO₂ in children with COVID-19 is unknown. However, there is no evidence that the target SpO₂ should differ from the 2015 PALICC recommendation.¹ An SpO₂ of 92% to 97% is recommended for most children with COVID-19 who require supplemental oxygen. The potential harm of hyperoxia in children was demonstrated in a recent meta-analysis of 11 observational studies of children without COVID-19.² The study demonstrated that critically ill children with hyperoxia had greater odds of mortality than those without hyperoxia (OR 1.59; 95% CI, 1.00–2.51). However, there was significant heterogeneity across the included studies for populations, definitions of hyperoxia, and the timing of assessments for mortality outcomes. For children with severe PARDS (i.e., those with an oxygenation index \geq 16 or SpO₂ index \geq 12.3),¹ an SpO₂ <92% can be considered to minimize exposure to a high FiO₂. Although no evidence clearly identifies a safe minimum SpO₂ in children, prolonged exposure to SpO₂ <88% should be avoided. When SpO₂ is <92%, monitoring oxygen delivery markers, including central venous SpO₂, is suggested.³

The limitations of currently available measurement devices should be considered when using pulse oximetry to manage children with COVID-19 or PARDS. Observational studies in children have reported that pulse oximetry may be inaccurate, particularly at lower oxygen saturations (\leq 90%) and for children who are Black.^{4,5} These reports are consistent with several adult observational studies that also identified inaccuracies in pulse oximetry measurements, particularly for patients with darker skin pigmentation.⁶⁻⁸ See <u>Clinical Spectrum of SARS-CoV-2 Infection</u> for more information.

Although procedures vary across institutions, the treatment of most children with PARDS who are critically ill is managed without the use of arterial lines or arterial blood gas testing, because arterial line placement in children, especially young children, can result in complications.⁹⁻¹¹ Clinicians should monitor for adequate delivery of oxygen or consider lowering the threshold for arterial line placement if a patient's SpO₂ measurements could be unreliable (e.g., for children who have darker skin or low SpO₂ levels). Monitoring methods could include observing the patient for altered mentation, measuring venous oxygen saturation, or using near-infrared spectroscopy.

High-Flow Nasal Cannula Oxygen and Noninvasive Ventilation for Children With COVID-19 and Acute Respiratory Failure

Recommendation

• For infants and children with COVID-19 and persistent respiratory failure despite conventional oxygen therapy who have no indicators for endotracheal intubation, a time-limited trial of noninvasive ventilation (NIV) or high-flow nasal cannula (HFNC) oxygen is recommended (AIIa). There is insufficient evidence for the Panel to recommend either for or against the use of HFNC oxygen over NIV or the use of NIV over HFNC oxygen in infants and children with COVID-19.

Rationale

No high-quality studies have evaluated the use of HFNC oxygen or NIV in children with COVID-19. Therefore, when choosing a mode of respiratory support for children with COVID-19, the principles of management used for patients without COVID-19 should be followed. Both the *Surviving Sepsis Campaign International Guidelines for the Management of Septic Shock and Sepsis-Associated Organ Dysfunction in Children* and PALICC recommend the use of NIV for children with respiratory failure who have no indication for intubation.^{12,13}

Furthermore, the response to NIV, particularly for children with more severe hypoxemia or high work of breathing, should be gauged early (within the first several hours). If the patient does not show improvement, intubation should be considered. To unload respiratory muscles, bilevel modes of NIV (with inspiratory pressure augmentation, such as BiPAP), if tolerated, are preferred over the use of continuous positive airway pressure (CPAP) alone, although CPAP is an alternative for children who cannot achieve an adequate seal with the NIV interface or who have significant patient-ventilator asynchrony.¹²

HFNC oxygen is a relatively new, but increasingly used, mode of respiratory support for infants and children with acute respiratory failure.¹⁴ Data from studies evaluating the effectiveness of HFNC oxygen relative to NIV or conventional oxygen are limited to studies of children with pneumonia in limited-resource settings and studies of children with bronchiolitis. Two randomized controlled trials of children with pneumonia were conducted in limited-resource settings. One study demonstrated a slightly lower relative risk of mortality with the use of HFNC oxygen when compared with conventional oxygen therapy (aHR 0.79; 95% CI, 0.54–1.16), although the results were not statistically significant.¹⁵ The other trial demonstrated that children treated with bubble CPAP ventilation had a lower risk of mortality than children who received low-flow oxygen (relative risk 0.25; 95% CI, 0.07–0.89; P = 0.02).¹⁶ The results also indicated that for the composite outcome of treatment failure, there was no difference between the use of HFNC oxygen and bubble CPAP (relative risk 0.50; 99.7% CI, 0.11–2.29).

A randomized, noninferiority trial compared HFNC oxygen (2 L/kg/min) and nasal CPAP among 142 infants aged <6 months with bronchiolitis not caused by COVID-19.¹⁷ The primary outcome was treatment failure within 24 hours, defined as an increase of \geq 1 point in the modified Wood's Clinical Asthma Score (M-WCAS) or Échelle Douleur Inconfort Nouveau-Né (EDIN) score (a neonatal pain and discomfort scale), a respiratory rate >60 breaths/min and an increase of >10 breaths/min from baseline, or >2 severe apnea episodes per hour. Treatment failure occurred more often in the HFNC oxygen arm than in the nasal CPAP arm (51% vs. 31%), a result that failed to meet the prespecified noninferiority margin. Notably, in the HFNC oxygen arm, 72% of the patients who had treatment failure were managed successfully with nasal CPAP, and there were no differences between the arms for intubation rates or length of stay in the pediatric intensive care unit (PICU).

A systematic review of the noninferiority trial and 2 smaller trials comparing HFNC oxygen to nasal CPAP summarized the results of 213 infants and children aged ≤ 2 years with bronchiolitis.¹⁸ Treatment failure in the 2 smaller trials was rare, and no differences were detected between the HFNC oxygen and nasal CPAP arms for any of the clinical outcomes.^{19,20}

In a study that assessed whether higher flow rates of HFNC oxygen improved outcomes, 286 infants aged ≤ 6 months and with severe bronchiolitis were randomized to receive HFNC oxygen 2 L/kg/min or HFNC oxygen 3 L/kg/min.²¹ The primary outcome of treatment failure (i.e., an increase of ≥ 1 point in M-WCAS or EDIN score, a respiratory rate ≥ 60 breaths/min and an increase of ≥ 10 breaths/min from baseline, or ≥ 2 severe apnea episodes per hour) occurred in 38.7% of the infants in the 2 L/kg/min arm and in 38.9% of the infants in the 3 L/kg/min arm (P = 0.98). Patient discomfort, as measured by EDIN score, occurred more often in the 3 L/kg/min arm than in the 2 L/kg/min arm (43% vs. 16%; P = 0.002).

HFNC oxygen is increasingly being used in children. These studies highlight the potential role of an HFNC oxygen trial in the management of children with acute respiratory failure due to COVID-19, particularly for infants and young children who may have NIV-related challenges, such as poor mask fit, discomfort, or patient-ventilator asynchrony. For the use of HFNC oxygen in children, consider flow rates of up to 2 L/kg/min, with a maximum of 60 L/min. If patients do not improve within the first few hours of receiving HFNC oxygen, their treatment should be escalated to NIV or intubation.

Awake Prone Positioning for Children Not Receiving Mechanical Ventilation

Recommendations

- There is insufficient evidence for the Panel to recommend either for or against a trial of awake prone positioning for children with persistent hypoxemia who require HFNC oxygen or NIV and do not require endotracheal intubation.
- For patients with refractory hypoxemia who meet the indications for intubation and mechanical ventilation, the Panel **recommends against** the use of awake prone positioning as a rescue therapy to avoid intubation (**AIII**).

Rationale

There are no high-quality pediatric data evaluating the effect of awake prone positioning on clinical outcomes in children with COVID-19 or non-COVID-19-related illness. Awake prone positioning may be considered for older children and adolescents (see <u>Oxygenation and Ventilation for Adults</u>). In addition, pediatric clinicians should consider a child's developmental stage and ability to comply with the protocols for awake prone positioning.

Intubation for Mechanical Ventilation in Children With Acute COVID-19

Recommendations

- If intubation becomes necessary, the Panel recommends that an experienced practitioner perform the procedure in a controlled setting due to the enhanced risk of exposing health care practitioners to SARS-CoV-2 during intubation (AIII).
- The Panel recommends using cuffed endotracheal tubes over uncuffed endotracheal tubes in children who require endotracheal intubation (AIIb).

Rationale

To optimize the safety of patients and health care workers and maximize first-attempt success, intubation should be performed in a controlled setting by an experienced practitioner. In addition, cuffed

endotracheal tubes are preferred for children of all ages to minimize leaks around the endotracheal tube, ensure delivery of ventilator pressure, decrease the risk of aspiration, reduce the need for endotracheal tube exchange, and reduce aerosolization of respiratory secretions during mechanical ventilation.^{3,22-24}

General Considerations for Children With COVID-19 and PARDS Who Require Mechanical Ventilation

Recommendations

For children with COVID-19 and PARDS who require mechanical ventilation:

- The Panel recommends using low tidal volume (VT) ventilation (VT 4–8 mL/kg of predicted body weight) over higher VT ventilation (VT >8 mL/kg) (AIIb).
- The Panel recommends targeting plateau pressures of ≤ 28 cm H₂O for children with normal chest wall compliance and ≤ 32 cm H₂O for those with impaired chest wall compliance (AIII).
- The Panel recommends using a higher positive end-expiratory pressure (PEEP) strategy (i.e., 10–15 cm H₂O or higher in patients with severe PARDS) over a lower PEEP strategy, titrated based on observed responses in oxygenation, hemodynamics, and respiratory system compliance (**BIIb**).
- The Panel recommends permissive hypercapnia (e.g., pH 7.15–7.30), if needed, to remain within lung-protective strategies and to minimize ventilator-associated lung injury, provided the patient does not have a coexisting condition that would be worsened by acidosis (e.g., severe pulmonary hypertension, ventricular dysfunction, intracranial hypertension) (AIII).
- The Panel **recommends against** the routine use of **inhaled nitric oxide (AIII)**.

Rationale

There is no evidence that ventilator management of children with PARDS due to COVID-19 should differ from ventilator management of patients with PARDS due to other causes. The Panel's recommendations are derived from the 2015 PALICC recommendations.^{1,3} Since the publication of the PALICC recommendations, no randomized trials have provided significant new evidence, although some observational data support some of the PALICC recommendations.

A large observational study conducted in 71 international PICUs reported that for patients with mild to moderate acute respiratory distress syndrome (ARDS), less adherence to the recommended VT of 5 mL/kg to 8 mL/kg (or 3 mL/kg to 6 mL/kg for patients with severe ARDS) was associated with higher mortality and with more time on ventilation.²⁵ In general, supraphysiologic VT ventilation (>8 mL/kg) should not be used in patients with PARDS, and VT should be adjusted within the acceptable range to maintain other lung-protective ventilation targets (e.g., maintaining ≤ 28 cm H₂O plateau pressure). The use of ultra-low VT ventilation (<4 mL/kg) has not been systematically studied in children, so it should be used with caution.

The ARDS Network established a ventilator protocol that includes suggested low PEEP/high FiO₂ levels.²⁶ The protocol suggests that for patients receiving FiO₂ \ge 0.6, a PEEP level of \ge 10 cm H₂O would be implemented, which aligns with recommendations from PALICC. Two observational studies have reported better clinical outcomes associated with use of the suggested (or higher) PEEP levels compared to lower PEEP levels.^{25,27} The multicenter studies, which included nearly 1,500 pediatric patients with ARDS, demonstrated that PEEP levels lower than those recommended by the ARDS Network were associated with higher mortality.

Inhaled nitric oxide can be considered as a rescue therapy for children with severe PARDS and COVID-19. In a small, randomized trial, the use of inhaled nitric oxide resulted in reduced use of *COVID-19 Treatment Guidelines*

extracorporeal membrane oxygenation (ECMO).²⁸ However, inhaled nitric oxide has a heterogeneous treatment effect, and many patients do not show improved gas exchange. Although adverse effects are rare, use of inhaled nitric oxide can have a substantial effect on health care costs. Therefore, inhaled nitric oxide should not be considered routine therapy for children with PARDS or COVID-19 who are receiving mechanical ventilation.

Fluid Management for Children With PARDS

Recommendation

• Following an initial resuscitation in children with PARDS due to COVID-19, clinicians should monitor and titrate fluid balance to maintain adequate intravascular volume while aiming to prevent positive fluid balance (**BIIb**).

Rationale

There is no evidence that fluid management in children with PARDS due to COVID-19 should differ from fluid management in patients with PARDS due to other causes. Therefore, the Panel's recommendation aligns with the PALICC recommendation.¹ No pediatric randomized trials have directly compared a liberal fluid strategy to a conservative fluid strategy in patients with PARDS of any etiology. Several observational studies have demonstrated an association between greater fluid overload and worse clinical outcomes, including fewer ventilator-free days and increased mortality.²⁹⁻³¹

In a multicenter study of 168 children with acute lung injury, daily and cumulative fluid balance were measured over the first 7 days after participants met the inclusion criteria. After adjusting for demographic characteristics, pediatric risk of mortality III (PRISM III) scores, vasopressor use, and the ratio of arterial partial pressure of oxygen to fraction of inspired oxygen, an increasing cumulative fluid balance on Day 3 was associated with fewer ventilator-free days, but no association with mortality was detected.²⁹

A more recent single-center study that included 732 children with acute lung injury demonstrated an association between higher cumulative fluid balance on Days 5 to 7 and increased mortality (for 100 mL/kg on Day 5, OR 1.34; 95% CI, 1.11–1.61) after adjusting for oxygenation index, the number of nonpulmonary organ failures, immunocompromised status, and vasopressor scores. Also, greater cumulative fluid balance on Days 4 to 7 was associated with a lower probability of successful extubation by Day 28.³¹ Collectively, the findings from these pediatric observational studies demonstrate the potential harm of fluid overload in children with PARDS, particularly after 3 to 4 days of illness.

These results are consistent with the findings from FACTT, a trial of conservative versus liberal fluid management strategies in adults.³² In adults, FACTT found no difference between the arms for 60-day mortality, but the conservative strategy arm demonstrated improved oxygenation and less time on mechanical ventilation and in the intensive care unit when compared with the liberal strategy arm. However, no analysis of data from prospective pediatric trials delineates a causal relationship between a specific, protocolized fluid management strategy, or the timing of such a strategy, and clinical outcomes. Therefore, an individualized fluid management approach that is titrated to maintain intravascular volume while preventing excessive positive fluid balance, as suggested by the 2015 PALICC recommendation, is appropriate.¹

Neuromuscular Blockade for Mechanically Ventilated Children With Severe PARDS

Recommendation

• For mechanically ventilated children with severe PARDS and COVID-19, the Panel recommends minimal yet effective use of neuromuscular blocking agents in conjunction with sedation, if sedation

COVID-19 Treatment Guidelines

alone is inadequate to achieve lung-protective ventilation (BIII).

Rationale

There is no evidence that the use of neuromuscular blockade in children with COVID-19 should differ from practices used for severe PARDS from other causes. Therefore, the Panel's recommendation aligns directly with the PALICC recommendation.¹ Since the publication of the 2015 PALICC recommendation, no new data support significant changes to the recommendation.

Therapies for Mechanically Ventilated Children With Severe PARDS and Refractory Hypoxemia

Recommendations

For children with severe PARDS and refractory hypoxemia after other oxygenation strategies have been optimized:

- The Panel recommends **inhaled nitric oxide** as a rescue therapy; if no rapid improvement in oxygenation is observed, inhaled nitric oxide should be discontinued (**BIIb**).
- The Panel recommends prone positioning for 12 to 16 hours per day over no prone positioning (**BIII**).
- There is insufficient evidence for the Panel to recommend either for or against the use of recruitment maneuvers, but if they are used in children, slow incremental and decremental adjustments in PEEP are preferred to sustained inflation maneuvers.
- There is insufficient evidence for the Panel to recommend either for or against the use of high-frequency oscillatory ventilation (HFOV) in children with PARDS.

Rationale

There is no evidence that the use of inhaled nitric oxide, prone positioning, or HFOV in children with COVID-19 should differ from practices used for severe PARDS from other causes. Therefore, the Panel's recommendations are largely based on PALICC recommendations.¹ Since the publication of the 2015 PALICC recommendations, many new trials evaluating these practices have been conducted.

One randomized controlled trial and 2 propensity-matched, observational studies have evaluated the use of inhaled nitric oxide in patients with PARDS since the publication of the PALICC recommendations. The randomized controlled trial included 55 patients and found that the use of inhaled nitric oxide resulted in no statistical difference between the arms for 28-day mortality (8% mortality in the inhaled nitric oxide arm vs. 28% in the placebo arm), although the trial was underpowered for this outcome.²⁸ However, the inhaled nitric oxide arm had approximately 5 more ventilator-free days than the placebo arm, a result that was primarily mediated by avoiding the use of ECMO. These results have been corroborated by observational studies, which also reported more ventilator-free days for patients who received inhaled nitric oxide.^{33,34} Although the evidence is insufficient to recommend the use of inhaled nitric oxide for all patients with ARDS, in cases of severe hypoxemia, it can be considered as a rescue therapy to potentially avoid the use of ECMO.

No new studies have evaluated the role of prone positioning in PARDS, although a large, multicenter trial is ongoing. Therefore, the Panel's recommendation to consider prone positioning in cases of severe PARDS aligns with the PALICC recommendation and is supported by adult data, primarily from PROSEVA, a trial on prone positioning in patients with ARDS.³⁵

The 2015 PALICC recommendations included the use of careful recruitment maneuvers with incremental and decremental adjustments in PEEP.¹ In children, this approach to recruitment maneuvers is preferred over sustained inflation maneuvers due to the increased risk of harm from barotrauma and hemodynamic compromise in patients with sustained inflation. Clinical trials in adults have highlighted the potential harm of applying recruitment maneuvers to patients who may not have recruitable lung.^{36,37} Therefore, although there is insufficient evidence to recommend either for or against the use of recruitment maneuvers in children with refractory hypoxemia, if recruitment maneuvers are used, the preferred strategy is slow, incremental and decremental adjustments in PEEP.

Since the publication of the 2015 PALICC recommendations, 2 small randomized controlled trials have examined the use of HFOV for PARDS.^{38,39} Neither study found a significant difference for mortality. Several observational studies using propensity matching have shown either no difference in outcomes between the HFOV and conventional ventilation arms or a potential for higher mortality or a longer ventilation time with the use of HFOV when compared with conventional ventilation.⁴⁰⁻⁴⁴ In some of these analyses, residual confounding has been a concern. A large, multicenter randomized controlled trial of HFOV for PARDS is ongoing. Therefore, the Panel has determined that there is insufficient evidence to recommend either for or against the use of HFOV in COVID-19-related PARDS. Some concerns have been raised about the use of HFOV and the aerosolization of COVID-19; however, adding a filter to the expiratory limb of the HFOV circuit may alleviate these concerns.

Multisystem Inflammatory Syndrome in Children

More than half of the patients with multisystem inflammatory syndrome in children (MIS-C) require mechanical ventilation or NIV.⁴⁵⁻⁴⁷ For patients with MIS-C, the indications for mechanical ventilation vary and include shock or cardiac dysfunction, pulmonary edema, procedural preparation (e.g., to facilitate sedation for central venous catheter placement), respiratory failure, or neurologic failure. The management of oxygenation and ventilation in children with MIS-C should follow the usual principles of shock management outlined in the Surviving Sepsis Campaign guidelines for children, as well as the principles for clinical management of heart failure and general critical care, as appropriate.¹³

References

- 1. Pediatric Acute Lung Injury Consensus Conference Group. Pediatric acute respiratory distress syndrome: consensus recommendations from the Pediatric Acute Lung Injury Consensus Conference. *Pediatr Crit Care Med.* 2015;16(5):428-439. Available at: https://www.ncbi.nlm.nih.gov/pubmed/25647235.
- Lilien TA, Groeneveld NS, van Etten-Jamaludin F, et al. Association of arterial hyperoxia with outcomes in critically ill children: a systematic review and meta-analysis. *JAMA Netw Open*. 2022;5(1):e2142105. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34985516</u>.
- 3. Rimensberger PC, Cheifetz IM, Pediatric Acute Lung Injury Consensus Conference Group. Ventilatory support in children with pediatric acute respiratory distress syndrome: proceedings from the Pediatric Acute Lung Injury Consensus Conference. *Pediatr Crit Care Med.* 2015;16(5 suppl 1):S51-S60. Available at: https://www.ncbi.nlm.nih.gov/pubmed/26035364.
- 4. Ross PA, Newth CJ, Khemani RG. Accuracy of pulse oximetry in children. *Pediatrics*. 2014;133(1):22-29. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/24344108</u>.
- 5. Andrist E, Nuppnau M, Barbaro RP, Valley TS, Sjoding MW. Association of race with pulse oximetry accuracy in hospitalized children. *JAMA Netw Open*. 2022;5(3):e224584. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35357460.
- 6. Chesley CF, Lane-Fall MB, Panchanadam V, et al. Racial disparities in occult hypoxemia and clinically based mitigation strategies to apply in advance of technological advancements. *Respir Care*. 2022;Published online ahead of print. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35679133.

- Valbuena VSM, Seelye S, Sjoding MW, et al. Racial bias and reproducibility in pulse oximetry among medical and surgical inpatients in general care in the Veterans Health Administration 2013–19: multicenter, retrospective cohort study. *BMJ*. 2022;378:e069775. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35793817.
- 8. Wong AI, Charpignon M, Kim H, et al. Analysis of discrepancies between pulse oximetry and arterial oxygen saturation measurements by race and ethnicity and association with organ dysfunction and mortality. *JAMA Netw Open*. 2021;4(11):e2131674. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34730820.
- Khemani RG, Smith L, Lopez-Fernandez YM, et al. Paediatric acute respiratory distress syndrome incidence and epidemiology (PARDIE): an international, observational study. *Lancet Respir Med.* 2019;7(2):115-128. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30361119</u>.
- Mahendra M, McQuillen P, Dudley RA, Steurer MA. Variation in arterial and central venous catheter use in pediatric intensive care units. *J Intensive Care Med*. 2021;36(11):1250-1257. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32969326</u>.
- Gleich SJ, Wong AV, Handlogten KS, Thum DE, Nemergut ME. Major short-term complications of arterial cannulation for monitoring in children. *Anesthesiology*. 2021;134(1):26-34. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33079134</u>.
- Essouri S, Carroll C, Pediatric Acute Lung Injury Consensus Conference Group. Noninvasive support and ventilation for pediatric acute respiratory distress syndrome: proceedings from the Pediatric Acute Lung Injury Consensus Conference. *Pediatr Crit Care Med.* 2015;16(5 suppl 1):S102-S110. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26035360</u>.
- 13. Weiss SL, Peters MJ, Alhazzani W, et al. Surviving Sepsis Campaign international guidelines for the management of septic shock and sepsis-associated organ dysfunction in children. *Pediatr Crit Care Med.* 2020;21(2):e52-e106. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32032273.
- 14. Willer RJ, Johnson MD, Cipriano FA, et al. Implementation of a weight-based high-flow nasal cannula protocol for children with bronchiolitis. *Hosp Pediatr*. 2021;11(8):891-895. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34234010.
- 15. Maitland K, Kiguli S, Olupot-Olupot P, et al. Randomised controlled trial of oxygen therapy and high-flow nasal therapy in African children with pneumonia. *Intensive Care Med.* 2021;47(5):566-576. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33954839.
- Chisti MJ, Salam MA, Smith JH, et al. Bubble continuous positive airway pressure for children with severe pneumonia and hypoxaemia in Bangladesh: an open, randomised controlled trial. *Lancet*. 2015;386(9998):1057-1065. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26296950</u>.
- Milesi C, Essouri S, Pouyau R, et al. High flow nasal cannula (HFNC) versus nasal continuous positive airway pressure (nCPAP) for the initial respiratory management of acute viral bronchiolitis in young infants: a multicenter randomized controlled trial (TRAMONTANE study). *Intensive Care Med.* 2017;43(2):209-216. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28124736</u>.
- Moreel L, Proesmans M. High flow nasal cannula as respiratory support in treating infant bronchiolitis: a systematic review. *Eur J Pediatr*. 2020;179(5):711-718. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32232547</u>.
- 19. Sarkar M, Sinha R, Roychowdhoury S, et al. Comparative study between noninvasive continuous positive airway pressure and hot humidified high-flow nasal cannulae as a mode of respiratory support in infants with acute bronchiolitis in pediatric intensive care unit of a tertiary care hospital. *Indian J Crit Care Med.* 2018;22(2):85-90. Available at: https://www.ncbi.nlm.nih.gov/pubmed/29531447.
- 20. Vahlkvist S, Jurgensen L, la Cour A, et al. High flow nasal cannula and continuous positive airway pressure therapy in treatment of viral bronchiolitis: a randomized clinical trial. *Eur J Pediatr*. 2020;179(3):513-518. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31828528</u>.
- 21. Milesi C, Pierre AF, Deho A, et al. A multicenter randomized controlled trial of a 3-L/kg/min versus 2-L/kg/ min high-flow nasal cannula flow rate in young infants with severe viral bronchiolitis (TRAMONTANE 2).

Intensive Care Med. 2018;44(11):1870-1878. Available at: https://www.ncbi.nlm.nih.gov/pubmed/30343318.

- 22. Weiss M, Dullenkopf A, Fischer JE, et al. Prospective randomized controlled multi-centre trial of cuffed or uncuffed endotracheal tubes in small children. *Br J Anaesth*. 2009;103(6):867-873. Available at: https://www.ncbi.nlm.nih.gov/pubmed/19887533.
- 23. Shi F, Xiao Y, Xiong W, Zhou Q, Huang X. Cuffed versus uncuffed endotracheal tubes in children: a metaanalysis. *J Anesth*. 2016;30(1):3-11. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26296534</u>.
- 24. Matava CT, Kovatsis PG, Lee JK, et al. Pediatric airway management in COVID-19 patients: consensus guidelines from the Society for Pediatric Anesthesia's Pediatric Difficult Intubation Collaborative and the Canadian Pediatric Anesthesia Society. *Anesth Analg.* 2020;131(1):61-73. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32287142.
- 25. Bhalla AK, Klein MJ, Emeriaud G, et al. Adherence to lung-protective ventilation principles in pediatric acute respiratory distress syndrome incidence and epidemiology study. *Crit Care Med.* 2021;49(10):1779-1789. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34259438.
- 26. Acute Respiratory Distress Syndrome Network, Brower RG, Matthay MA, et al. Ventilation with lower tidal volumes as compared with traditional tidal volumes for acute lung injury and the acute respiratory distress syndrome. *N Engl J Med*. 2000;342(18):1301-1308. Available at: https://www.ncbi.nlm.nih.gov/pubmed/10793162.
- 27. Khemani RG, Parvathaneni K, Yehya N, et al. Positive end-expiratory pressure lower than the ARDS network protocol is associated with higher pediatric acute respiratory distress syndrome mortality. *Am J Respir Crit Care Med.* 2018;198(1):77-89. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29373802</u>.
- 28. Bronicki RA, Fortenberry J, Schreiber M, Checchia PA, Anas NG. Multicenter randomized controlled trial of inhaled nitric oxide for pediatric acute respiratory distress syndrome. *J Pediatr*. 2015;166(2):365-369. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/25454942</u>.
- 29. Valentine SL, Sapru A, Higgerson RA, et al. Fluid balance in critically ill children with acute lung injury. *Crit Care Med.* 2012;40(10):2883-2889. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/22824936</u>.
- 30. Lima L, Menon S, Goldstein SL, Basu RK. Timing of fluid overload and association with patient outcome. *Pediatr Crit Care Med.* 2021;22(1):114-124. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32947381</u>.
- Black CG, Thomas NJ, Yehya N. Timing and clinical significance of fluid overload in pediatric acute respiratory distress syndrome. *Pediatr Crit Care Med.* 2021;22(9):795-805. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33965988</u>.
- 32. National Heart, Lung, and Blood Institute Acute Respiratory Distress Syndrome Clinical Trials Network, Wiedemann HP, Wheeler AP, et al. Comparison of two fluid-management strategies in acute lung injury. *N Engl J Med.* 2006;354(24):2564-2575. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/16714767</u>.
- 33. Gupta P, Richardson T, Hall M, et al. Effect of inhaled nitric oxide on outcomes in children with acute lung injury: propensity matched analysis from a linked database. *Crit Care Med.* 2016;44(10):1901-1909. Available at: https://www.ncbi.nlm.nih.gov/pubmed/27163193.
- 34. Bhalla AK, Yehya N, Mack WJ, et al. The association between inhaled nitric oxide treatment and ICU mortality and 28-day ventilator-free days in pediatric acute respiratory distress syndrome. *Crit Care Med*. 2018;46(11):1803-1810. Available at: https://www.ncbi.nlm.nih.gov/pubmed/30028363.
- 35. Guerin C, Reignier J, Richard JC, et al. Prone positioning in severe acute respiratory distress syndrome. *N Engl J Med.* 2013;368(23):2159-2168. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/23688302</u>.
- 36. Writing Group for the Alveolar Recruitment for Acute Respiratory Distress Syndrome Trial Investigators, Cavalcanti AB, Suzumura EA, et al. Effect of lung recruitment and titrated positive end-expiratory pressure (PEEP) vs low PEEP on mortality in patients with acute respiratory distress syndrome: a randomized clinical trial. *JAMA*. 2017;318(14):1335-1345. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/28973363</u>.
- 37. Hodgson CL, Cooper DJ, Arabi Y, et al. Maximal recruitment open lung ventilation in acute respiratory distress syndrome (PHARLAP): a Phase II, multicenter randomized controlled clinical trial. *Am J Respir Crit*

Care Med. 2019;200(11):1363-1372. Available at: https://www.ncbi.nlm.nih.gov/pubmed/31356105.

- 38. Samransamruajkit R, Rassameehirun C, Pongsanon K, et al. A comparison of clinical efficacy between high frequency oscillatory ventilation and conventional ventilation with lung volume recruitment in pediatric acute respiratory distress syndrome: a randomized controlled trial. *Indian J Crit Care Med.* 2016;20(2):72-77. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27076706</u>.
- 39. El-Nawawy A, Moustafa A, Heshmat H, Abouahmed A. High frequency oscillatory ventilation versus conventional mechanical ventilation in pediatric acute respiratory distress syndrome: a randomized controlled study. *Turk J Pediatr.* 2017;59(2):130-143. Available at: https://www.ncbi.nlm.nih.gov/pubmed/29276865.
- 40. Gupta P, Green JW, Tang X, et al. Comparison of high-frequency oscillatory ventilation and conventional mechanical ventilation in pediatric respiratory failure. *JAMA Pediatr*. 2014;168(3):243-249. Available at: https://www.ncbi.nlm.nih.gov/pubmed/24445980.
- 41. Guo YX, Wang ZN, Li YT, et al. High-frequency oscillatory ventilation is an effective treatment for severe pediatric acute respiratory distress syndrome with refractory hypoxemia. *Ther Clin Risk Manag*. 2016;12:1563-1571. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27799777</u>.
- 42. Bateman ST, Borasino S, Asaro LA, et al. Early high-frequency oscillatory ventilation in pediatric acute respiratory failure. A propensity score analysis. *Am J Respir Crit Care Med*. 2016;193(5):495-503. Available at: https://www.ncbi.nlm.nih.gov/pubmed/26492410.
- 43. Rowan CM, Loomis A, McArthur J, et al. High-frequency oscillatory ventilation use and severe pediatric ARDS in the pediatric hematopoietic cell transplant recipient. *Respir Care*. 2018;63(4):404-411. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29279362</u>.
- 44. Wong JJ, Liu S, Dang H, et al. The impact of high frequency oscillatory ventilation on mortality in paediatric acute respiratory distress syndrome. *Crit Care*. 2020;24(1):31. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32005285.
- 45. Yasuhara J, Watanabe K, Takagi H, Sumitomo N, Kuno T. COVID-19 and multisystem inflammatory syndrome in children: a systematic review and meta-analysis. *Pediatr Pulmonol*. 2021;56(5):837-848. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33428826</u>.
- 46. Feldstein LR, Tenforde MW, Friedman KG, et al. Characteristics and outcomes of US children and adolescents with multisystem inflammatory syndrome in children (MIS-C) compared with severe acute COVID-19. *JAMA*. 2021;325(11):1074-1087. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33625505</u>.
- Godfred-Cato S, Bryant B, Leung J, et al. COVID-19-associated multisystem inflammatory syndrome in children—United States, March–July 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(32):1074-1080. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32790663</u>.

Extracorporeal Membrane Oxygenation for Children

Last Updated: May 31, 2022

Recommendation

• The COVID-19 Treatment Guidelines Panel (the Panel) recommends that the use of extracorporeal membrane oxygenation (ECMO) should be considered for children with acute COVID-19 or multisystem inflammatory syndrome in children (MIS-C) who have refractory hypoxemia or shock when hemodynamic parameters cannot be maintained or lung-protective strategies result in inadequate gas exchange (CIII). Candidacy for ECMO should be determined on a case-by-case basis by the multidisciplinary team.

Rationale

ECMO is used as a rescue therapy for children with refractory hypoxemia or shock. Similar to outcomes for adults, outcomes for children managed with venovenous ECMO are variable and are influenced by the etiology and duration of respiratory failure and by underlying comorbid medical conditions.^{1,2} In addition, studies have shown that pediatric centers that treat fewer patients with ECMO have worse outcomes than facilities that treat a high volume of patients with ECMO.^{3,4} No randomized trials evaluate the efficacy or benefit of ECMO for hypoxemic respiratory failure in children without COVID-19 beyond the neonatal period. In an observational study of 122 children with severe pediatric acute respiratory distress syndrome (PARDS), 90-day mortality for children treated with ECMO and for those supported without ECMO was similar (25% vs. 30%).⁵

The Pediatric Acute Lung Injury Consensus Conference recommends considering ECMO for patients with severe PARDS from reversible causes or for children who are candidates for lung transplantation.⁶ The *Surviving Sepsis Campaign International Guidelines for the Management of Septic Shock and Sepsis-Associated Organ Dysfunction in Children* issued a weak recommendation, based on very low quality of evidence, to use venovenous ECMO for children with PARDS and refractory hypoxemia.⁷

Venoarterial ECMO has been used successfully for the treatment of refractory shock in children, although no trials evaluate this approach, and the potential benefits must be weighed against risks of bleeding or thromboembolic events.⁸⁻¹⁰ The Surviving Sepsis Campaign guidelines for children issued a weak recommendation, based on very low quality of evidence, for use of venoarterial ECMO in children with shock that is refractory to all other treatments; however, a standardized definition of refractory shock in children is not available.⁷

Studies evaluating data on the use of ECMO in children with COVID-19 and MIS-C are limited to case reports and case series.¹¹⁻¹³ A publicly available <u>registry for pediatric patients with COVID-19 on ECMO</u> is maintained by the multinational Extracorporeal Life Support Organization (ELSO). In-hospital mortality at 90 days was about 30%, which is similar to reports from non-COVID-19 ECMO cohorts.^{14,15} ELSO has published guidelines for use of ECMO in COVID-19.¹⁶ In general, ECMO candidacy for children with COVID-19 or MIS-C should be assessed using criteria similar to those used for other causes of severe respiratory failure or shock. Cannulation approaches and management principles should follow published <u>international guidelines</u> and local protocols for non-COVID-19 patients.

Pediatric clinicians should consider entering patients into clinical trials or registries to inform future

recommendations regarding use of ECMO in children with COVID-19. The following resources provide more information on an international ECMO registry and on clinical trials evaluating ECMO in children with COVID-19:

- The ELSO registry for ECMO in COVID-19
- <u>ClinicalTrials.gov</u>

References

- Zabrocki LA, Brogan TV, Statler KD, et al. Extracorporeal membrane oxygenation for pediatric respiratory failure: survival and predictors of mortality. *Crit Care Med.* 2011;39(2):364-370. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/20959787</u>.
- 2. Gow KW, Heiss KF, Wulkan ML, et al. Extracorporeal life support for support of children with malignancy and respiratory or cardiac failure: the extracorporeal life support experience. *Crit Care Med.* 2009;37(4):1308-1316. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/19242331</u>.
- Freeman CL, Bennett TD, Casper TC, et al. Pediatric and neonatal extracorporeal membrane oxygenation: does center volume impact mortality? *Crit Care Med.* 2014;42(3):512-519. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/24164955</u>.
- Gonzalez DO, Sebastiao YV, Cooper JN, Minneci PC, Deans KJ. Pediatric extracorporeal membrane oxygenation mortality is related to extracorporeal membrane oxygenation volume in US hospitals. *J Surg Res.* 2019;236:159-165. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30694751</u>.
- 5. Barbaro RP, Xu Y, Borasino S, et al. Does extracorporeal membrane oxygenation improve survival in pediatric acute respiratory failure? *Am J Respir Crit Care Med*. 2018;197(9):1177-1186. Available at: https://www.ncbi.nlm.nih.gov/pubmed/29373797.
- Dalton HJ, Macrae DJ, Pediatric Acute Lung Injury Consensus Conference Group. Extracorporeal support in children with pediatric acute respiratory distress syndrome: proceedings from the Pediatric Acute Lung Injury Consensus Conference. *Pediatr Crit Care Med.* 2015;16(5 Suppl 1):S111-S117. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26035361</u>.
- Weiss SL, Peters MJ, Alhazzani W, et al. Surviving Sepsis Campaign international guidelines for the management of septic shock and sepsis-associated organ dysfunction in children. *Pediatr Crit Care Med*. 2020;21(2):e52-e106. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32032273</u>.
- Schlapbach LJ, Chiletti R, Straney L, et al. Defining benefit threshold for extracorporeal membrane oxygenation in children with sepsis—a binational multicenter cohort study. *Crit Care*. 2019;23(1):429. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31888705</u>.
- 9. Ramanathan K, Yeo N, Alexander P, et al. Role of extracorporeal membrane oxygenation in children with sepsis: a systematic review and meta-analysis. *Crit Care*. 2020;24(1):684. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33287861.
- 10. Oberender F, Ganeshalingham A, Fortenberry JD, et al. Venoarterial extracorporeal membrane oxygenation versus conventional therapy in severe pediatric septic shock. *Pediatr Crit Care Med.* 2018;19(10):965-972. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30048365</u>.
- 11. Di Nardo M, Hoskote A, Thiruchelvam T, et al. Extracorporeal membrane oxygenation in children with coronavirus disease 2019: preliminary report from the collaborative european chapter of the extracorporeal life support organization prospective survey. ASAIO J. 2021;67(2):121-124. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33009172</u>.
- Alfoudri H, Shamsah M, Yousuf B, AlQuraini N. Extracorporeal membrane oxygenation and extracorporeal cardiopulmonary resuscitation for a COVID-19 pediatric patient: a successful outcome. *ASAIO J.* 2021;67(3):250-253. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33627597</u>.
- 13. Feldstein LR, Tenforde MW, Friedman KG, et al. Characteristics and outcomes of US children and adolescents

with multisystem inflammatory syndrome in children (MIS-C) compared with severe acute COVID-19. *JAMA*. 2021;325(11):1074-1087. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33625505</u>.

- 14. Extracorporeal Life Support Organization. Registry dashboard of ECMO-supported COVID-19 patient data. 2022. Available at: <u>https://www.elso.org/Registry/FullCOVID-19RegistryDashboard.aspx?</u> goHash=1&sO=1&all=true&NA= false&Eur=false&Asia=false&La=false&Africa=false&AA=false&Neo= true&Ped=true&Adlt=false&AllDts=true&YTD=false#TheFilter. Accessed May 18, 2022.
- 15. Extracorporeal Life Support Organization. ELSO live registry dashboard of ECMO patient data. 2022. Available at: <u>https://www.elso.org/Registry/ELSOLiveRegistryDashboard.aspx</u>. Accessed May 20, 2022.
- Badulak J, Antonini MV, Stead CM, et al. Extracorporeal membrane oxygenation for COVID-19: updated 2021 guidelines from the extracorporeal life support organization. *ASAIO J.* 2021;67(5):485-495. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33657573.

Antiviral Agents, Including Antibody Products

Last Updated: November 2, 2023

Remdesivir and ritonavir-boosted nirmatrelvir (Paxlovid) are approved by the Food and Drug Administration for the treatment of COVID-19.

Molnupiravir and high-titer COVID-19 convalescent plasma (CCP) are available only under Food and Drug Administration Emergency Use Authorizations for the treatment of COVID-19.

Summary Recommendations

Recommendations for Treating Nonhospitalized Adults

- The COVID-19 Treatment Guidelines Panel (the Panel) recommends the following anti-SARS-CoV-2 therapies as preferred treatments for COVID-19. These drugs are listed in order of preference:
 - Ritonavir-boosted nirmatrelvir (Paxlovid) (Alla)
 - Remdesivir (Blla)
- The Panel recommends **molnupiravir** as an alternative therapy when neither of the preferred therapies are available, feasible to use, or clinically appropriate **(Clla)**.

Recommendations for Treating Nonhospitalized Children

• For recommendations on using antiviral therapy in nonhospitalized children, see <u>Therapeutic Management of</u> <u>Nonhospitalized Children With COVID-19</u>.

Recommendations for Treating Hospitalized Adults or Children

 See <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> and <u>Therapeutic Management of Hospitalized</u> <u>Children With COVID-19</u> for recommendations on using remdesivir with or without immunomodulators in certain hospitalized patients.

Antiviral Treatments With Insufficient Evidence

- There is insufficient evidence for the Panel to recommend either for or against the use of high-titer CCP for the treatment of COVID-19 in hospitalized or nonhospitalized patients who are immunocompromised.
 - Some people who are immunocompromised have prolonged, symptomatic COVID-19 with evidence of ongoing SARS-CoV-2 replication. For the Panel's recommendations for managing these patients, see <u>Special Considerations in People Who Are Immunocompromised</u>.
- There is insufficient evidence for the Panel to recommend either for or against the use of high-titer CCP for the treatment of COVID-19 in nonhospitalized patients who are immunocompetent.

Antiviral Treatments That the Panel Recommends Against

- The Panel recommends against the use of the following drugs for the treatment of COVID-19, except in a clinical trial:
 - Interferon alfa or beta in nonhospitalized patients (Alla)
 - Interferon alfa in hospitalized patients (Alla)
 - Nitazoxanide (Blla)
- The Panel recommends against the use of the following drugs for the treatment of COVID-19:
 - Anti-SARS-CoV-2 monoclonal antibodies (AIII)
 - Chloroquine or hydroxychloroquine and/or azithromycin in hospitalized (AI) and nonhospitalized patients (Alla)
 - CCP in hospitalized patients who are immunocompetent (AI)
 - Lopinavir/ritonavir and other HIV protease inhibitors in hospitalized (AI) and nonhospitalized patients (AIII)
 - Systemic interferon beta in hospitalized patients (AI)

Summary Recommendations, continued

COVID-19 Pre-Exposure Prophylaxis

• The Panel **recommends against** the use of **tixagevimab plus cilgavimab (Evusheld)** as pre-exposure prophylaxis (PrEP) of COVID-19 (AIII).

The sections on <u>Chloroquine or Hydroxychloroquine and/or Azithromycin</u>, <u>Lopinavir/Ritonavir and Other HIV Protease</u> <u>Inhibitors</u>, and <u>Nitazoxanide</u> have been archived. The Panel will no longer be updating the information on these therapies.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Remdesivir

Last Updated: July 21, 2023

Remdesivir is a nucleotide prodrug of an adenosine analog. It binds to the viral RNA-dependent RNA polymerase and inhibits viral replication by terminating RNA transcription prematurely. Remdesivir has demonstrated in vitro and in vivo activity against SARS-CoV-2.¹

Intravenous remdesivir is approved by the Food and Drug Administration (FDA) for the treatment of COVID-19 in adults and pediatric patients aged ≥ 28 days and weighing ≥ 3 kg. In nonhospitalized patients with mild to moderate COVID-19 who are at high risk of progressing to severe disease, remdesivir should be started within 7 days of symptom onset and administered for 3 days. Hospitalized patients should receive remdesivir for 5 days or until hospital discharge, whichever comes first.² The FDA prescribing information for remdesivir indicates that if a patient does not clinically improve, clinicians may extend the treatment course for up to 5 additional days (for a total duration of 10 days). See <u>Table 4e</u> for more information.

Remdesivir has been studied in several clinical trials for the treatment of COVID-19. The recommendations from the COVID-19 Treatment Guidelines Panel (the Panel) are based on the results of these studies. See <u>Table 4a</u> for more information.

Recommendations

- For the Panel's recommendations and information on the clinical efficacy of using remdesivir to treat high-risk, nonhospitalized patients with mild to moderate COVID-19, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>.
- For the Panel's recommendations and information on the clinical efficacy of using remdesivir with or without immunomodulators to treat certain hospitalized patients, see <u>Therapeutic Management</u> of Hospitalized Adults With COVID-19.
- The data on using combinations of antiviral therapies for the treatment of COVID-19 are limited.³ Clinical trials are needed to determine the role of combination therapy in treating certain patients.

Monitoring, Adverse Effects, and Drug-Drug Interactions

Remdesivir can cause gastrointestinal symptoms (e.g., nausea), elevated transaminase levels, an increase in prothrombin time without a change in the international normalized ratio, and hypersensitivity reactions.

Before starting patients on remdesivir, the FDA recommends performing liver function and prothrombin time tests as clinically appropriate and repeating these tests during treatment as clinically indicated. Remdesivir may need to be discontinued if a patient's alanine transaminase (ALT) level increases to >10 times the upper limit of normal, and it should be discontinued if increases in ALT levels and signs or symptoms of liver inflammation are observed.²

Remdesivir should be administered in a setting where severe hypersensitivity reactions, such as anaphylaxis, can be managed. Patients should be monitored during the infusion and observed for at least 1 hour after the infusion as clinically appropriate.

Currently, no clinical drug-drug interaction studies of remdesivir have been conducted. In vitro, remdesivir is a minor substrate of cytochrome P450 (CYP) 3A4 and a substrate of the drug transporters

organic anion transporting polypeptide (OATP) 1B1 and P-glycoprotein. It is also an inhibitor of CYP3A4, OATP1B1, OATP1B3, and multidrug and toxin extrusion protein (MATE) 1.² See <u>Table 4e</u> for more information.

Patients Who Are Immunocompromised and Have Prolonged Symptoms and Evidence of Ongoing Viral Replication

Patients who are severely immunocompromised may have a prolonged duration of SARS-CoV-2 replication, which may lead to rapid viral evolution. There is concern that using a single antiviral agent in these patients may result in the emergence of resistant virus.⁴ Additional studies are needed to assess this risk. The role of combination antiviral therapy in the treatment of COVID-19 is not yet known.

For patients who are immunocompromised and have prolonged COVID-19 symptoms and evidence of ongoing viral replication (e.g., those with a low cycle threshold value, as measured by a reverse transcription polymerase chain reaction result or with a positive rapid antigen test result) despite receiving a course of antiviral therapy, the optimal management is unknown. Case reports and case series have documented the treatment of these patients with additional antiviral treatments, prolonged courses of antiviral treatments, high-titer COVID-19 convalescent plasma, or combination therapy.⁵⁻⁹ For a discussion of potential treatment options, see <u>Special Considerations in People Who Are Immunocompromised</u> and <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>.

Considerations in Patients With Renal Insufficiency

Remdesivir is formulated with sulfobutylether-beta-cyclodextrin (SBECD) sodium.² SBECD is a vehicle that is primarily eliminated through the kidneys. Accumulation of SBECD in patients with renal impairment may result in liver and renal toxicities.

Basing its decision on safety data primarily from the REDPINE clinical trial and pharmacokinetic data from a Phase 1 trial, the FDA updated the prescribing information for remdesivir to indicate that it can be used without dose adjustment in patients with an estimated glomerular filtration rate (eGFR) of <30 mL/min, including those receiving dialysis.²

Safety data for the use of remdesivir in patients with severely reduced kidney function are available from 2 randomized controlled trials:

- The REDPINE study was a manufacturer-sponsored, multinational, double-blind trial of remdesivir versus placebo in hospitalized adults with severe COVID-19 and an eGFR of <30 mL/ min.¹⁰ The trial was terminated due to low enrollment. Among 163 remdesivir and 80 placebo recipients with a mean age of 69 years, there were no statistically significant differences in treatment-emergent adverse events or serious treatment-emergent adverse events, including death. Among participants with baseline acute kidney injury or chronic kidney disease, there were no statistically significant differences in the progression of acute kidney injury, the need for renal replacement therapy, or death.
- The CATCO study was a multicenter, open-label trial that compared the use of remdesivir to standard of care in hospitalized adults with COVID-19.¹¹ A post hoc analysis was done for 59 patients with a baseline eGFR of <30 mL/min; 15 of these patients were on dialysis. The median age of the cohort was 74 years. Thirty-four patients received remdesivir for a median duration of 10 days, while 25 patients received standard of care. The standard of care patients had a lower median eGFR at baseline (12.4 mL/min) than patients treated with remdesivir (22.7 mL/min). There was no increased risk of renal toxicity at Day 5 among patients treated with remdesivir compared to standard of care, and there were no statistically significant differences in the need for

new dialysis, the need for mechanical ventilation, or mortality.

Although both the REDPINE and CATCO trials were underpowered to assess the clinical efficacy of remdesivir in patients with severely reduced kidney function, the available data suggest that remdesivir can be used safely in patients with an eGFR of <30 mL/min. These results are consistent with a systematic review of observational studies¹² and other retrospective studies that have reported that remdesivir was not associated with an increased incidence of adverse effects in patients with COVID-19 who had baseline eGFRs of <30 mL/min.¹³⁻¹⁵

Considerations in Pregnancy

See <u>Pregnancy</u>, <u>Lactation</u>, <u>and COVID-19 Therapeutics</u> for the Panel's guidance regarding the use of remdesivir during pregnancy and lactation.

Considerations in Children

See <u>Special Considerations in Children</u>, <u>Therapeutic Management of Nonhospitalized Children With</u> <u>COVID-19</u>, and <u>Therapeutic Management of Hospitalized Children With COVID-19</u>.

References

- 1. Wang M, Cao R, Zhang L, et al. Remdesivir and chloroquine effectively inhibit the recently emerged novel coronavirus (2019-nCoV) in vitro. *Cell Res.* 2020;30(3):269-271. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32020029</u>.
- 2. Remdesivir (Veklury) [package insert]. Food and Drug Administration. 2023. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2023/214787s019lbl.pdf.
- 3. Gliga S, Lübke N, Killer A, et al. Rapid selection of sotrovimab escape variants in severe acute respiratory syndrome coronavirus 2 Omicron-infected immunocompromised patients. *Clin Infect Dis.* 2023;76(3):408-415. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36189631</u>.
- Gandhi S, Klein J, Robertson AJ, et al. De novo emergence of a remdesivir resistance mutation during treatment of persistent SARS-CoV-2 infection in an immunocompromised patient: a case report. *Nat Commun.* 2022;13(1):1547. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35301314</u>.
- 5. Huygens S, Gharbharan A, Serroukh Y, et al. High-titer convalescent plasma plus nirmatrelvir/ritonavir treatment for non-resolving COVID-19 in six immunocompromised patients. *J Antimicrob Chemother*. 2023;78(7):1644-1648. Available at: https://www.ncbi.nlm.nih.gov/pubmed/37248664.
- Brosh-Nissimov T, Ma'aravi N, Leshin-Carmel D, et al. Combination treatment of persistent COVID-19 in immunocompromised patients with remdesivir, nirmaltrevir/ritonavir and tixegavimab/cilgavimab. *medRxiv*. 2023;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2023.04.07.23288144v1</u>.
- Mikulska M, Sepulcri C, Dentone C, et al. Triple combination therapy with two antivirals and monoclonal antibodies for persistent or relapsed SARS-CoV-2 infection in immunocompromised patients. *Clin Infect Dis*. 2023;Published online ahead of print. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36976301</u>.
- 8. Graziani L, Gori L, Manciulli T, et al. Successful use of nirmatrelvir/ritonavir in immunocompromised patients with persistent and/or relapsing COVID-19. *J Antimicrob Chemother*. 2023;78(2):555-558. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36544352</u>.
- 9. Trottier CA, Wong B, Kohli R, et al. Dual antiviral therapy for persistent coronavirus disease 2019 and associated organizing pneumonia in an immunocompromised host. *Clin Infect Dis*. 2023;76(5):923-925. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36281907</u>.
- Santos JR, Goldman JD, Tuttle KR, et al. The REDPINE study: efficacy and safety of remdesivir in people with moderately and severely reduced kidney function hospitalised for COVID-19 pneumonia. Presented at: 33rd European Congress of Clinical Microbiology and Infectious Diseases; April 15–18, 2023; Copenhagen,

Denmark. Available at: <u>https://www.askgileadmedical.com/docs/conference/JoseRamon_ECCMID2023_</u> <u>Redpine_P2635@pdf</u>.

- 11. Cheng M, Fowler R, Murthy S, et al. Remdesivir in patients with severe kidney dysfunction: a secondary analysis of the CATCO randomized trial. *JAMA Netw Open*. 2022;5(8):e2229236. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36036936.
- 12. Davoudi-Monfared E, Ahmadi A, Karimpour-Razkenari E, et al. Remdesivir administration in COVID-19 patients with renal impairment: a systematic review. *Am J Ther*. 2022;29(5):e520-e533. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35984955.
- Sunny S, Samaroo-Campbell J, Abdallah M, Luka A, Quale J. Is remdesivir safe in patients with renal impairment? Experience at a large tertiary urban medical center. *Infection*. 2023;51(1):247-252. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35616879</u>.
- 14. Aiswarya D, Arumugam V, Dineshkumar T, et al. Use of remdesivir in patients with COVID-19 on hemodialysis: a study of safety and tolerance. *Kidney Int Rep.* 2021;6(3):586-593. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33354635</u>.
- 15. Shah MK, Parikh M, Prajapati D, et al. Safety and tolerability of remdesivir in patients with end-stage renal disease on maintenance hemodialysis. *Indian J Crit Care Med.* 2022;26(5):619-625. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35719430</u>.

Table 4a. Remdesivir: Selected Clinical Trial Data

Last Updated: August 8, 2022

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for RDV. The studies summarized below are those that have had the greatest impact on the Panel's recommendations. Studies of hospitalized patients are listed first, followed by 1 study of nonhospitalized patients.

Methods	Results	Limitations and Interpretation	
ACTT-1: Multinational, Double-Blind, Placebo-Controlled Trial of Remdesivir in Hospitalized Patients With COVID-19 in 10 Countries ¹			
Key Inclusion Criteria	Participant Characteristics	Key Limitations	
 Laboratory-confirmed SARS-CoV-2 infection ≥1 of the following: 	 Mean age 59 years; 64% men; 53% White, 21% Black, 13% Asian, 24% Hispanic/Latinx 	 Wide range of disease severity among patients; study not powered to detect 	
Pulmonary infiltrates	 Coexisting conditions: 26% with 1; 55% with ≥2 	differences within subgroups	
• $\text{SpO}_2 \leq 94\%$ on room air	• 13% not on oxygen; 41% on supplemental oxygen; 18% on HFNC oxygen or NIV; 27% on MV or ECMO	 Study not powered to detect differences in mortality between arms 	
 Need for supplemental oxygen, HFNC oxygen, NIV, MV, or ECMO 	Median time from symptom onset to randomization: 9 days (IQR	 No data on longer-term morbidity 	
	6–12 days)	Interpretation	
 Key Exclusion Criteria ALT or AST >5 times ULN 	 23% received corticosteroids during study 	• In patients with severe COVID-19, RDV	
	Primary Outcomes	reduced the time to clinical recovery.	
• eGFR <30 mL/min	• Time to clinical recovery: 10 days in RDV arm vs. 15 days in	• The benefit was most apparent in	
Interventions	placebo arm (rate ratio for recovery 1.29; 95% Cl, 1.12–1.49; P	hospitalized patients who were receiving supplemental oxygen.	
• RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily for up to 9 more days (n = 541)	< 0.001)	 There was no observed benefit in those 	
• Placebo for up to 10 days ($n = 521$)	 Benefit of RDV greatest in patients randomized during first 10 days after symptom onset and those who required supplemental 	on HFNC oxygen, NIV, MV, or ECMO, but	
	oxygenation at enrollment	the study was not powered to detect	
 Primary Endpoint Time to clinical recovery 	 No difference in time to recovery for patients on HFNC oxygen, NIV, MV, or ECMO at enrollment 	differences within subgroups.	
Key Secondary Endpoints	Secondary Outcomes		
Clinical status at Day 15, as measured by an OSMortality by Day 29	 Improvement in clinical status at Day 15 more likely in RDV arm (OR 1.5; 95% Cl, 1.2–1.9; P < 0.001) 		
Occurrence of SAEs	 No difference between arms in mortality by Day 29 		
	Occurrence of SAEs: 25% in RDV arm vs. 32% in placebo arm		

COVID-19 Treatment Guidelines

Methods	Results	Limitations and Interpretation	
CATCO: Multicenter, Open-Label, Pragmatic RCT of Remdesivir in Hospitalized Patients With COVID-19 in Canada ²			
Key Inclusion Criterion	Participant Characteristics	Key Limitations	
 Laboratory-confirmed SARS-CoV-2 infection 	Median age 66 years; 60% men; 41% White	Open-label study	
Key Exclusion Criterion Already receiving RDV	Median time from symptom onset to randomization: 8 days	 Information on comorbidities was not available for 26% of patients. 	
Interventions	At entry:54% on low-flow oxygen	Interpretation	
• RDV 200 mg IV on Day 0, then RDV 100 mg IV once daily on Days 1–9 (n = 634)	• 24% on HFNC oxygen	RDV did not decrease in-hospital mortality among patients with	
• Local SOC (n = 647)	• 9% on MV	COVID-19 compared to SOC.	
Primary EndpointIn-hospital mortality	 Rates of comorbidities were similar between arms. 87% in both arms were receiving corticosteroids at baseline 	 Patients who received RDV were less likely to require MV than patients who received SOC. 	
Key Secondary Endpoints	Primary Outcome		
New need for MVHospital LOS	• In-hospital mortality: 19% in RDV arm vs. 23% in SOC arm (relative risk 0.83; 95% Cl, 0.67–1.03)		
Incidence of hepatic dysfunction, incidence of need for	Secondary Outcomes		
dialysis, and change in SCr at Day 5	 New need for MV: 8% in RDV arm vs. 15% in SOC arm (relative risk 0.53; 95% Cl, 0.38–0.75) 		
	No significant difference between arms in hospital LOS		
	• No difference between arms in incidence of new hepatic dysfunction, incidence of need for dialysis, or change in SCr at Day 5		

Methods	Results	Limitations and Interpretation
DisCoVeRy: Open-Label, Adaptive RCT of Remdesivir in Hospitalized Patients With Moderate or Severe COVID-19 in Europe ³		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Laboratory-confirmed SARS-CoV-2 infection 	 Median age 64 years; 70% men; 69% White 	 Open-label study
Illness of any duration	 74% with ≥1 coexisting condition 	• 440 participants in this study also
• Sp0 ₂ \leq 94% on room air or use of supplemental oxygen,	 40% received corticosteroids 	enrolled in the WHO Solidarity trial.
HFNČ oxygen, NIV, or MV	• Median time from symptom onset to randomization: 9 days	Interpretation
Key Exclusion Criteria	in both arms	• There was no clinical benefit of RDV
• ALT or AST >5 times ULN	61% with moderate disease; 39% with severe disease	in hospitalized patients who were
Severe chronic kidney disease	Primary Outcome	symptomatic for >7 days and who required supplemental oxygen.
Interventions	• No difference between arms in clinical status at Day 15 (OR 0.98; 95% Cl, $0.77-1.25$; $P = 0.85$)	roquioù ouppionontai oxygon.
• RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily for up to 9 days (n = 429)	 A prespecified subgroup analysis based on duration 	
• SOC (n = 428)	of symptoms found no significant difference in clinical	
	status between arms.	
Primary Endpoint	Secondary Outcomes	
Clinical status at Day 15, as measured by an OS	• Mortality by Day 29: 8% in RDV arm vs. 9% in SOC arm	
Key Secondary Endpoints	• Occurrence of SAEs: 33% in RDV arm vs. 31% in SOC arm	
Mortality by Day 29	(P = 0.48)	
Occurrence of SAEs		

Methods	Results	Limitations and Interpretation
WHO Solidarity Trial, Final Report: Multinational, Open-Label, Adaptive RCT in Hospitalized Patients With COVID-19 in 35 Countries ⁴		
Key Inclusion Criterion	Participant Characteristics	Key Limitations
 Not known to have received any study drug 	 46% aged 50–69 years; 22% aged ≥70 years; 63% men 	Open-label study
Interventions	 Rates of comorbidities were similar between arms At entry: 	 No data on time from symptom onset to enrollment
 RDV 200 mg IV on Day 0, then RDV 100 mg IV once daily on Days 1–9 (n = 4,146) Local SOC (n = 4,129) 	 71% on supplemental oxygen 9% on MV 	 Data analysis did not separate receipt of low-flow and high-flow oxygen
Primary Endpoint In-hospital mortality	 68% received corticosteroids during study; 4.6% received IL-6 inhibitors 	Interpretation
Key Secondary Endpoint	Primary Outcome	 There was no benefit of RDV in patients who were on MV at
Initiation of MV	 In-hospital mortality: 14.5% in RDV arm vs. 15.6% in SOC arm (rate ratio 0.91; 95% Cl, 0.82–1.02; P = 0.12) 	baseline.Compared to SOC, RDV had a
	 On MV: 42.1% vs. 38.6% (rate ratio 1.13; 95% Cl, 0.89–1.42; P = 0.32) 	modest but statistically significant effect on reducing the risk of death
	 Not on MV but receiving oxygen: 14.6% vs. 16.3% (rate ratio 0.87; 95% Cl, 0.76–0.99; P = 0.03) 	or progression to MV in hospitalized patients who required oxygen.
	 Not on oxygen initially: 2.9% vs. 3.8% (rate ratio 0.76; 95% Cl, 0.46–1.28; P = 0.30) 	
	Secondary Outcome	
	 Initiation of MV: 14.1% in RDV arm vs. 15.7% in SOC arm (rate ratio 0.88; 95% Cl, 0.77–1.00; P = 0.04) 	

Methods	Results	Limitations and Interpretation
<u>GS-US-540-5774 Study</u> : Multinational, Open-Label RCT of 10 Days or 5 Days of Remdesivir Compared With Standard of Care in Hospitalized Patients With Moderate COVID-19 in Asia, Europe, and the United States ⁵		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Laboratory-confirmed SARS-CoV-2 infection Pulmonary infiltrates 	• Demographic and baseline disease characteristics were similar across arms.	Open-label design may have affected decisions on concomitant mediaations (a.g., mars nations in
• $\text{SpO}_2 > 94\%$ on room air	Median age 57 years; 61% men; 58% White	medications (e.g., more patients in SOC arm received AZM, HCQ or CQ,
Key Exclusion Criteria	• 84% required no supplemental oxygen; 15% required low- flow oxygen; 1% required HFNC oxygen or NIV	and LPV/RTV) and time of hospital discharge.
 ALT or AST >5 times ULN CrCl <50 mL/min 	• Concomitant medication use in the 10-day RDV, 5-day RDV, and SOC arms:	 No data on time to return to activity for discharged patients
Interventions	• Steroids: 15%, 17%, 19%	· ·
• RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily for 9 days (n = 193)	 Tocilizumab: 1%, 1%, 5% HCQ or CQ: 11%, 8%, 45% 	InterpretationHospitalized patients with moderate
• RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily for 4 days (n = 191)	• LPV/RTV: 6%, 5%, 22%	COVID-19 who received 5 days of RDV had better clinical status at Day 11 than those who received SOC.
• Local SOC (n = 200)	• AZM: 21%, 18%, 31%	
Primary Endpoint	Median duration of therapy: 6 days in 10-day RDV arm vs. 5 days in 5-day RDV arm	 There was no difference in clinical status at Day 11 between patients who received 10 days of RDV and
Clinical status at Day 11, as measured by an OS	Primary Outcome	those who received SOC.
	Clinical status at Day 11:	
	 Significantly better in 5-day RDV arm than in SOC arm (OR 1.65; 95% Cl, 1.09–2.48; P = 0.02) 	
	• No difference between 10-day RDV arm and SOC arm ($P = 0.18$)	

Methods	Results	Limitations and Interpretation
<u>GS-US-540-5773 Study</u> : Multinational, Open-Label RCT of 10 Days or 5 Days of Remdesivir Compared With Standard of Care in Hospitalized Patients With Severe COVID-19 in Asia, Europe, and the United States ⁶		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Laboratory-confirmed SARS-CoV-2 infection 	• Median age: 61 years in 5-day RDV arm vs. 62 years in	 Open-label study
 Aged ≥12 years 	10-day RDV arm	 Lack of placebo arm
• Pulmonary infiltrates and $\text{SpO}_2 \leq 94\%$ on room air or receipt	• 60% men in 5-day RDV arm; 68% men in 10-day RDV arm	Baseline imbalances in clinical
of supplemental oxygen	Oxygen requirements at baseline for 5-day RDV arm and	status of patients in 5-day RDV and
Key Exclusion Criteria	10-day RDV arm:	10-day RDV arms
Need for MV or ECMO	• None: 17%, 11%	Interpretation
Multiorgan failure	 Low-flow oxygen: 56%, 54% 	In hospitalized patients with severe
• ALT or AST >5 times ULN	 HFNC oxygen or NIV: 24%, 30% 	COVID-19 who were not receiving MV or ECMO, using RDV for 5 or 10
• Estimated CrCl <50 mL/min	• MV or ECMO: 2%, 5%	days had similar clinical benefits.
Interventions	 Baseline clinical status worse in 10-day arm than in 5-day arm (P = 0.02) 	
• RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily for 4 days (n = 200)	Primary Outcome	
• RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily for	 After adjusting for baseline clinical status: 	
9 days (n = 197)	• Proportion with clinical improvement at Day 14: 65% in	
Primary Endpoint	5-day RDV arm vs. 54% in 10-day RDV arm ($P = 0.14$)	
Clinical status at Day 14, as measured by an OS		

Methods	Results	Limitations and Interpretation
<u>PINETREE</u> : Double-Blind, Placebo-Controlled Trial of Remdesivir for 3 Days in Nonhospitalized Patients With COVID-19 Who Were at High Risk of Disease Progression in Denmark, Spain, the United Kingdom, and the United States ⁷		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Laboratory-confirmed SARS-CoV-2 infection ≤4 days from screening 	 Mean age 50 years; 30% aged ≥60 years; 52% men; 80% White, 8% Black 	 Study halted early due to administrative issues.
 Aged ≥12 years 	• 62% with DM; 55% with obesity; 48% with HTN	Vaccinated individuals were
• \geq 1 risk factor for disease progression or aged \geq 60 years	Median duration of symptoms before first infusion: 5 days	excluded.
 Symptom onset ≤7 days from randomization 	(IQR 3–6 days)	Interpretation
 ≥1 ongoing COVID-19 symptom 	 Median time from RT-PCR confirmation: 2 days (IQR 1–4 days) 	 3 consecutive days of IV RDV resulted in an 87% relative
Key Exclusion Criteria	Primary Outcomes	reduction in the risk of
COVID-19 vaccination		hospitalization or death when
Receipt of supplemental oxygenPrevious hospitalization or treatment for COVID-19	 COVID-19-related hospitalization or death from any cause by Day 28: 2 (0.7%) in RDV arm vs. 15 (5.3%) in placebo arm (HR 0.13; 95% CI, 0.03–0.59; P = 0.008) 	compared to placebo.
Interventions	• Occurrence of AEs: 42% in RDV arm vs. 46% in placebo	
• RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily on	arm	
Days 2 and 3 (n = 279)	Secondary Outcome	
• Placebo (n = 283)	COVID-19-related, medically attended visit or death from	
Primary Endpoints	any cause by Day 28: 4 (1.6%) in RDV arm vs. 21 (8.3%) in	
 COVID-19-related hospitalization or death from any cause by Day 28 	placebo arm (HR 0.19; 95% Cl, 0.07–0.56)	
Occurrence of AEs		
Key Secondary Endpoint		
 COVID-19-related, medically attended visit or death from any cause by Day 28 		

Key: AE = adverse event; ALT = alanine transaminase; AST = aspartate aminotransferase; AZM = azithromycin; CQ = chloroquine; CrCl = creatinine clearance; DM = diabetes mellitus; ECMO = extracorporeal membrane oxygenation; eGFR = estimated glomerular filtration rate; HCQ = hydroxychloroquine; HFNC = high-flow nasal cannula; HTN = hypertension; IV = intravenous; IL = interleukin; LOS = length of stay; LPV/RTV = lopinavir/ritonavir; MV = mechanical ventilation; NIV = noninvasive ventilation; OS = ordinal scale; the Panel = the COVID-19 Treatment Guidelines Panel; RCT = randomized controlled trial; RDV = remdesivir; RT-PCR = reverse transcription polymerase chain reaction; SAE = serious adverse event; SCr = serum creatinine; SOC = standard of care; SpO₂ = oxygen saturation; ULN = upper limit of normal; WHO = World Health Organization

References

- 1. Beigel JH, Tomashek KM, Dodd LE, et al. Remdesivir for the treatment of COVID-19—final report. *N Engl J Med.* 2020;383(19):1813-1826. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32445440</u>.
- 2. Ali K, Azher T, Baqi M, et al. Remdesivir for the treatment of patients in hospital with COVID-19 in Canada: a randomized controlled trial. *CMAJ*. 2022;194(7):E242-E251. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35045989</u>.
- 3. Ader F, Bouscambert-Duchamp M, Hites M, et al. Remdesivir plus standard of care versus standard of care alone for the treatment of patients admitted to hospital with COVID-19 (DisCoVeRy): a Phase 3, randomised, controlled, open-label trial. *Lancet Infect Dis*. 2022;22(2):209-221. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34534511.
- 4. WHO Solidarity Trial Consortium. Remdesivir and three other drugs for hospitalised patients with COVID-19: final results of the WHO Solidarity randomised trial and updated meta-analyses. *Lancet*. 2022;399(10339):1941-1953. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35512728.
- 5. Spinner CD, Gottlieb RL, Criner GJ, et al. Effect of remdesivir vs standard care on clinical status at 11 days in patients with moderate COVID-19: a randomized clinical trial. *JAMA*. 2020;324(11):1048-1057. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32821939.
- 6. Goldman JD, Lye DCB, Hui DS, et al. Remdesivir for 5 or 10 days in patients with severe COVID-19. *N Engl J Med*. 2020:383(19):1827-1837. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32459919</u>.
- 7. Gottlieb RL, Vaca CE, Paredes R, et al. Early remdesivir to prevent progression to severe COVID-19 in outpatients. *N Engl J Med.* 2022;386(4):305-315. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34937145.

Ritonavir-Boosted Nirmatrelvir (Paxlovid)

Last Updated: November 2, 2023

Nirmatrelvir is an oral protease inhibitor that is active against M^{PRO}, a viral protease that plays an essential role in viral replication by cleaving the 2 viral polyproteins.¹ It has demonstrated antiviral activity against all coronaviruses that are known to infect humans.² Nirmatrelvir is packaged with ritonavir (as Paxlovid), a strong cytochrome P450 (CYP) 3A4 inhibitor and pharmacokinetic boosting agent that has been used to boost HIV protease inhibitors. Coadministration of ritonavir is required to increase nirmatrelvir concentrations to the target therapeutic range.

Ritonavir-boosted nirmatrelvir is approved by the Food and Drug Administration (FDA) for the treatment of mild to moderate COVID-19 in adults who are at high risk of progressing to severe COVID-19.³

Beginning November 1, 2023, distribution of Emergency Use Authorization (EUA)-labeled ritonavirboosted nirmatrelvir by the U.S. government will transition to distribution of commercially available, FDA-approved ritonavir-boosted nirmatrelvir by Pfizer. There will be a period of time during which both the EUA-labeled and FDA-approved packaged products will be available for use. For more information on the transition process, please refer to the <u>COVID-19 Therapeutics Commercialization</u> <u>Transition Guide</u>.

The EUA for ritonavir-boosted nirmatrelvir will continue to authorize the use of the EUA-labeled product for the treatment of adolescents aged 12 to 17 years and weighing \geq 40 kg who are at high risk of progressing to severe COVID-19.

Recommendations

- The COVID-19 Treatment Guidelines Panel (the Panel) recommends using **nirmatrelvir 300 mg with ritonavir 100 mg (Paxlovid)** orally (PO) twice daily for 5 days in nonhospitalized adults with mild to moderate COVID-19 who are at high risk of disease progression (AIIa). Treatment should be initiated as soon as possible and within 5 days of symptom onset. For information on medical conditions that confer high risk, see the Centers for Disease Control and Prevention webpage <u>People With Certain Medical Conditions</u>.
- Ritonavir-boosted nirmatrelvir is available through an FDA EUA for the treatment of mild to moderate COVID-19 in nonhospitalized adolescents aged 12 to 17 years and weighing ≥40 kg.⁴ For recommendations on using ritonavir-boosted nirmatrelvir in nonhospitalized children with COVID-19, see <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u>.
- There are no data from randomized clinical trials of ritonavir-boosted nirmatrelvir in hospitalized patients.
- For more information on ritonavir-boosted nirmatrelvir, see Table 4e.
- For a discussion of the treatment of prolonged, symptomatic COVID-19 in patients with evidence of ongoing SARS-CoV-2 replication, see the section titled Patients Who Are Immunocompromised and Have Prolonged COVID-19 Symptoms and Evidence of Ongoing Viral Replication below.

Drug-Drug Interactions

The FDA prescribing information for ritonavir-boosted nirmatrelvir includes a boxed warning about significant drug-drug interactions between ritonavir-boosted nirmatrelvir and other medications. These

interactions are primarily caused by the ritonavir component of the combination. Ritonavir, a strong CYP3A4 inhibitor and a P-glycoprotein inhibitor, may increase the blood concentration of certain concomitant medications and increase the potential for serious drug toxicities. Before prescribing ritonavir-boosted nirmatrelvir, clinicians should carefully review the patient's concomitant medications, including over-the-counter medications, herbal supplements, and recreational drugs, to evaluate potential drug-drug interactions. Clinicians should consider both the potential benefits of treatment with ritonavir-boosted nirmatrelvir and the potential risks related to drug-drug interactions. Many drug-drug interactions between ritonavir-boosted nirmatrelvir and concomitant medications **can be safely managed** (e.g., with certain statins, calcium channel blockers, or direct oral anticoagulants). For the Panel's recommendations on preferred and alternative antiviral therapies for outpatients with COVID-19, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>. Clinicians should be aware that the drug-drug interaction potential of ritonavir-boosted nirmatrelvir may change if it is used for extended durations.

The following resources provide information on identifying and managing drug-drug interactions.

- Quick reference lists:
 - Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications. Box 1 lists select outpatient medications that are not expected to have clinically relevant interactions with ritonavir-boosted nirmatrelvir. Box 2 lists select outpatient medications that have clinically relevant drug-drug interactions with ritonavir-boosted nirmatrelvir.
- Web-based drug-drug interaction checker:
 - <u>The Liverpool COVID-19 Drug Interactions website</u>
- Tables with guidance on managing specific drug-drug interactions:
 - The <u>University of Waterloo/University of Toronto drug interaction guide</u>
 - The FDA prescribing information for ritonavir-boosted nirmatrelvir

Rationale

The EPIC-HR trial enrolled nonhospitalized adults with mild to moderate COVID-19 who were not vaccinated and who were at high risk of progressing to severe disease. The trial demonstrated that starting ritonavir-boosted nirmatrelvir within 5 days of symptom onset in these patients reduced the risk of hospitalization or death through Day 28 by 89% compared to placebo.⁵ This efficacy is comparable to remdesivir (87% relative reduction)⁶ and greater than the efficacy reported for molnupiravir (31% relative reduction).⁷ However, these agents have not been directly compared in clinical trials.

Although ritonavir-boosted nirmatrelvir demonstrated a clinical benefit during the EPIC-HR trial, the benefits in unvaccinated people who are at low risk of progression to severe disease or in vaccinated people who are at high risk of progression to severe disease are unclear. The EPIC-SR trial, which included both of these populations, found that ritonavir-boosted nirmatrelvir did not reduce the duration of symptoms and did not have a statistically significant effect on the risk of hospitalization or death compared to placebo, although the event rates were low.⁸ Some observational studies have shown a benefit of ritonavir-boosted nirmatrelvir in vaccinated individuals who were at high risk of progressing to severe COVID-19.⁹⁻¹² However, observational studies have inherent limitations. In particular, the results of these studies may be affected by residual confounding. For information on treatment considerations for vaccinated individuals, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>.

Because of the potential for significant drug-drug interactions with concomitant medications, this regimen may not be the optimal choice for all patients. See <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications</u> for more information.

COVID-19 Treatment Guidelines

Patients Who Are Immunocompromised and Have Prolonged COVID-19 Symptoms and Evidence of Ongoing Viral Replication

For patients who are immunocompromised and have prolonged COVID-19 symptoms and evidence of ongoing viral replication (e.g., those with a low cycle threshold value, as measured by a reverse transcription polymerase chain reaction result or with a positive rapid antigen test result) despite receiving a course of antiviral therapy, the optimal management is unknown. Case reports and case series have documented the treatment of these patients with additional antiviral treatments, prolonged courses of antiviral treatments, high-titer COVID-19 convalescent plasma, or combination therapy.¹³⁻ ¹⁷ For information on potential treatment options, see <u>Special Considerations in People Who Are Immunocompromised</u> and <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>.

Viral Rebound and Symptom Recurrence

Observational studies and the EPIC-HR trial have described SARS-CoV-2 viral rebound and the recurrence of COVID-19 symptoms in some patients who have completed treatment with ritonavir-boosted nirmatrelvir.¹⁸⁻²¹ The frequency, mechanism, and clinical implications of these events are unclear. Viral rebound and the recurrence of COVID-19 symptoms can also occur in the absence of treatment with ritonavir-boosted nirmatrelvir.^{22,23}

The EPIC-HR trial demonstrated a clinical benefit of ritonavir-boosted nirmatrelvir in patients who were not vaccinated and who were at high risk of progressing to severe COVID-19. To date, the recurrence of COVID-19 symptoms following the use of ritonavir-boosted nirmatrelvir has not been associated with progression to severe COVID-19. Therefore, concerns about the recurrence of symptoms should not be a reason to avoid using ritonavir-boosted nirmatrelvir.^{22,24,25}

There are insufficient data on the efficacy of administering a second course of ritonavir-boosted nirmatrelvir to treat viral rebound or symptom recurrence. There are also insufficient data on whether a longer course of antiviral therapy will prevent viral rebound or symptom recurrence.

SARS-CoV-2 Resistance

Viral mutations that lead to substantial resistance to nirmatrelvir have been selected for in in vitro studies; the fitness of these mutations is unclear. Surveillance for the emergence of significant resistance to nirmatrelvir is critical, particularly in patients who are severely immunocompromised and who experience prolonged replication of SARS-CoV-2.

Additional Considerations

- Nirmatrelvir must be administered with ritonavir to achieve sufficient therapeutic plasma concentrations.
- Patients should complete the 5-day treatment course of ritonavir-boosted nirmatrelvir because there are concerns that a shorter treatment course may be less effective or may lead to the emergence of drug resistance.
- If a patient requires hospitalization after starting treatment, the full 5-day treatment course of ritonavir-boosted nirmatrelvir should be completed unless there are drug-drug interactions that preclude its use.
- There are very limited data on combining ritonavir-boosted nirmatrelvir with other antiviral therapies to treat nonhospitalized patients with COVID-19. Clinical trials are needed to determine whether combination therapy has a role in the treatment of COVID-19.
- The FDA prescribing information for ritonavir-boosted nirmatrelvir advise against crushing

COVID-19 Treatment Guidelines

nirmatrelvir and ritonavir tablets. However, some data indicate that the tablets can be split or crushed if necessary.²⁶

Monitoring and Adverse Effects

The most common adverse effects of ritonavir-boosted nirmatrelvir are dysgeusia, diarrhea, hypertension, and myalgia. Anaphylaxis, serious skin reactions, and other hypersensitivity reactions have also been reported.

There is no need to check a patient's renal function prior to prescribing ritonavir-boosted nirmatrelvir unless the patient is suspected to have moderate to severe renal impairment (i.e., those with an estimated glomerular filtration rate [eGFR] of <60 mL/min). For these patients, clinicians may consider checking the patient's renal function to inform the dosing of ritonavir-boosted nirmatrelvir. The dose should be reduced to nirmatrelvir 150 mg with ritonavir 100 mg twice daily in patients with moderate renal impairment (i.e., those with an eGFR of \geq 30 to <60 mL/min).

The FDA prescribing information states that ritonavir-boosted nirmatrelvir is not recommended for patients with an eGFR of <30 mL/min until more data are available to establish appropriate dosing.³ Additional information is available in the initial FDA Center for Drug Evaluation and Research review for the EUA of ritonavir-boosted nirmatrelvir.¹⁸ There is limited clinical experience with the use of ritonavir-boosted nirmatrelvir in patients with eGFR of <30 mL/min and in those who require hemodialysis.^{27,28} Based on limited data, some groups have proposed dosing adjustments for ritonavir-boosted nirmatrelvir in these patients.²⁹⁻³¹ A clinical trial (ClinicalTrials.gov Identifier <u>NCT05487040</u>) that will evaluate the use of ritonavir-boosted nirmatrelvir in patients with COVID-19 and severe renal impairment is currently underway.

Ritonavir-boosted nirmatrelvir **is not recommended** for patients with known or suspected severe hepatic impairment (i.e., Child-Pugh Class C), and it should be used with caution in patients with pre-existing liver diseases, liver enzyme abnormalities, or hepatitis. No pharmacokinetic or safety data are available for this patient population.

Considerations in Pregnant and Lactating People

See <u>Pregnancy</u>, <u>Lactation</u>, <u>and COVID-19 Therapeutics</u> for the Panel's guidance on the use of ritonavirboosted nirmatrelvir during pregnancy and lactation.

Considerations in Children

Ritonavir-boosted nirmatrelvir is available through an FDA EUA for the treatment of mild to moderate COVID-19 in nonhospitalized adolescents aged 12 to 17 years and weighing \geq 40 kg. For information on using ritonavir-boosted nirmatrelvir in pediatric patients, see <u>Special Considerations in Children</u>, <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u>, and <u>Therapeutic Management of Hospitalized Children With COVID-19</u>.

Clinical Data

The EPIC-HR study was a multinational randomized trial that compared the use of ritonavir-boosted nirmatrelvir PO twice daily for 5 days to placebo in nonhospitalized patients aged \geq 18 years with mild to moderate COVID-19 who were at high risk of clinical progression. Eligible patients were randomized within 5 days of symptom onset, were not vaccinated against COVID-19, and had at least 1 risk factor for progression to severe disease.⁵ Patients were excluded if they used medications that were either highly dependent upon CYP3A4 for clearance or strong inducers of CYP3A4.

A total of 2,246 patients enrolled in the trial. The mean age was 46 years, 51% of the patients were men, and 72% were White. Forty-seven percent of the patients tested negative for SARS-CoV-2 antibodies, and 66% started study treatment within 3 days of symptom onset.

Patients who were randomized within 3 days of symptom onset (n = 1,379) were included in the modified intention-to-treat (mITT) analysis. COVID-19-related hospitalizations or all-cause deaths occurred by Day 28 in 5 of 697 patients (0.72%) in the ritonavir-boosted nirmatrelvir arm and in 44 of 682 patients (6.5%) in the placebo arm. Among the 2,085 patients who were randomized within 5 days of symptom onset (mITT1 analysis), COVID-19–related hospitalizations and all-cause deaths occurred in 8 of 1,039 patients (0.77%) in the ritonavir-boosted nirmatrelvir arm and in 66 of 1,046 patients (6.3%) in the placebo arm (89% relative risk reduction; 5.6% estimated absolute reduction; 95% CI, 7.2% to 4.0%; P < 0.001). There were no deaths in the ritonavir-boosted nirmatrelvir arm, and 13 deaths occurred in the placebo arm.

A total of 2,224 patients who received at least 1 dose of either ritonavir-boosted nirmatrelvir or placebo were included in the EPIC-HR safety analysis set. Among these patients, dysgeusia and diarrhea occurred more frequently in ritonavir-boosted nirmatrelvir recipients than in placebo recipients (6% vs. 0.3% and 3% vs. 2%, respectively). Fewer ritonavir-boosted nirmatrelvir recipients discontinued the study drug due to an adverse event than placebo recipients (2% vs. 4%).

References

- Pillaiyar T, Manickam M, Namasivayam V, Hayashi Y, Jung SH. An overview of severe acute respiratory syndrome-coronavirus (SARS-CoV) 3CL protease inhibitors: peptidomimetics and small molecule chemotherapy. *J Med Chem.* 2016;59(14):6595-6628. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/26878082</u>.
- Owen DR, Allerton CMN, Anderson AS, et al. An oral SARS-CoV-2 MPRO inhibitor clinical candidate for the treatment of COVID-19. *Science*. 2021;374(6575):1586-1593. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34726479</u>.
- 3. Ritonavir-boosed nirmatrelvir (Paxlovid) [package insert]. Food and Drug Administration. 2023. Available at: <u>https://www.accessdata.fda.gov/drugsatfda_docs/label/2023/217188s000lbl.pdf</u>.
- 4. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Paxlovid. 2023. Available at: <u>https://www.fda.gov/media/155050/download</u>.
- Hammond J, Leister-Tebbe H, Gardner A, et al. Oral nirmatrelvir for high-risk, nonhospitalized adults with COVID-19. N Engl J Med. 2022;386(15):1397-1408. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35172054</u>.
- Gottlieb RL, Vaca CE, Paredes R, et al. Early remdesivir to prevent progression to severe COVID-19 in outpatients. *N Engl J Med.* 2022;386(4):305-315. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34937145</u>.
- Jayk Bernal A, Gomes da Silva MM, Musungaie DB, et al. Molnupiravir for oral treatment of COVID-19 in nonhospitalized patients. *N Engl J Med*. 2022;386(6):509-520. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34914868</u>.
- Pfizer. Pfizer reports additional data on PAXLOVID supporting upcoming new drug application submission to U.S. FDA. 2022. Available at: <u>https://www.pfizer.com/news/press-release/press-release-detail/pfizer-reports-additional-data-paxlovidtm-supporting</u>. Accessed July 20, 2023.
- 9. Dryden-Peterson S, Kim A, Kim AY, et al. Nirmatrelvir plus ritonavir for early COVID-19 in a large U.S. health system: a population-based cohort study. *Ann Intern Med.* 2023;176(1):77-84. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36508742</u>.
- 10. Arbel R, Wolff Sagy Y, Hoshen M, et al. Nirmatrelvir use and severe COVID-19 outcomes during

the Omicron surge. *N Engl J Med.* 2022;387(9):790-798. Available at: <u>https://www.ncbi.nlm.nih.gov/</u>pubmed/36001529.

- Ganatra S, Dani SS, Ahmad J, et al. Oral nirmatrelvir and ritonavir in nonhospitalized vaccinated patients with coronavirus disease 2019. *Clin Infect Dis*. 2023;76(4):563-572. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35986628.
- Shah MM, Joyce B, Plumb ID, et al. Paxlovid associated with decreased hospitalization rate among adults with COVID-19—United States, April–September 2022. *MMWR Morb Mortal Wkly Rep.* 2022;71(48):1531-1537. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36454693</u>.
- Huygens S, Gharbharan A, Serroukh Y, et al. High-titer convalescent plasma plus nirmatrelvir/ritonavir treatment for non-resolving COVID-19 in six immunocompromised patients. *J Antimicrob Chemother*. 2023;78(7):1644-1648. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37248664</u>.
- 14. Brosh-Nissimov T, Ma'aravi N, Leshin-Carmel D, et al. Combination treatment of persistent COVID-19 in immunocompromised patients with remdesivir, nirmatrelvir/ritonavir and tixagevimab/cilgavimab. J Microbiol Immunol Infect. 2023;Published online ahead of print. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/37805361/</u>.
- 15. Mikulska M, Sepulcri C, Dentone C, et al. Triple combination therapy with two antivirals and monoclonal antibodies for persistent or relapsed SARS-CoV-2 infection in immunocompromised patients. *Clin Infect Dis*. 2023;77(2):280-286. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36976301</u>.
- Graziani L, Gori L, Manciulli T, et al. Successful use of nirmatrelvir/ritonavir in immunocompromised patients with persistent and/or relapsing COVID-19. *J Antimicrob Chemother*. 2023;78(2):555-558. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36544352</u>.
- 17. Trottier CA, Wong B, Kohli R, et al. Dual antiviral therapy for persistent coronavirus disease 2019 and associated organizing pneumonia in an immunocompromised host. *Clin Infect Dis.* 2023;76(5):923-925. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36281907</u>.
- Food and Drug Administration. Emergency Use Authorization (EUA) for Paxlovid (nirmatrelvir tablets co-packaged with ritonavir tablets): Center for Drug Evaluation and Research (CDER) review. 2021. Available at: <u>https://www.fda.gov/media/155194/download</u>.
- Charness ME, Gupta K, Stack G, et al. Rebound of SARS-CoV-2 infection after nirmatrelvir-ritonavir treatment. *N Engl J Med.* 2022;387(11):1045-1047. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36069968.
- 20. Boucau J, Uddin R, Marino C, et al. Characterization of virologic rebound following nirmatrelvir-ritonavir treatment for coronavirus disease 2019 (COVID-19). *Clin Infect Dis.* 2023;76(3):e526-e529. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35737946</u>.
- 21. Anderson AS, Caubel P, Rusnak JM, EPIC-HR Trial Investigators. Nirmatrelvir-ritonavir and viral load rebound in COVID-19. *N Engl J Med.* 2022;387(11):1047-1049. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36069818</u>.
- 22. Soares H, Baniecki ML, Cardin R, et al. Viral load rebound in placebo and nirmatrelvir-ritonavir treated COVID-19 patients is not associated with recurrence of severe disease or mutations. *Res Sq.* 2022;Preprint. Available at: <u>https://www.researchsquare.com/article/rs-1720472/v1</u>.
- 23. Deo R, Choudhary MC, Moser C, et al. Symptom and viral rebound in untreated SARS-CoV-2 infection. *Ann Intern Med.* 2023;176(3):348-354. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36802755</u>.
- 24. Ranganath N, O'Horo JC, Challener DW, et al. Rebound phenomenon after nirmatrelvir/ritonavir treatment of coronavirus disease 2019 (COVID-19) in high-risk persons. *Clin Infect Dis*. 2023;76(3):e537-e539. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35698452.
- 25. Centers for Disease Control and Prevention. COVID-19 rebound after Paxlovid treatment. 2022. Available at: <u>https://emergency.cdc.gov/han/2022/han00467.asp</u>. Accessed July 13, 2023.
- 26. BC COVID Therapeutics Committee COVID Therapy Review and Advisory Working Group. Therapeutic brief: crushing nirmatrelvir/ritonavir (Paxlovid). 2022. Available at: <u>http://www.bccdc.ca/Health-</u>

COVID-19 Treatment Guidelines

Professionals-Site/Documents/COVID-treatment/Crushing_Paxlovid.pdf.

- 27. Hiremath S, Blake PG, Yeung A, et al. Early experience with modified dose nirmatrelvir/ritonavir in dialysis patients with coronavirus disease 2019. *Clin J Am Soc Nephrol*. 2023;18(4):485-490. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36723285</u>.
- Chan GCK, Lui GCY, Wong CNS, et al. Safety profile and clinical and virological outcomes of nirmatrelvirritonavir treatment in patients with advanced chronic kidney disease and coronavirus disease 2019 (COVID-19). *Clin Infect Dis.* 2023;Published online ahead of print. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37531093</u>.
- 29. University of Liverpool. Prescribing resources. 2022. Available at: <u>https://covid19-druginteractions.org/prescribing_resources</u>. Accessed July 13, 2023.
- 30. Ontario Health. COVID-19 supplemental clinical guidance #4: nirmatrelvir/ritonavir (Paxlovid) use in patients with advanced chronic kidney disease and patients on dialysis with COVID-19. 2022. Available at: <u>https://www.ontariohealth.ca/sites/ontariohealth/files/2022-04/PaxlovidClinicalGuide.pdf</u>.
- Hiremath S, McGuinty M, Argyropoulos C, et al. Prescribing nirmatrelvir/ritonavir for COVID-19 in advanced CKD. *Clin J Am Soc Nephrol*. 2022;17(8):1247-1250. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/35680135</u>.

Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications

Last Updated: November 2, 2023

Ritonavir, a strong cytochrome P450 (CYP) 3A4 inhibitor and a P-glycoprotein (P-gp) inhibitor, is coadministered with nirmatrelvir to increase the blood concentration of nirmatrelvir, thereby making it effective against SARS-CoV-2. Ritonavir may also increase blood concentrations of certain concomitant medications. The Food and Drug Administration (FDA) <u>prescribing information</u> includes a boxed warning about significant drug-drug interactions between ritonavir-boosted nirmatrelvir (Paxlovid) and other medications.

Before prescribing ritonavir-boosted nirmatrelvir to treat patients with mild to moderate COVID-19, carefully review the patient's concomitant medications, including over-the-counter medicines, herbal supplements, and recreational drugs. Clinicians should consider the potential benefits of treatment with ritonavir-boosted nirmatrelvir, the potential risks of drug-drug interactions, and whether any risks related to drug-drug interactions can be safely managed. Clinicians should be aware that many commonly used medications can be safely coadministered with ritonavir-boosted nirmatrelvir despite its drug-drug interaction potential. Box 1 includes commonly prescribed medications that are not expected to have clinically relevant interactions with ritonavir-boosted nirmatrelvir.

Because ritonavir-boosted nirmatrelvir is the only highly effective oral antiviral for the treatment of COVID-19, drug interactions that can be safely managed should not preclude the use of this medication.

Box 1. Select Outpatient Medications Not Expected to Have Clinically Relevant Interactions With Ritonavir-Boosted Nirmatrelvir (Paxlovid)

This list is primarily based on the most common medication searches by U.S. users on the Liverpool COVID-19 Drug Interactions website.

Medications Without Clinically Relevant Interactions						
	These medications may be coadministered without dose adjustment and without increased monitoring. This list is not inclusive of all noninteracting medications within each drug category.					
Acid Reducers • Famotidine • Omeprazole • Pantoprazole Allergy • Cetirizine • Diphenhydramine • Fexofenadine • Loratadine Anti-Infectives • Azithromycin • Cidofovir • Hydroxychloroquine • Tecovirimat • Valacyclovir	Cardiovascular • Aspirin • Atenolol • Carvedilol • Furosemide • Hydrochlorothiazide • Irbesartan • Isosorbide dinitrate • Lisinopril • Losartan • Metoprolol • Prasugrel Diabetes • Empagliflozin • Insulin • Metformin • Pioglitazone	Immunosuppressants Abrocitinib Baricitinib Methotrexate Mycophenolate Prednisone Lipid-Modifiers Ezetimibe Pitavastatin Pravastatin Migraine Frovatriptan Naratriptan Sumatriptan Zavegepant Neuropsychiatric Amitriptyline Bupropion 	Neuropsychiatric, cont'd Citalopram Duloxetine Escitalopram Fluoxetine Gabapentin Lorazepam Nortriptyline Olanzapine Paroxetine Sertraline Venlafaxine Pain Acetaminophen Aspirin Codeine Ibuprofen Meloxicam Naproxen	 Respiratory Corticosteroids (inhaled/nasal) Formoterol Montelukast Miscellaneous Allopurinol Contraceptives (PO)^a Cyclobenzaprine Donepezil Enoxaparin Finasteride Levothyroxine Most mAb products^b Ondansetron 		

COVID-19 Treatment Guidelines

Medications Without Clinically Relevant Interactions, continued

- ^a Coadministering contraceptive products that contain ethinyl estradiol with ritonavir-boosted nirmatrelvir may result in lower ethinyl estradiol concentrations. The FDA <u>prescribing information</u> for ritonavir-boosted nirmatrelvir suggests that individuals who use these types of contraceptive products should consider using an additional nonhormonal contraceptive method. However, the lower ethinyl estradiol concentrations are not expected to be clinically significant during the 5 days of therapy. The progestin concentration of a combined hormonal contraceptive is expected to remain similar or increase with coadministration, which would maintain the effectiveness of the PO contraceptive.
- ^b Ritonavir-boosted nirmatrelvir interacts with certain conjugated mAbs, such as ado-trastuzumab emtansine, mirvetuximab soravtansine, brentuximab vedotin, enfortumab vedotin, polatuzumab vedotin, and tisotumab vedotin.
 Before coadministering ritonavir-boosted nirmatrelvir and any of these conjugated mAbs, refer to the drug's FDA prescribing information and consult with the patient's specialist providers as needed.

Key: FDA = Food and Drug Administration; mAb = monoclonal antibody; PO = oral

Medications That Have Clinically Relevant Drug-Drug Interactions With Ritonavir-Boosted Nirmatrelvir (Paxlovid)

Clinicians should be aware that, in some cases, drug-drug interactions with ritonavir-boosted nirmatrelvir may lead to serious or life-threatening drug toxicities. The recommended treatment course of ritonavir-boosted nirmatrelvir for COVID-19 is 5 days. CYP3A4 inhibition occurs rapidly, with maximum inhibition occurring within 48 hours of ritonavir initiation.¹ After treatment is completed and ritonavir is discontinued, 70% to 90% of CYP3A4 inhibition resolves within 2 to 3 days.² The time to resolution of inhibition varies based on factors such as the patient's age; therefore, resolution may take longer in some individuals, such as in adults of advanced age.

Ritonavir is also an inhibitor of CYP2D6, P-gp, and organic anion transporting polypeptide (OATP) 1B1. When used for longer durations or chronically, ritonavir may induce CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, and uridine diphosphate-glucuronyltransferase (UGT). See below for more information.

Nirmatrelvir and ritonavir are CYP3A4 substrates. Ritonavir-boosted nirmatrelvir should not be given within 2 weeks of administering a strong CYP3A4 inducer (e.g., St. John's wort, rifampin). Ritonavir-boosted nirmatrelvir is **contraindicated** in this setting because the delayed offset of enzyme induction may reduce the concentrations of nirmatrelvir and ritonavir, rendering the treatment ineffective against SARS-CoV-2. An alternative treatment for COVID-19 should be prescribed.

Identifying Drug-Drug Interactions

Consult the following resources for information on identifying and managing drug-drug interactions.

- Quick reference lists:
 - Box 1 above lists select outpatient medications that are not expected to have clinically relevant interactions with ritonavir-boosted nirmatrelvir.
 - Box 2 below lists select outpatient medications that have clinically relevant drug-drug interactions with ritonavir-boosted nirmatrelvir.
- Web-based drug-drug interaction checker:
 - The Liverpool COVID-19 Drug Interactions website
- Tables with guidance on managing specific drug-drug interactions:
 - The University of Waterloo/University of Toronto drug interaction guide
 - The FDA prescribing information for ritonavir-boosted nirmatrelvir

Management Strategies for Drug-Drug Interactions

Consider the magnitude and significance of the potential drug-drug interaction when choosing management strategies for patients who will be receiving ritonavir-boosted nirmatrelvir. Potential strategies include:

- Increasing monitoring for potential adverse events to the concomitant medication.
- Adjusting the dose of the concomitant medication.
- Temporarily withholding the concomitant medication.
- Using an alternative to the concomitant medication.
- Using alternative COVID-19 therapies (see <u>Therapeutic Management of Nonhospitalized Adults</u> <u>With COVID-19</u>).

Use the chosen strategy for the 5-day duration of ritonavir-boosted nirmatrelvir treatment and for at least 2 to 3 days after treatment completion. The strategy may need to continue for a longer duration if ritonavir-boosted nirmatrelvir is initiated in an adult of advanced age or if the interacting medication has a long half-life.

Consider consulting with an expert (e.g., a pharmacist or the patient's specialist providers) when treating patients who are receiving highly specialized therapies or drugs that are prone to concentration-dependent toxicities, such as certain anticonvulsant, anticoagulant, immunosuppressant, antiarrhythmic, chemotherapeutic, and neuropsychiatric drugs.

The decision to prescribe ritonavir-boosted nirmatrelvir to patients who are receiving calcineurin and mammalian target of rapamycin inhibitors should always be made in consultation with the patient's specialist providers. Among reports submitted to the FDA Adverse Events Reporting System, the most commonly reported concomitant medications resulting in serious adverse reactions, including fatal events, were calcineurin inhibitors (e.g., tacrolimus).³ Ritonavir-boosted nirmatrelvir may be prescribed to select patients who are receiving these medications if an expert in managing the interaction is available and close therapeutic drug monitoring is logistically feasible. Otherwise, an alternative therapy for COVID-19 should be considered. See the <u>American Society of Transplantation statement</u> for more information.

Interactions between ritonavir-boosted nirmatrelvir and chemotherapeutic agents should also be managed in consultation with the patient's specialist providers. For guidance on managing these interactions, refer to the FDA prescribing information for ritonavir-boosted nirmatrelvir and the prescribing information for the chemotherapeutic agent. The <u>University Health Network/Kingston</u> <u>Health Sciences Centre</u> provides an additional resource for evaluating drug-drug interactions between ritonavir-boosted nirmatrelvir and chemotherapeutic agents.

Patients should be counseled about ritonavir-boosted nirmatrelvir's drug-drug interaction potential and the signs and symptoms of potential adverse effects. If ritonavir-boosted nirmatrelvir is prescribed to patients who take certain recreational drugs, those patients will require counseling and careful monitoring for adverse effects.

Box 2. Select Outpatient Medications That Have Clinically Relevant Drug-Drug Interactions With Ritonavir-Boosted Nirmatrelvir (Paxlovid)

The guidance in Box 2 is based on the drug-drug interaction potential of the FDA-approved 5-day course of ritonavir-boosted nirmatrelvir.

Not all medications that may interact with ritonavir-boosted nirmatrelvir are included in Box 2. Deviation from the recommended strategies may be appropriate in certain clinical scenarios.

	Prescribe Alte	ernative COVID-19 Therapy		
For these medications, management strategies are not possible or feasible, or the risks outweigh the potential benefits.				
Anticonvulsants • Carbamazepine • Phenobarbital • Phenytoin • Primidone Anti-Infectives • Glecaprevir/pibrentasvir • Rifampin • Rifapentine Immunosuppressants • Voclosporin	Cardiovascular • Amiodarone • Clopidogrel ^{a,b} • Disopyramide • Dofetilide • Dronedarone • Eplerenone • Flecainide • Ivabradine • Quinidine	Neuropsychiatric • Clozapine • Lurasidone • Midazolam (PO) • Pimozide Pulmonary Hypertension ^c • Sildenafil • Tadalafil • Vardenafil	Miscellaneous • Bosentan • Certain chemotherapeutic agents ^d • Ergot derivatives • Lumacaftor/ivacaftor • St. John's wort • Tolvaptan	
Temp	orarily Withhold Concon	nitant Medication, if Clinica	Ily Appropriate	
completion. They may need medication has a long half-li COVID-19 therapy. Anticoagulants • Rivaroxaban ^e Anti-Infectives • Erythromycin BPH • Alfuzosin • Silodosin Cardiovascular • Aliskiren • Ranolazine • Ticagrelor ^b • Vorapaxar	to be withheld for longer if ife. If withholding is not clir Immunosuppressants ^f Everolimus Sirolimus Tacrolimus Lipid-modifiers Atorvastatin ^g Lomitapide Lovastatin ^g Rosuvastatin ^g Simvastatin ^g	Migraine• Eletriptan• Rimegepant• UbrogepantNeuropsychiatric• Daridorexant• Lemborexant• Suvorexant• Triazolam ^h Erectile Dysfunction• Avanafil	 ced age or if the interacting native concomitant medication or Respiratory Salmeterol Miscellaneous Certain chemotherapeutic agents^d Colchicineⁱ Finerenone Flibanserin Naloxegol 	
Adju	st Concomitant Medicat	tion Dose and Monitor for A	dverse Effects	
Drug Interactions website o	r the <u>University of Waterloo</u> se of the concomitant med	lication cannot be adjusted, wi	onsult the Liverpool COVID-19 praction guide for specific dosing thhold the medication (if clinically Neuropsychiatric, cont'd Buspirone Cariprazine Chlordiazepoxide ^h Clobazam ^h Clorazepate ^h Diazepam ^h Estazolam ^h Iloperidone Lumateperone Pimavanserin Quetiapine Trazodone	

COVID-19 Treatment Guidelines

Adjust Concomitant Medication Dose and Monitor for Adverse Effects, continued			
Pain Fentanyl Hydrocodone Oxycodone Pulmonary Hypertension Riociguat 	 Miscellaneous Certain chemotherapeutic agents^d Darifenacin 	Miscellaneous, cont'd • Elexacaftor/ tezacaftor/ivacaftor • Eluxadoline • Ivacaftor	Miscellaneous, cont'dSolifenacinTezacaftor/ivacaftor
	ntinue Concomitant Medicat	tion and Monitor for Adv	verse Effects
patient's risk for AEs. Educa		. Consult the Liverpool CO	lividualized assessment of the <u>/ID-19 Drug Interactions website</u> ng guidance and dose adjustment
Anticoagulants • Warfarin Anti-Infectives • Brincidofovir ^I • Cobicistat- or ritonavir- boosted ARV drugs • Isavuconazole • Posaconazole • Voriconazole	 BPH Doxazosin Terazosin Diabetes Glyburide Cardiovascular Mexiletine Sacubitril Valsartan 	Migraine • Zolmitriptan Neuropsychiatric • Haloperidol • Hydroxyzine • Mirtazapine • Risperidone • Ziprasidone • Zolpidem	Pain Buprenorphine Hydromorphone Methadone Morphine Tramadol Miscellaneous Certain chemotherapeutic agents ^d Certain conjugated mAbs ^m Oxybutynin
 ^a Reduced effectiveness of clopidogrel is likely. It may be acceptable to continue clopidogrel if the benefits of using ritonavir-boosted nirmatrelvir outweigh the risk of reduced clopidogrel effectiveness. ^b For patients at very high risk of thrombosis (e.g., those who received a coronary stent within the past 6 weeks), consider prescribing an alternative antiplatelet (e.g., prasugrel, if clinically appropriate) or an alternative COVID-19 therapy. ^c Some PDE5 inhibitors are used to treat both PAH and erectile dysfunction; however, the doses used to treat PAH are higher than those used for erectile dysfunction. Because of this, and because PDE5 inhibitors are used chronically in patients with PAH, coadministration with ritonavir-boosted nirmatrelvir is contraindicated in these patients. PDE5 inhibitors can be coadministered with ritonavir-boosted nirmatrelvir in patients with erectile dysfunction, though the dose of the PDE5 inhibitor should be adjusted. ^d Ritonavir-boosted nirmatrelvir may increase concentrations of some chemotherapeutic agents, leading to an increased potential for drug toxicities. Some chemotherapeutic agents may decrease the effectiveness of ritonavir-boosted 			
nirmatrelvir. Please refer to the FDA prescribing information for ritonavir-boosted nirmatrelvir and the prescribing information for the chemotherapeutic agent and consult the patient's specialist provider. The <u>University Health Network/</u> <u>Kingston Health Sciences Centre</u> is an additional resource for evaluating drug-drug interactions for chemotherapeutic agents. ^e For patients who are at high risk of arterial or venous thrombosis (e.g., those who had a stroke within the past 3 months with a CHA ₂ DS ₂ -VASc score of 7–9 or a pulmonary embolism within the past month), consult the primary or specialty provider and consider using an alternative anticoagulant (e.g., LMWH) or an alternative COVID-19 therapy. For patients with a lower risk of arterial or venous thrombosis, clinicians may consider administering low-dose aspirin while rivaroxaban is being withheld.			
^f The use of another COVID-19 therapy may need to be considered. These immunosuppressants have significant drug-drug interaction potential with ritonavir, and they should not be used if close monitoring, including therapeutic drug monitoring (i.e., measuring drug concentrations), is not feasible. Consult a patient's specialist providers before coadministering these immunosuppressants with ritonavir-boosted nirmatrelvir. See the <u>American Society of</u> <u>Transplantation statement</u> for more information. ^g Withhold lovastatin and simvastatin for at least 12 hours before initiating ritonavir-boosted nirmatrelvir, during treatment, and for 5 days after treatment completion. Withhold atorvastatin and rosuvastatin at the beginning of treatment with ritonavir-boosted nirmatrelvir and resume after completing the 5-day course. If withholding a statin is not clinically appropriate (e.g., because the patient recently had a myocardial infarction), clinicians can reduce the doses of			

Continue Concomitant Medication and Monitor for Adverse Effects, continued

atorvastatin and rosuvastatin and continue treatment. However, lovastatin and simvastatin should be switched to an alternative statin.

- ^h The guidance on managing drug-drug interactions between certain benzodiazepines and ritonavir-boosted nirmatrelvir can vary significantly between product information resources. Note that abrupt discontinuation or rapid dose reduction of benzodiazepines may precipitate an acute withdrawal reaction.⁴ The risk is greatest for patients who have been using high doses of benzodiazepines over an extended period.
- Do not coadminister this medication with ritonavir-boosted nirmatrelvir in patients with hepatic or renal impairment.
- For medications that are not included on the Liverpool COVID-19 Drug Interactions website or in the University of Waterloo/University of Toronto drug interaction guide, refer to the FDA labels for information on coadministering these medications with ritonavir or other strong CYP3A4 and/or P-gp inhibitors (e.g., ketoconazole).
- Dexamethasone exposure is expected to increase 2.60-fold when dexamethasone is coadministered with ritonavirboosted nirmatrelvir.⁵ Clinicians should weigh the risks and benefits of continuing the patient's normal dose of dexamethasone (while monitoring for AEs) against the risks and benefits of decreasing the dose. Patients who are receiving higher doses of dexamethasone will be at a greater risk of AEs.
- Patients should take ritonavir-boosted nirmatrelvir at least 3 hours after taking brincidofovir.
- ^m Ritonavir-boosted nirmatrelvir interacts with certain conjugated mAbs, such as ado-trastuzumab emtansine, mirvetuximab soravtansine, brentuximab vedotin, enfortumab vedotin, polatuzumab vedotin, and tisotumab vedotin. Before coadministering ritonavir-boosted nirmatrelvir and any of these conjugated mAbs, refer to the drug's FDA prescribing information and consult with the patient's specialist providers as needed.

Key: AE = adverse effect: ARV = antiretroviral: BPH = benign prostatic hyperplasia: $CHA_{2}DS_{2}-VASc = congestive$ heart failure, hypertension, age, diabetes, stroke, vascular disease; CYP = cvtochrome P450; FDA = Food and Drug Administration; LMWH = low-molecular-weight heparin; mAb = monoclonal antibody; PAH = pulmonary arterial hypertension; PDE5 = phosphodiesterase 5; P-gp = P-glycoprotein; PO = oral

Drug-Drug Interaction Considerations When Using Extended Courses of Ritonavir-**Boosted Nirmatrelvir (Paxlovid)**

The guidance in this document is based on the drug-drug interaction potential of the FDA-approved 5-day course of ritonavir-boosted nirmatrelvir.

Longer treatment courses may be utilized in certain cases (see Special Considerations in People Who Are Immunocompromised). Clinicians should be aware that the drug-drug interaction potential of ritonavir may change based on duration of treatment. Clinicians should be aware that:

- Induction properties⁶ may become clinically relevant when ritonavir is used for longer durations (i.e., ≥ 10 days) or chronically (e.g., in people who take HIV protease inhibitors).⁷ For example, induction of CYP2C9 and CYP2C19 may decrease warfarin and voriconazole concentrations, and induction of glucuronidation may decrease lamotrigine or valproic acid concentrations.
- The management strategies listed in Box 2 are based on the drug-drug interaction potential of a 5-day treatment course of ritonavir-boosted nirmatrelvir. These strategies may need to be modified when using extended courses. For example, clinicians may need to decide whether to hold or reduce the dose of corticosteroids instead of continuing them as suggested in Box 2. Clinicians may need to adjust monitoring plans for adverse effects or therapeutic drug monitoring in certain patients (e.g., in those who are receiving tacrolimus). In other cases, the potential risks of holding certain agents (e.g., chemotherapeutic agents or statins in high-risk individuals) for extended periods to allow for safe coadministration of ritonavir-boosted nirmatrelvir may outweigh the potential benefits of treatment.
- After discontinuing longer courses of ritonavir-boosted nirmatrelvir, drug-drug interactions caused COVID-19 Treatment Guidelines

by CYP3A4 inhibition largely resolve within 2 to 3 days.² Drug-drug interactions caused by induction (e.g., CYP2C9, CYP2C19, UGT) resolve gradually and variably.^{8,9}

Clinicians should consult with an expert (e.g., pharmacists and physicians with HIV expertise) when using extended courses of ritonavir-boosted nirmatrelvir. The Liverpool COVID-19 Drug Interactions website also provides guidance for managing drug-drug interactions for extended courses (i.e., ≥ 10 days) of ritonavir-boosted nirmatrelvir.

References

- 1. Katzenmaier S, Markert C, Riedel KD, et al. Determining the time course of CYP3A inhibition by potent reversible and irreversible CYP3A inhibitors using a limited sampling strategy. *Clin Pharmacol Ther*. 2011;90(5):666-73. Available at: https://www.ncbi.nlm.nih.gov/pubmed/21937987.
- Stader F, Khoo S, Stoeckle M, et al. Stopping lopinavir/ritonavir in COVID-19 patients: duration of the drug interacting effect. *J Antimicrob Chemother*. 2020;75(10):3084-3086. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32556272</u>.
- 3. Food and Drug Administration Center for Drug Evaluation and Research. Antimicrobial drugs advisory committee meeting. 2023. Available at: <u>https://www.fda.gov/media/168508/download</u>.
- 4. Food and Drug Administration. FDA requiring Boxed Warning updated to improve safe use of benzodiazepine drug class. 2020. Available at: <u>https://www.fda.gov/media/142368/download</u>.
- Li M, Zhu L, Chen L, Li N, Qi F. Assessment of drug-drug interactions between voriconazole and glucocorticoids. *J Chemother*. 2018;30(5):296-303. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30843777</u>.
- 6. Foisy MM, Yakiwchuk EM, Hughes CA. Induction effects on ritonavir: implications for drug interactions. *Ann Pharmacother*. 2008;42(7):1048-1059. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/18577765</u>.
- 7. University of Liverpool. Evaluating the interaction risk of COVID-19 therapies. 2022. Available at: <u>https://covid19-druginteractions.org/prescribing_resources</u>. Accessed July 20, 2023.
- Ramsden D, Fung C, Hariparsad N, et al. Perspectives from the innovation and quality consortium induction working group on factors impacting clinical drug-drug interactions resulting from induction: focus on cytochrome 3A substrates. *Drug Metab Dispos*. 2019;47(10):1206-1221. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/31439574/</u>.
- Marzolini C, Kuritzkes DR, Marra F, et al. Recommendations for the management of drug-drug interactions between the COVID-19 antiviral nirmatrelvir/ritonavir (Paxlovid) and comedications. *Clin Pharmacol Ther*. 2022;112(6):1191-1200. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35567754</u>.

Molnupiravir

Last Updated: April 20, 2023

Molnupiravir is the oral prodrug of beta-D-N4-hydroxycytidine (NHC), a ribonucleoside that has shown antiviral activity against SARS-CoV-2 in vitro and in some clinical trials.^{1,2} NHC uptake by viral RNA-dependent RNA-polymerases results in viral mutations and lethal mutagenesis.^{3,4} On December 23, 2021, the Food and Drug Administration (FDA) issued an Emergency Use Authorization (EUA) for molnupiravir for the treatment of adults with mild to moderate COVID-19 who are within 5 days of symptom onset, who are at high risk of progressing to severe disease, and for whom alternative antiviral therapies are not accessible or clinically appropriate.^{5,6} Molnupiravir is expected to be active against the Omicron variant and its subvariants.⁶

As a mutagenic ribonucleoside antiviral agent, there is a theoretical risk that molnupiravir will be metabolized by the human host cell and incorporated into the host DNA, leading to mutations. Molnupiravir has been evaluated in 2 in vivo rodent mutagenicity assays. One study produced equivocal results. In the other study, there was no evidence for mutagenicity.⁶ The FDA concluded that, based on the available genotoxicity data and the 5-day duration of treatment, molnupiravir has a low risk for genotoxicity. In addition, there have been concerns about the potential effects of molnupiravir on SARS-CoV-2 mutation rates; the FDA has required that the manufacturer monitor genomic databases for the emergence of SARS-CoV-2 variants.

Recommendations

- The COVID-19 Treatment Guidelines Panel (the Panel) recommends using **molnupiravir 800 mg** orally (PO) twice daily for 5 days as an alternative therapy in nonhospitalized patients aged ≥18 years with mild to moderate COVID-19 who are at high risk of disease progression when ritonavir-boosted nirmatrelvir (Paxlovid) and remdesivir are not available, feasible to use, or clinically appropriate; treatment should be initiated as soon as possible and within 5 days of symptom onset (**CIIa**).
- The Panel **recommends against** the use of **molnupiravir** for the treatment of COVID-19 in pregnant patients unless there are no other options and therapy is clearly indicated (**AIII**). For more details, see Considerations in Pregnancy below.
- People who engage in sexual activity that may result in conception should use effective contraception during and following treatment with molnupiravir. For more details, see Considerations in Sexually Active Individuals below.

Molnupiravir may be used in patients who are hospitalized for a diagnosis other than COVID-19, provided they have mild to moderate COVID-19 and are at high risk of progressing to severe disease. For the Panel's recommendations on preferred and alternative antiviral therapies for outpatients with COVID-19, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>.

Rationale

The MOVe-OUT trial enrolled high-risk, unvaccinated, nonhospitalized adults and reported that molnupiravir reduced the rate of hospitalization or death among these patients by 31% compared to placebo.⁷ This trial was conducted in 2021 before the emergence of the Omicron variant and its subvariants. A secondary analysis of the patients who required hospitalization during the trial found a reduced need for respiratory interventions among those who received molnupiravir compared to those

who received placebo.⁸ Molnupiravir has shown activity against the Omicron subvariants in vitro and in animal studies.^{2,9-11}

The PANORAMIC trial enrolled participants during a period when the Omicron variant was circulating.¹² The participants were nonhospitalized adults with COVID-19 who were at high risk of progressing to severe disease, and 94% had received at least 3 doses of a COVID-19 vaccine. The study found that the use of molnupiravir plus usual care did not reduce the primary composite outcome of hospitalization or death compared to usual care alone. The rates of this composite outcome were low (1%) in both arms. Molnupiravir plus usual care was superior to usual care alone for several secondary clinical endpoints. For example, patients who received molnupiravir plus usual care reported recovering from COVID-19 an estimated 4 days earlier than those who received usual care alone. However, because the PANORAMIC trial was an open-label study and the patients knew whether they were receiving molnupiravir or not, this may have affected their reported symptoms. As a result, these findings are less reliable than those from a placebo-controlled trial.

Although the different COVID-19 treatment options have not been directly compared in clinical trials, the Panel recommends using **molnupiravir** only when ritonavir-boosted nirmatrelvir and remdesivir are not available, feasible to use, or clinically appropriate (**CIIa**). Molnupiravir appears to have lower clinical efficacy than these other treatment options.

Some observational studies have evaluated the use of molnupiravir in nonhospitalized or hospitalized adults who are at high risk of progressing to severe disease, including some patients who received COVID-19 vaccines, but these studies have limitations.¹³⁻¹⁵ For treatment considerations for vaccinated individuals, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>.

Additional Considerations

- Patients should complete the 5-day treatment course of molnupiravir. It is unknown whether a shorter course is less effective or associated with the emergence of molnupiravir-resistant mutations.
- If a patient requires hospitalization after starting treatment, the full treatment course of molnupiravir can be completed at the health care provider's discretion.
- The FDA EUA for molnupiravir provides instructions for preparing and administering capsule contents through orogastric or nasogastric tubes.⁶
- There are no data on using combination antiviral therapies for the treatment of nonhospitalized patients with COVID-19. Clinical trials are needed to determine whether combination therapy has a role in the treatment of SARS-CoV-2 infection.
- Patients who are severely immunocompromised can experience prolonged periods of SARS-CoV-2 replication, which may lead to rapid viral evolution. There are theoretical concerns that using a single antiviral agent in these patients may produce antiviral-resistant viruses. Additional studies are needed to assess this risk. The role of combination antiviral therapy in treating patients who are severely immunocompromised is not yet known. See <u>Special</u> <u>Considerations in People Who Are Immunocompromised</u> for more information.
- There are limited data on the frequency of SARS-CoV-2 rebound in patients who have completed treatment with molnupiravir. During the MOVe-OUT trial, rates of symptomatic SARS-CoV-2 rebound were low (approximately 1%) in both those who received molnupiravir and those who received placebo.⁶

Monitoring, Adverse Effects, and Drug-Drug Interactions

The most common adverse effects of molnupiravir are diarrhea, nausea, and dizziness. Based on in vitro studies, neither molnupiravir nor its active metabolite NHC are inhibitors or inducers of major drug-metabolizing enzymes or inhibitors of major drug transporters.

According to the FDA EUA, no drug-drug interactions have been identified for molnupiravir.

Considerations in Sexually Active Individuals

For individuals of childbearing potential, clinicians should assess the patient's pregnancy status before initiating molnupiravir.

Patients of childbearing potential should be counseled about abstaining from sex or using reliable contraception for the duration of therapy and for up to 4 days after taking molnupiravir. Reproductive toxicity has been reported in animal studies of molnupiravir, and molnupiravir may be mutagenic during pregnancy.

The FDA EUA states that men of reproductive potential who are sexually active with individuals of childbearing potential should be counseled to abstain from sex or use a reliable method of contraception for the duration of treatment **and for at least 3 months after the last dose of molnupiravir**.

Considerations in Pregnancy

The Panel **recommends against** the use of **molnupiravir** for the treatment of COVID-19 in pregnant patients unless there are no other options and therapy is clearly indicated (**AIII**). See <u>Pregnancy</u>, <u>Lactation</u>, and <u>COVID-19 Therapeutics</u> for more information.

Considerations in Lactating People

Because the risk of adverse effects in infants is currently unknown, the FDA EUA fact sheet **recommends against** feeding an infant breast milk from a patient who is taking molnupiravir for the duration of the treatment course and for 4 days after the final dose. See <u>Pregnancy, Lactation, and</u> <u>COVID-19 Therapeutics</u> for more information.

Considerations in Children

The MOVe-OUT and PANORAMIC trials excluded participants aged <18 years. There are no data available on the use of molnupiravir in children aged <18 years. Molnupiravir is not authorized for use in those aged <18 years due to potential effects on bone and cartilage growth.

Clinical Data

MOVe-OUT

MOVe-OUT was a multinational, Phase 3 trial that evaluated the use of molnupiravir in unvaccinated, nonhospitalized adults with mild to moderate COVID-19 who were at high risk of progressing to severe COVID-19 and enrolled within 5 days of symptom onset.⁷ The trial was conducted in 2021 before the emergence of the Omicron variant and its subvariants. Pregnant people, lactating people, and children were excluded from the study. Patients were randomized to receive molnupiravir 800 mg PO every 12 hours for 5 days or placebo.

The primary composite endpoint was all-cause hospitalization (defined as a hospital stay >24 hours) or death by Day 29.

Results

- The final analysis included 1,433 patients:
 - The median age was 43 years (with 17% aged >60 years); 49% of patients were men, 57% were White, 50% were Hispanic/Latinx, and 5% were Black or African American.
 - Four percent had a body mass index \geq 30, and 16% had diabetes.
- The time from the onset of COVID-19 symptoms to randomization was ≤3 days in 48% of patients.
- By Day 29, the use of molnupiravir reduced the risk of hospitalization or death by 31%.
 - Forty-eight of 709 patients (6.8%) in the molnupiravir arm and 68 of 699 patients (9.7%) in the placebo arm experienced hospitalization or death (adjusted difference -3.0%; 95% CI, -5.9% to -0.1%).
 - One death occurred in the molnupiravir arm and 9 deaths occurred in the placebo arm.
- There were no significant differences between the arms in the proportion of patients who experienced adverse events or serious adverse events.
- A secondary analysis of data from the patients who were hospitalized during the trial revealed that the use of molnupiravir reduced the risk of requiring respiratory interventions (conventional or high-flow oxygen delivery, noninvasive ventilation, or mechanical ventilation) by 21%.⁸

Limitations and Interpretation

• When compared with placebo, the use of molnupiravir had a modest benefit in reducing the risk of hospitalization or death in unvaccinated, nonpregnant, high-risk adults with mild to moderate COVID-19. Molnupiravir also reduced the risk of pulmonary complications in these patients. However, this study was conducted before the emergence of the Omicron variant and its subvariants.

PANORAMIC

PANORAMIC was a large, multicenter, open-label, adaptive platform trial that was conducted in the United Kingdom.¹² The study evaluated the use of molnupiravir in nonhospitalized adults who were at high risk of progressing to severe COVID-19. The participants were aged \geq 50 years or \geq 18 years with comorbid conditions, and they had either a positive SARS-CoV-2 reverse transcription polymerase chain reaction result or rapid antigen test result at baseline. Patients were enrolled within 5 days of symptom onset. Pregnant people, lactating people, children, and those of childbearing potential who were unwilling to use effective contraception were excluded from the study. Patients were randomized to receive molnupiravir 800 mg PO twice daily for 5 days plus usual care or usual care alone.

The primary endpoint was a composite of all-cause hospitalization (defined as ≥ 1 overnight hospital stay, ≥ 1 night at home with care and monitoring by hospital clinicians, or an overnight stay in an emergency room) or death within 28 days of randomization. The trial was conducted from December 8, 2021, to April 27, 2022, when the Omicron variant was the dominant variant in the United Kingdom.

Results

- The final analysis included 25,708 patients. The mean age was 56.6 years (with 26.5% aged ≥65 years), 94% of patients were White, and 59% were women.
- Ninety-four percent of the patients had received \geq 3 doses of a COVID-19 vaccine.
- Overall, 69% of patients had comorbidities, including 25% with lung disease, 15% with obesity, 12% with diabetes, 8% with heart disease, and 8.5% were immunocompromised.

- Twenty-four percent of patients were taking inhaled corticosteroids.
- The mean time from symptom onset to starting molnupiravir was 3 days (range 3–5 days). Among the patients who provided information on their molnupiravir use, 95% reported completing the 5-day treatment course.
- Data on the primary outcome was available for 25,054 patients (97%).
 - In both arms, approximately 1% of patients were hospitalized or died. There were 103 hospitalizations and 3 deaths in the molnupiravir arm compared with 96 hospitalizations and 5 deaths in the usual care alone arm (aOR 1.06; 95% CrI, 0.81–1.41; probability of superiority 0.33).
 - Subgroup analyses revealed no evidence for treatment interaction.
- Molnupiravir plus usual care was superior to usual care alone for several secondary clinical endpoints.
 - The time from randomization to self-reported first recovery was significantly shorter among those who received molnupiravir (median of 9 days; IQR 5–23) than those who received usual care alone (median of 15 days; IQR 7–not reached).
 - After adjusting for age and baseline comorbidities, molnupiravir significantly reduced the estimated median time to first recovery. The median time to first recovery was 10.4 days (95% CrI, 10.1–10.6) in the molnupiravir arm and 14.6 days (95% CrI, 14.2–15) in the usual care alone arm (HR 1.36; 95% BCI, 1.32–1.40; probability of superiority >0.99).
 - The use of molnupiravir also significantly reduced the time to early sustained recovery (defined as recovery by Day 14 that was sustained until Day 28), the time to sustained recovery, the time to alleviation of all symptoms, the time to sustained alleviation of all symptoms, and the time to initial reduction of symptom severity.
- Serious adverse events occurred in 0.4% of patients in the molnupiravir arm and 0.3% of patients in the usual care alone arm. No serious adverse events related to molnupiravir were reported; 145 patients (1.1%) withdrew because of adverse effects attributed to molnupiravir.

Limitations and Interpretation

• The use of molnupiravir did not reduce the rate of progression to hospitalization or death among vaccinated, nonpregnant, high-risk adults, but it did reduce the time to improvement of symptoms. However, because the PANORAMIC trial was an open-label study and the patients knew whether they were receiving molnupiravir or not, this may have affected their reported symptoms. As a result, these findings are less reliable than those from a placebo-controlled trial.

References

- Fischer WA II, Eron JJ Jr, Holman W, et al. A Phase 2a clinical trial of molnupiravir in patients with COVID-19 shows accelerated SARS-CoV-2 RNA clearance and elimination of infectious virus. *Sci Transl Med.* 2022;14(628):eabl7430. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34941423</u>.
- 2. Zou R, Peng L, Shu D, et al. Antiviral efficacy and safety of molnupiravir against Omicron variant infection: a randomized controlled clinical trial. *Front Pharmacol*. 2022;13:939573. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35784723</u>.
- 3. Zhou S, Hill CS, Sarkar S, et al. Beta-d-N4-hydroxycytidine inhibits SARS-CoV-2 through lethal mutagenesis but is also mutagenic to mammalian cells. *J Infect Dis.* 2021;224(3):415-419. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33961695.
- 4. Kabinger F, Stiller C, Schmitzová J, et al. Mechanism of molnupiravir-induced SARS-CoV-2 mutagenesis. *Nat Struct Mol Biol.* 2021;28(9):740-746. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34381216</u>.

COVID-19 Treatment Guidelines

- 5. Centers for Disease Control and Prevention. People with certain medical conditions. 2023. Available at: https://www.cdc.gov/coronavirus/2019-ncov/need-extra-precautions/people-with-medical-conditions.html. Accessed February 23, 2023.
- 6. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Lagevrio (molnupiravir) capsules. 2023. Available at: <u>https://www.fda.gov/media/155054/download</u>.
- Jayk Bernal A, Gomes da Silva MM, Musungaie DB, et al. Molnupiravir for oral treatment of COVID-19 in nonhospitalized patients. *N Engl J Med*. 2022;386(6):509-520. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34914868.
- Johnson MG, Puenpatom A, Moncada PA, et al. Effect of molnupiravir on biomarkers, respiratory interventions, and medical services in COVID-19: a randomized, placebo-controlled trial. *Ann Intern Med*. 2022;175(8):1126-1134. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35667065</u>.
- 9. Vangeel L, Chiu W, De Jonghe S, et al. Remdesivir, molnupiravir and nirmatrelvir remain active against SARS-CoV-2 Omicron and other variants of concern. *Antiviral Res.* 2022;198:105252. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35085683.
- Takashita E, Yamayoshi S, Simon V, et al. Efficacy of antibodies and antiviral drugs against Omicron BA.2.12.1, BA.4, and BA.5 subvariants. *N Engl J Med.* 2022;387(5):468-470. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35857646</u>.
- 11. Uraki R, Kiso M, Iida S, et al. Characterization and antiviral susceptibility of SARS-CoV-2 Omicron BA.2. *Nature*. 2022;607(7917):119-127. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35576972</u>.
- 12. Butler CC, Hobbs FDR, Gbinigie OA, et al. Molnupiravir plus usual care versus usual care alone as early treatment for adults with COVID-19 at increased risk of adverse outcomes (PANORAMIC): an open-label, platform-adaptive randomised controlled trial. *Lancet*. 2023;401(10373):281-293. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36566761.
- Flisiak R, Zarębska-Michaluk D, Rogalska M, et al. Real-world experience with molnupiravir during the period of SARS-CoV-2 Omicron variant dominance. *Pharmacol Rep.* 2022;74(6):1279-1285. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36001284</u>.
- Yip CF, Lui GCY, Man Lai MS, et al. Impact of the use of oral antiviral agents on the risk of hospitalization in community coronavirus disease 2019 patients (COVID-19). *Clin Infect Dis.* 2023;76(3):e26-e33. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36031408</u>.
- 15. Wong CKH, Au ICH, Lau KTK, et al. Real-world effectiveness of early molnupiravir or nirmatrelvir-ritonavir in hospitalised patients with COVID-19 without supplemental oxygen requirement on admission during Hong Kong's Omicron BA.2 wave: a retrospective cohort study. *Lancet Infect Dis.* 2022;22(12):1681-1693. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36029795</u>.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Anti-SARS-CoV-2 Monoclonal Antibodies

Last Updated: March 6, 2023

Monoclonal antibodies (mAbs) that target the SARS-CoV-2 spike protein have been shown to have clinical benefits in treating SARS-CoV-2 infection. However, laboratory studies have found that the activity of anti-SARS-CoV-2 mAbs against specific variants and subvariants can vary dramatically. Because of this, these products are not expected to be effective treatments or preventives for COVID-19 in areas where the circulating variants and subvariants are resistant to mAbs.

Recommendation

- The COVID-19 Treatment Guidelines Panel (the Panel) **recommends against** the use of anti-SARS-CoV-2 mAbs for the treatment or prevention of COVID-19 (**AIII**) because the dominant Omicron subvariants in the United States are not expected to be susceptible to these products.
- For the Panel's recommendations on treating nonhospitalized patients with COVID-19, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> and <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u>.

Anti-SARS-CoV-2 Monoclonal Antibodies That Have Received Emergency Use Authorizations

Four anti-SARS-CoV-2 mAb products (bamlanivimab plus etesevimab, casirivimab plus imdevimab, sotrovimab, and bebtelovimab) have received Emergency Use Authorizations (EUA) from the Food and Drug Administration (FDA) for the treatment of outpatients with mild to moderate COVID-19. However, they are not currently authorized for use in the United States because the dominant Omicron subvariants are not expected to be susceptible to these products. See the Centers for Disease Control and Prevention <u>COVID Data Tracker</u> for regular updates on the regional proportions of SARS-CoV-2 variants in the United States.

On December 8, 2021, tixagevimab plus cilgavimab (Evusheld) received an EUA from the FDA that allowed this combination to be used as COVID-19 pre-exposure prophylaxis (PrEP). These 2 recombinant human mAbs bind to nonoverlapping epitopes of the spike protein receptor-binding domain of SARS-CoV-2. However, because many Omicron subvariants, including the dominant Omicron subvariants in the United States, are not expected to be susceptible to tixagevimab plus cilgavimab, this product is not authorized for use as COVID-19 PrEP as of January 26, 2023. See <u>Prevention of SARS-CoV-2 Infection</u> for more information.

Table A. SARS-CoV-2 Variants Currently or Recently Circulating in the United States and Their Susceptibility to Anti-SARS-CoV-2 Monoclonal Antibodies

See the <u>Guidelines Archive</u> for information on bamlanivimab plus etesevimab, casirivimab plus imdevimab, and variants that were previously circulating in the United States.

WHO Label BEB		B TIX Plus CIL		SOT		
and Pango Lineage	In Vitro Susceptibilityª	Anticipated Clinical Activity	In Vitro Susceptibility ^a	Anticipated Clinical Activity	In Vitro Susceptibilityª	Anticipated Clinical Activity
Omicron BA.5	No change	Active	Moderate reduction	Active	Marked reduction	Unlikely to be active
Omicron BA.4.6/BF.7	No change	Active	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active
Omicron BQ.1	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active
Omicron BQ.1.1	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active
Omicron XBB	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active
Omicron XBB.1.5	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active	Marked reduction	Unlikely to be active

^a This information is based on the fold reduction in susceptibility reported in the FDA EUAs¹⁻³ and in vitro neutralization studies.⁴⁻⁹

Key: BEB = bebtelovimab; CIL = cilgavimab; EUA = Emergency Use Authorization; FDA = Food and Drug Administration; SOT = sotrovimab; TIX = tixagevimab; WHO = World Health Organization

References

- 1. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization (EUA) of sotrovimab. 2022. Available at: <u>https://www.fda.gov/media/149534/download</u>.
- Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Evusheld (tixagevimab co-packaged with cilgavimab). 2023. Available at: <u>https://www.fda.gov/media/154701/download</u>.
- 3. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for bebtelovimab. 2022. Available at: <u>https://www.fda.gov/media/156152/download</u>.
- 4. Imai M, Ito M, Kiso M, et al. Efficacy of antiviral agents against Omicron subvariants BQ.1.1 and XBB. *N Engl J Med.* 2023;388(1):89-91. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36476720</u>.
- 5. Wang Q, Iketani S, Li Z, et al. Alarming antibody evasion properties of rising SARS-CoV-2 BQ and XBB subvariants. *Cell*. 2023;186(2):279-286.e8. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36580913</u>.
- 6. Takashita E, Yamayoshi S, Halfmann P, et al. In vitro efficacy of antiviral agents against Omicron subvariant BA.4.6. *N Engl J Med*. 2022;387(22):2094-2097. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36383452/</u>.
- 7. Wang Q, Li Z, Ho J, et al. Resistance of SARS-CoV-2 Omicron subvariant BA.4.6. to antibody neutralisation. *Lancet Infect Dis.* 2022;22(12):1666-1668. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36328002/</u>.
- 8. Takashita E, Yamayoshi S, Simon V, et al. Efficacy of antibodies and antiviral drugs against Omicron BA.2.12.1, BA.4, and BA.5 subvariants. *N Engl J Med*. 2022;387(5):468-470. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35857646.
- 9. Wang Q, Guo Y, Iketani S, et al. Antibody evasion by SARS-CoV-2 Omicron subvariants BA.2.12.1, BA.4 and BA.5. *Nature*. 2022;608(7923):603-608. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35790190/</u>.

Table 4b. Anti-SARS-CoV-2 Monoclonal Antibodies: Selected Clinical Trial Data

Last Updated: April 29, 2022

This table describes only the clinical trials that have evaluated anti-SARS-CoV-2 mAbs for the treatment of COVID-19. Please see <u>Prevention</u> of <u>SARS-CoV-2 Infection</u> for a discussion of the clinical trials that have evaluated anti-SARS-CoV-2 mAbs for PEP of SARS-CoV-2 infection.

Methods	Results	Limitations and Interpretation		
BLAZE-1: Double-Blind RCT of Bamlanivimab 700 mg Plus Etesevimab 1,400 mg in Nonhospitalized Patients With Mild to Moderate COVID-19 in the United States and Puerto Rico ¹				
 Key Inclusion Criteria Aged ≥12 years At high risk for severe COVID-19 or hospitalization Interventions Within 3 days of a positive SARS-CoV-2 test result, single infusion of: BAM 700 mg plus ETE 1,400 mg (n = 511) Placebo (n = 258) 	 Participant Characteristics Median age 56 years; 30% aged ≥65 years; 53% women 87% White, 27% Hispanic/Latinx, 8% Black/African American Mean duration of symptoms was 4 days 76% with mild COVID-19, 24% with moderate COVID-19 Primary Outcomes COVID-19-related hospitalizations or all-cause deaths by Day 29: 4 (0.8%) in BAM plus ETE arm vs. 15 (5.8%) in placebo 	 Key Limitation Conducted before widespread circulation of the Omicron VOC Interpretation Compared to placebo, BAM plus ETE significantly reduced the number of COVID-19-related hospitalizations and all-cause deaths in high-risk 		
 Primary Endpoint COVID-19-related hospitalization (defined as ≥24 hours of acute care) or death from any cause by Day 29 	 arm (change of -5.0%; 95% Cl, -8.0% to -2.1%; P < 0.001) All-cause deaths by Day 29: 0 in BAM plus ETE arm vs. 4 (1.6%) in placebo arm 	patients. Iovimab Alone Versus Placebo in		
Key Inclusion Criteria	Participant Characteristics	Key Limitations		
Aged 18–64 years	Median age 35 years; 56% women	Only low-risk patients included		
 No risk factors for progression to severe COVID-19 Key Exclusion Criteria ≥1 of the following: Sp0₂ ≤93% on room air Respiratory rate ≥30 breaths/min Heart rate ≥125 bpm 	 36% Hispanic/Latinx, 19% Black/African American Mean duration of symptoms prior to enrollment was 3.6 days Primary Outcomes Proportion with PHVL: 13% in BAM plus ETE plus BEB arm vs. 21% in placebo arm (P = 0.098), with a relative reduction of 38% (95% Cl, -9% to 65%) 	 Not powered to assess hospitalizations and deaths Conducted before widespread circulation of the Omicron VOC Interpretations There were no differences in the proportion of patients with PHVL across the arms. 		

COVID-19 Treatment Guidelines

Methods	Results	Limitations and Interpretation			
BLAZE-4, Treatment Arms 9–11: Double-Blind RCT of Bamlanivimab Plus Etesevimab Plus Bebtelovimab Versus Bebtelovimab Alone Versus Placebo in Low-Risk, Nonhospitalized Patients With Mild to Moderate COVID-19 ² , continued					
 Interventions Within 3 days of a positive SARS-CoV-2 test result, single infusion of: BAM 700 mg plus ETE 1,400 mg plus BEB 175 mg (n = 127) BEB 175 mg (n = 125) Placebo (n = 128) Primary Endpoint Proportion of patients with PHVL (defined as SARS-CoV-2 VL >5.82 log₁₀ by Day 7) 	 14% in BEB arm vs. 21% in placebo arm (P = 0.147), with a relative reduction of 34% (95% Cl, -15% to 62%) Secondary Outcomes Mean decline in VL greater in mAb arms vs. placebo arm at Day 5 but not at Days 3, 7, or 11 COVID-19-related hospitalizations or all-cause deaths by Day 29: 3 (2.4%) in BAM plus ETE plus BEB arm, including 1 death 2 (1.6%) in BEB arm 2 (1.6%) in placebo arm 	 Few COVID-19-related hospitalizations or deaths from any cause occurred by Day 29 across the arms, as is expected for a population of individuals who were at low risk of severe COVID-19. Compared to placebo, the median time to sustained symptom resolution was shorter in the BEB arm. 			
 Key Secondary Endpoints Mean change in VL from baseline to Days 3, 5, 7, and 11 COVID-19-related hospitalization or death from any cause by Day 29 Time to sustained symptom resolution BLAZE-4, Treatment Arms 12 and 13: Open-Label Relation Nonhospitalized Patients With Mild to Moderate CO 	 Median time to sustained symptom resolution: 7 days in BAM plus ETE plus BEB arm vs. 8 days in placebo arm (P = 0.289) 6 days in BEB arm vs. 8 days in placebo arm (P = 0.003) CT of Bamlanivimab Plus Etesevimab Plus Bebtelovimab and plus Bebtelov	d Bebtelovimab Alone in High-Risk,			
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
 Aged ≥12 years Weight ≥40 kg ≥1 risk factor for progression to severe COVID-19 Key Exclusion Criteria ≥1 of the following: Sp0₂ ≤93% on room air Respiratory rate ≥30 breaths/min Heart rate ≥125 bpm Interventions Within 3 days of a positive SARS-CoV-2 test result, single infusion of: 	 Median age 50 years; 52% women 18% Hispanic/Latinx, 18% Black/African American Mean duration of symptoms prior to enrollment was 4.7 days 21% had at least 1 dose of COVID-19 vaccine Efficacy Outcomes COVID-19-related hospitalization or death from any cause by Day 29: 2 (4%) in BAM plus ETE plus BEB arm vs. 3 (3%) in BEB arm Mean decline in VL greater in BAM plus ETE plus BEB arm vs. BEB arm at Day 5 but not at Days 3, 7, or 11 	 Open-label study No placebo arm Not powered to assess hospitalizations and deaths Conducted before widespread circulation of the Omicron VOC Interpretation There was no difference in the proportion of patients who were hospitalized or who died between the arms. 			

COVID-19 Treatment Guidelines

Methods	Results	Limitations and Interpretation		
BLAZE-4, Treatment Arms 12 and 13: Open-Label RCT of Bamlanivimab Plus Etesevimab Plus Bebtelovimab and Bebtelovimab Alone in High-Risk, Nonhospitalized Patients With Mild to Moderate COVID-19 ² , continued				
 BAM 700 mg plus ETE 1,400 mg plus BEB 175 mg (n = 50) BEB 175 mg (n = 100) 				
Efficacy Endpoints				
 COVID-19-related hospitalization or death from any cause by Day 29 				
 Mean change in VL from baseline to Days 3, 5, 7, and 11 				
Double-Blind RCT of Casirivimab Plus Imdevimab in	Nonhospitalized Patients With Mild to Moderate COVID-19	3		
Key Inclusion Criteria	Participant Characteristics	Key Limitation		
 Aged ≥18 years 	Median age 50 years	Conducted before widespread circulation of		
 Laboratory-confirmed SARS-CoV-2 infection 	• 35% Hispanic/Latinx, 5% Black/African American	the Omicron VOC		
 Symptom onset within 7 days of randomization 	Median duration of symptoms prior to enrollment was 3	Interpretation		
• For patients included in the modified full analysis	days	• Compared to placebo, CAS 600 mg plus IMD		
only:	Primary Outcomes	600 mg and CAS 1,200 mg plus IMD 1,200 mg reduced the risk of COVID-19-related		
• ≥ 1 risk factor for severe COVID-19	COVID-19-related hospitalizations or all-cause deaths	hospitalizations or all-cause deaths in		
 Positive SARS-CoV-2 RT-PCR result at baseline 	through Day 29:	patients with mild to moderate COVID-19.		
Interventions	• 7 (1.0%) in CAS 600 mg plus IMD 600 mg arm vs. 24			
Single IV infusion of:	(3.2%) in placebo arm ($P = 0.002$)			
 CAS 600 mg plus IMD 600 mg (n = 736) or placebo (n = 748) 	 18 (1.3%) in CAS 1,200 mg plus IMD 1,200 mg arm vs. 62 (4.6%) in placebo arm (P < 0.001) 			
• CAS 1,200 mg plus IMD 1,200 mg (n = 1,355) or placebo (n = 1,341)	 All-cause deaths: 1 (0.1%) in CAS 600 mg plus IMD 600 mg arm vs. 1 			
Primary Endpoint	(0.1%) in placebo arm			
 ≥1 COVID-19-related hospitalization or death from any cause by Day 29 	• 1 (< 0.1%) in CAS 1,200 mg plus IMD 1,200 mg arm vs. 3 (0.2%) in placebo arm			

Methods	Results	Limitations and Interpretation	
COMET-ICE: Double-Blind RCT of Sotrovimab in Nonhospitalized Patients With Mild to Moderate COVID-19 in Brazil, Canada, Peru, Spain, and the United States⁴			
Key Inclusion Criteria	Participant Characteristics	Key Limitation	
 Aged ≥18 years ≥1 comorbidity or aged ≥55 years Pagitive SARS, Col(2 PT DCP or antigon test result 	 Median age 53 years; 20% aged ≥65 years; 54% women 65% Hispanic/Latinx, 8% Black/African American 62% with phasity 22% with DM: 17% with medarate to 	Conducted before widespread circulation of the Omicron VOC	
 Positive SARS-CoV-2 RT-PCR or antigen test result Symptom onset ≤5 days before enrollment Key Exclusion Criteria Hospitalized or required supplemental oxygen Severely immunocompromised Interventions SOT 500 mg IV (n = 528) Placebo (n = 529) Primary Endpoint 	 63% with obesity; 22% with DM; 17% with moderate to severe asthma Primary Outcome Hospitalizations or all-cause deaths by Day 29: 6 (1%) in SOT arm vs. 30 (6%) in placebo arm, including 2 deaths (adjusted relative risk 0.21; 95% Cl, 0.09–0.50; absolute difference -4.53%; 95% Cl, -6.70% to -2.37%; <i>P</i> < 0.001) 	 Interpretation Compared to placebo, SOT reduced the incidence of all-cause hospitalizations and deaths among patients with mild to moderate COVID-19. 	
Hospitalization or death from any cause by Day 29			

Key: BAM = bamlanivimab; bpm = beats per minute; BEB = bebtelovimab; CAS = casirivimab; DM = diabetes mellitus; ETE = etesevimab; IMD = imdevimab; IV = intravenous; mAb = monoclonal antibody; PEP = post-exposure prophylaxis; PHVL = persistently high viral load; RCT = randomized controlled trial; RT-PCR = reverse transcription polymerase chain reaction; SOT = sotrovimab; Sp0₂ = oxygen saturation; VL = viral load; VOC = variant of concern

References

- 1. Dougan M, Azizad M, Mocherla B, et al. A randomized, placebo-controlled clinical trial of bamlanivimab and etesevimab together in high-risk ambulatory patients with COVID-19 and validation of the prognostic value of persistently high viral load. *Clin Infect Dis*. 2021;Published online ahead of print. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34718468.
- 2. Food and Drug Administration. Fact sheet for healthcare providers: emergency use authorization for bebtelovimab. 2022. Available at: https://www.fda.gov/media/156152/download.
- 3. Weinreich DM, Sivapalasingam S, Norton T, et al. REGEN-COV antibody combination and outcomes in outpatients with COVID-19. *N Engl J Med*. 2021;385(23):e81. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34587383</u>.
- 4. Gupta A, Gonzalez-Rojas Y, Juarez E, et al. Effect of sotrovimab on hospitalization or death among high-risk patients with mild to moderate COVID-19: a randomized clinical trial. *JAMA*. 2022;327(13):1236-1246. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35285853</u>.

COVID-19 Convalescent Plasma

Last Updated: November 2, 2023

Plasma from donors who have recovered from COVID-19 (regardless of vaccination status) may contain antibodies to SARS-CoV-2 that could help suppress viral replication.¹ In August 2020, the Food and Drug Administration (FDA) issued an Emergency Use Authorization (EUA) for COVID-19 convalescent plasma (CCP) for the treatment of hospitalized patients with COVID-19. The EUA was subsequently revised. The current EUA limits the authorization to the use of CCP products that contain high levels of anti-SARS-CoV-2 antibodies (i.e., high-titer products) for the treatment of outpatients with COVID-19 who have immunosuppressive disease or who are receiving immunosuppressive treatment. The testing criteria used to identify high-titer CCP products was also revised.²

The use of CCP should be limited to high-titer products. Products that are not labeled "high titer" should not be used.

Recommendations

Patients Who Are Immunocompromised

- There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of high-titer CCP for the treatment of COVID-19 in hospitalized or nonhospitalized patients who are immunocompromised.
- Some people who are immunocompromised have prolonged, symptomatic COVID-19 with evidence of ongoing SARS-CoV-2 replication. Without definitive data, some Panel members would use 1 or more of the following treatment options:
 - Longer and/or additional courses of ritonavir-boosted nirmatrelvir (Paxlovid)
 - Longer and/or additional courses of remdesivir
 - High-titer CCP from a vaccinated donor who recently recovered from COVID-19 likely caused by a SARS-CoV-2 variant similar to the variant causing the patient's illness

See <u>Special Considerations in People Who Are Immunocompromised</u> for a broader discussion on the therapeutic management of COVID-19 in people who are immunocompromised.

Patients Who Are Immunocompetent

- The Panel **recommends against** the use of CCP for the treatment of COVID-19 in hospitalized patients who are immunocompetent (**AI**).
- There is insufficient evidence for the Panel to recommend either for or against the use of high-titer CCP for the treatment of COVID-19 in nonhospitalized patients who are immunocompetent.

Rationale

Patients Who Are Immunocompromised

This section pertains to people who are moderately or severely immunocompromised. For examples of moderately or severely immunocompromising conditions and for a broader discussion on the therapeutic management of COVID-19 in people who are immunocompromised, see <u>Special Considerations in</u> <u>People Who Are Immunocompromised</u>.

Patients who are immunocompromised are at risk of having reduced antibody responses to SARS-CoV-2 infection and COVID-19 vaccination, having suboptimal control of viral replication, and progressing to severe disease.^{3,4} Despite the lack of definitive evidence, there is a physiologic rationale for the use of SARS-CoV-2 antibody-based therapies in these patients.

Under the revised EUA issued on December 27, 2021, CCP is authorized for the treatment of COVID-19 in nonhospitalized or hospitalized patients who have immunosuppressive disease or are receiving immunosuppressive treatment.²

Evidence to support the use of CCP for the treatment of COVID-19 in patients who are immunocompromised is limited. No randomized, adequately powered trials evaluating CCP for the treatment of COVID-19 in these patients have been published. Some subgroup analyses generated from clinical trials that enrolled patients who were immunocompromised suggested a potential benefit from the use of CCP in this population.⁵⁻⁷ However, subgroup analyses need to be interpreted with caution. In the overall trial populations, there was no evidence of benefit from the use of CCP. Other clinical data from case series, retrospective case-control studies, and meta-analyses have been cited as support for the potential benefit of CCP in patients who are immunocompromised. However, the patient populations differed across studies, and the studies had design limitations, making the findings difficult to interpret.⁸⁻¹⁶

The emergence of SARS-CoV-2 variants further complicates assessment of benefit from the use of CCP. Although results from some in vitro studies suggest that CCP collected from vaccinated individuals who recovered from Omicron infection exhibits neutralizing activity against certain Omicron subvariants,¹⁷⁻²³ extrapolation of these results to the clinical setting is challenging for the following reasons:²⁴

- COVID-19 immune responses across donor populations are heterogeneous; thus, CCP products are variable.
- The tests used to qualify high-titer CCP measure anti-SARS-CoV-2 antibody titers. They do not directly measure neutralizing activity or account for currently circulating subvariants.
- Published in vitro studies that evaluated the virologic activity of CCP against the currently circulating variants used a variety of assays that are difficult to compare and interpret.^{20-23,25-27}
- The pharmacokinetics and pharmacodynamics of individual CCP products are not clearly understood; therefore, determining the clinical relevance of a degree of in vitro neutralization activity is difficult.

In this context, the Panel has concluded that there is insufficient evidence for a definitive recommendation for treatment of COVID-19 in people who are immunocompromised. For patients who are immunocompromised and have prolonged COVID-19 symptoms and evidence of ongoing viral replication (e.g., those with a low cycle threshold value, as measured by a reverse transcription polymerase chain reaction result or with a positive rapid antigen test result) despite receiving a course of antiviral therapy, the optimal management is unknown. Case reports and case series have documented the treatment of these patients with additional antiviral treatments, prolonged courses of antiviral treatments, high-titer CCP, or combination therapy.²⁸⁻³² The data for these approaches are not definitive, but some Panel members would use longer and/or additional courses of ritonavir-boosted nirmatrelvir or remdesivir, high-titer CCP, or combinations of these. If CCP is used, clinicians should attempt to obtain high-titer CCP from a vaccinated donor who recently recovered from COVID-19 likely caused by a SARS-CoV-2 variant similar to the variant causing the patient's illness.

Adequately powered, well-designed, and well-conducted randomized clinical trials are needed to provide more specific, evidence-based guidance on the role of CCP in the treatment of COVID-19 in patients who are immunocompromised.

COVID-19 Treatment Guidelines

Hospitalized Patients Who Are Immunocompetent

Under the revised EUA, the use of CCP is not authorized for hospitalized patients with COVID-19 who do not have immunosuppressive disease or who are not receiving immunosuppressive treatments.

Clinical data on the use of CCP for the treatment of COVID-19 in hospitalized patients who are immunocompetent, including data from several randomized trials and the U.S. Expanded Access Program for CCP, are summarized in <u>Table 4c</u>.

Results from the 3 largest randomized controlled trials that evaluated CCP in hospitalized patients— RECOVERY,³³ CONCOR-1,³⁴ and REMAP-CAP⁶—found no evidence of benefit from the use of high-titer CCP in hospitalized patients with COVID-19. All 3 were open-label trials that were stopped early due to futility.

The Panel **recommends against** the use of CCP for the treatment of COVID-19 in hospitalized patients who are immunocompetent (**AI**).

Nonhospitalized Patients Who Are Immunocompetent

CCP is not authorized for the treatment of nonhospitalized patients with COVID-19 who do not have immunosuppressive disease or who are not receiving immunosuppressive treatment.

Data from well-designed clinical trials that evaluated the use of CCP for the treatment of nonhospitalized patients with COVID-19 prior to the emergence of the Omicron variants are conflicting. These data are summarized in <u>Table 4c</u>. Differences in patient populations, the placebo used (e.g., some studies used saline, and some used non–SARS-CoV-2 plasma), and CCP manufacturing and testing methods may have contributed to the disparate outcomes and difficulty in reconciling results across these clinical trials. The emergence of SARS-CoV-2 variants further complicates the assessment of benefit from the use of CCP.

There is insufficient evidence for the Panel to recommend either for or against the use of high-titer CCP for the treatment of COVID-19 in nonhospitalized patients who are immunocompetent.

Considerations in Pregnancy

The safety and efficacy of using CCP during pregnancy have not been evaluated in clinical trials, and published data on its use in pregnant individuals with COVID-19 are limited to case reports.³⁵ Pathogen-specific immunoglobulins (Ig) are used clinically during pregnancy to prevent infection from varicella zoster virus and rabies virus and have been used in clinical trials of congenital cytomegalovirus infection.^{36,37} Pregnancy is not a reason to withhold CCP from a patient if it is otherwise indicated. The expected physiologic immunomodulation during pregnancy should not affect the decision to use CCP.

Considerations in Children

The safety and efficacy of CCP have not been systematically evaluated in pediatric patients. Published literature on its use in children is limited to case reports and case series. A few clinical trials evaluating the use of CCP in children are ongoing. The use of high-titer CCP may be considered on a case-by-case basis for hospitalized children who are immunocompromised and meet the EUA criteria for its use. CCP is not authorized by the FDA for use in patients who are immunocompetent.

Several antiviral therapies are available for the treatment of children with COVID-19 who are at high risk of progressing to severe disease. The use of these therapies in children may be considered on a caseby-case basis. See <u>Special Considerations in Children</u> and <u>Therapeutic Management of Hospitalized</u> <u>Children With COVID-19</u> for more information.

Monitoring and Adverse Effects

The available data suggest that serious adverse reactions following the administration of CCP are infrequent and consistent with the risks associated with plasma infusions for other indications. These risks include transfusion-transmitted infections (e.g., HIV, hepatitis B, hepatitis C), allergic reactions, anaphylactic reactions, febrile nonhemolytic reactions, transfusion-related acute lung injury, transfusion-associated circulatory overload, and hemolytic reactions. Hypothermia, metabolic complications, and post-transfusion purpura have also been described.^{2,33,38}

Additional risks of CCP transfusion include a theoretical risk of antibody-dependent enhancement of SARS-CoV-2 infection. In the CONCOR-1 trial, higher levels of full transmembrane spike IgG were associated with worse outcomes, suggesting that the use of CCP with nonfunctional anti-SARS-CoV-2 antibodies may be harmful.³⁴ A subgroup analysis in the REMAP-CAP trial showed potential harm in patients who received CCP transfusions more than 7 days after being hospitalized.⁶

When considering the use of CCP in patients with a history of severe allergic or anaphylactic transfusion reactions, consultation with a transfusion medicine specialist is advised.

References

- Wang X, Guo X, Xin Q, et al. Neutralizing antibody responses to severe acute respiratory syndrome coronavirus 2 in coronavirus disease 2019 inpatients and convalescent patients. *Clin Infect Dis*. 2020;71(10):2688-2694. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32497196</u>.
- 2. Food and Drug Administration. Fact sheet for health care providers: Emergency Use Authorization (EUA) of COVID-19 convalescent plasma for treatment of coronavirus disease 2019 (COVID-19). 2021. Available at: https://www.fda.gov/media/141478/download.
- 3. Cattaneo C, Masina L, Pagani C, et al. High mortality in fully vaccinated hematologic patients treated with anti-CD20 antibodies during the "Omicron wave" of COVID-19 pandemic. *Hematol Oncol.* 2023;41(1):205-207. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35933702</u>.
- 4. Shahzad M, Chaudhary SG, Zafar MU, et al. Impact of COVID-19 in hematopoietic stem cell transplant recipients: a systematic review and meta-analysis. *Transpl Infect Dis*. 2022;24(2):e13792. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35030267.
- 5. Denkinger CM, Janssen M, Schäkel U, et al. Anti-SARS-CoV-2 antibody-containing plasma improves outcome in patients with hematologic or solid cancer and severe COVID-19: a randomized clinical trial. *Nat Cancer*. 2023;4(1):96-107. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36581734</u>.
- 6. Writing Committee for the REMAP-CAP Investigators. Effect of convalescent plasma on organ support-free days in critically ill patients with COVID-19: a randomized clinical trial. *JAMA*. 2021;326(17):1690-1702. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34606578</u>.
- Lacombe K, Hueso T, Porcher R, et al. COVID-19 convalescent plasma to treat hospitalised COVID-19 patients with or without underlying immunodeficiency. *medRxiv*. 2022;Preprint. Available at: https://www.medrxiv.org/content/10.1101/2022.08.09.22278329v2.
- 8. Thompson MA, Henderson JP, Shah PK, et al. Association of convalescent plasma therapy with survival in patients with hematologic cancers and COVID-19. *JAMA Oncol*. 2021:1167-1175. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34137799</u>.
- 9. Lanza F, Monaco F, Ciceri F, et al. Lack of efficacy of convalescent plasma in COVID-19 patients with concomitant hematological malignancies: an Italian retrospective study. *Hematol Oncol*. 2022;40(5):857-863. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35932208</u>.
- Lang-Meli J, Fuchs J, Mathé P, et al. Case series: convalescent plasma therapy for patients with COVID-19 and primary antibody deficiency. *J Clin Immunol*. 2022;42(2):253-265. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34893946</u>.

- 11. Rodionov RN, Biener A, Spieth P, et al. Potential benefit of convalescent plasma transfusions in immunocompromised patients with COVID-19. *Lancet Microbe*. 2021;2(4):e138. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33817676.
- Franchini M, Focosi D, Percivalle E, et al. Variant of concern-matched COVID-19 convalescent plasma usage in seronegative hospitalized patients. *Viruses*. 2022;14(7):1443. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35891421</u>.
- Ljungquist O, Lundgren M, Iliachenko E, et al. Convalescent plasma treatment in severely immunosuppressed patients hospitalized with COVID-19: an observational study of 28 cases. *Infect Dis (Lond)*. 2022;54(4):283-291. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34878955</u>.
- 14. Ripoll JG, Gorman EK, Juskewitch JE, et al. Vaccine-boosted convalescent plasma therapy for patients with immunosuppression and COVID-19. *Blood Adv*. 2022;6(3):5951-5955. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36156121.
- Senefeld JW, Klassen SA, Ford SK, et al. Use of convalescent plasma in COVID-19 patients with immunosuppression. *Transfusion*. 2021;61(8):2503-2511. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34036587</u>.
- 16. Beraud M, Goodhue Meyer E, Lozano M, et al. Lessons learned from the use of convalescent plasma for the treatment of COVID-19 and specific considerations for immunocompromised patients. *Transfus Apher Sci.* 2022;61(3):103355. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35063360</u>.
- Lusvarghi S, Pollett SD, Neerukonda SN, et al. SARS-CoV-2 BA.1 variant is neutralized by vaccine booster-elicited serum but evades most convalescent serum and therapeutic antibodies. *Sci Transl Med.* 2022;14(645):eabn8543. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35380448</u>.
- Yu J, Collier AY, Rowe M, et al. Neutralization of the SARS-CoV-2 Omicron BA.1 and BA.2 variants. *N Engl J Med*. 2022;386(16):1579-1580. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35294809</u>.
- Richardson SI, Madzorera VS, Spencer H, et al. SARS-CoV-2 Omicron triggers cross-reactive neutralization and Fc effector functions in previously vaccinated, but not unvaccinated, individuals. *Cell Host Microbe*. 2022;30(6):880-886.e4. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35436444</u>.
- 20. Li M, Beck EJ, Laeyendecker O, et al. Convalescent plasma with a high level of virus-specific antibody effectively neutralizes SARS-CoV-2 variants of concern. *Blood Adv.* 2022;6(12):3678-3683. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35443020</u>.
- 21. Khan K, Karim F, Ganga Y, et al. Omicron BA.4/BA.5 escape neutralizing immunity elicited by BA.1 infection. *Nat Commun.* 2022;13(1):4686. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35948557</u>.
- 22. Wang W, Lusvarghi S, Subramanian R, et al. Antigenic cartography of well-characterized human sera shows SARS-CoV-2 neutralization differences based on infection and vaccination history. *Cell Host Microbe*. 2022;30(12):1745-1758.e7. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36356586</u>.
- 23. Di Germanio C, Simmons G, Thorbrogger C, et al. Vaccination of COVID-19 convalescent plasma donors increases binding and neutralizing antibodies against SARS-CoV-2 variants. *Transfusion*. 2022;62(3):563-569. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35129839</u>.
- Rijnders BJA, Huygens S, Mitjà O. Evidence-based dosing of convalescent plasma for COVID-19 in future trials. *Clin Microbiol Infect*. 2022;28(5):667-671. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/35150881</u>.
- 25. Qu P, Faraone J, Evans JP, et al. Neutralization of the SARS-CoV-2 Omicron BA.4/5 and BA.2.12.1 subvariants. *N Engl J Med.* 2022;386(26):2526-2528. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35704428</u>.
- 26. Richardson SI, Kgagudi P, Manamela NP, et al. Antibody-dependent cellular cytotoxicity against SARS-CoV-2 Omicron sub-lineages is reduced in convalescent sera regardless of infecting variant. *Cell Rep Med.* 2023;4(1):100910. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36603577</u>.
- 27. Tuekprakhon A, Nutalai R, Dijokaite-Guraliuc A, et al. Antibody escape of SARS-CoV-2 Omicron BA.4 and

BA.5 from vaccine and BA.1 serum. *Cell*. 2022;185(14):2422-2433.e13. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35772405</u>.

- 28. Huygens S, Gharbharan A, Serroukh Y, et al. High-titer convalescent plasma plus nirmatrelvir/ritonavir treatment for non-resolving COVID-19 in six immunocompromised patients. *J Antimicrob Chemother*. 2023;78(7):1644-1648. Available at: https://www.ncbi.nlm.nih.gov/pubmed/37248664.
- Brosh-Nissimov T, Ma'aravi N, Leshin-Carmel D, et al. Combination treatment of persistent COVID-19 in immunocompromised patients with remdesivir, nirmaltrevir/ritonavir and tixegavimab/cilgavimab. *medRxiv*. 2023;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2023.04.07.23288144v1</u>.
- 30. Mikulska M, Sepulcri C, Dentone C, et al. Triple combination therapy with two antivirals and monoclonal antibodies for persistent or relapsed SARS-CoV-2 infection in immunocompromised patients. *Clin Infect Dis*. 2023;Published online ahead of print. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36976301</u>.
- Graziani L, Gori L, Manciulli T, et al. Successful use of nirmatrelvir/ritonavir in immunocompromised patients with persistent and/or relapsing COVID-19. *J Antimicrob Chemother*. 2023;78(2):555-558. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36544352</u>.
- 32. Trottier CA, Wong B, Kohli R, et al. Dual antiviral therapy for persistent coronavirus disease 2019 and associated organizing pneumonia in an immunocompromised host. *Clin Infect Dis*. 2023;76(5):923-925. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36281907</u>.
- 33. RECOVERY Collaborative Group. Convalescent plasma in patients admitted to hospital with COVID-19 (RECOVERY): a randomised controlled, open-label, platform trial. *Lancet*. 2021;397(10289):2049-2059. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34000257</u>.
- 34. Bégin P, Callum J, Jamula E, et al. Convalescent plasma for hospitalized patients with COVID-19: an open-label, randomized controlled trial. *Nat Med.* 2021;27(11):2012-2024. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34504336.
- 35. Franchini M, Prefumo F, Grisolia G, et al. Convalescent plasma for pregnant women with COVID-19: a systematic literature review. *Viruses*. 2021;13(7):1194. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34206468.
- 36. Revello MG, Lazzarotto T, Guerra B, et al. A randomized trial of hyperimmune globulin to prevent congenital cytomegalovirus. *N Engl J Med.* 2014;370(14):1316-1326. Available at: https://www.ncbi.nlm.nih.gov/pubmed/24693891.
- 37. Hughes BL, Clifton RG, Rouse DJ, et al. A trial of hyperimmune globulin to prevent congenital cytomegalovirus infection. N Engl J Med. 2021;385(5):436-444. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34320288</u>.
- 38. Nguyen FT, van den Akker T, Lally K, et al. Transfusion reactions associated with COVID-19 convalescent plasma therapy for SARS-CoV-2. *Transfusion*. 2021;61(1):78-93. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33125158</u>.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Table 4c. COVID-19 Convalescent Plasma: Selected Clinical Trial Data

Last Updated: March 6, 2023

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for CCP. The studies summarized below are those that have had the greatest impact on the Panel's recommendations.

Methods	Results	Limitations and Interpretation	
REMAP-CAP: Multinational, Open-Label RCT of High-Titer CCP in Hospitalized Patients With Critical COVID-19 in Australia, Canada, the United Kingdom, and the United States ¹			
Key Inclusion Criterion	Participant Characteristics	Key Limitations	
Admitted to ICU while receiving respiratory support	Mean age 61 years; 68% men	Open-label study	
(HFNC oxygen, NIV, MV, ECMO) and/or vasopressor or	• 32% on MV	Not all patients in CCP arm received	
inotrope support	 29% SARS-CoV-2 antibody negative at baseline 	CCP (86% received CCP as per	
Key Exclusion Criteria	 94% received corticosteroids, 45% received RDV, 39% 	protocol and 95% received some CCP).	
CCP contraindicated	received IL-6 inhibitors	Interpretation	
Death imminent	Primary Outcome	There was no benefit of CCP in	
Interventions	• Median number of organ support-free days by Day 21: 0 days	hospitalized patients with critical COVID-19.	
• High-titer CCP (550 mL +/- 150 mL) within 48 hours of randomization (n = 1,084)	in CCP arm vs. 3 days in usual care arm (OR 0.97; 95% Crl, 0.82–1.14)		
• Usual care (n = 916)	Secondary Outcomes		
Primary Endpoint	No difference between arms in:		
Number of organ support-free days by Day 21	• In-hospital mortality: 37% in CCP arm vs. 38% in usual care		
Key Secondary Endpoints	arm		
In-hospital mortality	 Mortality by Day 28 or Day 90 		
Mortality by Day 28 and Day 90	Median number of respiratory support-free days: 0 days in		
 Number of respiratory support-free days 	CCP arm vs. 2 days in usual care arm		
ICU LOS	Median ICU LOS: 21 days in CCP arm vs. 17 days in usual care arm		
	care arm		

Methods	Results	Limitations and Interpretation		
CONCOR-1: Multinational, Open-Label RCT of CCP for Hospitalized Patients With COVID-19 in Canada, the United States, and Brazil ²				
Key Inclusion Criteria	Participant Characteristics	Key Limitations		
Receipt of supplemental oxygen	Mean age 68 years; 59% men	Open-label study		
Within 12 days of respiratory symptom onset	84% receiving systemic corticosteroids at enrollment	Trial stopped at 78% of planned		
Key Exclusion Criterion	Primary Outcome	enrollment after meeting prespecified futility criteria for early termination.		
 Imminent or current intubation Interventions 	 Intubation or death by Day 30: 32% in CCP arm vs. 28% in SOC arm (relative risk 1.16; 95% Cl, 0.94–1.43, P = 0.18) 	Interpretation		
 1–2 units of CCP (approximately 500 mL) from 1–2 donors (n = 625) SOC (n = 313) 	 Secondary Outcomes By Day 30, no difference between arms in: Time to intubation or death 	 There was no benefit of CCP in oxygen- dependent, hospitalized patients with COVID-19 who were within 12 days of symptom onset. 		
Primary Endpoint	 Mortality: 23% in CCP arm vs. 21% in SOC arm 			
Intubation or death by Day 30	Mean ICU LOS: 4.3 days in CCP arm vs. 3.7 days in SOC			
Key Secondary EndpointsTime to intubation or death by Day 30	arm Need for renal dialysis: 1.6% in CCP arm vs. 2.0% in SOC arm 			
Mortality by Day 30	• Frequency of SAEs by Day 30: 33% in CCP arm vs. 26% in			
ICU LOS by Day 30	SOC arm			
 Need for renal dialysis by Day 30 				
Frequency of SAEs by Day 30				
<u>RECOVERY</u>: Open-Label RCT of High-Titer CCP in Hospit		1		
Key Inclusion Criterion	Participant Characteristics	Key Limitation		
Clinically suspected or laboratory-confirmed SARS-	Mean age 64 years; 64% men	Open-label study		
CoV-2 infection	• 5% on MV	Interpretation		
Key Exclusion Criterion	92% received corticosteroids	• There was no benefit of CCP in		
CCP contraindicated	Primary Outcomes	hospitalized patients with COVID-19.		
 Interventions 2 units of high-titer CCP (approximately 275 mL per unit) with IgG against SARS-CoV-2 spike protein and sample to cutoff ratio ≥6.0. First unit administered ASAP after randomization, second unit administered ≥12 	 No difference between arms in: All-cause mortality by Day 28: 24% in each arm Mortality in patients without detectable SARS-CoV-2 antibodies: 32% in CCP arm vs. 34% in usual care arm 			
hours later (n = $5,795$)				

COVID-19 Treatment Guidelines

Methods	Results	Limitations and Interpretation			
RECOVERY: Open-Label RCT of High-Titer CCP in Hospitalized Patients in the United Kingdom ³ , continued					
• Usual care (n = 5,763)	Secondary Outcomes				
Primary Endpoint	No difference between arms in:				
All-cause mortality by Day 28	 Proportion discharged by Day 28: 66% in both arms 				
Key Secondary Endpoints	Proportion who progressed to MV or death by Day 28: OOV in COD array of 20% in yourd care array				
Time to hospital discharge by Day 28	29% in CCP arm vs. 29% in usual care arm				
 Among patients not receiving MV, progression to MV or death by Day 28 					
<u>RECOVER</u> : Open-Label RCT of High-Titer CCP in Hospital	ized Patients With Severe COVID-19 in 4 Risk Groups in Ge	rmany ⁴			
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
 PCR-confirmed SARS-CoV-2 infection 	• 136 participants were enrolled between September 2020	Open-label study			
• Hospitalized with SpO ₂ \leq 94% on room air or PaO ₂ /FiO ₂	and January 2022.	The live virus neutralizing assay used			
<300 mm Hg	Mean age 69 years; 68% men; 97% White	to select plasma for this trial may not produce the same results as the assays			
• ≥1 of the following criteria:	 Participants were enrolled from 4 mutually exclusive patient groups: 	used to qualify high-titer CCP in the			
 Hematologic cancer and/or receipt of active cancer therapy in past 24 months for any cancer 	• Patients with cancer ($n = 56$)	current FDA EUA.			
 Chronic immunosuppression due to medications and/ 	 Patients with immunosuppression who did not have 	Small sample size			
or underlying disease	cancer (n = 16, including 12 solid organ transplant	• Trial was terminated early because the			
• Aged >50 to \leq 75 years with ALC <0.8 x 10 ⁹ cells/L	recipients)	neutralizing activity of stored plasma against the Omicron variant was not			
and/or D-dimer >1 µg/mL	• Patients aged >50 to ≤75 years with lymphopenia and/	known.			
 Aged >75 years without other listed criteria 	or elevated D-dimer levels ($n = 36$)	 Low proportion of vaccinated 			
Key Exclusion Criterion	• Patients aged >75 years without other criteria (n = 26)	participants and limited use of current			
Requiring MV or NIV	• 11% were fully vaccinated	SOC therapies, such as antiviral or immunomodulatory agents			
Interventions	 8% received small-molecule antiviral drugs (12% in plasma arm vs. 5% in SOC arm); 37% received anti- 	 Subgroup analyses were not adjusted for 			
• 2 units (238–337 mL) of high-titer CCP (\geq 1:80) or	inflammatory drugs (40% in plasma arm vs. 33% in SOC	multiple comparisons.			
vaccinated donor plasma from 2 donors on Days 1 and $2 (n - 69)$	arm)	Interpretation			
2 (n = 68) • SOC (n = 66)	 60% received supplemental oxygen via nasal cannula; 21% received HFNC oxygen or NIV 	The trial did not demonstrate a benefit			
	Median 7 days between symptom onset and	of high-titer CCP or vaccinated donor			
Primary Endpoint	randomization	plasma in the overall study population.			
Time to 2-point improvement on a 7-point OS or hospital discharge					

COVID-19 Treatment Guidelines

Methods	Results	Limitations and Interpretation
<u>RECOVER</u>: Open-Label RCT of High-Titer CCP in Hospital	ized Patients With Severe COVID-19 in 4 Risk Groups in Ge	rmany ⁴ , continued
Key Secondary Endpoints • 28-day, 56-day, and 84-day overall survival rate	 Primary Outcome Median time to 2-point improvement on OS or hospital discharge: 13 days in plasma arm vs. 18 days in SOC arm (HR 1.29; 95% Cl, 0.86–1.93; P=0.205) Median time to improvement or hospital discharge among patients with cancer: 13 days in plasma arm vs. 31 days in SOC arm (HR 2.50; 95% Cl, 1.34–4.79; P=0.003) Key Secondary Outcomes 	• Results from the predefined subgroup analysis of patients with cancer suggest a potential benefit of CCP or vaccinated donor plasma. However, this analysis was conducted largely before the emergence of the Omicron subvariants, so the results should be interpreted with caution.
	 No difference between arms in overall survival; 27 patients (19.9%) died (HR for survival 0.72; 95% Cl, 0.33–1.55; P=0.403) Fewer patients with cancer died in plasma arm than in SOC arm (HR 0.28; 95% Cl, 0.06–0.96; P=0.042) 	
CSSC-004: RCT of Early Treatment With High-Titer CCP		1
Key Inclusion Criterion	Participant Characteristics	Key Limitation
 COVID-19 symptoms for <8 days Key Exclusion Criteria Prior or planned COVID-19–related hospitalization 	 Median age 44 years; 7% aged ≥65 years; 57% women; 79% White 8% with type 2 DM; 2% with CVD; 38% with BMI ≥30 	 Patients were at relatively low risk for disease progression. Interpretation
 Receipt of anti-SARS-CoV-2 mAbs Interventions Approximately 250 mL of CCP with SARS-CoV-2 spike- RBD lgG titer ≥1:320 (n = 592) Non-SARS-CoV-2 plasma (n = 589) Primary Endpoint COVID-19–related hospitalization or all-cause death 	 82% unvaccinated Median 6 days between symptom onset and transfusion Primary Outcomes COVID-19-related hospitalization within 28 days: 2.9% in CCP arm vs. 6.3% in control arm (absolute risk reduction 3.4 percentage points; 95% Cl, 1.0–5.8; <i>P</i> = 0.005) 53 of 54 hospitalizations occurred in unvaccinated individuals. None occurred in fully vaccinated individuals. 	 This trial demonstrated a benefit of CCP in unvaccinated outpatients with <8 days of COVID-19 symptoms.
within 28 days	 All-cause deaths within 28 days: 0 in CCP arm vs. 3 in control arm 	

Methods	Results	Limitations and Interpretation			
CONV-ERT: RCT of High-Titer, Methylene Blue-Treated CCP as an Early Treatment for Outpatients With COVID-19 in Spain ⁶					
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
 Aged ≥50 years 	Mean age 56 years; 54% men	Trial was underpowered because it			
• Mild or moderate COVID-19 symptoms for \leq 7 days	• 75% with \geq 1 risk factor for COVID-19 progression	was terminated early due to rising vaccination rates among the eligible			
Key Exclusion Criteria	 97% with mild COVID-19 	patient population.			
 Severe COVID-19 symptoms or requirement for 	 Median 4.4 days of symptoms prior to enrollment 	Methylene blue, which was used for			
hospitalization for any reason	Among 369 patients with available baseline serologic	pathogen inactivation in donor plasma,			
Previous SARS-CoV-2 infection	testing, 88% negative for both IgG anti-SARS-CoV-2 spike and IgM anti-SARS-CoV-2 S1-RBD	could have potentially impaired Fc-			
 Receipt of ≥1 COVID-19 vaccine 		region functionality of Ig and negatively impacted product efficacy and blinding.			
Interventions	Primary Outcomes	Interpretation			
• 250–300 mL of high-titer, methylene blue-treated CCP	 Hospitalization within 28 days: 12% in CCP arm vs. 11% in placebo arm (relative risk 1.05; 95% Crl, 0.78–1.41) 	 This trial did not demonstrate a benefit 			
(n = 188)	• Mean change in SARS-CoV-2 VL: $-2.41 \log_{10}$ copies/mL in	of CCP in unvaccinated outpatients with			
• 0.9% saline (n = 188)	CCP arm vs2.32 \log_{10} copies/mL in placebo arm	<7 days of COVID-19 symptoms.			
Primary Endpoints	Key Secondary Outcomes				
Hospitalization within 28 days	• Death by Day 60: 0 in CCP arm vs. 2 in placebo arm				
Mean change in SARS-CoV-2 VL from baseline to Day 7	(relative risk 0.20; 95% Cl 0.01–4.14)				
Key Secondary Endpoints	• No difference between arms in median time to symptom				
Death by Day 60	resolution: 12.0 days for both arms (HR 1.05; 95% Cl,				
Time to complete symptom resolution	0.85–1.30)				
Double-Blind RCT of Early High-Titer CCP Therapy to Pre	event Severe COVID-19 in Nonhospitalized Older Adults in /	Argentina ⁷			
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
• Aged \geq 75 years or aged 65–74 years with \geq 1 coexisting	Mean age 77 years; 38% men	Small sample size			
condition	Most with comorbidities	• Early termination because number of			
 Mild COVID-19 symptoms for <72 hours 	Primary Outcome	COVID-19 cases decreased			
Key Exclusion Criterion	• Severe respiratory disease by Day 15: 16% in CCP arm	Interpretation			
Severe respiratory disease	vs. 31% in placebo arm (relative risk 0.52; 95% Cl,	This trial demonstrated a benefit of CCP			
Interventions	0.29–0.94; <i>P</i> = 0.03)	in older adult outpatients with <72 hours of mild COVID-19 symptoms.			
 250 mL of CCP with IgG against SARS-CoV-2 spike protein >1:1,000 (n = 80) 					
• Saline (n = 80)					

Methods	Results	Limitations and Interpretation			
Double-Blind RCT of Early High-Titer CCP Therapy to Prevent Severe COVID-19 in Nonhospitalized Older Adults in Argentina ⁷ , continued					
Primary Endpoint					
 Severe respiratory disease, defined as respiratory rate ≥30 breaths/min and/or SpO₂ <93% on room air, by Day 15 					
SIREN-C3PO: Multicenter, Single-Blind RCT of High-Titer CC	P in Adults With COVID-19 in the United States ⁸				
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
• ED patient with \leq 7 days of symptoms	Median age 54 years; 46% men	• In the primary analysis, the number			
 PCR-confirmed SARS-CoV-2 infection 	• More patients with immunosuppression in CCP arm than in	of patients who required hospital			
• Aged \geq 50 years or aged \geq 18 years with \geq 1 risk factor for	placebo arm (13% vs. 7%)	admission during the index visit was not balanced across arms.			
disease progression	• More patients with ≥ 3 risk factors in CCP arm than in	• The CCP arm included more			
v Exclusion Criterion		patients with multiple risk factors,			
 Need for supplemental oxygen 	Primary Outcomes	including immunosuppression.			
Interventions	• No difference between arms in proportion with disease progression: 30% in CCP arm vs. 32% in placebo arm (risk	Interpretation			
• 250 mL of high-titer CCP (median titer 1:641) (n = 257)	difference 1.9%; 95% Crl, -6.0% to 9.8%)	The use of high-titer CCP within			
• Saline (n = 254)	• 25 patients (19 in CCP arm and 6 in placebo arm) required	1 week of symptom onset did not			
Primary Endpoint	hospitalization during index visit. In a post hoc analysis	prevent disease progression in outpatients with COVID-19 who			
 Disease progression, defined as hospital admission, death, or seeking emergency or urgent care within 15 days of randomization 	that excluded these patients, disease progression occurred in 24% in CCP arm vs. 30% in placebo arm (risk difference 5.8%; 95% Crl, -1.9% to 13.6%).	were at high risk of severe disease.			
Key Secondary Endpoints	Secondary Outcomes				
Severity of illness, as measured by an OS	• All-cause mortality within 30 days: 5 (1.9%) in CCP arm vs. 1 (0.4%) in placebo arm				
All-cause mortality within 30 days	No difference between arms in illness severity or mean				
 Number of hospital-free days by Day 30 	number of hospital-free days				

Methods	Results	Limitations and Interpretation			
CoV-Early: Double-Blind RCT of CCP in Nonhospitalized, High-Risk Adults With COVID-19 in the Netherlands ⁹					
 Key Inclusion Criteria Aged ≥70 years, aged ≥50 years with a comorbidity, or aged ≥18 years and severely immunocompromised Positive SARS-CoV-2 RT-PCR or antigen test result COVID-19 symptoms for ≤7 days 	 Participant Characteristics Median age 60 years; 22% women Median 5 days of symptoms Median 1 comorbidity Median Sp0₂ 97% at baseline 	 Key Limitations Study was discontinued after 421 of 690 planned participants were enrolled, resulting in decreased power. The CCP used was selected based on a PRNT50 assay and may not qualify as high-titer CCP per the current FDA EUA. Interpretation This trial did not demonstrate a benefit of CCP in nonhospitalized, high-risk patients with COVID-19. 			
 Key Exclusion Criteria Life expectancy <28 days History of TRALI IgA deficiency Interventions 300 mL of CCP with minimum PRNT50 titer of 1:160 (n = 207) Non-SARS-CoV-2 plasma collected prior to pandemic (n = 209) Primary Endpoint Improvement based on 5-point OS by Day 28 Secondary Endpoints Percentage of hospital admissions 	 7.9% SARS-CoV-2 IgG antibody negative at baseline 2.9% fully vaccinated; 5.0% received 1 vaccine Primary Outcome Odds of receiving highest score on 5-point OS by Day 28: OR 0.86; 95% Crl, 0.59–1.22 in CCP arm Secondary Outcomes Percentage of hospital admissions: 10 patients (4.8%) in CCP arm vs. 18 patients (8.6%) in non-SARS-CoV-2 arm (aHR 0.61; 95% Cl, 0.28–1.34) Number of days of symptoms: 13 days in CCP arm vs. 12 days in non-CCP arm (P = 0.99) 				
Number of days of symptoms					
Retrospective Evaluation of CCP Antibody Levels and the Ri	sk of Death From COVID-19 in the United States ¹⁰	1			
 Key Inclusion Criteria Severe or life-threatening COVID-19 Patients for whom samples of transfused CCP were available for retrospective analysis of antibody titer Interventions High-titer CCP (n = 515), medium-titer CCP (n = 2,006), or low-titer CCP (n = 561), characterized retrospectively Primary Endpoint Mortality by Day 30 after CCP transfusion 	 Participant Characteristics 31% aged ≥70 years; 61% men; 48% White, 37% Hispanic/Latinx 61% in ICU; 33% on MV 51% received corticosteroids, 31% received RDV Primary Outcomes Mortality by Day 30 after transfusion: 22% in high-titer CCP arm vs. 27% in medium-titer CCP arm vs. 30% in low-titer CCP arm 	 Key Limitation Lack of untreated control arm Interpretation The study data are not sufficient to establish the efficacy or safety of CCP. 			

Methods	Results	Limitations and Interpretation
Retrospective Evaluation of CCP Antibody Levels and the Ri	sk of Death From COVID-19 in the United States ¹⁰ , continued	
	• Lower risk of death in high-titer CCP arm than low-titer CCP arm (relative risk 0.75; 95% CI, 0.61–0.93)	
	 Lower mortality among patients not receiving MV before CCP transfusion (relative risk 0.66; 95% Cl, 0.48–0.91) 	
	 No difference in mortality between high-titer and low- titer arms among patients on MV before CCP transfusion (relative risk 1.02; 95% CI, 0.78–1.32) 	

Key: ALC = absolute lymphocyte count; ASAP = as soon as possible; BMI = body mass index; CCP = COVID-19 convalescent plasma; CVD = cardiovascular disease; DM = diabetes mellitus; ECMO = extracorporeal membrane oxygenation; ED = emergency department; EUA = Emergency Use Authorization; Fc = fragment crystallizable; FDA = Food and Drug Administration; HFNC = high-flow nasal cannula; ICU = intensive care unit; Ig = immunoglobulin; IL = interleukin; LOS = length of stay; mAb = monoclonal antibody; MV = mechanical ventilation; NIV = noninvasive ventilation; OS = ordinal scale; the Panel = the COVID-19 Treatment Guidelines Panel; PaO₂/FiO₂ = ratio of arterial partial pressure of oxygen to fraction of inspired oxygen; PCR = polymerase chain reaction; PRNT50 = 50% plaque reduction neutralization test; RBD = receptor-binding domain; RCT = randomized controlled trial; RDV = remdesivir; RT-PCR = reverse transcription polymerase chain reaction; SAE = serious adverse event; SOC = standard of care; SpO₂ = oxygen saturation; TRALI = transfusion-related acute lung injury; VL = viral load

References

- 1. Writing Committee for the REMAP-CAP Investigators. Effect of convalescent plasma on organ support-free days in critically ill patients with COVID-19: a randomized clinical trial. *JAMA*. 2021;326(17):1690-1702. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34606578</u>.
- 2. Bégin P, Callum J, Jamula E, et al. Convalescent plasma for hospitalized patients with COVID-19: an open-label, randomized controlled trial. *Nat Med*. 2021;27(11):2012-2024. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34504336</u>.
- 3. RECOVERY Collaborative Group. Convalescent plasma in patients admitted to hospital with COVID-19 (RECOVERY): a randomised controlled, open-label, platform trial. *Lancet*. 2021;397(10289):2049-2059. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34000257.
- 4. Denkinger CM, Janssen M, Schäkel U, et al. Anti-SARS-CoV-2 antibody-containing plasma improves outcome in patients with hematologic or solid cancer and severe COVID-19: a randomized controlled trial. *Nat Cancer*. 2023;4(1):96-107. Available at: https://pubmed.ncbi.nlm.nih.gov/36581734/.
- 5. Sullivan DJ, Gebo KA, Shoham S, et al. Early outpatient treatment for COVID-19 with convalescent plasma. *N Engl J Med.* 2022;386(18):1700-1711. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35353960</u>.
- Alemany A, Millat-Martinez P, Corbacho-Monné M, et al. High-titre methylene blue-treated convalescent plasma as an early treatment for outpatients with COVID-19: a randomised, placebo-controlled trial. *Lancet Respir Med.* 2022;10(3):278-288. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35150610</u>.
- 7. Libster R, Pérez Marc G, Wappner D, et al. Early high-titer plasma therapy to prevent severe COVID-19 in older adults. *N Engl J Med*. 2021;384(7):610-618. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33406353</u>.
- 8. Korley FK, Durkalski-Mauldin V, Yeatts SD, et al. Early convalescent plasma for high-risk outpatients with COVID-19. N Engl J Med.

COVID-19 Treatment Guidelines

2021;385(21):1951-1960. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34407339.

- 9. Gharbharan A, Jordans C, Zwaginga L, et al. Outpatient convalescent plasma therapy for high-risk patients with early COVID-19: a randomized placebo-controlled trial. *Clin Microbiol Infect*. 2023;29(2):208-214. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36007870.
- 10. Joyner MJ, Carter RE, Senefeld JW, et al. Convalescent plasma antibody levels and the risk of death from COVID-19. *N Engl J Med*. 2021;384(11):1015-1027. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33523609</u>.

Interferons

Last Updated: December 20, 2023

Interferons are a family of cytokines with in vitro and in vivo antiviral properties. Interferon beta-1a has been approved by the Food and Drug Administration (FDA) to treat relapsing forms of multiple sclerosis, and pegylated formulations of interferon alfa-2a and interferon alfa-2b have been approved by the FDA to treat hepatitis B and hepatitis C virus infections. Several interferons, including interferon alfa, beta, and lambda, have been evaluated for the treatment of COVID-19. Interferon lambda is not currently approved or authorized by the FDA for any use.

Recommendations

- For nonhospitalized patients with mild to moderate COVID-19, the COVID-19 Treatment Guidelines Panel (the Panel) **recommends against** the use of **interferon alfa** or **beta**, except in a clinical trial (**AIIa**).
- For hospitalized patients with COVID-19, the Panel **recommends against** the use of **systemic interferon alfa**, except in a clinical trial (**AIIa**).
- For hospitalized patients with COVID-19, the Panel **recommends against** the use of **systemic interferon beta** (**AI**).
- The Panel is unable to recommend either for or against the use of interferon lambda because this product is not currently available for clinical use.

Rationale

Interferon Alfa and Beta

Many of the studies that evaluated the use of systemic interferons for the treatment of hospitalized adults with COVID-19 were conducted in early 2020, before the widespread use of remdesivir or corticosteroids and other immunomodulators. In addition, these studies administered interferons with other drugs that have since been shown to have no clinical benefit in people with COVID-19, such as lopinavir/ritonavir and hydroxychloroquine.¹⁻³

More recent studies have shown no benefit of using interferon beta-1a to treat patients with COVID-19, and some of the trials have suggested that interferon beta-1a can cause harm in patients with severe disease, such as those who require high-flow oxygen, noninvasive ventilation, or mechanical ventilation.^{4,5} In a large randomized controlled trial of hospitalized patients with COVID-19, the combination of interferon beta-1a plus remdesivir showed no clinical benefit when compared to remdesivir alone.⁴ Similarly, the World Health Organization Solidarity trial did not show a benefit of administering interferon beta-1a to hospitalized patients, approximately 50% of whom were on corticosteroids.⁵

Systemic interferon alfa and inhaled interferons have also been evaluated in patients with COVID-19. The trials that have evaluated the use of interferon alfa have generally been small or moderate in size and have not been adequately powered to assess whether this agent provides a clinical benefit for patients with COVID-19.⁶⁻⁸

Interferon Lambda

Pegylated interferon lambda was studied in a randomized, double-blind, adaptive clinical trial that *COVID-19 Treatment Guidelines*

enrolled nonhospitalized patients with COVID-19 in Brazil and Canada.⁹ A total of 1,941 patients with risk factors for severe COVID-19 were randomized to receive either a single subcutaneous injection of pegylated interferon lambda 180 μ g or placebo. Eighty-three percent of these patients had received at least 1 dose of a COVID-19 vaccine. The primary outcome was a composite of observation in an emergency department for >6 hours or hospitalization, and 1 of the secondary outcomes was a composite of hospitalization or death. By Day 28 after randomization, the use of interferon lambda was associated with a 51% decrease in the occurrence of the primary outcome and a 39% decrease in the occurrence of this secondary outcome. Patients with a high baseline SARS-CoV-2 viral load who received interferon lambda were more likely to have cleared the virus by Day 7 than those who received placebo.

The drug was generally well tolerated. However, since pegylated interferon lambda is an investigational agent that is not currently available for clinical use, the Panel cannot make a recommendation for its use at this time.

Summaries of the studies that informed the Panel's recommendations can be found in Table 4d.

Considerations in Pregnant People

According to analyses of data from several large pregnancy registries, exposure to interferon beta-1b prior to conception or during pregnancy does not lead to an increased risk of adverse birth outcomes (e.g., spontaneous abortion, congenital anomaly).^{10,11} In a study that used data from pregnancy registries in Sweden and Finland, women who were exposed to interferon beta during pregnancy did not report significant changes in the birth weight, height, or head circumference of their infants.¹²

Considerations in Children

There are insufficient data on the use of interferons to treat respiratory viral infections in children to make any recommendations for treating children with COVID-19.

References

- 1. Alavi Darazam I, Hatami F, Mahdi Rabiei M, et al. An investigation into the beneficial effects of high-dose interferon beta 1-a, compared to low-dose interferon beta 1-a in severe COVID-19: The COVIFERON II randomized controlled trial. *Int Immunopharmacol*. 2021;99:107916. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34224994</u>.
- 2. Hung IF, Lung KC, Tso EY, et al. Triple combination of interferon beta-1b, lopinavir-ritonavir, and ribavirin in the treatment of patients admitted to hospital with COVID-19: an open-label, randomised, Phase 2 trial. *Lancet*. 2020;395(10238):1695-1704. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32401715.
- 3. Rahmani H, Davoudi-Monfared E, Nourian A, et al. Interferon beta-1b in treatment of severe COVID-19: a randomized clinical trial. *Int Immunopharmacol*. 2020;88:106903. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32862111.
- 4. Kalil AC, Mehta AK, Patterson TF, et al. Efficacy of interferon beta-1a plus remdesivir compared with remdesivir alone in hospitalised adults with COVID-19: a double-bind, randomised, placebo-controlled, Phase 3 trial. *Lancet Respir Med.* 2021;9(12):1365-1376. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34672949.
- 5. WHO Solidarity Trial Consortium. Repurposed antiviral drugs for COVID-19—interim WHO Solidarity trial results. *N Engl J Med.* 2021;384(6):497-511. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33264556</u>.
- 6. Yu J, Lu X, Tong L, et al. Interferon-alpha-2b aerosol inhalation is associated with improved clinical outcomes in patients with coronavirus disease–2019. *Br J Clin Pharmacol*. 2021;87(12):4737-4746. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/33982806</u>.
- 7. Wang B, Li D, Liu T, Wang H, Luo F, Liu Y. Subcutaneous injection of IFN alpha-2b for COVID-19:

an observational study. *BMC Infect Dis*. 2020;20(1):723. Available at: <u>https://pubmed.ncbi.nlm.nih.</u> gov/33008327.

- Pandit A, Bhalani N, Shashi Bhushan BL, et al. Efficacy and safety of pegylated interferon alpha-2b in moderate COVID-19: a phase II, randomized, controlled, open-label study. *Int J Infect Dis*. 2021;105:516-521. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/33713817</u>.
- 9. Reis G, Moreira Silva EAS, Medeiros Silva DC, et al. Early treatment with pegylated interferon lambda for COVID-19. *N Engl J Med*. 2023;388(6):518-528. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36780676</u>.
- Sandberg-Wollheim M, Alteri E, Moraga MS, Kornmann G. Pregnancy outcomes in multiple sclerosis following subcutaneous interferon beta-1a therapy. *Mult Scler*. 2011;17(4):423-430. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21220368</u>.
- Hellwig K, Duarte Caron F, Wicklein EM, Bhatti A, Adamo A. Pregnancy outcomes from the global pharmacovigilance database on interferon beta-1b exposure. *Ther Adv Neurol Disord*. 2020;13:1756286420910310. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32201504</u>.
- Burkill S, Vattulainen P, Geissbuehler Y, et al. The association between exposure to interferon-beta during pregnancy and birth measurements in offspring of women with multiple sclerosis. *PLoS One*. 2019;14(12):e0227120. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31887199</u>.

Table 4d. Interferons: Selected Clinical Trial Data

Last Updated: December 20, 2023

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for interferons. The studies summarized below are the randomized controlled trials that have had the greatest impact on the Panel's recommendations.

Methods	Results	Limitations and Interpretation
ACTT-3: Multinational, Double-Blind RCT of Interferon Be	19 ¹	
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Evidence of pneumonia (radiographic infiltrates, SpO₂ ≤94% on room air, or supplemental oxygen) 	 Mean age 59 years; 38% were aged ≥65 years 58% men; 32% Latinx, 60% White, 17% Black 	• After 270 patients were enrolled, OS6 patients were excluded because of an
No MV required	 Mean of 8.6 days of symptoms before enrollment 	increased frequency of AEs in this group.
Key Exclusion Criteria	• 90% had \geq 1 comorbidity; 58% with HTN; 58% with	Interpretation
 AST or ALT >5 times ULN 	obesity; 37% with DM	There was no clinical benefit of adding IFN bets 1s to PDV in beautifulized
Impaired renal function	Primary Outcome	IFN beta-1a to RDV in hospitalized patients with COVID-19.
Hospital discharge or transfer anticipated within 72 hours	 Median time to recovery: 5 days in both arms (rate ratio 0.99; 95% Cl, 0.87–1.13; P = 0.88) 	 The use of IFN beta-1a was associated with worse outcomes among patients
Interventions	 In patients on high-flow oxygen or NIV (OS6) at 	who were OS6 at baseline.
 RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily for 9 days plus IFN beta-1a 44 μg SUBQ every other day for up to 4 doses (n = 487) 	baseline, median time to recovery: >28 days in IFN beta-1a arm vs. 9 days in placebo arm (rate ratio 0.40; 95% Cl, 0.22–0.75; <i>P</i> = 0.0031)	
• RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily	Secondary Outcomes	
for 9 days plus placebo (n = 482)	No difference between arms in clinical status at Day 14	
Primary Endpoint	(OR 1.01; 95% Cl, 0.79–1.28)	
Time to recovery by Day 28	 No difference between IFN beta-1a arm and placebo arm in mortality by Day 28 in: 	
Key Secondary Endpoints	• All patients: 5% vs. 3% (HR 1.33; 95% Cl, 0.69–2.55)	
Clinical status at Day 14, as measured by an OS	• Patients who were OS6 at baseline: 21% vs. 12% (HR	
Mortality by Day 28	1.74; 95% Cl, 0.51–5.93)	

Methods	Results	Limitations and Interpretation		
WHO Solidarity Trial: Multinational, Open-Label, Adaptive RCT of IV or SUBQ Interferon Beta-1a or Other Repurposed Drugs in Hospitalized Adults With COVID-19 ²				
Key Inclusion Criteria	Participant Characteristics	Key Limitations		
 Diagnosis of COVID-19 	 35% aged <50 years; 19% aged ≥70 years; 63% men 	Open-label study		
 Not expected to be transferred elsewhere within 72 hours 	 70% on supplemental oxygen; 7% on ventilation Approximately 50% received corticosteroids during the 	 IFN beta-1a given as IV or SUBQ formulations at different doses. 		
Interventions	study.	Interpretation		
 IFN beta-1a 44 µg SUBQ on day of randomization, Day 3, and Day 6 (n = 1,656) IFN beta-1a 10 µg IV daily for 6 days for patients on high-flow oxygen, ventilation, or ECMO (n = 394) IFN beta-1a (either SUBQ or IV) and LPV/RTV 400 mg/50 mg twice daily for 14 days (n = 651) Local SOC (n = 2,050) Primary Endpoint In-hospital mortality 	 Primary Outcome In-hospital mortality: 11.9% in combined IFN beta-1a arms vs. 10.5% in SOC arm (rate ratio 1.16; 95% Cl, 0.96–1.39) For IFN beta-1a only (without LPV/RTV) recipients vs. SOC recipients, rate ratio was 1.12 (95% Cl, 0.83–1.51). Among those on ventilation at baseline, age-stratified rate ratio for in-hospital mortality was 1.40 (95% Cl, 0.93–2.11). 	 IFN beta-1a did not reduce in-hospital mortality in hospitalized patients with COVID-19. 		
Key Secondary Endpoint	Secondary Outcome			
Initiation of ventilation	 10% initiated ventilation in the combined IFN beta-1a arms and SOC arm. 			

Methods	Results	Limitations and Interpretation
DisCoVeRy Solidarity Trial Add-On: Open-Label, Adaptive Hospitalized Adults With COVID-19 in France ³	RCT of Interferon Beta-1a Plus Lopinavir/Ritonavir, Lopin	navir/Ritonavir, or Hydroxychloroquine in
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Positive SARS-CoV-2 PCR result 	Median age 63 years; 72% men	Open-label study
 Patients had pulmonary rales or crackles with Sp0₂ ≤94% on room air or required supplemental oxygen 	 29% with obesity; 26% with chronic cardiac disease; 22% with DM 	 Most patients had moderate disease. No IFN beta-1a arm without LPV/RTV
Interventions	36% had severe disease	• Study stopped early for futility.
 IFN beta-1a 44 µg SUBQ on Days 1, 3, and 6 plus LPV/ 	Median of 9 days of symptoms before randomization	Interpretation
RTV 400 mg/100 mg P0 twice daily for 14 days plus SOC	30% received steroids during the study.	Compared to SOC alone, the use of IFN-
(n = 145)	Primary Outcome	beta-1a plus LPV/RTV did not improve
• LPV/RTV 400 mg/100 mg P0 twice daily for 14 days plus S0C (n = 145)	No difference in clinical status at Day 15 for any intervention compared to SOC:	clinical status, rate of viral clearance, or time to viral clearance in hospitalized
• HCQ 400 mg twice on Day 1, then HCQ 400 mg daily for 9 days plus SOC (n = 145)	 IFN beta-1a plus LPV/RTV: aOR 0.69 (95% Cl, 0.45– 1.04; P = 0.08) 	patients with COVID-19.
SOC alone, which included corticosteroids,	• LPV/RTV: aOR 0.83 (95% Cl, 0.55–1.26; $P = 0.39$)	
anticoagulants, or immunomodulatory agents but not antivirals (n = 148)	 HCQ: a0R 0.93 (95% Cl, 0.62–1.41; P = 0.75) 	
Primary Endpoint	Secondary Outcomes	
Clinical status at Day 15, as measured by an OS	• No difference between arms in clinical status at Day 29	
Key Secondary Endpoints	No difference between arms in rate or time to SARS- CoV-2 viral clearance	
Clinical status at Day 29	• Time to improvement of 2 OS categories and hospital	
Rate of SARS-CoV-2 viral clearance	discharge by Day 29 was longer in LPV/RTV plus IFN	
Time to SARS-CoV-2 viral clearance by Day 29	beta-1a and LPV/RTV arms than in SOC arm.	
 Time to improvement of 2 OS categories by Day 29 		
 Time to hospital discharge by Day 29 		

Methods	Results	Limitations and Interpretation			
TOGETHER: Double-Blind, Adaptive RCT of Pegylated Interferon Lambda in Nonhospitalized Patients With COVID-19 in Brazil and Canada ⁴					
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
 Key Inclusion Criteria Positive SARS-CoV-2 antigen test result Within 7 days of symptom onset ≥1 high-risk factor for disease progression (e.g., age ≥50 years, comorbidities, immunosuppression) Up to 25% of patients could have no high-risk factors. Key Exclusion Criteria Need for hospitalization Sp0₂ ≤93% on room air Interventions Single dose of PEG-IFN lambda 180 µg SUBQ (n = 931) Placebo (n = 1,018; 825 received single SUBQ injection, 193 received PO placebo) Primary Endpoint Composite of ED observation >6 hours or hospitalization for COVID-19 by Day 28 Key Secondary Endpoints Composite of COVID-19–related hospitalization or death by Day 28 SARS-CoV-2 viral clearance at Day 7 Occurrence of AEs 	 Participant Characteristics Median age 43 years; 57.1% women; 95.1% self-identified as mixed race 1,919 (98.5%) from Brazil, 30 (1.5%) from Canada 50% with obesity 59.4% were randomized within 3 days of symptom onset. 83% received ≥1 COVID-19 vaccine dose. Primary Outcome Composite of ED observation >6 hours or hospitalization for COVID-19 by Day 28 (ITT): 25 (2.7%) in PEG-IFN lambda arm vs. 57 (5.6%) in placebo arm (relative risk 0.49; 95% Bayesian Crl, 0.30–0.76) 61 events (74%) were hospitalizations (ITT). Secondary Outcomes Composite of COVID-19–related hospitalization or death by Day 28: 22 (2.4%) in PEG-IFN lambda arm vs. 40 (3.9%) in placebo arm (relative risk 0.61; 95% Crl, 0.36–0.99) SARS-CoV-2 viral clearance at Day 7 among the 15% of patients with VL >192 million copies/mL at baseline: 50.5% in PEG-IFN lambda arm vs. 32.9% in placebo arm (OR 2.13; 95% Crl, 1.14–4.00) Occurrence of AEs: 141 (15.1%) in PEG-IFN lambda arm vs. 172 (16.9%) in placebo arm (relative risk 0.90; 95% 	 Key Limitations Health care facility capacity may have influenced the number and duration of ED observations. As this was an adaptive platform trial where multiple investigational treatments or placebos were being evaluated simultaneously, not all patients in the placebo arm received a placebo that was matched to PEG-IFN lambda. Interpretation In outpatients with COVID-19 who were within 7 days of symptom onset, PEG- IFN lambda reduced the need for ED observations >6 hours or hospitalization when compared with placebo. 			

Methods	Results	Limitations and Interpretation
Single-Blind RCT of Pegylated Interferon Lambda-1a for	Treatment of Outpatients With Uncomplicated COVID-19 i	n the United States⁵
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Aged 18–65 years Asymptomatic or symptomatic Positive SARS-CoV-2 RT-PCR result within 72 hours of enrollment 	 Median age 36 years; 42% women; 63% Latinx, 28% White 7% were asymptomatic. Median of 5 days of symptoms before randomization 	 Small sample size Interpretation PEG-IFN lambda-1a provided no virologic or clinical benefit compared to placebo
 Key Exclusion Criteria Current or imminent hospitalization Respiratory rate >20 breaths/min 	 Primary Outcome Median time to cessation of viral shedding: 7 days in both arms (aHR 0.81; 95% CI, 0.56–1.19; P = 0.29) 	among outpatients with uncomplicated COVID-19.
 Sp0₂ <94% on room air Decompensated liver disease Interventions Single dose of PEG-IFN lambda-1a 180 µg SUBQ (n = 60) Placebo (n = 60) Primary Endpoint 	 Secondary Outcomes No difference between PEG-IFN lambda-1a and placebo arms in: Proportion of patients hospitalized by Day 28: 3.3% for each arm Time to resolution of symptoms: 8 days vs. 9 days (HR 0.94; 95% Cl, 0.64–1.39) 	
 Time to first negative SARS-CoV-2 RT-PCR result Key Secondary Endpoints Hospitalization by Day 28 Time to complete symptom resolution 	 Other Outcome Patients who received PEG-IFN lambda-1a were more likely to have elevations of transaminase concentrations than patients who received placebo (25% vs. 8%; P = 0.027). 	

Methods	Results	Limitations and Interpretation	
Double-Blind RCT of Pegylated Interferon Lambda in Out	patients With Laboratory-Confirmed COVID-19 in Canada	6	
Key Inclusion Criteria	Participant Characteristics	Key Limitation	
Positive SARS-CoV-2 PCR result	Median age 46 years; 58% women; 52% White	Small sample size	
 Patients were within 7 days of symptom onset, or, if 	• 19% were asymptomatic.	Interpretation	
asymptomatic, were within 7 days of first positive SARS-	Mean of 4.5 days of symptoms before randomization	PEG-IFN lambda may accelerate VL	
CoV-2 test result.	Primary Outcome	decline and clearance in outpatients	
Key Exclusion Criterion	• 80% in PEG-IFN lambda arm vs. 63% in placebo arm	with COVID-19; however, the clinical significance of this finding is unclear.	
 Immunosuppression or condition that could be worsened by PEG-IFN lambda 	were negative for SARS-CoV-2 RNA at Day 7 ($P = 0.15$).	significance of this multiple unclear.	
Interventions	Secondary Outcomes		
 Single dose of PEG-IFN lambda 180 µg SUBQ (n = 30) 	• VL decline by Day 7 was greater in PEG-IFN lambda arm		
• Placebo $(n = 30)$	than in placebo arm ($P = 0.0041$).		
Primary Endpoint	• 1 participant in each arm hospitalized by Day 14		
	Other Outcome		
 Proportion of patients with negative SARS-CoV-2 test result on nasal mid-turbinate swab at Day 7 	 3 participants in each arm had mild elevations of aminotransferase concentrations. Increase was greater 		
Key Secondary Endpoints	in PEG-IFN lambda arm.		
Quantitative change in SARS-CoV-2 RNA over time			
 Hospitalization by Day 14 			

Key: AE = adverse event; ALT = alanine transaminase; AST = aspartate aminotransferase; DM = diabetes mellitus; ECMO = extracorporeal membrane oxygenation; ED = emergency department; HCQ = hydroxychloroquine; HTN = hypertension; IFN = interferon; ITT = intention-to-treat; IV = intravenous; LPV/RTV = lopinavir/ritonavir; MV = mechanical ventilation; NIV = noninvasive ventilation; OS = ordinal scale; the Panel = the COVID-19 Treatment Guidelines Panel; PCR = polymerase chain reaction; PEG-IFN = pegylated interferon; PO = oral; RCT = randomized controlled trial; RDV = remdesivir; RT-PCR = reverse transcription polymerase chain reaction; SOC = standard of care; SpO₂ = oxygen saturation; SUBQ = subcutaneous; ULN = upper limit of normal; VL = viral load; WHO = World Health Organization

References

- 1. Kalil AC, Mehta AK, Patterson TF, et al. Efficacy of interferon beta-1a plus remdesivir compared with remdesivir alone in hospitalised adults with COVID-19: a double-bind, randomised, placebo-controlled, Phase 3 trial. *Lancet Respir Med.* 2021;9(12):1365-1376. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34672949</u>.
- 2. WHO Solidarity Trial Consortium. Repurposed antiviral drugs for COVID-19—interim WHO Solidarity trial results. *N Engl J Med*. 2021;384(6):497-511. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33264556</u>.
- 3. Ader F, Peiffer-Smadja N, Poissy J, et al. An open-label randomized controlled trial of the effect of lopinavir/ritonavir, lopinavir/ritonavir plus

IFN-beta-1a and hydroxychloroquine in hospitalized patients with COVID-19. *Clin Microbiol Infect*. 2021;27(12):1826-1837. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34048876</u>.

- 4. Reis G, Moreira Silva EAS, Medeiros Silva DC, et al. Early treatment with pegylated interferon lambda for COVID-19. *N Engl J Med.* 2023;388(6):518-528. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36780676</u>.
- 5. Jagannathan P, Andrews JR, Bonilla H, et al. Peginterferon lambda-1a for treatment of outpatients with uncomplicated COVID-19: a randomized placebo-controlled trial. *Nat Commun*. 2021;12(1):1967. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33785743.
- 6. Feld JJ, Kandel C, Biondi MJ, et al. Peginterferon lambda for the treatment of outpatients with COVID-19: a Phase 2, placebo-controlled randomised trial. *Lancet Respir Med.* 2021;9(5):498-510. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33556319.

Table 4e. Characteristics of Antiviral Agents, Including Antibody Products

Last Updated: November 2, 2023

- This table contains drugs and products that have shown antiviral activity against SARS-CoV-2, including small-molecule antiviral drugs, CCP, and IFNs.
- RDV and RTV-boosted nirmatrelvir (Paxlovid) are approved by the FDA for the treatment of COVID-19.
- MOV and CCP have received EUAs from the FDA for the treatment of COVID-19.
- For drug-drug interaction information, please refer to product labels, EUA fact sheets, and <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications</u>.
- For the Panel's recommendations on using the drugs listed in this table, refer to to <u>Antiviral Agents, Including Antibody Products;</u> Therapeutic Management of Nonhospitalized Adults With COVID-19; Therapeutic Management of Hospitalized Adults With COVID-19; Therapeutic Management of Nonhospitalized Children With COVID-19; Therapeutic Management of Hospitalized Children With <u>COVID-19</u>; and <u>Pregnancy, Lactation, and COVID-19</u> Therapeutics.

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Anti-SARS-CoV-2	Antiviral Drugs (Small-Molecule Antivi	irals)			
Ritonavir- Boosted Nirmatrelvir (Paxlovid) Approved by the FDA for use in adults and authorized under an FDA EUA for use in children (aged ≥ 12 years and weighing ≥ 40 kg) for the treatment of mild to moderate COVID-19 in high- risk individuals.	 FDA Prescribing Information/EUA Dose for COVID-19^{1,2} eGFR ≥60 mL/min Nirmatrelvir 300 mg (two 150-mg tablets) with RTV 100 mg (one 100- mg tablet) twice daily for 5 days eGFR ≥30 to 60 mL/min Nirmatrelvir 150 mg (one 150-mg tablet) with RTV 100 mg (one 100- mg tablet) twice daily for 5 days eGFR <30 mL/min Not recommended (see comments) Severe Hepatic Impairment (Child- Pugh Class C) Not recommended 	 Dysgeusia Diarrhea Anaphylaxis, serious skin reactions, and other HSRs 	 Boxed warning: Monitor for potential AEs due to drug-drug interactions with concomitant medications. Weigh potential benefits of treatment against potential risks of drug-drug interactions. Use with caution in patients with pre-existing liver diseases, liver enzyme abnormalities, or hepatitis. Consider checking renal function in patients with suspected renal impairment. Monitor for HSRs. 	 RTV-boosted nirmatrelvir has significant drug- drug interactions. Before prescribing RTV-boosted nirmatrelvir, carefully review concomitant medications, including OTC medicines, herbal supplements, and recreational drugs. See <u>Drug-Drug Interactions</u> <u>Between Ritonavir-Boosted</u> <u>Nirmatrelvir (Paxlovid) and</u> <u>Concomitant Medications</u> for additional guidance and resources to assist with identifying drug-drug interactions. 	 The FDA prescribing information/EUA does not recommend using RTV-boosted nirmatrelvir in patients with eGFR <30 mL/min. Both nirmatrelvir and RTV tablets can be taken with or without food. The FDA prescribing information/EUA advises against crushing nirmatrelvir and RTV tablets. However, some data indicate that the tablets can be split or crushed if necessary.³

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Anti-SARS-CoV-	2 Antiviral Drugs (Small-Molecule Anti	virals), continued			
Remdesivir Approved by the FDA for the treatment of COVID-19 in individuals aged ≥28 days and weighing ≥3 kg.	Dose for Adults and Children Weighing ≥40 kg • RDV 200 mg IV on Day 1, then RDV 100 mg IV once daily from Day 2 Dose for Children Aged ≥28 Days and Weighing 3 kg to <40 kg	 Nausea ALT and AST elevations HSRs Increases in prothrombin time 	 Monitor patients for infusion-related reactions during the infusion and observe them for ≥1 hour after the infusion as clinically appropriate. Monitor renal function, hepatic function, and prothrombin time as clinically indicated. 	 Clinical drug-drug interaction studies of RDV have not been conducted. In vitro, RDV is a minor substrate of CYP3A4; a substrate of OATP1B1 and P-gp; and an inhibitor of CYP3A4, OATP1B1, OATP1B3, and MATE1.⁴ 	• RDV may be used without dose adjustment in patients with renal impairment, including those receiving dialysis. ⁴

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments	
Anti-SARS-CoV-2	Inti-SARS-CoV-2 Antiviral Drugs (Small-Molecule Antivirals), continued					
Molnupiravir Authorized under an FDA EUA for the treatment of mild to moderate COVID-19 in high- risk individuals aged ≥18 years.	 Dose Recommended in FDA EUA MOV 800 mg (four 200-mg capsules) PO every 12 hours for 5 days MOV is not authorized for use in people aged <18 years due to potential effects on bone and cartilage growth. 	 Diarrhea Nausea Dizziness Per the EUA, the 5-day course of MOV has a low risk for genotoxicity.⁵ See Molnupiravir for details. 	 Before initiating MOV, assess the patient's pregnancy status as clinically indicated. Monitor for potential AEs. 	 Clinical drug-drug interaction studies of MOV have not been conducted. Drug-drug interactions related to hepatic metabolism are not expected. 	 People of reproductive potential who are sexually active should use effective contraception during and after treatment with MOV. If MOV is prescribed for a pregnant person, the clinician should document that the risks and benefits were discussed with the patient and that the patient chose to receive MOV. Pregnant patients should also be offered the opportunity to participate in the MOV pregnancy surveillance program. Lactating people should not breastfeed their infants during treatment with MOV and for 4 days after treatment. MOV can be taken with or without food. The EUA provides instructions for preparing and administering MOV capsule contents through OG or NG tubes.⁵ 	

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments	
COVID-19 Convalesce	OVID-19 Convalescent Plasma					
High-Titer COVID-19 Convalescent Plasma Authorized under an FDA EUA for the treatment of COVID-19 in patients who are immunocompromised or who are receiving immunosuppressive treatment.	 Dose Recommended in FDA EUA Administer 1 high-titer CCP unit (about 200 mL) IV. Administer an additional CCP unit IV based on the prescribing provider's judgment and the patient's clinical response. 	 TRALI TACO Allergic reactions Anaphylactic reactions Febrile nonhemolytic reactions Hemolytic reactions Hypothermia Metabolic complications Transfusion-transmitted infections⁶ Thrombotic events Theoretical risk of antibody- mediated enhancement of infection and suppressed long-term immunity 	 Before administering CCP to patients with a history of severe allergic or anaphylactic transfusion reactions, consult a transfusion medicine specialist who is associated with the hospital's blood bank. Monitor for transfusion-related reactions. Monitor vital signs at baseline and during and after transfusion. 	• Drug products should not be added to the IV infusion line for the blood product.	• In patients with impaired cardiac function and heart failure, it may be necessary to reduce the CCP volume or decrease the transfusion rate.	
Interferons	1	1	1	1	1	
IFN Beta Not approved by the FDA and not recommended by the Panel for the treatment of COVID-19. Currently under investigation in clinical trials.	 Various doses and durations for IFN beta-1a and IFN beta-1b are being studied in clinical trials. 	 Flu-like symptoms (e.g., fever, fatigue, myalgia) Leukopenia, neutropenia, thrombocytopenia, lymphopenia Liver function abnormalities (ALT > AST) Injection site reactions Headache Hypertonia Pain Rash Worsening depression Induction of autoimmunity 	 Monitor CBC with differential and liver enzymes. Monitor for worsening CHF. Monitor for signs of depression and suicidal ideation. 	 Low potential for drug- drug interactions Use with caution with other hepatotoxic agents. Reduce dose if ALT is >5 times ULN. 	 Inhaled IFN beta-1a is not approved by the FDA for use in the United States. 	

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Interferons, continued					
PEG-IFN Lambda Not approved by the FDA and not recommended by the Panel for the treatment of COVID-19.	 Dose for COVID-19 in Clinical Trials Single dose of PEG-IFN lambda 180 μg SUBQ 	 Liver function abnormalities (ALT > AST) Injection site reactions 	 CBC with differential Liver enzymes Monitor for potential AEs. 	 Low potential for drug- drug interactions Use with caution with other hepatotoxic agents. 	• PEG-IFN lambda is not approved by the FDA for use in the United States.

Key: AE = adverse event; ALT = alanine transaminase; AST = aspartate aminotransferase; CBC = complete blood count; CCP = COVID-19 convalescent plasma; CHF = congestive heart failure; CYP = cytochrome P450; eGFR = estimated glomerular filtration rate; EUA = Emergency Use Authorization; FDA = Food and Drug Administration; HSR = hypersensitivity reaction; IFN = interferon; IV = intravenous; MATE = multidrug and toxin extrusion protein; MOV = molnupiravir; NG = nasogastric; OATP = organic anion transporting polypeptide; OG = orogastric; OTC = over-the-counter; the Panel = the COVID-19 Treatment Guidelines Panel; P-gp = P-glycoprotein; PEG-IFN = pegylated interferon; PO = oral; RDV = remdesivir; RTV = ritonavir; SUBQ = subcutaneous; TACO = transfusion-associated circulatory overload; TRALI = transfusion-related acute lung injury; ULN = upper limit of normal

References

- 1. Ritonavir-boosed nirmatrelvir (Paxlovid) [package insert]. Food and Drug Administration. 2023. Available at: <u>https://www.accessdata.fda.gov/drugsatfda_docs/label/2023/217188s000lbl.pdf</u>.
- 2. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Paxlovid. 2023. Available at: https://www.fda.gov/media/155050/download.
- 3. BC COVID Therapeutics Committee COVID Therapy Review and Advisory Working Group. Therapeutic brief: crushing nirmatrelvir/ritonavir (Paxlovid). 2022. Available at: <u>http://www.bccdc.ca/Health-Professionals-Site/Documents/COVID-treatment/Crushing_Paxlovid.pdf</u>.
- 4. Remdesivir (Veklury) [package insert]. Food and Drug Administration. 2023. Available at: <u>https://www.accessdata.fda.gov/drugsatfda_docs/label/2023/214787s019lbl.pdf</u>.
- 5. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Lagevrio (molnupiravir) capsules. 2023. Available at: <u>https://www.fda.gov/media/155054/download</u>.
- 6. Food and Drug Administration. Fact sheet for health care providers: Emergency Use Authorization (EUA) of COVID-19 convalescent plasma for treatment of coronavirus disease 2019 (COVID-19). 2021. Available at: https://www.fda.gov/media/141478/download.

Immunomodulators

Last Updated: October 10, 2023

Summary Recommendations

- The hyperactive inflammatory response to SARS-CoV-2 infection plays a central role in the pathogenesis of COVID-19. See <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> and <u>Therapeutic Management of Hospitalized</u> <u>Children With COVID-19</u> for the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations on the use of the following immunomodulators for hospitalized patients according to disease severity (listed in alphabetical order):
 - Abatacept
 - · Baricitinib (or tofacitinib)
 - Dexamethasone
 - Infliximab
 - Tocilizumab (or sarilumab)
- There is insufficient evidence for the Panel to recommend either for or against the use of the following immunomodulators for the treatment of COVID-19:
 - Anakinra
 - · Inhaled corticosteroids
 - Vilobelimab
- The Panel **recommends against** the use of **canakinumab** for the treatment of COVID-19, except in a clinical trial **(Blla)**.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Systemic Corticosteroids

Last Updated: July 21, 2023

Multiple randomized trials indicate that systemic corticosteroid therapy improves clinical outcomes and reduces mortality in hospitalized patients with COVID-19 who require supplemental oxygen,^{1,2} presumably by mitigating the COVID-19-induced systemic inflammatory response that can lead to lung injury and multisystem organ dysfunction. In contrast, in hospitalized patients with COVID-19 who do not require supplemental oxygen, the use of systemic corticosteroids provided no benefit and increased mortality.^{3,4} The COVID-19 Treatment Guidelines Panel's (the Panel) recommendations for the use of systemic corticosteroids in hospitalized patients with COVID-19 are based on results from these clinical trials (see Table 5a for more information). There are no data to support the use of systemic corticosteroids in nonhospitalized patients with COVID-19.

Recommendations

- The Panel recommends against the use of dexamethasone or other systemic corticosteroids in nonhospitalized patients in the absence of another indication (AIIb).
- See Therapeutic Management of Hospitalized Adults With COVID-19 for the Panel's recommendations on the use of dexamethasone or other systemic corticosteroids in certain hospitalized patients.
- Patients with COVID-19 who are receiving dexamethasone or another corticosteroid for an underlying condition should continue this therapy as directed by their health care provider (AIII).

Nonhospitalized Adults

There are no data to support the use of systemic corticosteroids in nonhospitalized patients with COVID-19. Therefore, the safety and efficacy of using systemic corticosteroids in this population have not been established. Generally, the use of systemic corticosteroids is associated with adverse events (e.g., hyperglycemia, neuropsychiatric symptoms, secondary infections), which may be difficult to detect and monitor in an outpatient setting. For more information, see General Management of Nonhospitalized Adults With Acute COVID-19.

Hospitalized Adults

The RECOVERY trial was a multicenter, open-label trial in the United Kingdom that randomly assigned 6,425 hospitalized patients to receive up to 10 days of dexamethasone 6 mg once daily plus standard care or standard care alone.³ Mortality at 28 days was lower among the patients who received dexamethasone than among those who received standard care alone. This benefit of dexamethasone was observed in patients who were mechanically ventilated or who required supplemental oxygen at enrollment. In contrast, no benefit was seen in patients who did not require supplemental oxygen at enrollment.

Among critically ill patients receiving supplemental oxygen with or without mechanical ventilation, several clinical trials, some of which were terminated early, demonstrated lower all-cause mortality at 28 days when systemic corticosteroids were compared with standard of care or placebo.¹

In addition to the randomized controlled trials, a large observational study evaluated the use of systemic corticosteroids in 15,404 hospitalized patients with positive SARS-CoV-2 polymerase chain reaction or antigen test results from a Department of Veteran Affairs database.⁴ Corticosteroids were administered to COVID-19 Treatment Guidelines 239

60% of the patients within 48 hours of admission, and 95% of the patients who received corticosteroids received dexamethasone. A total of 9,450 patients did not receive supplemental oxygen during the study. Of these patients, 3,514 (37%) received dexamethasone, administered for a median duration of 5 days (IQR 3–8 days). Using average treatment effect estimates, patients who received dexamethasone without supplemental oxygen had an increased risk of death within 90 days (HR 1.76; 95% CI, 1.47–2.12). Patients who received dexamethasone either without supplemental oxygen or with low-flow nasal cannula oxygen had a 60% higher risk of death. Although this study was observational, the investigators employed several statistical techniques to minimize potential bias, including propensity scoring and weighted analyses. Additionally, several subgroup and sensitivity analyses in this study confirmed the overall results.

Dexamethasone Dose

The RECOVERY platform trial studied the use of dexamethasone 6 mg once daily for up to 10 days,³ which is the currently recommended dose for hospitalized adults with COVID-19. Several other randomized controlled trials evaluated the role of higher doses of dexamethasone or other corticosteroids in hospitalized patients with different levels of respiratory support. The results of some key studies are summarized below.

Patients Who Received Conventional Oxygen or No Supplemental Oxygen

The RECOVERY platform trial included an additional study in which patients with COVID-19 and evidence of hypoxemia (i.e., receiving conventional supplemental oxygen or had oxygen saturation <92% on room air) were randomized to usual care plus high-dose dexamethasone (20 mg once daily for 5 days, then 10 mg once daily for 5 days or until hospital discharge, whichever came first) or usual care alone, which included low-dose dexamethasone (usually 6 mg once daily for 10 days).⁵ On May 11, 2022, the trial's independent data monitoring committee stopped enrolling participants receiving conventional oxygen therapy and those not receiving any supplemental oxygen. Among the 1,272 participants enrolled, 28-day mortality was higher in the high-dose dexamethasone arm than in the usual care arm (19% vs. 12%; rate ratio 1.59; 95% CI, 1.20–2.10; P = 0.0012).

Patients Who Received Noninvasive or Mechanical Ventilation

The COVID STEROID 2 trial investigated the use of different doses of corticosteroids in people with COVID-19 and severe hypoxemia.⁶ In this multicenter trial, hospitalized patients who required at least 10 L/min of oxygen or mechanical ventilation were randomized to receive up to 10 days of dexamethasone 6 mg once daily (n = 485) or dexamethasone 12 mg once daily (n = 497). The median number of days alive without life support at 28 days after randomization was 20.5 days in the dexamethasone 6 mg arm and 22.0 days in the dexamethasone 12 mg arm, yielding an adjusted mean difference of 1.3 days (95% CI, 0–2.6; P = 0.07). No differences between the arms were found for 28-or 90-day mortality. Although these conventional analyses did not quite reach statistical significance, a preplanned Bayesian analysis found that dexamethasone 12 mg had a higher probability of benefit and a lower probability of harm than dexamethasone 6 mg.⁷

In the COVIDICUS trial, patients with COVID-19 and acute hypoxemic respiratory failure were randomized to receive dexamethasone 6 mg once daily for 10 days (n = 276, of which 37 received placebo prior to release of results from the RECOVERY trial)³ or high-dose dexamethasone (i.e., 20 mg once daily for 5 days, then 10 mg once daily for 5 days; n = 270).⁸ At baseline, 98 patients were receiving mechanical ventilation, 114 were receiving continuous positive airway pressure, 10 were receiving noninvasive ventilation, 199 were receiving high-flow nasal cannula oxygen, and 125 were receiving standard oxygen therapy through a nonrebreather mask. There was no difference in 60-day mortality between the arms (HR 0.96, 95% CI, 0.69–1.33, P = 0.79).

The mixed results from these studies have led the Panel to continue to recommend 6 mg once daily as the preferred dose of dexamethasone in hospitalized patients with COVID-19 who require supplemental oxygen, including patients receiving noninvasive or mechanical ventilation. However, the Panel notes that both the conventional and Bayesian analyses conducted during the COVID STEROID 2 trial suggest that a dose of 12 mg might confer a benefit in patients who require noninvasive or mechanical ventilation.^{6,7}

Most patients in the COVID STEROID 2 trial did not receive additional immunomodulators beyond corticosteroids.⁶ Currently, there are no data from clinical trials that evaluated the safety and efficacy of using more or less than dexamethasone 6 mg once daily in combination with other immunomodulators to treat hospitalized adults with COVID-19.

Combination Immunomodulator Therapy

Using systemic corticosteroids in combination with other agents, including tocilizumab (see Interleukin-6 Inhibitors)^{9,10} or baricitinib (see Janus Kinase Inhibitors),¹¹ has been shown to have a clinical benefit in subsets of hospitalized patients with COVID-19, especially those with early critical illness and those with signs of systemic inflammation. For the Panel's recommendations on when to use dexamethasone with another immunomodulator, see <u>Therapeutic Management of Hospitalized Adults</u> With COVID-19.

See <u>Table 5a</u> for data from clinical trials that have evaluated the use of systemic corticosteroids in patients with COVID-19.

Systemic Corticosteroids Other Than Dexamethasone

Systemic corticosteroids other than dexamethasone, including hydrocortisone^{12,13} and methylprednisolone,^{14,15} have been studied for the treatment of COVID-19 in several randomized trials. Some of these trials were stopped early due to under-enrollment following the release of the RECOVERY trial results. Consequently, the sample size of many these trials was insufficient to assess efficacy (i.e., there were too few events to definitively confirm or exclude an effect, although many point estimates suggested a beneficial effect). Therefore, the evidence supporting the use of hydrocortisone or methylprednisolone for the treatment of COVID-19 is not as strong as the evidence supporting the use of dexamethasone. Based on the available evidence, the Panel has concluded the following:

- If dexamethasone is not available, alternative glucocorticoids (e.g., prednisone, methylprednisolone, hydrocortisone) can be used.
- For these drugs, the total daily dose equivalencies to dexamethasone 6 mg (orally or intravenously)¹⁶ are:
 - Prednisone 40 mg
 - Methylprednisolone 32 mg
 - Hydrocortisone 160 mg
- Half-life, duration of action, and frequency of administration vary among corticosteroids.
 - Long-acting corticosteroid: Dexamethasone; half-life 36 to 72 hours, administer once daily.
 - *Intermediate-acting corticosteroids:* Prednisone and methylprednisolone; half-life 12 to 36 hours, administer once daily or in 2 divided doses daily.
 - *Short-acting corticosteroid:* Hydrocortisone; half-life 8 to 12 hours, administer in 2 to 4 divided doses daily.

• Hydrocortisone is commonly used to manage septic shock in patients with COVID-19; see <u>Hemodynamics for Adults</u> for more information. Unlike other corticosteroids that have previously been studied in patients with acute respiratory distress syndrome, dexamethasone lacks mineralocorticoid activity and, thus, its effects on sodium balance and fluid volume are minimal.¹⁷

Monitoring, Adverse Effects, and Drug-Drug Interactions

- Clinicians should closely monitor patients with COVID-19 who are receiving dexamethasone for certain adverse effects (e.g., hyperglycemia, secondary infections, psychiatric effects, avascular necrosis).
- The use of systemic corticosteroids may increase the risk of opportunistic fungal infections (e.g., mucormycosis, aspergillosis) and reactivation of latent infections (e.g., hepatitis B virus infection, herpesvirus infections, strongyloidiasis, tuberculosis).¹⁸⁻²²
- Cases of severe and disseminated strongyloidiasis have been reported in patients with COVID-19 during treatment with tocilizumab and corticosteroids.^{23,24} Many clinicians would initiate empiric antiparasitic treatment (e.g., with the antiparasitic drug ivermectin) with or without serologic testing in patients who currently reside or who have previously resided in areas where *Strongyloides* is endemic (i.e., tropical, subtropical, or warm temperate areas).²⁵
- Using systemic corticosteroids with other immunosuppressants, such as tocilizumab or baricitinib, could theoretically increase the risk of secondary infections. However, clinical trials have reported no difference in the rates of secondary infections between patients who received corticosteroids in combination with another immunomodulatory agent and those who received corticosteroids alone.
- Dexamethasone is a moderate cytochrome P450 (CYP) 3A4 inducer. Therefore, it could reduce the concentration and potential efficacy of concomitant medications that are CYP3A4 substrates. Clinicians should carefully review a patient's concomitant medications to assess the potential for drug-drug interactions.

Considerations in Pregnancy

See <u>Pregnancy</u>, <u>Lactation</u>, <u>and COVID-19 Therapeutics</u> for the Panel's guidance regarding the use of dexamethasone during pregnancy and lactation.

Considerations in Children

Dexamethasone is recommended for hospitalized children with COVID-19 who require supplemental oxygen. See <u>Therapeutic Management of Hospitalized Children With COVID-19</u> for the Panel's recommendations. Methylprednisolone or another corticosteroid is recommended for the treatment of multisystem inflammatory syndrome in children (MIS-C). See <u>Therapeutic Management of Hospitalized Children With MIS-C, Plus a Discussion on MIS-A</u> for the Panel's recommendations.

References

- 1. WHO Rapid Evidence Appraisal for COVID-19 Therapies (REACT) Working Group. Association between administration of systemic corticosteroids and mortality among critically ill patients with COVID-19: a meta-analysis. *JAMA*. 2020;324(13):1330-1341. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32876694</u>.
- Li H, Yan B, Gao R, Ren J, Yang J. Effectiveness of corticosteroids to treat severe COVID-19: a systematic review and meta-analysis of prospective studies. *Int Immunopharmacol*. 2021;100:108121. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34492533</u>.
- 3. RECOVERY Collaborative Group. Dexamethasone in hospitalized patients with COVID-19. *N Engl J Med.* 2021;384(8):693-704. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32678530</u>.

- Crothers K, DeFaccio R, Tate J, et al. Dexamethasone in hospitalised COVID-19 patients not on intensive respiratory support. *Eur Respir J*. 2022;60(1):2102532. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34824060.
- RECOVERY Collaborative Group. Higher dose corticosteroids in patients admitted to hospital with COVID-19 who are hypoxic but not requiring ventilatory support (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2023;401(10387):1499-1507. Available at: https://pubmed.ncbi.nlm.nih.gov/37060915.
- 6. COVID STEROID 2 Trial Group. Effect of 12 mg vs 6 mg of dexamethasone on the number of days alive without life support in adults with COVID-19 and severe hypoxemia: the COVID STEROID 2 randomized trial. *JAMA*. 2021;326(18):1807-1817. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34673895.
- Granholm A, Munch MW, Myatra SN, et al. Dexamethasone 12 mg versus 6 mg for patients with COVID-19 and severe hypoxaemia: a pre-planned, secondary Bayesian analysis of the COVID STEROID 2 trial. *Intensive Care Med.* 2022;48(1):45-55. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34757439</u>.
- Bouadma L, Mekontso-Dessap A, Burdet C, et al. High-dose dexamethasone and oxygen support strategies in intensive care unit patients with severe COVID-19 acute hypoxemic respiratory failure: the COVIDICUS randomized clinical trial. *JAMA Intern Med.* 2022;182(9):906-916. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35788622</u>.
- 9. RECOVERY Collaborative Group. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2021;397(10285):1637-1645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33933206</u>.
- 10. REMAP-CAP Investigators. Interleukin-6 receptor antagonists in critically ill patients with COVID-19. *N Engl J Med.* 2021;384(16):1491-1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631065</u>.
- Marconi VC, Ramanan AV, de Bono S, et al. Efficacy and safety of baricitinib for the treatment of hospitalised adults with COVID-19 (COV-BARRIER): a randomised, double-blind, parallel-group, placebo-controlled Phase 3 trial. *Lancet Respir Med*. 2021;9(12):1407-1418. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34480861</u>.
- 12. Dequin PF, Heming N, Meziani F, et al. Effect of hydrocortisone on 21-day mortality or respiratory support among critically ill patients with COVID-19: a randomized clinical trial. *JAMA*. 2020;324(13):1298-1306. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32876689.
- Angus DC, Derde L, Al-Beidh F, et al. Effect of hydrocortisone on mortality and organ support in patients with severe COVID-19: the REMAP-CAP COVID-19 corticosteroid domain randomized clinical trial. *JAMA*. 2020;324(13):1317-1329. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32876697</u>.
- Corral-Gudino L, Bahamonde A, Arnaiz-Revillas F, et al. Methylprednisolone in adults hospitalized with COVID-19 pneumonia: an open-label randomized trial (GLUCOCOVID). *Wien Klin Wochenschr*. 2021;133(7-8):303-311. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33534047</u>.
- 15. Tang X, Feng YM, Ni JX, et al. Early use of corticosteroid may prolong SARS-CoV-2 shedding in nonintensive care unit patients with COVID-19 pneumonia: a multicenter, single-blind, randomized control trial. *Respiration*. 2021;100(2):116-126. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33486496</u>.
- Czock D, Keller F, Rasche FM, Häussler U. Pharmacokinetics and pharmacodynamics of systemically administered glucocorticoids. *Clin Pharmacokinet*. 2005;44(1):61-98. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/15634032</u>.
- Villar J, Ferrando C, Martínez D, et al. Dexamethasone treatment for the acute respiratory distress syndrome: a multicentre, randomised controlled trial. *Lancet Respir Med.* 2020;8(3):267-276. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32043986</u>.
- Garg D, Muthu V, Sehgal IS, et al. Coronavirus disease (COVID-19) associated mucormycosis (CAM): case report and systematic review of literature. *Mycopathologia*. 2021;186(2):289-298. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33544266</u>.

- Moorthy A, Gaikwad R, Krishna S, et al. SARS-CoV-2, uncontrolled diabetes and corticosteroids—an unholy trinity in invasive fungal infections of the maxillofacial region? A retrospective, multi-centric analysis. J Maxillofac Oral Surg. 2021;20(3):418-425. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33716414.
- Machado M, Valerio M, Álvarez-Uría A, et al. Invasive pulmonary aspergillosis in the COVID-19 era: an expected new entity. *Mycoses*. 2021;64(2):132-143. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33210776.
- 21. Chauvet P, Mallat J, Arumadura C, et al. Risk factors for invasive pulmonary aspergillosis in critically ill patients with coronavirus disease 2019-induced acute respiratory distress syndrome. *Crit Care Explor*. 2020;2(11):e0244. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33205046</u>.
- 22. Liu J, Wang T, Cai Q, et al. Longitudinal changes of liver function and hepatitis B reactivation in COVID-19 patients with pre-existing chronic hepatitis B virus infection. *Hepatol Res.* 2020;50(11):1211-1221. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32761993.
- 23. Lier AJ, Tuan JJ, Davis MW, et al. Case report: disseminated strongyloidiasis in a patient with COVID-19. *Am J Trop Med Hyg.* 2020;103(4):1590-1592. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32830642</u>.
- 24. Marchese V, Crosato V, Gulletta M, et al. Strongyloides infection manifested during immunosuppressive therapy for SARS-CoV-2 pneumonia. *Infection*. 2021;49(3):539-542. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32910321.
- 25. Stauffer WM, Alpern JD, Walker PF. COVID-19 and dexamethasone: a potential strategy to avoid steroid-related Strongyloides hyperinfection. *JAMA*. 2020;324(7):623-624. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32761166</u>.

Table 5a. Systemic Corticosteroids: Selected Clinical Trial Data

Last Updated: July 21, 2023

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for systemic corticosteroids. The studies summarized below are those that have had the greatest impact on the Panel's recommendations. Unless stated otherwise, the clinical trials listed below only included participants aged ≥ 18 years.

Methods	Results	Limitations and Interpretation			
RECOVERY: Open-Label RCT of Dexamethasone in Hospitalized Patients With COVID-19 in the United Kingdom ¹					
Key Inclusion Criterion	Participant Characteristics	Key Limitations			
 Hospitalized with suspected or laboratory-confirmed SARS-CoV-2 infection Key Exclusion Criteria Physician determination, based on patient's medical history, that risk of participation was too great An indication for corticosteroid theorem extends of the study. 	 Mean age 66 years; 64% men; 73% White 56% had ≥1 comorbidity; 24% with DM 89% had laboratory-confirmed SARS-CoV-2 infection Median of 7 days of DEX therapy At randomization: 16% received MV or ECMO 60% required supplemental oxygen but not MV 24% maximal as any lemental example 	 Open-label study Published data did not include results for key secondary endpoints (e.g., cause-specific mortality, need for renal replacement), AEs, and key subgroups (e.g., patients with comorbidities). Patients who required supplemental oxygen (but not MV) had variable severity of illness. It is unclear whether all patients in this group benefited from DEX or whether benefit was restricted to those 			
 therapy outside of the study Interventions DEX 6 mg IV or PO once daily plus SOC for up to 10 days or until discharge (n = 2,104) 	 24% required no supplemental oxygen Received RDV: <1% in both arms Received tocilizumab or sarilumab: 2% in DEX arm vs. 3% in SOC arm 	 requiring higher levels of supplemental oxygen. Patients aged >80 years were preferentially assigned to receive supplemental oxygen therapy (and not MV). 			
• SOC alone (n = $4,321$) Primary Endpoint	 Primary Outcome All-cause mortality at 28 days in DEX arm vs. SOC arm: All patients: 23% vs. 26% (age-adjusted rate ratio 0.83; 95%) 	 High mortality in this study may limit the generalizability of results to populations with lower baseline mortality. 			
All-cause mortality at 28 days	 Cl, 0.75–0.93; P < 0.001) Patients who required MV or ECMO at randomization: 29% vs. 41% (rate ratio 0.64; 95% Cl, 0.51–0.81) Patients who required supplemental oxygen but not MV at randomization: 23% vs. 26% (rate ratio 0.82; 95% Cl, 0.72–0.94) Patients who did not require supplemental oxygen at randomization: 18% vs. 14% (rate ratio 1.19; 95% Cl, 0.92–1.55) 	 Interpretation In hospitalized patients with severe COVID-19 who required supplemental oxygen, DEX reduced mortality at 28 days. The greatest benefit was seen in those receiving MV at randomization. There was no survival benefit for DEX in patients who did not require supplemental oxygen at randomization. 			

Methods	Results	Limitations and Interpretation		
CODEX: Open-Label RCT of Dexamethasone in Patients With Moderate or Severe ARDS and COVID-19 in Brazil ²				
Key Inclusion Criteria	Participant Characteristics	Key Limitations		
Confirmed or suspected SARS-CoV-2	Mean age 61 years; 63% men	Open-label study		
infection	Comorbidities in DEX arm vs. SOC arm:	Underpowered; enrollment stopped after release		
Received MV within 48 hours of meeting aritaria for moderate to source ADDC	 Obesity: 31% vs. 24% 	of data from the RECOVERY trial.		
criteria for moderate to severe ARDS (Pa0 ₂ /Fi0 ₂ \leq 200 mm Hg)	• DM: 38% vs. 47%	 Patients discharged before 28 days were not followed for rehospitalization or mortality. 		
Key Exclusion Criteria	 Vasopressor use: 66% in DEX arm vs. 68% in SOC arm 	 High mortality in this study may limit the 		
Received immunosuppressive drugs in past 21 days	 Mean Pa0₂/Fi0₂: 131 mm Hg in DEX arm vs. 133 mm Hg in SOC arm 	generalizability of results to populations with a lower baseline mortality.		
 Death expected within 24 hours 	 Median of 10 days of DEX therapy 	More than one-third of those randomized to		
Interventions	 No patients received RDV or tocilizumab 	receive SOC also received corticosteroids.		
• DEX 20 mg IV once daily for 5 days, then	 35% in SOC arm received corticosteroids for indications such as bronchospasm or septic shock 	InterpretationCompared with SOC alone, DEX increased the		
DEX 10 mg IV once daily for 5 days or until ICU discharge ($n = 151$)	Primary Outcome	number of days alive and free of MV over 28 days		
• SOC alone (n = 148)	• Mean number of days alive and free from MV by Day 28: 7 in	in patients with COVID-19 and moderate to severe ARDS.		
Primary Endpoint	DEX arm vs. 4 in SOC arm ($P = 0.04$)	Ando.		
Number of days alive and free from MV	Secondary Outcomes			
by Day 28	 No differences between arms by Day 28 in all-cause mortality (56% in DEX arm vs. 62% in SOC arm), number of ICU-free 			
Key Secondary Endpoints	days, or duration of MV or at Day 15 in score on 6-point OS			
All-cause mortality by Day 28	• Mean SOFA score at Day 7: 6.1 in DEX arm vs. 7.5 in SOC arm			
Number of ICU-free days by Day 28	(P = 0.004)			
Duration of MV by Day 28	Other Outcome			
Score on 6-point OS at Day 15SOFA score at Day 7	 Post hoc analysis of probability of death or MV by Day 15: 68% in DEX arm vs. 80% in SOC arm (OR 0.46) 			

Methods	Results	Limitations and Interpretation
Observational Cohort Study of Dexametha	Intensive Respiratory Support in the United States ³	
Key Inclusion Criterion	Participant Characteristics	Key Limitations
• Within 14 days of a positive SARS-CoV-2	Mean age 71 years; 95% men; 27% Black, 55% White	Retrospective observational study
test result	 77% did not receive IRS within 48 hours 	Because nearly all patients on MV or HFNC oxygen
Key Exclusion Criteria	83% admitted within 1 day after positive SARS-CoV-2 test	received DEX, analysis was restricted to patients
Recent receipt of corticosteroids	result	who did not receive IRS (i.e., those who received no supplemental oxygen or only low-flow nasal
• Receipt of IRS (defined as HFNC oxygen, NIV, or MV) within 48 hours	• Median duration of DEX for patients who did not receive IRS: 5 days for those not on supplemental oxygen at baseline vs. 6	cannula oxygen).There were differences between the arms in other
 Hospital LOS <48 hours 	days for those on low-flow nasal cannula oxygen	therapies received. The investigators attempted to
Interventions	 Received RDV: 43% of those who received DEX vs. 13% of those who did not 	account for this using different approaches (e.g.,
Corticosteroids (95% received DEX) administered within 48 hours of administered (n = 7.507)	 Received anticoagulants: 46% of those who received DEX vs. 10% of those who did not 	propensity scoring, weighted analyses, subgroup/ sensitivity analyses). Interpretation
admission (n = $7,507$)	Primary Outcome	 In hospitalized patients with COVID-19, the use
• No corticosteroids administered (n = 7,433)	• Risk of all-cause mortality at 90 days higher in those who	of DEX was not associated with a reduction in
Primary Endpoint	received DEX:	mortality among those who received low-flow
All-cause mortality at 90 days	those on low-flow nasal cannula oxygen: HR 1.59; 95% CI, 1.39–1.81	nasal cannula oxygen during the first 48 hours after admission, but it was associated with increased mortality among those who received
	 Those not on supplemental oxygen: HR 1.76; 95% CI, 1.47–2.12 	no supplemental oxygen during the first 48 hours after admission.
	 Those on low-flow nasal cannula oxygen: HR 1.08; 95% Cl, 0.86–1.36 	

Methods	Results	Limitations and Interpretation		
COVID STEROID 2: Blinded RCT of Dexamethasone 12 mg Versus 6 mg in Hospitalized Adults With COVID-19 and Severe Hypoxemia in Denmark, India, Sweden, and Switzerland ^{4,5}				
Key Inclusion Criteria	Participant Characteristics	Key Limitation		
Confirmed SARS-CoV-2 infection	 Median age 65 years; 31% women 	The randomized intervention period was		
• Requiring oxygen ≥10 L/min, NIV, CPAP,	• DM: 27% in 12 mg arm vs. 34% in 6 mg arm	<10 days for some patients because the trial		
or MV	• Median of 7 days from symptom onset to hospitalization in both	allowed up to 4 days of DEX before enrollment.		
Key Exclusion Criteria	arms	Interpretation		
• Treated with DEX >6 mg (or equivalent)	Received ICU care: 78% in 12 mg arm vs. 81% in 6 mg arm	• Among patients with COVID-19 and severe		
• Treated with corticosteroid within past	Oxygen requirements:	hypoxemia, the use of DEX 12 mg once daily did not result in more days alive without life		
5 days	• 54% on oxygen via nasal cannula or face mask (median flow rate	support at 28 days than DEX 6 mg once daily.		
Invasive fungal infection or active TB	23 L/min)	• A preplanned Bayesian analysis showed that		
Interventions	• 25% on NIV	DEX 12 mg had a higher probability of benefit		
• DEX 12 mg IV once daily for up to 10	• 21% on MV	and a lower probability of harm than DEX 6 mg. ⁵		
days (n = 497)	• 63% received RDV; 12% received IL-6 inhibitors or JAK inhibitors	ling.		
 DEX 6 mg IV once daily for up to 10 days (n = 485) 	Median of 7 days of DEX therapy in both arms			
- , ,	Primary Outcome			
Primary Endpoint	• Median number of days alive without life support at 28 days: 22.0			
 Number of days alive without life support (MV, circulatory support, or 	in 12 mg arm vs. 20.5 in 6 mg arm (adjusted mean difference 1.3 days; 95% Cl, 0.0–2.6; $P = 0.07$)			
kidney replacement therapy) at 28	 63.9% Bayesian probability of clinically important benefit and 			
days	0.3% Bayesian probability of clinically important bench and 0.3% Bayesian probability of clinically important harm for DEX 12			
Key Secondary Endpoints	mg			
Number of days alive without life	Secondary Outcomes			
support at 90 days	• At 90 days:			
 Number of days alive and out of hospital at 90 days 	 Median number of days alive without life support: 84 in 12 mg arm vs. 80 in 6 mg arm (P = 0.15) 			
Mortality at 90 days	• Median number of days alive and out of hospital: 62 in 12 mg arm			
Mortality at 28 days	vs. 48 in 6 mg arm ($P = 0.09$)			
SAEs at 28 days	 Mortality: 32% in 12 mg arm vs. 38% in 6 mg arm (adjusted relative risk 0.87; 99% Cl, 0.70–1.07; P = 0.09) 			
	At 28 days:			
	 Mortality: 27% in 12 mg arm vs. 32% in 6 mg arm (adjusted relative risk 0.86; 99% Cl, 0.68–1.08; P = 0.10) 			

Methods	Results	Limitations and Interpretation			
COVID STEROID 2: Blinded RCT of Dexamethasone 1 Sweden, and Switzerland ^{4,5} , continued	COVID STEROID 2: Blinded RCT of Dexamethasone 12 mg Versus 6 mg in Hospitalized Adults With COVID-19 and Severe Hypoxemia in Denmark, India, Sweden, and Switzerland ^{4,5} , continued				
	 SAEs, including septic shock and invasive fungal infections: 11% in 12 mg arm vs. 13% in 6 mg arm (adjusted relative risk 0.83; 99% Cl, 0.54–1.29; P = 0.27) 				
CAPE COVID : Double-Blind RCT of Hydrocortisone A	mong Critically III Patients With COVID-19 in France ⁶				
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
 Laboratory-confirmed SARS-CoV-2 infection or radiographically suspected COVID-19 with ≥1 of the following: MV with PEEP ≥5 cm H₂O PaO₂/FiO₂ <300 mm Hg and FiO₂ ≥50% on HFNC PaO₂/FiO₂ <300 mm Hg on reservoir mask oxygen Pulmonary severity index score >130 Key Exclusion Criteria 	 Mean age 62 years; 70% men; median BMI 28 96% had laboratory-confirmed SARS-CoV-2 infection Median symptom duration of 9–10 days 81% required MV at baseline Received vasopressors: 24% in hydrocortisone arm vs. 18% in placebo arm <5% received RDV or tocilizumab Median duration of treatment with study drug: 11 days in hydrocortisone arm vs. 12 days in placebo arm 	 Underpowered; enrollment stopped after release of data from the RECOVERY trial, resulting in limited power to detect differences between arms. Limited information about comorbidities Interpretation The use of hydrocortisone did not reduce the proportion of patients with COVID-19 and acute respiratory failure who 			
Septic shockDo-not-intubate orders	hydrocortisone arm vs. 13 days in placebo arm ($P = 0.25$) Primary Outcome	experienced treatment failure by Day 21.			
Continuous IV infusion of hydrocortisone 200 mg	• Treatment failure by Day 21: 42% in hydrocortisone arm vs. 51% in placebo arm ($P = 0.29$)				
 continuous iv infusion of hydrocortisone 200 mg per day for 7 days, then 100 mg per day for 4 days, then 50 mg per day for 3 days; if patient improved by Day 4, then IV infusion of hydrocortisone 200 mg per day for 4 days, then 100 mg per day for 2 days, then 50 mg per day for 2 days (n = 76) Placebo (n = 73) 	 Secondary Outcomes Need for intubation or prone positioning: no difference between arms (too few received ECMO or inhaled nitric oxide for comparison) Need for intubation in those not on MV at baseline: 50% in hydrocortisone arm vs. 75% in placebo arm 				
 Primary Endpoint Treatment failure (death or dependency on MV or high-flow oxygen) by Day 21 Key Secondary Endpoints 	 Proportion of patients with nosocomial infection by Day 28: no difference between arms Clinical status on Day 21: no difference between arms, but 15% died in hydrocortisone arm vs. 27% in placebo arm (P = 1000) 				
 Need for intubation, prone positioning, ECMO, or inhaled nitric oxide Nosocomial infection by Day 28 	 0.06) Discharged from ICU by Day 21: 57% in hydrocortisone arm vs. 44% in placebo arm; 23% in both arms still required MV 				

Methods	Results	Limitations and Interpretation			
CAPE COVID: Double-Blind RCT of Hydrocortisone Among Critically III Patients With COVID-19 in France ⁶ , continued					
 Clinical status on Day 21, as measured by a 5-item scale: Death In ICU and on MV Required high-flow oxygen therapy Required low-flow oxygen therapy Discharged from ICU 	 (P = 0.06) Discharged from ICU by Day 21: 57% in hydrocortisone arm vs. 44% in placebo arm; 23% in both arms still required MV 				
<u>REMAP-CAP</u>: Randomized, Open-Label, Adapt	ive Trial of Hydrocortisone in Patients With Severe COVID-19 ⁷				
 Key Inclusion Criteria Presumed or laboratory-confirmed SARS-CoV-2 infection ICU admission for respiratory or cardiovascular support Key Exclusion Criteria Presumed imminent death Systemic corticosteroid use >36 hours since ICU admission Interventions Hydrocortisone 50 mg IV every 6 hours for 7 days (n = 137) Shock-dependent hydrocortisone 50 mg IV every 6 hours for duration of shock for up to 28 days (n = 146) No hydrocortisone (n = 101) Primary Endpoint Number of days free of respiratory and 	 Participant Characteristics Mean age 60 years; 71% men; 53% White Mean BMI range of 29.7–30.9 for the 3 arms 50% to 64% required MV Primary Outcome Median number of days free of organ support by Day 21: no difference between arms (0 in each arm) Median adjusted ORs for hydrocortisone arms vs. no hydrocortisone arm: OR 1.43 (95% Crl, 0.91–2.27) with 93% Bayesian probability of superiority for fixed-dose hydrocortisone arm OR 1.22 (95% Crl, 0.76–1.94) with 80% Bayesian probability of superiority for shock-dependent hydrocortisone arm Key Secondary Outcome In-hospital mortality: no difference between arms (30% in fixed-dose hydrocortisone arm vs. 26% in shock-dependent hydrocortisone arm) 	 Key Limitations Open-label study Enrollment stopped after release of data from the RECOVERY trial. Interpretation The use of hydrocortisone did not increase the median number of days free of organ support in either the fixed-dose or the shock-dependent hydrocortisone arms, although early termination limited the power to detect differences between the arms. 			
cardiovascular organ support by Day 21 Key Secondary Endpoint • In-hospital mortality					

Methods	Results	Limitations and Interpretation				
Single-Blind RCT of Methylprednisolone i	ingle-Blind RCT of Methylprednisolone in Hospitalized Patients With COVID-19 Pneumonia in China ⁸					
Key Inclusion Criteria	Participant Characteristics	Key Limitations				
Laboratory-confirmed SARS-CoV-2	Mean age 56 years; 48% men	Small sample size				
infection	Median of 8 days from symptom onset to randomization	Terminated early because of decreasing incidence				
Pneumonia confirmed by chest CT scan	• At randomization, 71% receiving oxygen via nasal cannula	of COVID-19 pneumonia at study sites				
 Hospitalized on general ward for <72 	Primary Outcome	Interpretation				
hours Key Exclusion Criteria	 Clinical deterioration at 14 days: 4.8% in both arms (OR 1.0; 95% Cl, 0.134–7.442; P = 1.00) 	The incidence of clinical deterioration did not differ between the methylprednisolone and placebo				
Severe immunosuppression	Secondary Outcomes	arms.				
Corticosteroid use for other diseases	• No difference (all $P > 0.05$) between methylprednisolone arm					
Interventions	and placebo arm for:					
• Methylprednisolone 1 mg/kg per day IV	Clinical cure at 14 days: 51% vs. 58%					
for 7 days $(n = 43)$	Median number of days to clinical cure: 14 vs. 12					
• Saline $(n = 43)$	ICU admission: 4.8% in both arms					
Primary Endpoint	 In-hospital mortality: 0% vs. 2.3% 					
Clinical deterioration at 14 days	 Median number of days hospitalized: 17 vs. 13 					
Key Secondary Endpoints						
Clinical cure at 14 days						
Time to clinical cure						
ICU admission						
 In-hospital mortality 						
Number of days hospitalized						

Methods	Results	Limitations and Interpretation			
COVIDICUS: RCT of High-Dose Dexamethasone Versus Standard of Care Dexamethasone in Patients With COVID-19–Related Respiratory Failure in the ICU n France ⁹					
Key Inclusion Criteria	Participant Characteristics	Key Limitation			
 Laboratory-confirmed or suspected SARS-CoV-2 infection 	Median age 67 years; 76% menMedian of 9 days from symptom onset to randomization	Comparator arm was initially a placebo but was changed to a standard dose of DEX after the DECOVEDV trial results were released.			
ICU admission in past 48 hours	 81% with ≥1 comorbidity 	RECOVERY trial results were released.			
• Respiratory failure ($PaO_2 < 70 \text{ mm Hg}$,	 17% received RDV; <1% received tocilizumab 	Interpretation			
SpO ₂ <90% on room air, >30 breaths/ min, labored breathing, respiratory	Primary Outcome	Among ICU patients with COVID-19–related required to the patients with does DEX did not			
distress, or need for oxygen $\ge 6 \text{ L/min}$)	• All-cause mortality by Day 60: 26% in high-dose arm vs. 27%	respiratory failure, high-dose DEX did not significantly improve 60-day survival.			
Key Exclusion Criteria	in SOC arm (HR 0.96; 95% Cl, 0.69–1.33; <i>P</i> = 0.79)				
• Decision to limit life-sustaining treatment					
 Therapy with ≥0.5 mg/kg per day of prednisone equivalent for ≥3 weeks 					
 Active and untreated bacterial, fungal, or parasitic infection 					
Interventions					
• High dose: DEX 20 mg IV once daily for 5 days, then DEX 10 mg IV once daily for 5 days (n = 270)					
• SOC: DEX 6 mg IV once daily for 10 days (n = 239) or placebo (n = 37)					
Primary Endpoint					
All-cause mortality by Day 60					

Methods	Results	Limitations and Interpretation
RECOVERY: Open-Label RCT of Two Doses of Dexamethasone in Hospitalized Patients With COVID-19 in the United Kingdom, Asia, and Africa ¹⁰		
Key Inclusion Criteria	Note	Key Limitations
 Hospitalized with suspected or laboratory-confirmed SARS-CoV-2 infection 	 Enrollment for the subgroup of patients who received conventional oxygen or did not receive supplemental 	 Open-label study The larger RECOVERY trial stopped
 Sp0₂ <92% on room air 	oxygen was stopped prematurely due to safety	enrollment of patients in this subgroup
Key Exclusion Criteria	concerns. The results reported for this analysis only include patients from this subgroup.	(i.e., those who received conventional oxygen or did not receive supplemental
Physician determination, based on patient's medical	Participant Characteristics	oxygen) due to safety concerns.
history, that risk of participation was too great	• Mean age 61 years; 60% men; 54% Asian, 36% White	Interpretation
Contraindication to short-term corticosteroids	• 51% with ≥1 comorbidity; 19% with DM	In patients hospitalized with COVID-19
Suspected or confirmed influenza	 53% received ≥1 COVID-19 vaccine dose 	who had clinical hypoxemia (SpO ₂
 Current use of ritonavir-boosted nirmatrelvir (Paxlovid), ritonavir, or other potent CYP3A inhibitor 	 34% received RDV; 12% received tocilizumab or receipt of tocilizumab planned within 24 hours 	<92%) and did not require supplemental oxygen or required only conventional
Interventions	•	oxygen, use of high-dose DEX increased the risk of death and hyperglycemia
High-dose DEX 20 mg once daily plus SOC for 5 days	Primary Outcome	when compared with the use of standard
followed by 10 mg once daily for 5 days or until discharge, whichever came first (n = 659)	 All-cause mortality at 28 days: 19% in high-dose DEX arm vs. 12% in SOC arm (rate ratio 1.59; 95% Cl, 1.20–2.10; P = 0.0012) 	doses of corticosteroids.
 DEX 6 mg once daily plus SOC for 10 days or until discharge, whichever came first (n = 613) 	Secondary Outcomes	
	Time to discharge from hospital: 9 days in both arms	
Primary Endpoint	Composite of MV or death: 20% in high-dose DEX arm	
All-cause mortality at 28 days	vs. 13% in SOC arm (risk ratio 1.52; 95% Cl, 1.18–1.97)	
Key Secondary Endpoints	Safety Outcomes	
Time to discharge from hospital	Pneumonia not due to COVID-19: 10% in high-dose DEX	
 Composite of MV (including ECMO) or death 	arm vs. 6% in SOC arm (absolute difference 3.7%; 95%	
Key Safety Endpoints	CI, 0.7–6.6)	
 Infections other than COVID-19 	Hyperglycemia requiring new or increased insulin	
Metabolic complications	dose: 22% in high-dose DEX arm vs. 14% in SOC arm (absolute difference 7.4%; 95% Cl, 3.2–11.5)	

Key: AE = adverse event; ARDS = acute respiratory distress syndrome; BMI = body mass index; CPAP = continuous positive airway pressure; CT = computed tomography; CYP = cytochrome P450; DEX = dexamethasone; DM = diabetes mellitus; ECMO = extracorporeal membrane oxygenation; FiO_2 = fraction of inspired oxygen; HFNC = high-flow nasal cannula; ICU = intensive care unit; IL = interleukin; IRS = intensive respiratory support; IV = intravenous; JAK = Janus kinase; LOS = length of stay; MV = mechanical ventilation; NIV = noninvasive ventilation; OS = ordinal scale; the Panel = the COVID-19 Treatment Guidelines Panel; PaO_2 = arterial partial pressure of oxygen; PEEP = positive end-expiratory pressure; PO = oral; RCT = randomized controlled trial; RDV = remdesivir; SAE = serious adverse event; SOC = standard of care; SOFA = sequential organ failure assessment; SpO₂ = oxygen saturation; TB = tuberculosis

References

- 1. RECOVERY Collaborative Group. Dexamethasone in hospitalized patients with COVID-19. *N Engl J Med.* 2021;384(8):693-704. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32678530.
- Tomazini BM, Maia IS, Cavalcanti AB, et al. Effect of dexamethasone on days alive and ventilator-free in patients with moderate or severe acute respiratory distress syndrome and COVID-19: the CoDEX randomized clinical trial. *JAMA*. 2020;324(13):1307-1316. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32876695</u>.
- 3. Crothers K, DeFaccio R, Tate J, et al. Dexamethasone in hospitalised COVID-19 patients not on intensive respiratory support. *Eur Respir J*. 2022;60(1):2102532. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34824060.
- 4. COVID STEROID 2 Trial Group. Effect of 12 mg vs 6 mg of dexamethasone on the number of days alive without life support in adults with COVID-19 and severe hypoxemia: the COVID STEROID 2 randomized trial. *JAMA*. 2021;326(18):1807-1817. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34673895.
- Granholm A, Munch MW, Myatra SN, et al. Dexamethasone 12 mg versus 6 mg for patients with COVID-19 and severe hypoxaemia: a pre-planned, secondary Bayesian analysis of the COVID STEROID 2 trial. *Intensive Care Med.* 2022;48(1):45-55. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34757439.
- 6. Dequin PF, Heming N, Meziani F, et al. Effect of hydrocortisone on 21-day mortality or respiratory support among critically ill patients with COVID-19: a randomized clinical trial. *JAMA*. 2020;324(13):1298-1306. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32876689.
- Angus DC, Derde L, Al-Beidh F, et al. Effect of hydrocortisone on mortality and organ support in patients with severe COVID-19: the REMAP-CAP COVID-19 corticosteroid domain randomized clinical trial. *JAMA*. 2020;324(13):1317-1329. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32876697</u>.
- Tang X, Feng YM, Ni JX, et al. Early use of corticosteroid may prolong SARS-CoV-2 shedding in non-intensive care unit patients with COVID-19 pneumonia: a multicenter, single-blind, randomized control trial. *Respiration*. 2021;100(2):116-126. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33486496.
- Bouadma L, Mekontso-Dessap A, Burdet C, et al. High-dose dexamethasone and oxygen support strategies in intensive care unit patients with severe COVID-19 acute hypoxemic respiratory failure: the COVIDICUS randomized clinical trial. JAMA Intern Med. 2022;182(9):906-916. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35788622.
- RECOVERY Collaborative Group. Higher dose corticosteroids in patients admitted to hospital with COVID-19 who are hypoxic but not requiring ventilatory support (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2023;401(10387):1499-1507. Available at: https://pubmed.ncbi.nlm.nih.gov/37060915.

Inhaled Corticosteroids

Last Updated: December 20, 2023

Inhaled corticosteroids have been identified as potential COVID-19 therapeutic agents because of their targeted anti-inflammatory effects on the lungs. In addition, certain inhaled corticosteroids have been shown to impair viral replication of SARS-CoV-2¹ and downregulate the expression of the receptors used for cell entry.^{2,3} Several trials provide additional insights regarding the role of inhaled corticosteroids in treating outpatients with COVID-19. These trials are described below and in <u>Table 5b</u>.

Recommendations

- There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of inhaled corticosteroids for the treatment of COVID-19.
- There is insufficient evidence for the Panel to recommend either for or against the use of the combination of inhaled budesonide plus fluvoxamine for the treatment of COVID-19 in nonhospitalized patients.
- Patients with COVID-19 who are receiving an inhaled corticosteroid for an underlying condition should continue this therapy as directed by their health care provider (AIII).

Rationale

Compared to usual care, inhaled corticosteroid therapy decreased the time to recovery in 2 open-label randomized controlled trials in outpatients with mild symptoms of COVID-19.^{4,5} However, subsequent placebo-controlled, double-blind trials have shown that corticosteroid therapy did not reduce the duration of COVID-19 symptoms.⁶⁻⁸ The available evidence does not show that inhaled corticosteroid therapy reduces the risk of hospitalization or death due to COVID-19. However, the Panel acknowledges that there are areas of uncertainty. Studies conducted predominantly among unvaccinated patients have reported mixed results.

ACTIV-6 is the only randomized controlled trial of inhaled corticosteroid monotherapy that was conducted in a predominantly vaccinated population.⁸ In this study, treatment with inhaled fluticasone did not reduce the number of hospitalizations or health care visits or the time to sustained recovery. However, this study included patients who were at modest risk for complications from COVID-19. The median age of the patients was 45 years, and patients were not required to have a comorbidity to be included in the study.

The mixed results from these studies make it difficult to draw definitive conclusions about the benefit of using inhaled corticosteroids in people who are at high risk of disease progression. See <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for recommendations regarding therapies for high-risk outpatients.

The combination of inhaled budesonide plus oral fluvoxamine was studied in a large, double-blind, placebo-controlled, adaptive randomized trial in Brazil.⁹ Over 90% of the patients had received at least 2 doses of a COVID-19 vaccine. Treatment with this combination significantly reduced the incidence of the primary outcome, which was a composite of hospitalization or retention in an emergency setting for >6 hours. The proportion of patients who were hospitalized was the same in the treatment and placebo arms (0.9% vs. 1.1%), and the treatment did not significantly impact secondary outcomes such as health care attendance or the need for an emergency setting visit. It is unclear how the >6-hour emergency

setting outcome translates to other settings. In addition, the treatment with budesonide plus fluvoxamine was associated with significantly more adverse events.

For more information on these trials, see <u>Table 5b</u>.

No clinical trials have assessed the role of inhaled corticosteroids for the treatment of COVID-19 in hospitalized patients.

Monitoring, Adverse Effects, and Drug-Drug Interactions

Patients who are receiving inhaled corticosteroids may develop oral candidiasis.

Using a cytochrome P450 3A4 inhibitor, such as ritonavir-boosted nirmatrelvir (Paxlovid), with inhaled budesonide or fluticasone may lead to increased systemic absorption of the corticosteroid, which may result in systemic adverse effects from the corticosteroid.

Considerations in Pregnant People

There is insufficient evidence for the Panel to recommend either for or against the use of inhaled corticosteroids for the treatment of COVID-19 in people who are pregnant. Pregnant patients with COVID-19 who are receiving an inhaled corticosteroid for an underlying condition should continue this therapy as directed by their health care provider (**AIII**).

Considerations in Children

There is insufficient evidence for the Panel to recommend either for or against the use of inhaled corticosteroids for the treatment of COVID-19 in children. Children with COVID-19 who are receiving an inhaled corticosteroid for an underlying condition should continue this therapy as directed by their health care provider (AIII).

References

- 1. Matsuyama S, Kawase M, Nao N, et al. The inhaled steroid ciclesonide blocks SARS-CoV-2 RNA replication by targeting the viral replication-transcription complex in cultured cells. *J Virol*. 2020;95(1):e01648-20. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33055254.
- 2. Finney LJ, Glanville N, Farne H, et al. Inhaled corticosteroids downregulate the SARS-CoV-2 receptor ACE2 in COPD through suppression of type I interferon. *J Allergy Clin Immunol*. 2021;147(2):510-519.e5. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33068560.
- 3. Peters MC, Sajuthi S, Deford P, et al. COVID-19-related genes in sputum cells in asthma. Relationship to demographic features and corticosteroids. *Am J Respir Crit Care Med*. 2020;202(1):83-90. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32348692.
- 4. Ramakrishnan S, Nicolau DV Jr, Langford B, et al. Inhaled budesonide in the treatment of early COVID-19 (STOIC): a Phase 2, open-label, randomised controlled trial. *Lancet Respir Med*. 2021;9(7):763-772. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33844996.
- Yu LM, Bafadhel M, Dorward J, et al. Inhaled budesonide for COVID-19 in people at high risk of complications in the community in the UK (PRINCIPLE): a randomised, controlled, open-label, adaptive platform trial. *Lancet*. 2021;398(10303):843-855. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34388395.
- 6. Clemency BM, Varughese R, Gonzalez-Rojas Y, et al. Efficacy of inhaled ciclesonide for outpatient treatment of adolescents and adults with symptomatic COVID-19: a randomized clinical trial. *JAMA Intern Med.* 2022;182(1):42-49. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34807241.
- 7. Ezer N, Belga S, Daneman N, et al. Inhaled and intranasal ciclesonide for the treatment of COVID-19 in adult

outpatients: CONTAIN Phase II randomised controlled trial. *BMJ*. 2021;375:e068060. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34728476</u>.

- Boulware DR, Lindsell CJ, Stewart TG, et al. Inhaled fluticasone furoate for outpatient treatment of COVID-19. *N Engl J Med*. 2023;389(12):1085-1095. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37733308</u>.
- 9. Reis G, Dos Santos Moreira Silva EA, Medeiros Silva DC, et al. Oral fluvoxamine with inhaled budesonide for treatment of early-onset COVID-19: a randomized platform trial. *Ann Intern Med.* 2023;176(5):667-675. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37068273</u>.

Table 5b. Inhaled Corticosteroids: Selected Clinical Trial Data

Last Updated: December 20, 2023

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for inhaled corticosteroids. The studies summarized below are those that have had the greatest impact on the Panel's recommendations.

Methods	Results	Limitations and Interpretation
PRINCIPLE: Open-Label RCT of Inhaled Budesonide in Nonhospital	ized Patients With COVID-19 in the United Kingdo	om ¹
Key Inclusion Criteria	Participant Characteristics	Key Limitations
• Aged \geq 65 years or aged \geq 50 years with comorbidities	• Mean age 64.2 years; 52% women; 92%	Open-label trial
PCR-confirmed or suspected COVID-19	White	Primary endpoint of time to recovery
• ≤14 days of COVID-19 symptoms	81% with comorbidities	was based on patient self-report.
Key Exclusion Criteria	Median of 6 days from symptom onset to randomization	Interpretation
Already taking inhaled or systemic corticosteroids	randomization	• Inhaled budesonide reduced the time to
Unable to use an inhaler	Primary Outcomes	reported recovery but not the incidence of COVID-19–related hospitalization or
Contraindication for inhaled budesonide	• COVID-19–related hospitalization or death by Day 28: 6.8% in budesonide arm vs. 8.8% in	death.
Interventions	usual care arm (OR 0.75; 95% Crl, 0.55–1.03)	• The clinical significance of self-reported
• Usual care plus inhaled budesonide 800 μ g twice daily for 14 days (n = 1,069)	• Median time to reported recovery: 11.8 days in budesonide arm vs. 14.7 days in usual care	time to recovery in an open-label study is unclear.
• Usual care (n = 787)	arm (HR 1.21; 95% Crl, 1.08–1.36)	
Primary Endpoints		
COVID-19-related hospitalization or death by Day 28		
• Time to reported recovery up to 28 days from randomization		
STOIC: Open-Label, Phase 2 RCT of Inhaled Budesonide in Nonhos	pitalized Adults With Early COVID-19 in the Unite	d Kingdom ²
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Aged ≥18 years 	Mean age 45 years; 58% women	Small, open-label trial
• ≤7 days of COVID-19 symptoms	• 9% with CVD; 5% with DM	Trial was terminated early after
Key Exclusion Criteria	• 95% with positive SARS-CoV-2 RT-PCR result	statistical analysis determined that additional patients would not alter study
 Use of inhaled or systemic glucocorticoids in past 7 days Known allergy or contraindication to budesonide 	Median of 3 days from symptom onset to randomization	outcome.

Methods	Results	Limitations and Interpretation	
STOIC: Open-Label, Phase 2 RCT of Inhaled Budesonide in Nonhospitalized Adults With Early COVID-19 in the United Kingdom ² , continued			
Interventions Usual care plus inhaled budesonide 800 µg twice daily until 	 Primary Outcome COVID-19-related urgent care visit: 1% in 	• Secondary endpoint of time to recovery was based on patient self-report.	
symptom resolution (n = 70) • Usual care (n = 69)	budesonide arm vs. 14% in usual care arm (difference in proportion 0.131; 95% Cl, 0.043– 0.218; $P = 0.004$)	InterpretationIn adult outpatients with mild COVID-19,	
 Primary Endpoint COVID-19–related urgent care visit, including ED visit or hospitalization Key Secondary Endpoint 	 Secondary Outcome Median time to clinical recovery: 7 days in budesonide arm vs. 8 days in usual care arm 	 inhaled budesonide may reduce the need for urgent care, ED visit, or hospitalization. The clinical significance of self-reported time to recovery in an open-label study 	
Time to clinical recovery		is unclear.	
Phase 3, Double-Blind, Placebo-Controlled RCT of Inhaled Ci	clesonide in Nonhospitalized Patients With COVID-19 in	n the United States ³	
Key Inclusion Criteria	Participant Characteristics	Key Limitations	
 Aged ≥12 years Positive SARS-CoV-2 molecular or antigen diagnostic test result in previous 72 hours ≥1 symptoms of COVID-19 (i.e., fever, cough, dyspnea) 	 Mean age 43.3 years; 55.3% women; 86.3% White Mean BMI 29.4 22.3% with HTN; 7.5% with type 2 DM Higher rates of DM and asthma in ciclesonide arm 	 ED visit or hospital admission outcome was based on a small number of events. Primary endpoint of time to alleviation 	
Key Exclusion Criteria	Primary Outcome	of all symptoms was based on patient self-report.	
 Use of inhaled or intranasal corticosteroid within 14 days of enrollment or systemic corticosteroid within 90 days of enrollment 	 Median of 19 days in both arms for alleviation of all COVID-19–related symptoms (HR 1.08; 95% Cl, 0.84–1.38) 	 Interpretation Inhaled ciclesonide did not reduce the time to reported recovery in 	
Unable to use an inhaler	Secondary Outcomes	nonhospitalized patients with COVID-19.	
 Interventions Ciclesonide MDI 160 µg/actuation, administered as 2 actuations twice daily for 30 days (n = 197) 	 Alleviation of COVID-19–related symptoms by Day 30: 70.6% in ciclesonide arm vs. 63.5% in placebo arm (OR 1.28; 95% Cl, 0.84–19.7) 	• The robustness of the conclusion that inhaled ciclesonide reduced COVID-19- related ED visits or hospital admissions	
 Placebo MDI twice daily for 30 days (n = 203) Primary Endpoint 	• ED visit or hospital admission for COVID-19 by Day 30: 1.0% in ciclesonide arm vs. 5.4% in placebo arm (OR 0.18; 95% Cl, 0.04–0.85)	is uncertain. The small number of events is most likely due to the relatively low rate of comorbidities in the study population.	
 Time to alleviation of all COVID-19–related symptoms by Day 30 Key Secondary Endpoints 	 Hospital admission or death by Day 30: 1.5% in ciclesonide arm vs. 3.4% in placebo arm (OR 0.45; 95% Cl, 0.11–1.84) 	נווב סנעטאַ אָטאָטומעטוו.	
 Alleviation of COVID-19–related symptoms by Day 30 	• No deaths by Day 30 in either arm		

Methods	Results	Limitations and Interpretation
Phase 3, Double-Blind, Placebo-Controlled RCT of Inhaled Ciclesonide in Nonhospitalized Patients With COVID-19 in the United States ³ , continued		
ED visit or hospital admission for COVID-19 by Day 30		
Hospital admission or death by Day 30		
<u>CONTAIN</u> : Double-Blind RCT of Inhaled Ciclesonide Plus Intra	nasal Ciclesonide in Nonhospitalized Patients With (COVID-19 in Canada ⁴
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Aged ≥18 years 	Median age 35 years; 54% women; 61% White	Small study with a relatively young,
 Positive SARS-CoV-2 molecular diagnostic test result 	20% with comorbidities	healthy population
• \geq 1 symptoms of COVID-19 (i.e., fever, cough, shortness of	Primary Outcome	Interpretation
breath)	• Resolution of fever and all respiratory symptoms at	• The use of inhaled ciclesonide plus intranasal ciclesonide did not improve
 ≤5 days of COVID-19 symptoms 	Day 7: 40% in ciclesonide arm vs. 35% in placebo arm (adjusted risk difference 5.5%; 95% Cl, -7.8%	resolution of fever and respiratory
Key Exclusion Criteria	to 18.8%)	symptoms in nonhospitalized patients
 Receiving an inhaled corticosteroid or received a PO or IM corticosteroid within 7 days of enrollment 	Secondary Outcomes	with COVID-19.
Unable to use an inhaler	Resolution of fever and all respiratory symptoms	
 Has only nonrespiratory symptoms 	at Day 14: 66% in ciclesonide arm vs. 58% in	
Use of oxygen at home	placebo arm (adjusted risk difference 7.5%; 95% Cl, -5.9% to 20.8%)	
Vaccinated against COVID-19	Hospital admission by Day 14: 6% in ciclesonide	
Interventions	arm vs. 3% in placebo arm (adjusted risk	
 Ciclesonide MDI 600 µg/actuation plus intranasal ciclesonide 100 µg, both twice daily for 14 days (n = 105) 	difference 2.3%; 95% CI, -3.0% to 7.6%)	
- Saline placebo MDI plus intranasal saline, both twice daily for 14 days (n = 98)		
Primary Endpoint		
Resolution of fever and all respiratory symptoms at Day 7		
Key Secondary Endpoints		
Resolution of fever and all respiratory symptoms at Day 14		
Hospital admission by Day 14		

Methods	Results	Limitations and Interpretation
COVERAGE : Open-Label RCT of Inhaled Ciclesonide in Nonhospitalized Adults With COVID-19 in France ⁵		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Aged ≥60 years or aged ≥50 years with comorbidities 	Median age 63 years; 51% women	Small, open-label study
 Positive SARS-CoV-2 nasopharyngeal RT-PCR result or 	 72% with ≥1 comorbidities 	Interpretation
antigen test result	• 14% received ≥1 COVID-19 vaccine doses.	• In adult outpatients with mild COVID-19,
 ≤7 days of COVID-19 symptoms 	Primary Outcome	inhaled ciclesonide did not reduce the
Key Exclusion Criteria	Composite of hospitalization from any cause, need	proportion of patients who died, were
 Chronic use of inhaled corticosteroid therapy 	for COVID-19-related oxygen therapy at home, or	hospitalized, or required COVID-19– related oxygen therapy at home.
Unable to use an inhalation chamber	death by Day 14: 16% in ciclesonide arm vs. 12% in control arm	
 Ongoing therapy with a potent CYP3A4 inhibitor 		
Interventions	Secondary Outcome	
- Ciclesonide 160 μg via inhalation chamber, 2 puffs twice daily for 10 days (n = 110)	• Sustained alleviation of symptoms by Day 14: 54% in ciclesonide arm vs. 57% in control arm	
- Vitamin and trace element supplement, 2 capsules PO once or twice daily for 10 days (n = 107)		
Primary Endpoint		
 Composite of hospitalization from any cause, need for COVID-19–related oxygen therapy at home, or death by Day 14 		
Key Secondary Endpoint		
 Sustained alleviation of symptoms by Day 14 		

Methods	Results	Limitations and Interpretation
ACTIV-6: Decentralized, Placebo-Controlled, Platform RCT of Inhaled Fluticasone in Outpatients With COVID-19 in the United States ⁶		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Aged ≥30 years 	Median age 45 years; 63% women	• Low numbers of some clinical endpoints
 Positive SARS-CoV-2 nasopharyngeal RT-PCR result or 	 39% with BMI >30; 26% with HTN 	limited the ability to assess the effect of
antigen test result	• 65% received \geq 2 COVID-19 vaccine doses.	inhaled fluticasone on the key secondary endpoints.
• \leq 7 days of \geq 2 COVID-19 symptoms	Primary Outcome	Not all patients in the placebo arm
Key Exclusion Criterion	• No difference between arms in time to sustained	received a matched placebo.
Use of inhaled or systemic corticosteroids in preceding 30	recovery (HR 1.01; 95% Crl, 0.91–1.12)	Interpretation
days	Secondary Outcomes	• In adult outpatients with mild COVID-19,
Interventions	Hospitalization or death by Day 28: 0.5% in	inhaled fluticasone did not reduce the
 Inhaled fluticasone 200 µg once daily for 14 days (n = 656) 	fluticasone arm vs. 0.5% in placebo arm	time to sustained symptom recovery or
 Matching inhaled placebo (n = 350) or placebo from a different study (n = 271) 	• Urgent care visit, ED visit, or hospitalization by Day 28: 3.7% in fluticasone arm vs. in 2.1% placebo	the occurrence of urgent care visits, ED visits, or hospitalizations.
Primary Endpoint	arm (HR 1.9; 95% Crl, 0.8–3.5)	
• Time to sustained recovery (i.e., the last of 3 consecutive days without symptoms)	 Mean number of days unwell with ongoing symptoms: 11.2 in fluticasone arm vs. 11.3 in placebo arm 	
Key Secondary Endpoints		
Hospitalization or death by Day 28		
Urgent care visit, ED visit, or hospitalization by Day 28		
 Number of days unwell with ongoing symptoms 		

Methods	Results	Limitations and Interpretation
TOGETHER: Placebo-Controlled, Platform RCT of Oral Fluvoxamine and Inhaled Budesonide in Adults With Early-Onset COVID-19 in Brazil ⁷		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Aged ≥50 years or aged ≥18 years with comorbidities Laboratory-confirmed SARS-CoV-2 infection ≤7 days of COVID-19 symptoms 	 Median age 51 years; 61% women 42% with BMI >30 44% with HTN; 68% with ≥2 comorbidities 	 Multiple investigational treatments or placebos were evaluated simultaneously. Not all patients in the placebo arm received a matched placebo.
Key Exclusion Criteria	 94% received ≥2 COVID-19 vaccine doses. 	Interpretation
 Use of an SSRI Severe mental illness Cirrhosis, recent seizures, or severe ventricular cardiac arrythmia Interventions Fluvoxamine 100 mg PO twice daily plus inhaled budesonide 800 mcg twice daily for 10 days (n = 738) Placebo (n = 738; route, dosing frequency, and duration may have differed from fluvoxamine arm) Primary Endpoint Composite of ED observation >6 hours or hospitalization for COVID-19 by Day 28 Key Secondary Endpoints Hospitalization by Day 28 Health care attendance by Day 28 Any ED visit by Day 28 Occurrence of treatment-emergent AEs 	 Primary Outcome Composite of ED observation >6 hours or hospitalization by Day 28: 1.8% in fluvoxamine and budesonide arm vs. 3.7% in placebo arm (relative risk 0.50; 95% Crl, 0.25–0.92) Secondary Outcomes Hospitalization by Day 28: 0.9% in fluvoxamine plus budesonide arm vs. 1.1% in placebo arm Health care attendance by Day 28: 2.6% in fluvoxamine plus budesonide arm vs. 4.1% in placebo arm (relative risk 0.64; 95% Crl, 0.36– 1.11) Any ED visit by Day 28: 12.2% in fluvoxamine plus budesonide arm vs. 13.0% in placebo arm Occurrence of treatment-emergent AEs: 17.6% in fluvoxamine plus budesonide arm vs. 12.9% in placebo arm (relative risk 1.37; 95% Crl, 1.07– 1.75) Most AEs were grade 2 events. 	 Adult outpatients with mild COVID-19 who received a combination of fluvoxamine and inhaled budesonide had fewer ED observations >6 hours or hospitalizations for COVID-19 by Day 28 than those who received placebo. The use of fluvoxamine plus inhaled budesonide did not reduce the risk of hospitalization, health care attendance, or ED visits. It is difficult to define the clinical relevance of the >6-hour ED observation endpoint and apply it to practice settings in different countries. More AEs occurred with the use of fluvoxamine plus inhaled budesonide than with placebo.

Key: AE = adverse event; BMI = body mass index; CVD = cardiovascular disease; CYP = cytochrome P450; DM = diabetes mellitus; ED = emergency department; HTN = hypertension; IM = intramuscular; MDI = metered dose inhaler; the Panel = the COVID-19 Treatment Guidelines Panel; PCR = polymerase chain reaction; PO = oral; RCT = randomized controlled trial; RT-PCR = reverse transcription polymerase chain reaction; SSRI = selective serotonin reuptake inhibitor

References

- 1. Yu LM, Bafadhel M, Dorward J, et al. Inhaled budesonide for COVID-19 in people at high risk of complications in the community in the UK (PRINCIPLE): a randomised, controlled, open-label, adaptive platform trial. *Lancet*. 2021;398(10303):843-855. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34388395</u>.
- 2. Ramakrishnan S, Nicolau DV Jr, Langford B, et al. Inhaled budesonide in the treatment of early COVID-19 (STOIC): a Phase 2, open-label, randomised controlled trial. *Lancet Respir Med*. 2021;9(7):763-772. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33844996</u>.
- 3. Clemency BM, Varughese R, Gonzalez-Rojas Y, et al. Efficacy of inhaled ciclesonide for outpatient treatment of adolescents and adults with symptomatic COVID-19: a randomized clinical trial. *JAMA Intern Med.* 2022;182(1):42-49. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34807241.
- 4. Ezer N, Belga S, Daneman N, et al. Inhaled and intranasal ciclesonide for the treatment of COVID-19 in adult outpatients: CONTAIN Phase II randomised controlled trial. *BMJ*. 2021;375:e068060. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34728476</u>.
- 5. Duvignaud A, Lhomme E, Onaisi R, et al. Inhaled ciclesonide for outpatient treatment of COVID-19 in adults at risk of adverse outcomes: a randomised controlled trial (COVERAGE). *Clin Microbiol Infect*. 2022;28(7):1010-1016. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35304280</u>.
- 6. Boulware DR, Lindsell CJ, Stewart TG, et al. Inhaled fluticasone furoate for outpatient treatment of COVID-19. *N Engl J Med.* 2023;389(12):1085-1095. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37733308</u>.
- 7. Reis G, Dos Santos Moreira Silva EA, Medeiros Silva DC, et al. Oral fluvoxamine with inhaled budesonide for treatment of early-onset COVID-19: a randomized platform trial. *Ann Intern Med.* 2023;176(5):667-675. Available at: https://www.ncbi.nlm.nih.gov/pubmed/37068273.

Interleukin-6 Inhibitors

Last Updated: October 10, 2023

Interleukin (IL)-6 is a pleiotropic, proinflammatory cytokine produced by a variety of cell types, including lymphocytes, monocytes, and fibroblasts. Infection by SARS-CoV induces a dose-dependent production of IL-6 from bronchial epithelial cells.¹ COVID-19–associated systemic inflammation and hypoxemic respiratory failure can be associated with heightened cytokine release, as indicated by elevated blood levels of IL-6, C-reactive protein (CRP), D-dimer, and ferritin.²⁻⁴

The anti–IL-6 receptor monoclonal antibodies (mAbs) tocilizumab and sarilumab have been evaluated in hospitalized patients with COVID-19 who had systemic inflammation.

On December 21, 2022, the Food and Drug Administration (FDA) approved the use of intravenous (IV) tocilizumab for the treatment of COVID-19 in hospitalized adults who are receiving systemic corticosteroids and require supplemental oxygen, noninvasive ventilation (NIV), mechanical ventilation, or extracorporeal membrane oxygenation (ECMO).^{5,6}

Recommendations

• See <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> for the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations on the use of tocilizumab in hospitalized patients who require conventional oxygen, high-flow nasal cannula (HFNC) oxygen, NIV, or mechanical ventilation.

Additional Considerations

- If none of the recommended immunomodulatory therapies discussed in <u>Therapeutic Management</u> of <u>Hospitalized Adults With COVID-19</u> are available or feasible to use, **IV sarilumab** can be used in combination with dexamethasone (**CIIa**). Sarilumab is only commercially available as a subcutaneous (SUBQ) injection; see <u>Table 5e</u> for information regarding the preparation of an IV infusion using the SUBQ product.
- Tocilizumab and sarilumab should be used with caution in patients with COVID-19 who belong to populations that have not been adequately represented in clinical trials. This includes patients who are significantly immunosuppressed, such as those who have recently received other biologic immunomodulating drugs, and patients with any of the following:
 - Alanine transaminase levels >5 times the upper limit of normal
 - A high risk for gastrointestinal perforation
 - An uncontrolled serious bacterial, fungal, or non-SARS-CoV-2 viral infection
 - Absolute neutrophil counts $<500 \text{ cells}/\mu L$
 - Platelet counts <50,000 cells/µL
 - Known hypersensitivity to tocilizumab or sarilumab
- In both the REMAP-CAP and RECOVERY trials, 29% of patients received a second dose of tocilizumab at the discretion of their treating physician.^{7,8} However, there is insufficient evidence for the Panel to recommend either for or against the use of a second dose of tocilizumab for the treatment of COVID-19.
- Cases of severe and disseminated strongyloidiasis have been reported in patients with COVID-19

during treatment with tocilizumab and corticosteroids.^{9,10} Many clinicians would empirically initiate treatment for strongyloidiasis (e.g., with ivermectin), with or without serologic testing, in patients from areas where *Strongyloides* is endemic (i.e., tropical, subtropical, or warm temperate areas).¹¹

Rationale

The results of the RECOVERY and REMAP-CAP trials provide consistent evidence that tocilizumab, when administered as a second immunomodulatory agent in combination with a corticosteroid, offers a survival benefit in certain patients with COVID-19.^{7,8} Specifically, the patients who may benefit are those who are severely ill and require HFNC oxygen or NIV, those who are rapidly deteriorating with increasing oxygen needs, or those who are having a significant inflammatory response. In the REMAP-CAP trial, a long-term follow-up through 180 days confirmed that treatment with an anti–IL-6 receptor mAb improved survival among patients with severe to critical COVID-19.¹² However, the Panel found it challenging to determine which patients with COVID-19 who are receiving low-flow oxygen would benefit from receiving tocilizumab or sarilumab plus a corticosteroid (e.g., dexamethasone).

If none of the recommended immunomodulatory therapies are available or feasible to use, sarilumab may be used because the REMAP-CAP trial demonstrated that the use of tocilizumab and the use of sarilumab improved survival and reduced the duration of organ support.^{12,13} Sarilumab is currently only approved for use in the United States as a SUBQ injection.

Tocilizumab

Tocilizumab is a recombinant humanized anti–IL-6 receptor mAb approved by the FDA for use in certain hospitalized adults with COVID-19.⁵ It is also approved for use in patients with rheumatologic disorders and in patients with cytokine release syndrome induced by chimeric antigen receptor T cell therapy. Tocilizumab can be administered as an IV infusion or a SUBQ injection. Only the IV formulation of tocilizumab should be used for the treatment of COVID-19.

Clinical Data

Clinical data on the use of tocilizumab for the treatment of COVID-19, including data from several randomized trials and large observational studies, are summarized in <u>Table 5c</u>.

Two large randomized controlled trials, REMAP-CAP and RECOVERY, evaluated the use of tocilizumab in combination with standard-of-care corticosteroids.^{7,8} Both studies reported a statistically significant survival benefit from the use of tocilizumab in certain patients, including in patients who exhibited rapid respiratory decompensation associated with an inflammatory response.

REMAP-CAP enrolled critically ill patients who were within 24 hours of receiving respiratory support in an intensive care unit.⁷ At baseline, 29% of these patients were receiving HFNC oxygen, 42% were receiving NIV, and 29% were receiving mechanical ventilation. The patients were randomized to receive open-label tocilizumab or usual care. In-hospital mortality was 28% in the tocilizumab arm and 36% in the usual care arm. A follow-up analysis confirmed these findings.¹² At 180 days, mortality was 36% in the tocilizumab arm and 40% in the usual care arm.

The RECOVERY trial enrolled hospitalized patients with COVID-19 into an open-label platform trial that included several treatment options.⁸ A subset of all trial participants who had hypoxemia and CRP levels \geq 75 mg/L were offered enrollment into a second randomization that compared the use of tocilizumab to usual care. In this subgroup, the 28-day mortality was 31% in the tocilizumab arm and 35% in the usual care arm.

In contrast to the REMAP-CAP and RECOVERY trials, other randomized trials, including the

COVID-19 Treatment Guidelines

REMDACTA and EMPACTA trials, found that tocilizumab did not reduce all-cause mortality.^{14,15} In those trials, >80% of participants received corticosteroids as part of standard care, and most participants in the REMDACTA trial required NIV or HFNC oxygen.¹⁴

For additional findings from the REMAP-CAP and RECOVERY trials and the rationale for using tocilizumab in certain hospitalized patients who are exhibiting rapid respiratory decompensations due to COVID-19, see <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>.

Adverse Effects

The primary laboratory abnormalities reported in people receiving tocilizumab are elevated liver enzyme levels that appear to be dose dependent. Neutropenia or thrombocytopenia are uncommon. In randomized trials, no excess secondary infections were seen among patients who received combination therapy when compared with control patients. Additional adverse effects of tocilizumab, such as serious infections (e.g., tuberculosis, bacterial or fungal infections) and bowel perforation, have been reported.¹⁶⁻¹⁸

Considerations in Pregnant and Lactating People

See <u>Pregnancy</u>, <u>Lactation</u>, <u>and COVID-19 Therapeutics</u> for the Panel's guidance regarding the use of tocilizumab during pregnancy and lactation.

Considerations in Children

See <u>Therapeutic Management of Hospitalized Children With COVID-19</u> for the Panel's recommendations regarding the use of tocilizumab in children.

Drug Availability

On December 21, 2022, the FDA approved the use of IV tocilizumab for the treatment of COVID-19 in hospitalized adults who are receiving systemic corticosteroids and require supplemental oxygen, NIV, mechanical ventilation, or ECMO.⁵ In June 2021, the FDA issued an Emergency Use Authorization for the use of tocilizumab in combination with corticosteroids for the treatment of COVID-19 in hospitalized children aged ≥ 2 years who require supplemental oxygen, NIV, mechanical ventilation, or ECMO.⁶ If a patient's clinical signs or symptoms worsen or do not improve after the first dose of tocilizumab, 1 additional IV infusion of tocilizumab may be administered at least 8 hours after the initial infusion.

Sarilumab

Sarilumab is a recombinant humanized anti–IL-6 receptor mAb that is approved by the FDA for use in patients with rheumatoid arthritis. It is available as a SUBQ formulation and is not approved for the treatment of cytokine release syndrome.

Clinical Data

Clinical data on the use of sarilumab as a treatment for COVID-19 are summarized in Table 5c.

In the REMAP-CAP trial, the efficacy results for sarilumab were similar to those for tocilizumab.¹³ When compared with patients in the standard of care arm (n = 406), patients in the sarilumab arm (n = 485) had more organ support–free days (OR 1.50; 95% CrI, 1.13–2.00) and a greater likelihood of survival while hospitalized (OR 1.51; 95% CrI, 1.06–2.20). In-hospital mortality for the sarilumab arm and the standard of care arm was 33% and 37%, respectively, and mortality at 180 days was 33% and 40%, respectively.¹² A notable limitation to the sarilumab findings in the REMAP-CAP trial is that patients in the standard of care arm were enrolled earlier in the pandemic than those in the sarilumab arm.¹³ Randomization closed on November 2020 for the standard of care arm and continued through April 2021 for the sarilumab arm.

An adaptive, multinational, double-blind, randomized (2:2:1) trial compared the efficacy and safety of sarilumab 400 mg IV and sarilumab 200 mg IV to placebo in hospitalized patients with COVID-19.¹⁹ This trial did not show a clinical benefit of sarilumab in hospitalized patients who were receiving supplemental oxygen.

A similar adaptive study conducted in the United States in patients with severe and critical COVID-19 also failed to show a benefit of sarilumab.²⁰ In this placebo-controlled trial, mortality by Day 22 was reduced among the sarilumab recipients with critical COVID-19 pneumonia who required mechanical ventilation and received corticosteroids at baseline. However, due to the small sample size, this result was not statistically significant.

Adverse Effects

The primary laboratory abnormalities reported in people receiving sarilumab are transient or reversible elevations in liver enzyme levels that appear to be dose dependent and rare occurrences of neutropenia and thrombocytopenia.²¹ Additional adverse effects, such as serious infections (e.g., tuberculosis, bacterial or fungal infections) and bowel perforation have been reported, but only with long-term use of sarilumab.

Considerations in Pregnancy

There are insufficient data to determine whether the use of sarilumab is associated with an increased risk for major birth defects or miscarriage. As pregnancy progresses, mAbs are actively transported across the placenta (with the greatest transfer occurring during the third trimester), and immune responses in the exposed fetus may be affected.

Considerations in Children

See <u>Therapeutic Management of Hospitalized Children With COVID-19</u> for the Panel's recommendations regarding the use of sarilumab in children.

Drug Availability

IV administration of sarilumab is not approved by the FDA, but in clinical trials, single SUBQ sarilumab doses were modified to enable IV administration. See <u>Table 5e</u> for additional details.

References

- Yoshikawa T, Hill T, Li K, Peters CJ, Tseng CTK. Severe acute respiratory syndrome (SARS) coronavirusinduced lung epithelial cytokines exacerbate SARS pathogenesis by modulating intrinsic functions of monocyte-derived macrophages and dendritic cells. *J Virol.* 2009;83(7):3039-3048. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/19004938</u>.
- 2. Zhou F, Yu T, Du R, et al. Clinical course and risk factors for mortality of adult inpatients with COVID-19 in Wuhan, China: a retrospective cohort study. *Lancet*. 2020;395(10229):1054-1062. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32171076.
- 3. Huang C, Wang Y, Li X, et al. Clinical features of patients infected with 2019 novel coronavirus in Wuhan, China. *Lancet*. 2020;395(10223):497-506. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31986264</u>.
- Wang Z, Yang B, Li Q, Wen L, Zhang R. Clinical features of 69 cases with coronavirus disease 2019 in Wuhan, China. *Clin Infect Dis*. 2020;71(15):769-777. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32176772</u>.
- 5. Tocilizumab (Actemra) [package insert]. Food and Drug Administration. 2022. Available at: <u>https://www.accessdata.fda.gov/drugsatfda_docs/label/2022/125276s138lbl.pdf</u>.

- 6. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Actemra (tocilizumab). 2021. Available at: <u>https://www.fda.gov/media/150321/download</u>.
- 7. REMAP-CAP Investigators. Interleukin-6 receptor antagonists in critically ill patients with COVID-19. *N Engl J Med.* 2021;384(16):1491-1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631065</u>.
- 8. RECOVERY Collaborative Group. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2021;397(10285):1637-1645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33933206</u>.
- 9. Lier AJ, Tuan JJ, Davis MW, et al. Case report: disseminated strongyloidiasis in a patient with COVID-19. *Am J Trop Med Hyg.* 2020;103(4):1590-1592. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32830642</u>.
- Marchese V, Crosato V, Gulletta M, et al. Strongyloides infection manifested during immunosuppressive therapy for SARS-CoV-2 pneumonia. *Infection*. 2021;49(3):539-542. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32910321</u>.
- Stauffer WM, Alpern JD, Walker PF. COVID-19 and dexamethasone: a potential strategy to avoid steroidrelated Strongyloides hyperinfection. *JAMA*. 2020;324(7):623-624. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32761166</u>.
- Writing Committee for the REMAP-CAP Investigators. Long-term (180-day) outcomes in critically ill patients with COVID-19 in the REMAP-CAP randomized clinical trial. *JAMA*. 2023;329(1):39-51. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36525245</u>.
- 13. REMAP-CAP Investigators. Effectiveness of tocilizumab, sarilumab, and anakinra for critically ill patients with COVID-19: the REMAP-CAP COVID-19 immune modulation therapy domain randomized clinical trial. *medRxiv*. 2021;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2021.06.18.21259133v2</u>.
- Rosas IO, Diaz G, Gottlieb RL, et al. Tocilizumab and remdesivir in hospitalized patients with severe COVID-19 pneumonia: a randomized clinical trial. *Intensive Care Med.* 2021;47(11):1258-1270. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34609549</u>.
- Salama C, Han J, Yau L, et al. Tocilizumab in patients hospitalized with COVID-19 pneumonia. N Engl J Med. 2021;384(1):20-30. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33332779</u>.
- 16. Charan J, Dutta S, Kaur R, et al. Tocilizumab in COVID-19: a study of adverse drug events reported in the WHO database. *Expert Opin Drug Saf.* 2021;20(9):1125-1136. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34162299</u>.
- Peng J, Fu M, Mei H, et al. Efficacy and secondary infection risk of tocilizumab, sarilumab and anakinra in COVID-19 patients: a systematic review and meta-analysis. *Rev Med Virol*. 2022;32(3):e2295. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/34558756</u>.
- Shah R, Shah J, Gohil J, Revathi G, Surani S. Secondary infections in patients with COVID-19 pneumonia treated with tocilizumab compared to those not treated with tocilizumab: a retrospective study at a tertiary hospital in Kenya. *Int J Gen Med.* 2022;2022(15):2415-2425. Available at: https://pubmed.ncbi.nlm.nih.gov/35264878.
- Lescure FX, Honda H, Fowler RA, et al. Sarilumab in patients admitted to hospital with severe or critical COVID-19: a randomised, double-blind, placebo-controlled, Phase 3 trial. *Lancet Respir Med.* 2021;9(5):522-532. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33676590</u>.
- 20. Sivapalasingam S, Lederer DJ, Bhore R, et al. Efficacy and safety of sarilumab in hospitalized patients with COVID-19: a randomized clinical trial. *Clin Infect Dis*. 2022;75(1):e380-e388. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35219277.
- 21. Sarilumab (Kevzara) [package insert]. Food and Drug Administration. 2018. Available at: <u>https://www.accessdata.fda.gov/drugsatfda_docs/label/2018/761037s001lbl.pdf</u>.

COVID-19 Treatment Guidelines

Table 5c. Interleukin-6 Inhibitors: Selected Clinical Trial Data

Last Updated March 6, 2023

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for IL-6 inhibitors. The studies summarized below are those that have had the greatest impact on the Panel's recommendations.

Methods	Results	Limitations and Interpretation
<u>RECOVERY</u>: Open-Label RCT of Tocilizumab and Usual Care in Hospitalized Adults With COVID-19 in the United King		gdom ¹
Key Inclusion Criteria	Participant Characteristics	Key Limitations
• Evidence of COVID-19 progression ≤21 days after initial randomization to an intervention within the PECOV(EPV pretace) defined activity of the period activit	 Mean age 64 years; 67% men; 76% White 95% with PCR-confirmed SARS-CoV-2 infection 	 Arbitrary CRP ≥75 mg/L cutoff for enrollment
 within the RECOVERY protocol, defined as: SpO₂ <92% on room air or receipt of supplemental oxygen; and CRP ≥75 mg/L 	 At baseline: 45% on conventional oxygen 41% on HFNC oxygen or NIV 	 Difficult to define exact subset of patients in RECOVERY cohort who were subsequently selected for secondary randomization/tocilizumab trial
Key Exclusion Criterion Presence of non-SARS-CoV-2 infection	14% on MV82% on corticosteroids	InterpretationAmong hospitalized patients with
Interventions1 weight-based dose of tocilizumab	 Primary Outcomes 28-day all-cause mortality: 31% in tocilizumab arm vs. 35% in usual care arm (rate ratio 0.85; 95% Cl, 0.76–0.94; P = 0.003) 	COVID-19, hypoxemia, and elevated CRP levels, the use of tocilizumab was associated with a reduction in all-cause
 (maximum 800 mg) with possible second dose (n = 2,022) Usual care (n = 2,094) 	 28-day all-cause mortality among those who required MV at baseline: 49% in tocilizumab arm vs. 51% in usual care arm (risk ratio 0.93; 95% Cl, 0.74–1.18) 	mortality and a shorter time to hospital discharge.
Primary Endpoint	Secondary Outcomes	
 28-day all-cause mortality Key Secondary Endpoints Time to discharge from hospital within 28 	 Proportion discharged from hospital within 28 days: 57% in tocilizumab arm vs. 50% in usual care arm (rate ratio 1.22; 95% CI, 1.12–1.33; P < 0.0001) 	
 Among those not on MV at baseline, death or receipt of MV (including ECMO) within 28 days 	 Median time to hospital discharge: 19 days in tocilizumab arm vs. 28 days in usual care arm Proportion not on MV at baseline who died or required MV within 28 days: 35% in tocilizumab arm vs. 42% in usual care arm (rate ratio 0.84; 95% Cl, 0.77–0.92; P < 0.0001) 	

Methods	Results	Limitations and Interpretation
REMAP-CAP: Open-Label, Adaptive-Platform RCT of Tocilizumab and Sarilumab in Adults With COVID-19 in 21 Countries in		ries in Europe and North America ^{2,4} , cont'd
	 All-cause mortality at 180 days: 36% in tocilizumab arm vs. 40% in SOC arm (aHR 0.76; 95% Crl, 0.61–0.93) 	
	Sarilumab vs. SOC	
	 In-hospital survival: 67% in sarilumab arm vs. 63% in SOC arm (aOR 1.51; 95% Crl, 1.06–2.20) 	
	 All-cause mortality at 180 days: 33% in sarilumab arm vs. 40% in SOC arm (aHR 0.72; 95% Crl, 0.56–0.91) 	
	Pooled Tocilizumab and Sarilumab Arms vs. SOC Arm	
	 All-cause mortality at 180 days: 35% in pooled arms vs. 40% in SOC arm (aHR 0.74; 95% Crl, 0.61–0.90) 	
<u>COVACTA</u>: Double-Blind RCT of Tocilizumab i	n Hospitalized Adults With COVID-19 in 9 Countries in Europe and Nor	th America⁵
Key Inclusion Criteria	Participant Characteristics	Key Limitations
PCR-confirmed SARS-CoV-2 infection	Mean age 61 years; 70% men; 58% White	Modest power to detect differences in
• Hypoxemia	 30% on HFNC oxygen or NIV 	Day 28 clinical status
Bilateral chest infiltrates	• 38% on MV	 More patients received corticosteroids in placebo arm than tocilizumab arm.
Key Exclusion Criteria	 25% with multiorgan failure 	
Death imminent	Received corticosteroids at entry or during follow-up: 36% in	InterpretationThere was no difference between the
Presence of active non-SARS-CoV-2	tocilizumab arm vs. 55% in placebo arm	tocilizumab and placebo recipients in
infection	Primary Outcome	clinical status at Day 28 or survival.
Interventions	 No significant difference between arms in clinical status at Day 28 (P = 0.31) 	• The median time to hospital discharge
• 1 dose of tocilizumab 8 mg/kg with possible second dose, plus SOC (n = 294)	Secondary Outcomes	was significantly shorter in the tocilizumab arm than in the placebo arm.
• Placebo plus SOC (n = 144)	Median time to hospital discharge: 20 days in tocilizumab arm vs. 28	Although the result was not statistically
Primary Endpoint	days in placebo arm (HR 1.35; 95% Cl, 1.02–1.79)	significant, the tocilizumab arm had a shorter ICU LOS than the placebo arm.
Clinical status at Day 28, as measured by an OS	 Median ICU LOS: 9.8 days in tocilizumab arm vs. 15.5 days in placebo arm (difference 5.8 days; 95% Cl, –15.0 to 2.9) 	
Key Secondary Endpoints	• Mortality by Day 28: 20% in tocilizumab arm vs. 19% in placebo arm $(P = 0.94)$	
Time to hospital discharge	(1 - 0.07)	
• ICU LOS		
• Mortality by Day 28		

Methods	Results	Limitations and Interpretation
EMPACTA: Double-Blind RCT of Tocilizumab in Hospitalized Adults With COVID-19 in 6 Countries in North America, South America, and Africa ⁶		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 PCR-confirmed SARS-CoV-2 infection 	• Mean age 56 years; 59% men; 56% Hispanic/Latinx, 15% Black/	Moderate sample size
COVID-19 pneumonia	African American, 13% American Indian/Alaska Native	Interpretation
Key Exclusion Criteria	84% with elevated CRP	• In patients with COVID-19 pneumonia,
Death imminent	Concomitant medications:	tocilizumab reduced the likelihood of
 Receiving NIV or MV 	Corticosteroids: 80% in tocilizumab arm vs. 88% in placebo arm	progression to MV, ECMO, or death by Day 28 but did not reduce 28-day all-
Interventions	 RDV: 53% in tocilizumab arm vs. 59% in placebo arm 	cause mortality.
• 1 dose of tocilizumab 8 mg/kg with possible	Primary Outcome	
second dose, plus SOC ($n = 249$)	• Proportion who progressed to MV, ECMO, or death by Day 28: 12%	
• Placebo plus SOC (n = 128)	in tocilizumab arm vs. 19% in placebo arm (HR 0.56; 95% Cl, 0.33– 0.97; <i>P</i> = 0.04)	
Primary Endpoint	Secondary Outcomes	
 Progression to MV, ECMO, or death by Day 28 	 Median time to hospital discharge or readiness for discharge: 6.0 days in tocilizumab arm vs. 7.5 days in placebo arm (HR 1.16; 95%) 	
Key Secondary Endpoints	Cl, 0.91–1.48)	
Time to hospital discharge or readiness for discharge, as measured by an OS	 All-cause mortality by Day 28: 10.4% in tocilizumab arm vs. 8.6% in placebo arm (95% Cl, -5.2 to 7.8) 	
 All-cause mortality by Day 28 		

Methods	Results	Limitations and Interpretation
BACC Bay: Double-Blind RCT of Tocilizumab in Hospitalized Adults With COVID-19 in the United States ⁷		
 BACC Bay: Double-Blind RCT of Tocilizumab Key Inclusion Criteria Laboratory-confirmed SARS-CoV-2 infection ≥2 of the following conditions: Fever >38°C Pulmonary infiltrates Need for supplemental oxygen ≥1 of the following laboratory criteria: CRP ≥50 mg/L D-dimer >1,000 ng/mL LDH ≥250 U/L Ferritin >500 ng/mL Key Exclusion Criteria Receipt of supplemental oxygen at rate >10 L/min Recent use of biologic agents or small-molecule immunosuppressive therapy that increased risk for infection Interventions Tocilizumab 8 mg/kg plus usual care (n = 161) Placebo plus usual care (n = 81) 		 Limitations and Interpretation Key Limitations Wide confidence intervals due to small sample size and low event rates Few patients received RDV or corticosteroids Interpretation There was no benefit of tocilizumab in preventing MV or death, reducing the risk of clinical worsening, or reducing the time to discontinuation of oxygen. This could be due to the low rate of concomitant corticosteroid use among the study participants.
 Primary Endpoint Progression to MV or death by Day 28 Key Secondary Endpoints Clinical worsening by Day 28, as measured by an OS Discontinuation of supplemental oxygen among patients receiving it at baseline 		

Methods	Results	Limitations and Interpretation	
Double-Blind RCT of Sarilumab in Hospitalize	Double-Blind RCT of Sarilumab in Hospitalized Adults With Severe or Critical COVID-19 in 11 Countries in Europe, North America, South America, and Asia		
Key Inclusion Criteria	Participant Characteristics	Key Limitation	
COVID-19 pneumonia	• Median age 59 years; 63% men; 77% White, 36% Hispanic/Latinx	Moderate sample size	
Requirement for supplemental oxygen or	• 39% on HFNC oxygen, MV, or NIV	Interpretation	
intensive care	• 42% with BMI \geq 30; 43% with HTN; 26% with type 2 DM	Sarilumab did not reduce mortality	
Key Exclusion Criteria	• 20% received systemic corticosteroids before receiving intervention;	or time to clinical improvement in	
• Low probability of surviving or remaining at	63% received \geq 1 dose of corticosteroids during the study	hospitalized adults with COVID-19.	
study site	Primary Outcomes		
 Dysfunction of ≥2 organ systems and need for ECMO or renal replacement therapy 	 Median time to clinical improvement: 10 days in each sarilumab arm, 12 days in placebo arm 		
Interventions	• Sarilumab 200 mg arm vs. placebo arm: HR 1.03; 95% Cl, 0.75–		
• Sarilumab 400 mg IV (n = 173)	1.40; $P = 0.96$		
• Sarilumab 200 mg IV (n = 159)	• Sarilumab 400 mg arm vs. placebo arm: HR 1.14; 95% Cl, 0.84-		
• Placebo (n = 84)	1.54; $P = 0.34$		
Primary Endpoint	Secondary Outcome		
 Time to clinical improvement of ≥2 points on a 7-point OS 	 Survival to Day 29: 92% in placebo arm; 90% in sarilumab 200 mg arm (P = 0.63 vs. placebo); 92% in sarilumab 400 mg arm (P = 0.85 vs. placebo) 		
Key Secondary Endpoint			
Survival to Day 29			

Methods	Results	Limitations and Interpretation
REMDACTA : Double-Blind RCT of Tocilizumab and Remdesivir in Hospitalized Patients With Severe COVID-19 Pneumonia in Brazil, Russia, Spain, and the United States ⁹		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Aged ≥12 years PCR-confirmed SARS-CoV-2 infection Hospitalized with pneumonia confirmed by CXR or CT and requiring supplemental oxygen >6 L/min 	 Mean age 59 years, with 40% in tocilizumab arm and 34% in placebo arm aged ≥65 years; 63% men; 67% White Respiratory support: NIV or HFNC oxygen: 78% in tocilizumab arm vs. 83% in placebo arm MV or ECMO: 15% in tocilizumab arm vs. 11% in placebo arm 	 During the trial, primary outcome changed from clinical status at Day 28 to time to hospital discharge or readiness for discharge by Day 28 Imbalances in patient characteristics at baseline between arms
 Key Exclusion Criteria eGFR <30 mL/min ALT or AST >5 times ULN Presence of non-SARS-CoV-2 infection Treatment with antivirals, CCP, CQ, HCQ, or JAK inhibitors Interventions Up to 10 days of RDV plus: Tocilizumab 8 mg/kg IV with second dose within 8–24 hours if indicated (n = 434) Placebo (n = 215) Primary Endpoint Time to hospital discharge or readiness for discharge by Day 28 Key Secondary Endpoints Time to MV or death by Day 28 Clinical status at Day 14, as measured by an OS 	 • MV of ECMO. 15% in tocilizumab and vs. 11% in placebo and • Corticosteroid use: • At baseline: 83% in tocilizumab arm vs. 86% in placebo arm • During trial: 88% in each arm Primary Outcome • Time to hospital discharge or readiness for discharge by Day 28: 14 days in each arm (HR 0.97; 95% Cl, 0.78–1.19; <i>P</i> = 0.74) Secondary Outcomes • No difference between arms in: • Proportion who required MV or died by Day 28: 29% in each arm; time to death not evaluable (HR 0.98; 95% Cl, 0.72–1.34; <i>P</i> = 0.90) • Mean ordinal score for clinical status at Day 14: 2.8 in tocilizumab arm vs. 2.9 in placebo arm (<i>P</i> = 0.72) • Proportion who died by Day 28: 18% in tocilizumab arm vs. 20% in placebo arm; time to death not evaluable (HR 0.95; 95% Cl, 0.65–1.39; <i>P</i> = 0.79) 	 Possible underrepresentation of patients with rapidly progressive disease Interpretation Compared with placebo plus RDV, tocilizumab plus RDV did not shorten the time to discharge or readiness for discharge in patients with severe COVID-19 pneumonia. There was no difference in mortality between the arms.
• Time to death by Day 28		

Key: ALT = alanine transaminase; AST = aspartate transaminase; BMI = body mass index; CCP = COVID-19 convalescent plasma; CQ = chloroquine; CRP = C-reactive protein; CT = computed tomography; CXR = chest X-ray; DM = diabetes mellitus; ECMO = extracorporeal membrane oxygenation; eGFR = estimated glomerular filtration rate; HCQ = hydroxychloroquine; HFNC = high-flow nasal cannula; HTN = hypertension; ICU = intensive care unit; IL = interleukin; IV = intravenous; JAK = Janus kinase; LDH = lactate dehydrogenase; LOS = length of stay; MV = mechanical ventilation; NIV = noninvasive ventilation; OS = ordinal scale; the Panel = the COVID-19 Treatment Guidelines Panel; PCR = polymerase chain reaction; RCT = randomized controlled trial; RDV = remdesivir; SOC = standard of care; SpO₂ = oxygen saturation; ULN = upper limit of normal

References

- 1. RECOVERY Collaborative Group. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2021;397(10285):1637-1645. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33933206.
- 2. REMAP-CAP Investigators. Effectiveness of tocilizumab, sarilumab, and anakinra for critically ill patients with COVID-19: the REMAP-CAP COVID-19 immune modulation therapy domain randomized clinical trial. *medRxiv*. 2021;Preprint. Available at: https://www.medrxiv.org/content/10.1101/2021.06.18.21259133v2.
- 3. REMAP-CAP Investigators. Interleukin-6 receptor antagonists in critically ill patients with COVID-19. *N Engl J Med.* 2021;384(16):1491-1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631065</u>.
- 4. Writing Committee for the REMAP-CAP Investigators. Long-term (180-day) outcomes in critically ill patients with COVID-19 in the REMAP-CAP randomized clinical trial. *JAMA*. 2023;329(1):39-51. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36525245/</u>.
- 5. Rosas IO, Bräu N, Waters M, et al. Tocilizumab in hospitalized patients with severe COVID-19 pneumonia. *N Engl J Med.* 2021;384(16):1503-1516. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631066</u>.
- 6. Salama C, Han J, Yau L, et al. Tocilizumab in patients hospitalized with COVID-19 pneumonia. *N Engl J Med*. 2021;384(1):20-30. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33332779.
- 7. Stone JH, Frigault MJ, Serling-Boyd NJ, et al. Efficacy of tocilizumab in patients hospitalized with COVID-19. *N Engl J Med*. 2020;383(24):2333-2344. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33085857</u>.
- 8. Lescure FX, Honda H, Fowler RA, et al. Sarilumab in patients admitted to hospital with severe or critical COVID-19: a randomised, double-blind, placebo-controlled, Phase 3 trial. *Lancet Respir Med.* 2021;9(5):522-532. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33676590.
- 9. Rosas IO, Diaz G, Gottlieb RL, et al. Tocilizumab and remdesivir in hospitalized patients with severe COVID-19 pneumonia: a randomized clinical trial. *Intensive Care Med*. 2021;47(11):1258-1270. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34609549.

Janus Kinase Inhibitors

Last Updated: October 10, 2023

The primary mechanism of Janus kinase (JAK) inhibitors is interference with phosphorylation of the signal transducer and activator of transcription (STAT) proteins^{1,2} involved in vital cellular functions, including signaling, growth, and survival. JAK inhibitors are used as treatments for COVID-19 because they can prevent phosphorylation of key proteins involved in the signal transduction that leads to immune activation and inflammation (e.g., the cellular response to proinflammatory cytokines such as interleukin [IL]-6).³ Multiple JAK inhibitors are available, but only baricitinib and tofacitinib have been studied for the treatment of COVID-19.

In May 2022, the Food and Drug Administration (FDA) approved the use of baricitinib for the treatment of COVID-19 in hospitalized adults requiring supplemental oxygen, noninvasive ventilation (NIV), mechanical ventilation, or extracorporeal membrane oxygenation (ECMO).⁴

Recommendation

• See <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> for the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations on the use of baricitinib in hospitalized patients who require conventional oxygen, high-flow nasal cannula oxygen, NIV, or mechanical ventilation.

Additional Consideration

• If none of the recommended immunomodulatory therapies discussed in <u>Therapeutic Management</u> of <u>Hospitalized Adults With COVID-19</u> are available or feasible to use, **oral tofacitinib** can be used in combination with dexamethasone (**CIIa**).

Rationale

Several large randomized controlled trials have demonstrated that some patients who require supplemental oxygen and most patients who require oxygen through a high-flow device, NIV, or mechanical ventilation benefit from the use of dexamethasone in combination with a JAK inhibitor.

In the RECOVERY trial, baricitinib was associated with a survival benefit among hospitalized patients, with a treatment effect that was most pronounced among patients receiving NIV or oxygen supplementation through a high-flow device.⁵ The COV-BARRIER trial also demonstrated a survival benefit from baricitinib that was most pronounced among patients receiving high-flow oxygen or NIV.⁶ In the addendum to the COV-BARRIER trial, the benefit extended to patients receiving mechanical ventilation.⁷ Data from the ACTT-2⁸ and ACCT-4⁹ trials support the overall safety of baricitinib and the potential for benefit, but neither trial studied the drug in combination with dexamethasone as standard care.

The STOP-COVID study examined the use of tofacitinib in people with COVID-19 pneumonia who were not receiving NIV, mechanical ventilation, or ECMO at the time of enrollment.¹⁰ The study demonstrated a survival benefit in patients who received tofacitinib, nearly all of whom also received corticosteroids. If none of the other recommended immunomodulatory therapies are available or feasible to use, tofacitinib may be used as a substitute based on the findings from the STOP-COVID study.

Clinical trial data on the use of baricitinib and tofacitinib in patients with COVID-19 are summarized

below and in <u>Table 5d</u>.

Baricitinib

In May 2022, the FDA approved the use of baricitinib for the treatment of COVID-19 in hospitalized adults requiring supplemental oxygen, NIV, mechanical ventilation, or ECMO.⁴

Baricitinib is an oral JAK inhibitor that is selective for JAK1 and JAK2. It can modulate downstream inflammatory responses via JAK1/JAK2 inhibition and has exhibited dose-dependent inhibition of IL-6– induced STAT3 phosphorylation.¹¹ Baricitinib has postulated antiviral effects by blocking SARS-CoV-2 from entering and infecting lung cells.¹² See <u>Table 5d</u> for details on clinical trial data for baricitinib.

Tofacitinib

Tofacitinib is the prototypical JAK inhibitor, predominantly selective for JAK1 and JAK3, with modest activity against JAK2, and, as such, can block signaling from gamma-chain cytokines (e.g., IL-2, IL-4) and glycoprotein 130 proteins (e.g., IL-6, IL-11, interferons). It is an oral agent first approved by the FDA for the treatment of rheumatoid arthritis and has been shown to decrease levels of IL-6 in patients with this disease.¹³ Tofacitinib is also approved by the FDA for the treatment of psoriatic arthritis, juvenile idiopathic arthritis, and ulcerative colitis.¹⁴ See <u>Table 5d</u> for additional details on clinical trial data for tofacitinib.

Monitoring, Adverse Effects, and Drug-Drug Interactions

An FDA review of a large, randomized, safety clinical trial in people with rheumatoid arthritis compared tofacitinib to tumor necrosis factor inhibitors over 4 years and found that tofacitinib was associated with additional serious adverse events, including heart attack or stroke, cancer, blood clots, and death.¹⁵ Therefore, the FDA now requires new and updated warnings for drugs in the JAK inhibitor class, including baricitinib and tofacitinib. Data from randomized trials evaluating the safety of short-term use of JAK inhibitors in patients with COVID-19 have not revealed significant safety signals, including thrombosis.^{5,6,8-10} Because of the immunosuppressive effects of JAK inhibitors, all patients receiving either baricitinib or tofacitinib should be monitored for new infections.

Tofacitinib is a cytochrome P450 (CYP) 3A4 substrate. Dose modifications are required when the drug is administered with strong CYP3A4 inhibitors or when used with a moderate CYP3A4 inhibitor that is coadministered with a strong CYP2C19 inhibitor. Coadministration with a strong CYP3A4 inducer is not recommended. See <u>Table 5e</u> for kinase inhibitor drug characteristics and dosing information.

Considerations in Pregnant and Lactating People

See <u>Pregnancy</u>, <u>Lactation</u>, <u>and COVID-19 Therapeutics</u> for the Panel's guidance regarding the use of baricitinib during pregnancy and lactation. Pregnancy registries provide some outcome data on tofacitinib use during pregnancy for other conditions (e.g., ulcerative colitis, rheumatoid arthritis, psoriasis). Among the cases reported, pregnancy outcomes were similar to those among the general population.¹⁶⁻¹⁸

Considerations in Children

See <u>Therapeutic Management of Hospitalized Children With COVID-19</u> for the Panel's recommendations regarding the use of baricitinib or tofacitinib in children with COVID-19.

References

- 1. Babon JJ, Lucet IS, Murphy JM, Nicola NA, Varghese LN. The molecular regulation of Janus kinase (JAK) activation. *Biochem J*. 2014;462(1):1-13. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/25057888</u>.
- 2. Bousoik E, Montazeri Aliabadi H. "Do we know jack" about JAK? A closer look at JAK/STAT signaling pathway. *Front Oncol.* 2018;8:287. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30109213</u>.
- 3. Zhang W, Zhao Y, Zhang F, et al. The use of anti-inflammatory drugs in the treatment of people with severe coronavirus disease 2019 (COVID-19): the perspectives of clinical immunologists from China. *Clin Immunol*. 2020;214:108393. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32222466.
- 4. Baricitinib (Olumiant) [package insert]. Food and Drug Administration. 2022. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2022/207924s006lbl.pdf.
- RECOVERY Collaborative Group. Baricitinib in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial and updated meta-analysis. *Lancet*. 2022;400(10349):359-368. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35908569</u>.
- Marconi VC, Ramanan AV, de Bono S, et al. Efficacy and safety of baricitinib for the treatment of hospitalised adults with COVID-19 (COV-BARRIER): a randomised, double-blind, parallel-group, placebo-controlled Phase 3 trial. *Lancet Respir Med*. 2021;9(12):1407-1418. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34480861.
- Ely EW, Ramanan AV, Kartman CE, et al. Efficacy and safety of baricitinib plus standard of care for the treatment of critically ill hospitalised adults with COVID-19 on invasive mechanical ventilation or extracorporeal membrane oxygenation: an exploratory, randomised, placebo-controlled trial. *Lancet Respir Med.* 2022;10(4):327-336. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35123660.
- 8. Kalil AC, Patterson TF, Mehta AK, et al. Baricitinib plus remdesivir for hospitalized adults with COVID-19. *N Engl J Med.* 2021;384(9):795-807. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33306283</u>.
- Wolfe CR, Tomashek KM, Patterson TF, et al. Baricitinib versus dexamethasone for adults hospitalised with COVID-19 (ACTT-4): a randomised, double-blind, double placebo-controlled trial. *Lancet Respir Med*. 2022;10(9):888-899. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35617986</u>.
- Guimarães PO, Quirk D, Furtado RH, et al. Tofacitinib in patients hospitalized with COVID-19 pneumonia. N Engl J Med. 2021;385(5):406-415. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133856</u>.
- 11. McInnes IB, Byers NL, Higgs RE, et al. Comparison of baricitinib, upadacitinib, and tofacitinib mediated regulation of cytokine signaling in human leukocyte subpopulations. *Arthritis Res Ther.* 2019;21(1):183. Available at: https://www.ncbi.nlm.nih.gov/pubmed/31375130.
- 12. Richardson P, Griffin I, Tucker C, et al. Baricitinib as potential treatment for 2019-nCoV acute respiratory disease. *Lancet*. 2020;395(10223):e30-e31. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32032529.
- Migita K, Izumi Y, Jiuchi Y, et al. Effects of Janus kinase inhibitor tofacitinib on circulating serum amyloid A and interleukin-6 during treatment for rheumatoid arthritis. *Clin Exp Immunol*. 2014;175(2):208-214. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/24665995</u>.
- 14. Tofacitinib (Xeljanz) [package insert]. Food and Drug Administration. 2019. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2019/203214s024,208246s010lbl.pdf.
- 15. Food and Drug Administration. FDA requires warnings about increased risk of serious heart-related events, cancer, blood clots, and death for JAK inhibitors that treat certain chronic inflammatory conditions. 2021. Available at: <u>https://www.fda.gov/drugs/drug-safety-and-availability/fda-requires-warnings-about-increased-risk-serious-heart-related-events-cancer-blood-clots-and-death</u>. Accessed September 29, 2023.
- Clowse ME, Feldman SR, Isaacs JD, et al. Pregnancy outcomes in the tofacitinib safety databases for rheumatoid arthritis and psoriasis. *Drug Saf.* 2016;39(8):755-762. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/27282428</u>.
- 17. Mahadevan U, Dubinsky MC, Su C, et al. Outcomes of pregnancies with maternal/paternal exposure in the

COVID-19 Treatment Guidelines

tofacitinib safety databases for ulcerative colitis. *Inflamm Bowel Dis*. 2018;24(12):2494-2500. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29982686</u>.

 Wieringa JW, van der Woude CJ. Effect of biologicals and JAK inhibitors during pregnancy on healthrelated outcomes in children of women with inflammatory bowel disease. *Best Pract Res Clin Gastroenterol*. 2020;44-45:101665. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32359679</u>.

Table 5d. Janus Kinase Inhibitors: Selected Clinical Trial Data

Last Updated: August 8, 2022

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for kinase inhibitors. The studies summarized below are those that have had the greatest impact on the Panel's recommendations.

The information in this table may include data from preprints or articles that have not been peer reviewed. This section will be updated as new information becomes available. Please see <u>ClinicalTrials.gov</u> for more information on clinical trials evaluating kinase inhibitors.

Methods	Results	Limitations and Interpretation
<u>RECOVERY</u>: Open-Label RCT of Baricitinib Ve	rsus Usual Care in the United Kingdom ¹	
Key Inclusion Criterion	Participant Characteristics	Key Limitation
Hospitalized with suspected or laboratory-	Mean age 58 years; 66% men; 80% White	 Open-label study
confirmed SARS-CoV-2 infection	Median duration of symptoms at enrollment: 9 days	Interpretation
Key Exclusion Criteria	• 91% with laboratory-confirmed SARS-CoV-2 infection	 In patients hospitalized for COVID-19,
• eGFR <15 mL/min/1.73m ²	• At baseline:	BAR reduced the risk of death.
• ANC <500 cells/mm ³	95% received corticosteroids	
Evidence of active TB	 23% received tocilizumab 20% received remdesivir 	
Interventions	 42% received ≥1 COVID-19 vaccine 	
 BAR 4 mg PO daily for 10 days or until discharge, whichever comes first (n = 4,148) SOC (n = 4,008) 	 6% no supplemental oxygen required 68% simple oxygen 24% NIV 	
	• 3% MV	
Primary Endpoint28-day mortality	Primary Outcome	
	• 28-day mortality: 12% in BAR arm vs. 14% in SOC arm (age-adjusted rate ratio 0.87; 95% Cl, 0.77–0.98; <i>P</i> = 0.028)	
Key Secondary Endpoints		
• Time to discharge from hospital	Secondary Outcomes	
Composite of MV, ECMO, or death	 Discharge within 28 days: 80% in BAR arm vs. 78% in SOC arm (age-adjusted rate ratio 1.10; 95% CI, 1.04–1.15; P = 0.002) 	
	Median time to discharge: 8 days in both arms	
	• Composite of MV, ECMO, or death: 16% in BAR arm vs. 17% in SOC arm (age-adjusted risk ratio 0.89; 95% Cl, 0.81–0.98; $P = 0.016$)	

Methods	Results	Limitations and Interpretation	
<u>COV-BARRIER</u> : Double-Blind, Placebo-Controlled, Randomized Trial of Baricitinib in Hospitalized Adults in 12 Countries in Asia, Europe, North America, and South America ²			
Key Inclusion Criteria	Participant Characteristics	Key Limitation	
Laboratory-confirmed SARS-CoV-2 infection	 Mean age 58 years; 63% men 	• Results from the ACTT-2 trial prompted a	
• Evidence of pneumonia or active, symptomatic COVID-19	 79% received corticosteroids; 19% received RDV; 13% received oxygen but no steroids 	protocol amendment limiting enrollment to participants who required baseline	
• \geq 1 elevated inflammatory marker (CRP,	Primary Outcome	oxygen.	
D-dimer, LDH, or ferritin)	• Clinical progression or death by Day 28: 28% in BAR arm vs. 31% in	Interpretation	
Key Exclusion Criteria	placebo arm (OR 0.85; 95% Cl, 0.67–1.08; $P = 0.18$)	 Although the primary outcome of clinical progression or death was not 	
• MV or ECMO	Secondary Outcomes	significantly different between arms,	
 Receipt of immunosuppressants (including high-dose steroids) 	 Mortality by Day 28: 8% in BAR arm vs. 13% in placebo arm (HR 0.57; 95% Cl, 0.41–0.78; P = 0.0018) 	treatment with BAR plus SOC was associated with reduced mortality in	
Prior receipt of CCP or IVIG	• Mortality by Day 28 for those receiving corticosteroids at baseline: 9%	hospitalized adults with COVID-19 who	
• ANC <1,000 cells/µL	in BAR arm vs. 14% in placebo arm (HR 0.63; 95% Cl, 0.45–0.89)	were not receiving MV (see addendum below for results for patients who	
• ALC <200 cells/µL		required MV or ECMO).	
• ALT or AST >5 times ULN		• For patients receiving oxygen but not	
• eGFR <30 mL/min		steroids at baseline, the primary and	
Interventions		secondary outcomes were similar to the outcomes for the overall study	
• BAR 4 mg PO once daily for up to 14 days (n = 764)		population.	
• Placebo (n = 761)			
Primary Endpoint			
Clinical progression or death by Day 28			
Key Secondary Endpoint			
Mortality by Day 28			

Methods	Results	Limitations and Interpretation
<u>COV-BARRIER Addendum</u>: Double-Blind, Placebo-Controlled, Randomized Trial of Baricitinib in Hospitalized Adults on Mechanical Ventilation or Extracorporeal Membrane Oxygenation in Argentina, Brazil, Mexico, and the United States ³		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
• Laboratory-confirmed SARS-CoV-2 infection	• Mean age 59 years; 55% men	• Very small sample size, exploratory
• Evidence of pneumonia or active,	 86% received corticosteroids; 2% received RDV 	analysis
symptomatic COVID-19	Outcomes	High mortality in placebo arm
• ≥1 elevated inflammatory marker (CRP,	• Mortality at Day 28: 39% in BAR arm vs. 58% in placebo arm (HR	Interpretation
D-dimer, LDH, or ferritin) • MV or ECMO at baseline	0.54; 95% Cl, 0.31–0.96; <i>P</i> = 0.030)	In critically ill patients with COVID-19
	Number of ventilator-free days and duration of hospitalization: no	receiving MV or ECMO, treatment with BAR and SOC (including corticosteroids)
Key Exclusion Criteria	significant difference between arms	may decrease mortality.
 Receipt of immunosuppressants (including high-dose steroids) 		
 Prior receipt of CCP or IVIG 		
• ANC <1,000 cells/µL		
• ALC <200 cells/µL		
• ALT or AST >5 times ULN		
• eGFR <30 mL/min		
Interventions		
• BAR 4 mg PO once daily for up to 14 days $(n = 51)$		
• Placebo (n = 50)		
Key Endpoints		
Mortality at Day 28		
Number of ventilator-free days		
Duration of hospitalization		

Methods	Results	Limitations and Interpretation
ACTT-2: Double-Blind, Placebo-Controlled, Randomized Trial of Baricitinib Plus Remdesivir in Hospitalized Adults With COVID-19 in 8 Countries in Europe, North America, and Asia ⁴		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Positive SARS-CoV-2 PCR result 	 Mean age 55 years; 63% men; 48% White, 15% Black, 10% Asian 	 Not powered to detect difference in
• Radiographic infiltrates, $SpO_2 \le 94\%$ on	• At baseline:	mortality between arms
room air, or requiring supplemental oxygen,	 13% no supplemental oxygen required 	 Steroids not part of SOC
MV, or ECMO	 55% conventional oxygen 	Interpretation
Key Exclusion Criteria	 21% HFNC oxygen or NIV 	• Compared with RDV alone, BAR plus RDV
 Use of glucocorticoids for COVID-19 indications 	• 11% MV or ECMO	reduced recovery time and improved clinical status, particularly for patients
 ALT or AST >5 times ULN 	Primary Outcomes	who received HFNC oxygen or NIV at
Impaired renal function	 Median time to recovery: 7 days in BAR arm vs. 8 days in placebo arm (rate ratio 1.16; 95% Cl, 1.01–1.32; P = 0.03) 	baseline.
Interventions	 Median time to recovery for those receiving HFNC oxygen or NIV: 10 	
• BAR 4 mg PO once daily for 14 days or until discharge, plus RDV for 10 days or until	days in BAR arm vs. 18 days in placebo arm (rate ratio for recovery 1.51; 95% Cl, 1.10–2.08)	
discharge (n = 515)	Secondary Outcomes	
• Placebo plus RDV ($n = 518$)	 Improvement in clinical status at Day 15: greater in BAR arm vs. 	
Primary Endpoint	placebo arm (OR 1.3; 95% Cl, 1.0–1.6)	
Time to recovery by Day 28	• Mortality at Day 28: 5% in BAR arm vs. 8% in placebo arm (HR 0.65;	
Key Secondary Endpoints	95% Cl, 0.39–1.09)	
• Clinical status at Day 15 as measured by OS		
Mortality at Day 28		

Methods	Results	Limitations and Interpretation
ACTT-4: Double-Blind, Placebo-Controlled, Randomized Trial of Remdesivir With Baricitinib Versus Dexamethasone for Hospitalized Patients Requiring Supplemental Oxygen in Japan, Mexico, Singapore, South Korea, and the United States ⁵		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Hospitalized and requiring conventional oxygen, HFNC oxygen, or NIV Laboratory-confirmed SARS-CoV-2 infection 	 Median age 58 years; 58% men; 58% White, 34% Hispanic/Latinx At baseline: 85% low-flow oxygen 	 Study closed before completing enrollment of 1,500 as it was unlikely to show a difference between arms.
Key Exclusion Criterion	 15% HFNC oxygen or NIV 	 Not powered to analyze differences
 Receipt of CCP or >1 dose DEX 6 mg (or equivalent) or BAR before enrollment 	 Mean duration of symptoms at enrollment: 8 days 	between ordinal score subgroups HFNC oxygen or NIV at baseline.
Interventions	Primary Outcome	• Few patients died or required MV, which
 RDV IV for ≤10 days plus BAR 4 mg PO daily for ≤14 days plus DEX placebo IV (n = 516) 	 MV-free survival by Day 29: 87% in BAR arm vs. 88% in DEX arm (risk difference 0.6%; 95% Cl, -3.6% to 4.8%; P = 0.91) 	may have decreased the power to detect a difference between arms for MV-free
• RDV IV for ≤ 10 days plus BAR placebo PO	Secondary Outcomes	survival.
plus DEX 6 mg IV daily \leq 10 days (n = 494)	• Improved clinical status at Day 15: similar between arms (OR 1.01;	 Treatment-related differences in AEs for BAR vs. DEX were mainly related to
Primary Endpoint	95% Cl, 0.80–1.27)	laboratory abnormalities, not clinical
 MV-free survival by Day 29 	• For low-flow oxygen at baseline: OR 0.91; 95% Cl, 0.70–1.17	events. The clinical relevance of these
Key Secondary Endpoints	• For HFNC oxygen or NIV at baseline: OR 1.64; 95% CI, 0.92–2.90	differences in laboratory abnormalities is unclear.
Clinical status at Day 15 as measured by 0S	Median time to recovery: 6 days in BAR arm vs. 5 days in DEX arm (rete ratio 1.04: 05% CL 0.01, 1.10)	
Time to recovery	(rate ratio 1.04; 95% Cl, 0.91–1.19)	Interpretation
Key Safety Endpoints	Safety Outcomes	In hospitalized patients requiring
Occurrence of treatment-related AEs	 Occurrence of treatment-related AEs: 4% in BAR arm vs. 10% in DEX arm (risk difference 6.0%; 95% Cl, 2.8%–9.3%; P = 0.0004) 	conventional oxygen, HFNC oxygen, or NIV, the use of BAR or DEX resulted in
Occurrence of SAEs	 Occurrence of SAEs: 28% in BAR arm vs. 36% in DEX arm (risk difference 7.7%; 95% Cl, 1.8%–13.4%; P = 0.012) 	similar MV-free survival by Day 29.
	Most SAEs and treatment-related AEs were laboratory abnormalities.	

Methods	Results	Limitations and Interpretation
STOP-COVID: Double-Blind, Placebo-Controlled, Randomized Trial of Tofacitinib in Hospitalized Patients With COVID-19 Pneumonia in Brazil ⁶		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
• Laboratory-confirmed SARS-CoV-2 infection	 Mean age 56 years; 35% women 	Small sample size
COVID-19 pneumonia on CXR or CT	 Median 10 days symptom onset to randomization 	RDV not available during trial
 Hospitalized for <72 hours 	At baseline:	Interpretation
 Key Exclusion Criteria Receiving NIV, MV, or ECMO at baseline 	 75% supplemental oxygen 13% HFNC oxygen 	 Tofacitinib, when compared with placebo, led to a lower risk of mortality
History of or current thrombosis	• Use of glucocorticoids: 79% at baseline, 89% during hospitalization	or respiratory failure among hospitalized adults with COVID-19 pneumonia, most
 Immunosuppression or active cancer treatment 	 Primary Outcome Mortality or respiratory failure through Day 28: 18% in tofacitinib arm 	of whom received glucocorticoids.
Interventions	vs. 29% in placebo arm (risk ratio 0.63; 95% Cl, $0.41-0.97$; $P = 0.04$)	
 Tofacitinib 10 mg P0 twice daily for up to 14 days or until discharge (n = 144) Placebo (n = 145) 	 Secondary Outcome Mortality through Day 28: 2.8% in tofacitinib arm vs. 5.5% in placebo arm (HR 0.49; 95% Cl, 0.15–1.63) 	
Primary Endpoint		
Mortality or respiratory failure through Day 28		
Key Secondary Endpoint		
Mortality through Day 28		

Key: AE = adverse event; ALC = absolute lymphocyte count; ALT = alanine transaminase; ANC = absolute neutrophil count; AST = aspartate transaminase; BAR = baricitinib; CCP = COVID-19 convalescent plasma; CRP = C-reactive protein; CT = computed tomography; CXR = chest X-ray; DEX = dexamethasone; ECMO = extracorporeal membrane oxygenation; eGFR = estimated glomerular filtration rate; HFNC = high-flow nasal cannula; IVIG = intravenous immunoglobulin; LDH = lactate dehydrogenase; MV = mechanical ventilation; NIV = noninvasive ventilation; OS = ordinal scale; the Panel = the COVID-19 Treatment Guidelines Panel; PCR = polymerase chain reaction; PO = orally; RCT = randomized controlled trial; RDV = remdesivir; SAE = serious adverse event; SOC = standard of care; SpO₂ = oxygen saturation; TB = tuberculosis; ULN = upper limit of normal

References

- 1. RECOVERY Collaborative Group. Baricitinib in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial and updated meta-analysis. *Lancet*. 2022;400(10349):359-368. Available at: https://pubmed.ncbi.nlm.nih.gov/35908569.
- 2. Marconi VC, Ramanan AV, de Bono S, et al. Efficacy and safety of baricitinib for the treatment of hospitalised adults with COVID-19 (COV-BARRIER): a randomised, double-blind, parallel-group, placebo-controlled Phase 3 trial. *Lancet Respir Med.* 2021;9(12):1407-1418. Available at:

https://www.ncbi.nlm.nih.gov/pubmed/34480861.

- 3. Ely EW, Ramanan AV, Kartman CE, et al. Efficacy and safety of baricitinib plus standard of care for the treatment of critically ill hospitalised adults with COVID-19 on invasive mechanical ventilation or extracorporeal membrane oxygenation: an exploratory, randomised, placebo-controlled trial. *Lancet Respir Med.* 2022;10(4):327-336. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35123660.
- 4. Kalil AC, Patterson TF, Mehta AK, et al. Baricitinib plus remdesivir for hospitalized adults with COVID-19. *N Engl J Med.* 2021;384(9):795-807. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33306283.
- 5. Wolfe CR, Tomashek KM, Patterson TF, et al. Baricitinib versus dexamethasone for adults hospitalised with COVID-19 (ACTT-4): a randomised, double-blind, double placebo-controlled trial. *Lancet Respir Med.* 2022;10(9):888-899. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35617986.
- 6. Guimaraes PO, Quirk D, Furtado RH, et al. Tofacitinib in patients hospitalized with COVID-19 pneumonia. *N Engl J Med*. 2021;385(5):406-415. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133856</u>.

Abatacept

Last Updated: October 10, 2023

Cytotoxic T-lymphocyte-associated antigen 4 (CTLA-4) is a protein receptor that is expressed by activated T cells. By mediating inhibitory signals, this receptor can decrease T cell proliferation and cytokine production.^{1,2} Abatacept (CTLA-4-Ig) is a soluble fusion protein that contains CTLA-4 linked to human immunoglobulin, and it is used to block T cell activation. Because excessive T cell stimulation and proliferation is thought to propagate the pathogenesis of COVID-19,³ modulating this response may be a potential option for the treatment of COVID-19.⁴

Abatacept is approved by the Food and Drug Administration (FDA) for the treatment of inflammatory arthritis and for the prophylaxis of acute graft-versus-host disease.⁵ It is currently not approved for the treatment of COVID-19. Abatacept has been evaluated in clinical trials for the treatment of hospitalized patients with moderate to severe COVID-19.

Recommendation

See <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> for the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations on the use of abatacept in hospitalized patients who require conventional oxygen, high-flow nasal cannula (HFNC) oxygen, or noninvasive ventilation (NIV).

Rationale

The ACTIV-1 immune modulator trial was a double-blind, multi-arm, placebo-controlled, randomized trial in moderately to severely ill adults hospitalized with COVID-19.⁶ The trial separately evaluated treatment with abatacept, cenicriviroc, and infliximab versus placebo. All arms received standard care, and the separate analyses included data from a shared placebo arm. One substudy compared the use of a single dose of intravenous abatacept 10 mg/kg to placebo. The primary endpoint was time to recovery by Day 28. Key secondary endpoints included clinical status at Day 14 and mortality through Days 28 and 60.

The study concluded that use of abatacept in patients with COVID-19 did not have a significant effect on the time to recovery. A reduction in 28-day mortality, a secondary endpoint, was found. Patients who required mechanical ventilation or extracorporeal membrane oxygenation (ECMO) did not benefit from the use of abatacept.

Clinical Data

In the ACTIV-1 trial, the modified intention-to-treat analysis for the abatacept substudy included 509 patients in the abatacept arm and 510 patients in the placebo arm. At baseline, 53% of the patients required conventional oxygen supplementation, and 33% required HFNC oxygen or NIV. As part of their standard care before or during the study, 93% of the patients received remdesivir, and 91% received corticosteroids.

Results

• The use of abatacept did not reduce the median time to recovery, which was the primary endpoint. The median time to recovery was 9 days in both the infliximab and placebo arms (recovery rate ratio 1.12; 95% CI, 0.98–1.28; P = 0.09), and there was no differential effect across subgroups

based on disease severity (interaction P = 0.66).

- Mortality by Day 28 was lower among patients who received abatacept (56 of 509 patients [11.0%]) than among those who received placebo (77 of 510 patients [15.1%]; OR 0.62; 95% CI, 0.41–0.94).
- Subgroup analyses showed reduced mortality only among patients in the abatacept arm who required HFNC oxygen or NIV (OR 0.48; 95% CI, 0.28–0.84).
- Among patients who required mechanical ventilation or ECMO, there was no difference in mortality by Day 28 (OR 1.63; 95% CI, 0.66–4.05).
- There were no differences in secondary infections or in the number or severity of serious adverse events between the abatacept and placebo arms.

Limitations

- Each of the 3 active agents was compared to a shared placebo group without adjustment for multiple comparisons.
- Mortality was a secondary endpoint. Although the treatment difference found for mortality by Day 28 was nominally significant, no adjustment was made for having considered multiple outcomes (primary outcome and mortality).
- The study was not powered to analyze differences within disease severity subgroups.

Adverse Effects and Monitoring

Most of the data on the adverse effects of abatacept come from the chronic use of the agent for the treatment of autoimmune diseases and graft-versus-host disease. When abatacept is used for the prevention of acute graft-versus-host disease, the most commonly reported adverse effects include fever, anemia, hypertension, cytomegalovirus infection (or reactivation), pneumonia, epistaxis, CD4 lymphopenia, and acute kidney injury.⁵ Concomitant use with other immunomodulatory agents may increase the risk of serious infections. Due to its immunosuppressive effects, all patients who are receiving abatacept should also be monitored for new infections. In the ACTIV-1 trial, data on the safety of short-term use of abatacept in patients with COVID-19 did not reveal significant safety concerns.

Considerations in Pregnant and Lactating People

See <u>Pregnancy</u>, <u>Lactation</u>, <u>and COVID-19 Therapeutics</u> for the Panel's guidance regarding the use of abatacept during pregnancy and lactation.

Considerations in Children

The intravenous formulation of abatacept is approved by the FDA for the treatment of juvenile idiopathic arthritis and acute graft-versus-host disease in children aged ≥ 2 years. It is not approved for the treatment of COVID-19 in children, and there are no published reports on the efficacy of using abatacept in this population. No patients aged <18 years were included in the ACTIV-1 trial.

References

- 1. Alegre ML, Frauwirth KA, Thompson CB. T-cell regulation by CD28 and CTLA-4. *Nat Rev Immunol*. 2001;1(3):220-228. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/11905831/</u>.
- Oyewole-Said D, Konduri V, Vazquez-Perez J, et al. Beyond T-cells: functional characterization of CTLA-4 expression in immune and non-immune cell types. *Front Immunol*. 2020;11:608024. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/33384695/</u>.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

- 3. Chen Z, Wherry EJ. T cell responses in patients with COVID-19. *Nat Rev Immunol*. 2020;20:529-536. Available at: https://www.nature.com/articles/s41577-020-0402-6#citeas.
- Julià A, Bonafonte-Pardàs I, Gómez A, et al. Targeting of the CD80/86 proinflammatory axis as a therapeutic strategy to prevent severe COVID-19. *Sci Rep.* 2021;11(1):11462. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/34075090/</u>.
- 5. Abatacept (Orencia) [package insert]. Food and Drug Administration. 2021. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2021/125118s240lbl.pdf.
- 6. O'Halloran JA, Ko ER, Anstrom KJ, et al. Abatacept, cenicriviroc, or infliximab for treatment of adults hospitalized with COVID-19 pneumonia: a randomized clinical trial. *JAMA*. 2023;330(4):328-339. Available at: https://pubmed.ncbi.nlm.nih.gov/37428480.

Infliximab

Last Updated: October 10, 2023

Infliximab is a tumor necrosis factor–alpha (TNF-alpha) inhibitor that has been evaluated for the treatment of hospitalized patients with moderate to severe COVID-19. TNF-alpha is a pleiotropic proinflammatory cytokine mainly generated by activated macrophages, lymphocytes, and natural killer cells that plays a significant role in immune-mediated inflammatory diseases. Early in the COVID-19 pandemic, increased levels of interleukin (IL)-6 and TNF-alpha were identified as independent predictors of disease severity and death.¹ Furthermore, several cohort studies and registries noted that people with immune-mediated inflammatory diseases who were receiving TNF-alpha inhibitors were at lower risk for COVID-19–related hospitalizations and severe disease than people with immune-mediated inflammatory diseases who were receiving non–TNF-alpha biologic products.^{2,3} It has been hypothesized that modulating levels of TNF-alpha or its effects may reduce the duration or severity of COVID-19.

Recommendation

See <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> for the COVID-19 Treatment Guidelines Panel's (the Panel) recommendations on the use of infliximab in hospitalized patients who require conventional oxygen, high-flow nasal cannula (HFNC) oxygen, or noninvasive ventilation (NIV).

Rationale

The ACTIV-1 immune modulator trial was a double-blind, multi-arm, placebo-controlled, randomized trial in moderately to severely ill adults hospitalized with COVID-19.⁴ The trial separately evaluated treatment with abatacept, cenicriviroc, and infliximab versus placebo. All arms received standard care, and the separate analyses included data from a shared placebo arm. One substudy compared the use of a single dose of intravenous infliximab 5 mg/kg to placebo. The primary endpoint was time to recovery by Day 28. Key secondary endpoints included clinical status at Day 14 and mortality through Days 28 and 60.

The study concluded that use of infliximab in patients with COVID-19 did not have a significant effect on the time to recovery. A reduction in 28-day mortality, a secondary endpoint, was found. Patients who required mechanical ventilation or extracorporeal membrane oxygenation (ECMO) did not benefit from the use of infliximab.

Clinical Data

In the ACTIV-1 trial, the modified intention-to-treat analysis for the infliximab substudy included 517 patients in the infliximab arm and 516 patients in the placebo arm. At baseline, 52% of the patients required conventional oxygen supplementation, and 33% required HFNC oxygen or NIV. As part of their standard care before or during the study, 93% of the patients received remdesivir, and 92% received corticosteroids.

Results

• The use of infliximab did not reduce the median time to recovery, which was the primary endpoint. The median time to recovery was 8 days in the infliximab arm versus 9 days in the

placebo arm (recovery rate ratio 1.12; 95% CI, 0.99–1.28; P = 0.08), and there was no differential effect across subgroups based on disease severity (interaction P = 0.36).

- Mortality by Day 28 was lower among patients who received infliximab (52 of 517 patients [10.1%]) than among those who received placebo (75 of 516 patients [14%]; OR 0.59; 95% CI, 0.39–0.90).
- Subgroup analyses showed reduced mortality only among patients in the infliximab arm who required HFNC oxygen or NIV (OR 0.52; 95% CI, 0.29–0.91).
- Among patients who required mechanical ventilation or ECMO, there was no difference in mortality by Day 28 (OR 1.11; 95% CI, 0.45–2.72).
- There were no differences in secondary infections or in the number or severity of serious adverse events between the infliximab and placebo arms.

Limitations

- Each of the 3 active agents was compared to a shared placebo group without adjustment for multiple comparisons.
- Mortality was a secondary endpoint. Although the treatment difference found for mortality by Day 28 was nominally significant, no adjustment was made for having considered multiple outcomes (primary outcome and mortality).
- The study was not powered to analyze differences within disease severity subgroups.

Adverse Effects and Monitoring

Most of the data on adverse effects of infliximab come from the chronic use of the agent for the treatment of autoimmune diseases. Adverse effects include serious infections (including invasive fungal infections), infusion-related reactions and hypersensitivity, cytopenias, hepatotoxicity, and, rarely, cardiovascular and cerebrovascular events. Because of infliximab's immunosuppressive effects, all patients who receive it should be monitored for new infections. In the ACTIV-1 trial, data on the safety of short-term use of infliximab in patients with COVID-19 did not reveal significant safety concerns.

Considerations in Pregnant and Lactating People

See <u>Pregnancy</u>, <u>Lactation</u>, <u>and COVID-19 Therapeutics</u> for the Panel's guidance regarding the use of infliximab during pregnancy and lactation.

Considerations in Children

Infliximab is approved for the treatment of inflammatory bowel disease in children and is often used to treat juvenile idiopathic arthritis. The Food and Drug Administration has not approved the use of infliximab for the treatment of COVID-19 in children, and there are no published reports on the efficacy of infliximab in this population. No patients aged <18 years were included in the ACTIV-1 trial.

See <u>Therapeutic Management of Hospitalized Children With MIS-C, Plus a Discussion on MIS-A</u> for the Panel's recommendations regarding the use of infliximab in pediatric patients with multisystem inflammatory syndrome in children (MIS-C).

References

 Del Valle DM, Kim-Schulze S, Huang HH, et al. An inflammatory cytokine signature predicts COVID-19 severity and survival. *Nat Med.* 2020;26(10):1636-1643. Available at: <u>https://pubmed.ncbi.nlm.nih.</u> gov/32839624.

COVID-19 Treatment Guidelines

- Curtis JR, Zhou X, Rubin DT, et al. Characteristics, comorbidities, and outcomes of SARS-CoV-2 infection in patients with autoimmune conditions treated with systemic therapies: a population-based study. *J Rheumatol*. 2022;49(3):320-329. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/34782447</u>.
- 3. Izadi Z, Brenner EJ, Mahil SK, et al. Association between tumor necrosis factor inhibitors and the risk of hospitalization or death among patients with immune-mediated inflammatory disease and COVID-19. *JAMA Netw Open*. 2021;4(10):e2129639. Available at: https://pubmed.ncbi.nlm.nih.gov/34661663.
- 4. O'Halloran JA, Ko ER, Anstrom KJ, et al. Abatacept, cenicriviroc, or infliximab for treatment of adults hospitalized with COVID-19 pneumonia: a randomized clinical trial. *JAMA*. 2023;330(4):328-339. Available at: https://pubmed.ncbi.nlm.nih.gov/37428480.

Interleukin-1 Inhibitors

Last Updated: January 26, 2023

Endogenous interleukin (IL)-1 is elevated in patients with COVID-19.¹⁻³ In addition, SARS-CoV-2 infection causes epithelial damage that leads to the release of IL-1 beta, which recruits inflammatory cells and induces the release of IL-1 beta in monocytes. This in turn leads to the release of more IL-1 to recruit and activate additional innate immune cells. Drugs that block the IL-1 receptor (e.g., anakinra) or drugs that block IL-1 signaling (e.g., canakinumab) can potentially interrupt this autoinflammatory loop. These drugs are being investigated as potential treatments for COVID-19.

Anakinra is a recombinant human IL-1 receptor antagonist. It is approved by the Food and Drug Administration (FDA) to treat rheumatoid arthritis and cryopyrin-associated periodic syndromes, specifically neonatal-onset multisystem inflammatory disease.⁴ It is used off-label to treat severe chimeric antigen receptor T cell-mediated cytokine release syndrome and macrophage activation syndrome (MAS)/secondary hemophagocytic lymphohistiocytosis. On November 8, 2022, the FDA issued an Emergency Use Authorization (EUA) for anakinra. The EUA allows the use of anakinra to treat COVID-19 in certain hospitalized adults with pneumonia. These patients must have laboratory-confirmed SARS-CoV-2 infection, require supplemental oxygen (either low- or high-flow oxygen), be at risk of progressing to severe respiratory failure, and be likely to have elevated plasma levels of soluble urokinase plasminogen activator receptor (suPAR), a marker of inflammation.⁵

Canakinumab is a human monoclonal antibody that targets the beta subunit of IL-1 and is approved by the FDA for the treatment of systemic juvenile idiopathic arthritis and Still's disease.

Recommendations

- There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of anakinra for the treatment of COVID-19.
- The Panel **recommends against** the use of **canakinumab** for the treatment of COVID-19, except in a clinical trial (**BIIa**).

Rationale

In the SAVE-MORE trial, 594 hospitalized patients who had moderate or severe COVID-19 pneumonia and plasma suPAR levels \geq 6 ng/mL were randomized to receive either anakinra or placebo. The study found that patients who received anakinra had a lower risk of clinical progression of COVID-19 than those who received placebo.⁶ REMAP-CAP, an open-label, adaptive platform trial that evaluated the use of several immunomodulators in patients with COVID-19 who required organ support, found no clinical benefit of anakinra in these patients. In addition, among patients who received anakinra, no reduction in mortality was observed during a 180-day follow up.⁷ CORIMUNO-ANA-1 was a randomized controlled trial that compared the use of anakinra to usual care in 116 hospitalized patients who were hypoxemic but did not require high-flow oxygen or ventilation. This trial was stopped early due to futility.⁸

The SAVE-MORE study population was restricted to participants with high levels of suPAR (\geq 6 ng/mL), based on the hypothesis that this group is most likely to benefit from IL-1 inhibition. However, the laboratory assay that is used to assess suPAR levels is not currently available in many countries, including the United States. Using data from the SAVE-MORE and SAVE trials (both a priori, open-label, single-arm prospective studies), the FDA developed a scoring system that uses common clinical and laboratory factors to identify patients who are likely to have suPAR levels \geq 6 ng/mL.

The Panel has concluded that there is insufficient evidence to recommend either for or against the use of anakinra for the treatment of COVID-19 in hospitalized patients. Based on the available evidence, the Panel notes the following:

- Data from randomized trials has not consistently demonstrated a benefit of using anakinra to treat COVID-19.
- The suPAR assays that were used to identify patients for participation in the SAVE-MORE trial are not available in the United States.
- The scoring system that the FDA developed to identify patients who might have a high suPAR levels requires further validation.

Finally, CAN-COVID, a randomized controlled trial that evaluated the use of canakinumab in hospitalized patients with COVID-19 who were hypoxemic but did not require ventilatory support, reported that the use of canakinumab did not improve the likelihood of survival without mechanical ventilation.⁹ Therefore, the Panel **recommends against** the use of **canakinumab** for the treatment of COVID-19, except in a clinical trial (**BIIa**).

Clinical Data

SAVE-MORE

SAVE-MORE was a randomized controlled trial in 594 hospitalized patients with moderate or severe COVID-19 pneumonia and plasma suPAR levels \geq 6 ng/mL. Patients who required noninvasive ventilation or mechanical ventilation were excluded from the study. Patients were randomized 2:1 to receive anakinra 100 mg subcutaneously once daily for 10 days or placebo. The primary endpoint was clinical status at Day 28 on the 11-point World Health Organization Clinical Progression Scale (WHO-CPS).⁶ Additional analyses assessed outcomes at 60 and 90 days.¹⁰

Results

- Patients who were randomized to receive anakinra had a lower odds of a worse WHO-CPS score by Day 28 (OR 0.36; 95% CI, 0.26–0.50; *P* < 0.0001).
- The secondary endpoints also favored anakinra, including the absolute decrease in WHO-CPS scores from baseline at Days 14 and 28, the absolute decrease in Sequential Organ Failure Assessment (SOFA) scores from baseline at Day 7, the median time to hospital discharge, and the median duration of intensive care unit (ICU) stays.
- A smaller proportion of patients in the anakinra arm experienced secondary infections, including ventilator-associated pneumonias, than in the placebo arm (8.4% vs. 15.9%; P = 0.01).
- Twenty-eight-day mortality was lower among patients who received anakinra than those who received placebo (3.2% vs. 6.9%; HR 0.45; 95% CI, 0.21–0.98; P = 0.045).
- Additional analyses performed at 60 and 90 days showed a sustained survival benefit for anakinra.

Limitations

The laboratory assay that is used to assess suPAR levels is not currently available in many countries, including the United States. The FDA worked with the SAVE-MORE investigators to develop a scoring system that predicts whether a patient has suPAR levels ≥ 6 ng/mL using baseline data from patients who were randomized during the trial and a subset of patients who were screened but not randomized. The FDA's surrogate for suPAR levels ≥ 6 ng/mL is called SCORE 2, and it includes the following characteristics:

• Age \geq 75 years

COVID-19 Treatment Guidelines

- Severe pneumonia, as determined by WHO criteria
- Current or past smoker
- SOFA score ≥ 3
- Neutrophil to lymphocyte ratio ≥ 7
- Hemoglobin $\leq 10.5 \text{ g/dL}$
- Medical history of ischemic stroke
- Blood urea \geq 50 mg/dL and/or medical history of renal disease

Patients who met \geq 3 of these criteria were considered positive for SCORE 2 and likely to have a suPAR level \geq 6 ng/mL. SCORE 2 had a positive predictive value of 0.95, a sensitivity of 0.41, and specificity of 0.96 when retrospectively applied to the SAVE-MORE trial, and it had similar characteristics when applied to the SAVE trial, an open-label, single-arm prospective study that served as an external validation dataset. In the SAVE-MORE trial, a greater proportion of patients who were positive for SCORE 2 developed severe respiratory failure by Day 14 compared with those who met \leq 2 of the SCORE 2 criteria (41.4% vs. 8.0%).^{4,11}

REMAP-CAP

The REMAP-CAP trial is an open-label, adaptive platform trial in which eligible participants are randomized to several domains, including the Immune Modulation Therapy domain, which consists of 2 IL-6 inhibitors, anakinra, interferon beta-1a, and a control group. Participants are eligible for enrollment if they are within 24 hours of receiving respiratory or cardiovascular organ support in the ICU and they have suspected or microbiologically confirmed COVID-19. This population had more advanced disease than the population enrolled in the SAVE-MORE trial.

Anakinra 300 mg was given intravenously (IV) as a loading dose, followed by anakinra 100 mg IV every 6 hours for 14 days until patients were either free from mechanical ventilation for >24 hours or discharged from the ICU. The primary outcome was measured using an ordinal scale that included a composite of in-hospital mortality and duration of respiratory and cardiovascular organ support at 21 days; all deaths up to 90 days were assigned the worst outcome. The trial used a Bayesian design that allowed the authors to compare nonconcurrently randomized interventions across time periods.⁷ Additional analyses assessed outcomes at 180 days.³

Results

- Of the 2,274 participants who were randomized to 1 of the arms in the Immune Modulation Therapy domain, 365 individuals were assigned to receive anakinra and included in the analysis, 406 were assigned to the usual care (control) arm, 943 were assigned to receive tocilizumab, and 483 were assigned to receive sarilumab.
- Of those assigned to receive anakinra, 37% were receiving mechanical ventilation at study entry compared with 32% of patients in the other arms. The other patients received oxygen through a high-flow nasal cannula or noninvasive ventilation, with a few exceptions.
- The median number of organ support-free days was similar for patients who received anakinra and those who received usual care (0 days [IQR -1 to 15 days] vs. 0 days [IQR -1 to 15 days]). The aOR for organ support-free days was 0.99 for anakinra (95% CrI, 0.74–1.35), with a 47% posterior probability of superiority to control. Sixty percent of those who were assigned to receive anakinra survived compared with 63% of those who were assigned to the control arm, with a 44% posterior probability that anakinra was superior to usual care.
- Additional analyses performed at 180 days showed no reduction in mortality among patients who

COVID-19 Treatment Guidelines

received anakinra.3

• The risk of experiencing serious adverse events was similar between the arms.

Limitations

Patients were not randomized contemporaneously to receive anakinra or usual care; the treatment effect was estimated from an overarching model that mostly included patients who were randomized to receive an IL-6 inhibitor (tocilizumab or sarilumab) or usual care, and patients who were randomized to receive an IL-6 inhibitor or anakinra. Thus, the estimate of the treatment effect is not fully protected by randomization. This study also had an open-label design.

CORIMUNO-ANA-1

The CORIMUNO-ANA-1 trial randomized 116 hospitalized patients with COVID-19 pneumonia 1:1 to receive either usual care plus anakinra (200 mg IV twice a day on Days 1–3, 100 mg IV twice on Day 4, and 100 mg IV once on Day 5) or usual care alone. Patients were eligible for enrollment if they had laboratory-confirmed SARS-CoV-2 infection with COVID-19 pneumonia and they required >3 L/ min of supplemental oxygen. Patients who required high-flow oxygen, ventilation, or ICU admission were excluded. The 2 coprimary outcomes were the proportion of patients who had died or who needed noninvasive ventilation or mechanical ventilation by Day 4 (score of >5 on the WHO-CPS) and the proportion who survived without the need for noninvasive ventilation or mechanical ventilation (including high-flow oxygen) by Day $14.^8$

Results

- There was no difference between the anakinra plus usual care arm and the usual care alone arm in the coprimary outcomes: by Day 4, 36% of patients in the anakinra arm had died or required high-flow oxygen or ventilation compared with 38% in the usual care arm (90% CrI, -17.1 to 12.0, posterior probability of benefit 61%). By Day 14, 47% of patients in the anakinra arm had died or required noninvasive ventilation or mechanical ventilation compared with 51% in the usual care arm (median HR 0.97; 90% CrI, 0.62–1.52; posterior probability of benefit 55%).
- Fifty-two percent of patients received corticosteroids at study entry.
- Serious adverse events occurred in 46% of patients in the anakinra arm compared with 38% in the usual care arm; 11 of 59 patients (18.6%) in the anakinra arm experienced bacterial or fungal infections compared with 4 of 55 patients (7.3%) who received usual care.

Limitations

The limitations of this study include the small sample size, narrow eligibility criteria, and the fact that many patients did not receive current standard-of-care therapy (e.g., corticosteroids, remdesivir).

CAN-COVID

CAN-COVID was a double-blind, placebo-controlled randomized trial of 454 hospitalized patients with COVID-19 who were hypoxemic but not mechanically ventilated and had elevated C-reactive protein (\geq 20 mg/L) or ferritin (\geq 600 micrograms/L) levels. Patients were randomized 1:1 to receive a single dose of IV canakinumab (450 mg for a body weight of 40 kg to <60 kg, 600 mg for 60–80 kg, and 750 mg for >80 kg) or placebo. The primary outcome was survival without the need for mechanical ventilation from Days 3 through 29.⁹

Results

• There was no statistical difference between the canakinumab arm and placebo arm in the proportion of patients who survived without mechanical ventilation (88.8% vs. 85.7%; P = 0.29).

- The number of COVID-19-related deaths at 4 weeks was similar for the 2 arms (11 of 223 patients [4.9%] in the canakinumab arm vs. 16 of 222 patients [7.2%] in the placebo arm; OR 0.67; 95% CI, 0.30–1.50).
- Forty-one percent of patients in the canakinumab arm and 32% in the placebo arm received dexamethasone.
- Serious adverse events occurred in 16% of patients who received canakinumab and in 21% of patients who received placebo.

Limitations

The use of corticosteroids was unbalanced in this study, with more patients receiving dexamethasone at baseline in the canakinumab arm than in the placebo arm. More patients received dexamethasone after the trial was underway in the placebo arm than in the canakinumab arm (22.5% vs. 14.5%), and more patients received tocilizumab in the placebo arm than in the canakinumab arm (8.8% vs. 2.2%).

Other small cohort studies, case-control studies, and case series have reported mixed findings with regard to improvement in outcomes among patients who received anakinra for the treatment of COVID-19.¹²⁻¹⁵ The clinical implication of these findings is uncertain due to small sample sizes and unmeasured confounding factors. Therefore, these studies did not substantially influence the Panel's current recommendations for using IL-1 inhibitors.

Adverse Effects

Headache, nausea, vomiting, and liver enzyme elevations can occur with both anakinra and canakinumab.

Anakinra was not associated with any significant safety concerns when used in clinical trials for the treatment of sepsis.¹⁶⁻¹⁸ Increased rates of infection were reported with prolonged anakinra use in combination with tumor necrosis factor-alpha blockade, but not with short-term use.¹⁹

Considerations in Pregnancy

The data on using IL-1 inhibitors to treat COVID-19 in pregnant patients are currently limited. The American College of Rheumatology recommends against the use of anakinra during pregnancy.²⁰ Unintentional first-trimester exposure to anakinra is unlikely to be harmful, given the minimal transfer of monoclonal antibodies across the placenta early in pregnancy.²¹

Considerations in Children

Anakinra has been used in the treatment of severely ill children with rheumatologic conditions, including systemic juvenile idiopathic arthritis and MAS. The data on the use of anakinra in pediatric patients with acute respiratory distress syndrome or sepsis are limited. Anakinra is rarely used to treat pediatric patients with acute COVID-19, and it has been used in approximately 10% of cases of multisystem inflammatory syndrome in children (MIS-C).^{22,23} Anakinra is often included in institutional protocols for the treatment of MIS-C in the United States, and it is an option for second-line therapy for refractory MIS-C in national consensus guidelines, including the COVID-19 Treatment Guidelines.²⁴⁻ ²⁶ For more information, see <u>Therapeutic Management of Hospitalized Children With MIS-C</u>, Plus a <u>Discussion on MIS-A</u>.

Data on using canakinumab in pediatric patients are limited to use in patients with periodic fever syndromes and systemic juvenile idiopathic arthritis. There are no data on its use in pediatric patients with acute COVID-19 or MIS-C.

References

- 1. Shakoory B, Carcillo JA, Chatham WW, et al. Interleukin-1 receptor blockade is associated with reduced mortality in sepsis patients with features of macrophage activation syndrome: reanalysis of a prior Phase III trial. *Crit Care Med.* 2016;44(2):275-281. Available at: https://www.ncbi.nlm.nih.gov/pubmed/26584195.
- 2. Monteagudo LA, Boothby A, Gertner E. Continuous intravenous anakinra infusion to calm the cytokine storm in macrophage activation syndrome. *ACR Open Rheumatol*. 2020;2(5):276-282. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32267081.
- 3. Writing Committee for the REMAP-CAP Investigators. Long-term (180-day) outcomes in critically ill patients with COVID-19 in the REMAP-CAP randomized clinical trial. *JAMA*. 2023;329(1):39-51. Available at: https://pubmed.ncbi.nlm.nih.gov/36525245/.
- 4. Anakinra (Kineret) [package insert]. Food and Drug Administration. 2022. Available at: <u>https://www.fda.gov/media/163546/download</u>.
- 5. Food and Drug Administration. Anakinra (Kineret) EUA letter of authorization. 2022. Available at: <u>https://www.fda.gov/media/163081/download</u>.
- 6. Kyriazopoulou E, Poulakou G, Milionis H, et al. Early treatment of COVID-19 with anakinra guided by soluble urokinase plasminogen receptor plasma levels: a double-blind, randomized controlled phase 3 trial. *Nat Med.* 2021;27(10):1752-1760. Available at: https://pubmed.ncbi.nlm.nih.gov/34480127/.
- 7. REMAP-CAP Investigators. Effectiveness of tocilizumab, sarilumab, and anakinra for critically ill patients with COVID-19: the REMAP-CAP COVID-19 immune modulation therapy domain randomized clinical trial. *medRxiv*. 2021;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2021.06.18.21259133v2</u>.
- 8. CORIMUNO-19 Collaborative Group. Effect of anakinra versus usual care in adults in hospital with COVID-19 and mild-to-moderate pneumonia (CORIMUNO-ANA-1): a randomised controlled trial. *Lancet Respir Med.* 2021;9(3):295-304. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33493450</u>.
- Caricchio R, Abbate A, Gordeev I, et al. Effect of canakinumab vs placebo on survival without invasive mechanical ventilation in patients hospitalized with severe COVID-19: a randomized clinical trial. *JAMA*. 2021;326(3):230-239. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34283183</u>.
- Akinosoglou K, Kotsaki A, Gounaridi IM, et al. Efficacy and safety of early soluble urokinase plasminogen receptor plasma-guided anakinra treatment of COVID-19 pneumonia: a subgroup analysis of the SAVE-MORE randomised trial. *EClinicalMedicine*. 2023;56:101785. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36590789/</u>.
- 11. Kyriazopoulou E, Panagopoulos P, Metallidis S, et al. An open label trial of anakinra to prevent respiratory failure in COVID-19. *Elife*. 2021;10:e66125. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/33682678/</u>.
- 12. Aouba A, Baldolli A, Geffray L, et al. Targeting the inflammatory cascade with anakinra in moderate to severe COVID-19 pneumonia: case series. *Ann Rheum Dis*. 2020;79(10):1381-1382. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32376597.
- 13. Cavalli G, De Luca G, Campochiaro C, et al. Interleukin-1 blockade with high-dose anakinra in patients with COVID-19, acute respiratory distress syndrome, and hyperinflammation: a retrospective cohort study. *Lancet Rheumatol*. 2020;2(6):e325-e331. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32501454.
- 14. Huet T, Beaussier H, Voisin O, et al. Anakinra for severe forms of COVID-19: a cohort study. *Lancet Rheumatol*. 2020;2(7):e393-e400. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32835245</u>.
- 15. Kooistra EJ, Waalders NJB, Grondman I, et al. Anakinra treatment in critically ill COVID-19 patients: a prospective cohort study. *Crit Care*. 2020;24(1):688. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33302991.
- 16. Fisher CJ, Jr., Dhainaut JF, Opal SM, et al. Recombinant human interleukin 1 receptor antagonist in the treatment of patients with sepsis syndrome. Results from a randomized, double-blind, placebo-controlled trial. Phase III rhIL-1ra Sepsis Syndrome Study Group. *JAMA*. 1994;271(23):1836-1843. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/8196140</u>.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

- 17. Fisher CJ, Jr., Slotman GJ, Opal SM, et al. Initial evaluation of human recombinant interleukin-1 receptor antagonist in the treatment of sepsis syndrome: a randomized, open-label, placebo-controlled multicenter trial. *Crit Care Med.* 1994;22(1):12-21. Available at: https://www.ncbi.nlm.nih.gov/pubmed/8124953.
- Opal SM, Fisher CJ, Jr., Dhainaut JF, et al. Confirmatory interleukin-1 receptor antagonist trial in severe sepsis: a Phase III, randomized, double-blind, placebo-controlled, multicenter trial. The Interleukin-1 Receptor Antagonist Sepsis Investigator Group. *Crit Care Med.* 1997;25(7):1115-1124. Available at: https://www.ncbi.nlm.nih.gov/pubmed/9233735.
- Winthrop KL, Mariette X, Silva JT, et al. ESCMID Study Group for Infections in Compromised Hosts (ESGICH) Consensus Document on the safety of targeted and biological therapies: an infectious diseases perspective (soluble immune effector molecules [II]: agents targeting interleukins, immunoglobulins and complement factors). *Clin Microbiol Infect*. 2018;24 Suppl 2:S21-S40. Available at: https://www.ncbi.nlm.nih.gov/pubmed/29447987.
- 20. Sammaritano LR, Bermas BL, Chakravarty EE, et al. 2020 American College of Rheumatology guideline for the management of reproductive health in rheumatic and musculoskeletal diseases. *Arthritis Rheumatol*. 2020;72(4):529-556. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32090480</u>.
- Flint J, Panchal S, Hurrell A, et al. BSR and BHPR guideline on prescribing drugs in pregnancy and breastfeeding-part II: analgesics and other drugs used in rheumatology practice. *Rheumatology (Oxford)*. 2016;55(9):1698-1702. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26750125</u>.
- 22. Götzinger F, Santiago-García B, Noguera-Julián A, et al. COVID-19 in children and adolescents in Europe: a multinational, multicentre cohort study. *Lancet Child Adolesc Health*. 2020;4(9):653-661. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32593339.
- Derespina KR, Kaushik S, Plichta A, et al. Clinical manifestations and outcomes of critically ill children and adolescents with coronavirus disease 2019 in New York City. *J Pediatr*. 2020;226:55-63.e52. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32681989</u>.
- 24. Dove ML, Jaggi P, Kelleman M, et al. Multisystem inflammatory syndrome in children: survey of protocols for early hospital evaluation and management. *J Pediatr*. 2021;229:33-40. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33075369.
- 25. Henderson LA, Canna SW, Friedman KG, et al. American College of Rheumatology clinical guidance for multisystem inflammatory syndrome in children associated with SARS-CoV-2 and hyperinflammation in pediatric COVID-19: version 3. *Arthritis Rheumatol*. 2022;74(4):e1-e20. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35118829</u>.
- 26. Harwood R, Allin B, Jones CE, et al. A national consensus management pathway for paediatric inflammatory multisystem syndrome temporally associated with COVID-19 (PIMS-TS): results of a national Delphi process. *Lancet Child Adolesc Health*. 2021;5(2):133-141. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32956615</u>.

Vilobelimab

Last Updated: December 20, 2023

Vilobelimab is an anti-C5a monoclonal antibody. High concentrations of C5a have been reported in patients with severe COVID-19.¹ C5a activates innate immune system responses, including inflammation and the release of histamines, and can increase damage to local tissues.² A study in mice demonstrated that an anti-C5a monoclonal antibody reduced immune system activation and inhibited lung injury.³ Vilobelimab targets C5a, which is a product of complement activation, and preserves membrane attack complex function.⁴ Vilobelimab is not approved by the Food and Drug Administration (FDA) for any indication.

On April 4, 2023, the FDA issued an Emergency Use Authorization for the use of vilobelimab for the treatment of COVID-19 in hospitalized adults when it is administered within 48 hours of mechanical ventilation or extracorporeal membrane oxygenation.⁵

Recommendation

• There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of vilobelimab for the treatment of COVID-19.

Rationale

Results from the PANAMO trial were used to support the FDA Emergency Use Authorization.⁵ However, the prespecified analysis that stratified by study site showed that 28-day mortality among patients with COVID-19 who received vilobelimab was not significantly different from 28-day mortality among those who received placebo. The initially proposed primary study analysis did not stratify by study site. In the second phase of the study, the primary analysis was changed to stratify by site based on a recommendation from the FDA. The analysis that did not stratify by site demonstrated that all-cause mortality through Day 28 was significantly lower in the vilobelimab arm than in the placebo arm. Concomitant use of corticosteroids (97%) and antithrombotic agents (98%) was high in this study population. Prior or concomitant use of additional immunomodulators, such as tocilizumab (17% in the vilobelimab arm, 16% in the placebo arm) and baricitinib (3% in each arm), was low. The Panel determined that the results from the PANAMO trial were insufficient to recommend either for or against the use of vilobelimab for the treatment of COVID-19.

Monitoring, Adverse Effects, and Drug-Drug Interactions

Reports of adverse effects of vilobelimab are limited to a Phase 3 trial that included critically ill adult patients with COVID-19 who received intravenous vilobelimab 800 mg for up to 6 doses.^{5,6} Common adverse reactions (i.e., those with an incidence \geq 3% and that were observed at least 1% more frequently in the vilobelimab arm than in the placebo arm through Day 60) were pneumonia, sepsis, delirium, pulmonary embolism, hypertension, pneumothorax, deep vein thrombosis, herpes simplex, enterococcal infection, bronchopulmonary aspergillosis, increased hepatic enzymes, urinary tract infection, hypoxemia, thrombocytopenia, pneumomediastinum, respiratory tract infection, supraventricular tachycardia, constipation, and rash.

Vilobelimab is not expected to be associated with any pharmacokinetic drug-drug interactions.

Considerations in Pregnant People

There are no data on the use of vilobelimab during pregnancy, as pregnant individuals were excluded from the PANAMO trial.

Considerations in Children

There are no data on the use of vilobelimab in children. Vilobelimab is not authorized by the FDA for the treatment of COVID-19 in pediatric patients.

Clinical Data

The small (n = 30) Phase 2 portion of the Phase 2/3 PANAMO trial was too underpowered to draw any conclusions about study outcomes, including physiologic improvement at 5 days and mortality.⁷

The Phase 3 portion of the trial was a double-blind, randomized trial performed at 46 hospitals in Western Europe (i.e., Netherlands, France, Germany, Belgium), Brazil, Mexico, Russia, Peru, and South Africa from October 1, 2020, to October 4, 2021.⁶ The trial compared the use of vilobelimab plus standard of care with placebo plus standard of care in patients aged ≥ 18 years who had laboratory-confirmed SARS-CoV-2 infection, were receiving mechanical ventilation (and were within 48 hours of intubation), and had a ratio of arterial partial pressure of oxygen to fraction of inspired oxygen of 60 to 200 mm Hg at study entry. Vilobelimab 800 mg was administered intravenously on Days 1, 2, 4, 8, 15, and 22, if the patient remained hospitalized, for a maximum of 6 doses.

The primary outcome was all-cause mortality at 28 days. Secondary outcomes included all-cause mortality at 60 days, the proportion of patients who improved on a World Health Organization 8-point ordinal scale, the proportion of patients who developed acute kidney failure by Day 28, and the proportion of patients free from renal replacement therapy at Day 28.

Results

- The trial enrolled 369 patients; 368 patients were included in the analysis that did not stratify by study site (177 in the vilobelimab arm, 191 in the placebo arm).
- In the prespecified analysis that stratified by study site (n = 307), 28-day mortality was not significantly different between the vilobelimab and placebo arms (HR 0.73; 95% CI, 0.50–1.06; *P* = 0.094). The analysis for 28-day mortality that stratified by study site excluded the 61 patients (16.6%) from sites that had no deaths or had only 1 treatment group.
- In the analysis that did not stratify by study site (n = 368), 28-day mortality was lower in the vilobelimab arm than in the placebo arm (54 of 177 patients [31%] vs. 77 of 191 patients [44%]), and the difference between arms was statistically significant (HR 0.67; 95% CI, 0.48–0.96; P = 0.027).
- Prespecified subgroup analyses identified a significant reduction in 28-day mortality in the vilobelimab arm for subgroups of patients with severe acute respiratory distress syndrome (HR 0.55; 95% CI, 0.30–0.98; P = 0.044), patients with an estimated glomerular filtration rate of <60 mL/min (HR 0.55; 95% CI, 0.31–0.96; P = 0.036), and patients receiving mechanical ventilation and additional organ support (category 7 on the World Health Organization 8-point ordinal scale; HR 0.62; 95% CI, 0.40–0.95; P = 0.028).
- In a prespecified analysis of the Western Europe subgroup (i.e., Netherlands, France, Germany, Belgium), the vilobelimab arm had significantly lower 28-day mortality than the placebo arm (HR 0.51; 95% CI, 0.30–0.87; P = 0.014).

- For the secondary outcomes:
 - The analysis that stratified by study site showed no significant difference between the arms for all-cause mortality at 60 days (HR 0.74; 95% CI, 0.52-1.04; P = 0.082).
 - The vilobelimab arm had significantly fewer patients who required renal replacement therapy at Day 28 than the placebo arm (age-adjusted HR 0.54; 95% CI, 0.30–0.98; P = 0.042).

Limitations

- The results for the study's site-stratified, prespecified analysis were not significant.
- The analysis for 28-day mortality that stratified by study site excluded the 61 patients (16.6%) from sites that had no deaths or had only 1 treatment group.
- Very few patients received a second immunomodulator (tocilizumab or baricitinib), which makes the study results difficult to apply to current practice.
- Compared to other studies that have evaluated the use of immunomodulators for the treatment of COVID-19, Phase 3 of the PANAMO trial had a relatively small sample size.

References

- 1. Cugno M, Meroni PL, Gualtierotti R, et al. Complement activation in patients with COVID-19: a novel therapeutic target. *J Allergy Clin Immunol*. 2020;146(1):215-217. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32417135</u>.
- Manthey HD, Woodruff TM, Taylor SM, Monk PN. Complement component 5a (C5a). *Int J Biochem Cell Biol*. 2009;41(11):2114-2117. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/19464229</u>.
- 3. Carvelli J, Demaria O, Vély F, et al. Association of COVID-19 inflammation with activation of the C5a-C5aR1 axis. *Nature*. 2020;588(7836):146-150. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/32726800</u>.
- Chouaki Benmansour N, Carvelli J, Vivier E. Complement cascade in severe forms of COVID-19: recent advances in therapy. *Eur J Immunol*. 2021;51(7):1652-1659. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33738806</u>.
- 5. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Gohibic. 2023. Available at: <u>https://www.fda.gov/media/166824/download</u>.
- 6. Vlaar APJ, Witzenrath M, van Paassen P, et al. Anti-C5a antibody (vilobelimab) therapy for critically ill, invasively mechanically ventilated patients with COVID-19 (PANAMO): a multicentre, double-blind, randomised, placebo-controlled, Phase 3 trial. *Lancet Respir Med.* 2022;10(12):1137-1146. Available at: https://pubmed.ncbi.nlm.nih.gov/36087611.
- Vlaar APJ, de Bruin S, Busch M, et al. Anti-C5a antibody IFX-1 (vilobelimab) treatment versus best supportive care for patients with severe COVID-19 (PANAMO): an exploratory, open-label, Phase 2 randomised controlled trial. *Lancet Rheumatol*. 2020;2(12):e764-e773. Available at: <u>https://pubmed.ncbi.nlm.</u> <u>nih.gov/33015643</u>.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Table 5e. Characteristics of Immunomodulators

Last Updated: October 10, 2023

- The information in this table is derived from data on the use of these drugs for FDA-approved indications or from clinical trials that evaluated their use in patients with COVID-19.
- For dose modifications in patients with organ failure or those who require extracorporeal devices, please refer to product labels or EUAs, when available.
- There are currently not enough data to determine whether certain medications can be safely coadministered with therapies for the treatment of COVID-19. When using concomitant medications with similar toxicity profiles, consider performing additional safety monitoring.
- The potential additive, antagonistic, or synergistic effects and the safety of using certain combination therapies for the treatment of COVID-19 are unknown. Clinicians are encouraged to report AEs to the FDA *MedWatch* program.
- For drug-drug interaction information, please refer to product labels and visit the Liverpool COVID-19 Drug Interactions website.
- For the Panel's recommendations on using the drugs listed in this table, please refer to the drug-specific sections of the Guidelines; <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>; <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>;</u> <u>Therapeutic Management of Hospitalized Children With COVID-19</u>; and <u>Pregnancy, Lactation, and COVID-19</u> Therapeutics.

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Corticosteroid (S Recommended by	ystemic) the Panel for the treatment of CO	VID-19 in certain hospitalized pa	atients.		
Dexamethasone	 Dose for Adults With COVID-19 DEX 6 mg IV or PO once daily for up to 10 days or until hospital discharge, whichever comes first¹ 	 Hyperglycemia Secondary infections Reactivation of latent infections (e.g., HBV, HSV, strongyloidiasis, TB) Cases of disseminated strongyloidiasis have been reported in patients with COVID-19 during treatment with corticosteroids and tocilizumab. Psychiatric disturbances Avascular necrosis 	 Blood glucose BP Signs and symptoms of new infection 	 Moderate CYP3A4 inducer CYP3A4 substrate 	 If DEX is not available, an alternative corticosteroid (e.g., prednisone, methylprednisolone, hydrocortisone) can be used. The approximate total daily dose equivalencies for these glucocorticoids to DEX 6 mg (IV or PO) are: Prednisone 40 mg Methylprednisolone 32 mg Hydrocortisone 160 mg

COVID-19 Treatment Guidelines

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Corticosteroid (Systemic), continued	I		1	
		 Adrenal insufficiency Increased BP Peripheral edema Myopathy (particularly if used with NMBAs) 			
Janus Kinase In		ID-10 in certain beenitalized o	ationte		
Recommended b	y the Panel for the treatment of COVI FDA-Approved Doses for COVID-19 in Adults Aged ≥18 Years, per eGFR ² ≥60 mL/min/1.73 m ² • BAR 4 mg PO once daily for 14 days or until hospital discharge, whichever comes first 30 to <60 mL/min/1.73 m ² • BAR 2 mg PO once daily for 14 days or until hospital discharge, whichever comes first 15 to <30 mL/min/1.73 m ² • BAR 1 mg PO once daily for 14 days or until hospital discharge, whichever comes first <pre></pre>	 <i>D-19 in certain hospitalized pa</i> Lymphoma and other malignancies Thrombosis Gl perforation Treatment-related changes in lymphocytes, neutrophils, Hgb, liver enzymes HSV reactivation Herpes zoster Secondary infections Serious cardiac-related events (e.g., MI, stroke) 	 CBC with differential Renal function Liver enzymes Signs and symptoms of new infections 	Dose modification recommended when administering concurrently with a strong OAT3 inhibitor.	 See the FDA label² and EUA³ for dosing guidance for patients with: ALC <200 cells/µL ANC <500 cells/µL If increases in ALT or AST are observed and DILI is suspected, interrupt BAR treatment until the diagnosis of DILI is excluded. BAR tablets can be taken PO or crushed, dispersed in water, and given via gastrostomy tube.² Availability BAR is approved by the FDA for the treatment of COVID-19 in adults aged ≥18 years.² BAR is available through an FDA EUA for children aged 2–17 years who require supplemental oxygen, NIV, MV, or ECMO.³

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Janus Kinase In	hibitors, continued	I	I	,	
Baricitinib	days or until hospital discharge, whichever comes first				
	30 to <60 mL/min/1.73 m ²				
	BAR 1 mg PO once daily for 14 days or until hospital discharge, whichever comes first				
	<30 mL/min/1.73 m ²				
	Not recommended				
Tofacitinib	 Dose for COVID-19 in Clinical Trials Tofacitinib 10 mg P0 twice daily for up to 14 days or until hospital discharge, whichever comes first⁴ 	 Thrombotic events (e.g., PE, DVT, arterial thrombosis) Anemia Increased risk of infection Gl perforation Diarrhea Headache Herpes zoster Lipid elevations Liver enzyme elevations Lymphoma and other malignancies Serious cardiac-related events (e.g., MI, stroke) 	 CBC with differential Liver enzymes Signs and symptoms of new infections 	 Requires dose modification when administered with strong CYP3A4 inhibitors or when used with a moderate CYP3A4 inhibitor that is coadministered with a strong CYP2C19 inhibitor Coadministration with strong CYP3A4 inducers is not recommended. 	 Avoid use in patients with ALC <500 cells/mm³, ANC <1,000 cells/mm³, or Hgb <9 grams/dL. May require dose modification in patients with moderate or severe renal impairment or moderate hepatic impairment
Interleukin-6 Inl	hibitors (Anti-Interleukin-6 Recept	•			
	y the Panel for the treatment of COV	-	ients.		
Sarilumab	 Dose for COVID-19 in Clinical Trials 1 dose of sarilumab 400 mg IV^{5,6} 	 Neutropenia Thrombocytopenia Gl perforation HSRs Increased liver enzymes 	 HSRs Infusion-related reactions Neutrophils PLT 	Elevated IL-6 may downregulate CYP enzymes; thus, use of sarilumab may lead to increased metabolism of drugs that are CYP	• Sarilumab is not recommended in patients with ALT or AST >1.5 times the upper limit of the reference range, ANC <2,000 cells/mm ³ , or PLT <150,000

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Interleukin-6 Inf	nibitors (Anti-Interleukin-6 Re	ceptor Monoclonal Antibodie	es), continued		
Sarilumab		HBV reactivationInfusion-related reactions	Liver enzymes	• The effects of sarilumab on CYP enzymes may persist for weeks after the drug is stopped.	 Treatment with sarilumab may mask signs of acute inflammation or infection by suppressing fever and CRP levels.
					Availability
					 IV formulation of sarilumab is not approved by the FDA, but in clinical trials, a single SUBQ dose (using the prefilled syringes, not the prefilled pen) of sarilumab 400 mg was reconstituted in 100 cc 0.9% NaCl and given as an IV infusion over 1 hour.^{6,8}
					 IV infusion of sarilumab should occur within 4 hours of its preparation; it can be stored at room temperature until administered.
Tocilizumab	 FDA-Approved Dose for COVID-19 in Hospitalized Adults Tocilizumab 8 mg/kg (maximum 800 mg) by IV infusion over 1 hour FDA EUA Doses for COVID-19 in Hospitalized Children 	 HSRs Infusion-related reactions GI perforation Hepatotoxicity Treatment-related changes on laboratory tests for neutrophils, PLT, lipids, and liver enzymes HBV reactivation 	 HSRs Infusion-related reactions Neutrophils PLT Liver enzymes 	 Inhibition of IL-6 may lead to increased metabolism of coadministered drugs that are CYP450 substrates. The effects of tocilizumab on CYP enzymes may persist for weeks after the drug is stopped. 	 Tocilizumab is not recommended in patients with ALT or AST >10 times the upper limit of the reference range, ANC <1,000 cells/mm³, or PLT <50,000 cells/mm^{3.9} SUBQ formulation of tocilizumab is not intended for IV administration.
	Body Weight ≥30 kg • Tocilizumab 8 mg/kg by IV	 Secondary infections 			Availability
	infusion over 1 hour	 Cases of disseminated strongyloidiasis have been reported in patients 			 IV tocilizumab is approved by the FDA for the treatment of COVID-19 in hospitalized

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Interleukin-6 Inf	nibitors (Anti-Interleukin-6 Recep	otor Monoclonal Antibodies), continued		
Tocilizumab	 Body Weight <30 kg Tocilizumab 12 mg/kg by IV infusion over 1 hour For All Doses If clinical signs or symptoms worsen or do not improve following the first infusion, 1 additional dose may be administered at least 8 hours after the first dose. 	with COVID-19 during treatment with tocilizumab and corticosteroids.			 adults aged 18 years.¹⁰ Tocilizumab is available through an FDA EUA for the treatment of COVID-19 in certain hospitalized children aged 2–17 years.⁹
	phocyte-Associated Antigen 4 Ag y the Panel for the treatment of CON	•	l patients.		
Abatacept	 Dose for COVID-19 in Clinical Trials 1 dose of abatacept 10 mg/ kg (maximum 1,000 mg) by IV infusion over 30 minutes¹¹ 	 HSRs, including anaphylaxis HBV reactivation Secondary infections Patients with COPD may develop more frequent respiratory AEs. Headache Upper respiratory infection, nasopharyngitis Nausea Anemia HTN Decrease in CD4 count Hypermagnesemia Acute kidney injury¹² 	 HSRs Infusion-related reactions CBC with differential Electrolytes Renal function 	Drug-drug interactions are unlikely between abatacept and medications that are CYP substrates, inhibitors, or inducers.	 IV formulation of abatacept includes maltose, which may give falsely elevated blood glucose readings with certain blood glucose monitors (e.g., GDH-PQQ- based monitoring systems) on the day of infusion. In ACTIV-1, 1 case of anaphylaxis and 2 infusion- related reactions were reported among abatacept recipients.¹¹ Availability The IV formulation of abatacept is commercially available.

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
	Factor–Alpha Inhibitor				
•	y the Panel for the treatment of CO				
Infliximab	Dose for COVID-19 in Clinical Trials	 Infusion-related reactions 	• HSRs	Inhibition of cytokine	Availability
		, , , , , , , , , , , , , , , , , , , ,	 Infusion-related 	activity may lead to increased metabolism	Infliximab is available as
	 1 dose of infliximab 5 mg/kg by IV infusion over 2 hours¹¹ 	The following AEs are	reactions	of coadministered	an originator biologic or a biosimilar.
		associated with chronic use of infliximab:	 CBC with differential 	drugs that are CYP450 substrates.	biobirmut.
		 Hepatotoxicity 	• PLT	Cuboulatoon	
		Cytopenia (e.g.,	 Liver enzymes 		
		leukopenia, neutropenia, thrombocytopenia, pancytopenia)	 If infliximab is administered 		
		HBV reactivation	to patients with heart failure, they should be closely monitored.		
		 Secondary infections (e.g., invasive fungal infections, reactivation of latent TB) 			
		Heart failure			
		 CVA, MI, hypotension, hypertension, arrhythmias 			
		Transient vision loss			
		 Demyelinating disease 			
		 Lupus-like syndrome 			
		Headache			
		 Abdominal pain¹³ 			
	Ional Antibody EUA for the treatment of COVID-19 er for or against its use.	when initiated within 48 hours of	receiving MV or ECMO.	There is insufficient evidenc	e for the Panel to
Vilobelimab	FDA EUA Dose for COVID-19 in	Secondary infections	• CBC	None	Availability
	Hospitalized Adults Receiving MV or ECMO	Delirium	 Liver enzymes 		 Vilobelimab is not
		• PE	 Infusion-related 		approved by the FDA,
	Vilobelimab 800 mg by IV infusion after dilution, for a	• HTN	reactions		but it is commercially available for use in
	maximum of 6 doses; start	Pneumothorax	 Signs and 		hospitalized adults with

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Anti-C5a Monocl	onal Antibody, continued				
Vilobelimab	treatment within 48 hours of intubation (Day 1) followed by administration on Days 2, 4, 8, 15, and 22 if patient is still hospitalized (even if discharged from ICU)	 DVT Liver enzyme elevations Hypoxemia Thrombocytopenia Pneumomediastinum Supraventricular tachycardia Constipation Rash 	symptoms of new infections		COVID-19, as authorized in the EUA.
<u>Canakinumab: No</u> Anakinra	t recommended by the Panel for the FDA EUA Dose for COVID-19 in Hospitalized Patients Aged ≥18 Years • Anakinra 100 mg SUBQ once daily for 10 days Dose for CrCl <30 mL/min • Anakinra 100 mg SUBQ every other day for 5 total doses over 10 days ¹⁴	 Neutropenia (particularly when used concomitantly with other agents that can cause neutropenia) HSRs, including anaphylaxis and angioedema Secondary infections Injection site reactions 	 CBC with differential; assess neutrophils before starting treatment and during therapy. BMP Liver enzymes Renal function 	• Use with TNF- blocking agents is not recommended due to increased risk of infection.	 Contraindicated in patients with known hypersensitivity to proteins derived from <i>Escherichia coli</i>, anakinra, or any component of the product¹⁴ Patients with <1,500 neutrophils/mm³ were excluded from participation in the SAVE-MORE study.¹⁵ Availability
		 Liver enzyme elevations Hyperkalemia Hypernatremia Rash 			 SUBQ anakinra is available through an FDA EUA.¹⁴

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Interleukin-1 Inl	nibitors, continued			·	<u>.</u>
Canakinumab	 FDA-Approved Dose for Systemic JIA Canakinumab 4 mg/kg (maximum 300 mg) SUBQ every 4 weeks¹⁶ 	 HSRs Neutropenia Nasopharyngitis Diarrhea Respiratory tract infections Bronchitis Gastroenteritis Pharyngitis Musculoskeletal pain Vertigo Abdominal pain Injection site reactions Liver enzyme elevations 	 HSRs CBC with differential Liver enzymes 	 Binding of canakinumab to IL-1 may increase formation of CYP enzymes and alter metabolism of drugs that are CYP substrates. Use with TNF- blocking agents is not recommended due to potential increased risk of infection. 	 Availability IV canakinumab is not an approved formulation in the United States.¹⁶
Corticosteroids Not recommende	(Inhaled) In d by the Panel for the treatment	of COVID-19. Currently under i	investigation in clinical	trials.	
Budesonide (Inhaled)	 Dose for COVID-19 in Clinical Trials Budesonide 800 µg oral inhalation twice daily until symptom resolution or up to 14 days^{17,18} 	 Secondary infections Oral thrush Systemic AEs (less common) 	 Signs of AEs involving the oral mucosa or throat, including thrush Signs of systemic corticosteroid effects (e.g., adrenal suppression) 	 CYP3A4 substrate Do not use with strong CYP3A4 inhibitors. 	No comments

Drug Name	Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Corticosteroids (Inhaled), continued				
Ciclesonide (Inhaled)	 Dose for COVID-19 in Clinical Trials Ciclesonide 160 µg as 2 MDI inhalations twice daily for 30 days¹⁹ 	 Secondary infections Oral thrush Systemic AEs (less common) 	 Signs of AEs involving the oral mucosa or throat, including thrush Signs of systemic corticosteroid effects (e.g., adrenal suppression) 	 CYP3A4 substrate Strong CYP3A4 inhibitors are expected to have less effect on ciclesonide exposure than on budesonide exposure. 	No comments

Key: AE = adverse event; ALC = absolute lymphocyte count; ALT = alanine transaminase; ANC = absolute neutrophil count; AST = aspartate aminotransferase; BAR = baricitinib; BMP = basic metabolic panel; BP = blood pressure; CBC = complete blood count; CD4 = CD4 T lymphocyte; COPD = chronic obstructive pulmonary disease; CrCI = creatinine clearance; CRP = C-reactive protein; CVA = cerebral vascular accident; CYP = cytochrome P450; DEX = dexamethasone; DILI = drug-induced liver injury; DVT = deep vein thrombosis; ECMO = extracorporeal membrane oxygenation; eGFR = estimated glomerular filtration rate; EUA = Emergency Use Authorization; FDA = Food and Drug Administration; GDH-PQQ = glucose dehydrogenase pyrroloquinoline quinone; GI = gastrointestinal; HBV = hepatitis B virus; Hgb = hemoglobin; HSR = hypersensitivity reaction; HSV = herpes simplex virus; HTN = hypertension; ICU = intensive care unit; IL = interleukin; IV = intravenous; JIA = juvenile idiopathic arthritis; MDI = metered dose inhaler; MI = myocardial infarction; MV = mechanical ventilation; NaCI = sodium chloride; NIV = noninvasive ventilation; NMBA = neuromuscular blocking agent; OAT = organic anion transporter; the Panel = the COVID-19 Treatment Guidelines Panel; PE = pulmonary embolism; PLT = platelet count; PO = oral; SUBQ = subcutaneous; TB = tuberculosis; TNF = tumor necrosis factor

References

- 1. Randomised Evaluation of COVID-19 Therapy (RECOVERY). Low-cost dexamethasone reduces death by up to one third in hospitalised patients with severe respiratory complications of COVID-19. 2020. Available at: <u>https://www.recoverytrial.net/news/low-cost-dexamethasone-reduces-death-by-up-to-one-third-in-hospitalised-patients-with-severe-respiratory-complications-of-covid-19. Accessed June 7, 2023.</u>
- 2. Baricitinib (Olumiant) [package insert]. Food and Drug Administration. 2022. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2022/207924s006lbl.pdf.
- 3. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization (EUA) of baricitinib. 2022. Available at: https://www.fda.gov/media/143823/download.
- 4. Guimarães PO, Quirk D, Furtado RH, et al. Tofacitinib in patients hospitalized with COVID-19 pneumonia. *N Engl J Med*. 2021;385(5):406-415. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133856</u>.
- REMAP-CAP Investigators. Effectiveness of tocilizumab, sarilumab, and anakinra for critically ill patients with COVID-19: the REMAP-CAP COVID-19 immune modulation therapy domain randomized clinical trial. *medRxiv*. 2021;Preprint. Available at: <u>https://www.medrxiv.org/</u> <u>content/10.1101/2021.06.18.21259133v2</u>.
- 6. Sivapalasingam S, Lederer DJ, Bhore R, et al. Efficacy and safety of sarilumab in hospitalized patients with COVID-19: a randomized clinical trial.

COVID-19 Treatment Guidelines

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Clin Infect Dis. 2022;75(1):e380-e388. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35219277.

- 7. Sarilumab (Kevzara) [package insert]. Food and Drug Administration. 2018. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2018/761037s001lbl.pdf.
- 8. REMAP-CAP Investigators. Interleukin-6 receptor antagonists in critically ill patients with COVID-19. *N Engl J Med.* 2021;384(16):1491-1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631065</u>.
- 9. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Actemra (tocilizumab). 2021. Available at: https://www.fda.gov/media/150321/download.
- 10. Tocilizumab (Actemra) [package insert]. Food and Drug Administration. 2022. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2022/125276s138lbl.pdf.
- 11. O'Halloran JA, Ko ER, Anstrom KJ, et al. Abatacept, cenicriviroc, or infliximab for treatment of adults hospitalized with COVID-19 pneumonia: a randomized clinical trial. *JAMA*. 2023;330(4):328-339. Available at: https://pubmed.ncbi.nlm.nih.gov/37428480.
- 12. Abatacept (Orencia) [package insert]. Food and Drug Administration. 2021. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2021/125118s240lbl.pdf.
- 13. Infliximab (Remicade) [package insert]. Food and Drug Administration. 2021. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2021/103772s5401lbl.pdf.
- 14. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Kineret. 2022. Available at: https://www.fda.gov/media/163075/download.
- 15. Kyriazopoulou E, Poulakou G, Milionis H, et al. Early treatment of COVID-19 with anakinra guided by soluble urokinase plasminogen receptor plasma levels: a double-blind, randomized controlled Phase 3 trial. *Nat Med.* 2021;27(10):1752-1760. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34480127.
- 16. Canakinumab (Ilaris) [package insert]. Food and Drug Administration. 2020. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2020/125319s100lbl.pdf.
- 17. Ramakrishnan S, Nicolau DV Jr, Langford B, et al. Inhaled budesonide in the treatment of early COVID-19 (STOIC): a Phase 2, open-label, randomised controlled trial. *Lancet Respir Med*. 2021;9(7):763-772. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33844996</u>.
- 18. Yu LM, Bafadhel M, Dorward J, et al. Inhaled budesonide for COVID-19 in people at high risk of complications in the community in the UK (PRINCIPLE): a randomised, controlled, open-label, adaptive platform trial. *Lancet*. 2021;398(10303):843-855. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34388395.
- 19. Clemency BM, Varughese R, Gonzalez-Rojas Y, et al. Efficacy of inhaled ciclesonide for outpatient treatment of adolescents and adults with symptomatic COVID-19: a randomized clinical trial. *JAMA Intern Med.* 2022;182(1):42-49. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34807241.

Antithrombotic Therapy in Patients With COVID-19

Last Updated: October 10, 2023

Summary Recommendations

Chronic Anticoagulant and Antiplatelet Therapy

- The COVID-19 Treatment Guidelines Panel (the Panel) recommends that patients with COVID-19 who are receiving anticoagulant or antiplatelet therapies for underlying conditions continue these medications unless significant bleeding develops or other contraindications are present (AIII).
- Before prescribing ritonavir-boosted nirmatrelvir (Paxlovid) to patients who are receiving anticoagulant or antiplatelet therapy, clinicians should carefully review the patient's concomitant medications to evaluate potential drug-drug interactions. It may be necessary to modify the dosage of the antithrombotic agent, switch to another antithrombotic agent, or prescribe an alternative COVID-19 therapy. See <u>Drug-Drug Interactions Between Ritonavir-Boosted</u> <u>Nirmatrelvir (Paxlovid) and Concomitant Medications</u> for more information.

Screening and Evaluation for Venous Thromboembolism

- There is insufficient evidence for the Panel to recommend either for or against routine screening for venous thromboembolism (VTE) in patients with COVID-19 who do not have signs or symptoms of VTE, regardless of the status of their coagulation markers.
- For hospitalized patients with COVID-19 who experience rapid deterioration of pulmonary, cardiac, or neurological function or sudden, localized loss of peripheral perfusion, the Panel recommends evaluating the patients for thromboembolic disease (AIII).

Anticoagulant Treatment for Thrombosis

- When diagnostic imaging is not possible, the Panel recommends that patients with COVID-19 who are highly suspected to have thromboembolic disease be treated with therapeutic anticoagulation (AIII).
- The Panel recommends that patients with COVID-19 who require extracorporeal membrane oxygenation or continuous renal replacement therapy or who have thrombosis related to catheters or extracorporeal filters be treated with antithrombotic therapy as per the standard institutional protocols for those without COVID-19 (AIII).

Antithrombotic Therapy for Nonhospitalized Patients Without Evidence of Venous Thromboembolism

 In nonhospitalized patients with COVID-19, the Panel recommends against the use of anticoagulant and antiplatelet therapy (i.e., aspirin, P2Y12 inhibitors) for the prevention of VTE or arterial thrombosis, except in a clinical trial (Alla). This recommendation does not apply to patients with other indications for antithrombotic therapy.

Antithrombotic Therapy for Hospitalized, Nonpregnant Adults Without Evidence of Venous Thromboembolism

- The Panel **recommends agains**t using anticoagulant or antiplatelet therapy to prevent arterial thrombosis outside of the usual standard of care for patients without COVID-19 (AIII).
- In hospitalized patients, low-molecular-weight heparin (LMWH) or unfractionated heparin (UFH) is preferred over oral anticoagulants (AIII). Because these types of heparin have shorter half-lives, their effects can be reversed quickly. They can also be administered intravenously or subcutaneously, and they have fewer drug-drug interactions than oral anticoagulants.
- When heparin is used, LMWH is preferred over UFH.

For adults who require low-flow oxygen and do not require intensive care unit (ICU)-level care:

- The Panel recommends the use of a **therapeutic dose of heparin** for patients with D-dimer levels above the upper limit of normal who require low-flow oxygen and who do not have an increased risk of bleeding **(Clla)**.
 - Contraindications for the use of therapeutic anticoagulation in patients with COVID-19 include a platelet count <50 x 10⁹/L, hemoglobin <8 g/dL, the need for dual antiplatelet therapy, bleeding within the past 30 days that required an emergency department visit or hospitalization, a history of a bleeding disorder, or an inherited or active acquired bleeding disorder. This list is based on the exclusion criteria from clinical trials; patients with these conditions have an increased risk of bleeding.

Summary Recommendations, continued
• In patients without VTE who have started treatment with therapeutic doses of heparin, treatment should continue for 14 days or until they are transferred to the ICU or discharged from the hospital, whichever comes first.
• The Panel recommends the use of a prophylactic dose of heparin for patients who do not meet the criteria for receiving therapeutic heparin or are not receiving a therapeutic dose of heparin for other reasons, unless a contraindication exists (AI) .
 There is insufficient evidence for the Panel to recommend either for or against the use of a therapeutic dose of apixaban for VTE prophylaxis or the prevention of COVID-19 progression.
 The Panel recommends against the use of a therapeutic dose of rivaroxaban for VTE prophylaxis or the prevention of COVID-19 progression (Alla).
• There is insufficient avidence for the Danal to recommand either for or against the use of thromholytic agants for the

- There is insufficient evidence for the Panel to recommend either for or against the use of thrombolytic agents for the treatment of COVID-19.
- The Panel **recommends against** the use of antiplatelet therapy to prevent COVID-19 progression or death in noncritically ill patients (**Blla**).

For adults who require ICU-level care, including those receiving high-flow oxygen:

- The Panel recommends using a prophylactic dose of heparin as VTE prophylaxis, unless a contraindication exists (AI).
- For patients who start on a therapeutic dose of heparin in a non-ICU setting due to COVID-19 and then transfer to the ICU, the Panel recommends switching from the therapeutic dose to a **prophylactic dose of heparin**, unless VTE is confirmed **(BIII)**.
- The Panel **recommends against** the use of a therapeutic dose of anticoagulation for VTE prophylaxis, except in a clinical trial **(BI)**.
- There is insufficient evidence for the Panel to recommend either for or against the use of an intermediate dose of anticoagulation for VTE prophylaxis.
- There is insufficient evidence for the Panel to recommend either for or against the use of antiplatelet therapy in critically ill patients with COVID-19.

Antithrombotic Therapy for Patients Discharged From the Hospital

• The Panel **recommends against** routinely continuing VTE prophylaxis in patients with COVID-19 after hospital discharge unless they have another indication for anticoagulation (Alla).

Children With COVID-19 or MIS-C

• For the Panel's recommendations on the use of antithrombotic therapy in children, see <u>Therapeutic Management</u> of <u>Hospitalized Children With COVID-19</u> and <u>Therapeutic Management of Hospitalized Children With MIS-C, Plus a</u> <u>Discussion on MIS-A</u>.

Pregnant and Lactating Patients

- The Panel recommends that pregnant patients who are receiving anticoagulant or antiplatelet therapies for underlying conditions continue these medications after they receive a diagnosis of COVID-19 (AIII).
- The Panel recommends the use of a prophylactic dose of anticoagulation for pregnant patients who are hospitalized for manifestations of COVID-19, unless a contraindication exists (BIII).
- Because pregnant patients were not included in most of the clinical trials that evaluated the use of therapeutic anticoagulation in the setting of COVID-19, there is insufficient evidence for the Panel to recommend either for or against the use of therapeutic anticoagulation in pregnant patients with COVID-19 who do not have evidence of VTE.
- As in nonpregnant patients with COVID-19, VTE prophylaxis after hospital discharge is not routinely recommended for pregnant patients (BIII). Clinicians should consider an individual patient's VTE risk factors when making decisions about continuing VTE prophylaxis after discharge in pregnant or postpartum patients.
- The use of anticoagulation therapy during labor and delivery requires specialized care and planning. The management of anticoagulation therapy in pregnant patients with COVID-19 should be similar to the management used for pregnant patients with other conditions (AIII).
- UFH, LMWH, and warfarin do not accumulate in breast milk and do not induce an anticoagulant effect in the newborn; therefore, they can be used by breastfeeding individuals who require VTE prophylaxis or treatment (AIII).

Summary Recommendations, continued

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Association Between COVID-19 and Thromboembolism

COVID-19 has been associated with inflammation and a prothrombotic state, with increases in levels of fibrin, fibrin degradation products, fibrinogen, and D-dimer.^{1,2} In some studies, elevations in these markers have been associated with worse clinical outcomes.^{3,4}

Studies have reported varying incidences of venous thromboembolism (VTE) in patients with COVID-19. A meta-analysis of studies of hospitalized patients with COVID-19 who received VTE prophylaxis found an overall VTE prevalence of 14.1% (95% CI, 11.6–16.9).⁵ The VTE prevalence was higher in studies that used ultrasound screening (40.3%; 95% CI, 27.0–54.3) than in studies that did not (9.5%; 95% CI, 7.5–11.7). In randomized controlled trials conducted prior to the pandemic, the incidence of VTE in hospitalized patients who received VTE prophylaxis ranged from 0.3% to 1% for symptomatic VTE and from 2.8% to 5.6% for VTE overall.⁶⁻⁸ In randomized trials, the VTE incidence among critically ill patients without COVID-19 who received a prophylactic dose of anticoagulants ranged from 5% to 16%, and a prospective cohort study of critically ill patients with sepsis reported a VTE incidence of 37%.⁹⁻¹²

Guidelines for the use of antithrombotic therapy in patients with COVID-19 have been released by multiple organizations, including the American College of Chest Physicians,¹³ the American Society of Hematology,¹⁴ the Anticoagulation Forum,¹⁵ the International Society on Thrombosis and Haemostasis,¹⁶ the Italian Society on Thrombosis and Haemostasis,¹⁷ the National Institute for Health and Care Excellence (NICE),¹⁸ and the Royal College of Physicians.¹⁹ The American College of Chest Physicians also has guidance on the use of antithrombotic therapy to treat arterial thrombosis in people with COVID-19.²⁰

The guidelines referenced above agree that hospitalized, nonpregnant patients with COVID-19 should receive, at a minimum, a prophylactic dose of anticoagulation to prevent VTE. The <u>NICE guidelines</u> state: "Consider a treatment dose of a low-molecular-weight heparin (LMWH) for young people and adults with COVID-19 who need low-flow oxygen and who do not have an increased bleeding risk." Results from clinical trials have provided further information on the safety and efficacy of different antithrombotic strategies for patients with COVID-19.

Chronic Anticoagulant or Antiplatelet Therapy

The COVID-19 Treatment Guidelines Panel (the Panel) recommends that patients with COVID-19 who are receiving anticoagulant or antiplatelet therapies for underlying conditions continue these medications unless significant bleeding develops or other contraindications are present (**AIII**). Outpatients with COVID-19 who are receiving warfarin and are in isolation and unable to have international normalized ratio monitoring may be candidates for switching to direct oral anticoagulant therapy.²¹ Patients with a mechanical heart valve, ventricular assist device, valvular atrial fibrillation, or antiphospholipid antibody syndrome and patients who are lactating should not discontinue treatment with warfarin (**AIII**).

Before prescribing ritonavir-boosted nirmatrelvir (Paxlovid) to patients who are receiving anticoagulant or antiplatelet therapy, clinicians should carefully review the patient's concomitant medications to evaluate potential drug-drug interactions. It may be necessary to modify the dosage of the antithrombotic agent, switch to another antithrombotic agent, or prescribe an alternative COVID-19 therapy. See Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications for more information.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Screening and Evaluation for Venous Thromboembolism

VTE guidelines for patients without COVID-19 have recommended against performing routine screening ultrasounds in critically ill patients because no study has shown that this strategy reduces the rate of subsequent symptomatic thromboembolic complications.²² Although the incidence of thromboembolic events, especially pulmonary embolism, can be high among hospitalized patients with COVID-19, no published data demonstrate the clinical utility of using lower extremity ultrasounds as routine surveillance for deep vein thrombosis in this population.

There is insufficient evidence for the Panel to recommend either for or against routine screening for VTE in patients with COVID-19 who do not have signs or symptoms of VTE, regardless of the status of their coagulation markers. For hospitalized patients with COVID-19 who experience rapid deterioration of pulmonary, cardiac, or neurological function or sudden, localized loss of peripheral perfusion, the Panel recommends evaluating the patients for thromboembolic disease (AIII).

Managing Antithrombotic Therapy in Patients With COVID-19

When diagnostic imaging is not possible, the Panel recommends that patients with COVID-19 who are highly suspected to have thromboembolic disease be managed with therapeutic anticoagulation (AIII).

The Panel recommends that patients with COVID-19 who require extracorporeal membrane oxygenation (ECMO) or continuous renal replacement therapy or who have thrombosis related to catheters or extracorporeal filters be treated with antithrombotic therapy as per the standard institutional protocols for those without COVID-19 (AIII).

Selection of Anticoagulant or Antiplatelet Drugs

Whenever anticoagulant or antiplatelet therapy is used, potential drug-drug interactions with other concomitant medications must be considered. The University of Liverpool has collated a list of <u>drug-drug interactions</u>. In hospitalized patients, **LMWH** or **unfractionated heparin** (UFH) is preferred over oral anticoagulants (**AIII**). Because these types of heparin have shorter half-lives, their effects can be reversed quickly. They can also be administered intravenously or subcutaneously (SUBQ), and they have fewer drug-drug interactions than oral anticoagulants.

Management of Nonhospitalized Patients

ACTIV-4b was a placebo-controlled, randomized trial that evaluated the efficacy of using aspirin or prophylactic doses (2.5 mg) or therapeutic doses (5 mg) of apixaban in outpatients with COVID-19 aged >40 years.²³ After 657 outpatients were randomized, the trial was stopped in June 2021 due to a low event rate for the composite outcome of thromboembolic events, hospitalization, or death (1 patient each in the placebo, aspirin, and apixaban 2.5 mg arms and 2 patients in the apixaban 5 mg arm). The median time from randomization to receipt of treatment was 3 days, and 22 patients were hospitalized for COVID-19 prior to initiation of the study drugs.

Two trials evaluated the use of LMWH and its impact on hospitalization and mortality in outpatients with COVID-19. The ETHIC trial was a multicenter, open-label randomized controlled trial of unvaccinated outpatients with COVID- $19.^{24}$ Adults with at least 1 risk factor for severe disease were randomized to receive enoxaparin 40 mg SUBQ once daily (if they weighed <100 kg) or enoxaparin 40 mg SUBQ twice daily (if they weighed >100 kg) for 21 days or standard of care. The study was terminated early due to a low event rate and slow accrual of participants. There was no difference between the arms in the number of patients who met the composite endpoint of all-cause mortality or all-cause hospitalization (12 of 105 patients [11%] in the enoxaparin arm vs. 12 of 114 patients [11%] in the standard of care arm). Four of the 12 patients in the enoxaparin arm who were admitted to the *COVID-19 Treatment Guidelines*

hospital required acute medical care or intensive care unit (ICU) admission (3 required mechanical ventilation or ECMO). There were no hospitalizations in the standard of care arm. Bleeding events occurred in 2 patients who received enoxaparin and in 1 patient who received standard of care.

The OVID trial was a multicenter, open-label randomized controlled trial of 472 adults with COVID-19 aged >50 years who were randomized to receive enoxaparin 40 mg SUBQ once daily for 14 days or standard of care.²⁵ The study was terminated after recruiting 50% of the planned number of participants due to a low probability that enoxaparin would be superior to standard of care for the primary outcome. There was no difference between the arms in the number of patients who met the primary composite endpoint of all-cause hospitalization or mortality (8 of 234 patients [3%] in the enoxaparin arm vs. 8 of 238 patients [3%] in the standard of care arm). No major bleeding events occurred during the study.

In nonhospitalized patients with COVID-19, the Panel **recommends against** the use of anticoagulant and antiplatelet therapy (i.e., **aspirin**, **P2Y12 inhibitors**) for the prevention of VTE or arterial thrombosis, except in a clinical trial (**AIIa**). This recommendation does not apply to patients with other indications for antithrombotic therapy.

Management of Hospitalized Patients

Several studies have evaluated the risks and benefits of using prophylactic or therapeutic doses of anticoagulants in patients with COVID-19. Observational studies and clinical trials have examined the effects of anticoagulation on mortality, progression of COVID-19, thrombosis, and bleeding. Some of these studies are outlined below. Observational studies are included here only when evidence from clinical trials is not available.

Prophylactic Dose of Anticoagulation Versus No Anticoagulation-Observational Cohort

An observational study of 4,297 veterans hospitalized with COVID-19 evaluated the use of prophylactic anticoagulation.²⁶ A prophylactic dose of anticoagulation was administered to 3,627 patients with COVID-19 within 24 hours of hospital admission. An inverse probability of treatment weighted analysis showed a cumulative 30-day mortality of 14% among patients who received prophylactic anticoagulation and 19% among patients who were not treated with anticoagulation (HR 0.73; 95% CI, 0.66–0.81). Patients treated with the prophylactic dose did not have a significant difference in the risk of bleeding that required transfusion when compared with patients who were not treated (HR 0.87; 95% CI, 0.71–1.05). Overall, the study demonstrated that patients with COVID-19 may benefit from a prophylactic dose of anticoagulation.

Therapeutic Versus Prophylactic Doses of Heparin in Hospitalized Patients Who Do Not Require Intensive Care Unit-Level Care

Several randomized controlled trials have evaluated the role of therapeutic doses of heparin in reducing the risk of VTE events or death in patients hospitalized for COVID-19.

Four open-label randomized controlled trials (the large ATTACC/ACTIV-4a/REMAP-CAP multiplatform trial and the FREEDOM trial and the smaller RAPID and HEP-COVID trials) compared therapeutic doses of heparin to prophylactic or intermediate doses of the anticoagulant in selected hospitalized patients who did not require intensive care. Clinical data for these trials are summarized in Table 6a. The inclusion and exclusion criteria for these studies varied, but most of the studies included patients who required supplemental oxygen and had no risk of a major bleeding event. In the larger multiplatform trial, therapeutic doses of heparin increased the number of organ support-free days but did not significantly affect mortality or length of hospitalization when compared with prophylactic doses of heparin.²⁷ In the FREEDOM trial, there was no difference between the therapeutic and prophylactic anticoagulation arms in the occurrence of the 30-day primary composite outcome of all-cause mortality,

need for ICU-level care, systemic thromboembolism, or ischemic stroke. In a secondary analysis, 30-day mortality was significantly lower in patients who received therapeutic enoxaparin than in patients who received prophylactic enoxaparin.²⁸ However, only a small proportion of patients received concomitant corticosteroids or remdesivir as standard of care, and the trial was stopped early due to slow recruitment.

The RAPID trial enrolled patients with elevated D-dimer levels and hypoxemia.²⁹ The patients were randomized to receive therapeutic or prophylactic doses of heparin. There was no statistically significant difference between the arms for the primary endpoint, which was a composite of ICU admission, noninvasive ventilation (NIV) or mechanical ventilation, or death by Day 28. However, the therapeutic dose of heparin reduced the risk of all-cause death, a secondary outcome.

The HEP-COVID trial enrolled patients who required supplemental oxygen and had a D-dimer value >4 times the upper limit of normal (ULN) or a sepsis-induced coagulopathy score of \geq 4.³⁰ There were significantly fewer occurrences of the primary endpoint of VTE, arterial thromboembolism, or all-cause death within 32 days of randomization in the therapeutic LMWH arm than in the prophylactic LMWH arm, but there was no difference between arms for the outcome of death within 32 days.

Given the results of the ATTACC/ACTIV-4a/REMAP-CAP, FREEDOM, RAPID, and HEP-COVID trials, for hospitalized, nonpregnant adults with COVID-19 who do not require ICU-level care and have no evidence of VTE:

- The Panel recommends the use of a **therapeutic dose of heparin** for patients with D-dimer levels above the ULN who require low-flow oxygen and who do not have an increased risk of bleeding (**CIIa**).
 - Contraindications for the use of therapeutic anticoagulation in patients with COVID-19 include a platelet count <50 x 10⁹/L, hemoglobin <8 g/dL, the need for dual antiplatelet therapy, bleeding within the past 30 days that required an emergency department visit or hospitalization, a history of a bleeding disorder, or an inherited or active acquired bleeding disorder. This list is based on the exclusion criteria from clinical trials; patients with these conditions have an increased risk of bleeding.
- LMWH is preferred over UFH because of its ease of administration and because LMWH was the predominant form of heparin used in the clinical trials for COVID-19.
- In patients without VTE who have started treatment with therapeutic doses of heparin, treatment should continue for 14 days or until they are transferred to the ICU or discharged from the hospital, whichever comes first.
- Patients with predicted hospitalizations of <72 hours were excluded from the multiplatform ATTACC/ACTIV-4a/REMAP-CAP trial. It is currently unknown whether the benefits of using therapeutic doses of anticoagulation for short hospital stays outweigh the risks.
- The Panel recommends the use of a **prophylactic dose of heparin** for patients who do not meet the criteria for receiving therapeutic heparin or are not receiving a therapeutic dose of heparin for other reasons, unless a contraindication exists (**AI**).
- There is insufficient evidence for the Panel to recommend either for or against the use of thrombolytic agents for the treatment of COVID-19.

Prophylactic Versus Intermediate or Therapeutic Doses of Heparin in Hospitalized Patients Who Require Intensive Care Unit-Level Care

Several randomized controlled trials have evaluated the role of therapeutic doses of heparin in reducing the incidence of VTE events or death in patients in the ICU setting. Clinical data for these trials are summarized in <u>Table 6a</u>.

The INSPIRATION trial compared the use of an intermediate dose of enoxaparin (1 mg/kg SUBQ once daily) to a prophylactic dose of enoxaparin (40 mg/kg SUBQ once daily) in patients with COVID-19 who were in the ICU.³¹ The study reported no difference between the arms in the occurrence of the composite endpoint of adjudicated VTE, arterial thrombosis, ECMO, or all-cause mortality. Major bleeding occurred in 2.5% of patients in the intermediate-dose anticoagulation arm and in 1.4% of patients who received the prophylactic dose. Overall, there was no significant benefit of receiving an intermediate dose of anticoagulation for patients with COVID-19 who were in the ICU.

The ANTICOVID trial was an open-label study of hospitalized patients with COVID-19 who required oxygen therapy.³² Patients were randomized to receive a prophylactic dose of LMWH (n = 114), an intermediate dose of LWMH (n = 110), or a therapeutic dose of LMWH (n = 110). Patients in the study received either enoxaparin or tinzaparin. Patients underwent a computed tomography scan at baseline to ensure they did not have a pulmonary embolism. The study excluded patients weighing <40 kg or >100 kg.

The primary hierarchical outcome for this study was all-cause mortality or time to clinical improvement by Day 28. There was no difference between the arms for this outcome. The study also evaluated net clinical outcome, which was defined as a composite of venous and arterial thrombosis, major bleeding events (as defined by the International Society on Thrombosis and Hemostasis), or all-cause mortality by Day 28. A smaller percentage of patients who received intermediate-dose anticoagulation met the net clinical outcome criteria compared with those who received prophylactic-dose anticoagulation (16.4% vs. 29.8%; absolute difference -13.5%; P = 0.02). There was no statistically significant difference in the occurrence of the net clinical outcome between the therapeutic-dose anticoagulation arm and the prophylactic-dose or intermediate-dose arms. No difference in the occurrence of major bleeding events was seen among the study arms.

Tinzaparin is not available in the United States. This lack of availability, combined with the conflicting results of the INSPIRATION and ANTICOVID trials, has led the Panel to conclude that there is insufficient evidence to recommend either for or against the use of an intermediate dose of anticoagulation for VTE prophylaxis.

The multiplatform ATTACC/ACTIV-4a/REMAP-CAP trial compared the effectiveness of a therapeutic dose of heparin or LMWH with usual care in reducing the number of organ support-free days among critically ill patients with COVID-19.²⁷ All 3 trials were stopped for futility. Heparin doses in the usual care arm varied. The median number of organ support-free days and likelihood of survival to hospital discharge did not differ between the arms. Major bleeding occurred in 4% of patients who received therapeutic anticoagulation and in 2% of patients who received usual care. Therapeutic doses of heparin showed no significant benefit for patients with COVID-19 who were admitted to the ICU.

The COVID-PACT trial was a multicenter trial with a 2 x 2 factorial design. Critically ill patients with COVID-19 were randomized to receive a therapeutic dose or a prophylactic dose of anticoagulation. They were also randomized to receive either clopidogrel or no antiplatelet therapy.³³ The trial was stopped early because the decreasing number of ICU admissions for patients with COVID-19 made recruitment difficult. There was no difference between the arms in the occurrence of the primary endpoint (a composite of VTE or arterial thrombotic events at hospital discharge or Day 28). More moderate to severe bleeding events occurred among patients who were treated with therapeutic anticoagulation than among those who received prophylactic anticoagulation.

For hospitalized, nonpregnant adults with COVID-19 who require ICU-level care and who do not have documented or suspected VTE:

• The Panel recommends using a **prophylactic dose of heparin** as VTE prophylaxis, unless a contraindication exists (**AI**).

- For patients who start on a therapeutic dose of heparin in a non-ICU setting due to COVID-19 and then transfer to the ICU, the Panel recommends switching from the therapeutic dose to a **prophylactic dose of heparin**, unless VTE is confirmed (**BIII**).
- The Panel **recommends against** the use of a therapeutic dose of anticoagulation for VTE prophylaxis, except in a clinical trial (**BI**).
- There is insufficient evidence for the Panel to recommend either for or against the use of an intermediate dose of anticoagulation for VTE prophylaxis.

Apixaban or Rivaroxaban in Hospitalized Patients With COVID-19

The FREEDOM trial randomized patients 1:1:1 to receive a therapeutic dose of apixaban, a therapeutic dose of enoxaparin, or a prophylactic dose of enoxaparin.²⁸ The trial showed no difference in the occurrence of the primary composite endpoint between the therapeutic and prophylactic anticoagulation arms. In a secondary analysis, fewer deaths were reported at 30 days among patients who were treated with a therapeutic dose of apixaban than among those who received prophylactic enoxaparin (5% vs. 7%; HR 0.7; 95% CI, 0.49–0.99). Only a small proportion of patients were treated with dexamethasone or remdesivir as part of usual care; both of these drugs have been shown to have a benefit in this population. This open-label trial was also stopped early due to slow recruitment.

The FREEDOM trial is the only study that evaluated the use of therapeutic apixaban in patients with COVID-19; in contrast, 4 trials have evaluated the use of therapeutic heparin. Additionally, oral anticoagulants have the potential for drug-drug interactions and present unique challenges for managing hemorrhages. Due to these limitations, there is insufficient evidence for the Panel to recommend either for or against the use of a therapeutic dose of apixaban for VTE prophylaxis or the prevention of COVID-19 progression.

The ACTION trial randomized adults who were hospitalized with COVID-19 and elevated D-dimer levels (defined as levels that were above the laboratory ULN) to receive rivaroxaban 20 mg once daily for 30 days (n = 311) or usual care (n = 304).³⁴ A heterogenous population was included; 25% of patients did not require oxygen, 60% were treated with low-flow oxygen, and 15% needed high-flow oxygen, NIV, or mechanical ventilation. No statistical difference was found between the arms for the composite endpoint of time to death, hospitalization duration, or oxygen use duration (hierarchical analysis; win ratio 0.86; 95% CI, 0.59–1.22) or for the individual components of the composite endpoint. The probability of clinically relevant, nonmajor bleeding was greater in the rivaroxaban arm (5% in the rivaroxaban arm vs. 1% in the usual care arm; relative risk 5.23; 95% CI, 1.54–17.77), but for major bleeding events, the difference in probability between the arms was not significant (3% in the rivaroxaban arm vs. 1% in the usual care arm; relative risk 2.45; 95% CI, 0.78–7.73). Given the lack of benefit and the increased risk of bleeding events, the Panel **recommends against** the use of a **therapeutic dose of rivaroxaban** for VTE prophylaxis or the prevention of COVID-19 progression (**AIIa**).

Antiplatelet Therapy Versus Usual Care in Hospitalized Patients

Multiple retrospective cohort studies have suggested that the use of aspirin reduced in-hospital mortality in patients who were treated prior to hospital admission or within 24 hours of admission. These studies have been summarized in meta-analyses.³⁵⁻³⁸ These epidemiologic studies used propensity scoring or adjusted for potential confounders, but indication bias cannot be fully removed from these studies. Thus, randomized controlled trials are needed to further define the role of aspirin and other antiplatelet therapies as adjunctive treatments in the management of COVID-19.

The RECOVERY trial randomized hospitalized adults with COVID-19 to receive usual care plus aspirin 150 mg per day (n = 7,351) or usual care only (n = 7,541).³⁹ At enrollment, 38% of the patients required NIV or mechanical ventilation. Mortality at 28 days was 17% in both arms (rate ratio 0.96; 95% CI,

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

0.89–1.04). Among patients who were not receiving mechanical ventilation at baseline, there was no difference between the arms in the proportion of patients who progressed to requiring mechanical ventilation or who died (21% in the aspirin arm vs. 22% in the usual care arm; rate ratio 0.96; 95% CI, 0.90–1.03). Among those treated with aspirin, the incidence of thrombotic events was lower (4.6% vs. 5.3%; absolute difference 0.6%; SE 0.4%), and the incidence of major bleeding events was higher (1.6% vs. 1.0%; absolute difference 0.6%; SE 0.2%). Overall, in this large trial of hospitalized patients with COVID-19, the use of aspirin was associated with an increase in the incidence of major bleeding events and did not reduce the risk of death.

The ACTIV-4a trial compared the use of P2Y12 inhibitor therapy plus a therapeutic dose of heparin to a therapeutic dose of heparin alone in hospitalized patients with COVID-19. In this study, enrollment of noncritically ill patients was stopped early due to futility; the combination therapy did not increase the number of organ support-free days.⁴⁰ The limitations of this study include the open-label design, the use of different P2Y12 inhibitors, and the trial size.

Based on the findings of the ACTIV-4a and RECOVERY trials, the Panel **recommends against** the use of antiplatelet therapy to prevent COVID-19 progression or death in noncritically ill patients (**BIIa**).

The REMAP-CAP study team randomized critically ill patients with COVID-19 to receive aspirin (n = 565), a P2Y12 inhibitor (n = 455), or no antiplatelet therapy (n = 529).⁴¹ Treatment continued for 14 days or until hospital discharge, whichever came first. The aspirin and P2Y12 inhibitor arms were pooled for analysis because the criteria for equivalence were met. The trial was stopped early due to futility, as the median number of organ support-free days did not differ between the pooled antiplatelet arm and the control arm (7 days; IQR 1–16 days; 95.7% posterior probability of futility). There was no statistically significant difference between the arms in the number of patients who survived to hospital discharge (723 of 1,011 patients [71.5%] in the pooled antiplatelet arm vs. 354 of 521 patients [67.9%] in the control arm; median-adjusted OR 1.27; 95% CrI, 0.99–1.62). The pooled antiplatelet arm had improved survival by 90 days (median aHR 1.22; 95% CrI, 1.06–1.40). The use of antiplatelet therapy was associated with an increased incidence of major bleeding (2.1% in the pooled antiplatelet arm vs. 0.4% in the control arm; aOR 2.97; 95% CrI, 1.23–8.28; adjusted absolute risk difference of 0.8%; 95% CrI, 0.1% to 2.7%).

In the RECOVERY trial, the use of aspirin therapy was not associated with a reduction in mortality in the subgroups of patients who required NIV or mechanical ventilation at baseline. In the REMAP-CAP trial, administering antiplatelet therapy to critically ill patients with COVID-19 improved 90-day survival but did not increase the number of organ support-free days. In both studies, the use of antiplatelet therapy was associated with an increased risk of bleeding. The COVID-PACT trial randomized 292 adult patients with COVID-19 who required ICU-level care to receive either clopidogrel or no antiplatelet therapy.³³ There was no difference between the arms in the incidence of VTE, arterial thrombotic events, or bleeding.

Given the inconsistent results of these trials, there is insufficient evidence for the Panel to recommend either for or against the use of antiplatelet therapy in critically ill patients with COVID-19. Eligible patients should be encouraged to participate in clinical trials that are evaluating the use of antiplatelet therapy.

The clinical data for the trials discussed above are summarized in Table 6b.

Thrombolytic Therapy

Clinical trials are evaluating the effects of thrombolysis on mortality and the progression of COVID-19. There is insufficient evidence for the Panel to recommend either for or against the use of thrombolytic agents for VTE prophylaxis in hospitalized patients with COVID-19 outside of a clinical trial.

Patients Discharged From the Hospital

For patients with a high risk of VTE who do not have COVID-19, post-discharge prophylaxis has been shown to be beneficial. The Food and Drug Administration approved the use of rivaroxaban 10 mg once daily for 31 to 39 days in these patients.^{42,43} Inclusion criteria for the trials that studied post-discharge VTE prophylaxis included:

- A VTE risk score of ≥4 on the modified International Medical Prevention Registry on Venous Thromboembolism (IMPROVE) tool;⁴⁴ or
- A VTE risk score ≥ 2 on the modified IMPROVE tool⁴⁵ and a D-dimer level >2 times ULN.⁴²

The MICHELLE trial randomized 320 patients with COVID-19 and an IMPROVE score of \geq 4 or 2 to 3 with a D-dimer level >500 ng/mL to receive rivaroxaban 10 mg orally once daily or no anticoagulation for 35 days.⁴⁶ The primary outcome was a composite of symptomatic VTE, fatal pulmonary embolism, symptomatic arterial thromboembolism, cardiovascular death, or asymptomatic VTE detected on screening imaging at Day 35. Five patients (3%) who were treated with rivaroxaban and 15 patients (9%) who did not receive anticoagulation experienced a thrombotic event (relative risk 0.33; 95% CI, 0.13–0.9). One patient who received rivaroxaban and 10 patients who did not receive anticoagulation experienced symptomatic events. No major bleeding events occurred, and 2 patients in each arm had clinically relevant, nonmajor bleeding. The open-label design and the inclusion of asymptomatic events that were detected on screening ultrasounds and computed tomography scans may have biased the results. Additionally, two-thirds of the screened patients did not meet the eligibility criteria for the trial, which limits the generalizability of the results.

The ACTIV-4c trial randomized 1,217 patients who were hospitalized for symptomatic COVID-19 for >48 hours to receive apixaban 2.5 mg orally twice daily or placebo at hospital discharge.⁴⁷ The 30-day composite endpoint of all-cause mortality, venous thrombosis, or arterial thrombosis occurred in 2.13% of patients in the apixaban arm and in 2.31% of patients in the placebo arm. Major bleeding events were infrequent, occurring in 2 patients in the apixaban arm (0.4%) and in 1 patient in the placebo arm (0.2%). The trial's leadership and sponsors stopped the trial early because the event rate for the composite endpoint was lower than expected and the decreasing number of hospitalizations for people with COVID-19 made recruitment difficult. Based on the results of the MICHELLE and ACTIV-4c trials, the Panel **recommends against** routinely continuing VTE prophylaxis in patients with COVID-19 after hospital discharge unless they have another indication for anticoagulation (**AIIa**).

Although there is no clear benefit of administering anticoagulation after hospital discharge in all patients with COVID-19, results from the MICHELLE trial, which evaluated patients with COVID-19, and the MARINER trial, which evaluated patients who were hospitalized for other conditions and who had risk factors for VTE, suggest a possible benefit of using anticoagulation after discharge in patients who are at high risk of VTE. The need for VTE prophylaxis after a COVID-19-related hospital discharge should be assessed on a case-by-case basis. The criteria for assessing the risk of VTE in these patients are the same as the criteria used for patients who are hospitalized for other acute illnesses.

Children With COVID-19 or MIS-C

For the Panel's recommendations on the use of antithrombotic therapy in children, see <u>Therapeutic</u> <u>Management of Hospitalized Children With COVID-19</u> and <u>Therapeutic Management of Hospitalized</u> <u>Children With MIS-C, Plus a Discussion on MIS-A</u>.

Pregnant and Lactating Patients

Because pregnancy is a hypercoagulable state, the risk of thromboembolism is greater in pregnant individuals than in nonpregnant individuals.⁴⁸ It is not yet known whether COVID-19 increases this

risk, though some data do suggest that there is an increased risk. A cohort study in California compared perinatal outcomes among almost 44,000 pregnant people with and without COVID-19.⁴⁹ After adjusting for demographic factors and comorbidities, those with COVID-19 had a higher risk of severe maternal morbidity, preterm birth, and VTE.

In several other cohort studies of pregnant women with COVID-19 in the United States and Europe, VTE was not reported as a complication even among women with severe disease, although the receipt of prophylactic or therapeutic anticoagulation varied across the studies.⁵⁰⁻⁵² The <u>American College of</u> <u>Obstetricians and Gynecologists (ACOG)</u> advises that although there are not enough data to recommend either for or against the use of thromboprophylaxis, in the setting of COVID-19 during pregnancy, VTE prophylaxis can reasonably be considered for pregnant individuals hospitalized with COVID-19, particularly for those who have severe disease. If there are no contraindications, the Society for Maternal-Fetal Medicine recommends the use of heparin or LMWH in pregnant patients who are critically ill or receiving mechanical ventilation.⁵³ Several professional societies, including the American Society of Hematology and ACOG, have guidelines that specifically address the management of VTE in the context of pregnancy.^{54,55} If delivery is imminent, or if there are other risks for bleeding, the risk of bleeding may outweigh the potential benefit of using VTE prophylaxis in pregnant individuals.

Outside of pregnancy, D-dimer levels have been used to stratify VTE risk. However, physiologic increases in D-dimer levels may occur during pregnancy, making elevated D-dimer values an unreliable predictor that should not be used to evaluate VTE risk during pregnancy in the setting of COVID-19.⁵⁶⁻⁵⁸

In general, heparin compounds are the preferred anticoagulants to use during pregnancy. Because of its reliability and ease of administration, LMWH is recommended rather than UFH for the prevention and treatment of VTE in pregnant people.⁵⁵ Direct-acting anticoagulants are not routinely recommended for use during pregnancy because of a lack of safety data for pregnant individuals.⁵⁴ The use of warfarin to prevent or treat VTE should be avoided in pregnant individuals regardless of their COVID-19 status, especially during the first trimester, due to the concern for teratogenicity.

Specific recommendations for pregnant or lactating individuals with COVID-19 include:

- The Panel recommends that pregnant patients who are receiving anticoagulant or antiplatelet therapies for underlying conditions continue these medications after they receive a diagnosis of COVID-19 (AIII).
- The Panel recommends the use of a prophylactic dose of anticoagulation for pregnant patients who are hospitalized for manifestations of COVID-19, unless a contraindication exists (**BIII**).
- Because pregnant patients were not included in most of the clinical trials that evaluated the use of therapeutic anticoagulation in the setting of COVID-19, there is insufficient evidence for the Panel to recommend either for or against the use of therapeutic anticoagulation in pregnant patients with COVID-19 who do not have evidence of VTE.
- As in nonpregnant patients with COVID-19, VTE prophylaxis after hospital discharge is not routinely recommended for pregnant patients (**BIII**). Clinicians should consider an individual patient's VTE risk factors when making decisions about continuing VTE prophylaxis after discharge in pregnant or postpartum patients.
- The use of anticoagulation therapy during labor and delivery requires specialized care and planning. The management of anticoagulation therapy in pregnant patients with COVID-19 should be similar to the management used for pregnant patients with other conditions (AIII).
- UFH, LMWH, and warfarin do not accumulate in breast milk and do not induce an anticoagulant effect in the newborn; therefore, they can be used by breastfeeding individuals who require VTE prophylaxis or treatment (AIII).

References

- 1. Han H, Yang L, Liu R, et al. Prominent changes in blood coagulation of patients with SARS-CoV-2 infection. *Clin Chem Lab Med.* 2020;58(7):1116-1120. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32172226</u>.
- 2. Driggin E, Madhavan MV, Bikdeli B, et al. Cardiovascular considerations for patients, health care workers, and health systems during the COVID-19 pandemic. *J Am Coll Cardiol*. 2020;75(18):2352-2371. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32201335.
- 3. Guan WJ, Ni ZY, Hu Y, et al. Clinical characteristics of coronavirus disease 2019 in China. *N Engl J Med.* 2020;382(18):1708-1720. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32109013</u>.
- 4. Tang N, Bai H, Chen X, et al. Anticoagulant treatment is associated with decreased mortality in severe coronavirus disease 2019 patients with coagulopathy. *J Thromb Haemost*. 2020;18(5):1094-1099. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32220112.
- 5. Nopp S, Moik F, Jilma B, Pabinger I, Ay C. Risk of venous thromboembolism in patients with COVID-19: a systematic review and meta-analysis. *Res Pract Thromb Haemost*. 2020;4(7):1178-1191. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33043231.
- Cohen AT, Davidson BL, Gallus AS, et al. Efficacy and safety of fondaparinux for the prevention of venous thromboembolism in older acute medical patients: randomised placebo controlled trial. *BMJ*. 2006;332(7537):325-329. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/16439370</u>.
- 7. Leizorovicz A, Cohen AT, Turpie AG, et al. Randomized, placebo-controlled trial of dalteparin for the prevention of venous thromboembolism in acutely ill medical patients. *Circulation*. 2004;110(7):874-879. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/15289368</u>.
- Samama MM, Cohen AT, Darmon JY, et al. A comparison of enoxaparin with placebo for the prevention of venous thromboembolism in acutely ill medical patients. *N Engl J Med.* 1999;341(11):793-800. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/10477777</u>.
- Fraisse F, Holzapfel L, Couland JM, et al. Nadroparin in the prevention of deep vein thrombosis in acute decompensated COPD. *Am J Respir Crit Care Med.* 2000;161(4):1109-1114. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/10764298</u>.
- PROTECT Investigators for the Canadian Critical Care Trials Group, Australian and New Zealand Intensive Care Society Clinical Trials Group. Dalteparin versus unfractionated heparin in critically ill patients. N Engl J Med. 2011;364(14):1305-1314. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21417952</u>.
- Shorr AF, Williams MD. Venous thromboembolism in critically ill patients: observations from a randomized trial in sepsis. *Thromb Haemost*. 2009;101(1):139-144. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/19132200</u>.
- 12. Kaplan D, Casper TC, Elliott CG, et al. VTE incidence and risk factors in patients with severe sepsis and septic shock. *Chest.* 2015;148(5):1224-1230. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26111103</u>.
- Moores LK, Tritschler T, Brosnahan S, et al. Thromboprophylaxis in patients with COVID-19: a brief update to the CHEST guideline and expert panel report. *Chest.* 2022;162(1):213-225. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35167861</u>.
- American Society of Hematology. ASH guidelines on use of anticoagulation in patients with COVID-19. 2022. Available at: <u>https://www.hematology.org/education/clinicians/guidelines-and-quality-care/</u> <u>clinical-practice-guidelines/venous-thromboembolism-guidelines/ash-guidelines-on-use-of-anticoagulation-in-</u> <u>patients-with-covid-19</u>. Accessed September 14, 2023.
- Barnes GD, Burnett A, Allen A, et al. Thromboembolism and anticoagulant therapy during the COVID-19 pandemic: interim clinical guidance from the Anticoagulation Forum. *J Thromb Thrombolysis*. 2020;50(1):72-81. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32440883</u>.
- 16. Thachil J, Tang N, Gando S, et al. ISTH interim guidance on recognition and management of coagulopathy in COVID-19. *J Thromb Haemost*. 2020;18(5):1023-1026. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/32338827</u>.

COVID-19 Treatment Guidelines

- 17. Marietta M, Ageno W, Artoni A, et al. COVID-19 and haemostasis: a position paper from Italian Society on Thrombosis and Haemostasis (SISET). *Blood Transfus*. 2020;18(3):167-169. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32281926</u>.
- 18. National Institute for Health and Care Excellence. COVID-19 rapid guideline: managing COVID-19. 2023. Available at: <u>https://www.nice.org.uk/guidance/ng191</u>. Accessed June 8, 2023.
- 19. Royal College of Physicians. Clinical guide for the prevention, detection and management of thromboembolic disease in patients with COVID-19. 2020. Available at: <u>https://icmanaesthesiacovid-19.org/clinical-guide-prevention-detection-and-management-of-vte-in-patients-with-covid-19</u>. Accessed June 8, 2023.
- 20. Potpara T, Angiolillo DJ, Bikdeli B, et al. Antithrombotic therapy in arterial thrombosis and thromboembolism in COVID-19: an American College of Chest Physicians expert panel report. *Chest.* 2023;Published online ahead of print. Available at: https://pubmed.ncbi.nlm.nih.gov/37392958.
- 21. B.C. Provincial Academic Detailing Service. Switching from warfarin to a direct acting oral anticoagulant (DOAC): a practical guide for B.C. primary care clinicians (April 2020). 2020. Available at: https://www2.gov.bc.ca/assets/gov/health/health-drug-coverage/pharmacare/warfarin_to_doac_switch.pdf. Accessed June 12, 2023.
- 22. Kahn SR, Lim W, Dunn AS, et al. Prevention of VTE in nonsurgical patients: antithrombotic therapy and prevention of thrombosis, 9th ed: American College of Chest Physicians evidence-based clinical practice guidelines. *Chest.* 2012;141(2 Suppl):e195S-e226S. Available at: https://www.ncbi.nlm.nih.gov/pubmed/22315261.
- 23. Connors JM, Brooks MM, Sciurba FC, et al. Effect of antithrombotic therapy on clinical outcomes in outpatients with clinically stable symptomatic COVID-19: the ACTIV-4B randomized clinical trial. *JAMA*. 2021;326(17):1703-1712. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34633405.
- 24. Cools F, Virdone S, Sawhney J, et al. Thromboprophylactic low-molecular-weight heparin versus standard of care in unvaccinated, at-risk outpatients with COVID-19 (ETHIC): an open-label, multicentre, randomised, controlled, Phase 3b trial. *Lancet Haematol*. 2022;9(8):e594-e604. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35779560.
- 25. Barco S, Voci D, Held U, et al. Enoxaparin for primary thromboprophylaxis in symptomatic outpatients with COVID-19 (OVID): a randomised, open-label, parallel-group, multicentre, Phase 3 trial. *Lancet Haematol*. 2022;9(8):e585-e593. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35779558</u>.
- 26. Rentsch CT, Beckman JA, Tomlinson L, et al. Early initiation of prophylactic anticoagulation for prevention of coronavirus disease 2019 mortality in patients admitted to hospital in the United States: cohort study. *BMJ*. 2021;372:n311. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33574135.
- ATTACC, ACTIV-4a, and REMAP-CAP Investigators. Therapeutic anticoagulation with heparin in noncritically ill patients with COVID-19. *N Engl J Med.* 2021;385(9):790-802. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34351721</u>.
- 28. Stone GW, Farkouh ME, Lala A, et al. Randomized trial of anticoagulation strategies for noncritically ill patients hospitalized with COVID-19. *J Am Coll Cardiol*. 2023;81(18):1747-1762. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36889611</u>.
- 29. Sholzberg M, Tang GH, Rahhal H, et al. Effectiveness of therapeutic heparin versus prophylactic heparin on death, mechanical ventilation, or intensive care unit admission in moderately ill patients with COVID-19 admitted to hospital: RAPID randomised clinical trial. *BMJ*. 2021;375:n2400. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34649864.
- 30. Spyropoulos AC, Goldin M, Giannis D, et al. Efficacy and safety of therapeutic-dose heparin vs standard prophylactic or intermediate-dose heparins for thromboprophylaxis in high-risk hospitalized patients with COVID-19: the HEP-COVID randomized clinical trial. *JAMA Intern Med.* 2021;181(12):1612-1620. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34617959</u>.
- 31. INSPIRATION Investigators. Effect of intermediate-dose vs standard-dose prophylactic anticoagulation on thrombotic events, extracorporeal membrane oxygenation treatment, or mortality among patients

with COVID-19 admitted to the intensive care unit: the INSPIRATION randomized clinical trial. *JAMA*. 2021;325(16):1620-1630. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33734299</u>.

- 32. Labbé V, Contou D, Heming N, et al. Effects of standard-dose prophylactic, high-dose prophylactic, and therapeutic anticoagulation in patients with hypoxemic COVID-19 pneumonia: the ANTICOVID randomized clinical trial. JAMA Intern Med. 2023;183(6):520-531. Available at: https://pubmed.ncbi.nlm.nih.gov/36946232.
- 33. Bohula EA, Berg DD, Lopes MS, et al. Anticoagulation and antiplatelet therapy for prevention of venous and arterial thrombotic events in critically ill patients with COVID-19: COVID-PACT. *Circulation*. 2022;146(18):1344-1356. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36036760</u>.
- 34. Lopes RD, de Barros E Silva PGM, Furtado RHM, et al. Therapeutic versus prophylactic anticoagulation for patients admitted to hospital with COVID-19 and elevated D-dimer concentration (ACTION): an open-label, multicentre, randomised, controlled trial. *Lancet*. 2021;397(10291):2253-2263. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34097856</u>.
- 35. Chow JH, Yin Y, Yamane DP, et al. Association of prehospital antiplatelet therapy with survival in patients hospitalized with COVID-19: a propensity score-matched analysis. *J Thromb Haemost*. 2021;19(11):2814-2824. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34455688</u>.
- 36. Chow JH, Khanna AK, Kethireddy S, et al. Aspirin use is associated with decreased mechanical ventilation, intensive care unit admission, and in-hospital mortality in hospitalized patients with coronavirus disease 2019. *Anesth Analg.* 2021;132(4):930-941. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33093359</u>.
- 37. Chow JH, Rahnavard A, Gomberg-Maitland M, et al. Association of early aspirin use with in-hospital mortality in patients with moderate COVID-19. *JAMA Netw Open*. 2022;5(3):e223890. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35323950</u>.
- Abdi M, Lamardi ZH, Shirjan F, et al. The effect of aspirin on the prevention of pro-thrombotic states in hospitalized COVID-19 patients: systematic review. *Cardiovasc Hematol Agents Med Chem*. 2022;20(3):189-196. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35366783</u>.
- 39. RECOVERY Collaborative Group. Aspirin in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2022;399(10320):143-151. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34800427.
- Berger JS, Kornblith LZ, Gong MN, et al. Effect of P2Y12 inhibitors on survival free of organ support among non-critically ill hospitalized patients with COVID-19: a randomized clinical trial. *JAMA*. 2022;327(3):227-236. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35040887</u>.
- 41. REMAP-CAP Writing Committee for the REMAP-CAP Investigators. Effect of antiplatelet therapy on survival and organ support-free days in critically ill patients with COVID-19: a randomized clinical trial. *JAMA*. 2022;327(13):1247-1259. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35315874.
- 42. Spyropoulos AC, Lipardi C, Xu J, et al. Modified IMPROVE VTE risk score and elevated D-dimer identify a high venous thromboembolism risk in acutely ill medical population for extended thromboprophylaxis. *TH Open*. 2020;4(1):e59-e65. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32190813.
- 43. Cohen AT, Harrington RA, Goldhaber SZ, et al. Extended thromboprophylaxis with betrixaban in acutely ill medical patients. *N Engl J Med*. 2016;375(6):534-544. Available at: https://www.ncbi.nlm.nih.gov/pubmed/27232649.
- 44. Spyropoulos AC, Anderson FA Jr, FitzGerald G, et al. Predictive and associative models to identify hospitalized medical patients at risk for VTE. *Chest.* 2011;140(3):706-714. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21436241</u>.
- 45. Gibson CM, Spyropoulos AC, Cohen AT, et al. The IMPROVEDD VTE risk score: incorporation of D-dimer into the IMPROVE score to improve venous thromboembolism risk stratification. *TH Open*. 2017;1(1):e56-e65. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31249911</u>.
- 46. Ramacciotti E, Barile Agati L, Calderaro D, et al. Rivaroxaban versus no anticoagulation for post-discharge

thromboprophylaxis after hospitalisation for COVID-19 (MICHELLE): an open-label, multicentre, randomised, controlled trial. *Lancet*. 2022;399(10319):50-59. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34921756</u>.

- 47. Wang TY, Wahed AS, Morris A, et al. Effect of thromboprophylaxis on clinical outcomes after COVID-19 hospitalization. *Ann Intern Med.* 2023;176(4):515-523. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36940444</u>.
- 48. Heit JA, Kobbervig CE, James AH, et al. Trends in the incidence of venous thromboembolism during pregnancy or postpartum: a 30-year population-based study. *Ann Intern Med.* 2005;143(10):697-706. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/16287790</u>.
- Ferrara A, Hedderson MM, Zhu Y, et al. Perinatal complications in individuals in California with or without SARS-CoV-2 infection during pregnancy. *JAMA Intern Med.* 2022;182(5):503-512. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35311909</u>.
- 50. Breslin N, Baptiste C, Gyamfi-Bannerman C, et al. Coronavirus disease 2019 infection among asymptomatic and symptomatic pregnant women: two weeks of confirmed presentations to an affiliated pair of New York City hospitals. *Am J Obstet Gynecol MFM*. 2020;2(2):100118. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32292903.
- 51. Knight M, Bunch K, Vousden N, et al. Characteristics and outcomes of pregnant women admitted to hospital with confirmed SARS-CoV-2 infection in UK: national population based cohort study. *BMJ*. 2020;369:m2107. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32513659</u>.
- 52. Delahoy MJ, Whitaker M, O'Halloran A, et al. Characteristics and maternal and birth outcomes of hospitalized pregnant women with laboratory-confirmed COVID-19—COVID-NET, 13 states, March 1–August 22, 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(38):1347-1354. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32970655</u>.
- 53. Society for Maternal-Fetal Medicine. Management considerations for pregnant patients with COVID-19. 2021. Available at: <u>https://s3.amazonaws.com/cdn.smfm.org/media/2734/SMFM_COVID_Management_of_COVID_pos_preg_patients_2-2-21_(final).pdf</u>.
- 54. Bates SM, Rajasekhar A, Middeldorp S, et al. American Society of Hematology 2018 guidelines for management of venous thromboembolism: venous thromboembolism in the context of pregnancy. *Blood Adv*. 2018;2(22):3317-3359. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30482767</u>.
- 55. American College of Obstetricians and Gynecologists. ACOG practice bulletin no. 196 summary: thromboembolism in pregnancy. *Obstet Gynecol*. 2018;132(1):243-248. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29939933</u>.
- 56. Wang M, Lu S, Li S, Shen F. Reference intervals of D-dimer during the pregnancy and puerperium period on the STA-R evolution coagulation analyzer. *Clin Chim Acta*. 2013;425:176-180. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/23954836</u>.
- 57. Réger B, Péterfalvi A, Litter I, et al. Challenges in the evaluation of D-dimer and fibrinogen levels in pregnant women. *Thromb Res.* 2013;131(4):e183-e187. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/23481480</u>.
- 58. Hu W, Wang Y, Li J, et al. The predictive value of D-dimer test for venous thromboembolism during puerperium: a prospective cohort study. *Clin Appl Thromb Hemost*. 2020;26:1076029620901786. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32090610</u>.

Table 6a. Anticoagulant Therapy: Selected Clinical Trial Data

Last Updated: October 10, 2023

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for anticoagulant therapy. The studies summarized below are those that have had the greatest impact on the Panel's recommendations.

Methods	Results	Limitations and Interpretation	
ATTACC/ACTIV-4a/REMAP-CAP: Multiplatform, Open-Label RCT of Therapeutic Anticoagulation in Noncritically III, Hospitalized Patients With COVID-19 in 9 Countries ¹			
Key Inclusion Criterion	Participant Characteristics	Key Limitations	
 Key Inclusion Criterion Hospitalized with laboratory-confirmed SARS-CoV-2 infection without need for HFNC oxygen, NIV, MV, vasopressors, or inotropes Key Exclusion Criteria Hospital discharge expected in ≤72 hours Requirement for therapeutic anticoagulation or dual antiplatelet therapy High bleeding risk Interventions Therapeutic UFH or LMWH for 14 days or until hospital discharge, whichever came first (n = 1,190) SOC, which included prophylactic UFH or LMWH (n = 1,054) Primary Endpoint Organ support-free days at Day 21, as measured by an OS 	 Participant Characteristics Median age 59 years; 59% men; median BMI 30 52% with HTN; 30% with DM; 11% with CVD 66% required low-flow oxygen D-dimer: 48.4% <2 times ULN 28.4% ≥2 times ULN 23.1% unknown 62% on corticosteroids; 36% on RDV Primary Outcomes Therapeutic anticoagulation superior to SOC for organ support-free days (aOR 1.27; 95% Crl, 1.03–1.58; 99% posterior probability) 4% absolute difference in survival until hospital discharge without organ support that favored therapeutic arm (95% Crl, 0.5–7.2) Outcome consistent across D-dimer stratum Secondary Outcomes Survival until hospital discharge: 92% in both arms 	 Key Limitations Open-label study Anticoagulation dose varied in SOC arm (27% received intermediate-dose thromboprophylaxis) Inclusion criteria for hospital LOS and ICU-level care differed across trials. Only enrolled 17% of screened patients Interpretation Therapeutic heparin increased the number of organ support-free days and decreased the number of patients requiring organ support. Therapeutic heparin did not significantly affect hospital LOS or the number of major thrombosis events or deaths. Major bleeds occurred 1% more frequently in the therapeutic arm than in the SOC arm. 	
Key Secondary Endpoints	 No difference between arms in hospital LOS (aOR 1.03; 95% Crl, 0.94–1.13) 		
Survival until hospital dischargeHospital LOS	 Thrombosis: 1% in therapeutic arm vs. 2% in SOC arm Major bleeding events: 2% in therapeutic arm vs. 1% in SOC arm 		
Thrombosis or major bleeding events			

Methods	Results	Limitations and Interpretation	
RAPID: Open-Label RCT of Therapeutic Heparin in	RAPID: Open-Label RCT of Therapeutic Heparin in Moderately III, Hospitalized Patients With COVID-19 in 6 Countries ²		
Key Inclusion Criteria	Participant Characteristics	Key Limitations	
 Hospitalized with COVID-19 and D-dimer level ≥2 times ULN or any elevated D-dimer level and 	• Median age 60 years; 57% men; mean BMI 30	Open-label study	
$SpO_2 \le 93\%$ on room air	• 48% with HTN; 34% with DM; 7% with CVD	Only enrolled 12% of screened patients	
Hospitalized <5 days	91% had hypoxia; 6% received HFNC oxygen	Interpretation	
Key Exclusion Criteria	 D-dimer: 49% <2 times ULN 	 Compared to prophylactic heparin, therapeutic heparin reduced mortality (a secondary endpoint) 	
Indication for therapeutic anticoagulation	• 51% ≥2 times ULN	but had no effect on the composite primary	
Dual antiplatelet therapy	69% on corticosteroids	endpoint of ICU admission, the need for NIV or MV, or death up to 28 days.	
High bleeding risk	Primary Outcome	There were no differences between the arms in	
 Interventions Therapeutic UFH or LMWH for 28 days or until discharge or death (n = 228) 	 Composite of ICU admission, NIV or MV, or death up to 28 days: 16% in therapeutic arm vs. 22% in prophylactic arm (OR 0.69; 95% CI, 0.43–1.10) 	the percentages of patients who experienced VTE or major bleeding events.	
 Prophylactic UFH or LMWH for 28 days or until discharge or death (n = 237) 	Secondary Outcomes		
Primary Endpoint	 All-cause death: 2% in therapeutic arm vs. 8% in prophylactic arm (OR 0.22; 95% Cl, 0.07–0.65) 		
Composite of ICU admission, NIV or MV, or death up to 28 days	 Mean number of organ support-free days: 26 in therapeutic arm vs. 24 in prophylactic arm (OR 1.41; 		
Key Secondary Endpoints	95% Cl, 0.9–2.21)		
All-cause death	• No difference between arms for VTE (1% in therapeutic		
Mean number of organ support-free days	arm vs. 3% in prophylactic arm) or major bleeding events (1% in therapeutic arm vs. 2% in prophylactic		
• VTE	arm)		
Major bleeding events	• Mean number of hospital-free days alive: 20 in		
Mean number of hospital-free days alive	therapeutic arm vs. 18 in prophylactic arm (OR 1.09; 95% Cl, 0.79–1.50)		

Methods	Results	Limitations and Interpretation
HEP-COVID: Open-Label RCT of Therapeutic Heparin in High-Risk, Hospitalized Patients With COVID-19 in the United States ³		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Hospitalized with COVID-19 and required 	• Median age 67 years; 54% men; mean BMI 30	Open-label study
supplemental oxygen	 60% with HTN; 37% with DM; 75% with CVD 	Only enrolled 2% of screened patients
 D-dimer >4 times ULN or sepsis-induced coagulopathy score of ≥4 	 64% received oxygen via nasal cannula; 15% received high-flow oxygen or NIV; 5% received MV 	Interpretation
Hospitalized <72 hours	80% on corticosteroids	 Compared to usual care, therapeutic LMWH reduced the incidence of VTE, ATE, and death.
Key Exclusion Criteria	Primary Outcomes	Among patients who were not in the ICU,
 Indication for therapeutic anticoagulation 	• Composite of VTE, ATE, or death within 32 days: 29% in	therapeutic LMWH significantly reduced the
Dual antiplatelet therapy	therapeutic arm vs. 42% in usual care arm (relative risk	percentage of patients who experienced thrombotic events and did not increase the
High bleeding risk	0.68; 95% Cl, 0.49–0.96)	percentage of patients who experienced major
• CrCl <15 mL/min	 Death: 19% in therapeutic arm vs. 25% in usual care arm (relative risk 0.78; 95% Cl, 0.49–1.23) 	bleeding events.
Interventions	• Thrombotic events: 11% in therapeutic arm vs.	
• Therapeutic LMWH until hospital discharge or primary endpoint met (n = 129)	29% in usual care arm (relative risk 0.37; 95% Cl, 0.21–0.66)	
• Usual care of prophylactic or intermediate- dose LMWH until hospital discharge or primary endpoint met (n = 124)	• Non-ICU stratum composite of VTE, ATE, or death within 32 days: 17% in therapeutic arm vs. 36% in usual care arm (relative risk 0.46; 95% Cl, 0.27–0.81)	
Primary Endpoint	Safety Outcomes	
• Composite of VTE, ATE, or death from any cause within 32 days of randomization	 Major bleeding events within 32 days: 5% in therapeutic arm vs. 2% in usual care arm (relative risk) 	
Key Safety Endpoint	2.88; 95% Cl, 0.59–14.02)	
Major bleeding events within 32 days	 Non-ICU stratum major bleeding events within 32 days: 2% in both arms 	

Methods	Results	Limitations and Interpretation
ACTION: Open-Label RCT of Therapeutic Rivaroxaban in Hospitalized Patients With COVID-19 in Brazil ⁴		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Hospitalized for COVID-19 with elevated D-dimer level 	 Median age 57 years; 60% men; mean BMI 30 49% with HTN; 24% with DM; 5% with coronary 	 Open-label study Only enrolled 18% of screened patients
 Symptoms for ≤14 days 	disease	 Therapeutic rivaroxaban was administered for a
Key Exclusion CriteriaIndication for therapeutic anticoagulation	Critically ill: 7% in therapeutic arm vs. 5% in usual care arm	longer duration than prophylactic anticoagulation (30 days vs. a mean duration of 8 days).
 CrCl <30 mL/min P2Y12 inhibitor therapy or aspirin >100 mg High bleeding risk 	 75% required oxygen: 60% required low-flow oxygen; 8% required HFNC oxygen; 1% required NIV; 6% required MV 83% on corticosteroids 	 Interpretation When compared with usual care, therapeutic rivaroxaban did not reduce mortality, hospital duration or the persentence
Interventions	Primary Outcome	duration, oxygen use duration, or the percentage of patients who experienced thrombosis.
• Therapeutic anticoagulation for 30 days: rivaroxaban 15 mg or 20 mg once daily; if clinically unstable, enoxaparin 1 mg/kg twice daily or UFH (n = 311)	 No difference between arms in the composite of time to death, hospital duration, or oxygen use duration through Day 30 (win ratio 0.86; 95% Cl, 0.59–1.22) 	 Patients who received therapeutic rivaroxaban had more clinically relevant nonmajor bleeding events than those who received usual care.
 Usual care prophylactic anticoagulation with enoxaparin or UFH during hospitalization (n = 304) 	 Secondary Outcomes No difference between therapeutic and usual care arms in: 	 The longer duration of therapy in the rivaroxaban arm may have influenced the difference in bleeding events.
Primary Endpoint	Thrombosis: 7% vs. 10%	
• Hierarchical composite of time to death, hospital duration, or oxygen use duration through Day 30	 Mortality: 11% vs. 8% Any bleeding events: 12% in therapeutic arm vs. 3% in 	
Key Secondary Endpoints	usual care arm	
• Thrombosis, with and without all-cause death	 Major bleeding events: 3% in therapeutic arm vs. 1% in usual care arm 	
MortalityBleeding events	 Clinically relevant, nonmajor bleeding events: 5% in therapeutic arm vs. 1% in usual care arm 	

Methods	Results	Limitations and Interpretation
FREEDOM: RCT of Anticoagulation Strategies in Noncritically III Patients Who Were Hospitalized With COVID-19 in 10 Countries ⁵		
Key Inclusion Criterion	Participant Characteristics	Key Limitations
• Hospitalized for symptomatic COVID-19 for <48	Median age 52 years; 59% men; mean BMI 26	Open-label study
hours	• 32% with HTN; 19% with DM	• Terminated early due to slow recruitment (3,452
Key Exclusion Criteria	 22% on corticosteroids; 10% on RDV 	of 3,600 planned patients recruited)
Indication for therapeutic anticoagulation	Primary Outcome	 Minimal treatment with RDV or DEX as SOC for COVID-19
• CrCl <30 mL/min	30-day composite outcome: 11.3% in combined	
P2Y12 inhibitor therapy or aspirin >100 mg per	therapeutic arms vs. 13.2% in prophylactic arm (HR 0.85; 95% Cl, 0.69–1.04; $P = 0.11$)	 Interpretation When compared with prophylactic enoxaparin,
dayAnticipated hospitalization for <72 hours	 Primary endpoint was not statistically significant when 	therapeutic apixaban and therapeutic enoxaparin
	therapeutic enoxaparin or apixaban were compared to	did not reduce 30-day mortality, the need for ICU-
Interventions	prophylactic enoxaparin.	level care, or the occurrence of thromboembolism or ischemic stroke.
• Therapeutic apixaban 5 mg twice daily (n = 1,121)	Secondary Outcomes	 Fewer patients died in the therapeutic enoxaparin
 Therapeutic enoxaparin 1 mg/kg twice daily (n = 1,136) 	• All-cause mortality: 4.9% in therapeutic enoxaparin arm vs. 7.0% in prophylactic enoxaparin arm (HR 0.69;	and therapeutic apixaban arms than in the prophylactic enoxaparin arm.
• Usual care prophylactic enoxaparin ($n = 1,141$)	95% Cl, 0.49–0.99)	• There were no statistically significant differences
Primary Endpoint	 All-cause mortality: 5.0% in therapeutic apixaban arm vs. 7.0% in prophylactic enoxaparin arm (HR 0.7; 95%) 	between the arms in the percentages of patients
30-day composite of all-cause mortality, need	Cl, 0.49–0.99)	who experienced severe bleeding events.
for ICU-level care, systemic thromboembolism, or ischemic stroke. Endpoint assessed for the combined therapeutic arms vs. the prophylactic arm.	• BARC type 3 or 5 bleeding: 0.4% in combined therapeutic arms vs. 0.1% in prophylactic arm (IRR 3.96; 95% Cl, 0.50–31.27)	
Key Secondary Endpoints		
All-cause mortality		
Bleeding events (BARC type 3 or 5)		

Methods	Results	Limitations and Interpretation
COVID-PACT: Open-Label RCT of Full-Dose Versus Prophylactic-Dose Anticoagulation in Adults With COVID-19 Who Were Receiving Intensive Care Unit- Level Care in the United States ⁶		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Aged ≥18 years 	Median age 61 years; 41% women; 71% White	Open-label study (adjudication committee
Acute SARS-CoV-2 infection	• 99% required HFNC oxygen, NIV, or MV; 15%	members were blinded to the study arms)
• ICU-level care for \leq 96 hours prior to randomization	required MV (41% required MV during the study)	• Stopped early because the decreasing number
Key Exclusion Criteria	 31% to 37% crossed over to an alternative study treatment during the study 	of ICU admissions for patients with COVID-19 made recruitment difficult.
 Ongoing or planned use of full-dose anticoagulation or dual antiplatelet therapy 	Primary Outcome	• There was an unequal crossover between the arms, with a greater crossover from the
High bleeding risk	Composite of VTE or ATE events: 12% in full-	prophylactic anticoagulation arm to the full-
History of HIT	dose anticoagulation arm vs. 6% in prophylactic anticoagulation arm (win ratio 1.95; 95% Cl, 1.08–	dose anticoagulation arm.
Ischemic stroke within 2 weeks	3.55; P = 0.028)	Interpretation
Interventions	Secondary Outcome	Among patients with COVID-19 who required ICIL level core patients who required full does
 Full-dose anticoagulation until Day 28 or hospital discharge, whichever came first (n = 197) 	Clinically evident VTE or ATE: 10% in full-dose anticoagulation arm vs. 6% in prophylactic	ICU-level care, patients who received full-dose anticoagulation had fewer VTE or ATE events but no survival benefit compared to those who
 Prophylactic anticoagulation (n = 193) 	anticoagulation arm (win ratio 1.79; 95% Cl, 0.92–	received prophylactic anticoagulation.
• Eligible patients were also randomized 1:1 to receive	3.47; <i>P</i> = 0.087)	The prevalence of moderate or severe
clopidogrel or no antiplatelet therapy (n = 292)	Safety Outcomes	bleeding events was higher among patients
Primary Endpoint	No fatal bleeding events in either arm	who received full-dose anticoagulation than among those who received prophylactic
 Composite of VTE or ATE events, including death due to VTE or ATE, PE, clinically evident DVT, MI, ischemic stroke, systemic embolic event or acute limb ischemia, 	• Life-threatening bleeding events: 4 (2.1%) in full- dose anticoagulation arm vs. 1 (0.5%) in prophylactic anticoagulation arm ($P = 0.19$)	anticoagulation.
or clinically silent DVT through hospital discharge or Day 28	• Moderate or severe bleeding events: 15 (7.9%) in full-dose anticoagulation arm vs. 1 (0.5%) in	
Key Secondary Endpoint	prophylactic anticoagulation arm ($P = 0.002$)	
 Individual outcomes listed above, with the exception of clinically silent DVT 		
Key Safety Endpoints		
 Fatal or life-threatening bleeding events 		
Moderate or severe bleeding events		

Methods	Results	Limitations and Interpretation	
REMAP-CAP/ACTIV-4a/ATTACC : Multiplatform, Open-Label RCT of Therapeutic Anticoagulation in Critically III, Hospitalized Patients With COVID-19 in 20 Countries ⁷			
 Countries⁷ Key Inclusion Criteria Hospitalized with severe COVID-19 and receiving HFNC oxygen, NIV, MV, ECMO, vasopressors, or inotropes Hospitalized <72 hours (ACTIV-4a, ATTACC) or <14 days (REMAP-CAP) Key Exclusion Criteria Hospital discharge expected in ≤72 hours Requirement for therapeutic anticoagulation or dual antiplatelet therapy High bleeding risk Interventions Therapeutic UFH or LMWH for 14 days or until discharge, whichever came first (n = 534) Usual care (n = 564) Primary Endpoint Number of organ support-free days by Day 21 Key Secondary Endpoints Survival to hospital discharge Any thrombosis Composite of major thrombotic events or death 	 Participant Characteristics Median age 60 years; 70% men; median BMI 30 24% with chronic respiratory disease; 33% with DM; 10% with chronic kidney disease; 8% with severe CVD 32% required HFNC oxygen; 38% required NIV; 29% required MV 18% on vasopressors; 82% on corticosteroids; 32% on RDV Primary Outcome Median number of organ support-free days by Day 21: 4 in therapeutic arm vs. 5 in usual care arm (aOR 0.83; 95% Crl, 0.67–1.03; 99.9% posterior probability of futility; OR < 1.2) Secondary Outcomes No difference between therapeutic and usual care arms in: Survival to hospital discharge: 63% vs. 65% (aOR 0.84; 95% Crl, 0.64–1.11) Thrombosis: 6% vs. 10% Major thrombotic events or death: 41% in both arms Major bleeding events: 4% vs. 2% (aOR 1.48; 95% Crl, 0.75–3.04) 	 Key Limitations Open-label study Anticoagulation dose varied in usual care arm (i.e., 51% intermediate, 2% subtherapeutic, 5% therapeutic). Inclusion criteria for hospital LOS and ICU-level care differed across trials. Trial stopped for futility. Interpretation In patients who required ICU-level care, therapeutic heparin did not reduce the duration of organ support or mortality. Although the differences were not significant, patients who received therapeutic anticoagulation had more bleeding events and fewer thrombotic events than patients who received usual care. 	

Methods	Results	Limitations and Interpretation
INSPIRATION: Open-Label RCT of Intermediate-Dose Versus Prophylactic-Dose Anticoagulation in Patients With COVID-19 in Intensive Care Units in Iran ⁸		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
Admitted to ICU	• Median age 62 years; 58% men; median BMI 27	Open-label study
 Hospitalized <7 days 	 44% with HTN; 28% with DM; 14% with CAD 	Not all patients received ICU-level care.
Key Exclusion Criteria	 32% required NIV; 20% required MV 	Interpretation
 Life expectancy <24 hours Indication for therapeutic anticoagulation 	 23% on vasopressors; 93% on corticosteroids; 60% on RDV 	 Intermediate-dose anticoagulation did not significantly reduce the occurrence of VTE and
Bleeding or high bleeding risk	Primary Outcome	ATE, the need for ECMO, or mortality.
 Interventions Intermediate-dose anticoagulation: enoxaparin 1 mg/kg once daily (n = 276) 	 Composite of adjudicated acute VTE, ATE, the need for ECMO, or all-cause mortality: 46% in therapeutic arm vs. 44% in prophylactic arm (OR 1.06; 95% CI, 0.76–1.48) 	• Although the difference was not significant, patients who received intermediate-dose anticoagulation had more bleeding events than patients who received usual care.
 Prophylactic-dose anticoagulation (n = 286) 	Secondary Outcomes	
 Primary Endpoint Composite of adjudicated acute VTE, ATE, the need for ECMO, or all-cause mortality within 30 days 	 No difference between therapeutic and prophylactic arms in: All-cause mortality: 43% vs. 41% VTE: 3% in both arms 	
Key Secondary Endpoints	 Major bleeding events and clinically relevant 	
 All-cause mortality VTE	nonmajor bleeding events: 6.3% vs. 3.1% (OR 2.02; 95% Cl, 0.89–4.61)	
Bleeding events		

Methods	Results	Limitations and Interpretation
ANTICOVID: Open-Label RCT of Therapeutic-Dose Versus Intermediate-Dose Versus Prophylactic-Dose Anticoagulation in Patients With COVID-19 in Intensive Care Units in France ⁹		
Key Inclusion Criterion	Participant Characteristics	Key Limitations
• Hospitalized for <72 hours with hypoxemic	Mean age 58 years; 67% men; median BMI 27–28	Open-label study
COVID-19 pneumonia	 31% with HTN; 18% with DM; 4% with CAD 	Not all patients received ICU-level care.
Key Exclusion Criteria	• 23% required conventional oxygen; 61% required	 Study excluded patients weighing >100 kg.
• Weight <40 kg or >100 kg	HFNC; 7% required NIV; 10% required MV	• Tinzaparin is not available in the United States.
 Indication or contraindication for therapeutic anticoagulation 	 92% on corticosteroids; 0.6% on RDV; 34% on tocilizumab; 3% on vasopressors 	Interpretation
Bleeding or high bleeding risk	Primary Outcome	The use of intermediate doses of anticoagulants improved the not divised outcome by reducing the
Interventions	No difference between arms for hierarchical outcome	improved the net clinical outcome by reducing the number of thrombosis events.
Therapeutic-dose anticoagulation: tinzaparin 175 IU/kg once daily or enoxaparin 100 IU/kg twice	of all-cause mortality or time to clinical improvement by Day 28	 There was no difference between the arms in the occurrence of major bleeding events.
daily $(n = 110)$	Secondary Outcomes	
 Intermediate-dose anticoagulation: tinzaparin 7,000 IU once daily or enoxaparin 4,000 IU twice daily (n = 110) 	 Net clinical outcome by Day 28: 20.0% in therapeutic- dose arm vs. 16.4% in intermediate-dose arm vs. 29.8% in prophylactic-dose arm 	
 Prophylactic-dose anticoagulation: tinzaparin 3,500 IU once daily or enoxaparin 4,000 IU once daily (n = 114) 	 Venous or arterial thrombosis: 5% in therapeutic-dose arm vs. 5% in intermediate-dose arm vs. 20% in prophylactic-dose arm 	
Primary Endpoint	• Major bleeding events: 4% in therapeutic-dose arm vs.	
 Hierarchical outcome of all-cause mortality or time to clinical improvement of 2 points on a WHO scale by Day 28 	4% in intermediate-dose arm vs. 3% in prophylactic- dose arm	
Key Secondary Endpoints		
• Net clinical outcome by Day 28, defined as a composite of venous or arterial thrombosis, major bleeding events (as defined by ISTH), or all-cause death		
Major bleeding events		

Methods	Results	Limitations and Interpretation	
OVID: Open-Label RCT of Low-Molecular-Weight Switzerland ¹⁰	OVID: Open-Label RCT of Low-Molecular-Weight Heparin for Thromboprophylaxis in Symptomatic, Nonhospitalized Patients With COVID-19 in Germany and Switzerland ¹⁰		
Key Inclusion Criteria	Participant Characteristics	Key Limitations	
 Aged ≥50 years 	Median age 57 years; 46% women; 96% White	Open-label study	
 Positive SARS-CoV-2 test result within past 5 days 	Median time from COVID-19 diagnosis to randomization: 3 days	 Study terminated early due to a low probability that enoxaparin would be superior to the standard 	
• Respiratory symptoms or temperature ≥37.5 °C	• 24% with HTN; 8% with DM; 5% with CVD	of care for the primary outcome.	
Key Exclusion Criteria	• 9.5% received at least 1 dose of a COVID-19 vaccine	Interpretation	
Severe renal or hepatic dysfunction	Primary Outcome	Thromboprophylaxis with enoxaparin did not	
Severe anemia or recent major bleeding	Composite of any untoward hospitalization or all-	reduce the risk of hospitalization or death among nonhospitalized, symptomatic patients with	
Dual antiplatelet therapy	cause death by Day 30: 8 (3%) in enoxaparin arm vs. 8		
Interventions	(3%) in SOC arm (adjusted relative risk 0.98; 95% Cl, $0.37-2.56$; $P = 0.96$)		
• Enoxaparin 40 mg SUBQ once daily for 14 days (n = 234)	Secondary Outcomes		
• SOC (n = 238)	• Composite of major arterial and venous cardiovascular events by Day 20: 2 (1%) in an event in arm verse 4 (2%)		
Primary Endpoint	events by Day 30: 2 (1%) in enoxaparin arm vs. 4 (2%) in SOC arm (relative risk 0.51; 95% Cl, 0.09–2.74)		
 Composite of any untoward hospitalization or all-cause death by Day 30 	No major or clinically relevant nonmajor bleeding events occurred		
Key Secondary Endpoint			
 Composite of major arterial and venous cardiovascular events by Day 30 			
Bleeding events			

Methods	Results	Limitations and Interpretation
ETHIC: Open-Label RCT of Low-Molecular-Weight Heparin for Thromboprophylaxis in Symptomatic Outpatients With COVID-19 in Belgium, Brazil, India, South Africa, Spain, and the United Kingdom ¹¹		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Aged ≥30 years 	 Median age 59 years; 56% men 	Open-label study
 RT-PCR-confirmed SARS-CoV-2 infection, with symptoms for ≤9 days 	 Median time from first symptom to randomization: 5 days 	 Study terminated early because of low event rate and lack of efficacy.
 ≥1 risk factor for severe disease 	Primary Outcomes	Interpretation
Key Exclusion Criteria	Composite of all-cause hospitalization or all-cause	This study demonstrated no benefit of prophylaxis
 Receipt of COVID-19 vaccine 	mortality by Day 21: 12 (11%) in enoxaparin arm vs. 12 (11%) in SOC arm (HR 1.09; 95% Cl, 0.49–2.43; <i>P</i> =	with LMWH in outpatients with COVID-19 who were at risk of progressing to severe disease.
• eGFR <30 mL/min	(11%) in SOC and $(117.09, 95%)$ Ci, $0.49-2.43$, $P = 0.83$	were at risk of progressing to severe disease.
 Anticoagulant or antiplatelet therapy, except low- dose aspirin 	 Patients who required hospitalization: 12 in enoxaparin arm vs. 12 in SOC arm 	
Interventions	 Hospitalized patients who required acute medical 	
 Enoxaparin 40 mg SUBQ once daily (for patients weighing <100 kg) or enoxaparin 40 mg SUBQ 	care or ICU admission: 4 in enoxaparin arm vs. 0 in SOC arm	
twice daily (for patients weighing ≥100 kg), self-	Secondary Outcomes	
administered for 21 days (n = 105) • SOC (n = 114)	• VTE by Day 90: 1 (1%) in enoxaparin arm vs. 2 (2%) in SOC arm	
Primary Endpoint	• Bleeding events by Day 50: 2 (2%) in enoxaparin arm	
 Composite of all-cause hospitalization or all- cause mortality by Day 21 	vs. 2 (2%) in SOC arm	
Key Secondary Endpoints		
• VTE by Day 90		
Bleeding events by Day 50		

Methods	Results	Limitations and Interpretation
ACTIV-4C: Double-Blind RCT of 30 Days of Apixaban After Hospital Discharge in Patients With COVID-19 in the United States ¹²		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Hospitalized >48 hours with confirmed SARS- CoV-2 infection within 2 weeks of admission 	 Median age 54 years; 50% men; 27% Black, 17% Hispanic 	 Trial was terminated early due to a low event rate and because the decreasing number of
• PLT $>50 \text{ x} 10^{9} \text{ cells/L}$ and Hgb $>8 \text{ g/dL}$	 15% were receiving antiplatelet therapy 	hospitalizations for people with COVID-19 made recruitment difficult.
Key Exclusion Criteria	• At discharge, 16% were prescribed antiplatelet therapy;	
 Indication for therapeutic or prophylactic 	93% received aspirin.	Interpretation
anticoagulation at discharge	Primary Outcome	 Incidence of death or thromboembolism was low in this schort of notionto.
 Ischemic stroke, intracranial bleed, or neurosurgery within 3 months 	 Composite of death, ATE, or VTE by Day 30: 13 (2.1%) in apixaban arm vs. 14 (2.3%) in placebo arm (relative 	in this cohort of patients.Because the trial was terminated early, the results
Bleeding within past 30 days	risk 0.92; 95% Cl, 0.44–1.95; <i>P</i> = 0.85)	were imprecise, and the study was inconclusive.
 Major surgery within 14 days 	Safety Outcomes	
 Inherited or active acquired bleeding disorder 	• Major bleeding events: 2 (0.4%) in apixaban arm vs.	
Interventions	1 (0.2%) in placebo arm (relative risk 2.00; 95% Cl, 0.18–22.03)	
 Apixaban 2.5 mg twice daily for 30 days, starting at hospital discharge (n = 610) 	 Clinically relevant nonmajor bleeding events: 3 (0.6%) in apixaban arm vs. 6 (1.1%) in placebo arm (relative 	
• Placebo (n = 607)	risk 0.50; 95% Cl, 0.13–1.99)	
Primary Endpoint		
• Composite of death, ATE, or VTE by Day 30		
Key Safety Endpoint		
Bleeding events		

Methods	Results	Limitations and Interpretation
MICHELLE: Open-Label RCT of Using Rivaroxaban in Brazil ¹³	After Hospital Discharge in Patients With COVID-19 Wh	o Were at High Risk of Venous Thromboembolism
Key Inclusion Criteria	Participant Characteristics	Key Limitations
• Hospitalized for \geq 3 days with confirmed SARS-	Median age 57 years; 60% men	Open-label study with no placebo
CoV-2 infection	• While hospitalized, 86% received thromboprophylaxis	• Not all patients had the protocol-specified CTPA
 Increased risk of VTE, defined as an IMPROVE VTE score at hospital discharge of >4 or 2–3 	with enoxaparin, 14% received unfractionated heparin, and 5% received antiplatelet therapy.	or Doppler ultrasound during the study. However, a higher number of imaging evaluations occurred
with D-dimer level >500 ng/mL		among the patients in the rivaroxaban arm.
·	Primary Outcomes	
Key Exclusion Criterion	• Primary composite outcome by Day 35: 5 (3%) in	Interpretation
Suspicion or confirmation of a thrombotic event	rivaroxaban arm vs. 15 (9%) in no anticoagulation arm	 In patients who were at high risk of VTE, the use of thrombonrophylavia with riverovehan 10 mg P
Interventions	(relative risk 0.33; 95% Cl, 0.12–0.90; $P = 0.03$)	of thromboprophylaxis with rivaroxaban 10 mg P once daily for 35 days improved clinical outcome
 Rivaroxaban 10 mg PO once daily for 35 days, starting at hospital discharge (n = 159) 	 Difference driven mainly by incidence of PE (2 in rivaroxaban arm vs. 10 in no anticoagulation arm) 	when compared with no anticoagulation.
• No anticoagulation $(n = 159)$	Secondary Outcomes	
Primary Endpoint	• Symptomatic or fatal VTE: 1 (0.6%) in rivaroxaban arm	
Composite of symptomatic or fatal VTE,	vs. 8 (5.0%) in no anticoagulation arm (relative risk 0.13; 95% Cl, 0.02–0.99; <i>P</i> = 0.049)	
asymptomatic VTE on bilateral lower-limb venous	 Composite of symptomatic VTE, MI, stroke, or 	
ultrasound and CTPA, symptomatic ATE, or	cardiovascular death: 1 (0.6%) in rivaroxaban arm vs.	
cardiovascular death by Day 35	9 (5.7%) in no anticoagulation arm (relative risk 0.11;	
Key Secondary Endpoints	95% Cl, 0.01–0.87; <i>P</i> = 0.036)	
 Symptomatic or fatal VTE 	Safety Outcome	
 Composite of symptomatic VTE, MI, non- hemorrhagic stroke, or cardiovascular death 	• No major bleeding events occurred in either arm.	
Key Safety Endpoint		
Bleeding events		

Key: ATE = arterial thromboembolism; BARC = Bleeding Academic Research Consortium; BMI = body mass index; CAD = coronary artery disease; CrCI = creatinine clearance; CTPA = computed tomography pulmonary angiogram; CVD = cardiovascular disease; DEX = dexamethasone; DM = diabetes mellitus; DVT = deep vein thrombosis; ECMO = extracorporeal membrane oxygenation; eGFR = estimated glomerular filtration rate; HFNC = high-flow nasal cannula; Hgb = hemoglobin; HIT = heparin-induced thrombocytopenia; HTN = hypertension; ICU = intensive care unit; IMPROVE = International Medical Prevention Registry on Venous Thromboembolism; ISTH = International Society of Thrombosis and Hemostasis; LMWH = low-molecular-weight heparin; LOS = length of stay; MI = myocardial infarction; MV = mechanical ventilation; NIV = noninvasive ventilation; OS = ordinal scale; the Panel = the COVID-19 Treatment Guidelines Panel; PE = pulmonary embolism; PLT = platelet count; PO = oral; RCT = randomized controlled trial; RDV = remdesivir; RT-PCR = reverse transcriptase polymerase chain reaction; SOC = standard of care; SpO₂ = oxygen saturation; SUBQ = subcutaneous; UFH = unfractionated heparin; ULN = upper limit of normal; VTE = venous thromboembolism; WHO = World Health Organization

References

- 1. ATTACC, ACTIV-4a, and REMAP-CAP Investigators. Therapeutic anticoagulation with heparin in noncritically ill patients with COVID-19. *N Engl J Med.* 2021;385(9):790-802. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34351721</u>.
- 2. Sholzberg M, Tang GH, Rahhal H, et al. Effectiveness of therapeutic heparin versus prophylactic heparin on death, mechanical ventilation, or intensive care unit admission in moderately ill patients with COVID-19 admitted to hospital: RAPID randomised clinical trial. *BMJ*. 2021;375:n2400. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34649864.
- 3. Spyropoulos AC, Goldin M, Giannis D, et al. Efficacy and safety of therapeutic-dose heparin vs standard prophylactic or intermediate-dose heparins for thromboprophylaxis in high-risk hospitalized patients with COVID-19: the HEP-COVID randomized clinical trial. *JAMA Intern Med.* 2021;181(12):1612-1620. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34617959.
- 4. Lopes RD, de Barros ESPGM, Furtado RHM, et al. Therapeutic versus prophylactic anticoagulation for patients admitted to hospital with COVID-19 and elevated D-dimer concentration (ACTION): an open-label, multicentre, randomised, controlled trial. *Lancet*. 2021;397(10291):2253-2263. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34097856.
- 5. Stone GW, Farkouh ME, Lala A, et al. Randomized trial of anticoagulation strategies for noncritically ill patients hospitalized with COVID-19. *J Am Coll Cardiol*. 2023;81(18):1747-1762. Available at: https://pubmed.ncbi.nlm.nih.gov/36889611.
- 6. Bohula EA, Berg DD, Lopes MS, et al. Anticoagulation and antiplatelet therapy for prevention of venous and arterial thrombotic events in critically ill patients with COVID-19: COVID-PACT. *Circulation*. 2022;146(18):1344-1356. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36036760.
- REMAP-CAP, ACTIV-4a, and ATTACC Investigators. Therapeutic anticoagulation with heparin in critically ill patients with COVID-19. *N Engl J Med.* 2021;385(9):777-789. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34351722</u>.
- 8. INSPIRATION Investigators. Effect of intermediate-dose vs standard-dose prophylactic anticoagulation on thrombotic events, extracorporeal membrane oxygenation treatment, or mortality among patients with COVID-19 admitted to the intensive care unit: the INSPIRATION randomized clinical trial. *JAMA*. 2021;325(16):1620-1630. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33734299.
- Labbé V, Contou D, Heming N, et al. Effects of standard-dose prophylactic, high-dose prophylactic, and therapeutic anticoagulation in patients with hypoxemic COVID-19 pneumonia: the ANTICOVID randomized clinical trial. *JAMA Intern Med.* 2023;183(6):520-531. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36946232/</u>.
- Barco S, Voci D, Held U, et al. Enoxaparin for primary thromboprophylaxis in symptomatic outpatients with COVID-19 (OVID): a randomised, open-label, parallel-group, multicentre, Phase 3 trial. *Lancet Haematol*. 2022;9(8):e585-e593. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35779558</u>.
- 11. Cools F, Virdone S, Sawhney J, et al. Thromboprophylactic low-molecular-weight heparin versus standard of care in unvaccinated, at-risk outpatients with COVID-19 (ETHIC): an open-label, multicentre, randomised, controlled, Phase 3b trial. *Lancet Haematol*. 2022;9(8):e594-e604. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35779560.
- 12. Wang TY, Wahed AS, Morris A, et al. Effect of thromboprophylaxis on clinical outcomes after COVID-19 hospitalization. *Ann Intern Med.* 2023;176(4):515-523. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36940444</u>.
- 13. Ramacciotti E, Barile Agati L, Calderaro D, et al. Rivaroxaban versus no anticoagulation for post-discharge thromboprophylaxis after hospitalisation for COVID-19 (MICHELLE): an open-label, multicentre, randomised, controlled trial. *Lancet*. 2022;399(10319):50-59. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34921756</u>.

Table 6b. Antiplatelet Therapy: Selected Clinical Trial Data

Last Updated: December 1, 2022

The clinical trials described in this table do not represent all the trials that the Panel reviewed while developing the recommendations for antiplatelet therapy. The studies summarized below are those that have had the greatest impact on the Panel's recommendations.

Results	Limitations and Interpretation	
ACTIV-4a: Open-Label, Adaptive RCT of Adding a P2Y12 Inhibitor to Anticoagulant Therapy in Noncritically III Hospitalized Patients With COVID-19 in Brazil, Italy, Spain, and the United States ¹		
Participant Characteristics	Key Limitations	
 Mean age 53 years; 42% women; 62% White 	Open-label study	
• HTN: 43% in P2Y12 inhibitor arm vs. 55% in usual care	 Study stopped early for futility 	
arm	Different P2Y12 inhibitors used	
	Median duration of P2Y12 inhibitor use was 6	
•	days, which may not be sufficient to observe	
	effects.	
	Interpretation	
-	 Among hospitalized patients with COVID-19 who ware not originally ill adding a DOV10 inhibitor to 	
0 11 5 5 5	were not critically ill, adding a P2Y12 inhibitor to a therapeutic dose of heparin did not increase the	
	number of organ support-free days.	
	Major bleeding events occurred infrequently	
inhibitor arm vs. 0.7% in usual care arm (aOR 3.31;	during the study. The number of patients who	
95% Cl, 0.64–17.2; <i>P</i> = 0.15)	experienced a major bleeding event was not significantly different between the arms.	
Secondary Outcome	Significantly uncreate between the arms.	
Major thrombotic event or death by Day 28: 6.1% in		
1.42, 95% 61, 0.04–5.15)		
	 P2Y12 Inhibitor to Anticoagulant Therapy in Noncriticall Participant Characteristics Mean age 53 years; 42% women; 62% White HTN: 43% in P2Y12 inhibitor arm vs. 55% in usual care arm 65% on glucocorticoids; 52% on RDV; 3% on IL-6 inhibitors; 14% on aspirin Median duration of P2Y12 inhibitor treatment: 6 days 63% received ticagrelor; 37% received clopidogrel Primary Outcomes Median number of organ support-free days by Day 21: 21 in both arms (aOR 0.83; 95% Crl, 0.55–1.25; posterior probability of futility 96%) Major bleeding events by Day 28: 2.0% in P2Y12 inhibitor arm vs. 0.7% in usual care arm (aOR 3.31; 95% Cl, 0.64–17.2; P = 0.15) 	

Methods	Results	Limitations and Interpretation
RECOVERY: Open-Label RCT of Aspirin in Hospitalized Patients With COVID-19 in Indonesia, Nepal, and the United Kingdom ²		
Key Inclusion Criterion	Participant Characteristics	Key Limitation
Clinically suspected or laboratory-confirmed SARS-CoV-2 infection	 Mean age 59 years; 62% men; 75% White 97% had laboratory-confirmed SARS-CoV-2 infection 	 Because of open-label design, reporting of thrombotic and major bleeding events may have influenced treatment allocation.
Key Exclusion Criteria	• At baseline:	
 Hypersensitivity to aspirin 	33% on NIV or MV	Interpretation
 Recent history of major bleeding events 	• 34% on intermediate- or therapeutic-dose LMWH	 In hospitalized patients with COVID-19, the use
• Currently receiving aspirin or another antiplatelet	60% on standard-dose LMWH	of aspirin was not associated with reductions in 28-day mortality or the risk of progressing to MV
treatment	 7% received no thromboprophylaxis 	or death.
Interventions	94% on corticosteroids; 26% on RDV; 13% on	
• Aspirin 150 mg once daily until discharge (n =	tocilizumab; 6% on baricitinib	
7,351)	Primary Outcome	
• SOC alone $(n = 7,541)$	• All-cause mortality at 28 days: 17% in both arms (rate	
Primary Endpoint	ratio 0.96; 95% Cl, 0.89–1.04; P = 0.35)	
All-cause mortality at 28 days	Secondary Outcomes	
Key Secondary Endpoints	• Progression to MV or death at 28 days: 21% in aspirin	
Progression to MV or death at 28 days Major blooding or thrombatic events at 28 days	arm vs. 22% in SOC arm (risk ratio 0.96; 95% Cl, 0.90–1.03)	
Major bleeding or thrombotic events at 28 days	• Major bleeding events at 28 days: 1.6% in aspirin arm vs. 1.0% in SOC arm (<i>P</i> = 0.0028)	
	• Thrombotic events at 28 days: 4.6% in aspirin arm vs. 5.3% in SOC arm ($P = 0.07$)	

Results	Limitations and Interpretation	
REMAP-CAP: Open-Label, Adaptive RCT of Antiplatelet Therapy in Critically III Patients With COVID-19 in 8 Countries in Europe and Asia ³		
Participant Characteristics	Key Limitations	
Mean age 57 years; 34% women; 77% White	Open-label study	
• At baseline, 98% on LMWH:	Different P2Y12 inhibitors used	
19% on low-dose LMWH	• Trial stopped for futility. Because equivalence for	
 59% on intermediate-dose LMWH 	aspirin and P2Y12 inhibitor arms was reached,	
 12% on therapeutic-dose LMWH 	these arms were pooled for analyses.	
• 98% on steroids; 21% on RDV; 44% on tocilizumab;	Interpretation	
11% on sarilumab	• In critically ill patients with COVID-19, the use of	
• In P2Y12 inhibitor arm, 88.5% received clopidogrel,	aspirin or a P2Y12 inhibitor did not reduce the number of organ support-free days or in-hospital	
	mortality.	
	Patients in the pooled antiplatelet arm had more	
	major bleeding events than those in the control	
	arm, but they had improved survival over 90 days.	
21: 7 in pooled antiplatelet arm and control arm (aOR		
absolute difference 5%; 95% Crl, -0.2% to 9.5%; 97%		
posterior probability of efficacy)		
• Survival to Day 90: 72% in pooled antiplatelet arm vs.		
efficacy)		
• Major bleeding event by Day 14: 2.1% in pooled		
99.4%)		
	 atelet Therapy in Critically III Patients With COVID-19 in 8 Participant Characteristics Mean age 57 years; 34% women; 77% White At baseline, 98% on LMWH: 19% on low-dose LMWH 59% on intermediate-dose LMWH 12% on therapeutic-dose LMWH 98% on steroids; 21% on RDV; 44% on tocilizumab; 11% on sarilumab In P2Y12 inhibitor arm, 88.5% received clopidogrel, 1.3% received ticagrelor, 1.3% received prasugrel, and 8.8% received an unknown P2Y12 inhibitor Primary Outcome Data from aspirin and P2Y12 inhibitor arms were pooled and reported as "pooled antiplatelet arm" in final analysis: Median number of organ support-free days by Day 21: 7 in pooled antiplatelet arm and control arm (aOR 1.02; 95% Crl, 0.86–1.23; posterior probability of futility 96%) Secondary Outcomes Survival to hospital discharge: 71.5% in pooled antiplatelet arm vs. 67.9% in control arm (median-adjusted OR 1.27; 95% Crl, 0.99–1.62; adjusted absolute difference 5%; 95% Crl, -0.2% to 9.5%; 97% posterior probability of efficacy) Survival to Day 90: 72% in pooled antiplatelet arm vs. 68% in control arm (HR with pooled antiplatelets 1.22; 95% Crl, 1.06–1.40; 99.7% posterior probability of efficacy) Major bleeding event by Day 14: 2.1% in pooled antiplatelet arm vs. 0.4% in control arm (aOR 2.97; 95% Crl, 1.23–8.28; posterior probability of harm 	

Methods	Results	Limitations and Interpretation
COVID-PACT: Open-Label RCT of Clopidogrel in Adults With COVID-19 Who Were Receiving ICU-Level Care in the United States ⁴		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Aged ≥18 years 	Median age 58 years; 41% women; 71% White	Open-label study (adjudication committee
Acute SARS-CoV-2 infection	• At baseline, 99% required HFNC, NIV, or MV; 15%	members were blinded to the study arms)
 Required ICU-level care for ≤96 hours prior to randomization 	required MV (37% required MV during the study)	 Stopped early because decreasing number of ICU admissions for patients with COVID-19
	Primary Outcome	made recruitment difficult
Key Exclusion Criteria	 Composite of VTE or ATEs by hospital discharge or Day 28: 10% in both arms (win ratio 1.04; 95% Cl, 0.54– 	 31% discontinued clopidogrel
 Ongoing or planned use of a therapeutic dose of anticoagulation or dual antiplatelet therapy 	2.01; P = 0.90)	Interpretation
High risk of bleeding	Secondary Outcome	In patients with COVID-19 who required
History of HIT	• Composite of clinically evident VTE or ATEs by hospital	ICU-level care, clopidogrel did not reduce the incidence of thrombotic complications.
Ischemic stroke within 2 weeks	discharge or Day 28: 7% in clopidogrel arm vs. 9% in no clopidogrel arm (win ratio 0.79; 95% Cl, 0.38–1.65;	incluence of unormbolic complications.
Interventions	P = 0.53)	
 Clopidogrel 300 mg at randomization, then 	Safety Outcomes	
clopidogrel 75 mg once daily until hospital discharge or Day 28, whichever came first ($n = 152$)	• Fatal or life-threatening bleeding: 1.3% in clopidogrel	
 No clopidogrel therapy (n = 140) 	arm vs. 1.4% in no clopidogrel arm ($P = 1.00$)	
 Some patients also randomized to receive a therapeutic or prophylactic dose of anticoagulation (n = 290) 	• Moderate or severe bleeding: 4.0% in clopidogrel arm vs. 6.4% in no clopidogrel arm ($P = 0.83$)	
Primary Endpoint		
• Composite of VTE or ATEs by hospital discharge or Day 28. Events included death due to VTE or ATEs, PE, clinically evident DVT, MI, ischemic stroke, systemic embolic events or acute limb ischemia, and clinically silent DVT.		
Secondary Endpoint		
 Composite of clinically evident VTE or ATEs by hospital discharge or Day 28 		
Safety Endpoints		
 Fatal or life-threatening bleeding 		
Moderate or severe bleeding		

Key: ATE = arterial thrombotic event; BMI = body mass index; CrCI = creatinine clearance; CVD = cardiovascular disease; DM = diabetes mellitus; DVT = deep vein thrombosis; ECMO = extracorporeal membrane oxygenation; eGFR = estimated glomerular filtration rate; HFNC = high-flow nasal cannula; HIT = heparin-induced thrombocytopenia; HTN = hypertension; ICU = intensive care unit; IL = interleukin; LMWH = low-molecular-weight heparin; MI = myocardial infarction; MV = mechanical ventilation; NIV = noninvasive ventilation; NSAID = nonsteroidal anti-inflammatory drug; the Panel = the COVID-19 Treatment Guidelines Panel; PE = pulmonary embolism; RCT = randomized controlled trial; RDV = remdesivir; SOC = standard of care; ULN = upper limit of normal; VTE = venous thromboembolism

References

- 1. Berger JS, Kornblith LZ, Gong MN, et al. Effect of P2Y12 inhibitors on survival free of organ support among non-critically ill hospitalized patients with COVID-19: a randomized clinical trial. *JAMA*. 2022;327(3):227-236. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35040887.
- 2. RECOVERY Collaborative Group. Aspirin in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2022;399(10320):143-151. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34800427.
- 3. REMAP-CAP Writing Committee for the REMAP-CAP Investigators, Bradbury CA, Lawler PR, et al. Effect of antiplatelet therapy on survival and organ support-free days in critically ill patients with COVID-19: a randomized clinical trial. *JAMA*. 2022;327(13):1247-1259. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35315874.
- 4. Bohula EA, Berg DD, Lopes MS, et al. Anticoagulation and antiplatelet therapy for prevention of venous and arterial thrombotic events in critically ill patients with COVID-19: COVID-PACT. *Circulation*. 2022;146(18):1344-1356. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36036760.

Miscellaneous Drugs

Last Updated: December 20, 2023

Summary Recommendations

Fluvoxamine

- The COVID-19 Treatment Guidelines Panel (the Panel) recommends against the use of fluvoxamine for the treatment of COVID-19 in nonhospitalized patients (Alla).
- There is insufficient evidence for the Panel to recommend either for or against the use of the combination of inhaled budesonide plus fluvoxamine for the treatment of COVID-19 in nonhospitalized patients.
- Patients with COVID-19 who are receiving **fluvoxamine** for an underlying condition should continue this therapy as directed by their health care provider (**AIII**).

Intravenous Immunoglobulin

- The Panel **recommends against** the use of **intravenous immunoglobulin** (IVIG) for the treatment of acute COVID-19 in adults and children, except in a clinical trial **(AIII)**. This recommendation should not preclude the use of IVIG when it is otherwise indicated for the treatment of underlying conditions or complications that arise during the course of COVID-19.
- For the Panel's recommendations on the use of IVIG in people with multisystem inflammatory syndrome in children (MIS-C) or multisystem inflammatory syndrome in adults (MIS-A) and a discussion of the clinical data that support those recommendations, see <u>Therapeutic Management of Hospitalized Children With MIS-C</u>, <u>Plus a Discussion on MIS-A</u>.

Ivermectin

• The Panel recommends against the use of ivermectin for the treatment of COVID-19 (Alla).

Metformin

- There is insufficient evidence for the Panel to recommend either for or against the use of metformin for the treatment of COVID-19 in nonhospitalized patients.
- The Panel **recommends against** the use of **metformin** for the treatment of COVID-19 in hospitalized patients, except in a clinical trial **(BIII)**.
- Patients with COVID-19 who are receiving **metformin** for an underlying condition should continue this therapy as directed by their health care provider (AIII).

The Panel reviewed clinical trials that evaluated the use of the anti-inflammatory drug colchicine for the treatment of COVID-19; however, these trials failed to show a benefit of using colchicine in patients with COVID-19.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Fluvoxamine

Last Updated: December 20, 2023

Fluvoxamine is a selective serotonin reuptake inhibitor (SSRI) that is approved by the Food and Drug Administration (FDA) for the treatment of obsessive-compulsive disorder and is used for other conditions, including depression. Fluvoxamine is not approved by the FDA for the treatment of any infection.

In a murine sepsis model, fluvoxamine was found to bind to the sigma-1 receptor on immune cells, resulting in reduced production of inflammatory cytokines.¹ In an in vitro study of human endothelial cells and macrophages, fluvoxamine reduced the expression of inflammatory genes.²

Recommendations

- The COVID-19 Treatment Guidelines Panel (the Panel) **recommends against** the use of **fluvoxamine** for the treatment of COVID-19 in nonhospitalized patients (**AIIa**).
- There is insufficient evidence for the Panel to recommend either for or against the use of the combination of inhaled budesonide plus fluvoxamine for the treatment of COVID-19 in nonhospitalized patients.
- Patients with COVID-19 who are receiving **fluvoxamine** for an underlying condition should continue this therapy as directed by their health care provider (**AIII**).

Rationale

Six randomized trials have studied the use of fluvoxamine for the treatment of nonhospitalized patients with COVID-19.³⁻⁸ The TOGETHER and STOP COVID 2 trials enrolled unvaccinated patients with COVID-19 who had at least 1 risk factor for disease progression.^{3,5} These studies did not identify a consistent benefit of using fluvoxamine in these patients, although STOP COVID 2 was stopped early due to low primary outcome rates. Other outpatient therapies (i.e., ritonavir-boosted nirmatrelvir [Paxlovid], remdesivir) have been shown to be effective in preventing hospitalization or death in unvaccinated patients who are at high risk of disease progression. In subsequent trials where the majority of enrolled patients were vaccinated against COVID-19, fluvoxamine did not significantly reduce the risk of hospitalization or death, the time to recovery, or health care utilization.⁶⁻⁸ In several of these studies, fluvoxamine was associated with decreased adherence and/or an increase in the occurrence of nonserious adverse effects, primarily gastrointestinal symptoms.^{3,5,6}

The TOGETHER trial was a large, double-blind, placebo-controlled, adaptive randomized trial in Brazil that evaluated the use of inhaled budesonide plus oral fluvoxamine in patients with COVID-19.⁹ Over 90% of the patients had received at least 2 doses of a COVID-19 vaccine. Treatment with this combination significantly reduced the incidence of the primary outcome, which was a composite of hospitalization or retention in an emergency setting for >6 hours. The proportion of patients who were hospitalized was the same in the treatment and placebo arms (0.9% vs. 1.1%), and the treatment did not significantly impact secondary outcomes such as health care attendance or the need for an emergency setting visit. It is unclear how the >6-hour emergency setting outcome translates to other settings. In addition, treatment with budesonide plus fluvoxamine was associated with significantly more adverse events.

Summaries of the studies that informed the Panel's recommendations can be found in Table 7a.

Monitoring, Adverse Effects, and Drug-Drug Interactions

When fluvoxamine is used to treat psychiatric conditions, the most common adverse effect is nausea, but adverse effects can include other gastrointestinal effects (e.g., diarrhea, indigestion), neurologic effects (e.g., asthenia, insomnia, somnolence, anxiety, headache), and, rarely, suicidal ideation.

Fluvoxamine is a cytochrome P450 (CYP) 2D6 substrate, a potent inhibitor of CYP1A2 and CYP2C19, and a moderate inhibitor of CYP2C9, CYP2D6, and CYP3A4.¹⁰ Fluvoxamine can enhance the serotonergic effects of other SSRIs or monoamine oxidase inhibitors, resulting in serotonin syndrome; therefore, it should not be used within 2 weeks of receiving other SSRIs or monoamine oxidase inhibitors. Fluvoxamine may enhance the anticoagulant effects of antiplatelets and anticoagulants. Patients who are receiving these drugs should be closely monitored.

Considerations in Pregnant People

Fluvoxamine is not thought to increase the risk of congenital abnormalities; however, the data on its use in pregnant individuals are limited.^{11,12} An association between SSRI use in the late third trimester and a small increase in the risk of primary persistent pulmonary hypertension in newborns has not been excluded, although the absolute risk is likely low.¹³

Considerations in Children

Fluvoxamine is approved by the FDA for the treatment of obsessive-compulsive disorder in children aged \geq 8 years.¹⁴ The adverse effects of SSRI use seen in children are similar to those seen in adults, although children and adolescents appear to have higher rates of activation and vomiting than adults.¹⁵ There are no data on the use of fluvoxamine for the prevention or treatment of COVID-19 in children.

References

- 1. Rosen DA, Seki SM, Fernández-Castañeda A, et al. Modulation of the sigma-1 receptor-IRE1 pathway is beneficial in preclinical models of inflammation and sepsis. *Sci Transl Med*. 2019;11(478):eaau5266. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30728287</u>.
- Rafiee L, Hajhashemi V, Javanmard SH. Fluvoxamine inhibits some inflammatory genes expression in LPS/ stimulated human endothelial cells, U937 macrophages, and carrageenan-induced paw edema in rat. *Iran J Basic Med Sci.* 2016;19(9):977-984. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27803785</u>.
- 3. Reis G, Dos Santos Moreira-Silva EA, Medeiros Silva DC, et al. Effect of early treatment with fluvoxamine on risk of emergency care and hospitalisation among patients with COVID-19: the TOGETHER randomised, platform clinical trial. *Lancet Glob Health*. 2022;10(1):e42-e51. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34717820.
- 4. Lenze EJ, Mattar C, Zorumski CF, et al. Fluvoxamine vs placebo and clinical deterioration in outpatients with symptomatic COVID-19: a randomized clinical trial. *JAMA*. 2020;324(22):2292-2300. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33180097.
- Reiersen AM, Mattar C, Bender Ignacio RA, et al. The STOP COVID 2 study: fluvoxamine vs placebo for outpatients with symptomatic COVID-19, a fully remote randomized controlled trial. *Open Forum Infect Dis*. 2023;10(8):ofad419. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37622035</u>.
- 6. Stewart TG, Rebolledo PA, Mourad A, et al. Higher-dose fluvoxamine and time to sustained recovery in outpatients with COVID-19: the ACTIV-6 randomized clinical trial. *JAMA*. 2023;Published online ahead of print. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37976072</u>.
- 7. McCarthy MW, Naggie S, Boulware DR, et al. Effect of fluvoxamine vs placebo on time to sustained recovery in outpatients with mild to moderate COVID-19: a randomized clinical trial. *JAMA*. 2023;329(4):296-305. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36633838</u>.

- Bramante CT, Huling JD, Tignanelli CJ, et al. Randomized trial of metformin, ivermectin, and fluvoxamine for COVID-19. *N Engl J Med*. 2022;387(7):599-610. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36070710.
- 9. Reis G, Dos Santos Moreira Silva EA, Medeiros Silva DC, et al. Oral fluvoxamine with inhaled budesonide for treatment of early-onset COVID-19: a randomized platform trial. *Ann Intern Med.* 2023;176(5):667-675. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37068273</u>.
- Hemeryck A, Belpaire FM. Selective serotonin reuptake inhibitors and cytochrome P-450 mediated drug-drug interactions: an update. *Curr Drug Metab*. 2002;3(1):13-37. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/11876575</u>.
- Einarson A, Choi J, Einarson TR, Koren G. Incidence of major malformations in infants following antidepressant exposure in pregnancy: results of a large prospective cohort study. *Can J Psychiatry*. 2009;54(4):242-246. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/19321030</u>.
- Furu K, Kieler H, Haglund B, et al. Selective serotonin reuptake inhibitors and venlafaxine in early pregnancy and risk of birth defects: population based cohort study and sibling design. *BMJ*. 2015;350:h1798. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/25888213</u>.
- Huybrechts KF, Bateman BT, Palmsten K, et al. Antidepressant use late in pregnancy and risk of persistent pulmonary hypertension of the newborn. *JAMA*. 2015;313(21):2142-2151. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26034955</u>.
- 14. Fluvoxamine maleate tablets [package insert]. Food and Drug Administration. 2019. Available at: https://www.accessdata.fda.gov/drugsatfda_docs/label/2019/021519s012lbl.pdf.
- 15. Safer DJ, Zito JM. Treatment-emergent adverse events from selective serotonin reuptake inhibitors by age group: children versus adolescents. *J Child Adolesc Psychopharmacol*. 2006;16(1-2):159-169. Available at: https://www.ncbi.nlm.nih.gov/pubmed/16553536.

Table 7a. Fluvoxamine: Selected Clinical Trial Data

Last Updated: December 20, 2023

Methods	Results	Limitations and Interpretation	
ACTIV-6: Decentralized, Randomized, Placebo-Controlled,	ACTIV-6: Decentralized, Randomized, Placebo-Controlled, Platform Trial of Low-Dose Fluvoxamine in Patients With Mild to Moderate COVID-19 ¹		
Key Inclusion Criteria	Participant Characteristics	Key Limitation	
 Aged ≥30 years 	Mean age 47 years; 57% women; 81% White	• Low number of some clinical events, such as	
Positive SARS-CoV-2 nasopharyngeal RT-PCR result or	• 36% with BMI ≥30; 24% with HTN	hospitalization	
antigen test result	• 67% received \geq 2 doses of a SARS-CoV-2 vaccine.	Interpretation	
• \geq 2 COVID-19 symptoms for \leq 7 days	• Median of 5 days from symptom onset to receipt of	 In outpatients with mild to moderate 	
Key Exclusion Criterion	study drug	COVID-19, fluvoxamine 50 mg twice daily for	
 Receipt of fluvoxamine in past 14 days 	Primary Outcome	10 days did not reduce the time to recovery or the incidence of clinical events such as	
Interventions	Median time to recovery: 12 days in fluvoxamine	hospitalization, urgent care visits, or ED	
• Fluvoxamine 50 mg P0 twice daily for 10 days (n = 674)	arm vs. 13 days in placebo arm (HR 0.96; 95% Crl, 0.86–1.06)	visits.	
 Placebo (n = 614; 326 received matching placebo, 288 received placebo from another study arm) 	Secondary Outcomes		
Primary Endpoint	Hospitalization or death by Day 28: 0.2% in fluvoxamine arm vs. 0.3% in placebo arm (3 events)		
• Time to recovery, defined as time to third day of 3	total)		
consecutive days without symptoms	• Urgent care visit, ED visit, or hospitalization by Day		
Key Secondary Endpoints	28: 3.9% in fluvoxamine arm vs. 3.8% in placebo		
Hospitalization or death by Day 28	arm (HR 1.1; 95% Crl, 0.5–1.8)		
Urgent care visit, ED visit, or hospitalization by Day 28			

Methods	Results	Limitations and Interpretation
ACTIV-6: Decentralized, Randomized, Placebo-Controlled, Platform Trial of High-Dose Fluvoxamine in Patients With Mild to Moderate COVID-19 ²		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Aged ≥30 years 	• Median age 50 years; 66% women; 73% White	• Low number of some clinical events, such as
 Positive SARS-CoV-2 nasopharyngeal RT-PCR result or 	• 36% with BMI ≥30; 26% with HTN	hospitalization
antigen test result	• 77% received \geq 2 doses of a SARS-CoV-2 vaccine.	Interpretation
• \geq 2 COVID-19 symptoms for \leq 7 days	• Median of 5 days from symptom onset to receipt of	In outpatients with mild to moderate
Key Exclusion Criterion	study drug	COVID-19, fluvoxamine 100 mg twice daily
Receipt of fluvoxamine or other selective serotonin or	Primary Outcome	did not reduce the time to symptom recovery or the incidence of clinical events such as
norepinephrine reuptake inhibitors in past 14 days	Median time to recovery: 10 days in fluvoxamine	hospitalization, urgent care visits, or ED
Interventions	arm vs. 10 days in placebo arm (HR 0.99; 95% Crl,	visits.
 Fluvoxamine 50 mg PO twice daily for 1 day, then 	0.89–1.09)	
fluvoxamine 100 mg PO twice daily for 12 days (n = 589)	Secondary Outcomes	
• Placebo (n = 586)	Hospitalization or death by Day 28: 0.2% in	
Primary Endpoint	fluvoxamine arm vs. 0.3% in placebo arm (3 events total)	
 Time to recovery, defined as time to third day of 3 consecutive days without symptoms 	 Urgent care visit, ED visit, or hospitalization by Day 28: 2.4% in fluvoxamine arm vs. 3.6% in placebo 	
Key Secondary Endpoints	arm (HR 0.69; 95% Crl, 0.27–1.21)	
Hospitalization or death by Day 28		
• Urgent care visit, ED visit, or hospitalization by Day 28		

Methods	Results	Limitations and Interpretation
COVID-OUT: Randomized Trial of Metformin, Ivermectin, and Fluvoxamine in Patients With COVID-19 ³		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Aged 30–85 years BMI ≥25 or ≥23 if Asian or Latinx 	Median age 43–46 years; 54% women; 82% White	 In this trial, the study arms that did not include metformin were underpowered to detect differences in the primary endpoint.
Laboratory-confirmed SARS-CoV-2 infection within 3 days of randomization	 27% with CVD; 47% with BMI ≥30 56% received primary vaccination series. 	Interpretation
• <7 days of symptoms	Mean of 5 days from symptom onset to randomization	 Fluvoxamine did not impact the incidence of COVID-19–related complications such as
 Key Exclusion Criteria Immunocompromised Hepatic impairment, severe kidney disease Interventions Fluvoxamine 50 mg PO twice daily for 14 days (n = 334) Control (n = 327) Primary Endpoint Composite of hypoxemia (as measured by a home pulse oximeter), ED visit, hospitalization, or death by Day 14 Key Secondary Endpoints Individual components of the composite endpoint Total symptom severity score Drug interruption or discontinuation 	 Primary Outcome Composite of hypoxemia, ED visit, hospitalization, or death by Day 14: 24% in fluvoxamine arm vs. 25% in control arm (aOR 0.94; 95% Cl, 0.66–1.36) Secondary Outcomes Hospitalization by Day 14: 1.8% in fluvoxamine arm vs. 1.5% in control arm (aOR 1.11; 95% Cl, 0.33–3.76). Composite of ED visit, hospitalization, or death: 5.5% in fluvoxamine arm vs. 4.6% in control arm (aOR 1.17; 95% Cl, 0.57–2.40) No deaths occurred in either arm. No difference between arms in total symptom severity score over 14 days Drug interruption or discontinuation: 30% in those who only received fluvoxamine vs. 25% 	 Fluvoxamine did not impact symptom severity.

Methods	Results	Limitations and Interpretation	
TOGETHER: Double-Blind, Adaptive RCT of Fluvoxamine in Nonhospitalized Patients With COVID-19 in Brazil ⁴			
Key Inclusion Criteria	Participant Characteristics	Key Limitations	
 Aged ≥50 years or aged ≥18 years with comorbidities Laboratory-confirmed SARS-CoV-2 infection ≤7 days of symptoms 	 Median age 50 years; 58% women; 95% self-identified as mixed race 13% with uncontrolled HTN; 13% with type 2 DM; 50% with BMI ≥30 	 The >6-hour ED observation endpoint has not been used in other studies of interventions for nonhospitalized patients who are at high risk of hospitalization and death. 	
Key Exclusion CriteriaUse of an SSRI	Mean of 3.8 days from symptom onset to randomization	 Hospitalization or ED observation for >24 hours was analyzed in a post hoc analysis. 	
 Severe mental illness Cirrhosis, recent seizures, severe ventricular cardiac arrhythmia 	 Primary Outcome Composite of ED observation >6 hours or hospitalization by Day 28: 11% in fluvoxamine arm vs. 16% in placebo arm (relative risk 0.68; 	• As this was an adaptive platform trial where multiple investigational treatments or placebos were being evaluated simultaneously, not all patients in the placebo arm received a placebo	
• Fluvoxamine 100 mg PO twice daily for 10 days ($n = 741$)	95% Crl, 0.52–0.88) Secondary Outcomes	that was matched to fluvoxamine by route of administration, dosing frequency, or duration of therapy.	
 Placebo (n = 756; route, dosing frequency, and duration of placebo may have differed from fluvoxamine for some patients) Primary Endpoint 	 87% of clinical events were hospitalizations. No difference between arms in COVID-19– related hospitalizations: 10% in fluvoxamine arm vs. 13% in placebo arm (OR 0.77; 95% Cl, 	 PP analyses are not randomized comparisons, and they introduce bias when adherence is associated with factors that influence the outcome. Adherence was self-reported and not verified. 	
 Composite of ED observation >6 hours or hospitalization for COVID-19 by Day 28 	0.55–1.05)	Interpretation	
 Key Secondary Endpoints COVID-19–related hospitalization by Day 28 Composite of hospitalization or ED observation >24 hours 	 Lower risk of hospitalization or ED observation >24 hours in fluvoxamine arm than in placebo arm (relative risk 0.74; 95% Cl, 0.56–0.98)⁵ No difference between arms in time to 	 Fluvoxamine reduced the proportion of patients who met the composite endpoint of COVID-19– related hospitalization or retention in an ED for >6 hours. 	
 Time to symptom resolution Adherence to study drugs, defined as receiving >80% of possible doses Mathematical bath the primery ITT population and a PD. 	 symptom resolution Adherence: 74% in fluvoxamine arm vs. 82% in placebo arm (OR 0.62; 95% Cl, 0.48–0.81) 11% in fluvoxamine arm vs. 8% in 	 The use of fluvoxamine did not impact the incidence of COVID-19-related hospitalizations but did reduce the need for hospitalization or ED observations >24 hours. 	
 Mortality in both the primary ITT population and a PP population that included patients who took >80% of the study medication doses 	 Mortality (ITT): 2% in fluvoxamine arm vs. 3% in placebo arm (OR 0.68; 95% Cl, 0.36–1.27) 	 It is difficult to define the clinical relevance of the >6-hour ED observation endpoint and apply it to practice settings in different countries. Fluvoxamine did not have a consistent impact on 	
	 Mortality (PP): <1% in fluvoxamine arm vs. 2% in placebo arm (OR 0.09; 95% Cl, 0.01–0.47) 	 Fluvoxamine did not impact the time to symptom resolution. 	

Methods	Results	Limitations and Interpretation
STOP COVID 2: Fully Remote RCT of Fluvoxamine Versus Placebo in Outpatients With Symptomatic COVID-19 ⁶		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Aged ≥30 years 	• Median age 47 years; 62% women; 27%	Small sample size compared to other trials
Positive SARS-CoV-2 PCR result per patient self-report	non-White	Short follow-up period
 ≤7 days of symptoms 	• 44% with obesity; 21% with HTN	Interpretation
• ≥1 risk factor for clinical deterioration	Median of 5 days from symptom onset to randomization	• Fluvoxamine did not reduce the proportion of patients
Key Exclusion Criteria		who experienced clinical deterioration by Day 15.
Unstable medical comorbidities	Primary Outcome	
Significant interacting medications	Clinical deterioration: 4.8% in fluvoxamine arm vs. 5.5% in placebo arm	
Interventions	(absolute difference 0.68%; 95% Cl, -3.0	
• Fluvoxamine 50 mg P0 for 1 dose, then fluvoxamine 100	to 4.4)	
mg PO twice daily through Day 15 (n = 272)	Secondary Outcomes	
• Placebo (n = 275)	• GI AEs were significantly more common	
Primary Endpoint	in fluvoxamine arm	
Clinical deterioration within 15 days of randomization. Clinical deterioration was defined as:		
 Having dyspnea or being hospitalized for dyspnea or pneumonia; and 		
 Having Sp0₂ <92% on room air or requiring supplemental oxygen to attain Sp0₂ ≥92% 		
Key Secondary Endpoints		
Occurrence of AEs		

 Aged ≥18 years Positive SARS-CoV-2 PCR result ≤7 days of symptoms Key Exclusion Criteria Immunocompromised Unstable medical comorbidities Interventions Fluvoxamine 50 mg PO for 1 dose, then fluvoxamine 100 mg PO twice daily, then fluvoxamine 100 mg PO 3 times daily through Day 15 (n = 80) Placebo (n = 72) Mean age Black 56% with with astheted astronomy of the second and the second astronomy of the second as	Characteristics 46 years; 72% women; 25% obesity; 20% with HTN; 17% na 4 days from symptom onset to	 ted States⁷ Key Limitations Small sample size Short follow-up period Ascertaining clinical deterioration was challenging because all assessments were done remotely. 24% of patients stopped responding to follow-up prior to Day 15 but were included in the final analysis.
 Aged ≥18 years Positive SARS-CoV-2 PCR result ≤7 days of symptoms Key Exclusion Criteria Immunocompromised Unstable medical comorbidities Interventions Fluvoxamine 50 mg PO for 1 dose, then fluvoxamine 100 mg PO twice daily, then fluvoxamine 100 mg PO 3 times daily through Day 15 (n = 80) Placebo (n = 72) Primary Endpoint Clinical deterioration within 15 days of randomization. Mean age Black 56% with with asthetic the structure of the structure	46 years; 72% women; 25% obesity; 20% with HTN; 17% na 4 days from symptom onset to	 Small sample size Short follow-up period Ascertaining clinical deterioration was challenging because all assessments were done remotely. 24% of patients stopped responding to follow-up prior
 Positive SARS-CoV-2 PCR result ≤7 days of symptoms Key Exclusion Criteria Immunocompromised Unstable medical comorbidities Interventions Fluvoxamine 50 mg PO for 1 dose, then fluvoxamine 100 mg PO twice daily, then fluvoxamine 100 mg PO 3 times daily through Day 15 (n = 80) Placebo (n = 72) Primary Endpoint Clinical deterioration within 15 days of randomization. 	obesity; 20% with HTN; 17% na 4 days from symptom onset to	 Short follow-up period Ascertaining clinical deterioration was challenging because all assessments were done remotely. 24% of patients stopped responding to follow-up prior
 Fostive SARS-COV-2 PCR result ≤7 days of symptoms Key Exclusion Criteria Immunocompromised Unstable medical comorbidities Interventions Fluvoxamine 50 mg PO for 1 dose, then fluvoxamine 100 mg PO twice daily, then fluvoxamine 100 mg PO 3 times daily through Day 15 (n = 80) Placebo (n = 72) Primary Endpoint Clinical deterioration within 15 days of randomization. Secondary No patien and 4 pat hospitalize 	na 4 days from symptom onset to	 Ascertaining clinical deterioration was challenging because all assessments were done remotely. 24% of patients stopped responding to follow-up prior
 With asthetic structure Median of randomization 	na 4 days from symptom onset to	because all assessments were done remotely.24% of patients stopped responding to follow-up prior
 Immunocompromised Instable medical comorbidities Interventions Fluvoxamine 50 mg P0 for 1 dose, then fluvoxamine 100 mg P0 twice daily, then fluvoxamine 100 mg P0 3 times daily through Day 15 (n = 80) Placebo (n = 72) Primary Endpoint Clinical deterioration within 15 days of randomization. 		• 24% of patients stopped responding to follow-up prior
 Immunocompromised Unstable medical comorbidities Interventions Fluvoxamine 50 mg PO for 1 dose, then fluvoxamine 100 mg PO twice daily, then fluvoxamine 100 mg PO 3 times daily through Day 15 (n = 80) Placebo (n = 72) Primary Endpoint Clinical deterioration within 15 days of randomization. 		
 Interventions Fluvoxamine 50 mg PO for 1 dose, then fluvoxamine 100 mg PO twice daily, then fluvoxamine 100 mg PO 3 times daily through Day 15 (n = 80) Placebo (n = 72) Primary Endpoint Clinical deterioration within 15 days of randomization. Clinical deterioration within 15 days of randomization. 		
 Fluvoxamine 50 mg PO for 1 dose, then fluvoxamine 100 mg PO twice daily, then fluvoxamine 100 mg PO 3 times daily through Day 15 (n = 80) Placebo (n = 72) Primary Endpoint Clinical deterioration within 15 days of randomization. 	come	Interpretation
 Having dyspnea or being hospitalized for dyspnea or pneumonia; <i>and</i> Having SpO₂ <92% on room air or requiring supplemental oxygen to attain SpO₂ ≥92% Key Secondary Endpoint Hospitalization by Day 15 	terioration: 0% in fluvoxamine 3% in placebo arm (absolute 8.7%; 95% Cl, 1.8% to 16.4%) Dutcome s in fluvoxamine arm ents in placebo arm were	 Fluvoxamine reduced the proportion of patients who experienced clinical deterioration. Due to significant limitations, it is difficult to draw definitive conclusions about the efficacy of using fluvoxamine to treat COVID-19.

Methods	Results	Limitations and Interpretation
TOGETHER: Randomized Platform Trial of Oral Fluvoxamine Plus Inhaled Budesonide for the Treatment of Early Onset COVID-198		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
 Aged ≥50 years or aged ≥18 years with comorbidities Laboratory-confirmed SARS-CoV-2 infection ≤7 days of symptoms 	 Median age 51 years; 61% women 42% with BMI >30; 44% with HTN; 68% with multiple comorbidities 	 Multiple investigational treatments or placebos were evaluated simultaneously. Not all patients in the placebo arm received a matched placebo.
Key Exclusion Criteria	• 94% received \geq 2 doses of a COVID-19 vaccine.	Interpretation
 Use of an SSRI Severe mental illness Cirrhosis, recent seizures, severe ventricular cardiac arrhythmia 	 Primary Outcome Composite of ED observation >6 hours or hospitalization by Day 28: 1.8% in fluvoxamine plus inhaled budesonide arm vs. 3.7% in placebo arm (relative risk 0.50; 95% Crl, 0.25–0.92) 	 In adult outpatients with mild COVID-19, fluvoxamine plus inhaled budesonide reduced the need for ED observations >6 hours or hospitalization when compared with placebo. The use of fluvoxamine plus inhaled budesonide did not reduce hospitalization, health care attendance,
 Fluvoxamine 100 mg PO twice daily plus budesonide 800 µg inhaled twice daily for 10 days (n = 738) Placebo (n = 738; route, dosing frequency, and duration for some patients may have differed from treatment group) Primary Endpoint Composite of ED observation >6 hours or hospitalization for COVID-19 by Day 28 Key Secondary Endpoints Hospitalization by Day 28 Health care attendance by Day 28 Any ED visit by Day 28 Occurrence of AEs 	 0.25–0.92) Secondary Outcomes Hospitalization by Day 28: 0.9% in fluvoxamine plus inhaled budesonide arm vs. 1.1% in placebo arm Health care attendance by Day 28: 2.6% in fluvoxamine plus inhaled budesonide arm vs. 4.1% in placebo arm (relative risk 0.64; 95% Crl, 0.36–1.11) Any ED visit by Day 28: 12.2% in fluvoxamine plus inhaled budesonide arm vs. 13.0% in placebo arm Treatment-emergent AEs: 17.6% in fluvoxamine plus inhaled budesonide arm vs. 12.9% in placebo arm (relative risk 1.37; 95% Crl, 1.07–1.75) Most AEs were grade 2. 	 or the occurrence of any ED visit. It is difficult to define the clinical relevance of the >6-hour ED observation endpoint and apply it to practice settings in different countries. The use of fluvoxamine plus inhaled budesonide resulted in more AEs than placebo.

Key: AE = adverse event; BMI = body mass index; CVD = cardiovascular disease; DM = diabetes mellitus; ED = emergency department; GI = gastrointestinal; HTN = hypertension; ITT = intention-to-treat; the Panel = the COVID-19 Treatment Guidelines Panel; PCR = polymerase chain reaction; PO = oral; PP = per protocol; RCT = randomized controlled trial; RT-PCR = reverse transcription polymerase chain reaction; SpO₂ = oxygen saturation; SSRI = selective serotonin reuptake inhibitor

References

- 1. McCarthy MW, Naggie S, Boulware DR, et al. Effect of fluvoxamine vs placebo on time to sustained recovery in outpatients with mild to moderate COVID-19: a randomized clinical trial. *JAMA*. 2023;329(4):296-305. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36633838.
- 2. Stewart TG, Rebolledo PA, Mourad A, et al. Higher-dose fluvoxamine and time to sustained recovery in outpatients with COVID-19: the ACTIV-6 randomized clinical trial. *JAMA*. 2023;330(24):2354-2363. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37976072</u>.
- 3. Bramante CT, Huling JD, Tignanelli CJ, et al. Randomized trial of metformin, ivermectin, and fluvoxamine for COVID-19. *N Engl J Med*. 2022;387(7):599-610. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36070710</u>.
- 4. Reis G, Dos Santos Moreira-Silva EA, Medeiros Silva DC, et al. Effect of early treatment with fluvoxamine on risk of emergency care and hospitalisation among patients with COVID-19: the TOGETHER randomised, platform clinical trial. *Lancet Glob Health*. 2022;10(1):e42-e51. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34717820.
- 5. Reis G, Mills E. Fluvoxamine for the treatment of COVID-19 Author's reply. *Lancet Glob Health*. 2022;10(3):e333. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35180413.
- 6. Reiersen AM, Mattar C, Bender Ignacio RA, et al. The STOP COVID 2 study: fluvoxamine vs placebo for outpatients with symptomatic COVID-19, a fully remote randomized controlled trial. *Open Forum Infect Dis*. 2023;10(8):ofad419. Available at: https://www.ncbi.nlm.nih.gov/pubmed/37622035.
- 7. Lenze EJ, Mattar C, Zorumski CF, et al. Fluvoxamine vs placebo and clinical deterioration in outpatients with symptomatic COVID-19: a randomized clinical trial. *JAMA*. 2020;324(22):2292-2300. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33180097.
- 8. Reis G, Dos Santos Moreira Silva EA, Medeiros Silva DC, et al. Oral fluvoxamine with inhaled budesonide for treatment of early-onset COVID-19: a randomized platform trial. *Ann Intern Med.* 2023;176(5):667-675. Available at: https://www.ncbi.nlm.nih.gov/pubmed/37068273.

Intravenous Immunoglobulin

Last Updated: December 20, 2023

Recommendations

- The COVID-19 Treatment Guidelines Panel (the Panel) **recommends against** the use of **intravenous immunoglobulin** (**IVIG**) for the treatment of acute COVID-19 in adults and children, except in a clinical trial (**AIII**). This recommendation should not preclude the use of IVIG when it is otherwise indicated for the treatment of underlying conditions or complications that arise during the course of COVID-19.
- For the Panel's recommendations on the use of IVIG in people with multisystem inflammatory syndrome in children (MIS-C) or multisystem inflammatory syndrome in adults (MIS-A) and a discussion of the clinical data that support those recommendations, see <u>Therapeutic Management</u> of Hospitalized Children With MIS-C, Plus a Discussion on MIS-A.

Rationale

It is unknown whether IVIG products derived from pooled donor plasma contain high titers of antibodies that neutralize SARS-CoV-2. Information on SARS-CoV-2 antibody titer was not reported in the clinical trials that evaluated the use of IVIG for the treatment of COVID-19. The levels of SARS-CoV-2 antibodies in IVIG products likely vary depending on which SARS-CoV-2 variant was dominant when the plasma products were collected, and different lots of IVIG may have different titers of antibodies. Although IVIG preparations may have general immunomodulatory effects, these theoretical effects do not appear to benefit patients with COVID-19.¹

Considerations in Pregnant People

IVIG is commonly used during pregnancy for indications such as alloimmune thrombocytopenia.² However, because there is no clear evidence that IVIG is an effective treatment for acute COVID-19 in nonpregnant adults, the Panel **recommends against** the use of **IVIG** for the treatment of acute COVID-19 in pregnant individuals, except in a clinical trial (**AIII**). This recommendation should not preclude the use of IVIG when it is otherwise indicated for the treatment of underlying conditions or complications that arise during the course of COVID-19.

Considerations in Children

No comparative studies have evaluated the use of IVIG in pediatric patients with acute COVID-19. IVIG is used in combination with glucocorticoids to treat MIS-C in pediatric patients.³⁻⁶ However, because there is no clear evidence that IVIG is an effective treatment for acute COVID-19 in adults, the Panel **recommends against** the use of **IVIG** for the treatment of acute COVID-19 in children, except in a clinical trial (**AIII**). This recommendation should not preclude the use of IVIG when it is otherwise indicated.

For the Panel's recommendations for children with MIS-C, see <u>Therapeutic Management of Hospitalized</u> <u>Children With MIS-C, Plus a Discussion on MIS-A</u>.

Clinical Data

In a meta-analysis of 6 randomized controlled trials that enrolled hospitalized patients with COVID-19, the use of non-SARS-CoV-2–specific IVIG was not associated with a survival benefit.¹ All of the

included trials were conducted in 2020, when the presence of SARS-CoV-2 antibodies in blood donors was likely uncommon. None of the studies measured the titers of anti-SARS-CoV-2 antibodies. Blood supplies collected since that time likely have a higher level of these antibodies, and the IVIG derived from those supplies could be expected to have a higher level of SARS-specific antibodies. A British study performed in 2022 evaluated serum anti-SARS-CoV-2 spike antibody titers before and after IVIG infusion in 35 patients with primary immunodeficiencies who were receiving regular immunoglobulin replacement therapy.⁷ The study found that anti-SARS-CoV-2 spike antibody titers and the neutralization capacity of serum increased after IVIG infusion in most patients.

Different brands of commercially available IVIG products exhibit different levels of neutralizing activity against SARS-CoV-2 variants (e.g., BA.1, BA.4, BA.5, BQ.1.1, XBB). A study compared the anti-SARS-CoV-2 antibody levels in U.S. IVIG products that had expiration dates from 2020 to 2025.⁸ The study found that products with expiration dates in 2023 and 2024 were more likely to have higher levels of anti-SARS-CoV-2 antibodies than those with earlier expiration dates. In addition, the study reported an association between later expiration dates and increased inhibition of angiotensin-converting enzyme 2 binding activity. Preparations that were intended for intravenous administration had higher titers than those intended for subcutaneous administration. However, the neutralizing activity against the Omicron variant was lower than the activity against prior variants, and the efficacy of using IVIG for the treatment of COVID-19 remains uncertain.

These data do not provide clear evidence for a clinical benefit of administering IVIG to people with COVID-19. Randomized controlled trials are needed to further define the role of IVIG in the treatment of COVID-19. The use of non-SARS-CoV-2–specific IVIG for the treatment of COVID-19 should be limited to clinical trials.

Concentrated antibody preparations derived from pooled plasma collected from individuals who have recovered from COVID-19 can be manufactured as SARS-CoV-2 hyperimmunoglobulin (hIVIG). Treatment with SARS-CoV-2 hIVIG did not alter patient outcomes in a large randomized controlled trial of hospitalized patients with COVID-19, and hIVIG is not currently available for clinical use in the United States.⁹

References

- 1. Lai CC, Chen WC, Chen CY, Wei YF. The effect of intravenous immunoglobulins on the outcomes of patients with COVID-19: a systematic review and meta-analysis of randomized controlled trials. *Expert Rev Anti Infect Ther*. 2022;20(10):1333-1340. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35786174.
- Lakkaraja M, Berkowitz RL, Vinograd CA, et al. Omission of fetal sampling in treatment of subsequent pregnancies in fetal-neonatal alloimmune thrombocytopenia. *Am J Obstet Gynecol*. 2016;215(4):471.e1-471. e9. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27131591</u>.
- 3. Belhadjer Z, Auriau J, Méot M, et al. Addition of corticosteroids to immunoglobulins is associated with recovery of cardiac function in multi-inflammatory syndrome in children. *Circulation*. 2020;142(23):2282-2284. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33112651.
- 4. Ouldali N, Toubiana J, Antona D, et al. Association of intravenous immunoglobulins plus methylprednisolone vs immunoglobulins alone with course of fever in multisystem inflammatory syndrome in children. *JAMA*. 2021;325(9):855-864. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33523115.
- 5. Son MBF, Murray N, Friedman K, et al. Multisystem inflammatory syndrome in children—initial therapy and outcomes. *N Engl J Med*. 2021;385(1):23-34. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133855</u>.
- McArdle AJ, Vito O, Patel H, et al. Treatment of multisystem inflammatory syndrome in children. N Engl J Med. 2021;385(1):11-22. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34133854</u>.
- 7. Upasani V, Townsend K, Wu MY, et al. Commercial immunoglobulin products contain neutralising antibodies

against SARS-CoV-2 spike protein. *Clin Infect Dis*. 2023;77(7):950-960. Available at: <u>https://www.ncbi.nlm.</u> <u>nih.gov/pubmed/37338118</u>.

- Cousins K, Sano K, Lam B, et al. Detection of SARS-CoV-2 antibodies in immunoglobulin products. *J Allergy Clin Immunol Pract*. 2023;11(8):2534-2541.e2. Available at: https://www.ncbi.nlm.nih.gov/pubmed/37182564.
- ITAC Study Group. Hyperimmune immunoglobulin for hospitalised patients with COVID-19 (ITAC): a double-blind, placebo-controlled, Phase 3, randomised trial. *Lancet*. 2022;399(10324):530-540. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35093205.

Ivermectin

Last Updated: December 20, 2023

Ivermectin is a Food and Drug Administration (FDA)-approved antiparasitic drug used to treat several neglected tropical diseases, including onchocerciasis, helminthiases, and scabies.¹ For these indications, ivermectin has been widely used and is generally well tolerated.^{1,2} Ivermectin is not approved by the FDA for the treatment of any viral infection, including COVID-19. See the FDA webpage <u>Why You</u> <u>Should Not Use Ivermectin to Treat or Prevent COVID-19</u> for more information.

Ivermectin has been shown to inhibit the replication of SARS-CoV-2 in cell cultures.³ However, pharmacokinetic and pharmacodynamic studies suggest that achieving the plasma concentrations necessary for the antiviral efficacy detected in vitro would require administration of doses up to 100-fold higher than those approved for use in humans.^{4,5}

The safety and efficacy of ivermectin for the prevention and treatment of COVID-19 have been evaluated in clinical trials and observational cohorts. Summaries of the studies that informed the COVID-19 Treatment Guidelines Panel's (the Panel) recommendation can be found in <u>Table 7b</u>. The Panel reviewed additional studies, but these studies are not summarized in Table 7b because they have study design limitations or results that make them less definitive and informative.

Recommendation

• The Panel recommends against the use of ivermectin for the treatment of COVID-19 (AIIa).

Rationale

The Panel's recommendation is primarily informed by adequately powered, randomized trials of ivermectin that reported clinical outcomes. Studies that randomized participants to receive ivermectin or a matched placebo had the greatest impact on the Panel's recommendation.⁶⁻¹³

Trials have failed to find a clinical benefit of using ivermectin to treat COVID-19 in outpatients. In TOGETHER, an adaptive platform trial conducted in Brazil, there was no apparent difference between the ivermectin and placebo arms for the primary outcome of risk of emergency department visits or hospitalization (14.7% vs. 16.4%).¹⁴ In addition, there was no statistically significant difference between the ivermectin and placebo arms in mortality (3.1% vs. 3.5%). In COVID-OUT, a randomized factorial trial, the use of ivermectin did not reduce the occurrence of a composite outcome of emergency department visits, hospitalization, or death when compared with a matched control (5.7% vs. 4.1%).⁶

The ACTIV-6 trial was an adaptive platform trial conducted in outpatients with mild to moderate COVID-19 in the United States.^{15,16} Participants were randomized to receive an ivermectin regimen (either 400 μ g/kg for 3 days or 600 μ g/kg for 6 days) or a matching placebo. In the 400 μ g/kg phase of the study, the median time to sustained recovery was 12 days for the ivermectin arm and 13 days for the placebo arm. In the 600 μ g/kg phase of the study, the median time to sustained recovery was 11 days for both arms.

I-TECH, an open-label trial conducted in Malaysia, found no difference between the ivermectin and standard of care arms in the occurrence of the primary outcome of risk of progression to severe COVID-19 (21.6% vs. 17.3%).¹⁷ Patients in the ivermectin arm had a lower risk of mortality than those in the standard of care arm (relative risk 0.31; 95% CI, 0.09–1.11; P = 0.09), but this difference was not statistically significant.

The study populations in most of the reviewed trials were patients with mild to moderate COVID-19 who had a relatively low risk of disease progression, and the number of deaths was low (as expected). In these randomized trials, completely excluding an effect of ivermectin on COVID-19 disease progression is difficult because the trials were not powered to detect differences in secondary outcomes, such as death. However, data from these trials do not provide evidence that the use of ivermectin is effective for the treatment of COVID-19. For this reason, and because other medications now have demonstrated clear clinical benefits for the treatment of COVID-19, the Panel **recommends against** the use of **ivermectin** for the treatment of COVID-19 (**AIIa**).

See <u>Table 7b</u> for summaries of key studies that informed the Panel's recommendation.

References

- 1. Omura S, Crump A. Ivermectin: panacea for resource-poor communities? *Trends Parasitol*. 2014;30(9):445-455. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/25130507</u>.
- Kircik LH, Del Rosso JQ, Layton AM, Schauber J. Over 25 years of clinical experience with ivermectin: an overview of safety for an increasing number of indications. *J Drugs Dermatol*. 2016;15(3):325-332. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26954318</u>.
- Caly L, Druce JD, Catton MG, Jans DA, Wagstaff KM. The FDA-approved drug ivermectin inhibits the replication of SARS-CoV-2 in vitro. *Antiviral Res.* 2020;178:104787. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32251768</u>.
- Chaccour C, Hammann F, Ramón-García S, Rabinovich NR. Ivermectin and COVID-19: keeping rigor in times of urgency. *Am J Trop Med Hyg*. 2020;102(6):1156-1157. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32314704</u>.
- Guzzo CA, Furtek CI, Porras AG, et al. Safety, tolerability, and pharmacokinetics of escalating high doses of ivermectin in healthy adult subjects. *J Clin Pharmacol*. 2002;42(10):1122-1133. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/12362927</u>.
- Bramante CT, Huling JD, Tignanelli CJ, et al. Randomized trial of metformin, ivermectin, and fluvoxamine for COVID-19. *N Engl J Med*. 2022;387(7):599-610. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36070710</u>.
- Vallejos J, Zoni R, Bangher M, et al. Ivermectin to prevent hospitalizations in patients with COVID-19 (IVERCOR-COVID19) a randomized, double-blind, placebo-controlled trial. *BMC Infect Dis*. 2021;21(1):635. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34215210</u>.
- López-Medina E, López P, Hurtado IC, et al. Effect of ivermectin on time to resolution of symptoms among adults with mild COVID-19: a randomized clinical trial. *JAMA*. 2021;325(14):1426-1435. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33662102</u>.
- 9. Buonfrate D, Chesini F, Martini D, et al. High-dose ivermectin for early treatment of COVID-19 (COVER study): a randomised, double-blind, multicentre, phase II, dose-finding, proof-of-concept clinical trial. *Int J Antimicrob Agents*. 2022;59(2):106516. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34999239.
- Abd-Elsalam S, Noor RA, Badawi R, et al. Clinical study evaluating the efficacy of ivermectin in COVID-19 treatment: a randomized controlled study. *J Med Virol*. 2021;93(10):5833-5838. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34076901</u>.
- Ravikirti, Roy R, Pattadar C, et al. Evaluation of ivermectin as a potential treatment for mild to moderate COVID-19: a double-blind randomized placebo controlled trial in Eastern India. *J Pharm Pharm Sci.* 2021;24:343-350. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34265236</u>.
- Mohan A, Tiwari P, Suri TM, et al. Single-dose oral ivermectin in mild and moderate COVID-19 (RIVET-COV): a single-centre randomized, placebo-controlled trial. *J Infect Chemother*. 2021;27(12):1743-1749. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34483029</u>.
- 13. Bermejo Galan LE, Dos Santos NM, Asato MS, et al. Phase 2 randomized study on chloroquine, *COVID-19 Treatment Guidelines*

hydroxychloroquine or ivermectin in hospitalized patients with severe manifestations of SARS-CoV-2 infection. *Pathog Glob Health*. 2021;115(4):235-242. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33682640</u>.

- TOGETHER Investigators. Effect of early treatment with ivermectin among patients with COVID-19. N Engl J Med. 2022;386(18):1721-1731. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35353979</u>.
- 15. Naggie S, Boulware DR, Lindsell CJ, et al. Effect of ivermectin vs placebo on time to sustained recovery in outpatients with mild to moderate COVID-19: a randomized clinical trial. *JAMA*. 2022;328(16):1595-1603. Available at: https://pubmed.ncbi.nlm.nih.gov/36269852.
- Naggie S, Boulware DR, Lindsell CJ, et al. Effect of higher-dose ivermectin for 6 days vs placebo on time to sustained recovery in outpatients with COVID-19: a randomized clinical trial. *JAMA*. 2023;329(11):888-897. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36807465</u>.
- Lim SCL, Hor CP, Tay KH, et al. Efficacy of ivermectin treatment on disease progression among adults with mild to moderate COVID-19 and comorbidities: the I-TECH randomized clinical trial. *JAMA Intern Med*. 2022;182(4):426-435. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35179551</u>.

Table 7b. Ivermectin: Selected Clinical Trial Data

Last Updated: December 20, 2023

The clinical trials described in this table are the RCTs that had the greatest impact on the Panel's recommendation. The Panel reviewed other clinical studies that evaluated the use of IVM for the treatment of COVID-19.¹⁻²⁶ However, those studies have limitations that make them less definitive and informative than the studies summarized in this table.

Methods	Results	Limitations and Interpretation	
ACTIV-6: Double-Blind RCT of Ivermectin 600 µg/kg in Outpatients With Mild to Moderate COVID-19 in the United States ²⁷			
Key Inclusion Criteria	Participant Characteristics	Key Limitation	
 Aged ≥30 years 	Median age 48 years; 59.1% women	The low number of events limited	
Not hospitalized	• 38.1% with BMI >30; 9.2% with DM; 26.8% with HTN	the power to determine an effect on	
Positive SARS-CoV-2 test result within past 10 days	• 83.6% received ≥2 COVID-19 vaccine doses.	hospitalization and death.	
• \geq 2 COVID-19 symptoms for \leq 7 days	Median of 5 days from symptom onset to receipt of	Interpretation	
Key Exclusion Criteria	study drug	 Among outpatients with COVID-19, IVM 600 µg/kg PO once daily for 6 days did 	
End-stage kidney disease	Primary Outcome	not shorten time to sustained recovery	
Liver failure or decompensated cirrhosis	Median time to sustained recovery: 11 days in IVM arm	or reduce incidence of hospitalization or death.	
Interventions	vs. 11 days in placebo arm (HR 1.02; 95% Crl, 0.92– 1.13)		
• IVM 600 μ g/kg PO once daily for 6 days (n = 602)	Secondary Outcome		
• Placebo (n = 604)	Hospitalization or death by Day 28: 5 (0.8%) in IVM arm		
Primary Endpoint	vs. 2 (0.3%) in placebo arm		
• Time to sustained recovery (i.e., \geq 3 consecutive days	Safety Outcomes		
without symptoms)	• Occurrence of AEs: 52 of 566 patients (9.2%) in IVM arm		
Key Secondary Endpoint	vs. 41 of 576 patients (7.1%) in placebo arm		
Hospitalization or death by Day 28	Occurrence of SAEs: 5 of 566 patients (0.9%) in IVM arm		
Safety Endpoint	vs. 3 of 576 patients (0.5%) in placebo arm		
Occurrence of AEs and SAEs			

Methods	Results	Limitations and Interpretation		
ACTIV-6: Double-Blind RCT of Ivermectin 400 µg/kg Or	ACTIV-6: Double-Blind RCT of Ivermectin 400 µg/kg Once Daily in Outpatients With Mild to Moderate COVID-19 in the United States ²⁸			
Key Inclusion Criteria	Participant Characteristics	Key Limitation		
 Aged ≥30 years 	Mean age 48 years; 59% women	The low number of events limited		
Not hospitalized	• 41% with BMI >30; 11.5% with DM; 26% with HTN	the power to determine an effect on		
Positive SARS-CoV-2 test result within past 10 days	 47% received ≥2 COVID-19 vaccine doses. 	hospitalization and death.		
• \geq 2 COVID-19 symptoms for \leq 7 days	Median of 6 days from symptom onset to receipt of study	Interpretation		
Key Exclusion Criteria	drug	 Among outpatients with COVID-19, IVM 400 µg/kg PO once daily for 3 days did 		
End-stage kidney disease	Primary Outcome	not shorten time to sustained recovery		
Liver failure or decompensated cirrhosis	Median time to sustained recovery: 12 days in IVM arm	or reduce incidence of hospitalization or		
Interventions	vs. 13 days in placebo arm (HR 1.07; 95% Crl, 0.96– 1.17)	death.		
• IVM 400 μ g/kg PO once daily for 3 days (n = 817)	Secondary Outcome			
• Placebo (n = 774)	Hospitalization or death by Day 28: 10 (1.2%) in IVM arm			
Primary Endpoint	vs. 9 (1.2%) in placebo arm			
• Time to sustained recovery (i.e., \geq 3 consecutive days	Safety Outcomes			
without symptoms)	Occurrence of AEs: 24 of 766 patients (3.1%) in IVM arm			
Key Secondary Endpoint	vs. 27 of 724 patients (3.7%) in placebo arm			
 Hospitalization or death by Day 28 	Occurrence of SAEs: 9 of 766 patients (1.2%) in IVM arm			
Safety Endpoint	vs. 9 of 724 patients (1.2%) in placebo arm			
Occurrence of AEs and SAEs				

Methods	Results	Limitations and Interpretation
TOGETHER: Double-Blind, Adaptive RCT of Ivermectin in	Nonhospitalized Patients With COVID-19 in Brazil ²⁹	
Key Inclusion Criteria	Participant Characteristics	Key Limitations
 Positive SARS-CoV-2 antigen test result 	 Median age 49 years; 46% aged ≥50 years; 58% 	Health care facility capacity may have
Within 7 days of symptom onset	women; 95% self-identified as mixed race	influenced the number and duration of
• ≥1 high-risk factor for disease progression (e.g., aged	 Most prevalent risk factor: 50% with obesity 	ED visits and hospitalizations.
>50 years, comorbidities, immunosuppression)	44% within 3 days of symptom onset at enrollment	 No details on safety outcomes (e.g., type of treatment-emergent AEs) other than
Interventions	Primary Outcome	grading were reported.
• IVM 400 μ g/kg PO once daily for 3 days (n = 679)	• Composite of ED observation >6 hours or hospitalization	Interpretation
• Placebo (n = 679; not all patients received IVM placebo)	for COVID-19 by Day 28 (ITT): 100 (14.7%) in IVM arm vs.	 In outpatients with recent SARS-CoV-2
Primary Endpoint	111 (16.4%) in placebo arm (relative risk 0.90; 95% Crl, 0.70–1.16)	infection, IVM did not reduce the need
 Composite of ED observation >6 hours or hospitalization for COVID-19 by Day 28 	• 171 (81%) of events were hospitalizations (ITT)	for ED visits or hospitalization when compared with placebo.
	Secondary Outcomes	
Key Secondary Endpoints	• No difference between IVM arm and placebo arm in:	
Viral clearance at Day 7	• Viral clearance at Day 7 (relative risk 1.00; 95% Crl,	
All-cause mortality	0.68–1.46)	
Occurrence of AEs	 All-cause mortality: 21 (3.1%) vs. 24 (3.5%) (relative risk 0.88; Crl, 0.49–1.55) 	
	Occurrence of AEs	

Methods	Results	Limitations and Interpretation
COVID-OUT: RCT of Metformin, Ivermectin, and Fluvoxamine in Nonhospitalized Adults With COVID-19 in the United States ³⁰		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
Aged 30–85 years	Median age 46 years; 56% women; 82% White	 Study included Sp0₂ measurements
• BMI \geq 25 or \geq 23 if Asian or Latinx	Median BMI 30	using home pulse oximeters as 1 of
• Laboratory-confirmed SARS-CoV-2 infection within 3 days	• 27% with CVD	the composite measures of the primary endpoint. However, the FDA has issued
of randomization	 52% received primary COVID-19 vaccination series. 	a statement concerning the accuracy
 ≤7 days of COVID-19 symptoms 	 Mean of 4.8 days of symptoms 	of these home pulse oximeters, making
Key Exclusion Criteria	 Approximately 68% enrolled while Delta was the 	this study endpoint less reliable.
Immunocompromised	dominant variant; approximately 29% enrolled while Omicron was dominant.	• SpO ₂ data were incomplete or missing for 30% of the patients.
Hepatic impairment		The low number of events limited
• Stage 4–5 chronic kidney disease or eGFR <45 mL/	Primary Outcomes	the power to determine the effect on
min/1.73 m ²	 Composite of hypoxemia, ED visit, hospitalization, or death by Day 14: 105 (25.8%) in IVM arm vs. 96 (24.6%) 	hospitalization and death.
Interventions	in control arm (aOR 1.05; 95% Cl, $0.76-1.45$, $P = 0.78$)	Interpretations
• IVM 390–470 ug/kg PO once daily for 3 days (n = 410) in the following arms:	• No difference between IVM alone arm and placebo alone	IVM did not prevent the composite
the following arms:IVM alone (n = 206)	arm in occurrence of primary endpoint (aOR 1.06; 95%	endpoint of hypoxemia, ED visit,
• Metformin plus IVM (n = 204)	Cl, 0.67–1.67)	hospitalization, or death.No primary, secondary, or subgroup
 IVM control (n = 398), which included the following arms: 	 ED visit, hospitalization, or death by Day 14 in a prespecified secondary analysis: 23 (5.7%) in IVM arm 	analysis demonstrated a benefit for the
• Placebo alone ($n = 203$)	vs. 16 (4.1%) in control arm (aOR 1.39; 95% Cl, 0.72–	use of IVM over placebo.
• Metformin alone (n = 195)	2.69)	
	Hospitalization or death by Day 14 in a prespecified	
Primary Endpoints	secondary analysis: 4 (1.0%) in IVM arm vs. 5 (1.3%) in control arm (aOR 0.73; 95% Cl, 0.19–2.77); 1 death in	
 Composite of hypoxemia (SpO₂ ≤93%, as measured by a home pulse oximeter), ED visit, hospitalization, or death 	IVM arm vs. 0 deaths in control arm	
by Day 14	Secondary Outcomes	
 A prespecified secondary analysis evaluated the 	 No difference between arms in total symptom severity 	
occurrence of ED visits, hospitalization, or death by Day	score by Day 14	
14.	• Drug discontinuation or interruption: 20% in IVM arm vs.	
Key Secondary Endpoints	25% in placebo alone arm	
 Total symptom severity score by Day 14, as measured by a symptom severity scale 		
 Drug discontinuation or interruption 		

Methods	Results	Limitations and Interpretation
IVERCOR-COVID19: Double-Blind, Placebo-Controlled, Randomized Trial of Ivermectin in Nonhospitalized Patients With COVID-19 in Argentina ³¹		
Key Inclusion CriterionPositive SARS-CoV-2 RT-PCR result within 48 hours of	 Participant Characteristics Mean age 42 years; 8% aged ≥65 years; 47% women 	Key LimitationStudy enrolled a young population with
screening Key Exclusion Criteria	 24% with HTN; 10% with DM; 58% with ≥1 comorbidity Median of 4 days from symptom onset 	few of the comorbidities that predict disease progression.
 Required supplemental oxygen or hospitalization Concomitant use of CQ or HCQ 	 Primary Outcome Hospitalization for any reason: 5.6% in IVM arm vs. 8.3% in placebo arm (OR 0.65; 95% CI, 0.32–1.31; P = 0.23) 	 Interpretation Among patients who had recently acquired SARS-CoV-2 infection, there
 Weight-based dose of IVM PO at enrollment and 24 hours later for a maximum total dose of 48 mg (n = 250) 	 Secondary Outcomes Need for MV: 2% in IVM arm vs. 1% in placebo arm (P = 0.7) 	was no evidence that IVM provided any clinical benefit.
 Placebo (n = 251) Primary Endpoint 	• All-cause mortality: 2% in IVM arm vs. 1% in placebo arm ($P = 0.7$)	
 Hospitalization for any reason Key Secondary Endpoints 	• Occurrence of AEs: 18% in IVM arm vs. 21% in placebo arm ($P = 0.6$)	
Need for MV		
All-cause mortalityOccurrence of AEs		

Methods	Results	Limitations and Interpretation		
Double-Blind, Placebo-Controlled, Randomized Trial of	Double-Blind, Placebo-Controlled, Randomized Trial of Ivermectin in Patients With Mild COVID-19 in Colombia ³²			
Double-Blind, Placebo-Controlled, Randomized Trial of Key Inclusion Criteria Positive SARS-CoV-2 RT-PCR or antigen test result ≤7 days of COVID-19 symptoms Mild disease Key Exclusion Criteria Asymptomatic disease Severe pneumonia Hepatic dysfunction Interventions IVM 300 μg/kg PO once daily for 5 days (n = 200) Placebo PO (n = 198) Primary Endpoint Time to symptom resolution within 21 days Key Secondary Endpoints Clinical deterioration	 Participant Characteristics Median age 37 years; 4% aged ≥65 years in IVM arm, 8% in placebo arm; 39% men in IVM arm, 45% in placebo arm 79% with no known comorbidities Median of 5 days from symptom onset to randomization Primary Outcome Median time to symptom resolution: 10 days in IVM arm vs. 12 days in placebo arm (HR 1.07; <i>P</i> = 0.53) Symptoms resolved by Day 21: 82% in IVM arm vs. 79% in placebo arm Secondary Outcomes No difference between arms in proportion of patients who showed clinical deterioration or required escalation of care Occurrence of AEs: Discontinued treatment due to AEs: 8% in IVM arm vs. 	 Key Limitations Due to low event rates, the primary endpoint changed from the proportion of patients with clinical deterioration to the time to symptom resolution during the trial. The study enrolled younger, healthier patients, a population that does not typically develop severe COVID-19. Interpretation In patients with mild COVID-19, IVM 300 µg/kg once daily for 5 days did not improve the time to symptom resolution. 		
Occurrence of AEs	3% in placebo armNo SAEs related to intervention			

Methods	Results	Limitations and Interpretation
-TECH: Open-Label RCT of Ivermectin in Patients With Mild to Moderate COVID-19 in Malaysia ³³		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
Positive SARS-CoV-2 RT-PCR or antigen test result	Mean age 63 years; 55% women	Open-label study
 within 7 days of symptom onset Aged ≥50 years 	 68% received ≥1 COVID-19 vaccine dose; 52% received 2 doses. 	Interpretation
 ≥1 comorbidities 	• Most common comorbidities: 75% with HTN; 54% with	• In patients with mild to moderate COVID-19, there was no evidence
Key Exclusion CriteriaRequired supplemental oxygen	DM; 24% with dyslipidemia • Mean of 5 days symptom duration	that IVM provided any clinical benefit, including no evidence that IVM reduced
 Severe hepatic impairment (ALT >10 times the ULN) 	Primary Outcome	the risk of progression to severe disease.
 Interventions IVM 400 μg/kg PO once daily for 5 days plus SOC (n = 	 Progression to severe COVID-19 (mITT): 52 (21.6%) in IVM plus SOC arm vs. 43 (17.3%) in SOC alone arm (relative risk 1.25; 95% Cl, 0.87–1.80; P = 0.25) 	
241) • SOC (n = 249)	Secondary Outcomes	
Primary Endpoint	 No difference between IVM plus SOC arm and SOC alone arm in: 	
 Progression to severe COVID-19 (i.e., hypoxemia requiring supplemental oxygen to maintain SpO₂ ≥95%) 	 In-hospital, all-cause mortality by Day 28: 3 (1.2%) vs. 10 (4.0%) (relative risk 0.31; 95% Cl, 0.09–1.11; P = 	
Key Secondary Endpoints	0.09)	
 In-hospital, all-cause mortality by Day 28 MV or ICU admission 	 MV: 4 (1.7%) vs. 10 (4.0%) (relative risk 0.41; 95% Cl, 0.13–1.30; P = 0.17) 	
Occurrence of AEs	• ICU admission: 6 (2.5%) vs. 8 (3.2%) (relative risk 0.78; 95% CI, 0.27–2.20; <i>P</i> = 0.79)	
	• Occurrence of AEs: 33 (13.7%) in IVM plus SOC arm vs. 11 (4.4%) in SOC alone arm; most with diarrhea (14 vs. 4)	

Methods	Results	Limitations and Interpretation	
COVER: Phase 2, Double-Blind RCT of Ivermectin in Nonhospitalized Patients With COVID-19 in Italy ³⁴			
 COVER: Phase 2, Double-Bind RCT of ivermectin in Non Key Inclusion Criteria Asymptomatic or oligosymptomatic disease SARS-CoV-2 infection confirmed by RT-PCR result Not hospitalized or receiving supplemental oxygen Key Exclusion Criteria CNS disease Receiving dialysis Severe medical condition with <6 months survival prognosis Use of warfarin, antiviral agents, CQ, or HCQ Interventions IVM 1,200 µg/kg PO once daily for 5 days (n = 32) IVM 600 µg/kg plus placebo PO once daily for 5 days (n = 29) Placebo PO (n = 32) Primary Endpoints Number of SAEs Change in VL at Day 7 	 Participant Characteristics Median age 47 years; 58% men 86% with COVID-19 symptoms 2.2% received a COVID-19 vaccine. Primary Outcomes No SAEs related to intervention Mean log₁₀ reduction in VL at Day 7: 2.9 in IVM 1,200 µg/kg arm vs. 2.5 in IVM 600 µg/kg arm vs. 2.0 in placebo arm (IVM 1,200 µg/kg vs. placebo, <i>P</i> = 0.099; IVM 600 µg/kg vs. placebo, <i>P</i> = 0.122) Other Outcomes 14 (15.1%) discontinued treatment: 11 (34.4%) in IVM 1,200 µg/kg arm vs. 2 (6.9%) in IVM 600 µg/kg arm vs. 1 (3.1%) in placebo arm All discontinuations in IVM 1,200 µg/kg arm were due to tolerability 	 Key Limitations Small, Phase 2 study 90% of subjects screened were not enrolled for various reasons. Recruitment stopped early because of a decline in the number of COVID-19 cases. Interpretations A high dose of IVM (1,200 µg/kg) appears to be safe but not well tolerated; 34% of patients discontinued therapy due to AEs. There was no significant difference in reduction of VL between IVM and placebo arms. 	
• Drug discontinuation or interruption			

Methods	Results	Limitations and Interpretation		
Open-Label RCT of Ivermectin in Hospitalized Patients W	Dpen-Label RCT of Ivermectin in Hospitalized Patients With COVID-19 in Egypt ³⁵			
 Key Inclusion Criteria RT-PCR-confirmed SARS-CoV-2 infection by pharyngeal swab Hospitalized with mild to moderate COVID-19 Key Exclusion Criterion Cardiac problems Interventions IVM 12 mg PO once daily for 3 days (n = 82) SOC (n = 82) Primary Endpoint 	Articipant Characteristics Mean age 42 years for IVM arm, 39 years for SOC arm; 50% men 49% with ≥1 comorbidities imary Outcome All-cause mortality by 28 days: 3 (3.7%) in IVM arm ys. 4	Tey Limitation Small, open-label study nterpretation The use of IVM did not reduce all-cause mortality, hospital LOS, or the need for MV among patients with mild to moderate COVID-19.		
 Primary Endpoint All-cause mortality by 28 days 	$\operatorname{arm}(P = 0.085)$			

Methods	Results	Limitations and Interpretation		
Double-Blind, Placebo-Controlled, Randomized Trial of	Double-Blind, Placebo-Controlled, Randomized Trial of Ivermectin in Patients With Mild to Moderate COVID-19 in India ³⁶			
 Key Inclusion Criteria Positive SARS-CoV-2 RT-PCR or antigen test result Hospitalized with mild to moderate COVID-19 Interventions IVM 12 mg PO once daily for 2 days (n = 55) Placebo PO (n = 57) Primary Endpoint Negative SARS-CoV-2 RT-PCR result on Day 6 	 Participant Characteristics Mean age 53 years; 28% women 35% with HTN; 36% with DM 79% with mild COVID-19 Mean of 6.9 days from symptom onset 100% received HCQ, steroids, and antibiotics; 21% received RDV; 6% received tocilizumab. Primary Outcome Negative SARS-CoV-2 RT-PCR result on Day 6: 24% in 	 Key Limitations Although the primary endpoint was a negative SARS-CoV-2 RT-PCR result on Day 6, no RT-PCR result or an inconclusive RT-PCR result was reported for 42% of patients in the IVM arm and 23% in the placebo arm. The time to discharge was not reported, and outcomes after discharge were not evaluated. 		
 Key Secondary Endpoints Symptom resolution by Day 6 Discharge by Day 10 Need for ICU admission or MV In-hospital mortality 	 IVM arm vs. 32% in placebo arm (rate ratio 0.8; P = 0.348) Secondary Outcomes Symptom resolution by Day 6: 84% in IVM arm vs. 90% in placebo arm (rate ratio 0.9; P = 0.36) Discharge by Day 10: 80% in IVM arm vs. 74% in placebo arm (rate ratio 1.1; P = 0.43) No difference between arms in need for ICU admission or MV In-hospital mortality: 0 in IVM arm (0%) vs. 4 in placebo arm (7%) 	 Interpretation IVM provided no significant virologic or clinical benefit for patients with mild to moderate COVID-19. 		

Methods	Results	Limitations and Interpretation
RIVET-COV: Double-Blind, Placebo-Controlled, Randomized Trial of Ivermectin in Patients With Mild to Moderate COVID-19 in India ³⁷		
Key Inclusion Criteria	Participant Characteristics	Key Limitation
Positive SARS-CoV-2 RT-PCR or antigen test result	Mean age 35 years; 89% men	Small sample size
Nonsevere COVID-19	• 60% to 68% with mild COVID-19 (including asymptomatic	Interpretation
Key Exclusion Criteria	patients); 33% to 40% with moderate COVID-19	The use of IVM did not affect the
• CrCl <30 mL/min	 Median of 4–5 days symptom duration; similar across arms 	proportion of patients with negative
 Transaminases >5 times ULN 	 10% in each arm received concurrent antivirals (RDV, 	SARS-CoV-2 RT-PCR results at Day 5 or the clinical outcomes.
• MI, heart failure, QTc interval prolongation	favipiravir, or HCQ).	
Severe comorbidity	Primary Outcomes	
Interventions	Negative SARS-CoV-2 RT-PCR result at Day 5: 48% in	
• Single dose of IVM 24 mg PO (n = 51)	IVM 24 mg arm vs. 35% in IVM 12 mg arm vs. 31% in	
• Single dose of IVM 12 mg PO (n = 49)	placebo arm ($P = 0.30$)	
• Placebo (n = 52)	No significant difference between arms in decline of VL at Day 5	
Primary Endpoints		
Negative SARS-CoV-2 RT-PCR result at Day 5	Secondary Outcomes	
Decline of VL at Day 5	 No difference between arms in time to symptom resolution 	
Key Secondary Endpoints	• Clinical worsening at Day 14: 8% in IVM 24 mg arm vs.	
Time to symptom resolution	5% in IVM 12 mg arm vs. 11% in placebo arm ($P = 0.65$)	
Clinical worsening at Day 14	No difference between arms in number of hospital-free	
Number of hospital-free days at Day 28	days at Day 28	
Frequency of AEs	No difference between arms in frequency of AEs; no SAEs were reported	

Methods	Results	Limitations and Interpretation
Double-Blind RCT of Ivermectin, Chloroquine, or Hydroxychloroquine in Hospitalized Adults With Severe COVID-19 in Brazil ³⁸		
Key Inclusion Criteria	Participant Characteristics	Key Limitations
Hospitalized with laboratory-confirmed SARS-CoV-2	Mean age 53 years; 58% men	Small sample size
infection	Most common comorbidities: 43% with HTN; 28% with	No clearly defined primary endpoint
 ≥1 of the following severity criteria: 	DM; 38% with BMI >30	Interpretation
• Dyspnea	 76% with respiratory failure on admission 	 Compared to CQ or HCQ, IVM did not
 Tachypnea (>30 breaths/min) 	Outcomes	reduce the proportion of hospitalized
• SpO ₂ <93%	 No difference between IVM, CQ, and HCQ arms in: 	patients with severe COVID-19 who died
• $PaO_2/FiO_2 < 300 \text{ mm Hg}$	Need for supplemental oxygen: 88% vs. 89% vs. 90%	or who required supplemental oxygen, ICU admission, or MV
 Involvement of >50% of lungs confirmed by CXR or 	 Need for MV: 24% vs. 21% vs. 21% 	
CT scan	 ICU admission: 28% vs. 22% vs. 21% 	
Key Exclusion Criterion	 Mortality: 23% vs. 21% vs. 22% 	
Cardiac arrhythmia	Mean number of days of supplemental oxygen: 8 days	
Interventions	in each arm	
 IVM 14 mg once daily for 3 days (n = 53) 	No difference between arms in occurrence of AEs	
 CQ 450 mg twice daily on Day 0, then once daily for 4 days (n = 61) 	 Baseline characteristics significantly associated with mortality: 	
• HCQ 400 mg twice daily on Day 0, then once daily for 4	 Aged >60 years (HR 2.4) 	
days $(n = 54)$	• DM (HR 1.9)	
Endpoints	• BMI >33 (HR 2.0)	
• Need for supplemental oxygen, MV, or ICU admission	• Sp0 ₂ <90% (HR 5.8)	
Occurrence of AEs		
Mortality		

Key: AE = adverse event; ALT = alanine transaminase; BMI = body mass index; CNS = central nervous system; CQ = chloroquine; CrCI = creatinine clearance; CT = computed tomography; CVD = cardiovascular disease; CXR = chest X-ray; DM = diabetes mellitus; ED = emergency department; eGFR = estimated glomerular filtration rate; FDA = Food and Drug Administration; HCQ = hydroxychloroquine; HTN = hypertension; ICU = intensive care unit; ITT = intention-to-treat; IVM = ivermectin; LOS = length of stay; MI = myocardial infarction; mITT = modified intention-to-treat; MV = mechanical ventilation; the Panel = the COVID-19 Treatment Guidelines Panel; PaO₂/FiO₂ = ratio of arterial partial pressure of oxygen to fraction of inspired oxygen; PO = oral; RCT = randomized controlled trial; RDV = remdesivir; RT-PCR = reverse transcriptase polymerase chain reaction; SAE = severe adverse event; SOC = standard of care; SpO₂ = oxygen saturation; ULN = upper limit of normal; VL = viral load

References

- 1. Spoorthi V, Sasank S. Utility of ivermectin and doxycycline combination for the treatment of SARS-CoV-2. *Int Arch Integrated Med.* 2020;7(10):117-182. Available at: <u>https://iaimjournal.com/wp-content/uploads/2020/10/iaim_2020_0710_23.pdf</u>.
- 2. Camprubí D, Almuedo-Riera A, Martí-Soler H, et al. Lack of efficacy of standard doses of ivermectin in severe COVID-19 patients. *PLoS One*. 2020;15(11):e0242184. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33175880.
- 3. Bhattacharya R, Ray I, Mukherjee R, et al. Observational study on clinical features, treatment and outcome of COVID-19 in a tertiary care centre in India—a retrospective case series. Int J Sci Res. 2020;9(10):69-71. Available at: <u>https://www.worldwidejournals.com/international-journal-of-scientific-research-(IJSR)/article/observational-study-on-clinical-features-treatment-and-outcome-of-covid-19-in-a-tertiary-care-centre-in-india-andndash-a-retrospective-case-series/MzI0NTg=/?is=1&b1=141&k=36.</u>
- 4. Morgenstern J, Redondo JN, De León A, et al. The use of compassionate ivermectin in the management of symptomatic outpatients and hospitalized patients with clinical diagnosis of COVID-19 at the Medical Center Bournigal and the Medical Center Punta Cana, Rescue Group, Dominican Republic, from May 1 to August 10, 2020. *medRxiv*. 2020;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2020.10.29.20222505v1</u>.
- Cadegiani FA, Goren A, Wambier CG, McCoy J. Early COVID-19 therapy with azithromycin plus nitazoxanide, ivermectin or hydroxychloroquine in outpatient settings significantly improved COVID-19 outcomes compared to known outcomes in untreated patients. *New Microbes New Infect*. 2021;43:100915. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34249367</u>.
- 6. Carvallo H, Roberto H, Eugenia FM. Safety and efficacy of the combined use of ivermectin, dexamethasone, enoxaparin and aspirin against COVID 19. *medRxiv*. 2020;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2020.09.10.20191619v1</u>.
- 7. Bukhari KHS, Asghar A, Perveen N, et al. Efficacy of ivermectin in COVID-19 patients with mild to moderate disease. *medRxiv*. 2021;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2021.02.02.21250840v1</u>.
- 8. Elalfy H, Besheer T, El-Mesery A, et al. Effect of a combination of nitazoxanide, ribavirin, and ivermectin plus zinc supplement (MANS.NRIZ study) on the clearance of mild COVID-19. *J Med Virol.* 2021;93(5):3176-3183. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33590901</u>.
- 9. Chahla RE, Ruiz LM, Mena T, et al. Cluster randomised trials—ivermectin repurposing for COVID-19 treatment of outpatients with mild disease in primary health care centers. *Research Square*. 2021;Preprint. Available at: https://www.researchsquare.com/article/rs-495945/v1.
- 10. Tanioka H, Tanioka S, Kaga K. Why COVID-19 is not so spread in Africa: how does ivermectin affect it? *medRxiv*. 2021;Preprint. Available at: https://www.medrxiv.org/content/10.1101/2021.03.26.21254377v1.
- 11. Roy S, Samajdar SS, Tripathi SK, Mukherjee S, Bhattacharjee K. Outcome of different therapeutic interventions in mild COVID-19 patients in a single OPD clinic of West Bengal: a retrospective study. *medRxiv*. 2021;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2021.03.08.21252883v2</u>.
- 12. Shahbaznejad L, Davoudi A, Eslami G, et al. Effects of ivermectin in patients with COVID-19: a multicenter, double-blind, randomized, controlled clinical trial. *Clin Ther*. 2021;43(6):1007-1019. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34052007.
- 13. Roman YM, Burela PA, Pasupuleti V, et al. Ivermectin for the treatment of coronavirus disease 2019: a systematic review and meta-analysis of randomized controlled trials. *Clin Infect Dis*. 2022;74(6):1022-1029. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34181716.
- 14. Ahmed S, Karim MM, Ross AG, et al. A five-day course of ivermectin for the treatment of COVID-19 may reduce the duration of illness. *Int J Infect Dis*. 2021;103:214-216. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33278625</u>.

COVID-19 Treatment Guidelines

- 15. Chaccour C, Casellas A, Blanco-Di Matteo A, et al. The effect of early treatment with ivermectin on viral load, symptoms and humoral response in patients with non-severe COVID-19: a pilot, double-blind, placebo-controlled, randomized clinical trial. *EClinicalMedicine*. 2021;32:100720. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33495752.
- 16. Chachar AZK, Khan KA, Asif M, et al. Effectiveness of ivermectin in SARS-COV-2/COVID-19 patients. *Int J Sci.* 2020;9:31-35. Available at: https://www.ijsciences.com/pub/article/2378.
- 17. Gonzalez JLB, Gámez MG, Enciso EAM, et al. Efficacy and safety of ivermectin and hydroxychloroquine in patients with severe COVID-19: a randomized controlled trial. *Infect Dis Rep.* 2022;14(2):160-168. Available at: https://www.mdpi.com/2036-7449/14/2/20.
- 18. Hashim HA, Maulood MF, Ali CL, et al. Controlled randomized clinical trial on using ivermectin with doxycycline for treating COVID-19 patients in Baghdad, Iraq. *Iraqi J Med Sci.* 2021;29(1):107-115. Available at: https://www.iraqijms.net/index.php?do=view&type=article&id=779.
- 19. Khan MSI, Debnath CR, Nath PN, et al. Ivermectin treatment may improve the prognosis of patients with COVID-19. *Arch Bronconeumol (Engl Ed)*. 2020;56(12):828-830. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33293006</u>.
- 20. Krolewiecki A, Lifschitz A, Moragas M, et al. Antiviral effect of high-dose ivermectin in adults with COVID-19: a proof-of-concept randomized trial. *EClinicalMedicine*. 2021;37:100959. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34189446</u>.
- 21. Okumuş N, Demirtürk N, Çetinkaya RA, et al. Evaluation of the effectiveness and safety of adding ivermectin to treatment in severe COVID-19 patients. *BMC Infect Dis*. 2021;21(1):411. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33947344</u>.
- 22. Podder CS, Chowdhury N, Sina MI, Mohosin Ul Haque WM. Outcome of ivermectin treated mild to moderate COVID-19 cases: a single-centre, open-label, randomised controlled study. *IMC J Med Sci*. 2021;14(2). Available at: https://doi.org/10.3329/imcjms.v14i2.52826.
- 23. Soto-Becerra P, Culquichicón C, Hurtado-Roca Y, Araujo-Castillo RV. Real-world effectiveness of hydroxychloroquine, azithromycin, and ivermectin among hospitalized COVID-19 patients: results of a target trial emulation using observational data from a nationwide healthcare system in Peru. *medRxiv*. 2020;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2020.10.06.20208066v3</u>.
- 24. Rajter JC, Sherman MS, Fatteh N, et al. Use of ivermectin is associated with lower mortality in hospitalized patients with coronavirus disease 2019: the ivermectin in COVID nineteen study. *Chest*. 2021;159(1):85-92. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33065103.
- 25. Chowdhury ATMM, Shahbaz M, Karim MR, et al. A comparative study on ivermectin-doxycycline and hydroxychloroquine-azithromycin therapy on COVID-19 patients. *EJMO*. 2021;5(1):63-70. Available at: <u>https://www.ejmo.org/10.14744/ejmo.2021.16263/</u>.
- 26. Niaee MS, Namdar P, Allami A, et al. Ivermectin as an adjunct treatment for hospitalized adult COVID-19 patients: a randomized multi-center clinical trial. Asian Pac J Trop Med. 2021;14(6):266-273. Available at: https://journals.lww.com/aptm/Fulltext/2021/14060/Ivermectin as an adjunct treatment for.3.aspx.
- 27. Naggie S, Boulware DR, Lindsell CJ, et al. Effect of higher-dose ivermectin for 6 days vs placebo on time to sustained recovery in outpatients with COVID-19: a randomized clinical trial. *JAMA*. 2023;329(11):888-897. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36807465</u>.
- 28. Naggie S, Boulware DR, Lindsell CJ, et al. Effect of ivermectin vs placebo on time to sustained recovery in outpatients with mild to moderate COVID-19: a randomized clinical trial. *JAMA*. 2022;328(16):1595-1603. Available at: https://pubmed.ncbi.nlm.nih.gov/36269852.
- 29. TOGETHER Investigators. Effect of early treatment with ivermectin among patients with COVID-19. *N Engl J Med.* 2022;386(18):1721-1731. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35353979</u>.

COVID-19 Treatment Guidelines

- 30. Bramante CT, Huling JD, Tignanelli CJ, et al. Randomized trial of metformin, ivermectin, and fluvoxamine for COVID-19. *N Engl J Med*. 2022;387(7):599-610. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36070710</u>.
- 31. Vallejos J, Zoni R, Bangher M, et al. Ivermectin to prevent hospitalizations in patients with COVID-19 (IVERCOR-COVID19) a randomized, doubleblind, placebo-controlled trial. *BMC Infect Dis*. 2021;21(1):635. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34215210</u>.
- 32. López-Medina E, López P, Hurtado IC, et al. Effect of ivermectin on time to resolution of symptoms among adults with mild COVID-19: a randomized clinical trial. *JAMA*. 2021;325(14):1426-1435. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33662102.
- 33. Lim SCL, Hor CP, Tay KH, et al. Efficacy of ivermectin treatment on disease progression among adults with mild to moderate COVID-19 and comorbidities: the I-TECH randomized clinical trial. *JAMA Intern Med.* 2022;182(4):426-435. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35179551</u>.
- 34. Buonfrate D, Chesini F, Martini D, et al. High-dose ivermectin for early treatment of COVID-19 (COVER study): a randomised, double-blind, multicentre, phase II, dose-finding, proof-of-concept clinical trial. *Int J Antimicrob Agents*. 2022;59(2):106516. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34999239.
- 35. Abd-Elsalam S, Noor RA, Badawi R, et al. Clinical study evaluating the efficacy of ivermectin in COVID-19 treatment: a randomized controlled study. *J Med Virol*. 2021;93(10):5833-5838. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34076901</u>.
- 36. Ravikirti, Roy R, Pattadar C, et al. Evaluation of ivermectin as a potential treatment for mild to moderate COVID-19: a double-blind randomized placebo controlled trial in Eastern India. *J Pharm Pharm Sci.* 2021;24:343-350. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34265236.
- 37. Mohan A, Tiwari P, Suri TM, et al. Single-dose oral ivermectin in mild and moderate COVID-19 (RIVET-COV): a single-centre randomized, placebocontrolled trial. *J Infect Chemother*. 2021;27(12):1743-1749. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34483029.
- 38. Bermejo Galan LE, Dos Santos NM, Asato MS, et al. Phase 2 randomized study on chloroquine, hydroxychloroquine or ivermectin in hospitalized patients with severe manifestations of SARS-CoV-2 infection. *Pathog Glob Health*. 2021;115(4):235-242. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33682640</u>.

Metformin

Last Updated: December 20, 2023

Metformin has been identified as a potential COVID-19 therapeutic agent because of its possible action against the proteins that are involved in translation, its antiviral activity in vitro, and its anti-inflammatory and antithrombotic activities.¹⁻⁴ Data from observational studies have suggested that patients who were receiving metformin as treatment for diabetes at the time of their COVID-19 diagnosis had a lower risk of progressing to severe COVID-19.⁵⁻⁷ Randomized controlled trials have provided insight into the role of metformin in treating nonhospitalized patients with COVID-19. These trials are described below and in <u>Table 7c</u>.

Recommendations

- There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of metformin for the treatment of COVID-19 in nonhospitalized patients.
- The Panel **recommends against** the use of **metformin** for the treatment of COVID-19 in hospitalized patients, except in a clinical trial (**BIII**).
- Patients with COVID-19 who are receiving **metformin** for an underlying condition should continue this therapy as directed by their health care provider (**AIII**).

Rationale

Two randomized controlled trials (the TOGETHER and COVID-OUT trials) assessed the efficacy of using metformin in nonhospitalized patients with COVID-19. In these trials, the use of metformin did not reduce the risk of hospitalization or death in these patients. The Panel's recommendations are based on the results of these trials.

Other outpatient therapies (i.e., ritonavir-boosted nirmatrelvir [Paxlovid], remdesivir, molnupiravir) have been shown to be effective in preventing hospitalization or death in unvaccinated patients who are at high risk of disease progression.

Monitoring, Adverse Effects, and Drug-Drug Interactions

The most common adverse effects of metformin are nausea, vomiting, diarrhea, and headache. In rare cases, lactic acidosis may occur. The risk factors associated with lactic acidosis include older age, impaired renal or hepatic function, the use of iodinated contrast dye, cardiac dysfunction, metabolic disturbances, and excessive alcohol consumption. Metformin is not recommended for patients with an estimated glomerular filtration rate of <30 mL/min/1.73m².

Metformin is a substrate of the human organic cation transporters OCT1 and OCT2. Drugs that inhibit these transporters may increase the systemic exposure of metformin and increase the risk of metformin-related adverse effects.

Considerations in Pregnant People

Metformin is commonly used in pregnant people with type 2 diabetes mellitus. However, because clinical trials have not demonstrated a clear clinical benefit of using metformin in nonpregnant adults with COVID-19, there is no justification for administering it to pregnant people to treat COVID-19

outside of a clinical trial.

Considerations in Children

Although metformin is approved by the Food and Drug Administration for the treatment of type 2 diabetes mellitus in children aged >10 years, clinical trials that have evaluated its use for the treatment of COVID-19 have not included people aged <18 years. Given the lack of clear evidence of efficacy in adults, the Panel **recommends against** the use of **metformin** for the treatment of COVID-19 in pediatric patients, except in a clinical trial (**AIII**).

Clinical Data

TOGETHER Trial

The TOGETHER trial was a placebo-controlled platform clinical trial that was conducted in Brazil.⁸ The study enrolled nonhospitalized patients who had symptomatic SARS-CoV-2 infection for \leq 7 days, no history of COVID-19 vaccination, and an increased risk of progressing to severe disease. Patients were randomized to receive extended-release metformin 750 mg (n = 215) or placebo (n = 203) twice daily for 10 days.

The primary endpoint was a composite of retention in an emergency setting for >6 hours or hospitalization within 28 days of randomization. Secondary endpoints included viral clearance at Days 3 and 7, clinical improvement at Day 28, time to hospitalization or death, and the occurrence of adverse events. The study was stopped by the data and safety monitoring board for futility, as there was a low probability of demonstrating a difference between the study arms. Overall, there was no difference between the arms in the number of adverse events; however, the proportion of patients who experienced grade 3 events was higher in the metformin arm (9.8%) than in the placebo arm (4.4%).

COVID-OUT Trial

The COVID-OUT trial was a Phase 3, double-blind, placebo-controlled 2 x 3 factorial trial that evaluated the effectiveness of metformin, ivermectin, or fluvoxamine in patients with COVID-19.⁹ Patients were randomized to receive metformin or placebo in 1 factor and ivermectin, fluvoxamine, or placebo in the other factor. The study enrolled nonhospitalized adults within 3 days of a confirmed diagnosis of COVID-19 and \leq 7 days from symptom onset. Patients were aged 30 to 85 years and overweight. Those with stage 4 or 5 chronic kidney disease or an estimated glomerular filtration rate of <45 mL/min/1.73 m² were excluded. The metformin arm included those assigned to receive immediate-release oral metformin (titrated over several days to a final daily dose of 1,500 mg) alone or in combination with ivermectin or fluvoxamine.

The primary endpoint was a composite of development of hypoxemia (defined as oxygen saturation \leq 93%, as measured by a home pulse oximeter), emergency department visit, hospitalization, or death by Day 14. While this study was underway, the Food and Drug Administration raised concerns about the accuracy of home pulse oximeters. Approximately 50% of the patients received a primary COVID-19 vaccine series. The analyses showed no benefit for any of the 3 investigational agents in preventing the primary endpoint. In addition, the use of these agents did not lower the severity of COVID-19 symptoms over 14 days. A prespecified secondary analysis determined that, over 14 days of follow-up, those who received metformin had a lower risk of an emergency department visit, hospitalization, or death than those who did not receive metformin (adjusted OR 0.58; 95% CI, 0.35–0.94). A key secondary endpoint in the analysis was a composite of hospitalization or death by Day 28. Eight of 596 patients (1.3%) who received metformin met this endpoint compared with 19 of 601 patients (3.2%) who did not receive

metformin.

A secondary endpoint in the COVID-OUT trial assessed the impact of metformin on the development of long COVID. Since there is no standardized definition for long COVID, the endpoint was based on whether the patient had been given this diagnosis by a health care provider during the 10 months of follow-up. The study reported lower rates of long COVID in the metformin arm than in the control arm.¹⁰ However, providing treatment options for long COVID is beyond the scope of the Guidelines.

Although a secondary analysis of the COVID-OUT trial data demonstrated a benefit of metformin in patients with COVID-19, the results of the TOGETHER and COVID-OUT trials did not show a consistent benefit of metformin in these patients. Therefore, the Panel believes there is insufficient evidence to recommend either for or against the use of metformin for the treatment of COVID-19 in nonhospitalized patients. For more information on these trials, see <u>Table 7c</u>.

References

- 1. Karam BS, Morris RS, Bramante CT, et al. mTOR inhibition in COVID-19: a commentary and review of efficacy in RNA viruses. *J Med Virol*. 2021;93(4):1843-1846. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33314219</u>.
- 2. Del Campo JA, García-Valdecasas M, Gil-Gómez A, et al. Simvastatin and metformin inhibit cell growth in hepatitis C virus infected cells via mTOR increasing PTEN and autophagy. *PLoS One*. 2018;13(1):e0191805. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29385181</u>.
- 3. Postler TS, Peng V, Bhatt DM, Ghosh S. Metformin selectively dampens the acute inflammatory response through an AMPK-dependent mechanism. *Sci Rep.* 2021;11(1):18721. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34548527.
- 4. Xin G, Wei Z, Ji C, et al. Metformin uniquely prevents thrombosis by inhibiting platelet activation and mtDNA release. *Sci Rep.* 2016;6:36222. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27805009</u>.
- Li Y, Yang X, Yan P, Sun T, Zeng Z, Li S. Metformin in patients with COVID-19: a systematic review and meta-analysis. *Front Med (Lausanne)*. 2021;8:704666. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34490296</u>.
- Bramante CT, Buse J, Tamaritz L, et al. Outpatient metformin use is associated with reduced severity of COVID-19 disease in adults with overweight or obesity. *J Med Virol*. 2021;93(7):4273-4279. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33580540</u>.
- Luo P, Qiu L, Liu Y, et al. Metformin treatment was associated with decreased mortality in COVID-19 patients with diabetes in a retrospective analysis. *Am J Trop Med Hyg.* 2020;103(1):69-72. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32446312</u>.
- 8. Reis G, Dos Santos Moreira Silva EA, Medeiros Silva DC, et al. Effect of early treatment with metformin on risk of emergency care and hospitalization among patients with COVID-19: the TOGETHER randomized platform clinical trial. *Lancet Reg Health Am.* 2022;6:100142. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34927127.
- Bramante CT, Huling JD, Tignanelli CJ, et al. Randomized trial of metformin, ivermectin, and fluvoxamine for COVID-19. *N Engl J Med*. 2022;387(7):599-610. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36070710</u>.
- Bramante CT, Buse JB, Liebovitz DM, et al. Outpatient treatment of COVID-19 and incidence of post-COVID-19 condition over 10 months (COVID-OUT): a multicentre, randomised, quadruple-blind, parallel-group, Phase 3 trial. *Lancet Infect Dis.* 2023;23(10):1119-1129. Available at: <u>https://www.ncbi.nlm.</u> <u>nih.gov/pubmed/37302406</u>.

Table 7c. Metformin: Selected Clinical Trial Data

Last Updated: December 20, 2023

The Panel's recommendations for metformin are based on data from the clinical trials described in this table.

Methods	Results	Limitations and Interpretation			
TOGETHER: RCT of Metformin in Nonhospitalized Patients With COVID-19 in Brazil ¹					
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
 Aged ≥50 years or aged ≥18 years with ≥1 comorbidities 	 Median age 52 years; 57% women; 91% self- identified as mixed race 	 The >6-hour ED observation endpoint has not been used in other studies of interventions for 			
 Positive rapid antigen test result for SARS-CoV-2 infection 	 45% with BMI ≥30; 40% with HTN; 15% with DM 44% had COVID-19 symptoms for 0–3 days at 	nonhospitalized patients who are at high risk of hospitalization and death.			
 ≤7 days of COVID-19 symptoms 	enrollment	 Study was stopped early for futility. 			
Key Exclusion Criteria	Primary Outcome	Vaccinated individuals were excluded from trial.			
 Acute respiratory symptoms that required hospitalization 	• Study was stopped early by DSMB for futility. At the time the study was stopped, primary endpoint	InterpretationThis trial demonstrated no clinical benefit of			
Receipt of a COVID-19 vaccine	had occurred in 16% in metformin arm vs. 14% in	metformin in nonhospitalized patients with COVID-19.			
Interventions	placebo arm (relative risk 1.14; 95% Cl, 0.73–1.81; probability of superiority 28%).	 The use of metformin was associated with more 			
• Extended-release metformin 750 mg PO twice daily for 10 days (n = 215)	Secondary Outcomes	grade 3 AEs than placebo.			
 Placebo PO twice daily for 10 days (n = 203) 	No difference between arms in:				
Primary Endpoint	 Clinical improvement by Day 28 (OR 1.05; 95% Cl, 0.71–1.56) 				
 Composite of ED observation >6 hours or hospitalization for COVID-19 by Day 28 	 Viral clearance by Day 7 (OR 0.99; 95% Cl, 0.88–1.11) 				
Key Secondary Endpoints	• Time to hospitalization or death ($P = 0.53$)				
Clinical improvement by Day 28	Occurrence of treatment-emergent, grade 3 AEs:				
Viral clearance by Day 7	9.8% in metformin arm vs. 4.4% in placebo arm				
Time to hospitalization or death	(relative risk 2.11; 95% Cl, 1.05–4.61)Did not complete all phases of the study: 22% in				
Occurrence of AEs	metformin arm vs. 12% in placebo arm				
Study adherence	····				

Methods	Results	Limitations and Interpretation			
COVID-OUT: RCT of Metformin, Ivermectin, and Fluvoxamine in Nonhospitalized Adults With COVID-19 in the United States ²					
Key Inclusion Criteria	Participant Characteristics	Key Limitations			
 Aged 30–85 years BMI ≥25 or ≥23 if Asian or Latinx 	 Median age 46 years; 56% women; 82% White Median BMI 30 	 Analyses of secondary endpoints were not adjusted for multiple comparisons. 			
 Laboratory-confirmed SARS-CoV-2 infection within 3 days of randomization ≤7 days of COVID-19 symptoms 	 27% with CVD 52% received primary COVID-19 vaccination series Mean duration of symptoms was 4.8 days 	 Study included SpO₂ measurements using home pulse oximeters as 1 of the composite measures of the primary endpoint. However, the FDA has issued a statement concerning the accuracy of 			
Key Exclusion Criteria	 Approximately 66% enrolled while Delta was the 	these home pulse oximeters, making this study			
Immunocompromised	dominant variant; approximately 22% enrolled while	endpoint less reliable.			
Hepatic impairment	Omicron was dominant	Interpretation			
 Stage 4–5 chronic kidney disease or eGFR of <45 mL/min/1.73m² 	Primary OutcomesComposite of hypoxemia, ED visit, hospitalization, or	 The use of metformin did not prevent the occurrence of the primary composite endpoint of 			
Interventions	death by Day 14: 154 (24%) in metformin arm vs. 179	hypoxemia, ED visit, hospitalization, or death by			
 Immediate-release metformin 500 mg P0 on Day 1, 500 mg twice daily on Days 2–5, and 500 mg in morning and 1,000 mg in evening on Days 6–14 (n = 663) in the following arms: Metformin alone (n = 284) Metformin plus IVM 390–470 µg/kg P0 once daily for 3 days (n = 204) Metformin plus fluvoxamine 50 mg P0 twice daily for 14 days (n = 175) Control (n = 655), which included the following arms: 	 (27%) in control arm (aOR 0.84; 95% Cl, 0.66–1.09; P = 0.19) No difference between metformin alone arm and placebo alone arm in occurrence of primary endpoint (aOR 0.91; 95% Cl, 0.62–1.33) ED visit, hospitalization, or death by Day 14 in a prespecified secondary analysis: 27 (4.1%) in metformin arm vs. 48 (7.3%) in control arm (aOR 0.58; 95% Cl, 0.35–0.94) Hospitalization or death by Day 14 in a prespecified secondary analysis: 8 (1.2%) in metformin arm vs. 18 (2.7%) in control arm (aOR 0.47; 95% Cl, 0.20–1.11) 	 Day 14. Although the results of the prespecified secondary analyses of ED visits, hospitalization, or death by Day 14 and the secondary endpoint of hospitalization or death by Day 28 suggest a potential benefit of metformin, these results are not considered definitive. 			
 Placebo alone (n = 293) 	Secondary Outcomes				
• IVM or fluvoxamine alone (n = 362)	No difference between arms in total symptom severity				
Primary Endpoints	score by Day 14				
 Composite of hypoxemia (SpO₂ ≤93%, as measured by a home pulse oximeter), ED visit, hospitalization, or death by Day 14 	 Drug discontinuation or interruption: 29% in metformin arm vs. 25% in control arm Hospitalization or death by Day 28: 8 of 596 (1.3%) in 				
• A prespecified secondary analysis evaluated the occurrence of ED visits, hospitalization, or death by Day 14.	metformin arm vs. 19 of 601 (3.2%) in control arm				

Methods	Results	Limitations and Interpretation		
COVID-OUT: RCT of Metformin, Ivermectin, and Fluvoxamine in Nonhospitalized Adults With COVID-19 in the United States ² , continued				
Key Secondary Endpoints				
 Total symptom severity score by Day 14, as measured by a symptom severity scale 				
Drug discontinuation or interruption				
Hospitalization or death by Day 28				

Key: AE = adverse event; BMI = body mass index; CVD = cardiovascular disease; DM = diabetes mellitus; DSMB = data and safety monitoring board; ED = emergency department; eGFR = estimated glomerular filtration rate; FDA = Food and Drug Administration; HTN = hypertension; IVM = ivermectin; the Panel = the COVID-19 Treatment Guidelines Panel; PO = oral; RCT = randomized controlled trial; $SpO_2 =$ oxygen saturation

References

- Reis G, Dos Santos Moreira Silva EA, Medeiros Silva DC, et al. Effect of early treatment with metformin on risk of emergency care and hospitalization among patients with COVID-19: the TOGETHER randomized platform clinical trial. *Lancet Reg Health Am*. 2022;6:100142. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34927127</u>.
- 2. Bramante CT, Huling JD, Tignanelli CJ, et al. Randomized trial of metformin, ivermectin, and fluvoxamine for COVID-19. *N Engl J Med*. 2022;387(7):599-610. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36070710</u>.

Table 7d. Characteristics of Miscellaneous Drugs

Last Updated: December 20, 2023

- The information in this table is derived from data on the use of these drugs for FDA-approved indications or in investigational trials. It is supplemented with data on the use of these drugs in patients with COVID-19 or MIS-C, when available.
- For dose modifications for patients with organ failure or those who require extracorporeal devices, please refer to product labels, when available.
- There are currently not enough data to determine whether certain medications can be safely coadministered with therapies for the treatment of COVID-19. When using concomitant medications with similar toxicity profiles, consider performing additional safety monitoring.
- The potential additive, antagonistic, or synergistic effects and the safety of using certain combination therapies for the treatment of COVID-19 are unknown. Clinicians are encouraged to report AEs to the <u>FDA *MedWatch* program</u>.
- For drug-drug interaction information, please refer to product labels and visit the Liverpool COVID-19 Drug Interactions website.

Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments	
Fluvoxamine Not recommended by the Panel for the treatment of COVID-19 in nonhospitalized patients.					
 Doses for COVID-19 in Clinical Trials Fluvoxamine 50 mg P0 twice daily Fluvoxamine 100 mg P0 twice daily Fluvoxamine 100 mg P0 3 times daily 	 Nausea Diarrhea Dyspepsia Asthenia Insomnia Somnolence Sweating Suicidal ideation (rare) 	 Hepatic function Drug-drug interactions Withdrawal symptoms during dose tapering 	 CYP2D6 substrate Fluvoxamine inhibits several CYP isoenzymes (CYP1A2, CYP2C9, CYP3A4, CYP2C19, CYP2D6). Coadministration of tizanidine, thioridazine, alosetron, or pimozide with fluvoxamine is contraindicated. 	 Fluvoxamine may enhance the anticoagulant effects of antiplatelets and anticoagulants. Consider additional monitoring when these drugs are used concomitantly with fluvoxamine. The use of MAOIs concomitantly with fluvoxamine or within 14 days of treatment with fluvoxamine is contraindicated. 	

Dosing Regimens	Adverse Events	Monitoring Parameters	Drug-Drug Interaction Potential	Comments
Intravenous Immunoglobulin Primarily used for the treatment of MIS-C. Currently under investigation in clinical trials.				
 Dose for MIS-C 1 dose of IVIG 2 g/kg IBW IV, up to a maximum total dose of 100 g In the event of cardiac dysfunction or fluid overload, consider dividing the dose (IVIG 1 g/kg IBW/dose IV every 24 hours for 2 doses). 	 Allergic reactions, including anaphylaxis Renal failure Thromboembolic events Aseptic meningitis syndrome Hemolysis TRALI Transmission of infectious pathogens AEs may vary by formulation. Risk and severity of AEs may increase with high dose or rapid infusion. 	 Transfusion-related reactions Vital signs at baseline and during and after infusion Renal function; discontinue treatment if renal function deteriorates. 	 Not a CYP substrate; no drug- drug interactions expected 	 Rapid infusion should be avoided in patients with renal dysfunction or those who are at risk of thromboembolic events.
Metformin There is insufficient evidence for the Panel to recommend either for or against the use of metformin in nonhospitalized patients. Not recommended by the Panel for the treatment of COVID-19 in hospitalized patients, except in a clinical trial.				
 Doses for COVID-19 in Clinical Trials Immediate-release metformin 500 mg PO on Day 1; 500 mg twice daily on Days 2–5; and 500 mg in morning and 1,000 mg in evening on Days 6–14 Extended-release metformin 750 mg PO twice daily for 10 days 	 Diarrhea Nausea and vomiting Headache Lactic acidosis 	 Renal function Hepatic function Drug-drug interactions Alcohol use disorder 	 OCT1 and OCT2 substrate Drugs that interfere with OCT systems (e.g., cimetidine, dolutegravir, ranolazine, vandetanib) could increase systemic exposure to metformin. Concomitant use with carbonic anhydrase inhibitors (e.g., acetazolamide, topiramate, zonisamide) may increase the risk of lactic acidosis. 	• Alcohol intake may increase the risk of lactic acidosis.

Key: AE = adverse event; CYP = cytochrome P450; FDA = Food and Drug Administration; IBW = ideal body weight; IV = intravenous; IVIG = intravenous immunoglobulin; MAOI = monoamine oxidase inhibitor; MIS-C = multisystem inflammatory syndrome in children; OCT = organic cation transporter; the Panel = the COVID-19 Treatment Guidelines Panel; PO = oral; TRALI = transfusion-related acute lung injury

Supplements

Last Updated: December 20, 2023

Summary Recommendations

Vitamin C

- There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of vitamin C for the treatment of COVID-19 in nonhospitalized patients.
- The Panel recommends against the use of vitamin C for the treatment of COVID-19 in hospitalized patients (Alla).

Vitamin D

• There is insufficient evidence for the Panel to recommend either for or against the use of vitamin D for the prevention or treatment of COVID-19.

Zinc

- There is insufficient evidence for the Panel to recommend either for or against the use of zinc for the treatment of COVID-19.
- The Panel **recommends against** using zinc supplementation above the recommended dietary allowance (i.e., zinc 11 mg daily for men, zinc 8 mg daily for nonpregnant women) for the prevention of COVID-19, except in a clinical trial **(BIII)**.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Vitamin C

Last Updated: December 20, 2023

Vitamin C (ascorbic acid) is a water-soluble vitamin that has been considered for potential beneficial effects in patients with varying degrees of illness severity. It is an antioxidant and free radical scavenger that has anti-inflammatory properties, influences cellular immunity and vascular integrity, serves as a cofactor in endogenous catecholamine generation, and has been studied in many disease states, including COVID-19.^{1,2}

Recommendation for Nonhospitalized Patients With COVID-19

• There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of vitamin C for the treatment of COVID-19 in nonhospitalized patients.

Rationale

Because patients who are not critically ill with COVID-19 are less likely to experience oxidative stress or severe inflammation, the role of vitamin C in this setting is unknown.

Clinical Data for Nonhospitalized Patients With COVID-19

In an open-label trial conducted at 2 sites in the United States, outpatients with laboratory-confirmed SARS-CoV-2 infection were randomized to receive either 10 days of oral ascorbic acid 8,000 mg, zinc gluconate 50 mg, both agents, or standard of care.³ The primary endpoint was the number of days required to reach a 50% reduction in the patient's symptom severity score. The study was stopped early by an operational and safety monitoring board due to futility after 214 of the planned 520 participants were enrolled.

Patients who received standard of care achieved a 50% reduction in their symptom severity scores at a mean of 6.7 days (SD 4.4 days) compared with 5.5 days (SD 3.7 days) in the ascorbic acid arm, 5.9 days (SD 4.9 days) in the zinc gluconate arm, and 5.5 days (SD 3.4 days) in the arm that received both agents (overall P = 0.45).³ No serious adverse events related to the treatments were reported. Nonserious adverse events were experienced by 39.5% of patients in the ascorbic acid arm, 18.5% in the zinc gluconate arm, and 32.1% in the arm that received both agents, compared with 0% of patients in the standard of care arm (overall P < 0.001). The most common nonserious adverse effects in this study were gastrointestinal events.

The limitations of this study include the small sample size and the lack of a placebo control. In outpatients with COVID-19, treatment with high-dose zinc gluconate, ascorbic acid, or a combination of the 2 supplements, when compared with standard care, did not significantly decrease the number of days required to reach a 50% reduction in a symptom severity score.

Recommendation for Hospitalized Patients With COVID-19

• The Panel **recommends against** the use of vitamin C for the treatment of COVID-19 in hospitalized patients (**AIIa**).

Rationale

Randomized clinical trials have failed to demonstrate benefit from vitamin C as a therapeutic *COVID-19 Treatment Guidelines*

intervention for hospitalized patients with COVID-19. The data from these trials are summarized below.

Clinical Data for Hospitalized Patients With COVID-19

Two harmonized, randomized trials (LOVIT-COVID and REMAP-CAP) evaluated intravenous (IV) vitamin C versus a control in hospitalized patients with COVID-19 between July 2020 and July 2022.⁴ The studies enrolled patients from Asia, Australia, Europe, and North America, and data from the 2 studies were analyzed together. Patients in intensive care units who were critically ill and receiving organ support (1,568 patients from 90 sites) and hospitalized patients who were not critically ill (1,022 patients from 40 sites) were randomized to a vitamin C arm or a control arm. Patients in the intervention arm received IV vitamin C every 6 hours for 96 hours, for a maximum of 16 doses. Patients in the control arm received either no vitamin C or placebo. The composite primary outcome was a measure for days free of organ support up to 21 days and survival to hospital discharge. The study terminated enrollment after meeting criteria for harm and futility.

Among patients who were critically ill, the vitamin C arm (n = 1,037) had a median of 7 days free of organ support versus 10 days in the control arm (n = 531), with posterior probabilities of 8.6% for vitamin C efficacy and 99.9% for futility.⁴ Among patients who were not critically ill, both the vitamin C arm (n = 456) and the control arm (n = 566) had a median of 22 days free of organ support, with posterior probabilities of 2.9% for vitamin C efficacy and >99.9% for futility.

This study was limited by its use of combined data from 2 trials. The majority of patients enrolled were from an open-label study that used response-adaptive randomization.⁴ In addition, the precision of the treatment effect estimate in critically ill patients was limited because enrollment was stopped for harm. Data on individual vaccination status and the vitamin C product administered were unavailable. The study authors concluded that, in hospitalized patients with COVID-19, the probability that the use of vitamin C would increase the number of days free of organ support was low.

In a small, prospective, open-label randomized trial of hospitalized patients with severe COVID-19 in Pakistan, patients were randomized to receive vitamin C 50 mg/kg IV daily plus standard therapy (n = 75) or standard therapy alone (n = 75).⁵ Standard therapy included antipyretics, dexamethasone, and prophylactic antibiotics. Vitamin C recipients became symptom-free earlier (7.1 days vs. 9.6 days; P < 0.0001) and had a shorter duration of hospitalization (8.1 days vs. 10.7 days; P < 0.0001) than patients who received standard therapy alone. There were no significant differences between the arms for the outcomes of mortality and the need for mechanical ventilation. Limitations of this study include a small sample size, enrollment from only 1 hospital, and no clear method for recording symptoms.

In a pilot trial in China, 56 adults with COVID-19 who were in the intensive care unit were randomized to receive vitamin C 24 g IV daily for 7 days or placebo. The study was terminated early due to a reduction of cases of COVID-19 in China.⁶ Overall, the study found no differences between the arms for the outcomes of mortality, duration of mechanical ventilation, or change in median sequential organ failure assessment (SOFA) scores. The study reported improvements in oxygenation (as measured by the ratio of arterial partial pressure of oxygen to fraction of inspired oxygen) from baseline to Day 7 in the treatment arm that were statistically greater than those observed in the placebo arm (+20.0 vs. -51.9; P = 0.04).

In a randomized trial of 66 hospitalized patients with COVID-19 who required supplemental oxygen, treatment with vitamin C at doses escalating from 0.3 to 0.9 g/kg IV over 6 days (n = 44) was compared to standard of care (n = 22).⁷ The vitamin C did not improve the primary outcome of clinical status (defined as a composite of a 50% reduction in oxygen use, a 50% reduction in bronchodilator use, or

hospital discharge) at 72 hours after randomization.

Other Consideration

High concentrations of circulating vitamin C may affect the accuracy of point-of-care glucometers.^{8,9}

References

- 1. Fisher BJ, Seropian IM, Kraskauskas D, et al. Ascorbic acid attenuates lipopolysaccharide-induced acute lung injury. *Crit Care Med.* 2011;39(6):1454-1460. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21358394</u>.
- 2. Wei XB, Wang ZH, Liao XL, et al. Efficacy of vitamin C in patients with sepsis: an updated meta-analysis. *Eur J Pharmacol*. 2020;868:172889. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31870831</u>.
- 3. Thomas S, Patel D, Bittel B, et al. Effect of high-dose zinc and ascorbic acid supplementation vs usual care on symptom length and reduction among ambulatory patients with SARS-CoV-2 infection: the COVID A to Z randomized clinical trial. *JAMA Netw Open*. 2021;4(2):e210369. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33576820.
- LOVIT-COVID Investigators on behalf of the Canadian Critical Care Trials Group and the REMAP-CAP Investigators. Intravenous vitamin C for patients hospitalized with COVID-19: two harmonized randomized clinical trials. *JAMA*. 2023;330(18):1745-1759. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37877585</u>.
- 5. Kumari P, Dembra S, Dembra P, et al. The role of vitamin C as adjuvant therapy in COVID-19. *Cureus*. 2020;12(11):e11779. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33409026</u>.
- 6. Zhang J, Rao X, Li Y, et al. Pilot trial of high-dose vitamin C in critically ill COVID-19 patients. *Ann Intensive Care*. 2021;11(1):5. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33420963</u>.
- Coppock D, Violet PC, Vasquez G, et al. Pharmacologic ascorbic acid as early therapy for hospitalized patients with COVID-19: a randomized clinical trial. *Life (Basel)*. 2022;12(3):453. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35330204</u>.
- 8. Hager DN, Martin GS, Sevransky JE, Hooper MH. Glucometry when using vitamin C in sepsis: a note of caution. *Chest*. 2018;154(1):228-229. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30044741</u>.
- 9. Food and Drug Administration. Blood glucose monitoring devices. 2019. Available at: <u>https://www.fda.gov/</u> medical-devices/in-vitro-diagnostics/blood-glucose-monitoring-devices. Accessed November 21, 2023.

Vitamin D

Last Updated: December 20, 2023

Vitamin D is critical for bone and mineral metabolism. Because the vitamin D receptor is present on immune cells such as B cells, T cells, and antigen-presenting cells, and because these cells can synthesize the active vitamin D metabolite, vitamin D also has the potential to modulate innate and adaptive immune responses.¹ It is postulated that these immunomodulatory effects of vitamin D could potentially protect against SARS-CoV-2 infection or decrease the severity of COVID-19.

Vitamin D deficiency (defined as a serum concentration of 25-hydroxyvitamin D \leq 20 ng/mL) is common in the United States, particularly among people who identified as Hispanic or non-Hispanic Black.² These groups are overrepresented among cases of COVID-19 in the United States.³ Vitamin D deficiency is also more common in older patients and patients with obesity and hypertension; these factors have been associated with worse outcomes in patients with COVID-19.⁴ High levels of vitamin D may cause hypercalcemia and nephrocalcinosis.⁵

Recommendation

• There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of vitamin D for the prevention or treatment of COVID-19.

Rationale

The results from several cohort studies, clinical trials, and meta-analyses on the use of vitamin D for the prevention or treatment of COVID-19 have been published in peer-reviewed journals or have been made available as manuscripts ahead of peer review. However, most of these studies had significant limitations, such as small sample sizes or a lack of randomization and/or blinding. In addition, these studies used varying doses and formulations of vitamin D, enrolled participants with a range of COVID-19 severities, included different concomitant medications, and measured different study outcomes. All these factors make it difficult to compare results across studies. The studies summarized below are those that have had the greatest impact on the Panel's recommendations.

Although multiple observational cohort studies suggest that people with low vitamin D levels are at increased risk of SARS-CoV-2 infection and worse clinical outcomes after infection (e.g., higher mortality), clear evidence that vitamin D supplementation provides protection against infection or improves outcomes in patients with COVID-19 is still lacking.^{6,7}

Clinical Data on Vitamin D for Prevention

In a double-blind trial conducted at 4 hospitals in Mexico, frontline health care workers were randomized to receive vitamin D₃ 4,000 IU or placebo for 30 days.⁸ Participants were enrolled before COVID-19 vaccines became available. Over one-third of the enrolled participants dropped out before study completion. Of the 192 participants who completed follow-up, 6.4% of participants in the vitamin D₃ arm and 24.5% in the placebo arm acquired SARS-CoV-2 infection (relative risk 0.22; 95% CI, 0.08–0.59). At baseline, approximately 67% of participants had vitamin D deficiency, but this was not found to be an independent predictor of acquiring SARS-CoV-2 infection. The frequency of SARS-CoV-2 infection was very high in the placebo group, and it is unclear how these results translate to the use of vitamin D in vaccinated health care workers.

Clinical Data on Vitamin D for Treatment

In a double-blind trial conducted from June to October 2020 at 2 sites in Brazil, 240 hospitalized patients with moderate to severe COVID-19 were randomized to receive a single dose of vitamin D_3 200,000 IU or placebo.⁹ Patients were considered to have moderate to severe COVID-19 if they had a positive polymerase chain reaction (PCR) result for SARS-CoV-2 or compatible computed tomography scan findings and a respiratory rate >24 breaths/min or oxygen saturation <93% on room air. The primary outcome was length of hospital stay. The study found no significant difference in the median length of stay between the vitamin D_3 arm (7.0 days; IQR 4.0–10.0 days) and the placebo arm (7.0 days; IQR 5.0–13.0 days; log-rank *P*=0.59). No significant differences were observed between the arms in the proportion of patients who were admitted to the intensive care unit (ICU), the need for mechanical ventilation, or mortality. There were no significant safety concerns.

A randomized, double-blind, placebo-controlled study conducted in Argentina included 218 adult patients with COVID-19 who had been admitted to the hospital during the preceding 24 hours and who had oxygen saturation $\geq 90\%$ on room air and a risk factor for disease progression.¹⁰ Patients were randomized to receive a single oral dose of vitamin D₃ 500,000 IU or placebo. The primary outcome was the change in the respiratory sepsis-related organ failure assessment (rSOFA) score between baseline and the highest value recorded up to Day 7. There was no significant difference between the arms for this outcome, with a median change of 0 in both arms (P = 0.925). There were also no significant differences between the arms in the median length of hospital stay, the number of patients admitted to the ICU, or in-hospital mortality.

A randomized, open-label study conducted in France compared the effect of a high dose of vitamin D_3 (400,000 IU) to the standard dose of vitamin D_3 (50,000 IU) on mortality in 254 patients who were either hospitalized or living in nursing facilities near the study hospital sites.¹¹ Patients were aged \geq 65 years, had been diagnosed with SARS-CoV-2 infection within the preceding 3 days, and had at least 1 risk factor for disease progression (i.e., aged \geq 75 years, hypoxemia). Mortality was significantly different between the arms at 14 days, with 7 deaths (6%) among patients in the high-dose arm and 14 deaths (11%) among patients in the standard-dose arm (adjusted HR 0.33; 95% CI, 0.12–0.86; *P* = 0.02). However, mortality was not significantly different between the arms at 28 days (adjusted HR 0.70; 95% CI, 0.36–1.36; *P* = 0.29).

In an open-label pilot study, 50 hospitalized adults in New York with PCR-confirmed SARS-CoV-2 infection were randomized to receive calcitriol 0.5 μ g daily for 14 days or no treatment.¹² Calcitriol is the active metabolite of cholecalciferol or vitamin D₃ and is more commonly used to treat parathyroid disease. The study evaluated the change in oxygen saturation between patient admission and discharge. Additional outcomes were the length of hospital stay; mortality; and the need for endotracheal intubation, ICU admission, or hospital readmission within 30 days. Oxygen saturation was calculated using the ratio of peripheral oxygen saturation (measured by pulse oximetry) to fraction of inspired oxygen (SpO₂/FiO₂) as a surrogate for the ratio of arterial partial pressure of oxygen to fraction of inspired oxygen (PaO₂/FiO₂). Between admission and discharge, the patients who received no treatment had an average increase of 13.2 (SD 127.7) in the ratio, and those who received calcitriol had an increase of 91.04 (SD 119.08; *P* = 0.0305), implying an improvement in oxygenation.¹² There were no differences between the arms in the length of hospital stay, mortality, or the need for ICU admission or hospital readmission.

References

1. Aranow C. Vitamin D and the immune system. *J Investig Med*. 2011;59(6):881-886. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21527855</u>.

- Liu X, Baylin A, Levy PD. Vitamin D deficiency and insufficiency among US adults: prevalence, predictors and clinical implications. *Br J Nutr*. 2018;119(8):928-936. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/29644951</u>.
- 3. Forrest KY, Stuhldreher WL. Prevalence and correlates of vitamin D deficiency in US adults. *Nutr Res.* 2011;31(1):48-54. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21310306</u>.
- Centers for Disease Control and Prevention. People with certain medical conditions. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/need-extra-precautions/people-with-medical-conditions.html</u>. Accessed November 20, 2023.
- Committee to Review Dietary Reference Intakes for Vitamin D and Calcium, Food and Nutrition Board. Dietary Reference Intakes for Calcium and Vitamin D. National Academies Press; 2011. Available at: https://pubmed.ncbi.nlm.nih.gov/21796828.
- Chiodini I, Gatti D, Soranna D, et al. Vitamin D status and SARS-CoV-2 infection and COVID-19 clinical outcomes. *Front Public Health*. 2021;9:736665. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35004568</u>.
- Chen J, Mei K, Xie L, et al. Low vitamin D levels do not aggravate COVID-19 risk or death, and vitamin D supplementation does not improve outcomes in hospitalized patients with COVID-19: a meta-analysis and GRADE assessment of cohort studies and RCTs. *Nutr J*. 2021;20(1):89. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34719404</u>.
- 8. Villasis-Keever MA, López-Alarcón MG, Miranda-Novales G, et al. Efficacy and safety of vitamin D supplementation to prevent COVID-19 in frontline healthcare workers. A randomized clinical trial. *Arch Med Res*. 2022;53(4):423-430. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35487792</u>.
- Murai IH, Fernandes AL, Sales LP, et al. Effect of a single high dose of vitamin D₃ on hospital length of stay in patients with moderate to severe COVID-19: a randomized clinical trial. *JAMA*. 2021;325(11):1053-1060. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33595634</u>.
- Mariani J, Antonietti L, Tajer C, et al. High-dose vitamin D versus placebo to prevent complications in COVID-19 patients: multicentre randomized controlled clinical trial. *PLoS One*. 2022;17(5):e0267918. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35622854</u>.
- 11. Annweiler C, Beaudenon M, Gautier J, et al. High-dose versus standard-dose vitamin D supplementation in older adults with COVID-19 (COVIT-TRIAL): a multicenter, open-label, randomized controlled superiority trial. *PLoS Med*. 2022;19(5):e1003999. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35639792</u>.
- 12. Elamir YM, Amir H, Lim S, et al. A randomized pilot study using calcitriol in hospitalized COVID-19 patients. *Bone*. 2022;154:116175. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34508882.

Zinc

Last Updated: December 20, 2023

Increased intracellular zinc concentrations efficiently impair the replication of a number of RNA viruses.¹ Zinc has been shown to enhance cytotoxicity and induce apoptosis when used in vitro with a zinc ionophore (e.g., chloroquine). Chloroquine has also been shown to enhance intracellular zinc uptake in vitro.² Zinc levels are difficult to measure accurately, as zinc is distributed as a component of various proteins and nucleic acids.³

The recommended dietary allowance for elemental zinc is 11 mg daily for men and 8 mg daily for nonpregnant women.⁴ Long-term zinc supplementation can cause copper deficiency with subsequent reversible hematologic defects (i.e., anemia, leukopenia) and potentially irreversible neurologic manifestations (i.e., myelopathy, paresthesia, ataxia, spasticity).⁵⁻⁷ The use of zinc supplementation for durations as short as 10 months has been associated with copper deficiency.³ In addition, oral zinc can decrease the absorption of medications that bind with polyvalent cations (e.g., fluoroquinolones, HIV integrase inhibitors, tetracyclines).⁴

Recommendations

- There is insufficient evidence for the COVID-19 Treatment Guidelines Panel (the Panel) to recommend either for or against the use of zinc for the treatment of COVID-19.
- The Panel **recommends against** using zinc supplementation above the recommended dietary allowance (i.e., zinc 11 mg daily for men, zinc 8 mg daily for nonpregnant women) for the prevention of COVID-19, except in a clinical trial (**BIII**).

Rationale

The results from some cohort studies and clinical trials that evaluated the use of zinc in patients with COVID-19 have been published in peer-reviewed journals or have been made available as manuscripts ahead of peer review. However, most of these studies have significant limitations, such as small sample sizes or a lack of randomization or blinding. In addition, these studies used varying doses and formulations of zinc, enrolled participants with a range of COVID-19 severities, included different concomitant medications, and measured different study outcomes. All of these factors make it difficult to compare results across studies. Because zinc has not been shown to have a clinical benefit and may be harmful, the Panel **recommends against** using zinc supplementation above the recommended dietary allowance for the prevention of COVID-19, except in a clinical trial (**BIII**).

The studies summarized below are those that have had the greatest impact on the Panel's recommendations.

Clinical Data

In a double-blind, multicenter trial in Tunisia, nonhospitalized and hospitalized adults with COVID-19 were randomized within 7 days of symptom onset to receive elemental zinc 25 mg orally twice daily (n = 231) or matching placebo (n = 239) for 15 days.⁸ Approximately 20% of these patients had received a COVID-19 vaccine prior to enrollment. During the study, none of the patients received antiviral drugs, and <40% received corticosteroids.

The primary outcome in the study was a composite of death due to COVID-19 or intensive care unit

admission within 30 days of randomization.⁸ This study has several limitations. The study enrolled nonhospitalized and hospitalized patients, and comparing the results for these populations is difficult. In addition, only some patients received standard of care treatments. The data presented in the published paper had numerous and substantial inconsistencies.^{9,10} Together, these limitations make it difficult to interpret the results of this study or apply these findings to the current U.S. population with COVID-19.

In an open-label trial conducted at 2 sites in the United States, outpatients with laboratory-confirmed SARS-CoV-2 infection were randomized to receive either 10 days of zinc gluconate 50 mg, ascorbic acid 8,000 mg, both agents, or standard of care.¹¹ The primary endpoint was the number of days required to reach a 50% reduction in the patient's symptom severity score. The study was stopped early by an operational and safety monitoring board due to futility after 214 of the planned 520 participants were enrolled. Compared with standard of care, treatment with high-dose zinc gluconate, ascorbic acid, or a combination of the 2 supplements did not significantly decrease the number of days required to reach a 50% reduction in a symptom severity score. Patients who received standard of care achieved a 50% reduction in their symptom severity scores at a mean of 6.7 days (SD 4.4 days) compared with 5.5 days (SD 3.7 days) for the ascorbic acid arm, 5.9 days (SD 4.9 days) for the zinc gluconate arm, and 5.5 days (SD 3.4 days) for the arm that received both agents (overall P = 0.45).

Nonserious adverse effects occurred more frequently in patients who received supplements than in those who did not.¹¹ Nonserious adverse effects were experienced by 39.5% of patients in the ascorbic acid arm, 18.5% in the zinc gluconate arm, and 32.1% in the arm that received both agents, compared with 0% of patients in the standard of care arm (overall P < 0.001). The most common nonserious adverse effects in this study were gastrointestinal events.

In a randomized clinical trial conducted at 3 academic medical centers in Egypt, 191 patients with laboratory-confirmed SARS-CoV-2 infection were randomized to receive either zinc 220 mg twice daily plus hydroxychloroquine or hydroxychloroquine alone for a 5-day course.¹² The primary endpoints were recovery within 28 days, the need for mechanical ventilation, and death. The 2 arms were matched for age and gender. There were no significant differences between the arms in the percentages of patients who recovered within 28 days (79.2% in the zinc plus hydroxychloroquine arm vs. 77.9% in the hydroxychloroquine alone arm; P = 0.969), the number of patients who required mechanical ventilation (4 in the zinc plus hydroxychloroquine arm vs. 6 in the hydroxychloroquine alone arm; P = 0.537), or overall mortality (2 patients in each arm; P = 0.986). The only risk factors for mortality were age and the need for mechanical ventilation.

References

- 1. te Velthuis AJW, van den Worm SHE, Sims AC, et al. Zn(2+) inhibits coronavirus and arterivirus RNA polymerase activity in vitro and zinc ionophores block the replication of these viruses in cell culture. *PLoS Pathog.* 2010;6(11):e1001176. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21079686</u>.
- 2. Xue J, Moyer A, Peng B, et al. Chloroquine is a zinc ionophore. *PLoS One*. 2014;9(10):e109180. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/25271834</u>.
- 3. Hambridge K. The management of lipohypertrophy in diabetes care. *Br J Nurs*. 2007;16(9):520-524. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/17551441</u>.
- 4. Office of Dietary Supplements, National Institutes of Health. Zinc fact sheet for health professionals. 2022. Available at: <u>https://ods.od.nih.gov/factsheets/Zinc-HealthProfessional</u>. Accessed November 21, 2023.
- 5. Myint ZW, Oo TH, Thein KZ, Tun AM, Saeed H. Copper deficiency anemia: review article. *Ann Hematol*. 2018;97(9):1527-1534. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/29959467</u>.
- 6. Kumar N. Copper deficiency myelopathy (human swayback). *Mayo Clin Proc.* 2006;81(10):1371-1384. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/17036563</u>.

- Francis Z, Book G, Litvin C, Kalivas B. The COVID-19 pandemic and zinc-induced copper deficiency: an important link. *Am J Med.* 2022;135(8):e290-e291. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35367442</u>.
- 8. Abdallah SB, Mhalla Y, Trabelsi I, et al. Twice-daily oral zinc in the treatment of patients with coronavirus disease 2019: a randomized double-blind controlled trial. *Clin Infect Dis.* 2023;76(2):185-191. Available at: https://pubmed.ncbi.nlm.nih.gov/36367144.
- 9. Swindells S, Eschenauer GA, Nason M, Daar ES. Zinc and coronavirus disease 2019. *Clin Infect Dis*. 2023;77(4):662. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/37072885</u>.
- 10. Bel Haj Ali K, Sekma A, Trabelsi I, et al. Reply to Swindells et al. *Clin Infect Dis*. 2023;77(4):662-663. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/37073629</u>.
- 11. Thomas S, Patel D, Bittel B, et al. Effect of high-dose zinc and ascorbic acid supplementation vs usual care on symptom length and reduction among ambulatory patients with SARS-CoV-2 infection: the COVID A to Z randomized clinical trial. *JAMA Netw Open*. 2021;4(2):e210369. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33576820.
- 12. Abd-Elsalam S, Soliman S, Esmail ES, et al. Do zinc supplements enhance the clinical efficacy of hydroxychloroquine? A randomized, multicenter trial. *Biol Trace Elem Res*. 2021;199(10):3642-3646. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33247380</u>.

Considerations for Using Concomitant Medications in Patients With COVID-19

Last Updated: December 20, 2023

Summary Recommendations

- Patients with COVID-19 who are receiving concomitant medications (e.g., angiotensin-converting enzyme [ACE] inhibitors, angiotensin receptor blockers [ARBs], HMG-CoA reductase inhibitors [statins], systemic or inhaled corticosteroids, nonsteroidal anti-inflammatory drugs, acid-suppressive therapy) for underlying medical conditions should not discontinue ACE inhibitors and ARBs (Alla) or other medications (All) unless discontinuation is otherwise warranted by their clinical condition.
- The COVID-19 Treatment Guidelines Panel **recommends against** using medications off-label to treat COVID-19 if they have not been shown to be safe and effective for this indication in a clinical trial **(AIII)**.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Individuals with underlying medical conditions, such as cardiovascular disease, pulmonary disease, diabetes, and malignancy, and those who receive chronic immunosuppressive therapy are at higher risk of severe illness with COVID-19. These patients are often prescribed medications to treat their underlying medical conditions.

Early in the pandemic, some of these medications, such as angiotensin-converting enzyme (ACE) inhibitors, angiotensin receptor blockers (ARBs),¹ HMG-CoA reductase inhibitors (statins),^{2,3} and histamine-2 receptor antagonists,⁴ were hypothesized to offer potential as COVID-19 therapeutic agents. Others, such as nonsteroidal anti-inflammatory agents, were postulated to have negative impacts.⁵ Currently, there is no evidence that discontinuing medication for underlying medical conditions offers a clinical benefit for patients with COVID-19.⁶⁻⁸ For example, the Food and Drug Administration stated that there is no evidence linking the use of nonsteroidal anti-inflammatory agents with worsening of COVID-19 and advised patients to use them as directed.⁹ Additionally, the American Heart Association, the Heart Failure Society of America, and the American College of Cardiology issued a joint statement that renin-angiotensin-aldosterone system antagonists, such as ACE inhibitors and ARBs, should be continued as prescribed in those with COVID-19.¹⁰

Therefore, patients with COVID-19 who are treated with concomitant medications for an underlying medical condition **should not discontinue** ACE inhibitors and ARBs (**AIIa**) or other medications (**AIII**) unless discontinuation is otherwise warranted by their clinical condition. For patients with COVID-19 who require nebulized medications, precautions should be taken to minimize the potential for transmission of SARS-CoV-2 in the home and in health care settings.^{11,12}

The COVID-19 Treatment Guidelines Panel **recommends against** using medications off-label to treat COVID-19 if they have not been shown to be safe and effective for this indication in a clinical trial (**AIII**). Clinicians should refer to the <u>Therapies</u> section of the Guidelines for information on the medications that have been studied as potential therapeutic options for patients with COVID-19.

When prescribing medications to treat COVID-19, clinicians should always assess the patient's current medications for potential drug-drug interactions and additive adverse effects. The decision to continue or change a patient's medications should be individualized based on their specific clinical condition. Clinicians can refer to product labels and visit the <u>Liverpool COVID-19 Drug Interactions</u> website for guidance on identifying and managing drug-drug interactions. It is also worth noting that

ritonavir-boosted nirmatrelvir (Paxlovid), which is approved by the Food and Drug Administration for the treatment of mild to moderate COVID-19 in adults who are at high risk of progressing to severe COVID-19, has significant drug-drug interactions. See <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications</u> for more information.

References

- 1. Patel AB, Verma A. COVID-19 and angiotensin-converting enzyme inhibitors and angiotensin receptor blockers: what is the evidence? *JAMA*. 2020;323(18):1769-1770. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32208485</u>.
- 2. Lee KCH, Sewa DW, Phua GC. Potential role of statins in COVID-19. *Int J Infect Dis.* 2020;96:615-617. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32502659</u>.
- 3. Kashour T, Halwani R, Arabi YM, et al. Statins as an adjunctive therapy for COVID-19: the biological and clinical plausibility. *Immunopharmacol Immunotoxicol*. 2021;43(1):37-50. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33406943.
- 4. Mather JF, Seip RL, McKay RG. Impact of famotidine use on clinical outcomes of hospitalized patients with COVID-19. *Am J Gastroenterol*. 2020;115(10):1617-1623. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32852338.
- Yousefifard M, Zali A, Zarghi A, Madani Neishaboori A, Hosseini M, Safari S. Non-steroidal antiinflammatory drugs in management of COVID-19; a systematic review on current evidence. *Int J Clin Pract.* 2020;74(9):e13557. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32460369</u>.
- 6. Lopes RD, Macedo AVS, de Barros E Silva PGM, et al. Effect of discontinuing vs continuing angiotensinconverting enzyme inhibitors and angiotensin II receptor blockers on days alive and out of the hospital in patients admitted with COVID-19: a randomized clinical trial. *JAMA*. 2021;325(3):254-264. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33464336</u>.
- Cohen JB, Hanff TC, William P, et al. Continuation versus discontinuation of renin-angiotensin system inhibitors in patients admitted to hospital with COVID-19: a prospective, randomised, open-label trial. *Lancet Respir Med.* 2021;9(3):275-284. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33422263</u>.
- 8. Bauer A, Schreinlechner M, Sappler N, et al. Discontinuation versus continuation of renin-angiotensin-system inhibitors in COVID-19 (ACEI-COVID): a prospective, parallel group, randomised, controlled, open-label trial. *Lancet Respir Med.* 2021;9(8):863-872. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34126053</u>.
- 9. Food and Drug Administration. FDA advises patients on use of non-steroidal anti-inflammatory drugs (NSAIDs) for COVID-19. 2020. Available at: <u>https://www.fda.gov/drugs/drug-safety-and-availability/fda-advises-patients-use-non-steroidal-anti-inflammatory-drugs-nsaids-covid-19</u>. Accessed October 19, 2023.
- Bozkurt B, Kovacs R, Harrington B. Joint HFSA/ACC/AHA statement addresses concerns re: using RAAS antagonists in COVID-19. *J Card Fail*. 2020;26(5):370. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32439095</u>.
- 11. Cazzola M, Ora J, Bianco A, Rogliani P, Matera MG. Guidance on nebulization during the current COVID-19 pandemic. *Respir Med.* 2021;176:106236. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33248363</u>.
- 12. Sethi S, Barjaktarevic IZ, Tashkin DP. The use of nebulized pharmacotherapies during the COVID-19 pandemic. *Ther Adv Respir Dis.* 2020;14:1753466620954366. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33167796</u>.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Special Considerations in People Who Are Immunocompromised

Last Updated: November 2, 2023

Summary Recommendations

Prevention of COVID-19

- COVID-19 vaccination remains the most effective way to prevent serious outcomes and deaths from SARS-CoV-2 infection and should be considered the first line of prevention. The COVID-19 Treatment Guidelines Panel (the Panel) recommends COVID-19 vaccination for everyone who is eligible (AI), including those who are moderately or severely immunocompromised.
- Vaccine response rates may be lower in patients who are moderately or severely immunocompromised. Specific guidance on administering vaccines to these individuals is provided by the Centers for Disease Control and Prevention.
- All close contacts of people who are immunocompromised are strongly encouraged to stay up to date with COVID-19 vaccination (AI).
- There is insufficient evidence for the Panel to recommend either for or against the use of SARS-CoV-2 serologic testing to assess for immunity or to guide clinical decisions about using COVID-19 vaccines.

Management of Patients With COVID-19 Who Are Immunocompromised

- The Panel recommends consulting with the appropriate specialists when making decisions about stopping or adjusting the doses of immunosuppressive drugs in patients with COVID-19 (BIII).
- When selecting treatments for COVID-19, clinicians should consider factors such as the underlying disease; the specific immunosuppressants being used; the severity of COVID-19; and the potential for drug-drug interactions, overlapping toxicities, and secondary infections.
- For nonhospitalized patients with mild to moderate COVID-19 who are immunocompromised, the Panel recommends prompt treatment with antiviral drugs at the doses and durations recommended for the general population (AIII). For more information, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>.
- For most hospitalized patients with severe or critical COVID-19 who are immunocompromised, the Panel recommends using antiviral drugs and immunomodulatory therapies at the doses and durations recommended for the general population (AIII). For more information, see <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>.
- Some people who are immunocompromised have prolonged, symptomatic COVID-19 with evidence of ongoing SARS-CoV-2 replication. Without definitive data, some Panel members would use 1 or more of the following treatment options:
 - Longer and/or additional courses of ritonavir-boosted nirmatrelvir (Paxlovid)
 - · Longer and/or additional courses of remdesivir
 - High-titer COVID-19 convalescent plasma from a vaccinated donor who recently recovered from COVID-19 likely caused by a SARS-CoV-2 variant similar to the variant causing the patient's illness

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Introduction

Approximately 3% of people in the United States have immunocompromising conditions.¹ People who are immunocompromised are a heterogeneous population, and the severity of COVID-19 can vary significantly in this group. Some individuals who are immunocompromised may have a higher risk of hospitalization, complications, or death, and some may have outcomes that are comparable to those in the general population.

This section pertains to people who are moderately or severely immunocompromised, which includes those who:

- Are receiving active treatment for solid tumor and hematologic malignancies.
- Have hematologic malignancies (e.g., chronic lymphocytic lymphoma, non-Hodgkin lymphoma, multiple myeloma, acute leukemia) and are known to have poor responses to COVID-19 vaccines, regardless of the treatment status for the hematologic malignancy.
- Received a solid-organ or islet transplant and are receiving immunosuppressive therapy.
- Received chimeric antigen receptor T cell (CAR T-cell) therapy or a hematopoietic cell transplant (HCT) and are within 2 years of transplantation or are receiving immunosuppressive therapy.
- Have a moderate or severe primary immunodeficiency (e.g., severe combined immunodeficiency, DiGeorge syndrome, Wiskott-Aldrich syndrome, common variable immunodeficiency disease).
- Have advanced or untreated HIV infection (defined as people with HIV and CD4 T lymphocyte cell counts <200 cells/mm³, a history of an AIDS-defining illness without immune reconstitution, or clinical manifestations of symptomatic HIV).
- Are receiving active treatment with high-dose corticosteroids (i.e., ≥20 mg prednisone or equivalent per day for ≥2 weeks), alkylating agents, antimetabolites, transplant-related immunosuppressive drugs, cancer chemotherapeutic agents classified as severely immunosuppressive, or immunosuppressive or immunomodulatory biologic agents (e.g., B cell-depleting agents).

Analyses have found a higher risk of hospitalization or death from COVID-19 in those with a variety of immunocompromising conditions, including rheumatic diseases, hematological malignancies, solid organ transplants, and HIV.²⁻⁷ Factors that increase the risk of severe COVID-19 in the general population, such as older age, chronic kidney disease, cardiovascular disease, and other comorbidities, are significant drivers of risk in people who are immunocompromised. Moreover, some immunodeficiencies seem to increase the risk above and beyond traditional risk factors. For example, there is evidence that individuals who make autoantibodies to type I interferons (proteins that are critical to the protective immune response against viral infections) have a higher risk of severe COVID-19.⁸ Similarly, certain classes of medications, such as T cell–depleting or T cell–suppressing agents (e.g., antithymocyte globulin, calcineurin inhibitors, mycophenolate mofetil, belatacept) and B cell–depleting agents (e.g., rituximab, ocrelizumab, obinutuzumab), have been associated with more severe disease.^{9,10}

Prolonged shedding of SARS-CoV-2 has been reported in patients who are immunocompromised. A systematic review found that replication-competent virus could be detected for a median of 20 days in these patients, compared to 11 days in the general population.¹¹ Prolonged viral shedding may affect SARS-CoV-2 testing strategies and isolation durations for this group of patients. Moreover, case reports suggest that prolonged infections can create evolutionary pressure for the emergence of variants that resist therapies or evade vaccine-induced immunity.¹²⁻¹⁴

For any person who is eligible, clinicians should prescribe therapies for the treatment of COVID-19 as recommended in these Guidelines. However, at times during the pandemic, logistical constraints have limited the availability of therapies. In those cases, the COVID-19 Treatment Guidelines Panel (the Panel) suggests prioritizing the treatment of patients with COVID-19 who are at the highest risk of clinical progression (see Prioritization of Anti-SARS-CoV-2 Therapies for the Treatment of COVID-19 in Nonhospitalized Patients When There Are Logistical Constraints). Providers should use their clinical judgment when prioritizing patients for treatment and assess a patient's immunocompromised status, age, comorbidities, and vaccination status.

The sections below outline the Panel's rationale for the recommendations on preventing and managing COVID-19 in people who are immunocompromised. Some of the special considerations for patients

who are immunocompromised include the timing of COVID-19 vaccination, the management of immunosuppressive medications, and the strategies for treating COVID-19.

Prevention of COVID-19

Vaccination

COVID-19 vaccination remains the most effective way to prevent serious outcomes and death from SARS-CoV-2 infection. The Panel recommends COVID-19 vaccination for everyone who is eligible according to the guidance from the Centers for Disease Control and Prevention (CDC) (AI). This recommendation applies to:

- People who are moderately or severely immunocompromised
- People with active cancer and those receiving treatment for cancer
- Transplant and cellular immunotherapy candidates and recipients
- People with HIV
- All potential organ and hematopoietic cell donors
- Household members, close contacts, and health care workers who provide care for people who are immunocompromised

Authorized and approved COVID-19 vaccines in the United States are not live-virus vaccines and can be safely administered to patients who are immunocompromised. However, in people who are immunocompromised, the immune response to vaccination may be blunted, and the timing of vaccination requires special consideration. Nevertheless, vaccination is still recommended, as it may confer partial protection, including the protection provided by vaccine-induced, cell-mediated immunity.¹⁵

The Panel recommends following the current <u>COVID-19 vaccination guidance</u> from the CDC for people who are moderately or severely immunocompromised. This guidance includes information on the use of the updated 2023–2024 mRNA vaccines, which target the SARS-CoV-2 Omicron variant lineage XBB.1.5. The current CDC guidance also allows for the use of additional vaccine doses in people who are moderately or severely immunocompromised.¹⁶ There is a lack of data on the optimal timing for repeat vaccination in people who are immunocompromised, and the CDC recommends an interval of at least 2 months after the last dose. Other considerations may include the patient's current or expected level of immunosuppression, their age, comorbidities, and the time since their last vaccine dose. Clinicians should also take into account the prevalence of SARS-CoV-2 infection in the community and whether the patient intends to travel.

A preprint of a large observational study from Israel suggests a potential benefit of administering COVID-19 boosters every 6 months in groups with the highest risk of COVID-19–related hospitalization or death.¹⁷ The CDC-funded VISION Network evaluated the effectiveness of bivalent vaccines between September 13, 2022, and April 21, 2023, at 5 sites in 7 states.¹⁸ Among adults who were immunocompromised, a lower vaccine effectiveness (VE) was observed for the bivalent booster, but VE was sustained against critical COVID-19–associated outcomes, including intensive care unit admission and death. VE against hospitalization was 28% during the first 7 to 59 days after receipt of the bivalent dose and declined to 13% by 120 to 179 days; this indirectly supports using a 6-month interval for repeat vaccination.

The pivotal clinical trials that evaluated the safety and efficacy of the COVID-19 vaccines excluded people who were severely immunocompromised; therefore, the data for this population are less

robust.^{19,20} Observational data suggest that serological responses to vaccines may be blunted in patients who are immunocompromised.^{21,22} However, the MELODY trial reported detectable immunoglobulin G spike protein antibodies in approximately 80% of people in a large cohort of individuals in the United Kingdom who were immunocompromised and had received at least 3 doses of COVID-19 vaccines.²³ Those who had received anti-CD20 therapies within the past year were less likely than other groups in the study to have detectable anti-spike protein antibodies.

Vaccination of Close Contacts

Clinicians should strongly encourage all household members and close contacts of patients who are immunocompromised to be vaccinated against COVID-19 as soon as possible (**AI**). Before Omicron became the dominant circulating variant, a large cohort study of health care workers in Finland reported that COVID-19 vaccines were associated with a reduction in SARS-CoV-2 infections not only among vaccinated individuals but also among unvaccinated adult household members.²⁴ A 2022 systematic review and meta-analysis of 96 studies reported that people who received a complete primary COVID-19 vaccine series had reduced susceptibility and infectiousness. However, the vaccines were more effective against the Alpha variant than the Delta and Omicron variants.²⁵

Vaccination Timing and Immunosuppressive Therapies

If possible, COVID-19 vaccines should be administered at least 2 weeks before initiating or resuming immunosuppressive therapies. The timing of the vaccination should be determined based on the patient's current or planned immunosuppressive therapies, as well as the patient's medical condition and predicted response to the vaccine. Guidance about the timing of COVID-19 vaccination in solid organ transplant, HCT, and cellular immunotherapy candidates can be found in <u>Special Considerations</u> in <u>Solid Organ Transplant</u>, <u>Hematopoietic Cell Transplant</u>, and <u>Cellular Immunotherapy Candidates</u>, <u>Donors</u>, and <u>Recipients</u>. The CDC guidance allows the use of additional vaccine doses in people who are immunocompromised. Each additional dose should be administered at least 2 months after the last dose.

HCT and CAR T-cell recipients who received doses of COVID-19 vaccines prior to or during treatment with an HCT or CAR T-cell therapy should be revaccinated with the currently recommended primary vaccine series at least 3 months after the transplant or CAR T-cell therapy.²⁶ The American Society of Hematology has specific guidance about the timing of COVID-19 vaccination around cancer chemotherapy,²⁶ and the American College of Rheumatology also provides guidance for temporarily stopping immunosuppressive regimens during vaccination.²⁷

Polyethylene Glycol Allergies

The BNT162b2 (Pfizer-BioNTech) and mRNA-1273 (Moderna) mRNA bivalent vaccines contain polyethylene glycol (PEG), whereas the NVX-CoV2373 (Novavax) vaccine contains polysorbate 80. PEG and polysorbate are used in many products, including in agents used for cancer chemotherapy (e.g., PEG-asparaginase). PEG and polysorbate are structurally related, and cross-reactive hypersensitivity between these compounds might occur. The detection of PEG antibodies has not been shown to correlate with adverse reactions.²⁸ Therefore, testing for anti-PEG antibodies should not be used as a screening tool to assess the risk of allergic reactions²⁹ and should not replace an assessment by a specialist in those rare individuals with a history of anaphylaxis.³⁰ The <u>CDC has issued guidance</u> on triaging people with a history of allergic reactions to the components of COVID-19 vaccines.

Pre-Exposure Prophylaxis

Tixagevimab plus cilgavimab (Evusheld) is the only anti-SARS-CoV-2 monoclonal antibody (mAb) regimen that was shown to be effective for pre-exposure prophylaxis (PrEP) of COVID-19, and it was the only mAb regimen that was authorized by the Food and Drug Administration (FDA) for this use.

However, nearly all currently circulating Omicron subvariants in the United States are not susceptible to this combination. Therefore, tixagevimab plus cilgavimab is not currently authorized by the FDA for use as PrEP of COVID-19, and there are currently no other options for PrEP. The Panel **recommends against** the use of anti-SARS-CoV-2 mAbs such as **tixagevimab plus cilgavimab (Evusheld)** for PrEP of COVID-19 (**AIII**).

Serologic Testing to Guide Vaccination Strategies

There is insufficient evidence for the Panel to recommend either for or against the use of SARS-CoV-2 serologic testing to assess for immunity or to guide clinical decisions about using COVID-19 vaccines. More than 80 SARS-CoV-2 serologic tests, including quantitative, semiquantitative, neutralizing antibody, and point-of-care tests, have been issued Emergency Use Authorizations by the FDA to aid in detecting antibodies to SARS-CoV-2.³¹ However, these tests are not currently authorized for routine use in making individual medical decisions, and their ability to assess a person's level of immunity or protection from SARS-CoV-2 infection has not been evaluated.³² Most of these tests have not been calibrated to a reference standard, limiting the ability to compare and reproduce results from different tests.

Management of Patients With COVID-19 Who Are Immunocompromised

Adjusting Chronic Immunosuppressive Therapies

The Panel recommends consulting with the appropriate specialists when making decisions regarding stopping or adjusting the doses of immunosuppressive drugs in patients with COVID-19 (**BIII**). When selecting treatments for COVID-19, clinicians should consider factors such as the underlying disease; the specific immunosuppressants being used; the severity of COVID-19; and the potential for drug-drug interactions, overlapping toxicities, and secondary infections.

Early in the clinical course of COVID-19, the disease is primarily driven by the replication of SARS-CoV-2. Immunosuppressive medications can reduce the host immune responses that suppress viral replication, increasing the risk of prolonged viral shedding and infection.^{33,34} Clinicians should consider adjusting the doses of immunosuppressive medications or substituting certain immunosuppressive medications, if possible, to improve the patient's immune response to infection. When making decisions about stopping or reducing the dose of immunosuppressive drugs, clinicians should balance the potential benefit of enhancing the patient's immune response to COVID-19 with the risk of exacerbating the underlying condition. They should also consider the role of immunomodulation in the treatment of COVID-19.

Clinicians should be aware that many immunosuppressive drugs, particularly biologic agents, have long half-lives or prolonged periods of biologic activity. Patients may remain immunosuppressed long after the drugs are stopped. Care should be taken to not stop glucocorticoids abruptly, since this may result in adrenal insufficiency. For medications other than glucocorticoids, decisions about dose adjustments should be made on a case-by-case basis. For example, for some autoimmune diseases, temporary cessation of immunosuppression is often possible, and restarting medications 7 to 14 days after symptom resolution may be appropriate.^{27,35}

For solid organ transplant recipients, adjustments to immunosuppressive regimens should be individualized based on disease severity, the risk of graft rejection, the specific immunosuppressants being used, the type of transplant, the time since transplantation, the concentration of immunosuppressants, and the potential for drug-drug interactions.³⁶ See <u>Special Considerations in Solid</u> <u>Organ Transplant, Hematopoietic Cell Transplant, and Cellular Immunotherapy Candidates, Donors, and Recipients</u> for more information.

Therapeutic Management of Nonhospitalized Patients With COVID-19

For nonhospitalized patients with mild to moderate COVID-19 who are immunocompromised, the Panel recommends prompt treatment with antiviral drugs at the doses and durations recommended for the general population (AIII). See <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> to review the Panel's recommendations. Some special considerations for using these therapies in people who are immunocompromised are outlined below.

Ritonavir-Boosted Nirmatrelvir (Paxlovid)

In the EPIC-HR trial, the use of ritonavir-boosted nirmatrelvir reduced the risk of hospitalization or death when compared with placebo in nonhospitalized, unvaccinated adults who had laboratory-confirmed SARS-CoV-2 infection and a high risk of progressing to severe COVID-19.³⁷ Because the trial did not enroll many participants who were immunocompromised, the efficacy of ritonavir-boosted nirmatrelvir was not established for this population. In subsequent retrospective studies, some potential benefits of using ritonavir-boosted nirmatrelvir in people with various immunocompromising conditions have been observed.^{38,39}

Because ritonavir-boosted nirmatrelvir is the only highly effective oral antiviral therapy for COVID-19, it should be considered for patients who are immunocompromised if there are no potential drug-drug interactions or if the potential interactions can be safely managed. Clinicians should be aware of drug-drug interactions that may be life- or organ-threatening (see <u>Drug-Drug Interactions Between</u> <u>Ritonavir-Boosted Nirmatrelvir [Paxlovid] and Concomitant Medications</u>).⁴⁰ Notably, calcineurin inhibitors (e.g., tacrolimus, cyclosporine A) and mammalian target of rapamycin drugs (e.g., sirolimus, everolimus) have important drug-drug interactions with ritonavir. For this reason, the American Society of Transplantation recommends preferentially using other therapies, such as remdesivir, over ritonavir-boosted nirmatrelvir in people who are taking calcineurin inhibitors or mammalian target of rapamycin inhibitors.³⁶ Ritonavir can inhibit the metabolism of many cancer-directed therapies and should only be given after consulting with specialty pharmacists and other appropriate specialists.

Case reports have described reoccurring COVID-19 symptoms and positive SARS-CoV-2 test results in some patients who have completed treatment with ritonavir-boosted nirmatrelvir.⁴¹ A randomized trial is currently evaluating the effectiveness of longer courses or a second course of ritonavir-boosted nirmatrelvir (ClinicalTrials.gov Identifier NCT05438602). People with COVID-19 who are immunocompromised should not delay or avoid taking ritonavir-boosted nirmatrelvir due to concerns about the rebound of symptoms after treatment completion (see <u>Ritonavir-Boosted Nirmatrelvir</u> [Paxlovid]).

Remdesivir

Remdesivir was studied in nonhospitalized patients with mild to moderate COVID-19 who were at high risk of progressing to severe disease, and it was shown to be highly effective in reducing the risk of hospitalization and death.⁴² However, this trial only included a small number of participants who were immunocompromised. Because remdesivir treatment for nonhospitalized patients requires an intravenous infusion for 3 consecutive days, there may be logistical constraints to administering this drug in many settings. It can be considered for patients who are immunocompromised if other options, such as ritonavir-boosted nirmatrelvir, are not appropriate or available.

Molnupiravir

In the MOVe-OUT trial, molnupiravir reduced the rate of hospitalization or death when compared with placebo in nonhospitalized patients with COVID-19.⁴³ However, this trial only enrolled a small number of participants who were immunocompromised. The PANORAMIC trial enrolled a larger population of people who were immunocompromised, but this population was heterogeneous and the results of the

study were inconclusive.⁴⁴ Although the different treatment options have not been directly compared in clinical trials, the available evidence suggests that molnupiravir has a lower efficacy than the other options (see <u>Molnupiravir</u>). Other COVID-19 therapies should be prioritized over molnupiravir in patients who are immunocompromised.

COVID-19 Convalescent Plasma

There is insufficient evidence for the Panel to recommend either for or against the use of high-titer COVID-19 convalescent plasma (CCP) for the treatment of COVID-19 in nonhospitalized patients who are immunocompromised. The FDA issued an Emergency Use Authorization that allows the use of high-titer CCP for the treatment of COVID-19 in nonhospitalized or hospitalized patients who have immunosuppressive disease or are receiving immunosuppressive treatment.⁴⁵ However, the evidence generated from well-designed clinical trials that evaluated the use of CCP for the treatment of nonhospitalized patients is conflicting; these trials only enrolled a small number of patients who were immunocompromised.⁴⁶⁻⁴⁹

For a discussion on the potential use of high-titer CCP in patients who are immunocompromised and have prolonged viral replication, see the Therapeutic Management of Patients With COVID-19 Symptoms and Prolonged SARS-CoV-2 Replication section below.

Intravenous Immunoglobulin

Some individuals who are immunocompromised and have hypogammaglobulinemia are candidates for receiving supplemental antibodies in the form of intravenous immunoglobulin (IVIG) for the prevention of a variety of infections and in the setting of acute infections, including COVID-19. IVIG can be administered as outpatient or inpatient therapy. However, outside these specific circumstances, the Panel **recommends against** the use of **IVIG** for the prevention or treatment of acute COVID-19 in adults and children, except in a clinical trial (**AIII**). This recommendation should not preclude the use of IVIG when it is otherwise indicated for underlying conditions. See <u>Intravenous Immunoglobulin</u> for more information.

Therapeutic Management of Patients Who Are Hospitalized for COVID-19

For most hospitalized patients with severe or critical COVID-19 who are immunocompromised, the Panel recommends using antiviral drugs and immunomodulatory therapies at the doses and durations recommended for the general population (AIII). See <u>Therapeutic Management of Hospitalized</u> <u>Adults With COVID-19</u> for more information. The optimal management strategies and treatments for COVID-19 in hospitalized patients who are immunocompromised are unknown, since these individuals were either excluded from or poorly represented in major clinical trials. Nevertheless, clinical experience and retrospective data suggest that many patients who are immunocompromised have the expected responses to standard therapies for COVID-19.

Remdesivir

Case reports suggest that remdesivir can suppress, but does not always eliminate, viral replication in this population.^{50,51} In a large retrospective study of hospitalized patients who were immunocompromised, including patients who did not require supplemental oxygen, patients who received remdesivir had a lower risk of mortality at 14 days and 28 days than patients who did not receive remdesivir.⁵² The optimal duration of treatment with remdesivir in patients who are immunocompromised is unknown. Given the risk of prolonged viral replication in patients who are immunocompromised, some clinicians may choose to extend the course of antiviral therapy past 5 to 10 days. For patients receiving immunomodulatory therapy who have severe respiratory impairment due to COVID-19, clinicians may consider adding remdesivir treatment, although remdesivir has not been adequately studied in

prospective clinical trials to determine whether there is a benefit in these patients.

Corticosteroids

The RECOVERY trial reported a survival benefit for dexamethasone in inpatients with COVID-19 who were receiving oxygen, high-flow nasal cannula oxygen, noninvasive ventilation, or mechanical ventilation; however, specific data regarding the subgroup of patients who were immunocompromised are not available.⁵³ Unless otherwise indicated, corticosteroids should not be used for the treatment of COVID-19 in patients who are not receiving oxygen. In the RECOVERY trial, no survival benefit was observed for dexamethasone among the subset of patients with COVID-19 who did not require supplemental oxygen at enrollment. In an observational cohort study of U.S. veterans, the use of dexamethasone was associated with higher mortality in hospitalized patients with COVID-19 who did not require supplemental oxygen.

Patients who are immunocompromised may experience delayed development of favorable adaptive responses and a prolonged period of viral replication, as discussed above. For patients who are immunocompromised, are receiving minimal levels of conventional oxygen, and are earlier in the course of COVID-19 (e.g., those with <10 days of symptoms), the preferred approach may be emphasizing supportive care, using antiviral therapy, and avoiding corticosteroids. This strategy may reduce the duration of viral replication and the risk of secondary infections. Dexamethasone should be added if the patient has escalating oxygen requirements.

For patients who are immunocompromised and who were on chronic corticosteroids prior to hospitalization, the optimal dose of dexamethasone for the treatment of COVID-19 is unknown. The recommended dose of dexamethasone is 6 mg, which is equivalent to 40 mg of prednisone. This is the minimum dose of a steroid that should be used. Maintenance doses of corticosteroids should be discontinued while a patient is receiving dexamethasone, and the doses should be resumed as soon as possible after the patient recovers from COVID-19 or completes the course of dexamethasone.

Immunomodulators

Several randomized trials have shown that adding baricitinib or tocilizumab as a second immunomodulator to dexamethasone improves clinical outcomes in patients with severe or critical COVID-19.⁵⁵⁻⁵⁷ Another randomized trial that examined the use of infliximab, abatacept, or cenicriviroc in combination with dexamethasone in hospitalized adults with COVID-19 reported no differences between the study arms in the primary endpoint of time to recovery; however, patients who received infliximab or abatacept had a lower risk of mortality at 28 days.⁵⁸ These trials generally excluded patients who were immunocompromised or only included small numbers of these patients. For patients who are immunocompromised, the use of these agents may provide a clinical benefit similar to the benefit seen in the general population. However, it is not clear whether augmenting immunomodulation in this population increases the risk of serious bacterial, invasive fungal, or parasitic infections.

The Panel currently recommends adding another immunomodulator to dexamethasone in hospitalized patients with COVID-19 who are hypoxemic and experiencing clinical progression (see <u>Therapeutic</u> <u>Management of Hospitalized Adults With COVID-19</u>)</u>. This approach can also be used for most patients with COVID-19 who are immunocompromised. However, clinicians should consult with specialists to ensure that the risks of using additional immunosuppressive medications, including the risks of serious infections, do not outweigh the benefits. The patient should be closely monitored for infections.

COVID-19 Convalescent Plasma

There is insufficient evidence for the Panel to recommend either for or against the use of high-titer CCP for the treatment of COVID-19 in hospitalized patients who are immunocompromised. Three key randomized trials that evaluated the use of CCP for the treatment of COVID-19—RECOVERY, *COVID-19 Treatment Guidelines*

CONCOR-1, and REMAP-CAP—reported no evidence of a benefit of CCP in hospitalized patients with COVID-19. However, most of the patients enrolled in these trials were not immunocompromised.⁵⁹⁻⁶¹ Some of the subgroup analyses generated from clinical trials that enrolled patients who were immunocompromised suggested a potential benefit of CCP in this population,⁶¹⁻⁶³ but subgroup analyses need to be interpreted with caution (see <u>Table 4c</u>). Other clinical data from case series, retrospective case-control studies, and meta-analyses have been cited as support for the potential benefit of CCP in patients who are immunocompromised.⁶⁴⁻⁷² However, the patient populations differed across studies, and the studies had design limitations, making the findings difficult to interpret.

The RECOVER trial was a small, randomized trial that evaluated the use of plasma from donors who were convalescent and/or vaccinated against COVID-19 as a treatment for COVID-19 in hospitalized people with cancer, people with immunosuppression, people with lymphopenia and D-dimer levels $>1 \,\mu$ g/mL, and people aged >75 years. Only the subgroup of patients with cancer who received plasma treatment experienced a shorter median time to improvement and lower mortality when compared with the control arm.⁶²

For a discussion on the potential use of high-titer CCP in patients who are immunocompromised and have prolonged viral replication, see the Therapeutic Management of Patients With COVID-19 Symptoms and Prolonged SARS-CoV-2 Replication section below.

Therapeutic Management of Patients Who Are Hospitalized for Reasons Other Than COVID-19

People who are immunocompromised and have COVID-19 but were hospitalized for conditions other than COVID-19 should receive the same treatments as nonhospitalized patients (AIII). See <u>Therapeutic</u> <u>Management of Nonhospitalized Adults With COVID-19</u> for more information.

Therapeutic Management of Patients With COVID-19 Symptoms and Prolonged SARS-CoV-2 Replication

For patients who are immunocompromised and have prolonged COVID-19 symptoms and evidence of ongoing viral replication (e.g., those with a low cycle threshold value, as measured by a reverse transcription polymerase chain reaction result or with a positive rapid antigen test result) despite receiving a course of antiviral therapy, the optimal management is unknown. Case reports and case series have documented the treatment of these patients with additional antiviral treatments, prolonged courses of antiviral treatments, high-titer CCP, or combination therapy.⁷³⁻⁷⁷ The data for these approaches are not definitive, but some Panel members would use 1 or more of the following treatment options:

- Longer and/or additional courses of ritonavir-boosted nirmatrelvir
- Longer and/or additional courses of remdesivir
- High-titer CCP from a vaccinated donor who recently recovered from COVID-19 likely caused by a SARS-CoV-2 variant similar to the variant causing the patient's illness

Clinicians should be aware that the drug-drug interaction potential of ritonavir may change based on the duration of treatment. Even though cytochrome P450 (CYP) 3A4 inhibition by ritonavir is the primary concern when a 5-day course of ritonavir is used, clinicians should take into account that induction properties may become clinically relevant when ritonavir is used for 10 days or longer.⁷⁸

After discontinuing longer courses of ritonavir-boosted nirmatrelvir, drug-drug interactions due to CYP3A4 inhibition largely resolve within 2 to 3 days.⁷⁹ Drug-drug interactions that are caused by induction (e.g., CYP2C9, CYP2C19, uridine diphosphate-glucuronyltransferase) resolve gradually and variably.

Clinicians should consult experts (e.g., pharmacists and physicians with HIV expertise) for guidance on drug-drug interactions when using extended courses of ritonavir-boosted nirmatrelvir. For more information, see <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and</u> <u>Concomitant Medications</u>. The <u>Liverpool COVID-19 Drug Interactions website</u> provides guidance on managing drug-drug interactions in patients who are receiving for extended courses (i.e., ≥ 10 days) of ritonavir-boosted nirmatrelvir.

Considerations in Pregnant and Lactating People

Multiple studies have found that pregnant individuals have an increased risk of severe COVID-19 compared to age-matched controls, with increased rates of intensive care unit admission, mechanical ventilation, extracorporeal membrane oxygenation, and death.⁸⁰⁻⁸² Although hormonally mediated immunomodulation occurs during pregnancy, pregnancy is not a state of systemic immunosuppression. Changes in the immune response to certain infectious pathogens during pregnancy may increase the severity of respiratory illness in pregnant individuals. Physiologic changes, such as reduced pulmonary residual capacity, may also contribute to respiratory disease severity.⁸³⁻⁸⁶ Pregnant people who have underlying immunocompromising conditions or are receiving immunosuppressive medications likely have an even higher risk of severe disease. This patient group should be prioritized for the prevention and treatment of COVID-19.

Prevention

The Panel recommends COVID-19 vaccination as soon as possible for everyone who is eligible according to the CDC's Advisory Committee on Immunization Practices, including pregnant individuals (AI). COVID-19 vaccination is strongly recommended for pregnant individuals due to their increased risk for severe disease.^{87,88} Vaccination is especially important for pregnant people with concomitant risk factors such as underlying immunocompromising conditions (including those who are receiving immunosuppressive medications), as the risk for severe disease is likely additive.⁸¹

Treatment

Although pregnant patients have been excluded from the majority of the clinical trials that evaluated the use of COVID-19 therapeutics, pregnant patients with COVID-19 can be treated the same as nonpregnant patients, with a few exceptions. Pregnant patients who are immunocompromised or who have other risk factors likely have an even higher risk of severe COVID-19 and should be prioritized for treatment. Providers should refer to Pregnancy, Lactation, and COVID-19 Therapeutics for the Panel's guidance on treating COVID-19 in pregnant and lactating patients. Pregnant people who are immunocompromised comprise a heterogeneous group of patients, ranging from those who are mildly immunocompromised to those who are severely immunocompromised. Evaluating and managing pregnant patients require collaboration from a multidisciplinary team. This team should include a transplant or specialist, and a pharmacist.

Considerations in Children

Although the overall risk of critical illness and death related to COVID-19 is lower in children than adults, severe disease does occur, particularly in children with risk factors such as moderate to severe immunocompromising conditions. See <u>Special Considerations in Children</u> for a discussion of the risk factors for severe COVID-19 in children, and see <u>Therapeutic Management of Nonhospitalized</u> <u>Children With COVID-19</u> for the Panel's framework for assessing a child's risk of progression to severe COVID-19 based on vaccination status, comorbidities, and age.

Prevention

The Panel recommends COVID-19 vaccination as soon as possible for everyone who is eligible according to the CDC's Advisory Committee on Immunization Practices, including children (AI).

Treatment

The majority of children with mild to moderate COVID-19 will not progress to more severe illness; therefore, the Panel recommends managing these patients with supportive care alone (**AIII**). Few children, if any, have been enrolled in clinical trials of treatments for COVID-19. Among the children who were enrolled, very few were immunocompromised. Therefore, clinicians should be cautious when applying recommendations based on adult data to children. Clinicians need to consider the potential risks and benefits of therapy, the severity of the patient's disease, and underlying risk factors. See <u>Therapeutic Management of Hospitalized Children With COVID-19</u> and <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u> for the Panel's treatment recommendations in these scenarios.

References

- Wallace BI, Kenney B, Malani PN, et al. Prevalence of immunosuppressive drug use among commercially insured US adults, 2018–2019. *JAMA Netw Open*. 2021;4(5):e214920. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34014329</u>.
- 2. Vijenthira A, Gong IY, Fox TA, et al. Outcomes of patients with hematologic malignancies and COVID-19: a systematic review and meta-analysis of 3377 patients. *Blood.* 2020;136(25):2881-2892. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33113551.
- 3. Conway R, Grimshaw AA, Konig MF, et al. SARS-CoV-2 infection and COVID-19 outcomes in rheumatic diseases: a systematic literature review and meta-analysis. *Arthritis Rheumatol*. 2022;74(5):766-775. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34807517.
- 4. Song Q, Bates B, Shao YR, et al. Risk and outcome of breakthrough COVID-19 infections in vaccinated patients with cancer: real-world evidence from the National COVID Cohort Collaborative. *J Clin Oncol.* 2022;40(13):1414-1427. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35286152.
- 5. Ao G, Wang Y, Qi X, et al. The association between severe or death COVID-19 and solid organ transplantation: a systematic review and meta-analysis. *Transplant Rev (Orlando)*. 2021;35(3):100628. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34087553</u>.
- Wang Y, Feng R, Xu J, et al. An updated meta-analysis on the association between HIV infection and COVID-19 mortality. *AIDS*. 2021;35(11):1875-1878. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34397487</u>.
- MacKenna B, Kennedy NA, Mehrkar A, et al. Risk of severe COVID-19 outcomes associated with immunemediated inflammatory diseases and immune-modifying therapies: a nationwide cohort study in the OpenSAFELY platform. *Lancet Rheumatol*. 2022;4(7):e490-e506. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/35698725</u>.
- 8. Bastard P, Rosen LB, Zhang Q, et al. Autoantibodies against type I IFNs in patients with life-threatening COVID-19. *Science*. 2020;370(6515):eabd4585. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32972996</u>.
- Strangfeld A, Schäfer M, Gianfrancesco MA, et al. Factors associated with COVID-19-related death in people with rheumatic diseases: results from the COVID-19 Global Rheumatology Alliance physician-reported registry. *Ann Rheum Dis.* 2021;80(7):930-942. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33504483</u>.
- Sharifian-Dorche M, Sahraian MA, Fadda G, et al. COVID-19 and disease-modifying therapies in patients with demyelinating diseases of the central nervous system: a systematic review. *Mult Scler Relat Disord*. 2021;50:102800. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33578206</u>.

- 11. Qutub M, Aldabbagh Y, Mehdawi F, et al. Duration of viable SARS-CoV-2 shedding from respiratory tract in different human hosts and its impact on isolation discontinuation polices revision: a narrative review. *Clin Infect Pract.* 2022;13:100140. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35190799</u>.
- 12. Kemp SA, Collier DA, Datir RP, et al. SARS-CoV-2 evolution during treatment of chronic infection. *Nature*. 2021;592(7853):277-282. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33545711</u>.
- Corey L, Beyrer C, Cohen MS, et al. SARS-CoV-2 variants in patients with immunosuppression. N Engl J Med. 2021;385(6):562-566. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34347959</u>.
- Leung WF, Chorlton S, Tyson J, et al. COVID-19 in an immunocompromised host: persistent shedding of viable SARS-CoV-2 and emergence of multiple mutations: a case report. *Int J Infect Dis.* 2022;114:178-182. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34757008</u>.
- Ku JH, Sy LS, Qian L, et al. Vaccine effectiveness of the mRNA-1273 3-dose primary series against COVID-19 in an immunocompromised population: a prospective observational cohort study. *Vaccine*. 2023;41(24):3636-3646. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37173268</u>.
- Centers for Disease Control and Prevention. Interim clinical considerations for use of COVID-19 vaccines in the United States. 2023. Available at: <u>https://www.cdc.gov/vaccines/covid-19/clinical-considerations/interimconsiderations-us.html</u>. Accessed October 3, 2023.
- Yechezkel M, Samuel Faust J, Netzer D, et al. COVID-19 vaccine booster cadence by immunocompromised status. *medRxiv*. 2023;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2023.04.18.23288615v1</u>.
- Link-Gelles R, Weber ZA, Reese SE, et al. Estimates of bivalent mRNA vaccine durability in preventing COVID-19-associated hospitalization and critical illness among adults with and without immunocompromising conditions—VISION Network, September 2022–April 2023. MMWR Morb Mortal Wkly Rep. 2023;72(21):579-588. Available at: https://www.ncbi.nlm.nih.gov/pubmed/37227984.
- Baden LR, El Sahly HM, Essink B, et al. Efficacy and safety of the mRNA-1273 SARS-CoV-2 vaccine. N Engl J Med. 2021;384(5):403-416. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33378609</u>.
- 20. Polack FP, Thomas SJ, Kitchin N, et al. Safety and efficacy of the BNT162b2 mRNA COVID-19 vaccine. *N Engl J Med.* 2020;383(27):2603-2615. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33301246</u>.
- 21. Boyarsky BJ, Werbel WA, Avery RK, et al. Antibody response to 2-dose SARS-CoV-2 mRNA vaccine series in solid organ transplant recipients. *JAMA*. 2021;325(21):2204-2206. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33950155</u>.
- 22. Barrière J, Chamorey E, Adjtoutah Z, et al. Impaired immunogenicity of BNT162b2 anti-SARS-CoV-2 vaccine in patients treated for solid tumors. *Ann Oncol.* 2021;32(8):1053-1055. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33932508.
- 23. Pearce FA, Lim SH, Bythell M, et al. Antibody prevalence after three or more COVID-19 vaccine doses in individuals who are immunosuppressed in the UK: a cross-sectional study from MELODY. *Lancet Rheumatol*. 2023;5(8):e461-e473. Available at: <u>https://www.sciencedirect.com/science/article/pii/S2665991323001601</u>.
- 24. Salo J, Hägg M, Kortelainen M, et al. The indirect effect of mRNA-based COVID-19 vaccination on healthcare workers' unvaccinated household members. *Nat Commun.* 2022;13(1):1162. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35246536</u>.
- 25. Madewell ZJ, Yang Y, Longini IM Jr, Halloran ME, Dean NE. Household secondary attack rates of SARS-CoV-2 by variant and vaccination status: an updated systematic review and meta-analysis. *JAMA Netw Open*. 2022;5(4):e229317. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35482308</u>.
- 26. American Society of Hematology. ASH-ASTCT COVID-19 vaccination for HCT and CAR T cell recipients: frequently asked questions. 2022. Available at: <u>https://www.hematology.org/covid-19/ash-astct-covid-19-vaccination-for-hct-and-car-t-cell-recipients</u>. Accessed October 3, 2023.
- 27. American College of Rheumatology. COVID-19 vaccine clinical guidance summary for patients with rheumatic and musculoskeletal diseases. 2022. Available at: <u>https://www.rheumatology.org/Portals/0/Files/</u>

COVID-19-Vaccine-Clinical-Guidance-Rheumatic-Diseases-Summary.pdf.

- Carreño JM, Singh G, Tcheou J, et al. mRNA-1273 but not BNT162b2 induces antibodies against polyethylene glycol (PEG) contained in mRNA-based vaccine formulations. *Vaccine*. 2022;40(42):6114-6124. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36115801</u>.
- 29. Bavli Y, Chen BM, Gross G, et al. Anti-PEG antibodies before and after a first dose of Comirnaty (mRNA-LNP-based SARS-CoV-2 vaccine). *J Control Release*. 2023;354:316-322. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36549393</u>.
- 30. Brockow K, Mathes S, Fischer J, et al. Experience with polyethylene glycol allergy-guided risk management for COVID-19 vaccine anaphylaxis. *Allergy*. 2022;77(7):2200-2210. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34806775.
- 31. Food and Drug Administration. EUA authorized serology test performance. 2022. Available at: <u>https://www.fda.gov/medical-devices/coronavirus-disease-2019-covid-19-emergency-use-authorizations-medical-devices/eua-authorized-serology-test-performance</u>. Accessed October 3, 2023.
- 32. Food and Drug Administration. Antibody (serology) testing for COVID-19: information for patients and consumers. 2023. Available at: <u>https://www.fda.gov/medical-devices/coronavirus-covid-19-and-medical-devices/antibody-serology-testing-covid-19-information-patients-and-consumers</u>. Accessed October 3, 2023.
- 33. Aydillo T, Gonzalez-Reiche AS, Aslam S, et al. Shedding of viable SARS-CoV-2 after immunosuppressive therapy for cancer. *N Engl J Med.* 2020;383(26):2586-2588. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33259154</u>.
- 34. Tarhini H, Recoing A, Bridier-Nahmias A, et al. Long-term severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infectiousness among three immunocompromised patients: from prolonged viral shedding to SARS-CoV-2 superinfection. J Infect Dis. 2021;223(9):1522-1527. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33556961.
- 35. Rubin DT, Feuerstein JD, Wang AY, Cohen RD. AGA clinical practice update on management of inflammatory bowel disease during the COVID-19 pandemic: expert commentary. *Gastroenterology*. 2020;159(1):350-357. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32283100</u>.
- 36. American Society of Transplantation. COVID-19: FAQs for organ transplantation. 2023. Available at: <u>https://www.myast.org/sites/default/files/COVID%20FAQ%20for%20Tx%20professionals%202-2023%20FINAL.pdf</u>.
- 37. Hammond J, Leister-Tebbe H, Gardner A, et al. Oral nirmatrelvir for high-risk, nonhospitalized adults with COVID-19. N Engl J Med. 2022;386(15):1397-1408. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35172054.
- 38. Najjar-Debbiny R, Gronich N, Weber G, et al. Effectiveness of Paxlovid in reducing severe coronavirus disease 2019 and mortality in high-risk patients. *Clin Infect Dis*. 2023;76(3):e342-e349. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35653428</u>.
- 39. Qian G, Wang X, Patel NJ, et al. Outcomes with and without outpatient SARS-CoV-2 treatment for patients with COVID-19 and systemic autoimmune rheumatic diseases: a retrospective cohort study. *Lancet Rheumatol*. 2023;5(3):e139-e150. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36844970</u>.
- 40. University Health Network. Management of nirmatrelvir/ritonavir (Paxlovid) drug-drug interactions in oncology. 2022. Available at: <u>https://www.antimicrobialstewardship.com/paxlovid-ddi-oncology</u>.
- Charness ME, Gupta K, Stack G, et al. Rebound of SARS-CoV-2 infection after nirmatrelvir-ritonavir treatment. *N Engl J Med.* 2022;387(11):1045-1047. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36069968.</u>
- 42. Gottlieb RL, Vaca CE, Paredes R, et al. Early remdesivir to prevent progression to severe COVID-19 in outpatients. *N Engl J Med.* 2022;386(4):305-315. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34937145.
- 43. Jayk Bernal A, Gomes da Silva MM, Musungaie DB, et al. Molnupiravir for oral treatment of COVID-19 in

nonhospitalized patients. *N Engl J Med*. 2022;386(6):509-520. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34914868</u>.

- 44. Butler CC, Hobbs FDR, Gbinigie OA, et al. Molnupiravir plus usual care versus usual care alone as early treatment for adults with COVID-19 at increased risk of adverse outcomes (PANORAMIC): an open-label, platform-adaptive randomised controlled trial. *Lancet*. 2023;401(10373):281-293. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36566761.
- 45. Food and Drug Administration. Fact sheet for health care providers: Emergency Use Authorization (EUA) of COVID-19 convalescent plasma for treatment of coronavirus disease 2019 (COVID-19). 2021. Available at: https://www.fda.gov/media/141478/download.
- 46. Libster R, Pérez Marc G, Wappner D, et al. Early high-titer plasma therapy to prevent severe COVID-19 in older adults. *N Engl J Med*. 2021;384(7):610-618. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33406353.
- Korley FK, Durkalski-Mauldin V, Yeatts SD, et al. Early convalescent plasma for high-risk outpatients with COVID-19. *N Engl J Med*. 2021;385(21):1951-1960. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34407339</u>.
- 48. Alemany A, Millat-Martinez P, Corbacho-Monné M, et al. High-titre methylene blue-treated convalescent plasma as an early treatment for outpatients with COVID-19: a randomised, placebo-controlled trial. *Lancet Respir Med.* 2022;10(3):278-288. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35150610</u>.
- 49. Sullivan DJ, Gebo KA, Shoham S, et al. Early outpatient treatment for COVID-19 with convalescent plasma. *N Engl J Med.* 2022;386(18):1700-1711. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35353960</u>.
- 50. Helleberg M, Niemann CU, Moestrup KS, et al. Persistent COVID-19 in an immunocompromised patient temporarily responsive to two courses of remdesivir therapy. *J Infect Dis.* 2020;222(7):1103-1107. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32702095.
- 51. Gandhi S, Klein J, Robertson AJ, et al. De novo emergence of a remdesivir resistance mutation during treatment of persistent SARS-CoV-2 infection in an immunocompromised patient: a case report. *Nat Commun.* 2022;13(1):1547. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35301314</u>.
- 52. Mozaffari E, Chandak A, Gottlieb RL, et al. Remdesivir reduced mortality in immunocompromised patients hospitalized for COVID-19 across variant waves: findings from routine clinical practice. *Clin Infect Dis.* 2023;Published online ahead of print. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/37556727/</u>.
- 53. RECOVERY Collaborative Group. Dexamethasone in hospitalized patients with COVID-19. *N Engl J Med.* 2021;384(8):693-704. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32678530</u>.
- 54. Crothers K, DeFaccio R, Tate J, et al. Dexamethasone in hospitalised COVID-19 patients not on intensive respiratory support. *Eur Respir J*. 2022;60(1):2102532. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34824060.
- 55. Marconi VC, Ramanan AV, de Bono S, et al. Efficacy and safety of baricitinib for the treatment of hospitalised adults with COVID-19 (COV-BARRIER): a randomised, double-blind, parallel-group, placebo-controlled Phase 3 trial. *Lancet Respir Med*. 2021;9(12):1407-1418. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34480861</u>.
- 56. Kalil AC, Patterson TF, Mehta AK, et al. Baricitinib plus remdesivir for hospitalized adults with COVID-19. *N Engl J Med.* 2021;384(9):795-807. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33306283</u>.
- 57. RECOVERY Collaborative Group. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2021;397(10285):1637-1645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33933206</u>.
- 58. O'Halloran JA, Ko ER, Anstrom KJ, et al. Abatacept, cenicriviroc, or infliximab for treatment of adults hospitalized with COVID-19 pneumonia: a randomized clinical trial. *JAMA*. 2023;330(4):328-339. Available at: https://pubmed.ncbi.nlm.nih.gov/37428480.
- 59. RECOVERY Collaborative Group. Convalescent plasma in patients admitted to hospital with COVID-19

(RECOVERY): a randomised controlled, open-label, platform trial. *Lancet*. 2021;397(10289):2049-2059. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34000257</u>.

- 60. Bégin P, Callum J, Jamula E, et al. Convalescent plasma for hospitalized patients with COVID-19: an open-label, randomized controlled trial. *Nat Med.* 2021;27(11):2012-2024. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34504336.
- 61. Writing Committee for the REMAP-CAP Investigators. Effect of convalescent plasma on organ support-free days in critically ill patients with COVID-19: a randomized clinical trial. *JAMA*. 2021;326(17):1690-1702. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34606578</u>.
- 62. Denkinger CM, Janssen M, Schäkel U, et al. Anti-SARS-CoV-2 antibody-containing plasma improves outcome in patients with hematologic or solid cancer and severe COVID-19: a randomized clinical trial. *Nat Cancer*. 2023;4(1):96-107. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36581734</u>.
- 63. Lacombe K, Hueso T, Porcher R, et al. COVID-19 convalescent plasma to treat hospitalised COVID-19 patients with or without underlying immunodeficiency. *medRxiv*. 2022;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2022.08.09.22278329v2</u>.
- 64. Thompson MA, Henderson JP, Shah PK, et al. Association of convalescent plasma therapy with survival in patients with hematologic cancers and COVID-19. *JAMA Oncol.* 2021;7(8):1167-1175. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34137799.
- 65. Lanza F, Monaco F, Ciceri F, et al. Lack of efficacy of convalescent plasma in COVID-19 patients with concomitant hematological malignancies: an Italian retrospective study. *Hematol Oncol.* 2022;40(5):857-863. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35932208</u>.
- 66. Lang-Meli J, Fuchs J, Mathé P, et al. Case series: convalescent plasma therapy for patients with COVID-19 and primary antibody deficiency. *J Clin Immunol*. 2022;42(2):253-265. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34893946.
- 67. Rodionov RN, Biener A, Spieth P, et al. Potential benefit of convalescent plasma transfusions in immunocompromised patients with COVID-19. *Lancet Microbe*. 2021;2(4):e138. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33817676.
- Franchini M, Focosi D, Percivalle E, et al. Variant of concern-matched COVID-19 convalescent plasma usage in seronegative hospitalized patients. *Viruses*. 2022;14(7):1443. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35891421</u>.
- 69. Ljungquist O, Lundgren M, Iliachenko E, et al. Convalescent plasma treatment in severely immunosuppressed patients hospitalized with COVID-19: an observational study of 28 cases. *Infect Dis (Lond)*. 2022;54(4):283-291. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34878955</u>.
- 70. Ripoll JG, Gorman EK, Juskewitch JE, et al. Vaccine-boosted convalescent plasma therapy for patients with immunosuppression and COVID-19. *Blood Adv*. 2022;6(23):5951-5955. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36156121</u>.
- 71. Senefeld JW, Klassen SA, Ford SK, et al. Use of convalescent plasma in COVID-19 patients with immunosuppression. *Transfusion*. 2021;61(8):2503-2511. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34036587</u>.
- 72. Beraud M, Goodhue Meyer E, Lozano M, et al. Lessons learned from the use of convalescent plasma for the treatment of COVID-19 and specific considerations for immunocompromised patients. *Transfus Apher Sci.* 2022;61(3):103355. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35063360</u>.
- 73. Huygens S, Gharbharan A, Serroukh Y, et al. High-titer convalescent plasma plus nirmatrelvir/ritonavir treatment for non-resolving COVID-19 in six immunocompromised patients. *J Antimicrob Chemother*. 2023;78(7):1644-1648. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37248664</u>.
- 74. Brosh-Nissimov T, Ma'aravi N, Leshin-Carmel D, et al. Combination treatment of persistent COVID-19 in immunocompromised patients with remdesivir, nirmatrelvir/ritonavir and tixagevimab/cilgavimab. *medRxiv*. 2023;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2023.04.07.23288144v1</u>.

- 75. Mikulska M, Sepulcri C, Dentone C, et al. Triple combination therapy with two antivirals and monoclonal antibodies for persistent or relapsed SARS-CoV-2 infection in immunocompromised patients. *Clin Infect Dis*. 2023;77(2):280-286. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36976301</u>.
- 76. Graziani L, Gori L, Manciulli T, et al. Successful use of nirmatrelvir/ritonavir in immunocompromised patients with persistent and/or relapsing COVID-19. *J Antimicrob Chemother*. 2023;78(2):555-558. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36544352.
- 77. Trottier CA, Wong B, Kohli R, et al. Dual antiviral therapy for persistent coronavirus disease 2019 and associated organizing pneumonia in an immunocompromised host. *Clin Infect Dis*. 2023;76(5):923-925. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/36281907</u>.
- 78. Foisy MM, Yakiwchuk EM, Hughes CA. Induction effects on ritonavir: implications for drug interactions. *Ann Pharmacother*. 2008;42(7):1048-1059. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/18577765</u>.
- 79. Stader F, Khoo S, Stoeckle M, et al. Stopping lopinavir/ritonavir in COVID-19 patients: duration of the drug interacting effect. *J Antimicrob Chemother*. 2020;75(10):3084-3086. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32556272</u>.
- 80. Zambrano LD, Ellington S, Strid P, et al. Update: characteristics of symptomatic women of reproductive age with laboratory-confirmed SARS-CoV-2 infection by pregnancy status—United States, January 22–October 3, 2020. MMWR Morb Mortal Wkly Rep. 2020;69(44):1641-1647. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33151921</u>.
- Ellington S, Strid P, Tong VT, et al. Characteristics of women of reproductive age with laboratory-confirmed SARS-CoV-2 infection by pregnancy status—United States, January 22–June 7, 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(25):769-775. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32584795</u>.
- Allotey J, Stallings E, Bonet M, et al. Clinical manifestations, risk factors, and maternal and perinatal outcomes of coronavirus disease 2019 in pregnancy: living systematic review and meta-analysis. *BMJ*. 2020;370:m3320. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32873575</u>.
- Denney JM, Nelson EL, Wadhwa PD, et al. Longitudinal modulation of immune system cytokine profile during pregnancy. *Cytokine*. 2011;53(2):170-177. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21123081</u>.
- 84. Kourtis AP, Read JS, Jamieson DJ. Pregnancy and infection. *N Engl J Med.* 2014;370(23):2211-2218. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/24897084</u>.
- 85. Mosby LG, Rasmussen SA, Jamieson DJ. 2009 pandemic influenza A (H1N1) in pregnancy: a systematic review of the literature. *Am J Obstet Gynecol*. 2011;205(1):10-18. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/21345415</u>.
- 86. Siston AM, Rasmussen SA, Honein MA, et al. Pandemic 2009 influenza A(H1N1) virus illness among pregnant women in the United States. *JAMA*. 2010;303(15):1517-1525. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/20407061</u>.
- 87. Centers for Disease Control and Prevention. COVID-19 vaccines while pregnant or breastfeeding. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/vaccines/recommendations/pregnancy.html</u>. Accessed October 3, 2023.
- American College of Obstetricians and Gynecologists. COVID-19 vaccination considerations for obstetricgynecologic care. 2023. Available at: <u>https://www.acog.org/clinical/clinical-guidance/practice-advisory/</u> <u>articles/2020/12/covid-19-vaccination-considerations-for-obstetric-gynecologic-care</u>. Accessed October 3, 2023.

COVID-19 Treatment Guidelines

Special Considerations in Adults and Children With Cancer

Last Updated: November 2, 2023

Ourseen De commendations
Summary Recommendations
• COVID-19 vaccination remains the most effective way to prevent SARS-CoV-2 infection and should be considered the first line of prevention. The COVID-19 Treatment Guidelines Panel (the Panel) recommends COVID-19 vaccination as soon as possible for everyone who is eligible (AI) , including patients with active cancer and patients receiving treatment for cancer (AIII) .
• Because vaccine response rates may be lower in people with cancer, specific guidance on administering vaccines to these individuals is provided by the Centers for Disease Control and Prevention. For people with cancer, the Panel recommends following the most current <u>COVID-19 vaccination schedule</u> for people who are moderately or severely immunocompromised (AIII).
• Vaccinating household members, close contacts, and health care providers who provide care to patients with cancer is important to protect these patients from infection. All close contacts are strongly encouraged to get vaccinated against COVID-19 as soon as possible (AIII).
• The Panel recommends performing diagnostic molecular or antigen testing for SARS-CoV-2 in patients with cancer who develop signs and symptoms that suggest acute COVID-19 (AIII). The Panel also recommends performing diagnostic molecular testing in asymptomatic patients prior to procedures that require anesthesia and before initiating cytotoxic chemotherapy and long-acting biologic therapy (BIII).
• The recommendations for treating COVID-19 in patients with cancer are the same as those for the general population (AIII). See <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> and <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> for more information.
• Decisions about administering cancer-directed therapy to patients with acute COVID-19 and those who are recovering from COVID-19 should be made on a case-by-case basis; clinicians should consider the indication for chemotherapy, the goals of care, and the patient's history of tolerance to the treatment (BIII) .
 Clinicians who are treating COVID-19 in patients with cancer should consult a hematologist or oncologist before adjusting cancer-directed medications (AIII).
• Clinicians should pay careful attention to potential overlapping toxicities and drug-drug interactions between drugs used to treat COVID-19 (e.g., ritonavir-boosted nirmatrelvir [Paxlovid], dexamethasone) and cancer-directed therapies, prophylactic antimicrobials, and other medications (AIII).
Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.
People being treated for cancer may be at increased risk of severe COVID-19, and clinical outcomes of

People being treated for cancer may be at increased risk of severe COVID-19, and clinical outcomes of COVID-19 are generally worse in people with cancer than in people without cancer.¹⁻⁴ A meta-analysis of 46,499 patients with COVID-19 showed that all-cause mortality (risk ratio 1.66; 95% CI, 1.33–2.07) was higher in patients with cancer, and that patients with cancer were more likely to be admitted to intensive care units (risk ratio 1.56; 95% CI, 1.31–1.87).⁵ A patient's risk of immunosuppression and susceptibility to SARS-CoV-2 infection depend on the type of cancer, the treatments administered, and the stage of disease (e.g., patients actively being treated compared to those in remission). In a study that used data from the COVID-19 and Cancer Consortium Registry, patients with cancer who were in remission or who had no evidence of disease had a lower risk of death from COVID-19 than those who were receiving active treatment.⁶ It is unclear whether cancer survivors have an increased risk for severe COVID-19 and its complications when compared with people without a history of cancer.

This section of the COVID-19 Treatment Guidelines focuses on testing for SARS-CoV-2, managing COVID-19 in patients with cancer, and managing cancer-directed therapies during the COVID-19 *COVID-19 Treatment Guidelines*

pandemic. The optimal management and therapeutic approach to COVID-19 in this population has not yet been defined.

COVID-19 Vaccination in Patients With Cancer

The clinical trials that evaluated the COVID-19 vaccines that received Emergency Use Authorizations or approvals from the Food and Drug Administration (FDA) excluded severely immunocompromised patients. The COVID-19 vaccines authorized for use in the United States are not live vaccines; therefore, they can be safely administered to people who are immunocompromised.

Given the effectiveness of COVID-19 vaccines in the general population and the increased risk of severe COVID-19 and mortality in patients with cancer, the COVID-19 Treatment Guidelines Panel (the Panel) recommends COVID-19 vaccination as soon as possible for patients with active cancer and for patients receiving treatment for cancer (AIII).

For people with cancer, the Panel recommends following the most current <u>COVID-19 vaccination</u> <u>schedule</u> for people who are moderately or severely immunocompromised (**AIII**).

Observational data suggest that serological responses to vaccines may be blunted in patients who are immunocompromised.^{7,8} However, vaccination is still recommended for these patients because it may provide partial protection, including protection from vaccine-induced, cell-mediated immunity. See the Centers for Disease Control and Prevention (CDC) website <u>COVID-19 Vaccines for People Who Are</u> <u>Moderately or Severely Immunocompromised</u> for the current COVID-19 vaccination schedule for these individuals.

Vaccinating household members, close contacts, and health care providers who provide care to patients with cancer is important to protect these patients from infection. All close contacts are strongly encouraged to get vaccinated against COVID-19 as soon as possible (**AIII**). There is evidence that vaccinated individuals infected with SARS-CoV-2 have lower viral loads than unvaccinated individuals^{9,10} and that COVID-19 vaccines reduce the incidence of SARS-CoV-2 infections not only among vaccinated individuals but also among their household contacts.¹¹⁻¹³

When determining the timing of COVID-19 vaccination in patients with cancer, clinicians should consider the following factors:

- If possible, patients planning to receive chemotherapy should receive vaccinations for COVID-19 at least 2 weeks before starting chemotherapy.^{14,15}
- Hematopoietic cell and chimeric antigen receptor T cell recipients can be offered COVID-19 vaccination starting at least 3 months after therapy.¹⁵

It is unknown whether the immune response to COVID-19 vaccination can increase the risk of graft-versus-host disease. No immune-related adverse events were reported after COVID-19 vaccination in 2 studies of patients with cancer who received immune checkpoint inhibitors.^{16,17}

Decreased immunologic responses to COVID-19 vaccination have been reported in patients receiving treatment for solid tumors and hematologic malignancies.^{8,18} The type of therapy has been shown to influence the patient's response to vaccination. For example, people with chronic lymphocytic leukemia who were treated with Bruton's tyrosine kinase inhibitors or venetoclax with or without anti-CD20 antibodies had extremely low response rates (16.0% and 13.6%, respectively).¹⁸ In comparison, approximately 80% to 95% of patients with solid tumors showed immunologic responses.^{8,19,20} Several observational studies support the use of a third vaccine dose in patients with cancer, even though vaccine failure may still occur.²¹⁻²³ See the CDC website <u>COVID-19</u> Vaccines for People Who Are Moderately or

Severely Immunocompromised for guidance on vaccine dosing.

Polyethylene Glycol Allergies

Polyethylene glycol (PEG) and polysorbate are used in many products, including cancer treatments (e.g., PEG-asparaginase). The BNT162b2 (Pfizer-BioNTech) and mRNA-1273 (Moderna) mRNA vaccines contain PEG, and the NVX-CoV2373 (Novavax) vaccine contains polysorbate 80. PEG and polysorbate are structurally related, and cross-reactive hypersensitivity could occur.²⁴ These COVID-19 vaccines should not be given to individuals with a history of severe allergic reactions (e.g., anaphylaxis) to any component of COVID-19 vaccines, including PEG.

Testing for SARS-CoV-2 in Patients With Cancer

The Panel recommends performing diagnostic molecular or antigen testing for SARS-CoV-2 in patients with cancer who develop signs and symptoms that suggest acute COVID-19 (AIII).

Patients with cancer who are receiving chemotherapy are at risk of developing neutropenia. The National Comprehensive Cancer Network (NCCN) Guidelines for Hematopoietic Growth Factors categorizes cancer treatment regimens based on the patient's risk of developing neutropenia.²⁵ A retrospective study suggests that patients with cancer and neutropenia have a higher mortality rate if they develop COVID-19.²⁶ Studies have reported an increased risk of poor clinical outcomes for patients with COVID-19 in the setting of neutropenia and/or during the perioperative period.^{27,28} Because of this, the Panel recommends performing diagnostic molecular testing for SARS-CoV-2 in asymptomatic patients prior to procedures that require anesthesia and before initiating cytotoxic chemotherapy and long-acting biologic therapy (**BIII**).

General Guidance for Patients With Cancer

Patients with cancer frequently engage with the health care system to receive treatment and supportive care for cancer or treatment-related complications. Nosocomial transmission of SARS-CoV-2 to patients and health care workers has been reported.²⁹⁻³¹ Health care providers and patients should take precautions to reduce the risk of SARS-CoV-2 exposure and infection, including wearing a mask, maintaining a distance of 6 feet from others, and practicing good hand hygiene.³² Telemedicine can minimize the need for in-person services and reduce the risk of SARS-CoV-2 exposure. For medically or socially vulnerable populations, telemedicine may improve access to providers, but it could worsen disparities if these populations have limited access to technology.

Decisions about treatment regimens, surgery, and radiation therapy for the underlying malignancy should be made on a case-by-case basis, and clinicians should consider the biology of the cancer, the need for hospitalization, the number of clinic visits required, and the anticipated degree of immunosuppression. Additional factors that should be considered include the following:

- If possible, avoid treatment delays for curable cancers that have been shown to have worse outcomes when treatment is delayed (e.g., pediatric acute lymphoblastic leukemia).
- When the available treatment regimens are equally effective, regimens that can be administered orally or those that require fewer infusions are preferred.³³
- The potential risks of drug-related lung toxicity (e.g., from using bleomycin or PD-1 inhibitors) must be balanced with the clinical efficacy of alternative regimens or the risk of delaying care.³⁴
- Preventing neutropenia can decrease the risk of neutropenic fever and the need for emergency department evaluation and hospitalization. Granulocyte colony-stimulating factor (G-CSF) should be given with chemotherapy regimens that have an intermediate (10% to 20%) or high (>20%) risk of febrile neutropenia.³⁵

- Cancer treatment regimens that do not affect the outcomes of COVID-19 in patients with cancer may not need to be altered. In a prospective observational study, receipt of immunotherapy, hormonal therapy, or radiotherapy in the month prior to SARS-CoV-2 infection was not associated with an increased risk of mortality among patients with cancer and COVID-19.³⁶
- Radiation therapy guidelines suggest increasing the dose per fraction and reducing the number of daily treatments to minimize the number of hospital visits.³⁷

Febrile Neutropenia

Patients with cancer and febrile neutropenia should undergo diagnostic molecular or antigen testing for SARS-CoV-2 and evaluation for other infectious agents. They should also be given empiric antibiotics.³⁸ Low-risk febrile neutropenia patients should be treated at home with oral antibiotics or intravenous infusions of antibiotics to limit nosocomial exposure to SARS-CoV-2. Patients with high-risk febrile neutropenia should be hospitalized per standard of care. Empiric antibiotics should be continued per standard of care in patients who test positive for SARS-CoV-2. Clinicians should also continuously evaluate neutropenic patients for emergent infections.

Treating COVID-19 and Managing Chemotherapy in Patients With Cancer and COVID-19

Retrospective studies suggest that patients with cancer who were admitted to the hospital with SARS-CoV-2 infection have a high case-fatality rate, with higher rates observed in patients with hematologic malignancies than in those with solid tumors.^{39,40}

The recommendations for treating COVID-19 in patients with cancer are the same as those for the general population (AIII). See <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> and <u>Therapeutic Management of Hospitalized Adults With COVID-19</u> for more information. Patients with cancer are at high risk of progressing to severe COVID-19 and are eligible to receive anti-SARS-CoV-2 therapies in the outpatient setting if they develop mild to moderate COVID-19.

In patients with COVID-19 who required supplemental oxygen or mechanical ventilation, the use of dexamethasone has been associated with lower mortality than standard of care treatment alone.⁴¹ In patients with cancer, dexamethasone is commonly used to prevent chemotherapy-induced nausea, as a part of tumor-directed therapy, and to treat inflammation associated with brain metastasis. The side effects of dexamethasone are expected to be the same in patients with cancer as in those without cancer. If possible, treatments not currently recommended for SARS-CoV-2 infection should be administered as part of a clinical trial, since the safety and efficacy of these agents have not been well defined in patients with cancer.

Tocilizumab or baricitinib used in combination with dexamethasone is recommended for some patients with severe or critical COVID-19 who exhibit rapid respiratory decompensation (see <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>).⁴²⁻⁴⁴ The risks and benefits of using dexamethasone in combination with tocilizumab or baricitinib in patients with cancer who recently received chemotherapy is unknown. Because dexamethasone, tocilizumab, and baricitinib are immunosuppressive agents, patients who receive these medications should be closely monitored for secondary infections.

Therapeutic anticoagulation for patients with cancer who are hospitalized for COVID-19 should be managed similarly to anticoagulation for other hospitalized patients. Patients with platelet counts <50,000 cells/µL should not receive therapeutic anticoagulation to treat COVID-19. Clinicians should follow hospital protocols for managing anticoagulation in patients with thrombocytopenia.

The NCCN recommends against using G-CSF and granulocyte-macrophage colony-stimulating factor in patients with cancer and acute COVID-19 who do not have bacterial or fungal infections to avoid the hypothetical risk of increasing inflammatory cytokine levels and pulmonary inflammation.^{45,46} Secondary infections (e.g., invasive pulmonary aspergillosis) have been reported in critically ill patients with COVID-19.^{47,48}

Decisions about administering cancer-directed therapy to patients with acute COVID-19 and those who are recovering from COVID-19 should be made on a case-by-case basis; clinicians should consider the indication for chemotherapy, the goals of care, and the patient's history of tolerance to the treatment (**BIII**). The optimal time to initiate or restart cancer-directed therapies after the infection has resolved is unclear. If possible, clinicians should withhold treatment until COVID-19 symptoms have resolved. Prolonged viral shedding may occur in patients with cancer,² although it is unknown how this relates to infectious virus and how it impacts outcomes. The decision to restart cancer treatments in this setting should be made on a case-by-case basis. Clinicians who are treating COVID-19 in patients with cancer should consult a hematologist or oncologist before adjusting cancer-directed medications (**AIII**).

Medication Interactions

The use of antiviral or immune-based therapies to treat COVID-19 can present additional challenges in patients with cancer. Clinicians should pay careful attention to potential overlapping toxicities and drug-drug interactions between drugs used to treat COVID-19 (e.g., ritonavir-boosted nirmatrelvir [Paxlovid], dexamethasone) and cancer-directed therapies, prophylactic antimicrobials, and other medications (AIII).

A 5-day course of ritonavir-boosted nirmatrelvir is 1 of the preferred therapies for treating mild to moderate COVID-19 in nonhospitalized patients who are at risk for disease progression. However, this regimen has the potential for significant and complex drug-drug interactions with concomitant medications, primarily due to the ritonavir component of the combination. Boosting with ritonavir, a strong cytochrome P450 (CYP) 3A inhibitor, is required to increase the exposure of nirmatrelvir to a concentration that is effective against SARS-CoV-2. Ritonavir may also increase concentrations of certain concomitant medications, including certain chemotherapeutic agents and immunotherapies used to treat cancer. Significant increases in the concentrations of these drugs may lead to serious and sometimes life-threatening drug toxicities. Additionally, ritonavir is an inhibitor, inducer, and substrate of various other drug-metabolizing enzymes and drug transporters.

Before prescribing ritonavir-boosted nirmatrelvir, clinicians should carefully review the patient's concomitant medications. Clinicians should refer to resources such as the <u>Liverpool COVID-19 Drug</u> <u>Interactions website</u>, <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and</u> <u>Concomitant Medications</u>, and the FDA <u>prescribing information</u> for ritonavir-boosted nirmatrelvir for guidance on identifying and managing potential drug-drug interactions. If significant interactions prohibit the concomitant use of ritonavir-boosted nirmatrelvir, another COVID-19 treatment option should be used.

Dexamethasone is commonly used as an antiemetic for patients with cancer and is recommended for the treatment of certain patients with COVID-19 (see <u>Therapeutic Management of Hospitalized Adults With</u> <u>COVID-19</u>). Dexamethasone is a weak to moderate CYP3A4 inducer; therefore, interactions with any CYP3A4 substrates need to be considered.

Special Considerations in Children

Preliminary published reports suggest that pediatric patients with cancer may have milder manifestations of COVID-19 than adult patients with cancer, although larger studies are needed.⁴⁹⁻⁵¹ Guidance on

managing children with cancer during the COVID-19 pandemic is available from an international group that received input from the International Society of Paediatric Oncology, the Children's Oncology Group, St. Jude Global, and Childhood Cancer International.⁵² Two publications provide guidance on managing specific malignancies and supportive care and a summary of weblinks from groups of experts that are relevant to the care of pediatric oncology patients during the COVID-19 pandemic.^{52,53} Special considerations for using antiviral drugs in children who are immunocompromised, including those with malignancy, are available in a multicenter guidance statement.⁵⁴

References

- 1. Dai M, Liu D, Liu M, et al. Patients with cancer appear more vulnerable to SARS-CoV-2: a multicenter study during the COVID-19 outbreak. *Cancer Discov*. 2020;10(6):783-791. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32345594</u>.
- Shah V, Ko Ko T, Zuckerman M, et al. Poor outcome and prolonged persistence of SARS-CoV-2 RNA in COVID-19 patients with haematological malignancies; King's College Hospital experience. *Br J Haematol*. 2020;190(5):e279-e282. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32526039</u>.
- 3. Yang K, Sheng Y, Huang C, et al. Clinical characteristics, outcomes, and risk factors for mortality in patients with cancer and COVID-19 in Hubei, China: a multicentre, retrospective, cohort study. *Lancet Oncol.* 2020;21(7):904-913. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32479787</u>.
- 4. Robilotti EV, Babady NE, Mead PA, et al. Determinants of COVID-19 disease severity in patients with cancer. *Nat Med.* 2020;26(8):1218-1223. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32581323</u>.
- Giannakoulis VG, Papoutsi E, Siempos II. Effect of cancer on clinical outcomes of patients with COVID-19: a meta-analysis of patient data. *JCO Glob Oncol*. 2020;6:799-808. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32511066</u>.
- Kuderer NM, Choueiri TK, Shah DP, et al. Clinical impact of COVID-19 on patients with cancer (CCC19): a cohort study. *Lancet*. 2020;395(10241):1907-1918. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32473681</u>.
- Herzog Tzarfati K, Gutwein O, Apel A, et al. BNT162b2 COVID-19 vaccine is significantly less effective in patients with hematologic malignancies. *Am J Hematol.* 2021;96(10):1195-1203. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34185336</u>.
- Barrière J, Chamorey E, Adjtoutah Z, et al. Impaired immunogenicity of BNT162b2 anti-SARS-CoV-2 vaccine in patients treated for solid tumors. *Ann Oncol.* 2021;32(8):1053-1055. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33932508.
- Levine-Tiefenbrun M, Yelin I, Katz R, et al. Initial report of decreased SARS-CoV-2 viral load after inoculation with the BNT162b2 vaccine. *Nat Med.* 2021;27(5):790-792. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33782619</u>.
- Petter E, Mor O, Zuckerman N, et al. Initial real world evidence for lower viral load of individuals who have been vaccinated by BNT162b2. *medRxiv*. 2021;Preprint. Available at: <u>https://www.medrxiv.org/content/10.1101/2021.02.08.21251329v1</u>.
- 11. Salo J, Hägg M, Kortelainen M, et al. The indirect effect of mRNA-based COVID-19 vaccination on healthcare workers' unvaccinated household members. *Nat Commun.* 2022;13(1):1162. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35246536.
- de Gier B, Andeweg S, Backer JA, et al. Vaccine effectiveness against SARS-CoV-2 transmission to household contacts during dominance of Delta variant (B.1.617.2), the Netherlands, August to September 2021. *Euro Surveill*. 2021;26(44):2100977. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34738514</u>.
- Madewell ZJ, Yang Y, Longini IM Jr, Halloran ME, Dean NE. Household secondary attack rates of SARS-CoV-2 by variant and vaccination status: an updated systematic review and meta-analysis. *JAMA Netw Open.* 2022;5(4):e229317. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35482308</u>.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

- American Society of Hematology. General principles of COVID-19 vaccines for immunocompromised patients. 2022. Available at: <u>https://www.hematology.org/covid-19/ash-astct-covid-19-and-vaccines</u>. Accessed May 30, 2023.
- 15. American Society of Hematology. ASH-ASTCT COVID-19 vaccination for HCT and CAR T cell recipients: frequently asked questions. 2022. Available at: <u>https://www.hematology.org/covid-19/ash-astct-covid-19-vaccination-for-hct-and-car-t-cell-recipients</u>. Accessed May 30, 2023.
- Chen YW, Tucker MD, Beckermann KE, et al. COVID-19 mRNA vaccines and immune-related adverse events in cancer patients treated with immune checkpoint inhibitors. *Eur J Cancer*. 2021;155:291-293. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34400057</u>.
- Waissengrin B, Agbarya A, Safadi E, Padova H, Wolf I. Short-term safety of the BNT162b2 mRNA COVID-19 vaccine in patients with cancer treated with immune checkpoint inhibitors. *Lancet Oncol.* 2021;22(5):581-583. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33812495</u>.
- Herishanu Y, Avivi I, Aharon A, et al. Efficacy of the BNT162b2 mRNA COVID-19 vaccine in patients with chronic lymphocytic leukemia. *Blood.* 2021;137(23):3165-3173. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/33861303/</u>.
- Massarweh A, Eliakim-Raz N, Stemmer A, et al. Evaluation of seropositivity following BNT162b2 messenger RNA vaccination for SARS-CoV-2 in patients undergoing treatment for cancer. *JAMA Oncol.* 2021;7(8):1133-1140. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34047765</u>.
- 20. Shroff RT, Chalasani P, Wei R, et al. Immune responses to two and three doses of the BNT162b2 mRNA vaccine in adults with solid tumors. *Nat Med.* 2021;27(11):2002-2011. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34594036.
- 21. Mair MJ, Berger JM, Mitterer M, et al. Third dose of SARS-CoV-2 vaccination in hemato-oncological patients and health care workers: immune responses and adverse events—a retrospective cohort study. *Eur J Cancer*. 2022;165:184-194. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35248840</u>.
- 22. Abid MB, Rubin M, Ledeboer N, et al. Efficacy of a third SARS-CoV-2 mRNA vaccine dose among hematopoietic cell transplantation, CAR T cell, and BiTE recipients. *Cancer Cell*. 2022;40(4):340-342. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35202585</u>.
- 23. Re D, Seitz-Polski B, Brglez V, et al. Humoral and cellular responses after a third dose of SARS-CoV-2 BNT162b2 vaccine in patients with lymphoid malignancies. *Nat Commun.* 2022;13(1):864. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35165284</u>.
- 24. Centers for Disease Control and Prevention. Interim clinical considerations for use of COVID-19 vaccines: appendices, references, and previous updates. 2023. Available at: <u>https://www.cdc.gov/vaccines/covid-19/</u> <u>clinical-considerations/interim-considerations-us-appendix.html</u>. Accessed May 30, 2023.
- 25. Becker PS, Griffiths EA, Alwan LM, et al. NCCN guidelines insights: hematopoietic growth factors, version 1.2020. J Natl Compr Canc Netw. 2020;18(1):12-22. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31910384</u>.
- 26. Yarza R, Bover M, Paredes D, et al. SARS-CoV-2 infection in cancer patients undergoing active treatment: analysis of clinical features and predictive factors for severe respiratory failure and death. *Eur J Cancer*. 2020;135:242-250. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32586724.
- 27. American Society of Clinical Oncology. ASCO special report: a guide to cancer care delivery during the COVID-19 pandemic. 2021. Available at: <u>https://www.asco.org/sites/new-www.asco.org/files/content-files/2020-ASCO-Guide-Cancer-COVID19.pdf</u>.
- 28. American Society of Anesthesiologists. The ASA and APSF joint statement on perioperative testing for the COVID-19 virus. 2020. Available at: <u>https://www.asahq.org/about-asa/newsroom/news-releases/2020/06/asa-and-apsf-joint-statement-on-perioperative-testing-for-the-covid-19-virus</u>. Accessed May 30, 2023.
- 29. Wang X, Zhou Q, He Y, et al. Nosocomial outbreak of COVID-19 pneumonia in Wuhan, China. *Eur Respir J*. 2020;55(6):2000544. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32366488</u>.

- Luong-Nguyen M, Hermand H, Abdalla S, et al. Nosocomial infection with SARS-CoV-2 within departments of digestive surgery. *J Visc Surg.* 2020;157(3S1):S13-S18. Available at: <u>https://www.ncbi.nlm.nih.gov/ pubmed/32381426</u>.
- Rivett L, Sridhar S, Sparkes D, et al. Screening of healthcare workers for SARS-CoV-2 highlights the role of asymptomatic carriage in COVID-19 transmission. *Elife*. 2020;9:e58728. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32392129</u>.
- 32. Centers for Disease Control and Prevention. How to protect yourself and others. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/prevent-getting-sick/prevention.html</u>. Accessed May 30, 2023.
- 33. American Society of Clinical Oncology. Cancer treatment and supportive care. 2022. Available at: <u>https://old-prod.asco.org/covid-resources/patient-care-info/cancer-treatment-supportive-care</u>. Accessed May 30, 2023.
- 34. American Society of Hematology. COVID-19 and Hodgkin lymphoma: frequently asked questions. 2022. Available at: <u>https://www.hematology.org/covid-19/covid-19-and-hodgkin-lymphoma</u>. Accessed May 30, 2023.
- 35. Griffiths EA, Alwan LM, Bachiashvili K, et al. Considerations for use of hematopoietic growth factors in patients with cancer related to the COVID-19 pandemic. *J Natl Compr Canc Netw.* 2020;19(13):18-21. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32871558</u>.
- 36. Lee LY, Cazier JB, Angelis V, et al. COVID-19 mortality in patients with cancer on chemotherapy or other anticancer treatments: a prospective cohort study. *Lancet*. 2020;395(10241):1919-1926. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32473682</u>.
- 37. Yahalom J, Dabaja BS, Ricardi U, et al. ILROG emergency guidelines for radiation therapy of hematological malignancies during the COVID-19 pandemic. *Blood.* 2020;135(21):1829-1832. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32275740</u>.
- 38. Zimmer AJ, Freifeld AG. Optimal management of neutropenic fever in patients with cancer. *J Oncol Pract.* 2019;15(1):19-24. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30629902</u>.
- 39. Mehta V, Goel S, Kabarriti R, et al. Case fatality rate of cancer patients with COVID-19 in a New York hospital system. *Cancer Discov*. 2020;10(7):935-941. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32357994</u>.
- 40. Meng Y, Lu W, Guo E, et al. Cancer history is an independent risk factor for mortality in hospitalized COVID-19 patients: a propensity score-matched analysis. *J Hematol Oncol*. 2020;13(1):75. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32522278</u>.
- 41. RECOVERY Collaborative Group. Dexamethasone in hospitalized patients with COVID-19. *N Engl J Med.* 2021;384(8):693-704. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32678530</u>.
- 42. RECOVERY Collaborative Group. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet.* 2021;397(10285):1637-1645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33933206</u>.
- 43. REMAP-CAP Investigators. Interleukin-6 receptor antagonists in critically ill patients with COVID-19. *N Engl J Med*. 2021;384(16):1491-1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631065</u>.
- 44. Kalil AC, Patterson TF, Mehta AK, et al. Baricitinib plus remdesivir for hospitalized adults with COVID-19. *N Engl J Med.* 2021;384(9):795-807. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33306283</u>.
- 45. Nawar T, Morjaria S, Kaltsas A, et al. Granulocyte-colony stimulating factor in COVID-19: is it stimulating more than just the bone marrow? *Am J Hematol*. 2020;95(8):E210-E213. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32419212.
- 46. National Comprehensive Cancer Network. NCCN hematopoietic growth factors: short-term recommendations specific to issues with COVID-19 (SARS-CoV-2). 2020. Available at: <u>https://www.iononline.com/-/media/assets/ion/pdf/covid19-resources/nccn_hgf_covid-19_19may20.pdf</u>.
- 47. van Arkel ALE, Rijpstra TA, Belderbos HNA, et al. COVID-19-associated pulmonary aspergillosis. *Am J Respir Crit Care Med.* 2020;202(1):132-135. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32396381</u>.
- 48. Alanio A, Dellière S, Fodil S, Bretagne S, Mégarbane B. Prevalence of putative invasive pulmonary

aspergillosis in critically ill patients with COVID-19. *Lancet Respir Med*. 2020;8(6):e48-e49. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32445626</u>.

- 49. Hrusak O, Kalina T, Wolf J, et al. Flash survey on severe acute respiratory syndrome coronavirus-2 infections in paediatric patients on anticancer treatment. *Eur J Cancer*. 2020;132:11-16. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32305831.
- 50. André N, Rouger-Gaudichon J, Brethon B, et al. COVID-19 in pediatric oncology from French pediatric oncology and hematology centers: high risk of severe forms? *Pediatr Blood Cancer*. 2020;67(7):e28392. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32383827</u>.
- 51. de Rojas T, Peréz-Martínez A, Cela E, et al. COVID-19 infection in children and adolescents with cancer in Madrid. *Pediatr Blood Cancer*. 2020;67(7):e28397. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32383819.
- 52. Sullivan M, Bouffet E, Rodriguez-Galindo C, et al. The COVID-19 pandemic: a rapid global response for children with cancer from SIOP, COG, SIOP-E, SIOP-PODC, IPSO, PROS, CCI, and St Jude Global. *Pediatr Blood Cancer*. 2020;67(7):e28409. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32400924.
- 53. Bouffet E, Challinor J, Sullivan M, et al. Early advice on managing children with cancer during the COVID-19 pandemic and a call for sharing experiences. *Pediatr Blood Cancer*. 2020;67(7):e28327. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32239747.
- 54. Chiotos K, Hayes M, Kimberlin DW, et al. Multicenter initial guidance on use of antivirals for children with coronavirus disease 2019/severe acute respiratory syndrome coronavirus 2. *J Pediatric Infect Dis Soc*. 2020;9(6):701-715. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32318706</u>.

Special Considerations in Solid Organ Transplant, Hematopoietic Cell Transplant, and Cellular Immunotherapy Candidates, Donors, and Recipients

Last Updated: November 2, 2023

Summary Recommendations

Vaccination for COVID-19

- COVID-19 vaccination remains the most effective way to prevent SARS-CoV-2 infection and should be considered the first line of prevention. Given the effectiveness of COVID-19 vaccines in the general population and the increased risk of worse clinical outcomes of COVID-19 in transplant and cellular immunotherapy candidates and recipients, the COVID-19 Treatment Guidelines Panel (the Panel) recommends COVID-19 vaccination for these patients (AIII).
- Because vaccine response rates may be lower in moderately or severely immunocompromised patients, specific guidance on administering vaccines to these individuals is provided by the Centers for Disease Control and Prevention (CDC). See the CDC webpage <u>COVID-19 Vaccines for People Who Are Moderately or Severely Immunocompromised</u> for the current vaccination schedule for this population.
- Vaccinating household members, close contacts, and health care providers who provide care to transplant and cellular immunotherapy candidates and recipients is important to protect these patients from infection. All close contacts are strongly encouraged to get vaccinated against COVID-19 as soon possible (AI).
- Clinicians should strongly encourage all potential organ and hematopoietic cell donors to get vaccinated against COVID-19 (AI).

Potential Transplant and Cellular Immunotherapy Candidates

- The Panel recommends performing diagnostic molecular or antigen testing for SARS-CoV-2 for all potential solid organ transplant, hematopoietic cell transplant (HCT), and cellular immunotherapy candidates with signs and symptoms that suggest acute COVID-19 (AIII). Additional guidance is available from medical professional organizations. See the text below for more information.
- If SARS-CoV-2 is detected or if infection is strongly suspected in a potential transplant or cellular immunotherapy candidate, transplantation or immunotherapy should be deferred, if possible (**BIII**).
- The optimal management and therapeutic approach to COVID-19 in these populations is unknown. At this time, the procedures for evaluating and managing COVID-19 in transplant candidates are the same as those for nontransplant patients (AIII).

Potential Transplant Donors

• The Panel recommends performing diagnostic molecular testing for SARS-CoV-2 and assessing for symptoms of COVID-19 in all potential solid organ transplant and HCT donors prior to donation (AIII). Additional guidance is available from medical professional organizations. See the text below for more information.

Transplant and Cellular Immunotherapy Recipients With COVID-19

- Clinicians should follow the guidelines for evaluating and managing COVID-19 in nontransplant patients when treating transplant and cellular immunotherapy recipients (AIII). See <u>Therapeutic Management of Hospitalized Adults With</u> <u>COVID-19</u> and <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for more information.
- Immunocompromised patients with mild to moderate COVID-19 are at high risk of progressing to severe disease and should receive anti-SARS-CoV-2 therapies for treatment.
- Clinicians who are treating COVID-19 in transplant and cellular immunotherapy patients should consult a transplant specialist before adjusting immunosuppressive medications (AIII).
- When treating COVID-19, clinicians should pay careful attention to potential overlapping toxicities and drugdrug interactions between drugs used to treat COVID-19 (e.g., ritonavir-boosted nirmatrelvir [Paxlovid]) and immunosuppressants, prophylactic antimicrobials, or other medications (AIII).

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Introduction

Treating COVID-19 in solid organ transplant, hematopoietic cell transplant (HCT), and cellular immunotherapy recipients can be challenging due to the presence of coexisting medical conditions, the potential for transplant-related cytopenias, and the need for chronic immunosuppressive therapy to prevent graft rejection and graft-versus-host disease. Transplant recipients may also have a higher risk of exposure to SARS-CoV-2 given their frequent contact with the health care system. Since immunosuppressive agents modulate several aspects of immune response, the severity of COVID-19 could potentially be affected by the type and intensity of the immunosuppressive effect of the agent, as well as by specific combinations of immunosuppressive agents. Some transplant recipients have medical comorbidities that have been associated with more severe cases of COVID-19 and a greater risk of mortality, which makes the impact of transplantation on disease severity difficult to assess.

The International Society for Heart and Lung Transplantation, the American Society of Transplantation (AST), the American Society for Transplantation and Cellular Therapy, and the European Society for Blood and Marrow Transplantation provide guidance for clinicians who are caring for transplant recipients with COVID-19 and guidance on screening potential donors and transplant or cellular immunotherapy candidates. In addition, the American Society of Hematology offers guidance regarding COVID-19 vaccination for transplant and cellular immunotherapy recipients.

This section of the COVID-19 Treatment Guidelines complements these sources and focuses on considerations for managing COVID-19 in solid organ transplant, HCT, and cellular immunotherapy recipients. The optimal management and therapeutic approach to COVID-19 in these populations is unknown. At this time, the procedures for evaluating and managing COVID-19 in transplant recipients are the same as those for nontransplant patients (AIII). See <u>Therapeutic Management of Hospitalized</u> <u>Adults With COVID-19</u> and <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for more information. The risks and benefits of each medication used to treat COVID-19 may be different for transplant patients and nontransplant patients.

Vaccination for COVID-19

The clinical trials that evaluated the safety and efficacy of the COVID-19 vaccines excluded patients who were severely immunocompromised.^{1,2} The currently authorized or approved COVID-19 vaccines are not live vaccines; therefore, they can be safely administered to people who are immunocompromised. However, solid organ transplant recipients have reduced immunological antibody responses following a primary 2-dose or 3-dose series of the mRNA COVID-19 vaccines.³⁻⁶

Given the effectiveness of COVID-19 vaccines in the general population and the increased risk of worse clinical outcomes of COVID-19 in transplant and cellular immunotherapy recipients, the COVID-19 Treatment Guidelines Panel (the Panel) recommends COVID-19 vaccination for potential transplant and cellular immunotherapy candidates and recipients (**AIII**). See the Centers for Disease Control and Prevention (CDC) website <u>COVID-19 Vaccines for People Who Are Moderately or Severely</u> <u>Immunocompromised</u> for the current COVID-19 vaccination schedule for transplant and cellular immunotherapy recipients.

When determining the timing of COVID-19 vaccination (including booster doses) in solid organ transplant, HCT, and cellular immunotherapy recipients, clinicians should consider the following factors:

- Ideally, solid organ transplant candidates should receive COVID-19 vaccines (either the primary series or booster doses) while they are awaiting transplant.
- In general, the last vaccine should be administered at least 2 weeks prior to a solid organ transplant, or vaccination should be started 1 month after a solid organ transplant.

COVID-19 Treatment Guidelines

- In certain situations, it may be appropriate to delay the primary series of vaccinations or booster doses until 3 months after a solid organ transplant, such as when T cell– or B cell–ablative therapy (with antithymocyte globulin or rituximab) is used at the time of transplant.⁷
- Reducing the dose of immunosuppressants and withholding immunosuppressants prior to vaccination **are not recommended**.
- COVID-19 vaccines can be offered as early as 3 months after a patient receives HCT or chimeric antigen receptor T cell therapy, although the vaccines may be less effective in these patients than in the general population.⁸⁻¹⁰ If possible, patients who are scheduled to receive cytotoxic or B cell-depleting therapies should receive their COVID-19 vaccination before initiating these therapies or between cycles of these therapies.
- After receiving the primary series of vaccinations or booster doses,¹¹ people who are immunocompromised should be advised to continue to exercise precautions to reduce their risk of SARS-CoV-2 exposure and infection (e.g., they should wear a mask, maintain a distance of 6 feet from others, and avoid crowds and poorly ventilated spaces).

There is insufficient evidence for the Panel to recommend either for or against the use of SARS-CoV-2 serologic testing to assess for immunity or to guide clinical decisions about using COVID-19 vaccines. For people who received COVID-19 vaccines during treatment with immunosuppressive drugs, it is currently unknown whether revaccination offers a clinical benefit.

Vaccinating household members, close contacts, and health care providers who provide care to transplant and cellular immunotherapy candidates and recipients is important to protect these patients from infection. All close contacts are strongly encouraged to get vaccinated against COVID-19 as soon as possible (AI). There is evidence that vaccinated individuals who are infected with SARS-CoV-2 have lower viral loads than unvaccinated individuals^{12,13} and that COVID-19 vaccines reduce the incidence of SARS-CoV-2 infections not only among vaccinated individuals but also among their household contacts.¹⁴⁻¹⁶ Clinicians should strongly encourage all potential organ and hematopoietic cell donors to get vaccinated against COVID-19 (AI).

Assessing SARS-CoV-2 Infection

The risk of transmission of SARS-CoV-2 from donors to candidates is unknown. The probability that a donor or candidate may have SARS-CoV-2 infection can be estimated by considering the epidemiologic risk, obtaining a clinical history, and testing with molecular techniques. No current testing strategy is sensitive enough or specific enough to totally exclude active infection.

Assessing Transplant and Cellular Immunotherapy Candidates

The Panel recommends performing diagnostic molecular or antigen testing for SARS-CoV-2 in all potential solid organ transplant, HCT, and cellular immunotherapy candidates with signs and symptoms that suggest acute COVID-19 (**AIII**). The CDC testing algorithm recommends performing additional confirmatory testing with a laboratory-based nucleic acid amplification test (NAAT) when a person who is strongly suspected of having SARS-CoV-2 infection (i.e., a person who is symptomatic) receives a negative result on an antigen test.¹⁷ Shortly before solid organ transplant, HCT, or cellular immunotherapy, all candidates should undergo diagnostic molecular testing for SARS-CoV-2 and assessment for symptoms of COVID-19 (**AIII**).

Assessing Donors

The Panel recommends performing diagnostic molecular testing for SARS-CoV-2 and assessing for symptoms of COVID-19 in all potential solid organ transplant and HCT donors prior to donation

(AIII). Additional guidance is available from medical professional organizations, such as the <u>Organ</u> <u>Procurement and Transplantation Network (OPTN)</u> and the <u>AST</u>.

Living donors should undergo a SARS-CoV-2 NAAT using a specimen collected from the respiratory tract within 3 days of donation. Deceased donors should be tested for SARS-CoV-2 infection using a NAAT with a specimen taken from the upper respiratory tract within 72 hours of death; ideally, the test should be performed as close to organ recovery as possible. Lower respiratory sampling for COVID-19 testing is required for potential lung transplant donors by the United Network for Organ Sharing.¹⁸ The OPTN and AST provide information to help guide the decision-making process when managing solid organ transplant donors with a history of COVID-19.

If SARS-CoV-2 Infection Is Detected or Strongly Suspected in Transplant and Cellular Immunotherapy Candidates

If SARS-CoV-2 is detected or if infection is strongly suspected in a potential transplant or cellular immunotherapy candidate, transplantation or immunotherapy should be deferred, if possible (**BIII**). The optimal disease-free interval before transplantation or immunotherapy is not known. In this situation, decisions about the appropriate timing for transplantation or cellular immunotherapy should be made on a case-by-case basis. Clinicians should consider both the risk of viral transmission and the risks of delaying or altering therapy, which may include progression of the underlying disease or death.

Transplant Recipients With COVID-19

Solid organ transplant recipients receiving immunosuppressive therapy should be considered at increased risk for severe COVID-19.^{19,20} Initial reports of transplant recipients hospitalized with COVID-19 suggest mortality rates of up to 28%.²¹⁻²⁵

Risk of Graft Rejection

There are concerns that COVID-19 itself may increase the risk for acute rejection. Acute cellular rejection should not be presumed in solid organ transplant recipients without biopsy confirmation, regardless of whether the individual has COVID-19. Similarly, immunosuppressive therapy should be initiated in recipients with or without COVID-19 who have rejection confirmed by a biopsy.¹⁹

There are limited data on the incidence and clinical characteristics of SARS-CoV-2 infection in HCT26 and cellular immunotherapy recipients.²⁷ Data from the Center for International Blood and Marrow Transplant Research demonstrated that approximately 30% of a cohort of 318 HCT recipients died within 30 days of COVID-19 diagnosis.²⁶ This probability of mortality was observed in both allogeneic and autologous recipients. Older age (\geq 50 years), male sex, and receipt of a COVID-19 diagnosis within 12 months of transplantation were associated with a higher risk of mortality among allogeneic recipients. In autologous recipients, patients with lymphoma had a higher risk of mortality than patients who had plasma cell disorder or myeloma.

A smaller study demonstrated slightly lower mortality among HCT and cellular immunotherapy recipients than earlier reports. This study found that the number of comorbidities, the presence of infiltrates on initial chest imaging, and neutropenia were predictors for increased disease severity.²⁸ Additional factors that have been used to determine the clinical severity of other respiratory viral infections include the degree of cytopenia, the intensity of the conditioning regimen, the graft source, the degree of mismatch, and the need for further immunosuppression to manage graft-versus-host disease. Prolonged viral shedding has been described in solid organ transplant and HCT recipients; this can have implications for preventing infection and for the timing of therapeutic interventions.²⁹

Treating COVID-19 in Transplant Recipients

Outpatient transplant recipients who are immunosuppressed or who have certain underlying comorbidities are candidates for several other therapeutic agents that are available through Food and Drug Administration (FDA) Emergency Use Authorizations (EUAs). See <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u> for more information.

When treating hospitalized patients with mild to moderate, symptomatic COVID-19, clinicians should consider administering the therapeutics used in nonhospitalized patients with similar disease severity. Data from a large randomized controlled trial found that a short course of dexamethasone (6 mg once daily for up to 10 days) improved survival in hospitalized people with severe COVID-19 who were mechanically ventilated or who required supplemental oxygen.³⁰ Tocilizumab or baricitinib used in combination with dexamethasone is recommended for some patients with severe or critical COVID-19.³⁰⁻³² Because dexamethasone, tocilizumab, and baricitinib are immunosuppressive agents, patients who receive these medications should be closely monitored for secondary infections.

Therapeutic anticoagulation for transplant recipients who are hospitalized for COVID-19 should be managed similarly to anticoagulation for other hospitalized patients. Patients with platelet counts <50,000 cells/µL should not receive therapeutic anticoagulation to treat COVID-19. Clinicians should follow hospital protocols for managing anticoagulation in patients with thrombocytopenia.

The Panel's recommendations for the use of remdesivir, dexamethasone, tocilizumab, baricitinib, and anticoagulation in hospitalized patients with COVID-19 can be found in <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>.

Concomitant Medications

Clinicians should pay special attention to the potential for drug-drug interactions and overlapping toxicities between treatments for COVID-19 and concomitant medications, such as immunosuppressants used to prevent allograft rejection and antimicrobials used to prevent or treat opportunistic infections. Dose modifications may be necessary for drugs used to treat COVID-19 in transplant recipients with pre-existing organ dysfunction. Adjustments to the immunosuppressive regimen should be individualized based on disease severity, the specific immunosuppressants used, the type of transplant, the time since transplantation, the drug concentration, and the risk of graft rejection.²² Clinicians who are treating COVID-19 in transplant and cellular immunotherapy patients should consult a transplant specialist before adjusting immunosuppressive medications (**AIII**).

Drug-Drug Interactions

Calcineurin inhibitors (e.g., cyclosporine, tacrolimus) and mammalian target of rapamycin (mTOR) inhibitors (e.g., everolimus, sirolimus), which are commonly used to prevent allograft rejection, have narrow therapeutic indices. Medications that inhibit or induce cytochrome P450 (CYP) enzymes or P-glycoprotein may put patients who receive these drugs at risk of clinically significant drug-drug interactions, increasing the need for therapeutic drug monitoring and the need to assess for signs of toxicity or rejection.³³

A 5-day course of ritonavir-boosted nirmatrelvir (Paxlovid) is 1 of the preferred therapies for treating mild to moderate COVID-19 in nonhospitalized patients who are at risk for disease progression. However, this regimen has the potential for significant and complex drug-drug interactions with concomitant medications, primarily due to the ritonavir component of the combination. Boosting with ritonavir, a strong CYP3A inhibitor, is required to increase the exposure of nirmatrelvir to a concentration that is effective against SARS-CoV-2. Ritonavir may also increase concentrations of

certain concomitant medications, including calcineurin and mTOR inhibitors, during the treatment course and for \geq 3 days after ritonavir is discontinued. Significant increases in the concentrations of these drugs may lead to serious and sometimes life-threatening drug toxicities.

If remdesivir is not available or feasible to use, ritonavir-boosted nirmatrelvir may be used with caution and only when close therapeutic drug monitoring of the antirejection therapy is possible. Clinicians should consult with transplant specialists during the treatment course. General guidance for coadministering ritonavir-boosted nirmatrelvir with concomitant medications includes temporarily withholding certain immunosuppressive agents (e.g., tacrolimus, everolimus, sirolimus) or reducing the dosage of certain immunosuppressive agents (e.g., cyclosporine), monitoring the patient closely for toxicities, and performing therapeutic drug monitoring during and after the 5-day treatment course of ritonavir-boosted nirmatrelvir.^{34,35}

Some small case series have reported success using these recommendations to manage patients;^{36,37} however, cases of significant toxicities due to supratherapeutic tacrolimus concentrations have also been reported.³⁸ Therapeutic drug monitoring should be used to guide the process of reintroducing or modifying the doses of calcineurin and mTOR inhibitors in patients who have completed a course of ritonavir-boosted nirmatrelvir. Clinicians should also consult with a specialist who has experience with dose management. Clinicians should take additional precautions when treating transplant recipients who are also receiving other concomitant medications (e.g., certain triazole antifungals) that may interact with ritonavir, the immunosuppressants, or both. The extent and significance of multiple drug-drug interactions are much more complex and unpredictable.

Clinicians should refer to resources such as the <u>Liverpool COVID-19 Drug Interactions website</u>, <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant</u> <u>Medications</u>, and the FDA <u>prescribing information</u> on ritonavir-boosted nirmatrelvir for guidance on identifying and managing potential drug-drug interactions. If significant interactions prohibit the concomitant use of ritonavir-boosted nirmatrelvir, another COVID-19 treatment option should be used (see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>).

Among the drugs that are commonly used to treat hospitalized patients with COVID-19, dexamethasone is a moderate inducer of CYP3A4, and interleukin-6 inhibitors may lead to increased metabolism of CYP substrates. Clinicians should closely monitor the serum concentrations of calcineurin and mTOR inhibitors when these drugs are used.

Additional details about the adverse effects and drug-drug interactions of antiviral medications and immune-based therapies for COVID-19 are noted in <u>Tables 4e</u> and <u>5e</u>.

References

- 1. Baden LR, El Sahly HM, Essink B, et al. Efficacy and safety of the mRNA-1273 SARS-CoV-2 vaccine. *N Engl J Med.* 2021;384(5):403-416. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33378609</u>.
- Polack FP, Thomas SJ, Kitchin N, et al. Safety and efficacy of the BNT162b2 mRNA COVID-19 vaccine. N Engl J Med. 2020;383(27):2603-2615. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33301246</u>.
- Boyarsky BJ, Werbel WA, Avery RK, et al. Antibody response to 2-dose SARS-CoV-2 mRNA vaccine series in solid organ transplant recipients. *JAMA*. 2021;325(21):2204-2206. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33950155.
- 4. Hallett AM, Greenberg RS, Boyarsky BJ, et al. SARS-CoV-2 messenger RNA vaccine antibody response and reactogenicity in heart and lung transplant recipients. *J Heart Lung Transplant*. 2021;40(12):1579-1588. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34456108</u>.
- 5. Mazzola A, Todesco E, Drouin S, et al. Poor antibody response after two doses of severe acute respiratory

COVID-19 Treatment Guidelines

syndrome coronavirus 2 (SARS-CoV-2) vaccine in transplant recipients. *Clin Infect Dis*. 2022;74(6):1093-1096. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34166499</u>.

- Kamar N, Abravanel F, Marion O, et al. Three doses of an mRNA COVID-19 vaccine in solid-organ transplant recipients. *N Engl J Med.* 2021;385(7):661-662. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34161700</u>.
- 7. American Society of Transplantation. COVID-19 vaccine FAQ sheet. 2021. Available at: <u>https://www.myast.org/sites/default/files/2021_08_13%20COVID%20VACCINE%20FAQ-Prof8132021_FINAL.pdf</u>.
- American Society of Hematology. ASH-ASTCT COVID-19 vaccination for HCT and CAR T cell recipients: frequently asked questions. 2022. Available at: <u>https://www.hematology.org/covid-19/ash-astct-covid-19-vaccination-for-hct-and-car-t-cell-recipients</u>. Accessed February 15, 2023.
- 9. Ljungman P, Avetisyan G. Influenza vaccination in hematopoietic SCT recipients. *Bone Marrow Transplant*. 2008;42(10):637-41. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/18724396</u>.
- Ram R, Hagin D, Kikozashvilli N, et al. Safety and immunogenicity of the BNT162b2 mRNA COVID-19 vaccine in patients after allogeneic HCT or CD19-based CART therapy—a single-center prospective cohort study. *Transplant Cell Ther*. 2021;27(9):788-794. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34214738.
- Centers for Disease Control and Prevention. Stay up to date with COVID-19 vaccines including boosters. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/vaccines/stay-up-to-date.html</u>. Accessed February 15, 2023.
- Levine-Tiefenbrun M, Yelin I, Katz R, et al. Initial report of decreased SARS-CoV-2 viral load after inoculation with the BNT162b2 vaccine. *Nat Med.* 2021;27(5):790-792. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33782619</u>.
- Petter E, Mor O, Zuckerman N, et al. Initial real world evidence for lower viral load of individuals who have been vaccinated by BNT162b2. *medRxiv*. 2021;Preprint. Available at: https://www.medrxiv.org/content/10.1101/2021.02.08.21251329v1.
- Salo J, Hägg M, Kortelainen M, et al. The indirect effect of mRNA-based COVID-19 vaccination on healthcare workers' unvaccinated household members. *Nat Commun.* 2022;13(1):1162. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35246536.
- 15. Madewell ZJ, Yang Y, Longini IM Jr, Halloran ME, Dean NE. Household secondary attack rates of SARS-CoV-2 by variant and vaccination status: an updated systematic review and meta-analysis. *JAMA Netw Open.* 2022;5(4):e229317. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35482308</u>.
- 16. de Gier B, Andeweg S, Backer JA, et al. Vaccine effectiveness against SARS-CoV-2 transmission to household contacts during dominance of Delta variant (B.1.617.2), the Netherlands, August to September 2021. Euro Surveill. 2021;26(44):2100977. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34738514</u>.
- 17. Centers for Disease Control and Prevention. Guidance for antigen testing for SARS-CoV-2 for healthcare providers testing individuals in the community. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/lab/resources/antigen-tests-guidelines.html</u>. Accessed February 15, 2023.
- United Network for Organ Sharing. Lower respiratory testing of all potential lung donors for SARS-CoV-2 now required. 2021. Available at: <u>https://unos.org/news/sars-cov-2-lower-respiratory-testing-potential-lungdonors-may-27</u>. Accessed February 15, 2023.
- Fix OK, Hameed B, Fontana RJ, et al. Clinical best practice advice for hepatology and liver transplant providers during the COVID-19 pandemic: AASLD expert panel consensus statement. *Hepatology*. 2020;72(1):287-304. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32298473</u>.
- 20. Centers for Disease Control and Prevention. Underlying medical conditions associated with higher risk for severe COVID-19: information for healthcare professionals. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/clinical-care/underlyingconditions.html</u>. Accessed February 15, 2023.
- 21. Akalin E, Azzi Y, Bartash R, et al. COVID-19 and kidney transplantation. N Engl J Med. 2020;382(25):2475-

2477. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32329975.

- 22. Pereira MR, Mohan S, Cohen DJ, et al. COVID-19 in solid organ transplant recipients: initial report from the US epicenter. *Am J Transplant*. 2020;20(7):1800-1808. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32330343.
- 23. Alberici F, Delbarba E, Manenti C, et al. A single center observational study of the clinical characteristics and short-term outcome of 20 kidney transplant patients admitted for SARS-CoV2 pneumonia. *Kidney Int.* 2020;97(6):1083-1088. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32354634</u>.
- 24. Montagud-Marrahi E, Cofan F, Torregrosa JV, et al. Preliminary data on outcomes of SARS-CoV-2 infection in a Spanish single center cohort of kidney recipients. *Am J Transplant*. 2020;20(10):2958-2959. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32368838.
- 25. Kates OS, Haydel BM, Florman SS, et al. Coronavirus disease 2019 in solid organ transplant: a multicenter cohort study. *Clin Infect Dis*. 2021;73(11):e4090-e4099. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32766815</u>.
- 26. Sharma A, Bhatt NS, St Martin A, et al. Clinical characteristics and outcomes of COVID-19 in haematopoietic stem-cell transplantation recipients: an observational cohort study. *Lancet Haematol*. 2021;8(3):e185-e193. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33482113</u>.
- 27. Gold JAW, Wong KK, Szablewski CM, et al. Characteristics and clinical outcomes of adult patients hospitalized with COVID-19—Georgia, March 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(18):545-550. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32379729</u>.
- 28. Shah GL, DeWolf S, Lee YJ, et al. Favorable outcomes of COVID-19 in recipients of hematopoietic cell transplantation. J Clin Invest. 2020;130(12):6656-6667. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32897885</u>.
- 29. Aydillo T, Gonzalez-Reiche AS, Aslam S, et al. Shedding of viable SARS-CoV-2 after immunosuppressive therapy for cancer. *N Engl J Med*. 2020;383(26):2586-2588. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33259154</u>.
- 30. RECOVERY Collaborative Group. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2021;397(10285):1637-1645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33933206</u>.
- 31. REMAP-CAP Investigators. Interleukin-6 receptor antagonists in critically ill patients with COVID-19. *N Engl J Med.* 2021;384(16):1491-1502. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33631065</u>.
- 32. Kalil AC, Patterson TF, Mehta AK, et al. Baricitinib plus remdesivir for hospitalized adults with COVID-19. *N Engl J Med.* 2021;384(9):795-807. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33306283</u>.
- 33. Elens L, Langman LJ, Hesselink DA, et al. Pharmacologic treatment of tansplant recipients infected with SARS-CoV-2: considerations regarding therapeutic drug monitoring and drug-drug interactions. *Ther Drug Monit*. 2020;42(3):360-368. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32304488</u>.
- 34. Lange NW, Salerno DM, Jennings DL, et al. Nirmatrelvir/ritonavir use: managing clinically significant drug-drug interactions with transplant immunosuppressants. Am J Transplant. 2022;22(7):1925-1926. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35015924</u>.
- 35. American Society of Transplantation. COVID-19: FAQs for Organ Transplantation. 2023. Available at: <u>https://www.myast.org/sites/default/files/COVID%20FAQ%20for%20Tx%20professionals%202-2023%20FINAL.pdf</u>.
- 36. Salerno DM, Jennings DL, Lange NW, et al. Early clinical experience with nirmatrelvir/ritonavir for the treatment of COVID-19 in solid organ transplant recipients. *Am J Transplant*. 2022;22(8):2083-2088. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35278260</u>.
- Wang AX, Koff A, Hao D, Tuznik NM, Huang Y. Effect of nirmatrelvir/ritonavir on calcineurin inhibitor levels: early experience in four SARS-CoV-2 infected kidney transplant recipients. *Am J Transplant*. 2022;22(8):2117-2119. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35158412</u>.

38. Rose DT, Gandhi SM, Bedard RA, et al. Supratherapeutic tacrolimus concentrations with nirmatrelvir/ ritonavir in solid organ transplant recipients requiring hospitalization: a case series using rifampin for reversal. *Open Forum Infect Dis.* 2022;9(7):ofac238. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35854994</u>.

Special Considerations During Pregnancy and After Delivery

Last Updated: July 21, 2023

Summary Recommendations

Current guidance from the <u>Centers for Disease Control and Prevention</u>, the <u>American College of Obstetricians and</u> <u>Gynecologists</u>, and the <u>Society for Maternal-Fetal Medicine</u> details the management of pregnant patients with COVID-19. This section of the COVID-19 Treatment Guidelines complements that guidance. The following are key considerations regarding the management of COVID-19 in pregnancy:

- Pregnant people should be counseled about the increased risk for severe disease from SARS-CoV-2 infection and receive recommendations on ways to protect themselves and their families from infection.
- If hospitalization for COVID-19 is indicated for a pregnant patient, care should be provided in a facility that can conduct maternal and fetal monitoring, when appropriate.
- General management of COVID-19 in pregnant patients should include:
 - · Fetal and uterine contraction monitoring based on gestational age, when appropriate
 - · Individualized delivery planning
 - A multispecialty, team-based approach that may include consultation with obstetric, maternal-fetal medicine, infectious disease, pulmonary-critical care, and pediatric specialists, as appropriate
- The COVID-19 Treatment Guidelines Panel (the Panel) **recommends against** withholding COVID-19 treatments or vaccination from pregnant or lactating individuals specifically because of pregnancy or lactation (AIII).
- In general, the therapeutic management of pregnant patients with COVID-19 should be the same as for nonpregnant patients, with a few exceptions (AIII). Notable exceptions include:
 - The Panel **recommends against** the use of **molnupiravir** for the treatment of COVID-19 in pregnant patients unless there are no other options and therapy is clearly indicated **(AIII)**.
 - There is insufficient evidence for the Panel to recommend either for or against the use of therapeutic anticoagulation in pregnant patients with COVID-19 who do not have evidence of venous thromboembolism. See <u>Antithrombotic</u> <u>Therapy in Patients With COVID-19</u> for more information.
- For details regarding therapeutic recommendations and pregnancy considerations, see <u>Therapeutic Management of</u> <u>Nonhospitalized Adults With COVID-19</u>; <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>; <u>Pregnancy</u>, <u>Lactation</u>, and <u>COVID-19 Therapeutics</u>; and the individual drug sections.
- There are limited data on the use of COVID-19 therapeutic agents in pregnant and lactating people. When making decisions about treatment, pregnant or lactating people and their clinical teams should use a shared decision-making process and consider several factors, including the severity of COVID-19, the risk of disease progression, and the safety of specific medications for the fetus, infant, or pregnant or lactating individual. For detailed guidance on using the Panel-recommended COVID-19 therapeutic agents during pregnancy, see <u>Pregnancy, Lactation, and COVID-19</u>. <u>Therapeutics</u>.
- The decision to feed the infant breast milk while the lactating patient is receiving therapeutic agents for COVID-19 should be a collaborative effort between the patient and the clinical team, including infant care providers. The patient and the clinical team should discuss the potential benefits of the therapeutic agent and evaluate the potential risk of pausing lactation on future breast milk delivery to the infant. For more information, see <u>Pregnancy, Lactation, and COVID-19 Therapeutics</u>.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Epidemiology of COVID-19 in Pregnancy

Although the overall risk of severe illness is low, pregnant people with COVID-19 are at a higher risk of severe disease than nonpregnant people. After adjustments have been made for age, race/ethnicity, and underlying medical conditions, pregnant women have significantly higher rates of intensive care unit

(ICU) admission (10.5 vs. 3.9 cases per 1,000 cases; adjusted risk ratio [aRR] 3.0; 95% CI, 2.6–3.4), mechanical ventilation (2.9 vs. 1.1 cases per 1,000 cases; aRR 2.9; 95% CI, 2.2–3.8), extracorporeal membrane oxygenation (0.7 vs. 0.3 cases per 1,000 cases; aRR 2.4; 95% CI, 1.5–4.0), and death (1.5 vs. 1.2 cases per 1,000 cases; aRR 1.7; 95% CI, 1.2–2.4).¹

An ongoing systematic review and meta-analysis of 149 studies also described increased odds of ICU admission and mechanical ventilation among pregnant and recently pregnant patients with COVID-19 when compared with nonpregnant patients of reproductive age.^{2,3} Compared with pregnant women and recently pregnant women without COVID-19, pregnant women with COVID-19 were at a higher risk of preterm birth and stillbirth.

Obstetric and Perinatal Outcomes in Patients With COVID-19

An observational cohort study of all pregnant patients at 33 U.S. hospitals with a singleton gestation and a positive result on a SARS-CoV-2 virologic test evaluated maternal characteristics and outcomes across disease severity.⁴ The data suggested that adverse perinatal outcomes were more common in patients with severe or critical disease than in asymptomatic patients with SARS-CoV-2 infection, including an increased incidence of cesarean delivery (59.6% vs. 34.0% of patients; aRR 1.57; 95% CI, 1.30–1.90), hypertensive disorders of pregnancy (40.4% vs. 18.8%; aRR 1.61; 95% CI, 1.18–2.20), and preterm birth (41.8% vs. 11.9%; aRR 3.53; 95% CI, 2.42–5.14). The perinatal outcomes for those with mild to moderate illness were similar to those observed among asymptomatic patients with SARS-CoV-2 infection.

Among 1,249,634 delivery hospitalizations in the United States from March 2020 through September 2021, women with COVID-19 had an increased risk of stillbirth, which was defined as fetal death at >20 weeks' gestation (aRR 1.90; 95% CI, 1.69–2.15).⁵ The risk of stillbirth was higher during the time period that the Delta variant was the dominant variant in the United States (aRR 4.04; 95% CI, 3.28–4.97) than during the pre-Delta period (aRR 1.47; 95% CI, 1.27–1.71).

A retrospective cohort analysis collected data from 14,104 pregnant or recently postpartum individuals who delivered at U.S. hospitals that participated in the Gestational Research Assessments for COVID-19 (GRAVID) study.⁶ Compared with pregnant individuals who did not have SARS-CoV-2 infection, patients with COVID-19 during pregnancy had an increased risk of meeting the composite endpoint of maternal death or severe morbidity related to hypertensive disorders of pregnancy, postpartum hemorrhage, or infection. Eighty percent of the patients in this cohort tested positive for SARS-CoV-2 infection during the third trimester. The primary composite endpoint occurred in 13.4% of patients with COVID-19 during pregnancy or within 6 weeks postpartum and in 9.2% of those without COVID-19 (aRR 1.41; 95% CI, 1.23–1.61).

When compared with those who did not have a positive SARS-CoV-2 test result, pregnant patients who had SARS-CoV-2 infection prior to 28 weeks' gestation had a subsequent increased risk of fetal/neonatal death (aRR 1.97; 95% CI, 1.01–3.85), preterm birth at <37 weeks (aRR 1.29; 95% CI, 1.02–1.63), and hypertensive disorders of pregnancy with delivery at <37 weeks' gestation (aRR 1.74; 95% CI, 1.19–2.55).⁷ There were no significant differences between these groups of patients in the risk of preterm birth at <34 weeks, any major congenital abnormalities, or a size for gestational age of less than the fifth or tenth percentiles. There were also no significant differences between these groups in the rates of gestational hypertension overall or preeclampsia with severe features. These data suggest that those with SARS-CoV-2 infection early in gestation may also have an increased risk of subsequent adverse pregnancy outcomes.

Vertical Transmission of COVID-19

Although vertical transmission of SARS-CoV-2 is possible, current data suggest that it is rare.⁸ A review

of 101 infants born to 100 women with SARS-CoV-2 infection at a single U.S. academic medical center found that 2 infants (2%) had indeterminate SARS-CoV-2 polymerase chain reaction (PCR) results, which were presumed to be positive; however, the infants exhibited no evidence of clinical disease. It is reassuring that the majority of the infants received negative PCR results after rooming with their mothers and breastfeeding directly (the mothers in this study practiced appropriate hand and breast hygiene).

Data collected by the Centers for Disease Control and Prevention (CDC) as part of the Surveillance for Emerging Threats to Mothers and Babies Network showed that among 4,038 infants born to people with COVID-19, for whom laboratory testing information was available and who were tested during the delivery hospitalization, 227 infants (5.6%) had positive PCR results for SARS-CoV-2.⁹

The published data to date were largely collected prior to the emergence of the Omicron variants. The risk of vertical transmission may vary based on viral dynamics and the transmissibility of the circulating variants in a community; however, the variant-specific factors that are associated with vertical transmission have not been determined. For additional information on vertical transmission and infants born to people with SARS-CoV-2 infection, see <u>Special Considerations in Children</u>.

Racial and Ethnic Disparities Among Pregnant People With COVID-19

Between January 22 and June 7, 2020, 8,207 pregnant women with COVID-19 were reported to CDC. Among these women, 46% were reported to be Hispanic and 22% were reported to be Black.¹⁰ Those proportions were higher than the proportions of Hispanic and Black women who gave birth in 2019 (24% and 15%, respectively), suggesting that pregnant people who are Hispanic or Black may be disproportionately affected by SARS-CoV-2 infection. It is important to note that these disparities are related to social determinants of health, current and historic inequities in access to health care and other resources, and structural racism. The American College of Obstetricians and Gynecologists (ACOG) has published guidance on addressing health equity during the COVID-19 pandemic.

Prevention of COVID-19 in Pregnancy

Pregnant people should be counseled about the increased risk for severe disease from SARS-CoV-2 and the measures they can take to protect themselves and their families from infection. Nonpharmacologic measures include practicing physical distancing, washing hands regularly, and wearing a face covering as per guidance from the CDC.

COVID-19 Vaccines

The COVID-19 Treatment Guidelines Panel (the Panel) **recommends against** withholding COVID-19 vaccination from pregnant or lactating individuals specifically because of pregnancy or lactation (**AIII**).

Pregnant people should be counseled about the benefits of COVID-19 vaccination, which include a decreased risk of severe disease and hospitalization for the pregnant person and a decreased risk of hospitalization for the infant in the first 6 months of life.¹¹ The Society for Maternal-Fetal Medicine, the ACOG, and the CDC recommend that all eligible persons, including pregnant and lactating individuals and those planning to become pregnant, receive a COVID-19 vaccine or vaccine series.¹²⁻¹⁴ This includes booster doses, if the person is eligible. The CDC has published up-to-date guidance regarding COVID-19 vaccination, including guidance for administering vaccines to pregnant and lactating individuals.¹⁵ COVID-19 vaccines can be administered regardless of trimester and in concert with other vaccines recommended during pregnancy.¹³

Pregnant people were not included in the initial COVID-19 vaccine studies. However, a growing body of observational data supports the efficacy and safety of administering COVID-19 vaccines to this population. At this time, the mRNA vaccines BNT162b2 (Pfizer-BioNTech) and mRNA-1273

(Moderna) are recommended for pregnant or lactating individuals. The adjuvanted vaccine NVX-CoV2373 (Novavax) can also be used.^{13,14} For the most up-to-date clinical recommendations, see the <u>CDC guidelines on using COVID-19 vaccines</u>. The ACOG and the Society for Maternal-Fetal Medicine provide guidance for counseling pregnant and lactating patients about COVID-19 vaccination.^{12,13}

Efficacy

A prospective cohort study of 131 subjects at 2 academic medical centers compared the immunogenicity and reactogenicity of the mRNA COVID-19 vaccines in pregnant and lactating women and nonpregnant controls. The study also compared vaccine-generated immunity to the immune response to natural SARS-CoV-2 infection among pregnant participants.¹⁶ Maternal immunoglobulin (Ig) G antibody levels were similar after vaccination in pregnant and lactating women and in nonpregnant controls, and the antibody response did not differ by trimester of vaccination. Vaccinated pregnant women had significantly higher levels of antibodies than pregnant women who had had natural SARS-CoV-2 infection during the previous 4 to 12 weeks. In addition, maternal receipt of a COVID-19 vaccine series was protective against infant hospitalization for COVID-19 in the first 6 months of life.¹¹

Antibody Transfer to the Neonate

The available data indicate that vaccine-derived antibodies are passively transferred to the neonate during pregnancy and lactation.¹⁷ A case control study that was conducted at 20 pediatric hospitals in 17 states in the United States from July 1, 2021, to January 17, 2022, assessed the relationship between maternal vaccination with a 2-dose mRNA COVID-19 vaccine during pregnancy and pediatric hospitalization for COVID-19.¹¹ In this study, 379 infants aged <6 months were hospitalized. Among these infants, 176 had COVID-19 and were considered case infants; the remaining 203 infants did not have COVID-19 and were considered control infants. Sixteen percent of the mothers of the case infants had received 2 doses of COVID-19 vaccine during pregnancy compared with 32% of the mothers of control infants.

Maternal completion of a 2-dose primary mRNA COVID-19 vaccination series during pregnancy led to a decrease in the number of infant hospitalizations for COVID-19 during the first 6 months of life (61% decrease; 95% CI, 31% to 78%).¹¹ There were no statistically significant differences between the case infants and control infants in the presence of underlying medical conditions or the occurrence of premature birth. Of the 43 case infants who were admitted to the ICU, 88% had mothers who were unvaccinated. These data further support the CDC's recommendation for COVID-19 vaccination in people who are pregnant, breastfeeding, or trying to become pregnant or who might become pregnant in the future.¹⁵

Safety

A study that used data from 3 vaccine safety reporting systems in the United States reported that the frequency of adverse events among 35,691 vaccine recipients who identified as pregnant was similar to the frequency observed among nonpregnant patients. Local injection site pain, nausea, and vomiting were reported slightly more frequently in pregnant people than in nonpregnant people.¹⁸ Other systemic reactions were reported more frequently among nonpregnant vaccine recipients, but the overall reactogenicity profile was similar for pregnant and nonpregnant patients.

The CDC is enrolling pregnant patients in the v-safe COVID-19 Vaccine Pregnancy Registry to collect and analyze data related to COVID-19 vaccination in pregnant people and their infants.¹⁸ As of May 2, 2022, a total of 23,779 pregnant people in the United States have been enrolled. Surveillance data from 3,958 pregnant patients enrolled in the registry showed that, among 827 people who completed their pregnancies, there were no safety signals among obstetric or neonatal outcomes when rates of pregnancy

loss (spontaneous abortion or stillbirth), preterm birth, congenital anomalies, infants who were small for gestational age, and neonatal death were compared to historic incidences in the peer-reviewed literature.

Managing COVID-19 in Pregnancy

As in nonpregnant patients, SARS-CoV-2 infection in pregnant patients can present as asymptomatic/ presymptomatic disease or with a wide range of clinical manifestations, from mild symptoms that can be managed with supportive care at home to severe disease and respiratory failure that requires ICU admission. The illness severity, underlying comorbidities, and clinical status of pregnant patients who have symptoms compatible with COVID-19 should be assessed to determine whether in-person evaluation for potential hospitalization is needed.

If hospitalization is indicated, care should be provided in a facility that can conduct maternal and fetal monitoring, when appropriate. General management of COVID-19 in pregnant patients should include:

- Fetal and uterine contraction monitoring based on gestational age, when appropriate
- Individualized delivery planning
- A multispecialty, team-based approach that may include consultation with obstetric, maternal-fetal medicine, infectious disease, pulmonary-critical care, and pediatric specialists, as appropriate

In general, the recommendations for managing COVID-19 in nonpregnant patients also apply to pregnant patients.

Therapeutic Management of COVID-19 in the Setting of Pregnancy

To date, most SARS-CoV-2-related clinical trials have excluded individuals who are pregnant or lactating. In cases where lactating and pregnant individuals have been included in studies, only a small number have been enrolled. This makes providing evidence-based recommendations on the use of anti-SARS-CoV-2 therapies in these vulnerable patients difficult and potentially limits their treatment options. When possible, pregnant and lactating individuals should not be excluded from clinical trials of COVID-19 therapeutic agents or vaccines.

The Panel **recommends against** withholding COVID-19 treatments from pregnant or lactating individuals specifically because of pregnancy or lactation (**AIII**). For details regarding therapeutic recommendations and pregnancy and lactation considerations, see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>; <u>Therapeutic Management of Hospitalized Adults With COVID-19</u>; <u>Pregnancy, Lactation, and COVID-19</u> Therapeutics; and the individual drug sections.

There are limited data on the use of COVID-19 therapeutic agents in pregnant and lactating people. When making decisions about treatment, pregnant or lactating people and their clinical teams should use a shared decision-making process and consider several factors, including the severity of COVID-19, the risk of disease progression, and the safety of specific medications for the fetus, infant, or pregnant or lactating individual.

In general, the therapeutic management of pregnant patients with COVID-19 should be the same as for nonpregnant patients, with a few exceptions (AIII). Notable exceptions include:

- The Panel **recommends against** the use of **molnupiravir** for the treatment of COVID-19 in pregnant patients unless there are no other options and therapy is clearly indicated (**AIII**). For more information regarding the use of molnupiravir in pregnant patients, see <u>Pregnancy, Lactation, and COVID-19 Therapeutics</u>.
- Pregnant patients were not included in most of the clinical trials that evaluated therapeutic

COVID-19 Treatment Guidelines

anticoagulation in the setting of COVID-19, and there is a potential for increased maternal risks if bleeding occurs during pregnancy. Therefore, there is insufficient evidence for the Panel to recommend either for or against the use of therapeutic anticoagulation in pregnant patients with COVID-19 who do not have evidence of venous thromboembolism.

Timing of Delivery

The ACOG provides <u>detailed guidance</u> on the timing of delivery and the risk of vertical transmission of SARS-CoV-2.

In most cases, the timing of delivery should be dictated by obstetric indications rather than maternal diagnosis of COVID-19. For people who had suspected or confirmed COVID-19 early in pregnancy and who recovered, no alteration to the usual timing of delivery is indicated.

After Delivery

Therapeutic management in postpartum patients should follow guidelines for nonpregnant patients. However, the use of anticoagulation therapy in the immediate postpartum period should be individualized, as there may be an increased risk of bleeding, especially after an operative delivery.

The majority of studies have not demonstrated the presence of SARS-CoV-2 in breast milk; therefore, breastfeeding is not contraindicated for people with laboratory-confirmed or suspected SARS-CoV-2 infection.¹⁹ Precautions should be taken to avoid transmission to the infant, including practicing good hand hygiene, wearing face coverings, and performing proper pump cleaning before and after breast milk expression.

The decision to feed the infant breast milk while the lactating patient is receiving therapeutic agents for COVID-19 should be a collaborative effort between the patient and the clinical team, including infant care providers. The patient and the clinical team should discuss the potential benefits of the therapeutic agent and evaluate the potential risk of pausing lactation on future breast milk delivery to the infant.

Specific guidance on the postdelivery management of infants born to individuals with known or suspected SARS-CoV-2 infection, including breastfeeding recommendations, is provided by the <u>American Academy of Pediatrics</u>.

References

- 1. Zambrano LD, Ellington S, Strid P, et al. Update: characteristics of symptomatic women of reproductive age with laboratory-confirmed SARS-CoV-2 infection by pregnancy status—United States, January 22–October 3, 2020. *MMWR Morb Mortal Wkly Rep.* 2020;69(44):1641-1647. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33151921</u>.
- 2. Allotey J, Stallings E, Bonet M, et al. Clinical manifestations, risk factors, and maternal and perinatal outcomes of coronavirus disease 2019 in pregnancy: living systematic review and meta-analysis. *BMJ*. 2020;370:m3320. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32873575.
- 3. Allotey J, Stallings E, Bonet M, et al. Update to living systematic review on COVID-19 in pregnancy. *BMJ*. 2022;377:o1205. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35636775</u>.
- 4. Metz TD, Clifton RG, Hughes BL, et al. Disease severity and perinatal outcomes of pregnant patients with coronavirus disease 2019 (COVID-19). *Obstet Gynecol*. 2021;137(4):571-580. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33560778.
- 5. DeSisto CL, Wallace B, Simeone RM, et al. Risk for stillbirth among women with and without COVID-19 at delivery hospitalization—United States, March 2020–September 2021. *MMWR Morb Mortal Wkly Rep.* 2021;70(47):1640-1645. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34818318</u>.

- Metz TD, Clifton RG, Hughes BL, et al. Association of SARS-CoV-2 infection with serious maternal morbidity and mortality from obstetric complications. *JAMA*. 2022;327(8):748-759. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35129581</u>.
- Hughes BL, Sandoval GJ, Metz TD, et al. First- or second-trimester SARS-CoV-2 infection and subsequent pregnancy outcomes. *Am J Obstet Gynecol*. 2023;228(2):226.e1-226.e9. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35970201</u>.
- Dumitriu D, Emeruwa UN, Hanft E, et al. Outcomes of neonates born to mothers with severe acute respiratory syndrome coronavirus 2 infection at a large medical center in New York City. *JAMA Pediatr*. 2021;175(2):157-167. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33044493</u>.
- 9. Centers for Disease Control and Prevention. Data on COVID-19 during pregnancy: birth and infant outcomes. 2022. Available at: <u>https://stacks.cdc.gov/view/cdc/122064</u>.
- Ellington S, Strid P, Tong VT, et al. Characteristics of women of reproductive age with laboratory-confirmed SARS-CoV-2 infection by pregnancy status—United States, January 22–June 7, 2020. MMWR Morb Mortal Wkly Rep. 2020;69(25):769-775. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32584795</u>.
- Halasa NB, Olson SM, Staat MA, et al. Effectiveness of maternal vaccination with mRNA COVID-19 vaccine during pregnancy against COVID-19-associated hospitalization in infants aged <6 months—17 states, July 2021–January 2022. *MMWR Morb Mortal Wkly Rep*. 2022;71(7):264-270. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35176002</u>.
- Society for Maternal-Fetal Medicine. Provider considerations for engaging in COVID-19 vaccine counseling with pregnant and lactating patients. 2022. Available at: <u>https://www.smfm.org/covidclinical</u>. Accessed February 13, 2023.
- American College of Obstetricians and Gynecologists. COVID-19 vaccination considerations for obstetricgynecologic care. 2023. Available at: <u>https://www.acog.org/clinical/clinical-guidance/practice-advisory/</u> <u>articles/2020/12/covid-19-vaccination-considerations-for-obstetric-gynecologic-care</u>. Accessed February 9, 2023.
- 14. Centers for Disease Control and Prevention. Summary document for interim clinical considerations for use of COVID-19 vaccines currently authorized or approved in the United States. 2022. Available at: https://www.cdc.gov/vaccines/covid-19/downloads/summary-interim-clinical-considerations.pdf.
- 15. Centers for Disease Control and Prevention. COVID-19 vaccines while pregnant or breastfeeding. 2022. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/vaccines/recommendations/pregnancy.html</u>. Accessed February 9, 2023.
- 16. Gray KJ, Bordt EA, Atyeo C, et al. Coronavirus disease 2019 vaccine response in pregnant and lactating women: a cohort study. *Am J Obstet Gynecol*. 2021;225(3):303e1-303e17. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33775692</u>.
- 17. Yang YJ, Murphy EA, Singh S, et al. Association of gestational age at coronavirus disease 2019 (COVID-19) vaccination, history of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection, and a vaccine booster dose with maternal and umbilical cord antibody levels at delivery. *Obstet Gynecol.* 2022;139(3):373-380. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34963127.
- Shimabukuro TT, Kim SY, Myers TR, et al. Preliminary findings of mRNA COVID-19 vaccine safety in pregnant persons. *N Engl J Med*. 2021;384(24):2273-2282. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33882218</u>.
- 19. American College of Obstetricians and Gynecologists. COVID-19 FAQs for obstetrician-gynecologists, obstetrics. 2023. Available at: <u>https://www.acog.org/clinical-information/physician-faqs/covid-19-faqs-for-ob-gyns-obstetrics</u>. Accessed February 9, 2023.

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Pregnancy, Lactation, and COVID-19 Therapeutics

Last Updated: October 10, 2023

General Considerations

The COVID-19 Treatment Guidelines Panel (the Panel) **recommends against** withholding COVID-19 treatments or vaccination from pregnant or lactating individuals specifically because of pregnancy or lactation (**AIII**).

The decision to feed the infant breast milk while the lactating patient is receiving therapeutic agents for COVID-19 should be a collaborative effort between the patient and the clinical team, including infant care providers. The patient and the clinical team should consider the benefits of breastfeeding, the postnatal age of the infant, the need for the medication, any underlying risks of exposing the infant to the drug, and the potential adverse outcomes of COVID-19.

If a patient is receiving a treatment for COVID-19 that prevents them from feeding breast milk to their infant for a limited time, the clinical team should still encourage pumping breast milk and provide lactation support. This ensures that lactation can continue after the patient stops receiving the treatment.

While a person with COVID-19 is breastfeeding, prevention measures should be taken to avoid transmitting SARS-CoV-2 to the infant. These measures include practicing good hand hygiene, wearing face coverings, and performing proper pump cleaning before and after breast milk expression.

Table A: Recommendations for the Use of COVID-19 Therapeutics in Pregnant and Lactating People

For the Panel's recommendations on when to use the medications listed below, refer to <u>Therapeutic</u> <u>Management of Nonhospitalized Adults With COVID-19</u> and <u>Therapeutic Management of Hospitalized</u> <u>Adults With COVID-19</u>.

Drug Name	Pregnancy	Lactation
Abatacept	Recommended in hospitalized patients, if indicated. Pregnant patients and their health care providers should jointly decide whether to use abatacept during pregnancy, and the decision-making process should include a discussion of the potential risks and benefits.	Should be offered to patients who qualify for this therapy. There is minimal data on the transmission of abatacept to breastmilk. Breastfeeding may be considered while a patient receives abatacept.
Baricitinib	Recommended in hospitalized patients, if indicated. Pregnant patients and their health care providers should jointly decide whether to use baricitinib during pregnancy, and the decision-making process should include a discussion of the potential risks and benefits.	Feeding breast milk should be avoided while taking baricitinib and for 4 days after the last dose. Lactation support should be provided during this time. ^a
Dexamethasone	Recommended in hospitalized patients, if indicated.	Should be offered to patients who qualify for this therapy. Breastfeeding can continue while a patient receives dexamethasone.
Heparin (LMWH and UFH)	Recommended in hospitalized patients if indicated and if the patient does not have an obstetric-related bleeding risk (e.g., imminent delivery, bleeding complications of pregnancy) that would preclude use. See <u>Antithrombotic Therapy in Patients With COVID-19</u> for more information.	Should be offered to patients who qualify for this therapy. Breastfeeding can continue while a patient receives LMWH or UFH.

COVID-19 Treatment Guidelines

Drug Name	Pregnancy	Lactation
Infliximab	Recommended in hospitalized patients, if indicated. Pregnant patients and their health care providers should jointly decide whether to use infliximab during pregnancy, and the decision-making process should include a discussion of the potential risks and benefits.	Should be offered to patients who qualify for this therapy. The available data show that the amount of infliximab that transfers through breast milk is negligible. Breastfeeding can continue while a patient receives infliximab.
Molnupiravir	Recommended against , unless there are no other options and therapy is clearly indicated.	Breastfeeding is not recommended while a patient is taking molnupiravir and for 4 days after the last dose. ¹ Lactation support should be provided during this time. ^a
Remdesivir	Recommended, if indicated.	Should be offered to patients if indicated. Breastfeeding can continue while a patient receives remdesivir.
Ritonavir- Boosted Nirmatrelvir (Paxlovid)	Recommended, if indicated.	Should be offered to patients who qualify for this therapy. Breastfeeding can continue while a patient receives ritonavir-boosted nirmatrelvir.
Tocilizumab	Recommended in hospitalized patients, if indicated. Pregnant patients and their health care providers should jointly decide whether to use tocilizumab during pregnancy, and the decision-making process should include a discussion of the potential risks and benefits.	Should be offered to patients who qualify for this therapy. Breastfeeding can continue while a patient receives tocilizumab.

^a If a patient is receiving a treatment for COVID-19 that prevents them from feeding breast milk for a limited time, the clinical team should still encourage pumping breast milk and provide lactation support. This ensures that lactation can resume after the patient stops receiving the treatment.

Key: LMWH = low-molecular-weight heparin; the Panel = the COVID-19 Treatment Guidelines Panel; UFH = unfractionated heparin

Rationale

Abatacept

Pregnancy

As there are no data on the use of abatacept during pregnancy in hospitalized patients with COVID-19, this drug should be used only if baricitinib and tocilizumab are not available or feasible to use. When deciding whether to prescribe abatacept to a pregnant individual, clinicians need to consider the severity of the patient's COVID-19, the patient's comorbidities, and the gestational age of the fetus.

There is a paucity of data on the use of abatacept in pregnant individuals. It is currently not known whether abatacept can cross the human placenta; however, abatacept has crossed the placenta in animal studies. One study reported alterations to the immune systems of the offspring of animals that received supratherapeutic doses of abatacept throughout pregnancy.² It is not known whether the immune systems of infants who were exposed to a single dose of abatacept in utero might be impacted. Abatacept should only be used during pregnancy if the benefits clearly outweigh the potential risks. If abatacept exposure occurs during pregnancy, clinicians are encouraged to submit patient-specific information to existing pregnancy registries.

Lactation

Abatacept should be offered to patients who qualify for this therapy. It is not known whether abatacept is transferred to breast milk during lactation or whether it is absorbed systemically by the infant. Because

abatacept is a large molecule, only small amounts are thought to be transferred to breast milk. Patients who are receiving abatacept may consider breastfeeding.

Baricitinib

Pregnancy

When deciding whether to prescribe baricitinib to a pregnant individual, clinicians need to consider the severity of the patient's COVID-19, the patient's comorbidities, and the gestational age of the fetus.

Baricitinib is a Janus kinase (JAK) inhibitor. As a small-molecule drug, baricitinib is likely to pass through the placenta; therefore, fetal risk cannot be ruled out.³ In animal studies, baricitinib doses that exceeded the therapeutic human dose were associated with embryofetal developmental abnormalities. Pregnancy registries provide some data on the use of tofacitinib, another JAK inhibitor, during pregnancy for other conditions (e.g., ulcerative colitis, rheumatoid arthritis, psoriasis). Pregnancy outcomes among the participants who received tofacitinib were similar to those among the general population.⁴⁻⁶

Lactation

There is no information on the use of baricitinib in lactating people or on the effects of baricitinib on breastfed infants; however, baricitinib has been detected in the breast milk of lactating rats.⁷ Feeding breast milk should be avoided for 4 days (approximately 5–6 elimination half-lives) after baricitinib is discontinued.

Dexamethasone

Pregnancy

A short course of betamethasone or dexamethasone, which are both known to cross the placenta, is routinely used to decrease neonatal complications of prematurity in people who are at risk of imminent preterm birth.^{8,9} Treating COVID-19 with a short course of dexamethasone can lower the risk of death in pregnant individuals. In addition, dexamethasone carries a low risk of fetal adverse effects.

Lactation

Dexamethasone should be offered to lactating patients with COVID-19 who qualify for this therapy. Breast milk can be fed to the infant while the lactating patient is receiving dexamethasone. Although there are limited data on the use of dexamethasone in lactating patients, some published reports about a related antenatal corticosteroid (betamethasone) reported a time-limited decrease in the volume of breast milk production.^{10,11} Given the benefits of breast milk, additional lactation support has been recommended if needed.

Heparin (Low-Molecular-Weight and Unfractionated)

Pregnancy

In general, the preferred anticoagulants during pregnancy are heparin compounds. Because of its reliability and ease of administration, low-molecular-weight heparin is recommended rather than unfractionated heparin for the prevention and treatment of venous thromboembolism in pregnant people.

The use of anticoagulation therapy during labor and delivery requires specialized care and planning. The management of anticoagulation therapy in pregnant patients with COVID-19 should be similar to the management used for pregnant patients with other conditions.

Lactation

Low-molecular-weight heparin, unfractionated heparin, and warfarin do not accumulate in breast milk

COVID-19 Treatment Guidelines

and do not induce an anticoagulant effect in the newborn; therefore, they can be used by breastfeeding individuals who require venous thromboembolism prophylaxis or treatment.

Infliximab

Pregnancy

As there are no data on the use of infliximab during pregnancy in hospitalized patients with COVID-19, infliximab should be used only if baricitinib and tocilizumab are not available or feasible to use. When deciding whether to prescribe infliximab to a pregnant individual, clinicians need to consider the severity of the patient's COVID-19, the patient's comorbidities, and the gestational age of the fetus.

There are limited data on the use of infliximab to treat COVID-19 in pregnant patients. It has been used to treat autoimmune diseases in pregnant individuals when the benefits outweigh the potential risks. Infliximab crosses the placenta and has been detected in the serum of infants born to patients treated with infliximab during pregnancy. No adverse effects have been reported in these infants.

Lactation

Infants who are breastfed by people receiving infliximab show minimal absorption of this agent. No adverse effects have been reported in these infants.¹² Therefore, infliximab should be offered to patients who qualify. Breastfeeding can continue while a patient receives infliximab.

Molnupiravir

Pregnancy

The Panel **recommends against** the use of **molnupiravir** for the treatment of COVID-19 in pregnant patients unless there are no other options and therapy is clearly indicated (**AIII**).

The Food and Drug Administration (FDA) Emergency Use Authorization states that molnupiravir is not recommended for use in pregnant patients because fetal toxicity has been reported in animal studies of molnupiravir. However, when other therapies are not available, pregnant people with COVID-19 who are at high risk of progressing to severe disease may reasonably choose molnupiravir therapy after being fully informed of the potential risks, particularly if they are beyond the time of embryogenesis (i.e., >10 weeks' gestation). The patient should also be informed about the pregnancy surveillance program and offered the opportunity to participate.

Lactation

There is no data on the use of molnupiravir in lactating people; however, molnupiravir has been detected in the offspring of lactating rats. Molnupiravir is not authorized for use in children aged <18 years. Because the risk of adverse effects in infants is currently unknown, the FDA Emergency Use Authorization fact sheet does not recommend feeding an infant breast milk from a patient who is taking molnupiravir for the duration of the treatment course and until 4 days after the final dose.

Remdesivir

Pregnancy

While pregnant individuals were excluded from the clinical trials that evaluated the safety and efficacy of remdesivir for the treatment of COVID-19, subsequent reports on the use of remdesivir in pregnant individuals have been reassuring. Among 95 pregnant patients with moderate, severe, or critical COVID-19 who were included in a secondary analysis of data from a COVID-19 pregnancy registry in Texas, the composite maternal and neonatal outcomes were similar between those who received remdesivir (n = 39) and those who did not.¹³

A systematic review of 13 observational studies that included 113 pregnant people also reported few adverse effects of remdesivir in pregnant patients with COVID-19. The most common adverse effect was a mild elevation in transaminase levels.¹⁴

Lactation

Remdesivir is approved by the FDA for use in pediatric patients aged ≥ 28 days and weighing ≥ 3 kg. Limited data have suggested that the drug is poorly absorbed via the oral route; therefore, the levels of the drug that are absorbed when the infant ingests breast milk are low.^{15,16} One case report described a patient with COVID-19 who received remdesivir during the immediate postpartum period.¹⁶ Based on the concentration of remdesivir in the maternal serum and breast milk, the calculated milk-to-serum ratio was low. Therefore, the levels of remdesivir that would have reached a breastfed infant were estimated to be low.

Ritonavir-Boosted Nirmatrelvir (Paxlovid)

Pregnancy

Ritonavir has been used extensively during pregnancy in people with HIV and has a favorable safety profile during pregnancy. The mechanisms of action for both nirmatrelvir and ritonavir and the results of animal studies and case series suggest that this regimen can be used safely in pregnant individuals.

Two descriptive case series evaluated outcomes among pregnant patients with COVID-19 who received ritonavir-boosted nirmatrelvir. One case series included 47 patients with COVID-19 and a median gestational age of 28.4 weeks. These patients started taking ritonavir-boosted nirmatrelvir after a median duration of 1 day of COVID-19 symptoms. Thirty (64%) patients in the cohort had clinical characteristics in addition to pregnancy that increased their risk of progressing to severe COVID-19. The patients tolerated ritonavir-boosted nirmatrelvir well, with no serious adverse effects noted in either the pregnant patients or the neonates during the study period.¹⁷ The other case series included 7 patients with a mean gestational age of 26.4 weeks who initiated ritonavir-boosted nirmatrelvir after approximately 2 days of COVID-19 symptoms. One patient developed dysgeusia and stopped treatment, but the remaining 6 patients completed 5 days of treatment. Six of the patients were fully vaccinated, and 4 of these patients had also received a booster dose. All the patients reported resolution of their COVID-19 symptoms, and no fetal or neonatal adverse effects were observed during the study period.¹⁸

Ritonavir-boosted nirmatrelvir should be offered to pregnant and recently pregnant patients with COVID-19 who qualify for this therapy based on the results of a risk-benefit assessment. The risk-benefit assessment may include factors such as medical comorbidities, body mass index, vaccination status, and the number and severity of the risk factors for severe disease.

Obstetricians should be aware of potential drug-drug interactions when prescribing this agent. See <u>Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant</u> <u>Medications</u> for more information.

Lactation

Studies of infants who were exposed to ritonavir through breast milk suggest that the amount of ritonavir that transfers through breast milk is negligible and not considered clinically significant.²⁴

There are no data on the use of nirmatrelvir in lactating people. However, a prebirth-to-lactation study performed in rats reported an 8% decrease in body weight on Postnatal Day 17 in the offspring of rats that received nirmatrelvir and had systemic exposures that were 8 times higher than the clinical exposures at the authorized human dose. This reduction in body weight was not seen in the offspring of rats that had exposures that were 5 times higher than the clinical exposures at the authorized

human dose.³ Because the overall oral absorption of nirmatrelvir is poor, it is unlikely that the levels of nirmatrelvir absorbed from breast milk ingestion would be clinically relevant or expected to cause adverse effects in an infant.¹⁹

Tocilizumab

Pregnancy

Pregnant individuals have been excluded from clinical trials that evaluated the use of the antiinterleukin-6 receptor monoclonal antibody tocilizumab for the treatment of COVID-19. An analysis of data from a global safety database reported pregnancy outcomes from 288 women who were exposed to tocilizumab during their pregnancies. Eighty-nine percent of these women received tocilizumab as ongoing treatment for rheumatoid arthritis, and most were exposed to tocilizumab during their first trimester. The rates of congenital abnormalities among the infants born to these women were not higher than the rates seen in the general population. However, an increased rate of preterm birth was observed among these individuals when compared with the general population. A retrospective report of 61 pregnant women who were exposed to tocilizumab at conception or during their first trimesters showed no increased rates of congenital abnormalities or spontaneous abortion.²⁰

As pregnancy progresses, monoclonal antibodies are actively transported across the placenta, with the greatest transfer occurring during the third trimester. This may affect immune responses in the exposed fetus. If a pregnant patient receives tocilizumab after 20 weeks' gestation, clinicians should delay administering live viral vaccines to the infant for at least 6 months.

Lactation

There is limited information on the use of tocilizumab in lactating patients. Based on case report data, the amount of tocilizumab transferred to the infant via breast milk appears to be very low, with no reports of adverse effects.²¹

References

- 1. Food and Drug Administration. Fact sheet for healthcare providers: Emergency Use Authorization for Lagevrio (molnupiravir) capsules. 2023. Available at: <u>https://www.fda.gov/media/155054/download</u>.
- 2. Abatacept (Orencia) [package insert]. Food and Drug Administration. 2021. Available at: <u>https://www.accessdata.fda.gov/drugsatfda_docs/label/2021/125118s240lbl.pdf</u>.
- 3. Sammaritano LR, Bermas BL, Chakravarty EE, et al. 2020 American College of Rheumatology guideline for the management of reproductive health in rheumatic and musculoskeletal diseases. *Arthritis Rheumatol*. 2020;72(4):529-556. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32090480</u>.
- Clowse ME, Feldman SR, Isaacs JD, et al. Pregnancy outcomes in the tofacitinib safety databases for rheumatoid arthritis and psoriasis. *Drug Saf.* 2016;39(8):755-762. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/27282428</u>.
- 5. Mahadevan U, Dubinsky MC, Su C, et al. Outcomes of pregnancies with maternal/paternal exposure in the tofacitinib safety databases for ulcerative colitis. *Inflamm Bowel Dis*. 2018;24(12):2494-2500. Available at: https://www.ncbi.nlm.nih.gov/pubmed/29982686.
- 6. Wieringa JW, van der Woude CJ. Effect of biologicals and JAK inhibitors during pregnancy on healthrelated outcomes in children of women with inflammatory bowel disease. *Best Pract Res Clin Gastroenterol*. 2020;44-45:101665. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32359679.
- 7. Baricitinib (Olumiant) [package insert]. Food and Drug Administration. 2022. Available at: <u>https://www.accessdata.fda.gov/drugsatfda_docs/label/2022/207924s006lbl.pdf</u>.
- 8. Liggins GC, Howie RN. A controlled trial of antepartum glucocorticoid treatment for prevention of the respiratory distress syndrome in premature infants. *Pediatrics*. 1972;50(4):515-525. Available at: https://www.

ncbi.nlm.nih.gov/pubmed/4561295.

- Gyamfi-Bannerman C, Thom EA, Blackwell SC, et al. Antenatal betamethasone for women at risk for late preterm delivery. *N Engl J Med*. 2016;374(14):1311-1320. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/26842679</u>.
- Henderson JJ, Hartmann PE, Newnham JP, Simmer K. Effect of preterm birth and antenatal corticosteroid treatment on lactogenesis II in women. *Pediatrics*. 2008;121(1):e92-e100. Available at: https://pubmed.ncbi.nlm.nih.gov/18166549.
- Henderson JJ, Newnham JP, Simmer K, Hartmann PE. Effects of antenatal corticosteroids on urinary markers of the initiation of lactation in pregnant women. *Breastfeed Med.* 2009;4(4):201-206. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/19772378</u>.
- Ben-Horin S, Yavzori M, Kopylov U, et al. Detection of infliximab in breast milk of nursing mothers with inflammatory bowel disease. *J Crohns Colitis*. 2011;5(6):555-558. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/22115374/</u>.
- Gutierrez R, Mendez-Figueroa H, Biebighauser JG, et al. Remdesivir use in pregnancy during the SARS-CoV-2 pandemic. *J Matern Fetal Neonatal Med.* 2022;35(25):9445-9451. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35168447</u>.
- 14. Budi DS, Pratama NR, Wafa IA, et al. Remdesivir for pregnancy: a systematic review of antiviral therapy for COVID-19. *Heliyon*. 2022;8(1):e08835. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35128114</u>.
- 15. National Library of Medicine. Drugs and lactation database (LactMed) [Internet]. Remdesivir. 2023. Available at: <u>https://www.ncbi.nlm.nih.gov/books/NBK556881</u>. Accessed April 19, 2023.
- 16. Wada YS, Saito J, Hashii Y, et al. Remdesivir and human milk: a case study. *J Hum Lact*. 2022;38(2):248-251. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35189734</u>.
- 17. Garneau WM, Jones-Beatty K, Ufua MO, et al. Analysis of clinical outcomes of pregnant patients treated with nirmatrelvir and ritonavir for acute SARS-CoV-2 infection. *JAMA Netw Open*. 2022;5(11):e2244141. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36445705</u>.
- Loza A, Farias R, Gavin N, et al. Short-term pregnancy outcomes after nirmatrelvir-ritonavir treatment for mild-to-moderate coronavirus disease 2019 (COVID-19). *Obstet Gynecol*. 2022;140(3):447-449. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36356238</u>.
- Eng H, Dantonio AL, Kadar EP, et al. Disposition of nirmatrelvir, an orally bioavailable inhibitor of SARS-CoV-2 3c-like protease, across animals and humans. *Drug Metab Dispos*. 2022;50(5):576-590. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35153195</u>.
- 20. Nakajima K, Watanabe O, Mochizuki M, et al. Pregnancy outcomes after exposure to tocilizumab: a retrospective analysis of 61 patients in Japan. *Mod Rheumatol*. 2016;26(5):667-671. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/26873562</u>.
- 21. Saito J, Yakuwa N, Kaneko K, et al. Tocilizumab during pregnancy and lactation: drug levels in maternal serum, cord blood, breast milk and infant serum. *Rheumatology (Oxford)*. 2019;58(8):1505-1507. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/30945743</u>.

COVID-19 Treatment Guidelines

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Influenza and COVID-19

Last Updated: December 20, 2023

Summary Recommendations

Influenza Vaccination

- People with acute COVID-19 who have not received an influenza vaccine during influenza season should be vaccinated after they recover from acute illness and are no longer in isolation (BIII).
 - Patients may be vaccinated while they are still in isolation if they are in a health care setting.
- An influenza vaccine and a COVID-19 vaccine may be administered concurrently at different injection sites. The <u>Advisory Committee on Immunization Practices</u> and the <u>Centers for Disease Control and Prevention</u> (CDC) provide more information on COVID-19 and influenza vaccines.

Diagnosis of Influenza and COVID-19 When Influenza Viruses and SARS-CoV-2 Are Cocirculating

- Only testing can distinguish between SARS-CoV-2 and influenza virus infections and identify SARS-CoV-2 and influenza virus coinfection.
- The COVID-19 Treatment Guidelines Panel (the Panel) recommends performing influenza testing in addition to SARS-CoV-2 testing in outpatients with acute respiratory illness if the results will change the clinical management strategy for the patient (e.g., administering antiviral treatment for influenza) (BIII).
- The Panel recommends testing for both viruses in all hospitalized patients with acute respiratory illness (AIII).
- Clinicians should consider performing additional testing in specific clinical circumstances. Secondary bacterial infection is more common with influenza than with COVID-19, so additional testing for bacterial pathogens is important in patients with influenza who have clinical signs that suggest bacterial superinfection, especially for those who are immunocompromised or intubated.
- See the CDC webpage <u>Information for Clinicians on Influenza Virus Testing</u> and the Infectious Diseases Society of America (IDSA) <u>clinical practice guidelines</u> for more information.

Antiviral Treatment of Influenza When Influenza Viruses and SARS-CoV-2 Are Cocirculating

- Antiviral treatment for influenza is the same for all patients regardless of SARS-CoV-2 coinfection (AIII).
 - For information on using antiviral drugs to treat influenza in hospitalized and nonhospitalized patients, see the <u>CDC</u> and <u>IDSA</u> recommendations.
 - There are no clinically significant drug-drug interactions between the antiviral agents used to treat influenza and the antiviral agents or immunomodulators used to treat COVID-19.
- The Panel recommends starting hospitalized patients who are suspected of having influenza on empiric treatment for influenza with **oseltamivir** as soon as possible regardless of their COVID-19 status and without waiting for influenza test results (AIIb).
 - Oseltamivir treatment should be continued until nucleic acid detection assay results rule out influenza. For patients
 who are not intubated, assays should be performed on upper respiratory tract specimens. For patients who are
 intubated, assays should be performed on both upper and lower respiratory tract specimens.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Introduction

Clinicians should monitor local influenza and SARS-CoV-2 activities during influenza season to inform the evaluation and management of patients with acute respiratory illness. This can be done by tracking local and state public health surveillance data, assessing the results of testing performed at health care facilities, and reviewing the Centers for Disease Control and Prevention (CDC) <u>Weekly U.S. Influenza</u> <u>Surveillance Report</u>.

COVID-19 Treatment Guidelines

Influenza Vaccination

For Patients With Acute COVID-19 or Those Recovering From COVID-19

The Advisory Committee on Immunization Practices (ACIP) recommends offering an influenza vaccine by the end of October to all people aged ≥ 6 months in the United States.¹ Unvaccinated persons can still benefit from influenza vaccination after October as long as influenza viruses are still circulating in the community. People with acute COVID-19 who have not received an influenza vaccine should be vaccinated after they recover from acute illness and are no longer in isolation (**BIII**). Patients may be vaccinated while they are still in isolation if they are in a health care setting.

There are currently no data on the safety, immunogenicity, or efficacy of administering influenza vaccines to patients with acute COVID-19 or those who are recovering from COVID-19. Vaccination in people who have mild illness is safe and effective.² Clinicians should consider deferring influenza vaccination for symptomatic patients with moderate or severe COVID-19 until they have recovered and completed their COVID-19 isolation period. It is not known whether administering dexamethasone or other immunomodulatory therapies to patients with severe COVID-19 will affect the immune response to the influenza vaccine. People with asymptomatic SARS-CoV-2 infection or mild COVID-19 should seek influenza vaccination when they no longer require isolation. They may be vaccinated sooner if they are in a health care setting for other reasons. See the influenza vaccine recommendations from the <u>CDC</u> and the <u>American Academy of Pediatrics</u>.

Coadministration of COVID-19 Vaccines and Influenza Vaccines

Coadministration of a COVID-19 vaccine and an influenza vaccine at different injection sites has been shown to be safe.³⁻⁶ Providers and patients should be aware of a potential increase in reactogenicity when both vaccines are administered concurrently. The <u>CDC</u> and <u>ACIP</u> provide more information on coadministering influenza and COVID-19 vaccines.

Clinical Presentation of Influenza Versus COVID-19

The signs and symptoms of uncomplicated, clinically mild influenza overlap with those of mild COVID-19. Loss of taste and smell can occur with both diseases, but these symptoms are more common with COVID-19 than with influenza. Fever is not always present in patients with either disease, particularly in young infants, adults of advanced age, and patients who are immunosuppressed. Complications of influenza and COVID-19 can be similar, but the onset of influenza complications and severe disease typically occurs within a week of illness, whereas the onset of severe COVID-19 usually occurs in the second week of illness.

Because of the overlap in signs and symptoms, when SARS-CoV-2 and influenza viruses are cocirculating, diagnostic testing for both viruses is needed to distinguish between SARS-CoV-2 and influenza virus infection and to identify coinfection in people with an acute respiratory illness. Coinfection with influenza virus and SARS-CoV-2 has been described in case reports and case series,⁷⁻¹¹ but it is uncommon.¹² Observational studies have reported greater disease severity in adult patients with influenza virus and SARS-CoV-2 coinfection than in those with SARS-CoV-2 infection alone.^{13,14} In pediatric patients, coinfection with the 2 viruses was associated with greater disease severity than infection with influenza virus alone.¹⁵

Testing for SARS-CoV-2 and Influenza

The COVID-19 Treatment Guidelines Panel (the Panel) recommends performing influenza testing in addition to SARS-CoV-2 testing in outpatients with acute respiratory illness if the results will change the clinical management strategy for the patient (e.g., administering antiviral treatment for influenza) (**BIII**).

The Panel recommends testing for both viruses in all hospitalized patients with acute respiratory illness (AIII).

Several multiplex molecular assays and multiplex antigen assays that detect SARS-CoV-2 and influenza A and B viruses have received Food and Drug Administration Emergency Use Authorizations or De Novo classifications and can provide results in 15 minutes to 8 hours using a single respiratory specimen.¹⁶⁻¹⁸ For more information, see the CDC webpage Information for Clinicians on Influenza Virus Testing and the recommendations from the Infectious Diseases Society of America (IDSA) on the use of influenza tests and the interpretation of test results.

Treating Influenza With Antiviral Agents

Antiviral treatment for influenza is the same for all patients regardless of SARS-CoV-2 coinfection (AIII). There are no clinically significant drug-drug interactions between the antiviral agents used to treat influenza and the antiviral agents or immunomodulators used to treat COVID-19. The IDSA recommends administering antiviral treatment for influenza to all hospitalized patients with influenza.¹⁹

The Panel recommends starting hospitalized patients who are suspected of having influenza on empiric treatment for influenza with **oseltamivir** as soon as possible regardless of their COVID-19 status and without waiting for influenza test results (**AIIb**). Oseltamivir has no activity against SARS-CoV-2.²⁰ The standard dose of oseltamivir is absorbed well, even in critically ill patients. For patients who cannot tolerate oral or enterically administered oseltamivir (e.g., because of gastric stasis, malabsorption, or gastrointestinal bleeding), intravenous peramivir is an option.¹⁹ There are no data on the activity of peramivir against SARS-CoV-2.

See the CDC webpage <u>Influenza Antiviral Medications: Summary for Clinicians</u> for clinical algorithms for using antiviral agents in patients with suspected or laboratory-confirmed influenza, including pregnant people and other people who are at high risk for influenza complications. The IDSA clinical practice guidelines also provide recommendations on using antiviral agents to treat influenza,¹⁹ and the American Academy of Pediatrics provides recommendations on the antiviral treatment of influenza in children.²¹

When the result of an influenza nucleic acid detection assay from an upper respiratory tract specimen is negative in a patient who is receiving antiviral treatment for influenza:

- In a patient who is not intubated: Antiviral treatment for influenza can be stopped.
- *In a patient who is intubated:* Antiviral treatment for influenza should be continued, and a lower respiratory tract specimen (e.g., endotracheal aspirate) should be collected and tested using an influenza nucleic acid detection assay. If the lower respiratory tract specimen is also negative, antiviral treatment for influenza can be stopped.

COVID-19 Treatment Considerations for Hospitalized Patients With Suspected or Confirmed Influenza Virus Coinfection

Corticosteroids, which are used to treat patients with severe COVID-19, may prolong influenza viral replication and may be associated with poor outcomes for influenza.^{19,22} Currently, no data are available on the use of corticosteroids in patients with SARS-CoV-2 and influenza virus coinfection. However, because dexamethasone has demonstrated substantial benefits in patients with COVID-19 who require supplemental oxygen, the benefits of using corticosteroids in patients with severe SARS-CoV-2 and influenza virus coinfection likely outweigh any potential harm.

Although severe influenza may be associated with a dysregulated innate immune response, there are

no data on the use of immunomodulatory therapies, such as interleukin-6 inhibitors (e.g., tocilizumab, sarilumab) or Janus kinase inhibitors (e.g., baricitinib, tofacitinib), for the treatment of severe influenza. There are also no data on the effects these therapies may have on influenza virus infection, such as potentially prolonging viral replication. These immunomodulators have demonstrated a clinical benefit in certain patients with COVID-19. When considering using these drugs in patients with COVID-19 who have suspected or laboratory-confirmed influenza, clinicians should carefully weigh the known benefits for treatment of severe COVID-19 against the unknown theoretical risks for patients with influenza.

Observational studies have reported that co-occurrence of community-acquired secondary bacterial pneumonia appears to be infrequent in people with COVID-19; it is more common in people who have influenza.²³⁻²⁸ Typical bacterial causes of community-acquired pneumonia with severe influenza are *Staphylococcus aureus* (both methicillin-resistant *S. aureus* [MRSA] and methicillin-susceptible *S. aureus* [MSSA]), *Streptococcus pneumoniae*, and group A *Streptococcus*.¹⁹

Patients with COVID-19 who develop new respiratory symptoms with or without fever or respiratory distress and who do not have a clear diagnosis should be evaluated for the possibility of nosocomial influenza.

References

- 1. Grohskopf LA, Blanton LH, Ferdinands JM, et al. Prevention and control of seasonal influenza with vaccines: recommendations of the Advisory Committee on Immunization Practices—United States, 2023–24 influenza season. *MMWR Recomm Rep.* 2023;72(2):1-25. Available at: <u>https://www.cdc.gov/mmwr/volumes/72/rr/rr7202a1.htm</u>.
- 2. Centers for Disease Control and Prevention. Contraindications and precautions. 2023. Available at: <u>https://www.cdc.gov/vaccines/hcp/acip-recs/general-recs/contraindications.html</u>. Accessed December 18, 2023.
- 3. Izikson R, Brune D, Bolduc JS, et al. Safety and immunogenicity of a high-dose quadrivalent influenza vaccine administered concomitantly with a third dose of the mRNA-1273 SARS-CoV-2 vaccine in adults aged ≥65 years: a Phase 2, randomised, open-label study. *Lancet Respir Med.* 2022;10(4):392-402. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35114141.
- Lazarus R, Baos S, Cappel-Porter H, et al. Safety and immunogenicity of concomitant administration of COVID-19 vaccines (ChAdOx1 or BNT162b2) with seasonal influenza vaccines in adults in the UK (ComFluCOV): a multicentre, randomised, controlled, Phase 4 trial. *Lancet*. 2021;398(10318):2277-2287. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34774197</u>.
- Hause AM, Zhang B, Yue X, et al. Reactogenicity of simultaneous COVID-19 mRNA booster and influenza vaccination in the US. *JAMA Netw Open*. 2022;5(7):e2222241. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35838667</u>.
- Janssen C, Mosnier A, Gavazzi G, et al. Coadministration of seasonal influenza and COVID-19 vaccines: a systematic review of clinical studies. *Hum Vaccin Immunother*. 2022;18(6):2131166. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36256633</u>.
- Hashemi SA, Safamanesh S, Ghasemzadeh-Moghaddam H, Ghafouri M, Azimian A. High prevalence of SARS-CoV-2 and influenza A virus (H1N1) coinfection in dead patients in Northeastern Iran. *J Med Virol*. 2021;93(2):1008-1012. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32720703</u>.
- Huang BR, Lin YL, Wan CK, et al. Co-infection of influenza B virus and SARS-CoV-2: a case report from Taiwan. *J Microbiol Immunol Infect*. 2021;54(2):336-338. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32646801</u>.
- 9. Yue H, Zhang M, Xing L, et al. The epidemiology and clinical characteristics of co-infection of SARS-CoV-2 and influenza viruses in patients during COVID-19 outbreak. *J Med Virol*. 2020;92(11):2870-2873. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32530499.

- 10. Cuadrado-Payán E, Montagud-Marrahi E, Torres-Elorza M, et al. SARS-CoV-2 and influenza virus coinfection. *Lancet*. 2020;395(10236):e84. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32423586</u>.
- 11. Wu X, Cai Y, Huang X, et al. Co-infection with SARS-CoV-2 and influenza A virus in patient with pneumonia, China. *Emerg Infect Dis*. 2020;26(6):1324-1326. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32160148</u>.
- Pawlowski C, Silvert E, O'Horo JC, et al. SARS-CoV-2 and influenza coinfection throughout the COVID-19 pandemic: an assessment of coinfection rates, cohort characteristics, and clinical outcomes. *PNAS Nexus*. 2022;1(3):pgac071. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35860600</u>.
- Stowe J, Tessier E, Zhao H, et al. Interactions between SARS-CoV-2 and influenza, and the impact of coinfection on disease severity: a test-negative design. *Int J Epidemiol*. 2021;50(4):1124-1133. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33942104</u>.
- Swets MC, Russell CD, Harrison EM, et al. SARS-CoV-2 co-infection with influenza viruses, respiratory syncytial virus, or adenoviruses. *Lancet*. 2022;399(10334):1463-1464. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35344735</u>.
- 15. Adams K, Tastad KJ, Huang S, et al. Prevalence of SARS-CoV-2 and influenza coinfection and clinical characteristics among children and adolescents aged <18 years who were hospitalized or died with influenza—United States, 2021–22 influenza season. *MMWR Morb Mortal Wkly Rep.* 2022;71(50):1589-1596. Available at: https://www.ncbi.nlm.nih.gov/pubmed/36520656.
- 16. Food and Drug Administration. In vitro diagnostics EUAs—molecular diagnostic tests for SARS-CoV-2. 2023. Available at: <u>https://www.fda.gov/medical-devices/coronavirus-disease-2019-covid-19-emergency-use-authorizations-medical-devices/in-vitro-diagnostics-euas-molecular-diagnostic-tests-sars-cov-2</u>. Accessed December 18, 2023.
- Food and Drug Administration. In vitro diagnostics EUAs—antigen diagnostic tests for SARS-CoV-2. 2023. Available at: <u>https://www.fda.gov/medical-devices/coronavirus-disease-2019-covid-19-emergency-use-authorizations-medical-devices/in-vitro-diagnostics-euas-antigen-diagnostic-tests-sars-cov-2</u>. Accessed December 18, 2023.
- Food and Drug Administration. Evaluation of automatic class III designation (De Novo) summaries. 2023. Available at: <u>https://www.fda.gov/about-fda/cdrh-transparency/evaluation-automatic-class-iii-designation-de-novo-summaries</u>. Accessed December 18, 2023.
- Uyeki TM, Bernstein HH, Bradley JS, et al. Clinical practice guidelines by the Infectious Diseases Society of America: 2018 update on diagnosis, treatment, chemoprophylaxis, and institutional outbreak management of seasonal influenza. *Clin Infect Dis*. 2019;68(6):895-902. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/30834445</u>.
- 20. Choy KT, Wong AY, Kaewpreedee P, et al. Remdesivir, lopinavir, emetine, and homoharringtonine inhibit SARS-CoV-2 replication in vitro. *Antiviral Res.* 2020;178:104786. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32251767.
- Committee on Infectious Diseases. Recommendations for prevention and control of influenza in children, 2023–2024. *Pediatrics*. 2023;152(4):e2023063772. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/37641879</u>.
- 22. Zhou Y, Fu X, Liu X, et al. Use of corticosteroids in influenza-associated acute respiratory distress syndrome and severe pneumonia: a systemic review and meta-analysis. *Sci Rep.* 2020;10(1):3044. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32080223.
- 23. Vaughn VM, Gandhi TN, Petty LA, et al. Empiric antibacterial therapy and community-onset bacterial coinfection in patients hospitalized with coronavirus disease 2019 (COVID-19): a multi-hospital cohort study. *Clin Infect Dis.* 2021;72(10):e533-e541. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32820807</u>.
- 24. Adler H, Ball R, Fisher M, Mortimer K, Vardhan MS. Low rate of bacterial co-infection in patients with COVID-19. *Lancet Microbe*. 2020;1(2):e62. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32835331</u>.

- 25. Russell CD, Fairfield CJ, Drake TM, et al. Co-infections, secondary infections, and antimicrobial use in patients hospitalised with COVID-19 during the first pandemic wave from the ISARIC WHO CCP-UK study: a multicentre, prospective cohort study. *Lancet Microbe*. 2021;2(8):e354-e365. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34100002</u>.
- 26. Hedberg P, Johansson N, Ternhag A, et al. Bacterial co-infections in community-acquired pneumonia caused by SARS-CoV-2, influenza virus and respiratory syncytial virus. *BMC Infect Dis*. 2022;22(1):108. Available at: https://www.ncbi.nlm.nih.gov/pubmed/35100984.
- 27. Pandey M, May A, Tan L, et al. Comparative incidence of early and late bloodstream and respiratory tract coinfection in patients admitted to ICU with COVID-19 pneumonia versus influenza A or B pneumonia versus no viral pneumonia: Wales multicentre ICU cohort study. *Crit Care*. 2022;26(1):158. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/35655224</u>.
- 28. Rouzé A, Martin-Loeches I, Povoa P, et al. Early bacterial identification among intubated patients with COVID-19 or influenza pneumonia: a European multicenter comparative clinical trial. Am J Respir Crit Care Med. 2021;204(5):546-556. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34038699</u>.

Special Considerations in People With HIV

Last Updated: November 2, 2023

Summary Recommendations

Prevention of COVID-19

- The COVID-19 Treatment Guidelines Panel (the Panel) recommends that people with HIV receive COVID-19 vaccines, regardless of their CD4 T lymphocyte (CD4) cell count or HIV viral load, because the potential benefits outweigh the potential risks (AIIb).
- People with HIV should receive booster doses of COVID-19 vaccines as recommended by the Centers for Disease Control and Prevention (CDC).
- For people with untreated or advanced HIV, the Panel recommends following the most recent <u>COVID-19 vaccination</u> <u>schedule</u> from the CDC for people who are moderately or severely immunocompromised. Advanced HIV is defined by the CDC as people with CD4 counts <200 cells/mm³, a history of an AIDS-defining illness without immune reconstitution, or clinical manifestations of symptomatic HIV.

Diagnosis of SARS-CoV-2 Infection

• The Panel recommends using the same approach for diagnosing SARS-CoV-2 infection in people with HIV as in people without HIV (AIII).

Management of COVID-19

- The recommendations for the triage, management, and treatment of COVID-19 in people with HIV are the same as those for the general population (AIII).
- Nonhospitalized people with HIV and mild to moderate COVID-19 may be eligible to receive the therapies that are currently recommended for treatment (see <u>Therapeutic Management of Nonhospitalized Adults With COVID-19</u>). However, in situations where there are logistical constraints for administering these therapies, priority should be given to those with untreated or advanced HIV (AIII). See <u>Prioritization of Anti-SARS-CoV-2 Therapies for the Treatment of COVID-19</u> in Nonhospitalized Patients When There Are Logistical Constraints for details.
- People with HIV who are receiving ritonavir-based or cobicistat-based antiretroviral therapy (ART) can receive the 5-day course of ritonavir-boosted nirmatrelvir (Paxlovid) to treat COVID-19 without altering or interrupting their ART (i.e., they can continue using the ritonavir or cobicistat dose associated with their ART in addition to the dose of ritonavir used with nirmatrelvir).
- In patients with advanced HIV who have suspected or laboratory-confirmed SARS-CoV-2 infection, clinicians should consider HIV-associated opportunistic infections in the differential diagnosis of clinical symptoms and consider consulting an HIV specialist.
- When starting treatment for COVID-19 in patients with HIV, clinicians should pay careful attention to potential drug-drug interactions and overlapping toxicities among COVID-19 treatments, antiretroviral (ARV) medications, antimicrobial therapies, and other medications.

Management of HIV

- People with HIV who develop COVID-19, including those who require hospitalization, should continue their ART and opportunistic infection treatment and prophylaxis whenever possible.
- Clinicians treating COVID-19 in people with HIV should consult an HIV specialist before adjusting or switching a patient's ARV medications.
- An ARV regimen should not be modified for the purpose of preventing or treating SARS-CoV-2 infection.
- Clinicians should consult an HIV specialist to determine the optimal time to initiate ART in people who present with COVID-19 and untreated HIV.

Each recommendation in the Guidelines receives a rating for the strength of the recommendation (A, B, or C) and a rating for the evidence that supports it (I, IIa, IIb, or III). See <u>Guidelines Development</u> for more information.

Introduction

Approximately 1.2 million people in the United States are living with HIV. Most of these individuals are in care, and many are receiving antiretroviral therapy (ART) and have well-controlled diseaspanele.¹ Similar to COVID-19, HIV disproportionately affects racial and ethnic minorities and people living in low-income settings in the United States; these demographic groups also appear to have a higher risk of poor outcomes for COVID-19.² Many people with HIV have 1 or more comorbidities or conditions that may put them at higher risk of severe COVID-19.³

Information on SARS-CoV-2/HIV coinfection is evolving. The sections below outline the current knowledge regarding preventing, diagnosing, and treating SARS-CoV-2 infection in people with HIV and managing HIV during the COVID-19 pandemic.

Clinical Outcomes of COVID-19 in People With HIV

In some of the initial case series of people with COVID-19 in Europe and the United States, no significant differences were observed in the clinical outcomes of COVID-19 between people with HIV and people who did not have HIV.⁴⁻¹¹ Several subsequent studies have reported worse outcomes for patients with HIV and COVID-19, especially in patients with CD4 T lymphocyte (CD4) cell counts <200 cells/mm³.¹²⁻¹⁸ Many of these studies were done before the widespread use of COVID-19 vaccines; however, people with advanced HIV may have a suboptimal response to vaccines.^{19,20}

Prevention of COVID-19 in People With HIV

People with HIV should be advised to use the same strategies for preventing SARS-CoV-2 infection that are recommended for people without HIV (**AIII**). There is currently no clear evidence that antiretroviral (ARV) medications can prevent SARS-CoV-2 infection. Some studies suggested that tenofovir disoproxil fumarate/emtricitabine may play a role in preventing SARS-CoV-2 acquisition or hospitalization or death associated with COVID-19; however, the significance of these findings is unclear. These studies may not have adequately controlled for confounding variables such as age and comorbidities. In addition, most of these studies were conducted in unvaccinated patients.²¹⁻²³

The COVID-19 Treatment Guidelines Panel (the Panel) recommends that people with HIV receive COVID-19 vaccines, regardless of their CD4 count or HIV viral load, because the potential benefits outweigh the potential risks (**AIIb**). People with HIV were included in the clinical trials of the 2 mRNA vaccines (Pfizer and Moderna) and the glycoprotein vaccine (Novavax) that are currently available through Emergency Use Authorizations and/or approval from the Food and Drug Administration (FDA).²⁴⁻²⁶ Typically, people with HIV who are receiving ART and who have achieved virologic suppression respond well to licensed vaccines. Data from studies that used COVID-19 vaccines in people with HIV confirm that people who are receiving ART and have normal CD4 counts have good immunologic responses to the vaccines.²⁷⁻²⁹ However, vaccine response rates are generally lower in people with lower CD4 counts (e.g., <200 cells/mm³).^{19,20,30}

For people with untreated or advanced HIV, the Panel recommends following the most recent <u>COVID-19 vaccination schedule</u> from the Centers for Disease Control and Prevention (CDC) for people who are moderately or severely immunocompromised. Advanced HIV is defined by the CDC as people with CD4 counts <200 cells/mm³, a history of an AIDS-defining illness without immune reconstitution, or clinical manifestations of symptomatic HIV. Patients who have poor adherence or who experience virologic failure while on ART may have a similar risk of severe COVID-19 as those with untreated HIV. For additional considerations regarding vaccination in people who are immunocompromised, see <u>Special Considerations in People Who Are Immunocompromised</u>.

Diagnostic and Laboratory Testing for COVID-19 in People With HIV

Diagnosis of SARS-CoV-2 Infection in People With HIV

The Panel recommends using the same approach for diagnosing SARS-CoV-2 infection in people with HIV as in people without HIV (**AIII**). See <u>Testing for SARS-CoV-2 Infection</u> for more information. There is currently no evidence that the performance characteristics of nucleic acid amplification tests (NAATs) and antigen tests differ in people with and without HIV when diagnosing acute SARS-CoV-2 infection. The Panel **recommends against** the use of serologic testing as the sole basis for diagnosis of acute SARS-CoV-2 infection (**AIII**). However, if diagnostic serologic testing is performed in a patient with HIV, the results should be interpreted with caution because cross-reactivity between antibodies to SARS-CoV-2 and HIV has been reported.³¹

Correlation of CD4 Count in People With HIV and COVID-19

The normal range for CD4 counts in healthy adults is about 500 to 1,600 cells/mm³. People with HIV who have a CD4 count of \geq 500 cells/mm³ have similar cellular immune function to those without HIV. In people with HIV, a CD4 count <200 cells/mm³ meets the definition for AIDS. For patients receiving ART, the hallmark of treatment success is a plasma HIV RNA measurement below the level of detection by a polymerase chain reaction assay. Lymphopenia is a common laboratory finding in patients with COVID-19; therefore, in patients with HIV, clinicians should note that CD4 counts obtained during acute COVID-19 may not accurately reflect the patient's HIV disease stage.

There have been some reports of people with advanced HIV who have presented with COVID-19 and another coinfection, including *Pneumocystis jirovecii* pneumonia and other opportunistic infections.³²⁻³⁶ In patients with advanced HIV who have suspected or laboratory-confirmed SARS-CoV-2 infection, clinicians should consider HIV-associated opportunistic infections in the differential diagnosis of clinical symptoms and consider consulting an HIV specialist.

Clinical Presentation of COVID-19 in People With HIV

It is currently unknown whether people with HIV have a higher incidence of SARS-CoV-2 infection or a higher rate of progression to symptomatic disease than the general population. Approximately 50% of people with HIV in the United States are aged >50 years,³⁷ and many have comorbidities that are associated with more severe COVID-19. These comorbidities include hypertension, diabetes mellitus, cardiovascular disease, a history of smoking, chronic lung disease, chronic liver disease, and cancer.³⁸

There are a number of case reports and case series that describe the clinical presentation of COVID-19 in people with HIV.^{4-11,21,39} These studies indicate that the clinical presentation of COVID-19 is similar in people with and without HIV. Most of the published reports describe populations in which the majority of individuals with HIV are receiving ART and have achieved virologic suppression. Consequently, the current understanding of the impact of COVID-19 in people with advanced HIV and low CD4 counts or persistent HIV viremia is limited.

Managing COVID-19 in People With HIV

The recommendations for the triage and management of COVID-19 in people with HIV are the same as those for the general population (AIII).

The treatment of COVID-19 in people with HIV is the same as for those without HIV (**AIII**). Nonhospitalized people with HIV and mild to moderate COVID-19 may be eligible to receive the therapies that are currently recommended for treatment (see <u>Therapeutic Management of</u> <u>Nonhospitalized Adults With COVID-19</u>). However, in situations where there are logistical constraints

for administering these therapies, priority should be given to those with untreated or advanced HIV infection (AIII).

When starting treatment for COVID-19 in patients with HIV, clinicians should pay careful attention to potential drug-drug interactions and overlapping toxicities among COVID-19 treatments, ARV medications, antimicrobial therapies, and other medications. The therapeutic options for nonhospitalized patients with HIV who present with mild to moderate COVID-19 include ritonavir-boosted nirmatrelvir (Paxlovid), intravenous remdesivir, and molnupiravir.

Drug-drug interactions are a special concern with ritonavir-boosted nirmatrelvir. People with HIV who are receiving ritonavir-based or cobicistat-based ART can receive the 5-day course of ritonavir-boosted nirmatrelvir to treat COVID-19 without altering or interrupting their ART (i.e., they can continue using the ritonavir or cobicistat dose associated with their ART in addition to the dose of ritonavir used with nirmatrelvir). Before prescribing ritonavir-boosted nirmatrelvir for a patient who is not already on a ritonavir-based or cobicistat-based regimen, clinicians should carefully review the patient's concomitant medications, including over-the-counter medicines and herbal supplements, and evaluate the potential for drug-drug interactions. Clinicians should utilize resources such as Drug-Drug Interactions Between Ritonavir-Boosted Nirmatrelvir (Paxlovid) and Concomitant Medications, the FDA prescribing information for ritonavir-boosted nirmatrelvir, and the Liverpool COVID-19 Drug Interactions website for additional guidance on identifying and managing drug-drug interactions.

In hospitalized patients, the appropriate treatment strategy depends on disease severity (see <u>Therapeutic</u> <u>Management of Hospitalized Adults With COVID-19</u>)</u>. Dexamethasone, which is recommended for use in combination with baricitinib or tocilizumab for some patients with severe or critical COVID-19, is an immunosuppressive agent. The safety of using this drug in patients who are immunocompromised, including those with advanced HIV, has not been studied. Therefore, patients with advanced HIV who are receiving dexamethasone should be closely monitored for secondary infections.

Dexamethasone is a dose-dependent inducer of cytochrome P450 3A4 and could potentially lower the levels of certain co-administered ARV drugs. More than a single dose of dexamethasone **is not recommended** for patients receiving rilpivirine as part of their ARV regimen. Clinicians should consult an HIV specialist before administering dexamethasone to these patients. It is currently unknown whether administering ≤ 10 days of dexamethasone impacts the clinical efficacy of other ARV drugs. Patients with HIV who are receiving dexamethasone as treatment for COVID-19 should follow up with their HIV providers to assess their virologic response.

Although some ARV drugs were studied early in the pandemic for the treatment of COVID-19, none of these agents have been shown to be effective.

Managing HIV in People With COVID-19

People with HIV who develop COVID-19, including those who require hospitalization, should continue their ART and their medications for the treatment or prevention of opportunistic infections whenever possible. If a patient with HIV needs to receive the next dose of the long-acting injectables cabotegravir/ rilpivirine, ibalizumab, or lenacapavir while hospitalized for COVID-19, clinicians should make arrangements with the patient's hospital provider to continue administering the medication without interruption. ART interruption may lead to rebound viremia, and, in some cases, the emergence of drug resistance. If the appropriate ARV drugs are not on the hospital's formulary, administer medications from the patient's home supplies, if available.

Clinicians treating COVID-19 in people with HIV should consult an HIV specialist before adjusting or switching a patient's ARV medications. An ARV regimen should not be modified for the purpose

of preventing or treating SARS-CoV-2 infection. Many drugs, including some ARV agents (e.g., lopinavir/ritonavir, boosted darunavir, tenofovir disoproxil fumarate/emtricitabine), have been or are being evaluated in clinical trials or are prescribed off-label to treat or prevent SARS-CoV-2 infection. Lopinavir/ritonavir and darunavir/cobicistat have not been found to be effective for the treatment of COVID-19.^{40,41}

For patients receiving an investigational ARV medication as part of their ARV regimen, arrangements should be made with the investigational study team to continue the medication, if possible.

For critically ill patients who require tube feeding, some ARV medications are available in liquid formulations, and some ARV pills may be crushed. Clinicians should consult an HIV specialist and/or pharmacist to assess the best way to continue an effective ARV regimen for a patient with a feeding tube. Information may be available in the drug product label or from the <u>Toronto General Hospital</u>.

For people who present with COVID-19 and have either a new diagnosis of HIV or a history of HIV but are not receiving ART, the optimal time to start or restart ART is currently unknown. For people with HIV who have not initiated ART or who have been off therapy for >2 weeks before presenting with COVID-19, an HIV specialist should be consulted about initiating or reinitiating ART as soon as clinically feasible. If ART is initiated, maintaining treatment and linking patients to HIV care upon hospital discharge is critical. If an HIV specialist is not available, clinical consultation is available by phone through the <u>National Clinician Consultation Center</u>, Monday through Friday, 9 am to 8 pm EST.

Considerations in Pregnant and Lactating People

Pregnant or recently pregnant individuals are at a higher risk of severe illness and death from COVID-19 than nonpregnant individuals (see <u>Special Considerations During Pregnancy and After Delivery</u>). Although the data on pregnancy and maternal outcomes in individuals who have COVID-19 and HIV are limited, a prospective meta-analysis demonstrated that individuals with COVID-19 and HIV had a 67% greater risk of being admitted to the intensive care unit and a 72% greater risk of needing critical care.⁴² An observational study from Botswana found that offspring who were exposed to both HIV and SARS-CoV-2 had a high prevalence of adverse birth outcomes.⁴³

Given the severity of COVID-19 in pregnant or recently pregnant individuals, COVID-19 vaccines should be offered to all pregnant and lactating individuals and to those who are planning to become pregnant, including those who are also living with HIV. Pregnant individuals with HIV who have COVID-19 should be triaged, managed, and treated the same way as pregnant individuals without HIV. Clinicians should consider any additional comorbidities when assessing the risk of severe COVID-19 in these patients. See Pregnancy, Lactation, and COVID-19 Therapeutics for information regarding the therapies recommended for the treatment of COVID-19.

Pregnant individuals with HIV who are hospitalized for COVID-19 should continue their ART and opportunistic infection treatment and prophylaxis. Clinicians should consult an HIV specialist if any changes to ARV regimens are needed.

Considerations in Children

In general, children appear less likely to become severely ill with COVID-19 than adults. In the few publications that have described cases of COVID-19 among children or adolescents with HIV, most cases were mild, and HIV did not appear to be an independent predictor of severe COVID-19.⁴⁴⁻⁴⁷ Children with HIV who are eligible should receive COVID-19 vaccines and booster doses regardless of their CD4 count or viral load. Children with HIV and COVID-19 or MIS-C should receive the same treatment as children without HIV. See <u>Therapeutic Management of Hospitalized Children With</u>

<u>COVID-19</u>, <u>Therapeutic Management of Nonhospitalized Children With COVID-19</u>, and <u>Therapeutic</u> <u>Management of Hospitalized Children With MIS-C</u>, <u>Plus a Discussion on MIS-A</u> for more information.

Parents of children with HIV and COVID-19 should be advised to continue their child's ART without interruption if the child is being managed at home. For children with HIV who are hospitalized for COVID-19, ART should be continued for the duration of hospitalization.

References

- Harris NS, Johnson AS, Huang YA, et al. Vital signs: status of human immunodeficiency virus testing, viral suppression, and HIV preexposure prophylaxis—United States, 2013-2018. *MMWR Morb Mortal Wkly Rep.* 2019;68(48):1117-1123. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/31805031</u>.
- Meyerowitz EA, Kim AY, Ard KL, et al. Disproportionate burden of coronavirus disease 2019 among racial minorities and those in congregate settings among a large cohort of people with HIV. *AIDS*. 2020;34(12):1781-1787. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32604138</u>.
- Centers for Disease Control and Prevention. Underlying medical conditions associated with higher risk for severe COVID-19: information for healthcare professionals. 2023. Available at: <u>https://www.cdc.gov/coronavirus/2019-ncov/hcp/clinical-care/underlyingconditions.html</u>. Accessed July 6, 2023.
- 4. Gervasoni C, Meraviglia P, Riva A, et al. Clinical features and outcomes of patients with human immunodeficiency virus with COVID-19. *Clin Infect Dis*. 2020;71(16):2276-2278. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32407467</u>.
- Härter G, Spinner CD, Roider J, et al. COVID-19 in people living with human immunodeficiency virus: a case series of 33 patients. *Infection*. 2020;48(5):681-686. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32394344</u>.
- Karmen-Tuohy S, Carlucci PM, Zervou FN, et al. Outcomes among HIV-positive patients hospitalized with COVID-19. *J Acquir Immune Defic Syndr*. 2020;85(1):6-10. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32568770</u>.
- Patel VV, Felsen UR, Fisher M, et al. Clinical outcomes and inflammatory markers by HIV serostatus and viral suppression in a large cohort of patients hospitalized with COVID-19. *J Acquir Immune Defic Syndr*. 2021;86(2):224-230. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33433966</u>.
- 8. Shalev N, Scherer M, LaSota ED, et al. Clinical characteristics and outcomes in people living with human immunodeficiency virus hospitalized for coronavirus disease 2019. *Clin Infect Dis.* 2020;71(16):2294-2297. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32472138</u>.
- 9. Sigel K, Swartz T, Golden E, et al. Coronavirus 2019 and people living with human immunodeficiency virus: outcomes for hospitalized patients in New York City. *Clin Infect Dis*. 2020;71(11):2933-2938. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32594164</u>.
- 10. Stoeckle K, Johnston CD, Jannat-Khah DP, et al. COVID-19 in hospitalized adults with HIV. *Open Forum Infect Dis.* 2020;7(8):ofaa327. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32864388</u>.
- Vizcarra P, Pérez-Elías MJ, Quereda C, et al. Description of COVID-19 in HIV-infected individuals: a singlecentre, prospective cohort. *Lancet HIV*. 2020;7(8):e554-e564. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32473657</u>.
- 12. Western Cape Department of Health in collaboration with the National Institute for Communicable Diseases, South Africa. Risk factors for coronavirus disease 2019 (COVID-19) death in a population cohort study from the Western Cape Province, South Africa. *Clin Infect Dis.* 2021;73(7):e2005-e2015. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32860699.
- 13. Bhaskaran K, Rentsch CT, MacKenna B, et al. HIV infection and COVID-19 death: a population-based cohort analysis of UK primary care data and linked national death registrations within the OpenSAFELY platform. *Lancet HIV.* 2021;8(1):e24-e32. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33316211</u>.

- 14. Geretti AM, Stockdale AJ, Kelly SH, et al. Outcomes of coronavirus disease 2019 (COVID-19) related hospitalization among people with human immunodeficiency virus (HIV) in the ISARIC World Health Organization (WHO) clinical characterization protocol (UK): a prospective observational study. *Clin Infect Dis*. 2021;73(7):e2095-e2106. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33095853</u>.
- 15. Dandachi D, Geiger G, Montgomery MW, et al. Characteristics, comorbidities, and outcomes in a multicenter registry of patients with human immunodeficiency virus and coronavirus disease 2019. *Clin Infect Dis*. 2021;73(7):e1964-e1972. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32905581</u>.
- Hoffmann C, Casado JL, Härter G, et al. Immune deficiency is a risk factor for severe COVID-19 in people living with HIV. *HIV Med*. 2021;22(5):372-378. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33368966</u>.
- 17. Tesoriero JM, Swain CE, Pierce JL, et al. COVID-19 outcomes among persons living with or without diagnosed HIV infection in New York State. *JAMA Netw Open*. 2021;4(2):e2037069. Available at: https://www.ncbi.nlm.nih.gov/pubmed/33533933.
- Sun J, Patel RC, Zheng Q, et al. COVID-19 disease severity among people with HIV infection or solid organ transplant in the United States: a nationally-representative, multicenter, observational cohort study. *medRxiv*. 2021;Preprint. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34341798</u>.
- Chun HM, Milligan K, Agyemang E, et al. A systematic review of COVID-19 vaccine antibody responses in people with HIV. *Open Forum Infect Dis*. 2022;9(11):ofac579. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36438620/</u>.
- 20. Haidar G, Agha M, Bilderback A, et al. Prospective evaluation of coronavirus disease 2019 (COVID-19) vaccine responses across a broad spectrum of immunocompromising conditions: the COVID-19 vaccination in the immunocompromised study (COVICS). *Clin Infect Dis.* 2022;75(1):e630-e644. Available at: https://pubmed.ncbi.nlm.nih.gov/35179197/.
- 21. Del Amo J, Polo R, Moreno S, et al. Incidence and severity of COVID-19 in HIV-positive persons receiving antiretroviral therapy: a cohort study. *Ann Intern Med.* 2020;173(7):536-541. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32589451</u>.
- 22. Lea AN, Leyden WA, Sofrygin O, et al. Human immunodeficiency virus status, tenofovir exposure, and the risk of poor coronavirus disease 19 outcomes: real-world analysis from 6 United States cohorts before vaccine rollout. *Clin Infect Dis.* 2023;76(10):1727-1734. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36861341/</u>.
- 23. Rombini MF, Cecchini D, Menendez SD, et al. Tenofovir-containing antiretroviral therapy and clinical outcomes of SARS-CoV-2 infection in people living with HIV. *Viruses*. 2023;15(5):1127. Available at: https://pubmed.ncbi.nlm.nih.gov/37243213/.
- 24. Baden LR, El Sahly HM, Essink B, et al. Efficacy and safety of the mRNA-1273 SARS-CoV-2 vaccine. *N Engl J Med.* 2021;384(5):403-416. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33378609</u>.
- 25. Polack FP, Thomas SJ, Kitchin N, et al. Safety and efficacy of the BNT162b2 mRNA COVID-19 vaccine. *N Engl J Med.* 2020;383(27):2603-2615. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33301246</u>.
- 26. Food and Drug Administration. Fact sheet for healthcare providers administering vaccine (vaccination providers): Emergency Use Authorization (EUA) of the Janssen COVID-19 vaccine to prevent coronavirus disease 2019 (COVID-19). 2022. Available at: https://www.fda.gov/media/146304/download.
- 27. Levy I, Wieder-Finesod A, Litchevsky V, et al. Immunogenicity and safety of the BNT162b2 mRNA COVID-19 vaccine in people living with HIV-1. *Clin Microbiol Infect*. 2021;27(12):1851-1855. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/34438069</u>.
- 28. Woldemeskel BA, Karaba AH, Garliss CC, et al. The BNT162b2 mRNA vaccine elicits robust humoral and cellular immune responses in people living with human immunodeficiency virus (HIV). *Clin Infect Dis*. 2022;74(7):1268-1270. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34293114.
- 29. Frater J, Ewer KJ, Ogbe A, et al. Safety and immunogenicity of the ChAdOx1 nCoV-19 (AZD1222) vaccine against SARS-CoV-2 in HIV infection: a single-arm substudy of a phase 2/3 clinical trial. *Lancet HIV*.

2021;8(8):e474-e485. Available at: https://www.ncbi.nlm.nih.gov/pubmed/34153264.

- 30. Zhou Q, Liu Y, Zeng F, et al. Correlation between CD4 T-cell counts and seroconversion among COVID-19 vaccinated patients with HIV: a meta-analysis. *Vaccines (Basel)*. 2023;11(4):789. Available at: https://pubmed.ncbi.nlm.nih.gov/37112701/.
- 31. Tan SS, Chew KL, Saw S, Jureen R, Sethi S. Cross-reactivity of SARS-CoV-2 with HIV chemiluminescent assay leading to false-positive results. *J Clin Pathol*. 2021;74(9):614. Available at: https://www.ncbi.nlm.nih.gov/pubmed/32907911.
- 32. Blanco JL, Ambrosioni J, Garcia F, et al. COVID-19 in patients with HIV: clinical case series. *Lancet HIV*. 2020;7(5):e314-e316. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32304642</u>.
- 33. Coleman H, Snell LB, Simons R, Douthwaite ST, Lee MJ. Coronavirus disease 2019 and Pneumocystis jirovecii pneumonia: a diagnostic dilemma in HIV. AIDS. 2020;34(8):1258-1260. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32501852</u>.
- 34. Yanes RR, Malijan GMB, Escora-Garcia LK, et al. Detection of SARS-CoV-2 and HHV-8 from a large pericardial effusion in an HIV-positive patient with COVID-19 and clinically diagnosed Kaposi sarcoma: a case report. *Top Med Health*. 2022;50(1):72. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36153612/</u>.
- 35. Basso RP, Poester VR, Benelli JL, et al. COVID-19-associated histoplasmosis in an AIDS patient. *Mycopathologia*. 2021;186(1):109-112. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/33156463/</u>.
- 36. Anggraeni AT, Soedarsono S, Soeprijanto B. Concurrent COVID-19 and Pneumocystis jirovecii pneumonia: the importance of radiological diagnostic and HIV testing. *Radiol Case Rep.* 2021;16(12):3685-3689. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/34630801/</u>.
- 37. Centers for Disease Control and Prevention. Estimated HIV incidence and prevalence in the United States 2014–2018. *HIV Surveillance Rep.* 2020;25(1):1-78. Available at: <u>https://www.cdc.gov/hiv/pdf/library/reports/</u>surveillance/cdc-hiv-surveillance-supplemental-report-vol-25-1.pdf.
- 38. Kong AM, Pozen A, Anastos K, Kelvin EA, Nash D. Non-HIV comorbid conditions and polypharmacy among people living with HIV age 65 or older compared with HIV-negative individuals age 65 or older in the United States: a retrospective claims-based analysis. *AIDS Patient Care STDS*. 2019;33(3):93-103. Available at: https://www.ncbi.nlm.nih.gov/pubmed/30844304.
- 39. Byrd KM, Beckwith CG, Garland JM, et al. SARS-CoV-2 and HIV coinfection: clinical experience from Rhode Island, United States. *J Int AIDS Soc.* 2020;23(7):e25573. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32657527</u>.
- 40. RECOVERY Collaborative Group. Lopinavir-ritonavir in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. *Lancet*. 2020;396(10259):1345-1352. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/33031764</u>.
- 41. Chen J, Xia L, Liu L, et al. Antiviral activity and safety of darunavir/cobicistat for the treatment of COVID-19. *Open Forum Infect Dis.* 2020;7(7):ofaa241. Available at: <u>https://www.ncbi.nlm.nih.gov/pubmed/32671131</u>.
- 42. Smith ER, Oakley E, Grandner GW, et al. Clinical risk factors of adverse outcomes among women with COVID-19 in the pregnancy and postpartum period: a sequential, prospective meta-analysis. *Am J Obstet Gynecol*. 2023;228(2):161-177. Available at: https://pubmed.ncbi.nlm.nih.gov/36027953/.
- 43. Jackson-Gibson M, Diseko M, Caniglia EC, et al. Association of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection with maternal mortality and neonatal birth outcomes in Botswana by human immunodeficiency virus status. *Obstet Gynecol.* 2023;141(1):135-143. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/36701614/</u>.
- 44. Berzosa Sánchez A, Epalza C, Navarro ML, et al. SARS-CoV-2 infection in children and adolescents living with HIV in Madrid. *Pediatr Infect Dis J*. 2022;41(10):824-826. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35796220/</u>.
- 45. Vanetti C, Trabattoni D, Stracuzzi M, et al. Immunocological characterization of HIV and SARS-CoV-2 coinfected young individuals. *Cells*. 2021;10(11):3187. Available at: <u>https://pubmed.ncbi.nlm.nih.</u>

gov/34831410/.

- 46. van der Zalm MM, Lishman J, Verhagen LM, et al. Clinical experience with severe acute respiratory syndrome coronavirus 2-related illness in children: hospital experience in Cape Town, South Africa. *Clin Infect Dis.* 2021;72(12):e938-e944. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/33170927/</u>.
- Nachega JB, Sam-Agudu NA, Machekano RN, et al. Assessment of clinical outcomes among children and adolescents hospitalized with COVID-19 in 6 Sub-Saharan African countries. *JAMA Pediatr*. 2022;176(3):e216436. Available at: <u>https://pubmed.ncbi.nlm.nih.gov/35044430/</u>.

Appendix A, Table 1. COVID-19 Treatment Guidelines Panel Members

Last Updated: October 10, 2023

Co-Chairs Co-Chairs Roy M. Gulick, MD, MPH Weill Cornell Medicine, New York, NY H. Clifford Lane, MD National Institutes of Health, Bethesda, MD Henry Masur, MD National Institutes of Health, Bethesda, MD Executive Secretary Julith Aberg, MD Julith Aberg, MD Icahn School of Medicine at Mount Sinai, New York, NY Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennepin Healthcare/University of Mimesota, Minneapolis, MN Lisa Baumann Kreuziger, ND, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles, Los Angeles, CA Kathleen Chickos, MD, MSCE Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA Craig Coopersmith, MD Ernory University School of Medicine, Atlanta, GA Eric Daar, MD Harbor-UCLA Medical Center, Torrance, CA Army L, Dzierba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschemaer, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschemaer, PharmD New Y	Name	Affiliation	
H. Cifford Lane, MD National Institutes of Health, Bethesda, MD Henry Masur, MD National Institutes of Health, Bethesda, MD Executive Secretary National Institutes of Health, Bethesda, MD Members Judith Aberg, MD Judith Aberg, MD Icahn School of Medicine at Mount Sinai, New York, NY Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennerpin Healthcace/University of Minnesota, Minneapolis, MN Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukee, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSE Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA Craig Coopersmith, MD Emory University School of Medicine, Atlanta, GA Eric Daar, MD Harbor-UCLA Medical Center, Torrance, CA Army L. Dizerba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD Minversity of Minhesota, Minneapolis, MN Laura Evans, MD, MSc University of Minnesota, Mi	Co-Chairs		
H. Cifford Lane, MD National Institutes of Health, Bethesda, MD Henry Masur, MD National Institutes of Health, Bethesda, MD Executive Secretary National Institutes of Health, Bethesda, MD Members Judith Aberg, MD Judith Aberg, MD Icahn School of Medicine at Mount Sinai, New York, NY Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennerpin Healthcace/University of Minnesota, Minneapolis, MN Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukee, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSE Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA Craig Coopersmith, MD Emory University School of Medicine, Atlanta, GA Eric Daar, MD Harbor-UCLA Medical Center, Torrance, CA Army L. Dizerba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD Minversity of Minhesota, Minneapolis, MN Laura Evans, MD, MSc University of Minnesota, Mi	Roy M. Gulick, MD, MPH	Weill Cornell Medicine, New York, NY	
Henry Masur, MD National Institutes of Health, Bethesda, MD Executive Secretary Alice K. Pau, PharmD National Institutes of Health, Bethesda, MD Members Judith Aberg, MD Icahn School of Medicine at Mount Sinai, New York, NY Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennepin Healthcare/University of Minnesota, Minneapolis, NN Lisa Baumann Kreuziger, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSC University of California, Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSC University of California, Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSC Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA Craig Coopersmith, MD Ermory University School of Medicine, Atlanta, GA Freo Dar, MD Harbor-UCLA Medical Center, Torrance, CA Arny L. Dzierba, PharmD New York-Presbyterina Hospital/Narvard Medical School, Boston, MA David V. Glidenen, PhD University of Minesota, Minneapolis, MN			
Executive Secretary National Institutes of Health, Bethesda, MD Members Icahn School of Medicine at Mount Sinai, New York, NY Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennepin Healthcare/University of Minnesota, Minneapolis, MN Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukee, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles, Los Angeles, CA Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kara Chew, MD, MSC Children's Hospital New York, NY Greigory Eschanauer, PharmD New York-Presbytainal New York, NY Greigory Eschanauer, PharmD New York-Presbyterian Hospital, New York, NY			
Alice K. Pau, PharmD National Institutes of Health, Bethesda, MD Members Judith Aberg, MD Icahn School of Medicine at Mount Sinai, New York, NY Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennepin Heatthcare/University of Minnesota, Minneapolis, MN Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukee, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSCE Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA Craig Coopersmith, MD Emory University of Medicine, Atlanta, GA Eric Daar, MD Harbor-UCLA Medical Center, Torrance, CA Army L. Dzierba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD University of Minnesota, Minneapolis, MN <td></td> <td></td>			
Members Judith Aberg, MD Icahn School of Medicine at Mount Sinai, New York, NY Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennepin Healthcare/University of Minnesota, Minneapolis, MN Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukke, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles/University of Southern California, Los Angeles, CA Kathleen Chiotos, MD, MSC University of California, Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSCE Children's Hospital Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSCE Children's Hospital Josh Angeles, Los Angeles, CA Kathleen Chiotos, MD Harbor-UCLA Medical Center, Torrance, CA Army L. Dzierba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD University of Wichigan, Ann Arbor, MI Laura Evans, MD, MSC University of California San Francisco, San Francisco, CA Neil Gidenberg, MD, PhD University of California San Francisco, San Francisco, CA </td <td></td> <td>National Institutes of Health, Bethesda, MD</td>		National Institutes of Health, Bethesda, MD	
Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennepin Healthcare/University of Minnesota, Minneapolis, MN Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukee, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles/University of Southern California, Los Angeles, CA Stephen V. Cantrill, MD Denver Health, Denver, CO Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kathieen Chiotos, MD, MSCE Children's Hospital Los Angeles, Los Angeles, CA Kathieen Chiotos, MD, MSCE Children's Hospital Center, Torrance, CA Army L. Dzierba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD University of Washington, Seattle, WA John J. Gallagher, DNP, RN Case Western Reserve University School of Nursing, Cleveland, OH Rajesh Gandhi, MD Massachusetts General Hospital/Harvard Medical School, Boston, MA David V. Glidden, PhD University of Minnesota, Minneapolis, MN Eric Das, MD, PhD University of Minnesota, Minneapolis, MN			
Adaora Adimora, MD, MPH University of North Carolina School of Medicine, Chapel Hill, NC Jason Baker, MD, MS Hennepin Healthcare/University of Minnesota, Minneapolis, MN Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukee, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles/University of Southern California, Los Angeles, CA Stephen V. Cantrill, MD Denver Health, Denver, CO Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kathieen Chiotos, MD, MSCE Children's Hospital Cos Angeles, Los Angeles, CA Kathieen Chiotos, MD, MSCE Children's Hospital Center, Torrance, CA Army L. Dzierba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD University of Mashington, Seattle, WA John J. Gallagher, DNP, RN Case Western Reserve University School of Nursing, Cleveland, OH Rajesh Gandhi, MD Massachusetts General Hospital/Harvard Medical School, Boston, MA David V. Gildden, PhD University of Minnesota, Minneapolis, MN Eric Das, MD, PhD Johns Hopkins All Children's Institute for Clinical and Tran	Judith Aberg, MD	Icahn School of Medicine at Mount Sinai, New York, NY	
Jason Baker, MD, MS Hennepin Healthcare/University of Minnesota, Minneapolis, MN Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukee, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles/University of Southern California, Los Angeles, CA Stephen V. Cantrill, MD Denver Health, Denver, CO Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSCE Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA Craig Coopersmith, MD Emory University School of Medicine, Atlanta, GA Eric Daar, MD Harbor-UCLA Medical Center, Torrance, CA Army L. Dzierba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD University of Washington, Seattle, WA John J. Gallagher, DNP, RN Case Western Reserve University School of Nursing, Cleveland, OH Rajesh Gandhi, MD Massachusetts General Hospital/Harvard Medical School, Boston, MA David V. Gildden, PhD University of California San Francisco, CA Neil Goldenberg, MD,			
Lisa Baumann Kreuziger, MD, MS Versiti/Medical College of Wisconsin, Milwaukee, WI Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles/University of Southern California, Los Angeles, CA Stephen V. Cantrill, MD Denver Health, Denver, CO Kata Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Children's Hospital Cos Angeles, Los Angeles, CA Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA Craig Coopersmith, MD Emory University School of Medicine, Atlanta, GA Eric Daar, MD Harbor-UCLA Medical Center, Torrance, CA Amy L. Dzierba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD University of Michigan, Ann Arbor, MI Laura Evans, MD, MSC University of California San Francisco, San Francisco, CA John J. Gallagher, DNP, RN Case Western Reserve University School of Nursing, Cleveland, OH Rajesh Gandhi, MD University of California San Francisco, San Francisco, CA Neil Goldenberg, MD, PhD Johns Hopkins All Children's Institute for Clinical and Translational Research, St. Pettersburg, FL Birg			
Roger Bedimo, MD, MS University of Texas Southwestern/Veterans Affairs North Texas Health Care System, Dallas, TX Pamela S. Belperio, PharmD Department of Veterans Affairs, Los Angeles, CA Anoopindar Bhalla, MD Children's Hospital Los Angeles/University of Southern California, Los Angeles, CA Stephen V. Cantrill, MD Denver Health, Denver, CO Kara Chew, MD, MS University of California, Los Angeles, Los Angeles, CA Kathleen Chiotos, MD, MSCE Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA Craig Coopersmith, MD Emory University School of Medicine, Atlanta, GA Fric Daar, MD Harbor-UCLA Medical Center, Torrance, CA Army L. Dzierba, PharmD New York-Presbyterian Hospital, New York, NY Gregory Eschenauer, PharmD University of Michigan, Ann Arbor, MI Laura Evans, MD, MSC University of Washington, Seattle, WA John J. Gallagher, DNP, RN Case Western Reserve University School of Nursing, Cleveland, OH Rajesh Gandhi, MD Massachusetts General Hospital/Harvard Medical School, Boston, MA David V. Glidden, PhD University of Minnesota, Minneapolis, MN Erica J. Hardy, MD, MMSC Warren Alpert Medical School of Brown University, Providence, RI Lauren Henderson, MD, MMSC Duke Univer			
Anoopindar Bhalla, MDChildren's Hospital Los Angeles/University of Southern California, Los Angeles, CAStephen V. Cantrill, MDDenver Health, Denver, COKara Chew, MD, MSUniversity of California, Los Angeles, Los Angeles, CAKathleen Chiotos, MD, MSCEChildren's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PACraig Coopersmith, MDEmory University School of Medicine, Atlanta, GAEric Daar, MDHarbor-UCLA Medical Center, Torrance, CAAmy L. Dzierba, PharmDNew York-Presbyterian Hospital, New York, NYGregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MSCUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMSCWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScDuiversity School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine, Murrora, COMarla J. Keller, MDAlbert Einstein College of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine, Atlanta, GAMitchell M. Levy, MDMassachusetts General Hospital/Harvard Medical School, Boston, M		University of Texas Southwestern/Veterans Affairs North Texas Health Care	
Stephen V. Cantrill, MDDenver Health, Denver, COKara Chew, MD, MSUniversity of California, Los Angeles, Los Angeles, CAKathleen Chiotos, MD, MSCEChildren's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PACraig Coopersmith, MDEmory University School of Medicine, Atlanta, GAEric Daar, MDHarbor-UCLA Medical Center, Torrance, CAAmy L. Dzierba, PharmDNew York-Presbyterian Hospital, New York, NYGregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MScUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta,	Pamela S. Belperio, PharmD	Department of Veterans Affairs, Los Angeles, CA	
Kara Chew, MD, MSUniversity of California, Los Angeles, Los Angeles, CAKathleen Chiotos, MD, MSCEChildren's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PACraig Coopersmith, MDEmory University School of Medicine, Atlanta, GAEric Daar, MDHarbor-UCLA Medical Center, Torrance, CAAmy L. Dzierba, PharmDNew York-Presbyterian Hospital, New York, NYGregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MScUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical Scho	Anoopindar Bhalla, MD	Children's Hospital Los Angeles/University of Southern California, Los Angeles, CA	
Kathleen Chiotos, MD, MSCEChildren's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PACraig Coopersmith, MDEmory University School of Medicine, Atlanta, GAEric Daar, MDHarbor-UCLA Medical Center, Torrance, CAAmy L. Dzierba, PharmDNew York-Presbyterian Hospital, New York, NYGregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MScUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RI Jonthan Li, MD, MMScBrenna Li, MD, MMScBrenra Hospital/Harvard Medical School, Boston, MA	Stephen V. Cantrill, MD	Denver Health, Denver, CO	
Craig Coopersmith, MDEmory University School of Medicine, Atlanta, GAEric Daar, MDHarbor-UCLA Medical Center, Torrance, CAAmy L. Dzierba, PharmDNew York-Presbyterian Hospital, New York, NYGregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MScUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medical School, Boston, MAArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Autora, GOMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, G	Kara Chew, MD, MS	University of California, Los Angeles, Los Angeles, CA	
Eric Daar, MDHarbor-UCLA Medical Center, Torrance, CAAmy L. Dzierba, PharmDNew York-Presbyterian Hospital, New York, NYGregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MScUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RI Lauren Henderson, MD, MMScBrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Brown University, Providence, RIJohnson, MDWarren Alpert Medical School of Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Brown University, Providence, RIJonsthan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Kathleen Chiotos, MD, MSCE	Children's Hospital of Philadelphia/University of Pennsylvania, Philadelphia, PA	
Amy L. Dzierba, PharmDNew York-Presbyterian Hospital, New York, NYGregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MScUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RI Lauren Henderson, MD, MMScBernna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Brown University, Providence, RIJohnsch MDWarren Alpert Medical School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Craig Coopersmith, MD	Emory University School of Medicine, Atlanta, GA	
Amy L. Dzierba, PharmDNew York-Presbyterian Hospital, New York, NYGregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MScUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RI Lauren Henderson, MD, MMScBernna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Brown University, Providence, RIJohnsch MDWarren Alpert Medical School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Eric Daar, MD	Harbor-UCLA Medical Center, Torrance, CA	
Gregory Eschenauer, PharmDUniversity of Michigan, Ann Arbor, MILaura Evans, MD, MScUniversity of Washington, Seattle, WAJohn J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RI Lauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RI Jonathan Li, MD, MMSc		New York-Presbyterian Hospital, New York, NY	
John J. Gallagher, DNP, RNCase Western Reserve University School of Nursing, Cleveland, OHRajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RI Lauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MAProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medical School, Boston, MAArthur Kim, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Gregory Eschenauer, PharmD	University of Michigan, Ann Arbor, MI	
Rajesh Gandhi, MDMassachusetts General Hospital/Harvard Medical School, Boston, MADavid V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RI Lauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medical Center, Bronx, NYArthur Kim, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Laura Evans, MD, MSc	University of Washington, Seattle, WA	
David V. Glidden, PhDUniversity of California San Francisco, San Francisco, CANeil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	John J. Gallagher, DNP, RN	Case Western Reserve University School of Nursing, Cleveland, OH	
Neil Goldenberg, MD, PhDJohns Hopkins All Children's Institute for Clinical and Translational Research, St. Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Rajesh Gandhi, MD	Massachusetts General Hospital/Harvard Medical School, Boston, MA	
Petersburg, FLBirgit Grund, PhDUniversity of Minnesota, Minneapolis, MNErica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	David V. Glidden, PhD	University of California San Francisco, San Francisco, CA	
Erica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Neil Goldenberg, MD, PhD		
Erica J. Hardy, MD, MMScWarren Alpert Medical School of Brown University, Providence, RILauren Henderson, MD, MMScBoston Children's Hospital/Harvard Medical School, Boston, MACarl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Birgit Grund, PhD	University of Minnesota, Minneapolis, MN	
Carl Hinkson, MSRCProvidence Health & Services, Everett, WABrenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Erica J. Hardy, MD, MMSc		
Brenna L. Hughes, MD, MScDuke University School of Medicine, Durham, NCSteven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Lauren Henderson, MD, MMSc	Boston Children's Hospital/Harvard Medical School, Boston, MA	
Steven Johnson, MDUniversity of Colorado School of Medicine, Aurora, COMarla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Carl Hinkson, MSRC	Providence Health & Services, Everett, WA	
Marla J. Keller, MDAlbert Einstein College of Medicine/Montefiore Medical Center, Bronx, NYArthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Brenna L. Hughes, MD, MSc	Duke University School of Medicine, Durham, NC	
Arthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Steven Johnson, MD	University of Colorado School of Medicine, Aurora, CO	
Arthur Kim, MDMassachusetts General Hospital/Harvard Medical School, Boston, MAJeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Marla J. Keller, MD	Albert Einstein College of Medicine/Montefiore Medical Center, Bronx, NY	
Jeffrey L. Lennox, MDEmory University School of Medicine, Atlanta, GAMitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA			
Mitchell M. Levy, MDWarren Alpert Medical School of Brown University, Providence, RIJonathan Li, MD, MMScBrigham and Women's Hospital/Harvard Medical School, Boston, MA	Jeffrey L. Lennox, MD		
Jonathan Li, MD, MMSc Brigham and Women's Hospital/Harvard Medical School, Boston, MA			
	Christine MacBrayne, PharmD, MSCS Children's Hospital Colorado, Aurora, CO		

COVID-19 Treatment Guidelines

Name	Affiliation
Members, continued	
Gregory Martin, MD, MSc	Emory University School of Medicine, Atlanta, GA
Nandita R. Nadig, MD	Northwestern University School of Medicine, Chicago, IL
Andrew T. Pavia, MD	University of Utah School of Medicine, Salt Lake City, UT
Grant Schulert, MD, PhD	Cincinnati Children's Hospital Medical Center/University of Cincinnati College of Medicine, Cincinnati, OH
Nitin Seam, MD	National Institutes of Health, Bethesda, MD
Steven Q. Simpson, MD	University of Kansas Medical Center, Kansas City, KS
Susan Swindells, MBBS	University of Nebraska Medical Center, Omaha, NE
Pablo Tebas, MD	University of Pennsylvania, Philadelphia, PA
Phyllis Tien, MD, MSc	University of California, San Francisco/San Francisco VA Healthcare System, San Francisco, CA
Alpana A. Waghmare, MD	Seattle Children's Hospital, Seattle, WA
Cameron R. Wolfe, MBBS	Duke University School of Medicine, Durham, NC
Jinoos Yazdany, MD, MPH	University of California, San Francisco, San Francisco, CA
Community Members	
Danielle M. Campbell, MPH	University of California, Los Angeles, Los Angeles, CA
Carly Harrison	LupusChat, New York, NY
Richard Knight, MBA	American Association of Kidney Patients, Bowie, MD
Pharmacology Consultants	
Sarita Boyd, PharmD	Food and Drug Administration, Silver Spring, MD
Jomy George, PharmD	National Institutes of Health, Bethesda, MD
Kimberly Scarsi, PharmD	University of Nebraska Medical Center, Omaha, NE
Ex Officio Members, U.S. Government Repre	
Timothy Burgess, MD	Department of Defense, Bethesda, MD
Derek Eisnor, MD	Biomedical Advanced Research and Development Authority, Washington, DC
Joseph Francis, MD, MPH	Department of Veterans Affairs, Washington, DC
Pragna Patel, MD, MPH, DTM&H	Centers for Disease Control and Prevention, Atlanta, GA
Virginia Sheikh, MD, MHS	Food and Drug Administration, Silver Spring, MD
Timothy M. Uyeki, MD, MPH	Centers for Disease Control and Prevention, Atlanta, GA
U.S. Government Support Team	
Richard T. Davey, Jr., MD	National Institutes of Health, Bethesda, MD
Alison Han, MD (Co-Team Coordinator)	National Institutes of Health, Bethesda, MD
Elizabeth S. Higgs, MD, DTM&H, MIA	National Institutes of Health, Bethesda, MD
Martha C. Nason, PhD (Biostatistics Support)	National Institutes of Health, Bethesda, MD
Michael Proschan, PhD (Biostatistics Support)	National Institutes of Health, Bethesda, MD
Renee Ridzon, MD	National Institutes of Health, Bethesda, MD
Kanal Singh, MD, MPH (Co-Team Coordinator)	National Institutes of Health, Bethesda, MD
Assistant Executive Secretaries	
Safia Kuriakose, PharmD	Frederick National Laboratory for Cancer Research, in support of NIAID, Frederick, MD
Andrea M. Lerner, MD, MS	National Institutes of Health, Bethesda, MD

Appendix A, Table 2. COVID-19 Treatment Guidelines Panel Financial Disclosure for Companies Related to COVID-19 Treatment or Diagnostics

Last Updated: October 10, 2023

Reporting Period: April 1, 2022, to March 31, 2023

B	Financial Disclosure	
Panel Member	Company	Relationship
Judith Aberg, MD	Emergent	Research support
	Gilead	Research support
	GSK/ViiV Healthcare	Advisory board, research support
	Janssen	Research support
	Kintor	DSMB chair/member
	Merck	Advisory board, research support
	Pfizer	Research support
	Regeneron	Research support
Adaora Adimora, MD, MPH	Gilead	Consultant, honoraria
	Merck	Advisory board, consultant, honoraria, research support
Jason Baker, MD, MS	None	N/A
Lisa Baumann Kreuziger, MD, MS	3M	Stockholder, spouse is an employee
	Versiti	Employee
Roger Bedimo, MD, MS	Gilead	Advisory board
	GSK/ViiV Healthcare	Advisory board
	Janssen	Advisory board
	Merck	Advisory board, research support
	Shionogi	Advisory board
	Theratechnologies	Advisory board
Pamela S. Belperio, PharmD	None	N/A
Anoopindar Bhalla, MD	None	N/A
Timothy Burgess, MD	AstraZeneca	Research support
Danielle M. Campbell, MPH	Gilead	Advisory board
	GSK/ViiV Healthcare	Attendee at a community stakeholder meeting
Stephen V. Cantrill, MD	None	N/A
Kara Chew, MD, MS	Pardes Biosciences	Consultant
	Merck	Research support
Kathleen Chiotos, MD, MSCE	None	N/A
Craig Coopersmith, MD	None	N/A
Eric Daar, MD	Gilead	Consultant, research support
	GSK/ViiV Healthcare	Research support
	Merck	Consultant
	Theratechnologies	Consultant

COVID-19 Treatment Guidelines

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

Devel Merchan	Financial Disclosure	
Panel Member	Company	Relationship
Richard T. Davey, Jr., MD	None	N/A
Amy L. Dzierba, PharmD	None	N/A
Derek Eisnor, MD	None	N/A
Gregory Eschenauer, PharmD	None	N/A
Laura Evans, MD, MSc	None	N/A
Joseph Francis, MD, MPH	None	N/A
John J. Gallagher, DNP, RN	None	N/A
Rajesh Gandhi, MD	None	N/A
David V. Glidden, PhD	Gilead	Consultant
Neil Goldenberg, MD, PhD	Anthos Therapeutics	Advisory board, pediatric clinical trial design
	Bayer	Advisory board, pediatric clinical trial design
	Boehringer-Ingelheim	Advisory board, pediatric clinical trial design
	CPC Clinical Research	DSMB chair/member
	Daiichi Sankyo	Steering committee chair, pediatric clinical trial design and oversight
	Novartis	DSMB chair/member
Birgit Grund, PhD	None	N/A
Roy M. Gulick, MD, MPH	None	N/A
Alison Han, MD	None	N/A
Erica J. Hardy, MD, MMSc	None	N/A
Carly Harrison	Aurinia	Advisory board, consultant
	GSK	Advisory board, consultant
Lauren Henderson, MD, MMSc	Adaptive Biotechnologies	Research support
	Bristol Myers Squibb	Research support
	Pfizer	External panel for grant reviews
	Sobi	Consultant, research support
	SkyGenic	Consultant, spouse is an employee
Elizabeth S. Higgs, MD, DTM&H, MIA	None	N/A
Carl Hinkson, MSRC	None	N/A
Brenna L. Hughes, MD, MSc	None	N/A
Steven Johnson, MD	None	N/A
Marla J. Keller, MD	None	N/A
Arthur Kim, MD	Kintor Pharmaceuticals	DSMB chair/member
Richard Knight, MBA	None	N/A
Safia Kuriakose, PharmD	None	N/A
H. Clifford Lane, MD	None	N/A
Jeffrey L. Lennox, MD	GSK/ViiV Healthcare	Research support
	Pfizer	Spouse is an employee of Pfizer contract
Andrea M. Lerner, MD, MS	None	N/A
Mitchell M. Levy, MD	Endpoint	Advisory board
	Inotrem	Advisory board

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024

David Mansher	Financial Disclosure	
Panel Member	Company	Relationship
Jonathan Li, MD, MMSc	Merck	Research support
Christine MacBrayne, PharmD, MSCS	None	N/A
Gregory Martin, MD, MSc	Grifols	Research grants review panel
Henry Masur, MD	None	N/A
Nandita R. Nadig, MD	None	N/A
Martha C. Nason, PhD	Bristol Myers Squibb	Stockholder
	Medtronic	Stockholder
Pragna Patel, MD, MPH, DTM&H	None	N/A
Alice K. Pau, PharmD	None	N/A
Andrew T. Pavia, MD	GSK	Consultant
	Haleon	Honoraria
Michael Proschan, PhD	None	N/A
Renee Ridzon, MD	None	N/A
Grant Schulert, MD, PhD	Sobi	Consultant
Nitin Seam, MD	None	N/A
Virginia Sheikh, MD, MHS	None	N/A
Steven Q. Simpson, MD	None	N/A
Kanal Singh, MD, MPH	None	N/A
Susan Swindells, MBBS	GSK/ViiV Healthcare	Research support
Pablo Tebas, MD	None	N/A
Phyllis Tien, MD, MSc	Merck	Research support
Timothy M. Uyeki, MD, MPH	None	N/A
Alpana A. Waghmare, MD	AlloVir	Research support
	Ansun BioPharma	Research support
	Pfizer	Research support
	Vir Biotechnology	Research support
Cameron R. Wolfe, MBBS	Adamis Pharmaceuticals Corporation	DSMB chair
	Atea Pharmaceuticals	DSMB chair/member
	Biogen	DSMB chair
Jinoos Yazdany, MD, MPH	AstraZeneca	Consultant, research support
	Aurinia	Research support
	Bristol Myers Squibb	Research support
	Gilead	Research support

Key: DSMB = data and safety monitoring board

Downloaded from https://www.covid19treatmentguidelines.nih.gov/ on 2/5/2024