RAPID RECOMMENDATIONS

A living WHO guideline on drugs for covid-19

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ABSTRACT

UPDATES

This is the twelfth version (eleventh update) of the living guideline, replacing earlier versions (available as data supplements). New recommendations will be published as updates to this guideline.

CLINICAL QUESTION

What is the role of drugs in the treatment of patients with covid-19?

CONTEXT

The evidence base for therapeutics for covid-19 is evolving with numerous randomised controlled trials (RCTs) recently completed and under way. The emerging SARS-CoV-2 variants (such as omicron) and subvariants are also changing the role of therapeutics. This update provides updated recommendations for remdesivir, addresses the use of combination therapy with corticosteroids, in patients with severe or critical covid-19, and modifies previous recommendations for the neutralising monoclonal antibodies sotrovimab and casirivimab-imdevimab in patients with non-severe covid-19.

NEW OR UPDATED RECOMMENDATIONS

• Remdesivir: a conditional recommendation for its use in patients with severe covid-19; and a conditional recommendation against its use in patients with critical covid-19.

• Concomitant use of IL-6 receptor blockers (tocilizumab or sarilumab) and the JAK inhibitor baricitinib: these drugs may now be combined, in addition to corticosteroids, in patients with severe or critical covid-19.

• Sotrovimab and casirivimab-imdevimab: strong recommendations against their use in patients with covid-19, replacing the previous conditional recommendations for their use.

UNDERSTANDING THE NEW RECOMMENDATIONS

When moving from new evidence to updated recommendations, the Guideline Development Group (GDG) considered a combination of evidence assessing relative benefits and harms, values and preferences, and feasibility issues.

For remdesivir, new trial data were added to a previous subgroup analysis and provided sufficiently trustworthy evidence to demonstrate benefits in patients with severe covid-19, but not critical covid-19. The GDG considered benefits of remdesivir to be modest and of moderate certainty for key outcomes such as mortality and mechanical ventilation, resulting in a conditional recommendation.

For baricitinib, the GDG considered clinical trial evidence (RECOVERY) demonstrating reduced risk of death in patients already receiving corticosteroids and IL-6 receptor blockers. The GDG acknowledged that the clinical trials were not representative of the world population and that the risk-benefit balance may be less advantageous, particularly in patients who are immunosuppressed at higher risk of opportunistic infections (such as serious fungal, viral, or bacteria), those already deteriorating where less aggressive or stepwise addition of immunosuppressive medications may be preferred, and in areas where certain pathogens such as HIV or tuberculosis, are of concern. The panel anticipated that there would be situations where clinicians may opt for less aggressive immunosuppressive therapy or to combine medications in a stepwise fashion in patients who are deteriorating. The decision to combine the medications will depend on their availability, and the treating clinician’s perception of the risk-benefit balance associated with combination immunosuppressive therapy, particularly in patient populations at risk of opportunistic
infections who may have been under-represented in clinical trials. When making a strong recommendation against the use of monoclonal antibodies for patients with covid-19, the GDG considered in vitro neutralisation data demonstrating that sotrovimab and casirivimab-imdevimab evaluated in clinical trials have meaningfully reduced neutralisation activity of the currently circulating variants of SARS-CoV-2 and their subvariants. There was consensus among the panel that the absence of in vitro neutralisation activity strongly suggests absence of clinical effectiveness of these monoclonal antibodies. However, there was also consensus regarding the need for clinical trial evidence in order to confirm clinical efficacy of new monoclonal antibodies that reliably neutralise the circulating strains in vitro. Whether emerging new variants and subvariants might be susceptible to sotrovimab, casirivimab-imdevimab, or other anti-SARS-CoV-2 monoclonal antibodies cannot be predicted.

PRIOR RECOMMENDATIONS

- Recommended for patients with severe or critical covid-19—strong recommendations for systemic corticosteroids; IL-6 receptor blockers (tocilizumab or sarilumab) in combination with corticosteroids; and baricitinib as an alternative to IL-6 receptor blockers, in combination with corticosteroids.
- Recommended for patients with non-severe covid-19 at highest risk of hospitalisation—a strong recommendation for nirmatrelvir/ritonavir; conditional recommendations for molnupiravir and remdesivir.
- Not recommended for patients with non-severe covid-19—a conditional recommendation against systemic corticosteroids; a strong recommendation against convalescent plasma; a recommendation against fluvoxamine, except in the context of a clinical trial; and a strong recommendation against colchicine.
- Not recommended for patients with non-severe covid-19 at low risk of hospitalisation—a conditional recommendation against nirmatrelvir/ritonavir.
- Not recommended for patients with severe or critical covid-19—a recommendation against convalescent plasma except in the context of a clinical trial; and a conditional recommendation against the JAK inhibitors ruxolitinib and tofacitinib.
- Not recommended, regardless of covid-19 disease severity—a strong recommendations against hydroxychloroquine and against lopinavir/ritonavir; and a recommendation against ivermectin except in the context of a clinical trial.

ABOUT THIS GUIDELINE

This living guideline from the World Health Organization (WHO) incorporates new evidence to dynamically update recommendations for covid-19 therapeutics. The GDG typically evaluates a therapy when the WHO judges sufficient evidence is available to make a recommendation. While the GDG takes an individual patient perspective in making recommendations, it also considers resource implications, acceptability, feasibility, equity, and human rights. This guideline was developed according to standards and methods for trustworthy guidelines, making use of an innovative process to achieve efficiency in dynamic updating of recommendations. The methods are aligned with the WHO Handbook for Guideline Development and according to a pre-approved protocol (planning proposal) by the Guideline Review Committee (GRC). A box at the end of the article outlines key methodological aspects of the guideline process. MAGIC Evidence Ecosystem Foundation provides methodological support, including the coordination of living systematic reviews with network meta-analyses to inform the recommendations. The full version of the guideline is available online in MAGICapp and in PDF, with a summary version here in The BMJ. These formats should facilitate adaptation, which is strongly encouraged by WHO to contextualise recommendations in a healthcare system to maximise impact.

FUTURE RECOMMENDATIONS

Recommendations on anticoagulation are planned for the next update to this guideline.

Vaccines are linked to limiting hospitalisations, but limitations in global access and residual uncertainties remain regarding the duration of protection following vaccination or natural infection, and their efficacy against emerging SARS-CoV-2 variants and subvariants. Therefore, the potential for drugs to treat people infected with covid-19 remains of interest. This living guideline responds to emerging evidence from randomised controlled trials (RCTs) on existing and new drug treatments for covid-19.

More than 5000 trials investigating covid-19 interventions have been registered or are ongoing (see section on emerging evidence). Although most of these studies are small and of variable methodological quality, some large, international platform trials have provided robust evidence. Such trials can also adapt their design, recruitment strategies, and selection of interventions based on new insights. Examples include ACCT, RECOVERY, WHO SOLIDARITY, REMAP-CAP, and ACTIV, which recruit large numbers of patients in many countries. An overview of ongoing trials is available from the Infectious Diseases Data Observatory, through their living systematic review of covid-19 clinical trial registrations and the World Health Organization (WHO) website (https://www.covid-nma.com/dataviz/).

However, existing and emerging evidence demonstrates remaining uncertainties concerning treatment effects for all outcomes of importance to patients. There is also a need for better evidence on values and preferences of patients with covid-19. Moreover, the rapidly evolving evidence landscape requires trustworthy interpretation and expeditious clinical practice guidelines to inform clinicians and health care decision-makers.

Several living network meta-analyses associated with this guideline incorporate emerging trial data and allow for analysis of comparative effectiveness of multiple covid-19 treatments. 

Box 1: Linked resources in this BMJ Rapid Recommendations cluster

**Versions of this guideline**
- MAGICapp (https://app.magicapp.org/#/guideline/nBko1z)
  - Expanded version of the guideline, including methods, processes, and results with multi-layered recommendations, evidence summaries, and decision aids for use on all devices
  - MATCH-IT interactive decision support incorporating multiple treatment comparisons for recommended drugs in non-severe covid-19 at highest risk of hospitalisation: https://magicevidence.org/match-it/220404dist-covid-meds/

**Linked research**

- The BMJ 2020;370:m3379 | doi: 10.1136/bmj.m3379
- BMJ: first published as 10.1136/bmj.m3379 on 4 September 2020. Downloaded from http://www.bmj.com on 4 December 2022 by guest. Protected by copyright.
What triggered this version of the guideline and what is coming next?

This twelfth version of the WHO living guideline was triggered by:

- The publication of a large RCT (RECOVERY), confirming the benefits of the JAK inhibitor baricitinib in patients with severe or critical covid-19, also in patients receiving corticosteroids and IL-6 receptor blockers
- The availability of data from in vitro SARS-CoV-2 neutralisation assays, suggesting that monoclonal antibodies (sotrovimab and casirivimab-idecemab) are not effective for patients with non-severe covid-19 in the current stage of the pandemic with the predominant omicron variants
- New RCTs on the effects of remdesivir in patients with severe and critical covid-19.
- The WHO has a standing steering committee to evaluate possibilities for new drug recommendations and updates to existing drug recommendations. The WHO considers multiple factors, including the extent of available evidence, and whether and when additional evidence might be anticipated, to make decisions. Other therapeutics in progress for this WHO living guideline include therapeutic anticoagulation.

How to use this guideline and associated resources

This is a living guideline. The recommendations and evidence included here will be updated, and new recommendations will be added for other treatments for covid-19. The infographic provides a summary of the recommendations. Readers can find more detailed information in the full version of the WHO guideline (see box 1 for links to MAGICapp and the PDF version).

Who do the recommendations apply to?

This guideline applies to all patients with covid-19. Recommendations may differ based on the severity of covid-19, according to WHO severity definitions (box 2). These definitions avoid reliance on access to healthcare to define patient subgroups.

**Box 2: WHO definitions of illness severity for covid-19**

- **Critical covid-19**—Defined by the criteria for acute respiratory distress syndrome (ARDS), sepsis, septic shock, or other conditions that would normally require the provision of life-sustaining therapies such as mechanical ventilation (invasive or non-invasive) or vasopressor therapy.
- **Severe covid-19**—Defined by any of:
  - Oxygen saturation 90% on room air*
  - Signs of pneumonia
  - Signs of severe respiratory distress (in adults, accessory muscle use, inability to complete full sentences, respiratory rate ≥ 30 breaths per minute; and, in children, very severe chest wall indrawing, grunting, central cyanosis, or presence of any other general danger signs including inability to breastfeed or drink, lethargy, convulsions, or reduced level of consciousness).
- **Non-severe covid-19**—Defined as the absence of any criteria for severe or critical covid-19.

*The Guideline Development Group (GDG) noted that the oxygen saturation threshold of 90% to define severe covid-19 was arbitrary and should be interpreted cautiously when defining illness severity. For example, clinicians must use their judgment to determine whether a low oxygen saturation is a sign of severity or is normal for a given patient with chronic lung disease. Similarly, clinicians may interpret a saturation of 90-94% on room air as abnormal in a patient with normal lungs, or as an early sign of severe disease in a patient with a downward clinical trajectory. Generally, in cases where there is any doubt, the GDG suggested erring on the side of considering disease as severe.
How to use the recommendations
Selecting therapeutic agents

Over two years into the pandemic, there are several effective treatments to choose between. When moving from evidence to recommendations for these drugs, the Guideline Development Group (GDG) considered a combination of the evidence regarding relative benefits and harms, values and preferences, practical issues, resource considerations, and feasibility and equity considerations (box 3). The GDG notes that these issues have to be considered when choosing between therapeutic agents, and when re-using or adapting the recommendations in national or local contexts.
For drugs recommended in non-severe illness: the lack of
therapeutics:

WHO aims to provide a stimulus to engage all possible mechanisms to
improve global access to diagnostic tests and effective interventions and
how countries can address such challenges; such as the integration of
a covid-19 clinical pathway and establishing services to offer
intravenous treatments.

At a time of drug shortage, it may be necessary to prioritise use through
clinical triage such as selecting patients with the highest baseline risk
for mortality (for instance, those with critical illness over those with severe
illness), in whom the absolute benefit of treatment is greatest. Other
suggestions for prioritisation, which lack direct evidence, include focusing
on patients with an actively deteriorating clinical course and avoiding
treatment in patients with established multi-organ failure (in whom the
benefit is likely to be small).

Some therapies can be used in combination (such as those for severe
or critical covid-19) while others are to be used as alternatives.
Recommended combinations of treatments are based on direct
comparisons from trials demonstrating additional benefit, such as
adding the JAK inhibitor baricitinib to IL-6 receptor blockers and
to systemic corticosteroids in patients with severe or critical covid-19.

An interactive decision support tool incorporates multiple treatment
comparisons to inform the use of one drug over another with related
mechanisms of action for patients with non-severe covid-19 at
highest risk of hospitalisation: https://magicevidence.org/match-
lt/220404dist-covid-meds/.

Identifying patients with non-severe illness at highest risk of
hospitalisation

Several recommendations for therapeutics in patients with
non-severe covid-19 apply only to those at highest risk of
hospitalisation, defined as beyond 10% risk of being hospitalised
for covid-19. These patients should achieve what the GDG agreed
would represent what most patients would value as an important
benefit: a 6% absolute reduction in hospital admission (see box on
how this guideline was created). Reliably identifying those at highest
risk is challenging because of the changing global context, with
evolution of the virus and patterns of vaccination, thus raising the
importance of validation of models to local contexts. A living
systematic review of 232 risk prediction models for covid-19, recently
updated, identified two promising risk prediction tools before
circulation of the omicron variant: these tools concur that typical
characteristics of people at highest risk include older age,
immunosuppression, and/or presence of chronic diseases, with
lack of vaccination as an additional risk factor.

Uncertainties

Uncertainties exist regarding covid-19 therapeutics and emerging
evidence which may both inform clinical decision-making as well as
future research. The recommendations therefore need to be used
in light of these uncertainties. Specific uncertainties are listed with
the relevant drug, but many uncertainties are common across
therapeutics:

- For drugs recommended in non-severe illness: the lack of
  accurate clinical prediction guides to establish the individual
  patient risk of hospitalisation in order to best identify patients
  that would most benefit from interventions; data regarding
  emergence of resistance and efficacy against new variants; safety
  and efficacy in children and in immunocompromised, vaccinated,
or pregnant patients and other specific subgroups of patients;
  optimal duration of therapies; head-to-head comparisons of
  recommended treatments; and relative effectiveness of
  combination therapy.

- For drugs recommended in severe or critical illness: safety and
  efficacy in children and in immunocompromised, vaccinated,
or pregnant patients and other specific subgroups of patients;
  long term mortality and functional outcomes in covid-19
  survivors; and immunity and the risk of a subsequent infection,
  which may affect the risk of death after 28 days.

The recommendations

Remdesivir (Update 11, published 16 September 2022)

Overview

Remdesivir was developed for treatment of hepatitis C virus
infection, and was also studied in Ebola and Marburg virus
infections before being repurposed for SARS-CoV-2. Remdesivir is a
nucleoside analogue which interacts with the SARS-CoV-2
polymerase to elicit delayed chain termination during RNA genome
synthesis; its mechanism of action is further described in the full
version of the guideline (box 1). Resistance to remdesivir has been
reported in vitro and in a case report, but the clinical significance of
these observations remain unclear.

Update—An initial conditional (weak) recommendation was made
on 20 November 2020, suggesting not to use remdesivir for patients
with covid-19 regardless of illness severity. This was based on data
from the four RCTs available at the time, with 7333 participants
hospitalised for covid-19. In the tenth iteration of the guideline, a
new recommendation was made for the use of remdesivir for patients
with non-severe illness. In this twelfth iteration of the guideline, a
new recommendation for patients with severe or critical covid-19
are provided, given new trial data providing sufficiently trustworthy
evidence for a subgroup effect demonstrating modest benefit in
patients with severe, but not critical, covid-19.

Evidence—The clinical evidence underpinning the recommendations
(focused on the benefits and short-term harms from trial data) is
outlined in box 4. Evidence in severe and critical illness are
presented separately, based on new data providing sufficiently
trustworthy evidence for a subgroup effect demonstrating modest benefit
in patients with severe, but not critical covid-19.

Box 4: Remdesivir data for severe or critical covid-19 (https://app.mag-
icapp.org/#/guideline/nBkO1E/rec/noRN6w and https://app.mag-
icapp.org/#/guideline/nBkO1E/rec/nB6Gy)

The living network meta-analysis for remdesivir was informed by five
trials, which enrolled 7643 patients with severe or critical covid-19. All
trials were published in peer-reviewed journals, and none included
children or pregnant women. See more trial details in appendix 12 on
bmj.com.

For patients with severe covid-19, remdesivir possibly reduces mortality
(odds ratio (OR) 0.89 (95% confidence interval (CI) 0.78 to 1.02); absolute
difference 13 fewer deaths per 1000 patients (95% CI 26 fewer to 2 more);
low certainty), probably reduces the need for mechanical ventilation (OR
0.87 (0.77 to 0.99); absolute difference 14 fewer per 1000 patients (24
fewer to 1 fewer); moderate certainty), and probably has little or no impact
on time to symptom improvement (absolute difference 0.7 fewer days
(1.8 fewer to 0.6 more); moderate certainty).

For patients with critical covid-19, remdesivir possibly has little or no
effect on mortality (OR 1.15 (0.89 to 1.51); absolute difference 34 more
fewer to 17 more); low certainty).
Recommendation 1: For patients with severe covid-19, we suggest treatment with remdesivir (weak or conditional recommendation).

Understanding the recommendation
When moving from evidence to the conditional recommendation to use remdesivir in patients with severe covid-19, the GDG emphasised the benefits on survival and reduction in need for invasive mechanical ventilation, and the likelihood of little or no serious adverse events attributable to the drug. The GDG acknowledged that some serious adverse events may not have been accurately captured during the relatively short follow-up period in the included trials. Of note, although the GDG has recommended for other antiviral drugs in patients with non-severe illness, remdesivir is the only one with a recommendation for use in patients with severe covid-19.

The GDG did not anticipate important variability in patient values and preferences, although the low certainty of evidence and ongoing uncertainty in effect contributed to the conditional recommendation. There was insufficient trial level data to examine subgroups based on age or to consider patients requiring non-invasive ventilation (those on bilevel ventilation or high flow nasal cannula) as a separate subgroup of interest.

When making the recommendation for treatment with remdesivir, the GDG carefully considered the credibility of subgroup findings based on severity of disease, where remdesivir demonstrated a possible survival benefit in patients with severe covid-19, while possibly having no impact on mortality in patients with critical covid-19. The GDG used the ICEMAN tool to assess the credibility of subgroup effects, and ultimately decided the credibility of the observed subgroup finding based on severity of illness was moderate, therefore warranting separate recommendations for each, while recognising residual uncertainties (see MAGICapp for full details regarding ICEMAN assessments).

Balance of benefits and harms—There was low certainty evidence suggesting remdesivir possibly reduces mortality, and moderate certainty evidence suggesting probable reduction in need for mechanical ventilation with probably little or no impact on time to symptom improvement. The drug is well tolerated, and adverse events are rare.

Values and preferences—The GDG inferred that the majority of well informed patients with severe covid-19 would choose to receive remdesivir due to the possible reduction in mortality and need for mechanical ventilation and the safety of the drug. The GDG anticipated little variation in values and preferences between patients for this intervention.

Applicability—Insufficient evidence exists to inform a recommendation around use in children. Decisions regarding its use in pregnant or breastfeeding women should, in the absence of trials enrolling such participants, be made between the pregnant person and their healthcare provider while discussing whether the potential benefit justifies the potential risk to the mother and fetus. See MAGICapp for additional guidance.

Practical issues—Remdesivir is administered as one intravenous infusion daily over 10 consecutive days. The recommended dose is 200 mg intravenously on day 1, followed by 100 mg intravenously on days 2 to 10. Shorter regimens of five days are described in the smaller trials, and local practices may vary. Administration should be as early as possible in the time course of the disease. Patients with severe liver or kidney disease warrant additional caution. See MAGICapp for additional guidance.

Resource implications, acceptability, feasibility, equity, and human rights—Given the intravenous administration of remdesivir daily over 10 days, this is more easily done for hospitalised patients with severe disease, as opposed to the outpatient setting. Obstacles to access in low and middle income countries due to cost, feasibility, and availability are of concern (see box 3 for more details).

Recommendation 2: For patients with critical covid-19, we suggest not to use remdesivir (weak or conditional recommendation).

Understanding the recommendation
When moving from evidence to the conditional recommendation not to use remdesivir in patients with critical covid-19, the GDG emphasised the lack of benefit on survival or other patient-important outcomes as demonstrated in the subgroup analysis judged to be of moderate credibility. The GDG recognised there is ongoing uncertainty, and there may still be a subset of patients who would benefit (for example, immunocompromised, persistent viraemia), but there is insufficient evidence to make recommendations specific to these subsets of critical patients.

The GDG did not anticipate important variability in patient values and preferences, although the low certainty of evidence and ongoing uncertainty in effect contributed to the conditional recommendation. There was insufficient trial level data to examine subgroups based on age, or to consider patients requiring non-invasive ventilation (those on bilevel ventilation or high flow nasal cannula) as a separate subgroup of interest.

Balance of benefits and harms—Low certainty evidence suggests remdesivir possibly has little or no effect on mortality and need for mechanical ventilation, and an uncertain effect on time to symptom improvement. The drug is well tolerated, and adverse events are rare.

Values and preferences—The GDG inferred that the majority of well informed patients with critical covid-19 would choose not to receive remdesivir due to little or no impact on patient-important outcomes. The GDG anticipated little variation in values and preferences between patients for this intervention.

Applicability, practical issues, resource implications, acceptability, feasibility, equity, and human rights—Similar issues exist as for patients with severe illness. Such considerations are less relevant for patients with critical illness, given the weak or conditional recommendation against use.

Recommendation 3: We suggest treatment with remdesivir for patients with non-severe covid-19, conditional to those at highest risk of hospitalisation (conditional or weak recommendation).

Understanding the recommendation
The GDG emphasised the benefits of decreased need for hospitalisation, along with little or no serious adverse effects.
attributable to the drug. Feasibility, costs, access, and complexity of administration were also carefully considered and led to the conditional recommendation for use only in patients at highest risk of hospitalisation.

**Balance of benefits and harms**—In highest risk patients with non-severe illness, moderate certainty evidence showed that remdesivir probably provides an important reduction in hospital admissions and may have little or no effect on mortality. The impact of remdesivir on mechanical ventilation and time to symptom resolution is very uncertain. Treatment probably does not increase the likelihood of serious adverse effects leading to drug discontinuation.

**Values and preferences**—The GDG inferred that almost all well informed patients with a low risk of hospitalisation would decline remdesivir, and only those at highest risk who would choose to receive treatment.

**Applicability**—Only one included trial enrolled children (aged ≥12 years) with small numbers included; the applicability of this recommendation to children therefore remains uncertain. In the absence of trial data for children aged <12 years with weight <40 kg, the use of remdesivir in these children is not recommended. Uncertainty also remains with regard to administration of remdesivir to pregnant or lactating women. The decision regarding use should be made between the pregnant individual and their healthcare provider while discussing whether the potential benefit justifies the potential risk to the mother and fetus.

The GDG also had concerns regarding whether the drug would retain efficacy against emerging variants of concern, such as omicron BA-1 or BA-2. Surveillance is needed for SARS-CoV-2 strains with reduced susceptibility to remdesivir, and further research is needed to examine the role of combination therapy in severely immunocompromised patients. In the absence of further data, the GDG did not have reason to believe the activity against known variants would be diminished.

**Practical issues**—Remdesivir is administered via intravenous infusion as a three-day regimen, in keeping with the large trials informing the recommendation; 200 mg is administered intravenously on day 1, followed by 100 mg given intravenously on days 2 and 3. Administration should be as early as possible in the course of the disease, with monitoring for allergic, infusion related, or other adverse outcomes. In the included studies, remdesivir was administered within seven days of disease onset. Additional considerations regarding practical issues are summarised in MAGICapp.

**Resource implications, acceptability, feasibility, equity, and human rights**—The infusion schedule represents a feasibility challenge in the outpatient settings. Furthermore, remdesivir is unlikely to be available for all individuals who, given the option, would choose to receive the treatment. See box 3 for concerns about costs, availability, and health inequities.

**Janus kinase (JAK) inhibitors (Update 11, published 16 September 2022)**

**Overview**

JAK inhibitors inhibit intracellular signalling in response to numerous interleukins, interferons, colony stimulating factors, and hormones. As a consequence, they interfere with many cellular responses, including antiviral responses, angiotensin-converting enzyme 2 (ACE2) expression, T cell function and differentiation, and macrophage activation. Baricitinib, ruxolitinib, and tofacitinib are three of at least nine JAK inhibitors. Their inherent differences, as well as variation in dosing and administration and pharmacokinetics, limit class-wide recommendations, and the GDG decided to make separate recommendations for individual drugs.

**Update**—The existing strong recommendation concerning baricitinib for patients with severe or critical covid-19 was updated by the GDG in this twelfth version of the living guideline. This follows the availability of new clinical trial evidence for baricitinib administered in combination with corticosteroids and IL-6 receptor blockers suggesting that the incremental survival benefit afforded by baricitinib exists even among patients also treated with corticosteroids and IL-6 receptor blockers.

**Evidence**—For patients with covid-19, data were derived from four trials that enrolled 8151 patients for baricitinib, two trials that enrolled 475 inpatients for ruxolitinib, and one trial that enrolled 289 patients for tofacitinib. See MAGICapp for detailed description of the mechanism of action, evidence, and subgroup analyses underpinning the recommendation (https://app.magicapp.org/#/guideline/nBkOtiE/rec/E5A0aN).

**Recommendation 1: We recommend treatment with baricitinib for patients with severe or critical covid-19 (strong recommendation).**

**Understanding the recommendation**

In this update, the GDG confirmed the existing strong recommendation to use baricitinib in patients with severe or critical covid-19. This update was based on additional data from 8156 patients enrolled in the RECOVERY trial, which confirmed a survival benefit (now high certainty evidence) and other benefits, with little or no serious adverse events, of a drug that may be administered easily. The GDG acknowledged that some serious adverse events, such as fungal infections, may not have been accurately captured during the relatively short follow-up period in the included trials. Because of different mechanisms of action, the GDG considered baricitinib separately from other JAK inhibitors.

Costs and access remain important considerations, and the GDG recognises that this recommendation could exacerbate health inequities. This strong recommendation further strengthens the impetus to address these concerns and maximise access across regions and countries. The GDG did not anticipate important variability in patient values and preferences, and judged that other contextual factors would not alter the recommendation.

The GDG had previously made a strong recommendation for use of IL-6 receptor blockers (tocilizumab and sarilumab) or baricitinib as alternative agents administered in addition to corticosteroids for patients with severe or critical covid-19. The GDG had elected to refrain from recommending the combination of these three immunosuppressive drugs until clear evidence of incremental benefit emerged. The RECOVERY trial has now provided this evidence, demonstrating that combining corticosteroids, IL-6 receptor blockers, and baricitinib provides incremental survival benefit.

In RECOVERY, 2659 patients received baricitinib along with corticosteroids and IL-6 receptor blockers. The effect of baricitinib in this subgroup was consistent with the beneficial effect of baricitinib in patients who were not treated with IL-6 receptor blockers.

Although these three immunosuppressive drugs are recommended and may be administered jointly, the panel anticipated that there would be situations where clinicians may opt for less aggressive immunosuppressive therapy or choose to combine medications in a stepwise fashion in patients who are deteriorating. However, since
the drugs have not undergone direct comparisons, the GDG felt that clinicians should choose between baricitinib and IL-6 receptor blockers on the basis of experience and comfort using the drugs, local institutional policies, route of administration (baricitinib is oral; IL-6 receptor blockers are intravenous), and cost.

**Balance of benefits and harms**—In patients with severe or critical illness, baricitinib reduces mortality (high certainty), and probably reduces duration of mechanical ventilation and hospital length of stay (both moderate certainty). Treatment probably results in little or no increase in serious adverse events leading to drug discontinuation (moderate certainty). Some serious adverse events such as severe infections which may arise from immunosuppressive therapy like baricitinib may not have been accurately captured during the relatively short follow-up in the included trials. This risk may vary in different parts of the world according to the local prevalence of infections such as tuberculosis. This risk may also be less pertinent, given the short course of baricitinib used for the treatment of covid-19.

Subgroup analyses were undertaken for JAK inhibitors as a class (rather than on individual drugs) and revealed no evidence of a subgroup effect on relative risk in younger (<70 years old) versus older patients, those with critical versus severe covid-19, those receiving or not receiving corticosteroids at baseline, and those receiving or not receiving remdesivir or IL-6 blockers at baseline.

**Values and preferences**—The GDG inferred that almost all well informed patients with severe or critical covid-19 would choose to receive baricitinib due to the likely reduction in mortality, and moderate certainty evidence of little or no increase in serious adverse events. The GDG anticipated little variation in values and preferences between patients for this intervention.

**Applicability**—None of the included RCTs for baricitinib enrolled children, or pregnant or lactating women; therefore, the applicability of this recommendation to these groups remains uncertain. The GDG did not have reason to believe that patients in these groups with covid-19 would respond differently; decisions regarding the use of JAK inhibitors in these groups should be guided by discussion between the individual and their healthcare provider.

**Practical issues**—Baricitinib is administered orally once daily as tablets; it can be crushed, dispersed in water, or given via a nasogastric tube. Based on trials informing the recommendation, the recommended dose is 4 mg daily orally in adults with normal renal function for a duration of 14 days or until hospital discharge, whichever is first. The optimal duration of treatment is unknown. Dose adjustments may be needed for patients with leucopenia, renal impairment, or hepatic impairment, all of which should be monitored during treatment, and for patients taking strong organic anion transporter 3 (OAT3) inhibitors such as probenecid, where drug interactions warrant dose reductions.

Baricitinib, like IL-6 receptor blockers, should be initiated at the same time as systemic corticosteroids; there are currently no data to suggest that specific timing during hospitalisation or the course of illness is beneficial.

See MAGICapp for more information regarding practical issues.

**Resource implications, feasibility, equity, and human rights**—Compared with some other candidate treatments for covid-19, baricitinib is expensive. The recommendation does not take into account cost effectiveness. See box 3 for related considerations. As baricitinib is administered orally once daily, hospitalised patients should find it easy to accept this treatment.

**Recommendation 2:** We suggest not to use ruxolitinib or tofacitinib for patients with severe or critical covid-19 (conditional or weak recommendation).

**Understanding the recommendation**

Low to very low certainty evidence for mortality and duration of mechanical ventilation and a possible increase in serious adverse events, particularly for tofacitinib, drove the weak recommendation not to use ruxolitinib or tofacitinib in patients with severe or critical covid-19. Clinicians should consider using ruxolitinib or tofacitinib only if neither baricitinib nor IL-6 receptor blockers (tocilizumab or sarilumab) are available. The GDG emphasised the need for more trial evidence to better inform the recommendations; this is anticipated through ongoing trials for these JAK inhibitors.

**Benefits and harms**—Low to very low certainty evidence from small trials failed to demonstrate benefits for mortality or duration of mechanical ventilation, and suggested tofacitinib may increase adverse events leading to drug discontinuation. When more evidence is available, the GDG acknowledged that these drugs may prove to have similar benefits as baricitinib.

**Values and preferences**—Most well informed patients would decline ruxolitinib or tofacitinib. However, a minority might choose to receive one or the other drug if neither baricitinib nor IL-6 receptor blockers are available, given that the possibility of benefit has not been excluded and a class effect of JAK inhibitors might exist.

**Applicability**—None of the included RCTs enrolled children; therefore, the applicability of this recommendation to children remains uncertain. Uncertainty also remains with regards to the administration of ruxolitinib or tofacitinib to pregnant or lactating women.

**Practical issues**—Both drugs are administered orally twice daily as tablets and can be dispersed in water or administered via nasogastric tube.

The GDG referred to treatment regimens in the included trials, available via MAGICapp, in the absence of other available information. If ruxolitinib or tofacitinib is administered, like with IL-6 receptor blockers, it should be given with systemic corticosteroids; specific timing during hospitalisation or in the context of the course of illness is not specified.

**Resource implications, equity, and human rights**—Efforts to ensure access to drugs should focus on those that are currently recommended.

**Specific uncertainties, emerging evidence, and future research (for all JAK inhibitors)**

- Safety and efficacy of combination therapy of baricitinib with corticosteroids and IL-6 receptor blockers on longer term outcomes
- Safety and efficacy in areas where certain infections (such as HIV infection, tuberculosis, and some fungal infections) are endemic
- Impact of tofacitinib and ruxolitinib relative to that of baricitinib.

**Sotrovimab (Update 11, published 16 September 2022)**

**Overview**

Sotrovimab is a single human monoclonal antibody that binds to a highly conserved epitope in the SARS-CoV-2 spike protein, preventing the virus from entering cells.
Balance of benefits and harms
been rigorously evaluated in clinical trials. Neutralisation would not be sufficient to confirm clinical effectiveness. Notwithstanding, proof of potent in vitro neutralisation activity, the GDG made a strong recommendation against the use of sotrovimab. See MAGICapp for detailed description of the mechanism of action, detailed in vitro neutralisation activity data, and rationale underpinning the recommendation (https://app.magicapp.org/#/guideline/nBkO1E/rec/LA69PM).

Recommendation: We recommend not to use sotrovimab for patients with non-severe covid-19 (strong recommendation).

Understanding the recommendation
Although previous clinical trial evidence available via the LNMA remains accurate,\(^1\) the panel concluded that it is no longer applicable to covid-19 caused by the SARS-CoV-2 variants that are currently circulating globally. The panel surmised that the likelihood of covid-19 caused by former variants was extremely low and that, accordingly, evidence of sotrovimab’s clinical effectiveness for covid-19 was inexistent.

Of note, the panel applied the same rationale to the recommendation for casirivimab-imdevimab.

The GDG agreed that large, high quality clinical trials generally provide the best evidence of clinical effectiveness for therapeutic interventions. The GDG also continues to base its recommendations strictly on critically important outcomes. From the perspective of clinical guidelines, mechanistic studies and surrogate outcomes are useful to identify candidate therapies for clinical trials but are of no use in the evaluation of clinical effectiveness. The panel concluded that the emerging evidence demonstrating that sotrovimab did not neutralise current variants in vitro would have justified not launching clinical trials and now renders the results of previous trials inapplicable. In vitro assays were deemed sufficient to rule out a clinical effect. Notwithstanding, proof of potent in vitro neutralisation would not be sufficient to confirm clinical effectiveness. Therefore, the GDG will only consider making recommendations for new monoclonal antibodies once they have been rigorously evaluated in clinical trials.

Balance of benefits and harms—There was consensus among the panel that it is highly unlikely that the clinical effectiveness of sotrovimab would persist in the absence of adequate in vitro neutralisation of the circulating variants. Accordingly, the panel concluded that the evidence upon which hinged the previous recommendation was no longer applicable.

Values and preferences—The GDG inferred that, in the absence of compelling evidence of clinical effectiveness for the currently circulating SARS-CoV-2 variants, almost all well informed patients would not choose to receive sotrovimab.

Applicability—Given the updated recommendation against treatment, issues pertaining to applicability were felt to be less relevant.

Practical issues—Given the updated recommendation against treatment, related practical issues were felt to be less relevant.

Resource implications, equity, human rights, acceptability, and feasibility—The strong recommendation against the use of sotrovimab is further supported by their challenges with availability and feasibility, such as limited production, intravenous administration, and requirement for expertise to offer such treatment while oral antiviral therapies are available.

Specific uncertainties, emerging evidence, and future research

- Efficacy and safety for severe or critical seronegative covid-19 patients

Casirivimab-imdevimab (neutralising monoclonal antibodies)
(Update 11, published 16 September 2022)

Overview
Casirivimab and imdevimab are two fully human antibodies (REGN10933 and REGN10987) that bind to the SARS-CoV-2 spike protein and have demonstrated antiviral activity in animal models. It has been postulated that administration of a combination of casirivimab and imdevimab might have differential effects in patients who have produced their own anti-SARS-CoV-2 spike protein antibodies (hereafter seropositive) compared with those who have not (hereafter seronegative); it was hypothesised that effects might be larger for, or restricted to, seronegative individuals who have not yet mounted an effective natural antibody response.

Update
An updated recommendation concerning the neutralising antibodies casirivimab-imdevimab for patients with covid-19 was published in this twelfth version of the WHO living guideline. Previously, a conditional recommendation was provided for patients with non-severe covid-19 at highest risk of hospitalisation, and for patients with severe or critical illness with seronegative status. Following the emergence of the currently circulating SARS-CoV-2 variants and subvariants (such as omicron) now dominating worldwide, and availability of in vitro data showing lack of neutralisation activity, the GDG made a strong recommendation against the use of casirivimab-imdevimab for all patients with covid-19. See MAGICapp for detailed description of the mechanism of action, detailed in vitro neutralisation activity data, and rationale underpinning the recommendation (https://app.magicapp.org/#/guideline/nBkO1E/section/LG5NRE).

Recommendation: We recommend not to use casirivimab-imdevimab for patients with covid-19, regardless of illness severity (strong recommendation).

Understanding the recommendation
Although previous clinical trial evidence available via the LNMA remains accurate,\(^1\) the GDG concluded that it is no longer applicable to covid-19 caused by the SARS-CoV-2 variants that are currently circulating globally. The panel surmised that the likelihood of covid-19 caused by former variants was extremely low and that, accordingly, evidence of casirivimab-imdevimab clinical effectiveness for covid-19 was inexistent.

Of note, the panel applied the same rationale to the recommendation for sotrovimab.

The GDG agreed that large, high quality clinical trials generally provide the best evidence of clinical effectiveness for therapeutic interventions. The GDG also continues to base its recommendations strictly on predefined patient-important outcomes. From the perspective of clinical practice guidelines, mechanistic studies and surrogate outcomes are useful to identify candidate therapies for clinical trials but are of no use in the evaluation of clinical effectiveness. The panel concluded that the emerging evidence demonstrating that casirivimab-imdevimab did not neutralise
current variants in vitro would have justified not launching clinical trials and now renders the results of previous trials inapplicable. In vitro assays were deemed sufficient to rule out a clinical effect. Notwithstanding, proof of potent in vitro neutralisation would not be sufficient to confirm clinical effectiveness. Therefore, the GDG will only consider making recommendations for new monoclonal antibodies once they have been rigorously evaluated in clinical trials.

Balance of benefits and harms—There was consensus among the panel that it is highly unlikely that the clinical effectiveness of casirivimab-imdevimab would persist in the absence of adequate in vitro neutralisation of the circulating variants. Accordingly, the panel concluded that the evidence upon which hinged the previous recommendations was no longer applicable.

Values and preferences—The GDG inferred that, in the absence of compelling evidence of clinical effectiveness for the currently circulating SARS-CoV-2 variants, almost all well informed patients would not choose to receive casirivimab-imdevimab.

Applicability—Given the updated recommendation against treatment, issues pertaining to applicability were felt to be less relevant.

Practical issues—Given the updated recommendation against treatment, related practical issues were felt to be less relevant.

Resource implications, acceptability, feasibility, equity, and human rights—The strong recommendation against the use of casirivimab-imdevimab is further supported by their challenges with availability and feasibility, such as limited production, intravenous administration and requirement for expertise to offer such treatment while oral options are available.

Specific uncertainties, emerging evidence, and future research
No specific uncertainties: see uncertainties section in “How to use this guideline” (above)

Fluvoxamine (Update 10, published 14 July 2022)

Overview
Fluvoxamine is a selective serotonin reuptake inhibitor (SSRI) approved as an antidepressant. It increases concentrations of serotonin in the synaptic cleft. Indirect evidence from non-covid-19 disease models has suggested possible anti-inflammatory properties, and mechanistic studies have suggested host-directed antiviral properties. There are currently no published preclinical studies that directly demonstrate a mechanism in covid-19.

Evidence—For patients with non-severe covid-19, data were derived from three trials that enrolled 2208 non-hospitalised patients. See MAGICapp for detailed description of the mechanism of action, evidence, and subgroup analyses underpinning the recommendation (https://app.magicapp.org/#/guideline/nBkO1E/rec/jmYxpV).

Recommendation 1: For patients with non-severe covid-19, we recommend not to use fluvoxamine, except in the context of a clinical trial (recommended only in a research setting).

Understanding the recommendation
Insufficient evidence of benefit, coupled with a lack of a clear mechanism of action and known drug interactions, drove the recommendation against use in clinical care.

Balance of benefits and harms—There was low to moderate certainty evidence suggesting little or no effect on hospitalisation, mortality, or mechanical ventilation, and an absence of reliable data on serious adverse effects attributable to the drug. Fluvoxamine is known for substantial pharmacological interactions. In the largest trial, markedly more patients discontinued treatment in the fluvoxamine group than in the placebo group. Acknowledging that its evaluation of the certainty of the evidence may differ from other published meta-analyses, GDG members pointed out that early stopping due to apparent benefit may have biased the results of the largest trial. They argued that, although the stopping rules were pre-specified, the decision to stop the trial was based on the effect estimate on a composite outcome of questionable importance; meanwhile the number of important events was lower and vulnerable to bias. The GDG also raised concerns regarding the uncertain applicability of this trial conducted in a single country.

Values and preferences—The GDG inferred that almost all well informed patients would choose not to receive fluvoxamine therapy for covid-19 based on available evidence. The GDG did not believe that other considerations, such as feasibility, acceptability, equity, and cost, would affect this specific recommendation. Specifically, the GDG did not consider the potential role of fluvoxamine as an antidepressant for this guideline of medications for covid-19.

Applicability—None of the included studies enrolled children, and the applicability of this recommendation to children is therefore uncertain. However, the GDG did not see a reason to assume that children with covid-19 would respond any differently to treatment with fluvoxamine.

Practical issues—The GDG made a recommendation against using fluvoxamine for treatment of patients with covid-19 outside the setting of a clinical trial, and therefore practical considerations are less relevant for this drug.

Resource implications, acceptability, feasibility, equity, and human rights—Fluvoxamine is relatively inexpensive compared with other drugs used for covid-19, and widely available, including in low income settings. Its use would risk diverting attention and resources away from interventions that are more likely to provide a benefit.

Specific uncertainties, emerging evidence, and future research—The current evidence does not justify using fluvoxamine to treat covid-19. However, the recommendation does not imply that fluvoxamine is proven to be ineffective. Decisions to further investigate the effects of fluvoxamine for covid-19 will likely hinge on how stakeholders perceive the opportunity cost of investigating the effects of fluvoxamine over other candidate therapies. The GDG discussions highlighted the following knowledge gaps:

- What are the effects of fluvoxamine in patients who have a combination of non-severe covid-19 that is at risk of deteriorating and significant symptoms of anxiety?
- What are the side effects of fluvoxamine therapy in patients with non-severe covid-19 at risk of deteriorating?
- What proportion of patients with non-severe covid-19 at risk of deteriorating would be ineligible due to risk of pharmacological interactions?

The GDG surmised that, in the future, investigators would have to carefully consider whether fluvoxamine could still be compared with placebo or no treatment given that effective treatments are available. If the rationale to further investigate fluvoxamine is its advantageous cost and availability, non-inferiority designs may be considered.
Colchicine (Update 10, published 14 July 2022)

Overview

Colchicine is an anti-inflammatory drug used to treat gout, recurrent pericarditis, familial Mediterranean fever, and other inflammatory conditions. Proposed mechanisms for its anti-inflammatory effect include a reduction in neutrophil chemotaxis, inflammasome signalling inhibition, and decreased production of cytokines such as interleukin 1β. No published animal models evaluating colchicine in SARS-CoV-2 infection are published at the time of guideline publication.

Evidence—For patients with non-severe covid-19, data were derived from 10 trials that enrolled 17 914 non-hospitalised patients. See MAGICapp for detailed description of the mechanism of action, evidence, and subgroup analyses underpinning the recommendation.

Recommendation 1: For patients with non-severe covid-19, we recommend against treatment with colchicine (strong recommendation).

Understanding the recommendation

The lack of benefits on hospitalisations, mortality, and mechanical ventilation, combined with possible harms and toxicity, drove the strong recommendation against the use of colchicine in patients with non-severe covid-19.

Balance of benefits and harms—In patients with non-severe covid-19, colchicine has little or no impact on mortality or mechanical ventilation (moderate certainty). It is unclear whether it affects hospitalisations or adverse effects leading to drug discontinuation. The GDG discussed the risk of drug interactions and colchicine’s narrow therapeutic window, particularly in patients with or at risk of hepatic and renal failure. Colchicine toxicity can be severe and sometimes fatal. The planned subgroup analyses for colchicine versus standard care did not show different relative effects for disease severity or age (children, adults, older adults), with no data reported from illness onset.

Values and preferences—The GDG inferred that almost all well informed patients would choose not to receive colchicine.

Applicability—The applicability of this recommendation to children is currently uncertain because none of the included studies enrolled children. However, the GDG did not consider that children with covid-19 would respond any differently to treatment with colchicine.

Practical issues—The GDG made a strong recommendation against using colchicine for treatment of patients with non-severe covid-19, and therefore practical considerations are less relevant.

Resource implications, acceptability, feasibility, equity, and human rights—These considerations did not affect this specific recommendation. Although colchicine is relatively inexpensive compared with other drugs used for covid-19, and widely available, including in low income settings, the evidence does not justify the use of colchicine for non-severe covid-19 anywhere. Although the cost of colchicine may be low, the GDG raised concerns regarding the risk of diverting attention and resources away from interventions that are more likely to provide a benefit.

Specific uncertainties, emerging evidence, and future research—The GDG considered it unlikely that future studies would identify subgroups of patients who would benefit from colchicine.

Nirmatrelvir/ritonavir (Update 9, published 22 April 2022)

Overview

Nirmatrelvir is a SARS-CoV protease inhibitor which prevents viral replication. Nirmatrelvir/ritonavir is administered orally in combination with ritonavir, a HIV protease inhibitor, which boosts its pharmacokinetics. Through its impact on metabolism and clearance, ritonavir is a perpetrator of many drug-drug interactions during active treatment and possibly for several days after treatment completion. While these may be more easily managed with short durations of treatment, twice daily administration involves doubling ritonavir dose relative to most modern antiretroviral regimens.

The antiviral effect of nirmatrelvir has been demonstrated through in vitro animal studies and human models. Nirmatrelvir retains activity against the omicron BA1 variant in vitro, but clinical data are currently unavailable. There remains uncertainty regarding risk of emergence of resistance; in vitro studies have suggested acquired mutations in the protease sequence may significantly reduce nirmatrelvir activity, though variably so depending on type and number of mutations.

Evidence—For patients with non-severe covid-19, data were derived from two trials (EPIC-SR and EPIC-HR) that enrolled 3100 non-hospitalised patients. See MAGICapp for detailed description of the mechanism of action, evidence, and subgroup analyses underpinning the recommendations (https://app.magicapp.org/#/guideline/nBkO1E/rec/LwrMyv and https://app.magicapp.org/#/guideline/nBkO1E/rec/Lrv08R).

Recommendation 1: For patients with non-severe covid-19 at highest risk of hospitalisation, we recommend treatment with nirmatrelvir/ritonavir (strong recommendation).

Understanding the recommendation

Only a minority of patients who are at highest risk are likely to achieve sufficient benefit in reduced admission to hospital. Nirmatrelvir/ritonavir likely represents a superior choice for these patients because it may have greater efficacy in preventing hospitalisation than available alternatives, has fewer concerns with respects to harms than molnupiravir, and is easier to administer than intravenous remdesivir and antibodies.

Balance of benefits and harms—Beyond the important benefits in reducing hospital admission in patients at highest risk of hospital admission, treatment may have little or no impact on mortality; data regarding effects on time to symptom resolution or mechanical ventilation are limited. Treatment does not increase the likelihood of serious adverse effects leading to drug discontinuation, though diarrhoea and altered taste were noted to occur more often with treatment relative to placebo. The GDG also acknowledged the paucity of information related to emergence of resistance.

Values and preferences—The GDG inferred that almost all well informed patients at highest risk of hospitalisation would choose to receive nirmatrelvir/ritonavir.

Applicability—Given included RCTs enrolled only non-pregnant adults, the GDG concluded that nirmatrelvir/ritonavir should not be offered to children or to breastfeeding or pregnant women with covid-19.

Practical issues—As per the large trials informing the recommendation, nirmatrelvir/ritonavir is administered as 300 mg/100 mg orally every 12 hours for five days. Administration should be as early as possible in the course of the disease. In the included studies, nirmatrelvir/ritonavir was administered within five days.
of disease onset. Through its impact on metabolism and clearance, ritonavir is a perpetrator of many drug-drug interactions, warranting serious consideration by clinicians. The Liverpool covid-19 drug interaction checker may constitute a valuable tool for management of drug interactions with nirmatrelvir/ritonavir. Additional considerations regarding practical issues are summarised in MAGICapp.

**Resource implications, acceptability, feasibility, equity, and human rights—**Nirmatrelvir/ritonavir is unlikely to be available for all individuals who, given the option, would choose to receive the treatment.

**Recommendation 2:** For patients with non-severe covid-19 at low risk of hospitalisation, we suggest no nirmatrelvir/ritonavir (conditional or weak recommendation).

**Understanding the recommendation**

Most patients with non-severe covid-19 at low risk of hospitalisation will experience trivial benefits with use of nirmatrelvir/ritonavir. The GDG inferred that most such patients would be uninterested in taking the drug for these trivial benefits. However, there are likely to be an appreciable number of individuals who place a high value on very small reductions in the risk of hospitalisation and would thus choose to use nirmatrelvir/ritonavir; therefore, a conditional (rather than a strong) recommendation was made. The benefits and harms and other factors the GDG considered are comprehensively described in the full version of the guideline (see box 1).

**Specific uncertainties, emerging evidence, and future research (across disease severities)**

No drug-specific uncertainties; see broadly applicable uncertainties section in “How to use this guideline” (above).

**Molnupiravir (Update 8, published 01 March 2022)**

**Overview**

Molnupiravir is an antiviral administered orally. It was re-purposed as an antiviral for covid-19 because it inhibits replication of SARS-CoV-2 with an in vitro potency broadly similar to remdesivir. This inhibitory effect has been shown in animal studies, both at higher and lower doses, with possibly greater efficacy when combined with favipiravir (compared with either drug alone). The drug is active against alpha and beta variants in vivo based on studies in hamsters and human cell models, and delta and omicron variants in vitro (no data in vivo). In vitro and animal studies have suggested the possibility of carcinogenesis; no human data with long term follow-up are available regarding this. There is also residual uncertainty regarding other long term harms; the efficacy of the drug against variants, particularly those with higher replication or transmission rates; the possibility of a selective pressure for resistant mutations at an individual level, with the potential to spread at a population level; and the emergence of new variants related to random mutagenesis arising from molnupiravir’s mechanism of action. These issues are comprehensively described in the full version of the guideline (see box 1).

**Evidence—**For patients with non-severe covid-19, data were derived from six trials that enrolled 4827 patients, of which the LNMA team had access to data for 4796 patients. See MAGICapp for detailed description of the mechanism of action, evidence, and subgroup analyses underpinning the recommendation (https://app.magicapp.org/#/guideline/nBkO1e/rec/EBs5WNb).

**Recommendation 1:** We suggest treatment with molnupiravir for patients with non-severe covid-19, conditional to those at highest risk of hospitalisation (conditional or weak recommendation).

**Understanding the recommendation**

Only a minority of patients who are at highest risk are likely to achieve sufficient benefit. Especially given the safety concerns related to molnupiravir, the WHO and the GDG recognise the need to mitigate risks, both for individual patients and at the population level.

**Balance of benefits and harms—**Molnupiravir probably provides benefits on admission to hospital and time to symptom resolution (both moderate certainty) and may have a small effect on mortality (low certainty), without an increase in short term adverse effects (high certainty). Absolute benefits depend on the prognosis of the individual patient. The GDG also considered potential long term harms of molnupiravir, including risk of malignancy based on preclinical data (very low certainty), in what they ultimately considered to be a close balance between benefits and harms.

**Values and preferences—**The GDG inferred that almost all well informed patients with a low risk of hospitalisation would decline molnupiravir, and only those at highest risk would choose to receive treatment.

**Applicability—**

- Children. Due to evidence of impact on growth plate thickness and decreased bone formation in some animal studies, molnupiravir should not be used in children.
- Pregnancy, breastfeeding, and conception. Since molnupiravir elicited embryo-fetal lethality and teratogenicity in offspring when given to pregnant animals, it should not be used in pregnant or breastfeeding women. If pregnancy status is unclear, one should perform a pregnancy test before starting molnupiravir treatment. Women and people who can get pregnant should be counselled regarding reducing the risk of conception (such as using birth control) during treatment and for at least four days after the last dose of molnupiravir.
- Uncertainty remains regarding consequences to children conceived by fathers receiving or having recently received molnupiravir, and whether spermatogenesis may be especially prone to mutagenic effects. Men who might father a child should use birth control during treatment and for at least three months after the last dose of molnupiravir.
- Mitigation strategies at the population level include active sequence monitoring of SARS-CoV-2 detected in clinical respiratory samples for patients receiving therapy and active pharmacovigilance programmes.

**Practical issues—**As per large trials informing the recommendation, molnupiravir is dosed as 800 mg orally every 12 hours for five days. Administration should be as early as possible in the course of the disease. In the included studies, molnupiravir was administered within five days of symptom onset.

**Resource implications, feasibility, equity, and human rights—**Molnupiravir is unlikely to be available for all individuals who, given the option, would choose to receive the treatment.

**Specific uncertainties, emerging evidence, and future research**

- Need for clinical data to investigate safety and applicability concerns (including in children, lactating or pregnant women, and men; and long term impact on mutagenesis and cancer risk).
Convalescent plasma (Update 6, published 6 December 2021)

Overview

Treatment with convalescent plasma involves the transfer of endogenously produced neutralising antibodies present within the plasma from previously infected and recovered patients into patients with active infection. The concentrations (titre) of neutralising antibodies present within convalescent plasma are highly variable between donors, and various methodologies to measure antibody levels are available.

Evidence—Data were derived from 16 trials enrolling 16,236 patients across illness severities, of which four RCTs with 1,602 patients informed estimates for outcomes in non-severe illness. See MAGICapp for detailed description of the mechanism of action, evidence, and subgroup analyses underpinning the recommendation (https://app.magicapp.org/#/guideline/nBoKo1E/section/LG5NRE).

Recommendation 1: We recommend not to use convalescent plasma for patients with non-severe covid-19 (strong recommendation).

Understanding the recommendation

The GDG noted that, although not demonstrated in the evidence summary, there remains a potential for harms with blood product transfusion. Most importantly, given there was no benefit demonstrated for any of the critical or important outcomes for non-severe covid-19, the GDG did not see any justification for the resources (including time and cost) that would be associated with administration of convalescent plasma.

Balance of benefits and harms—In patients with non-severe illness, convalescent plasma does not have an important impact on mortality (high certainty). Convalescent plasma probably does not affect mechanical ventilation (moderate certainty). There were no data evaluating the risk of hospitalisation with convalescent plasma; the impact is therefore very uncertain. Convalescent plasma probably does not result in important increases in risks of transfusion-related acute lung injury, transfusion-associated circulatory overload (both moderate certainty), or allergic reactions (low certainty).

Values and preferences—Applying the agreed values and preferences, the GDG inferred that almost all well informed patients with non-severe covid-19 would choose against receiving convalescent plasma.

Acceptability and applicability—Although blood transfusion is acceptable to most, there is a subset of the population who will not accept allogenic blood transfusions. There are also regulatory challenges in most jurisdictions related to blood product transfusions. The included RCTs enrolled non-pregnant women and men. The GDG did not have reason to believe that children or pregnant women with covid-19 would respond any differently to treatment with convalescent plasma; the GDG therefore inferred that children and pregnant women should not receive the intervention either.

Practical issues—Issues include, though are not limited to, the identification and recruitment of potential donors, collection of plasma, storage and distribution of plasma, and infusion of convalescent plasma into recipients.

Resource implications, feasibility, equity, and human rights—The GDG noted that convalescent plasma use is associated with significant resource requirements, including identification of potential donors, testing of donors to ensure adequate titres of anti-SARS-CoV-2 antibodies, collection of donor plasma, storage of plasma, transportation of plasma to recipient location, and administration of plasma. These resources and feasibility issues are compounded for those with non-severe illness, who are most often outpatients. Also, this process is costly and time consuming. Given the number of patients with non-severe illness and the low event rate in this subgroup of patients, mobilising the use of convalescent plasma on a large scale would be of questionable feasibility.

Recommendation 2: We recommend not to use convalescent plasma for patients with severe or critical covid-19, except in the context of a clinical trial (recommended only in research settings).

Understanding the recommendation

Given relative benefits and harms, the GDG agreed further research addressing these patient-important outcomes would be valuable for patients with severe or critical illness. A recommendation to use a drug only in the setting of clinical trials is appropriate when there is low certainty evidence, and future research has potential to reduce uncertainty about the effects of the intervention, and for doing so at a reasonable cost.

Balance of benefits and harms—In patients with severe or critical covid-19, convalescent plasma may not result in an important impact on mortality, mechanical ventilation, time to symptom improvement, length of hospital stay, or ventilator-free days (all low or very low certainty). Convalescent plasma probably does not result in important increases in risks of transfusion-related acute lung injury, transfusion-associated circulatory overload (both moderate certainty), or allergic reactions (low certainty). However, there is always potential for harms with blood product transfusions.

Values and preferences—Applying the agreed values and preferences, the GDG inferred that almost all well informed patients would choose against receiving convalescent plasma outside the research setting.

Specific uncertainties, emerging evidence, and future research

- Effects of high titre convalescent plasma on mortality and other patient-important outcomes.
- Effects in patients with seronegative antibody status.

Interleukin-6 (IL-6) receptor blockers (Update 4, published 6 July 2021)

Overview

IL-6 receptor blockers tocilizumab and sarilumab are monoclonal antibodies approved for use in rheumatoid arthritis. Elevated IL-6 concentrations are associated with severe outcomes in covid-19, including respiratory failure and death. IL-6 receptor blockers antagonise membrane-bound and soluble forms of the IL-6 receptor, blocking the cytokine’s activation and regulation of the immune response to infection.

Update—WHO has made a strong recommendation for JAK inhibitors, specifically baricitinib, in patients with severe or critical covid-19. As per guidance updated in this twelfth iteration of the guideline, an IL-6 receptor blocker and baricitinib may now be given as combination therapy (see discussion for JAK inhibitors above).

Evidence—In addition to the linked network meta-analysis, this recommendation was also informed by an independent prospective meta-analysis from the WHO Rapid Evidence Appraisal for covid-19 group. The network meta-analysis included 30 RCTs with 10,618 participants, and these data were used by the GDG for all outcomes other than mortality. We used the prospective meta-analysis for
mortality because it included additional data that was unpublished at the time. The prospective meta-analysis pooled data from 22 RCTs with 10 156 participants.24 See MAGICapp for detailed description of the mechanism of action, evidence, and subgroup analyses underpinning the recommendation (https://app.magicapp.org/#/guideline/nBkO1E/section/LG5NRE).

Recommendation: We recommend treatment with IL-6 receptor blockers (tocilizumab or sarilumab) for patients with severe or critical covid-19 (strong recommendation).

Understanding the recommendation

Of note, corticosteroids have previously been strongly recommended in patients with severe or critical covid-19, and we recommend that patients meeting these severity criteria should now receive both corticosteroids and IL-6 receptor blockers, possibly with baricitinib as combination therapy (see above).

Balance of benefits and harms—IL-6 receptor blockers reduce mortality and need for mechanical ventilation (both high certainty), and may reduce durations of mechanical ventilation and hospitalisation (both low certainty).

There was uncertainty about the risk of serious adverse effects (very low certainty). There may be little or no increased risk of bacterial infections. However, the GDG had some concerns that, given the short term follow-up of most trials and the challenges associated with accurately capturing adverse events such as bacterial or fungal infections, that the evidence summary may under-represent the risks of treatment with IL-6 receptor blockers. Furthermore, the trials of IL-6 receptor blockers that inform this recommendation were mostly performed in high-income countries, where the risk of infectious complications may be less than in some other parts of the world; the generalisability of the data on these adverse events is therefore unclear.

Values and preferences—The GDG inferred that almost all well informed patients with severe or critical covid-19 infection would want to receive IL-6 receptor blockers, given the reduction in mortality and mechanical ventilation, despite low certainty around evidence for serious adverse events. A minority of the GDG felt that a significant proportion of patients might decline the intervention due to the uncertainties around harms, and taking into account the small reduction in mortality.

Applicability—None of the included RCTs enrolled children or pregnant women. Although this resulted in uncertain applicability, the GDG did not have reason to believe that children or pregnant women with COVID-19 would respond any differently to treatment with IL-6 receptor blockers.

Practical issues—IL-6 receptor blockers require intravenous administration but only require one, or at most two, doses. See MAGICapp for practical information, including considerations if IL-6 receptor blockers are considered in children and pregnant women.

Resource implications, acceptability, feasibility, equity, and human rights—Compared with other treatments for covid-19, IL-6 receptor blockers are expensive and may be inaccessible. The recommendation does not consider cost effectiveness. Given limited availability of the drug, one may consider the relative effects (odds ratio 0.85) for reduction in mortality with IL-6 receptor blockers result in 28 fewer deaths per 1000 patients (95% confidence interval 9 to 47 fewer deaths) in critically ill patients, compared with 12 fewer deaths per 1000 patients (4 to 19 fewer deaths) in severely ill patients.

Finally, sarilumab is not indicated for use in children; therefore, there could be a preference for tocilizumab in this subgroup.

Specific uncertainties, emerging evidence, and future research

- Safety data, including nosocomial infections.
- Immunity and the risk of subsequent infection, which may affect the risk of death after 28 days.
- Outcomes by different IL-6 receptor blocker dosing, and optimal timing of drug initiation.

Ivermectin (Update 3, published 31 March 2021)

Overview

Ivermectin is an antiparasitic agent that interferes with nerve and muscle function of helminths through binding glutamate-gated chloride channels. The treatment is relatively inexpensive and accessible internationally. We currently lack persuasive evidence of a mechanism of action for ivermectin in covid-19; any observed clinical benefit would be unexplained.

We are aware of a few new, relatively small trials published since our recommendation was made, and that one key trial has since been retracted, given concerns about research fraud.25 26 However, the updated evidence summary from the living network meta-analysis is consistent with our previous recommendation. This updated evidence summary will be fully considered by the GDG in an upcoming iteration of the guideline.

Evidence—The living systematic review and network meta-analysis pooled data from 16 trials with 2407 participants. Of the included trials, 75% examined patients with non-severe illness, and 25% included patients with both severe and non-severe illness. See MAGICapp for detailed description of the mechanism of action, evidence, and subgroup analyses underpinning the recommendation (https://app.magicapp.org/#/guideline/nBkO1E/section/LG5NRE).

Recommendation: We recommend not to use ivermectin for patients with covid-19, regardless of illness severity, except in the context of a clinical trial (recommended only in research settings).

Understanding the recommendation

Very low certainty evidence was a critical factor in the recommendation.

Balance of benefits and harms—Certainty of evidence for mortality was deemed very low, despite a point estimate and confidence interval that seemed to suggest benefit with ivermectin; similar judgments were made for other outcomes, including mechanical ventilation, hospital admission, duration of hospitalisation, and viral clearance.

Ivermectin may have little or no effect on time to clinical improvement (low certainty) and may increase the risk of adverse events leading to drug discontinuation (low certainty). A recommendation to only use a drug in the setting of clinical trials is appropriate when there is very low certainty evidence, and when future research has large potential for reducing uncertainty about the effects of the intervention and at a reasonable cost.

Subgroup analyses indicated no effect modification based on dose. We were unable to examine subgroups based on age or severity of illness due to insufficient trial data. Therefore, we assumed similar effects across all subgroups.

Values and preferences—The GDG inferred that almost all well informed patients would not want to receive ivermectin, given...
available evidence left a very high degree of uncertainty in effects on critical outcomes and the possibility of harms, such as adverse events associated with treatment.

**Applicability**—None of the included trials enrolled children or pregnant women; the applicability of the evidence to these subgroups is therefore uncertain, though there is no rationale to suggest they would respond differently.

**Resource implications, acceptability, feasibility, equity, and human rights**—Although the cost of ivermectin may be low per patient, the GDG raised concerns about diverting attention and resources away from care likely to provide a benefit, such as corticosteroids in patients with severe covid-19, and other supportive care interventions. Also, use of ivermectin for covid-19 would divert supply away from pathologies for which it is clearly indicated, potentially contributing to drug shortages, especially for helminth control and elimination programmes. If corticosteroids are used in the treatment of covid-19, empiric treatment with ivermectin may still be considered in strongyloidiasis-endemic areas, albeit not for treatment of covid-19 itself.

**Specific uncertainties, emerging evidence, and future research**

No specific uncertainties: see uncertainties section in “How to use this guideline” (above).

**Hydroxychloroquine (Update 2, published 17 December 2020)**

**Evidence**—The recommendation addressing hydroxychloroquine was informed by results from the living network meta-analysis, pooling data from 30 RCTs with 10 921 participants. See MAGICapp for detailed description of the evidence and subgroup analyses underpinning the recommendation (https://app.magicapp.org/#/guideline/nBkOhE/section/j197zj).

**Recommendation:** We recommend not to use hydroxychloroquine or chloroquine for patients with covid-19, regardless of illness severity (strong recommendation).

**Understanding the recommendation**

**Balance of benefits and harms**—Hydroxychloroquine and chloroquine probably do not reduce mortality or mechanical ventilation (both moderate certainty) and may have no effect on duration of hospitalisation (low certainty). The evidence does not exclude the potential for a small increased risk of death and mechanical ventilation with hydroxychloroquine. The effect on other less important outcomes, including time to symptom resolution, admission to hospital, and viral clearance, remains uncertain.

Hydroxychloroquine may increase the risk of diarrhoea and nausea or vomiting (both low certainty), a finding consistent with evidence from its use in other conditions. Diarrhoea and vomiting may increase the risk of hypovolaemia, hypotension, and acute kidney injury, especially in settings where healthcare resources are limited. Whether and to what degree hydroxychloroquine increases the risk of cardiac toxicity, including life-threatening arrhythmias, when used in patients with covid-19 is uncertain (very low certainty). Subgroup analyses indicated no effect modification based on severity of illness, age, cumulative dose, or predicted day 3 serum trough concentrations. Therefore, we assumed similar effects in all subgroups.

We also reviewed evidence comparing the use of hydroxychloroquine plus azithromycin versus hydroxychloroquine alone. There was no evidence that the addition of azithromycin modified the effect of hydroxychloroquine for any outcome (very low certainty).

**Values and preferences**—Applying the agreed values and preferences, the GDG inferred that almost all well informed patients would not want to receive hydroxychloroquine.

**Applicability**—None of the included trials enrolled children or adolescents; the applicability to this subgroup is therefore uncertain.

**Resource implications, feasibility, equity, and human rights**—Hydroxychloroquine and chloroquine are relatively inexpensive compared with other drugs used for covid-19 and are already widely available, including in low-income settings. Although the cost may be low per patient, the GDG raised concerns about diverting attention and resources away from care likely to provide a benefit such as corticosteroids in patients with severe covid-19 and other supportive care interventions.

**Specific uncertainties, emerging evidence, and future research**

Although some uncertainty remains, the GDG felt that further research was unlikely to uncover a subgroup of patients who would benefit from hydroxychloroquine on the most important outcomes (mortality, mechanical ventilation) given the consistent results in trials across illness severity and location.

**Lopinavir-ritonavir (Update 2, published 17 December 2020)**

**Evidence**—The recommendation was informed by data from seven RCTs with 7429 participants. See MAGICapp for detailed description of the evidence and subgroup analyses underpinning the recommendation (https://app.magicapp.org/#/guide-line/nBkOhE/section/EgybK).

**Recommendation:** We recommend not to use lopinavir-ritonavir for patients with covid-19, regardless of illness severity (strong recommendation).

**Balance of benefits and harms**—Lopinavir-ritonavir probably has little or no effect on mortality and mechanical ventilation (both moderate certainty); effects on time to clinical improvement and other patient-important outcomes were uncertain (low or very low certainty). Treatment probably increases the risk of diarrhoea and nausea or vomiting (both moderate certainty), a finding consistent with the indirect evidence evaluating its use in patients with HIV infection. Diarrhoea and vomiting may increase the risk of hypovolaemia, hypotension, and acute kidney injury, especially in settings where healthcare resources are limited. There was an uncertain effect on viral clearance and acute kidney injury.

Subgroup analysis indicated no effect modification based on severity of illness or age. As there was no evidence of a statistical subgroup effect, we did not formally evaluate credibility. Although the trials did not report subgroup effects by time from symptom onset, many of the trials enrolled patients early in the illness course. The GDG therefore felt that the evidence applies to all patients with covid-19.

**Values and preferences**—Applying the agreed values and preferences, the GDG inferred that almost all well informed patients would not want to receive lopinavir-ritonavir given that the evidence suggested there was probably no effect on mortality or need for mechanical ventilation and there was a risk of adverse events including diarrhoea and nausea or vomiting.

**Resource implications, feasibility, equity, and human rights**—Although the cost of lopinavir-ritonavir is not as high as some other investigational drugs for covid-19 and the drug is generally available in most healthcare settings, the GDG raised concerns about opportunity costs and the importance of not drawing attention and
resources away from best supportive care or the use of corticosteroids in severe covid-19.

Specific uncertainties, emerging evidence, and future research
Although some uncertainty remains, the GDG felt that further research was unlikely to uncover a subgroup of patients who would benefit from lopinavir-ritonavir on the most important outcomes (mortality, mechanical ventilation) given the consistent results in trials across illness severity and location.

Systemic corticosteroids (Original publication, published 4 September 2020)

Evidence—The GDG reviewed evidence from eight RCTs (7184 patients) evaluating systemic corticosteroids versus usual care in treatment of covid-19, seven of which reported mortality data by subgroup of illness severity. Mortality data from one trial, GLUCOCOVID, were not incorporated in the summary of finding for mortality because the mortality outcome data were not available by subgroup. See MAGICapp for detailed description of the evidence and subgroup analyses underpinning the recommendation and practical information on how to administer systemic corticosteroids (https://app.magicapp.org/#/guideline/nBk0zE/section/nByvRL).

Update—Whereas the recommendations remain unchanged, the evidence summary available via MAGICapp for corticosteroids was updated before the fifth iteration of the living guideline. The baseline risk estimates for mortality are now based on the WHO SOLIDARITY trial (as for other drugs in this guideline) rather than the initial ISARIC cohort study that likely overestimates current mortality risks at the global level. This update was also needed to inform the baseline risk for mortality in the evidence summary informing the strong recommendation for IL-6 inhibitors in addition to standard care for patients with severe or critical covid-19, where corticosteroids provide a relative reduction in mortality by 21%.

Recommendation 1: We recommend treatment with systemic corticosteroids for patients with severe or critical covid-19 (strong recommendation).

Balance of benefits and harms—Ultimately, the GDG made its recommendation on the basis of a 28-day mortality reduction of 3.4% in severe or critical covid-19 combined (moderate certainty). Systemic corticosteroids probably reduce the need for mechanical ventilation (moderate certainty).

Overall, the GDG has reasonable certainty that the adverse effects, when considered together, are sufficiently limited in importance and frequency, and suggested that corticosteroids administered in these doses for 7-10 days are not associated with an increased risk of adverse events, beyond likely increasing the incidence of hyperglycaemia and hyponatremia (both moderate certainty). In contrast with new agents proposed for covid-19, clinicians have vast experience administering systemic corticosteroids, and the GDG was reassured by their overall safety profile.

Values and preferences—The GDG took an individual patient perspective to values and preferences but, given the burden of the pandemic for healthcare systems globally, also placed a high value on resource allocation and equity. The benefits of corticosteroids on mortality were deemed of critical importance to patients, with little or no anticipated variability in their preference to be offered treatment if severely ill from covid-19.

Applicability—Applicability is less clear for populations that were under-represented in the considered trials, such as children, patients with tuberculosis, and those who are immunocompromised. In considering potential contraindications to short term systemic corticosteroids in such patients, clinicians must determine if they warrant depriving a patient of a potentially lifesaving therapy. Clinicians should exercise caution in use of corticosteroids in patients with diabetes or underlying immunocompromise. The GDG was confident that clinicians using these guidelines would be aware of additional potential side effects and contraindications to systemic corticosteroid therapy, which may vary geographically in function of endemic microbiological flora.

Acceptability and practical issues—The relative short duration of a course of systemic corticosteroid therapy, and the generally benign safety profile of systemic corticosteroids administered for up to 7-10 days led the GDG to conclude that the acceptability of this intervention was high. Practical issues are summarised in detail on MAGICapp.

Resource implications, feasibility, equity, and human rights—Systemic corticosteroids are low cost, easy to administer, and readily available globally. Dexamethasone and prednisolone are among the most commonly listed medicines in national essential medicines lists; listed by 95% of countries. Accordingly, systemic corticosteroids are among a relatively small number of interventions for covid-19 that have the potential to reduce inequities and improve equity in health. Those considerations influenced the strength of this recommendation.

Recommendation 2: We suggest not to use systemic corticosteroids for patients with non-severe covid-19 (conditional or weak recommendation).

Balance of benefits and harms—Systemic corticosteroids may increase the risk of 28-day mortality (low certainty).

Values and preferences—The conditional recommendation was driven by likely variation in patient values and preferences. The GDG judged that most individuals with non-severe illness would decline systemic corticosteroids. However, many may want them after shared decision making with their treating physician.

Applicability—Several specific circumstances were considered.

- Systemic corticosteroids should not be stopped for patients with non-severe covid-19 who are already treated with systemic corticosteroids for other reasons (such as patients with chronic obstructive pulmonary disease or chronic autoimmune disease).
- If the clinical condition of patients with non-severe covid-19 worsens (that is, increase in respiratory rate, signs of respiratory distress or hypoxaemia) they should receive systemic corticosteroids (see recommendation 1).
- Pregnancy: antenatal corticosteroid therapy may be administered for pregnant women at risk of preterm birth from 24 to 34 weeks’ gestation when there is no clinical evidence of maternal infection and adequate childbirth and newborn care are available. In cases where the woman presents with mild or moderate covid-19, the clinical benefits of antenatal corticosteroid might outweigh the risks of potential harm to the mother. In this situation, the balance of benefits and harms for the woman and the preterm newborn should be discussed with the woman to ensure an informed decision, as this assessment may vary depending on the woman’s clinical condition, her wishes and those of her family, and available healthcare resources.
- Endemic infections that may worsen with corticosteroids should be considered. For example, for Strongyloides stercoralis hyper-infection associated with corticosteroid therapy, diagnosis or empiric treatment may be considered in endemic areas if steroids are used.
Resource implications, feasibility, equity, and human rights—To help guarantee access to systemic corticosteroids for patients with severe or critical covid-19, it is reasonable to avoid their administration to patients who, given the current evidence, do not seem to derive any benefit from this intervention.

Specific uncertainties, emerging evidence, and future research

Remaining uncertainties include effects on:

- Patients with non-severe covid-19 (that is, pneumonia without hypoxaemia).
- Immunity and the risk of a subsequent infection, which may affect the risk of death after 28 days.
- By different steroid preparation, dosing, and optimal timing of drug initiation.

Sources of evidence

To create recommendations, the GDG relied on evidence synthesised in two living network meta-analyses coordinated by MAGIC.6,7

Derivation of absolute effects for drug treatments

For patients with non-severe illness, we used the median of the control arm of the RCTs that contributed to the evidence. For patients with severe or critical illness, the GDG identified the control arm of the WHO SOLIDARITY trial, performed across a wide variety of countries and geographical regions, as representing the most relevant source of evidence for baseline risk estimates for mortality and mechanical ventilation.4 Systemic corticosteroids now represent standard of care in patients with severe or critical covid-19 (see strong recommendation issued by WHO in September 2020). Therefore, the baseline risk estimates in the evidence summaries for JAK inhibitors, convalescent plasma and IL-6 receptor blockers were adjusted for treatment effects of corticosteroids for the outcome of mortality and mechanical ventilation.4 For other outcomes, we used the median of the control arm of the RCTs that contributed to the evidence. Baseline risks, and thus absolute effects, may vary significantly geographically and over time. Thus, users of this guideline may prefer estimating absolute effects by using local event rates. Recommended combinations of treatments are based on direct comparisons from trials demonstrating additional benefit, such as adding baricitinib or interleukin-6 receptor blockers to systemic corticosteroids in patients with severe or critical covid-19. In patients with non-severe covid-19 the absence of direct comparisons from RCTs necessitate indirect comparisons from the living network meta-analysis to inform judgments made about alternative treatment options.
We thank all the following collaborators who contributed to this endeavour, as detailed in the WHO guidance (see link in box 1).

Post-publication review through rapid responses on bmj.com and through MAGICapp.

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- World Health Organisation (WHO) Secretariat for Therapeutics and COVID-19
- External reviewers for WHO

We also thank:

- The living systematic review and network meta-analysis team, led by investigators Reed Siemieniuk and Romina Brignardello-Petersen at McMaster University, Canada
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Main infographic: Summary of recommendations and evidence


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