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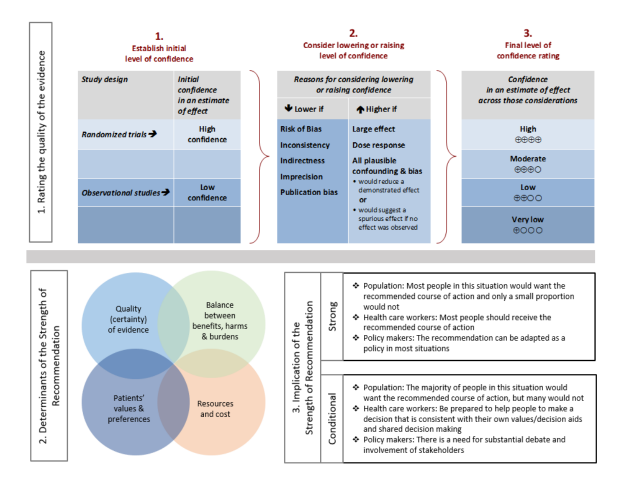
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# **Methods**

• Approach and implications to rating the quality of evidence and strength of recommendations using GRADE methodology

**Figure 1.** Approach and implications to rating the quality of evidence and strength of recommendations using GRADE methodology (unrestricted use of figure granted by the U.S. GRADE Network)



# Hydroxychloroquine/chloroquine & hydroxychloroquine/chloroquine + azithromycin

## **Evidence profiles**

- Hydroxychloroquine compared to no hydroxychloroquine for hospitalized patients with COVID-
- Hydroxychloroquine and azithromycin compared to no hydroxychloroquine/azithromycin for hospitalized patients with COVID-19

**Tables and Figures** 

Table 1. GRADE evidence profile, Recommendation 1

Question: Hydroxychloroquine compared to no hydroxychloroquine for hospitalized patients with COVID-19

Last reviewed and updated 12/23/2020

			Certainty as	sessment			<b>№</b> of p	atients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	hydroxy- chloroquine	no hydroxy- chloroquine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(RCTs) (follow	up: ran	ge 22 days to 49	days)								
5 <sup>1-5</sup>	randomized trials	not serious a	not serious	not serious <sup>b</sup>	serious <sup>c</sup>	none	561/2976 (18.9%)	908/4532 (20.0%)	<b>RR 1.08</b> (0.99 to 1.19)	16 more per 1,000 (from 2 fewer to 38 more)	⊕⊕⊕○ MODERATE	CRITICAL
Clinical s	tatus (assesse	d with:	 7-point scale; hiç	l gher signifies w	 /orsening seve	rity)						
12	randomized trials	serious d	not serious	not serious	serious <sup>e</sup>	none	159	173	-	median <b>1.21 higher</b> (0.69 higher to 2.11 higher)	ФФОО	CRITICAL
	. ,											
			nical ventilation				402/0400	004/0447	DD 4.40	0 4 000		ODITION
2 <sup>1,3</sup>	randomized trials	serious f	not serious	not serious	serious <sup>c</sup>	none	193/2162 (8.9%)	281/3447 (8.2%)	<b>RR 1.10</b> (0.92 to 1.31)	8 more per 1,000 (from 7 fewer to 25 more)	⊕⊕⊖⊖ LOW	CRITICAL
Arrhythm	ias											
16	observational studies	very serious g	not serious	not serious	very serious e,h	none	44/271 (16.2%)	23/221 (10.4%)	<b>RR 1.56</b> (0.97 to 2.50)	58 more per 1,000 (from 3 fewer to 156 more)	⊕⊖⊖⊖ VERY LOW	CRITICAL
Adverse	events, any											
4 2,7-9	randomized trials	serious i	not serious	not serious	serious <sup>e</sup>	none	94/315 (29.8%) <sup>j</sup>	18/176 (10.2%) <sup>k</sup>	<b>RR 2.36</b> (1.49 to 3.75)	139 more per 1,000 (from 50 more to 281 more)	ФФО LOW	IMPORTANT

**Tables and Figures** 

			Certainty as	sessment			<b>№</b> of p	atients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	hydroxy- chloroquine	no hydroxy- chloroquine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Severe a		(assess	ed with: untowar	d medical even	t leading to de	ath, a life-threater	ning experienc	e, prolongatio	n of hospitaliza	ation, or persistent o	r significant disal	oility or
14	randomized trials	not serious	not serious	not serious	very serious <sup>e</sup>	none	14/242 (5.8%)	11/237 (4.6%)	OR 1.26 (0.56 to 2.84) <sup>1</sup>	11 more per 1,000 (from 20 fewer to 75 more)	ФФОО LOW	CRITICAL
QT prolo	l ngation (RCTs	)										
1 2	randomized trials	not serious	not serious	not serious	very serious <sup>h</sup>	none	13/89 (14.6%)	1/58 (1.7%)	RR 8.47 (1.14 to 63.03)	<b>129 more per 1,000</b> (from 2 more to 1,000 more)	ФФОО LOW	IMPORTANT
QT prolo	ngation (NRS)											
2 6,10	observational studies	very serious g,m	not serious	not serious	serious <sup>h</sup>	none	46/355 (13.0%)	13/311 (4.2%)	<b>RR 2.89</b> (1.62 to 5.16)	<b>79 more per 1,000</b> (from 26 more to 174 more)	⊕⊖⊖⊖ VERY LOW	IMPORTANT
High cert Moderate Low cert Very low Risk of b	e certainty: We a ainty: Our confid certainty: We ha ias: Study limitat tency: Unexplain	ry confide re modera ence in the ave very lit ions ed heterog	nt that the true effect ately confident in the e effect estimate is li	effect estimate: T mited: The true ef effect estimate: T findings	he true effect is li fect may be subs		the estimate of the	ne effect		it is substantially differer	ıt	
Imprecis		nce in the	estimate of an effect		cular decision							

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio; OR: Odds ratio

#### **Explanations**

- a. Co-interventions were provided to patients in both studies but balanced across arms.
- b. Cavalcanti 2020 excludes persons receiving supplemental oxygen at a rate of more than 4 liters per minute.
- c. The 95% CI cannot exclude the potential for no benefit or harm.
- d. Cavalcanti was an open-label trial.

**Tables and Figures** 

- e. The 95% CI includes the potential for both benefit and harm. Few events suggest the potential for fragility in the estimate.
- Few events suggest the potential for fragility in the estimate.
- g. Concerns with unmeasured and residual confounding. Multiple co-interventions received across arms.
- h. Few events reported do not meet the optimal information size and suggest fragility in the estimate.
- i. Did not report on blinding (including outcome adjudication committee), sequence generation or allocation concealment; Chen J 2020: all patients received nebulized alpha-interferon, 80% vs. 67.7% of subjects received Abidiol in the hydroxychloroquine vs. placebo arm, respectively. Two subjects in the control arm received lopinavir/ritonavir.
- j. Chen J 2020: 4 AEs include diarrhea, fatigue and transient AST elevation. Chen Z 2020: 1 rash, 1 headache. Tang 2020: 21 AEs include disease progression (1%), URI (1%), diarrhea (10%), vomiting (3%).
- k. Three AEs reported in two patients include: AST elevation, creatinine elevation and anemia
- I. aOR: age, sex, baseline COVID Outcome Scale category, baseline Sequential Organ Failure Assessment score, and duration of acute respiratory infection symptoms prior to randomization
- m. Mahevas 2020 does not report on AEs in the comparator arm.

- 1. RECOVERY Collaborative Group, Horby P, Mafham M, et al. Effect of Hydroxychloroguine in Hospitalized Patients with Covid-19. N Engl J Med 2020; 383(21): 2030-40.
- 2. Cavalcanti AB, Zampieri FG, Rosa RG, et al. Hydroxychloroquine with or without Azithromycin in Mild-to-Moderate Covid-19. N Engl J Med 2020; 383: 2041-52.
- 3. WHO Solidarity Trial Consortium, Pan H, Peto R, et al. Repurposed Antiviral Drugs for Covid-19 Interim WHO Solidarity Trial Results. N Engl J Med 2021; 384: 497-511.
- 4. Self WH, Semler MW, Leither L, et al. Effect of hydroxychloroquine on clinical status at 14 days in hospitalized patients with COVID-19: A randomized clinical trial. JAMA **2020**; 324(21): 2165-76.
- 5. Ulrich RJ, Troxel AB, Carmody E, et al. Treating COVID-19 With Hydroxychloroquine (TEACH): A Multicenter, Double-Blind Randomized Controlled Trial in Hospitalized Patients Open Forum Infect Dis **2020**; 7(10): ofaa446.
- 6. Rosenberg ES, Dufort EM, Udo T, et al. Association of treatment with hydroxychloroquine or azithromycin with in-hospital mortality in patients with COVID-19 in New York state. JAMA **2020**; 323(4): 2493:502.
- 7. Chen J, Liu D, Liu L, et al. A pilot study of hydroxychloroquine in treatment of patients with moderate COVID-19. Journal of Zhejiang University (Medical Sciences) **2020**; 49(2): 215-9.
- 8. Chen Z, Hu J, Zhang Z, et al. Efficacy of hydroxychloroquine in patients with COVID-19: results of a randomized clinical trial. medRxiv **2020**; Available at: https://doi.org/10.1101/2020.03.22.20040758 [Preprint 10 April 2020].
- 9. Tang W, Cao Z, Han M, et al. Hydroxychloroquine in patients with mainly mild to moderate coronavirus disease 2019: open label, randomised controlled trial. BMJ **2020**; 369: m1849.
- 10. Mahevas M, Tran V-T, Roumier M, et al. No evidence of clinical efficacy of hydroxychloroquine in patients hospitalized for COVID-19 infection with oxygen requirement: results of a study using routinely collected data to emulate a target trial. medRxiv 2020; Available at: https://doi.org/10.1101/2020.04.10.20060699 [Preprint 14 April 2020].

**Tables and Figures** 

 Table 2. GRADE evidence profile, Recommendation 2

Question: Hydroxychloroquine and azithromycin compared to no hydroxychloroquine/azithromycin for hospitalized patients with COVID-19

Last updated 8/20/2020; last reviewed 12/23/2020

			Certainty as	sessment			<b>№</b> of p	atients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	hydroxy- chloroquine	no hydroxy- chloroquine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(RCTs) (follow	-up: rang	e 22 days to 49 d	lays)								
11	randomized trials	not serious a	not serious	not serious <sup>b</sup>	very serious	none	5/172 (2.9%)	6/173 (3.5%)	HR 0.64 (0.18 to 2.21)	12 fewer per 1,000 (from 28 fewer to 40 more)	ФФОО LOW	CRITICAL
Mortality	(NRS)											
3 2-4	observational studies	very serious e	not serious	not serious	serious <sup>d</sup>	none	between person reported an adju reported an adju adjustment of 0.	s treated with H0 usted HR of 0.98 usted HR in a sul	CQ + AZ and (95% CI: 0.75) oset after prop 5, 1.77); Rose	5, 1.28); Magagnoli pensity score nberg 2020	⊕⊖⊖⊖ VERY LOW	CRITICAL
Clinical s	tatus (assesse	d with: 7	point scale, high	ner values repre	esent worse cli	inical outcomes)						
11	randomized trials	serious <sup>f</sup>	not serious	not serious <sup>b</sup>	serious <sup>d,g</sup>	none	172	173	-	MD <b>0.99 higher</b> (0.57 higher to 1.73 higher)	ФФОО	CRITICAL
Virologic	failure (follow	up: rang	e 5 days to 6 day	s: assessed wi	th: PCR test)							
2 5-7	observational studies	very serious	serious <sup>i</sup>	serious <sup>j</sup>	serious <sup>c</sup>	none	29/71 (40.8%)	12/12 (100.0%) <sup>1</sup>	not estimable		⊕⊖⊖⊖ VERY LOW	IMPORTANT
QT proloi	l ngation (RCTs)	)										
11	randomized trials	not serious	not serious	serious <sup>m,n</sup>	serious <sup>c</sup>	none	17/116 (14.7%)	1/58 (1.7%)	RR 8.50 (1.16 to 62.31)	129 more per 1,000 (from 3 more to 1,000 more)	ФФСС	IMPORTANT

Tables and Figures

			Certainty as	sessment			№ of p	atients	I	Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	hydroxy- chloroquine	no hydroxy- chloroquine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
QT prolor	ngation (NRS)											
27,8	observational studies	very serious h	not serious	serious <sup>n</sup>	serious <sup>c</sup>	none	10/95 (10.5%)	-	-	-	⊕⊖⊖⊖ VERY LOW	IMPORTANT
Serious a	dverse events	i										
11	randomized	serious f	not serious	not serious °	serious c,d	none	5/239 (2.1%)	0/50 (0.0%)	RR 2.34	0 fewer per	$\Phi\Phi \cap \cap$	CRITICAL

#### GRADE Working Group grades of evidence

trials

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio

#### **Explanations**

- a. Co-interventions were provided to patients but balanced across arms. Cavalcanti 2020 was open label; however, likely did not influence the outcome of mortality.
- b. Cavalcanti 2020 excludes persons receiving supplemental oxygen at a rate of more than 4 liters per minute.
- c. A very small number of events. Optimal information size not met.
- d. The 95% CI includes the potential for both benefit and harm.
- e. Concerns with unmeasured and residual confounding. Multiple co-interventions received across arms.
- f. Cavalcanti was an open-label trial.
- g. Optimal information size not met.
- h. No contemporaneous control groups; no adjustment for baseline severity, resulting in high risk for residual confounding
- i. Two case series from France showed divergent results
- Surrogate marker for mortality or resolution of COVID-19.
- k. Gautret reported 21/61 patients as positive at day 6 (estimate from supplied graph); Molina reported 8/10 patients positive at day 5 or 6. Pooled rates of virologic failure using fixed effects inverse variance method resulted in a 43% failure rate (95% CI, 32% to 54%)

 $\Theta\Theta \cup \cup$ 

LOW

1.000

(from 0 fewer to 0 fewer)

(0.13 to

41.61)

**Tables and Figures** 

- I. Gautret reported on a historical viral clearance rate in symptomatic patients from a separate hospital. Criteria for selection of patients remains unclear, as presumably a sizable number of untreated patients could have been available with data on viral clearance.
- m. Indirect measure of arrhythmia-specific mortality.
- n. Azithromycin and hydroxychloroquine can independently cause QT prolongation. Used together there can be an additive effect. Caution should be exercised with other agents known to prolong the QT interval.
- o. Molina 2020: 1/11 leading to treatment discontinuation; Chorin 2020: 9/84 with significant QTc prolongation of more than 500 ms.
- p. Cavalcanti 2020 serious adverse events included pulmonary embolism, Qtc prolongation, myocardial infarction, abdominal-wall hemorrhage.

- 1. Cavalcanti AB, Zampieri FG, Rosa RG, et al. Hydroxychloroquine with or without Azithromycin in Mild-to-Moderate Covid-19. N Engl J Med 2020; 383: 2041-52.
- 2. Rosenberg ES, Dufort EM, Udo T, et al. Association of treatment with hydroxychloroquine or azithromycin with in-hospital mortality in patients with COVID-19 in New York state. JAMA **2020**; 323(4): 2493:502.
- 3. Magagnoli J, Narendran S, Pereira F, et al. Outcomes of hydroxychloroguine usage in United States veterans hospitalized with Covid-19. Med 2020; 1(1): 114-27.e3.
- 4. Ip A, Berry DA, Hansen E, et al. Hydroxychloroquine and Tocilizumab Therapy in COVID-19 Patients-An Observational Study. PloS One 2020; 15(8): e0237693.
- 5. Gautret P, Lagier JC, Parola P, et al. Hydroxychloroquine and azithromycin as a treatment of COVID-19: results of an open-label non-randomized clinical trial. Int J Antimicrob Agents **2020**: 56(1): 105949.
- 6. Gautret P, Lagier JC, Parola P, et al. Clinical and microbiological effect of a combination of hydroxychloroquine and azithromycin in 80 COVID-19 patients with at least a six-day follow up: A pilot observational study. Travel Med Infect Dis **2020**; 34: 101663.
- 7. Molina JM, Delaugerre C, Goff J, et al. No Evidence of Rapid Antiviral Clearance or Clinical Benefit with the Combination of Hydroxychloroquine and Azithromycin in Patients with Severe COVID-19 Infection. Médecine et Maladies Infectieuses **2020**; 50(4): 384.
- 8. Chorin E, Dai M, Shulman E, et al. The QT Interval in Patients with SARS-CoV-2 Infection Treated with Hydroxychloroquine/Azithromycin. medRxiv **2020**; Available at: https://doi.org/10.1101/2020.04.02.20047050 [Preprint 3 April 2020].

# **Hydroxychloroquine as post-exposure prophylaxis**

## **Evidence profiles**

 Hydroxychloroquine compared to no hydroxychloroquine for post-exposure prophylaxis of COVID-19

**Tables and Figures** 

Table 3. GRADE evidence profile, Recommendation 3

Question: Hydroxychloroquine compared to no hydroxychloroquine for post-exposure prophylaxis of COVID-19

New evidence profile developed 9/23/2021

			Certainty a	ssessment			Nº of p	atients	E	ffect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	hydroxy- chloroquine	no hydroxy- chloroquine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
ympton	natic SARS-C	oV-2 infe	ction (follow-up:	14 days) <sup>a</sup>								
3 1,2,3	randomized trials	not serious	not serious	not serious	serious <sup>b</sup>	none	166/1883 (8.8%)	177/1941 (9.1%)	<b>RR 0.95</b> (0.77 to 1.16)	5 fewer per 1,000 (from 21 fewer to 15 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
lospitali	zation (follow	v-up: 14 d	ays)									
3 1,2,3	randomized trials	not serious	not serious	not serious	very serious <sup>b</sup>	none	13/2018 (0.6%)	14/2129 (0.7%)	<b>RR 1.00</b> (0.47 to 2.12)	0 fewer per 1,000 (from 3 fewer to 7 more)	⊕⊕⊖⊖ Low	CRITICAL
ortality	(follow-up: 1	4 days)										
3 1,2,3	randomized trials	not serious	not serious	not serious	very serious <sup>b</sup>	none	5/2018 (0.2%)	12/2129 (0.6%)	<b>RR 0.45</b> (0.16 to 1.28)	3 fewer per 1,000 (from 5 fewer to 2 more)	ФФОО LOW	CRITICAL
Serious a	dverse even	ts (follow-	-up: 14 days)							l		
3 1,2,3	randomized trials	not serious	not serious	not serious	very serious <sup>b</sup>	none	16/2018 (0.8%)	19/2129 (0.9%)	<b>RR 0.91</b> (0.47 to 1.76)	1 fewer per 1,000 (from 5 fewer to 7 more)	ФФСС	CRITICAL
ligh certa Moderate ow certai Yery low c	certainty: We a inty: Our confid certainty: We has: ss: Study limitat	ery confident are moderate ence in the ave very litter ions	t that the true effect ely confident in the e effect estimate is lim	effect estimate: The hited: The true effect effect estimate: The	true effect is likel ct may be substan	y to be close to the estially different from the	stimate of the effect, be estimate of the effect different from the estir	ot .	ty that it is substant	ially different		

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

Publication bias: Selective publication of studies

Imprecision: The confidence in the estimate of an effect to support a particular decision

#### **Explanations**

a. Boulware included both laboratory-confirmed COVID-19 as well as probable COVID-19; 11/49 patients receiving HCQ were laboratory confirmed and 9/58 receiving placebo were laboratory confirmed.

**Tables and Figures** 

b. The 95% CI includes both the potential of benefit and the risk of harm.

- 1. Barnabas RV, Brown ER, Bershteyn A, et al. Hydroxychloroquine as Postexposure Prophylaxis to Prevent Severe Acute Respiratory Syndrome Coronavirus 2 Infection : A Randomized Trial. Ann Intern Med **2021**; 174(3): 344-52.
- 2. Boulware DR, Pullen MF, Bangdiwala AS, et al. A Randomized Trial of Hydroxychloroquine as Postexposure Prophylaxis for Covid-19. N Engl J Med 2020; 383(6): 517-25.
- 3. Mitja O, Corbacho-Monne M, Übals M, et al. A Cluster-Randomized Trial of Hydroxychloroquine for Prevention of Covid-19. N Engl J Med 2021; 384(5): 417-27.

# Lopinavir/ritonavir

## **Evidence profiles**

- Prophylactic lopinavir/ritonavir compared to no prophylactic lopinavir/ritonavir for persons exposed to COVID-19
- Lopinavir/ritonavir compared to no lopinavir/ritonavir for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease
- Lopinavir/ritonavir compared to no lopinavir/ritonavir for hospitalized patients with severe COVID-19

Tables and Figures

Table 4. GRADE evidence profile, Recommendation 4

Question: Prophylactic lopinavir/ritonavir compared to no prophylactic lopinavir/ritonavir for persons exposed to COVID-19

New evidence profile developed 2/16/2022

Symptomatic SARS-COV-2 infection  11 randomized not trials serious  Symptomatic SARS-COV-2 infection	n (COVID-19) regar not serious no n (COVID-19), nega	ardless of bar	serious <sup>a</sup>	Other considerations rology (follow-up: none	prophylactic lopinavir/ ritonavir  21 days)  35/209 (16.7%)	no prophylactic lopinavir/ ritonavir 13/109 (11.9%)	Relative (95% CI) HR 0.60 (0.29 to 1.26) b	Absolute (95% CI) 46 fewer per 1,000 (from 83 fewer to	Certainty  ⊕⊕⊕○	Importance
trials serious  Symptomatic SARS-COV-2 infection  1 randomized not n	not serious no	not serious	serious <sup>a</sup>			13/109 (11.9%)			_	CRITICAL
trials serious  Symptomatic SARS-COV-2 infection  1 randomized not n	n (COVID-19), nega			none	35/209 (16.7%)	13/109 (11.9%)			_	CRITICAL
Tanadinizad not		gative PCR ar						` 29 more)	MODERATE	
Tanadinizad not			na serology at	baseline (follow-	up: 21 days)					
liidis Selious	not serious no	not serious	serious <sup>a</sup>	none	8/159 (5.0%)	7/90 (7.8%)	HR 0.59 (0.17 to 2.02)	31 fewer per 1,000 (from 64 fewer to 73 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Adverse events (follow-up: 29 days)	)									
11 randomized serious on trials	not serious no	not serious	not serious	none	175/207 (84.5%)	33/107 (30.8%)	<b>RR 2.74</b> (2.05 to 3.66)	537 more per 1,000 (from 324 more to 820 more)	⊕⊕⊕○ MODERATE	CRITICAL

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that has not been peer reviewed or published.

CI: Confidence interval; HR: Hazard ratio; PCR: Polymerase chain reaction; RR: Risk ratio

#### **Explanations**

- a. Few events, unable to exclude benefits as well as harms
- b. This pre-specified primary endpoint adjusted analysis is a mixed model analysis adjusted for baseline imbalance

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

- c. Participants not blinded to lopinavir/ritonavir
- d. Two serious adverse events occurred and both judged by the author as unrelated to lopinavir/ritonavir

**Tables and Figures** 

1. Labhardt ND, Smit M, Petignat I, et al. Post-exposure Lopinavir-Ritonavir Prophylaxis versus Surveillance for Individuals Exposed to SARS-CoV-2: The COPEP Pragmatic Open-Label, Cluster Randomized Trial. EClinicalMedicine **2021**; 42: 101188.

**Tables and Figures** 

Table 5. GRADE evidence profile, Recommendation 5

Question: Lopinavir/ritonavir compared to no lopinavir/ritonavir for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease

#### New evidence profile developed 2/16/2022

			Certainty as	sessment			Nº of p	atients	E	ffect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	lopinavir/ ritonavir	no lopinavir/ ritonavir	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow-up:	90 days)										
11	randomized trials	not serious	not serious	not serious	very serious <sup>a</sup>	none	2/244 (0.8%)	1/227 (0.4%)	<b>RR 1.86</b> (0.17 to 20.40)	4 more per 1,000 (from 4 fewer to 85 more)	$\bigoplus_{LOW}\bigcirc$	CRITICAL
COVID-19	9-related hos	spitalizatio	ns (follow-up: 90	days)								
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	serious <sup>a</sup>	none	14/244 (5.7%)	11/227 (4.8%)	HR 1.16 (0.53 to 2.56)	8 more per 1,000 (from 22 fewer to 71 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Serious a	adverse ever	nts (follow-	up: 90 days)									
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	serious <sup>a</sup>	none	20/232 (8.6%)	12/220 (5.5%)	<b>RR 1.58</b> (0.79 to 3.16)	32 more per 1,000 (from 11 fewer to 118 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
High certa Moderate Low certa	certainty: We inty: Our confid	ery confiden are moderat dence in the	t that the true effect ely confident in the effect estimate is lin	effect estimate: Th nited: The true eff	ne true effect is lik ect may be subst		the estimate of th	e effect		s substantially different		
Inconsiste Indirectne Imprecision	ss: Applicabilit	ned heterogo by or general ence in the e	eneity across study a zability to the resea stimate of an effect on of studies	rch question	cular decision							

NB: Certainty ratings may be derived from evidence that has not been peer reviewed or published.

CI: Confidence interval; HR: Hazard ratio; RR: Risk ratio

#### **Explanations**

a. Sparse data, few events, unable to excluded harms as well as benefits

#### References

1. Reis G, Moreira Silva E, Medeiros Silva DC, et al. Effect of Early Treatment With Hydroxychloroquine or Lopinavir and Ritonavir on Risk of Hospitalization Among Patients With COVID-19: The TOGETHER Randomized Clinical Trial. JAMA Netw Open **2021**; 4(4): e216468.

Tables and Figures

Table 6. GRADE evidence profile, Recommendation 6

Question: Lopinavir/ritonavir compared to no lopinavir/ritonavir for hospitalized patients with severe COVID-19

## Last reviewed and updated 11/22/2020

serious <sup>a,d</sup>	not serious  follow up: 28 day not serious  ment discontinuat not serious	not serious	serious b	Other considerations  none  none	166/1655 (10.0%)	938/4896 (19.2%)	RR 1.00 (0.89 to 1.13)  RR 1.12 (0.93 to	Absolute (95% CI)  0 fewer per 1,000 (from 21 fewer to 25 more)  11 more per 1,000 (from 6 fewer to 30	Certainty  MODERATE	CRITICAL
not serious <sup>a</sup> ventilation ( serious <sup>a,d</sup>	follow up: 28 day not serious nent discontinua	ys) not serious			(17.3%) °	(19.2%) 297/3380	(0.89 to 1.13)	(from 21 fewer to 25 more)	MODERATE	
ventilation ( serious a.d	follow up: 28 day not serious nent discontinua	ys) not serious			(17.3%) °	(19.2%) 297/3380	(0.89 to 1.13)	(from 21 fewer to 25 more)	MODERATE	
serious a,d	not serious	not serious	serious <sup>b</sup>	none						CRITICAL
ng to treatm	nent discontinua	tion	serious <sup>b</sup>	none						CRITICAL
		I .					1.34)	more)	LOW	
serious <sup>a</sup>	not serious	not serious								
			very serious <sup>e</sup>	none	Nearly 14% of lopinavir–ritonavir recipients were unable to complete the full 14-day course of administration. This was due primarily to gastrointestinal adverse events, including anorexia, nausea, abdominal discomfort, or diarrhea, as well as two serious adverse events, both acute gastritis. Two recipients had self-limited skin eruptions. Such side effects, including the risks of hepatic injury, pancreatitis, more severe cutaneous eruptions, and QT prolongation, and the potential for multiple drug interactions due to CYP3A inhibition, are well documented with this drug combination. The side-effect profile observed in the current trial arouses concern about the use of higher or more prolonged lopinavir–ritonavir dose regimens in efforts to improve outcomes.				⊕⊖⊖⊖ VERY LOW	IMPORTANT
rovement a	t 14 days (follow	up: 14 days)								
serious <sup>a</sup>	not serious	not serious	very serious <sup>f</sup>	none	54/99 (54.5%)	70/100 (70.0%)	<b>RR 0.78</b> (0.62 to 0.97)	154 fewer per 1,000 (from 266 fewer to 21 fewer)	⊕⊖⊖⊖ VERY LOW	CRITICAL
g	serious a grades of every confident	serious <sup>a</sup> not serious  grades of evidence  ry confident that the true effect li	grades of evidence  ry confident that the true effect lies close to that o	serious a not serious not serious very serious f  grades of evidence  ry confident that the true effect lies close to that of the estimate of the	serious a not serious not serious very serious f none  grades of evidence  rry confident that the true effect lies close to that of the estimate of the effect	recipients had so including the risk cutaneous erupt for multiple drug well documented profile observed use of higher or regimens in efform at 14 days (follow up: 14 days)  serious a not serious not serious very serious form none form the following serious form the following serious form none form the following serious form none form the following serious form the follo	recipients had self-limited ski including the risks of hepatic cutaneous eruptions, and QT for multiple drug interactions well documented with this drup profile observed in the currer use of higher or more prolong regimens in efforts to improve regimens in efforts to improve serious a not serious not serious very serious for none for the serious for the effect serious for the effect for the effect for confident that the true effect lies close to that of the estimate of the effect for cutaneous eruptions, and QT for multiple drug interactions well documented with this drup profile observed in the currer use of higher or more prolong regimens in efforts to improve the cutaneous eruptions, and QT for multiple drug interactions well documented with this drup profile observed in the currer use of higher or more prolong regimens in efforts to improve the cutaneous for multiple drug interactions well documented with this drup profile observed in the currer use of higher or more prolong regimens in efforts to improve the cutaneous for multiple drug interactions well documented with this drup profile observed in the currer use of higher or more prolong regimens in efforts to improve the cutaneous for multiple drug interactions well documented with this drup profile observed in the currer use of higher or more prolong regimens in efforts to improve the cutaneous for multiple drug interactions well documented with the cutaneous for multiple drug interactions well documented with the cutaneous for multiple drug interactions well documented with the cutaneous for multiple drug interactions well documented with the cutaneous for multiple drug interactions well documented to improve the cutaneous for multiple drug interactions well documented to improve the cutaneous for multiple drug interactions and the cutaneous for multiple drug	recipients had self-limited skin eruptions. including the risks of hepatic injury, panor cutaneous eruptions, and QT prolongatio for multiple drug interactions due to CYP3 well documented with this drug combination profile observed in the current trial arouse use of higher or more prolonged lopinavir regimens in efforts to improve outcomes.  Tovement at 14 days (follow up: 14 days)  serious a not serious not serious very serious for none for the effect lies close to that of the estimate of the effect for the effect lies close to that of the estimate of the effect for the effect for the effect lies close to that of the estimate of the effect for the effect lies close to that of the estimate of the effect for the effect lies close to that of the estimate of the effect for the effect lies close to that of the estimate of the effect for the effect lies close to that of the estimate of the effect for the effect lies close to that of the estimate of the effect for the effect lies close to that of the estimate of the effect lies close to that of the estimate of the effect lies close to that of the estimate of the effect lies close to the estimate of the effect lies close to the effect lies close to that of the estimate of the effect lies close to the effect	recipients had self-limited skin eruptions. Such side effects, including the risks of hepatic injury, pancreatitis, more severe cutaneous eruptions, and QT prolongation, and the potential for multiple drug interactions due to CYP3A inhibition, are well documented with this drug combination. The side-effect profile observed in the current trial arouses concern about the use of higher or more prolonged lopinavir—ritonavir dose regimens in efforts to improve outcomes.  **rovement at 14 days (follow up: 14 days)**  **serious a not serious not serious very serious f none 54/99 (54.5%) 70/100 RR 0.78 (0.62 to 0.97) 154 fewer per 1,000 (from 266 fewer to 21 fewer)  **grades of evidence**  **ry confident that the true effect lies close to that of the estimate of the effect**	recipients had self-limited skin eruptions. Such side effects, including the risks of hepatic injury, pancreatitis, more severe cutaneous eruptions, and QT prolongation, and the potential for multiple drug interactions due to CYP3A inhibition, are well documented with this drug combination. The side-effect profile observed in the current trial arouses concern about the use of higher or more prolonged lopinavir–ritonavir dose regimens in efforts to improve outcomes.  **Tovement at 14 days (follow up: 14 days)**  **Serious a not serious not serious very serious formula provided in the current trial arouses concern about the use of higher or more prolonged lopinavir–ritonavir dose regimens in efforts to improve outcomes.  **Tovement at 14 days (follow up: 14 days)**  **Serious a not serious not serious very serious formula provided in the current trial arouses concern about the use of higher or more prolonged lopinavir–ritonavir dose regimens in efforts to improve outcomes.  **Tovement at 14 days (follow up: 14 days)**  **Serious a not serious not serious very serious formula provided in the current trial arouses concern about the use of higher or more prolonged lopinavir–ritonavir dose regimens in efforts to improve outcomes.  **Tovement at 14 days (follow up: 14 days)**  **Serious a not serious not serious very serious formula provided in the current trial arouses concern about the use of higher or more prolonged lopinavir–ritonavir dose regimens in efforts to improve outcomes.  **Tovement at 14 days (follow up: 14 days)**  **Serious a not serious not serious very serious formula provided in the current trial arouses concern about the use of higher or more prolonged lopinavir–ritonavir dose regimens in efforts to improve outcomes.  **Tovement at 14 days (follow up: 14 days)**  **Serious a not serious

**Tables and Figures** 

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

**Imprecision:** The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

#### **Explanations**

- a. Unblinded studies which can affect outcomes that require judgment, such as how investigators judge clinical improvement or decide to stop the treatment in patients with side effects.
- b. 95% CI may not include a meaningful difference.
- c. Modified intention to treat data from Cao 2020 used for this outcome; some deaths were excluded when drug was not given.
- d. One patient randomized to the lopinavir-ritonavir arm in Cao 2020 was mechanically ventilated at baseline.
- e. Small number of events making estimates highly uncertain
- f. The upper boundary of the 95% confidence interval crosses the threshold of meaningful improvement as the worst case estimate is a 3% RRR.

- 1. Cao B, Wang Y, Wen D, et al. A Trial of Lopinavir-Ritonavir in Adults Hospitalized with Severe Covid-19. N Engl J Med 2020; 382(19): 1787-99.
- 2. WHO Solidarity Trial Consortium, Pan H, Peto R, et al. Repurposed Antiviral Drugs for Covid-19 Interim WHO Solidarity Trial Results. N Engl J Med 2021; 384: 497-511.
- 3. RECOVERY Collaborative Group, Horby PW, Mafham M, et al. Lopinavir–ritonavir in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. The Lancet **2020**; 396(10259): 1345-52.

# **Glucocorticoids**

## **Evidence profiles**

- Glucocorticoids compared to no glucocorticoids for critically ill patients with COVID-19
- Glucocorticoids compared to no glucocorticoids for hospitalized patients with severe but not critical COVID-19
- Glucocorticoids compared to no glucocorticoids for hospitalized patients with COVID-19 not receiving supplemental oxygen

**Tables and Figures** 

**Table 7.** GRADE evidence profile, Recommendation 7

Question: Glucocorticoids compared to no glucocorticoids for critically ill patients with COVID-19

#### Last reviewed and updated 9/25/2020

			Certainty a	ssessment			Nº of p	atients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	cortico- steroids	no cortico- steroids	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow up: 28	8 days)										
8 1	randomized trials	not serious	not serious	not serious	not serious	none	280/749 (37.4%)	485/1095 (44.3%)	OR 0.66 (0.54 to 0.82)	99 fewer per 1,000 (from 143 fewer to 48 fewer)	⊕⊕⊕ ніGн	CRITICAL
Hospital o	discharge (fo	llow up: 2	8 days)									
1 2 randomized trials randomized a rot serious a not serious serious b not serious none 1360/2104 (64.6%) RR 1.11 (1.04 to 1.19) 67 more per 1,000 (from 24 more to 116 more)										IMPORTANT		
Serious a	dverse event	s										
Serious adverse events    6												
High certa Moderate of Low certai	certainty: We a nty: Our confide	ry confident re moderate ence in the	t that the true effect li ely confident in the ef effect estimate is limi	ffect estimate: The tr ted: The true effect	rue effect is likely t may be substantia		stimate of the e	ffect	•	t is substantially different	,	

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; OR: Odds ratio; RR: Risk ratio

#### **Explanations**

- a. Analysis adjusted for baseline age.
- b. Indirectness due to different health care system (allocation of intensive care resources in an unblinded study). Indirectness to other corticosteroids.
- c. The 95% CI includes the potential for both harm as well as benefit. Few events reported do not meet the optimal information size and suggest fragility in the estimate.

- 1. WHO Rapid Evidence Appraisal for COVID-19 Therapies Working Group, Sterne JAC, Murthy S, et al. Association Between Administration of Systemic Corticosteroids and Mortality Among Critically III Patients With COVID-19: A Meta-analysis. JAMA 2020; 324(13): 1330-41.
- 2. RECOVERY Collaborative Group, Horby P, Lim WS, et al. Dexamethasone in Hospitalized Patients with Covid-19. N Engl J Med 2021; 384: 693-704.

**Tables and Figures** 

Table 8. GRADE evidence profile, Recommendation 8

Question: Glucocorticoids compared to no glucocorticoids for hospitalized patients with severe but not critical COVID-19

Last reviewed and updated 9/25/2020

			Certainty as	sessment			Nº of ∣	patients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	gluco- corticoids	no gluco- corticoids	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow up: 2	8 days)										
11	randomized trials	not serious <sup>a</sup>	not serious	serious <sup>b</sup>	not serious	none	454/2104 (21.6%)	1065/4321 (24.6%)	RR 0.83 (0.74 to 0.92)	42 fewer per 1,000 (from 64 fewer to 20 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
Hospital	discharge (fo	ollow up: 2	8 days)									
1 1	randomized trials	not serious <sup>a</sup>	not serious	serious <sup>b</sup>	not serious	none	1360/2104 (64.6%)	2639/4321 (61.1%)	RR 1.11 (1.04 to 1.19)	67 more per 1,000 (from 24 more to 116 more)	⊕⊕⊕⊖ MODERATE	IMPORTANT
Adverse	events											
							hyperglycemia agitation/confu	ring a short cours , neurological sid usion), adrenal su Henzen 2000; Si	de effects (e.g ippression, an	., id risk of infection	-	CRITICAL
High certa Moderate Low certal Very low c Risk of bia Inconsiste Indirectne Imprecisio	certainty: We a inty: Our confic certainty: We h as: Study limita cncy: Unexplair ss: Applicability	ery confident are moderate dence in the ave very little tions ned heteroge y or generalize nce in the es	that the true effect lely confident in the e effect estimate is lime e confidence in the e eneity across study fi zability to the resear- stimate of an effect to	ffect estimate: The ited: The true effe iffect estimate: Th ndings ch question	e true effect is like ct may be substa e true effect is like		ne estimate of the	effect	•	is substantially differe	ent	

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

## **Explanations**

- a. Analysis adjusted for baseline age.
- b. Indirectness due to different health care system (allocation of intensive care resources in an unblinded study). Indirectness to other corticosteroids.

#### Reference

1. RECOVERY Collaborative Group, Horby P, Lim WS, et al. Dexamethasone in Hospitalized Patients with Covid-19. N Engl J Med 2021; 384: 693-704.

**Tables and Figures** 

Table 9. GRADE evidence profile, Recommendation 9

Question: Glucocorticoids compared to no glucocorticoids for hospitalized patients with COVID-19 not receiving supplemental oxygen

Last reviewed and updated 9/25/2020

Certainty assessment							<b>№</b> of	patients		Effect			
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	gluco- corticoids	no gluco- corticoids	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance	
Mortality	lortality (follow up: 28 days)												
1 <sup>1</sup>	randomized trials	serious a	not serious	not serious	serious <sup>b</sup>	none	85/501 (17.0%)	137/1034 (13.2%)	<b>RR 1.22</b> (0.93 to 1.61)	29 more per 1,000 (from 9 fewer to 81 more)	ФФОО	CRITICAL	
Hospital	discharge (fo	ollow up: 2	28 days)										
1 1	randomized trials	serious a	not serious	not serious	serious <sup>c</sup>	none	366/501 (73.1%)	791/1034 (76.5%)	<b>RR 0.99</b> (0.87 to 1.12)	8 fewer per 1,000 (from 99 fewer to 92 more)	ФФОО	IMPORTANT	
Adverse	events	I					1			I			
							hyperglycemia	ing a short course , neurological side nd risk of infectior	-	CRITICAL			
High certa Moderate Low certa Very low o Risk of bia Inconsiste	certainty: We a inty: Our confid certainty: We h as: Study limita ency: Unexplain	ery confiden are moderate dence in the nave very little tions ned heteroge	t that the true effect ely confident in the e effect estimate is lin	effect estimate: The hited: The true effe effect estimate: The indings	e true effect is like ect may be substa	ne effect ly to be close to the es ntially different from the ely to be substantially d	estimate of the e	ffect	sibility that it is substa	ntially different			

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

Publication bias: Selective publication of studies

#### **Explanations**

- a. Risk of bias due to post hoc subgroup effect among persons not receiving supplemental oxygen.
- b. The 95% CI includes the potential for appreciable harm and cannot exclude the potential for benefit. Few events reported do not meet the optimal information size and suggest fragility in the estimate.
- c. The 95% CI cannot exclude the potential for either appreciable harm or benefit.

Imprecision: The confidence in the estimate of an effect to support a particular decision

#### Reference

1. RECOVERY Collaborative Group, Horby P, Lim WS, et al. Dexamethasone in Hospitalized Patients with Covid-19. N Engl J Med **2021**; 384: 693-704.

## **Inhaled corticosteroids**

## **Evidence profiles**

• Inhaled corticosteroids compared to no inhaled corticosteroids for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease

**Tables and Figures** 

**Table 10.** GRADE evidence profile, Recommendation 10

**Question:** Inhaled corticosteroids compared to no inhaled corticosteroids for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease **Last reviewed and updated 10/10/2022** 

Certainty assessment							Nº of p	Effect				
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	inhaled corticosteroids	no inhaled corticosteroids	Relative (95% CI)	Absolute (95% CI)		Importance
Mortality	(follow-up: r	ange 14 da	ys to 30 days)									
7 <sup>1-7</sup>	randomized trials	not serious <sup>a</sup>	not serious	not serious <sup>b</sup>	serious <sup>c</sup>	none	7/1951 (0.4%)	13/1925 (0.7%)	<b>RR 0.58</b> (0.24 to 1.44)	3 fewer per 1,000 (from 5 fewer to 3 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Hospitali	zations (follo	w-up: rang	ge 14 days to 30 o	days)								
6 1-3,5,7,8	randomized trials	serious <sup>a</sup>	not serious	not serious d	serious <sup>c</sup>	none	95/1928 (4.9%)	122/1906 (6.4%)	<b>RR 0.81</b> (0.52 to 1.27)	12 fewer per 1,000 (from 31 fewer to 17 more)	ФФОО	CRITICAL
Serious a	dverse even	ts (follow-	up: range 14 days	s to 30 days)								
5 1,3-5,7	randomized trials	not serious <sup>a</sup>	not serious	not serious	serious <sup>c</sup>	none	36/1671 (2.2%)	26/1727 (1.5%)	<b>RR 1.14</b> (0.32 to 3.99)	2 more per 1,000 (from 10 fewer to 45 more)	⊕⊕⊕⊖ MODERATE	CRITICAL

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that has not been peer reviewed or published.

CI: confidence interval; RR: risk ratio

#### **Explanations**

- a. Agusti 2022, Duvignaud 2022, Ramakrishnan 2021, Yu 2021 were open-label trials, which may introduce bias into outcomes subjectively measured, such as COVID-19-related hospitalizations and SAEs.
- b. 8/35 patients in Song 2021 received HCQ in addition to ciclesonide. All patients in Song 2021 had mild-to-moderate COVID-19 and were hospitalized.
- c. Sparse data, few events, unable to excluded harms as well as benefits

**Tables and Figures** 

d. In Yu 2021 the following patients were admitted to hospital without need for supplemental oxygen: budesonide 17/787 (2%) placebo 21/799 (3%).

- 1. Yu LM, Bafadhel M, Dorward J, et al. Inhaled budesonide for COVID-19 in people at high risk of complications in the community in the UK (PRINCIPLE): a randomised, controlled, open-label, adaptive platform trial. Lancet **2021**; 398(10303): 843-55.
- 2. Clemency BM, Varughese R, Gonzalez-Rojas Y, et al. Efficacy of Inhaled Ciclesonide for Outpatient Treatment of Adolescents and Adults With Symptomatic COVID-19: A Randomized Clinical Trial. JAMA Intern Med 2022; 182(1): 42-9.
- 3. Ezer N, Belga S, Daneman N, et al. Inhaled and intranasal ciclesonide for the treatment of covid-19 in adult outpatients: CONTAIN phase II randomised controlled trial. BMJ **2021**; 375: e068060.
- 4. Song JY, Yoon JG, Seo YB, et al. Ciclesonide Inhaler Treatment for Mild-to-Moderate COVID-19: A Randomized, Open-Label, Phase 2 Trial. J Clin Med 2021; 10(16): 3545.
- 5. Accelerating Covid-19 Therapeutic I, Vaccines -6 Study G, Naggie S. Inhaled Fluticasone for Outpatient Treatment of Covid-19: A Decentralized, Placebo-controlled, Randomized, Platform Clinical Trial. medRxiv 2022.
- 6. Agusti A, De Stefano G, Levi A, et al. Add-on inhaled budesonide in the treatment of hospitalised patients with COVID-19: a randomised clinical trial. Eur Respir J 2022; 59(3).
- 7. Duvignaud A, Lhomme E, Onaisi R, et al. Inhaled ciclesonide for outpatient treatment of COVID-19 in adults at risk of adverse outcomes: a randomised controlled trial (COVERAGE). Clin Microbiol Infect 2022; 28(7): 1010-6.
- 8. Ramakrishnan S, Nicolau DV, Jr., Langford B, et al. Inhaled budesonide in the treatment of early COVID-19 (STOIC): a phase 2, open-label, randomised controlled trial. Lancet Respir Med **2021**; 9(7): 763-72.

## Interleukin-6 inhibitors

## **Evidence profiles**

- Tocilizumab compared to no tocilizumab for hospitalized patients with COVID-19
- Sarilumab compared to no sarilumab for hospitalized patients with COVID-19

**Tables and Figures** 

**Table 11.** GRADE evidence profile, Recommendation 11

Question: Tocilizumab compared to no tocilizumab for hospitalized patients with COVID-19

Last updated 2/17/2021; last reviewed 9/14/2021

			Certainty as	sessment		№ of patients		Ef	fect			
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	tocilizumab	no tocilizumab	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow-up: ra	ange 28 days	s to 30 days)									
8 1-8	randomized trials	not serious a	not serious	not serious	serious <sup>b</sup>	none	810/3280 (24.7%)	893/3054 (29.2%)	<b>RR 0.91</b> (0.79 to 1.04)	26 fewer per 1,000 (from 61 fewer to 12 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Clinical d	eterioration (	follow-up: ra	ange 14 days to 3	30 days)								
7 1-6,8	randomized trials	serious <sup>c</sup>	not serious	not serious d	not serious	none	799/2712 (29.5%)	939/2503 (37.5%)	<b>RR 0.83</b> (0.77 to 0.89)	64 fewer per 1,000 (from 86 fewer to 41 fewer)	⊕⊕⊕⊜ MODERATE	CRITICAL
Serious a	dverse event	s							1			
<b>7</b> 1-7,e	randomized trials	serious <sup>c</sup>	not serious	not serious	serious <sup>f</sup>	none	210/1249 (16.8%)	141/946 (14.9%)	<b>RR 0.89</b> (0.74 to 1.07)	16 fewer per 1,000 (from 39 fewer to 10 more)	ФФОО LOW	CRITICAL
High certal Moderate of Low certal Very low con Risk of bia Inconsiste Indirectnes Imprecisio	certainty: We a nty: Our confide ertainty: We ha s: Study limitati ncy: Unexplain ss: Applicability	ry confident the re moderately ence in the effective very little commons ed heterogene or generalization in the estimate.	at the true effect lies confident in the effe ect estimate is limite onfidence in the effe ity across study find polity to the research mate of an effect to s	ct estimate: The tr d: The true effect ct estimate: The tr ings question	ue effect is likely may be substantia rue effect is likely	effect to be close to the estim. ally different from the es to be substantially diffe	timate of the effe	ct	ssibility that it is s	ubstantially differe	nt	

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

#### **Explanations**

- a. Although some studies did not blind participants or investigators, this is unlikely to affect the mortality outcome.
- b. 95% CI includes benefits as well as harms.
- c. Some studies lacked blinding and due to the mechanism of tocilizumab (reduction in inflammatory marker), unblinding likely occurred in the blinded studies.

**Tables and Figures** 

- d. Definition of clinical deterioration varied, with all studies including need for ventilation and death, but other studies included need for ICU admission (2 studies) or PaO<sub>2</sub>/FiO<sub>2</sub> ratio of less than 150 mmHg (1 study).
- e. The 95% CI includes both potential for harm as well as benefit; Few events reported do not meet the optimal information size and suggest fragility in the estimate.

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**Tables and Figures** 

**Table 12.** GRADE evidence profile, Recommendation 12

Question: Sarilumab compared to no sarilumab for hospitalized patients with COVID-19

New evidence profile developed 9/14/2021

Certainty assessment								№ of patients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	sarilumab	no sarilumab	Relative (95% CI)	Absolute (95% CI)		Importance
Mortality	(assessed w	ith: indirect	estimate from no	etwork meta-an	alysis)							
18 <sup>1,a</sup>	randomized trials	not serious	not serious	not serious	very serious <sup>b</sup>	none	Network estimate: <b>OR: 0.80</b> ; 95%: CI: 0.61, 1.04 Direct estimate: <b>OR: 0.98</b> ; 95% CI: 0.62, 1.56 Indirect estimate: <b>OR: 0.72</b> ; 95% CI: 0.52, 0.99			ФФОО LOW	CRITICAL	
Clinical d	eterioration	(follow-up: 2	21 days; assesse	d with: progres	sion to intuba	tion, ECMO, or deat	h)					
2 <sup>2,3</sup>	randomized trials	serious <sup>c</sup>	not serious <sup>d</sup>	not serious <sup>e</sup>	very serious <sup>f</sup>	none	72/305 (23.6%)	157/341 (46.0%) <sup>9</sup>	<b>RR 0.67</b> (0.42 to 1.05)	<b>152 fewer per 1,000</b> (from 267 fewer to 23 more)	⊕⊖⊖⊖ VERY LOW	CRITICAL
Serious a	dverse even	ts (follow-up	o: 21 days)									
4 2-4	randomized trials	serious <sup>c</sup>	not serious	not serious	serious <sup>h</sup>	none	566/1520 (37.2%)	158/795 (19.9%)	RR 1.03 (0.89 to 1.18)	6 more per 1,000 (from 22 fewer to 36 more)	ФФОО LOW	CRITICAL
High certa Moderate Low certai Very low c Risk of bia Inconsiste Indirectne Imprecisio	certainty: We a inty: Our confic ertainty: We h is: Study limital ncy: Unexplair ss: Applicability	ery confident the remoderately lence in the eff ave very little of tions ned heterogener or generalizance in the estir	nat the true effect lie confident in the effect estimate is limite confidence in the effect estimate is sometimate and the effect estimate of an effect to some the effect to some effect	ect estimate: The ted: The true effect ect estimate: The fact estimate: The fact estimate ings	rue effect is likely may be substanti true effect is likely	effect to be close to the estim ally different from the es to be substantially diffe	stimate of the effe	ect	esibility that it is s	ubstantially dil	fferent	

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; OR: Odds ratio; RR: Risk ratio

#### **Explanations**

- a. 18 trials included in the network.
- b. The direct network estimate crosses the line of no effect; however, the indirect estimate in the network demonstrates a trend toward mortality reduction when sarilumab + corticosteroids rather than corticosteroids alone is given. Few events reported in the direct network estimate suggesting fragility.
- c. Lack of blinding of study personnel, participants, and outcome assessors.

**Tables and Figures** 

- d. Substantial heterogeneity present (I<sup>2</sup>=57%); however, likely contributes to the wide CI and accounted for within imprecision.
- e. Definition of clinical deterioration varied, with all studies including need for ventilation; however, one study included ECMO and death and the other study included use of high-flow cannula.
- f. 95% CI cannot exclude the possibility of harm. Few events suggest fragility of the estimate.
- g. Analysis includes participants free of invasive mechanical ventilation at baseline for Gordon and patients free of high-flow cannula at baseline.
- h. 95% CI cannot exclude the possibility of harms.

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# **Convalescent plasma**

## **Evidence profiles**

- Convalescent plasma compared to no convalescent plasma for hospitalized patients with COVID-
- Convalescent plasma compared to no convalescent plasma for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease

Tables and Figures

 Table 13. GRADE evidence profile, Recommendation 13

Question: Convalescent plasma compared to no convalescent plasma for hospitalized patients with COVID-19

Last reviewed and updated 11/4/2021

			Certainty asse	ssment	Nº of p	atients		Effect				
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	convalescent plasma	no convalescent plasma	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(RCTs) (follow-	up: range 1	5 days to 60 days)									
18 <sup>1-18</sup>	randomized trials	not serious a,b	not serious	not serious	serious <sup>c</sup>	none	2163/9082 (23.8%)	2007/8150 (24.6%)	RR 0.98 (0.93 to 1.03)	5 fewer per 1,000 (from 17 fewer to 7 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Need for	mechanical ver	ntilation						•	•	•		
4 3,6,9,14	randomized trials	serious d	not serious	not serious	serious <sup>e</sup>	none	184/581 (31.7%)	166/471 (35.2%)	RR 1.10 (0.94 to 1.29)	35 more per 1,000 (from 21 fewer to 102 more)	ФФСС	CRITICAL
Serious a	dverse events	(transfusion	-associated circul	latory overload,	transfusion-re	elated acute lung i	njury, severe alle	ergic transfusion	reaction) (	follow-up: 4 ho	urs)	
1 <sup>19</sup>	observational studies	extremely serious <sup>f</sup>	not serious	not serious	not serious	none	the SAEs, 63 dea and 13 of those of related to the train There were 83 not transfusion-assore reports of transfusion	O transfused patie aths were reported deaths were judge insfusion of COVID on-death SAEs repetiated circulatory cision-related acute ere allergic transfu	⊕⊖⊖ VERY LOW	CRITICAL		
Serious a	dverse events	(mortality, c	ardiac, thrombotic	c, sustained hyp	otensive even	ts requiring interv	rention) (follow-u	ıp: 7 days)				
1 <sup>19</sup>	observational studies	extremely serious <sup>f</sup>	not serious	not serious	not serious	none	transfusion, 1711 events (5.68%) w 643 cardiac even transfusion); 406 intravenous pres	10 transfused patie I deaths (8.56%) a vere reported. Non its (569 judged as sustained hypote sor support; and 8 s (55 judged as ur	⊕⊖⊖⊖ VERY LOW	CRITICAL		
Any adve	rse events (RC	Ts)										
11 3,4,6,8,11- 13,15-18	randomized trials	serious d	not serious	not serious <sup>9</sup>	serious <sup>h</sup>	none	574/2843 (20.2%)	307/1959 (15.7%)	RR 1.08 (0.94 to 1.26)	13 more per 1,000 (from 9 fewer to 41 more)	ФФСС	IMPORTANT

**Tables and Figures** 

#### **GRADE Working Group grades of evidence**

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio; HR: Hazard Ratio; OR: Odds ratio; SAEs: Serious adverse events

#### **Explanations**

- a. Li 2020 time between symptom onset and randomization was over 14 days for >90% (median 30 days), no adjustment for co-interventions, allocation concealment methods not reported and participants and healthcare professionals not blinded.
- b. Many trials had concerns due to open-label trial, allocation concealment not reported, and no adjustments for co-interventions. Differences between study protocol and published report (e.g., inclusion criteria, outcomes, intervention groups) noted for Pouladzadeh 2021.
- c. The 95% CI includes the potential for appreciable benefit; however, cannot exclude the potential for no effect.
- d. Concerns include open-label trial design and assessment of outcome.
- e. The 95% CI may not include a clinically meaningful reduction in need for mechanical ventilation.
- f. No comparative effects available. Some subjectivity in classification of outcomes as transfusion related.
- g. Lack standard definition for adverse events. Studies report on mild to severe events.
- h. The 95% CI includes the potential for both increased harms, as well as no increased harms.

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- 3. AlQahtani M, Abdulrahman A, AlMadani A, et al. Randomized controlled trial of convalescent plasma therapy against standard therapy in patients with severe COVID-19 disease. Sci Rep **2021**; 11: 9927.
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**Tables and Figures** 

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**Tables and Figures** 

**Table 14.** GRADE evidence profile, Recommendation 14

**Question:** Convalescent plasma compared to no convalescent plasma for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease **Last reviewed and updated 1/21/2022** 

			Certainty as	sessment			Nº of p	patients	Eff	fect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	convalescent plasma	no convalescent plasma	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
All-cause	mortality (fo	llow-up: ra	ange 15 days to 2	8 days) <sup>a</sup>								
3 <sup>1-3</sup>	randomized trials	not serious	not serious	not serious	very serious <sup>b</sup>	none	3/929 (0.3%)	7/923 (0.8%)	<b>RR 0.53</b> (0.14 to 1.98)	4 fewer per 1,000 (from 7 fewer to 7 more)	ФФОО	CRITICAL
COVID-19	related hos	pitalization	s, ED/urgent care	e visits, or deat	h (follow-up: 1	5 days)						
2 <sup>1,3</sup>	randomized trials	not serious	not serious	not serious	serious <sup>c</sup>	none	94/849 (11.1%)	118/843 (14.0%)	<b>RR 0.79</b> (0.62 to 1.00)	29 fewer per 1,000 (from 53 fewer to 0 fewer)	⊕⊕⊕○ MODERATE	CRITICAL
Hospitali	zations (all-c	ause) (follo	ow-up: range 15 c	lays to 28 days	)							
2 1,3	randomized trials	not serious	not serious	not serious	serious <sup>d</sup>	none	73/867 (8.4%)	98/869 (11.3%)	<b>RR 0.74</b> (0.56 to 0.98)	29 fewer per 1,000 (from 50 fewer to 2 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
Progress	ion to severe	respirator	y disease (follow	-up: 15 days; a	ssessed with:	defined as a resp	iratory rate of ≥	30 breaths per m	inute, SaO₂ < 93	% on room air, or	both)	
12	randomized trials	not serious <sup>e</sup>	not serious	serious <sup>f</sup>	serious <sup>g</sup>	none	13/80 (16.3%)	25/80 (31.3%)	<b>RR 0.52</b> (0.29 to 0.94)	150 fewer per 1,000 (from 222 fewer to 19 fewer)	ФФОО	CRITICAL
Serious a	dverse even	ts: serious	transfusion read	tions (requiring	treatment or	admission) (follow	v-up: 15 days)					
2 1,3	randomized trials	not serious	not serious	not serious	very serious <sup>c</sup>	none	5/849 (0.6%)	0/843 (0.0%)	<b>RR 5.95</b> (0.72 to 49.29) h	6 more per 1,000 (from 1 more to 11 more)	ФФОО	CRITICAL
Any adve	erse events (f	ollow-up: 1	I5 days)									
2 1,3	randomized trials	not serious	not serious	not serious	serious <sup>c</sup>	none	127/849 (15.0%)	147/843 (17.4%)	<b>RR 0.86</b> (0.70 to 1.05)	24 fewer per 1,000 (from 52 fewer to 9 more)	⊕⊕⊕⊖ MODERATE	IMPORTANT

**Tables and Figures** 

#### **GRADE Working Group grades of evidence**

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; ED: Emergency department; RR: Risk ratio; SaO2: Saturated oxygen

## **Explanations**

- a. Deaths beyond 15 days and up to 30 days: an additional 5 deaths occurred in the plasma group and 1 death in placebo (normal saline) group.
- b. Only one event.
- c. 95% CI includes benefits as well as harms; OIS not met.
- d. Few events reported. 95% CI may not include clinically meaningful benefit.
- e. Trial was terminated early due to futility.
- f. Oxygenation and respiration rates are surrogate measures of need for ventilation, morbidity and death.
- g. Few events reported do not meet the optimal information size and suggest fragility of the estimate.
- h. Using 0.5 event continuity correction.
- i. Zero events in the control group. Absolute risk difference not informed by relative risk

#### References

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# Remdesivir

## **Evidence profiles**

- Remdesivir compared to no remdesivir for ambulatory patients at high risk for severe COVID-19
- Remdesivir 5 days compared to remdesivir 10 days for hospitalized patients with severe but not critical COVID-19
- Remdesivir compared to no antiviral treatment for hospitalized patients with severe COVID-19
- Remdesivir compared to no antiviral treatment for hospitalized patients with critical COVID-19 (IV/ECMO)

**Tables and Figures** 

Table 15. GRADE evidence profile, Recommendation 15

Question: Remdesivir compared to no remdesivir for ambulatory patients at high risk for severe COVID-19

Last updated 12/23/2021; last reviewed 2/7/2022

			Certainty ass	sessment			Nº of p	atients	E	ffect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	remdesivir	no remdesivir	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow-up:	28 days)					•					
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	very serious a	none	0/279 (0.0%)	0/283 (0.0%)	not estimable		⊕⊕⊖⊖	CRITICAL
Hospitali	zation (all-ca	ause) (follow	-up: 28 days)	•	<u> </u>		•		•		1	
11	randomised trials	not serious	not serious	not serious	very serious b	none	5/279 (1.8%)	18/283 (6.4%)	HR 0.28 (0.10 to 0.75)	45 fewer per 1,000 (from 57 fewer to 16 fewer)	ФФОО	CRITICAL
COVID-19	9-related me	dically atten	ded visits (follov	v-up: 28 days)	l		•		l		<u> </u>	
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	very serious <sup>b</sup>	none	4/246 (1.6%)	21/252 (8.3%)	HR 0.19 (0.07 to 0.56)	67 fewer per 1,000 (from 77 fewer to 36 fewer)	ФФСО	IMPORTANT
Serious a	dverse ever	nts		l	l		•		l		<u> </u>	
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	serious <sup>b</sup>	none	5/279 (1.8%)	19/283 (6.7%)	<b>RR 0.27</b> (0.10 to 0.70)	49 fewer per 1,000 (from 60 fewer to 20 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
High certa Moderate Low certai	certainty: We inty: Our confi	ery confident the are moderately dence in the eff	nat the true effect lie confident in the eff fect estimate is limit	ect estimate: The ed: The true effect	true effect is likel t may be substan		ne estimate of th	e effect		at it is substantially di	fferent	
nconsiste ndirectne mprecisio	ss: Applicabilit	ned heterogene y or generaliza ence in the estil	eity across study fin bility to the research mate of an effect to of studies	n question	ar decision	· · · · · ·						

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; HR: Hazard ratio; RR: Risk ratio

**Explanations** 

**Tables and Figures** 

- a. Zero events and relatively small sample size (less than half the patients of the planned sample size were enrolled).
- b. Few events do not meet the optimal information size and suggest fragility in the estimate (less than half the patients of the planned sample size were enrolled).

#### Reference

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**Tables and Figures** 

Table 16. GRADE evidence profile, Recommendation 16

Question: Remdesivir 5 days compared to remdesivir 10 days for hospitalized patients with severe but not critical COVID-19

Last updated 9/10/2020; last reviewed 5/16/2021

			Certainty ass	essment			Nº of pa	atients	E	ffect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	remdesivir 5 days	remdesivir 10 days	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
/lortality												
1 <sup>1</sup>	randomized trials	serious <sup>b</sup>	not serious	not serious	serious <sup>a</sup>	none	16/200 (8.0%)	21/197 (10.7%)	HR 0.75 (0.40 to 1.39)	27 fewer per 1,000 (from 64 fewer to 42 more)	ФФОО Low	CRITICAL
linical i	mprovement	at 14 days										
<b>1</b> ¹	randomized trials	serious <sup>b</sup>	not serious	not serious	serious <sup>c</sup>	none	129/200 (64.5%)	107/197 (54.3%)	RR 1.19 (1.01 to 1.40)	103 more per 1,000 (from 5 more to 217 more)	ФФОО	CRITICAL
Serious a	adverse even	ts								•	•	
1 <sup>1</sup>	randomized trials	serious <sup>b</sup>	not serious	not serious	serious <sup>c</sup>	none	42/200 (21.0%)	68/197 (34.5%)	RR 0.61 (0.44 to 0.85)	135 fewer per 1,000 (from 193 fewer to 52 fewer)	ФФОО LOW	CRITICAL
Adverse	events leadii	ng to treatm	ent discontinuati	on								
1 <sup>1</sup>	randomized trials	serious b,d	not serious	not serious	serious <sup>c</sup>	none	9/200 (4.5%)	20/197 (10.2%)	RR 0.44 (0.21 to 0.95)	57 fewer per 1,000 (from 80 fewer to 5 fewer)	ФФОО Low	CRITICAL
	orking Group		dence hat the true effect lie	a alaaa ta that of t	ha antimata of th	o officet						
loderate ow certa	certainty: We a	are moderately dence in the ef	y confident in the effe fect estimate is limite	ect estimate: The ed: The true effect	true effect is likely may be substant	e effect  to be close to the estimate  itally different from the y to be substantially o	e estimate of the et	ffect	ossibility that it	is substantially diff	erent	
nconsiste ndirectne nprecisio	ss: Applicabilit	ned heterogen y or generaliza nce in the esti	eity across study find ability to the research mate of an effect to s	question	ar decision							

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

**Tables and Figures** 

CI: Confidence interval; RR: Risk ratio

## **Explanations**

- a. The 95% CI includes the potential for both appreciable benefit, as well as appreciable harm. Few events reported do not meet the optimal information size and suggest fragility in the estimate.
- b. Goldman 2020 did not blind participants, healthcare workers or outcome assessors. After randomization, disease severity was greater in the 10-day arm; while the analysis adjusted for baseline characteristics including disease severity, there is still the potential for residual confounding.
- c. The lower boundary of the 95% CI may not include a clinically meaningful effect. Few events reported do not meet the optimal information size and suggest fragility in the estimate
- d. Goldman stratified adverse events by days 1-5, 6-10. AEs leading to treatment discontinuation during days 1-5 were 9 (4%) in the 5-day arm and 14 (7%) in the 10-day arm.

#### Reference

1. Goldman JD, Lye DCB, Hui DS, et al. Remdesivir for 5 or 10 Days in Patients with Severe Covid-19. N Engl J Med 2020; 383: 1827-37.

**Tables and Figures** 

Table 17a. GRADE evidence profile, Recommendation 17a

Question: Remdesivir compared to no antiviral treatment for hospitalized patients with severe COVID-19

Last reviewed and updated 5/16/2021

			Certainty ass	sessment			№ of p	atients	E	ffect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	remdesivir	no remdesivir	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow-up:	range 28 da	ys to 29 days)									
3 1-3	randomized trials	serious a,b,c	not serious	not serious	serious <sup>d</sup>	none	369/2726 (13.5%)	374/2593 (14.4%)	<b>RR 0.92</b> (0.77 to 1.10)	12 fewer per 1,000 (from 33 fewer to 14 more)	ФФОО LOW	CRITICAL
Time to I	recovery (fol	low-up: 29 d	lays)									
12	randomized trials	serious <sup>c</sup>	not serious	not serious	not serious	none	345/486 (71.0%)	306/471 (65.0%)	Rate ratio 1.31 (1.12 to 1.52)	97 more per 1,000 (from 41 more to 147 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Clinical i	mprovement	t (follow-up:	28 days)									
11	randomized trials	not serious a,b	not serious	not serious	very serious <sup>d</sup>	none	103/158 (65.2%)	45/78 (57.7%)	<b>RR 1.13</b> (0.91 to 1.41)	75 more per 1,000 (from 52 fewer to 237 more)	ФФСС	CRITICAL
Need for	mechanical	ventilation (	(follow-up: 29 da	ıys)	•		•	•	•			
12	randomized trials	not serious	not serious	not serious	serious <sup>e</sup>	none	52/402 (12.9%)	82/364 (22.5%)	<b>RR 0.57</b> (0.42 to 0.79)	<b>97 fewer per 1,000</b> (from 131 fewer to 47 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
Serious	adverse ever	nts (grade 3/	(4)									
2 1,2	randomized trials	not serious	not serious	not serious	serious <sup>f</sup>	none	44/632 (7.0%)	53/545 (8.9%)	<b>RR 0.79</b> (0.54 to 1.16)	20 fewer per 1,000 (from 45 fewer to 16 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Hospitali	ization											
11	randomized trials	not serious a,b	not serious	not serious	very serious <sup>d</sup>	none	158	78	-	MD 1 days higher (0.12 higher to 1.88 higher)	ФФОО LOW	IMPORTANT

**Tables and Figures** 

			Certainty ass	sessment			Nº of pa	atients	E	fect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	remdesivir	no remdesivir	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Duration	of mechanic	al ventilatio	n									
1 1	randomized trials	not serious	not serious	not serious	serious <sup>d</sup>	none	158	78	-	MD <b>8.5 days</b> lower (9.14 lower to 7.86 lower)	⊕⊕⊕⊖ MODERATE	IMPORTANT

#### **GRADE Working Group grades of evidence**

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

**Imprecision:** The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; HR: Hazard Ratio; RR: Risk ratio; OR: Odds ratio; MD: Mean difference

#### **Explanations**

- a. Co-interventions received in Wang 2020 include: interferon alpha-2b, lopinavir/ritonavir, vasopressors, antibiotics, corticosteroid therapy and were balanced between arms.
- b. Wang 2020 stopped early due to lack of recruitment. Trial initiated after reduction in new patient presentation (most patients enrolled later in the disease).
- c. Post hoc analysis of patients with severe disease from Pan 2020 and Beigel 2020 may introduce bias.
- d. The 95% CI may not include a clinically meaningful effect.
- e. Few events do not meet the optimal information size and suggest fragility in the estimate.
- f. The 95% CI cannot exclude the potential for benefit or harm. Also, few events do not meet the optimal information size.

#### References

- 1. Wang Y, Zhang D, Du G, et al. Remdesivir in adults with severe COVID-19: a randomised, double-blind, placebo-controlled, multicentre trial. Lancet **2020**; 395(10236): 1569-78.
- 2. Beigel JH, Tomashek KM, Dodd LE, et al. Remdesivir for the Treatment of Covid-19 Final Report. N Engl J Med 2020; 383(19): 1813-26.
- 3. WHO Solidarity Trial Consortium, Pan H, Peto R, et al. Repurposed Antiviral Drugs for Covid-19 Interim WHO Solidarity Trial Results. N Engl J Med 2021; 384: 497-511.

**Tables and Figures** 

Table 17b. GRADE evidence profile, Recommendation 17b

Question: Remdesivir compared to no antiviral treatment for hospitalized patients with critical COVID-19 (IV/ECMO)

Last updated 4/5/2021; last reviewed 5/16/2021

			Certainty as	sessment			Nº of p	atients	Effe	ct		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	remdesivir	no remdesivir	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow-up:	range 28 day	ys to 29 days)									
2 1,2	randomized trials	serious <sup>a</sup>	not serious	not serious	serious <sup>b,c</sup>	none	126/385 (32.7%)	100/387 (25.8%)	RR 1.23 (0.99 to 1.53)	59 more per 1,000 (from 3 fewer to 137 more)	ФФОО LOW	CRITICAL
Time to r	ecovery (fol	low-up: 29 d	lays)									
11	randomized trials	very serious <sup>a</sup>	not serious	not serious	very serious <sup>d</sup>	none	63/131 (48.1%)	77/154 (50.0%)	HR 0.98 (0.70 to 1.36)	7 fewer per 1,000 (from 116 fewer to 110 more)	⊕⊖⊖⊖ VERY LOW	CRITICAL
Serious a	adverse ever	nts (grade 3/	(4)	•			•		•			
2 1,3	randomized trials	not serious	not serious	not serious e	serious <sup>d</sup>	none	44/632 (7.0%)	53/545 (9.7%)	<b>RR 0.79</b> (0.54 to 1.16)	20 fewer per 1,000 (from 45 fewer to 16 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
High certa Moderate Low certa Very low of Risk of bia Inconsiste Indirectne	certainty: We inty: Our confi- certainty: We has: Study limita- ency: Unexplai ss: Applicabilit	ery confident to are moderated dence in the enave very little stions ned heterogen y or generaliza	that the true effect li y confident in the effect estimate is limi	fect estimate: The ted: The true effect ffect estimate: The ndings th question	e true effect is like ct may be substar e true effect is like	ne effect ly to be close to the es ntially different from the ely to be substantially d	estimate of the	effect		is substantial	ly different	

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio; HR: Hazard Ratio

#### **Explanations**

- a. Post hoc analysis of patients with severe disease from Pan 2020 and Beigel 2020 may introduce bias.
- b. The 95% CI may not include a clinically meaningful effect.
- c. OIS for mortality: 1682

Publication bias: Selective publication of studies

d. The 95% CI cannot exclude the potential for benefit or harm. Also, few events do not meet the optimal information size.

**Tables and Figures** 

e. Serious adverse events calculated from severe study groups in Beigel 2020 & Wang 2020, not invasive mechanical ventilation/ECMO subgroup.

#### References

- 1. Beigel JH, Tomashek KM, Dodd LE, et al. Remdesivir for the Treatment of Covid-19 Final Report. N Engl J Med 2020; 383(19): 1813-26.
- 2. WHO Solidarity Trial Consortium, Pan H, Peto R, et al. Repurposed Antiviral Drugs for Covid-19 Interim WHO Solidarity Trial Results. N Engl J Med 2021; 384: 497-511.
- 3. Wang Y, Zhang D, Du G, et al. Remdesivir in adults with severe COVID-19: a randomised, double-blind, placebo-controlled, multicentre trial. Lancet **2020**; 395(10236): 1569-78.

**Tables and Figures** 

# **Famotidine**

## **Evidence profiles**

• Famotidine compared to no famotidine for ambulatory patients with mild-to-moderate COVID-19

**Tables and Figures** 

Table 18. GRADE evidence profile, Recommendation 18

Question: Famotidine compared to no famotidine for ambulatory patients with mild-to-moderate COVID-19

New evidence profile developed 5/17/2022

			Certainty ass	essment			Nº of pa	atients	E	ffect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	high-dose famotidine (80 mg tid)	no famotidine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Sympton	n resolution (fo	llow-up: 28	B days) <sup>a</sup>									
11	randomized trials	not serious	not serious	not serious	very serious	none	19/27 (70.4%) °	18/28 (64.3%)	<b>RR 1.10</b> (0.76 to 1.58)	64 more per 1,000 (from 154 fewer to 373 more)	ФФОО LOW	CRITICAL
Adverse	events d											
11	randomized trials	not serious	not serious	not serious	very serious	none	2/27 (7.4%)	3/28 (10.7%)	<b>RR 0.69</b> (0.13 to 3.80)	33 fewer per 1,000 (from 93 fewer to 300 more)	ФФОО LOW	IMPORTANT

#### **GRADE Working Group grades of evidence**

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

## **Explanations**

- a. Time to symptom resolution was the primary end point. However, the authors reported a faster (earlier) rate of symptom resolution with famotidine. No deaths were encountered.
- b. Sparse data, few events and small sample size
- c. Only p-value reported; number of events estimated from survival curve graph.
- d. No serious adverse events were encountered. Transaminase elevation in 1 patient in both arms; nausea / vomiting in 1 patient with famotidine; thrombocytopenia and hives in 1 patient each in the placebo group.

#### Reference

1. Brennan CM, Nadella S, Zhao X, et al. Oral famotidine versus placebo in non-hospitalised patients with COVID-19: a randomised, double-blind, data-intense, phase 2 clinical trial. Gut **2022**; 71(5): 879-88.

**Tables and Figures** 

Table 19. GRADE evidence profile, Recommendation 19

Question: Famotidine compared to no famotidine for hospitalized patients with severe COVID-19

Last reviewed and updated 5/17/2022

			Certainty asso	essment			Nº of p	oatients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	famotidine	no famotidine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	1											
11	randomized trials	serious a	not serious	not serious	serious <sup>b</sup>	none	8/89 (9.0%)	9/89 (10.1%)	RR 0.89 (0.36 to 2.20)	11 fewer per 1,000 (from 65 fewer to 121 more)	ФФОО	CRITICAL
Mechani	cal ventilation											
11	randomized trials	serious a	not serious	not serious	serious <sup>b</sup>	none	21/89 (23.6%)	24/89 (27.0%)	RR 0.88 (0.53 to 1.45)	32 fewer per 1,000 (from 127 fewer to 121 more)	ФФОО	CRITICAL
ICU care												
11	randomized trials	serious a	not serious	not serious	serious <sup>b</sup>	none	18/89 (20.2%)	20/89 (22.5%)	<b>RR 0.90</b> (0.51 to 1.58)	22 fewer per 1,000 (from 110 fewer to 130 more)	ФФОО	CRITICAL
Time to	symptom-free						•					
11	randomized trials	serious a	not serious	not serious	serious <sup>b</sup>	none	89	89	-	MD 0.9 days fewer (1.44 fewer to 0.36 fewer)	ФФОО	IMPORTANT
Length o	of hospital stay					•	•	•		•	•	
11	randomized trials	serious a	not serious	not serious	serious <sup>b</sup>	none	89	89	-	MD 1.7 days fewer (2.77 fewer to 1.13 fewer)	ФФСС	IMPORTANT

Serious adverse events

**Tables and Figures** 

			Certainty asso	essment			<b>№</b> of p	atients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	famotidine	no famotidine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
0	observational studies						adverse eve diarrhea (1.7 4.7%), but o serious adve Johnson syr necrotizing e rhabdomyoly	ents include co 7%), dizziness overall famotidi erse events (< ndrome, toxic o enterocolitis, a	enstipation (1 s (1.3%) and ine is well to 1%) include: epidermal ne anaphylaxis, hospital-acqu	headache (1%- lerated. Rare but Stevens- ecrolysis,	-	CRITICAL

## **GRADE Working Group grades of evidence**

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

**Imprecision:** The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; MD: Mean difference; RR: Risk ratio

## **Explanations**

- a. Unclear allocation concealment in an unblinded study
- b. Sparse data, small number of events or patients

#### Reference

1. Pahwani S, Kumar M, Aperna F, et al. Efficacy of Oral Famotidine in Patients Hospitalized With Severe Acute Respiratory Syndrome Coronavirus 2. Cureus 2022; 14(2): e22404

Tables and Figures

# Janus kinase inhibitors

## **Evidence profiles**

- Baricitinib compared to no baricitinib for hospitalized patients receiving standard of care for severe COVID-19
- Baricitinib compared to no baricitinib for critically ill (OS-7) patients with COVID-19 pneumonia requiring invasive mechanical ventilation
- Baricitinib with remdesivir compared to remdesivir for hospitalized patients with COVID-19
- Tofacitinib compared to no tofacitinib for hospitalized patients with COVID-19

**Tables and Figures** 

Table 20. GRADE evidence profile, Recommendation 20

Question: Baricitinib compared to no baricitinib for hospitalized patients receiving standard of care for severe COVID-19

Last reviewed and updated 4/29/2022

			Certainty as	ssessment			№ of p	atients	Effec	ot		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	baricitinib	no baricitinib	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow-up: ı	range 28 day	s to 60 days)									
21,2	randomized trials	not serious	not serious	not serious	serious <sup>a</sup>	none	592/4912 (12.1%)	662/4769 (13.9%)	RR 0.87 (0.78 to 0.96)	18 fewer per 1,000 (from 31 fewer to 6 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
Mechanic	cal ventilatio	n (follow-up	: 28 days)									
12	randomized trials	not serious	not serious	not serious	serious <sup>a</sup>	none	283/4014 (7.1%)	322/3891 (8.3%)	<b>RR 0.85</b> (0.73 to 0.99)	<b>12 fewer per 1,000</b> (from 22	⊕⊕⊕⊖ MODERATE	CRITICAL
										fewer to 1 more)		
Disease p	progression	(follow-up: 2	28 days; assesse	ed with: progre	ssion to high-f	low oxygen, non-inv	vasive ventilati	on oxygen, in	vasive mechan	more)	ion, or death)	
	progression randomized trials		28 days; assesse not serious	ed with: progre	ssion to high-f	low oxygen, non-inv none	212/764 (27.7%)	on oxygen, in 232/761 (30.5%)	OR 0.85 (0.67 to 1.08) <sup>b</sup>	more)	ion, or death)	IMPORTANT
13	randomized trials		not serious				212/764	232/761	<b>OR 0.85</b> (0.67 to	more) ical ventilat 33 fewer per 1,000 (from 78 fewer to	$\oplus \oplus \oplus \bigcirc$	IMPORTANT

Tables and Figures

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

**Imprecision:** The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; HR: Hazard Ratio; OR: Odds ratio; RR: Risk ratio

#### **Explanations**

- a. 95% CI cannot exclude no benefit.
- b. Multiple imputation includes N=756 for placebo and N=762 for baricitinib
- c. Number of events does not meet optimal information size
- d. 95% CI cannot exclude no harm.
- e. Non-comparative serious adverse events were reported in the RECOVERY 2022 trial (baricitinib N=4,148): 13 total (5 serious infections, 3 bowel perforations, 2 pulmonary embolisms, 1 each of ischemic colitis, elevated transaminases and seizure)

#### References

- 1. Marconi VC, Ramanan AV, de Bono S, et al. Efficacy and safety of baricitinib for the treatment of hospitalised adults with COVID-19 (COV-BARRIER): a randomised, double-blind, parallel-group, placebo-controlled phase 3 trial. Lancet Respir Med 2021; 9(12): 1407-18.
- 2. RECOVERY Collaborative Group, Horby PW, Emberson JR, et al. Baricitinib in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial and updated meta-analysis. medRxiv 2022: Available at: <a href="https://doi.org/10.1101/2022.03.02.22271623">https://doi.org/10.1101/2022.03.02.22271623</a> [Preprint 3 March 2022].
- 3. Marconi VC, Ramanan AV, de Bono S, et al. Baricitinib plus Standard of Care for Hospitalized Adults with COVID-19. medRxiv **2021**: Available at: <a href="https://doi.org/10.1101/2021.04.30.21255934">https://doi.org/10.1101/2021.04.30.21255934</a> [Preprint 3 May 2021].

**Tables and Figures** 

Table 21. GRADE evidence profile, Recommendation 20

Question: Baricitinib compared to no baricitinib for critically ill (OS-7) patients with COVID-19 pneumonia requiring invasive mechanical ventilation

Last reviewed and updated 4/29/2022

			Certainty as	ssessment			Nº of p	oatients	Ef	fect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	baricitinib	no baricitinib	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
ortality	(HR) (follow	-up: 60 da	ys)									
21,2	randomized trials	not serious	not serious	not serious	serious <sup>a</sup>	none	61/185 (33.0%)	75/167 (44.9%)	<b>RR 0.74</b> (0.57 to 0.97)	117 fewer per 1,000 (from 193 fewer to 13 fewer)	⊕⊕⊕ MODERATE	CRITICAL
nvasive	mechanical	ventilation	free days (follow	<i>ı</i> -up: 60 days)								
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	very serious a,b	none	51	50	-	MD 2.36 vent free days more (6.1 more to 1.4 fewer) °	ФФОО LOW	IMPORTANI
Days of h	nospitalizatio	n (follow-	up: 60 days)		•				•			
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	very serious	none	51	50	-	MD 2.3 days fewer (4.6 fewer to 0	ФФОО	CRITICAL
Serious a	adverse even	ts (follow	-up: 28 days)									
11	randomized trials	not serious	not serious	not serious	serious ª	none	25/50 (50.0%)	35/49 (71.4%)	<b>RR 0.70</b> (0.50 to 0.97)	214 fewer per 1,000 (from 357 fewer to 21 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
	Norking Group	-		18	f the confirmation of	l			ı	•		
Moderate Low cert	e certainty: We ainty: Our conf	are modera	e effect estimate is li	effect estimate: Th mited: The true effe	e true effect is like ect may be substa	rie effect ely to be close to the ntially different from t ely to be substantially	he estimate of t	he effect		t it is substantially o	lifferent	

Tables and Figures

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

**Imprecision:** The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; HR: Hazard Ratio; MD: Mean difference; RR: Risk ratio

#### **Explanations**

a. Few number of events, does not meet optimal information size

- b. Pooled mortality event data RR: 0.73 (95% CI: 0.50, 1.06) cannot exclude no meaningful benefit and therefore suggests fragility when compared with the HR.
- c. 95% CI includes both the possibility of benefit and risk of harm
- d. Adjusted for age (<65, ≥65) and region (U.S., rest of the world)
- e. 95% CI cannot exclude no benefit

#### Reference

- 1. Ely EW, Ramanan AV, Kartman CE, et al. Efficacy and safety of baricitinib plus standard of care for the treatment of critically ill hospitalised adults with COVID-19 on invasive mechanical ventilation or extracorporeal membrane oxygenation: an exploratory, randomised, placebo-controlled trial. Lancet Respir Med 2022; 10(4): 327-36.
- 2. RECOVERY Collaborative Group, Horby PW, Emberson JR, et al. Baricitinib in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial and updated meta-analysis. medRxiv 2022: Available at: <a href="https://doi.org/10.1101/2022.03.02.22271623">https://doi.org/10.1101/2022.03.02.22271623</a> [Preprint 3 March 2022].

**Tables and Figures** 

Table 22. GRADE evidence profile, Recommendation 21

Question: Baricitinib with remdesivir compared to remdesivir for hospitalized patients with COVID-19

Last updated 5/16/2021; last reviewed 10/11/2021

			Certainty ass	essment			Nº of p	atients	Effe	ect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	baricitinib + RDV	RDV	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow-up: 2	28 days)										
1 1	randomized trials	not serious	not serious	not serious	serious <sup>a</sup>	none	24/515 (4.7%)	37/518 (7.1%)	HR 0.65 (0.39 to 1.09)	24 fewer per 1,000 (from 43 fewer to 6 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Clinical r	ecovery - ho	spitalized re	quiring suppleme	ental O₂/receivi	ng noninvasiv	e ventilation or hi	gh-flow O <sub>2</sub> (o	rdinal 5+6) (a	assessed with: (	Ordinal scale <	4)	
1 <sup>1</sup>	randomized trials	serious <sup>b</sup>	not serious	not serious	serious <sup>c</sup>	none	344/391 (88.0%)	316/389 (81.2%)	RR 1.08 (1.02 to 1.15)	<b>65 more per</b> <b>1,000</b> (from 16 more to 122 more)	ФФОО	CRITICAL
Clinical r	ecovery - rec	eiving nonir	vasive ventilatio	on or high-flow	O <sub>2</sub> , invasive m	echanical ventilat	ion or ECMO	(ordinal 6+7	; stratified) (ass	essed with: Or	dinal scale <4)	
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	serious <sup>e</sup>	none	122/176 (69.3%)	114/191 (59.7%)	HR 1.29 (1.00 to 1.66) <sup>d</sup>	93 more per 1,000 (from 0 fewer to 182 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
New use	of mechanic	al ventilatior	or ECMO (follow	w-up: 29 days)								
1 1	randomized trials	serious <sup>f</sup>	not serious	not serious	serious <sup>g</sup>	none	46/461 (10.0%)	70/461 (15.2%)	<b>RR 0.66</b> (0.46 to 0.93)	52 fewer per 1,000 (from 82 fewer to 11 fewer)	ФФОО LOW	CRITICAL
Serious a	dverse even	ts (follow-up	o: 28 days)		-				•			
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	serious <sup>g</sup>	none	81/507 (16.0%)	107/509 (21.0%)	RR 0.76 (0.59 to 0.99) h	50 fewer per 1,000 (from 86 fewer to 2 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL

**Tables and Figures** 

#### **GRADE Working Group grades of evidence**

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio; HR: Hazard Ratio; OR: Odds ratio; RDV: Remdesivir

#### **Explanations**

- a. 95% CI includes substantial benefits as well as substantial harms
- b. Non-stratified subgroup post hoc analysis.
- c. Lower boundary of the 95% CI crosses our threshold for a meaningful difference.
- d. Data from table S6. Although described as "analysis as randomized" in this stratum of severe COVID-19 patients, the analysis included moving patient from a baseline of "moderate" to "severe" post hoc (19 in the baricitinib group vs 21 in the placebo group), thus altering the original stratification. However, re-analysis using to original strata data (ordinal scale 6 and 7 from table 2) and 28-day cutoff (as a binary, non-time to event analysis) produce a similar result (RR 1.2, 95% CI 1.005 to 1.43). Not rated down for post hoc analysis concerns.
- e. 95% CI includes substantial benefits as well as no effect
- f. Not a predefined stratum. Secondary analysis.
- g. Less than 300 events; concern for fragility
- h. SAEs in 5 or more participants in any preferred term by treatment group. 6/507 were thought related to study drug in the baricitinib group; 5/509 were thought to be related to the study drug in the placebo group.

#### Reference

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**Tables and Figures** 

Table 23. GRADE evidence profile, Recommendation 22

Question: To facitinib compared to no to facitinib for hospitalized patients with COVID-19

New evidence profile developed 8/21/2021

Certainty assessment								atients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	tofacitinib	no tofacitinib	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Death or	respiratory fa	ilure (follow-	up: 28 days)									
<b>1</b> ¹	randomized trials	not serious	not serious	not serious	very serious a,b	none	26/144 (18.1%)	42/145 (29.0%)	<b>RR 0.63</b> (0.41 to 0.97)	107 fewer per 1,000 (from 171 fewer to 9 fewer)	ФФОО LOW	CRITICAL
/lortality	(follow-up: 28	days)					l .					
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	very serious a,c	none	4/144 (2.8%)	8/145 (5.5%)	<b>RR 0.49</b> (0.15 to 1.63)	28 fewer per 1,000 (from 47 fewer to 35 more)	ФФОО LOW	CRITICAL
rogress	ion to mechai	nical ventilati	on or ECMO (follo	ow-up: 28 days)			l	l	I			
<b>1</b> ¹	randomized trials	not serious	not serious	not serious	very serious <sup>a</sup>	none	1/144 (0.7%)	4/145 (2.8%)	RR 0.25 (0.03 to 2.20)	21 fewer per 1,000 (from 27 fewer to 33 more)	ФФОО	CRITICAL
Serious a	dverse event	s (follow-up:	28 days)									
1 <sup>1</sup>	randomized trials	not serious	not serious	not serious	very serious a,c	none	20/142 (14.1%) <sup>d</sup>	17/142 (12.0%)	<b>RR 1.18</b> (0.64 to 2.15)	22 more per 1,000 (from 43 fewer to 138 more)	ФФОО	CRITICAL
High certa Moderate Low certa Very low Risk of bi Inconsist Indirectne Imprecisi	certainty: We a sinty: Our confic certainty: We h as: Study limital ency: Unexplair ess: Applicability	ery confident that are moderately of lence in the effect ave very little continuous decimals are the second of the	at the true effect lies of confident in the effect ct estimate is limited onfidence in the effect ty across study findinality to the research quate of an effect to su	t estimate: The true : The true effect ma t estimate: The true gs uestion	e effect is likely to b ay be substantially e effect is likely to l	ect be close to the estima different from the est be substantially differ	imate of the effec	ct .	ssibility that it is s	ubstantially different		

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

CI: Confidence interval; ECMO: Extracorporeal mechanical oxygenation; RR: Risk ratio

## **Explanations**

a. Small number of events; fragility present.

**Tables and Figures** 

- b. Upper boundary of the 95% CI crosses a threshold of meaningful effect.
- c. 95% CI cannot exclude no harm.
- d. One DVT was observed in the tofacitinib group vs zero in the placebo group.

## Reference

1. Guimaraes PO, Quirk D, Furtado RH, et al. Tofacitinib in Patients Hospitalized with Covid-19 Pneumonia. N Engl J Med 2021; 385(5): 406-15.

# **Ivermectin**

## **Evidence profiles**

- Ivermectin compared to no ivermectin for patients hospitalized with COVID-19
- Ivermectin compared to no ivermectin for ambulatory persons for management of COVID-19

**Tables and Figures** 

Table 24. GRADE evidence profile, Recommendation 23

Question: Ivermectin compared to no ivermectin for patients hospitalized with COVID-19

## Last reviewed and updated 10/10/2022

			Certainty assess	sment			№ of patients		Effect			
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ivermectin	no ivermectin	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
lortality (follow-	up: range 14	days to 28	days)									
11 <sup>1-11</sup>	randomized trials	not serious <sup>a</sup>	not serious <sup>b</sup>	not serious	serious <sup>c</sup>	none	66/1033 (6.4%)	53/937 (5.7%)	<b>RR 0.85</b> (0.40 to 1.84)	8 fewer per 1,000 (from 34 fewer to 48 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
leed for mechan	ical ventilation	on (follow-ເ	ıp: 28 days)									
3 7.8.11	randomized trials	serious <sup>d</sup>	not serious	not serious	very serious °	none	13/594 (2.2%)	28/583 (4.8%)	<b>RR 0.45</b> (0.24 to 0.86)	26 fewer per 1,000 (from 37 fewer to 7 fewer)	⊕⊖⊖⊖ VERY LOW	CRITICAL
Symptom resolut	tion (follow-u	p: 7 days)										
1 <sup>12</sup>	randomized trials	serious <sup>d</sup>	not serious	not serious	very serious <sup>c</sup>	none	16/25 (64.0%)	15/25 (60.0%)	<b>RR 1.07</b> (0.69 to 1.65)	<b>42 more per 1,000</b> (from 186 fewer to 390 more)	⊕⊖⊖⊖ VERY LOW	CRITICAL
iral clearance a	t day 7 (RCT)	(follow-up:	: range 7 days to	29 days)								
6 4,5,8,10,13,14	randomized trials	serious e	serious <sup>f</sup>	serious <sup>g</sup>	very serious °	none	77/202 (38.1%)	55/158 (34.8%)	<b>RR 1.06</b> (0.74 to 1.52)	21 more per 1,000 (from 91 fewer to 181 more)	⊕⊖⊖⊖ VERY LOW	IMPORTAN <sup>*</sup>
Serious adverse	events (follow	v-up: 28 da	ıys)									
6 2,4,7,8,9,11	randomized trials	not serious	not serious	not serious	serious <sup>c</sup>	none	38/734 (5.2%)	52/712 (7.3%)	<b>RR 1.03</b> (0.32 to 3.34)	2 more per 1,000 (from 50 fewer to 171 more)	⊕⊕⊕⊖ MODERATE	CRITICAL

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

**Tables and Figures** 

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

**Imprecision:** The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that has not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

#### **Explanations**

- a. Hashim 2021 allocated patients based on odd/even days of recruitment.
- b. Substantial heterogeneity observed (I2=68%) and introduced by Elshafie 2022 in which mortality events were reported at day 14 instead of 28 days.
- c. The 95% CI cannot exclude no meaningful effect. Few events reported do not meet the optimal information size and suggest fragility of the estimate
- d. Open label trial may lead to bias with measurement of subjective outcomes.
- e. Podder 2020 assigns participants based on odd or even registration numbers, also, 20 patients were excluded following randomization without sensitivity analysis to explore imbalance across treatment arms.
- f. Some heterogeneity observed (12=53%). Possibly explained by the longer duration of treatment (5 days compared to 1 day) in Ahmed 2021.
- g. Viral clearance is a surrogate for clinical improvement, such as hospitalization, need for ICU care and mechanical ventilation.

#### References

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**Tables and Figures** 

 Table 25. GRADE evidence profile, Recommendation 24

Question: Ivermectin compared to no ivermectin for ambulatory persons for management of COVID-19

Last reviewed and updated 10/10/2022

		С	ertainty assessm	ent			Nº of p	atients	Ef	fect		Importance
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	ivermectin	no ivermectin	Relative (95% CI)	Absolute (95% CI)	Certainty	
ortality												
14 1-14	randomized trials	not serious <sup>a</sup>	not serious	not serious	not serious	none	29/3580 (0.8%)	37/3393 (1.1%)	<b>RR 0.86</b> (0.53 to 1.40)	2 fewer per 1,000 (from 5 fewer to 4 more)	⊕⊕⊕ ніGH	CRITICAL
ogression to seve	re disease (as	sessed wi	th: need for inva	sive ventilation	1)			T		T		•
7 1,2,4,5,7,8,12	randomized trials	not serious	not serious	not serious	serious <sup>b</sup>	none	31/1505 (2.1%)	43/1375 (3.1%)	RR 0.70 (0.44 to 1.11)	9 fewer per 1,000 (from 18 fewer to 3 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
ospitalization (follo	w-up: 28 days	s)										
7 8,10-15	randomized trials	not serious	not serious	not serious	serious <sup>c</sup>	none	134/2714 (4.9%)	141/2517 (5.6%)	RR 0.88 (0.71 to 1.11)	7 fewer per 1,000 (from 16 fewer to 6 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
ral clearance at da	y 7 (RCT) (follo	ow-up: rai	nge 6 days to 29	days)	•				•			•
6 2-4,8,13,15	randomized trials	not serious	not serious	serious <sup>d,e</sup>	very serious °	none	178/574 (31.0%)	193/281 (68.7%)	RR 1.01 (0.78 to 1.31)	7 more per 1,000 (from 151 fewer to 213 more)	⊕⊖⊖⊖ VERY LOW	IMPORTAN'
me to recovery (as	sessed with: o	days)										
4 1.5.6,12	randomized trials	very serious a,f	serious <sup>g</sup>	not serious <sup>h</sup>	not serious	none	709	576	-	MD 2.99 days fewer (4.76 fewer to 1.22 fewer)i	⊕⊖⊖⊖ VERY LOW	IMPORTAN'

Serious adverse events (respiratory failure, sepsis, multiorgan failure, etc.)

**Tables and Figures** 

7 2,3,5,8,10,11,16	randomized trials	not serious	not serious	not serious	serious <sup>j</sup>	none	31/1973 (1.6%)	40/1933 (2.1%)	<b>RR 0.81</b> (0.51 to 1.30)	4 fewer per 1,000 (from 10 fewer to 6 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
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#### **GRADE Working Group grades of evidence**

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

**Imprecision:** The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that has not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio; MD: Mean difference

#### **Explanations**

- a. Concerns with unmeasured and residual confounding. Hashim 2021 allocated patients based on odd/even days of recruitment.
- b. The 95% CI cannot exclude no benefit from treatment.
- c. The 95% CI includes the potential for both appreciable benefit as well as the potential for harm. Few events reported do not meet the optimal information size and suggest fragility of the estimate
- d. Viral clearance is a surrogate for clinical improvement, such as hospitalization, need for ICU care and mechanical ventilation.
- e. Ravikirti 2021 reported viral clearance at day 6.
- f. Open label trial may lead to bias with measurement of subjective outcomes.
- g. High heterogeneity I2=90% introduced by Hashim 2021.
- h. Ivermectin was combined with doxycycline.
- i. The binary endpoint of time to recovery from the ACTIV-6 trial could not be combined with pooled continuous analysis of days to recovery; however, did not show a reduction with a HR: 1.09 (0.98, 1.22).
- j. The 95% CI cannot exclude the potential of increased SAEs in the treatment arm. Few events suggest fragility in the estimate.

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**Tables and Figures** 

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# **Fluvoxamine**

# **Evidence profiles**

• Fluvoxamine compared to no fluvoxamine for ambulatory patients with COVID-19

Tables and Figures

Table 26. GRADE evidence profile, Recommendation 25

Question: Fluvoxamine compared to no fluvoxamine for ambulatory patients with COVID-19

New evidence profile developed 10/22/2021; last updated 11/8/2021

			Certainty as	ssessment			Nº of p	atients		Effect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	fluvoxamine	no fluvoxamine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow up: 28	days) a										
2 1,2	randomized trials	not serious	not serious	not serious	very serious b	none	17/821 (2.1%)	25/828 (3.0%)	RR 0.69 (0.38 to 1.27)	9 fewer per 1,000 (from 19 fewer to 8 more)	ФФОО LOW	CRITICAL
Hospitali	zation, emerge	ency roon	n visits (>6 hours	s), or oxygen sa	turation <92% (f	ollow up: 28 days	) a					
<b>2</b> 1,2	randomized trials	not serious	not serious	serious <sup>c</sup>	serious <sup>b</sup>	none	79/821 (9.6%)	125/828 (15.1%)	<b>RR 0.64</b> (0.50 to 0.84)	54 fewer per 1,000 (from 75 fewer to 24 fewer)	ФФОО	CRITICAL
Hospitali	zation for COV	/ID-19 (fo	llow up: 28 days)	a								
21,2	randomized trials	not serious	not serious	not serious	very serious b	none	76/821 (9.3%)	103/828 (12.4%)	RR 0.75 (0.57 to 0.99)	31 fewer per 1,000 (from 53 fewer to 1 fewer)	ФФОО	CRITICAL
Viral clea	rance (follow	up: 7 day	s)									
12	randomized trials	serious d	not serious	serious <sup>e</sup>	very serious <sup>b</sup>	none	40/207 (19.3%)	58/221 (26.2%)	<b>RR 0.74</b> (0.52 to 1.05)	68 fewer per 1,000 (from 126 fewer to 13 more)	⊕⊖⊖⊖ VERY LOW	IMPORTANT
Serious a	adverse events	s <sup>a</sup>										
2 1,2	randomized trials	not serious	not serious	not serious	very serious f	none	60/821 (7.3%)	75/828 (9.1%)	RR 0.81 (0.59 to 1.12)	17 fewer per 1,000 (from 37 fewer to 11 more)	ФФОО LOW	CRITICAL

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that includes pre-print articles, which have not been peer reviewed or published.

**Tables and Figures** 

CI: Confidence interval; RR: Risk ratio; MD: Mean difference

## **Explanations**

- a. Lenze et al had a 15-day follow-up period; Reis et al had a 28 day follow up period; Serious adverse events for Reis et al included only the non-mortal grade 4 and grade 3 treatment emergent adverse events.
- b. 95% CI includes both the potential for benefit and the risk of harms; few events suggest fragility of the estimate.
- c. Hospitalization, emergency room visits are surrogate marker for clinical deterioration leading to ICU care, ventilation and mortality. In addition, best supportive care may have been substantially different in Brazil at that time compared to the U.S. health system.
- d. Data available for approximately 1/3 of study population per treatment group.
- e. Viral clearance is a surrogate for clinical improvement, such as hospitalization, need for ICU care, and mechanical ventilation.
- f. 95% CI cannot exclude the possibility of meaningful harm.

#### References

- 1. Lenze EJ, Mattar C, Zorumski CF, et al. Fluvoxamine vs Placebo and Clinical Deterioration in Outpatients With Symptomatic COVID-19: A Randomized Clinical Trial. JAMA 2020; 324(22): 2292-300.
- 2. Reis G, dos Santos Moreira Silva EA, Medeiros Silva DC, et al. Effect of early treatment with fluvoxamine on risk of emergency care and hospitalisation among patients with COVID-19: the TOGETHER randomised, platform clinical trial. Lancet **2021**; S2214-109X(21): 00448-4.

**Tables and Figures** 

# Nirmatrelvir/ritonavir

## **Evidence profiles**

• Nirmatrelvir/ritonavir compared to no nirmatrelvir/ritonavir for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease

## **FDA Emergency Use Authorization criteria**

• FDA EUA criteria for the use of nirmatrelvir/ritonavir co-packaged as Paxlovid™

## **Contraindications**

- Nirmatrelvir/ritonavir is contraindicated with drugs that are highly dependent on CYP3A for clearance and for which elevated concentrations are associated with serious and/or lifethreatening reactions
- Nirmatrelvir/ritonavir is contraindicated with drugs that are potent CYP3A inducers where significantly reduced nirmatrelvir or ritonavir plasma concentrations may be associated with the potential for loss of virologic response and possible resistance

**Tables and Figures** 

Table 27. GRADE evidence profile, Recommendation 26

Question: Nirmatrelvir/ritonavir compared to no nirmatrelvir/ritonavir for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease

New evidence profile developed 12/23/2021; last updated 2/3/2022

			Certainty as	sessment			<b>№</b> of p	patients	Ef	fect		
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	nirmatrelvir/ ritonavir	no nirmatrelvir/ ritonavir	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
All-cause	mortality (f	ollow-up:	28 days)				•					
1 <sup>1</sup>	randomized trials	serious <sup>a</sup>	not serious	not serious <sup>b</sup>	serious <sup>c</sup>	none	0/1039 (0.0%)	12/1046 (1.1%)	<b>RR 0.04</b> (0.00 to 0.68)	11 fewer per 1,000 (from 18 fewer to 5 fewer) d	ФФОО	CRITICAL
COVID-19	9-related hos	spitalizatio	ns (follow-up: 2	8 days)								
1 <sup>1</sup>	randomized trials	serious <sup>a</sup>	not serious	not serious b,e	serious <sup>c</sup>	none	8/1039 (0.8%)	65/1046 (6.2%)	<b>RR 0.12</b> (0.06 to 0.26)	55 fewer per 1,000 (from 58 fewer to 46 fewer)	ФФОО	CRITICAL
COVID-19	9-related hos	spitalizatio	n or all-cause d	eath (follow-up	: 28 days)							
1 <sup>1</sup>	randomized trials	serious <sup>a</sup>	not serious	not serious <sup>b</sup>	serious <sup>c</sup>	none	8/1039 (0.8%)	66/1046 (6.3%)	<b>RR 0.12</b> (0.06 to 0.25)	56 fewer per 1,000 (from 59 fewer to 47 fewer)	ФФОО	CRITICAL
Serious a	dverse ever	nts - not re	ported									
0	-	-	-	-	-	-	-	-	-	-	-	CRITICAL
High certa Moderate Low certa Very low c Risk of bia Inconsiste Indirectne Imprecisio	certainty: We inty: Our confi certainty: We has: Study limita ency: Unexplai ss: Applicabiliti	rery confider are moderated dence in the nave very litt stions ned heterogry by or general ence in the e	at that the true effectely confident in the effect estimate is like confidence in the eneity across study izability to the reseastimate of an effect	effect estimate: Ti mited: The true eff effect estimate: T findings arch question	he true effect is lifect may be subs	f the effect ikely to be close to th tantially different fror ikely to be substantia	n the estimate of th	e effect	ossibility that it is so	ubstantially different		

NB: Certainty ratings are derived from evidence that has not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

**Explanations** 

Tables and Figures

- a. Evidence profile based on information reported in FDA EUA and due to limited available study details, unable to exclude potential risks of bias. Concerns about selective outcome reporting as hospitalization or death from any cause and all-cause mortality are reported out of 10 outcome measures identified in the trial protocol, including SAEs and adverse events.
- b. The primary SARS-CoV-2 variant across both treatment arms was Delta (98%), including clades 21J, 21A, and 21I.
- c. Small number of events; fragility present
- d. Recalculated due to zero events in the intervention arm.
- e. COVID-19 related hospitalizations is a surrogate for ICU admission, mechanical ventilation and death. Not rated down.

#### Reference

U.S. Food and Drug Administration. Fact Sheet for Healthcare Providers: Emergency Use Authorization for Paxlovid™. Available at: <a href="https://www.fda.gov/media/155050/download">https://www.fda.gov/media/155050/download</a>. Accessed 3 February 2022.

Figure 2. FDA EUA criteria for the use of nirmatrelvir/ritonavir co-packaged as Paxlovid™ 1

Paxlovid is authorized for the treatment of mild-to-moderate COVID-19 in adults and pediatric patients (12 years of age and older weighing at least 40 kg) with positive results of direct SARS-CoV-2 viral testing, and who are at high risk for progression to severe COVID-19, including hospitalization or death.

#### Reference

1. U.S. Food and Drug Administration. Fact Sheet for Health Care Providers: Emergency Use Authorization (EUA) for Paxlovid™ Available at: <a href="https://www.fda.gov/media/155050/download">https://www.fda.gov/media/155050/download</a>. Accessed 22 December 2021.

**Figure 3.** Nirmatrelvir/ritonavir is contraindicated with drugs that are highly dependent on CYP3A for clearance and for which elevated concentrations are associated with serious and/or life-threatening reactions <sup>1\*</sup>

- Alpha1-adrenoreceptor antagonist: alfuzosin
- Antianginal: ranolazine
- Antiarrhythmic: amiodarone, dronedarone, flecainide, propafenone, quinidine
- Anti-gout: colchicine
- Antipsychotics: lurasidone, pimozide
- Benign prostatic hyperplasia agents: silodosin
- Cardiovascular agents: eplerenone, ivabradine
- Ergot derivatives: dihydroergotamine, ergotamine, methylergonovine
- HMG-CoA reductase inhibitors: lovastatin, simvastatin
- Immunosuppressants: voclosporin
- Microsomal triglyceride transfer protein inhibitor: lomitapide
- Migraine medications: eletriptan, ubrogepant
- Mineralocorticoid receptor antagonists: finerenone
- Opioid antagonists: naloxegol
- PDE5 inhibitor: sildenafil (Revatio®) when used for pulmonary arterial hypertension (PAH)
- Sedative/hypnotics: triazolam, oral midazolam
- Serotonin receptor 1A agonist/serotonin receptor 2A antagonist: flibanserin
- Vasopressin receptor antagonists: tolvapta

#### Reference

1. U.S. Food and Drug Administration. Fact Sheet for Health Care Providers: Emergency Use Authorization (EUA) for Paxlovid™ Available at: https://www.fda.gov/media/155050/download. Accessed 3 November 2022.

<sup>\*</sup>Please check drug interactions before initiating nirmatrelvir/ritonavir as the table above does not list all therapeutic agents or classes with potential interactions; see <u>Liverpool COVID-19 interactions website</u>.

**Tables and Figures** 

**Figure 4.** Nirmatrelvir/ritonavir is contraindicated with drugs that are potent CYP3A inducers where significantly reduced nirmatrelvir or ritonavir plasma concentrations may be associated with the potential for loss of virologic response and possible resistance  $^1$ 

- Anticancer drugs: apalutamide
- Anticonvulsant: carbamazepine, phenobarbital, primidone, phenytoin
- Cystic fibrosis transmembrane conductance regulator potentiators: lumacaftor/ivacaftor
- Antimycobacterials: rifampin
- Herbal products: St. John's Wort (Hypericum perforatum)

#### Reference

1. U.S. Food and Drug Administration. Fact Sheet for Health Care Providers: Emergency Use Authorization (EUA) for Paxlovid™ Available at: <a href="https://www.fda.gov/media/155050/download">https://www.fda.gov/media/155050/download</a>. Accessed 3 November 2022.

**Tables and Figures** 

## Molnupiravir

## **Evidence profiles**

• Molnupiravir compared to no molnupiravir for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease

## **FDA Emergency Use Authorization criteria**

• FDA EUA criteria for the use of molnupiravir

Tables and Figures

 Table 28. GRADE evidence profile, Recommendation 27

Question: Molnupiravir compared to no molnupiravir for ambulatory patients with mild-to-moderate COVID-19 at high risk for progression to severe disease

Last reviewed and updated 2/8/2023

	Certainty assessment						№ of patients		Effect			
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	molnupiravir	no molnupiravir	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	(follow-up: ra	ange 28 days	s to 29 days)									
<b>3</b> <sup>1-3</sup>	randomized trials	not serious	not serious	serious <sup>a,b</sup>	serious <sup>c</sup>	none	4/13328 (0.0%)	14/13314 (0.1%)	<b>RR 0.28</b> (0.09 to 0.86)	1 fewer per 1,000 (from 1 fewer to 0 fewer)	ФФОО LOW	CRITICAL
Hospitaliz	zations (follo	w-up: 29 day	rs)									
2 2,3	randomized trials	not serious	not serious	serious <sup>b,d</sup>	not serious	none	103/12619 (0.8%)	100/12615 (0.8%)	<b>RR 1.03</b> (0.78 to 1.35)	0 fewer per 1,000 (from 2 fewer to 3 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Hospitaliz	zation or dea	th (all-cause	) (follow-up: 29 d	ays)								
2 1,2	randomized trials	not serious	not serious	serious °	not serious	none	153/13238 (1.2%)	166/13224 (1.3%)	<b>RR 0.92</b> (0.74 to 1.14)	1 fewer per 1,000 (from 3 fewer to 2 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Serious a	dverse event	s (follow-up	: range 28 days to	o 29 days)								
5 <sup>1-5</sup>	randomized trials	not serious	not serious	not serious <sup>b</sup>	serious <sup>c,f</sup>	none	57/13706 (0.4%)	67/13827 (0.5%)	<b>RR 0.57</b> (0.22 to 1.52)	2 fewer per 1,000 (from 4 fewer to 3 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Adverse 6	events											
4 1,3-5	randomized trials	not serious	not serious	not serious <sup>b</sup>	serious c,f	none	97/932 (10.4%)	106/884 (12.0%)	<b>RR 0.81</b> (0.47 to 1.40)	23 fewer per 1,000 (from 64 fewer to 48 more)	⊕⊕⊕⊖ MODERATE	IMPORTANT

**Tables and Figures** 

#### **GRADE Working Group grades of evidence**

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Risk of bias: Study limitations

**Inconsistency:** Unexplained heterogeneity across study findings **Indirectness:** Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings are derived from evidence that has not been peer reviewed or published.

CI: Confidence interval; HR: Hazard ratio; RR: Risk ratio

#### **Explanations**

- a. In Bernal 2021, after day 29, one additional death resulting from adverse events occurred in the molnupiravir group and three additional deaths occurred in the placebo group.
- b. Participants included in recent large trials may not represent the population at high risk for developing severe disease.
- c. Small number of events.
- d. COVID-19 related hospitalizations is a surrogate for ICU admission, mechanical ventilation and death. Not rated down.
- e. All 10 patients reported as died at day 29 had been hospitalized.
- f. 95% CI cannot exclude the possibility of harms.

- 1. Jayk Bernal A, Gomes da Silva MM, Musungaie DB, et al. Molnupiravir for Oral Treatment of Covid-19 in Nonhospitalized Patients. N Engl J Med **2021**: Available at: https://doi.org/10.1056/nejmoa2116044 [Epub ahead of print 16 December 2021].
- 2. Butler CC, Hobbs FDR, Gbinigie OA, et al. Molnupiravir plus usual care versus usual care alone as early treatment for adults with COVID-19 at increased risk of adverse outcomes (PANORAMIC): an open-label, platform-adaptive randomised controlled trial. Lancet **2023**; 401(10373): 281-93.
- 3. Khoo SH, FitzGerald R, Saunders G, et al. Molnupiravir versus placebo in unvaccinated and vaccinated patients with early SARS-CoV-2 infection in the UK (AGILE CST-2): a randomised, placebo-controlled, double-blind, phase 2 trial. Lancet Infect Dis 2023; 23(2): 183-95.
- 4. Fischer WA, 2nd, Eron JJ, Jr., Holman W, et al. A Phase 2a clinical trial of Molnupiravir in patients with COVID-19 shows accelerated SARS-CoV-2 RNA clearance and elimination of infectious virus. Sci Transl Med **2021**: eabl7430. Available at: https://doi.org/10.1126/scitranslmed.abl7430 [Epub ahead of print 23 December 2021].
- 5. Zou R, Peng L, Shu D, et al. Antiviral Efficacy and Safety of Molnupiravir Against Omicron Variant Infection: A Randomized Controlled Clinical Trial. Front Pharmacol **2022**; 13: 939573.

**Tables and Figures** 

Figure 5. FDA EUA criteria for the use of molnupiravir 1

Molnupiravir may only be used for the treatment of mild-to-moderate COVID-19 in adults who are at high-risk for progression to severe COVID, including hospitalization or death, and for whom alternative COVID-19 treatment options authorized by FDA are not accessible or clinically appropriate.

### Reference

 U.S. Food and Drug Administration. Fact Sheet for Patients And Caregivers: Emergency Use Authorization (EUA) Of Molnupiravir For Coronavirus Disease 2019 (COVID-19). Available at: <a href="https://www.fda.gov/media/155055/download">https://www.fda.gov/media/155055/download</a>. Accessed 13 February 2023. **Tables and Figures** 

## **Colchicine**

## **Evidence profiles**

- Colchicine compared to no colchicine for hospitalized patients with COVID-19
- Colchicine compared to no colchicine for ambulatory persons with mild-to-moderate COVID-19

**Tables and Figures** 

Table 29. GRADE evidence profile, Recommendation 28

Question: Colchicine compared to no colchicine for hospitalized patients with COVID-19

Last reviewed and updated 6/13/2022

	Certainty assessment						№ of patients		Effect			
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	colchicine	no colchicine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality												
10 1-10	randomized trials	not serious	not serious	not serious	serious <sup>a</sup>	none	1335/6684 (20.0%)	1385/6810 (20.3%)	RR 0.99 (0.92 to 1.06)	2 fewer per 1,000 (from 16 fewer to 12 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Mechanica	al ventilation				•	•	•			•		
5 <sup>4-8</sup>	randomized trials	not serious <sup>b</sup>	not serious	not serious	not serious	none	652/6242 (10.4%)	651/6370 (10.2%)	RR 1.02 (0.90 to 1.16)	2 more per 1,000 (from 10 fewer to 16 more)	⊕⊕⊕ HIGH	CRITICAL
Length of	hospital stay											
4 1-3,9	randomized trials	serious <sup>c</sup>	serious <sup>d</sup>	not serious	serious <sup>a,e</sup>	none	134	132	-	MD 1.77 days fewer (3.69 fewer to 0.15 more)	⊕⊖⊖⊖ VERY LOW	CRITICAL
Adverse e	vents			l	•							
3 8-10	randomized trials	serious <sup>c</sup>	not serious	not serious	serious <sup>e,f</sup>	none	41/148 (27.7%)	20/151 (13.2%)	RR 2.04 (1.07 to 3.91)	138 more per 1,000 (from 9 more to 385 more)	$\bigoplus_{LOW} \bigcirc$	IMPORTANT
	orking Group gra								l .			
Moderate c Low certair	ertainty: We are	moderately once in the effe	ct estimate is limited:	estimate: The true The true effect may	effect is likely to be y be substantially of	ct e close to the estimate different from the estin e substantially differe	nate of the effect		y that it is subs	stantially different		
Inconsister Indirectnes Imprecisior	s: Applicability o	I heterogenei r generalizab e in the estim	ty across study findin ility to the research quate of an effect to sup f studies	uestion	cision							

Tables and Figures

NB: Certainty ratings may be derived from evidence that has not been peer reviewed or published.

CI: Confidence interval; MD: Mean difference; RR: Risk ratio

#### **Explanations**

- a. 95% CI cannot exclude the potential for both meaningful benefit or harm.
- b. Largest trial was not blinded.
- Subjectively measured outcome with >50% of studies in analysis with unclear or unreported methods for randomization and lack of blinding.
- d. High I2 (97%). One study had an imbalance of patients receiving dexamethasone (23% vs 45% in intervention vs placebo arm) possibly contributing to shorter duration of hospitalization in placebo arm.
- e. Few events suggest fragility of the estimate.
- f. 95% CI cannot exclude the potential for no meaningful harm.

- 1. Mareev VY, Orlova YA, Plisyk AG, et al. Proactive anti-inflammatory therapy with colchicine in the treatment of advanced stages of new coronavirus infection. The first results of the COLORIT study. Kardiologiia **2021**; 61(2): 15-27.
- 2. Alsultan M, Obeid A, Alsamarrai O, et al. Efficacy of Colchicine and Budesonide in Improvement Outcomes of Patients with Coronavirus Infection 2019 in Damascus, Syria: A Randomized Control Trial. Interdiscip Perspect Infect Dis 2021; 2021: 2129006.
- 3. Lopes MI, Bonjorno LP, Giannini MC, et al. Beneficial effects of colchicine for moderate to severe COVID-19: a randomised, double-blinded, placebo-controlled clinical trial. RMD Open **2021**; 7(1): e001455.
- 4. Diaz R, Orlandini A, Castellana N, et al. Effect of Colchicine vs Usual Care Alone on Intubation and 28-Day Mortality in Patients Hospitalized With COVID-19: A Randomized Clinical Trial. JAMA Netw Open **2021**; 4(12): e2141328.
- Deftereos SG, Giannopoulos G, Vrachatis DA, et al. Effect of Colchicine vs Standard Care on Cardiac and Inflammatory Biomarkers and Clinical Outcomes in Patients Hospitalized With Coronavirus Disease 2019: The GRECCO-19 Randomized Clinical Trial. JAMA Netw Open 2020; 3(6): e2013136.
- 6. RECOVERY Collaborative Group. Colchicine in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. Lancet Respir Med **2021**; 9(12): 1419-26.
- 7. Gaitán-Duarte HG, Álvarez-Moreno C, Rincón-Rodríguez CJ, et al. Effectiveness of Rosuvastatin plus Colchicine, Emtricitabine/Tenofovir and a combination of them in Hospitalized Patients with SARS Covid-19. EClinicalMedicine **2022**; 43: 101242.
- 8. Pascual-Figal DA, Roura-Piloto AE, Moral-Escudero E, et al. Colchicine in Recently Hospitalized Patients with COVID-19: A Randomized Controlled Trial (COL-COVID). Int J Gen Med **2021**; 14: 5517-26.
- 9. Absalón-Aguilar A, Rull-Gabayet M, Perez-Fragoso A, et al. Colchicine Is Safe Though Ineffective in the Treatment of Severe COVID-19: a Randomized Clinical Trial (COLCHIVID). J Gen Intern Med 2022; 37(1): 4-14.
- 10. Gorial FI, Maulood MF, Abdulamir AS, Alnuaimi AS, Abdulrrazaq MK, Bonyan FA. Randomized controlled trial of colchicine add on to the standard therapy in moderate and severe corona virus Disease-19 infection. Ann Med Surg (Lond) **2022**; 77: 103593.

**Tables and Figures** 

Table 30. GRADE evidence profile, Recommendation 29

Question: Colchicine compared to no colchicine for ambulatory persons with mild-to-moderate COVID-19

Last reviewed and updated 6/13/2022

	Certainty assessment						№ of patients		Effect			
№ of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	colchicine	no colchicine	Relative (95% CI)	Absolute (95% CI)	Certainty	Importance
Mortality	ortality											
3 1-3	randomized trials	not serious <sup>a</sup>	not serious	not serious	serious <sup>b</sup>	none	5/2431 (0.2%)	11/2426 (0.5%)	RR 0.50 (0.19 to 1.33)	2 fewer per 1,000 (from 4 fewer to 1 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Hospitali	zation											
21,3	randomized trials	not serious ª	not serious	not serious <sup>c</sup>	serious <sup>d</sup>	none	107/2391 (4.5%)	131/2386 (5.5%)	RR 0.82 (0.64 to 1.05)	10 fewer per 1,000 (from 20 fewer to 3 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Need for	mechanical v	entilation							•			
2 1,3	randomized trials	not serious	not serious	not serious	serious <sup>b</sup>	none	10/2230 (0.4%)	20/2204 (0.9%)	<b>RR 0.50</b> (0.24 to 1.07)	5 fewer per 1,000 (from 7 fewer to 1 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Serious a	dverse event	s					<u>'</u>		ľ	•		
11	randomized trials	not serious	not serious	not serious	serious <sup>b,e</sup>	none	108/2195 (4.9%)	139/2217 (6.3%)	<b>RR 0.78</b> (0.61 to 1.00)	14 fewer per 1,000 (from 24 fewer to 0 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
GRADE W	orking Group g	rades of evi	<u>dence</u>				ı	1				

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

**Tables and Figures** 

Risk of bias: Study limitations

Inconsistency: Unexplained heterogeneity across study findings Indirectness: Applicability or generalizability to the research question

Imprecision: The confidence in the estimate of an effect to support a particular decision

Publication bias: Selective publication of studies

NB: Certainty ratings may be derived from evidence that has not been peer reviewed or published.

CI: Confidence interval; RR: Risk ratio

#### **Explanations**

- a. Potential bias due to unclear or unreported details of randomization or deviations from intended interventions; however, low risk of bias for these domains within the study carrying the largest weight in the analysis and findings are not inconsistent.
- b. Few events suggests fragility of the estimate.
- c. Hospital admission is an intermediary outcome for morbidity, ICU admission, and need for ventilation. Not rated down.
- d. 95% CI cannot exclude no meaningful benefit.
- e. 95% CI cannot exclude no meaningful difference.

- Tardif J-C, Bouabdallaoui N, L'Allier PL, et al. Efficacy of colchicine in non-hospitalized patients with COVID-19. medRxiv 2021: Available at: https://doi.org/10.1101/2021.01.26.21250494 [Preprint 27 January 2021].
- 2. Gorial FI, Maulood MF, Abdulamir AS, Alnuaimi AS, Abdulrrazaq MK, Bonyan FA. Randomized controlled trial of colchicine add on to the standard therapy in moderate and severe corona virus Disease-19 infection. Ann Med Surg (Lond) 2022; 77: 103593.
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## How to approach a patient when considering pharmacologic treatments for COVID-19

- Assessment of clinical severity of COVID-19 to target treatments
- Precautions with therapeutic agents used in treating COVID-19
- COVID-19 therapies by disease severity and care location

Table 31. Assessment of clinical severity of COVID-19 to target treatments

## **Severity of COVID-19**

Mild-to-moderate COVID-19 (SpO<sub>2</sub>  $\geq$ 94% on room air and not needing supplemental oxygen) with risk factors for progression to severe disease, hospitalization or death <sup>a</sup>

Severe but not critical COVID-19 (SpO<sub>2</sub><94% on room air or needing low-flow supplemental oxygen)

Critical COVID-19 needing high-flow oxygen/ or non-invasive ventilation

Critical COVID-19 needing mechanical ventilation or ECMO

ECMO: Extracorporeal membrane oxygenation; SpO2: Oxygen saturation

a. A few of the risk factors are: age >60 years, BMI >25, diabetes, hypertension, cardiovascular disease, chronic lung disease, cancer, or immunocompromised patients. Risk factors for progression are changing as the epidemic evolves with new variants, vaccination, and previous infection rates.

 Table 32. Precautions with therapeutic agents used in treating COVID-19

Characteristic or concern	Therapeutic agents
Reduced eGFR/ increased creatinine (specific cut-offs to be mentioned for each agent)	<ul> <li>Remdesivir- Use with caution when CrCl &lt;30 mL/min</li> <li>Baricitinib- dose adjustment when CrCl &lt;60 mL/min; not recommended for eGFR, 15 mL/min</li> <li>Tofacitinib- dose adjustment when CrCl &lt;50 mL/min</li> <li>Nirmatrelvir/ritonavir- dose adjustment when eGFR &lt;60 mL/min; not recommended for eGFR</li> <li>30 mL/min</li> </ul>
Increased AST or ALT (specific cut offs to be mentioned for each agent)	<ul> <li>Baricitinib- discontinue if ALT or AST increases due to treatment</li> <li>Remdesivir- consider discontinuation if ALT/AST increases to &gt;10x the upper limit of normal</li> <li>Tofacitinib- reduce dose for moderate hepatic impairment</li> <li>Tocilizumab- may cause hepatic injury</li> <li>Sarilumab- warning to avoid when ALT/AST are &gt;1.5x ULN; discontinue if ALT/AST become 5x ULN during therapy</li> </ul>
Cytopenias a (specific cut-offs to be mentioned for each agent)	<ul> <li>Tofacitinib- warning to avoid when lymphocytes &lt;500 cells/mm3, neutrophils &lt;1000 cells/mm³, or hemoglobin &lt;9 g/dL</li> <li>Baricitinib- warning to avoid when lymphocytes &lt;500 cells/mm³, neutrophils &lt;1000 cells/mm³, or hemoglobin &lt;8 g/dL</li> <li>Tocilizumab- associated with neutropenia and thrombocytopenia; warning to avoid for chronic use when ANC &lt;2000 cells/mm³ or platelets &lt;100,000 per mm³</li> <li>Sarilumab- associated with neutropenia and thrombocytopenia; warning to avoid for chronic use when ANC &lt;2000 cells/mm³ or platelets &lt;150,000 per mm³</li> </ul>
Anti-rejection medications	Nirmatrelvir/ritonavir significantly increases concentrations of tacrolimus, cyclosporine, and sirolimus. Dose modification or temporary discontinuation of these agents are required during concomitant use.

Characteristic or concern	Therapeutic agents
Age (pediatric and adolescent) b	Molnupiravir is suggested for patients ≥18 years
	<ul> <li>Tocilizumab is suggested for patients ≥2 years</li> </ul>
	Sarilumab is suggested for patients ≥18 years
	<ul> <li>Baricitinib is suggested for patients ≥2 years</li> </ul>
	<ul> <li>Tofacitinib is suggested for patients ≥2 years</li> </ul>
	<ul> <li>Neutralizing antibodies are suggested for patients ≥12 years</li> </ul>
	<ul> <li>Nirmatrelvir/ritonavir is suggested for patients &gt;12 years</li> </ul>
	Remdesivir is indicated for all ages
	Dexamethasone is indicated for all ages
Reproductive concerns and pregnancy	Molnupiravir is not recommended during pregnancy
	Females: Advise individuals of childbearing potential to use a reliable method of contraception for the duration of treatment and for 4 days after the last dose of molnupiravir
	Males: Advise sexually active individuals with partners of childbearing potential to use a reliable method of contraception during treatment and for at least 3 months after the last dose of molnupiravir

**ALT:** Alanine transaminase; **ANC:** Absolute neutrophil count; **AST:** Aspartate transaminase; **CrCl:** Creatinine clearance; **eGFR:** Estimated glomerular filtration rate; **ULN:** Upper limit of normal

- a. Warnings come from chronic use of these medications for rheumatological disease. Patients with COVID-19 may have cytopenias, particularly lymphocytopenia, due to the viral infection. Using these agents in that situation may be indicated.
- b. Most pediatric data is derived from adult patients or other indications for these drugs.

**Table 33.** COVID-19 therapies by disease severity and care location

Care location and COVID-19 severity	Pharmacologic treatments available in the United States
Ambulatory mild-to- moderate disease (not hypoxemic) with high risk for progression to severe disease, hospitalization or death (see individual drug section for specific considerations for each of these agents)	<ul> <li>Nirmatrelvir/ritonavir X 5 days (oral)</li> <li>Remdesivir x 3 days (intravenous)</li> <li>Anti-SARS-CoV-2 monoclonal antibodies a</li> <li>If other treatment options are not available then consider Molnupiravir x 5 days (oral) or, if immunocompromised, high-titer convalescent plasma with activity against circulating variant (intravenous).</li> </ul>
Can be considered in patients with mild-moderate COVID-19 hospitalized for other reasons	<ul> <li>Systemic steroids have no demonstrated benefit and may harm.</li> <li>No benefit demonstrated for hydroxychloroquine, azithromycin, lopinavir/ritonavir, or ivermectin.</li> </ul>
Hospitalized for mild-to- moderate COVID-19 (not hypoxemic)	<ul> <li>If at high risk for progression and within 7 days of symptom onset, remdesivir x 3 days.</li> <li>Systemic steroids have no demonstrated benefit and may harm.</li> <li>No benefit demonstrated in RCTs for convalescent plasma, hydroxychloroquine, azithromycin, lopinavir/ritonavir, or ivermectin.</li> </ul>
Hospitalized for severe, but not critical COVID-19 (hypoxemic needing low flow supplemental oxygen)	<ul> <li>Corticosteroids (dexamethasone 6 mg/d x 10 days or until discharge or an equivalent dose of another agent).</li> <li>Remdesivir x 5 days</li> <li>Tocilizumab or Sarilumab in progressive disease with elevated inflammatory makers.</li> <li>Baricitinib or tofacitinib in patients with elevated inflammatory markers.</li> <li>No benefit demonstrated in RCTs for convalescent plasma, hydroxychloroquine, azithromycin, lopinavir/ritonavir, or ivermectin.</li> </ul>
Hospitalized for critically ill COVID-19, needing non-invasive ventilation or Hi flow oxygen	<ul> <li>Corticosteroids (dexamethasone 6 mg/d x 10 days or until discharge or an equivalent dose of hydrocortisone or methylprednisolone).</li> <li>Tocilizumab or Sarilumab in patients with elevated inflammatory makers</li> </ul>

Care location and COVID-19 severity	Pharmacologic treatments available in the United States
	<ul> <li>Baricitinib or tofacitinib in patients with elevated inflammatory markers</li> <li>No benefit demonstrated in RCTs for remdesivir, convalescent plasma, hydroxychloroquine, azithromycin, lopinavir/ritonavir, or ivermectin.</li> </ul>
Hospitalized for critically ill COVID-19, needing invasive mechanical ventilation or ECMO	<ul> <li>Corticosteroids (dexamethasone 6 mg/d x 10 days or until discharge or an equivalent dose of hydrocortisone or methylprednisolone).</li> <li>Tocilizumab or sarilumab in patients with elevated inflammatory makers</li> <li>Baricitinib or tofacitinib in patients with elevated inflammatory markers</li> <li>No benefit demonstrated in RCTs for remdesivir, convalescent plasma, hydroxychloroquine, azithromycin, lopinavir/ritonavir, or ivermectin.</li> </ul>

ECMO: Extracorporeal membrane oxygenation; RCTs: Randomized controlled trials

a. Neutralizing antibodies that are active against prevalent variants should be utilized. For example, at present (04/2022) bebtelovimab has *in vitro* activity against Omicron BA.2 subvariant and should be utilized, but casirivimab/imdevimab, bamlanivimab/etesevimab and sotrovimab do not have reliable activity against circulating omicron BA.2 variant and should be avoided.

# Pediatric considerations for treatment of SARS-CoV-2 infection and multisystem inflammatory syndrome in children

• Case definitions for Multisystem Inflammatory Syndrome in Children (MIS-C) and Paediatric inflammatory multisystem syndrome temporally associated with COVID-19 (PIMS-TC, also called pediatric multisystem inflammatory disorder [PMIS])

**Table 34.** Case definitions for Multisystem Inflammatory Syndrome in Children (MIS-C) and Paediatric inflammatory multisystem syndrome temporally associated with COVID-19 (PIMS-TC, also called pediatric multisystem inflammatory disorder [PMIS])

	MIS-C (CDC 2020) <sup>1</sup>	PIMS-TS or PMIS (Royal College of Paediatrics and Child Health 2020) <sup>2</sup>
Includes	<ul> <li>Age &lt;21 years presenting with:         <ul> <li>Fever (&gt;38.0°C for ≥24 hours, or report of subjective fever lasting ≥24 hours)</li> <li>Laboratory evidence of inflammation (including, but not limited to, one or more of the following: an elevated C-reactive protein, erythrocyte sedimentation rate, fibrinogen, procalcitonin, d-dimer, ferritin, lactic acid dehydrogenase, or interleukin 6, elevated neutrophils, reduced lymphocytes and low albumin),</li> <li>Evidence of clinically severe illness requiring hospitalization, with multisystem (&gt;2) organ involvement (cardiac, renal, respiratory, hematologic, gastrointestinal, dermatologic or neurological)</li> </ul> </li> </ul>	<ul> <li>A child presenting with:         <ul> <li>Persistent fever &gt;38.5°C</li> </ul> </li> <li>Laboratory evidence of inflammation (neutrophilia, elevated CRP and lymphopenia)</li> <li>Evidence of single or multi-organ dysfunction (shock, cardiac, respiratory, renal, gastrointestinal or neurological disorder) with additional features (listed in Appendix of reference)</li> </ul>
Excludes	Patients with alternative plausible diagnoses	Patients with any other microbial cause, including bacterial sepsis, staphylococcal or streptococcal shock syndromes, infections associated with myocarditis such as enterovirus
Other criteria	Positive for current or recent SARS-CoV-2 infection by RT-PCR, serology, or antigen test; OR COVID-19 exposure within the 4 weeks prior to the onset of symptoms	SARS-CoV-2 PCR testing may be positive or negative

- 1. Centers for Disease Control and Prevention. Multisystem Inflammatory Syndrome in Children (MIS-C) Associated with Coronavirus Disease 2019 (COVID-19). Available at: <a href="https://emergency.cdc.gov/han/2020/han00432.asp">https://emergency.cdc.gov/han/2020/han00432.asp</a>. Accessed 23 November 2021.
- 2. Royal College of Paediatrics and Child Health. Guidance: Paediatric multisystem inflammatory syndrome temporally associated with COVID-19, **2020**.

Tables and Figures