ONGOING LIVING UPDATE OF COVID-19 THERAPEUTIC OPTIONS

Summary of Evidence • Rapid Review, 3 October 2022





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Ongoing Living Update of COVID-19 Therapeutic Options: Summary of Evidence, Rapid Review. 3 October 2022

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Disclaimer

This document includes the results of a rapid systematic review of current available literature. The information included in this review reflects the evidence as of the date posted in the document. In recognition of the fact that there are numerous ongoing clinical studies, PAHO will periodically update this review and corresponding recommendations as new evidence becomes available.

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Executive summary

Background

The urgent need for evidence on measures to respond to the COVID-19 pandemic had led to a rapid escalation in numbers of studies testing potential therapeutic options. The vast amount of data generated by these studies must be interpreted quickly so that physicians have the information to make optimal treatment decisions and manufacturers can scale-up production and bolster supply chains. Moreover, obtaining a quick answer to the question of whether or not a particular intervention is effective can help investigators involved in the many ongoing clinical trials to change focus and pivot to more promising alternatives. It is crucial for healthcare workers to have access to the most up-to-date research evidence to inform their treatment decisions.

To address this evidence gap, we compiled the following database of evidence on potential therapeutic options for COVID-19. We hope this information will help investigators, policy makers, and prescribers navigate the flood of relevant data to ensure that management of COVID-19, at both individual and population levels, is based on the best available knowledge. We will endeavor to continually update this resource as more research is released into the public space.

Summary of evidence

Tables 1 and 2, which divide the total group of identified studies into randomized (Table 1) and non-randomized (Table 2) designs, indicate the primary outcome measures used for each investigation and the level of certainty. <u>A living interactive version of Tables 1 and 2 is available here.</u> Table 3 summarizes the status of evidence for the 237 potential therapeutic options for COVID-19 for which studies were identified through our systematic review.



		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		Invasive	1	and the second		
		Overall number of studies including the	Mortality	mechanical ventilation	Symptom resolution	Prevention of infection	Adverse events	Hospitalization
Intervention		intervention, n=730	(n of studies)	(n of studies)	(n of studies)	(n of studies)	(n of studies)	(n of studies)
Hydroxychloroguine or Chloroguine		61		10		7(*)	20	13
Convalescent plasma	NEW	58	50	22	13		17	4 (5)
Ivermectin		49	12(*)		7(*)	1(*)	8	
Favipiravir	NEW	29			4(*)		8	
Tocilizumab		29			12		17	
Corticosteroids		23			6		6	
Lopinavir-Ritonavir		21	4				3	
	NICTAL			4	1			
Anticoagulants	NEW		11(@@)	3	1		11 (*)	4
Vitamin D	NEW					2(@@)	2	3
Sofosbuvir +/- Daclatasvir or others		16						
Colchicine		15	12(**)	6(**)	5(**)		3	2
Mouthwash		14	1	1	2			
ACEIs or ARBs		12	8(*)	9	3		1	1
REGEN-COV (casirivimab and imdevimab)		12	2(##)	2(##)	3(##)	3	6	4
Azithromycin		11	6	5	6			2
Molnupiravir		10	4		2		4	6
Remdesivir		10	8	7	4		4	1
Sarilumab		10	-				6	
Bamlanivimab +/- etesevimab		9			3	1	6	
							And and a second se	
Corticosteroids (inhaled)		9			8		4	5
Mesenchimal cell tranplantation	NEW	9	7				1	
Vitamin C		-9			4		1	
Melatonin		8	3		3			
Zinc		8	2	1	2	2	1	1
Baricitinib		7	5		3		3	
Interferon beta-1a		7	6	4	2		2	
Nitazoxanide		7	2	1	1		3	2
Umifenovir		7	1	2			1	
Anakinra		6					5	
							Taxan and tax	
IVIG		6					1	
Bromhexine Hydrochloride		5				2		
Camostat mesilate		5			3		2	2
Probiotics		5			1			
Tenofovir • emtricitabine		5	2	2	1	1	3	2
Aspirin		4	3	3	1		1	
Doxycycline		4	2		2	1		1
Fluvoxamine		4	1				2	3 (§)
Hyperimmune anti-COVID-19 IVIG		4			1		2	107
Nasal hypertonic saline		4			1			
Nitric oxide		4	2	2				
Proxalutamide	10.1	4	3	3	2			2
Peg-IFN lambda	NEW	4			1		1	1
Quercetin		4	3		2		1	1
Cofactors		3	1		1		1	
Famotidine		3	2	2	1			
Hyperbaric oxygen		3	3	2	.1		1	
Interferon beta-1b		3	2	3	1			
Low-dose radiation therapy		3	2					
Metformin		3	2				1	2
N-acetylcysteine		3	2			1	1	
		3						
Omega-3 fatty acids Ruxolitinib							1.0	
1 (m. harren ille)		3					3	
Sotrovimab		3	1				1	1
Statins		3	2					
Tixagevimab-Cilgavimab		3	3		1	1	3	1
Beta glucans		2					1	
Canakinumab		2	2	4	4		1	
Dutasteride		2			1			
Electrolyzed salme		2			1		4	
lota-Carrageenan		2					2	4
Leflunomide		2	-				-	
Levamisole		2			1			2
Linagliptin		2						
Niclosamide		2		1			1	1
Nigella sativa +/- Honey		2			1			1
Opaganib		2	2				2	
P2Y12		2	2	1	1		2	

Table 1. List of RCTs of interventions for COVID-19 with primary outcome measures and certainty (n=730) (interactive online version)



	Overall number of	1	Invasive mechanical		Prevention of	1 Contractor	
ntervention	studies including the intervention, n=730	Mortality (n of studies)	ventilation (n of studies)	Symptom resolution (n of studies)	infection (n of studies)	Adverse events (n of studies)	Hospitalization (n of studies)
Peg-IFN alfa			2	2		(n or studies)	(in or studies)
Pentoxifylline		PC	2 2				-
Regdanvimab		2		2		2	
Resveratrol			3 3			4	3
Spironolactone			1 1				
Thalidomide		2	1 3				
Tissue-plasminogen activator (tPA)	NEW	2	2			1	
Tofacitinib		2	1	1		1	
Vilobelimab		i and i a	2				
99mTc-MDP		1					
Adalimumab			1 1				
Alpha-1 antitrypsin		1	1				
Amiodarone			1 1			19	
Ammonium chloride		1	1 1	The second se		11	
AMP5A (inhaled)		1	1			1	
APMV2020 (aspirin, promethazine, micronutrients)		1	1			1	1
Aprepitant		1					
Aprotinin		-	1				
		1	1				
Arbidol							
ArtemiC			1	1			
Artemisinin		1		1			
Atazanavir-ritonavir		1	1 1	1			
Atovaquone		1	1			5	
Auxora		1	1	1			
Avdoralimab		1	1			1	
			1	1			
Aviptadil			1	1			2
Ayush-64		1		1		1	
AZD1656	NEW	1	1	1			1
Azelastine (inhaled)		1		1		1	1
Azvudine		1					
Baloxavir		1		1			
BCG		1	4				
			1				
Bebtelovimab							
Bioven		1	1				
Bicarbonate (inhaled)		1	1				
Boswellia extract		1		1			1
Calcitriol		1	1				1
Cannabidiol		1	1 1	1			
CD24Fc		1	1 1				
CERC-002			1				0.
			-1			A Real Property lies and a	
Chloroquine nasal drops		1					
CIGB-325		1		1			
Clarithromycin		1					
Clazakizumab		1	1 1	1			
Clevudine		1				1 ×	
Colchicine + rosuvastatin		1	1 1			3	
Corticosteroids (nasal)		1					
Crizanlizumab		1	1 1	1			
			4				
Curcumin + Piperine		1		1		1	
Curcumin + Quercetin + Vitamin D		1					
Darunavir-Cobicistat		1					
Dapagliflozin		1	1	1		1	1
Degarelix		1	1 1				
DFV890	NEW		1 1	1			
Dimethyl sulfoxide (DSMO)	and the second se	1				1	
		2					
Dornase alfa (inh)		1		1			
Dupilumab		1	1				
Edaravone		1	1 7				
Endothelial dysfunction protocol		1	1 1			1 8	
Enisamium		1		1			
Ensitrelvir			1				
				-			
Ensovibep			1	1		-	
Enzalutamide			1 1				
Ethanol (inhaled)		1	1	1		3	
Febuxostat		1					1
Fenofibrate		1	1 1				





hereit.	Overall number of studies including the	Mortality	Invasive mechanical ventilation	Symptom resolution	Prevention of Infection	Adverse events	Hospitalization
Intervention	intervention, n=730	(n of studies)	(n of studies)	(n of studies)	(n of studies)	(n of studies)	(n of studies)
Gabapentin +/- Montelukast		1		1		1	
GB0139 (inhaled)		1	1			1	
Gimsilumab (Anti-GM-CSF Monoclonal Antibody)		1	1	1		1	
Helium (inhaled)		1					
Hemadsorption		1	1	t			
Hesperidin		1	-	1 1			-
Hypertonic saline (inhaled)			2				
hzVSF-v13							
Ibrutinib		1	1	1		1	
Icatibant/ iC1e/K		1	1				
Icosapent ethyl	1.1	1		1			
IFN-alpha2b + IFN-gamma		1					
Imatinib			1	t		1	
Indomethacin			1			-	
Infliximab		-	1	1		1	
INM005 (equine antibodies)			1	1 1		1	
Interferon beta-1a (inhaled)			1	1 1		1	
Interferon gamma		1					
Interferon kappa + TFF2		t	1			1	
Interferon-2		1	1	1		1	
Isothymol		1	1				
Itolizumab		1	1	r l		1	
			1			1	
Ivermectin (inhaled)				1			
Ixekizumab		1	1				
KB109		1	1	1		1	
L-arginine		1	1			1	
Lactococcus Lactis (intranasal)		t		1		1	
Lactoferrin		t		1			
Lenzilumab			1				
Levilimab			1	1 1		1	
Lincomycin							
Lithium		1	1			1	
Mavrilimumab		1	1 -	1 1		1	
Mefenamic acid		1	1			1	4
Metisoprinol		t de la companya de la					
Methylene blue		1	Ť				
Metoprolol			1				
Metronidazole							
				-			
Montelukast			1				
Mupadolimab						1	
Mycobacterium w		F	1				
N-acetylcysteine (inhaled)	1.1	1	1				
Nafamostat mesylate		t •	1			- 1	
Namilumab		1	1	1		1	
Nano-curcumin		t Barris t	M			1	
Neem (Azadirachta Indica A. Juss)							
			1		-	1	
Nicotine patches		-				1 m	
Nirmatrelvir-ritonavir		-	1			1	
Norelgestromin and Ethinylestradiol		1					
Novaferon		1					
NSAIDS		1	1	1		1	
Nutritional support		1	1	1			
OP-101			1	1		1	
Oblimab			1			-	
						1	
Palmitoylethanolamide NEV							
Pembrolizumab				1 1		1	
Pirfenidone NEV	/	t in the second s	1	1 1		.1	
Plitidepsin		1	1	1		1	
PNB001 (CCK-A antagonist)		t	1	1			
Polymerized type I collagen (PT1C)		t I and a second se					
Potassium Canrenoate		1	1			-	
			4			-	
Povidone iodine			1			1	
Progesterone		1		1		1	
Prolectin-M		1		1		1	
Propolis		t	1	1 1			
Prostacyclin		1	1			1	
Prostacyclin (inhaled)			1				
(minutes)			1				





ntervention	Overall number of studies including the intervention, n=730	Mortality (n of studies)	Invasive mechanical ventilation (n of studies)	Symptom resolution (n of studies)	Prevention of infection (n of studies)	Adverse events (n of studies)	Hospitalizatior (n of studies)
Pyridostigmine			1	1 1	(ii oi otadico)		(in or statutes)
Raloxifene		1	1				
Ramipril		1	1			1	
RD-X19 (light therapy)		1		1			
Recombinant Super-Compound IFN		1	1	1			
Remdesivir (inhaled)		1					
leparixin		1	1	1			1
libavirin							
Ribavirin + Interferon beta-1b		1					
hG-CSF		1	1	1			1
hG-CSF (inhaled)		1	1	1 1		1	
hu-pGSN		1	1	1			
abizabulin		1	1				
ecukinumab		1	1	1			
enicapoc		1	1				
entinox						1	1
hort-wave diathermy		1	1	1		1	
ildenafil		1	1	1		1	1
ilymarin		1		1		1	1
iltuximab		1	1	1			
itagliptin		1	1	1			
tem-cell nebulization		1	1	1		1	1
ulodexide		1	1	1		1	1
afenoquine		1		1		1	1
D-0903 (inhaled JAK-inhibitor)		1	1			1	1
hymoQuinone		1				1	1
ranilast		1	1	1			
riazavirin		1	1	1		1	1
XA-127		1	1	1			
Itraviolet light phototherapy		1	1			1	1
erapamil		1	1	1		1	1
tamin B	NEW	1					
AV-19 (swine polyclonal antibodies)		1	1			1	
lucoplan		1	1			1	
-Lipoic acid		1	1				

The COLCORONA trial that included patients with recent onset mild disease showed a tendency to less hospitalizations, less mortality and less mechanical ventilation requirements. However the certainty on those potential benefits was low because of very serious imprecision as the number of events was low; (##) Subgroup of seronegative patients; (@) High dose schemes (i.e dexamethasone 12 mg a day) may be more effective than standard dose schemes (i.e dexamethasone 6 mg a day); (@@) Excluding high risk of bias studies; (§) Observed effects would probably be considered important in patients with very high hospitalization risk (>10%).

	GRADE High- Moderate certainty	GRADE Low certainty
Beneficial effect		and the second
No significant effect		
Harmful effect		
Uncertain effect		
No evidence or no estimable effect		

Table 2. List of non-RCTs of interventions for COVID-19 with primary outcome measures and certainty (n=7). (interactive online version)

Intervention	Overall number of studies including t intervention		Mechanical ventilation (n of studies)	Symptom resolution (n of studies)	Prevention of infection (n of studies)	Adverse events (n of studies)
NSAID		7	7			
Dana finial affinat	GRADE High- Moderate certainty	GRADE Low certa	inty			
Beneficial effect No significant effect		_				
Harmful effect						
Uncertain effect	1. Contract (1. Co					
No evidence or no estimable effect						



Table 3. Summary of findings on potential therapeutic options for COVID-19 (n=237), as at 3 October 2022

	8Intervention	Summary of findings
1	99mTc-MDP	Uncertainty in potential benefits and harms. Further research is needed.
2	Adalimumab	Uncertainty in potential benefits and harms. Further research is needed.
3	ACEIs or ARBs	Continuing or initiating ACEIs or ARBs in patients with COVID-19 may increase mortality. However, the certainty of the evidence was low. Further research is needed.
4	Alpha-1 antitrypsin	Uncertainty in potential benefits and harms. Further research is needed.
5	Amiodarone	Uncertainty in potential benefits and harms. Further research is needed.
6	Ammonium chloride	Uncertainty in potential benefits and harms. Further research is needed.
7	AMP5A (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
8	Anakinra	Anakinra may not reduce mortality or increase severe adverse events. However, the certainty of the evidence was low because of risk of bias and imprecision. Further research is needed.
9	Anticoagulants	There are specific recommendations on the use of antithrombotic agents for thromboprophylaxis in hospitalized patients with COVID-19. Regarding the best thromboprophylactic scheme, anticoagulants in intermediate (i.e., enoxaparin 1 mg/kg a day) or full dose (i.e., enoxaparin 1 mg/kg twice a day) probably does not decrease mortality in comparison with prophylactic dose (i.e., enoxaparin 40 mg a day). Anticoagulants in full dose decrease venous thromboembolic events but increase major bleeding in comparison with prophylactic dose. In mild ambulatory patients, anticoagulants in prophylactic dose, may not importantly improve time to symptom resolution or reduce hospitalizations.
10	APMV2020 (aspirin, promethazine, micronutrients)	Uncertainty in potential benefits and harms. Further research is needed.



	8Intervention	Summary of findings
_		
11	Aprepitant	Uncertainty in potential benefits and harms. Further research is needed.
12	Aprotinin	Uncertainty in potential benefits and harms. Further research is needed.
13	Arbidol	Uncertainty in potential benefits and harms. Further research is needed.
14	ArtemiC (artemisinina, curcumina, frankincense, and vitamin C):	Uncertainty in potential benefits and harms. Further research is needed.
15	Artemisinin	Uncertainty in potential benefits and harms. Further research is needed.
16	Aspirin	Aspirin probably does not reduce mortality, or mechanical ventilation and probably does not increase symptom resolution or improvement.
17	Atazanavir/ritonavir	Uncertainty in potential benefits and harms. Further research is needed.
18	Atovaquone	Uncertainty in potential benefits and harms. Further research is needed.
19	Auxora	Auxora may not increase severe adverse events. The effects of auxora on other important outcomes are uncertain. Further research is needed.
20	Avdoralimab	Uncertainty in potential benefits and harms. Further research is needed.
21	Aviptadil	Uncertainty in potential benefits and harms. Further research is needed.
22	Ayush-64	Uncertainty in potential benefits and harms. Further research is needed.
23	AZD1656	AZD1656 may improve time to symptom resolution. The effects of AZD 1656 on other important outcomes are uncertain. Further research is needed.





	8Intervention	Summary of findings
24	Azelastine	Uncertainty in potential benefits and harms. Further research is needed.
25	Azithromycin	Azithromycin probably does not reduce mortality or mechanical ventilation and does not improve time to symptom resolution.
26	Azvudine	Uncertainty in potential benefits and harms. Further research is needed.
27	Baricitinib	In patients with moderate to critical disease, baricitinib reduces mortality, probably reduces mechanical ventilation requirements, and probably improves time to symptom resolution, without increasing severe adverse events.
28	Baloxavir	Uncertainty in potential benefits and harms. Further research is needed.
29	Bamlanivimab +/- etesevimab (monoclonal antibody)	Bamlanivimab probably reduces hospitalizations in patients with COVID-19 and it probably reduces symptomatic infections in exposed individuals. It is uncertain if it affects mortality or mechanical ventilation requirements. Further research is needed.
30	BCG	Uncertainty in potential benefits and harms. Further research is needed.
31	Bebtelovimab	Uncertainty in potential benefits and harms. Further research is needed.
32	Beta-glucans	Uncertainty in potential benefits and harms. Further research is needed.
33	Bicarbonate (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
34	Bioven	Uncertainty in potential benefits and harms. Further research is needed.
35	Boswellia extract	Uncertainty in potential benefits and harms. Further research is needed.
36	Bromhexine hydrochloride	Bromhexine may reduce symptomatic infections in exposed individuals. Its effects on other clinical important outcomes are uncertain. Further research is needed.





	8Intervention	Summary of findings
37	Calcitriol	Uncertainty in potential benefits and harms. Further research is needed.
38	Camostat mesilate	Camostat mesilate may not improve time to symptom resolution. Further research is needed.
39	Canakinumab	Uncertainty in potential benefits and harms. Further research is needed.
40	Cannabidiol	Uncertainty in potential benefits and harms. Further research is needed.
41	CD24Fc (soluble CD24 appended to heavy chains 2 and 3 of human immunoglobulin G1)	CD24Fc may reduce mechanical ventilation and increase symptom resolution or improvement. However, certainty of the evidence was low for imprecision. Further research is needed.
42	CERC-002	Uncertainty in potential benefits and harms. Further research is needed.
43	Chloroquine nasal drops	Uncertainty in potential benefits and harms. Further research is needed.
44	CIGB-325	Uncertainty in potential benefits and harms. Further research is needed.
45	Clarithromycin	Uncertainty in potential benefits and harms. Further research is needed.
46	Clazakizumab	Clazakizumab may reduce mechanical ventilation and improve time to symptoms resolution. However, certainty of the evidence was low. Further research is needed.
47	Clevudine	Uncertainty in potential benefits and harms. Further research is needed.
48	Cofactors (L-carnitine, N-acetylcysteine, nicotinamide, serine)	Uncertainty in potential benefits and harms. Further research is needed.
49	Colchicine	Colchicine probably does not reduce mortality, mechanical ventilation requirements or increase symptom resolution or improvement with moderate certainty. In patients with mild recent onset COVID-19 colchicine probably





	8Intervention	Summary of findings
		
		does not have an important effect on hospitalizations. However, the certainty of the evidence was low because of imprecision.
50	Colchicine + rosuvastatin	Uncertainty in potential benefits and harms. Further research is needed.
51	Convalescent plasma	Convalescent plasma does not reduce mortality or reduces mechanical ventilation requirements or improves time to symptom resolution with moderate to high certainty of the evidence. In patients with recent onset mild COVID-19 convalescent plasma probably does not have an important effect on hospitalizations. Convalescent plasma may not increase severe adverse events.
52	Crizanlizumab	Uncertainty in potential benefits and harms. Further research is needed.
53	Curcumin + piperine	Uncertainty in potential benefits and harms. Further research is needed.
54	Curcumin + quercetin + vitamin D	Uncertainty in potential benefits and harms. Further research is needed.
55	Dapagliflozin	Dapagliflozin may reduce mortality but probably does not increase symptom resolution. Further research is needed.
56	Darunavir-cobicistat	Uncertainty in potential benefits and harms. Further research is needed.
57	Degarelix	Uncertainty in potential benefits and harms. Further research is needed.
58	DFV890	DVF890 may improve time to symptom resolution. The effects of AZD 1656 on other important outcomes are uncertain. Further research is needed.
59	Dimethyl sulfoxide (DSMO)	Uncertainty in potential benefits and harms. Further research is needed.
60	Dornase alfa (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.





	8Intervention	Summary of findings
61	Doxycycline	Doxycycline does not increase symptom resolution or improvement and may not reduce hospitalizations.
62	Dutasteride	Uncertainty in potential benefits and harms. Further research is needed.
63	Dupilumab	Uncertainty in potential benefits and harms. Further research is needed.
64	Edaravone	Uncertainty in potential benefits and harms. Further research is needed.
65	Electrolyzed saline	Uncertainty in potential benefits and harms. Further research is needed.
66	Endothelial dysfunction protocol	Uncertainty in potential benefits and harms. Further research is needed.
67	Enisamium	Uncertainty in potential benefits and harms. Further research is needed.
68	Ensovibep	Uncertainty in potential benefits and harms. Further research is needed.
69	Ensitrelvir	Uncertainty in potential benefits and harms. Further research is needed.
70	Enzalutamide	Uncertainty in potential benefits and harms. Further research is needed.
71	Ethanol (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
72	Famotidine	Uncertainty in potential benefits and harms. Further research is needed.
73	Favipiravir	Favipiravir may increase mortality and mechanical ventilation requirements; it may not reduce hospitalizations and it does not improve symptom resolution. Further research is needed.



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	8Intervention	Summary of findings
		a manual y a amanga
74	Febuxostat	Uncertainty in potential benefits and harms. Further research is needed.
75	Fenofibrate	Fenofibrate may not increase severe adverse events. The effects of fenofibrate on other important outcomes are uncertain. Further research is needed.
76	Finasteride	Uncertainty in potential benefits and harms. Further research is needed.
77	Fluvoxamine	In patients with recent onset mild COVID-19 fluvoxamine probably does not have an important effect on hospitalizations and may not increase severe adverse events. Certainty of the evidence was low to moderate. Further research is needed.
78	Fostamatinib	Uncertainty in potential benefits and harms. Further research is needed.
79	Gabapentin +/- montelukast	Uncertainty in potential benefits and harms. Further research is needed.
80	GB0139 (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
81	Gimsilumab (anti-GM-CSF monoclonal antibody)	Gimsilumab may not reduce mortality nor increase symptom resolution. Further research is needed.
82	Helium (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
83	Hemadsorption	Uncertainty in potential benefits and harms. Further research is needed.
84	Hesperidin	Hesperidin may not improve symptom resolution; however, the certainty of the evidence was low. Further research is needed.
85	Hydroxychloroquine and chloroquine	Hydroxychloroquine or chloroquine probably increases mortality, and probably does not reduce invasive mechanical ventilation or significantly improve time to symptom resolution with moderate certainty. When used prophylactically in persons exposed to COVID-19 it may not have an important effect on the risk of infection and in patients with mild, recent onset disease, and it may not have an important effect on hospitalizations. However, certainty of the evidence is low because of risk of bias and imprecision.



	8Intervention	Summary of findings
	ontervention	Summary or minings
86	Hyperbaric oxygen	Uncertainty in potential benefits and harms. Further research is needed.
87	Hyperimmune anti-COVID-19 intravenous immunoglobulin (C-IVIG)	Uncertainty in potential benefits and harms. Further research is needed.
88	Hypertonic saline (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
89	hzVSF-v13	Uncertainty in potential benefits and harms. Further research is needed.
90	Ibrutinib	Uncertainty in potential benefits and harms. Further research is needed.
91	Icatibant/iC1e/K	Uncertainty in potential benefits and harms. Further research is needed.
92	Icosapent ethyl	Uncertainty in potential benefits and harms. Further research is needed.
93	Imatinib	Imatinib may not increase severe adverse events. The effects of imatinib on other importan outcomes are uncertain. Further research is needed.
94	Indomethacin	Uncertainty in potential benefits and harms. Further research is needed.
95	Infliximab	Uncertainty in potential benefits and harms. Further research is needed.
96	INM005 (polyclonal fragments of equine antibodies)	Uncertainty in potential benefits and harms. Further research is needed.
97	Interferon alpha-2b and interferon gamma	Uncertainty in potential benefits and harms. Further research is needed.
98	Interferon beta-1a	IFN beta-1a probably does not reduce mortality, invasive mechanical ventilation requirements or improve symptom resolution. Further research is needed.





	8Intervention	Summary of findings
99	Interferon beta-1a (inhaled)	Inhaled interferon beta-1a may improve time to symptom resolution. Further research is needed.
100	Interferon beta-1b	Uncertainty in potential benefits and harms. Further research is needed.
101	Interferon gamma	Uncertainty in potential benefits and harms. Further research is needed.
102	Interferon kappa and TFF2	Uncertainty in potential benefits and harms. Further research is needed.
103	Interleukin-2	Uncertainty in potential benefits and harms. Further research is needed.
104	Iota-carrageenan	Uncertainty in potential benefits and harms. Further research is needed.
105	Itolizumab	Uncertainty in potential benefits and harms. Further research is needed.
106	Ivermectin	Although pooled estimates suggest significant benefits with ivermectin, included studies' methodological limitations and a small overall number of events result in very low certainty of the evidence. Based on the results reported by the RCTs classified as low risk of bias, ivermectin probably does not reduce mortality or improve time to symptom resolution. In patients with recent onset of the disease, ivermectin probably does not have an important effect on hospitalizations and may not increase severe adverse events. It is uncertain if it reduces symptomatic infections when used as prophylaxis.
107	Ivermectin (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
108	IVIG (intravenous immunoglobulin)	Uncertainty in potential benefits and harms. Further research is needed.
109	Ixekizumab	Uncertainty in potential benefits and harms. Further research is needed.
110	KB109	Uncertainty in potential benefits and harms. Further research is needed.





	8Intervention	Summary of findings
111	L-arginine	Uncertainty in potential benefits and harms. Further research is needed.
112	Lactococcus lactis (intranasal)	Uncertainty in potential benefits and harms. Further research is needed.
113	Lactoferrin	Uncertainty in potential benefits and harms. Further research is needed.
114	Leflunomide	Uncertainty in potential benefits and harms. Further research is needed.
115	Lenzilumab	Lenzilumab may reduce mechanical ventilation requirements and may not increase severe adverse events. The effects of lenzilumab on other important outcomes are uncertain. Further research is needed.
116	Levamisole	Uncertainty in potential benefits and harms. Further research is needed.
117	Levilimab	Levilimab may improve time to symptom resolution; however, the certainty of the evidence was low. The effects of levilimab on other important outcomes are uncertain. Further research is needed.
118	Linagliptin	Uncertainty in potential benefits and harms. Further research is needed.
119	Lincomycin	Uncertainty in potential benefits and harms. Further research is needed.
120	Lithium	Uncertainty in potential benefits and harms. Further research is needed.
121	Lopinavir-ritonavir	Lopinavir-ritonavir probably does not reduce mortality with moderate certainty. Lopinavir-ritonavir may not be associated with a significant increase in severe adverse events. However, the certainty is low because of risk of bias and imprecision.
122	Low-dose radiation therapy	Uncertainty in potential benefits and harms. Further research is needed.



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	8Intervention	Summary of findings
123	Mavrilimumab	Uncertainty in potential benefits and harms. Further research is needed.
124	Mefenamic acid	Uncertainty in potential benefits and harms. Further research is needed.
125	Melatonin	Uncertainty in potential benefits and harms. Further research is needed.
126	Mesenchymal stem-cell transplantation	Mesenchymal stem-cell transplantation may reduce mortality. However, the certainty of the evidence is low. Further research is needed.
127	Metformin	Metformin may not reduce hospitalizations in patients with recent onset mild disease. However, certainty of the evidence is low because of imprecision. Further research is needed.
128	Methylene blue	Uncertainty in potential benefits and harms. Further research is needed.
129	Metisoprinol	Uncertainty in potential benefits and harms. Further research is needed.
130	Metoprolol	Uncertainty in potential benefits and harms. Further research is needed.
131	Metronidazole	Uncertainty in potential benefits and harms. Further research is needed.
132	Molnupiravir	In patients with recent onset mild COVID-19 molnupiravir reduces hospitalizations, it may improve symptom resolution and may not increase severe adverse events.





	8Intervention	Summary of findings
133	Montelukast	Uncertainty in potential benefits and harms. Further research is needed.
134	Mouthwash	Mouthwash may improve time to symptom resolution. Uncertainty in potential benefits and harms on other outcomes. Further research is needed.
135	Mupadolimab	Uncertainty in potential benefits and harms. Further research is needed.
136	Mycobacterium w	Uncertainty in potential benefits and harms. Further research is needed.
137	N-acetylcysteine	Uncertainty in potential benefits and harms. Further research is needed.
138	N-acetylcysteine (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
139	Nafamostat mesylate	Uncertainty in potential benefits and harms. Further research is needed.
140	Namilumab	Uncertainty in potential benefits and harms. Further research is needed.
141	Nano-curcumin	Uncertainty in potential benefits and harms. Further research is needed.
142	Nasal hypertonic saline	Uncertainty in potential benefits and harms. Further research is needed.
143	Neem (<i>Azadirachta indica</i> A. Juss)	Uncertainty in potential benefits and harms. Further research is needed.
144	Niclosamide	Uncertainty in potential benefits and harms. Further research is needed.



	8Intervention	Summary of findings
145	Nicotine patches	Uncertainty in potential benefits and harms. Further research is needed.
146	<i>Nigella sativa</i> +/- honey	Uncertainty in potential benefits and harms. Further research is needed.
147	Nirmatrelvir-ritonavir	Nirmatrelvir-ritonavir probably reduces hospitalizations in patients with mild recent onset COVID-19 and risk factors for severity, and it probably does not increase severe adverse events.
148	Nitazoxanide	Uncertainty in potential benefits and harms. Further research is needed.
149	Nitric oxide	Uncertainty in potential benefits and harms. Further research is needed.
150	Non-steroidal anti-inflammatory drugs (NSAIDs)	Current best evidence suggests no association between NSAIDs consumption and COVID-19 related mortality. However, the certainty of the evidence is very low because of the risk of bias. Further research is needed.
151	Norelgestromin and ethinylestradiol	Uncertainty in potential benefits and harms. Further research is needed.
152	Novaferon	Uncertainty in potential benefits and harms. Further research is needed.
153	Nutritional support	Uncertainty in potential benefits and harms. Further research is needed.
154	Omega-3 fatty acids	Uncertainty in potential benefits and harms. Further research is needed
155	OP-101	Uncertainty in potential benefits and harms. Further research is needed
156	Opaganib	Opaganib may not reduce mortality or mechanical ventilation, it may not increase severe adverse events but it may increase symptom resolution or improvement. Further research is needed.



	8Intervention	Summary of findings
157	Otilimab	Uncertainty in potential benefits and harms. Further research is needed
158	Ozone	Uncertainty in potential benefits and harms. Further research is needed.
159	P2Y12 inhibitors	P2Y12 inhibitors may increase mortality, may not improve time to symptom resolution and may increase severe adverse events. However, certainty of the evidence was low because of imprecision. Further research is needed.
160	Palmitoylethanolamide	Uncertainty in potential benefits and harms. Further research is needed.
161	Peg-interferon alfa	Uncertainty in potential benefits and harms. Further research is needed.
162	Peg-interferon lamda	Uncertainty in potential benefits and harms. Further research is needed.
163	Pembrolizumab	Uncertainty in potential benefits and harms. Further research is needed.
164	Pentoxifylline	Uncertainty in potential benefits and harms. Further research is needed.
165	Pirfenidone	Uncertainty in potential benefits and harms. Further research is needed.
166	Plitidepsin	Uncertainty in potential benefits and harms. Further research is needed.
167	PNB001 (CCK-A antagonist)	Uncertainty in potential benefits and harms. Further research is needed.
168	Polymerized type I collagen (PT1C)	Uncertainty in potential benefits and harms. Further research is needed.
169	Potassium canrenoate	Uncertainty in potential benefits and harms. Further research is needed.





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	8Intervention	Summary of findings
170	Povidone iodine (nasal spray)	Uncertainty in potential benefits and harms. Further research is needed.
171	Probiotics	Uncertainty in potential benefits and harms. Further research is needed.
172	Progesterone	Uncertainty in potential benefits and harms. Further research is needed
173	Prolectin-M	Uncertainty in potential benefits and harms. Further research is needed
174	Propolis	Uncertainty in potential benefits and harms. Further research is needed
175	Prostacyclin	Uncertainty in potential benefits and harms. Further research is needed
176	Prostacyclin (inhaled)	Uncertainty in potential benefits and harms. Further research is needed
177	Proxalutamide	Uncertainty in potential benefits and harms. Further research is needed
178	Pyridostigmine	Uncertainty in potential benefits and harms. Further research is needed
179	Quercetin	Uncertainty in potential benefits and harms. Further research is needed
180	Raloxifene	Uncertainty in potential benefits and harms. Further research is needed
181	Ramipril	Uncertainty in potential benefits and harms. Further research is needed.
182	RD-X19 (light therapy)	Uncertainty in potential benefits and harms. Further research is needed.



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	8Intervention	Summary of findings
183	Recombinant super-compound interferon	Uncertainty in potential benefits and harms. Further research is needed.
184	REGEN-COV (casirivimab and imdevimab)	In seronegative patients with severe to critical disease, REGEN-COV probably reduces mortality and increases symptom resolution and improvement. In patients with recent onset mild disease, REGEN-COV probably reduces hospitalizations and time to symptom resolution without increasing severe adverse events, and in asymptomatic exposed individuals REGEN-COV reduces symptomatic infections.
185	Regdanvimab	Regdanvimab may improve time to symptom resolution in mild to moderate patients. Its effects on mortality and mechanical ventilation are uncertain. Further research is needed.
186	Remdesivir	In hospitalized patients with moderate to critical disease, remdesivir probably reduces mortality and mechanical ventilation, and it may improve time to symptom resolution without increasing severe adverse events. In patients with recent onset mild COVID-19, it may reduce hospitalizations. However, the certainty is low because of risk of bias and imprecision.
187	Remdesivir (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
188	Reparixin	Uncertainty in potential benefits and harms. Further research is needed.
189	Resveratrol	Uncertainty in potential benefits and harms. Further research is needed.
190	rhG-CSF (in patients with lymphopenia)	Uncertainty in potential benefits and harms. Further research is needed.
191	rhG-CSF (inhaled)	Uncertainty in potential benefits and harms. Further research is needed.
192	rhu-pGSN	Uncertainty in potential benefits and harms. Further research is needed.
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	8Intervention	Summary of findings
193	Ribavirin	Uncertainty in potential benefits and harms. Further research is needed.
194	Ribavirin + interferon beta-1b	Uncertainty in potential benefits and harms. Further research is needed.
195	Ruxolitinib	Ruxolitinib may reduce mortality; however, the certainty of the evidence was low. Further research is needed.
196	Sabizabulin	Uncertainty in potential benefits and harms. Further research is needed.
197	Sarilumab	Sarilumab may not reduce mortality and probably does not improve time to symptom resolution but may decrease mechanical ventilation requirements without increasing severe adverse events. However, the certainty is low because of imprecision and inconsistency.
198	Secukinumab	Uncertainty in potential benefits and harms. Further research is needed.
199	Senicapoc	Uncertainty in potential benefits and harms. Further research is needed.
200	Sentinox	Uncertainty in potential benefits and harms. Further research is needed.
201	Short-wave diathermy	Uncertainty in potential benefits and harms. Further research is needed.
202	Sildenafil	Uncertainty in potential benefits and harms. Further research is needed.
203	Siltuximab	Uncertainty in potential benefits and harms. Further research is needed.
204	Silymarin	Uncertainty in potential benefits and harms. Further research is needed.
205	Sitagliptin	Uncertainty in potential benefits and harms. Further research is needed.



	8Intervention	Summary of findings
206	Sofosbuvir +/- daclatasvir, ledipasvir, velpatasvir, or ravidasvir	Sofosbuvir with or without daclatasvir or ledipasvir may increase mortality and not reduce mechanical ventilation requirements, and it probably does not improve time to symptom resolution. Further research is needed to confirm these findings.
207	Sotrovimab	Sotrovimab may probably reduce hospitalizations in patients with recent onset mild COVID-19.
208	Spironolactone	Uncertainty in potential benefits and harms. Further research is needed.
209	Statins	Statins may reduce mortality; however, certainty of the evidence was low. Further research is needed.
210	Stem-cell nebulization	Uncertainty in potential benefits and harms. Further research is needed.
211	Steroids (corticosteroids)	Corticosteroids reduce mortality and probably reduce invasive mechanical ventilation requirements in patients with severe COVID-19 infection with moderate certainty. Corticosteroids may not significantly increase the risk of severe adverse events. Higher-dose schemes (i.e., dexamethasone 12 mg a day) may not be more effective than standard dose schemes (i.e., dexamethasone 6 mg a day).
212	Steroids (corticosteroids, inhaled)	Inhaled corticosteroids may improve time to symptom resolution but probably does not have an important effect on hospitalizations. Its effects on other important outcomes are uncertain. Further research is needed.
213	Steroids (corticosteroids, nasal)	Uncertainty in potential benefits and harms. Further research is needed.
214	Sulodexide	Uncertainty in potential benefits and harms. Further research is needed.
215	Tafenoquine	Uncertainty in potential benefits and harms. Further research is needed.



	8Intervention	Summary of findings
216	TD-0903 (inhaled JAK-inhibitor)	Uncertainty in potential benefits and harms. Further research is needed.
217	Tenofovir + emtricitabine	Tenofovir + emtricitabine may not reduce mortality but may reduce mechanical ventilation. However, certainty of the evidence was low. Further research is needed.
218	Thalidomide	Uncertainty in potential benefits and harms. Further research is needed.
219	Thymoquinone	Uncertainty in potential benefits and harms. Further research is needed.
220	Tissue-plasminogen activator (tPA)	Uncertainty in potential benefits and harms. Further research is needed.
221	Tixagevimab–cilgavimab	Tixagevimab-cilgavimab probably reduces mortality, hospitalizations, and SARS-COV-2 infections in exposed individuals and may not increase severe adverse events.
222	Tocilizumab	Tocilizumab reduces mortality and reduces mechanical ventilation requirements without possibly increasing severe adverse events.
223	Tofacitinib	Tofacitinib may increase symptom resolution or improvement and severe adverse events. Certainty of the evidence was low. Further research is needed.
224	Tranilast	Uncertainty in potential benefits and harms. Further research is needed.
225	Triazavirin	Uncertainty in potential benefits and harms. Further research is needed.
226	TXA-127	Uncertainty in potential benefits and harms. Further research is needed.



	8Intervention	Summary of findings
227	Ultraviolet light phototherapy	Uncertainty in potential benefits and harms. Further research is needed.
228	Umifenovir	Uncertainty in potential benefits and harms. Further research is needed.
229	Verapamil	Uncertainty in potential benefits and harms. Further research is needed.
230	Vilobelimab	Vilobelimab probably reduces mortality and probably does not increase severe adverse events.
231	Vitamin B	
232	Vitamin C	Vitamin C may increase symptom resolution or improvement. Its effects on other clinical important outcomes are uncertain. Further research is needed.
233	Vitamin D	Vitamin D does no reduce infections in exposed individuals and probably not reduce hospitalizations. Vitamin D effect on other important outcomes is uncertain. Further research is needed.
234	XAV-19 (swine glyco-humanized polyclonal antibodies)	Uncertainty in potential benefits and harms. Further research is needed.
235	Zilucoplan	Uncertainty in potential benefits and harms. Further research is needed.
236	Zinc	Zinc may not improve symptom resolution. However, the certainty of the evidence was low because of imprecision. Its effects on other clinical important outcomes are uncertain. Further research is needed.
237	α-lipoic acid	Uncertainty in potential benefits and harms. Further research is needed.



Key findings

• **Therapeutic options:** According to WHO International Clinical Trials Registry Platform (ICTRP), hundreds of potential interventions are being assessed in more than 10,000 clinical trials and observational studies. In this review, we identified and examined 237 therapeutic options.

• **Corticosteroids:** The body of evidence on corticosteroids, which includes 24 RCTs, shows that low- or moderate-dose treatment schemes (RECOVERY trial dose was 6 mg of oral or intravenous preparation once daily for 10 days) are probably effective in reducing mortality in patients with severe COVID-19 infection. These results remained robust after including studies in which patients with acute respiratory distress syndrome (ARDS) secondary to alternative etiologies (not COVID-19 related) were randomized to corticosteroids or placebo/no corticosteroids. Higher-dose schemes (i.e., dexamethasone 12 mg a day) may not be more effective than standard dose schemes (i.e., dexamethasone 6 mg a day).

• **Remdesivir:** The results of 10 RCTs, including the final results of the SOLIDARITY trial, show that in hospitalized patients with moderate to critical disease, remdesivir probably reduces mortality and mechanical ventilation, and it may improve time to symptom resolution. Certainty of the evidence was moderate because of imprecision. In patients with recent onset mild COVID-19 remdesivir may reduce hospitalizations; however, the certainty of the evidence is low because of imprecision. Further research is needed.

• Hydroxychloroquine, lopinavir–ritonavir, and interferon beta-1a: The body of evidence on hydroxychloroquine, lopinavir-ritonavir, and interferon beta-1a, including anticipated findings from the RECOVERY and SOLIDARITY trials, showed no benefit in terms of mortality reduction, invasive mechanical ventilation requirements or time to clinical improvement. Furthermore, the analysis showed probable mortality increment in those patients treated with hydroxychloroquine. Seven studies with low risk of bias that assessed hydroxychloroquine in exposed individuals showed a modest reduction in symptomatic infections, but certainty of the evidence was low because of imprecision and inconsistency. Further research is needed to confirm these findings.

• Antibiotics: The body of evidence on azithromycin and doxycycline shows no significant benefits in patients with mild to moderate or severe to critical COVID-19.

• **Convalescent plasma:** The results of 58 RCTs assessing convalescent plasma in COVID-19, including the RECOVERY trial with 11,558 hospitalized patients, showed no mortality reduction, significant mechanical ventilation requirement reduction or time to symptom resolution improvement with moderate to high certainty of the evidence. In mild patients, convalescent plasma probably does not have an important effect on hospitalizations with moderate certainty. Convalescent plasma may not increase severe adverse events with low certainty. No significant



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differences were observed between patients treated early (< 4 days since symptom onset) or with more advanced disease in a subgroup analysis from the RECOVERY trial.

• **Tocilizumab:** The results of 28 RCTs assessing tocilizumab show that, in patients with severe or critical disease, tocilizumab reduces mortality and mechanical ventilation requirements without significantly increasing severe adverse events.

• **Clazakizumab:** The results of one RCT suggests that, in patients with severe or critical disease, clazakizumab may mechanical ventilation requirements and improve time to symptom resolution. However, certainty of the evidence was low because of imprecision. Further research is needed.

• Sarilumab: The results of 10 RCTs assessing sarilumab show that, in patients with severe or critical disease, sarilumab may not reduce mortality and probably does not improve time to symptom resolution but may reduce mechanical ventilation requirements without significantly increasing severe adverse events. However, certainty of the evidence was low and further research is needed to confirm these findings.

• Anakinra: The results of six RCTs assessing anakinra in hospitalized patients with non-severe disease, show inconsistent results on mortality and symptom resolution and suggest that anakinra may not reduce mortality or increase severe adverse events. Certainty of the evidence was low and further research is needed.

• **Tofacitinib:** The results of two RCTs assessing tofacitinib in hospitalized patients with moderate to severe disease, suggest possible increase in symptom resolution or improvement and possible increase in severe adverse events with tofacitinib. Certainty of the evidence was low and further research is needed.

• Vilobelimab: The results of two RCTs assessing vilobelimab show that, in patients with severe or critical disease, vilobelimab probably reduces mortality without significantly increasing severe adverse events.

• **Colchicine:** The results of 15 RCTs assessing colchicine, including the COLCORONA study that recruited 4,488 patients with recent COVID-19 diagnosis and risk factors for severity and the RECOVERY trial that recruited 11,340 hospitalized patients, show that colchicine probably does not reduce mortality, mechanical ventilation requirements, improve time to symptom resolution, or reduce hospitalizations. These findings are mainly driven by the RECOVERY study. The COLCORONA study that included outpatients with mild early COVID-19 suggest possible reduction in hospitalizations, mechanical ventilation requirements and mortality in this subgroup. However, certainty of the evidence was low because of very severe imprecision due to a small number of events.

• **Ivermectin:** Pooled estimates of 49 RCTs suggest mortality reduction with ivermectin, but the certainty of the evidence was very low because of methodological limitations and small number of events. Based on the results reported by the subgroup RCTs classified as low risk of bias,



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ivermectin probably does not reduce mortality or improve time to symptom resolution, and probably does not have an important effect on hospitalizations. Further research is needed to confirm these findings.

• **Favipiravir:** Twenty-nine RCTs assessed favipiravir vs SOC or other interventions. Their results suggest that favipiravir may increase mortality and mechanical ventilation requirements, it may not reduce hospitalizations and it does not improve symptom resolution. Further research is needed to confirm these findings.

• **Sofosbuvir** +/- **daclatasvir**, **ledipasvir**, **velpatasvir**, **or ravidasvir**: Sixteen RCTs assessed sofosbuvir with or without daclatasvir, ledipasvir, or velpatasvir against standard of care or other interventions. Subgroup analysis showed significant differences between low risk of bias and high risk of bias studies. The results of the two studies classified as low risk of bias suggest that sofosbuvir alone or in combination may increase mortality and not reduce mechanical ventilation requirements, and it probably does not improve time to symptom resolution. Further research is needed to confirm these findings.

• **Tenofovir** + **emtricitabine:** Five RCTs assessed tenofovir + emtricitabine against standard of care or other interventions. Their results suggest that tenofovir + emtricitabine may not reduce mortality and may decrease mechanical ventilation requirements. However, certainty of the evidence was low because of imprecision and risk of bias. Further research is needed to confirm these findings.

• **Baricitinib:** The results of seven RCTs show that, in patients with moderate to critical disease, baricitinib reduces mortality, probably reduces mechanical ventilation requirements, and probably improves time to symptom resolution, without increasing severe adverse events.

• **Ruxolitinib:** The results of three RCTs show that, in patients with moderate to critical disease, ruxolitinib may reduce mortality. However, the certainty of the evidence was low because of imprecision and inconsistency. Further research is needed.

• CD24Fc (soluble CD24 appended to heavy chains 2 and 3 of human immunoglobulin G1): The results of one RCT shows that in patients with severe disease, CD24Fc may reduce mechanical ventilation and increase symptom resolution. However, the certainty of the evidence was low because of imprecision. Further research is needed.

• **REGEN-COV** (casirivimab and imdevimab): The results of 12 RCTs suggest that, in patients with severe to critical disease, overall REGEN-COV may reduce mortality and mechanical ventilation, or increase symptom resolution or improvement. However, the certainty of the evidence was low. A subgroup analysis suggests a differential effect on seronegative patients in which REGEN-COV probably reduces mortality and mechanical ventilation requirements and increases symptom resolution or improvement. In patients with recent onset mild COVID-19, REGEN-COV probably reduces hospitalizations and improves time to symptom resolution without increasing severe adverse events, and in exposed asymptomatic individuals REGEN-COV



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reduces symptomatic infections. One study that compared REGEN-COV (casirivimab and imdevimab) against bamlanivimab +/- etesevimab in non-severe patients with risk factors for severity, reported no important differences in hospitalizations.

• **Bamlinivimab** +/- **etesevimab:** The results of six RCTs suggest that bamlinivimab probably decreases hospitalizations in patients with COVID-19 and probably decreases symptomatic infection in exposed individuals. Its effects on other clinical important outcomes are uncertain. Further research is needed. One study that compared bamlanivimab +/- etesevimab against REGEN-COV (casirivimab and imdevimab) in non-severe patients with risk factors for severity, reported no important differences in hospitalizations.

• **Sotrovimab:** The results of two RCTs show that, in patients with recent onset mild COVID-19, sotrovimab probably reduces hospitalizations and improves time to symptom resolution without increasing severe adverse events. The certainty of the evidence was moderate because of imprecision but with evidence of equipoise between sotrovimab and REGEN-COV.

• **Regdanvimab:** The results of two RCTs show that, in patients with mild to moderate disease, regdanvimab may improve time to symptom resolution. However, the certainty of the evidence was low because of imprecision. Its effects on other important outcomes are uncertain. Further research is needed to confirm or discard these findings.

• **Tixagevimab–cilgavimab:** The results of three RCTs show that, in individuals with COVID-19, tixagevimab–cilgavimab probably reduces mortality and hospitalizations, and in those exposed to SARS-COV-2 tixagevimab–cilgavimab probably reduces symptomatic infections without increasing severe adverse events.

• **Proxalutamide:** The results of four RCTs suggest that proxalutamide may result in important benefits. However, the certainty of the evidence was very low because of very serious risk of bias, imprecision, and indirectness. Further research is needed to confirm or discard these findings.

• **Dapagliflozin:** The results of one RCT suggests that, in patients with cardiometabolic risk factors hospitalized with moderate COVID-19, dapagliflozin may reduce mortality, but probably does not increase symptom resolution. However, the certainty of the evidence was low because of imprecision. Further research is needed to confirm or discard these findings.

• **Mesenchymal stem-cell transplantation:** The results of nine RCTs show that, in patients with severe to critical, mesenchymal stem-cell transplantation may reduce mortality. However, the certainty of the evidence was low because of imprecision. Further research is needed to confirm or discard these findings.

• **Inhaled corticosteroids:** The results of nine RCTs show that inhaled corticosteroids may improve time to symptom resolution but probably does not have an important effect on hospitalizations. Its effects on other relevant outcomes are uncertain. Further research is needed.





• **Fluvoxamine:** The results of four RCTs suggest that in patients with mild disease, fluvoxamine probably does not have an important effect on hospitalizations and may not increase adverse events. The certainty of the evidence was moderate to low because of imprecision. Further research is needed.

• Lenzilumab: The results of one RCT suggests that lenzilumab may reduce invasive mechanical ventilation requirements in severe patients without increasing severe adverse events. However, the certainty of the evidence was low because of imprecision. Further research is needed.

• **INM005 (polyclonal fragments of equine antibodies):** Currently, there is very low certainty about the effects of INM005 on clinically important outcomes.

• **Famotidine:** Currently, there is very low certainty about the effects of famotidine on clinically important outcomes.

• Anticoagulants: Thromboembolic complications in patients infected with COVID-19 are relatively frequent. As for hospitalized patients with severe medical conditions current guidelines recommend thromboprophylactic measures to be adopted for inpatients with COVID-19 infection. Regarding the best thromboprophylactic scheme, excluding four studies classified as with high risk of bias, the results of ten RCTs that compared anticoagulants in intermediate (i.e., enoxaparin 1 mg/kg a day) or full dose (i.e., enoxaparin 1 mg/kg twice a day) versus prophylactic dose (i.e., enoxaparin 40 mg a day) showed no differences in mortality with moderate certainty (imprecision). In mild ambulatory patients four RCTs suggest that rivaroxaban or enoxaparin in prophylactic dose may not importantly improve time to symptom resolution or reduce hospitalizations.

• Aspirin: Results of four RCTs inform that aspirin probably does not reduce mortality or mechanical ventilation and probably does not increase symptom resolution or improvement.

• **P2Y12 inhibitors:** The results of two RCTs suggest that P2Y12 in combination with anticoagulants in prophylactic or full dose may not reduce mortality, may not improve time to symptom resolution, and may increase severe adverse events. However, the certainty of the evidence was low because of imprecision and the effects on other important outcomes are uncertain. Further research is needed.

• **NSAIDs:** No association between NSAIDs exposure and increased mortality was observed. However, certainty of the evidence is very low and further research is needed to confirm these findings.

• ACEIs or ARBs: The results of eight low-risk of bias RCTs suggest that initiating or continuing ACEIs or ARBs in patients with COVID-19 may increase mortality. However, certainty of the evidence is low because of imprecision and further research is needed to confirm these findings.

• **Molnupiravir:** The results of 10 RCTs show that molnupiravir reduces hospitalizations in patients with recent onset mild to moderate disease, and may not increase severe adverse events.





• **Nirmatrelvir-ritonavir:** The results of one RCT shows that nirmatrelvir-ritonavir probably reduces hospitalizations in patients with recent onset mild to moderate disease, and probably does not increase severe adverse events.

• Vitamin D: The results of 20 RCTs show that vitamin D does not reduce symptomatic infections and probably does not reduce hospitalizations. Vitamin D effects on other important outcomes are uncertain. Further research is needed.

• Vitamin C: The results of nine RCTs suggest that vitamin C may increase symptom resolution or improvement. However, the certainty of the evidence was low and vitamin C effects on other important outcomes are uncertain. Further research is needed.

• **Probiotics:** The results of four RCTs suggest that probiotics may improve time to symptom resolution. However, the certainty of the evidence was low because of imprecision and the effects on other important outcomes are uncertain. Further research is needed.

• **Mouthwash:** The results of 14 RCTs suggest that mouthwashes may improve time to symptom resolution. However, the certainty of the evidence was low because of imprecision and the effects on other important outcomes are uncertain. Further research is needed.

• **Camostat mesilate:** The results of five RCTs suggest that camostat mesilate may not improve time to symptom resolution. However, the certainty of the evidence was low because of imprecision and indirectness, furthermore the effects on other important outcomes are uncertain. Further research is needed.

• **Opaganib:** The results of two RCTs suggest that opaganib may not reduce mortality or mechanical ventilation, it may not increase severe adverse events but it may increase symptom resolution or improvement. However, certainty of the evidence was low because of imprecision. Further research is needed.

Changes since previous edition

• **Prifenidone:** New evidence included affecting results interpretation and/or certainty of the evidence judgments.

- Peg-interferon (IFN) lamda: New evidence included without significant changes.
- Mesenchymal stem-cell transplantation: New evidence included without significant changes.
- Tissue plasminogen activator (tPA): New evidence included without significant changes.
- Anticoagulants: New evidence included without significant changes.



• Vitamin D: New evidence included affecting results interpretation and/or certainty of the evidence judgments.

• Favipitavir: New evidence included affecting results interpretation and/or certainty of the evidence judgments.

• Vilobelimab: New evidence included affecting results interpretation and/or certainty of the evidence judgments.

• **Palmitoylethanolamide:** New evidence included affecting results interpretation and/or certainty of the evidence judgments.

• AZD1656: New evidence included affecting results interpretation and/or certainty of the evidence judgments.

• **DFV890:** New evidence included affecting results interpretation and/or certainty of the evidence judgments.

• Vitamin B: New evidence included without significant changes.

• Convalescent plasma: New evidence included without significant changes.





Concluding remarks

• The Pan American Health Organization (PAHO) is continually monitoring ongoing research on any possible therapeutic options. As evidence emerges, then PAHO will immediately assess and update its position, particularly as it applies to any special subgroup populations such as children, expectant mothers, and those with immune conditions.

• PAHO is also mindful of the emerging differential impact of COVID-19 on ethnic and minority groups and is continuously seeking data that could help in mitigating excess risk of severe illness or death in minority subgroups. These groups are plagued by social and structural inequities that bring to bear a disproportionate burden of COVID-19 illness.

• The safety of the patient suffering from COVID-19 is a key priority to improve the quality of care in the provision of health services.

• There remains an urgent need for additional high-quality randomized controlled trials that include patients with COVID-19 before most therapeutic options can be administered with any confidence. Adequately designed and reported clinical trials are crucial for the practice of evidence-based medicine. Most of the research to date on COVID-19 has very poor methodology that is hidden and very difficult to validate. Greater transparency and better designed studies are urgently needed.





Hallazgos clave

Opciones terapéuticas: Según el portal de búsqueda de la Plataforma de Registros Internacionales de Ensayos Clínicos de la Organización Mundial de la Salud, se están investigando cientos de posibles tratamientos o sus combinaciones en más de 10.000 ensayos clínicos y estudios observacionales. En esta revisión, examinamos 237 opciones terapéuticas potenciales.

• **Corticosteroides:** El conjunto de evidencia sobre los corticoesteroides incluye 24 ensayos clínicos controlados aleatorizados (ECCA) y muestra que la administración de dosis bajas y moderadas (la dosis utilizada en el estudio RECOVERY fue de 6 mg diarios de dexametasona por vía oral o intravenosa durante 10 días) probablemente reduce la mortalidad en pacientes con infección grave por SARS-CoV-2. Los resultados se mantuvieron uniformes tras agregar al análisis estudios en los que pacientes con síndrome de dificultad respiratoria aguda de otras etiologías recibieron corticosteroides o manejo estándar de forma aleatoria. Esquemas con dosis más altas (por ejemplo, 12 mg de dexametasona por día) podrían no resultar más efectivos que los esquemas habituales (por ejemplo, 6 mg de dexametasona por día).

• **Remdesivir:** Los resultados de 10 ECCA, incluidos los resultados finales del ensayo Solidaridad, muestran que en pacientes hospitalizados con enfermedad de moderada a critica, el remdesivir probablemente reduce la mortalidad y la necesidad de ventilación mecánica invasiva, y podría mejorar el tiempo de resolución de los síntomas. La certeza de la evidencia es moderada por imprecisión. En pacientes con enfermedad leve de comienzo reciente, el remdesivir podría reducir las hospitalizaciones, pero la certeza de la evidencia es baja por imprecisión. Se necesita más información.

• Hidroxicloroquina, interferón beta 1-a y lopinavir-ritonavir: El conjunto de evidencia sobre la hidroxicloroquina, el interferón beta 1-a y el lopinavir-ritonavir, incluidos los resultados preliminares de los estudios RECOVERY y Solidaridad, no muestra beneficios en la reducción de la mortalidad, la necesidad de ventilación mecánica invasiva o el plazo necesario para la mejoría clínica. Incluso la evidencia sobre la hidroxicloroquina sugiere que su utilización probablemente genere un incremento en la mortalidad. Siete estudios con riesgo bajo de sesgo que evaluaron la hidroxicloroquina en personas expuestas a la COVID-19 sugieren una reducción modesta del riesgo de infección, pero la certeza de la evidencia es baja por inconsistencia (falta de congruencia (*inconsistency*) e imprecisión. Se necesita más información para confirmar estas conclusiones.

• Antibióticos: El conjunto de evidencia identificado sobre la azitromicina y la doxiciclina no muestra beneficios significativos en pacientes con COVID-19 de leve a moderada, o grave a crítica.

• **Plasma de convalecientes:** Los resultados de 58 ECCA que evaluaron el uso de plasma de convalecientes en pacientes con COVID-19, incluido el estudio RECOVERY que incorpora 11.558 pacientes, no mostraron reducción de la mortalidad, disminución de la necesidad de ventilación mecánica invasiva ni mejoría en el tiempo de resolución de los síntomas con certeza



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moderada. En pacientes leves, el plasma de convalecientes probablemente no tenga ningún efecto importante sobre las hospitalizaciones con certeza moderada. El plasma de convalecientes podría no asociarse a un aumento de los eventos adversos graves con certeza baja. En un análisis de subgrupo del estudio RECOVERY, no se observó ningún efecto diferencial entre los pacientes tratados con rapidez (menos de 4 días desde el inicio de los síntomas) y los que presentaban enfermedad más avanzada al iniciar dicho tratamiento.

• **Tocilizumab:** Los resultados de 28 ECCA muestran que el tocilizumab reduce la mortalidad y la necesidad de ventilación invasiva sin un incremento importante de los efectos adversos graves en pacientes con enfermedad grave o crítica.

• **Clazakizumab:** Los resultados de un ECCA sugieren que el clazakizumab podría reducir la necesidad de ventilación mecánica invasiva y mejorar el tiempo de resolución de los síntomas. Sin embargo, la certeza en la evidencia es baja y se necesita más información.

• Sarilumab: Los resultados de diez ECCA muestran que el sarilumab podría no reducir la mortalidad y probablemente no mejore el tiempo de resolución de los síntomas, aunque sí podría reducir la necesidad de ventilación invasiva sin un incremento importante de los efectos adversos graves en pacientes con enfermedad grave o crítica. Sin embargo, la certeza de la evidencia es baja y se necesita más información para confirmar estas conclusiones.

• Anakinra: Los resultados de seis ECCA que evaluaron la anakinra en pacientes hospitalizados con enfermedad no grave muestran resultados incongruentes en la mortalidad y la resolución de los síntomas y sugieren que podría no reducir la mortalidad ni aumentar los eventos adversos graves. La certeza de la evidencia es baja y se necesita más información.

• **Tofacitinib:** Los resultados dos ECCA que evaluaron el tofacitinib en pacientes hospitalizados con enfermedad de moderada a grave indican una posible mejora de la resolución de los síntomas, aunque con un posible aumento de los eventos adversos graves. La certeza de la evidencia es baja y se necesita más información.

• Vilobelimab: Los resultados de 2 ECCA muestran que el vilobelimab probablemente reduzca la mortalidad sin un incremento importante de los efectos adversos graves en pacientes con enfermedad grave o crítica.

• Colchicina: Los resultados de quince ECCA —entre los que se encuentra el estudio COLCORONA, que incluyó 4488 pacientes con diagnóstico reciente de COVID-19 y factores de riesgo para enfermedad grave, y el estudio RECOVERY, que incorpora 11.340 pacientes hospitalizados— muestran que la colchicina probablemente no reduzca la mortalidad o la necesidad de ventilación mecánica, no mejore la velocidad de resolución de los síntomas ni reduzca las hospitalizaciones. Estos resultados se sustentan fundamentalmente en el estudio RECOVERY. El estudio COLCORONA, que incluyó pacientes ambulatorios con enfermedad leve, apunta una posible reducción de las hospitalizaciones, de la necesidad de ventilación



mecánica y de la mortalidad en este subgrupo. Sin embargo, la certeza de la evidencia es baja por imprecisión muy grave, ya que el número de eventos fue reducido.

• **Ivermectina:** Los resultados combinados de 49 ECCA indican una reducción de la mortalidad con la ivermectina. Sin embargo, la certeza de la evidencia es muy baja por limitaciones metodológicas y un número de eventos reducido. Con base en la información facilitada por los estudios con riesgo bajo de sesgo, la ivermectina probablemente no reduzca la mortalidad ni se asocie a una mejoría en el tiempo de resolución de los síntomas, ni tenga un efecto importante sobre las hospitalizaciones. Se necesita más información para confirmar estas conclusiones.

• Favipiravir: Veintinueve ECCA evaluaron el favipiravir en comparación con la prestación de cuidados estándares u otras intervenciones. Los resultados sugieren que el favipiravir podría aumentar la mortalidad y la necesidad de ventilación invasiva mecánica, podría no reducir las hospitalizaciones y no mejora la resolución de los síntomas. Se necesita más información para confirmar estas conclusiones.

• Sofosbuvir con o sin daclatasvir, ledipasvir, velpatasvir o ravidasvir: Dieciséis ECCA evaluaron el sofosbuvir solo o en combinación con daclatasvir, ledipasvir o velpatasvir en comparación con la prestación de cuidados estándares u otras intervenciones. Los resultados de los estudios con un riesgo alto de sesgo y de los estudios con un riesgo bajo de sesgo fueron sustancialmente diferentes. Los resultados de los dos estudios clasificados con riesgo bajo de sesgo sugieren que el sofosbuvir solo o en combinación podría aumentar la mortalidad y no reducir la necesidad de ventilación invasiva mecánica, y probablemente no mejore el tiempo de resolución de los síntomas. Se necesita más información para confirmar estas conclusiones.

• **Tenofovir y emtricitabina:** Los resultados de cinco ECCA sugieren que el tenofovir y la emtricitabina podrían no reducir la mortalidad, pero probablemente reduzcan la necesidad de ventilación mecánica invasiva. Sin embargo, la certeza de la evidencia es baja por imprecisión y riesgo de sesgo. Se necesita más información para confirmar estas conclusiones.

• **Baricitinib:** Los resultados de siete ECCA muestran que, en pacientes con enfermedad de moderada a crítica, el baricitinib reduce la mortalidad, y probablemente reduzca la necesidad de ventilación mecánica invasiva y mejore el tiempo de resolución de síntomas sin aumentar los eventos adversos graves.

• **Ruxolitinib:** Los resultados de tres ECCA sugieren que, en pacientes con enfermedad de moderada a grave, el ruxolitinib podría reducir la mortalidad. Sin embargo, la certeza de la evidencia es baja por inconsistencia (falta de congruencia) e imprecisión. Se necesita más información.

• CD24Fc (cadenas pesadas 2 y 3 de inmunoglobulina humana G1 anexadas a CD24): Los resultados de un ECCA muestran que, en pacientes con enfermedad grave, el CD24Fc podría reducir la necesidad de ventilación mecánica invasiva y mejorar la resolución de síntomas. Sin embargo, la certeza de la evidencia es baja por imprecisión. Se necesita más información.



• **REGEN-COV** (casirivimab e imdevimab): Los resultados de 12 ECCA muestran que, en pacientes con enfermedad grave o crítica, el REGEN-COV podría reducir la mortalidad y la necesidad de ventilación mecánica invasiva y mejorar la velocidad de resolución de los síntomas. Sin embargo, la certeza de la evidencia es baja. Un análisis de subgrupo mostró un efecto diferencial en pacientes con anticuerpos negativos. En este subgrupo, el REGEN-COV probablemente reduzca la mortalidad y la necesidad de ventilación mecánica e incremente la resolución de los síntomas. En pacientes con enfermedad leve de comienzo reciente, el REGEN-COV probablemente reduzca las hospitalizaciones y mejore el tiempo de resolución de los síntomas sin aumentar el riesgo de eventos adversos graves; y en personas asintomáticas, expuestas a SARS-CoV-2, el REGEN-COV reduce las infecciones sintomáticas. La certeza de la evidencia es alta para infecciones sintomáticas y de baja a moderada por información indirecta e imprecisión para los desenlaces restantes. Un estudio que comparó el REGEN-COV (casirivimab e imdevimab) con el bamlanivimab con o sin etesevimab en pacientes con síntomas leves y factores de riesgo para enfermedad grave notificó ausencia de diferencias importantes en las hospitalizaciones.

• **Bamlinivimab con o sin etesevimab:** Los resultados de seis ECCA indican que el bamlanivimab probablemente reduzca las hospitalizaciones en pacientes con COVID-19 y probablemente disminuya las infecciones sintomáticas en personas expuestas. Sus efectos sobre otros desenlaces importantes son inciertos. Se necesita más información. Un estudio que comparó el bamlanivimab con o sin etesevimab con el REGEN-COV (casirivimab e imdevimab) en pacientes con síntomas leves y factores de riesgo para enfermedad grave notificó ausencia de diferencias importantes en las hospitalizaciones.

• **Sotrovimab:** Los resultados de dos ECCA muestran que, en pacientes con enfermedad leve de comienzo reciente, el sotrovimab probablemente reduzca las hospitalizaciones y mejore el tiempo de resolución de los síntomas sin aumentar el riesgo de eventos adversos graves. La certeza de la evidencia es moderada por imprecisión, pero incluye hallazgos de eficacia similar entre el sotrovimab y el REGEN-COV.

• **Regdanvimab:** Los resultados de dos ECCA muestran que, en pacientes con enfermedad leve a moderada, el regdanivimab podría mejorar el tiempo de resolución de los síntomas. Sin embargo, la certeza de la evidencia es baja por imprecisión. Sus efectos sobre otros desenlaces importantes son inciertos. Se necesita más información para confirmar o descartar estas conclusiones.

• **Tixagevimab y cilgavimab:** Los resultados de tres ECCA muestran que el tixagevimab y el cilgavimab probablemente reduzcan la mortalidad, las hospitalizaciones y las infecciones sintomáticas en personas expuestas al SARS-CoV-2 y podrían no aumentar los eventos adversos graves.

• **Proxalutamida:** Los resultados de cuatro ECCA sugieren un efecto favorable asociado a la proxalutamida. Sin embargo, la certeza de la evidencia es muy baja por riesgo muy grave de sesgo, imprecisión e información indirecta. Se necesita más información para confirmar o descartar estas conclusiones.



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• **Dapagliflozina:** Los resultados de un ECCA muestran que, en pacientes con factores de riesgo cardiometabólicos hospitalizados por COVID-19 moderada, la dapagliflozina podría reducir la mortalidad, pero probablemente no mejore la resolución de los síntomas. Sin embargo, la certeza de la evidencia es baja por imprecisión. Se necesita más información para confirmar o descartar estas conclusiones.

• **Trasplante de células madre mesenquimatosas:** Los resultados de ocho ECCA apuntan que, en pacientes con enfermedad de grave a crítica, el trasplante de células madre mesenquimatosas podría reducir la mortalidad. Sin embargo, la certeza de la evidencia es baja por imprecisión. Se necesita más información para confirmar o descartar estas conclusiones.

• **Corticosteroides inhalados:** Los resultados de nueve ECCA muestran que los corticosteroides inhalados podrían mejoran el tiempo de resolución de los síntomas, pero probablemente no afecten las hospitalizaciones de forma considerable. Sus efectos sobre otros desenlaces importantes son inciertos. Se necesita más información.

• Fluvoxamina: Los resultados de cuatro ECCA sugieren que, en pacientes con enfermedad leve, la fluvoxamina probablemente no tenga un efecto importante sobre las hospitalizaciones y podría no incrementar los eventos adversos. La certeza de la evidencia es de baja a moderada por imprecisión. Se necesita más información.

• Lenzilumab: Los resultados de un ECCA sugieren que el lenzilumab podría reducir la necesidad de ventilación mecánica invasiva en pacientes graves sin aumentar los eventos adversos graves. Sin embargo, la certeza de la evidencia es baja por imprecisión. Se necesita más información.

• **INM005 (fragmentos policionales de anticuerpos equinos):** Por el momento, la certeza de la evidencia sobre los efectos del INM005 en desenlaces críticos es muy baja.

• Famotidina: Por el momento, la certeza de la evidencia sobre los efectos de la famotidina en desenlaces clínicamente importantes es muy baja.

• Anticoagulantes: Las complicaciones tromboembólicas en pacientes con COVID-19 son relativamente frecuentes. Al igual que en pacientes hospitalizados por afecciones médicas graves, las directrices vigentes indican que los pacientes hospitalizados por COVID-19 sean tratados con medidas tromboprofilácticas. En relación con el mejor esquema tromboprofiláctico, excluyendo cuatro estudios clasificados con riesgo alto de sesgo, los resultados de diez ECCA que compararon los anticoagulantes en dosis intermedias (p. ej., 1 mg/kg de enoxaparina por día) o dosis completas (p. ej., 1 mg/kg de enoxaparina cada 12 h por día) frente a dosis profilácticas (p. ej., 40 mg de enoxaparina por día) no mostraron diferencias en la mortalidad con certeza moderada (imprecisión). Los resultados de cuatro ECCA sugieren que, en pacientes ambulatorios con enfermedad leve, el rivaroxabán o la enoxaparina en dosis profilácticas podría no mejorar el tiempo de resolución de los síntomas de forma considerable ni reducir las hospitalizaciones.



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• Aspirina: Los resultados de cuatro ECCA informan que la aspirina probablemente no reduzca la mortalidad o la necesidad de ventilación mecánica ni mejore la velocidad de resolución de los síntomas.

• Inhibidores P2Y12: Los resultados de dos ECCA sugieren que el tratamiento con P2Y12 combinado con anticoagulantes en dosis profilácticas o completas podría no reducir la mortalidad ni mejorar el tiempo de resolución de los síntomas, y podría aumentar los eventos adversos graves. Sin embargo, la certeza de la evidencia es baja y los efectos sobre otros desenlaces importantes son inciertos. Se necesita más información.

• Antiinflamatorios no esteroideos (AINE): Hasta el momento, el uso de los AINE no está asociado con un incremento de la mortalidad. Sin embargo, la certeza de la evidencia es muy baja, por lo que se necesita más información para confirmar estas conclusiones.

• **IECA y ARB:** Los resultados de ocho ECCA con riesgo bajo de sesgo sugieren que el inicio o continuación de los IECA y los ARB en pacientes con COVID-19 podría aumentar la mortalidad. Sin embargo, la certeza de la evidencia es baja, por lo que se necesita más información para confirmar estas conclusiones.

• **Molnupiravir:** Los resultados de diez ECCA muestran que el tratamiento con molnupiravir reduce las hospitalizaciones y podría no aumentar los eventos adversos graves en pacientes con enfermedad de leve a moderada de comienzo reciente.

• Nirmatrelvir y ritonavir: Los resultados de un ECCA muestran que el tratamiento con nirmatrelvir y ritonavir probablemente reduzca las hospitalizaciones y no aumente los eventos adversos graves en pacientes con enfermedad de leve a moderada de comienzo reciente.

• Vitamina D: Los resultados de 20 ECCA muestran que el tratamiento con vitamina D no reduce las infecciones y probablemente no reduzca las hospitalizaciones. Los efectos de la vitamina D sobre otros desenlaces importantes son inciertos. Se necesita más información.

• Vitamina C: Los resultados de nueve ECCA sugieren que el tratamiento con vitamina C podría mejorar la resolución de los síntomas. Sin embargo, la certeza de la evidencia es baja y el efecto sobre otros desenlaces importantes es incierto. Se necesita más información.

• **Probióticos:** Los resultados de cuatro ECCA sugieren que el tratamiento con probióticos podría mejorar el tiempo de resolución de los síntomas. Sin embargo, la certeza de la evidencia es baja por imprecisión y los efectos sobre otros desenlaces importantes son inciertos. Se necesita más información.

• Enjuague bucal: Los resultados de 14 ECCA sugieren que el tratamiento con enjuagues bucales podría mejorar el tiempo de resolución de los síntomas. Sin embargo, la certeza de la evidencia es baja por imprecisión y los efectos sobre otros desenlaces importantes son inciertos. Se necesita más información.





• Mesilato de camostat: Los resultados de cinco ECCA sugieren que el tratamiento con mesilato de camostat podría no mejorar el tiempo de resolución de los síntomas. Sin embargo, la certeza de la evidencia es baja por imprecisión e información indirecta, y los efectos sobre otros desenlaces importantes son inciertos. Se necesita más información.

• **Opaganib:** Los resultados de dos ECCA sugieren que el opaganib podría no reducir la mortalidad ni la necesidad de ventilación mecánica invasiva y probablemente no incremente los eventos adversos graves, pero podría mejorar el tiempo de resolución de los síntomas. Sin embargo, la certeza de la evidencia es baja por imprecisión. Se necesita más información.

Cambios respecto a la versión anterior

• **Pirfenidiona:** La evidencia nueva incluida modifica la interpretación de los resultados o la certeza de la evidencia.

• **Peg-interferon (IFN) lamda:** La evidencia nueva incluida no modifica la interpretación de los resultados ni la certeza de la evidencia.

• **Trasplante de células madre mesenquimatosas:** La evidencia nueva incluida no modifica la interpretación de los resultados ni la certeza de la evidencia.

• Activador del plasminógeno tisular (APt): La evidencia nueva incluida no modifica la interpretación de los resultados ni la certeza de la evidencia.

• Anticoagulantes: La evidencia nueva incluida no modifica la interpretación de los resultados ni la certeza de la evidencia.

• Vitamina D: La evidencia nueva incluida modifica la interpretación de los resultados o la certeza de la evidencia.

• Favipiravir: La evidencia nueva incluida modifica la interpretación de los resultados o la certeza de la evidencia.

• Vilobelimab: La evidencia nueva incluida modifica la interpretación de los resultados o la certeza de la evidencia.

• **Palmitoiletanolamida:** La evidencia nueva incluida modifica la interpretación de los resultados o la certeza de la evidencia.

• AZD1656: La evidencia nueva incluida modifica la interpretación de los resultados o la certeza de la evidencia.





• **DFV890:** La evidencia nueva incluida modifica la interpretación de los resultados o la certeza de la evidencia.

• Vitamina B: La evidencia nueva incluida no modifica la interpretación de los resultados ni la certeza de la evidencia.

• Plasma de convalescientes: La evidencia nueva incluida no modifica la interpretación de los resultados ni la certeza de la evidencia.





Conclusiones

• La Organización Panamericana de la Salud (OPS) hace seguimiento en todo momento de la evidencia en relación con cualquier posible intervención terapéutica. A medida que se disponga de evidencia nueva, la OPS la incorporará con rapidez y actualizará sus recomendaciones, especialmente si dicha evidencia se refiere a grupos en situación de vulnerabilidad como los niños y niñas, las mujeres embarazadas, las personas mayores o los pacientes inmunocomprometidos, entre otros.

• La OPS también tiene en cuenta las diferencias en el impacto de la COVID-19 sobre las minorías y los diferentes grupos étnicos. En consecuencia, la Organización recopila constantemente información que pueda servir para mitigar el exceso de riesgo de enfermedad grave o muerte de estas minorías. Estos grupos sufren inequidades sociales y estructurales que conllevan una carga de enfermedad desproporcionada.

• La seguridad de los pacientes afectados por la COVID-19 es una prioridad clave de la mejora de la calidad de la atención y los servicios de salud.

• Sigue siendo apremiante la necesidad de elaborar ensayos clínicos aleatorizados de alta calidad que incluyan pacientes con COVID-19 a fin de poder desarrollar estrategias de manejo confiables. La importancia de los ensayos clínicos controlados aleatorizados con un diseño adecuado es fundamental en la toma de decisiones basadas en la evidencia. Hasta el momento, la mayoría de la investigación en el campo de la COVID-19 tiene muy baja calidad metodológica, lo que dificulta su identificación y validación. Urge incrementar la transparencia y plantear estudios de más calidad.



Systematic review of therapeutic options for treatment of COVID-19

Background

The vast amount of data generated by clinical studies of potential therapeutic options for COVID-19 presents important challenges. This new information must be interpreted quickly so that prescribers can make optimal treatment decisions with as little harm to patients as possible, and so that medicines manufacturers can scale-up production rapidly and bolster their supply chains. Interpreting new data quickly will save lives by ensuring that reportedly successful drugs can be administered to as many patients as possible as quickly as possible. Moreover, if evidence indicates that a medication is not effective, then ongoing clinical trials could change focus and pivot to more promising alternatives. Since many physicians are currently using treatments that rely on compassionate-use exemptions or off-label indications to treat patients with COVID-19,¹ it is crucial that they have access to the most up-to-date research evidence to inform their treatment decisions.

To address this evidence gap, we compiled the following database of evidence on potential therapeutic options for COVID-19. We hope this information will help investigators, policy makers, and prescribers navigate the flood of relevant data to ensure that management of COVID-19 at both individual and population levels is based on the best available knowledge. We will endeavor to continually update this resource as more research is released into the public space.

Methods

We used the Living OVerview of Evidence (L·OVE; https://iloveevidence.com) platform to identify studies for inclusion in this review. This platform is a system that maps PICO (Patient–Intervention–Comparison–Outcome) questions to a repository developed by Epistemonikos Foundation. This repository is continuously updated through searches in electronic databases, preprint servers, trial registries, and other resources relevant to COVID-19. The latest version of the methods, the total number of sources screened, and a living flow diagram and report of the project is updated regularly on the L·OVE website.²

Search strategy

We systematically searched in L·OVE for COVID-19. The search terms and databases covered are described on the L·OVE search strategy methods page available at: <u>https://app.iloveevidence.com/loves/5e6fdb9669c00e4ac072701d?question_domain=undefined& section=methods</u>. The repository is continuously updated, and the information is transmitted in real-time to the L·OVE platform. It was last checked for this review on 3 October 2022. The





searches covered the period from the inception date of each database, and no study design, publication status or language restriction was applied.

Study selection

The results of the searches in the individual sources were de-duplicated by an algorithm that compares unique identifiers (database identification number, digital object identifier (DOI), trial registry identification number), and citation details (i.e., author names, journal, year of publication, volume, number, pages, article title, and article abstract). Then, the information matching the search strategy was sent in real-time to the L·OVE platform where at least two authors independently screened the titles and abstracts yielded against the inclusion criteria. We obtained the full reports for all titles that appeared to meet the inclusion criteria or required further analysis and then decided about their inclusion.

Inclusion criteria

We aimed to find all available RCTs for potential therapeutic pharmacological interventions for COVID-19 with study designs that included head-to-head comparisons, or control groups with no intervention or a placebo. Target patient populations included both adults and children exposed to or with confirmed or suspected COVID-19. We focused on comparative effectiveness studies that provide evidence on outcomes of crucial importance to patients (mortality, invasive mechanical ventilation, symptom resolution or improvement, infection [prophylaxis studies] and severe adverse events).³ In addition to RCTs, we included comparative non-RCTs that report on effects of NSAID consumption on mortality. We only incorporated non-RCTs that included at least 100 patients. We presented results of RCTs and non-RCTs separately.⁴

Living evidence synthesis

An artificial intelligence algorithm deployed in the Coronavirus/COVID-19 topic of the L·OVE platform provides instant notification of articles with a high likelihood of being eligible. The authors review them, decide upon inclusion, and update the living web version of the review accordingly. If meta-analytical pooling is possible from retrieved evidence, we will do this to derive more precise estimates of effect and derive additional statistical power.

The focus has been on RCTs studies for all included therapeutic pharmacological interventions (adults and children). Adults and children exposed to or with confirmed or suspected COVID-19 were and will be included. Trials that compare interventions head-to-head or against no intervention or placebo is the focus. We have focused on comparative effectiveness studies that provide evidence on patient-important outcomes (mortality, invasive mechanical ventilation, symptom resolution or improvement, infection (prophylaxis studies), hospitalization (studies that included patients with non-severe disease) and severe adverse events).³ For studies that assessed thromboprophylactic interventions we also assessed venous thromboembolic events and major bleeding. For the outcome "hospitalization" we included information from studies reporting the



World Health Organization

number of hospitalizations or the number of hospitalizations combined with the number of deaths without hospitalization. We did not include information from studies reporting a combination of hospitalizations and medical consultations. No electronic database search restrictions were imposed.

For any meta-analytical pooling, if and when data allow, we pool all studies and present the combined analysis with relative and absolute effect sizes. To assess interventions' absolute effects, we applied relative effects to baseline risks (risks with no intervention). We extracted mortality and invasive mechanical ventilation baseline risks from the ISARIC cohort as of 18 December 2020.^{5,6} For baseline infection risk in exposed to COVID-19 we used estimates from a SR on physical distancing and mask utilization,⁷ and for adverse events and symptom resolution/improvement we used the mean risk in the control groups from included RCTs until 18 December 2020. For venous thromboembolic events and major bleeding baseline risk we used the mean risk in the control groups from included RCTs until 23 December 2021. We continuously monitor baseline risks by assessing the mean risk of every outcome in the control groups of included RCTs. When substantial changes to baseline risks are detected, we update the estimates used for absolute effects calculations. For mortality, there were some drug instances whereby we provide systematic-review (meta-analysis) evidence indirectly related to patients with COVID-19, e.g., corticosteroids in patients with ARDS.

For result interpretations and imprecision assessment we used a minimally contextualized approach which considers whether the 95%CI includes the null effect, or, when the point estimate is close to the null effect, whether the 95%CI lies within the boundaries of small but important benefit and harm that corresponds to every outcome assessed.^{8,9}

We used the following thresholds to define important benefits and harms: Mortality, +/-1%; Mechanical ventilation, +/-2%; Symptom resolution or improvement, +/-5%; Symptomatic infection in exposed individuals, +/-5%; Hospitalization in patients with mild recent COVID-19, +/-1.9%; Severe adverse events, +/-3%.

For some interventions when we found significant heterogeneity, we performed subgroup analysis considering: 1) risk of bias (high/moderate vs low risk of bias); 2) disease severity (mild, moderate, severe, or critical); and 3) intervention's characteristics (i.e., different doses or administration schemes). When we observed significant differences between subgroups, we presented individual subgroup's estimates of effect and certainty of the evidence assessment.

A risk of bias assessment was applied to RCTs focusing on randomization, allocation concealment, blinding, attrition, or other biases relevant to the estimates of effect (Table 4).¹⁰ For non-RCTs, potential residual confounding was assumed in all cases and certainty of the evidence was downgraded twice for risk of bias. The GRADE approach was used to assess the certainty on the body of evidence for every comparison on an outcome basis (Table 5).¹¹ Risk of bias judgments were compared against other similar projects (Drug treatments for covid-19: living systematic



<u>review and network meta-analysis</u> and <u>The COVID-NMA initiative</u>). Significant discrepancies were discussed until a final decision was reached.

We used MAGIC authoring and publication platform (https://app.magicapp.org/) to generate the tables summarizing our findings, which are included in Appendix 1.

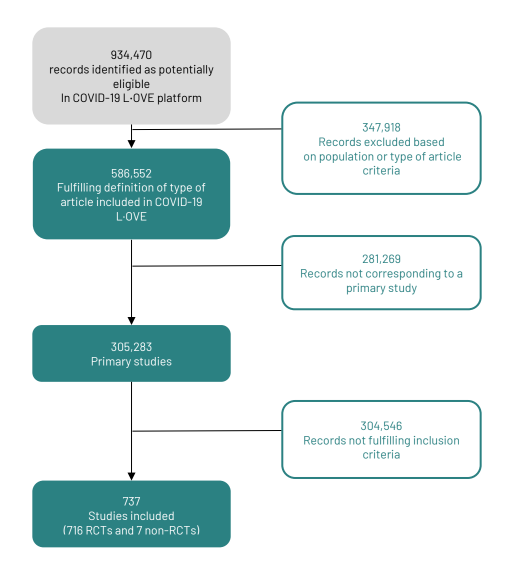


Results

Studies identified and included

Study identification and selection process is described in Figure 1. A total of 737 studies were selected for inclusion, 730 RCTs and 7 non-RCTs. A list of excluded studies is available upon request.

Figure 1. Study identification and selection process





Risk of bias

Overall, our risk of bias assessment for the limited reported RCTs resulted in high risk of bias due to suboptimal randomization, allocation concealment, and blinding (as well as other methodological and reporting concerns). Most RCTs were also very small in size and had small event numbers. The methods were very poor overall, and the reporting was suboptimal. For the observational studies, we had concerns with the representativeness of study groups (selection bias) and imbalance of the known and unknown prognostic factors (confounding). Many studies are also at risk of being confounded by indication. Most are not prospective in nature and the outcome measures are mainly heterogeneous with wide variation in reporting across the included studies. In general, follow-up was short and as mentioned, confounded potentially by the severity of disease, comorbidities, and previous or concomitant COVID-19 treatment. The risk of bias assessment of each RCT is presented in Table 4.

Table 4. Risk of bias of included RCTs



Study	Risk-of-bias arising fror randomization process	n Risk-of-bias due to deviations from the intended interventions	Risk-of-bias due to misssing outcome data	Risk-of-bias in measurement of the outcome	Risk-of-bias in selection of the reported result	Overall Risk-of-bias judg Mortality and Invasive mechanical ventilation	Symptoms, infection an adverse events
RECOVERY - Dexa	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
RECOVERY - Hydroxychloroquine	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
SCN PEP CoV-2	Low	Some Concerns	Some Concerns	Some Concerns	Low	LOW	Some Concerns
CTT-1	Low	Low	Low	Some Concerns	Low	Low	Low
				Low		LOW	
COVID-19 PEP	Low	Low	High	and the second second	Low	1.00	High
Cavalcanti et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Camran SM et al	High	Some Concerns	Low	High	Low	1 St. 1	High
COVID-19 PET	Low	Low	Low	Low	Low	Low	Low
SIMPLE	Low	Some Concerns	Low	Some Concerns	Low	Low	High
SCN PEP CoV-2	High	Some Concerns	Low	High	Low		High
Chen C et al	High	Some Concerns	Low	Some Concerns	Low	High	High
CAP-China remdesivir 2	Low	Low	Low	Low	Low	Low	Low
OTUS China	Low	Some Concerns	Low	Some Concerns	Low	Low	High
and et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
lung IF et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
RECCO-19	Low	Some Concerns	Low	Some Concerns	Low	Low	High
i Let al	High	Some Concerns	Low	Some Concerns	Low	High	High
ASTAVI	Low	Some Concerns	Low	High	Low	right	High
		and the second sec					
chen, Zeng et al	High	Some Concerns	Low	Some Concerns	Low	High	High
chuan Li C et al	High	Some Concerns	Low	Some Concerns	Low	High	High
heng et al	High	Some Concerns	Low	Some Concerns	Low	High	High
LACOI	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ONCOVID	Low	Some Concerns	Low	Some Concerns	Low	Some Concerns	High
LUCOCOVID	High	Some Concerns	Low	Low	Low	High	High
loroCOVID19	Low	Low	Low	Some Concerns	Low	Low	Low
avoudi-Monfared et al	High	Some Concerns	Low	Low	Low	High	High
hen et al	High	Some Concerns	Low	Low	Low	High	High
avoodi L et al	High	Some Concerns	Low	Low	Low	High	High
rashchenko AA et al		Some Concerns	Low	Low	Low		
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hen PC et al	High	Some Concerns	Low	Low	Low	High	High
IC-nCoV	High	Some Concerns	Low	Low	Low	High	High
ou Y et al	High	Some Concerns	Low	Low	Low	High	High
laar APJ et al	High	Some Concerns	Low	Some Concerns	Low	High	High
IC-COVID-19	High	Some Concerns	Low	Some Concerns	Low	High	High
uvenmez O et al	High	Some Concerns	Low	Some Concerns	Low	High	High
luang et al	High	Some Concerns	Low	Some Concerns	Low	High	High
uan et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ten Z et al	High	Some Concerns	Low	Some Concerns	Low	High	High
fehboob R et al	High	Some Concerns	Low	Some Concerns	Low	High	High
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Chong et al	Low	Some Concerns	Low	Low	Low	Low	High
akoulas et al	High	Some Concerns	Low	Some Concerns	Low	High	High
lu K, Wang M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
SPERANZA	High	Some Concerns	Low	Some Concerns	Low	High	High
opes et al	High	Low	Low	Low	Low	High	High
Duarte M et al	High	High	High	Some Concerns	Some Concerns	High	High
fetcovid	Low	Low	Low	Low	Low	Low	Low
Mansour E et al	Low	Low	Low	Some Concerns	Low	Low	High
hang J et al	High	Some Concerns	Low	Some Concerns	Low	High	High
RECOVERY - Lopinavir-ritonavir	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
ARDEA	Low	Low	Low	Low	Low	Low	Low
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bbaspour Kasgari H et al	High	Some Concerns	Low	Some Concerns	Low	High	High
iadeghi A et al	High	Some Concerns	Low	Low	Low	High	High
ihu L et al	High	Some Concerns	Low	Some Concerns	Low	High	High
IMPLE 2	Low	Some Concerns	Low	Some Concerns	Low	Some Concerns	High
bd-Elsalam S et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ekhavati E et al	High	Some Concerns	Low	Some Concerns	Low	High	High
houman et al	High	Some Concerns	Low	Some Concerns	Low	High	High
tahmani H et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ConPlas-19	Low	Some Concerns	Low	Some Concerns	Low	Some Concerns	High
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EMAP-CAP	Low	Some Concerns	Low	Some Concerns	Low	Low	High
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CODEX	Low	Some Concerns	Low	Some Concerns	Low	Low	High
OVIDIOL	High	Some Concerns	Low	Some Concerns	Low	High	High
APE COVID	Low	Low	Low	Low	Low	Low	Low
OVACTA	Low	Low	Low	Low	Low	Low	Low
OALITION II	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Tetal	High	Some Concerns	Low	Some Concerns	Low	High	High
/ang D et al	High	Some Concerns	Low	Some Concerns	Low	High	High
howdhury et al	High	Some Concerns	Low	Some Concerns	Low	High	High
LACID	Low	Some Concerns	Low	Some Concerns	Low	Some Concerns	High
harebachi N et al		Low	Low	Low	Low	Some Concerns	Some Concerns
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X-COVID19	High	Some Concerns	Low	Some Concerns	Low	High	High
heng LL et al	High	Some Concerns	Low	Some Concerns	Low	High	High
arahani R et al	High	Some Concerns	Low	Some Concerns	Low	High	High
imura KS et al	High	Some Concerns	Low	Some Concerns	Low	High	High
TENEA-Co-300	High	Some Concerns	Low	Some Concerns	Low	High	High
/u X et al	Low	Low	Low	Low	Low	Low	Low
alcells ME et al (Pontificia Universidad Catolica de Chile)	Low	Some Concerns	Low	Some Concerns	Low	Low	High
dalatifard M et al (Tehran University of Medical Sciences)	High	Some Concerns	Low	Some Concerns	Low	High	High
OVID-19 PREP		Low	Low		Low	a state of the second se	
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loi Y et al (Fujita Health University Hospital)	High	Some Concerns	Low	Some Concerns	Low	High	High
odder et al	High	Some Concerns	Low	Some Concerns	Low	High	High
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PCF19Muthar K et alHighSome ConcernsLowSome ConcernsLowHighHighAthattar K et alHighLowLowLowLowLowHighHighAthattar K et alHighSome ConcernsLowSome ConcernsLowHighHighFDL21616224.00HighSome ConcernsLowSome ConcernsLowHighHighAdeChalainsHighSome ConcernsLowSome ConcernsLowHighHighProlectindHighSome ConcernsLowSome ConcernsLowHighHighAdeChalainsHighSome ConcernsLowSome ConcernsLowHighHighCARSUESHighSome ConcernsLowSome ConcernsLowLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLow		Low		Low	Some Concerns	Low	Low	High
MathemHighSome ConcernsLowSome ConcernsLowLowHighHighAhmed et alHighSome ConcernsLowSome ConcernsLowHighHighHighAbd-Blaum S et al (Tartia University)HighSome ConcernsLowSome ConcernsLowHighHighAdd-Blaum S et al (Tartia University)HighSome ConcernsLowSome ConcernsLowHighHighAdd-Blaum S et al (Tartia University)HighSome ConcernsLowSome ConcernsLowHighHighAdd-Blaum S et al (Tartia University)HighSome ConcernsLowSome ConcernsLowHighHighCARGLESHighSome ConcernsLowSome ConcernsLowLowSome ConcernsLowNoreLowSome ConcernsLowNoreLowSome ConcernsLowL	Niaee et al	Some Concerns	Some Concerns	Low	Some Concerns	Low	High	High
Ahmed taiHighLowLowLowLowLowLowHighHighTOLL-19-22-200HighSome ConcernsLowSome ConcernsLowHighHighProtectin-MHighSome ConcernsLowSome ConcernsLowHighHighProtectin-MHighSome ConcernsLowSome ConcernsLowHighHighCARDELSHighSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowCARDELSHighSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowCor	PICP19	High	Some Concerns	Low	Some Concerns	Low	High	High
IPOLE-19-02-LODHighSome ConcernsLowSome ConcernsLowHighHighAdd-Baalam S et al (Tanta University)HighSome ConcernsLowSome ConcernsLowHighHighAdd-Baalam S et al (Tanta University)HighSome ConcernsLowSome ConcernsLowHighHighMadonado V et alHighSome ConcernsLowSome ConcernsLowHighHighCARGLESHighSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowLowLowLowLowLowLowLowLowLowLowLowLowLowSome ConcernsLowLowLowLowSome ConcernsRecordernsRecordernsLowLowLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowLowSome ConcernsLowLowLowSome ConcernsLowLowSome ConcernsLowLowLowSome ConcernsLowLowLowSome ConcernsLow <td>Mukhtar K et al</td> <td>High</td> <td>Some Concerns</td> <td>Low</td> <td>Some Concerns</td> <td>Low</td> <td>High</td> <td>High</td>	Mukhtar K et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Abd-Bisdam S et al (Tanta University)HighSome ConcernsLowSome ConcernsLowHighHighHighProtectin-MHighSome ConcernsLowSome ConcernsLowHighHighHighCARSLESHighSome ConcernsLowSome ConcernsLowHighHighHighGARSLESHighSome ConcernsLowSome ConcernsLowHighHighHighCARSLESLowLowSome ConcernsLowLowSome ConcernsCowCowCome ConcernsCowCowCowCome ConcernsCow<	Ahmed et al	High	Low	Low	Low	Low	High	High
prodecimal Maidonado Vet al Maidonado Vet al GARGLESHighSome ConcernsLowSome ConcernsLowHighHighMaidonado Vet al CARGLESHighSome ConcernsLowSome ConcernsLowHighHighERSULowLowSome ConcernsLowLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowLowSome ConcernsLowLowLowLowLowLowLowLowSome ConcernsLow<	ITOLI-C19-02-I-00	High	Some Concerns	Low	Some Concerns	Low	High	High
National Vet al GARGLESHigh HighSome Concerns LowLowSome Concerns Some ConcernsLowHigh HighHigh HighGARGLESLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowSome ConcernsLowLowSome ConcernsLowLowSome ConcernsLowLowLowSome ConcernsLow	Abd-Elsalam S et al (Tanta University)	High	Some Concerns	Low	Some Concerns	Low	High	High
LARGLESHighSome ConcernsLowSome ConcernsLowHighHighHighERSulLowLowSome ConcernsLowLowLowSome ConcernsLowLowLowLowLowLowLowLowSome ConcernsRecovernsLowLowLowSome ConcernsLowLowSome ConcernsLowLowLowSome ConcernsSome ConcernsLow <td>Prolectin-M</td> <td>High</td> <td>Some Concerns</td> <td>Low</td> <td>Some Concerns</td> <td>Low</td> <td>High</td> <td>High</td>	Prolectin-M	High	Some Concerns	Low	Some Concerns	Low	High	High
ERSulLowLowLowLowLowLowLowSome ConcernsSome ConcernsConcernsSome ConcernsConcernsSome Concerns	Maldonado V et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Chaccour et alLowLowLowLowLowLowLowLowLowLowLowLowLowLowLowLowConcorrensSome ConcernsLowLowSome ConcernsElDO-2801-1001LowLowLowLowSome ConcernsLowLowLowLowLowSome ConcernsElDO-2801-1001Low <td>GARGLES</td> <td>High</td> <td>Some Concerns</td> <td>Low</td> <td>Some Concerns</td> <td>Low</td> <td>High</td> <td>High</td>	GARGLES	High	Some Concerns	Low	Some Concerns	Low	High	High
ACTT-2LowLowSome ConcernsLowLowLowSome ConcernsLowLowLowSome ConcernsSome ConcernsLowLowLowLowCowLowCowLowCowLowCowLowCowLow <td>ERSul</td> <td>Low</td> <td>Low</td> <td>Some Concerns</td> <td>Low</td> <td>Low</td> <td>Some Concerns</td> <td>Some Concerns</td>	ERSul	Low	Low	Some Concerns	Low	Low	Some Concerns	Some Concerns
RECOVERY EJDD-2801-1001LowLowLowLowLowLowLowLowConcenseLow<	Chaccour et al	Low	Low	Low	Low	Low	Low	Low
EID-2801-1001LowLowLowLowLowLowLowLowWeinreichLowL		Low	Low	Some Concerns	Low	Low	Some Concerns	Some Concerns
WeinreichLow		Low	Some Concerns	Low	Low	Low	Low	Some Concerns
Roozbeh F et alLowSome ConcernsLowSome ConcernsLow <td></td> <td></td> <td>Low</td> <td></td> <td></td> <td>Low</td> <td>Low</td> <td>Low</td>			Low			Low	Low	Low
ACTIV-3/TICO Low Some Concerns Low Low Low Low High High Chachar et al High Some Concerns Low Some Concerns Low High High Babylova Let al Low Some Concerns Low Some Concerns Low High High Babslova Let al Low Some Concerns Low Low High High REPLACE COVID Low Low Some Concerns Low Some Concerns Low Low High High Kiff et al Low Low Some Concerns Low	Weinreich	Low		Low	Low	Low	Low	Low
Chachar et al High Some Concerns Low Some Concerns Low High High Balykova LA et al High Some Concerns Low Some Concerns Low High High Balakola et al Low Low Low Low Low Low Low Low Low Magnetic High		Low	Some Concerns	Low	Some Concerns	Low	Low	High
Bakkova LA et al High Some Concerns Low Some Concerns Low			Low	Some Concerns	Low	Low	Low	High
Babakia et al Low Some Concerns Low Some Concerns Low Low <t< td=""><td></td><td></td><td>Some Concerns</td><td>Low</td><td>Some Concerns</td><td>Low</td><td>High</td><td>High</td></t<>			Some Concerns	Low	Some Concerns	Low	High	High
REMAP-CAP - todiizumab Low Some Concerns Low Some Concerns Low Low High Abdemakoud AA et al High Some Concerns Low Some Concerns Low High High REPLACE COVID Low Some Concerns Low Some Concerns Low Englished High Some Concerns Low Some Concerns Low Some Concerns Low Low Some Concerns Low Low Some Concerns Low Low Some Concerns Low Low Low Low	Balykova LA et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Abdeimaksoud AA et al High Some Concerns Low Some Concerns Low Some Concerns Low High High REFLACE COVID Low Some Concerns Low Some Concerns Low Low High REFLACE COVID Low High		Low	Low	Low		Low	Low	Low
REPLACE COVID Low Some Concerns Low Some Concerns Low High High Courcers Low Low Low Some Concerns Low Some Concerns Low Low Low Migh High COV/FERON Low Some Concerns Low Some Concerns Low				Low	Some Concerns			
REPLACE COVID Low Some Concerns Low Some Concerns Low Some Concerns Low Low Low Some Concerns Low Low Low Low Some Concerns Low	Abdelmaksoud AA et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Kumari P et al High Some Concerns Low Some Concerns Low High High FK/FA/0DA-CoV/2D20 High Low Low Low Low Low High High Cohalia et al High Some Concerns Low Some Concerns Low High High COVIFER/ON Low Some Concerns Low Some Concerns Low Main High COVIFER/ON Low Some Concerns Low Some Concerns Low High	REPLACE COVID	Low	Some Concerns	Low	Some Concerns			High
FK/FAV00A-CoV/2020 High Low Low Low Low High High Chalke tal High Some Concerns Low Some Concerns Low High High COVIFER/ON Low Some Concerns Low Some Concerns Low Low Some Concerns RECOVERY-Plasma Low Some Concerns Low Low Low Some Concerns Interferon in COVID (Aisy Darazam 1 et al) Low Some Concerns Low Some Concerns Low Low Low High JamailMoghadma/Sinkkall S et al High Some Concerns Low Some Concerns Low High High Sedighiyan M et al High Some Concerns Low Some Concerns Low High High Roostaci A et al High Some Concerns Low Some Concerns Low High High SecOvid Low Some Concerns Low Some Concerns Low High High High High	Kirti et al	Low	Low	Low	Low	Low	Low	Low
Chahla et al High Some Concerns Low Low Low Migh RECOVER/Veisam Low Some Concerns Low Low Low Low More Concerns Low Low More Concerns Low Migh High	Kumari P et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Chalase al High Some Concerns Low Some Concerns Low Some Concerns Low High High COVIFERON Low Some Concerns Low Some Concerns Low Low Main High RECOVER/Plasma Low Some Concerns Low Low Low Low Some Concerns Low Low Main Main Main Some Concerns Low Low Low High	FK/FAV00A-CoV/2020	High	Low	Low	Low	Low	High	High
RECOVERY-Plasma Low Some Concerns Low Low Low Some Concerns Interferon in COVID (Javi Darazam 1 et al) Low Some Concerns Low Some Concerns Low Low High AB-DRUG-SARS-004 (Cadegiani FA et al) High Some Concerns Low Some Concerns Low High High JamailMoghadmSinklail S et al High Some Concerns Low Some Concerns Low High High Sedighiyan M et al High Some Concerns Low Some Concerns Low High High Roostaci A et al High Low Low Low Low Low High High See-Covid Low Some Concerns Low Low Low Low High High SecTor High Some Concerns Low Some Concerns Low Low High High			Some Concerns	Low	Some Concerns	Low		
Interferon in COVID (Alavi Darazam I et al) Low Some Concerns Low Some Concerns Low Low High AB-DRUG-SARS-004 (Cadegiani FA et al) High Some Concerns Low Some Concerns Low High High JamailMoghandamSishikali S et al High Some Concerns Low Some Concerns Low High High Sedighiyan M et al High Some Concerns Low Some Concerns Low High High Rootsack A et al High Low Low Some Concerns Low Low High High Sec-Oxid Low Some Concerns Low Low Low High High Sec-Oxid Low Some Concerns Low Some Concerns Low High High Sec-Oxid Low Some Concerns Low Some Concerns Low High High	COVIFERON	Low	Some Concerns	Low	Some Concerns	Low	Low	High
AB-DRUG-SARS-004 (Cadegiani FA et al) High Some Concerns Low Some Concerns Low High High JamailMoghadamSiahkali S et al High Some Concerns Low Some Concerns Low High High Sedighyan M et al High Some Concerns Low Some Concerns Low High High Roostaei A et al High Low Low Some Concerns Low High High Bee-Covid Low Some Concerns Low Some Concerns Low High High SEOT High Some Concerns Low Some Concerns Low High High	RECOVERY-Plasma	Low	Some Concerns	Low	Low	Low	Low	
AB-DRUG-SARS-004 (Cadegiani FA et al) High Some Concerns Low Some Concerns Low High High JamailMoghadamSiahkali S et al High Some Concerns Low Some Concerns Low High High Sedighyan M et al High Some Concerns Low Some Concerns Low High High Roostaei A et al High Low Low Some Concerns Low High High Bee-Covid Low Some Concerns Low Some Concerns Low High High SEOT High Some Concerns Low Some Concerns Low High High								
JamailMoghadamSiahkali S et al High Some Concerns Low Some Concerns Low High High Sedghyan M et al High Some Concerns Low Some Concerns Low High High Rootsaei A et al High Low Low Low Low Low High High Bee-Covid Low Some Concerns Low Some Concerns Low High High SEOT High Some Concerns Low Some Concerns Low High High		High		Low	Some Concerns	Low	High	
Sedigitiyan M et al High Some Concerns Low Some Concerns Low High High Rootsaic A et al High Low Low Low Low High High Bee-Covid Low Some Concerns Low Some Concerns Low High SEOT High Some Concerns Low Some Concerns Low High				Low		Low		
Roostaei A et al High Low Low Low Low High High Bee-Covid Low Some Concerns Low Some Concerns Low Low High SEOT High Some Concerns Low Some Concerns Low High High						Low		
Bee-Covid Low Some Concerns Low Some Concerns Low Low High SEOT High Some Concerns Low Some Concerns Low High High								
SEOT High Some Concerns Low Some Concerns Low High High								-
		Low	Low	Low	Low	Low	Low	Low
Shahbaznejad et al Low Low Low Low Low Low Low Low								
Sporth i al High Some Concerns Low Some Concerns Low High High								
المهنا المراجع المستعلمين المراجع	1 · · · · · · · · · · · · · · · · · · ·	-		l.			-	-



Samaha et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Bukhari el al	High	Some Concerns	Low	Some Concerns	Low	High	High
Okumus et al /eiga	High Low	Some Concerns Some Concerns	Low Low	Some Concerns Low	Low	High Low	High Some Concerns
Sottlieb	Low	Low	Low	Low	Low	Low	Low
BRACE CORONA	Low	Some Concerns	Some Concerns	Low	Low	Low	High
CORIMUNO-ANA-1	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Thakar A et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Onal H et al	High	High	Low	Some Concerns	Low	High	High
Tang X et al	Low	Some Concerns	Low	Low	Low	Low	Low
COLCORONA	Low	Some Concerns	Low	Low	Low	Low	Low
.opardo Dabbous HM et al	Low High	Low Some Concerns	Low	Low Some Concerns	Low	Low High	Low High
ATTRACT	Low	Some Concerns	Low	Low	Low	Low	Low
Ranibar K et al	Some Concerns	Low	Low	Low	Low	Some Concerns	Some Concerns
EAT-DUTA AndroCoV	Low	Low	High	Low	Low	High	High
arnoosh G et al	Some Concerns	Some Concerns	High	Some Concerns	Low	High	High
Ghalili H et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Baklaushev VP et al	High	Some Concerns	Low	Some Concerns	Low	High	High
GLLER	High	Some Concerns	Low	Some Concerns	Low	High	High
IYDRA Sali S et al	Low	Some Concerns	Low	Low	Low	Low	Low
all S et al	High High	Some Concerns	Low	Some Concerns	Low	High High	High High
SVU-MED-CHT019-420860	High	Some Concerns	Low	Some Concerns	Low	High	High
TOIC	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Borges M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
RECOVERY-TCZ	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
COVIDAtoZ -Zinc	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
COVID-19 Early Treatment	Low	Some Concerns	Low	Low	Low	Low	Low
Shogenova LV et al	High	Some Concerns	Low	Some Concerns	Low	High	High
FC16844	Low	Some Concerns	Low	Low	Low	Low	Low
ARTI-19	High	Some Concerns	Low	Some Concerns	Low	High	High
Purwati	High	Some Concerns	Low	Some Concerns	Low	High	High
/B-N-IVIG-COVID-19/2020-CT2	High	Some Concerns	Low	Some Concerns	Low	High	High
amaati H et al Seltran-HCQ	High High	Some Concerns	Low	Some Concerns	Low	High High	High High
ZINC COVID	Low	Some Concerns Some Concerns	Low	Some Concerns	Low	Low	Low
ATCH 1	Low	Some Concerns	Low	Some Concerns	Low	Low	High
B-DRUG-SARS-004-2	High	Some Concerns	Low	Some Concerns	Low	High	High
Nouri-Vaskeh M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
opez-Medina et al	Low	Low	Low	Low	Low	Low	Low
akkireddy M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Silva	High	Some Concerns	Low	Some Concerns	Low	High	High
RINCIPLE	Low	Some Concerns	Some Concerns	Some Concerns	Low	Some Concerns	High
Bermejo Galan et al	Low	Low	Low	Low	Low	Low	Low
Pott-Junior et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Mikhaylov	Low	Some Concerns	Low	Some Concerns	Low	Low	High
2GAMMACOVID-19	High	Some Concerns	Low	Some Concerns	Low	High	High
AAAS9924 Tolouian et al	Low	Low Some Concerns	Some Concerns	Some Concerns	Low	Some Concerns	Some Concerns High
ElZein R et al	High	Some Concerns	Low	Some Concerns	Low	High	High
PEGI.20.002	High	Some Concerns	Low	Some Concerns	Low	High	High
MASH-COVID	Low	Some Concerns	Low	Low	Low	Low	Low
NSPIRATION	Low	Some Concerns	Low	Low	Low	Some Concerns	Some Concerns
Zarychanski	Low	Some Concerns	Low	Low	Low	Some Concerns	Some Concerns
Santos PSS et al	Low	Some Concerns	Low	Low	Low	Low	Low
Solaymani-Dodaran M et al	Low	Some Concerns	Low	Low	Low	Low	Low
D-0903-0188	High	Some Concerns	Low	Some Concerns	Low	High	High
DISCOVER	Low	Some Concerns	Low	Low	Low	Low	Low
SURG-2020-28683	Low	Some Concerns	Low	Low	Low	Low	Low
Vavi-Moghaddam M et al CT-P59 3.2	High	Some Concerns Some Concerns	Low	Some Concerns	Low	High	High
/adollahzadeh M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
BCovid	Low	Some Concerns	Low	Low	Low	Low	Low
lanna Huang Y et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Saynitdinova VV et al	High	Some Concerns	Low	Some Concerns	Low	High	High
031-120	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Seltran Gonzalez JL et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Doaei S et al	Low	Some Concerns	Some Concerns	Some Concerns	Low	Some Concerns	High
COVID-AIV	High	Some Concerns	Low	Some Concerns	Low	High	High
Amra Betal	High	Some Concerns	Low	Some Concerns	Low	High	High
Ribakov AR et al	High	Some Concerns Some Concerns	Low	Some Concerns	Low	High	High
üshoria N et al CERC-002-CVID-201	Low High	Low	High	Some Concerns	Low	High	High High
Mahajan L et al	High	Some Concerns	Low	Some Concerns	Low	High	High
PRINCIPLE	Low	Some Concerns	Some Concerns	Some Concerns	Low	Some Concerns	Some Concerns
Pouladzadeh M et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
IBOTCOVID19	High	Some Concerns	Low	Some Concerns	Low	High	High
RESIST	High	Some Concerns	Low	Some Concerns	Low	High	High
RESIST	High	Some Concerns	Low	Some Concerns	Low	High	High
CARR-COV-02	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Seet	Low	Some Concerns	Low	Some Concerns	Low	Low	High
BU-COVID19-ConvalescentPlasma	Low	Some Concerns	Low	Low	Low	Low	Low
TOGETHER	Low	Some Concerns	Low	Low	Low	Low	Low
hao H et al	High	Some Concerns	Low	Some Concerns	Low	High	High
DSCAR	Low	Some Concerns	Low	Low	Low	Low	Low
POLYCOR	Low	Some Concerns	Low	Low	Low	Low	Low
/anguard Samimagham HR et al	Low	Some Concerns Some Concerns	Low	Low Some Concerns	Low	Low	Low High
Samimagnam HR et al CamoCO-19	Low	Some Concerns	Low	Low	Low	Low	Low
BCR-PNB-001	High	Some Concerns	Low	Some Concerns	Low	High	High
		Control Control 110					1.

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Siami Z et al	High	Some Concerns	Low	Some Concerns	Low	High	High
CLOROTRIAL	High	Some Concerns	Low	Some Concerns	Low	High	High
ROBCO	High	Some Concerns	Low	Some Concerns	Low	High	High
		and the second sec	2000	Contraction of the second second	12.2.12		
esari TM et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ISCO	High	Some Concerns	Low	Some Concerns	Low	High	High
INS-COVID-PK	Low	Some Concerns	Low	Low	Low	Low	Low
tashad A et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Noni M et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ACCT	Low	Some Concerns	Low	Some Concerns	Low	Low	High
COV-BARRIER	Low	Some Concerns	Low	Low	Low	Low	Low
IVE-AIR	Low	Some Concerns	Low	Low	Low	Low	Low
PreToVid	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Mahmoudi M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
AGIL F	Low	Some Concerns	Low	Some Concerns	Low	Low	High
lamdy Salman O et al	Low	Some Concerns	Low	Low	Low	Low	Low
COVID-RT-01	Low	Some Concerns		Low		Low	
COVID-ARB	- 2.2		Low	1	Low		Low
Perepu II et al	High	Some Concerns	Low	Some Concerns	Low	High	High
	High	Some Concerns	Low	Some Concerns	Low	High	High
arychanski-Non-critical	Low	Some Concerns	Low	Low	Low	Some Concerns	Some Concerns
arilumab-COVID19 Study	Low	Some Concerns	Low	Low	Low	Low	Low
APSID	High	Some Concerns	Low	Some Concerns	Low	High	High
HEER	High	Some Concerns	Low	Some Concerns	Low	High	High
ECOVERY - Colchicine	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
ilvia Mendez-Flores S et al	High	Low	Low	Low	Low	High	High
AVE-MORE	Low	Some Concerns	Low	Low	Low	Low	Low
vinchester S et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Igohary MAS et al	High	Some Concerns	Low	Some Concerns	Low	High	High
RMY-1	Low	Some Concerns	Low	Low	Low	Low	Low
amidi-Alamdari D et al				and the second se			
	High	Some Concerns	Low	Some Concerns	Low	High	High
arehoseinzade E et al	High	Some Concerns	Low	Some Concerns	Low	High	High
bd-Elsalam S et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
liber et al	Low	Low	Some Concerns	Low	Low	Low	Low
aisal et al	High	Some Concerns	Low	Some Concerns	Low	High	High
OVECOD	Low	Some Concerns	Low	Some Concerns	Low	Low	High
CTION	Low	Some Concerns	Low	Low	Low	Some Concerns	Some Concerns
BLAZE-2	Low	Low	Low	Low	Low	Low	Low
roPAC-COVID	Low	Low	Low	Low	Low	Low	Low
ian F et al	High	Some Concerns	Low	Some Concerns	Low	High	High
RECOVERY - ASA	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
IONEST	Low	Some Concerns	Low	Some Concerns	Low	Low	High
COMET-ICE	Low	Low	Low	Low	Low	Low	Low
SMMSCCOVID19	Low	Low	Low	Low	Low	Low	Low
SENTAD-COVID	Low	Some Concerns	Low	Some Concerns	Low	Low	High
SEV-COVID	High	Some Concerns	Low	Some Concerns	Low	High	High
CATALYST	High	Some Concerns	Low	Some Concerns	Low	High	High
NiSet al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
RECOVERY - REGEN-COV	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
Taher A et al	High	Low	Low	Low	Low	High	High
ACEI-COVID	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
Covid-19 Phase 3 Prevention Trial	Low	Low	Low	Low	Low	Low	Low
EIDD-2801-2003	Low	Low	Low	Low	Low	Low	Low
REMAP-CAP	Low	Some Concerns	Low	Low	Low	Low	Some Concerns
							Contraction of the second second
STOP-COVID	Low	Low	Low	Low	Low	Low	Low
/allejos et al	Low	Low	Low	Low	Low	Low	Low
CONCOR-1	Low	Some Concerns	Low	Some Concerns	Low	Low	High
LBERTA HOPE-Covid 19	Low	Low	Low	Low	Low	Low	Low
lamed DM et al	High	Some Concerns	Low	Some Concerns	Low	High	High
COUNTER-COVID	Low	Low	Low	Low	Low	Low	Low
bdulamir AS et al	High	Some Concerns	Low	Some Concerns	Low	High	High
P-DRUG-SARS-003	High	Low	Low	Low	Low	High	High
vref ZF et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Pierro F et al	High	Some Concerns	Low	Some Concerns	Low	High	High
R0-CORONA	Low	Some Concerns	Low	Some Concerns	Low	Low	High
RCHITECTS	Low	Low	Low	Low	Low	Low	Low
CORIMUNO-TOCI ICU	Low	Some Concerns	Low	Some Concerns	Low	Low	High
		and the second second		and the second sec			
COV-AID	Low	Low	Low	Low	Low	Low	Low
OVIDOSE-2	Low	Low	Low	Low	Low	Low	Low
OVIDSTORM	Low	Low	Low	Low	Low	Low	Low
OVITOZ-01	Low	Low	Low	Low	Low	Low	Low
IMO-0224-20	High	Low	Low	Low	Low	High	High
EMDACTA	Low	Low	Low	Low	Low	Low	Low
mmCoVA	Low	Low	Low	Low	Low	Low	Low
avoudian N et al	Low	Low	Low	Low	Low	Low	Low
OCOVID	Low	Low	Low	Low	Low	Low	Low
OVINTOC	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ORIMUNO-SARI	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ORIMUNO-SARI ICU	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ARCOVID		the second se	and the			a benefit of the second s	
	Low	Low Common	Low	Low	Low	Low	Low
ARICOR	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ARTRE	Low	Some Concerns	Low	Some Concerns	Low	Low	High
:OV-AID-2	Low	Low	Low	Low	Low	Low	Low
REGENERON Sari P3	Low	Some Concerns	Low	Low	Low	Low	Low
OPEP	Low	Some Concerns	Low	Some Concerns	Low	Low	High
APID	Low	Some Concerns	Low	Low	Low	Some Concerns	Some Concerns
Vang Q et al	High	Some Concerns	Low	Some Concerns	Low	High	High
		Washington and an and a second	10.00	Succession in the second second			1 S 1 2 1
losseinzadeh A et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ILAZE-1	Low	Low	Low	Low	Low	Low	Low
lajmeddin F et al	Low	Low	Low	Low	Low	Low	Low
AN-COVID	Low	Low	Low	Low	Low	Low	Low
duardo FP et al	Low	Low	Low	Low	Low	Low	Low
B-DRUG-SARS-005	High	Low	Low	Low	Low	High	High
B-DRUG-SARS-005						riigh	



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ACTION	Low	Low	High	Low	Low	Some Concerns	Some Concerns
Saitan-Duarte HG et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
abico S et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
LACOVID	Low	Some Concerns	Low	Some Concerns	Low	Some Concerns	High
AIIC	Low	Some Concerns	Low	Some Concerns	Low	Low	High
SHOP	Low	Some Concerns	Low	Some Concerns	Low	Low	High
sadipooya K et al	High	Some Concerns	Low	Some Concerns	Low	High	High
avichandran et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ARE-19	Low	Low	Low	Low	Low	Low	Low
OXYCOV	Low	Some Concerns	Low	Some Concerns	Low	Low	High
RINCIPLE	Low	Low	Low	Low	Low	Low	Low
arikh D et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ovid-19 Phase 3 Prevention Trial - Exposed	Low	Low	Low	Low	Low	Low	Low
hree C	Low	Low	Low	Low	Low	Low	Low
OVIDIT	Low	Some Concerns	Low	Some Concerns	Low	Low	High
UMC-COVID-19	High	Some Concerns	Low	Some Concerns	Low	High	High
bbass S et al	High	Some Concerns	Low	Some Concerns	Low	High	High
3PO	Low	Low	Low	Low	Low	Low	Low
osak et al	High	Some Concerns	Low	Some Concerns	Low	High	High
OGHETER-Fluvoxamine	Low	Low	Low	Low	Low	Low	Low
OCIDEX	Low	Some Concerns	Low	Some Concerns	Low	Low	High
akharian A et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ERO-HCQ	Low	Low	Low	Low	Low	Low	Low
lizadeh Z et al	High	Some Concerns	Low	Some Concerns	Low	High	High
hushan S et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ASCEPA COVID-19 CARDIOLINK-9	High	Some Concerns	Low	Some Concerns	Low	High	High
hinkai M et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
odrigues C et al	Low	Low	Low	Low	Low	Low	Low
ousavi SA et al	High	Some Concerns	Low	Some Concerns	Low	High	High
trich	Low	Low	Low	Low	Low	Low	Low
ADRID-COVID	Low	Some Concerns	Low	Some Concerns	Low	Low	High
2W-MC-PYAA	Low	Low	Low	Low	Low	Low	Low
AWn-Plasma	Low	Some Concerns	Low	Some Concerns	Low	Some Concerns	High
PTIMISE-C19	Low	Low	Low	Low	Low	Low	Low
oppola	High	Low	Low	Low	Low	High	High
LV-020-001	Low	Low	Low	Low	Low	Low	Low
ates MRI RESPOND-1	Low	Low	Low	Low	Low	Low	Low
CTIV-2	High	Some Concerns	Low	Some Concerns	Low	Low	Low
ARVIN	Low	Low	Low	Low	Low	Low	Low
uonfrate et al	Low	Low	Low	Low	Low	Low	Low
IcCreary M et al	Low	Low	Low	Low	Low	Low	Low
ihanei M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
naner wetai laskin etal	Low	Low	Low	Low	Low	Low	Low
OL-COVID	High	Some Concerns		Some Concerns		High	High
RINCIPLE - Colchicine	Low	Some Concerns	Low	Some Concerns	Low	Low	High
lassaniazad M et al	High	Low	Low	Low	Low	High	High
tamachandran R et al	Low	Low	Low	Low	Low	Low	Low
PI-006-002	High	Low	Low	Low	Low	High	High
N-Domênico MB et al	High	Low	Some Concerns	Low	Low	High	High
T-P59 1.2	Low	Low	Low	Low	Low	Low	Low
BC-110	Low	Low	Low	Low	Low	Low	Low
ORONA			Contraction of the second s	Low			
	Low	Low	Low		Low	Low	Low
TARS	High	Some Concerns	Low	Some Concerns	Low	High	High
RTAN-C19	High	Low	High	Low	Low	High	High
labalola OE et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ESPERIDIN	Low	Low	Low	Low	Low	Low	Low
teszinate	Low	Low	Low	Low	Low	Low	Low
zizi H et al	High	Low	High	Low	Low	High	High
IGHT-COVID-19	High	Some Concerns	Low	Some Concerns	Low	High	High
ANDIDATE	Low	Low	Low	Low	Low	Low	Low
EMICOP	High	Some Concerns	Low	Some Concerns	Low	High	High
EP-COVID	Low	Low	Low	Low	Low	Some Concerns	Some Concerns
CTIV-4B	Low	Low	Low	Low	Low	Low	Low
OV-BARRIER-IMV	Low	Low	Low	Low	Low	Low	Low
EFINE	Low	Some Concerns	Low	Some Concerns	Low	Low	High
EV-COVID	High	Some Concerns	Low	Some Concerns	Low	High	High
ARPAC	High	Some Concerns	Low	Some Concerns	Low	High	High
amir YM et al	High	Some Concerns	Low	Some Concerns	Low	High	High
bd-Elsalam S et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ROCOV-19-2020	High	Some Concerns	Low	Some Concerns	Low	High	High
aghighi S et al	High	Some Concerns	Low	Some Concerns	Low	High	High
UXCOVID	Low	Low	Low	Low	Low	Low	Low
CTT-3	Low	Low	Low	Low	Low	Low	Low
meri A et al	High	Some Concerns	Low	Some Concerns	Low	High	High
aghbooli Z et al	High	Low	Low	Low	Low	High	High
ITEREST	Low	Low	Low	Low	Low	Low	Low
liynyk O et al	High	Some Concerns	Low	Some Concerns	Low	High	High
B-P12-01	Low	Low	Low	Low	Low	Low	Low
obarak S et al	Low	Low	Low	Low	Low	Low	Low
al Fetal	High	Some Concerns	Low	Some Concerns	Low	High	High
hu R et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ONTAIN	Low	Low	Low	Low	Low	Low	Low
OV-AID-3	Low	Some Concerns	Low	Some Concerns	Low	Low	High
omersan-Karakaya	Low	Low	Low	Low	Low	Low	Low
OVID-19-MCS	High	Low	Low	Low	Low	High	High
ldiz E et al	High	Some Concerns	Low	Some Concerns	Low	High	High
YTOCOV-19	High	Some Concerns	Low	Some Concerns	Low	High	High
	High	Some Concerns	Low	Some Concerns	Low		
ashtani ED et al	Inigh	Some Concerns	1. The second			High	High
	Low	1 mm	Low	Low			
LPS-COVID	Low	Low	Low	Low	Low	Low	Low
Igahtani FD et al LPS-COVID t10933-10987-COV-20145 ICACS	Low Low High	Low Low Some Concerns	Low Low Low	Low Low Some Concerns	Low	Low Low High	Low High



PennCCP2	High	Some Concerns	Low	Some Concerns	Low	High	High
Toroghi N et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Isa F et al	Low	Low	Low	Low	Low	Low	Low
MOVe-OUT	Low	Low	Low	Low	Low	Low	Low
Weinreich_2	Low	Low	Low	Low	Low	Low	Low
Beigmohammadi MT et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Sarhan RM et al	High	Some Concerns	Low	Some Concerns	Low	High	High
AP-014	High	Some Concerns	Low	Some Concerns	Low	High	High
Asgardoon M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Kharazmi AB et al	High	Some Concerns	Low	Some Concerns	Low	High	High
COMBAT-COVID	Low	Low	Low	Low	Low	Low	Low
ACPREGCOV	Low	Low	Low	Low	Low	Low	Low
X-Covid 19	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Holubar M et al	Low	Low	Low	Low	Low	Low	Low
Malaysian Favipiravir Study	Low	Some Concerns	Low	Some Concerns	Low	Low	High
George C et al TSUNAMI	Low	Low Some Concerns	Low	Low Some Concerns	Low	Low	Low
	SI C P	Low	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Low	Low	Low	High Low
COnV-ert & CoV-Early Raghavan K et al	Low High	Some Concerns	Low	Some Concerns	Low	High	High
Shohan et al	High	Some Concerns	Low	Some Concerns	Low	High	High
CSSC-004	Low	Low	Low	Low	Low	Low	Low
Cannellotto M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
CRITICAL	Low	Low	Low	Low	Low	Low	Low
Regkirona_Part2							
PINETREE	Low	Low	Low	Low	Low	Low	Low
BUCOSARS	Low	Low	Low	Low	Low	Low	Low
BK-CLV-201	High	Some Concerns	Low	Some Concerns	Low	High	High
HIGHLOWDEXA	High	Some Concerns	Low	Some Concerns	Low	High	High
DEFINE	High	Some Concerns	Low	Some Concerns	Low	High	High
Ahmad B et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Pushkala et al.	High	Some Concerns	Low	Some Concerns	Low	High	High
Baxter AL et al	High	Some Concerns	Low	Some Concerns	Low	High	High
FAVI-COV-US201	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Kazempour et al.	High	Some Concerns	Low	Some Concerns	Low	High	High
Kerget B et al	High	Some Concerns	Low	Some Concerns	Low	High	High
WINCOVID	High	Some Concerns	Low	Some Concerns	Low	High	High
Poleti ML et al	Low	Low	High	Low	Low	High	High
COP20	Low	Some Concerns	Low	Some Concerns	Low	Low	High
WHIP COVID-19	Low	Low	Low	Low	Low	Low	Low
TOGETHER 2	Low	Low	Low	Low	Low	Low	Low
CONTAIN COVID-19	Low	Low	Low	Low	Low	Low	Low
COVIDENZA	Low	Some Concerns	Low	Some Concerns	Low	Low	High
COLCOVID	Low	Low	Low	Low	Low	Low	Low
Alsultan M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
OPTIMISE-C19	Low	Low	Low	Low	Low	Low	Low
COVID-Omega-F	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Majidi N et al	High	Low	Low	Low	Low	High	High
ICU-VR	High	Some Concerns	Low	Some Concerns	Low	High	High
ALLIANCE	High	Some Concerns	Low	Some Concerns	Low	High	High
PROTECT-EHC	Low	Low	Low	Low	Low	Low	Low
UNAB-003	High	Some Concerns	Low	Some Concerns	Low	High	High
Tolouian R et al	Low	Low	Low	Low	Low	Low	Low
INSPIRATION/INSPIRATION-S	Low	Low	Low	Low	Low	Low	Low
Abuhasira R et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Hu Q et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Avi-Mild	Low	Low	Low	Low	Low	Low	Low
APLICOV-PC	Low	Low	Low	Low	Low	Low	Low
MARIPOSA	Low	Some Concerns	Low	Some Concerns	Low	Low	High
IMPACT Covid19DPP4i	High	Some Concerns	Low	Some Concerns Some Concerns	Low	High	High
	High	Some Concerns	Low	and the second se	Low	High	High
ABB-COVID19 COVID MED	Low	Low	Low	Low	Low	Low	Low
COVID MED Naik NB et al	Low High	Some Concerns	Low	Low Some Concerns	Low	High	Low High
ACTIV-4a	Low	Low	Low	Low	Low	Low	Low
CATCO	Low	Some Concerns	Low	Some Concerns	Low	Low	High
MEFECOVID-19	Low	Low	Low	Low	Low	Low	Low
Rondanelli M et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
De Santis GC et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Murugesan H et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Manomaipiboon A et al	Low	Low	Low	Low	Low	Low	Low
DOXPREVENT.ICU	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Pourdowlat G et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Chupp G et al	Low	Low	Low	Low	Low	Low	Low
NACOVID	Low	Some Concerns	Low	Some Concerns	Low	Low	High
MEDIC-LAUMC	High	Low	Low	Low	Low	High	High
REsCue	Low	Low	Low	Low	Low	Low	Low
ITAC	Low	Low	Low	Low	Low	Low	Low
EPIC-HR	Low	Low	Low	Low	Low	Low	Low
I-TECH	Low	Some Concerns	Low	Some Concerns	Low	Low	High
FORCE	Low	Low	Low	Low	Low	Low	Low
Caims DM et al	Low	Low	Low	Low	Low	Low	Low
PHYDRA	Low	Low	Low	Low	Low	Low	Low
Nekoukar Z et al	Low	Low	Low	Low	Low	Low	Low
RAAS-COVID-19	High	Some Concerns	Low	Some Concerns	Low	High	High
SpiroCOVID19	Low	Low	Low	Low	Low	Low	Low
CR216-21	Low	Some Concerns	Low	Some Concerns	Low	Low	High
EPICOS	Low	Low	Low	Low	Low	Low	Low
COPERNICO	Low	Some Concerns	Low	Some Concerns	Low	Low	High
PROTECT-Patient trial	High	Some Concerns	Low	Some Concerns	Low	High	High
Singh H et al	Low	Low	Low	Low	Low	Low	Low
Barzin Tond S et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
RECOVERY	High	Some Concerns	Low	Some Concerns	Low	High	High

RUXCOVID-DEVENT	Low	Low	Low	Low	Low	Low	Low
SAC-COVID	Low	Low	Low	Low	Low	Low	Low
323Oct2020	Low	Low	Low	Low	Low	Low	Low
hafoon M et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ORTIVID	Low	Low	Low	Low	Low	Low	Low
OVERAGE	Low	Some Concerns	Low	Some Concerns	Low	Low	High
assaniazad M et al	Low	Some Concerns	Low	Some Concerns	Low	Low	High
REATHE	Low	Low	Low	Low	Low	Low	Low
aronova TL et al	High	Some Concerns	Low	Some Concerns	Low	High	High
eCOVID	Low	Low	Some Concerns	Low	Low	Some Concerns	Some Concerns
OVID-VIT-D	High	Some Concerns	Low	Some Concerns	Low	High	High
DGHETER - Ivermectin	Low	Low	Low	Low	Low	Low	Low
LARE	Low	Low	Low	Low	Low	Low	Low
rennan CM et al	Low	Low	Some Concerns	Low	Low	High	High
B 3305	Low	Low	Low	Low	Low	Low	Low
barsi P et al	High	Some Concerns	Low	Some Concerns	Low	High	High
thi-Kazerooni M et al belatto CK et al	High Some Concerns	Low	Low	Low	Low	High Some Concerns	High
		Low	Low	Low	Low		Some Concerns
FESAVER	Low	Low	Low	Low	Low	Low	Low
	1000	Low		Low		Low	
ACCPT	Low	Low	Low	Low	Low	Low	Low
PC-SARS	Low	Low	Low	Low	Low	Low	Low
errick J et al	Low	Low	Low	Low	Low	Low	Low
tem G et al	Low	Low	Low	Low	Low	Low	Low
owdhury FR et al	Low	Low	Low	Low	Low	Low	Low
ACO-COVID	Low	Low	Low	Low	Low	Low	Low
COT	Low	Low	Low	Low	Low	Low	Low
-CLARITY	Low	Low	Low	Low	Low	Low	Low
go EM et al RUCONPLASMA	Low	Low	Low	Low	Low	Low	Low
	Low	Low	Low	Low	Low	Low	Low
P-COVID-19	Low	Low	Low	Low	Low	Low	Low
ONFIDENT	Low	Low	Low	Low	Low	Low	Low
C/COVID-19	Low	Low	Low	Low	Low	Low	Low
DP-COVID-19	Some Concerns	Low	Low	Low	Low	Some Concerns	Some Concerns
CAP	Low	Low	Low	Low	Low	Low	Low
	Low	Some Concerns	Low	Some Concerns	Low	Low	High
EMAP-CAP	Low	Some Concerns	Low	Some Concerns	Low	Low	High
OPE - Coalition V	Low	Low	Low	Low	Low	Low	Low
Qahtani M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
mehecati	High	Some Concerns	Low	Some Concerns	Low	High	High
	Low	Some Concerns	Low	Some Concerns	Low	Low	High
o Hetal	High	Some Concerns	Low	Some Concerns	Low	High	High
orial Fletal	High	Some Concerns	Low	Some Concerns	Low	High	High
IpaCt-RT	High	Some Concerns	Low	Some Concerns	Low	High	High
OVIPOC	High	Some Concerns	Low	Some Concerns	Low	High	High
afeDrop	Some Concerns	Low	Low	Low	Low	Some Concerns	Some Concerns
edondo-Calvo FJ et al	Low	Low	Some Concerns	Low	Low	High	High
ANDLE	Low	Low	Low	Low	Low	Low	Low
OVID-Compromise	Low	Low	Low	Low	Low	Low	Low
тсн	Low	Low	Low	Low	Low	Low	Low
imar D et al	High	Some Concerns	Low	Some Concerns	Low	High	High
OVID-19-HBO	Low	Some Concerns	Low	Some Concerns	Low	Low	High
OVASE	High	Some Concerns	Low	Some Concerns	Low	High	High
CT-MP-COVID-19	Low	Low	Low	Low	Low	Low	Low
OPLA-II	Low	Some Concerns	Low	Some Concerns	Low	Low	High
oppock D et al	High	Some Concerns	Low	Some Concerns	Low	High	High
adavi M et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ROVENT	Low	Some Concerns	Low	Some Concerns	Low	Low	High
ahwani S et al	High	Some Concerns	Low	Some Concerns	Low	High	High
ostafaie A et al	7.4					NA	NA
LVERBULLET						NA	NA
-2020-785-176						NA	NA
S-US-553-9020						NA	NA
AWn-AZITHRO	Low	Some Concerns	Low	Some Concerns	Low	Low	High
W-MSC	Low	Low	Low	Low	Low	Low	Low
oVIP	Low	Low	Low	High	High	High	High
izadeh N et al	Some Concerns	Low	Low	Low	Low	Some Concerns	Some Concerns
llo	Low	Low	Low	Low	Low	Low	Low
		Low	Low	Low	Low	Low	Low
211-4	Low	LOW				1 1 m m	Low
castri E et al	Low	Low	Low	Low	Low	Low	
castri E et al ROTHROMCOVID			Low Low	Low Some Concerns	Low	Low	High
castri E et al ROTHROMCOVID DVID-HEP	Low	Low					
castri E et al ROTHROMCOVID DVID-HEP FU-2020-0707	Low Low	Low Some Concerns	Low	Some Concerns	Low	Low	High High Low
eastri E et al KOTHROMCOVID SVID-HEP U-2020-0707 ANTICO	Low Low Low	Low Some Concerns Some Concerns Low Some Concerns	Low Low Low Low	Some Concerns Some Concerns Low Some Concerns	Low Low	Low Low	High High
eastri E et al KOTHROMCOVID SVID-HEP U-2020-0707 ANTICO	Low Low Low Low	Low Some Concerns Some Concerns Low	Low Low Low	Some Concerns Some Concerns Low	Low Low Low	Low Low Low	High High Low
castri E et al ROTHROMCOVID DVID-HEP TU-2020-0707 ANTICO SSC-001	Low Low Low Low Low	Low Some Concerns Some Concerns Low Some Concerns Low Low	Low Low Low Low	Some Concerns Some Concerns Low Some Concerns	Low Low Low	Low Low Low Low	High High Low High
castri E et al XOTHROMCOVID 2010-HEP TU-2020-0707 ANTICO SSC-001 Make H et al	Low Low Low Low Low	Low Some Concerns Some Concerns Low Some Concerns Low	Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low	Low Low Low Low Low	Low Low Low Low Low	High High Low High Low
eastri E et al IOTHROMCOVID VVID-HEP U-2020-0707 INITICO ISC-001 Make H et al U-COV	Low Low Low Low Low Low Low	Low Some Concerns Some Concerns Low Some Concerns Low Low	Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Low	Low Low Low Low Low Low	Low Low Low Low Low Low	High High Low High Low Low
aastri E et al KOTHROMCOVID SVID-HEP U-2020-0707 NITICO SSC-001 Akae H et al LU-COV	Low Low Low Low Low Low High	Low Some Concerns Low Some Concerns Low Low Some Concerns	Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Low Some Concerns	Low Low Low Low Low Low Low	Low Low Low Low Low High	High High Low High Low Low High
eastri E et al XOTHROMCOVID 2010-HEP TU-2020-0707 ANTICO SSC-001 Auke H et al LU-COV Ahman SMA et al CCTG-COVID	Low Low Low Low Low Low High High	Low Some Concerns Low Some Concerns Low Low Some Concerns Low	Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Low Some Concerns Low	Low Low Low Low Low Low Low	Low Low Low Low Low Low High High	High High Low High Low Low High High
castri E et al X0THROMCOVID XVID-HEP U-2220-0707 ANTICO SSC-001 Muse H et al LU-COV shman SMA et al CCTD-COVID SPIRE	Low Low Low Low Low Low Low High High Low	Low Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low	Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Low Some Concerns Low Low	Low Low Low Low Low Low Low Low Low	Low Low Low Low Low High High	High High Low High Low Low High High Low
castri E et al NOTH:ROMCOVID SVID-HEP TU-2020-0707 ANTICO SSC-001 Jukae H et al LU-COV Johnan SMA et al LCTIC-COVID SPIRE SC-006	Low Low Low Low Low Low High High Low Low	Low Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Low	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low	Low Low Low Low Low Low Low Low Low	Low Low Low Low Low Low High High Low Low	High High Low High Low High High Low Low
castri E et al X0TH:ROMCOVID VUID-HEP FU-2020-0707 ANTICO SSC-001 ukae H et al LLCOV Maman SMA et al XCTIC-COVID SPIRE GC-006 EPAVID-19	Low Low Low Low Low Low Low High High Low Low Low Low	Low Some Concerns Low Some Concerns Low Low Some Concerns Low Low Low Low Low Some Concerns	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Low Some Concerns Low Low Low Low Some Concerns	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low Low High High Low Low Low Low Low	High High Low High Low High High Low Low Low Low High
castri E et al X0TH:ROMCOVID 2010-HEP TU-2020-0707 ANTICO SSC-001 Kushe H et al LLU-COV Anhman SMA et al CCTIC-COVID SPIRE GC-006 EPAVID-19 0 COV/ED	Low Low Low Low Low Low High High High Low Low Low Low Low High	Low Some Concerns Low Some Concerns Low Low Some Concerns Low Low Low Low Some Concerns Some Concerns Some Concerns	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Some Concerns Some Concerns	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low High High Low Low Low Low Low Low High	High High Low High Low Low High High Low Low High High
castri E et al XOTH:ROMCOVID VID-HEP TU-2020-0707 ANTICO SSC-001 ukae H et al LU-COV Ahman SMA et al VCTC-COVID SPIRE GC-006 EPAVID-19 0 COV-ED Iasia-Keever MA et al	Low Low Low Low Low Low High High Low Low Low Low High High	Low Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Low Low Low Some Concerns Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Low Low Low Low Low Low Some Concerns Some Concerns Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low High High High Low Low Low Low High High	High High Low High Low Low High High Low Low High High High
cashi E et al XOTHROMCOVID XVID-HEP TU-2020-0707 ANTICO SSC-001 Jakae H et al LU-COV Jahman SMA et al CTD-COVID SSPIRE SC-006 SSVID-19 D COV-ED Jasia-Keever MA et al RED-TRIAL	Low Low Low Low Low Low Low High High Low Low Low Low Low Low Low	Low Some Concerns Low Some Concerns Low Low Some Concerns Low Low Low Low Low Some Concerns Some Concerns Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Some Concerns Some Concerns Low Low	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low Low High High Low Low Low Low Low High High High	High High Low High Low Low High High Low Low Low High High High High Low
castri E et al X0TH:ROMCOVID 2010-HEP TU-2020-0707 ANTICO SSC-001 Kuke H et al LLU-COV ahman SMA et al CCTC-C:OVID SPIRE GC-006 EPAVID-19 0 COV-ED Ilasia-Kever MA et al ARED-TRIAL mas BE et al	Low Low Low Low Low Low High High Low Low Low High High High Low Low	Low Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Low Come Concerns Some Concerns Some Concerns Low Low Low Low	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Some Concerns Some Concerns Low Low Low	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low Low High High Low Low High High Low Low Low Low	High High Low High Low High High Low Low Low High High High Low Low Low
castri E et al NOTH:ROMCOVID SVID-HEP TU-2020-0707 ANTICO SSC-001 ukae H et al LU-COV Minama SMA, et al LCTIC-COVID SPIRE SC-006 EPAVID-19 O COV-ED Iassis-Keever MA et al RRED-TIKIAL nize BE et al RUCK	Low Low Low Low Low High High Low Low Low High High High High	Low Some Concerns Low Some Concerns Low Low Some Concerns Low Low Low Low Some Concerns Some Concerns Low Low Some Concerns Low Low Low Low	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Low Low Low Low Low Some Concerns Some Concerns Low Low Some Concerns Some Concerns Some Concerns	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low High High High Low Low High High Low High	High High Low High Low Low High High Low Low High High High High Low Low High
castri E et al XOTH:ROMCOVID VUID-HEP TU-2020-0707 ANTICO SSC-001 ukae H et al LU-COV ahman SMA et al XCTIC-COVID SPIRE 6C-006 EPAVID-19 0 COV-ED Ilinais-Keever MA et al ARED-TRIAL ARED-TRIAL TRUCK	Low Low Low Low Low Low Low High High Low Low Low Low High High High Low High Low High Low	Low Some Concerns Low Some Concerns Low Low Some Concerns Low Low Low Some Concerns Some Concerns Low Low Low Low Low Low Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Some Concerns Some Concerns Low Low Some Concerns Low Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low High High Low Low Low Low High High High High Low Low High Low Low Low Low	High High Low High Low High High Low Low Low Low High High High High Low Low Low Low
icastri E et al NOTH:ROMCOVID VID-HEP TU-2020-0707 ANTICO SSC-001 Urkae H et al LLU-COV Ahman SMA et al ACTIC-COVID SIGNIE GC-008 EPAVID-19 O COV-ED Itasia-Keever MA et al ARED-TRIAL Noze BE et al TRUICK CTIV-6 ZEIJ, MIG	Low Low Low Low Low Low High High Low Low Low Low High High High High Low Low Low Low	Low Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Low Some Concerns Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Low Some Concerns Some Concerns Low Low Some Concerns Low Low Some Concerns Low Low	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low High High How Low Low Low Low Low Low Low Low High High Low Kigh Low Low	High High Low High Low Low High High Low Low High High High Low High Low High
CTT-4 Lisaht E et al ROTHROMCOVID OVID-HEP TU-2020-0707 ANTICO SSC-001 Uut-COV Juna H et al LU-COV Juna H et al LU-COV Juna H et al CCC-COVID SPIRE GC-006 EPANID-18 O COV-ED Illasis-Kever MA et al ARED-TRIAL SPRE BE et al RED-TRIAL SPRE BE et al CTV-6 ezal_Mid Ezal_Severe Spissewkinai_Treat	Low Low Low Low Low Low Low High High Low Low Low Low High High High Low High Low High Low	Low Some Concerns Low Some Concerns Low Low Some Concerns Low Low Low Low Some Concerns Some Concerns Low Low Low Low Low Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Some Concerns Some Concerns Low Low Some Concerns Low Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low High High Low Low Low Low High High High High Low Low High Low Low Low Low	High High Low High Low High High Low Low Low Low High High High High Low Low Low Low



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Mirahmadiza	adeb et al	Low	Low	Low	Low	Low	Low	Low
George et al		Low	Some Concerns	Low	Some Concerns	Low	Low	High
	1							
Rojas et al		Low	Some Concerns	Low	Some Concerns	Low	Low	High
Bargay-Lleor		High		Low	Some Concerns	Low	High	High
ETHIC		High		Low	Some Concerns	Low	High	High
OVID		Low	Some Concerns	Low	Some Concerns	Low	Low	High
Mukae H et a	al	Low	Low	Low	Low	Low	Low	Low
Khan et al		High	Some Concerns	Low	Some Concerns	Low	High	High
Moslemi et a	al	High	Some Concerns	Low	Some Concerns	Low	High	High
Stambouli et	tal	Low	Low	Low	Low	Low	Low	Low
Stambouli et	tal	Low	Low	Low	Low	Low	Low	Low
Alemany et a	al	Low	Low	Low	Low	Low	Low	Low
McMahon et	tal	Low	Low	Low	Low	Low	Low	Low
Karampitsak	ios et al	High	Some Concerns	Low	Some Concerns	Low	High	High
Carvalho Ne	uenschwander et al	Low	Low	Low	Low	Low	Low	Low
Amoushahi e	et al	High	Low	Low	Low	Low	High	High
Castro-Rodri	iquez et al	High	Some Concerns	High	Some Concerns	Low	High	High
Terada et al	-	High		Low	Some Concerns	Low	High	High
Medhat et al		Low	Some Concerns	Low	Some Concerns	Low	Low	High
Prasenohadi		Low	Low	Low	Low	Low	Low	Low
TACKLE		Low	Low	Low	Low	Low	Low	Low
TICO		Low		Low	Low	Low	Low	Low
Labro et al		Low	Low	Low	Low	Low	Low	Low
Askari rt al		Low	Low	Low	Low	Low	Low	Low
Dow et al		High	Low	Low	Low	Low	High	High
Cecconi et al		Low		Low	Some Concerns			
						Low	Low	High
Tirupakuzhi e	etai	Low	Some Concerns	Low	Some Concerns	Low	Low	High
Lau et al		Low	Low	Low	Low	Low	Low	Low
COVIT-TRIA	AL	High		Low	Some Concerns	Low	High	High
Karonova		High		Low	Some Concerns	Low	High	High
Bencheqrour	n	Low	Low	Low	Low	Low	Low	Low
Panatto		High	Some Concerns	Low	Some Concerns	Low	High	High
UW 20-535		High	Some Concerns	Low	Some Concerns	Low	High	High
Barnette		High	Low	Low	Low	Low	High	High
Saviano		High	Some Concerns	Low	Some Concerns	Low	High	High
Tobback		Low	Low	Low	Low	Low	Low	Low
Barrueco		Low	Low	Low	Low	Low	Low	Low
Zeyad		High		Low	Some Concerns	Low	High	High
Self		Low		Low	Low	Low	Low	Low
Kumar		High	Some Concerns	Low	Some Concerns	Low	High	High
Zou		High		Low	Some Concerns	Low	High	High
Tandon		Low	Low	Low	Low	Low	Low	Low
COVIDICUS		Low		Low	Low	Low	Low	Low
Dastenae Rabbani		High	Some Concerns	Low	Some Concerns	Low	High	High
		High	Some Concerns	Low		Low	High	High
Bharti								
		Low		Some Concerns	Low	High	High	High
Ojeda		High	Low	Low	Low	Low	High	High
Bozorgmehr		High High	Low Some Concerns	Low Low	Low Some Concerns		High High	High High
		High	Low	Low	Low	Low	High	High
Bozorgmehr	rguengoitia	High High	Low Some Concerns	Low Low	Low Some Concerns	Low Low	High High	High High
Bozorgmehr Romero-Ibar	rguengoitia	High High High	Low Some Concerns Some Concerns Low	Low Low Low	Low Some Concerns Some Concerns	Low Low Low	High High High	High High High
Bozorgmehr Romero-Ibar ACTIV-6 - Fli	rguengoitia	High High High Low	Low Some Concerns Some Concerns Low Low	Low Low Low Low	Low Some Concerns Some Concerns Low	Low Low Low Low	High High High Low	High High High Low
Bozorgmehr Romero-Ibar ACTIV-6 - Fli BLAZE-4	rguengoitia	High High High Low Low	Low Some Concerns Some Concerns Low Low	Low Low Low Low Low	Low Some Concerns Some Concerns Low Low	Low Low Low Low Low	High High High Low Low	High High High Low Low
Bozorgmehr Romero-Ibar ACTIV-6 - Flu BLAZE-4 PRANA	rguengoitia	High High High Low Low	Low Some Concerns Some Concerns Low Low Low Low	Low Low Low Low Low Low	Low Some Concerns Some Concerns Low Low Low	Low Low Low Low Low	High High High Low Low Low	High High Ligh Low Low
Bozorgmehr Romero-Ibar ACTIV-6 - Flu BLAZE-4 PRANA Aryan	rguengoitia	High High Low Low Low High High	Low Some Concerns Low Low Low Low Some Concerns	Low Low Low Low Low Low Low	Low Some Concerns Some Concerns Low Low Low Low	Low Low Low Low Low Low	High High Low Low Low High High	High High Low Low Low High
Bozorgmehr Romero-Ibar ACTIV-6 - Fli BLAZE-4 PRANA Aryan Cervero Abroug	rguengoitia luticazone	High High Low Low Low High High	Low Some Concerns Low Low Low Low Some Concerns Low	Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns	Low Low Low Low Low Low Low Low Low	High High Low Low High High High	High High Low Low High High High
Bozorgmehr Romero-Ibar ACTIV-6 - Fli BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - I	guengoitia luticazone	High High Low Low Low High High High Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns	Low Low Low Low Low Low Low Low Low	Low Some Concerns Some Concerns Low Low Low Some Concerns Low Some Concerns	Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High Low	High High Low Low High High High
Bozorgmehr Romero-Ibar ACTIV-6 - Fil BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - I PLATCOV - I	rguengoitia uticazone Iver Regen	High High Low Low High High High Low Low	Low Some Concerns Low Low Low Some Concerns Low Some Concerns Some Concerns	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Some Concerns	Low Low Low Low Low Low Low Low Low Low	High High Low Low High High High Low Low	High High Low Low Low High High High High
Bozorgmehr Romero-Ibar ACTIV-6 - Fli BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 Fogleman C	rguengoitia luticazone Iver Regen et al	High High Low Low High High Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	High High Low Low High High Low Low Low Low	High High Low Low Low High High High High Low
Bozorgmehr Romero-Ibar ACTIV-6 - Fil BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - I PLATCOV - I Fogleman C PanCOVID1	rguengoitia luticazone Iver Regen et al	High High Low Low High High Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Some Concerns Some Concerns Low Some Concerns	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Low Some Concerns Some Concerns Low Some Concerns	Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High Low Low Low Low	High High Low Low High High High High High Low High
Bozorgmehr Romero-Ibar ACTIV-6 - FH BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 Fogleman C PanCOVID 11 AGILE	rguengoitia luticazone Iver Regen et al	High High Low Low High High Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Low Some Concerns Some Concerns Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Some Concerns Some Concerns Some Concerns Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High Low Low Low Low Low Low	High High Low Low Low High High High High Low High Low
Bozorgmehr Romero-Ibar ACTIV-6 - Fil BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 PLATCOV - 1 Fogleman C PanCOVID AGILE D-COVID	rguengoitia luticazone Iver Regen et al	High High Low Low High High Low Low Low Low Low	Low Some Concerns Low Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High Low Low Low Low Low Low Low	High High Low Low High High High High Low High Low High
Bozorgmehr Romero-Ibar ACTIV-6 - FII BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - I PLATCOV - I PLATCOV - I Fogleman C PanCOVID1 AGILE D-COVID IRICT	rguengoitia luticazone Iver Regen et al 9	High High Low Low High High Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Some Concerns Some Concerns Low Some Concerns Low Low	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High Low Low Low Low Low Low Low Low Low	High High Low Low Low High High High High Low High Low Low
Bozorgmehr Romero-Ibar ACTIV-8 – Fil BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV – I Fogleman C PanCOVID 1 AGILE D-COVID IRICT Choudhary F	rguengoitia luticazone Iver Regen et al 9 R et al	High High Low Low Low High High Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Low Some Concerns Some Concerns Low Some Concerns Low Low Low Low Some Concerns	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Low Some Concerns Some Concerns Low Some Concerns Low Low Low	Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High Low Low Low Low Low Low Low Low Low	High High Low Low Low High High High Low High Low High Low High
Bozorgmehr Romero-Ibar ACTIV-6 - Fil BLAZE 4 PRANA Aryan Cervero Abroug PLATCOV - I PLATCOV - I PLATCO	rguengoitia luticazone Iver Regen et al 9 R et al	High High Low Low High High Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Low Some Concerns Some Concerns Some Concerns Some Concerns Some Concerns Some Concerns	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Low Some Concerns Some Concerns Some Concerns Some Concerns Some Concerns Some Concerns Some Concerns Some Concerns	Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High High High Low High Low High Low High
Bozorgmehr Romero-Ibar ACTIV-6 - Fih BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 PLATCOV - 1 PLATCOV - 1 PATCOV - 1 PATCOV - 1 PATCOV - 1 PATCOV - 1 Fogleman C PanCOVID 11 AGILE D-COVID IRICT Choudhary F Khodashahi AAAT0535	rguengoitia luticazone liver Regen et al 9 R et al R et al	High High Low Low High High Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Low	Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High High High Low High Low High Low High Low High
Bozorgmehr Romero-Ibar ACTIV-6 - Fil BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 PLATCOV - 1 PanCOVID AGILE D-COVID IRICT Choudhary Fi Khodashahi AAT0535	rguengoitia luticazone lver Regen et al 9 R et al R et al	High High Low Low High High Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Low Low Low Low Low Low Low Low Low	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low	Low	High High Low Low Low High Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High High High Low High Low High Low Low Low
Bozorgmehr Romero-Ibar ACTIV-6 - Fil BLAZE 4 PRANA Aryan Cervero Abroug PLATCOV - I PLATCOV - I SOUGHAN F COVUD I RICT Choudhany F Khodashahi AATOSS ACTIV-3/TIC SOITAI R et al RICVIDE I PLATCOV - I	rguengoitia luticazone lver Regen et al 9 ₹ et al R et al 20 al	High High Low Low High High Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Some Concerns	Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Low Some Concerns Low Low Low Low Low Some Concerns Low Some Concerns Low Low Some Concerns Low Some Concerns Some Concerns Some Concerns Some Concerns Low Some Concerns Low Some Concerns Low Low Some Concerns Some Concerns Some Concerns Low Low Low Some Concerns Some Concerns Some Concerns Some Concerns Low Low Low Some Concerns Some Concerns Some Concerns Some Concerns Some Concerns Some Concerns Low Some Concerns Some Con	Low	High High Low Low Low High High Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High High High Low High Low High Low Low High Low Low High
Bozorgmehr Romero-Ibar ACTIV-6 - Fin BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV-1 PLATCOV-1 PLATCOV-1 PLATCOV-1 PLATCOV-1 PACOVID IRICT Choudhary Fi Khodashahi AAT0535 ACTIV-3TIC Soltani R et a ANACONDA	rguengoitia luticazone lver Regen et al 9 ₹ et al R et al 20 al	High High Low Low High High Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Low Low Some Concerns Low	Low	Low Some Concerns Low Low Low Low Some Concerns Low Low Cow Low Some Concerns Some C	Low	High High Low Low Low High High Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High High High Low High Low High Low High Low High Low High High
Bozorgmehr Romero-Iba- BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 PLATCOV - 1 PanCOVID AGILE D-COVID IRICT Choudhary F Khodashahi AAT0535 ACTIV-3TIC Soltani R et a ANACONDA BTI-202	rguengoitia luticazone lver Regen et al 9 R et al R et al 20 al	High High Low Low High High High Low Low Low Low Low Low Low Low Low High Low Low High Low Low Low	Low Some Concerns Low Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low	Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low	Low	High High High Low Low Low High Low Low Low Low Low Low Low Low Low Low	High High Low Low Low High High High High Low High Low High Low High Low High Low High Low Low Low Low
Bozorgmehr Romero-Ibar ACTIV-6 - Fil BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 PLATCOV - 1 PLATCOV - 1 PLATCOV - 1 PATCOV - 1 PATCOV - 1 PLATCOV - 1 Fogleman C PanCOVID1 RICT Choudhary F Khodashahi AAT0535 ACTIV-3/TIC Soltani R et a ANACONDA BTI-202 ReCOVery-5	rguengoitia luticazone lver Regen et al 9 R et al R et al 20 al	High High Low Low High High Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Low Cow Some Concerns Some Concerns Some Concerns Some Concerns Low Some Concerns Some Concerns Low Some Concerns Some Conce	Low	Low Some Concerns Low Low Low Some Concerns Low Low Some Concerns Low Low Some Concerns Some Concerns Low Low Some Concerns Some Concerns Low Some Concerns Low Some Concerns Some Concerns Low Some Concerns Low Some Concerns	Low	High High Low Low Low High High Low Low Low Low Low Low Low Low Low Low	High High High Low Low High High High High High Low High Low High Low High Low High Low Low High Low Low Low
Bozorgmehr Romero-Ibar ACTIV-6 - Fin BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 PLATCOV - 1 PLATCO	rguengoitia luticazone Iver Regen et al 9 R et al R et al NO al N	High High Low Low High High Low Low Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Come Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low Low Com Some Concerns Low Low Com Com Some Concerns Low Low Low Com	Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low Low Some Concerns Low Low Some Concerns Some Concerns Some Concerns Low Low Some Concerns Low Low Low Some Concerns Low Low Some Concerns Low	Low	High High Low Low Low High Low Low Low Low Low Low Low Low Low Low	High High High Low Low High High High High Low High Low High Low High Low High Low High Low Low Low Low Low Low Low Low
Bozorgmehr Romero-Iba- BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV -1 PLATCOV -1 PanCOVID AGILE D-COVID ROUGHIN AGILE D-COVID ROUGHIN AAT0535 ACTIV-3TIC Soltani R et a ANCONDA BTI-202 ReCOVerJS MOVe-0UT-	rguengoitia luticazone Iver Regen et al 9 R et al R et al NO al N	High High Low Low High High High Low Low Low Low Low Low Low Low Low High Low Low High Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low	Low	Low Some Concerns Low Low Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low	Low	High High High Low Low High High Low Low Low Low Low Low Low Low High Low High Low High Low Low High Low Low Low Low Low Low Low Low Low Low	High High High Low Low High High High High Low High Low High Low High Low High Low High Low High Low High Low High Low Low Low Low
Bozorgmehr Romero-Ibar ACTIV-6 - Fit BLAZE-4 PRANA Aryan Cervero Abroug PLATCOV - 1 PLATCOV - 1 PLATCO	rguengoitia luticazone Iver Regen et al 9 R et al R et al SIRIO 5IRIO	High High Low Low Low High High Low Low Low Low Low Low Low Low Low High Low Low High Low Low Low Low Low Low Low Low	Low Some Concerns Low Low Low Some Concerns Some Concerns Low Some Concerns Some Concerns Low Some Concerns Low Low Some Concerns Low	Low	Low Some Concerns Low Low Low Low Commons Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Low Some Concerns Some Concerns Low	Low	High High Low Low Low High High Low Low Low Low Low Low Low Low Low Low	High High High Low Low High High High High Low Low High Low High Low High Low Low Low Low Low Low Low Low Low
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Main findings

Corticosteroids

<u>See Summary of findings Table 1, Appendix 1</u>

We identified 17 RCTs including 9,485 participants in which systemic corticosteroids (dexamethasone, methylprednisolone, or hydrocortisone) were compared against standard of care or other treatments. Thirteen of these trials provided information on mortality for the corticosteroids against standard of care comparison. The RECOVERY trial was the biggest with 2,104 patients assigned to dexamethasone and 4,321 to standard of care. Sixteen studies included patients with severe to critical disease, as shown by the fact that mortality in the control groups ranged from 14.2% to 61.4%, and one study included hospitalized patients without respiratory failure. In the RECOVERY trial, a subgroup analysis which stratified patients by the amount of baseline respiratory support they received, showed significant differences favoring those with oxygen requirements. However, as mortality was high in the subgroup of patients that did not receive baseline oxygen treatment (14%), we decided to adopt a conservative approach and include the primary analysis considering all randomized patients. In addition, we identified eight studies including 2,490 patients in which different corticosteroid dosage schemes were compared and one study including 42 patients in which high dose steroids were compared to tocilizumab. Our results showed:

- Corticosteroids probably reduce mortality, RR 0.90 (95%CI 0.80 to 1.01); RD -1.6% (95%CI -3.2% to 0.2%); Moderate certainty ⊕⊕⊕○ (Figure 2)
- Corticosteroids probably reduce invasive mechanical ventilation requirement, RR 0.87 (95%CI 0.73 to 1.04); RD -2.2% (95%CI -4.7% to 0.7%); Moderate certainty ⊕⊕⊕○
- Corticosteroids may improve time-to-symptom resolution, RR 1.19 (95%CI 0.95 to 1.5); RD 11.5% (95%CI -3% to 30%); Low certainty ⊕⊕○○
- Corticosteroids may not significantly increase the risk of severe adverse events, RR 0.89 (95%CI 0.68 to 1.17); RD -1.1% (95%CI -3.3% to 1.7%); Low certainty ⊕⊕⊖○
- Results were consistent with trials in which corticosteroids were used to treat non COVID-19 patients with ARDS. No significant differences between subgroups of studies using different corticosteroids were observed. (Figures 3 and 4)
- High-dose corticosteroids (i.e., dexamethasone 12 mg a day) may not reduce mortality compared to standard-dose corticosteroids (i.e., dexamethasone 6 mg a day), RR 0.97 (95%CI 0.78 to 1.21); RD -0.5% (95%CI -3.5% to 3.4%); Low certainty ⊕⊕○○ (Figure 5) (based on low risk of bias studies)
- It is uncertain if high-dose corticosteroids (i.e., dexamethasone 12 mg a day) increase or reduce mechanical ventilation compared to standard-dose corticosteroids (i.e., dexamethasone 6 mg a day), RR 0.94 (95%CI 0.41 to 2.11); RD -1% (95%CI -10.2% to 19.2%); Very low certainty ⊕○○○



- High-dose corticosteroids (i.e., dexamethasone 12 mg a day) may not increase symptom resolution or improvement compared to standard-dose corticosteroids (i.e., dexamethasone 6 mg a day), RR 0.99 (95%CI 0.9 to 1.08); RD -0.6% (95%CI -5.5% to 4.9%); Low certainty ⊕⊕○○
- High-dose corticosteroids (i.e., dexamethasone 12 mg a day) may not increase severe adverse events compared to standard-dose corticosteroids (i.e., dexamethasone 6 mg a day), RR 0.82 (95%CI 0.6 to 1.11); RD -1.8% (95%CI -4.1% to 1.1%); Low certainty ⊕⊕○○

Figure 2. All-cause mortality in RCTs comparing corticosteroids with standard of care for treatment of patients with COVID-19

Study	TE seTE	Risk Ratio	RR 95%-C	Weight (fixed)	Weight (random)
RECOVERY - Dexa	-0.11 0.0476	10	0.89 [0.81; 0.98	63.3%	38.8%
GLUCOCOVID	0.15 0.5290		1.16 [0.41; 3.27	0.5%	1.1%
Metcovid	-0.03 0.1299	+	0.97 [0.75; 1.25	8.5%	14.2%
DEXA-COVID19	0.54 0.8797		1.71 [0.31; 9.61	0.2%	0.4%
REMAP-CAP	-0.17 0.1715	-4-	0.84 [0.60; 1.18	4.9%	9.2%
Steroids-SARI	-0.04 0.2621		0.96 [0.57; 1.60	2.1%	4.4%
COVID STEROID	1.03 0.7270		2.80 [0.67; 11.64	0.3%	0.6%
CoDEX	-0.09 0.0968	+	0.92 [0.76; 1.11	15.3%	21.1%
CAPE COVID	-0.64 0.3377		0.53 [0.27; 1.02	1.3%	2.7%
Edalatifard M et al (Tehran University of Medical Scier	nces) -1.99 0.7199	I	0.14 [0.03; 0.56	0.3%	0.6%
Tang X et al	-1.10 1.6187 —		0.33 [0.01; 7.96	0.1%	0.1%
Jamaati H et al	0.06 0.2217	- -	1.07 [0.69; 1.65	2.9%	5.9%
Ghanei M et al	-0.46 0.6316		0.63 [0.18; 2.18	0.4%	0.8%
Fixed effect model		6	0.90 [0.83; 0.97		
Random effects model		¢	0.90 [0.80; 1.01		100.0%
Heterogeneity: $I^2 = 17\%$, $\tau^2 = 0.0062$, $p = 0.27$		1 1 1 1 1			
		0.1 0.51 2 10			

Figure 3. All-cause mortality in RCTs comparing corticosteroids with standard of care for treatment of patients with COVID-19 or ARDS without COVID-19



Study	TE seTE	Risk Ratio	RR S	Weig 95%-Cl (fixe	
Population = COVID-19 pat RECOVERY - Dexamethaso GLUCOCOVID Metcovid DEXA-COVID19 REMAP-CAP Steroids-SARI COVID STEROID CoDEX CAPE COVID Edalatifard Tang Jamaati H et al Ghanei M et al Fixed effect model Random effects model			0.89 [0.81 1.24 [0.48 0.97 [0.75 1.71 [0.31 0.84 [0.60 0.96 [0.57 2.80 [0.67; 0.92 [0.76 0.53 [0.27 0.14 [0.03 0.33 [0.11 1.07 [0.69 0.63 0.90 [0.83 0.90 [0.83	; 3.19] 0.5 ; 1.25] 7.5 ; 9.61] 0.2 ; 1.18] 4.3 ; 1.60] 1.8 ; 1.60] 1.8 ; 1.102] 1.1 ; 0.56] 0.2 ; 7.96] 0.0 ; 1.65] 2.6 ; 2.18] 0.3 ; 0.97] 87.8	% 1.1% % 11.2% % 0.3% % 7.3% % 3.5% % 0.5% % 16.4% % 2.2% % 0.5% % 0.1% % 0.1% % 0.1% % 0.7%
Heterogeneity: $l^2 = 18\%$, $\tau^2 = 0.5$ Population = ARDS patient Meduri 2007 Rezk 2013 Steinberg 2006 Liu 2012 Tangyuo 2016 Villar 2020 Zhao 2014 Fixed effect model Random effects model Heterogeneity: $l^2 = 0\%$, $\tau^2 = 0$, l^2	-0.58 0.3147 -2.53 2.4204 0.02 0.2330 -1.11 0.7132 -0.15 0.1831 -0.42 0.1906 -0.17 0.3368		0.56 [0.30 0.08 [0.00 1.02 [0.65 0.33 [0.08 0.86 [0.60 0.66 [0.45 0.84 [0.43 0.77 [0.63 0.77 [0.63	; 1.04] 1.3 ; 9.19] 0.0 ; 1.61] 2.3 ; 1.34] 0.2 ; 1.23] 3.8 ; 0.96] 3.5 ; 1.63] 1.1 ; 0.94] 12.2	% 2.5% % 0.0% % 4.4% % 0.5% % 6.6% % 6.2% % 2.2%
Fixed effect model Random effects model Heterogeneity: $I^2 = 16\%$, $\tau^2 = 0$. Residual heterogeneity: $I^2 = 12\%$		0.1 1 10 10	0.88 [0.82 0.87 [0.79	; 0.95] 100.0 ; 0.96]	% 100.0%

Figure 4. All-cause mortality by type of corticosteroids in RCTs using comparison with standard of care for treatment of patients with COVID-19 or ARDS without COVID-19



Study	TE	seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
Drug = Dexamethasone RECOVERY - Dexamethasor			Į.		[0.81; 0.98]		29.0%
DEXA-COVID19 CoDEX		0.8797 0.0968	1.		[0.31; 9.61] [0.76; 1.11]		0.3% 16.4%
Villar 2020		0.1906	-		[0.45; 0.96]		6.2%
Jamaati H et al		0.2217	4		[0.69; 1.65]		4.8%
Fixed effect model			à		[0.82; 0.96]		
Random effects model			0		[0.82; 0.96]		56.6%
Heterogeneity: $I^2 = 0\%, \tau^2 = 0, p$	= 0.44						
Drug = Methylprednisone							
GLUCOCOVID	0.22	0.4806	_ 	1.24	[0.48; 3.19]	0.5%	1.1%
Metcovid		0.1299	Ť		[0.75; 1.25]		11.2%
Steroids-SARI		0.2621			[0.57; 1.60]		3.5%
Meduri 2007		0.3147			[0.30; 1.04]		2.5%
Rezk 2013 Steinborg 2006		2.4204 - 0.2330			[0.00; 9.19]		0.0% 4.4%
Steinberg 2006 Edalatifard		0.2330			[0.65; 1.61] [0.03; 0.56]		4.4%
Tang		1.6187			[0.01; 7.96]		0.1%
Fixed effect model	1.10	1.0107	4		[0.75; 1.09]		
Random effects model			4		[0.61; 1.13]		23.4%
Heterogeneity: $I^2 = 40\%$, $\tau^2 = 0.0$	0657, p =	0.11					
Drug = Hydrocortisone							
REMAP-CAP	-0.17	0.1715	4	0.84	[0.60; 1.18]	4.3%	7.3%
COVID STEROID	1.03	0.7270		2.80	[0.67; 11.64]		0.5%
CAPE COVID	-0.64	0.3377		0.53	[0.27; 1.02]		2.2%
Liu 2012		0.7132	<u>+</u>		[0.08; 1.34]		0.5%
Tangyuo 2016	-0.15	0.1831	1		[0.60; 1.23]		6.6%
Fixed effect model			2		[0.65; 1.01]		47 40/
Random effects model Heterogeneity: $I^2 = 36\%$, $\tau^2 = 0.0$	0464 p =	0.18		0.79	[0.57; 1.10]		17.1%
	0404, p -	0.10					
Drug = Budesonide	0.47	0.2200		0.04	10 42 4 601	4 40/	0.00/
Zhao 2014	-0.17	0.3368	Ţ		[0.43; 1.63]		2.2%
Fixed effect model Random effects model			X		[0.43; 1.63] [0.43; 1.63]		2.2%
Heterogeneity: not applicable			Ť	0.04	[0.45, 1.05]		2.2/0
Drug = Prednisolone							
Ghanei M et al	-0.46	0.6316		0.63	[0.18; 2.18]	0.3%	0.7%
Fixed effect model			\Rightarrow		[0.18; 2.18]	0.3%	
Random effects model			\rightarrow	0.63	[0.18; 2.18]		0.7%
Heterogeneity: not applicable							
Fixed effect model					[0.82; 0.95]		-
Random effects model			((0.87	[0.79; 0.96]		100.0%
Heterogeneity: $I^2 = 16\%$, $\tau^2 = 0.0$ Residual heterogeneity: $I^2 = 31\%$	0069, p =	0.25	01 01 1 10 40				
Residual neterogeneity: I ⁻ = 31%	₀, p = 0.1	∠ 0.0	001 0.1 1 10 10	00			



Figure 5. All-cause mortality in RCTs comparing high-dose corticosteroids (i.e., dexamethasone 12 mg a day) with standard-dose corticosteroids (i.e., dexamethasone 6 mg a day) in patients with COVID-19

Study	TE	seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
RoB = Moderate - His	in .		£1				
Ranjbar K et al	-0.68	0.3810		0.51	[0.24; 1.07]	3.3%	5.7%
HIGHLOWDEXA	-0.09	0.4978			[0.34; 2.42]	2.0%	3.5%
Dastenae	-0.32	0.4082 -			[0.33; 1.62]	2.9%	5.0%
Fixed effect model					[0.41; 1.07]	8.2%	
Random effects mod	el				[0.41; 1.07]		14.1%
Heterogeneity $T^2 = DW_0$		0.62					
RoB = Low							
COVID STEROID 2	-0.18	0.0995		0.84	[0.69; 1.02]	49.1%	33.5%
Maskin et al	0,00	0.2148		1.00	[0.66; 1.52]	10.5%	14.4%
Toroghi N et al	0.75	0.3526	· · · · ·	2.12	[1.06; 4.23]	3.9%	6.5%
RCT-MP-COVID-19	-0.19	0.3301		0.83	[0.43; 1.58]	4.5%	7.3%
COVIDICUS	-0.03	0.1431		0.97	[0.73; 1.28]	23.7%	24.2%
Fixed effect model			\diamond	D,92	[0.80; 1.06]	91.8%	
Random effects mod Helenogeneity, / ² = 42%		A1. p = 0.14			[0.78; 1.21]		85.9%
Fixed effect model			-0	0.90	[0.78; 1.03]	100.0%	
Random effects mod	el		\diamond		[0.76; 1.10]		100.0%
Heterogeneity: /2 = 26%		75. p = 0.22		212.0	and the second		A231242
Residual heterogeneity:			0.5 1 2				

In addition, one study that compared high dose corticosteroids (dexamethasone 20 mg a day) to tocilizumab reported higher mortality in patients treated with high dose corticosteroids.

Remdesivir

See Summary of findings Table 2, Appendix 1

We identified ten RCTs including 11,814 patients in which remdesivir was compared against standard of care or other treatments. In addition, we identified one study that compared different remdesivir dosage schemes. The WHO SOLIDARITY trial was the biggest with 4,146 patients assigned to remdesivir and 4,129 to standard of care. Five studies included patients with severe disease as shown by the fact that mortality in the control groups ranged from 8.3% to 12.6%, and three studies included non-severe patients with 2% or less mortality in the control arm. Our results showed:

• Remdesivir probably reduces mortality, RR 0.93 (95%CI 0.89 to 1.03); RD -1.1% (95%CI -1.8% to 0.5%); Moderate certainty ⊕⊕⊕○ (Figure 6)

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- Remdesivir probably reduces invasive mechanical ventilation requirement, RR 0.76 (95%CI 0.56 to 1.04); RD -4.2% (95%CI -7.6% to 0.7%); Moderate certainty ⊕⊕⊕○ (Figure 7)
- Remdesivir may improve time to symptom resolution, RR 1.1 (95%CI 0.96 to 1.28); RD 6% (95%CI -2.4% to 17%); Low certainty ⊕⊕○○ (Figure 8)
- Remdesivir may reduce hospitalizations in patients with recent onset mild, RR 0.28 (95%CI 0.11 to 0.75); RD -3.4% (95%CI -4.3% to -1.2%); Low certainty ⊕⊕○○
- Remdesivir may not increase the risk of severe adverse events, RR 0.77 (95%CI 0.46 to 1.29); RD -2.3% (95%CI -5.5% to 3%); Low certainty ⊕⊕○○

Figure 6. All-cause mortality with remdesivir use vs. standard of care in randomized control trials including COVID-19 patients

										Weight	Weight
Study	TE	seTE		Ris	sk Ra	tio		RR	95%-CI	(fixed)	(random)
ACTT-1	-0.34 0	0.1948		-				0.71	[0.49; 1.04]	6.1%	6.1%
CAP-China remdesivir 2	0.08 0	0.3554		-	-i+-	-		1.09	[0.54; 2.18]	1.8%	1.8%
SIMPLE 2	-0.43 (0.6651	-		+	-		0.65	[0.18; 2.40]	0.5%	0.5%
WHO SOLIDARITY - Remdes	ivir -0.07 (0.0523			10			0.93	[0.84; 1.03]	84,1%	84.1%
Mahajan L et al	0.57 0	0.6900		-			_	1.76	[0.46; 6.82]	0.5%	0.5%
Abd-Elsalam S et al	0.25 (0.4837		-	-			1.29	[0.50; 3.32]	1.0%	1.0%
Sarhan RM et al	0.30 (0.3360		-		-		1.35	[0.70; 2.60]	2.0%	2.0%
CATCO	0.03 0	0.2385		~	+	->		1.03	[0.65; 1.65]	4.0%	4.0%
Fixed effect model					-			0.93	[0.85; 1.03]	100.0%	
Random effects model Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0$, p	= 0.65		-	1	4	1	-	0.93	[0.85; 1.03]	-	100.0%
(1010)0g0(1010) / = 0.0, / = 0, p	0.00		0.2	0.5	đ	2	5				

Figure 7. Invasive mechanical ventilation requirements in RCTs comparing remdesivir with standard of care for treatment of patients with COVID-19



Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
ACTT-1	-0.55 0.1618		0.57	[0.42: 0.79]	9.7%	25.5%
CAP-China remdesivir 2	-0.61 0.4144		0.54	[0.24; 1.22]		10.3%
SIMPLE 2	-2.26 1.0920		0.10	[0.01; 0.89]	0.2%	2.0%
WHO SOLIDARITY - Remdesit	/ir -0.11 0.0549		0.89	[0.80; 1.00]	83.9%	33.4%
Mahajan L et al	0.75 0.8324		2.12	[0.41; 10.82]	0.4%	3.3%
Abd-Elsalam S et al	0.32 0.4426		1.38	[0.58: 3.27]	1.3%	9.4%
CATCO	-0.25 0.2881		0.78	[0.44; 1.37]	3.1%	16.2%
Fixed effect model		2	0.85	10.77; 0.941	100.0%	
Random effects model Heterogeneity: $l^2 = 57\%$, $\tau^2 = 0.07$	$20, \rho = 0.03$		0.76	[0.56; 1.04]	-	100.0%
Construction of the second s		0.1 0.51 2 10				

Figure 8. Symptom resolution or improvement in RCTs comparing remdesivir with standard of care for treatment of patients with COVID-19

Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
ACTT-1	0.28 0.0829	÷ <u> </u>	— 1.32	[1.12; 1.55]	27.9%	27.3%
CAP-China remdesivir 2	0.05 0.1159			[0.84; 1.32]	14.3%	20.6%
SIMPLE 2	0.11 0.0671	- <u>*</u>	1.12	[0.98; 1.28]	42.6%	30.9%
Sarhan RM et al	-0.10 0.1125		0.91	[0.73; 1.13]	15.2%	21.2%
Fixed effect model Random effects model Heterogeneity: $I^2 = 62\%$, τ^2		0.05		[1.03; 1.23] [0.96; 1.28]		 100.0%
		0.75 1	1.5			

Hydroxychloroquine and Chloroquine

See Summary of findings Table 3, Appendix 1

We identified 61 RCTs including 25,977 patients in which hydroxychloroquine or chloroquine were compared against standard of care or other treatments. The RECOVERY trial was the biggest with 1,561 patients assigned to dexamethasone and 3,155 to standard of care. In both the RECOVERY and SOLIDARITY trials, patients had severe disease as shown by the high mortality risk in control arms (24.9% and 9.2%, respectively). The remaining studies included patients with non-severe disease, as shown by the lower mortality risk in control arms, ranging from 0 to 5.2%.



Additionally, we identified nine studies in which hydroxychloroquine was used in healthy persons to prevent COVID-19 infection. Our results showed:

- Hydroxychloroquine or chloroquine probably does not increase mortality, RR 1.09 (95%CI 1 to 1.19); RD 1.4% (95%CI 0% to 3%); Moderate certainty ⊕⊕⊕○ (Figure 9)
- Hydroxychloroquine or chloroquine probably does not reduce invasive mechanical ventilation requirement; RR 1.08 (95%CI 0.93 to 1.25); RD 1.4% (95%CI -1.2% to 4.3%); Moderate certainty ⊕⊕⊕○
- Hydroxychloroquine or chloroquine probably does not improve time to symptom resolution, RR 1.01 (95%CI 0.93 to 1.1); RD 0.6% (95%CI -4.2% to 6.1%); Moderate certainty ⊕⊕⊕○
- Hydroxychloroquine or chloroquine may not have an important effect on COVID-19 symptomatic infection in exposed individuals, RR 0.87 (95%CI 0.65 to 1.15); RD 2.2% (95%CI -6.1% to 2.7%); Low certainty ⊕⊕○○ (Figure 10) (based on low risk of bias studies)
- Hydroxychloroquine or chloroquine may not significantly increase the risk of severe adverse events, RR 0.90 (95%CI 0.66 to 1.22); RD -1% (95%CI -3.5% to 2.2%); Low certainty ⊕⊕○○
- Hydroxychloroquine or chloroquine may not have an important effect on hospitalizations in patients with mild COVID-19, RR 0.82 (95%CI 0.61 to 1.1); RD 0.9% (95%CI -1.9% to 0.5%); Low certainty ⊕⊕○○

Figure 9. All-cause mortality in RCTs comparing hydroxychloroquine or chloroquine with standard of care in patients with COVID-19

Study	TE	seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
RECOVERY Hidrowichlarosting	0.07	0.0518		1 00	10.07: 4.101	67.1%	67.1%
RECOVERY - Hydroxychloroquine			И.		[0.97; 1.19]		
Cavalcanti et al		0.5454			[0.44; 3.70]		0.6%
COVID-19 PET		1,4109			[0.06; 15.81]		0.1%
Abd-Elsalam S et al	0.18	0.5883		1.20	[0.38; 3.80]	0.5%	0.5%
TEACH	0.06	0.5275		1.06	[0.38; 2.99]	0.6%	0.6%
WHO SOLIDARITY - HCQ	0.10	0.1367	+	1.11	[0.85; 1.45]	9.6%	9.6%
PETAL	-0.02	0.2677		0.98	[0.58; 1.65]	2.5%	2.5%
HYCOVID	-0.61	0.4913		0.54	[0.21; 1.42]	0.7%	0.7%
HYDRA	-0.08	0.1704		0.93	[0.66; 1.29]	6.2%	6.2%
Beltran-HCQ	-0.98	0.7806		0.37	[0.08; 1.73]	0.3%	0.3%
CLOROTRIAL	0.45	0.3527		1.57	[0.79; 3.13]	1.4%	1.4%
ProPAC-COVID	-0.78	1.2107	· · · · · · · · · · · · · · · · · · ·	0.46	[0.04; 4.92]	0.1%	0.1%
SEV-COVID	-0.62	0.6693		0.54	[0.15; 2.01]	0.4%	0.4%
COPE - Coalition V	-0.01	0.6301		0.99	[0.29; 3.41]	0.5%	0.5%
IRICT	0.38	0.1407	-	1.47	[1.11; 1.93]	9.1%	9.1%
Choudhary R et al	-1.10	1.6268 -		0.33	[0.01; 8.08]	0.1%	0.1%
Fixed effect model				1.09	[1.00; 1.19]	100.0%	
Random effects model			ĺ.		[1.00; 1.19]		100.0%
Heterogeneity: $l^2 = 0\%$, $\tau^2 = 0$, $p = 0.6$	1				Acres Cours		Contractor
and the second	-		0.1 0.51 2 10				



Figure 10. Symptomatic infection in RCTs comparing hydroxychloroquine or chloroquine with no prophylaxis among individuals exposed to COVID-19

Study	TE	seTE	Risk Ratio	RR	9	5%-CI	Weight (fixed)	Weight (random)
RoB = High/Some concerns			1					
BCN PEP CoV-2	-0.12	0.2537		0.89	[0.54;	1.461	10.0%	10.0%
COVID-19 PEP	-0.19			0.83		1.18]	19.6%	19.6%
Seet et al	-0.43	0.2149		0.65		0.991		13.9%
CHEER	0.40	0.4144		1.49	11 C	3.37]		3.7%
NA	-0.55	0.7242		0.58	10.14;			1.2%
Tirupakuzhi et al	-0.14	0.4057	-	0.87	[0.39;			3.9%
Fixed effect model			4		[0.66;			
Random effects model. Helerogenelly: $t^2 = 0\%$, $\tau^2 = 0$, $\mu = 0.60$			•		[0.66;			52,4%
RoB = Low								
COVID-19 PREP	-0.30	0.1996		0.74	[0.50;	1.10]	16.1%	16.1%
PrEP COVID	-1.21	1.6284 -	· · · ·	0.30	[0.01;	7.25]	0.2%	0.2%
PATCH	0.65	0.8473		1.91	[0.36;	10.03]	0.9%	0.9%
COVID-19 PEP (University of Washington)	0.22	0.2185	-	1.24	[0.81;	1.90]	13.5%	13.5%
HERO-HCQ	-0.27	0.2008		0.77	[0.52;	1.13]	15.9%	15.9%
WHIP COVID-19	0.01	1.2217		1.01	[0.09;	11.02]	0.4%	0.4%
PHYDRA	-1.74	1.0654	+ +	0.17	[0.02;	1.41]	0.6%	0.6%
Fixed effect model			4	0.67	[0.69;	1.09]	47.6%	
Random effects model			4	0,87	[0.65;	1.15]	~	47.6%
Helerogeneity: $l^2 = 17\% \pi^2 = 0.0241$, $p = 0.30$								
Fixed effect model			4	0.84	[0.72;	0.98]	100.0%	
Random effects model Heterogeneity: $l^2 = 0\%$, $\tau^2 = 0$, $p = 0.53$				0.84	[0.72;	0.98]	in a c	100.0%
Residual heterogeneity: $I^2 = 0\%$, $p = 0.45$			0.1 0.51 2 10					

In addition, we identified a systematic review¹² that included 12 unpublished studies providing information on mortality outcome. Overall pooled estimates did not differ when including unpublished information (OR 1.08, 95%CI 0.99 to 1.18).

Lopinavir-ritonavir

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See Summary of findings Table 4, Appendix 1

We identified 21 RCTs including 10,697 patients in which lopinavir-ritonavir was compared against standard of care or other treatments. The RECOVERY trial was the biggest with 1,616 patients assigned to dexamethasone and 3,424 to standard of care. Three studies provided information on mortality outcome, all of which included patients with severe disease, as shown by the mortality risk in control arms, which ranged from 10.6% to 25%. Our results showed:

- Lopinavir-ritonavir probably does not reduce mortality, RR 1.01 (95%CI 0.92 to 1.11); RD 0.2% (95%CI -1.3% to 1.8%); Moderate certainty ⊕⊕⊕○ (Figure 11)
- Lopinavir-ritonavir does not reduce invasive mechanical ventilation requirement; RR 1.07 (95%CI 0.98 to 1.17); RD 1.2% (95%CI -0.3% to 2.9%); High certainty ⊕⊕⊕⊕





- Lopinavir-ritonavir probably does not improve symptom resolution or improvement; RR 1.03 (95%CI 0.92 to 1.15); RD 1.8% (95%CI -4.8% to 9%); Moderate certainty ⊕⊕⊕○
- Lopinavir-ritonavir may not increase the risk of severe adverse events, RR 0.6 (95%CI 0.37 to 0.98); RD -4.1% (95%CI -6.5% to -0.2%); Low certainty ⊕⊕○○
- It is uncertain if lopinavir-ritonavir increases or decreases symptomatic infections in exposed individuals, RR 1.40 (95%CI 0.78 to 2.54); RD 1.8% (95%CI -3.8% to -26.8%); Very low certainty ⊕○○○
- It is uncertain if lopinavir-ritonavir increases or decreases hospitalizations, RR 1.22 (95%CI 0.61 to 2.47); RD 1.1% (95%CI -1.9% to -7.1%); Very low certainty ⊕○○○

Figure 11. All-cause mortality in RCTs comparing lopinavir–ritonavir with standard of care for treatment of patients with COVID-19

Study	TE	seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
LOTUS China	-0.26	0.2693		0.77	[0.45; 1.30]	3.2%	3.2%
RECOVERY - Lopinavir-ritonavir	0.03	0.0554		1.03	[0.93; 1.15]	76.1%	76.1%
WHO SOLIDARITY - Lopinavir-Ritonavir	-0.04	0.1082			[0.78; 1.19]		19.9%
NA	-0.18	0.5323 -		0.83	[0.29; 2.37]	0.8%	0.8%
Fixed effect model			\$	1.01	[0.92; 1.11]	100.0%	-
Random effects model Heterogeneity: $l^2 = 0\%$, $\tau^2 = 0$, $p = 0.67$			· · · · · ·	1.01	[0.92; 1.11]	- H - #	100.0%
			0.5 1 2				

Convalescent plasma See summary of findings Table 5 in appendix 1

We identified 58 RCTs including 24,753 patients in which convalescent plasma was compared against standard of care or other treatments. RECOVERY was the largest study including 11,588 patients. Most studies (52/58) included severely ill patients, as shown by the mortality rate in the control arms, ranging from 5.5% to 53%. The remaining studies included patients with recent onset symptoms and reported a control-arm mortality rate of 0.4% to 6.6%, or non-infected exposed individuals. Convalescent plasma was administered in one to three infusions to symptomatic patients in all cases. Our results showed:

- Convalescent plasma does not reduce mortality, RR 0.98 (95%CI 0.93 to 1.03); RD -0.3% (95%CI -1.1% to 0.5%); High certainty ⊕⊕⊕⊕ (Figure 12)
- Convalescent plasma does not significantly reduce invasive mechanical ventilation requirements, RR 1.03 (95% CI 0.94 to 1.11); RD 0.5% (95%CI -1% to 1.9%); High certainty ⊕⊕⊕⊕
- Convalescent plasma probably does not improve symptom resolution or improvement, RR 0.99 (95% CI 0.95 to 1.02); RD -0.6% (95% CI -3% to 1.2); High certainty ⊕⊕⊕⊕



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- It is uncertain if convalescent plasma reduces symptomatic infections in exposed individuals, RR 0.92 (95% CI 0.32 to 2.62); RD -1.4% (95%CI -11.8% to 28.2); Very low certainty ⊕○○○
- Convalescent plasma may not increase severe adverse events, RR 1.05 (95% CI 0.90 to 1.22); RD 0.5% (95%CI -1% to 2.2%); Low certainty ⊕⊕⊖○
- Convalescent plasma probably has no important effect on hospitalizations, RR 0.77 (95% CI 0.57 to 1.03); RD -1.1% (95%CI -2.1% to 0.1%); Moderate certainty ⊕⊕⊕○ (Figure 13). The observed effect would probably be considered important in patients with very high hospitalization risk.

Figure 12. All-cause mortality in RCTs comparing convalescent plasma with standard of care for treatment of patients with COVID-19



Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
RoB2 = High/Moderate						
Li L et al	-0.42 0.4117		0.65	[0.29; 1.47]	0.4%	0.9%
CONCOVID	-0.61 0.4594		0.55	[0,22; 1.34]	0.3%	0.7%
ConPlas-19	-2.07 1.4740			[0.01; 2.26]	0.0%	0.1%
PLACID	0.07 0.2303	+		[0.68; 1.68]	1.3%	2.6%
ILBS-COVID-02	1.17 1.0933	_		[0.38; 27.40]	0.1%	0.1%
AlQahtani M et al	-0.69 1.1832			[0.05; 5.08]		0.1%
PICP19	-0.34 0.3485			[0.36; 1.41]		1.2%
Baklaushev VP et al	-0.83 0.9635			[0.07; 2.87]	0.1%	0.2%
AAAS9924	-0.67 0.2963			[0.29: 0.92]		1.6%
CAPSID	-0.45 0.3341			[0.33; 1.22]	0.6%	1.3%
PLACOVID	0.33 0.3278			[0.73; 2.63]	0.6%	1.3%
DAWn-Plasma	Carlanzi, Thursday, and					1.5%
	0.05 0.3109			[0.57: 1.94]		
PennCCP2	-1.63 0.7412			[0.05; 0.83]		0.3%
IMPACT	-0.13 0.4470			[0.37; 2.11]		0.7%
COP-COVID-19	-0.04 0.5019			[0.36; 2.57]		0.6%
CAPRI	0.12 1.3718			[0.08; 16.55]		0.1%
Fixed effect model		4		[0.667 0.99]	6.2%	
Random effects model	0.351	9	0.79	[0.64; 0.99]		13,2%
RoB2 = Low						
PLASM-AR	-0.04 0.3308		0.06	[0.50: 1.83]	0.6%	1.3%
FundacionINFANT-Plasma	-0.69 0.8515			[0.09; 2.65]		0.2%
RECOVERY-Plasma	0.00 0.0358	T		[0.93; 1.07]		27.0%
Pouladzadeh M et al	-0.51 0.6831			[0.16; 2.29]		0.3%
SBU-COVID19-ConvalescentPlasma				[0.36; 1.86]		0.8%
REMAP-CAP	-0.03 0.0578	7		[0.87; 1.09]		19.7%
CONCOR-1	0.12 0.1266	+		[0.88; 1.45]		7.4%
COVIDIT	0.19 0.4422			[0.51; 2.89]	0.3%	0.7%
C3PO	1.60 1.0919		4.94	[0.58; 42.00]	0.1%	0.1%
TSUNAMI	-0.27 0.3399		0.77	[0.39; 1.49]	0.6%	1.2%
COnV-ert & CoV-Early	-0.69 1.2227		0.50	[0.05; 5.52]	0.0%	0.1%
CSSC-004	-1.95 1.5107		0.14	[0.01; 2.75]	0.0%	0.1%
COP20	-0.60 0.8385			[0.11; 2.84]		0.2%
CONTAIN COVID-19	-0.02 0.1967	+		[0.67; 1.44]	1.7%	3.5%
De Santis GC et al	-0.14 0.2984		0.87	[0.48; 1.56]	0.8%	1.6%
PROTECT-Patient trial	-0.19 0.3592			[0.41: 1.68]		1.1%
LIFESAVER	0.69 1.2748			[0.16; 24.33]	0.0%	0.1%
RECOVER	0.09 0.5374			[0.38; 3.13]		0.5%
LACCPT	0.15 0.3574	_		[0.58; 2.35]	0.5%	1.1%
CPC-SARS	-1.76 0.4904			[0.07; 0.45]	0.3%	0.6%
Herrick J et al	-1.39 1.5411 -			[0.01; 5.13]	0.0%	0.1%
Tatem G et al	-0.29 0.8266			[0.15; 3.79]	0.1%	0.2%
Chowdhury FR et al	-0.51 0.7638			[0.13; 2.68]	0.1%	0.2%
PLACO-COVID	0.54 0.4392			[0.72; 4.05]	0.4%	0.8%
ASCOT	-0.51 1.1738			[0.06; 5.99]		0.1%
PERUCONPLASMA	-1.02 1.0831			[0.04; 3.02]	0.1%	0.1%
CP-COVID-19	1.14 0.7916	1		[0.66; 14.73]	0.1%	0.2%
CONFIDENT	-0.12 0.1689	1		[0.64; 1.24]	2.4%	4.5%
PC/COVID-19	-0.46 0.8827			[0.11; 3.56]	0.1%	0.2%
CCAP-2	0.57 0.5336			[0.62; 5.01]		0.5%
COOPCOVID	0.15 0.2432	+		[0.72; 1.87]	1.1%	2.3%
GOPLA-II	0.13 0.2021	+		[0.76; 1.69]	1.7%	3.3%
Rojas et al	1.08 0.7891		2.93	[0.62; 13.78]	0.1%	0.2%
Self	0.07 0.1397	t	1.07	[0.82; 1.41]	3.5%	6.3%
Fixed effect model			1.00	[0.94; 1.05]	93.8%	
Random effects model Helensenney $t^2 = 0$, $\sigma = 0.0 = 0.48$				(0.94; 1.05)		86.8%
				a lashish	a h	
Fixed effect model				[0.93; 1.03]		400 00
Random effects model Heterogeneity: $l^2 = 8\%$, $\tau^2 = 0.0042$, $p = 1$	· · ·	1	0.97	[0.90; 1.04]		100.0%





Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
C3PO	-0.11 0.1722		0.90	[0.64; 1.26]	49.6%	43.6%
COnV-ert & CoV-Early	-0.14 0.2269	*	0.87	[0.56; 1.36]	28.5%	30.7%
CSSC-004	-0.65 0.2631		0.52	[0.31; 0.87]	21.2%	24.8%
CSSC-001	-1.54 1.5415 —		0.21	[0.01; 4.41]	0.6%	0.9%
Fixed effect model		4	0.79	[0.62; 1.00]	100.0%	
Random effects mode		A	0.77	[0.57; 1.03]	-	100.0%
Heterogeneity: $I^2 = 24\%$,	$r^2 = 0.0223, p = 0.27$	1 1 1 1 - 1				
		0.1 0.51 2 10				

Figure 13. Hospitalizations comparing convalescent plasma with standard of care for treatment of patients with COVID-19

In one of the studies, 58 patients were randomized to early administration of convalescent plasma (at the time they were randomized) or late administration (only if clinical deterioration was observed). All patients in the early arm received the treatment, while just 43.3% of patients received it in the late arm. Results showed no mortality reduction (OR 4.22, 95%CI 0.33 to 53.57) or reduction in the need for invasive mechanical ventilation requirement reduction (OR 2.98, 95%CI 0.41 to 21.57) with early infusion. However, the certainty of the evidence was very low $\oplus \bigcirc \bigcirc$ because of imprecision. In addition, no significant differences were observed in the subgroup of patients treated early (< 4 days since the beginning of symptoms) versus late (> 4 days since the beginning of symptoms) versus late (> 4 days since the beginning of symptoms) with convalescent plasma, in the RECOVERY trial.

Tocilizumab

See Summary of findings Table 6 in Appendix 1

We identified 29 RCTs including 9,466 patients in which tocilizumab was compared against standard of care or other interventions. Twenty studies reported on the mortality outcome, including the RECOVERY study that recruited 4,116 patients. All studies included severe patients, but some excluded critical patients. The proportion of critical patients in those studies that included them was 16.5% to 47.5%. Our results showed:

- Tocilizumab reduces mortality, RR 0.86 (95%CI 0.79 to 93); RD -2.2% (95%CI -3.4% to -1.1%); High certainty ⊕⊕⊕⊕ (Figure 14)
- Tocilizumab reduces invasive mechanical ventilation requirements, RR 0.84 (95%CI 0.79 to 0.91); RD -2.8% (95%CI -3.6% to -1.6%); High certainty ⊕⊕⊕⊕ (Figure 15)
- Tocilizumab may improve time to symptom resolution, RR 1.08 (95%CI 1.02 to 1.14); RD 4.8% (95%CI 1.2% to 8.5%); Low certainty ⊕⊕⊖○
- Tocilizumab probably does not significantly increase severe adverse events at 28-30 days, RR 0.95 (95%CI 0.87 to 1.04); RD -0.5% (95%CI -1.3% to 0.4%); Moderate certainty ⊕⊕⊕○





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Figure 14. All-cause mortality in RCTs comparing tocilizumab with standard of care for treatment of patients with COVID-19

Study	TE	seTE		R	isk Rati	0		RR	9	5%-CI	Weight (fixed)	Weight (random)
COVACTA	-0.02	0.1770			+			0.98	[0.69;	1.39]	5.6%	5.6%
RCT-TCZ-COVID-19	0.79	1.2117		-				2.20	[0.20; 1	23.65]	0.1%	0.1%
BACC Bay Tocilizumab Trial	0.41	0.6526			- <u>i</u> +	-		1.51	[0.42;	5.42]	0.4%	0.4%
CORIMUNO-TOCI 1	-0.07	0.4869			-			0.93	[0.36;	2.42]	0.7%	0.7%
EMPACTA	0.19	0.3428			- +			1.22	[0.62;	2.38]	1.5%	1.5%
REMAP-CAP - tocilizumab	-0.24	0.1090			-			0.78	[0.63;	0.97]	14.8%	14.8%
Veiga	0.83	0.4551			++	-		2.30	[0.94;	5.61]	0.8%	0.8%
RECOVERY-TCZ	-0.16	0.0542						0.85	[0.76;	0.95]	59.6%	59.6%
PreToVid	-0.45	0.2564			-+			0.64	[0.39;	1.06]	2.7%	2.7%
Mahmoudi et al	0.33	0.5818			- +			1.40	[0.45;	4.37]	0.5%	0.5%
Hamed DM et al	0.82	1.1908		-				2.26	[0.22; 1	23.33]	0.1%	0.1%
ARCHITECTS	-1.51	1.4863		+				0.22	[0.01;	4.05]	0.1%	0.1%
CORIMUNO-TOCI ICU	-0.21	0.3415			- <u>i</u> -			0.81	[0.41;	1.58]	1.5%	1.5%
COV-AID	0.13	0.4772			<u>+</u>			1.14	[0.45;	2.91]	0.8%	0.8%
COVIDOSE-2	-2.53	1.4916						0.08	[0.00;	1.49]	0.1%	0.1%
COVIDSTORM	0.42	1.6170					-	1.53	[0.06; 3	36.31]	0.1%	0.1%
HMO-0224-20	-0.46	0.3606						0.63	[0.31;	1.28]	1.3%	1.3%
REMDACTA	-0.07	0.1736			+			0.93	[0.66;	1.31]	5.8%	5.8%
ImmCoVA	0.20	0.9579		_	<u>.</u>	_		1.23	[0.19;	8.02]	0.2%	0.2%
COVINTOC	-0.34	0.3677						0.71	[0.34;	1.46]	1.3%	1.3%
TOCIDEX	-0.28	0.2972						0.76	[0.42;	1.35]	2.0%	2.0%
Fixed effect model Random effects model Heterogeneity: $l^2 = 0\%$, $\tau^2 = 0$,	p = 0.69	9			0				[0.79; [0.79;		100.0% 	 100.0%
, , , , , , , , , , , , , , , , , , ,			0.01	0.1	1	10	100					

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Figure 15. Mechanical ventilation requirement in RCTs comparing tocilizumab with standard of care for treatment of patients with COVID-19

Study	TE	seTE		R	isk Rati	0		RR	9	5%-CI	Weight (fixed)	Weight (random)
,									_		. ,	(,
COVACTA	+	0.1826							[0.53;			4.0%
RCT-TCZ-COVID-19	+ +	0.2930			+			1.10		1.95]		1.5%
BACC Bay Tocilizumab Trial									[0.29;	-		0.7%
CORIMUNO-TOCI 1		0.4905		_	•				[0.15;			0.5%
EMPACTA	-0.44	0.3173							[0.35;			1.3%
REMAP-CAP - tocilizumab		0.1128			1				[0.65;	-		10.4%
Veiga	+ +	0.2990			1				[0.44;		1.5%	1.5%
RECOVERY-TCZ	-0.17	0.0454			+			0.84	[0.77;	0.92]	64.1%	64.1%
PreToVid		0.2851							[0.39;			1.6%
Hamed DM et al	1.22	0.7647							[0.76;	-		0.2%
CORIMUNO-TOCI ICU	-0.08	0.4160							[0.41;			0.8%
COV-AID	++	0.3306			∦ ⊷				[0.68;			1.2%
COVIDOSE-2	-2.47	1.4908							[0.00;			0.1%
COVIDSTORM	-0.20	0.6929		-					[0.21;			0.3%
COVITOZ-01	0.46	1.5801					-	1.59	[0.07;	35.15]	0.1%	0.1%
HMO-0224-20	0.08	0.4067			- 				[0.49;			0.8%
REMDACTA	-0.02	0.1320			H			0.98	[0.76;	1.28]	7.6%	7.6%
ImmCoVA	-0.49	0.6461		_				0.61	[0.17;	2.18]	0.3%	0.3%
TOCOVID	-1.11	1.1483			<u>⊷</u> ∦			0.33	[0.03;	3.12]	0.1%	0.1%
COVINTOC	-0.22	0.4225						0.80	[0.35;	1.83]	0.7%	0.7%
TOCIDEX	-0.16	0.2437			+			0.85	[0.53;	1.37]	2.2%	2.2%
Fixed effect model					Ŷ						100.0%	
Random effects model					0			0.84	[0.79;	0.91]		100.0%
Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0$,	p = 0.7	4	1	I	I	I	1					
			0.01	0.1	1	10	100					

A subgroup analysis, performed in the RECOVERY trial, comparing the effect of tocilizumab in severe and critical patients, did not suggest a subgroup modification effect according to baseline disease severity (p=0.52).

In addition, one study that compared standard dose (4 mg/kg) versus high dose (8 mg/kg) found no significant differences and one study that compared baricitinib versus tocilizumab reported no significant differences in mortality or mechanical ventilation. However, the certainty of the evidence was low because of imprecision.

Anticoagulants

See Summary of findings Table 7, Appendix 1

Thromboembolic complications in patients infected with COVID-19 are relatively frequent.¹³ As for hospitalized patients with severe medical conditions, current guidelines recommend thromboprophylaxis measures should be used for inpatients with COVID-19 infection.¹⁴ Regarding the best thromboprophylactic scheme, we identified 20 RCTs including 8,131 patients that compared anticoagulants in intermediate (i.e., enoxaparin 1 mg/kg a day) or full dose (i.e.,



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enoxaparin 1 mg/kg twice a day) versus prophylactic dose (i.e., enoxaparin 40 mg a day), or anticoagulants versus standard of care in patients with mild ambulatory disease. In addition we identified one study that compared rivaroxaban and enoxaparin in hospitalized patients. All studies included hospitalized patients with COVID-19. Our results showed:

- In moderate to critical patients, anticoagulants in intermediate dose or full dose may not reduce mortality in comparison with prophylactic dose, RR 0.99 (95%CI 0.83 to 1.19); RD -0.2% (95%CI -2.7% to 3%); Moderate certainty ⊕⊕⊕○ (excluding high risk of bias studies) (Figure 16)
- In moderate to critical patients, anticoagulants in intermediate dose may reduce venous thromboembolic events in comparison with prophylactic dose, RR 0.82 (95%CI 0.43 to 1.59); RD -1.3% (95%CI -4% to 4.1%); Low certainty ⊕⊕○○
- In moderate to critical patients, anticoagulants in full dose reduce venous thromboembolic events in comparison with prophylactic dose, RR 0.56 (95%CI 0.44 to 0.71); RD -3.1% (95%CI -3.9% to -2%); High certainty ⊕⊕⊕⊕
- In moderate to critical patients, anticoagulants in intermediate dose or full dose probably increase major bleeding in comparison with prophylactic dose, RR 1.56 (95%CI 1.08 to 2.25); RD 1.1% (95%CI 0.2% to 2.4%); Moderate certainty ⊕⊕⊕○
- In mild ambulatory patients, anticoagulants in prophylactic dose may not improve time to symptom resolution, RR 1.08 (95%CI 0.92 to 1.27); RD 4.8% (95%CI -4.8% to 16.4%); Low certainty ⊕⊕○○
- In mild ambulatory patients, anticoagulants in prophylactic dose may not reduce hospitalizations, RR 0.94 (95%CI 0.55 to 1.59); RD -0.3% (95%CI -2.2% to 2.8%); Low certainty ⊕⊕○○
- In mild ambulatory patients it is uncertain if anticoagulants in prophylactic dose increase or decrease clinically important bleeding and hospitalization; Very low certainty ⊕○○○



Figure 16. All-cause mortality in RCTs using anticoagulants in therapeutic dose, intermediate dose or prophylactic dose for treatment of hospitalized patients with COVID-19

						Weight	Weight
Study	TE	seTE	Risk Ratio	RR	95%-CI	(fixed)	(random)
RoB = Some concern	19		1				
HESACOVID	-1.10	1.0646		0.33	[0.04; 2.69]	0.2%	0.7%
INSPIRATION	0.05	0.0991	60	1.05	[0.87; 1.28]	27.5%	21.2%
Zarychanski-Critical	0.05	0.0799	60	1.05	[0.90; 1.23]	42.3%	23.3%
Zarychanski-Non-critic	al -0.11	0.1465	+	0.89	[0.67; 1.19]	12.6%	16.1%
ACTION	0.40	0.2560	+	1.49	[0.90; 2.46]	4.1%	8.4%
RAPID	-1.47	0.5449		0.23	[0.08; 0.67]	0.9%	2.4%
HEP-COVID	-0.25	0.2376		0.78	[0.49; 1.23]	4.8%	9.4%
X-Covid 19	1.62	1.0854		5.05	[0.60; 42.43]	0.2%	0.6%
PROTHROMCOVID		0.9023			[0.26; 9.05]		0.9%
PROTHROMCOVID		0.9016			[0.30; 10.23]		0.9%
COVID-HEP	0.01	0.8009			[0.21; 4.87]	0.4%	1.2%
Fixed effect model	1.1		\$		[0.92: 1.13]	93.7%	
Random effects mod	6)		\$		(0.83; 1.19)	1000	85.2%
Helerogeneity: $t^{1} = 97\%$	ч ^а = 10.05	44 g = 0 1	D				
RoB = High							
Perepu U et al	-0.34	0.3307	-++	0.71	[0.37; 1.37]	2.5%	5.7%
BEMICOP	0.66	1.1994		- 1.94	[0.18; 20.35]	0.2%	0.5%
Oliynyk O et al	-0.50	0.3075		0.61	[0.33; 1.11]	2.9%	6.4%
TACOVID	0.69	0.5916		2.00	[0.63; 6.38]	0.8%	2.1%
Fixed effect model			\diamond	0.78	(0.52; 1.16)	6.3%	
Random effects mod	01		4	0.82	(0.50; 1.34)		14.8%
Heterogeneity: $l^2 = 22\%$	7 0,05	62 p=0.2	B				
Fixed effect model				1.00	[0.90; 1.11]	100.0%	
Random effects mod	el		\$	0.96	[0.81; 1.14]		100.0%
Heterogeneity: 12 = 34%,	$\tau^2 = 0.02$	72, p = 0.0	9 [[] []]		4 12 Y Y Y		
Residual heterogeneity:			0.1 0.5 1 2 10				

NSAIDs

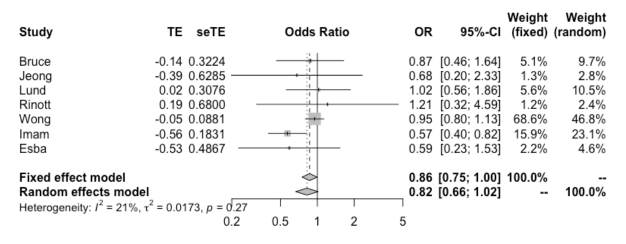
See Summary of findings Table 8, Appendix 1

We identified seven non-RCTs including at least 100 patients in which COVID-19 mortality risk was compared between groups of patients exposed to NSAIDs and those that were not. Populations varied between studies. For example, Wong et al. included individuals exposed to COVID-19 (living in a region affected by the pandemic) while other studies included only patients with confirmed COVID-19 infection. Our results showed:

 No association between NSAID exposure and mortality, OR 0.82 (95%CI 0.66 to 1.02); Very low certainty ⊕○○○ (Figure 17)



Figure 17. All-cause mortality in non-RCTs comparing exposure to NSAIDs with no exposure in individuals exposed to or infected with COVID-19



Interferon Beta-1a

See Summary of findings Table 9, Appendix 1

We identified seven RCTs including 7,017 patients in which interferon beta-1a was compared against standard of care or other treatments and informed on mortality outcome. The WHO SOLIDARITY trial was the biggest, with 2,144 patients assigned to intervention and 2,147 to control. The studies included severe patients, as shown by the fact that mortality in the control arms ranged from 10.5% to 45%. Our results showed:

- Interferon beta-1a (subcutaneous) probably does not reduce mortality, RR 0.99 (95%CI 0.75 to 1.31); RD -0.2% (95%CI -4% to 5%); Moderate certainty ⊕⊕⊕○ (Figure 18)
- Interferon beta-1a (subcutaneous) probably does not reduce invasive mechanical ventilation requirements, RR 1.01 (95%CI 0.87 to 1.18); RD 0.2% (95%CI -2.2% to 3.1%); Moderate certainty ⊕⊕⊕○
- Interferon beta-1a (subcutaneous) probably does not increase symptom resolution or improvement; RR 0.96 (95%CI 0.92 to 0.99); RD -2.6% (95%CI -4.8% to -3.2%); Moderate certainty ⊕⊕⊕○
- Interferon beta-1a probably does not increase severe adverse events, RR 1.03 (95%CI 0.85 to 1.24); RD 0.3% (95%CI -1.5% to 2.4%); Moderate certainty ⊕⊕⊕○
- Interferon beta-1a (inhaled) may improve time to symptom resolution, HR 2.19 (95%CI 1.03 to 4.69); RD 26.4% (95%CI 1.1% to 38.1%); Low certainty ⊕⊕○○



Figure 18. All-cause mortality with IFN beta-1a vs. standard of care in randomized studies including COVID-19 patients

Study	TE	seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
Davoudi-Monfared et al	-0.83	0.3666		0.44	[0.21: 0.90]	3.3%	11.5%
WHO SOLIDARITY - Interferon	0.17	0.0774	100	1.19	[1.02; 1.38]	75.1%	40.4%
COVIFERON	-0.41	0.5627		0.67	[0.22; 2.01]	1.4%	5.7%
ACTT-3	0.26	0.3256		1.30	[0.69; 2.46]	4.2%	13.7%
INTEREST	0.03	0.1691	+	1.03	[0.74; 1.44]	15.7%	28.0%
Castro-Rodriguez et al	-1.17	1.6319 —	· · · ·	0.31	[0.01; 7.60]	0.2%	0.8%
Fixed effect model			-	1.12	[0.98; 1.27]	100.0%	-
Random effects model Heterogeneity: $l^2 = 45\%$, $\tau^2 = 0.04$	52, p =	0.11	0.1 0.51 2 10	0.99	[0.75; 1.31]	-	100.0%

Bamlanivimab +/- etesevimab (monoclonal antibody)

See Summary of findings Table 10, Appendix 1

We identified nine RCTs including 5,939 patients in which bamlanivimab was compared against standard of care or other treatments. Eight studies included patients with mild to moderate COVID-19 and one included exposed individuals and assessed bamlanivimab as a prophylactic intervention. Our results showed:

- It is uncertain if bamlanivimab reduces mortality or mechanical ventilation requirements; RR 0.68 (95%CI 0.17 to 2.8); RD -5.1% (95%CI -13.2% to 2.8%); Very low certainty ⊕○○○
- Bamlanivimab probably does not significantly improve time to symptom resolution, RR 1.02 (95%CI 0.99 to 1.06); RD 1.2% (95%CI 3.6% to 5.4%); Moderate certainty ⊕⊕⊕○
- Bamlanivimab probably decreases symptomatic infection in exposed individuals, RR 0.56 (95%CI 0.39 to 0.81); RD -7.6% (95%CI -10.6% to -3.6%); Moderate certainty ⊕⊕⊕○
- Bamlanivimab may not increase severe adverse events; RR 1.12 (95%CI 0.75 to 1.66); RD 1.2% (95%CI -2.5% to -6.7%); Low certainty ⊕⊕○○
- Bamlanivimab probably reduces hospitalizations in patients with non-severe disease; RR 0.37 (95%CI 0.21 to 0.65); RD -3% (95%CI -3.8% to -1.7%); Moderate certainty ⊕⊕⊕○ (Figure 19)

Figure 19. Hospitalizations with bamanivimab vs. standard of care in randomized studies including COVID-19 patients





Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
BLAZE-1 BLAZE-1 ACTIV-2	-1.36 0.5485 -1.19 0.3389 -0.29 0.5283		0.30	[0.09; 0.75] [0.16; 0.59] [0.26; 2.10]	21.3% 55.8% 22.9%	24.1% 50.3% 25.6%
Fixed effect model Random effects mode Heterogeneity: $I^2 = 20\%$,		0.5 1 2		[0.22; 0.59] [0.21; 0.65]	100.0% 	 100.0%

In addition, one study that compared bamlanivimab +/- etesevimab against REGEN-COV (casirivimab and imdevimab) in non-severe patients with risk factors for severity reported no important differences in hospitalizations.

Favipiravir

See Summary of findings Table 11, Appendix 1

We identified 29 RCTs including 4,624 patients in which favipiravir was compared against standard of care or other treatments. Seventeen studies reported on favipiravir with or without HCQ versus standard of care, two studies reported on favipiravir vs HCQ or CQ, two study reported on favipiravir vs lopinavir ritonavir and the remaining studies compared favipiravir against other active interventions. As there is moderate to high certainty that HCQ and lopinavir-ritonavir are not related to significant benefits, we assumed those interventions as equivalent to standard of care. Our results showed:

- Favipiravir may increase mortality; RR 1.08 (95%CI 0.77 to 1.52); RD 1.3% (95%CI 3.7% to 8.3%); Low certainty ⊕⊕○○
- Favipiravir may increase mechanical ventilation requirements; RR 1.27 (95%CI 0.91 to 1.76); RD 4.7% (95%CI -1.6% to 13.1%); Low certainty ⊕⊕⊖○
- Favipiravir probably does not increase symptom resolution or improvement, RR 1.01 (95%CI 0.97 to 1.05); RD 0.6% (95%CI -1.8% to 3%); High certainty ⊕⊕⊕⊕ (Figure 20) (based on low risk of bias studies)
- It is uncertain if favipiravir increases the risk of severe adverse events; RR 0.92 (95%CI 0.56 to 1.52); RD -0.8% (95%CI -4.5% to 5.3%); Very low certainty ⊕○○○
- Favipiravir may not reduce hospitalizations in patients with non-severe disease; RR 1.33 (95%CI 0.64 to 1.78); RD 1.6% (95%CI -1.7% to 3.7); Low certainty ⊕⊕○○



Figure 20. Symptom resolution at 7-28 days in randomized studies comparing favipiravir with standard of care in patient with COVID-19

						Weight	Weight
Study	TE seT	E	Risk Ratio	RR	95%-CI	(fixed)	(random)
RoB = High			H				
Ivashchenko AA et al	-0.07 0.225	1		0.93	[0.60; 1.45]	0.7%	4.3%
Lou Y et al	0.11 0.434	6	<u>k</u>	1.11	[0.47; 2.60]	0.2%	1.4%
Ruzhentsova T et al (R-Pharm) 0.39 0.200	4	<u>}</u> ∔⊷	1.48	[1.00; 2.18]	0.8%	5.1%
FAV052020 (Promomed, LLC)	0.59 0.289	3		1.80	[1.02; 3.17]	0.4%	2.9%
Udwadia ZF et al	0.20 0.111	2	<u>+-</u>	1.22	[0.98; 1.52]	2.7%	9.7%
Balykova LA et al	0.59 0.289	3	<u> </u>	1.80	[1.02; 3.17]	0.4%	2.9%
FACCT	-0.07 0.096	5	-4	0.93	[0.77; 1.13]	3.6%	10.8%
Shinkai M et al	0.28 0.135	3	<u>+</u>	1.32	[1.02; 1.73]	1.8%	8.2%
FAVI-COV-US201	0.00 0.294	4	- -	1.00	[0.56; 1.78]	0.4%	2.9%
Rahman SMA et al	1.79 0.555	8		- 6.00	[2.02; 17.83]	0.1%	0.9%
Sirijatuphat	0.90 0.268	4	li ——	2.45	[1.45; 4.15]	0.5%	3.3%
Fixed effect model			\$	1.21	[1.09; 1.34]	11.5%	
Random effects model			\diamond	1.36	[1.10; 1.68]		52.6%
Heterogeneity: $I^2 = 66\%, \tau^2 = 0.06$	696, <i>p</i> < 0.01						
RoB = Low							
Solaymani-Dodaran M et al	-0.01 0.047		ŧ		[0.90; 1.09]	14.8%	14.4%
CVD-04-CD-001	0.05 0.146		- 		[0.79; 1.40]		7.5%
Holubar M et al	0.15 0.111	5	H+	1.16	[0.94; 1.45]	2.7%	9.7%
Golan	0.01 0.021	9	÷.		[0.96; 1.05]	69.5%	15.8%
Fixed effect model			2	1.01	[0.97; 1.05]	88.5%	
Random effects model			ę:	1.01	[0.97; 1.05]		47.4%
Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0$, $p =$	= 0.59						
Fixed effect model			P		[0.99; 1.07]	100.0%	
Random effects model		_	♦	1.16	[1.05; 1.30]		100.0%
Heterogeneity: $I^2 = 66\%$, $\tau^2 = 0.07$		1		1			
Residual heterogeneity: $I^2 = 59\%$, <i>p</i> < 0.01	0.1	0.5 1 2 1	0			

Ivermectin

See Summary of findings Table 12, Appendix 1

We identified 49 RCTs including 13,326 patients in which ivermectin was compared against standard of care or other treatments. Studies included patients with mild to severe disease, as shown by the mortality rates in the control arms, which ranged from 0% to 42%. Most studies did not report on clinical important outcomes and most of the ones that did have important methodological limitations including inappropriate randomization process and lack or unclear report of allocation concealment. Our results showed:

- Ivermectin probably does not reduce mortality, RR 1 (95%CI 0.8 to 1.24); RD -0% (95%CI -3.2% to 3.8%); Moderate certainty ⊕⊕⊕○ (Figure 21) (based on low risk of bias studies)
- It is uncertain if ivermectin affects mechanical ventilation, RR 0.82 (95%CI 0.58 to 1.17); RD -3.1% (95%CI -7.3% to 2.9%); Very low certainty ⊕○○○ (based on low risk of bias studies)



- Ivermectin probably does not improve symptom resolution or improvement, RR 1.04 (95%CI 0.98 to 1.1); RD 2.4% (95%CI -1.2% to 6%); Moderate certainty ⊕⊕⊕○ (Figure 22) (based on low risk of bias studies).
- It is uncertain if ivermectin affects symptomatic infection, RR 1.01 (95%CI 0.54 to 1.89); RD 0.2% (95%CI -8% to 15.5%); Very low certainty ⊕○○○ (based on low risk of bias studies)
- Ivermectin may not increase severe adverse events, RR 1.05 (95%CI 0.69 to 1.62); RD 0.5% (95%CI -3.2% to 6.3%); Low certainty ⊕⊕○○
- Ivermectin probably does not have an important effect on hospitalizations in patients with recent onset non-severe disease, RR 0.90 (95%CI 0.74 to 1.1); RD -0.5% (95%CI -1.2% to 0.5%); Moderate certainty ⊕⊕⊕○ (based on low risk of bias studies). The observed effect would probably be considered important in patients with very high hospitalization risk (>10%).





Figure 21. Mortality in randomized studies comparing ivermectin with standard of care or other treatments in patients with COVID-19

Study	Experin Events		Co Events	ontrol Total		Ris	sk Rat	io		RR	9	5%-CI	Weight (fixed)	Weight (random)
RoB2 = High/Some con	cerns						1							
Mahmud et al	0	183	3	180			+			0.14	[0.01;	2.70]	1.8%	1.1%
Hashim HA et al	2	70	6	70			+			0.33	[0.07;	1.60]	3.0%	3.3%
Elgazzar et al (mild)	0	100	4	100			+			0.11	[0.01;	2.04]	2.2%	1.1%
Elgazzar et al (severe)	2	100	20	100		- 10				0.10	[0.02;	0.42]	9.9%	3.9%
Niaee et al	4	120		60			-31				[0.06;		7.3%	5.6%
Okumus et al	6	30	-	30		-	+				[0.27;		4.5%	7.1%
Beltran-IVER	5	36		70				-		1.22	[0.43;	3.45]	2.7%	6.0%
R-2020-785-176	2	65		46						1.42	[0.13;	15.15]	0.6%	1.6%
Rezai_Severe	13		18	298		-	*				[0.35;		9.1%	9.2%
Fixed effect model		1015		954		<	>			0.42	[0.29;	0.61]	41.0%	
Random effects model						<	\geq			0.42	[0.23;	0.79]		38.9%
Heterogeneity: $I^2 = 48\%$, τ^2	= 0.3821	p = 0.	05				4.4.4							
RoB2 = Low														
Kirti et al	0	55		57		+	1				[0.01;		2.2%	1.1%
Shahbaznejad et al	1	35		34			1 +				[0.12;		0.3%	1.0%
Lopez-Medina et al	0	200		198			1				[0.01;		0.7%	1.0%
Bermejo Galan et al	12	53		115			+				[0.57;		7.8%	10.2%
Abd-Elsalam et al	3	82		82		_	1				[0.17;		2.0%	3.7%
Vallejos et al	4	250		251		-	1.	_			[0.30;		1.5%	3.6%
I-TECH	3	241	10	249			1				[0.09;		4.9%	4.5%
TOGHETER - Ivermectin		679	24	679			71				[0.49;		11.9%	10.6%
ACTIV-6	1	817	0	774							[0.12;		0.3%	0.9%
Rezai_Mild	1	268	1	281							[0.07;		0.5%	1.2%
George et al	13	73	-	39			西.				[0.39;		5.2%	8.2%
IRICT	49	104	43	102			÷.				[0.82;		21.5%	14.1%
COVID-OUT - Ivermectir	1	408	0	396			1 1				[0.12;	•	0.3%	0.9%
Fixed effect model		3265		3257			1				[0.75;		59.0%	
Random effects model							1			1.00	[0.80;	1.24]		61.1%
Heterogeneity: $I^2 = 0\%$, τ^2	= 0, p = 0.	74					4							
Fixed effect model		4280		4211			\$						100.0%	
Random effects model							\diamond			0.69	[0.50;	0.95]		100.0%
Heterogeneity: $I^2 = 39\%$, τ^2			03		1	1	1	1	1					
Residual heterogeneity: I ²	= 17%, p =	= 0.24			0.01	0.1	1	10	100					



	Experin	nental	C	ontrol				Weight	Weight
Study	Events	Total	Events	Total	Risk Ratio R	R	95%-CI	(fixed)	(random)
Ro82 = High/Same car	CENTS				1.1				
Chowdhury et al	50	60	40	56	1:	17 [0.95; 1.43]	3.2%	5.9%
Mahmud et al	141	183	113	180			1.07; 1.41]		7.3%
Elgazzar et al (mild)	99	100	74	100			1.19; 1.51]		7.8%
Elgazzar et al (severe)	94	100	50	100			1.54; 2.30]		5.8%
Chachar et al	16	25	15	25			0.69; 1.65]		2.4%
Okumus et al	22	30	16	30			0.92; 2.05]		2.8%
NA	32	36	64	70			0.85: 1.111		7.4%
Kishoria et al	8	19	6	16		12 1	0.49; 2.56]	0.5%	0.8%
Faisal et al	48	50	42	50			1.00; 1.31]		7.4%
I-TECH	122	241	131	249			0.81; 1.14]		6.5%
Fixed effect model		844		876			1.13: 1.28)	41.1%	100
Random effects model							1.05: 1.37]	11112	54.2%
Heterogenery, $\vec{r}^2 = 77^4 a_i \hat{j}$	= 0.028	10-0	1.Q1		1				
RoB2 = Low									
Kirti et al	46	55	51	57		93 [0.81; 1.08]	3.9%	7.1%
Mohan et al	74	80	39	45	1.0	07 [0.94; 1.22]	3.9%	7.5%
Lopez-Medina et al	164	200	156	198	1.	04 [0.94; 1.15]	12.1%	8.3%
Manomaipiboon A et al	15	36	11	36		36 [0.73; 2.55]	0.9%	1.4%
ACTIV-6	176	817	137	774	1.:	22 [1.00; 1.49]	10.9%	5.9%
Rezai_Mild	249	268	256	281	1.0	02 [0.97; 1.07]	19.3%	9.1%
Mirahmadizadeh et al	160	261	77	130		03 [0.87; 1.23]	8.0%	6.5%
Fixed effect model		1717		1521	Ø 11	17 I	1.01: 1.13]	58.9%	1.1.1.4
Random effects model					0	14 I	0.98; 1,10)	1.1	45,8%
Hélerogeneny (* 31 o. r	2 = 0.004	7 11 -0	18						
Fixed effect model		2561		2397	å 1. ⁻	12 [1.08; 1.17]	100.0%	
Random effects model	D				♦ 1. ♦ 1.	13 I	1.05; 1.22]	12	100.0%
Heterogeneity: $I^2 = 77\%$, τ	$^{2} = 0.017$	0, p < 0	0.01		1 1 1		a series and		
Residual heterogeneity: 12					0.5 1 2				

Figure 22. Symptom resolution or improvement in randomized studies comparing ivermectin with standard of care or other treatments in patients with COVID-19

Although pooled estimates suggest significant benefits with ivermectin for some critical outcomes, these are mainly driven by studies with important methodological limitations. Furthermore, results of the studies classified as low risk of bias significantly differ from those classified as high risk of bias which results in significant uncertainty about ivermectin effects. Further research is needed to confirm or discard those findings.

Baricitinib

See Summary of findings Table 13, Appendix 1

We identified seven RCTs including 12,363 patients in which baricitinib was compared against standard of care or other treatments. All studies included moderate to severe hospitalized patients. Critical patients were excluded. Our results showed:





- Baricitinib reduces mortality, RR 0.73 (95%CI 0.57 to 0.92); RD -4.3% (95%CI -6.9% to -1.3%); High certainty ⊕⊕⊕⊕ (Figure 23)
- Baricitinib probably reduces mechanical ventilation, RR 0.83 (95%CI 0.66 to 1.04); RD 2.9% (95%CI -5.9% to 0.7%); Moderate certainty ⊕⊕⊕○
- Baricitinib probably improves time to symptom resolution, RR 1.27 (95%CI 1.13 to 1.42); RD 16.4% (95%CI 7.9% to 25.5%); Moderate certainty ⊕⊕⊕○
- Baricitinib probably does not increase severe adverse events, RR 0.78 (95%CI 0.64 to 0.95); RD -2.2% (95%CI -3.7% to -0.5%); Moderate certainty ⊕⊕⊕○

Figure 23. Mortality in randomized studies comparing baricitinib with standard of care in patients with COVID-19

Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
ACTT-2	-0.43 0.2546		0.65	[0.40; 1.07]	4.0%	14.8%
COV-BARRIER	-0.48 0.1533		0.62	[0.46; 0.83]	11.0%	25.2%
COV-BARRIER-IMV	-0.39 0.2118		0.68	[0.45; 1.02]	5.8%	18.5%
RECOVERY	-0.10 0.0574		0.91	[0.81; 1.02]	78.7%	38.5%
PanCOVID19	-0.87 0.6799 —	+ 1 1 1	0.42	[0.11; 1.59]	0.6%	3.0%
Fixed effect model		-	0.84	[0.76; 0.93]	100.0%	-
Random effects mode Heterogeneity: $l^2 = 55\%$,			0.73	[0.57; 0.92]		100.0%
	0.	.2 0.5 1 2 5	5			

In addition one study that compared baricitinib versus tocilizumab reported no significant differences in mortality or mechanical ventilation. However, the certainty of the evidence was low because of imprecision.

Azithromycin

PAHO®

See Summary of findings Table 14, Appendix 1

We identified 11 RCTs including 10,612 patients in which azithromycin was compared against standard of care or other treatments. RECOVERY trial was the biggest study including 7,762 patients with severe disease (mortality in the control arm 19%). Our results showed:

- Azithromycin probably does not reduce mortality, RR 1.01 (95%CI 0.92 to 1.1); RD 0.2% (95%CI -1.3% to 1.6%); Moderate certainty ⊕⊕⊕○ (Figure 24)
- Azithromycin probably does not reduce mechanical ventilation requirements, RR 0.92 (95%CI 0.77 to 1.1); RD -1.4% (95%CI -4% to 1.7%); Moderate certainty ⊕⊕⊕○
- Azithromycin does not improve time to symptom resolution, RR 1.02 (95%CI 0.99 to 1.04); RD 1.2% (95%CI -0.6% to 2.4%); High certainty ⊕⊕⊕⊕
- It is uncertain if azithromycin increases severe adverse events, RR 1.23 (95%CI 0.51 to 2.96); RD 2.4% (95%CI -5% to 19.9%); Very low certainty ⊕○○○





Azithromycin may not reduce hospitalizations, RR 0.98 (95%CI 0.52 to 1.86); RD -0.1% (95%CI -2.3% to 4.1%); Low certainty ⊕⊕○○



Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
Sekhavati E et al	-1.12 1.6219 -		0.33	Percise states		0.1%
COALITION II	0.05 0.1211	Ť	1.05	Address The 14		
RECOVERY	-0.00 0.0494	100	1.00	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1 A 1 A 1 A 1 A 1 A 1 A 1 A 1 A 1 A 1 A	84.5%
ATOMIC2	0.01 1.4094		1.01	[0.06; 16.05]	0.1%	0.1%
Ghanei M et al	0.00 0.5614		1.00	[0.33; 3.01]	0.7%	0.7%
DAWn-AZITHRO	0.19 0.5806		1.21	[0.39; 3.78]	0.6%	0.6%
Fixed effect model		•	1.01	[0.92; 1,10]	100.0%	
Random effects mo Heterogeneity: $I^2 = 0\%$	P. 9.1	r 1 1 1	1.01	[0.92; 1.10]		100.0%
2005	and the second	0.1 0.51 2 10				

Figure 24. Mortality in randomized studies comparing azithromycin with standard of care in patients with COVID-19

ACEI/ARB initiation or continuation

We identified 12 RCTs including 1,812 patients in which patients with COVID-19 were randomized to initiate or continue ACEI/ARB treatment and compared to standard of care or discontinue ACEI/ARB. Our results showed:

- ACEI/ARB initiation or continuation may increase mortality, RR 1.13 (95%CI 0.77 to 1.64); RD 2.1% (95%CI -3.7% to 10.2%); Low certainty ⊕⊕○○ (Figure 25) (based on low risk of bias studies)
- ACEI/ARB discontinuation may reduce mechanical ventilation requirements, RR 0.89 (95%CI 0.66 to 1.22); RD -1.9% (95%CI -5.9% to 3.8%); Low certainty ⊕⊕○○



Figure 25. Mortality in randomized studies comparing initiation or continuation vs standard of care o discontinuation of ACEI/ARB in patients with COVID-19

Study	TE	seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
RoB = High Duarte M et al Nouri-Vaskeh M et al COVID-ARB Fixed effect model Random effects model Heterogeneity: $l^2 = 0\%$, τ^2	-0.97 -0.06	0.6062 0.8062 1.3678		0.38 0.94 0.28	[0.06; 0.61] [0.08; 1.85] [0.06; 13.68] [0.11; 0.68] [0.11; 0.68]		9.3% 5.7% 2.2% 17.2%
RoB = Low REPLACE COVID BRACE CORONA ATTRACT ACEI-COVID Najmeddin F et al ALPS-COVID COVID MED RAAS-COVID-19 Fixed effect model Random effects model	-0.03 -1.02 0.44 0.26 0.03 0.80 -0.87	0.4057 0.4649 1.1382 0.4344 0.6163 0.4029 0.9690 1.1884		0.97 0.36 1.56 1.29 1.03 2.22 0.42 1.13	[0.51; 2.50] [0.39; 2.42] [0.04; 3.35] [0.67; 3.66] [0.39; 4.33] [0.47; 2.27] [0.33; 14.84] [0.04; 4.31] [0.77; 1.64] [0.77; 1.64]	8.2% 19.2% 3.3%	17.0% 14.1% 3.0% 15.5% 9.1% 17.2% 4.1% 2.8%
Heterogeneity: $l^2 = 0\%$, τ^2 Fixed effect model Random effects model Heterogeneity: $l^2 = 19\%$, τ Residual heterogeneity: l^2	² = 0.08	26, p = 0	.27 0.1 0.5 1 2 10		[0.65; 1.29] [0.59; 1.32]	100.0% 	 100.0%

Colchicine

See Summary of findings Table 15, Appendix 1

We identified 15 RCTs including 18,605 patients in which colchicine was compared against standard of care or other treatments. The COLCORONA trial was the biggest including mild ambulatory patients, with 2,235 patients assigned to intervention and 2,253 to control, and the RECOVERY trial was the biggest including moderate to critical hospitalized patients, with 5,610 patients assigned to intervention and 5,730 assigned to control. Our results showed:

- Colchicine probably does not reduce mortality, RR 0.99 (95%CI 0.92 to 1.05); RD -0.2% (95%CI -1.3% to 0.8%); Moderate certainty ⊕⊕⊕○ (Figure 26)
- Colchicine probably does not reduce mechanical ventilation requirements, RR 0.98 (95%CI 0.89 to 1.07); RD -0.3% (95%CI -1.9% to 1.2%); Moderate certainty ⊕⊕⊕○ (Figure 27)
- Colchicine does not increase symptom resolution or improvement, RR 1 (95%CI 0.98 to 1.02); RD 0% (95%CI -1.2% to 1.2%); High certainty ⊕⊕⊕⊕



85



- Colchicine does not significantly increase severe adverse events, RR 0.78 (95%CI 0.61 to 0.99); RD -2.2% (95%CI -4% to -0.1%); High certainty ⊕⊕⊕⊕
- Colchicine may not significantly increase pulmonary embolism, RR 5.55 (95%CI 1.23 to 25); RD 0.4% (95%CI 0.02% to 2.2%); Low certainty ⊕○○○
- Colchicine probably has no important effect on hospitalizations in patients with recent onset disease, RR 0.81 (95%CI 0.63 to 1.04); RD -0.9% (95%CI -1.8% to 0.2%); Moderate certainty ⊕⊕⊕○

Figure 26. Mortality in randomized studies comparing colchicine vs standard of care in patients with COVID-19

Study	TE	seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)	
Severity = Moderale to (critical							
GRECCO-19		1.1008		0.28	[0.03; 2.38]	0.1%	0.1%	
Lopes et al	-1.61	1.5312			[0.01; 4.02]		0.1%	
RECOVERY - Colchicine	0.01	0.0366	10.		[0.94; 1.08]	87.6%	87.6%	
COL-COVID	-1.63	1.5366		0.20	[0.01; 3.99]	0.0%	0.0%	
COLCOVID	-0.08	0.1075	+		[0.75; 1.14]		10.2%	
Alsultan M et al	-0.44	0.5976			[0.20; 2.07]		0.3%	
Gorial FI et al	-1.10	1.1438		0.33	[0.04; 3.14]	0.1%	0.1%	
Mostafaie A et al	-1.79	1.0646 -		0.17	[0.02; 1.34]	0.1%	0.1%	
STRUCK	-1.48	1.5053			[0.01; 4.37]		0.1%	
Cecconi et al	-0.35	0.4755			[0.28; 1.79]			
Rabbani	0.22	0.4986		1.25	[0.47; 3.32]	0.5%	0.5%	
Fixed effect model			•		[0.92; 1.06]		-	
Random effects model			\$		[0.91: 1.06]	1.1.1	99.6%	
Haterogeneity, $t^2 = 1\%, \tau^2$	0:0006	$\mu = 0.43$			4 200 mese			
Severity = Mild			1.1					
COLCORONA	-0.58	0.5570		0.56	[0.19; 1.67]	0.4%	0.4%	
PRINCIPLE - Colchicine	-1.26	1.6287		0.28	[0.01; 6.92]	0.0%	0.0%	
Fixed effect model			\sim	0.52	[0.19; 1.47]	0.4%	11 T 17	
Random effects model			\sim	0.52	[0.19; 1.47]		0.4%	
Heterogeneity, $l^2 = 0\%$, τ^2 ($0, \rho =$	0.69						
Fixed effect model				0.99	[0.92; 1.05]	100.0%	A	
Random effects model		C + + +			[0.92; 1.05]		100.0%	
Heterogeneity: $I^2 = 0\%$, $\tau^2 =$			1 1 1	1				
Residual heterogeneity: 12 =			0.1 1 10	100				

PAHO

care in patients with COVID-19 Weight Weight Study TE seTE **Risk Ratio** 95%-Cl (fixed) (random) RR Severity = Moderate to critical 0.2% 1.1% GRECCO-19 -1.51 1.0779 0.22 [0.03; 1.82] 74.1% RECOVERY - Colchicine 0.04 0.0547 1.04 [0.93; 1.16] 46.3% COL-COVID -1.12 1.1378 0.33 [0.04; 3.04] 0.2% 1.0%

0.86 [0.71; 1.05]

0.50 [0.18; 1.43]

1.87 [0.36; 9.74]

0.99 (0.90; 1.08)

0.93 [0.76; 1.13]

0.53 [0.26; 1.09]

0.53 [0.26; 1.09]

0.53 [0.26; 1.09]

0.98 [0.89; 1.07] 100.0%

22.8%

0.8%

0.3%

1.6%

1.6%

98.4%

Figure 27. Mechanical ventilation in randomized studies comparing colchicine vs standard of

Random effects model Heterogeneity: $l^2 = 43\%$, $\tau^2 = 0.0254$, p = 0.10Residual heterogeneity: $l^2 = 36\%$, p = 0.17 0.1 0.5 1 2 10 Observed results apply mostly to hospitalized patients with moderate to critical disease. The COLCORONA trial that included patients with recent onset mild disease showed a tendency to less hospitalizations less mortality and less mechanical ventilation requirements. However, the

COLCORONA trial that included patients with recent onset mild disease showed a tendency to less hospitalizations, less mortality and less mechanical ventilation requirements. However, the certainty on those potential benefits was low because of very serious imprecision because of a small number of events.

Sofosbuvir +/- daclatasvir, ledipasvir, or velpatasvir

-0.15 0.0986

-0.68 0.5322

0.63 0.8408

-0.64 0.3710

See Summary of findings Table 16, Appendix 1

We identified 16 RCTs including 3,061 patients in which sofosbuvir alone or in combination with daclatasvir or ledipasvir was compared against standard of care or other treatments. Two studies compared sofosbuvir alone vs. standard of care, one study compared sofosbuvir alone vs. lopinavir-ritonavir, seven studies compared sofosbuvir + daclatasvir vs. standard of care, three studies compared sofosbuvir + daclatasvir vs. lopinavir-ritonavir, and three studies compared sofosbuvir + ledipasvir vs. standard of care. As there is moderate to high certainty that lopinavir-ritonavir is not related to significant benefits, we assumed that intervention as equivalent to standard of care. The DISCOVER trial was the biggest, with 1,083 patients and the only one categorized as with low risk of bias. Studies included patients with mild to severe disease. Our results showed:



COLCOVID

Cecconi et al

Fixed effect model

Severity = Mild COLCORONA

Fixed effect model

Fixed effect model

Random effects model

Heterogeneity, not applicable

Random effects model

Helengeneity /* = 26% +* = 0.0165, p = 0.17

Rabbani

37.4%

4.3%

1.8%

91.9%

8.1%

B.1%

- Sofosbuvir +/- daclatasvir or ledipasvir may increase mortality, RR 1.14 (95%CI 0.83 to 1.56); RD 2.2% (95%CI -2.7% to 9%); Low certainty ⊕⊕○○ (Figure 28) (based on low risk of bias studies)
- Sofosbuvir +/- daclatasvir or ledipasvir may not reduce mechanical ventilation requirements, RR 1.02 (95%CI 0.59 to 1.76); RD 0.3% (95%CI -7.1% to 13.1%); Low certainty ⊕⊕⊖○ (based on low risk of bias studies)
- Sofosbuvir +/- daclatasvir or ledipasvir probably does not improve time to symptom resolution, RR 1.01 (95%CI 0.95 to 1.08); RD 0.6% (95%CI -3% to 4.8%); Moderate certainty ⊕⊕⊕○ (based on low risk of bias studies)
- It is uncertain if sofosbuvir +/- daclatasvir or ledipasvir affects symptomatic infections in exposed individuals, RR 0.52 (95%CI 0.30 to 0.89); RD -8.3% (95%CI -12.1% to -1.9%); Very low certainty ⊕○○○
- It is uncertain if sofosbuvir +/- daclatasvir or ledipasvir increases severe adverse events, RR 0.35 (95%CI 0.06 to 2.19); RD -6.6% (95%CI -9.6% to 12.1%); Very low certainty ⊕○○○

Figure 28. Mortality in randomized studies comparing sofosbuvir +/- daclatasvir or ledipasvir vs standard of care in patients with COVID-19

Study	TE seTE	Risk Ratio	RR 95%	Weight Cl (fixed)	Weight (random)
RoB = High Abbaspour Kasgari H et al Sadeghi A et al Yakoot M et al (Pharco Corporate Khalili H et al Sali S et al Alavi-Moghaddam M et al Yadollahzadeh M et al Elgohary MAS et al El Bendary et al Abbass S et al Fixed effect model Random effects model Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0$, $p = 0$	-0.05 0.7860 -0.03 0.8698 -1.77 0.7117 0.33 0.8931 -2.56 1.4621 -0.42 0.3409 -0.69 0.5439		0.14 [0.01; 2. 0.60 [0.16; 2. 0.41 [0.08; 2. 0.95 [0.20; 4. 0.97 [0.18; 5. 0.17 [0.04; 0. 1.40 [0.24; 8. 0.08 [0.00; 1. 0.66 [0.34; 1. 0.50 [0.17; 1. 0.55 [0.36; 0. 0.55 [0.36; 0.	31] 3.4% 00] 2.5% 45] 2.6% 33] 2.1% 69] 3.2% 04] 2.0% 35] 0.8% 29] 13.9% 45] 5.4% 33] 36.6%	1.8% 7.1% 5.4% 5.7% 4.8% 6.7% 4.6% 1.9% 17.7% 10.1%
RoB = Low DISCOVER SOVECOD Fixed effect model Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0$, $p = 0$ Fixed effect model Bandom effects model	0.14 0.1628 0.00 0.7853 86		1.15 [0.83; 1. 1.00 [0.21; 4. 1.14 [0.83; 1. 1.14 [0.83; 1. 0.87 [0.68; 1.	66] 2.6% 66] 63.4% 66]	28.5% 5.7%
Random effects model Heterogeneity: $I^2 = 29\%$, $\tau^2 = 0.1213$ Residual heterogeneity: $I^2 = 0\%$, $p =$		0.01 0.1 1 10	0.68 [0.46; 1.0 100	.2]	100.0%



REGEN-COV (casirivimab and imdevimab)

See Summary of findings Table 17, Appendix 1

We identified 12 RCTs including 25,207 patients in which REGEN-COV (casirivimab and imdevimab) was compared against standard of care, or other treatments, in patients with recent onset COVID-19. The RECOVERY trial was the biggest, included severe to critical patients and reported differential effect in seronegative patients at baseline. Eight of the other nine studies included mild patients with recent onset disease or exposed individuals with negative PCR. Our results showed:

- Overall REGEN-COV may decrease mortality, RR 0.83 (95%CI 0.63 to 1.09); RD -2.7% (95%CI -5.9% to 1.4%); Low certainty ⊕⊕○○
- In seronegative patients REGEN-COV probably decreases mortality, RR 0.79 (95%CI 0.71 to 0.89); RD -3.4% (95%CI -4.6% to -1.8%); Moderate certainty ⊕⊕⊕○ (Figure 29)
- Overall REGEN-COV may decrease mechanical ventilation, RR 0.79 (95%CI 0.54 to 1.14); RD -3.6% (95%CI -8% to 2.4%); Low certainty ⊕⊕○○
- In seronegative patients REGEN-COV probably reduces mechanical ventilation, RR 0.82 (95%CI 0.74 to 0.9); RD -3.1% (95%CI -4.5% to -1.7%); Moderate certainty ⊕⊕⊕○
- Overall REGEN-COV may increase symptom resolution, RR 1.06 (95%CI 1 to 1.12); RD 3.6% (95%CI 0% to 7.2%); Low certainty ⊕⊕⊕○
- In seronegative patients REGEN-COV probably increases symptom resolution, RR 1.1 (95%CI 1.06 to 1.14); RD 6% (95%CI 3.6% to 8.5%); Moderate certainty ⊕⊕⊕○
- REGEN-COV reduces symptomatic infections in exposed individuals, RR 0.24 (95%CI 0.08 to 0.76); RD -13.2% (95%CI -16% to -4.2%); High certainty ⊕⊕⊕
- REGEN-COV probably does not increase severe adverse events, RR 0.51 (95%CI 0.38 to 0.67); RD -5% (95%CI -6.3% to -3.4%); Moderate certainty ⊕⊕⊕○
- REGEN-COV probably reduces hospitalization, RR 0.28 (95%CI 0.19 to 0.42); RD -3.5% (95%CI -3.9% to -2.8%); Moderate certainty ⊕⊕⊕○ (Figure 30)

Figure 29. Mortality in randomized studies comparing REGEN-COV vs standard of care in seronegative patients with COVID-19





Study	TE	seTE	F	Risk Rati	o	RR	95%-CI	Weight (fixed)	Weight (random)
RECOVERY - REGEN-COV Somersan-Karakaya		0.0589 0.2726		÷			[0.73; 0.92] [0.26; 0.76]	95.5% 4.5%	59.6% 40.4%
Fixed effect model Random effects model Heterogeneity: I^2 = 79%, τ^2 = 0.	1453, <i>µ</i>	o = 0.03	0.5	1	2		[0.71; 0.89] [0.36; 1.14]	100.0% 	 100.0%

Figure 30. Hospitalization in randomized studies comparing REGEN-COV vs standard of care in patients with COVID-19

Study	TE	seTE		Ri	sk Rat	lio		RR	95%-CI	Weight (fixed)	Weight (random)
Weinreich	-1.24	0.2251		1.18	ÐĐ.			0.29	[0.19; 0.45]	87.6%	87.6%
Covid-19 Phase 3 Prevention Trial - Asymptomatic	-1.91	1.5054	-					0.15	[0.01; 2.84]	2.0%	2.0%
Covid-19 Phase 3 Prevention Trial - Exposed	-2.56	1.4668	-	-+	-			0.08	[0.00; 1.36]	2.1%	2.1%
Weinreich_2	-1.14	0.7257		+	-			0.32	[0.08; 1.32]	8.4%	8.4%
Fixed effect model				0	>			0.28	[0.19; 0.42]	100.0%	
Random effects model				0	>			0.28	[0.19; 0.42]		100.0%
Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0$, $p = 0.80$			1	1		1	1				
			0.01	0.1	1	10	100				

In addition, two studies that compared REGEN-COV (casirivimab and imdevimab) against bamlanivimab +/- etesevimab and sotrovimab in non-severe patients with risk factors for severity reported no important differences in hospitalizations.

Aspirin

We identified four RCTs including 16,696 patients in which aspirin was compared against standard of care in patients with COVID-19. Our results showed:

- Aspirin probably does not reduce mortality, RR 0.95 (95%CI 0.89 to 1.02); RD -0.8% (95%CI -1.8% to 0.3; Moderate certainty ⊕⊕⊕○ (Figure 31)
- Aspirin probably does not reduce mechanical ventilation, RR 0.94 (95%CI 0.84 to 1.05); RD -1% (95%CI -2.8% to 0.9%); Moderate certainty ⊕⊕⊕○
- Aspirin probably does not increase symptom resolution or improvement, RR 1.02 (95%CI 1.0 to 1.04); RD 1% (95%CI -0.1% to 2.2%); Moderate certainty ⊕⊕⊕○

Figure 31. Mortality in randomized studies comparing aspirin vs standard of care in patients with COVID-19





Study	TE seTE	R	isk Ra	atio		RR	95%-CI	Weight (fixed)	Weight (random)
RESIST	-0.86 0.6834			_		0.42	[0.11; 1.62]	0.2%	0.3%
RECOVERY - ASA	-0.04 0.0363		+			0.97	[0.90; 1.04]	86.4%	85.3%
REMAP-CAP - ASA	-0.11 0.0922		-			0.89	[0.74; 1.07]	13.4%	14.4%
Fixed effect model			\$			0.95	[0.89; 1.02]	100.0%	
Random effects mode	el		¢.			0.95	[0.89; 1.02]		100.0%
Heterogeneity: I ² = 1%, a	$r^2 = 0.0001, p = 0.36$								
o y .		0.2 0.5	5 1	2	5				

Sotrovimab

See Summary of findings Table 18, Appendix 1

We identified three RCTs including 4,934 patients with recent onset mild COVID-19 and risk factors for severe disease, in which sotrovimab was compared against standard of care or other interventions. Our results showed:

- Sotrovimab probably reduces hospitalizations, RR 0.20 (95%CI 0.08 to 0.48); RD -3.8% (95%CI -4.6% to -2.5%); Moderate certainty ⊕⊕⊕○ (certainty upgraded because of evidence of equipoise of sotrovimab and REGEN-COV)
- Severe adverse events, RR 0.34 (95%CI 0.16 to 0.68); RD -6.7% (95%CI -8.6% to -3.3%); Moderate certainty ⊕⊕⊕○

One study that compared REGEN-COV and sotrovimab in mild to moderate patients showed similar hospitalization rates (RR 0.93 95%CI, 0.77 to 1.13)

Mesenchymal stem-cell transplantation

We identified nine RCTs including 338 patients with severe to critical COVID-19, in which mesenchymal stem-cell transplantation was compared against standard of care. Our results showed:

Mesenchymal stem-cell transplantation may reduce mortality, RR 0.61 (95%CI 0.43 to 0.88); RD -6.2% (95%CI -9.1% to -1.9%); Low certainty ⊕⊕⊖⊖ (Figure 32)

Figure 32. Mortality in randomized studies comparing mesenchymal stem-cell transplantation vs standard of care in patients with COVID-19





Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)
Shu L et al	-1.06 1.4724		0.35	[0.02; 6.19]	1.6%	1.6%
Lanzoni G et al	-0.92 0.7303				6.3%	6.3%
ISMMSCCOVID19	-0.47 0.2500	-	0.62	[0.38; 1.02]	54.2%	54.2%
Zhu R et al	-1.61 1.5268 -		0.20	[0.01; 3.99]	1.5%	1.5%
Fathi-Kazerooni M et al	-0.62 0.3345		0.54	[0.28; 1.03]	30.3%	30.3%
Rebelatto CK et al	1.00 0.9708		2.73	[0.41; 18.28]	3.6%	3.6%
Tavakol ASJ et al	0.69 1.1402		2.00	[0.21; 18.69]	2.6%	2.6%
Fixed effect model Random effects model Heterogeneity: $l^2 = 0\%$, τ^2		0.1 0.51 2 10		[0.43; 0.88] [0.43; 0.88]	100.0% 	 100.0%

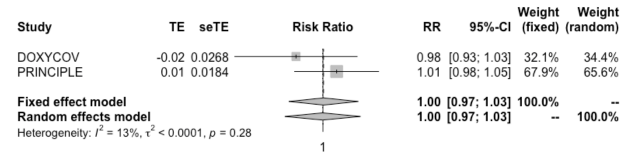


Doxycycline

We identified four RCTs including 2,415 patients with mild COVID-19, in which doxycycline was compared against standard of care. Our results showed:

- It is uncertain if doxycycline reduce or increase mortality, RR 1.10 (95%CI 0.63 to 1.93); RD 1.6% (95%CI -5.9% to 14.9%); Very low certainty ⊕○○○
- Doxycycline does not increase symptom resolution or improvement, RR 1 (95%CI 0.97 to 1.03); RD -0% (95%CI -91.8% to -1.8%); High certainty ⊕⊕⊕⊕ (Figure 33)
- Doxycycline may not reduce hospitalizations, RR 1.16 (95%CI 0.76 to 1.76); RD 0.7% (95%CI -1.1% to 3.6%); Low certainty ⊕⊕○○

Figure 33. Symptom resolution or improvement in randomized studies comparing doxycycline vs standard of care in patients with COVID-19



Inhaled corticosteroids

See Summary of findings Table 19, Appendix 1

We identified nine RCTs including 4,309 patients with mild COVID-19, in which inhaled coticosteroids were compared against standard of care. Our results showed:

- It is uncertain if inhaled corticosteroids reduce or increase mortality, RR 0.82 (95%CI 0.44 to 1.53); RD -2.8% (95%CI -9% to 8.5%); Very low certainty ⊕○○○
- It is uncertain if inhaled corticosteroids reduce or increase mechanical ventilation, RR 0.94 (95%CI 0.44 to 1.98); RD -1% (95%CI -9.6% to 17%); Very low certainty ⊕○○○
- Inhaled corticosteroids probably increase symptom resolution or improvement, RR 1.09 (95%CI 0.99 to 1.2); RD 5.5% (95%CI -0.6% to 12.1%); Low certainty ⊕⊕○○ (Figure 34)
- Inhaled corticosteroids probably does not have an important effect on hospitalizations, RR 0.9 (95%CI 0.7 to 1.15); RD -0.5% (95%CI -1.4% to 0.7%); Moderate certainty ⊕⊕⊕○
- It is uncertain if inhaled corticosteroids reduce or increase severe adverse events, RR 0.5 (95%CI 0.23 to 1.12); RD -5.1% (95%CI -7.9% to 1.2%); Very low certainty ⊕○○○





Figure 34. Symptom resolution or improvement in randomized studies comparing inhaled corticosteroids vs standard of care in patients with COVID-19

Study	TE se	TE	Risk Ratio	, j	RR	95%-CI	Weight (fixed)	Weight (random)
STOIC	0.09 0.10	001	-11-	1	.09	[0.90; 1.33]	1.9%	12.6%
PRINCIPLE	0.18 0.04		1			[1.10; 1.32]	8.8%	22.8%
KUMC-COVID-19	-0.06 0.22	286				[0.60; 1.47]	0.4%	3.7%
ALV-020-001	0.10 0.07	703	1			[0.97; 1.27]	3.9%	17.8%
CONTAIN	0.19 0.14	433				[0.91; 1.60]	1.0%	7.9%
NA	-0.21 0.3	174		0.	81	[0.43: 1.50]	0.2%	2.0%
COVERAGE	0.15 0.20	021		- 1.	16	[0.78; 1.73]	0.5%	4.5%
ACTIV-6 - Fluticazone	0.00 0.01	153		1.	.00	[0.97; 1.03]	83.3%	28.7%
Fixed effect model			0	1.	.02	[1.00; 1.05]	100.0%	
Random effects mode Heterogeneity: $I^2 = 62\%$,		n=0.01	\$	-1 1	.09	[0.99; 1.20]	- · ·	100.0%
, lotor og on only . 1 - oz /o,		0.5	1	2				

Fluvoxamine

See Summary of findings Table 20, Appendix 1

We identified four RCTs including 2,356 patients with COVID-19, in which fluvoxamine was compared against standard of care. Our results showed:

- It is uncertain if fluvoxamine reduces or increase mortality, RR 0.69 (95%CI 0.36 to 1.27); RD -5% (95%CI -10.2% to 4.3%); Very low certainty ⊕○○○
- It is uncertain if fluvoxamine reduces or increase mechanical ventilation, RR 0.77 (95%CI 0.45 to 1.3); RD -3.7% (95%CI -8.8% to 4.8%); Very low certainty ⊕○○○
- Fluvoxamine probably does not have an important effect on hospitalizations in patients with recent onset disease, RR 0.79 (95%CI 0.6 to 1.03); RD -1% (95%CI -1.9% to 0.1%); Moderate certainty ⊕⊕⊕○ (Figure 35). The observed effect would probably be considered important in patients with very high hospitalization risk.
- Fluvoxamine may not increase severe adverse events, RR 0.81 (95%CI 0.54 to 1.22); RD -1.9% (95%CI -4.7% to 2.2%); Low certainty ⊕⊕○○



Figure 35. Hospitalizations in randomized studies comparing fluvoxamine vs standard of care in patients with COVID-19

Study	TE seTE	Ri	sk Rati	0	RR	95%-CI	Weight (fixed)	Weight (random)
Lenze E et al TOGETHER-Fluvoxamine COVID-OUT - Fluvoxamine	-2.30 1.4818 -0.24 0.1435 e 0.17 0.6005	•			0.78	[0.01; 1.83] [0.59; 1.04] [0.36; 3.83]	93.8%	3.1% 80.7% 16.3%
Fixed effect model Random effects model Heterogeneity: I^2 = 17%, τ^2 =	0.0652, <i>p</i> = 0.30	l 0.1	1	10		[0.60; 1.03] [0.47; 1.32]	100.0% 	 100.0%

Molnupiravir

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See Summary of findings Table 21, Appendix 1

We identified ten RCTs including 4,532 patients with COVID-19, in which molnupiravir was compared against standard of care. Our results showed:

- It is uncertain if molnupiravir reduces or increase mortality, RR 0.35 (95%CI 0.06 to 2.19); RD -1.4% (95%CI -15% to 19.4%); Very low certainty ⊕○○○
- It is uncertain if molnupiravir reduces or mechanical ventilation, RR 0.36 (95%CI 0.11 to 1.12); RD -11.1% (95%CI -15.4% to 2.1%); Very low certainty ⊕○○○
- Molnupiravir reduces hospitalizations in patients with recent onset disease, RR 0.6 (95%CI 0.44 to 0.81); RD -1.9% (95%CI -2.7% to -0.9%); High certainty ⊕⊕⊕⊕ (Figure 36)
- Molnupiravir may increase symptom resolution, RR 1.17 (95%CI 1.1 to 1.3); RD 10.3% (95%CI 3.6% to -18.2%); Low certainty ⊕⊕○○
- Molnupiravir may not increase severe adverse events, RR 0.75 (95%CI 0.48 to 1.19); RD -2.6% (95%CI -5.3% to -1.9%); Low certainty ⊕⊕○○

Figure 36. Hospitalizations in randomized studies comparing molnupiravir vs standard of care in patients with COVID-19

Study	TE	seTE	Ri	sk Ratio	D		RR	95%-CI	Weight (fixed)	Weight (random)
EIDD-2801-2003	0.28 1	.1446	_				1.33	[0.14; 12.52]	1.7%	1.8%
MOVe-OUT	-0.36 0	.1808		+			0.70	[0.49; 0.99]	68.6%	66.8%
HCR/III/MOLCOV/04/2021-01	-1.19 0	.4254		<u>⊢</u> †			0.30	[0.13; 0.70]	12.4%	13.0%
CR216-21	-0.62 0	.4653	_	-			0.54	[0.22; 1.34]	10.4%	10.9%
AGILE	-2.20 1	.4832 —		++-			0.11	[0.01; 2.03]	1.0%	1.1%
MOVe-OUT - ph2	-0.55 0	.6123		+			0.58	[0.17; 1.91]	6.0%	6.4%
Fixed effect model Random effects model Heterogeneity: $I^2 = 1\%$, $\tau^2 = 0.0$	033, p = 0			\$	1			[0.45; 0.80] [0.44; 0.81]		 100.0%
		0.01	0.1	1	10	100				

Nirmatrelvir-ribavirin

See Summary of findings Table 22, Appendix 1

We identified one RCT including 2,085 patients with COVID-19, in which nirmatrelvir-ritonavir was compared against standard of care. Our results showed:

- It is uncertain if nirmatrelvir-ritonavir reduces or increase mortality, RR 0.04 (95%CI 0.002 to 0.68); RD -15.3% (95%CI -15.9% to -5.1%); Very low certainty ⊕○○○
- Nirmatrelvir-ritonavir probably reduces hospitalizations in patients with recent onset disease, RR 0.12 (95%CI 0.06 to 0.25); RD -5.2% (95%CI -7.1% to -2%); Moderate certainty ⊕⊕⊕○
- Nirmatrelvir-ritonavir probably does not increase severe adverse events, RR 0.49 (95%CI 0.30 to 0.80); RD -5.2% (95%CI -7.8% to 0.5%); Moderate certainty ⊕⊕⊕○

Ruxolitinib

See Summary of findings Table 23, Appendix 1

We identified three RCTs including 686 patients with COVID-19, in which ruxolitinib was compared against standard of care. RUXOCOVID-DEVENT was the biggest trial including 211 patients with critical COVID-19. Our results showed:

- Ruxolitinb may reduce mortality, RR 0.72 (95%CI 0.59 to 0.89); RD -4.5% (95%CI -6.5% to -1.7%); Low certainty ⊕⊕○○ (Figure 37)
- It is uncertain if ruxolitinib increases or decreses mechanical ventilation, RR 0.99 (95%CI 0.49 to 1.99); RD -0.1% (95%CI -8.8% to 17.%); Very low certainty ⊕○○○
- Ruxolitinib may not improve time to symptom resolution, RR 1.05 (95%CI 0.89 to 1.24); RD 3% (95%CI -6.7% to 14.5%); Low certainty ⊕⊕○○
- It is uncertain if ruxolitinib incresses or decreases severe adverse events, RR 1.12 (95%CI 0.69 to 1.82); RD 1.2% (95%CI -3.7% to 8.4%); Very low certainty ⊕○○○

Figure 37. Mortality in randomized studies comparing ruxolitinib vs standard of care in patients with COVID-19





Study	TE seTE	Risk Ratio	F	RR 95%-C	Weight (fixed)	Weight (random)
Cao Y et al	-1.99 1.4808		0.	14 [0.01; 2.48]	0.5%	3.9%
RUXCOVID	0.42 0.6588		1.	52 [0.42; 5.51]	2.6%	16.6%
RUXCOVID-DEVENT	-0.33 0.1073	+	0.	72 [0.58; 0.88	96.9%	79.5%
Fixed effect model		ف	0.	72 [0.59; 0.89]	100.0%	
Random effects mode		\Rightarrow	0.	76 [0.42; 1.36]		100.0%
Heterogeneity: $I^2 = 21\%$,	$\tau^2 = 0.1002, p = 0.28$					
	0.01	0.1 1	10 100			

CD24Fc

See Summary of findings Table 24, Appendix 1

We identified one RCT including 234 patients with COVID-19, in which CD24Fc was compared against standard of care. Our results showed:

- It is uncertain if CD24Fc reduces or increases mortality, RR 0.9 (95%CI 0.49 to 1.69); RD -1.5% (95%CI -8.2% to 11%); Very low certainty ⊕○○○
- CD24Fc may decrease mechanical ventilation, RR 0.57 (95%CI 0.34 to 0.96); RD -7.4% (95%CI -11.4% to -0.7%); Low certainty ⊕⊕○○
- CD24Fc may increase symptom resolution, RR 1.18 (95%CI 1 to 1.39); RD 10.7% (95%CI -0.2% to 23.4%); Low certainty ⊕⊕⊖⊖
- It is uncertain if CD24Fc increases or decreases severe adverse events, RR 0.98 (95%CI 0.61 to 1.57); RD -0.2% (95%CI -4% to 5.8%); Very low certainty ⊕○○○

Vitamin D

See Summary of findings Table 25, Appendix 1

We identified 20 RCTs including 44,071 patients with COVID-19, in which Vitamin D was compared against standard of care or other treatments. Our results showed:

- It is uncertain if vitamin D reduces or increases mortality, RR 1.24 (95%CI 0.8 to 1.91); RD 3.8% (95%CI -3.2% to 14.4%); Very low certainty ⊕○○○
- It is uncertain if vitamin D reduces or increases mechanical ventilation, RR 0.5 (95%CI 0.25 to 1); RD -8.6% (95%CI -13% to 0%); Very low certainty ⊕○○○
- It is uncertain if vitamin D reduces or increases symptom resolution or improvement, RR 1.78 (95%CI 1.1 to 2.94); RD 39.4.6% (95%CI 4.6% to 39.4%); Very low certainty ⊕○○○
- Vitamin D does not reduce symptomatic infections in exposed individuals, RR 1.06 (95%CI 0.91 to 1.24); RD 1% (95%CI -1.6% to 4.2%); High certainty ⊕⊕⊕⊕ (excluding high risk of bias studies) (Figure 38)



- Vitamin D probably does not reduce hospitalizations, RR 1.26 (95%CI 0.84 to 1.89); RD 1.2% (95%CI -0.8% to 4.3%); Moderate certainty ⊕⊕⊕○
- Vitamin D may not increase severe adverse events, RR 1.03 (95%CI 0.84 to 1.26); RD 0.3% (95%CI -1.6% to 2.7%); Low certainty ⊕⊕○○

Figure 38. Symptomatic infections in randomized studies comparing vitamin D vs standard of care in persons exposed to COVID-19

Study	TE seTE	Risk Ratio	RR	95%-CI	Weight (fixed)	Weight (random)	
RoB = Low		11					
CORONAVIT	0.22 0.1488	#	1.25	[0.93; 1.67]	26.6%	30.2%	
Brunvoll	-0.00 0.0931			[0.83; 1.20]		31.6%	
Fixed effect model		1.5		[0.91; 1.24]	94.6%	1000	
Random offects mod	el	•	1.08	[0.88; 1.34]	1.11	61.8%	
Helerogeneity: $l^2 = 3.6\%$				1.0.00.014			
RoB = High							
Villasis-Keever MA et	al -1.30 0.4100	-+	0.27	[0.12; 0.61]	3.5%	20.2%	
Romero-Ibarguengoitia	a -1.56 0.5981			[0.06; 0.68]		14.1%	
Van Helmond				[0.00; 0.43]		3.9%	
Fixed effect model		\Diamond		[0,12; 0.42]		-	
Random effects mod	el	0		[0,10: 0.46]	1.1.1.2	36,2%	
Haterogeneity: $I^2 = 20\%$		1					
Fixed effect model		Å	0.98	[0.84; 1.13]	100.0%		
Random effects mod	el	\diamond		[0.32; 1.03]		100.0%	
Heterogeneity: $I^2 = 84\%$	$\tau^2 = 0.2721, p < 0.01$		0			N 2124 2 9 2	
Residual heterogeneity:		01 0.1 1 10 10	00				
CALIFICATION OF STATES	C. M. C. LANK, C.	201 2010 0 001 95					

In addition one study that compared high dose vitamin D supplementation (cholecalciferol 400,000 IU) versus standard dose (cholecalciferol 50,000 IU) reported no significant differences in mortality at 28 days (HR 0.7 95%CI 0.36 to 1.36) in patients hospitalized for COVID-19.

Tixagevimab–Cilgavimab

See Summary of findings Table 26, Appendix 1

We identified three RCT including 7,492 individuals with COVID-19 or exposed to SARS-COV-2, in which Tixagevimab–cilgavimab was compared against standard of care. Our results showed:

• Tixagevimab–cilgavimab probably reduces mortality, RR 0.72 (95%CI 0.54 to 0.96); RD -4.5% (95%CI -7.4% to -0.6%); Moderate certainty ⊕⊕⊕○ (Figure 39)



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- Tixagevimab–cilgavimab probably does not increase symptom resolution or improvement, RR 1.03 (95%CI 0.99 to 1.08); RD 2% (95%CI -0.6% to 4.7%); Moderate certainty ⊕⊕⊕○
- Tixagevimab–cilgavimab probably reduces symptomatic infections in exposed individuals, RR 0.18 (95%CI 0.09 to 0.35); RD -14.2% (95%CI -15.8% to -11.2%); Moderate certainty ⊕⊕⊕○
- Tixagevimab–cilgavimab may not increase severe adverse events, RR 0.95 (95%CI 0.69 to 1.31); RD -0.5% (95%CI -3.2% to 3.2%); Low certainty ⊕⊕○○
- Tixagevimab–cilgavimab probably reduces mortality, RR 0.42 (95%CI 0.24 to 0.74); RD -2.8% (95%CI -3.6% to 1.3%); Moderate certainty ⊕⊕⊕○

Figure 39. Mortality in randomized studies comparing Tixagevimab–cilgavimab vs standard of care in patients with COVID-19

Study	TE seTE	Risk Ratio	RR	95%-Cl	Weight (fixed)	Weight (random)
PROVENT	-0.44 0.5031			[0.24; 1.73] [0.32; 3.07]	8.5% 6.5%	8.5% 6.5%
TICO	-0.35 0.1587		0.71	[0.52; 0.96]	85.0%	85.0%
Fixed effect mode Random effects m Heterogeneity: $l^2 = 0$	nodel			[0.54; 0.96] [0.54; 0.96]	100.0%	100.0%
		0.5 1 2				

Vilobelimab

See Summary of findings Table 27, Appendix 1

We identified two RCT including 398 individuals with severe to critical COVID-19 in which vilobelimab was compared against standard of care. Our results showed:

- Vilobelimab probably reduces mortality, RR 0.76 (95%CI 0.6 to 0.98); RD -3.8% (95%CI -6.4% to -0.3%); Moderate certainty ⊕⊕⊕○ (Figure 40)
- Tixagevimab–cilgavimab may not increase severe adverse events, RR 0.94 (95%CI 0.8 to 1.11); RD -0.6% (95%CI -2% to 1.1%); Moderate certainty ⊕⊕⊕○

Figure 40. Mortality in randomized studies comparing vilobelimab vs standard of care in patients with COVID-19





Study	TE	seTE		Ris	k Ra	tio		RR	95%-CI	Weight (fixed)	Weight (random)
Vlaar APJ et al PANAMO vilobelimab		0.8272	-		H	_	5		[0.12; 3.04] [0.60; 0.99]	2.4% 97.6%	2.4% 97.6%
Fixed effect model Random effects mode Heterogeneity: $l^2 = 0\%$, τ^2		410000	Ē		1	1	-1	0.76	[0.60; 0.98] [0.60; 0.98]		100.0%
			0.2	0.5	1	2	5				



Full description of included studies

Table 5, below, lists all the identified studies that were included in this systematic review by intervention. The treatments are arranged in alphabetical order. Study or author names, publication status, patient populations, interventions, sources of bias, outcomes, effect sizes and certainty are listed for each study.





99mTc-MDP Uncertainty in potential benefits and harms. Further research is needed.						
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (SOC) and GRADE certainty of the evidence	
RCT						
<u>Yuan et al</u> ; ¹⁵ preprint; 2020	Patients with mild COVID-19 infection. 10 assigned to 99mTc- MDP 5/ml once a day for 7 days and 11 assigned to standard of care.	Median age 61 ± 20, male 42.9%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information	

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Table 5. Description of included studies and interventions effects





Adalimumab Uncertainty in potential benefits and harms. Further research is needed.						
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence	
RCT	•				-	
Fakharian A et al trial; ¹⁶ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 34 assigned to adalimumab 40 mg once and 34 assigned to SOC	Mean age 54.6 ± 12, male 58.8%, hypertension 29.4%, diabetes 27.9%, COPD 1.5%, CHD 4.4%, CKD 1.5%, cancer 1.5%	Corticosteroids 100%, remdesivir 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕⊕○○Invasive mechanical ventilation: Very low certainty ⊕⊕○○Symptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: No informationHospitalization: No information	
	Uncerta	Alpha-1 inty in potential benefits a	antitrypsin and harms. Further rese	arch is needed.		
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence	



RCT	RCT							
McElvaney et al; ¹⁷ peer reviewed; 2021	Patients with critical COVID-19 infection. 25 assigned to alpha-1 antitrypsin 120 mg/kg once a week and 11 assigned to SOC	Mean age 58.4 ± , male 61.1%, hypertension 44.4%, diabetes 27.7%, COPD 30.5%, CHD 16.6%, CKD 27.7%, obesity 66.6%	Corticosteroids 72.2%, remdesivir 0%, hydroxychloroquine 0%, tocilizumab 0%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty $\oplus \oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationSymptomatic informationSymptomatic informationSymptomatic informationSymptomatic informationHospitalization: No information			
	Amiodarone Uncertainty in potential benefits and harms. Further research is needed.							
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence			
RCT								
ReCOVery-SIRIO trial; ¹⁸ Navarese et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 71 assigned to amiodarone 200 to 400 mg a day and 72 assigned to SOC	Median age 61.3 , male 62.3%, diabetes 23.7%, COPD 6.5%, cancer 7%,	Remdesivir 1.9%, hydroxychloroquine 2.3%, azithromycin 6%, convalescent plasma 1.9%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of	Mortality: Very low certainty $\oplus \oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\oplus \oplus \bigcirc \bigcirc$			





	Uncertai	Ammoni inty in potential benefits a	um chloride and harms. Further resea	allocation probably inappropriate. arch is needed.	Symptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕⊕○○Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
<u>Siami et al</u> ; ¹⁹ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 60 assigned to ammonium chloride 125 mg and 60 assigned to SOC	NR	Corticosteroids 100%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Blinding and concealment probably inappropriate.	Mortality: Very low certainty ⊕⊕○○ Invasive mechanical ventilation: Very low certainty ⊕⊕○○ Symptom resolution or improvement: No information Symptomatic





					infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncertai	AMP5A inty in potential benefits a	(inhaled) nd harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
<u>AP-014 trial</u> ; ²⁰ Roshon et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 19 assigned to AMP5A (inhaled) four nebulization a day for 5 days and 21 assigned to SOC	Mean age 64 ± 15, male 62.5%	Corticosteroids 78%, remdesivir 40%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕⊕○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationinformationAdverse events: Very low certainty ⊕⊕○○

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					Hospitalization: No information
Anakinra may not	reduce mortality or inc	rease severe adverse even	akinra ts. However the cerrtaint rther research is needed.	y of the evidence was low	because of risk of bias
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT				L	
CORIMUNO- <u>ANA-1 tria</u> l; ²¹ Bureau et al; Peer reviewed; 2020	Patients with mild to moderate COVID-19. 59 assigned to anakinra 400 mg a day for 3 days followed by 200 mg for 1 day followed by 100 mg for 1 day and 55 assigned to SOC	Median age 66 ± 17, male 70%, diabetes 29.8%, COPD 7.9%, asthma 7%, CHD 31.6%, cancer 9.6%,	Corticosteroids 46.5%, hydroxychloroquine 5.3%, lopinavir- ritonavir 3.5%, tocilizumab 0.8%, azithromycin 24.6%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: RR 0.96 (95%CI 0.57 to 1.6); RD -0.6% (95%CI - 6.9% to 9.6%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
<u>SAVE-MORE</u> <u>trial</u> ; ²² Kyriazopoulou et al; preprint; 2021	Patients with moderate to severe COVID-19 infection. 405 assigned to anakinra 100 mg SC a day for 7 to 10 days and 189 assigned to SOC	Mean age 61.9 ± 12.1, male 57.9%, diabetes 15.8%, COPD 4%, asthma %, CHD 3%, CKD 1.7%	Corticosteroids 86.2%, remdesivir 71.9%, azithromycin 18.7%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection





COV-AID-3 trial; ²³ Declercq et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 112 assigned to anakinra 100 mg a day for 28 days and 230 assigned to SOC	Mean age 65.5, male 77.4%, hypertension 46.4%, diabetes 27.7%, COPD %, CHD 20.5%, CKD 10.8%	Corticosteroids 62.3%, remdesivir 5%, hydroxychloroquine 11.7%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	(prophylaxis studies): No information Adverse events: RR 0.98 (95%CI 0.78 to 1.24); RD -0.2% (95%CI -2.2% to 2.5%); Low certainty ⊕⊕○○
Kharazmi et al; ²⁴ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 15 assigned to anakinra 100 mg a day for up to 14 days and 15 assigned to SOC	Mean age 54.1, male 63.3%, hypertension 33.3%, diabetes 36.6%, CHD 26.6%	Corticosteroids 63.3%, remdesivir 20%, lopinavir-ritonavir 63.3%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Hospitalization: No information
<u>Zeyad et al</u> ; ²⁵ preprint; 2022	Patients with severe to critical COVID-19 infection. 40 assigned to Anakinra 200 mg a day for 3 days and 40 assigned to SOC	Mean age 49.9 ± 11.7, male 82.5%, diabetes 43.8%, COPD 1.3%, CHD 8.8%, CKD 1.3%	Corticosteroids 100%, remdesivir 83.8%, azithromycin 78.8%, convalescent plasma 67.5%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
ANACONDA trial; ²⁶ Audemard- Verger et al; peer reviewed; 2022	Patients with severe COVID-19 infection. 36 assigned to anakinra 400 mg a day for 3 days followed by 200 mg a day for 7 days and 34 assigned to SOC	Mean age 70.6 , male 73.2%, hypertension 49.3%, diabetes 21.1%, COPD 9.9%, asthma 4.2%, CHD 12.7%, CKD 9.9%	Corticosteroids 63.4%, hydroxychloroquine 1.5%, azithromycin 12.6%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	





Angiotensin-converting enzyme inhibitors (ACEIs) or angiotensin receptor blockers (ARBs) Continuing or initiating ACEIs or ARBs may not reduce mortality. Further research is needed to confirm or discard these findings						
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence	
RCT						
<u>REPLACE</u> <u>COVID trial</u> ; ²⁷ Cohen et al; Peer reviewed; 2020	Patients with mild to severe COVID-19 previously treated with ACEI/ARB. 75 assigned to continuation of ACEI/ARB and 77 assigned to discontinuation of ACEI/ARB	55.5%, hypertension	NR	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: RR 1.13 (95%CI 0.77 to 1.64); RD 2.1% (95%CI - 3.7% to 10.2%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: RR 0.89 (95%CI 0.66 to 1.22); RD -1.9% (95%CI - 5.9% to 3.8%); Low certainty $\oplus \oplus \bigcirc \bigcirc$	
BRACE CORONA trial; ²⁸ Lopes et al; Peer reviewed; 2020	Patients with mild to moderate COVID-19. 334 assigned to continuation of ACEI/ARB and 325 assigned to discontinuation of ACEI/ARB	Median age 55.5 ± 19, male 59.6%, hypertension 100%, diabetes 31.9%, COPD %, asthma 3.9%, CHD 4.6%, CKD 1.4%, cancer 1.5%,	Corticosteroids 49.5%, hydroxychloroquine 19.7%, tocilizumab 3.6%, azithromycin 90.6%, convalescent plasma %, antivirals 42%	Some concerns for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Open label study with blinded outcome assessment. Significant number of patients excluded after randomization.	Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: Very low certainty	



ACEI-COVID trial; ²⁹ Bauer et al; peer reviewed; 2021		Mean age 72 ± 11, male 63%, hypertension 98%, diabetes 33%, CHD 22%	Remdesivir 6.8%	Low for mortality and mechanical ventilation; some Concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	000
<u>ATTRACT trial</u> ; ³⁰ Tornling et al; peer reviewed; 2020	Patients with moderate to severe COVID-19. 51 assigned to C21 (ARB) 200 mg a day for 7 days and 55 assigned to SOC	Mean age 52.6 ± 10.3, male 75.5%, hypertension 30.2%, diabetes 34%	Corticosteroids 84.9%, remdesivir 67%, hydroxychloroquine 13.2%	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	
<u>Nouri-Vaskeh et</u> <u>al;</u> ³¹ Peer reviewed; 2020	Patients with mild to severe COVID-19 infection and non- treated hypertension. 41 assigned to losartan 50 mg a day for 14 days and 39 assigned to Amlodipine 5 mg a day for 14 days	Mean age 63.5 ± 16, male 51.2%, diabetes 23.7%, COPD 15%, asthma %, CHD 18.7%,	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
SURG-2020-28683 trial; ³² Puskarich et al; Preprint; 2021	Patients with mild to moderate COVID-19 infection. 58 assigned to losartan 25 mg a day for 10 days and 59 assigned to SOC	Age (35-54) 46%, male 51.4%, hypertension 7.7%, diabetes 6%, COPD %, asthma 10.2%	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	





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<u>COVID-ARB</u> <u>trial</u> ; ³³ Geriak et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 16 assigned to losartan 25 mg a day for 10 days and 15 assigned to SOC	Median age 53, male %, hypertension 38.7%, diabetes 25.8%, CHD 3.2%, obesity 41.9%	Corticosteroids 22.6%, remdesivir 29%, hydroxychloroquine 9.7%, , azithromycin 16.1%, convalescent plasma 6.5%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Duarte et al</u> ; ³⁴ peer reviewed; 2020	Patients with moderate to severe COVID-19 infection. 71 assigned to telmisartan 80 mg twice daily and 70 assigned to SOC	Mean age 66 ± 17, male 53.2%, hypertension 44.3%, diabetes 19%, chronic lung disease 11.4%, asthma 1.3%, CHD NR%, CKD 3.2%, cerebrovascular disease 6.9%, obesity 15.2%	Corticosteroids 50.6%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate. Significant number of exclusions post randomization. Stop early for benefit in the context of multiple interim analysis.	
<u>Najmeddin et al;</u> ³⁵ peer reviewed; 2021	Patients with severe COVID-19 infection. 28 assigned to continuation of ACEI/ARB and 29 assigned to discontinuation of ACEI/ARB	Mean age 66.3 ± 9.9, male 46.9%, diabetes 50%, COPD 1.6%, CHD 25%, CKD 1.6%, cancer 4.7%,	Corticosteroids 42.2%, remdesivir 10.9%, , azithromycin 9.4%,	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events Notes: 10.9% lost to follow-up	
<u>ALPS-COVID</u> <u>trial</u> ; ³⁶ Puskarich et al; peer reviewed; 2021	Patients with moderate COVID-19 infection. 101 assigned to ACEI/ARB losartan 100 mg a day and 104 assigned to SOC	Mean age 55, male 60%, hypertension 42%, diabetes 22.9%, COPD 11.7%, asthma 13.2%, CHD 7.8%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	





<u>COVID MED</u> <u>trial;</u> ³⁷ Freilich et al; preprint; 2021	Patients with moderate to severe COVID-19 infection. 9 assigned to losartan 25 mg and 5 assigned to SOC	Mean age 63, male 64.2%, diabetes 7.1%, COPD 42.9%, asthma %, CHD 42.9%, CKD 0%, immunosuppression 35.7%, obesity 14.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>RAAS-COVID-19</u> <u>trial</u> ; ³⁸ Sharma et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 25 assigned to continuation of ACEI/ARB and 21 assigned to discontinuation of ACEI/ARB	Mean age 71.5 ± 12.9, male 56.5%, hypertension 100%, diabetes 43.5%, COPD 4.4%, CKD 19.6%, cerebrovascular disease 6.5%, cancer 6.5%,	Corticosteroids 47.8%,	Low for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
There are specifi	c recommendations on t		Dagulants gents ⁸ for thrombonron	ylaxis in hospitalized pati	ents with COVID-19.
1 mg/kg twice a day	y) probably do not decrea	ase mortality in comparise	on with prophylactic dose	arin 1 mg/kg a day) or ful e (i.e., enoxaparin 40 mg a r bleeding in comparison y	day). Anticoagulants in
1 mg/kg twice a day	y) probably do not decrea	ase mortality in comparise	on with prophylactic dose		day). Anticoagulants in
1 mg/kg twice a day intermediate or ful Study; publication	 probably do not decrea l dose decrease venous the second sec	ase mortality in comparise hromboembolic events bu	on with prophylactic dos t probably increase majo Additional	e (i.e., enoxaparin 40 mg a r bleeding in comparison Risk of bias and study	day). Anticoagulants in with prophylactic dose. Interventions effects vs standard of care and GRADE certainty of the
1 mg/kg twice a day intermediate or ful Study; publication status	 probably do not decrea l dose decrease venous the second sec	ase mortality in comparise hromboembolic events bu	Additional interventions Corticosteroids 70%, hydroxy-chloroquine	e (i.e., enoxaparin 40 mg a r bleeding in comparison Risk of bias and study	day). Anticoagulants in with prophylactic dose. Interventions effects vs standard of care and GRADE certainty of the





<u>REMAP-CAP</u> , <u>ACTIV-4a</u> , <u>ATTACC trial</u> ; ⁴⁰ Zarychanski et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 534 assigned low molecular weight heparin therapeutic dose (i.e., enoxaparin 1 mg/kg twice a day) and 564 assigned to prophylactic dose (i.e., enoxaparin 40 mg a day)	Mean age 61 ± 12.5, male 70%, diabetes 32.7%, COPD 24.1%, CHD 6.9%, CKD 9.6%,	Corticosteroids 79.3%, remdesivir 30.8%, tocilizumab 1.8%,	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Open-label study but outcome assessors were blinded.	resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Venous
INSPIRATION trial; ⁴¹ Sadeghipour et al; peer reviewed; 2021	Patients with moderate to critical COVID-19 infection. 276 assigned to low molecular weight heparin intermediate dose (i.e., enoxaparin 1 mg/kg a day) and 286 assigned to low molecular weight heparin prophylactic dose (i.e., enoxaparin 40 mg a day)	Median age 62 ± 21, male 57.8%, hypertension 44.3%, diabetes 27.7%, COPD 6.9%, CHD 13.9%, CKD %, cerebrovascular disease 3%	Corticosteroids 93.2%, remdesivir 60.1%, lopinavir-ritonavir 1%, tocilizumab 13.2%	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Open-label study but outcome assessors were blinded.	thromboembolic events (intermediate dose): RR 0.82 (95%CI 0.43 to 1.59); RD -1.3% (95%CI -4% to 4.1%); Low ⊕⊕○○ Venous thromboembolic events (therapeutic dose): RR 0.56 (95%CI 0.44 to 0.71); RD -3.1% (95%CI -
Perepu et al; ⁴² preprint; 2021	Patients with severe to critical COVID-19 infection. 87 assigned to low molecular weight heparin intermediate dose (i.e., enoxaparin 1 mg/kg a day) and 86 assigned to low molecular weight heparin prophylactic dose (i.e., enoxaparin 40 mg a day)	Median age 64 ± 62, male 56%, hypertension 60%, diabetes 37%, COPD 23%, CHD 31%, cancer 12%, obesity 49%	Corticosteroids 75%, remdesivir 61%, azithromycin 21%, convalescent plasma 27%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	RD -3.1% (95%CI - 3.9% to -2%); High ⊕⊕⊕⊕ Major bleeding: RR 1.56 (95%CI 1.08 to 2.25); RD 1.1% (95%CI 0.2% to 2.4%); Moderate ⊕⊕⊕⊖ Hospitalization: No information
<u>REMAP-CAP,</u> <u>ACTIV-4a,</u> <u>ATTACC trial</u> ; ⁴³ Zarychanski et al; preprint; 2021	Patients with moderate to severe COVID-19 infection. 1171 assigned to enoxaparin 1 mg/kg twice a day and 1048	Mean age 59 ± 14, male 58.7%, hypertension 51.8%, diabetes 29.7%, COPD 21.7%, CHD 10.6%, CKD 6.9%, immunosuppressive	Corticosteroids 61.7%, remdesivir 36.4%, tocilizumab 0.6%,	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse	





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	assigned to low molecular weight heparin prophylactic dose (i.e., enoxaparin 40 mg a day)	therapy 9.7%		events Notes: Open-label study but outcome assessors were blinded.
ACTION trial; ⁴⁴ Lopes et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 311 assigned to enoxaparin 1 mg/kg twice a day or rivaroxaban 20 mg a day and 304 assigned to low molecular weight heparin prophylactic dose (i.e., enoxaparin 40 mg a day) or unfractionated heparin prophylactic dose		Corticosteroids 83%	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Although patients and careers were aware of the intervention arm assigned, outcome assessors were blinded.
<u>RAPID trial</u> ; ⁴⁵ Sholzberg et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 228 assigned to therapeutic anticoagulation (i.e., enoxaparin 1 mg/kg) twice a day and 237 assigned to low molecular weight heparin prophylactic dose (i.e., enoxaparin 40 mg a day) or unfractionated heparin prophylactic dose	Mean age 60 ± 14.5, male 56.8%, hypertension 43.8%, diabetes 34.4%, COPD 13.5%, asthma %, CHD 7.3%, CKD 7.1%, cerebrovascular disease 4.1%, cancer 6.9%,	Corticosteroids 69.4%	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Open-label study but outcome assessors were blinded.
HEP-COVID trial; ⁴⁶ Spyropoulos et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 129 assigned to enoxaparin 1 mg/kg twice a day and 124 assigned to low molecular weight		Corticosteroids 81%, remdesivir 70.6%,	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events



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	heparin prophylactic dose (i.e., enoxaparin 40 mg a day) or unfractionated heparin prophylactic dose			
BEMICOP trial; ⁴⁷ Marcos et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 33 assigned to bemiparin 115 IU/kg once daily and 32 assigned to low molecular weight heparin prophylactic dose (i.e., enoxaparin 40 mg a day) or unfractionated heparin prophylactic dose	Mean age 62.7 ± 13, male 63.1%, hypertension 33.8%, diabetes 7.7%, COPD 16.9%, asthma %, CHD 6.2%, cancer 3.1%,	Corticosteroids 95.4%, remdesivir 13.8%, tocilizumab 23.1%	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>Oliynyk et al</u> ; ⁴⁸ peer reviewed; 2021	Patients with severe COVID-19 infection. 84 assigned to enoxaparin 100 anti- Xa IU/kg twice a day or unfractionated heparin 80 U/kg/h intravenously, followed by a maintenance dose of 18 U/kg/h and 42 assigned to enoxaparin enoxaparin 50 anti-Xa IU/kg a day	Mean age 70.6, male 60.3%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>X-Covid 19 trial</u> ; ⁴⁹ Morici et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 91 assigned to enoxaparin 40 mg twice a day and 92 assigned to low	Mean age 59 ± 21, male 62.8%, hypertension 36.1%, diabetes 13.7%, COPD 5.5%, CKD 1.6%, cerebrovascular disease 2.7%	Corticosteroids 45.9%, remdesivir 21.8%, tocilizumab 1.1%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded

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	molecular weight heparin prophylactic dose (i.e., enoxaparin 40 mg a day) or unfractionated heparin prophylactic dose			study which might have introduced bias to symptoms and adverse events outcomes results.	
PROTHROMCO VID trial; ⁵⁰ Muñoz- Rivas et al; preprint; 2021	Patients with severe COVID-19 infection. 103 assigned to tinzaparin 175 IU/kg once daily, 91 assigned to tinzaparin 100 IU/kg once daily and 106 assigned to tinzaparin 4500 IU once daily	Mean age 56.3, male 60.6%, hypertension 33%, diabetes 16.7%, COPD 4%, CHD 3.3%, CKD 2%, cerebrovascular disease 1.3%	Corticosteroids 89.3%, remdesivir 18%, tocilizumab 15%; Vaccinated 23%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
COVID-HEP trial; ⁵¹ Blondon et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 79 assigned to enoxaparin 1 mg/kg twice daily and 80 assigned to enoxaparin 20 to 60 mg once daily. Critically ill patients received enoxaparin 40 mg twice daily.	Mean age 62 ± 12, male 66%, hypertension 36.5%, diabetes 18.9%, COPD 11.9%, CHD 9.4%, cancer 6.3%	Corticosteroids 94.3%, tocilizumab 11.9%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>TACOVID trial</u> ; ⁵² Rashidi et al; peer reviewed; 2022	Patients with severe to critical COVID-19 infection. 5 assigned to UFH 80 IU/kg and 5 assigned to UFH 15000 IU a day	Mean age 61.5, male 60%, hypertension 40%, diabetes 30%, CHD 10%, CKD 0%, cancer 0%, obesity 20%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	

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<u>Kumar et al</u> ; ⁵³ peer reviewed ; 2021	Patients with moderate COVID-19 infection. 115 assigned to rivaroxaban 10 to 15 mg a day and 113 assigned to LMWH-P	Mean age 53 ± , male 71.3%, hypertension 26.6%, diabetes 30.3%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
ACTIV-4B trial; ⁵⁴ Connors et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 278 assigned to apixaban 2.5 to 5 mg twice a day and 136 assigned to SOC	Median age 54 ± 13, male 40.9%, hypertension 35.3%, diabetes 18.3%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: No information Invasive mechanical ventilation: No information
<u>Gates MRI</u> <u>RESPOND-1</u> <u>trial</u> , ⁵⁵ Ananworanich et al; peer reviewed; 2021	Patients with mild covid-19 and risk factors for severity. 222 assigned to rivaroxaban 10 mg a day and 222 assigned to SOC	Median age 49, male 39.3%, hypertension 51.8%, diabetes 27.7%, COPD 6.1%, immunosuppressive therapy 3.4%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Symptom resolution or improvement: RR 1.08 (95%CI 0.92 to 1.27); RD 4.8% (95%CI -4.8% to 16.4%); Low
OVID trial; ⁵⁶ Barco et al; peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 234 assigned to LMWH-P enoxaparin 40 mg a day for 14 days and 238 assigned to SOC	Mean age 56.5 ± , male 54%, hypertension 24.4%, diabetes 8%, COPD 2%, asthma %, CHD %, CKD %, cerebrovascular disease %, immunosuppresive therapy %, cancer %, obesity %	lopinavir-ritonavir %, tocilizumab %, azithromycin %,	Low for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	 ⊕⊕○○ Symptomatic infection (prophylaxis studies): No information Venous thromboembolic events
ETHIC trial; ⁵⁷ Cools et al; peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 105 assigned to enoxaparin 40 mg a day for 21 days and 114 assigned to SOC	Mean age 59 ± , male 55.7%, hypertension 70.4%, diabetes 30.8%, COPD 12.3%, cerebrovascular disease 1.8%, immunosuppression 2.5%, cancer 1.2%	Vaccinated 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably	<pre>(intermediate dose): No information Clinically important bleeding: Very low certainty ⊕○○○ Hospitalization: RR</pre>

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				inappropriate.	0.94 (95%CI 0.55 to 1.59); RD -0.3% (95%CI -2.2% to 2.8%); Low ⊕⊕⊖⊖
		20 (aspirin, proi inty in potential benefits		micronutrients) esearch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT		F			
Kumar et al; ⁵⁸ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 99 assigned to APMV2020 (aspirin 150 mg, promethazine 5 mg, vit D 2000 IU, vit C 750 mg, niacinamide 80 mg, zinc 15 mg, potassium 100 micrograms, sodioum selenate 82.5 micrograms) twice a day for 10 days and 93 assigned to SOC	Mean age 37 ± , male 55.5%	Vaccinated 95%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: Very low certainty ⊕○○○



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	Uncerta	Api inty in potential benefits :	•epitant and harms. Further r	esearch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT	•	•		• •	
Mehboob et al; ⁵⁹ preprint; 2020	Patients with mild to critical COVID-19 infection. 10 assigned to aprepitant 80 mg once a day for 3–5 days and 8 assigned to standard of care	Mean age 54.2 ± 10.91, male 61.1%,	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanica ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No
	Uncerta	Ap inty in potential benefits	rotinin and harms. Further r	esearch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence



Redondo-Calvo et al; ⁶⁰ peer reviewed; 2021	Patients with severe COVID-19 infection. 28 assigned to aprotinin 500 KIU a day for 11 days and 32 assigned to SOC	Mean age 55, male 65%, hypertension 47.4%, diabetes 29.8%, COPD 10.8%, CHD 17%	Corticosteroids 96.5%, remdesivir 12%, tocilizumab 10.5%, Vaccinated 35.1%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Significant loss to follow up.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncertai	${f A}{f i}$ inty in potential benefits a	•bidol and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT			J	<u> </u>	
<u>Khodashahi et al</u> ; ⁶¹ peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 50 assigned to arbidol 600 mg a day for 7 days and 50 assigned to SOC	Mean age 60.6 ± 19, male 55.6%, hypertension 13%, diabetes 12%	Hydroxychloroquine 100%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No





				events outcomes results.	information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
			nin, frankincens and harms. Further resea	e, and vitamin C) arch is needed.)
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT				<u>I</u>	<u>.</u>
MGC-006 trial; ⁶² Hellou et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 33 assigned to ArtemiC (artemisinin, curcumin, frankincense and vitamin C) oral spray twice a day and 17 assigned to SOC	Mean age 52 ± , male 50%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events:

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					Very low certainty ⊕○○○ Hospitalization: No information
	Uncertai	Arte inty in potential benefits a	e misinin and harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT				-	
<u>ARTI-19 trial</u> ; ⁶³ Tieu et al; Preprint; 2020	Patients with mild to moderate COVID-19. 39 assigned to artemisinin 500 mg for 5 days and 21 assigned to SOC	Mean age 43.3 ± 11.9, male 63.3%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
Aspirin probably	does not reduce mortali		Spirin on and probably does no	ot increase symptom resolu	tion or improvement.
Study; publication	Patients and	Comorbidities	Additional	Risk of bias and study	Interventions effects





status	interventions analyzed		interventions	limitations	vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
<u>RESIST trial;</u> ⁶⁴ Ghati et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 221 assigned to aspirin 75 mg once a day for 10 days and 219 assigned to SOC	Mean age 53.1 ± 9.2, male 73.3%, hypertension 28.6%, diabetes 27.7%, CHD 1.1%, CKD 2.4%	Corticosteroids 27.3%, remdesivir 20.6%, hydroxychloroquine 9.9%, tocilizumab 0.6%, convalescent plasma 0.2%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Blinding and concealment probably inappropriate.	Mortality: RR 0.95 (95%CI 0.89 to 1.02); RD -0.8% (95%CI - 1.8% to 0.3%); Moderate certainty ⊕⊕⊕○ Invasive mechanical ventilation: RR 0.94
RECOVERY - <u>ASA trial;</u> ⁶⁵ Horby et al; peer reviewed; 2021	Patients with moderate to critical COVID-19 infection. 7351 assigned to aspirin 150 mg a day and 7541 assigned to SOC	Median age 59.2 ± 14.2, male 61.5%, diabetes 22%, COPD 19%, asthma %, CHD 10.5%, CKD 3%,	Corticosteroids 94%	Low for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	(95%CI 0.84 to 1.05); RD -1% (95%CI - 2.8% to 0.9%); Moderate certainty ⊕⊕⊕○ Symptom resolution or improvement: RR 1.02 (95%CI 1.0 to 1.04); RD 1% (95%CI -0.1% to 2.2%); Moderate certainty
<u>ACTIV-4B trial</u> ; ⁵⁴ Connors et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 144 assigned to aspirin 81 mg a day and 136 assigned to SOC	Median age 54 ± 13, male 40.9%, hypertension 35.3%, diabetes 18.3%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	⊕⊕⊕○ Symptomatic infection (prophylaxis studies): No information
<u>REMAP-CAP -</u> <u>ASA trial</u> ; ⁶⁶ Bradbury et al; peer reviewed; 2021	critical COVID-19	Median age 57, male 65%, hypertension %, diabetes 22.7%, CHD 4.2%, CKD 3.4%	Corticosteroids 98.1%, remdesivir 22%, tocilizumab 42.9%	Low for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to	Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: Very low certainty \oplus \bigcirc \bigcirc

			vir/ritonavir	symptoms and adverse events outcomes results.	
Study; publication status	Uncertai Patients and interventions analyzed	inty in potential benefits a	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
Nekoukar et al; ⁶⁷ peer reviewed; 2021	Patients with severe COVID-19 infection. 62 assigned to atazanavir/ritonavir 300/100 mg a day for 5 to 10 days and 62 assigned to lopinavir- ritonavir 200/50 mg a day for 5 to 10 days	Mean age 49.9 ± 12.6, male 55.6%, hypertension 16.9%, diabetes 27.4%, COPD 0.8%, asthma 1.6%,	Corticosteroids 42.7%, remdesivir 13.7%, tocilizumab 3.2%, azithromycin 50.8%,	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanica ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○
	Theorem	Atov inty in potential benefits a	vacuone	wah is peopled	Hospitalization: No information





Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
<u>STU-2020-0707</u> <u>trial;</u> ⁶⁸ Jain et al; preprint; 2021	Patients with moderate to severe COVID-19 infection. 41 assigned to atovacuone 3000 mg a day for 10 days and 19 assigned to SOC	Mean age 50.9, male 63%, hypertension 63%, diabetes 63%, COPD 20%, asthma %, CHD 12%, CKD 33%, cancer 10%, obesity 38%	remdesivir 60%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
Auxora may not i	ncrease severe adverse (events. The effects of au	IXOra uxora on other importa eeded.	n outcomes are uncertai	n. Further research is
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
<u>CARDEA trial</u> ; ⁶⁹	Patients with severe	Mean age 60, male	Steroids 100%,	Low for mortality and	Mortality: RR 0.68



Bruen et al; Preprint; 2020	COVID-19 infection. 130 assigned to auxora initial dose 2.0 mg/kg (max 250 mg), followed by 1.6 mg/kg (max 200 mg) at 24 and 48 h and 131 assigned to SOC	67.4%, hypertension 62.8%, diabetes 41.8%	remdesivir 77.6%, tocilizumab 2.8%	mechanical ventilation; low for symptom resolution, infection and adverse events	(95%CI 0.39 to 1.17); RD -5.1% (95%CI - 9.8% to 2.7%); Low certainty $\oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: No information Symptom resolution or improvement: RR 1.07 (95%CI 0.94 to 1.22); RD 4.2% (95%CI -3.6% to 13.3%); Very low certainty $\oplus \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No information Adverse events: RR 0.69 (95%CI 0.48 to 1); RD -3.2% (95%CI -5.3% to 0%); Low certainty $\oplus \bigcirc \bigcirc$ Hospitalization: No information
	Uncerta	Avdo inty in potential benefits 2	ralimab and harms. Further resea	urch is needed	_
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
FORCE trial ; ⁷⁰ Carvelli et al ;	Patients with severe to critical COVID-19	Mean age 63.6, male 71%, hypertension 51%,	Corticosteroids 85%,	Low for mortality and mechanical ventilation;	Mortality: RR 1.68 (95%CI 0.87 to 3.26);





preprint ; 2021	infection. 103 assigned to avdoralimab 500 mg once followed by 200 mg every 48 hours and 104 assigned to SOC			low for symptom resolution, infection and adverse events	RD 10.9% (95%CI - 2.1% to 36.2%); Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: RR 1.15 (95%CI 0.85 to 1.55); RD 1.5% (95%CI -1.5% to 5.6%); Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
					Hospitalization: No information
	Uncertai	${f Av}$ in potential benefits a	ptadil nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
COVID-AIV trial ^{;71} Jihad et al; preprint (now retracted); 2021	Patients with severe to critical COVID-19 infection. 136 assigned to aviptadil three	Mean age 61 ± NR, male 69%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection,	Mortality: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical





	infusions of 50, 100 and 150 pmol/kg/hr and 67 assigned to SOC	Ayu inty in potential benefits a	ush-64	and adverse events Notes: Blinding and concealment probably inappropriate.	ventilation: No informationSymptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
Singh et al; ⁷² peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 37 assigned to Ayush-64 1500 mg a day for 30 days and 37 assigned to SOC	Mean age 35.89, male 62.1%, comorbidities 0%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very

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					Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: Very low certainty \oplus \bigcirc \bigcirc
AZD1656 may imp	prove time to symptom re	esolution. The effects of A		tant outcomes are uncerta	in. Further research is
Study; publication status	Patients and interventions analyzed	n Comorbidities	eeded. Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
ARCADIA trial; ⁷³ Chorlton et al; peer reviewed; 2022	Diabetic patients with moderate to severe COVID-19 infection. 80 assigned to AZD1656 200 mg a day for 21 days and 73 assigned to SOC	Mean age 64, male 63.4%, hypertension %, diabetes 100%,	Corticosteroids 73.2%, tocilizumab 3.9%, anakinra 0.7%, sarilumab 0.7%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanical ventilation: No information Symptom resolution or improvement: RR 1.18 (95%CI 0.9 to 1.62); RD 11% (95%CI -8.4% to 37.5%); Low certainty $\bigoplus \bigoplus \bigcirc \bigcirc$
					Symptomatic infection (prophylaxis studies): No



				information
				Adverse events: Very low certainty ⊕○○○
				Hospitalization: No information
cin probably does not re			does not improve time to sympt	tom resolution.
Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
-				
Patients with mild COVID-19 infection. 56 assigned to azelastine (inhaled) 0.02 to 0.1% twice a day for 11 days and 28 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: No information Invasive mechanica ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○
	Patients and interventions analyzed Patients with mild COVID-19 infection. 56 assigned to azelastine (inhaled) 0.02 to 0.1% twice a day for 11 days and 28	Patients and interventions analyzed Comorbidities Patients with mild COVID-19 infection. 56 assigned to azelastine (inhaled) 0.02 to 0.1% twice a day for 11 days and 28 NR	Patients and interventions analyzed Comorbidities Additional interventions Patients with mild COVID-19 infection. NR NR S6 assigned to azelastine (inhaled) NR NR 0.02 to 0.1% twice a day for 11 days and 28 Image: Comorbidities Image: Comorbidities	Patients and interventions analyzed Comorbidities Additional interventions Risk of bias and study limitations Patients with mild COVID-19 infection. 56 assigned to azelastine (inhaled) 0.02 to 0.1% twice a day for 11 days and 28 NR NR



Azithromyo	Azithromycin Azithromycin probably does not reduce mortality or mechanical ventilation and does not improve time to symptom resolution.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence				
RCT									
peer-reviewed; 2020 to severe C infection. 5 to azithron mg twice d	to severe COVID-19 infection. 56 assigned to azithromycin 500 mg twice daily and 55 assigned to standard of	Mean age 57.1 ± 15.73, male 45.9%	Hydroxychloroquine 100%, lopinavir- ritonavir 100%	High for mortality and invasive mechanical ventilation; High for symptom resolution, infection, and adverse events	Mortality: RR 1.01 (95%CI 0.92 to 1.1); RD 0.2% (95%CI - 1.3% to 1.6%); Moderate certainty ⊕⊕⊕⊖				
				Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Invasive mechanical ventilation: RR 0.92 (95%CI 0.77 to 1.1); RD -1.4% (95%CI -				
<u>Guvenmez et al</u> ; ⁷⁶ peer-reviewed; 2020	Patients with moderate COVID-19 infection. 12 assigned to lincomycin 600 mg twice a day for 5 days	Mean age 58.7 ± 16, male 70.8%,	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse	4% to 1.7%); Moderate certainty ⊕⊕⊕⊖ Symptom				
	and 12 assigned to azithromycin 500 mg on first day followed by 250 mg a day for 5 days			events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	resolution or improvement: RR 1.02 (95%CI 0.99 to 1.04); RD 1.2% (95%CI -0.6% to 2.4%); High certainty ⊕⊕⊕⊕				
	Patients with severe COVID-19. 214 assigned to azithromycin 500 mg once a day for 10 days and 183 assigned to	Median age 59.8 ± 19.5, male 66%, hypertension 60.7%, diabetes 38.2%, chronic lung disease 6%, asthma %, coronary heart disease 5.8%,	Corticosteroids 18.1%, lopinavir-ritonavir 1%, oseltamivir 46%, ATB 85%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events	Symptomatic infection (prophylaxis studies): No information				
	standard of care	chronic kidney disease 11%, cerebrovascular disease 3.8%, immunosuppression %,		Notes: Non-blinded study which might have introduced bias to symptoms and adverse	Adverse events: RR 1.23 (95%CI 0.51 to 2.96); RD 2.4%				



		cancer 3.5%, obesity %		events outcomes results.	(95%CI -5% to 19.9%); Very low
<u>RECOVERY trial</u> ⁷⁸ Horby et al; preprint; 2020	Patients with moderate to critical COVID-19. 2582 assigned to azithromycin 500 mg a day for 10 days and 5182 assigned to standard of care	male 62%, diabetes 27.5%, COPD 24.5%,	Corticosteroids 61%,	Low for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	certainty ⊕○○○ Hospitalization: RR 0.98 (95%CI 0.52 to 1.86); RD -0.1% (95%CI -2.3% to 4.1%); Low certainty ⊕⊕○○
<u>Rashad et al;</u> ⁷⁹ preprint ; 2020	Patients with mild to moderate COVID-19. 107 assigned to AZT 500 mg a day for 7 days, 99 assigned to Clarithromycin 1000 mg a day for 7 days and 99 assigned to SOC	Mean age 44.4 ± 18, male 29.8%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
PRINCIPLE trial; ⁸⁰ Butler et al; peer reviewed; 2021	Patients with mild to severe COVID-19 infection. 500 assigned to azithromycin 500 mg a day for 3 days and 629 assigned to SOC	Mean age 60.7 ± 7.8, male 43%, hypertension 42%, diabetes 18%, COPD 38%, asthma %, CHD 15%, cerebrovascular disease 6%,	NR	Some concerns for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Significant loss to follow-up.	
<u>ATOMIC2 trial</u> ; ⁸¹ Hinks et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 145 assigned	Mean age 45.9 ± 14.8, male 51.5%, hypertension 17.6%,	NR	Low for mortality and mechanical ventilation; high for symptom	





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	to azithromycin 500 mg a day for 14 days and 147 assigned to SOC	diabetes 8.5%, COPD 4.1%, asthma 18%, CHD 4.1%, cancer 0.3%,		resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
ACTION trial; ⁸² Oldenburg et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 131 assigned to azithromycin 1.2 g once and 70 assigned to SOC	Median age 43, male 44%, hypertension 12.2%, diabetes 3.8%, COPD 1.5%, asthma 12%, CKD 1%, cerebrovascular disease 1%, cancer 0.4%,	NR	Some concerns for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Significant loss to follow-up.	
<u>Ghanei et al</u> ; ⁸³ peer reviewed; 2021	Patients with severe COVID-19 infection. 110 assigned to lopinavir-ritonavir 200/50 mg twice a day for 7 days and 110 assigned to azithromycin 500 mg once followed by 250 mg a day for 5 days	Mean age 58.1 ± 16.3, male 51.5%, hypertension 24.7%, diabetes 12.2%, asthma 4.5%, CHD 8.9%, CKD 1.2%,	Convalescent plasma 1.8%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>trial</u> ; ⁸⁴ Gyselinck et	Patients with sevre to critical COVID-19 infection. 119 assigned to AZT 500 mg a day for 5 days and 64 assigned to SOC	61.8%, hypertension	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	

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Azvudine Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence			
RCT								
Ren et al; ⁸⁵ peer- reviewed; 2020	Patients with mild to moderate COVID-19 infection. 10 assigned to azvudine 5 mg once a day and 10 assigned to standard of care	Median age 52 ± 59, male 60%, hypertension 5%, diabetes 5%, coronary heart disease 5%	Antivirals 100%, antibiotics 40%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No			
	Uncertai	Bal inty in potential benefits a	OXAVIT and harms. Further re	esearch is needed.				
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence			



Lou et al; ⁸⁶ preprint; 2020	Patients with mild to severe COVID-19 infection. 10 assigned to baloxavir 80 mg a day on days 1, 4 and 7, 9 assigned to favipiravir and 10 assigned to standard of care	Mean age 52.5 ± 12.5, male 72.4%, hypertension 20.7%, diabetes 6.9%, coronary heart disease 13.8%	Antivirals 100%, interferon 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No informationAdverse events: No informationHospitalization: No information
Bamlanivimab may				al antibody) in if it affects mortality or	mechanical ventilation
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT		r			
<u>BLAZE-1 trial</u> ; ⁸⁷ Chen et al; peer- reviewed; 2020	Patients with mild to moderate COVID-19. 309 assigned to bamlanivimab 700 mg, 2800 mg, or 7000 mg once and 143 assigned to standard of care	Mean age 45 ± 68, male 55%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or

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ACTIV-3/TICO trial; ⁸⁸ Lundgren et al; Peer reviewed; 2020	Patients with moderate to severe COVID-19. 163 assigned to bamlanivimab 7000 mg once and 151 assigned to SOC	Median age 71 ± 22, male 66%, hypertension 49%, diabetes 29%, COPD %, asthma 9%, CHD 4%, CKD 11%, obesity 52%	Corticosteroids 49%, remdesivir 95%,	Low for mortality and adverse events; high for symptom resolution. Notes: Significant loss to follow-up for symptom improvement/resolution outcome.	improvement: RR 1.02 (95%CI 0.99 to 1.06); RD 1.2% (95%CI 3.6% to 5.4%); Moderate certainty ⊕⊕⊕⊖ Symptomatic
<u>Gottlieb et al</u> , ⁸⁹ Peer reviewed; 2020	Patients with mild to moderate COVID-19. 309 assigned to bamlanivimab 700- 7000 mg once, 112 assigned to bamlanivimab + etesevimab and 156 assigned to SOC	Mean age 44.7 ± 15.7, male 45.4%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	infection (prophylaxis studies): RR 0.56 (95%CI 0.39 to 0.81); RD -7.6% (95%CI - 10.6% to -3.6%); Moderate certainty ⊕⊕⊕○ Adverse events: RR
<u>BLAZE-2 trial</u> ; ⁹⁰ Cohen et al; peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 484 assigned to bamlanivimab 4200 mg once and 482 assigned to SOC	Median age 53	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	1.12 (95%CI 0.75 to 1.66); RD 1.2% (95%CI -2.5% to - 6.7%); Low certainty ⊕⊕○○ Hospitalization: RR
<u>BLAZE-1 trial</u> ; ⁹¹ Dougan et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 518 assigned to bamlanivimab + etesevimab 2800/2800 mg and 517 assigned to SOC	Mean age 53.8 ± 16.8, hypertension 33.9%, diabetes 27.5%, COPD %, CHD 7.4%, CKD 3.5%, immunosuppressive therapy 4.9%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	0.37 (95%CI 0.21 to 0.65); RD -3% (95%CI -3.8% to - 1.7%); Moderate certainty ⊕⊕⊕⊖
J2W-MC-PYAA trial; ⁹² Chen et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 18 assigned to bamlanivimab 700 to 7000 mg once and 6 assigned to SOC	Mean age 53.9, male 54.2%, hypertension 33.3%, diabetes 25%, asthma 25%, CHD 12.5%, CKD 4%, obesity 8.3%	Corticosteroids 29.1%, remdesivir 50%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
OPTIMISE-C19 trial; ⁹³ McCreary et al; peer reviewed; 2022	Patients with mild COVID-19 infection disease and risk factors for severity. 922	Mean age 56 ± 16, male 46%, hypertension 53%, diabetes 25%, COPD 19%, asthma %, CHD	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and	





	assigned to REGN- CoV2 (Regeneron) and 1013 assigned to bamlanivimab +/- etesevimab	18%, CKD 6.5%, immunosuppresive therapy 27%, obesity 48%		adverse events	
<u>ACTIV-2 trial</u> ; ⁹⁴ Chew et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 159 assigned to bamlanivimab 700 to 7000 mg and 158 assigned to SOC	Mean age 46.2 ± , male 48.9%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>OPTIMISE-C19</u> <u>trial;</u> ⁹⁵ Huang et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 2454 assigned to REGN- COV2 (Regeneron) one infusion and 1104 assigned to sotrovimab one infusion	Mean age 54 ± 18, male %, hypertension 30%, diabetes 12%, CHD 16%, CKD 4.7%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
MANTICO trial; ⁹⁶ Mazzaferri et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 107 assigned to sotrovomab 500 mg once and 106 assigned to bamlanivimab + etesevimab 700/1400 mg once and 106 assigned to REGEN- COV2 600/600 mg once	Mean age 65 ± 15, male 57.2%, diabetes 2.9%, COPD 16.7%, asthma %, CHD 37.9%, CKD 5.1%, immunosuppression19. 6%, obesity 25.4%	Vaccinated 28.6%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>BLAZE-4 trial;</u> ⁹⁷ Dougan et al; preprint; 2022	Patients with mild to moderate COVID-19 infection. 225 assigned to bebtelovimab 175 mg once and 175 assigned to bebtelovimab 175 mg + bamlanivimab 700 mg + etesevimab 1400 mg mg once	Median age 35 ± , male 44.5%	Vaccinated 20.7%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	



Baricitinib reduce	Baricitinib Baricitinib reduces mortality and probably reduces mechanical ventilation requirements and improves time to symptom resolution, without increasing severe adverse events.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence				
RCT									
<u>ACTT-2 trial</u> ; ⁹⁸ Kalil et al; peer- reviewed; 2020	Patients with moderate to severe COVID-19. 515 assigned to baricitinib + remdesivir 4 mg a day for 14 days + 200 mg once followed by 100 mg a day for 10 days and 518 assigned to remdesivir	Mean age 55.4 ± 15.7, male 63.1%, comorbidities 84.4%	Corticosteroids 11.9%	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Significant loss to follow-up.	Mortality: RR 0.73 (95%CI 0.57 to 0.92); RD -4.3% (95%CI - 6.9% to -1.3%); High certainty ⊕⊕⊕⊕ Invasive mechanical ventilation: RR 0.83 (95%CI 0.66 to 1.04); RD -2.9% (95%CI -				
COV-BARRIER trial; ⁹⁹ Marconi et al; peer reviewed; 2021	0	Mean age 57.6 ± 14.1, male 63.1%, hypertension 47.9%, diabetes 30%, COPD 4.6%, obesity 33%	Corticosteroids 79.3%, remdesivir 18.9%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	5.9% to 0.7%); Moderate certainty ⊕⊕⊕⊖ Symptom resolution or improvement: RR				
<u>COV-BARRIER-</u> <u>IMV trial</u> ; ¹⁰⁰ Wesley et al; preprint; 2021	Patients with critical COVID-19 infection. 51 assigned to baricitinib 4 mg a day for 14 days and 50 assigned to SOC	Mean age 58.6 ± 13.8, male 54.5%, hypertension 54.5%, diabetes 35.6%, COPD 3%, obesity 56.4%	Corticosteroids 86.1%, remdesivir 2%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	1.27 (95%CI 1.13 to 1.42); RD 16.4% (95%CI 7.9% to 25.5%); Moderate certainty ⊕⊕⊕⊖ Symptomatic				
RECOVERY trial; ¹⁰¹ Horby et al; preprint; 2021	Patients with severe to critical COVID-19 infection. 4148 assigned to baricitinib 4 mg a day for 10 days and 4008 assigned to SOC	Mean age 58.1± 15.5, male 66%, hypertension %, diabetes 23%, COPD 20.4%, asthma %, CHD 18.2%, CKD 2%,	Corticosteroids 95.2%, remdesivir 20.4%, tocilizumab 23%, Regeneron 11%; Vaccinated42%	Low for mortality and mechanical ventilation; some Concerns for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to	infection (prophylaxis studies): No information Adverse events: RR 0.78 (95%CI 0.64 to 0.95); RD -2.2% (95%CI -3.7% to - 0.5%); Moderate				



				symptoms and adverse events outcomes results.	certainty ⊕⊕⊕⊖ Hospitalization: No
ACTT-4 trial; ¹⁰² Wolfe et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 516 assigned to baricitinib 4 mg a day for 14 days and 494 assigned to dexamethasone 6 mg a day for 10 days	Mean age 58.3 ± 14, male 58%, hypertension 59.2%, diabetes 39.6%, COPD 9%, asthma 11%, CHD 9.6%, CKD 9.3%, immunosuppression 3.4%, cancer 5.6%, obesity 61.9%	Remdesivir 100%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	information
Karampitsakos et al; ¹⁰³ preprint; 2022	Patients with severe COVID-19 infection. 125 assigned to baricitinib 4 mg a day for 14 days and 126 assigned to TCZ 8 mg/kg once	Mean age 72.5, male 59.4%, hypertension 53.8%, cancer 9.2%, obesity 8%	Corticosteroids 100%, remdesivir 100%; Vaccinated 20.3%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
PanCOVID19 trial; ¹⁰⁴ Montejano et al; peer reviewed; 2022	Patients with severe COVID-19 infection. 145 assigned to baricitinib 2 to 4 mg a day for 14 days and 142 assigned to SOC	Median age 67, male 65.5%, hypertension 57.5%, diabetes 29.6%, obesity 18.8%	Corticosteroids 100%, remdesivir 15.3%, Vaccinated 91%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
	Uncerta	E inty in potential benefits a	BCG nd harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence



Padmanabhan et al; ¹⁰⁵ preprint; 2020	Patients with severe COVID-19. 30 assigned to BCG 0.1 ml once and 30 assigned to standard of care Uncerta	Mean age 45.2 ± 36.5, male 60%, obesity 23% Bebte inty in potential benefits a	Remdesivir 6.6%, elovimab	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
BLAZE-4 trial; ⁹⁷ Dougan et al; preprint; 2022	Patients with mild to moderate COVID-19 infection. 252 assigned to bebtelovimab 175 +/- bamlanivimab/etesevi mab mg once and 128 assigned to SOC	Median age 35 ± , male 44.5%	Vaccinated 20.7%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies):





					No information
					Adverse events: Very low certainty ⊕○○○ Hospitalization: Very
					low certainty ⊕○○○
	Uncertai	Beta inty in potential benefits a	glucans and harms. Further res	earch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
<u>Raghavan et al</u> ; ¹⁰⁶ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 16 assigned to beta glucans 3 to 13 gr a day and 8 assigned to SOC	Mean age 41.2	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	Mortality: No information Invasive mechanical ventilation: No information
				Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Symptom resolution or improvement: No information
<u>Pushkala et al</u> ; ¹⁰⁷ preprint; 2021	Patients with mild to moderate COVID-19 infection. 21 assigned to beta glucans 19 gr a day and assigned to	Mean age 44 ± , male 65%, hypertension 10%, diabetes 37.5%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events	Symptomatic infection (prophylaxis studies): No information
	SOC			Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
		Bicarbon	ate (inhaled)		
	Uncerta	inty in potential benefits a		earch is needed.	
Study; publication	Patients and	Comorbidities	Additional	Risk of bias and study	Interventions effects vs





status	interventions analyzed		interventions	limitations	standard of care (standard of care) and GRADE certainty of the evidence
RCT					
Delic et al, ¹⁰⁸ peer reviewed; 2022	Patients with critical COVID-19 infection. 42 assigned to bicarbonate (inhaled) twice a day and 52 assigned to SOC	Mean age 66, male 79.8%, hypertension 57.4%, diabetes 33%, CHD 5.3%, cerebrovascular disease 5.3%	Corticosteroids 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncertai	B inty in potential benefits a	ioven and harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
<u>Rybakov et al</u> ; ¹⁰⁹ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 32 assigned to bioven 0.8-1 g/kg once a day for 2 days and 34 assigned to SOC	NA	NA	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution



		Boswel	lia extract	allocation is probably inappropriate.	or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncertai	inty in potential benefits a		rch is needed.	
	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
Barzin Tond et al; ¹¹⁰ peer reviewed; 2021		Mean age 53.8, male 52%, hypertension 22%, diabetes 28%, COPD 2%, asthma 2%, CHD 2%, obesity 24%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information

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Bromhexine may re	Bromhexine hydrochloride Bromhexine may reduce symptomatic infections in exposed individuals. Its effects on other clinical important outcomes are uncertain. Further research is needed.							
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence			
RCT								
<u>Li T et al</u> ; ¹¹¹ peer- reviewed; 2020	Patients with severe to critical COVID-19. 12 assigned to bromhexine hydrochloride 32 mf three times a day for 14 days and 6 assigned to standard of care	Median age 52 ± 15.5, male 77.8%, hypertension 33.3%, diabetes 11.1%	Corticosteroids 22.2%, interferon 77.7%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: Very			
<u>Ansarin et al</u> ; ¹¹² peer-reviewed; 2020	Patients with mild to critical COVID-19. 39 assigned to bromhexine 8 mg three time a day for 14 days and 39 assigned to standard of care	Mean age 59.7 ± 14.9, male 55.1%, hypertension 50%, diabetes 33.3%	Hydroxychloroquine 100%	High for mortality and invasive mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): RR 0.38 (95%CI 0.13 to 1.09); RD -10.8% (95%CI - 15.1% to 1.6%); Low certainty $\oplus \oplus \bigcirc \bigcirc$			
<u>Mikhaylov et al</u> ; ¹¹³ Peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 25 assigned to bromhexine 12 mg a day and 25 assigned to SOC	Mean age 40.6 ± 7.6, male 42%, comorbidity 6%	NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to	Adverse events: Very low certainty ⊕○○○ Hospitalization: No information			



<u>Tolouian et al</u> ; ¹¹⁴ Peer reviewed; 2021	to critical COVID-19 infection. 48 assigned to bromhexine 32 mg a	Mean age 52 ± 16, male 46%, hypertension 39%, diabetes 33%, COPD 7%, asthma 6%, CHD 9%, CKD 5%, cerebrovascular disease 2%, cancer 6%	Lopinavir-ritonavir 100%, interferon 100%	symptoms and adverse events outcomes results. Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse	
<u>Tolouian et al</u> ; ¹¹⁵ preprint; 2021	Individuals exposed to SARS-CoV-2 infection. 187 assigned to bromhexine 24 mg a day for 14 days and 185 assigned to SOC	53.2%, hypertension 6.2%, diabetes 9.1%,	NR	events outcomes results. Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
	Uncertai	Cal inty in potential benefits a	citriol nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT	l				
<u>Elamir et al</u> ; ¹¹⁶ peer reviewed; 2022	Patients with moderate COVID-19 infection. 25 assigned to calcitriol 0.5 μg daily for 14 days and 25 assigned to SOC	Mean age 66.5, male 30%, hypertension 60%, diabetes 40%, COPD 16%, cancer 4%, obesity 20%	Corticosteroids 50%, remdesivir 52%, convalescent plasma 12%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No





	Camostat mesi	Camost late may not increase sym	at mesilate	r research is needed.	information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
CamoCO-19 trial; ¹¹⁷ Gunst et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 137 assigned to camostat mesilate 200 mg a day for 5 days and 68 assigned to SOC	male 60%, hypertension 34%, diabetes 17%, COPD 10%, asthma 13%, CHD 19%, cancer	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$
<u>Chupp et al</u> ; ¹¹⁸ preprint; 2021	Patients with mild COVID-19 infection. 35 assigned to camostat mesilate 800 mg a day for 7 days and 35 assigned to SOC	Mean age 44.1 ± 13.3, male 60%, hypertension 20%, diabetes 5.7%, CKD 2.9%, obesity 68.6%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Symptom resolution or improvement: RR 1.02 (95%CI 0.94 to 1.11); RD 1.2% (95%CI -3.6% to 6.6%); Low certainty
<u>CANDLE trial</u> ; ¹¹⁹ Kinoshita et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 78 assigned	Mean age 55.9 ± 18.4, male 50.3%, hypertension 28.4%,	NR	Low for mortality and mechanical ventilation; low for symptom	⊕⊕○○ Symptomatic

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<u>Terada et al</u> ; ¹²⁰ peer reviewed; 2022	to camostat mesilate 2400 mg a day for 14 days and 77 assigned to SOC Patients with mild to severe COVID-19 infection. 56 assigned to camostat 600 mg +	diabetes 17.4%, COPD 16.1%, asthma %, CHD 5.2%, CKD 5.8%, obesity 9.7% Mean age 58.3, male 64.9%, diabetes 24.8%, COPD 9.4%, CHD 2.6%	NR	resolution, infection and adverse events High for mortality and mechanical ventilation; high for symptom resolution, infection and	infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○
	ciclesonide (inhaled) 1200 µg a day and 61 assigned to SOC			adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Hospitalization: Very low certainty ⊕○○○
<u>Tobback et al</u> ; ¹²¹ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 61 assigned to camostat mesilate 300 mg a day for 5 days and 29 assigned to SOC	Median age 40, male 45.6%, diabetes 1.1%, cancer 6.7%, obesity 6.7%	Vaccinated 7.8%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
	Uncertai	Cana inty in potential benefits a	kinumab and harms. Further resea	nrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
CAN-COVID trial; ¹²² Cariccchio et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 223 assigned to canakinumab 450– 750 mg/kg once and 223 assigned to SOC	Median age 59, male 58.8%, hypertension 55.7%, diabetes 36.1%, COPD 7.3%, asthma 7.7%, CHD 20.3%, CKD 8.8%, cerebrovascular disease 5.9%	Corticosteroids 36.3%, remdesivir 20.7%, hydroxychloroquine 13.2%, azithromycin 37.4%, convalescent plasma 3.5%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$
Three C trial; ¹²³	Patients with moderate	Mean age 68.8 ± 13.2,	Steroids 46.7%,	Low for mortality and	Symptom resolution or



Cremer et al; peer reviewed; 2021	to severe COVID-19 infection. 29 assigned to canakinumab 300 to 600 mg once and 16 assigned to SOC	male 73.3%, hypertension 71.1%, diabetes 46.7%, COPD 17.8% CHD 22.2%, CKD 33.3%, cerebrovascular disease 4.4%	remdesivir 46.7%, convalescent plasma 9%	mechanical ventilation; low for symptom resolution, infection, and adverse events	<pre>improvement: No information Symptomatic infection (prophylaxis studies): Very low certainty ⊕○○○ Adverse events: Very low certainty ⊕○○○ Hospitalization: No information</pre>
	Uncertai	Cani inty in potential benefits a	nabidiol and harms. Further resea	rrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT			1		
	Patients with mild to moderate COVID-19 infection. 49 assigned to cannabidiol 300 mg a day for 14 days and 42 assigned to SOC	Mean age 39.7, male 32.7%, hypertension 4.4%, diabetes 2.2%, COPD %, asthma 3.3%, cancer 1.1%, obesity 6.6%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty \bigoplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \bigoplus \bigcirc \bigcirc Symptom resolution or improvement: Very low certainty \bigoplus \bigcirc \bigcirc Symptomatic infection (prophylaxis studies): No



		luble CD24 appe human immu	unoglobulin G1)		information Adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Hospitalization: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
	critical COVID-19 infection. 116 assigned to CD24Fc 480 mg	Mean age 57.8 ± 14, male 74.8%, hypertension 54.7%, diabetes 21.4%, COPD 1.7%, asthma 9.4%, obesity 15.4%	Corticosteroids 83.3%, remdesivir 68.4%, hydroxychloroquine 1.3%, convalescent plasma 54.3%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: RR 0.57 (95%CI 0.34 to 0.96); RD -7.4% (95%CI - 11.4% to -0.7%); Low certainty $\bigoplus \bigcirc \bigcirc$ Symptom resolution or
					improvement: RR 1.18 (95%CI 1 to 1.39); RD 10.7% (95%CI -0.2% to 23.4%); Low certainty ⊕⊕○○ Symptomatic infection (prophylaxis studies): No

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					information Adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Hospitalization: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
	Uncertai	CERC-002 (mo inty in potential benefits :	noclonal antibo	dy) arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
Perlin et al; ¹²⁶ preprint; 2021	Patients with mild to moderate COVID-19 infection. 31 assigned to CERC-002 16 mg/kg once and 31 assigned to SOC	Mean age 58.5 ± 14, male 69.5%	Corticosteroids 91.5%, remdesivir 68.2%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate. Significant loss to follow-up.	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: No information

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Chloroquine nasal drops Uncertainty in potential benefits and harms. Further research is needed.								
Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence				
Patients with mild COVID-19. 30 assigned to chloroquine nasal drops 0.03% six times a day for 10 days and 30 assigned to SOC	Mean age 34.9 ± 10.35, male 78.3%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information				
Uncerta			arch is needed.					
Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence				
	Patients and interventions analyzed Patients with mild COVID-19. 30 assigned to chloroquine nasal drops 0.03% six times a day for 10 days and 30 assigned to SOC Signed to SOC Uncertain Patients and interventions	Uncertainty in potential benefits a Patients and interventions analyzed Comorbidities Patients with mild COVID-19. 30 assigned to chloroquine nasal drops 0.03% six times a day for 10 days and 30 assigned to SOC Mean age 34.9 ± 10.35, male 78.3% Value Mean age 34.9 ± 10.35, male 78.3% Comorbidities Patients with mild COVID-19. 30 Mean age 34.9 ± 10.35, male 78.3% Comorbidities Image: Social data and the second data and t	Patients and interventions Patients and interventions analyzed Comorbidities Additional interventions Patients with mild COVID-19, 30 assigned to chloroquine nasal drops 0.03% six times a day for 10 days and 30 assigned to SOC Mean age 34.9 ± 10.35, male 78.3% NR Signed to SOC Mean age 34.9 ± 10.35, male 78.3% NR COVID-19, 30 assigned to solve a days of 10 days and 30 assigned to SOC Solve a days of 10 days and 30 assigned to SOC NR CIGB-325 Uncertainty in potential benefits and harms. Further reserver Comorbidities Additional interventions	Uncertainty in potential benefits and barms. Further research is needed. Patients and interventions analyzed Comorbidities Additional interventions Risk of bias and study limitations Patients with mild COVID-19, 30 assigned to chloroquine nasal drops 0.03% six times a day for 10 days and 30 assigned to SOC Mean age 34.9 ± 10.35, assigned to SOC NR High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate. Notes: Non-blinded study. Concealment of allocation is probably inappropriate. CIGB-325 Uncertainty in potential benefits and harms. Further research is needed. Patients and interventions Comorbidities Additional interventions Risk of bias and study limitations				



ATENEA-Co-300 trial; ¹²⁸ Cruz et al; preprint; 2020	Patients with mild to moderate COVID-19. 10 assigned to CIGB- 325 2.5 mg/kg/day during 5-consecutive days) and 10 assigned to standard of care	Mean age 45.3 ± 12, male 70%, hypertension 25%, diabetes 0%, cancer 5%, obesity 25%	Hydroxychloroquine 100%, lopinavir- ritonavir 100%, IFN 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: Very low certainty $\oplus \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty $\oplus \bigcirc \bigcirc$ Hospitalization: No information
	Uncertai	Clarit inty in potential benefits a	hromycin nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
<u>Rashad et al</u> ; ⁷⁹ preprint; 2020	Patients with mild to moderate COVID-19. 107 assigned to AZT 500 mg a day for 7 days, 99 assigned to clarithromycin 1000 mg a day for 7 days and 99 assigned to SOC	Mean age 44.4 ± 18, male 29.8%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or

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				inappropriate.	<pre>improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information</pre>
	Uncerta	Claza	kizumab and harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT					
Lonze et al; ¹²⁹ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 78 assigned to clazakizumab 12.5 to 25 mg a day and 74 assigned to SOC	Mean age 61.8 ± 12.2, male 70.4%, hypertension 63.2%, diabetes 42.4%, COPD 16.4%, asthma %, CHD 34.2%, immunosuppresive therapy 7.2%, cancer 8.6%, obesity 11.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: No information Invasive mechanical ventilation: RR 0.66 (95%CI 0.43 to 1.01); RD -7.6% (95%CI - 9.8% to 1.7%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Symptom resolution or improvement: RR 1.23 (95%CI 0.87 to 1.76); RD 13.9% (95%CI -7.9% to 46%); Low certainty $\oplus \oplus \bigcirc \bigcirc$

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					Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncertai	Cle inty in potential benefits a	vudine Ind harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT BK-CLV-201 trial; ¹³⁰ Song et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 41 assigned to clevudine 120 mg a day for 14 days and 20 assigned to SOC	Mean age 59.9 ± 12.8, male 49.2%, hypertension 45.9%, diabetes 26.2%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or information Symptomatic information Symptomatic information Adverse events: Very low certainty ⊕ ○ ○ ○



					Hospitalization: No information
		carnitine, N-ace nty in potential benefits a		otinamide, serine) arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT	L		ł		
COVID-19-MCS trial; ¹³¹ Altay et al; preprint; 2020	Patients with mild to moderate COVID-19. 71 assigned to cofactors (L-carnitine, N-acetylcysteine, nicotinamide, serine) and 22 assigned to	Mean age 35.6 ± 47, male 60%	Hydroxychloroquine 100%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events	Mortality: No information Invasive mechanical ventilation: No information
	standard of care		assessors not l	Notes: Outcome assessors not blinded. Possible reporting bias.	Symptom resolution or improvement: Very
COVID-19-MCS t <u>rial</u> , ¹³² Altay et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 229 assigned to cofactors (L- carnitine, N- acetylcysteine, nicotinamide, serine) and 75 assigned to SOC	Mean age 36.3 , male 57.6%, hypertension 9.2%, diabetes 6.2%	Hydroxychloroquine 81.9%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events:
<u>Hu et al;</u> ¹³³ preprint; 2021	Patients with moderate to severe with diabetes COVID-19 infection. 12 assigned to nicotinamide 500 mg a day and 12 assigned to SOC	Mean age 69.5, male 45.8%, hypertension 33.3%, diabetes 16.6%, COPD 0%, CHD 8.3%, CKD 4.2%, cerebrovascular disease 8.3%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Very low certainty ⊕○○○ Hospitalization: No information



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care (standard of care) and GRADE certainty of the evidence
RCT				·	
<u>GRECCO-19</u> <u>tria</u> l; ¹³⁴ Deftereos et al; peer-reviewed; 2020	Patients with severe COVID-19 infection. 50 assigned to colchicine 1.5 mg once followed by 0.5 mg twice daily until hospital discharge or 21 days and 55 assigned to standard of	Median age 64 ± 11, male 58.1%, hypertension 45%, diabetes 20%, chronic lung disease 4.8%, coronary heart disease 13.3%, immunosuppression 3.75%	Hydroxychloroquine 98%, lopinavir- ritonavir 31.4%, tocilizumab 3.8%, azithromycin 92%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have	Mortality: RR 0.99 (95%CI 0.92 to 1.05) RD -0.2% (95%CI - 1.3% to 0.8%); Moderate certainty ⊕⊕⊕⊖ Invasive mechanica ventilation: RR 0.9
	care			introduced bias to symptoms and adverse events outcomes results.	(95%CI 0.89 to 1.02 RD -0.3% (95%CI - 1.9% to 1.4%); Moderate certainty
<u>Lopes et al</u> ; ¹³⁵ preprint; 2020	Patients with moderate to severe COVID-19 infection. 19 assigned to colchicine 0.5 mg three times a day, for 5 days followed by 0.5 mg twice daily for 5 days and 19 assigned to	Median age 50.75 ± 26.2, male 40%, diabetes 31.4%, chronic lung disease 14.2%, coronary heart disease 40%	Corticosteroids 40%, hydroxychloroquine 100%, azithromycin 100%, heparin 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded	 ⊕⊕⊕○ Symptom resolution or improvement: RR (95%CI 0.98 to 1.02 RD 0% (95%CI -1.2)
	standard of care			study. Concealment of allocation is probably inappropriate.	to 1.2%); High certainty ⊕⊕⊕⊕ Symptomatic
<u>Salehzadeh et al</u> ; ¹³⁶ preprint; 2020	to critical COVID-19. 50 assigned to colchicine 1 mg a day for 6 days and 50	Mean age 56, male 41%, hypertension 11%, diabetes 11%, chronic lung disease 4%, coronary heart disease	Hydroxychloroquine 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse	infection (prophylaxis studies): No information Adverse events: RI
	assigned to standard of care	15%, chronic kidney disease 5%		events Notes: Non-blinded study. Concealment of	0.78 (95%CI 0.61 tc 0.99); RD -2.2% (95%CI -4% to -0.19





				allocation is probably inappropriate.	High certainty ⊕⊕⊕⊕
<u>Tardif et al</u> , ¹³⁷ peer- reviewed; 2020	Patients recently diagnosed mild COVID-19 and risk factors for severe disease. 2235 assigned to colchicine 1 mg a day for 3 days followed by 0.5 mg for a total of 27 days and 2253 assigned to SOC	Mean age 54.3, male 46%, hypertension 36.3%, diabetes 19.9%, COPD 26.5%, CHD 5.4%, obesity 45.7%	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	Pulmonary embolism: RR 5.55 (95%CI 1.23 to 25); RD 0.4% (95%CI 0.02% to 2.2%); Low certainty ⊕⊕○○ Hospitalization: RR 0.81 (95%CI 0.63 to 1.04); RD -0.9%
<u>RECOVERY -</u> <u>Colchicine trial;</u> ¹³⁸ Horby et al; peer reviewed; 2021	Patients with moderate to critical COVID-19 infection. 5610 assigned to colchicine 500 mg twice a day for 10 days and 5730 assigned to SOC	Mean age 63.4 ± 13.8, male 69.5%, diabetes 25.5%, COPD 21.5%, asthma %, CHD 21%, CKD 3%	Corticosteroids 94%	Low for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	to 1.04); RD -0.9% (95%CI -1.8% to 0.2%); Low certainty ⊕⊕⊖⊖
COL-COVID <u>trial</u> ; ¹³⁹ Figal et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 52 assigned to colchicine 1.5 gr once followed by 1 gr a day for 7 days and 51 assigned to SOC	Mean age 51 ± 12, male 52.4%, hypertension 27.2%, diabetes 14.6%, COPD 1%, CHD 2.9%, CKD 6.8%, cerebrovascular disease 1.9%, immunosuppresive therapy %, cancer %, obesity 21.4%	Corticosteroids 74.8%, remdesivir 32%, lopinavir-ritonavir 1%, tocilizumab 9.7%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
PRINCIPLE - Colchicine trial; ¹⁴⁰ Dorward et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 156 assigned to colchicine 500 μg a day for 14 days and 133 assigned to SOC	Mean age 61, male 50%, hypertension 19.5%, diabetes 10.9%, COPD or asthma 32.2%, CHD 8%, cerebrovascular disease, or other neurological diseases	NR	Low for mortality and mechanical ventilation; high for symptom resolution, hospitalization, and adverse events	

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		5.2%		Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>COLCOVID</u> <u>trial;¹⁴¹ Diaz et al;</u> peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 640 assigned to colchicine 1.5 mg once followed by 1 mg a day for 14 days and 639 assigned to SOC	Mean age 62 ± 14, male 64.9%, hypertension 47.7%, diabetes 22.7%, COPD 9.6%, CHD 7.1%, CKD 2.3%, cerebrovascular disease 2%, cancer 2.3%	Corticosteroids 91.5%, hydroxychloroquine 0.3%, lopinavir- ritonavir 0.2%, convalescent plasma 7.3%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>Alsultan et al</u> ; ¹⁴² peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 14 assigned to colchicine 1.5 mg once followed by 1 mg a day for 5 days and 21 assigned to SOC	Age 60 to 80 65.3, male 38.8%, diabetes 53.1%, CKD 8.2%, cerebrovascular disease 4.1%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
Pourdowlat et al; ¹⁴³ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 89 assigned to colchicine 0.5 mg for 3 days and then continued 1 mg/day for 12 days and 63 assigned to SOC	Mean age 55, male 56.4%, hypertension 12.7%, diabetes 14.5%, COPD %, asthma 3.6%, CHD 5.4%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>Gorial et al</u> ; ¹⁴⁴ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 80 assigned to colchicine 1 mg a day for 7 days followed by 0.5 mg a day for 14 days and 80 assigned to SOC	53.1%, hypertension 41.2%, diabetes 20.6%, COPD %, asthma 1.2%, cancer 2.5%, obesity 35%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably	



				inappropriate.
Mostafaie et al; <u>NCT04392141,</u> other; 2021	Patients with moderate to severe COVID-19 infection. 60 assigned to colchicine and 60 assigned to SOC	Mean age 53.5 ± 15.1, male 54.2%, hypertension 26.7%, diabetes 7.5%, cancer 5.8%,	NR	NA
<u>STRUCK trial</u> ; ¹⁴⁵ Pimenta Bonifácio et al; preprint; 2021	Patients with severe to critical COVID-19 infection. 14 assigned to colchicine 1 mg a day for 4 weeks and 16 assigned to SOC	Mean age 48.9 ± 12.2, male 61.7%, hypertension 45%, diabetes 21.7%, COPD 6.7%, CHD 5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>Cecconi et al</u> ; ¹⁴⁶ peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 119 assigned to colchicine 1 mg once followed by 0.5 mg a day for 5 days and 120 assigned to SOC	Mean age 65.1 ± 16, male 59%, hypertension 40%, diabetes 16%, COPD 4%, asthma 5%, CHD 7%	Corticosteroids 98%, remdesivir 15.5%, hydroxychloroquine 0%, lopinavir-ritonavir 0.8%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>Rabbani et al</u> ; ¹⁴⁷ peer reviewed; 2022	Patients with moderate to severe with cardiac injury COVID-19 infection. 48 assigned to colchicine 1.2 mg a day for 30 days and 45 assigned to SOC	Mean age 71, male 67.7%, hypertension 78.5%, diabetes 26.9%, COPD 10.8%, CKD 28%,	Corticosteroids 62.4%, remdesivir 69.9%, hydroxychloroquine 1.1%, convalescent plasma 14%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.

Uncertainty in potential benefits and harms. Further research is needed.



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Gaitan-Duarte et al; ¹⁴⁸ preprint; 2021	Patients with moderate to severe COVID-19 infection. 153 assigned to colchicine + rosuvastatin 1 mg + 40 mg a day for 14 days and 161 assigned to SOC	male 68%, hypertension	Corticosteroids 98%,	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: No information
		ality or mechanical ventils		nprove time to symptom re ncrease severe adverse eve	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence



<u>Li et al</u> ; ¹⁴⁹ peer- reviewed; 2020	Patients with moderate to critical COVID-19 infection. 52 assigned to convalescent plasma 4 to 13 mL/kg of recipient body weight and 51 assigned to standard of care	Median age 70 ± 8, male 58.3%, hypertension 54.3%, diabetes 10.6%, coronary heart disease 25%, chronic kidney disease 5.8%, cerebrovascular disease 17.45%, cancer 2.9%, liver disease 10.7%	Corticosteroids 39.2%, antivirals 89.3%, ATB 81%, IFN 20.2%, IVIG 25.4%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: RR 0.98 (95%CI 0.93 to 1.03); RD -0.3% (95%CI - 1.1% to 0.5%); High certainty ⊕⊕⊕ Invasive mechanical ventilation: RR 1.03 (95% CI 0.94 to 1.11); RD 0.5%	
CONCOVID trial; Gharbharan et al; ¹⁵⁰ preprint; 2020	Patients with moderate to critical COVID-19 infection. 43 assigned to convalescent plasma 300 ml once or twice and 43 assigned to standard of care	Median age 62 ± 18, male 72%, hypertension 26%, diabetes 24.4%, chronic lung disease 26.7%, coronary heart disease 23.2%, chronic kidney disease 8.1%, immunosuppression 12.8%, cancer 9.3%	NR	Low for mortality and invasive mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	1.11); RD 0.5% (95%CI -1% to 1.9%); High certainty ⊕⊕⊕⊕ Symptom resolution or improvement: RR 0.99 (95% CI 0.95 to 1.02); RD -0.6% (95%CI -3% to 1.2%); High certainty ⊕⊕⊕⊕	
<u>Avendaño-Solá</u> et al; ¹⁵¹ preprint; 2020	Patients with severe COVID-19. 38 assigned to convalescent plasma 250-300 ml once and 43 assigned to standard of care	Mean age 60.8 ± 15.5, male 54.3%, hypertension 39.5%, diabetes 20.9%, chronic lung disease 12.3%, asthma NR%, coronary heart disease 18.5%, chronic kidney disease 4.9%	Corticosteroids 56.8%, remdesivir 4.94%, hydroxychloroquine 86.4%, lopinavir- ritonavir 41.9%, tocilizumab 28.4%, azithromycin 61.7%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Symptomatic infection (prophylaxis studies): Very low certainty ⊕○○○ Adverse events: RR 1.05 (95% CI 0.9 to 1.22); RD 0.5% (95% CI -1% to 2.2%); Low certainty ⊕⊕○○	
<u>PLACID trial</u> ; ¹⁵² Agarwal et al; preprint; 2020	Patients with severe COVID-19. 235 assigned to convalescent plasma 200 ml twice in 24 h and 229 assigned to standard of care	Median age 52 ± 18, male 76.3%, hypertension 37.3%, diabetes 43.1%, chronic lung disease 3.2%, coronary heart disease 6.9%, chronic kidney	Corticosteroids 64.4%, remdesivir 4.3%, hydroxychloroquine 67.7%, lopinavir- ritonavir 14.2%, tocilizumab 9%, azithromycin 63.8%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events	Hospitalization: RR 0.77 (95% CI 0.57 to 1.03); RD -1.1% (95%CI -2.1% to 0.1%); Moderate certainty ⊕⊕⊕⊖	





		disease 3.7%, cerebrovascular disease 0.9%, cancer 0.2%, obesity 7.1%		Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>PLASM-AR trial</u> ; ¹⁵³ Simonovich et al; peer-reviewed; 2020	Patients with severe to critical COVID-19. 228 assigned to convalescent plasma and 105 assigned to standard of care	Mean age 62 ± 20, male 67.6%, hypertension 47.7%, diabetes 18.3%, COPD 7.5%, asthma 4.2%, coronary heart disease 3.3%, chronic kidney disease 4.2%	Corticosteroids 93.3%, hydroxychloroquine 0.3%, lopinavir- ritonavir 3%, tocilizumab 4.2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
ILBS-COVID-02 <u>trial</u> ; ¹⁵⁴ Bajpai et al; preprint; 2020	Patients with severe to critical COVID-19. 14 assigned to convalescent plasma 500 ml twice and 15 assigned to standard of care	0	Hydroxychloroquine 100%, azithromycin 100%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>AlQahtani et al</u> ; ¹⁵⁵ preprint; 2020	Patients with severe to critical COVID-19. 20 assigned to convalescent plasma 200 ml twice and 20 assigned to standard of care	Mean age 51.6 ± 13.7, male 80%, hypertension 25%, diabetes 30%, COPD 7.5%, asthma %, coronary heart disease 10%, chronic kidney disease 5%	Corticosteroids 12.5%, hydroxychloroquine 92.5%, lopinavir- ritonavir 85%, tocilizumab 30%, azithromycin 87.5%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Fundacion</u> <u>INFANT-Plasma</u> <u>tria</u> l; ¹⁵⁶ Libster et al; preprint; 2020	Patients with mild to moderate COVID-19. 80 assigned to convalescent plasma 250 ml and 80 assigned to standard of care	Mean age 77.1 ± 8.6, male 47.5%, hypertension 71.2%, diabetes 22.5%, COPD 4.4%, asthma 3.8%, coronary heart disease 13.1%, chronic kidney disease 2.5%, cancer	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	





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		3.8%, obesity 7.5%		
<u>PICP19 trial;</u> ¹⁵⁷ Ray et al; peer reviewed; 2020	Patients with severe COVID-19. 40 assigned to convalescent plasma 200 ml and 40 assigned to standard of care	Mean age 61 ± 11.5, male 71.2%, hypertension 43.7%, diabetes 58.7%, COPD 6.2%, CHD 10%, cerebrovascular disease 2.5%	Steroids 50%, remdesivir 31.2%, hydroxychloroquine 37.5%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>RECOVERY-</u> <u>Plasma trial</u> ; ¹⁵⁸ Horby et al; Other; 2020	Patients with severe to critical COVID-19 infection. 5795 assigned to CP 275 ml a day for two days and 5763 assigned to SOC	Median age 63.5 ± 14.7, male 64.2%, diabetes 26%, COPD 24%, CHD 22%	Corticosteroids <1%, lopinavir-ritonavir <1%, azithromycin 10%, colchicine 14%	Low for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to
				symptoms and adverse events outcomes results.
<u>Baklaushev et al</u> ; ¹⁵⁹ peer reviewed; 2020	Patients with moderate to severe COVID-19. 46 assigned to CP 640 ml divided in two infusions and 20 assigned to SOC	Age 56.3 ± 11, male 60.6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events
				Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>O'Donnell et al</u> ; ¹⁶⁰ Peer-reviewed; 2021	Patients with severe to critical COVID-19 infection. 150 assigned to CP one infusion and 73 assigned to SOC	Median age 61 ± 23, male 65.9%, hypertension 33.6%, diabetes 36.8%, COPD 9%, CHD 37.7%, CKD 9.4%, obesity 48.8%	Corticosteroids 81%, remdesivir 6%, hydroxychloroquine 6%	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events





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				Notes: Sensitivity analysis including loss to follow-up patients significantly modified results. At the time mortality was measured the number of patients on IMV was significantly higher in the intervention arm.	
Beltran Gonzalez et al; ¹⁶¹ preprint; 2021	Patients with severe to critical COVID-19 infection. 130 assigned to CP 200 ml a day for 2 days and 60 assigned to IVIG	Mean age 58 ± 25, male 62.6%, hypertension 35.2%, diabetes 34.7%, COPD 4.7%, CHD 3.1%, CKD 3.1%, cerebrovascular disease 1.05%, cancer 0.53%, obesity 41.5%	Corticosteroids 82.6%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
Pouladzadeh et al; ¹⁶² peer reviewed; 2021	Patients with severe COVID-19 infection. 30 assigned to CP 500 ml once or twice and 30 assigned to SOC	Mean age 55.3 ± 13.6, male 55%, comorbidities 50%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>SBU-COVID19 -</u> <u>Convalescent</u> <u>Plasma trial</u> ; ¹⁶³ Bennett-Guerrero et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 59 assigned to CP 480 ml once and 15 assigned to SOC	Mean age 65.5 ± 16.6, male 59.5%, hypertension 68.9%, diabetes 33.7%, COPD 12.1%, CHD 17.6%, CKD 9.5%, cerebrovascular disease 14.8%, immunosuppressive therapy 8.1%	Corticosteroids 60.8%, remdesivir 24.3%, hydroxychloroquine 31%, tocilizumab 21.6%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	

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<u>Salman et al</u> ; ¹⁶⁴ peer reviewed; 2021	Patients with severe COVID-19 infection. 15 assigned to CP 250 ml once and 15 assigned to SOC	Median age 57 ± 10, male 70%, diabetes 30%, asthma 16.6%, cerebrovascular disease 43.3%	Corticosteroids 76.6%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>CAPSID trial</u> ; ¹⁶⁵ Koerper et al; preprint; 2021	Patients with severe to critical COVID-19 infection. 53 assigned to CP 850 ml in three infusions and 52 assigned to SOC	Mean age 60 ± 13, male 73.3%, hypertension 56.2%, diabetes 31.4%, COPD 16.2%, CHD 21.9%, cancer 4.7%, obesity 54.2%	Corticosteroids 89.5%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>REMAP-CAP</u> <u>trial</u> ; ¹⁶⁶ Green et al; 2021	Patients with moderate to critical COVID-19 infection. 1075 assigned to CP 550- 700 ml and 904 assigned to SOC	Mean age 62 ± 12.9, male 67.6%, diabetes 30.9%, COPD 23.2%, asthma 19.4%, CHD 8.1%, CKD 10.4%, immunosuppressive therapy 6.4%, cancer 1.4%	Corticosteroids 93.4%, remdesivir 45.1%, tocilizumab 2%	Low for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>CONCOR-1</u> <u>trial</u> ; ¹⁶⁷ Bégin et al; preprint; 2021	Patients with severe COVID-19 infection. 614 assigned to CP 500 ml and 307 assigned to SOC	Mean age 67.5 ± 15.6, male 59.1%, diabetes 35%, COPD 24.1%, CHD 62%	Corticosteroids 80.4%, azithromycin 44.3%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>PLACOVID</u> <u>trial</u> ; ¹⁶⁸ Sekine et al;	Patients with severe to critical COVID-19	Median age 60.5 ± 20, male 58.1%,	Corticosteroids 98.8%	Low for mortality and mechanical ventilation;





peer reviewed; 2021	infection. 80 assigned to CP 300 ml twice and 80 assigned to SOC	hypertension 61.3%, diabetes 39.4%, COPD 13.8%, CHD 21.9%, obesity 56.9%		high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>COVIDIT trial</u> ; ¹⁶⁹ Kirenga et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 69 assigned to CP 150 -300 ml twice and 67 assigned to SOC	Mean age 50 ± 23.5, male 71.3%, hypertension 36%, diabetes 32%, asthma 3.7%, obesity 33.3%	Corticosteroids 58.8%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse	
<u>C3PO trial</u> ; ¹⁷⁰ Korley et al; peer reviewed; 2021	Patients with early mild to moderate COVID-19 infection with risk factors for severe disease. 257 assigned to CP 250 ml and 254 assigned to SOC	Median age 54 ± 21, male 46%, hypertension 42.3%, diabetes 27.8%, COPD 6.1%, CHD 10%, CKD 5.3%, cancer 0.8%, obesity %	NR	events outcomes results. Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
DAWn-Plasma trial; ¹⁷¹ Devos et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 320 assigned to CP 200 to 250 ml once or twice and 163 assigned to SOC	Mean age 62 ± 14, male 68.7%, hypertension %, diabetes 29.6%, COPD 9.4%, asthma 10.1%, CHD 14.1%, CKD 13.4%,	Corticosteroids 66.4%, remdesivir 14.8%, hydroxychloroquine 1.4%, lopinavir- ritonavir 0.4%, tocilizumab 0.6%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>PennCCP2 trial</u> ; ¹⁷² Bar et al; peer	Patients with severe COVID-19 infection.	Mean age 63 , male 45.6%, hypertension	Corticosteroids 83.5%, remdesivir 81%,	High for mortality and mechanical ventilation;	





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reviewed; 2021	40 assigned to CP two units and 39 assigned to SOC	67.1%, diabetes 40.5%, COPD 29.1%, CHD 29.1%, CKD 32.9%, immunosuppression 13.9%, cancer 26.6%, obesity 45.6%	hydroxychloroquine 2.5%,	high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
Manichetti et al;	Patients with moderate to severe COVID-19 infection. 231 assigned to CP 200 ml a day for 1 to 3 days and 239 assigned to SOC	male 64.3%,	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
	Patients with mild to moderate COVID-19 infection. 390 assigned to CP 200 to 300 ml once and 392 assigned to SOC	Median age 58 ± 11, male 66.8%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>CSSC-004 trial;</u> ¹⁷⁵ Sullivan et al; peer reviewed; 2022	Patients with mild COVID-19 infection. 592 assigned to CP 250 ml and 589 assigned to SOC	Median age 44, male 43%, hypertension 23.3%, diabetes 8.4%, asthma 11.2%, CHD 2%, CKD 0.9%, cerebrovascular disease 0.2%, cancer 0.5%, obesity 17.3%	Vaccinated 17.5%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>COP20 trial</u> ; ¹⁷⁶ Holm et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 17 assigned to CP 200 to 250 ml on three consecutive days and 14 assigned to SOC	Mean age 73.2 ± , male 61.3%, hypertension 41.9%	Corticosteroids 71%, remdesivir 10%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have	





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				introduced bias to symptoms and adverse events outcomes results.
<u>CONTAIN</u> <u>COVID-19 trial</u> ; ¹⁷⁷ Ortigoza et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 463 assigned to CP 250 ml once and 463 assigned to SOC	Median age 63, male 59.1%, hypertension 60.7%, diabetes 35.3%, COPD %, asthma 11.7%, CHD 42.9%, CKD 10.5%, cancer 11.3%,	Corticosteroids 76.6%, remdesivir 57.1%, hydroxychloroquine 3.5%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
IMPACT trial; ¹⁷⁸ <u>Baldeón et al</u> ; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 63 assigned to CP 5 ml/kg and 95 assigned to SOC	Mean age 55.5, male 67.7%, hypertension 22.2%, diabetes 19.6%, obesity 24.7%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>De Santis et al</u> ; ¹⁷⁹ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 36 assigned to CP 600 ml a day for 3 days and 71 assigned to SOC	Mean age 59.8, male 62.6%, hypertension 56%, diabetes 38.3%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
	Patients with severe COVID-19 infection. 52 assigned to CP 200- 250 ml once and 51 assigned to SOC	Median age 56, male 40.8%, hypertension 54.4%, diabetes 38.8%, COPD 3.9%, CHD 2.9%, CKD 2.9%, cancer 1.9%, obesity 47.6%	Corticosteroids 94.2%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
LIFESAVER <u>trial;¹⁸¹ et al; other;</u> 2021	Patients with severe to critical COVID-19 infection. 4 assigned to	NR	NR	Low for mortality and mechanical ventilation; low for symptom





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extracted from systematic review		critical COVID-19 infection. 8 assigned to CP and 6 assigned to	NR	NR	mechanical ventilation; low for symptom resolution, infection and adverse events	
Tatem G et al: ¹⁸¹ Patients with severe to NR Low for mortality and					extracted from	
	<u>Tatem G et al</u> ; ¹⁸¹	Patients with severe to	NR	NR	Low for mortality and	



other; 2021	critical COVID-19 infection. 20 assigned to CP and 10 assigned to SOC			mechanical ventilation; low for symptom resolution, infection and adverse events Notes: RoB assessment extracted from systematic review	
<u>Chowdhury FR et</u> <u>al</u> ; ¹⁸¹ other; 2021	Patients with severe to critical COVID-19 infection. 20 assigned to CP and 10 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes: RoB assessment extracted from systematic review	
<u>PLACO-COVID</u> <u>trial</u> ; ¹⁸¹ other; 2021	Patients with severe to critical COVID-19 infection. 60 assigned to CP and 60 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes: RoB assessment	
				extracted from systematic review	
ASCOT trial; ¹⁸¹ other; 2021	Patients with moderate to severe COVID-19 infection. 15 assigned to CP and 18 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
				Notes: RoB assessment extracted from systematic review	
Co-CLARITY trial; ¹⁸¹ other; 2021	Patients with moderate to severe COVID-19 infection. 13 assigned to CP and 12 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
	1				





			1	
				Notes: RoB assessment extracted from systematic review
<u>Rego EM et al</u> ; ¹⁸¹ other; 2021	Patients with moderate to severe COVID-19 infection. 9 assigned to CP and 8 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
				Notes: RoB assessment extracted from systematic review
PERUCONPLAS MA trial; ¹⁸¹ other; 2021	Patients with severe to critical COVID-19 infection. 12 assigned to CP and 13 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
				Notes: RoB assessment extracted from systematic review
<u>CP-COVID-19</u> <u>trial</u> ; ¹⁸¹ other; 2021	Patients with severe to critical COVID-19 infection. 49 assigned to CP and 51 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
				Notes: RoB assessment extracted from systematic review
CONFIDENT <u>trial;¹⁸¹ other;</u> 2021	Patients with severe to critical COVID-19 infection. 150 assigned to CP and 151 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
				Notes: RoB assessment extracted from systematic review
PC/COVID-19	Patients with severe to	NR	NR	Low for mortality and

<u>trial</u> ; ¹⁸¹ other; 2021	critical COVID-19 infection. 38 assigned to CP and 36 assigned to SOC			mechanical ventilation; low for symptom resolution, infection and adverse events Notes: RoB assessment extracted from systematic review	
<u>COP-COVID-19</u> <u>trial;¹⁸¹ other; 2021</u>	Patients with severe to critical COVID-19 infection. 20 assigned to CP and 11 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes: RoB assessment extracted from systematic review	
<u>CCAP-2 trial</u> ; ¹⁸³ peer reviewed; 2022	critical COVID-19 infection. 98 assigned to CP 600 ml once and	Mean age 65.3, male 72.2%, hypertension 28.5%, diabetes 22.2%, COPD 11.1%, cancer 6.9%,	Corticosteroids 88.9%, remdesivir 86.1%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>COOPCOVID</u> <u>trial</u> ; ¹⁸⁴ Song et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 87 assigned to CP 200 to 400 ml once and 42 assigned to SOC	Median age 61 ± , male 68%, one or more comorbidities 92%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>COPLA-II trial</u> ; ¹⁸⁵ Bajpai et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 200 assigned to CP 250 ml twice and 200 assigned to SOC	Mean age 55.5 ± 1.17, male 67.3%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	



				Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
CAPRI trial; <u>NCT</u> <u>04421404;</u> other; 2021	Patients with moderate to severe COVID-19 infection. 16 assigned to CP 250 ml once and 18 assigned to SOC	Median age 57, male 44.1%	NR	NA
<u>CoVIP trial;</u> ¹⁸⁶ Bartelt et al; preprint; 2021	Patients with moderate to critical COVID-19 infection. 14 assigned to CP (high titer) 200 to 300 ml twice and 41 assigned to CP (normal titer) 200 to 300 ml twice	Median age 61, male 64%, hypertension 20%, diabetes 43.6%, COPD 16.3%, CHD 12.7%, immunosuppressive therapy 29.1%, cancer 5.5%, obesity 58.2%	Corticosteroids 90.9%, remdesivir 92.7%	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Significant cross- over which affected blinding. No intention to treat analysis estimates provided.
<u>CSSC-001 trial</u> ; ¹⁸⁷ Shoham et al; peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 81 assigned to CP one unit once and 87 assigned to SOC	Median age 47, male 55%, diabetes 6.1%, asthma 5%, CHD 2.2%, immunosuppresive therapy 0.5%, cancer 1.1%	Vaccinated 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
<u>Rojas et al</u> ; ¹⁸⁸ peer reviewed; 2022	Patients with severe COVID-19 infection. 46 assigned to CP 250 ml twice and 45 assigned to SOC	Mean age 55, male 70.3%, hypertension 25.3%, diabetes 16.5%, COPD %, asthma 4.4%, CKD 5.5%	Corticosteroids 96.7%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
Bargay-Lleonart et al; ¹⁸⁹ peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 37 assigned	Mean age 58.2, male 61.1%	NR	High for mortality and mechanical ventilation; high for symptom

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	to CP 300 ml twice and 17 assigned to SOC			resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>Self et al</u> ; ¹⁹⁰ peer reviewed; 2022	Patients with moderate to critical COVID-19 infection. 487 assigned to CP 200 to 400 ml once and 473 assigned to SOC	57.3%, hypertension	Corticosteroids 86.7%, remdesivir 70.8%, Vaccinated 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	
Balcells et al; ¹⁹¹ peer reviewed; 2020	Patients with moderate to severe COVID-19. 28 assigned to convalescent plasma at enrolment, 200 mg twice and 30 assigned to convalescent plasma when clinical deterioration was observed (43.3% received CP in this arm)	male 50%, hypertension 67.2%, diabetes 36.2%, chronic lung disease %, asthma 5.1%, coronary heart disease %, chronic	Corticosteroids 51.7%, hydroxychloroquine 12%, lopinavir- ritonavir 1.7%, tocilizumab 3.4%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty ()Invasive mechanical ventilation: Very low certainty ()Symptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ()Very low certainty ()Hospitalization: No information
Non-RCT				1	
<u>Joyner et al</u> ; ¹⁹² peer-	Patients with moderate	Median age 62.3 ± 79.3 ,	NR	Low for specific	Adverse events:





reviewed; 2020	to critical COVID-19 infection. 20000 received CP	male 60.8%		transfusion related adverse events	Transfusion related circulatory overload 0.18%; Transfusion related lung injury 0.10%; Severe allergic transfusion reaction 0.10%
	Uncerta	Criza inty in potential benefits a	nlizumab and harms. Further resea	nrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
CRITICAL trial; ¹⁹³ Leucker et al; peer reviewed; 2021	Patients with severe to critical COVID- 19 infection. 22 assigned to crizanlizumab 5 mg/kg once and 20 assigned to SOC	Mean age 56.6, male 54.5%, hypertension 70.4%, diabetes 43.1%, COPD 9.1%, asthma 6.8%, CHD 11.3%, CKD 11.3%, cerebrovascular disease 2.2%,	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	Mortality: Very low certainty \oplus ()Invasive mechanical ventilation: Very low certainty \oplus ()Symptom resolution or improvement: Very low certainty \oplus ()Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty \oplus ()Hospitalization: No



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					information
	Uncerta	Curcumi inty in potential benefits a	in + Piperine and harms. Further rese	arch is needed.	l
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT		•	•	•	
<u>Askari et al</u> ; ¹⁹⁴ peer reviewed; 2022	Patients with mild to moderate COVID- 19 infection. 23 assigned to curcumin + piperine 1000/10 mg a day for 14 days and 23 assigned to SOC	Mean age 47.6 ± 13.9, male 58.7%, hypertension 23.9%, diabetes 26.1%, CHD 15.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
		Curcumin + Qu inty in potential benefits			
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the



					evidence
RCT					
Khan et al; ¹⁹⁵ peer reviewed; 2022	Patients with moderate COVID- 19 infection. 25 assigned to curcumin + quercetin + Vit D 168 mg + 260 mg + 360 IU and 25 assigned to SOC	Mean age 43.9, male 50%, hypertension 28%, diabetes 34%	Vaccinated 52%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: Very low certainty ⊕○○○Symptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
Dapaş	gliflozin may reduce mor		ngliflozin 10t increase symptom res	solution. Further research	is needed.
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
DARE-19 trial; ¹⁹⁶ Kosiborod et al; peer reviewed; 2021	Patients with moderate COVID-19 infection and cardiometabolic risk factors. 625	Mean age 61.4 ± 13.5, male 57.4%, hypertension 84.8%, diabetes 50.9%, COPD	Corticosteroids 28.4%, remdesivir 18%	Low for mortality and mechanical ventilation; low for symptom resolution, infection,	Mortality: RR 0.76 (95%Cl 0.51 to 1.12); RD -3.8% (95%Cl -7.8% to

	assigned to dapagliflozin 10 mg for 30 days and 625 assigned to SOC	4.6%, CHD 7.2%, CKD 6.6%, obesity 48.1%		and adverse events	1.9%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: No information Symptom resolution or improvement: RR 1.02 (95%CI 0.98 to 1.06); RD 1.2% (95%CI -1.2% to 3.6%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Hospitalization: No information
	Uncertai	Darunav inty in potential benefits a	ir-cobicistat nd harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
DC-COVID-19 <u>trial</u> ; ¹⁹⁷ Chen et al; peer-reviewed; 2020	Patients with mild COVID-19 infection. 15 assigned to darunavir-cobicistat 800 mg/150 mg once a	Mean age 47.2 ± 2.8, male NR, diabetes 6.6%, coronary heart disease 26.6%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse	Mortality: No information Invasive mechanical ventilation: No



	day for 5 days and 15 assigned to standard of care		garelix	events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncerta	inty in potential benefits	and harms. Further rese	earch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>HITCH trial</u> , ¹⁹⁸ Nickols et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 62 assigned to degarelix 240 mg once and 34 assigned to SOC	Mean age 68.5 ± 8.4, male 100%, hypertension 78.1%, diabetes 51%, COPD 15.6%, asthma 12.5%, CHD 28.1%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis



					studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
DFV890 may imj	prove time to sympton	resolution. The effect	FV890 s of AZD 1656 on othe h is needed.	r important outcomes ar	e uncertain. Further
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Madurka et al; ¹⁹⁹ peer reviewed; 2022	Patients with severe COVID-19 infection. 70 assigned to DFV890 100 mg a day for 14 days and 72 assigned to SOC	Mean age 61, male 67.6%, hypertension 60.6%, diabetes 26.1%, COPD 9.9%, CHD 12%, CKD 2.1%, cerebrovascular disease 4.9%, cancer 6.4%,	Corticosteroids 71.1%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Significant loss to follow-up.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: RR 1.15 (95%CI 0.96 to 1.36); RD 9.1% (95%CI 2.4% to 21.8%); Low certainty ⊕⊕○○ Symptomatic infection (prophylaxis studies): No information Adverse events:

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Study; publication status		methyl sulfoxide inty in potential benefits a Comorbidities			Very low certainty ⊕○○○ Hospitalization: No information Interventions effects vs standard of care and GRADE certainty of the evidence
RCT <u>Hosseinzadeh et</u> al; ²⁰⁰ preprint; 2021	Individuals exposed to SARS-CoV-2 infection. 116 assigned to DSMO three applications a day for one month and 116 assigned to SOC			Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationGymptomatic infection (prophylaxis studies): Very low certainty ⊕○○○Adverse events: No informationHospitalization: No information
Study; publication status	Doxycycline doo Patients and interventions analyzed	es not improve time to syn	alfa (inhaled) mptom resolution. Furth Additional interventions	er research is needed. Risk of bias and study limitations	Interventions effects vs standard of care





					and GRADE certainty of the evidence
RCT				·	
COVASE trial; ²⁰¹ Porter et al; preprint; 2021	Patients with severe COVID-19 infection. 30 assigned to inhaled dornase alfa 5 mg a day for 7 days and 9 assigned to SOC	Mean age 56, male 76.9%, any commorbiditie 51.2%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Doxycycline do	Doxy es not improve time to syr	y cycline nptom resolution. Furtl	er research is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
DOXYCOV trial; ²⁰² Sobngwi et al; preprint; 2021	Patients with mild COVID-19 infection. 92 assigned to doxycycline 200 mg a	Mean age 39 ± 13, male 52.4%, hypertension 1.1%, asthma 1.6%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection,	Mortality: Very low certainty ⊕○○○ Invasive mechanical

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	day for 7 days and 95 assigned to SOC Patients with mild COVID-19 infection. 780 assigned to doxycycline 200 mg once followed by 100 mg a day for 7 days and 948 assigned to SOC Patients with moderate to severe COVID-19	63.8%, hypertension	NR Corticosteroids 81.4%, tocilizumab 1.3%,	and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Low for mortality and mechanical ventilation;	ventilation: No information Symptom resolution or improvement: RR 1 (95%CI 0.97 to 1.03); RD 0% (95%CI -1.8% to 1.8%); High certainty ⊕⊕⊕⊕ Symptomatic infection (prophylaxis studies): Very low certainty ⊕○○○ Adverse events: Very low certainty
al; preprint; 2021	to doxycycline 200 mg	53.2%, diabetes 35.7%, COPD 9%, asthma 7.5%, CHD 13.4%, cancer 1.3%,		high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	 ⊕○○○ Hospitalization: RR 1.16 (95%CI 0.76 to 1.76); RD 0.7% (95%CI -1.1% to 3.6%); Low certainty ⊕⊕○○
Stambouli et al; ²⁰⁵ peer reviewed; 2022	Individuals exposed to SARS-CoV-2 infection. 56 assigned to doxycycline 100 mg a day for 6 weeks and 57 assigned to SOC	male 61%, hypertension 4.1%, diabetes 2.3%, COPD 0.6%, asthma	Vaccinated 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
	Uncertai	Dup inty in potential benefits a	ilumab nd harms. Further resea	irch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					



SafeDrop trial; ²⁰⁶ Sasson et al; preprint; 2021	Patients with severe COVID-19 infection. 19 assigned to dupilumab 600 mg once followed by 300 mg on days 14 and 28 and 21 assigned to SOC	Mean age 61, male 57.5%, hypertension 45%, diabetes 37.5%, COPD 12.5%, asthma 20%, CHD 22.5%, CKD 25%, cancer 17.5%, obesity 72.5%	Corticosteroids 97.5%, remdesivir 85%, tocilizumab 0%; Vaccinated 65%	Some Concerns for mortality and mechanical ventilation; some Concerns for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: Very low certainty @OOOInvasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationAdverse events: No informationAdverse events: No informationHospitalization: No information
	Uncerta	Dut: inty in potential benefits a	isteride nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	-	-			
<u>AB-DRUG-SARS-</u> <u>004 tria</u> l; ²⁰⁷ Cadegiani et al; preprint; 2020	Patients with mild COVID-19. 64 assigned to dutasteride (dosage not reported) and 66 assigned to standard of care	Mean age 42 ± 12, male 100 %, diabetes 11%, COPD 0%, asthma 1%, coronary heart disease 1%, cancer 0%, obesity 15.4%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or





EAT-DUTA AndroCoV trial; ²⁰⁸ Cadegiani et al; Peer reviewed; 2020	moderate COVID-19.	Mean age 41.9 ± 12.4, male 100%, hypertension 21.8%, diabetes 9.2%, COPD 0%, asthma 1.1%, CHD 1.1%, cancer 0%, obesity 10.3%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Significant lost to follow-up.	improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: No informationHospitalization: Very low certainty ⊕○○○
	Uncertai	Eda inty in potential benefits a	ravone nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Moslemi et al; ²⁰⁹ peer reviewed; 2022	Patients with severe COVID-19 infection. 19 assigned to edaravone 30 mg a day for 3 days and 19 assigned to SOC	Mean age 60.5, male 47.3%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information

					Adverse events: No information Hospitalization: No information
	Uncertai	Electro inty in potential benefits a	lyzed saline and harms. Further resea	rrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			•		
<u>TX-COVID19</u> <u>trial</u> ; ²¹⁰ Delgado- Enciso et al; preprint; 2020	Patients with mild to moderate COVID-19. 45 assigned to electrolyzed saline nebulizations 4 times a day for 10 days and 39 assigned to standard of care	Mean age 47 ± 14.6, male 53.5%, hypertension 18.9%, diabetes 11.9%	Corticosteroids 3.65%, hydroxychloroquine 7.5%, ivermectin 9.4%, ATB 30.6%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information
ICU-VR trial; Gutiérrez-García et al; ²¹¹ peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 79 assigned to electrolyzed saline nasal sprays and gargles three times a day and 84 assigned to SOC	Mean age 42 ± , male 26.4%, hypertension 6.7%, diabetes 4.9%, obesity 13.5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Symptomatic infection (prophylaxis studies): Very low certainty ⊕○○○ Adverse events: No information Hospitalization: Very low certainty ⊕○○○

Uncertainty in potential benefits and harms. Further research is needed.



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Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
MEDIC-LAUMC trial; ²¹² Matli et al; peer reviewed; 2022	Patients with mild to severe COVID-19 infection. 17 assigned to nicorandil 20 mg a day, L-arginine 3 gr a day, folate 5 mg a day, nebivolol 2.5 to 5 mg a day, and atorvastatin 40 mg a day for 14 days, and 20 assigned to SOC	Mean age 56.6, male 81.8%, hypertension 27%, diabetes 21.6%, asthma 10.8%, CHD 5.4%, CKD 2.7%, cancer 2.7%,	Corticosteroids 91.9%, remdesivir 59.5%, tocilizumab 8.1%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncertai	Enis inty in potential benefits a	amium nd harms. Further resea	irch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Holubovska et al</u> ; ²¹³ Preprint; 2020	Patients with moderate to severe COVID-19.	NR	NR	High for mortality and mechanical ventilation;	Mortality: No information



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	assigned to enisamium 500 mg 4 times a day for 7 days or SOC. Number of patients in each arm not reported.			High for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information				
	Ensitrelvir Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence				
RCT									



Mukae et al-2; ²¹⁴ preprint; 2021	Patients with mild to moderate COVID-19 infection. 30 assigned to ensitrelvir 125 to 250 mg a day for 5 days and 17 assigned to SOC	Mean age 38.9, male 61.7%, Ens	Vaccinated 80.8%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
Ensovibep may	not improve time to sy		e effectos of ensovibep o earch is needed	on other importan outco	mes are uncertain.
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
ACTIV-3/TICO trial; ²¹⁵ Barkauskas et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 247 assigned to ensovibep 600 mg once and 238 assigned to SOC	Median age 57 ± , male 56.7%, hypertension 39.4%, diabetes 23.5%, COPD 6.2%, asthma 9.3%, CHD %, CKD 9.5%, cerebrovascular disease %, immunosuppresive therapy 6.2%, cancer %,	Corticosteroids 72.9%, remdesivir 68.7%, hydroxychloroquine %, lopinavir-ritonavir %, tocilizumab %, azithromycin %, convalescent plasma %; Vaccinated 31.6%	Low for mortality and mechanical ventilation; Low for symptom resolution, infection and adverse events Notes:	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or



		obesity 13.4%	utamide		improvement: RR 0.95 (95% CI 0.8 to) 1.16); RD -2.8% (95% CI -13.1% to) 9.7%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Hospitalization: No information
	Uncertai	EIIZAI inty in potential benefits a		rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
COVIDENZA trial; ²¹⁶ Welen et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 30 assigned to enzalutamide 160 mg a day for 5 days and 12 assigned to SOC	Median age 64.9, hypertension 45.2%, diabetes 19%, asthma 14.3%, CHD 9.5%, cancer 11.9%,	Corticosteroids 85.7%, remdesivir 28.6%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or improvement: No information





		Ethano	l (inhaled)		(prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncertai	inty in potential benefits a		rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
Non-RCT			·		
Amoushahi et al; ²¹⁷ preprint; 2022	Patients with moderate to severe COVID-19 infection. 44 assigned to ethanol (inhaled) 3 sprays, four times a day for 7 days and 55 assigned to SOC	Mean age 46.4 ± 12.8, male 43.7%,	Corticosteroids 100%, remdesivir 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No



					information			
Famotidine Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
Non-RCT	•			· · · · · · · · · · · · · · · · · · ·				
Samimagham et al; ²¹⁸ preprint; 2021	Patients with moderate to severe COVID-19 infection. 10 assigned to famotidine 160 mg for up to 14 days and 10 assigned to SOC	Mean age 47.5 ± 13, male 60%,	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty ⊕○○○ Invasive mechanica ventilation: No information Symptom			
<u>Brennan et al</u> ; ²¹⁹ peer reviewed; 2021	Patients with mild recent onset COVID- 19 infection. 27 assigned to famotidine 60 mg a day for 14 days and 28 assigned to SOC	Mean age 35 ± 20, male 36.4%	Vaccinated 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Significant loss to follow up.	resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: Na information			
Pahwani et al; ²²⁰ beer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 89 assigned to famotidine 40 mg a day and 89 assigned to SOC	Mean age 51.5 ± 11.5, male 68.5%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.				



	symptom resolution. Further research is needed.						
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence		
RCT							
<u>Chen et al;</u> preprint; ²²¹ 2020	Patients with moderate to critical COVID-19 infection. 116 assigned to favipiravir 1600 mg twice the first day followed by 600 mg twice daily for 7 days and 120 assigned to umifenovir 200 mg three times daily for 7 days	Mean age not reported male 46.6%, hypertension 27.9%, diabetes 11.4%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: RR 1.08 (95%Cl 0.77 to 1.52); RD 1.3% (95%Cl -3.7% to 8.3%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: RR 1.27 (95%Cl 0.91 to 1.76); RD 4.7% (95%Cl - 1.6% to 13.1%); Low certainty $\oplus \oplus \bigcirc \bigcirc$		
<u>Ivashchenko et al;</u> ²²² peer-reviewed; 2020	Patients with moderate COVID-19 infection. 20 assigned to favipiravir 1600 mg once followed by 600 mg twice a day for 12 days, 20 assigned to favipiravir and 20 assigned to standard of care	Mean age not reported	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Symptom resolution or improvement: RR 1.01 (95%CI 0.97 to 1.05); RD 0.6% (95%CI -1.8% to 3%); High certainty ⊕⊕⊕⊕ Symptomatic		
<u>Lou et al</u> ; ⁸⁶ preprint; 2020	Patients with mild to severe COVID-19 infection. 10 assigned to baloxavir 80 mg a day on days 1, 4 and 7, 9 assigned to favipiravir and 10 assigned to standard of care	Mean age 52.5 ± 12.5, male 72.4%, hypertension 20.7%, diabetes 6.9%, coronary heart disease 13.8%,	Antivirals 100%, IFN 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	infection (prophylaxis studies): No information Adverse events: RR 0.92 (95%CI 0.56 to 1.52); RD -0.8% (95%CI -4.5% to 5.3%); Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$		



Doi et al; ²²³ peer- reviewed; 2020	Patients with mild COVID-19. 44 assigned to favipiravir (early) 1800 mg on day 1 followed by 800 mg twice daily for 10 days and 45 assigned to favipiravir (late) 1800 mg on day 6 followed by 800 mg twice daily for 10 days	Median age 50 ± 26.5, male 61.4%, comorbidities 39%	Corticosteroids 2.3%, ATB 12.5%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Hospitalization: RR 1.33 (95%CI 0.64 to 1.78); RD 1.6% (95%CI -1.7% to 3.7%); Low certainty ⊕⊕⊖⊖
Dabbous et al; ²²⁴ preprint (now retracted); 2020	Patients with mild to moderate COVID-19. 50 assigned to favipiravir 3200 mg once followed by 1200 mg a day for 10 days and 50 assigned to hydroxychloroquine + oseltamivir 800 mg once followed by 400 mg a day for 10 days + 75 mg a day for 10 days	Mean age 36.3 ± 12, male 50%, any comorbidities 15%	NR	High for mortality and invasive mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Zhao et al</u> ; ²²⁵ peer- reviewed; 2020	Patients with moderate to critical COVID-19 infection. 13 assigned to favipiravir 3200 mg once followed by 600 mg twice a day for 7 days, 7 assigned to TCZ 400 mg once or twice and 5 assigned to favipiravir + TCZ	Mean age 72 ± 40, male 54%, hypertension 42.3%, diabetes 11.5%, coronary heart disease 23.1%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Khamis et al</u> ; ²²⁶ peer-reviewed; 2020	Patients with moderate to severe COVID-19. 44 assigned to favipiravir + inhaled interferon beta-1B 1600 mg once followed by 600 mg	Mean age 55 ± 14, male 58%, hypertension 54%, diabetes 45%, COPD 5.6%, coronary heart disease 15%, chronic kidney disease 20%	Corticosteroids 67%, tocilizumab 35%, convalescent plasma 58%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events	



	twice a day for 10 days + 8 million UI for 5 days and 45 assigned to standard of care			Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Ruzhentsova et</u> <u>al</u> ; ²²⁷ preprint; 2020	Patients with mild to moderate COVID-19. 112 assigned to favipiravir 1800 mg once followed by 800 mg twice a day for 10 days and 56 assigned to standard of care	Mean age 42 ± 10.5, male 47%	NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Promomed;</u> NCT04542694; Other; 2020	Patients with moderate COVID-19. 100 assigned to favipiravir 3200 mg once followed by 600 mg twice a day for 14 days and 100 assigned to standard of care	Mean age 49.68 ± 13.09, male 48.5%,	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
Udwadia et al; ²²⁸ peer-reviewed; 2020	Patients with mild to moderate COVID-19. 72 assigned to favipiravir 3600 mg once followed by 800 mg twice a day for 14 days and 75 assigned to standard of care	Mean age 43.4 ± 11.7, male 73.5%, comorbidities 25.9%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Balykova et al</u> ; ²²⁹ peer-reviewed; 2020	Patients with moderate to severe COVID-19. 100 assigned to favipiravir 3200 mf once followed	Mean age 49.7 ± 13, male 50%, hypertension 28.5%, diabetes 9%, COPD 5%, asthma %, CHD 6%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events	



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	by 1200 mg a day for 14 days and 100 assigned to SOC			Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>Solaymani-Dodaran</u> <u>et al;²³⁰ p</u> eer- reviewed; 2021	critical COVID-19 infection. 190 assigned	Mean age 57.6 ± 17.3, male 55%, hypertension 34.9%, diabetes 25.7%, COPD 3.5%, asthma 3.8%, CHD 10.7%, CKD 1.6%	Corticosteroids 27.6%, remdesivir 1.1%,	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events
<u>Zhao et al</u> ; ²³¹ peer reviewed; 2021	Patients with COVID- 19 infection who were discharged from hospital. 36 assigned to favipiravir 3200 mg once followed by 1200 mg a day for 7 days and 19 assigned to SOC	male 45.5%, hypertension 30.9%,	Corticosteroids 3.6%, remdesivir 0%, hydroxychloroquine 5.5%, lopinavir- ritonavir 16.4%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
FACCT trial; ²³² Bosaeed et al; preprint; 2021	critical COVID-19 infection. 125 assigned	Mean age 52 ± 13, male 59%, hypertension 40.9%, diabetes 42.1%, asthma 11.8%, CKD 2.4%	Corticosteroids 88.6%, tocilizumab 9%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
Shinkai et al; ²³³ peer reviewed; 2021	Patients with moderate COVID-19 infection. 107 assigned to favipiravir 3200 mg once followed by 1600 mg a day for 14 days and 49 assigned to SOC	Mean age 46.2, any comorbidities 75.6%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have





				introduced bias to symptoms and adverse events outcomes results.
FIGHT-COVID- 19 trial; ²³⁴ Atipornwanich et al; preprint; 2021	Patients with mild to severe COVID-19 infection. 320 assigned to favipiravir 6000 mg once followed by 2400 mg a day + lopinavir ritonavir 800/200 mg or lopinavir ritonavir 800/200 mg a day or HCQ 800 mg a day or darunavir ritonavir 1200/200 mg a day + HCQ 400 mg a day or favipiravir 6000 mg followed by 2400 mg + darunavir ritonavir 1200/200 mg a day + HCQ 400 mg a day for 7 to 14 days.	Mean age 42 ± 15.7, male 47.8%, obesity 24.6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>CVD-04-CD-001</u> <u>trial</u> ; ²³⁵ Shenoy et al; preprint; 2021	Patients with moderate to severe COVID-19 infection. 175 assigned to favipiravir 3600 mg on day 1 followed by 1600 mg a day for 10 days and 178 assigned to SOC	Mean age 51.9 ± 12.5, male 67.4%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>Holubar et al</u> ; ²³⁶ preprint; 2021	Patients with mild to moderate COVID-19 infection. 59 assigned to favipiravir 3600 mg once followed by 1600 mg a day for 10 days and 57 assigned to SOC	Mean age 43 ± 12, male 51.9%, hypertension 8.6%, diabetes 8.6%, COPD 4.3%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
<u>Malaysian</u> <u>Favipiravir Study</u>	Patients with mild to moderate COVID-19	Mean age 62.5 ± 8, male 48.4%, hypertension	Corticosteroids 24.6%, tocilizumab 2%,	Low for mortality and mechanical ventilation;





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	infection. 250 assigned to favipiravir 3601 mg once followed by 1600 mg a day for 5 days and 250 assigned to SOC	80.2%, diabetes 49.8%, COPD 1.4%, asthma 7.4%, CHD 15%, CKD 1.4%, immunocompromised therapy 0.4%, cancer 1.4%, obesity 20.6%	vaccinated 0.4%	high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>trial</u> ; ²³⁸ Finberg et al;	Patients with moderate to severe COVID-19 infection. 25 assigned to favipiravir 3600 mg once folowed by 2000 mg a day for 14 days and 25 assigned to SOC	Mean age 57.2 ± 13.14, male 60%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Avi-Mild trial</u> ; ²³⁹ Bosaeed et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 112 assigned to favipiravir 3600 mg once followed by 1600 mg a day for 5 to 7 days and 119 assigned to SOC	Median age 37, male 67%, hypertension 6%, diabetes 10.8%, COPD %, asthma 3.4%, CHD 0.4%, obesity 16.8%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>Hassaniazad et al</u> ; ²⁴⁰ peer reviewed; 2021	Patients with severe COVID-19 infection. 32 assigned to favipiravir 3200 mg once followed by 1200 mg for 5 days and 31 assigned to lopinavir- ritonavir 400/100 mg a day for 7 days	Mean age 53.7 ± 13.5, male 57.1%, hypertension 27%, diabetes 20.6%, COPD 1.6%, CHD 14.2%, obesity 7.9%	Interferon beta 100%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>FLARE trial</u> ; ²⁴¹ Lowe et al; preprint;	Patients with recent onset mild COVID-19	Mean age 40 ± 12, male 51.2%, obesity 16.7%,	Vaccinated 51.2%	Low for mortality and mechanical ventilation;	



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2021	infection. 59 assigned to favipiravir 3600 mg once followed by 1600 mg a day for 7 days and 60 assigned to SOC	any comorbidity 15%		low for symptom resolution, infection and adverse events	
<u>Tabarsi et al</u> ; ²⁴² peer reviewed; 2021	COVID-19 infection. 32 assigned to favipiravir 3200 mg	Median age 57, male 58.1%, hypertension 12.9%, diabetes 21%, COPD %, asthma 3.2%, CHD 14.5%, CKD 3.2%, therapy %, cancer 4.8%, obesity 3.2%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>AlQahtani et al</u> ; ²⁴³ peer reviewed; 2021	moderate COVID-19 infection. 54 assigned	Mean age 44, male 47.1%, diabetes 26.1%, COPD 7.6%, asthma %, CHD 1.3%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>Rahman et al</u> ; ²⁴⁴ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 25 assigned to favipiravir 1200 mg a day for 5 days and 25 assigned to SOC	Mean age 37.8 ± 10.7, male 66%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	
<u>McMahon et al</u> ; ²⁴⁵ preprint; 2022	Patients with mild to moderate COVID-19 infection. 95 assigned to favipiravir 1800 mg once followed by 1600 mg a day for 14 days	Mean age 36, male 54.8%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	





	and 95 assigned to SOC				
<u>Golan et al</u> ; ²⁴⁶ peer reviewed; 2022	Patients with mild COVID-19 infection. 599 assigned to favipiravir 3600 mg once followed by 1600 mg a day for 10 days and 588 assigned to SOC	Age >60 14.7%, male 45.7%, any comorbidities 17.9%	Vaccinated 11%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<mark>Sirijatuphat et al</mark> , ²⁴⁷ preprint; 2022	Patients with mild to moderate COVID-19 infection. 62 assigned to favipiravir 3600 mg once followed by 1600 mg a day for 5 to 14 days and 31 assigned to SOC	Median age 30, male 35.5%, obesity 28%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
	Uncerta	Feb inty in potential benefits a	uxostat and harms. Further resea	rrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Davoodi et al; ²⁴⁸ peer-reviewed; 2020	Patients with moderate to severe COVID-19 infection. 30 assigned to febuxostat 80 mg per day and 30 assigned to HCQ	Mean age 57.7 ± 8.4, male 59%, hypertension NR%, diabetes 27.8%, chronic lung disease 1.9%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information





					Symptomaticinfection(prophylaxisstudies): NoinformationAdverse events: NoinformationHospitalization:Very low certainty⊕○○○Hospitalization: Noinformation
	·	Fend	ofibrate		<u></u>
Fenofibrate may	not increase severe ad	verse events. The effects		r importan outcomes are	e uncertain. Further
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	•		•		
FERMIN trial; ²⁴⁹ Chirinos et al; preprint; 2022	Patients with mild to moderate COVID-19 infection. 350 assigned to fenofibrate 145 mg a day for 10 days and 351 assigned to SOC	Mean age 49 ± 16, male 53%, hypertension 27%, diabetes 15%, COPD 12%, CHD 7%,	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information





					Adverse events: RR 0.76 (95% CI 0.53 to) 1.08); RD - 2.5% (95% CI - 4.8% to) 0.8%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Hospitalization: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
	Uncerta	Fina inty in potential benefits a	I steride and harms. Further resea	irch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT		<u> </u>	L	L	
Zarchoseinzade et al; ²⁵⁰ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 40 assigned to finasteride 5 mg a day for 7 days and 40 assigned to SOC	Mean age 72 ± 14, male 100%, hypertension 66.3%, diabetes 25%, COPD 12.5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation and blinding probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty



					⊕ ○○○ Hospitalization: No information
Fluvoxamine prob: Study; publication status	ably does not have an im Patients and interventions analyzed		Dxamine izations and may not inc Additional interventions	rease adverse events. Furtl Risk of bias and study limitations	ner research is needed. Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Lenze et al</u> ; ²⁵¹ peer- reviewed; 2020	Patients with mild to moderate COVID-19. 80 assigned to fluvoxamine incremental dose to 100 mg three times a day for 15 days and 72 assigned to standard of care	Median age 45.5 ± 20.5, male 28.2%, hypertension 19.7%, diabetes 11%, asthma 17.1%, obesity 56.6%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanica ventilation: Very low certainty ⊕○○○ Symptom
TOGETHER- Fluvoxamine trial; ²⁵² Reis et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 741 assigned to fluvoxamine 100 mg a day for 10 days and 756 assigned to SOC	Median age 50 ± 18, male 42.5%, hypertension 13.2%, diabetes 16.5%, COPD 0.6%, asthma 1.9%, CHD 1.1%, CKD 0.3%, obesity 0.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes:	resolution or improvement: No information Symptomatic infection (prophylaxis studies): No
<u>Seo et al</u> ; ²⁵³ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 26 assigned to fluvoxamine 200 mg a day for 10 days and 26 assigned to SOC	Mean age 53, male 59.6%, hypertension 26.9%, diabetes 7.7%, COPD 3.8%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	information Adverse events: RR 0.81 (95% CI 0.54 to) 1.22); RD -1.9% (95% CI -4.7% to) 2.2%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Hospitalization: RF





<u>COVID-OUT</u> <u>trial</u> ; ²⁵⁴ Bramante et al; peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 334 assigned to fluvoxamine 100 mg a day for 14 days and 327 assigned to SOC	Median age 44.5, male 45.8%, hypertension 26.9%, diabetes 1.1%, obesity 47.2%		Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	0.79 (95%CI 0.6 to 1.03); RD -1% (95%CI -1.9% to 0.1%); Moderate certainty ⊕⊕⊕⊖
	Uncertai	Fosta inty in potential benefits a	amatinib and harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	l				
Strich et al; ²⁵⁵ peer- reviewed; 2021	Patients with severe to critical COVID-19 infection. 30 assigned to fostamatinib 300 mg a day for 14 days and 29 assigned to SOC	Mean age 55.6 ± 13.7, male 79.7%, hypertension 54.2%, diabetes 37.3%, asthma 11.9%, CHD 13.6%, obesity 57.6%	Corticosteroids 100%, remdesivir 100%, convalescent plasma 42.4%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
		Gabapentin	+/- Montelukast		



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT				· ·	
Soltani et al; ²⁵⁶ peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 127 assigned to gabapentin +/- montelukast 900 mg a day +/- 10 mg a day for 5 days and 53 assigned to dextromethorphan	Mean age 56.7, male 56.1%, hypertension 22.2%, diabetes 16.1%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncerta	GB013 inty in potential benefits	9 (inhaled) and harms. Further	research is needed.	
Study; publication tatus	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence





DEFINE trial; ²⁵⁷ Gaughan et al; preprint; 2021	Patients with severe COVID-19 infection. 20 assigned to GB0139 (inhaled) and 21 assigned to SOC	Mean age 65, male 56%, hypertension 39%, diabetes 17%, asthma 14.6%, CHD 24.4%, CKD 7.3%, cancer 9.7%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
		mab (Anti-GM-(l Antibody) Further research is neede	d.
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
BREATHE trial; ²⁵⁸ Criner et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 113 assigned to gimsilumab 400 mg on day 1 and 200 mg on day 8 and 112 assigned to SOC	Mean age 60 ± 14, male 68.4%, hypertension 46.2%, diabetes 20.9%, COPD 7.6%, asthma %, CHD 8%, CKD %, cerebrovascular disease %, immunosuppresive therapy %, cancer %, obesity 26.7%	Corticosteroids 87.5%, remdesivir 50.6%, hydroxychloroquine 4%, ltocilizumab 7.6%, azithromycin 32.4%, convalescent plasma 0.4%;	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: RR 1.02 (95%CI 0.67 to 1.56); RD 0.3% (95%CI - 5.3% to 6%); Low certainty ⊕⊕○○ Invasive mechanical ventilation: No information

	Uncertai	Helium	(inhaled)	rch is needed.	Symptom resolution or improvement: RR $0.98 (95\% CI 0.82 to)$ $1.16); RD -1.2\%$ $(95\% CI -10.9\% to)$ $9.7\%); Low certainty\oplus \oplus \bigcirc \bigcircSymptomaticinfection(prophylaxisstudies): NoinformationAdverse events:Very low certainty\oplus \bigcirc \bigcirc \bigcircHospitalization: Noinformation$
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
	Patients with severe to critical COVID-19. 38 assigned to helium 50% to 79% mixed with oxygen and 32 assigned to SOC		NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic





Hesperidin	may not improve sympt		peridin the certainty of the evic	ience was low. Further resea	infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
HESPERIDIN trial; ²⁶⁰ Dupuis et al; preprint; 2021	COVID-19 infection. 104 assigned to hesperidin 1000 mg	Mean age 41 ± 12.1, male 44.9%, hypertension 10.6%, diabetes 3.2%, COPD 0.9%, asthma 13.5%, CHD 0%, cerebrovascular disease 0%,	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: RR 0.87 (95%CI 0.57 to 1.34); RD -7.9% (95%CI -26.1% to 20.6%); Low certainty ⊕⊕○○ Symptomatic infection (prophylaxis studies): No information Adverse events:





					Very low certainty ⊕○○○ Hospitalization: Very low certainty ⊕○○○
	Uncertai	Hema inty in potential benefits a	dsorption and harms. Further resea	urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
CYTOCOV-19 trial; ²⁶¹ Jarczak et al; preprint; 2021	Patients with critical COVID-19 infection. 12 assigned to hemadsorption and 12 assigned to SOC	Mean age 64.5 , male 75%, hypertension 66.6%, diabetes 33.3%, CHD 4%, CKD 25%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty \bigcirc Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty \bigcirc Symptomatic infection (prophylaxis studies): No informationAdverse events: No informationHospitalization: No information

Hydroxychloroquine and chloroquine

Hydroxychloroquine or chloroquine probably increases mortality, and probably does not reduce invasive mechanical ventilation or significantly improve time to symptom resolution with moderate certainty. When used prophylactically in persons exposed to COVID-19, it may not have an important effect on the risk of infection; and in patients with mild, recent onset disease, it may not have an important effect on hospitalizations. However, certainty of the evidence is low because of risk of bias and imprecision.



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>CloroCOVID19</u> <u>trial</u> ; ²⁶² Borba et al; peer-reviewed; 2020	Patients with severe COVID-19 infection. 41 assigned to chloroquine 600 mg twice a day for 10 days and 40 assigned to chloroquine 450 mg twice on day 1 followed by 450 mg once a day for 5 days	Mean age 51.1 ± 13.9, male 75.3%, hypertension 45.5%, diabetes 25.5%, chronic lung disease NR%, asthma 7.4%, coronary heart disease 17.9%, chronic kidney disease 7.4%, alcohol use disorder 27.5%, HIV 1.8%, tuberculosis 3.6%,	Azithromycin 100%, oseltamivir 89.7%	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: RR 1.09 (95%CI 1 to 1.19); RD 1.4% (95%CI 0% to 3%); Moderate certainty ⊕⊕⊕○ Invasive mechanical ventilation: RR 1.08 (95%CI 0.93 to 1.25); RD 1.4% (95%CI - 1.2% to 4.3%);
<u>Huang et al</u> ; ²⁶³ peer- reviewed; 2020	Patients with moderate to severe COVID-19 infection. 10 assigned to chloroquine 500 mg twice a day for 10 days and 12 assigned to lopinavir-ritonavir 400/100 mg twice a day for 10 days	Mean age 44 ± 21, male 59.1%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Moderate certainty $\oplus \oplus \oplus \bigcirc$ Symptom resolution or improvement: RR 1.01 (95%CI 0.93 to 1.1); RD 0.6% (95%CI -4.2% to 6.1%); Moderate certainty $\oplus \oplus \oplus \bigcirc$
<u>RECOVERY -</u> <u>Hydroxychloroquin</u> <u>e trial;²⁶⁴ Horby et</u> al; preprint; 2020	Patients with Mild to critical COVID-19 infection. 1561 assigned to hydroxychloroquine 800 mg once followed by 400 mg twice a day for 9 days and 3155 assigned to standard of care	Mean age 65.3 ± 15.3, male %, diabetes 26.9%, chronic lung disease 21.9%, asthma NR%, coronary heart disease 25.4%, chronic kidney disease 7.8%, HIV 0.4%	NR	Low for mortality and invasive mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Symptomatic infection (prophylaxis studies): RR 0.87 (95%CI 0.65 to 1.15); RD -2.2% (95%CI - 6.1% to 2.7%); Low certainty ⊕⊕○○ Severe Adverse events: RR 0.90 (95%CI 0.66 to 1.22); RD -1% (95%CI -
BCN PEP CoV-2	Individuals exposed to	Mean age 48.6 ± 19,	NR	Some concerns for	3.5% to 2.2%); Low certainty $\bigoplus \bigoplus \bigcirc \bigcirc$





<u>trial</u> ; ²⁶⁵ Mitja et al; preprint; 2020	SARS-CoV-2 infection. 1116 assigned to hydroxychloroquine 800 mg once followed by 400 mg x once a day for 6 days and 1198 assigned to standard of care	male 27%, diabetes 8.3%, chronic lung disease 4.8%, coronary heart disease 13.3%, Nervous system disease 4.1%		mortality and invasive mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Significant number of patients excluded from analysis.	Hospitalization: RR 0.82 (95%CI 0.61 to 1.1); RD -0.9% (95%CI -1.9% to 0.5%); Low certainty ⊕⊕○○
COVID-19 PEP trial; ²⁶⁶ Boulware et al; peer-reviewed; 2020	Individuals exposed to SARS-CoV-2 infection. 414 assigned to hydroxychloroquine 800 mg once followed by 600 mg daily for a total course of 5 days and 407 assigned to standard of care	male 48.4%,	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Significant loss of information that might have affected the study's results.	
Cavalcanti et al <u>trial</u> ; ²⁶⁷ Cavalcanti et al; peer-reviewed; 2020	moderate to severe	heart disease 0.8%, chronic kidney disease	Corticosteroids 1.5%, ACE inhibitors 1.2%, ARBs 17.4%, NSAID 4.4%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Kamran SM et al</u> <u>trial;²⁶⁸ Kamran et</u> al; preprint; 2020		Mean age 36 ± 11.2, male 93.2%, diabetes 3%, comorbidities 7.6%	NR	High for symptom resolution, infection, and adverse events	



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	400 mg twice a day once then 200 mg twice a day for 4 days and 151 assigned to standard of care			Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>COVID-19 PET</u> <u>trial</u> ; ²⁶⁹ Skipper et al; peer-reviewed; 2020		Median age 40 ± 9, male 44%, hypertension 11%, diabetes 4%, chronic lung disease %, asthma 11%,	NR	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>BCN PEP CoV-2</u> <u>trial</u> ; ²⁷⁰ Mitja et al; preprint; 2020	Patients with mild COVID-19 infection. 136 assigned to hydroxychloroquine 800 mg once followed by 400 mg a day for 6 days and 157 assigned to standard of care	Mean age 41.6 ± 12.6, male 49%, comorbidities 53.2%	NR	High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>Tang et al;</u> peer- reviewed; ²⁷¹ 2020	Patients with mild to moderate COVID-19 infection. 75 assigned to hydroxychloroquine 1200 mg daily for three days followed by 800 mg daily to complete 7 days and 75 assigned to standard of care	Mean age 46.1 ± 14.7, male 54.7%, hypertension 6%, diabetes 14%, other comorbidities 31%	Corticosteroids 7%, lopinavir-ritonavir 17%, umifenovir 47%, oseltamivir 11%, entecavir 1%, ATB 39%, ribavirin 47%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcome results.
<u>Chen et al.</u> ²⁷² preprint; 2020	Patients with moderate COVID-19 infection. 31 assigned to hydroxychloroquine 200 mg twice a day for	Mean age 44 ± 15.3, male 46.8%,	ATB 100%, IVIG 100%, antivirals 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events





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	5 days and 31 assigned to standard of care			Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>Chen et al</u> ; ²⁷³ preprint; 2020	Patients with moderate COVID-19 infection. 18 assigned to hydroxychloroquine 200 mg twice a day for 10 days, 18 assigned to chloroquine and 12 assigned to standard of care	Mean age 47.4 ± 14.46, male 45.8%, hypertension 16.7%, diabetes 18.7%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>Chen et al</u> , ²⁷⁴ preprint; 2020	Patients with mild to severe COVID-19 infection. 21 assigned to hydroxychloroquine 400 mg twice on day one followed by 200 mg twice a day for 6 days and 12 assigned to standard of care	Mean age 32.9 ± 10.7, male 57.6%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>HC-nCoV trial;</u> ²⁷⁵ Jun et al; peer- reviewed; 2020	Patients with mild to severe COVID-19 infection. 15 assigned to hydroxychloroquine 400 mg once a day for 5 days and 15 assigned to standard of care	Mean age 48.6 ± 3.7, male 0.7%, hypertension 26.6%, diabetes 6.6%, chronic lung disease 3.3%	Lopinavir-ritonavir 6.6%, umifenovir 73.3%, IFN 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>Abd-Elsalam et al</u> ; ²⁷⁶ peer-reviewed; 2020		Mean age 40.7 ± 19.3, male 58.8%, chronic kidney disease 3.1%,	NR	High for mortality and invasive mechanical ventilation; high for



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	to hydroxychloroquine 400 mg twice on day one followed by 200 mg tablets twice daily for 15 days and 97 assigned to standard of care	obesity 61.9%, comorbidities 14.3%, liver disease 1%		symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
COVID-19 PREP trial; ²⁷⁷ Rajasingham et al; peer-reviewed; 2020	Individuals exposed to SARS-CoV-2 infection. 989 assigned to hydroxychloroquine 400 mg twice in one day followed by 400 mg once weekly for 12 weeks or 400 mg twice weekly for 12 weeks and 494 assigned to standard of care	Median age 41 ± 15, male 49%, hypertension 14%, asthma 10%	NR	Low for infection, and adverse events	
TEACH trial; ²⁷⁸ Ulrich et al; peer- reviewed; 2020	Patients with mild to moderate COVID-19. 67 assigned to hydroxychloroquine 800 mg on day 1 followed by 200 mg twice a day for 2 to 5 days and 61 assigned to standard of care	Mean age 66 ± 16.2, male 59.4%, hypertension 57.8%, diabetes 32%, chronic lung disease 7%, asthma 15.6%, coronary heart disease 26.6%, chronic kidney disease 7.8%, cerebrovascular disease 6.2%	Corticosteroids 10.2%, remdesivir 0.8%, lopinavir-ritonavir 0.8%, azithromycin 23.4%, convalescent plasma 13.3%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	
PrEP_COVID trial; ²⁷⁹ Grau-Pujol et al; preprint; 2020	Individuals exposed to SARS-CoV-2 infection. 142 assigned to hydroxychloroquine 400 mg daily for four days followed by 400 mg weekly for 6 months and 127 assigned to standard of care	male 26.8%,	NR	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	





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<u>PATCH trial</u> ; ²⁸⁰ Abella et al; peer- reviewed; 2020	Individuals exposed to SARS-CoV-2 infection. 64 assigned to hydroxychloroquine 600 mg a day for 8 weeks and 61 assigned to standard of care	Median age 33 ± 46, male 31%, hypertension 21%, diabetes 3%, asthma 17%	NR	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	
WHO SOLIDARITY; ²⁸¹ Pan et al; Preprint; 2020	Patients with moderate to critical COVID-19 infection. 948 assigned to HCQ 800 mg once followed by 200 mg twice a day for 10 days and 900 assigned to SOC	Age range 50 – 69 43.5% years old, male 59.8%, diabetes 21.9%, COPD 6.9%, asthma 4.9%, CHD 14.1%	Steroids 20.9%, convalescent plasma 1.4%, Anti IL6 2.1%	Low for mortality and mechanical ventilation; some Concerns for symptom resolution, infection and adverse events Notes: Non-blinded study wich might have introduced bias to symptoms and adverse events outomes results.	
Davoodi et al; ²⁴⁸ peer-reviewed; 2020	Patients with moderate to severe COVID-19 infection. 30 assigned to febuxostat 80 mg per day and 30 assigned to hydroxychloroquine	Mean age 57.7 ± 8.4, male 59%, hypertension NR%, diabetes 27.8%, chronic lung disease 1.9%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
COVID-19 PEP (University of Washington) trial; Barnabas et al; ²⁸² Abstract; 2020	Individuals exposed to SARS-CoV-2 infection. 381 assigned to hydroxychloroquine 400 mg for three days followed by 200 mg for 11 days and 400 assigned to standard of care	male 40%	NR	Low for symptom resolution, infection, and adverse events	

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<u>PETAL trial</u> ; ²⁸³ Self et al; peer-reviewed; 2020	Patients with moderate to severe COVID-19. 242 assigned to hydroxychloroquine 800 mg on day 1 followed for 200 mg twice a day for 5 days and 237 assigned to standard of care	Median age 58.5 ± 24.5, male 56%, hypertension 52.8%, diabetes 34.6%, COPD 8.1%, asthma %, coronary heart disease %, chronic kidney disease 8.8%,	Corticosteroids 18.4%, remdesivir 21.7%, azithromycin 19%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>HAHPS trial</u> ; ²⁸⁴ Brown et al; peer- reviewed; 2020	Patients with moderate to critical COVID-19. 42 assigned to hydroxychloroquine 800 mg once followed by 200 mg twice a day for 5 days and 43 assigned to azithromycin	Median age 55 ± 23, male 61%, diabetes 26%, coronary heart disease 11%, chronic kidney disease 9%, cerebrovascular disease 8%, cancer 2%	Corticosteroids 15%, remdesivir 11%, lopinavir-ritonavir 1%, tocilizumab 24%, convalescent plasma 24%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Co-interventions were not balanced between study arms	
HYCOVID trial; ²⁸⁵ Dubee et al; peer reviewed; 2020	Patients with mild to moderate COVID-19. 124 assigned to hydroxychloroquine 800 mg once followed by 400 mg a day for 8 days and 123 assigned to standard of care	Median age 77 ± 28, male 48.4%, hypertension 53.4%, diabetes 17.3%, COPD 11.2%, cerebrovascular disease 17.3%, obesity 27.7%	Corticosteroids 9.6%, lopinavir-ritonavir 1.2%, azithromycin 8.4%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>Q-PROTECT</u> <u>trial</u> ; ²⁸⁶ Omrani et al; peer-reviewed; 2020	Patients with mild COVID-19. 152 assigned to hydroxychloroquine 600 mg daily for 7 days and 152 assigned to hydroxychloroquine + azithromycin	Mean age 41 ± 16, male 98.4%,	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>Dabbous et al;</u> ²⁸⁷ peer reviewed; 2020	Patients with mild to moderate COVID-19. 44 assigned to favipiravir 3200 mg once followed by 600	Mean age 35.5 ± 16.8, male 48.9%, comorbidities 18.4%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events	



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	mg twice a day for 10 days and 48 assigned to CQ			Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>HYDRA trial</u> ; ²⁸⁸ Hernandez- Cardenas et al; Preprint; 2020	Patients with severe to critical COVID-19. 106 assigned to hydroxychloroquine 400 mg a day for 10 days and 108 assigned to SOC	Mean age 49.6 ± 12, male 75%, hypertension 16%, diabetes 47%, CHD 11%, CKD 0%, obesity 66%	Corticosteroids 52.4%, lopinavir-ritonavir 30.4%, tocilizumab 2.5%, azithromycin 24.5%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
COVID-19 Early <u>Treatment trial</u> ; ²⁸⁹ Johnston et al; peer- reviewed; 2020	Patients with mild COVID-19. 60 assigned to hydroxychloroquine 800 mg once followed by 400 mg a day for 10 days, 65 assigned to HCQ + AZT 500 mg once followed by 250 mg a day for 5 days and 65 assigned to SOC	Median age 37 ±, male 43.3%, hypertension 20.9%, diabetes 11.6%, COPD 9.3%, asthma 1.6%, immunosuppressive therapy 0.8%, obesity 76%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>Purwati et al</u> ; ²⁹⁰ peer reviewed; 2020	Patients with mild to moderate COVID-19. 128 assigned to lopinavir-ritonavir 500/100 a day, 123 assigned to hydroxychloroquine 200 mg a day and 119 to SOC	Median age 36.5 ± NR, male 95.3%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
Beltran et al; ²⁹¹ peer reviewed; 2020	Patients with moderate to severe COVID-19. 33 assigned to hydroxychloroquine 800 mg once followed by 400 mg a day for 5	Mean age 54 ± 23.5, male 46.8%, hypertension 19.1%, diabetes 9.6%, COPD 1%, CHD 7.4%, cerebrovascular disease 5.3%	Corticosteroids 9.6%, lopinavir-ritonavir 44.7%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded





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	days and 37 assigned to SOC			study. Concealment of allocation is probably inappropriate.
<u>PATCH 1 trial</u> ; ²⁹² Amaravadi et al; preprint; 2020	Patients with mild COVID-19 infection. 17 assigned to hydroxychloroquine 400 mg a day and 17 assigned to SOC	Median age 53 ± 37, male 26%, hypertension 18%, diabetes 9%, , asthma 12%,	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>Bermejo Galan et</u> <u>al</u> ; ²⁹³ peer reviewed; 2021	critical COVID-19	Mean age 53.4 ± 15.6, male 58.2%, hypertension 43.4%, diabetes 28.1%, COPD 5.3%, CKD 2.5%, cancer 3%, obesity 37.5%	Corticosteroids 98%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>Seet et al</u> ; ²⁹⁴ peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 432 assigned to hydroxychloroquine 400 mg once followed by 200 mg a day for 42 days and 619 assigned to SOC (vitamin C)	Mean age 33, male 100%, hypertension 1%, diabetes 0.3%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
TOGETHER <u>trial</u> ; ²⁹⁵ Reis et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 214 assigned to hydroxychloroquine 800 mg once followed by 400 mg a day for 9 days and 227 assigned to SOC	Mean age 53, male 45%, hypertension 49.3%, diabetes 19.4%, COPD 2.5%, asthma 8.6%, CHD 3.9%, CKD 0.7%, cancer 1.2%, obesity 34.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events





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CLOROTRIAL trial; ²⁹⁶ Réa-Neto et al; peer reviewed; 2021		Median age 53 ±, male 66.7%, hypertension 38.1%, diabetes 25.7%, COPD 8.6%, immunosuppressive therapy 5.7%	Corticosteroids 72.4%, azithromycin 89.5%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>CHEER trial</u> ; ²⁹⁷ Syed et al; peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 154 assigned to hydroxychloroquine 200-400 mg once a week to three weeks and 46 assigned to SOC	Mean age 30.6 ± 8, male 54.5%, hypertension 4.5%, diabetes 3.5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
ProPAC-COVID trial; ²⁹⁸ Sivapalan et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 61 assigned to hydroxychloroquine + AZT 400 mg plus 500 to 250 mg a day and 56 assigned to SOC	Median age 65 ± 25, male 56%, hypertension 38%, diabetes 24%, COPD 9%, asthma 22%, CHD 7%, CKD 7%	Corticosteroids 32%, remdesivir 25%,	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	
Byakika-Kibwika et	moderate COVID-19	Median age 32 ± 27, male 72%, hypertension 2.8%, diabetes 2.8%, COPD %, CHD 0.9%,	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>ALBERTA HOPE-</u> <u>Covid19 trial</u> ; ³⁰⁰	Patients with mild COVID-19 infection.	Mean age 46.8 ± 11.2, male 55.4%,	NR	Low for mortality and mechanical ventilation;	



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Schwartz et al; peer reviewed; 2021	111 assigned to hydroxychloroquine 800 mg once followed by 400 mg for 5 days and 37 assigned to SOC	hypertension 27.8%, diabetes 19.6%, asthma 13.5%		Low for symptom resolution, infection, and adverse events	
HERO-HCQ trial _. ; ³⁰¹ Naggie et al ; preprint ; 2021	SARS-CoV-2 infection. 683 assigned to	Mean age 43.6 ± , male 44.7%, hypertension 14.6%, diabetes 4%, COPD 0.2%, asthma 9.9%, CHD 0.8%, obesity 33.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
Rodrigues et al; ³⁰² peer reviewed; 2021	Patients with mild COVID-19 infection. 42 assigned to hydroxychloroquine + azithromycin 400/500 mg a day for 7 days and 42 assigned to SOC	Mean age 36.5 ± 9.6, male 40.5%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>Babalola et al;</u> ³⁰³ preprint; 2021	Patients with mild to severe COVID-19 infection. 31 assigned to hydroxychloroquine + AZT 200/500 mg a day for 3 days and 30 assigned to SOC	Mean age 40.4 ± 1.9, male 63%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
FIGHT-COVID- <u>19 trial</u> ; ²³⁴ Atipornwanich et al; preprint; 2021	Patients with mild to severe COVID-19 infection. 320 assigned to favipiravir 6000 mg once followed by 2400 mg a day + lopinavir ritonavir 800/200 mg or lopinavir ritonavir	Mean age 42 ± 15.7, male 47.8%, obesity 24.6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of	





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	800/200 mg a day or hydroxychloroquine 800 mg a day or Darunavir ritonavir 1200/200 mg a day + hydroxychloroquine 400 mg a day or favipiravil 6000 mg followed by 2400 mg + darunavir ritonavir 1200/200 mg a day + HCQ 400 mg a day for 7 to 14 days.			allocation probably inappropriate.	
SEV-COVID <u>trial</u> ; ³⁰⁴ Panda et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 37 assigned to hydroxychloroquine 400 mg twice on first day followed by 400 mg per oral daily for 10 days + ribavirin (1.2 g orally as a loading dose followed by 600 mg orally every 12 hours) for 10 days and 40 assigned to SOC	Mean age 49.1, male 75%, hypertension 32.7%, diabetes 27.7%, COPD 7.9%, asthma %, CHD 11.9%, cancer 1%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>Ahmad et al</u> ; ³⁰⁵ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 100 assigned to hydroxychloroquine 800 once followed by 400 mg a day for 5 days or chloroquine 500 mg a day for 7 days and 50 assigned to SOC	Mean age 37.6, male 95.3%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
WHIP COVID-19 trial; ³⁰⁶ McKinnon et al; peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 398 assigned to	Mean age 44.9 ± 11.9, male 42%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and	

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	hydroxychloroquine 400 mg a week or 400 mg once followed by 200 mg a day and 200 assigned to SOC			adverse events	
)	Individuals exposed to SARS-CoV-2 infection. 62 assigned to hydroxychloroquine 200 mg a day for 60 days and 65 assigned to SOC	Mean age 31.1, male 42.5%, obesity 18.5%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
EPICOS trial; ³⁰⁸ Polo et al; preprint; 2021	Individuals exposed to SARS-CoV-2 infection. 231 assigned to hydroxychloroquine 200 mg a day and 223 assigned to SOC	Mean age 38, male 38.5%, hypertension 5%, diabetes 0.8%, COPD 0%, asthma 6.4%, CHD 0.7%, cancer 0.6%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	
<u>COPE – Coalition</u> <u>V trial</u> ; ³⁰⁹ Avezum et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 689 assigned to hydroxychloroquine 800 mg once followed by 400 mg a day for 7 days and 683 assigned to SOC	Median age 45 ± 20, male 46.9%, hypertension 53.4%, diabetes 16.2%, asthma 13%, CHD 3.4%, obesity 54.8%	Azithromycin 19%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
AlQahtani et al; ²⁴³ peer reviewed; 2021	Patients with moderate COVID-19 infection. 51 assigned to HCQ 800 mg once followed by 400 mg a day for 10 days and 52 assigned to SOC	Mean age 44, male 47.1%, diabetes 26.1%, COPD 7.6%, asthma %, CHD 1.3%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	





<u>Omehecatl trial</u> ; ³¹⁰ Roy-García et al; preprint; 2021	Patients with moderate COVID-19 infection. 61 assigned to HCQ 400 mg +/- AZT 500 mg a day for 5 days and 31 assigned to SOC	Mean age 37 ± , male 48.9%, commorbidities 27.2%	NR; Vaccinated 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>Tirupakuzhi et al</u> ; ³¹¹ peer reviewed; 2022	Individuals exposed to SARS-CoV-2 infection. 213 assigned to HCQ 800 mg once followed by 400 mg a week for 12 weeks and 203 assigned to SOC	male 52.6%,	Vaccinated 76.3%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
IRICT trial; ³¹² Elshafie et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 97 assigned to HCQ 400 mg once followed by 200 mg a day for 5 days and 102 assigned to SOC	Mean age 60, male 54.3%, hypertension 40.7%, diabetes 30.1%, CKD 10.6%, obesity 20.6%	Corticosteroids 100%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
<u>Choudhary et al</u> ; ³¹³ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 99 assigned to HCQ 1400 mg once followed by 600 mg a day for 5 days and 99 assigned to SOC	Mean age 43, male 48%, hypertension 24%, diabetes 3.5%, asthma 7.8%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.

Uncertainty in potential benefits and harms. Further research is needed.





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Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT				• •	
<u>Hadanny et al</u> ; ³¹⁴ preprint; 2021	Patients with severe to critical COVID-19 infection. 20 assigned to hyperbaric oxygen two sessions a day for 4 days and 9 assigned to SOC	Median age 65.4 ± 7.8, male 60%, hypertension 72%, diabetes 60%, COPD %, asthma 8%, CHD 24%, cancer 4%, obesity 8%	Corticosteroids 92%, tocilizumab 24%, convalescent plasma 80%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Blinding and concealment are probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very
<u>Cannellotto et al</u> , ³¹⁵ peer reviewed; 2021	Patients with severe COVID-19 infection. 20 assigned to hyperbaric oxygen 5 sesions (90 minutes duration each) and 20 assigned to SOC	Mean age 55.2 ± 9.2, male 65%, hypertension 32.5%, diabetes 17.5%, COPD 5%, asthma 5%, CHD %, CKD 5%, cancer 5%, obesity 35%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate. The study was stopped early for benefit.	<pre>ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: Not information</pre>
COVID-19-HBO trial; ³¹⁶ Kjellberg et al; preprint; 2021	Patients with severe COVID-19 infection. 15 assigned to hyperbaric oxygen 60 minutes at 2.4 ATA for up tp 5 sesions and 15 assigned to SOC	Mean age 64, male 56.6%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	

Hyperimmune anti-COVID-19 intravenous immunoglobulin (C-IVIG) Uncertainty in potential benefits and harms. Further research is needed.



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Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence		
RCT							
<u>Ali et al</u> ; ³¹⁷ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 40 assigned to C-IVIG 0.15- 0.3 g/kg once and 10 assigned to SOC	Mean age 56.5 ± 13.1, male 70%, hypertension 52%, diabetes 36%, COPD 10%, CHD 8%	Corticosteroids 100%, remdesivir 94%, tocilizumab 6%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information		
Parikh et al; ³¹⁸ preprint; 2021	Patients with moderate to severe COVID-19 infection. 30 assigned to C-IVIG 30 ml twice and 30 assigned to SOC	Mean age 52 ± 10.1, male 73.3%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No		
<u>ITAC trial;</u> <u>Polizzotto et al;</u> ³¹⁹ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 295 assigned to C- IVIG 400 mg/kg and 284 assigned to SOC	Mean age 59 ± 21, male 57%, hypertension 43%, diabetes 28%, COPD 7%, asthma 10%, CHD 5%, CKD 7%, immunosuppression 5%	Corticosteroids 56%; Vaccinated 2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	information Adverse events: Very low certainty ⊕○○○ Hospitalization: No		
<u>COVID-</u> <u>Compromise</u> <u>trial</u> ; ³²⁰ Huygens et al; preprint; 2021	Immunocompromised patients with moderate to severe COVID-19 infection. 10 assigned to C-IVIG 15 gr once and 8 assigned to IVIG		Corticosteroids 77.7%; Vaccinated 72.2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	information		
	Hypertonic saline (inhaled) Uncertainty in potential benefits and harms. Further research is needed.						



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					<u>.</u>
Delic et al; ¹⁰⁸ peer reviewed; 2022	Patients with critical COVID-19 infection. 42 assigned to hypertonic saline (inhaled) twice a day and 52 assigned to SOC	Mean age 65.7 , male 68%, hypertension 60.6%, diabetes 30.9%, CHD 7.4%, cerebrovascular disease 2.1%	Corticosteroids 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncerta	hzV inty in potential benefits a	SF-v13 and harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Prasenohadi et al</u> ; ³²¹ peer reviewed; 2022	Patients with moderate to severe	Mean age 50.8 ± , male 61.3%, obesity 22.6%	NR	Low for mortality and mechanical ventilation;	Mortality: Very low certainty ⊕○○○





	COVID-19 infection. 43 assigned to hzVSF- v13 200 to 400 mg once followed by two infusions of 100 to 200 mg and 19 assigned to SOC			low for symptom resolution, infection and adverse events	Invasive mechanical ventilation: No informationSymptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
	Uncerta	Ibr inty in potential benefits a	r utinib and harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
iNSPIRE trial; ³²² Coutre et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 22 assigned to ibrutinib 420 mg a day for 14 to 28 days and 24 assigned to SOC	Median age 51.5, male 70%, hypertension 39%, diabetes 43%, COPD 2%, asthma 9%, CHD 2%, CKD 4%, obesity 24%	Corticosteroids 63%, remdesivir 72%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty \oplus ()Invasive mechanical ventilation: Very low certainty \oplus ()Symptom resolution or





					<pre>improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information</pre>				
	Icatibant / iC1e/K Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence				
RCT				L					
Mansour et al, ³²³ preprint; 2020	Patients with moderate to severe COVID-19 infection. 10 assigned to icatibant 30 mg every 8 hours for 4 days, and 10 assigned to iC1e/K	Mean age 51.6 ± 11.5, male 53.3%, hypertension 50%, diabetes 46.7%, asthma 3.3%, obesity 43.3%	NR	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No				





	Uncertai	Icosap Inty in potential benefits a	Dent ethyl nd harms. Further resea	arch is needed.	information Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			•	•	
VASCEPA COVID-19 CARDIOLINK-9 trial; ³²⁴ kosmopoulos et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 46 assigned to icosapent ethyl 8 g a day for three days followed 4 g a day for 11 days and 49 assigned to SOC	NR	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
Imatinib <u>may not in</u>	crease sever <u>e adverse ev</u>		atinib 11b on other importan ou	tcomes are uncertain. Fur	ther research is needed
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE



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					certainty of the evidence
RCT	<u>.</u>				- -
COUNTER- COVID trial; ³²⁵ Aman et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 197 assigned to imatinib 800 mg once followed by 400 mg a day for 10 days and 188 assigned to SOC	Median age 64 ± 17, male 69%, hypertension 37.6%, diabetes 25%, COPD 18.4%, asthma 18%, CHD 22%, obesity 38%	Corticosteroids 72%, remdesivir 21%	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: RR 1.05 (95%CI 0.84 to 1.32); RD 0.5% (95%CI -1.6% to 3.3%); Low certainty \oplus \bigcirc \bigcirc Hospitalization: No information
	Uncerta	Indor inty in potential benefits a	nethacin and harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT				•	
<u>Ravichandran et</u> <u>al;</u> ³²⁶ preprint; 2021	Patients with moderate COVID-19	Mean age 47 ± 16, male 56.2%, hypertension	NR	High for mortality and mechanical ventilation;	Mortality: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$

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	infection. 102 assigned to indomethacin 75 mg a day and 108 assigned to SOC	19%, diabetes 29%		high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Invasive mechanical ventilation: Very low certainty⊕○○○Symptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
	Uncerta	Infl inty in potential benefits a	iximab Ind harms. Further resea	nrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>CATALYST</u> <u>trial</u> ; ³²⁷ Fisher et al; peer reviewed; 2021	Patients with moderate to critical COVID-19 infection. 29 assigned to infliximab and 34 assigned to SOC	Median age 64.5 ± 20, male 61.8%	Corticosteroids 94.3%, remdesivir 61.8%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty



					 ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
INM005 may n	INM005 ot improve symptom res		gments of equine case severe adverse event ler research is needed.	e antibodies) s. Its effects on other impo	ortant outcomes are
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	1	1		1	1
Lopardo et al; ³²⁸ peer reviewed; 2020	Patients with moderate to severe COVID-19. 118 assigned to INM005 4 mg/kg in two doses on days 1 and 3 and 123 assigned to SOC	Mean age 53.8 ± 12.5, male 65.1%, comorbidities 80%	Corticosteroids 57.2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Symptom resolution or improvement: RR 1.06 (95%Cl 0.96 to 1.66); RD 3.6% (95%Cl -2.4% to 10.3%); Low certainty $\bigoplus \bigcirc \bigcirc$ Symptomatic

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					(prophylaxis studies): No information Adverse events: RR 0.66 (95%Cl 0.37 to 1.18); RD -3.5% (95%Cl -6.4% to 1.8%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Hospitalization: No information
		erferon alpha-2b			
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	<u></u>			<u>.</u>	<u></u>
ESPERANZA trial; ³²⁹ Esquivel- Moynelo et al; preprint; 2020	Patients with mild to moderate COVID-19 infection. 30 assigned to interferon alpha-2b plus interferon gamma twice a week for two weeks (standard care) and 33 assigned to interferon alpha-2b three times a week (IM)	Median age 38 ± 63, male 54%, hypertension 22.2%, diabetes 4.7%, asthma 6.3%, coronary heart disease 6.3%, any comorbidities 50.8%	Hydroxychloroquine 100%, lopinavir- ritonavir 100%, antibiotics 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information

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					Hospitalization: No information				
Interferon beta-1a IFN beta-1a probably does not reduce mortality or invasive mechanical ventilation requirements. Inhaled interferon beta-1a may improve time to symptom resolution.									
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence				
RCT	•		•	•					
Davoudi-Monfared et al; ³³⁰ preprint; 2020	Patients with severe COVID-19 infection. 42 assigned to interferon beta-1a 44 µg subcutaneous, three times a week and 39 assigned to standard of care	asthma 1.2%, coronary	Corticosteroids 53%, hydroxychloroquine 97.5%, azithromycin 14.8%, ATB 81%, immunoglobulin 30.8%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: RR 0.99 (95%Cl 0.75 to 1.31); RD -0.2% (95%Cl -4% to 5%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ Invasive mechanical ventilation: RR 1.01 (95%Cl 0.87 to 1.18); RD 0.2% (95%Cl -2.2% to 3.1%); Moderate				
WHO SOLIDARITY trial; ²⁸¹ Pan et al; peer reviewed; 2020	Patients with moderate to critical COVID-19 infection. 2144 assigned to interferon beta-1a three doses over six days of 44µg and 2147 assigned to SOC	Age range 50-69 years old 46.3%, male 62.3%, diabetes 25.2%, COPD 5.4%, asthma 4.3%, CHD 22%	Steroids 58.7%, convalescent plasma 2.4%, Anti IL6 3.6%	Low for mortality and mechanical ventilation; some Concerns for symptom resolution, infection and adverse events Notes: Non-blinded study wich might have introduced bias to symptoms and adverse events outomes results.	certainty $\oplus \oplus \oplus \bigcirc$ Symptom resolution or improvement: RR 0.96 (95%CI 0.92 to 0.99); RD -2.6% (95%CI -4.8% to - 3.2%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ Symptomatic infection				
COVIFERON <u>trial</u> ; ³³¹ Darazam et al; Preprint; 2020	Patients with severe to critical COVID-19 infection. 20 assigned to interferon beta-1a 44 micrograms on days	Mean age 69 ± 27, male 51.7%, hypertension 33.3%, diabetes 23.3%, CHD 16.3%, CKD 8.3%, cancer 1.7%,	Hydroxychloroquine 100%, lopinavir- ritonavir 100%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events	(prophylaxis studies): Very low certainty ⊕○○○ Adverse events: RR				

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<mark>Darazam et al</mark> ; ³³² Preprint; 2020	1, 3 and 6, 20 assigned to interferon beta-1b 0.25 mg on days 1, 3 and 6 and 20 assigned to SOC Patients with severe to critical COVID-19. 85 assigned to interferon beta-1a 88 micrograms on days 1, 3 and 6 and 83 assigned to interferon beta-1a 44 micrograms on days 1, 3 and 6	Mean age 59.8 ± 16.5, male 61.9%, hypertension 37.3%, diabetes 26.8%, COPD 1.2%, asthma 1.8%, CHD 18.7%, CKD 8.3%, cerebrovascular disease 5.4%, cancer 0.6%	Corticosteroids 1.1%, lopinavir-ritonavir 100%	Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	1.03 (95%CI 0.85 to 1.24); RD 0.3% (95%CI -1.5% to 2.4%); Moderate certainty $\oplus \oplus \oplus$ Hospitalization: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
ACTT-3 trial; ³³³ Kalil et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 487 assigned to interferon beta-1a 44 µg a day for up to four days and 482 assigned to SOC	Mean age 58.7 ± 15.9, male 58%, hypertension 58%, diabetes 37%, COPD 11%, asthma 13%, CKD 12%, obesity 58%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>INTEREST trial</u> ; ³³⁴ Ranieri et al; peer reviewed; 2021	Patients with critical COVID-19 infection. 144 assigned to interferon beta-1a 10 μg a day for 6 days and 152 assigned to SOC	Mean age 58, male 65.8%	Corticosteroids 35.1%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>Castro-Rodriguez et</u> <u>al</u> ; ³³⁵ preprint; 2022	Individuals exposed to SARS-CoV-2 infection. 607 assigned to interferon beta-1a 125µg three time and 565 assigned to SOC	Mean age 34 ± , male 47.3%, diabetes 3.9%, COPD 0.1%, asthma 5.6%, CHD 5.1%, CKD 0.3%, cancer 1.2%	Corticosteroids %, Vaccinated 23.2%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	



				Significant loss to follow-up.	
Monk P et al; ³³⁶ et al; peer-reviewed; 2020	Patients with mild to severe COVID-19. 48 assigned to interferon beta-1a nebulized once a day for 15 days and 50 assigned to standard of care	Mean age 57.1 ± 13.2, male 59.2%, hypertension 54.7%, diabetes 22.6%, COPD 44.2%, asthma %, coronary heart disease 24.5%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or improvement: HR 2.19 (95%Cl 1.03 to 4.69); RD 26.4% (95%Cl 1.1% to 38.1%); Low certainty \oplus \bigcirc \bigcirc Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: No information
	Uncerta	Interfer inty in potential benefits a	on beta-1b	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Rahmani et al</u> ; ³³⁷ peer-reviewed; 2020	Patients with severe COVID-19. 33	Median age 60 ± 10.5, male 59%, hypertension	Corticosteroids 21.2%, ATB 51.5%, antivirals	High for mortality and invasive mechanical	Mortality: Very low certainty ⊕○○○





	assigned to interferon beta-1b 250 mcg subcutaneously every other day for two consecutive weeks and 33 assigned to standard of care	40.9%, diabetes 31.8%, chronic lung disease 4.5%, asthma NR%, coronary heart disease 30.3%, chronic kidney disease NR%, cerebrovascular disease NR%, immunosuppression NR%, cancer 3%, obesity NR%	100%	ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: Very low certainty ⊕○○○
<u>COVIFERON</u> <u>trial</u> ; ³³¹ Darazam et al; Preprint; 2020	Patients with severe to critical COVID-19 infection. 20 assigned to interferon beta-1a 44 micrograms on days 1, 3 and 6, 20 assigned to interferon beta-1b 0.25 mg on days 1, 3 and 6 and 20 assigned to SOC	Mean age 69 ± 27, male 51.7%, hypertension 33.3%, diabetes 23.3%, CHD 16.3%, CKD	Hydroxychloroquine 100%, lopinavir- ritonavir 100%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
<u>UW 20-535 trial</u> ; ³³⁸ Tam et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 51 assigned to interferon beta-1b 16 million IU a day for 5 days and 49 assigned to SOC	Mean age 65, male 52.8%, hypertension 42.3%, diabetes 22.6%, COPD %, asthma 3.8%, CHD 9.4%, CKD 4.2%, cerebrovascular disease 2.4%, cancer 8.5%, obesity 4.7%	Corticosteroids 29.2%, remdesivir 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
	Uncertai	Interfer inty in potential benefits a	On gamma nd harms. Further resea	nrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence





<u>Myasnikov et al</u> ; ³³⁹ Peer reviewed; 2021	Patients with moderate COVID-19 infection. 18 assigned to interferon gamma 500000 IU a day for 5 days and 18 assigned to SOC	Mean age 63 ± 12, male 44%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationSymptomatic informationAdverse events: No informationHospitalization: No information
	Uncertai	Interferon ka inty in potential benefits a	appa plus TFF2		
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Fu et al</u> ; ³⁴⁰ peer- reviewed; 2020	moderate COVID-19.	Mean age 35.2 ± 11.2, male 63.7%, hypertension 5%, diabetes 3.7%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: No information





					Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncertai	Inter inty in potential benefits a	leukin-2 and harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT		L			
STRUCK trial; ¹⁴⁵ Pimenta Bonifácio et al; preprint; 2021	critical COVID-19 infection. 14 assigned to IL-2 1.5 million IU	Mean age 48.9 ± 12.2, male 61.7%, hypertension 45%, diabetes 21.7%, COPD 6.7%, CHD 5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty





					⊕ ○○○ Hospitalization: No information
	Uncertai	Iota-ca inty in potential benefits a	rrageenan and harms. Further re	search is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
IVERCAR-TUC <u>trial</u> ; ³⁴¹ Chahla et al; Preprint; 2020	Individuals exposed to SARS-CoV-2 infection. 117 assigned to ivermectin + iota- carrageenan 12 mg a week + 6 sprays a day for 4 weeks and 117 assigned to SOC	Median age 38 ± 12.5, male 42.7%, hypertension 9%, diabetes, 7.3%, CKD 2.1%, obesity 11.9%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No
CARR-COV-02 trial; ³⁴² Figueroa et al; preprint; 2021	Individuals exposed to SARS-CoV-2 infection. 196 assigned to Iota-carrageenan 1 puff four times a day for 21 days and 198 assigned to SOC	Mean age 38.6 ± 9.6, male 24.8%, hypertension 4.8%, diabetes 0.2%, COPD 3.3%, cancer 0%, obesity 5%	NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	information Symptomatic infection (prophylaxis studies): Very low certainty \oplus \bigcirc \bigcirc Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: Very low certainty \oplus \bigcirc \bigcirc

Uncertainty in potential benefits and harms. Further research is needed.





Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Ojeda et al</u> ; ³⁴³ preprint; 2022	Patients with moderate to critical COVID-19 infection. 300 assigned to isothymol 6 mg until discharge and 300 assigned to SOC	Mean age 54, male 48.8%, hypertension 60.6%, diabetes 13.2%, asthma 24%, CHD 10.8%, CKD 5%, obesity 16.8%	Corticosteroids 12.5%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate. Unbalanced baseline risk (16% of included patients in intervention on mechanical ventilation vs. 9% in placebo).	Mortality: Very low certainty () () ()Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationoffection (prophylaxis studies): No informationAdverse events: No informationHospitalization: No information
	Uncertai	Itoli inty in potential benefits a	zumab nd harms. Further resea	urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>ITOLI-C19-02-I-00</u> <u>trial;³⁴⁴ Kumar et al;</u> preprint; 2020	COVID-19. 20	Mean age 49 ± 13, male 86.6%, hypertension 20%,	Nr	High for mortality and mechanical ventilation; high for symptom resolution, infection,	Mortality: Very low certainty ⊕○○○ Invasive mechanical



		•	•		
Ivermectin prol probably does	followed by 0.8 mg/kg weekly and 10 assigned to standard of care	ortality or improve time to ffect on hospitalizations a	•mectin o symptom resolution. In nd may not increase seve s when used as prophyla	and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Zagazig University <u>trial</u> ; ³⁴⁵ Shouman et al; peer-reviewed; 2020	Individuals exposed to SARS-CoV-2 infection. 203 assigned to ivermectin 15 to 24 mg and 101 assigned to standard of care	Mean age 38.72 ± 15.94, male 51.3%, hypertension 10.2%, diabetes 8.1%, CKD 1%, asthma 2.7%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded	Mortality: RR 1 (95%CI 0.8 to 1.24); RD -0% (95%CI - 3.2% to 3.8%); Moderate certainty ⊕⊕⊕○ Invasive mechanical



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<u>Chowdhury et al</u> ; ³⁴⁶ preprint; 2020	Patients with mild to moderate COVID-19. 60 assigned to ivermectin plus doxycycline 200 µgm/kg single dose + 100 mg BID for 10days and 56 assigned to hydroxychloroquine plus azithromycin	Mean age 33.9 ± 14.1, male 72.4%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	7.3% to 2.9%); Very Low certainty ⊕○○○ Symptom resolution or improvement: RR 1.04 (95%CI 0.98 to 1.1); RD 2.4% (95%CI -1.2% to 6%); Moderate certainty
Podder et al; ³⁴⁷ peer- reviewed; 2020	Patients with mild to moderate COVID-19. 32 assigned to ivermectin 200 µgm/kg once and 30 assigned to standard of care	Mean age 39.16 ± 12.07, male 71%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	<pre> ⊕⊕⊕○ Symptomatic infection (prophylaxis studies): RR 1.01 (95%CI 0.54 to 1.89); RD 0.2% (95%CI -8% to 15.5%); Very low certainty ⊕○○○ Adverse events: RR</pre>
<u>Hashim et a</u> l; ³⁴⁸ preprint; 2020	Patients with mild to critical COVID-19. 70 assigned to ivermectin plus doxycycline 200 µgm/kg two or three doses + 100 mg twice a day for 5 to 10 days and 70 assigned to standard of care	Mean age 48.7 ± 8.6, male %	Corticosteroids 100%, azithromycin 100%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	1.05 (95%CI 0.69 to 1.62); RD 0.5% (95%CI -3.2% to 6.3%); Low certainty ⊕⊕○○ Hospitalization: RR 0.90 (95%CI 0.74 to 1.1); RD -0.5% (95%CI -1.2% to 0.5%); Moderate certainty ⊕⊕⊕
<u>Mahmud et al</u> ; ³⁴⁹ peer-reviewed; 2020	Patients with mild to moderate COVID-19. 183 assigned to ivermectin plus doxycycline 12 mg once + 100 mg twice a day for 5 days and 180 assigned to standard of care	Mean age 39.6 ± 13.2, male 58.8%,	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events. Notes: 8% of patients were lost to follow-up.	certainty ⊕⊕⊕⊖





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Elgazzar et al (mild); ³⁵⁰ preprint (now retracted); 2020	Patients with mild to moderate COVID-19. 100 assigned to ivermectin 400 µgm/kg once for 4 days and 100 assigned to hydroxychloroquine	Mean age 55.2 ± 19.8, male 69.5%, hypertension 11.5%, diabetes 14.5%, COPD %, asthma 5.5%, coronary heart disease 4%, chronic kidney disease %	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Elgazzar et al</u> (severe); ³⁵⁰ preprint (now retracted); 2020	Patients with severe COVID-19. 100 assigned to ivermectin 400 µgm/kg once for 4 days and 100 assigned to hydroxychloroquine	Mean age 58.9 ± 19.5, male 71%, hypertension 16%, diabetes 20%, COPD %, asthma 13%, coronary heart disease 7.5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Elgazzar et al</u> (prophylaxis); ³⁵⁰ preprint (now retracted); 2020	Individuals exposed to SARS-CoV-2 infection. 100 assigned to ivermectin 400 µgm/kg twice (second dose after one week) and 100 assigned to standard of care	NR	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Krolewiecki et al</u> , ³⁵¹ peer-reviewed; 2020		Mean age 40.2 ± 12, male 55.5%, hypertension 13.3%, diabetes 15.5%, COPD 11.1%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	



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<u>Niace et al</u> ; ³⁵² preprint; 2020	Patients with mild to severe COVID-19. 120 assigned to ivermectin 200-800 microg/kg and 60 assigned to standard of care	Median age 67 ± 22, male 50%	NR	Some concerns for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Concealment of allocation possibly inappropriate.	
<u>Ahmed et al</u> ; ³⁵³ peer-reviewed; 2020	Patients with mild COVID-19. 55 assigned to ivermectin 12 mg a day for 5 days +/- doxycycline and 23 assigned to standard of care	Mean age 42, male 46%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	
<u>SAINT trial</u> ; ³⁵⁴ Chaccour et al; peer-reviewed; 2020	Patients mild (early within 3 days of onset) COVID-19. 12 assigned to ivermectin 400 microg/kg and 12 assigned to SOC	Median age 26 ± 36, male 50%,	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>Cachar et al</u> ; ³⁵⁵ peer- reviewed; 2020	Patients with mild COVID-19. 25 assigned to ivermectin 36 mg once and 25 assigned to SOC	Mean age 40.6 ± 17, male 62%, hypertension 26%, diabetes 40%, obesity 12%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Babalola et al;</u> ³⁵⁶ peer-reviewed; 2020	Patients with mild to moderate COVID-19 infection. 42 assigned to ivermectin 12 to 24 mg a week for 2	Mean age 44.1 ± 14.7, male 69.4%, hypertension 14.5%, diabetes 3.2%,	Corticosteroids 3.2%,	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	





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	weeks and 20 assigned to lopinavir-ritonavir			
	Patients with mild to moderate COVID-19. 55 assigned to ivermectin 24 mg divided in two doses and 57 assigned to SOC	Mean age 52.5 ± 14.7, male 72.3%, hypertension 34.8%, diabetes 35.7%, COPD 0.9%, asthma 0.9%, CHD 8.9%, CKD 2.7%, cerebrovascular disease 0%, cancer 5.4%, obesity %	Corticosteroids 100%, remdesivir 20.5%, hydroxychloroquine 100%, tocilizumab 6.3%, convalescent plasma 13.4%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>IVERCAR-TUC</u> <u>trial</u> ; ³⁴¹ Chahla et al; Preprint; 2020	Individuals exposed to SARS-CoV-2 infection. 117 assigned to ivermectin + iota- carrageenan 12 mg a week + 6 sprays a day for 4 weeks and 117 assigned to SOC	Median age 38 ± 12.5, male 42.7%, hypertension 9%, diabetes, 7.3%, CKD 2.1%, obesity 11.9%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>Mohan et al</u> ; ³⁵⁸ preprint; 2020	Patients with mild to moderate COVID-19 infection. 80 assigned to ivermectin 12 to 24 mg once and 45 assigned to SOC	Mean age 35.3 ± 10.4, male 88.8%, hypertension 11.2%, diabetes 8.8%, CHD 0.8%,	Corticosteroids 14.4%, remdesivir 1.6%, hydroxychloroquine 4%, azithromycin 11.2%,	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events
<u>Shahbaznejad et</u> <u>al;</u> ³⁵⁹ peer-reviewed; 2020	Patients with moderate to severe COVID-19 infection. 35 assigned to ivermectin 0.2 mg/kg once and 34 assigned to SOC	Mean age 46.4 ± 22.5, male 50.7%	Chloroquine 75.4%, lopinavir-ritonavir 79.7%, azithromycin 57.9%,	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events
<u>Spoorthi et al</u> ; ³⁶⁰ Unpublished; 2020	Patients with mild to moderate COVID-19 assigned to ivermectin 0.2 mg/kg once or SOC	NR	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events





				Notes: Non-blinded study. Concealment of allocation is probably inappropriate. RoB assessment from secondary sources as publication not available.	
<u>Samaha et al</u> ; ³⁶¹ peer-reviewed (now retracted); 2020	Patients with mild (asymptomatic) COVID-19 infection. 50 assigned to ivermectin 9 to 12 mg or 150 µg/kg once and 50 assigned to SOC	Mean age 31.6 ± 7.7, male 50%, hypertension 8%, diabetes 6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Randomization process and concealment of allocation is probably inappropriate.	
<u>Bukhari et al</u> ; ³⁶² Preprint; 2020	Patients with mild to moderate COVID-19. 45 assigned to ivermectin 12 mg once and 41 assigned to SOC	NR	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Okumus et al;</u> ³⁶³ peer-reviewed; 2021	Patients with severe COVID-19. 30 assigned to ivermectin 0.2 mg/kg for 5 days and 30 assigned to SOC	Mean age 62 ± 12, male 66%, hypertension 21.6%, diabetes 45%, COPD 1.6%, CHD 1.6%, cancer 1.6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Beltran et al;</u> ²⁹¹ peer reviewed; 2021	Patients with moderate to severe	Mean age 54 ± 23.5, male 46.8%,	Corticosteroids 9.6%, lopinavir-ritonavir	High for mortality and mechanical ventilation;	





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	COVID-19. 36 assigned to ivermectin 12–18 mg once and 37 assigned to SOC	hypertension 19.1%, diabetes 9.6%, COPD 1%, CHD 7.4%, cerebrovascular disease 5.3%	44.7%	high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	
Lopez-Medina et al; ³⁶⁴ peer-reviewed; 2021	Patients with mild to moderate COVID-19 infection. 200 assigned to ivermectin 300 µg/kg a day for 5 days and 198 assigned to SOC	Median age 37 ± 19, male 42%, hypertension 13.4%, diabetes 5.5%, COPD 3%, CHD 1.7%, cancer %, obesity 18.9%	Corticosteroids 4.5%	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	
<u>Bermejo Galan et</u> <u>al</u> ; ²⁹³ peer-reviewed; 2021	Patients with severe to critical COVID-19 infection. 53 assigned to ivermectin 42 mg and 115 assigned to HCQ or CQ	Mean age 53.4 ± 15.6, male 58.2%, hypertension 43.4%, diabetes 28.1%, COPD 5.3%, CKD 2.5%, cancer 3%, obesity 37.5%	Corticosteroids 98%	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	
Pott-Junior et al; ³⁶⁵ peer-reviewed (now retracted); 2021	Patients with moderate to critical COVID-19 infection. 27 assigned to ivermectin 100 to 400 mcg/kg and 4 assigned to SOC	Mean age 49.4 ± 14.6, male 45.2%	Corticosteroids 32.3%,	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Kishoria et al</u> ; ³⁶⁶ peer-reviewed; 2021	Patients with moderate to severe COVID-19 infection. 19 assigned to ivermectin 12 mg and 16 assigned to SOC	Mean age 38, male 66%	Hydroxychloroquine 100%	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse	



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				events outcomes results.	
<u>Seet et al</u> , ²⁹⁴ peer- reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 617 assigned to ivermectin 12 mg once and 619 assigned to SOC (vitamin C)	Mean age 33, male 100%, hypertension 1%, diabetes 0.3%	NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Abd-Elsalam et al</u> ; ³⁶⁷ peer-reviewed; 2021		Mean age 40.8 ± 16.5, male 50%, hypertension 19.5%, diabetes 16.4%	NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Biber et al;</u> ³⁶⁸ peer- reviewed; 2021	Patients with mild recent onset COVID- 19 infection. 47 assigned to ivermectin 48 to 55 mg administered for three days and 42 assigned to SOC	Mean age 35 ± 19, male 78.4%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes: 5.2% of patients lost to follow-up.	
Faisal et al; ³⁶⁹ peer- reviewed; 2021	Patients with mild COVID-19 infection. 50 assigned to ivermectin 12 mg a day for 5 days and 50 assigned to SOC	Mean age 46 ± 3, male 80%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably	





				inappropriate.
<u>Vallejos et al</u> ; ³⁷⁰ peer reviewed; 2021	Patients with mild COVID-19 infection. 250 assigned to ivermectin 24-36 mg and 251 assigned to SOC	Mean age 42.5 ± 15.5, male 52.7%, hypertension 23.8%, diabetes 9.6%, COPD 2.8%, asthma 7.2%, CHD 1.8%, cancer 1.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>COVER trial</u> ; ³⁷¹ Buonfrate et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 61 assigned to ivermectin 600 to 1200 µg/kg once a day for 5 days and 32 assigned to SOC	Median age 47 ± 27, male 58.1%, diabetes 9.7%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>Manomaipiboon et</u> <u>al;</u> ³⁷² preprint; 2021	Patients with mild COVID-19 infection. 36 assigned to ivermectin 12 mg a day for 5 days and 36 assigned to SOC	Mean age 48.6 ± 14.8, male 37.5%, hypertension 40.3%, diabetes 23.6%, CHD 2.8%, CKD 6.9%, cerebrovascular disease 2.8%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events
<u>I-TECH trial</u> ; ³⁷³ Chee Loon Lim et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 241 assigned to ivermectin 6 to 12 mg a day for 5 days and 249 assigned to SOC	Mean age 62.5, male 49.5%, hypertension 82%, diabetes 58.2%, COPD 8.4%, CHD 12.6%, CKD 15.7%, cerebrovascular disease 4.2%, immunosuppressive therapy 0.2%, cancer 3.1%, obesity 26%	Corticosteroids 28.9%, tocilizumab 0.9%, Baricitinib 2.4%; Vaccinated 56.4%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>TOGHETER</u> <u>trial</u> ; ³⁷⁴ Reis et al; peer reviewed; 2021	Patients with recent onset mild COVID-19 infection. 679 assigned to ivermectin 400	Median age 49, male 41.8%, hypertension 8.4%, diabetes 12.9%, COPD 3%, asthma	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and

	μg/kg once a day for 3 days and 679 assigned to SOC	8.4%, CHD 1.8%, CKD 0.5%, obesity 49.7%		adverse events
	Patients with mild COVID-19 infection. 33 assigned to ivermectin and 33 assigned to soc	Mean age 38.5 ± 14.6, male 27.3%, hypertension 8.9%, diabetes 5.3%, CHD 7.1%, CKD 1.8%, obesity 19.6%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
Cruz Arteaga et al; <u>NCT04673214</u> ; other; 2021	Patients with mild COVID-19 infection. 65 assigned to ivermectin adjusted to body weight and 46 assigned to SOC	Age (18 – 65 years old) 96.4% , male 47.7%,	NR	NA
ACTIV-6 trial; ³⁷⁶ Naggie et al; preprint; 2022	Patients with mild to moderate COVID-19 infection. 817 assigned to ivermectin 400 µg/kg for three days and 774 assigned to SOC	Median age 47, male 46.6%, diabetes 11.8%, COPD 3.65%, asthma 15.5%, CHD 4.5%, CKD 0.77%, cancer 3.02%, obesity 40.8%	Remdesivir 0.3%, Vaccinated 48.8%	Low for mortality and mechanical ventilation; Low for symptom resolution, infection and adverse events
<u>Rezai_Mild trial;</u> ³⁷⁷ Rezai et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 268 assigned to ivermectin 0.4 mg/kg a day for 3 days and 281 assigned to SOC	Mean age 35.4 ± 17.4, male 53.4%, hypertension 7.8%, diabetes 7.3%, asthma 2.4%, CHD 2.7%, cancer 0.6%, obesity 21.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:
<u>Rezai_Severe</u> <u>trial</u> ; ³⁷⁷ Rezai et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 311 assigned to ivermectin 0.4 mg/kg a day for 3 days and 298 assigned to SOC	Mean age 53.8, male 47.8%, hypertension 28.4%, diabetes 31.7%, COPD %, asthma 3%, CHD 12.2%, obesity 73.3%	Corticosteroids 90.7%, remdesivir 98.2%, hydroxychloroquine 35%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Significant loss to follow up.
<u>Angkasekwinai</u> <u>treatement trial</u> ; ³⁷⁸ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 233 assigned	Mean age 39.5 ± 12.1, male 43.2%, hypertension 11.2%,	Vaccinated 74.9%	Low for mortality and mechanical ventilation; low for symptom



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	to ivermectin 400–600 μg/kg/d and 214 assigned to SOC	diabetes 6.9%, COPD 0.2%, CHD 1.8%, CKD 0.4%, cerebrovascular disease 0.2%, cancer 0.2%,		resolution, infection and adverse events Notes:
<u>Angkasekwinai</u> <u>prevention trial</u> ; ³⁷⁸ peer reviewed; 2022	Individuals exposed to SARS-CoV-2 infection. 259 assigned to ivermectin 400–600 µg/kg/d and 277 assigned to SOC	Mean age 37.6 ± 12, male 42.2%, hypertension 8.8%, diabetes 4.7%, COPD 0.2%, CHD 1.1%, cerebrovascular disease 0.4%, cancer 1.3%	Vaccinated 84.1%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:
<u>Mirahmadizadeh et</u> <u>al</u> ; ³⁷⁹ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 261 assigned to ivermectin 12 to 24 mg once and 130 assigned to SOC	Mean age 39.3, male 53.9%, hypertension 6.1%, diabetes 3.8%, COPD 0.8%, CHD 0.8%, CKD 0.5%, cancer 0.3%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:
<u>George et al</u> ; ³⁸⁰ peer reviewed; 2022	Patients with hematological disorders and mild to moderate COVID-19 infection. 73 assigned to ivermectin 12 to 24 mg once and 39 assigned to SOC	Mean age 41.2 ± , male 70.5%, cancer 75.9%	Corticosteroids 62.5%, remdesivir 18.7%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>PLATCOV - Iver</u> <u>trial</u> ; ³⁸¹ Schilling et al; peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 45 assigned to ivermectin 600µg/kg daily for seven days and 41 assigned to SOC	Mean age 28, male 45.5%	NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>IRICT trial</u> ; ³¹² Elshafie et al; peer	Patients with moderate to severe	Mean age 59.4 ± , male 53.4%, hypertension	Corticosteroids 100%	Low for mortality and mechanical ventilation;



reviewed; 2022	COVID-19 infection. 104 assigned to ivermectin 36 mg on days 1, 3 and 6 and 102 assigned to SOC	38.3%, diabetes 27.7%, CKD 9.2%, obesity 19.9%		low for symptom resolution, infection and adverse events	
Nimitvilai et al; ³⁸² peer reviewed; 2022	Patients with mild COVID-19 infection. 57 assigned to ivermectin 0.6 mg/kg for 3 days and 56 assigned to HCQ 200 mg a day + darunavir/ritonavir 400/100 mg a day for 5 days	Mean age 40, male 45.1%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
COVID-OUT trial; ²⁵⁴ Bramante et al; peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 410 assigned to Ivermectin 390 to 470 µg/kg a day for 3 days and 398 assigned to SOC	Median age 45.5, male 45.3%, hypertension 22.8%, diabetes 1.6%, obesity 47.4%	monoclonal antibodies	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
	Uncertai	Ivermect inty in potential benefits a	in (inhaled) Ind harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Aref et al</u> ; ³⁸³ peer reviewed; 2021	Patients with mild COVID-19 infection. 57 assigned to inhaled (inh) ivermectin and 57 assigned to SOC	Mean age 45 ± 19, male 71.9%, hypertension 17.5%, diabetes 12.3%, COPD 0.9%, cerebrovascular disease 3.5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Randomization	Mortality: No information Invasive mechanical ventilation: No information





		Intravenous imm	unoglobulin (IV	and concealment of allocation is probably inappropriate.	Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
		inty in potential benefits a			
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT				•	
<u>Sakoulas et al</u> ; ³⁸⁴ preprint; 2020	Patients with severe COVID-19 infection. 16 assigned to IVIG 0.5 g/kg/day for 3 days and 17 assigned to standard of care	Mean age 54 ± NR, male 60.6%, hypertension 33.3%, diabetes 36.3%, chronic lung disease 12%, coronary heart disease 3%, chronic kidney disease 3%, immunosuppression 3%	Corticosteroids 78.7%, remdesivir 51.5%, convalescent plasma 15.2%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No
<u>Gharebaghi et al</u> ; ³⁸⁵ preprint; 2020	Patients with severe to critical COVID-19. 30 assigned to IVIG 5 g a day for 3 days and 29	Mean age 56 ± 16, male 69.5%, hypertension 22%, diabetes 27.1%, chronic lung disease	NR	Some concerns for mortality and invasive mechanical ventilation; some concerns for	information Symptomatic infection



Tabarsi et al; ³⁸⁶ peer-reviewed; 2020 Raman et al; ³⁸⁷ Peer reviewed; 2020	assigned to standard of care Patients with severe COVID-19. 52 assigned to IVIG 400 mg/kg daily for three doses and 32 assigned to standard of care Patients with moderate to severe COVID-19. 50 assigned to IVIG 0.4 g/kg for 5 days and 50 assigned to SOC	3.3%, Mean age 53 ± 13, male 77.4%, hypertension 20.2%, diabetes 21.4%, COPD 1.2%, asthma %, coronary heart disease %, chronic kidney disease 4.7%, cancer 1.2%, Mean age 48.7 ± 12, male 33%, hypertension 31%, obesity 16%	NR	symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate. High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate. High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of	(prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
				allocation is probably inappropriate.	
	Uncertai	Ixek inty in potential benefits a	izumab Ind harms. Further resea	urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>STRUCK trial</u> ; ¹⁴⁵ Pimenta Bonifácio et al; preprint; 2021	Patients with severe to critical COVID-19 infection. 16 assigned to ixekizumab 80 mg once and 16 assigned	Mean age 48.9 ± 12.2, male 61.7%, hypertension 45%, diabetes 21.7%, COPD 6.7%, CHD 5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No



	to SOC	KB109 (microb inty in potential benefits a	iome modificate		informationSymptom resolution or improvement: Very low certainty
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Haran et al</u> , ³⁸⁸ preprint; 2021	Patients with mild to moderate COVID-19 infection. 169 assigned to KB109 9-36 g twice a day for 14 days and 172 assigned to SOC		NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic

Haran et al; ³⁸⁸	Patients with mild to	Median age 36 ± 56,	NR	Low for mortality and	Mortality: Very low
preprint; 2021	moderate COVID-19	male 40.8%,		mechanical ventilation;	certainty $\oplus \bigcirc \bigcirc \bigcirc$
-	infection. 169 assigned	hypertension 18%,		High for symptom	
	to KB109 9-36 g twice	diabetes 2.5%, COPD		resolution, infection,	Invasive mechanica
	a day for 14 days and	8.8%, cerebrovascular		and adverse events	ventilation: No
	172 assigned to SOC	disease 2.3%, cancer			information
		0.8%, obesity 3.7%		Notes: Non-blinded	
				study which might have	Symptom
				introduced bias to	resolution or
				symptoms and adverse	improvement: Very
				events outcomes results.	low certainty
					$\oplus 000$
					Symptomatic

					<pre>infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information</pre>
	Uncerta	<i>L-a</i> L-a inty in potential benefits a	r ginine nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Coppola et al; ³⁸⁹ peer reviewed; 2021	COVID-19 infection. 45 assigned to L-	Mean age 61.6, male 81.2%, hypertension 36.7%, diabetes 10%, CHD 14.5%, obesity 10%	Corticosteroids 100%, remdesivir 27.8%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
PROBCO trial; ³⁹⁰ Endam et al; preprint; 2021	Patients with mild recently diagnosed COVID-19 infection. 12 assigned to <i>Lactococcus lactis</i> (intranasal) two nasal irrigations a day and 11 assigned to SOC	Mean age 30.4 ± 9.1, male 30%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncerta		actoferrin its and harms. Further	research is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence



<u>Algahtani et al</u> ; ³⁹¹ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 36 assigned to lactoferrin 200 to 400 mg a day and 18 assigned to SOC	Mean age 48.6, male 60.3%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncerta	Leflu inty in potential benefits a	Inomide Ind harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	•			•	
<u>Hu et al</u> ; ³⁹² peer- reviewed; 2020	Patients with mild to critical COVID-19 infection. 5 assigned to Leflunomide 50 mg every 12 h (three doses) followed by 20 mg a day for 10 days and 5 assigned to standard of care	Mean age 52.5 ± 11.5, male 30%, hypertension 60%, chronic lung disease 10%	Umifenovir 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No





	Uncertai	Lev: inty in potential benefits a	amisole nd harms. Further rese	arch is needed.	infection (prophylaxis studies): No information Adverse events: RR 0.82 (95%CI 0.62 to 1.07); RD -1.8% (95%CI -3.9% to 0.7%); Low certainty ⊕⊕⊕○ Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Roostaei et al</u> ; ³⁹⁵ Preprint; 2020	Patients with mild to moderate COVID-19. 25 assigned to levamisole 150 mg a day for 3 days and 25 assigned to SOC	Mean age 36.6 ± 13.7, male 60%,	Hydroxychloroquine 100%,	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or
<u>Asgardoon et al</u> ; ^{3%} preprint; 2021	Patients with mild to moderate COVID-19 infection. 185 assigned to levamisole 50 mg a day for 10 days and 180 assigned to SOC	Median age 40 ± 18.75, male 56.1%, hypertension 8.8%, diabetes 9.4%, CHD 1.6%	Hydroxychloroquine 11.2%,	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	<pre>improvement: Mortality: Very low certainty ⊕○○○</pre> Symptomatic infection (prophylaxis studies): No information Adverse events: No

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					information
					Hospitalization: Very low certainty ⊕○○○
					Hospitalization: No information
Levilimab may imp	rove time to symptom re	solution; however, the ce	vilimab rtainty of the evidence w 1. Further research is new	as low. The effects of levilined and the second s	nab on other importar
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	1				
CORONA trial; ³⁹⁷ Lomakin et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 103 assigned to levilimab 364 mg once (subcutaneous) and 103 assigned to SOC	Mean age 58.3 ± 11.8, male 52.9%, CHD 15.5%,	Corticosteroids 7.3%, hydroxychloroquine 67.4%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanica ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Symptom resolution or improvement: Mortality: RR 1.48 (95%CI 1.13 to 1.93) RD 29.1% (95%CI - 7.9% to 56.4%); Low certainty $\bigoplus \bigcirc \bigcirc$
					Symptomatic infection (prophylaxis studies): No information
					Adverse events: No information



					Hospitalization: No information
	Uncerta	Lin: inty in potential benefits :	agliptin and harms. Further resea	urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	1				
<u>Abuhasira et al</u> ; ³⁹⁸ peer reviewed; 2021	Patients with moderate to severe with diabetes COVID- 19 infection. 32 assigned to linagliptin 5 mg a day and 32 assigned to SOC	Mean age 66.9 ± 13.9, male 59.4%, diabetes 100%,	Corticosteroids 82.8%, remdesivir 50%, convalescent plasma 10.9%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty $\oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Symptom resolution or improvement: No
<u>Covid19DPP4i</u> <u>trial</u> ; ³⁹⁹ Guardado- Mendoza et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 34 assigned to linagliptin 5 mg a day and 35 assigned to SOC	Mean age 58.5, male 63.7%, hypertension %, diabetes 66.6%, CHD 5.8%, CKD 14.5%, cerebrovascular disease 2.9%,	Corticosteroids 100%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information

Uncertainty in potential benefits and harms. Further research is needed.



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Guvenmez et al</u> ; ⁷⁶ peer-reviewed; 2020	Patients with moderate COVID-19 infection. 12 assigned to lincomycin 600 mg twice a day for 5 days and 12 assigned to azithromycin 500 mg on first day followed by 250 mg a day for 5 days	Mean age 58.7 ± 16, male 70.8%,	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
		T:	thium		
	Uncerta	inty in potential benefits a		arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	·	·	·	·	
<u>Spuch et al</u> , ⁴⁰⁰ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 15 assigned to lithium 400 mg a	Mean age 58.6, male 56.7%, hypertension 30%, diabetes 3.3%, COPD %, CHD 6.7%,	Corticosteroids 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and	Mortality: Very low certainty ⊕○○○ Invasive mechanical





	day and 15 assigned to SOC		ir-ritonavir	adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	ventilation: No informationSymptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
Lopinavir-ritonavi		uce mortality with moder:	ate certainty. Lopinavir-	ritonavir may not be assoc f risk of bias and imprecisi	
Lopinavir-ritonavi Study; publication status		uce mortality with moder:	ate certainty. Lopinavir-		
Study; publication	increase in severe adver Patients and interventions	uce mortality with modera se events. However, the c	ate certainty. Lopinavir- ertainty is low because o Additional	f risk of bias and imprecisi Risk of bias and study	Interventions effects vs standard of care and GRADE certainty of the



ELACOI trial; ⁴⁰² Li et al; peer-reviewed; 2020	Patients with moderate to severe COVID-19 infection. 34 assigned to lopinavir-ritonavir 200/50 mg twice daily for 7-14 days, 35 assigned to umifenovir and 17 assigned to standard of care	Mean age 49.4 ± 14.7, male 41.7%	Corticosteroids 12.5%, intravenous immunoglobulin 6.3%	Low for mortality and invasive mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	certainty ⊕⊕⊕⊕ Symptom resolution or improvement: RR 1.03 (95%CI 0.92 to 1.15); RD 1.8% (95%CI -4.8% to 9%); Moderate certainty ⊕⊕⊕○ Symptomatic infection
RECOVERY - Lopinavir-ritonavir trial; ⁴⁰³ Horby et al; other; 2020	Patients with mild to critical COVID-19 infection. 1616 assigned to lopinavir- ritonavir 400/100 mg twice a day for 10 days and 3424 assigned to standard of care	Mean age 66.2 ± 15.9, male 60.5%, diabetes 27.5%, chronic lung disease 23.5%, coronary heart disease 26%	NR	Low for mortality and invasive mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	(prophylaxis studies): Very low certainty $\bigoplus \bigcirc \bigcirc$ Severe Adverse events: RR 0.6 (95%CI 0.37 to 0.98); RD -4.1% (95%CI - 6.5% to -0.2%); Low certainty $\bigoplus \bigcirc \bigcirc$ Hospitalization:
<u>Huang et al</u> ; peer- reviewed; ²⁶³ 2020	Patients with moderate to severe COVID-19 infection. 10 assigned to CQ 500 mg twice a day for 10 days and 12 assigned to lopinavir-ritonavir 400/100 mg twice a day for 10 days	Mean age 44 ± 21, male 59.1%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Very low certainty ⊕○○○
<u>Zheng et al;</u> preprint; ⁴⁰⁴ 2020	Patients with moderate to severe COVID-19 infection. 30 assigned to novaferon 40 microg twice a day (inh), 30 assigned to novaferon	Median age 44.5 ± NR, male 47.1%	NR	High for mortality and invasive mechanical ventilation; High for symptom resolution, infection, and adverse events	



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	plus lopinavir- ritonavir 40 mg twice a day (inh) + 400/100 mg a day and 29 assigned to lopinavir- ritonavir			Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Chen et al;</u> preprint; ⁴⁰⁵ 2020	Patients with mild to moderate COVID-19 infection. 33 assigned to ribavirin 2 g IV loading dose followed by orally 400-600 mg every 8 hours for 14 days, 36 assigned to lopinavir-ritonavir and 32 assigned to ribavirin plus lopinavir- ritonavir	Mean age 42.5 ± 11.5, male 45.5%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
WHO SOLIDARITY trial; ²⁸¹ Pan et al; peer reviewed; 2020	Patients with moderate to critical COVID-19 infection. 1404 assigned to lopinavir-ritonavir 200/50MG twice a day for 14 days and 1368 assigned to SOC	Age range 50-69 years old 43.1%, male 59.6%, diabetes 24.2%, COPD 6.5%, asthma 4.9%, CHD 21%	Steroids 27.2%, convalescent plasma 1.4%, anti IL6 3%	Low for mortality and mechanical ventilation; some Concerns for symptom resolution, infection and adverse events Notes: Non-blinded study wich might have introduced bias to symptoms and adverse events outomes results.	
Sali et al; ⁴⁰⁶ Peer reviewed; 2020	Patients with moderate to severe COVID-19. 22 assigned to sofosbuvir 400 mg a day and 32 assigned to lopinavir- ritonavir 400/100 mg every 12 hours	Mean age 56.5 ± 14, male 53.7%, diabetes 33%,	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	



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<u>Purwati et al</u> ; ⁴⁰⁷ Peer reviewed; 2020	Patients with mild to moderate COVID-19. 128 assigned to lopinavir-ritonavir 500/100 a day, 123 assigned to HCQ 200 mg a day and 119 to SOC	Median age 36.5 ± NR, male 95.3%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Kasgari et al</u> ; ⁴⁰⁸ peer- reviewed; 2020	Patients with moderate COVID-19 infection. 24 assigned to sofosbuvir/daclatasvir 400/60 mg twice daily and 24 assigned to hydroxychloroquine plus lopinavir- ritonavir	Median age 52.5 ± NR, male 37.5%, hypertension 35.4%, diabetes 37.5%, chronic lung disease 2%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Yadollahzadeh et</u> <u>al;</u> ⁴⁰⁹ Preprint; 2021	Patients with mild to moderate COVID-19 infection. 58 assigned to sofosbuvir/ daclatasvir 400/60 mg a day for 10 days and 54 assigned to lopinavir-ritonavir 400/100 mg twice a day for 7 days	Mean age 57.4 ± 15, male 44.6%, hypertension 25%, diabetes 21.4%, COPD 3.6%, CHD 15.2%, CKD 6.2%, immunosuppression 3.6%, cancer 10.7%	Hydroxychloroquine 100%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
TOGETHER trial; ²⁹⁵ Reis et al; peer reviewed; 2021	to lopinavir-ritonavir	Mean age 53 ± 76, male 45%, hypertension 49.3%, diabetes 19.4%, COPD 2.5%, asthma 8.6%, CHD 3.9%, CKD 0.7%, cancer 1.2%, obesity 34.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	



<u>COPEP trial</u> ; ⁴¹⁰ Labhardt et al; preprint; 2021	Individuals exposed to SARS-CoV-2 infection. 209 assigned to lopinavir-ritonavir 400/10 mg a day for 5 days and 109 assigned to SOC	Median age 39 ± 22, male 50.6%, hypertension 8.2%, diabetes 3.1%, COPD 7.8%, CHD 2.5%, cancer 0.6%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>Ghanei et al</u> ; ⁸³ peer reviewed; 2021	Patients with severe COVID-19 infection. 110 assigned to lopinavir-ritonavir 200/50 mg twice a day for 7 days and 110 assigned to azithromycin 500 mg once followed by 250 mg a day for 5 days	Mean age 58.1 ± 16.3, male 51.5%, hypertension 24.7%, diabetes 12.2%, asthma 4.5%, CHD 8.9%, CKD 1.2%	Convalescent plasma 1.8%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
FIGHT-COVID- 19 trial; ²³⁴ Atipornwanich et al; preprint; 2021	Patients with mild to severe COVID-19 infection. 320 assigned to favipiravir 6000 mg once followed by 2400 mg a day + lopinavir ritonavir 800/200 mg or lopinavir ritonavir 800/200 mg a day or HCQ 800 mg a day or darunavir ritonavir 1200/200 mg a day + HCQ 400 mg a day or favipiravil 6000 mg followed by 2400 mg + darunavir ritonavir 1200/200 mg a day + HCQ 400 mg a day for 7 to 14 days.		NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.





SEV-COVID trial; ³⁰⁴ Panda et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 24 assigned to lopinavir ritonavir + ribavirin lopinavir (200 mg) + ritonavir (50 mg) two tablets twice daily + ribavirin (1.2 g orally as a loading dose followed by 600 mg orally every 12 hours) for 10 days and 24 assigned to SOC	Mean age 49.1, male 75%, hypertension 32.7%, diabetes 27.7%, COPD 7.9%, asthma %, CHD 11.9%, cancer 1%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>Nekoukar et al</u> ; ⁶⁷ peer reviewed; 2021	Patients with severe COVID-19 infection. 62 assigned to atazanavir/ritonavir 300/100 mg a day for 5 to 10 days and 62 assigned to lopinavir- ritonavir 200/50 mg a day for 5 to 10 days	Mean age 49.9 ± 12.6, male 55.6%, hypertension 16.9%, diabetes 27.4%, COPD 0.8%, asthma 1.6%	Corticosteroids 42.7%, remdesivir 13.7%, tocilizumab 3.2%, azithromycin 50.8%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
Hassaniazad et al; ²⁴⁰ peer reviewed; 2021	Patients with severe COVID-19 infection. 32 assigned to favipiravir 3200 mg once followed by 1200 mg for 5 days and 31 assigned to lopinavir- ritonavir 400/100 mg a day for 7 days	Mean age 53.7 ± 13.5, male 57.1%, hypertension 27%, diabetes 20.6%, COPD 1.6%, CHD 14.2%, obesity 7.9%	Interferon beta 100%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>FLARE trial</u> ; ²⁴¹ Lowe et al; preprint; 2021	Patients with mild recento onset COVID- 19 infection. 60 assigned to lopinavir- ritonavir 800/200 mg a	Mean age 40 ± 12, male 51.2%, obesity 16.7%, any comorbidity 15%	Vaccinated 51.2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	





<u>Tabarsi et al</u> ; ²⁴² peer reviewed; 2021	day for 7 days and 60 assigned to SOC Patients with severe COVID-19 infection. 32 assigned to favipiravir 3200 mg once followed by 1200 mg a day for 7 days and 30 assigned to lopinavir-ritonavir 400/100 mg a day for 7 days	Median age 57, male 58.1%, hypertension 12.9%, diabetes 21%, COPD %, asthma 3.2%, CHD 14.5%, CKD 3.2%, therapy %, cancer 4.8%, obesity 3.2%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
	Uncerta	Low-dose ra inty in potential benefits a	diation therapy nd harms. Further resea		
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>COVID-RT-01</u> <u>trial</u> , ⁴¹¹ Papachristofilou et al; peer reviewed; 2021	Patients with critical COVID-19 infection. 11 assigned to low- dose radiation therapy 0.5 to 1.0 Gy and 11 assigned to SOC	Mean age 75, male 77.3%, diabetes 54.6%, COPD 22.7%, asthma %, CHD 40.9%, cancer 18.2%,	Corticosteroids 100%, remdesivir 50%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty
WINCOVID trial; ⁴¹² Ganesan et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 34 assigned to low- dose radiation therapy 0.5 Gy single session and 17 assigned to SOC	Age (>56) 58.8% , male 66.6%, hypertension 35.3%, diabetes 68.6%, asthma 2%	Corticosteroids 100%, remdesivir 50.9%, tocilizumab 21.6%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	 ⊕○○○ Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No
<u>IMpaCt-RT trial</u> ; ⁴¹³ Singh et al; peer	Patients with severe COVID-19 infection.	Median age 56 ± , male 53.8%	Corticosteroids 100%, remdesivir 46.1%,	High for mortality and mechanical ventilation;	information



reviewed; 2021	7 assigned to low-dose radiation therapy 0.7 Gy and 6 assigned to SOC		azithromycin 100%,	high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Adverse events: No information Hospitalization: No information
	Uncerta	Mavri inty in potential benefits a	limumab nd harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
MASH-COVID trial; ⁴¹⁴ Cremer et al; peer reviewed; 2021		Mean age 56.7 ± 23.8, male 65%, hypertension 55%, diabetes 43%, COPD 8%, CKD 8%, cerebrovascular disease 3%	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: Very low certainty ⊕○○○Symptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information

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	Melatonin Uncertainty in potential benefits and harms. Further research is needed.							
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT		•		•				
Farnoosh et al; ⁴¹⁵ peer reviewed; 2020	Patients with mild to moderate COVID-19. 24 assigned to melatonin 9 mg a day for 14 days and 20 assigned to SOC	Mean age 51.85 ± 14.25, male 59.1%, hypertension 25%, diabetes 22.7%, CHD 6.8%, cancer 6.8%,	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Concealment of allocation is probably inappropriate. Significant loss to follow-up.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information			
<u>Davoodian et al</u> ; ⁴¹⁶ preprint; 2021	Patients with severe COVID-19 infection. 41 assigned to melatonin 6 mg a day for 14 days and 39 assigned to SOC	Median age 56 ± 40, male 56.8%, hypertension 18.5%, diabetes 14.8%, CHD 19.8%, CKD 3.7%	Corticosteroids 12.3%, hydroxychloroquine 69%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Symptom resolution or improvement: Very low certainty ⊕○○○			
Alizadeh et al; ⁴¹⁷ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 14 assigned to melatonin 6 mg a day for 14 days and 17 assigned to SOC	Mean age 36 ± 8.2, male 64.3%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Symptomatic infection (prophylaxis studies): Very low certainty ⊕○○○ Adverse events: No information Hospitalization: No information			
Mousavi et al; ⁴¹⁸ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 48 assigned to melatonin 3 mg a day	Mean age 52.9, male 44.8%, hypertension 30.2%, diabetes 28.1%, COPD 3.1%, asthma 5.2%, CHD 15.6%,	Corticosteroids 82.3%, hydroxychloroquine 97.9%, lopinavir- ritonavir 2.1%, azithromycin 100%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events				





	for 10 days and 48 assigned to SOC	CKD 5.2%,		Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>Hasan et al</u> ; ⁴¹⁹ peer reviewed; 2021	Patients with severe COVID-19 infection. 82 assigned to melatonin 10 mg a day for 14 days and 76 assigned to SOC	Mean age 56.3 ± 7.7, male 72.2%, hypertension 53.2%, diabetes 29.7%, asthma 10.1%, cerebrovascular disease 15.2%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>MeCOVID trial</u> ; ⁴²⁰ García-García et al; peer reviewed; 2021	Healthcare workers exposed to SARS- COV-2. 151 assigned to melatonin 2 mg a day for 12 weeks and 163 assigned to SOC	Median age 40, male 18.8%, hypertension 3.2%, CHD 0.3%, cancer 2.5%, obesity 0.3%	NR	Some Concerns for mortality and mechanical ventilation; some Concerns for symptom resolution, infection and adverse events
				Notes: Significant loss to follow up.
<u>Alizadeh et al</u> ; ⁴²¹ peer reviewed; 2021	Patients with critical COVID-19 infection. 33 assigned to melatonin 21 mg a day and 34 assigned to SOC	Mean age 63.5, male 64%	NR	Some Concerns for mortality and mechanical ventilation; some Concerns for symptom resolution, infection and adverse events
				Notes: Concealment of allocation probably inappropriate.
Fogleman C et al rrial; ⁴²² peer eviewed; 2022	Patients with mild to moderate COVID-19 infection. 32 assigned to melatonin 10 mg a day for 14 days and 34	Median age 52, male 44.9%, hypertension 26.5%, diabetes 16.3%	Vaccinated 2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events



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	assigned to SOC							
Mefenamic acid Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT					*			
MEFECOVID-19 trial; ⁴²³ Guzman- Esquivel et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 19 assigned to mefenamic acid 1500 mg a day for 7 days and 17 assigned to SOC	Mean age 39.5 ± 15.4, male 33.3%, diabetes 5.6%, asthma 2.8%, obesity 47.2%	Corticosteroids 2.8%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanica ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Hospitalization: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$			
		lesenchymal ster senchymal stem-cell trar						
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the			



					evidence
RCT					
<u>Shu et al</u> ; ⁴²⁴ peer- reviewed; 2020	Patients with severe COVID-19 infection. 12 assigned to mesenchymal stem cell 2 × 10^6 cells/kg one infusion and 29 assigned to standard of care	Median age 61 ± 10, male 58.5%, hypertension 22%, diabetes 19.5%	Corticosteroids 100%, antibiotics 87.8%, antivirals 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: RR 0.61 (95%CI 0.43 to 0.88); RD -6.2% (95%CI - 9.1% to -1.9%); Low certainty ⊕⊕◯◯
<u>Shi et al</u> ; ⁴²⁵ preprint; 2020	Patients with severe COVID-19. 65 assigned to mesenchymal stem cell three infusions with 4.0 × 107 cells each and 35 assigned to standard of care	Mean age 60.3 ± 8.4, male 56%, hypertension 27%, diabetes 17%, COPD 2%	Corticosteroids 22%	Low for mortality and mechanical ventilation	Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: Very
<u>Lanzoni et al</u> ; ⁴²⁶ preprint; 2020	Patients with severe to critical COVID-19. 12 assigned to mesenchymal stem cell 100±20 ×106 UC- MSC twice and 12 assigned to standard of care	Mean age 58.7 ± 17.5, male 54.1%, hypertension 66.7%, diabetes 45.8%, coronary heart disease 12.5%, , cancer 4.2%, obesity 66.6%	Corticosteroids 90.4%, remdesivir 66.7%, hydroxychloroquine 12.5%, tocilizumab 20.8%, convalescent plasma 29.1%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	
<u>Dilogo et al</u> ; ⁴²⁷ peer reviewed; 2021	Patients with critical COVID-19 infection. 20 assigned to mesenchymal stem cell one 100 ml infusion and 20 assigned to SOC	age >60, 45%, male 75%, hypertension 42.5%, diabetes 50%, CHD 25%, CKD 17.5%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	information Hospitalization: No information
<u>Zhu et al;</u> ⁴²⁸ peer reviewed; 2021	Patients with severe COVID-19 infection.	Median age 65, male 37.9%, hypertension	Corticosteroids 67.2%	High for mortality and mechanical ventilation;	

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	29 assigned to mesenchymal stem cell 1 × 106 cells per kilogram body weight, once and 29 assigned to SOC	25.8%, diabetes 13.8%, COPD 1.7%, CHD 10.3%, cerebrovascular disease 8.6%		high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
Fathi-Kazerooni et al; ⁴²⁹ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 14 assigned to mesenchymal stem cell 5 ml a day for 5 days and 15 assigned to SOC	Mean age 50 ± , male 65.5%, hypertension 31%, diabetes 24.1%	Corticosteroids 100%, remdesivir 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	
<u>Rebelatto et al</u> ; ⁴³⁰ peer reviewed; 2021	Patients with critical COVID-19 infection. 11 assigned to mesenchymal stem cell three doses of 5 × 105 cells/kg UC-MSCs and 6 assigned to SOC	Mean age 56, male 70.5%, hypertension 52.9%, diabetes 41.2%, COPD 5.9%, CKD 5.9%, obesity 52.9%		Some Concerns for mortality and mechanical ventilation; Some Concerns for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	
DW-MSC trial; ⁴³¹ Karyana et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 6 assigned to mesenchymal stem cell 5.0 × 10 ⁷ cells to 1.0 × 10 ⁸ cells and 3 assigned to SOC	Age range 31 to 47, male 66.6%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>Farkhad et al</u> ; ⁴³² preprint; 2022	Patients with severe COVID-19 infection. 10 assigned to mesenchymal stem cell 3 intravenous infusions of UC- MSCs (1 × 10^6	Mean age 61.7, male 65%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded	



	cells/kg BW per injection) every other day and 10 assigned to SOC			study. Concealment of allocation probably inappropriate.	
	Metforn	Met nin may not reduce hospit	formin alizations. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>TOGETHER 2</u> <u>trial</u> ; ⁴³³ Reis et al; peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 215 assigned to MTF 1500 mg a day and 203 assigned to SOC	Median age 52, male 42.8%, hypertension 40%, diabetes 14.6%, COPD 1.2%, asthma 8.1%, CHD 3%, CKD 0.5%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information
DMMETCOV19-2 <u>trial</u> ; ⁴³⁴ Ventura- López et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 10 assigned to metformin 1240 mg a day for 14 days and 10 assigned to SOC	Mean age 47.5, male 85%, hypertension 20%, diabetes 20%, COPD 10%,	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection and adverse events	Symptom resolution or improvement: No information Symptomatic infection
COVID-OUT trial; ²⁵⁴ Bramante et al; peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 663 assigned to metformin 1500 mg a day for 14 days and 398 assigned to SOC	Median age 45.5, male 44%, hypertension 26.7%, diabetes 2%, obesity 48.8%	Corticosteroids 1.5%, monoclonal antibodies 4.2%; Vaccinated 52.2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	(prophylaxis studies): No information Adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Hospitalization: RR 0.92 (95%CI 0.61 to 1.37); RD -0.4% (95%CI -1.9% to 1.8%); Low certainty $\oplus \bigcirc \bigcirc$
		Methy	lene blue		



	Uncerta	inty in potential benefits a	and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Hamidi-Alamdari et al; ⁴³⁵ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 40 assigned to methylene blue 1 mg/kg every 12 to 8 h for 14 days and 40 assigned to SOC	Mean age 54 ± 13, male 52.5%, hypertension 17.5%, diabetes 10%	Corticosteroids 87.5%, azithromycin 92.5%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No
	Uncerta	Meti inty in potential benefits a	soprinol and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Borges et al; ⁴³⁶ peer reviewed; 2020	Patients with mild to moderate COVID-19.	Mean age 33.2 ± 16, male 53.3%, COPD	NR	High for mortality and mechanical ventilation;	Mortality: No information



	30 assigned to metisoprinol 1500 mg/kg/day for 14 days and 30 assigned to SOC	10%, CKD 16.6%, cancer 3.3%		High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information						
	Uncerta		Metoprolol Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence						
		Comorbidities			vs standard of care and GRADE certainty of the						



					Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncerta	Metro inty in potential benefits a	onidazole and harms. Further resea	nrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Kazempour et al; ⁴³⁸ peer reviewed; 2021	Patients with moderate COVID-19 infection. 20 assigned to metronidazole 1 g a day for 7 days and 24 assigned to SOC	Mean age 63 ± 16.3, male 59.1%, hypertension 47.7%, diabetes 18.2%, COPD 6.8%, asthma %, CHD 4.5%	Hydroxychloroquine 59%, lopinavir- ritonavir 43.2%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No





					information			
Molnupiravir Molnupiravir reduces hospitalizations in patients with recent onset mild to moderate disease and may improve symptom resolution. It may not increase severe adverse events.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT								
<u>Painter et al</u> ; ⁴³⁹ Preprint; 2020	Healthy volunteers. 64 assigned to molnupiravir 80 to 1600 mg twice a day for 5.5 days	Mean age 39.6 ± 39, male 82.8%,	NR	Low for adverse events	Mortality: RR 0.35 (95%CI 0.06 to 2.19); RD -1.4% (95%CI - 15% to 19.4%); Very low certainty			
AGILE trial; ⁴⁴⁰ Khoo et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 12 assigned to molnupiravir 600- 1600 mg a day and 6 assigned to SOC	Median age 56 ± 58, male 27.8%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	$ \begin{array}{c} \bigoplus \bigcirc \bigcirc \bigcirc \\ \hline \\ Invasive mechanical \\ ventilation; RR 0.36 \\ (95\% CI 0.11 to 1.12); \\ RD -11.1\% (95\% CI - \\ 15.4\% to -2.1\%); Very \\ low certainty \\ \bigoplus \bigcirc \bigcirc \bigcirc \\ \hline \\ Symptom \\ resolution or \\ \end{array} $			
Fischer et al; ⁴⁴¹ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 140 assigned to molnupiravir 200 to 800 mg twice a day for 5 days and 62 assigned to SOC	Age >65 6%±, male 48.6%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	resolution or improvement: RR 1.17 (95%CI 1.1 to 1.3); RD 10.3% (95%CI 3.6% to - 18.2%); Low certainty ⊕⊕⊖⊖			
<u>MOVe-OUT trial;</u> <u>et al</u> ; ⁴⁴² Bernal et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 709 assigned to molnupiravir 1600 mg a day for 5 days	Median age 43, male 48.7%, diabetes 15.9%, COPD 4%, asthma %, CHD 11.7%, CKD 5.9%, cancer 2%, obesity 73.7%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Symptomatic infection (prophylaxis studies): No information Adverse events: RR			



	and 699 assigned to SOC				0.75 (95%CI 0.48 to 1.19); RD -2.6% (95%CI -5.3% to -
Hetero et al; other; 2021	Patients with mild COVID-19 infection. 371 assigned to molnupiravir 1600 mg a day and 370 assigned to SOC	NR	NR	Not assessed	1.9%); Low certainty ⊕⊕○○ Hospitalization: RR 0.6 (95%CI 0.44 to 0.81); RD -1.9% (95%CI -2.7% to -
Tippabhotla et al; preprint; 2021	Patients with mild COVID-19 infection. 610 assigned to molnupiravir 800 mg a day for 5 days and 610 assigned to SOC	Mean age 36.5 ± 11, male 61.6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	0.9%); High certainty ⊕⊕⊕⊕
reviewed; 2022	Patients with mild to moderate COVID-19 infection. 76 assigned to molnupiravir 1600 mg a day for 5 days and 31 assigned to SOC	Median age 39.8 ± , male 55.5%	Vaccinated 91.7%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
Khoo et al; preprint; 2022	Patients with mild to moderate COVID-19 infection. 90 assigned to molnupiravir 1600 mg a day for 5 days and 90 assigned to SOC	Mean age 42.5 ± , male 42.8%	Vaccinated 50%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
Ariibas et al; peer	Patients with moderate to severe COVID-19 infection. 226 assigned to	Mean age 57, male 66.6%	Corticosteroids 67.1%, remdesivir 23.7%; Vaccinated 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and	



		Mean age 52.6, male 49.2%, diabetes 16.6%, COPD 3.6%, asthma %, CHD 8.3%, CKD 2.3%, immunosuppression 0%, cancer 1%, obesity 48.7%	Corticosteroids %, remdesivir %, hydroxychloroquine %, lopinavir-ritonavir %, tocilizumab %, azithromycin %, convalescent plasma %; Vaccinated %	adverse events Low for mortality and mechanical ventilation; Low for symptom resolution, infection and adverse events Notes:	
	Uncertai	Mon inty in potential benefits a	telukast and harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Kerget et al, ⁴⁴⁸ peer reviewed; 2021	Patients with moderate COVID-19 infection. 120 assigned to montelukast 10 to 20 mg a day and 60 assigned to SOC	Mean age 54.6 ± 15.3, male 42.2%, hypertension 30%, diabetes 19%, asthma 1.7%, CHD 1.1%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information



					Hospitalization: No information			
Mouthwash Mouthwash may improve time to symptom resolution. Uncertainty in potential benefits and harms on other outcomes. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT								
<u>Mukhtar et al</u> ; ⁴⁴⁹ preprint ; 2020	Patients with mild to critical COVID-19. 46 assigned to mouthwash with hydrogen peroxide 2% and chlorhexidine gluconate mixed solution three times a day and 46 assigned to standard of care	Mean age 49, male 78.2%, hypertension 37%, diabetes 41.3%, coronary heart disease 6.5%, chronic kidney disease 12%, c obesity 31.5%	Corticosteroids 53.2%, remdesivir 26%, hydroxychloroquine 21.7%, lopinavir- ritonavir 54.3%, azithromycin 57.6%, convalescent plasma 13%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or			
GARGLES trial; ⁴⁵⁰ Mohamed et al; preprint; 2020	Patients with COVID- 19. 10 assigned to mouthwash with povidone iodine or essential oils 3 times a day and 10 assigned to mouthwash with water or no mouthwash	Median age 28.9, male 80%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	resolution or improvement: RR 1.36 (95%CI 1.04 to 1.78); RD 21.8% (95%CI 2.4% to 47.3%); Low certainty ⊕⊕○○ Symptomatic infection (prophylaxis studies): No			
KILLER trial; ⁴⁵¹ Guenezan et al; peer reviewed; 2020	Patients with mild COVID-19. 12 assigned to mouthwash with 25 ml of 1% povidone iodine and 12 assigned to SOC	Mean age 45 ± 23, male 33%, hypertension 12.5%, diabetes 4%,	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	information Adverse events: No information Hospitalization: No information			



<u>Elzein et al</u> ; ⁴⁵² preprint; 2021	Patients with mild to severe COVID-19 infection. 52 assigned to mouthwash with povidone or chlorhexidine and 9 assigned to SOC	Mean age 45.3 ± 16.7, male 40.9%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Santos et al</u> , ⁴⁵³ preprint; 2021	Patients with mild to moderate COVID-19 infection. 20 assigned to mouthwash with anionic iron tetracarboxyphthalocy anine derivative 5 times a day and 21 assigned to SOC	Mean age 53.7 ± 44.5, male 63%	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	
<u>BBCovid trial</u> , ⁴⁵⁴ Carrouel et al; preprint; 2021	Patients with mild COVID-19 infection. 76 assigned to mouthwash with ß- cyclodextrin-citrox three times a day and 78 assigned to SOC	Mean age 43.8 ± 15.5, male 45.7%,	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	
<u>Huang et al</u> ; ⁴⁵⁵ peer reviewed; 2021	Patients with moderate to critical COVID-19 infection. 66 assigned to mouthwash chlorhexidine 0.12% 15 ml twice a day for 4 days and 55 assigned to SOC	Median age 62 ± 66, male 58%	Corticosteroids 100%, remdesivir 100%,	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Eduardo et al</u> ; ⁴⁵⁶ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 34 assigned to mouthwash	Mean age 54.7, male 74.4%, hypertension 30.2%, diabetes 23.2%, COPD 11.6%, CHD 18.6%, CKD 11.6%,	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	





	cetylpyridinium chloride, zinc, chlorhexidine, hydrogen peroxide and 9 assigned to SOC	obesity 13.9%			
<u>Di-Domênico et</u> <u>al;</u> ⁴⁵⁷ peer reviewed; 2021	to mouthwash with	Age >60 17%, male 39.6%, hypertension 22.6%, diabetes 11.3%, COPD 5.7%, CHD 3.8%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Significant number of patients excluded post- randomization resulting in potential inbalances in baseline risks	
ACPREGCOV trial; ⁴⁵⁸ Damião Costa et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 50 assigned to mouthwash 15 mL of 0.12% chlorhexidine gluconate and 50 assigned to SOC	Mean age 39 ± 12, male 50%, hypertension 17%, diabetes 4%, obesity 25%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
BUCOSARS <u>trial</u> ; ⁴⁵⁹ Ferrer et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 54 assigned to mouthwash with povidone-iodine, hydrogen peroxide, cetylpyridinium chloride or chlorhexidine and 13 assigned to SOC	Mean age 54 - 55 ± , male 67%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>Poleti ML et al</u> <u>trial;</u> ⁴⁶⁰ Poleti et al; ; 2021	Patients with mild COVID-19 infection. 59 assigned to mouthwash with antimicrobial phthalocyanine	Mean age 34 ± 21, male 38%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	



Alemany et al; ⁴⁶¹ peer reviewed; 2022	derivative and 75 assigned to SOC Patients with mild COVID-19 infection. 60 assigned to mouthwash with 0.07% cetylpyridinium and 58 assigned to SOC	Mean age 46, male 41.5%	NR	Notes: Significant loss to follow-up. Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	
Barrueco et al; ⁴⁶² peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 35 assigned to mouthwash with povidone-iodine 2%, hydrogen peroxide 1%, cetylpyridinium chloride 0.07% or chlorhexidine 0.12% and 10 assigned to SOC	Mean age 62.4 ± , male 54.5%,	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
	Uncerta	Mupa inty in potential benefits a	adolimab and harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Miller et al</u> ; ⁴⁶³ preprint; 2021	Patients with moderate to severe COVID-19 infection. 29 assigned to mupadolimab 1-2 mg/kg and 11 assigned to SOC	Median age 55, male 57.5%, any comorbidities 45%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information



					Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○
	Uncerta	Mycoba inty in potential benefits a	acterium W and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT		• •	•	•	•
Sehgal et al; peer eviewed; 2021	Patients with severe to critical COVID-19 infection. 22 assigned to Mycobacterium w 0.3 ml SC once a day for 3 days and 20 assigned to SOC	Mean age 56 ± 15, male 69%, hypertension 31%, diabetes 33.3%, COPD 4.8%, asthma 4.8%	Corticosteroids 100%, hydroxychloroquine 26.2%, tocilizumab 12%, convalescent plasma 7%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			•		
<u>de Alencar et al</u> ; ⁴⁶⁵ peer-reviewed; 2020	Patients with severe COVID-19. 68 assigned to NAC 21 g once and 67 assigned to standard of care	Mean age 58.5 ± 22.5, male 59.2%, hypertension 46.6%, diabetes 37.7%, cancer 12.6%,	NR	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty
<u>Gaynitdinova et</u> al; ⁴⁶⁶ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 24 assigned to NAC 1200- 1500 mg once and 22 assigned to SOC	Mean age 57.9 ± 12.7	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	 ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis)
<u>Taher et al</u> ; ⁴⁶⁷ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 47 assigned to NAC 40 mg/kg a day for 3 days and 45 assigned to SOC	Mean age 57.6 ± 18.7, male 58.7%, diabetes 23.9%, COPD 15.2%, asthma %, CHD 28.2%,	Corticosteroids 69.6%, hydroxychloroquine 90.2%, azithromycin 51.1%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	 studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: N information
	Uncerta	N-acetylcys inty in potential benefits a	teine (inhaled) and harms. Further resea	irch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence



RCT					
Delic et al; ¹⁰⁸ peer reviewed; 2022	Patients with critical COVID-19 infection. 39 assigned to N- acetylcysteine (inhaled) twice a day and 52 assigned to SOC	Mean age 68.3 ± , male 74.8%, hypertension 61.5%, diabetes 27.5%, COPD %, asthma %, CHD 7.7%, CKD %, cerebrovascular disease 4.4%	Corticosteroids 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No
	Uncerta	Nafamos inty in potential benefits a	t at mesylate and harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
DEFINE trial; ⁴⁶⁸ Quinn et al; preprint; 2021	Patients with moderate to severe COVID-19 infection. 21 assigned to nafamostat 0.2 mg/kr/hr for 7 days and 21 assigned to SOC	Mean age 63.6, male 59.5%, hypertension 38.1%, diabetes 21.4%, COPD %, asthma 9.5%, CHD 14.3%, CKD 4.8%, immunosuppression 7.1%, cancer 9.5%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or

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				symptoms and adverse events outcomes results.	<pre>improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information</pre>
	Uncerta	Nam inty in potential benefits a	iilumab nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
CATALYST trial; ³²⁷ Fisher et al; preprint; 2021	Patients with moderate to critical COVID-19 infection. 55 assigned to namilumab and 54 assigned to SOC	Median age 62.8 ± 18, male 68.5%	Corticosteroids 90.7%, remdesivir 53.7%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information





					Adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
					Hospitalization: No information
	Uncertai		-Curcumin and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			-		
Hassaniazad et al; ⁴⁶⁹ peer reviewed; 2021	Patients with mild to severe COVID-19 infection. 20 assigned to nano-curcumin 160 mg a day for 14 days and 20 assigned to SOC	Mean age 48.5 ± 10.9, male 55%	Corticosteroids 87.5%, hydroxychloroquine 45%, lopinavir- ritonavir 52.5%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Kimura et al</u> ; ⁴⁷⁰ peer-reviewed; 2020	Patients with mild to moderate COVID-19. 14 assigned to nasal hypertonic saline 250 cc twice daily, 14 assigned to nasal hypertonic saline plus surfactant and 17 assigned to standard of care	Mean age 37.9 ± 15.7, male 53.3%, hypertension 24.4%, diabetes 6.6%, chronic lung disease 15.5%, coronary heart disease 4.4%,	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No
<u>Yildiz et al</u> ; ⁴⁷¹ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 50 assigned to nasal hypertonic saline and 50 assigned to SOC	Mean age 38.8 ± , male 58%, hypertension 12%, diabetes 6%, COPD/asthma 4%, CHD 15%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection
<u>George et al</u> ; ⁴⁷² peer reviewed; 2021	Patients with mild COVID-19 infection. 20 assigned to nasal hypertonic saline (Caclium rich hypertonic salts) and 20 assigned to SOC	Age range 22-45		Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	 (prophylaxis studies): No information Adverse events: No information Hospitalization: No
<u>Baxter et al</u> ; ⁴⁷³ preprint; 2021	Patients with mild to moderate COVID-19 infection. 37 assigned to nasal saline 240 ml + povidone-iodine twice a day for 14 days and 42 assigned to	Mean age 64 ± 7.9 , male 54.4%, hypertension 43.4%, diabetes 11.3%, COPD %, asthma 5.7%, immunocompromised 3.8%, obesity 45%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded	information



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE
	Uncertai	Niclo inty in potential benefits a	osamaide and harms. Further rese	arch is needed.	
					Hospitalization: No information
					Adverse events: No information
					Symptomatic infection (prophylaxis studies): Very low certainty ⊕○○○
				study. Concealment of allocation is probably inappropriate. Significant loss to follow-up.	Symptom resolution or improvement: No information
	to neem 50 mg for 28 days and 84 assigned to SOC			resolution, infection, and adverse events Notes: Non-blinded	Invasive mechanical ventilation: No information
<u>Nesari et al</u> ; ⁴⁷⁴ other; 2021	Individuals exposed to SARS-CoV-2 infection. 70 assigned	Mean age 37, male %	NR	High for mortality and mechanical ventilation; high for symptom	Mortality: No information
RCT		L			<u>.</u>
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
	Uncertai	Neem (<i>Azadirac</i> inty in potential benefits a	c <i>hta indica</i> A. Ju and harms. Further rese		
	bicarbonate twice a day for 14 days			allocation probably inappropriate.	
	nasal saline 240 ml +2.5 mL sodium			study. Concealment of	





					certainty of the evidence
RCT					
<u>Abdulamir et al</u> ; ⁴⁷⁵ preprint; 2021	Patients with mild to critical COVID-19 infection. 75 assigned to niclosamaide 4 g once followed by 3 g a day for 7 days and 75 assigned to SOC	Mean age 49.3 ± 16, male 53.3%, hypertension 12.7%, diabetes 8%, asthma 0.7%, cancer 0.7%, obesity 0.7%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Symptom resolution or
<u>Cairns et al</u> ; ⁴⁷⁶ peer reviewed; 2021	Patients with mild COVID-19 infection. 33 assigned to niclosamide 2 g a day for 7 days and 34 assigned to SOC	Mean age 36.4 ± 13, male 61.2%, hypertension 7.5%, asthma 7.5%, CHD 1.5%, obesity 7%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	resolution orimprovement: NoinformationSymptomaticinfection(prophylaxisstudies): NoinformationAdverse events:Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Hospitalization:Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
	Uncerta	Nicotin inty in potential benefits a	ne patches and harms. Further resea	urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Labro et al</u> ; ⁴⁷⁷ peer reviewed; 2022	Patients with critical COVID-19 infection. 106 assigned to	Mean age 61, male 69.7%, hypertension 58.7%, diabetes 41.4%,	Corticosteroids 64.5%, tocilizumab 0.5%	Low for mortality and mechanical ventilation; low for symptom	Mortality: RR 1.02 (95%CI 0.67 to 1.57); RD 0.3% (95%CI -





	nicotine patches 14 mg a day for a maximum of 30 days and 112 assigned to SOC	COPD 3.2%, cerebrovascular disease 8.3%, immunosuppresion 6%,		resolution, infection and adverse events	5.2% to 5.7%); Low certainty ⊕⊕○○ Invasive mechanical ventilation: No information
					Symptom resolution or improvement: No information
					Symptomatic infection (prophylaxis studies): No information
					Adverse events: Very low certainty ⊕○○○
					Hospitalization: No information
	Uncertai	<i>Nigella sat</i> inty in potential benefits a	<i>iva</i> +/- Honey and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
HNS-COVID-PK trial; ⁴⁷⁸ Ashraf et al; preprint; 2021	Patients with moderate to severe COVID-19 infection. 157 assigned to honey + <i>Nigella sativa</i> 1 g + 80 mg/kg three times a day for 13 days and 156 assigned to SOC	> 60 age 52 ±, male 56.8%, hypertension 31.6%, diabetes 36.7%	Corticosteroids 26.5%, azithromycin 73.8%, ivermectin 36.4%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom





Koshak et al; ⁴⁷⁹ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 91 assigned to <i>Nigella sativa</i> 500 mg twice a day for 10 days and 92 assigned to SOC	Mean age 36 ± 11, male 53%, hypertension 9%, diabetes 8%, asthma 4%, CHD 0.5%, obesity 25%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: Very low certainty ⊕○○○				
	Nirmatrelvir-ritonavir Nirmatrelvir-ritonavir probably reduces hospitalizations. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence				
RCT									
EPIC-HR trial; ⁴⁸⁰ Hammond et al; peer reviewed; 2021	Patients with COVID- 19 infection. 1039 assigned to nirmatrelvir/ritonavir 600/200 mg a day for 5 days and 1046 assigned to SOC	Median age 46, male 51.1%, hypertension 32.9%, diabetes 12.1%, obesity 35.6%	NR; vaccinated 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No				





					information Adverse events: RR 0.49 (95%CI 0.30 to) 0.80); RD -5.2% (95%CI -7.1% to -2%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ Hospitalization: RR 0.12 (95%CI 0.06 to) 0.25); RD -4.2% (95%CI -4.5% to -3.5%); Moderate certainty $\oplus \oplus \oplus \bigcirc$
	Uncertai	Nitaz inty in potential benefits a	OXANIDE nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT				•	
<u>SARITA-2 trial</u> ; ⁴⁸¹ Rocco et al; preprint; 2020	Patients with mild COVID-19. 194 assigned to nitazoxanide 500 mg three times a day for 5 days and 198 assigned to standard of care	Age range 18 - 77, male 47%, comorbidities 13.2%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Significant loss to follow-up.	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Symptom resolution or improvement: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$
<u>Fontanesi et a</u> l; ⁴⁸² preprint ; 2020	Patients with mild to critical COVID-19. 25 assigned to	Age > 65 46%, male 30%	NR	High for mortality and mechanical ventilation; High for symptom	Symptomatic infection

PAHO Pan American Brealth Organization

	nitazoxanide 1200 mg a day for 7 days and 25 assigned to SOC			resolution, infection, and adverse events Notes: Concealment of allocation and blinding probably inappropriate.	(prophylaxis studies): Very low certainty ⊕○○○ Adverse events: Very low certainty
<u>Silva et al</u> ; ⁴⁸³ preprint; 2021	Patients with mild to moderate COVID-19 infection. 23 assigned to nitazoxanide 2-3 g a day for 14 days and 13 assigned to SOC	Male 72.2%,	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	 ⊕○○○ Hospitalization: Very low certainty ⊕○○○
<u>Vanguard trial;</u> ⁴⁸⁴ Rossignol et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 184 assigned to nitazoxanide 600 mg a day for 5 days and 195 assigned to SOC	Mean age 40.3 ± 15.4, male 43.5%, comorbidities 34%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
NACOVID trial; ⁴⁸⁵ Fowotade et al; preprint; 2021	Patients with mild to severe COVID-19 infection. 31 assigned to nitazoxanide 2000 mg plus atazanavir/ritonavir 300/100 mg a day and 26 assigned to SOC	Mean age 38 ± 16, male 67%, obesity 19%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Medhat et al</u> ; ⁴⁸⁶ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 77 assigned to nitazoxanide 2000 mg a day for 14 days and 73 assigned to SOC	Mean age 45, male 45.3%, hypertension 21.3%, diabetes 19.3%	Corticosteroids 44%, hydroxychloroquine 7.3%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded	





COVER HCW trial; ⁴⁸⁷ Sokhela et al; peer reviewed; 2022	Patients with exposed to COVID-19 infection. 280 assigned to nitazoxanide 1000 mg a day for 1 week followed by 2000 mg a day for 24 weeks and 283 assigned to SOC	Median age 24, male 51.9%, hypertension 8.2%, diabetes 1.1%, COPD 2.2%	Vaccinated 0%	study which might have introduced bias to symptoms and adverse events outcomes results. Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Significant loss to follow-up.	
	Uncertai	Nitr nty in potential benefits a	ic oxide nd harms. Further resea	rrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Moni et al</u> ; ⁴⁸⁸ preprint; 2021	Patients with severe COVID-19 infection. 14 assigned to inhaled nitric oxide (iNO) pulses of 30 min for 3 days and 11 assigned to SOC	Mean age 59.8 ± 10, male 72%, hypertension 44%, diabetes 56%, COPD 12%, CHD 24%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: Very
Winchester et al; ⁴⁸⁹ peer-reviewed; 2021	Patients with mild COVID-19 infection. 40 assigned to nitric oxide nasal spray	Mean age 44, male 36.7%, hypertension 6.3%, diabetes 6.3%, COPD 1.2%, CHD 0%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection,	$\begin{array}{c} \text{improvement: Very} \\ \text{low certainty} \\ \oplus \bigcirc \bigcirc \bigcirc \\ \\ \text{Symptomatic} \end{array}$





NO COV-ED trial, ⁴⁹⁰ Strickland et al; peer reviewed; 2021	(NONS) 4 sprays 5 to 6 times a day for 9 days and 40 assigned to SOC Patients with moderate COVID-19 infection. 19 assigned to inhaled nitric oxide (iNO) 5 liters per minute and 15 assigned to SOC	Mean age 41, male 53.2%, hypertension 12.8%, diabetes 6.4%, COPD 14.9%, CHD 2.1%, immunosuppression 4.3%	NR	and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate. High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
<u>Tandon et al</u> ; ⁴⁹¹ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 64 assigned to nitric oxide nasal spray (NONS) 0.45 mL/dose six times a day for 8 days and 69 assigned to SOC	Mean age 37.8, male 64.4%, any commorbidities 12.1%	Vaccinated 46.1%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
Current best evid	ence suggests no associat	eroidal anti-infl ion between NSAID consi s very low because of the	umption and COVID-19	related mortality. Howeve	r, the certainty of the
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Mobarak et al</u> ; ⁴⁹² peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 39 assigned to naproxen 1000 mg a day and 38 assigned to SOC	Mean age 47 , male 55.8%, hypertension 9%, diabetes 17%, CHD 13%, CKD 5.2%, obesity 1.3%	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection and adverse events Notes:	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information





Non-RCT					Symptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
<u>Eilidh et al</u> ; ⁴⁹³ peer- reviewed; 2020	Patients with moderate to severe COVID-19 infection. 54 received NSAID and 1168 received alternative treatment schemes	Age < 65 31.7%, male 56.5%, hypertension 50.3%, diabetes 27%, coronary heart disease 22.3%, chronic kidney disease 38.7%,	NR	High for mortality Notes: Non-randomized study with retrospective design. Regression was implemented to adjust for potential confounders (age, sex, smoking status, CRP levels, diabetes, hypertension, coronary artery disease, reduced renal function).	Mortality: OR 0.82 (95%CI 0.66 to 1.02); Very low certainty
Jeong et al; ⁴⁹⁴ preprint; 2020	Patients with moderate to severe COVID-19 infection. 354 received NSAID and 1470 received alternative treatment schemes	Age >65 36%, male 41%, hypertension 20%, diabetes 12%, chronic lung disease 16%, asthma 6%, chronic kidney disease 2%, cancer 6%	NR	High for mortality and invasive mechanical ventilation Notes: Non-randomized study with retrospective design. Propensity score and IPTW were implemented to adjust for potential	⊕○○○





				confounders (age, sex, health insurance type, hypertension, hyperlipidemia, diabetes mellitus, malignancy, asthma, chronic obstructive pulmonary disease, atherosclerosis, chronic renal failure, chronic liver disease, rheumatoid arthritis, osteoarthritis, gastrointestinal, conditions, and use of co-medications).	
Lund et al; ⁴⁹⁵ peer- reviewed; 2020	Patients with mild to severe COVID-19 infection. 224 received NSAID and 896 received alternative treatment schemes	Median age 54 ± 23, male 41.5%, chronic lung disease 3.9%, asthma 5.4%, coronary heart disease 10.2%, cerebrovascular disease 3.4%, cancer 7.1%, obesity 12.5%	Corticosteroids 7.1%	High for mortality and invasive mechanical ventilation Notes: Non-randomized study with retrospective design. Propensity score and matching were implemented to adjust for potential confounders (age, sex, relevant comorbidities, use of selected prescription drugs, and phase of the outbreak.	
Rinott et al; ⁴⁹⁶ peer- reviewed; 2020	Patients with moderate to critical COVID-19 infection. 87 received NSAID and 316 received alternative treatment schemes	Median age 45 ± 37, male 54.6%, diabetes 9.4%, coronary heart disease 12.9%,	NR	High for mortality and invasive mechanical ventilation Notes: Non-randomized study with retrospective design. No adjustment for potential confounders.	
<u>Wong et a</u> l; ⁴⁹⁷ preprint; 2020	Individuals exposed to SARS-CoV-2 infection. 535519	Median age 51 ± 23, male 42.7%, hypertension 19.6%,	Corticosteroids 2.2%, hydroxychloroquine 0.6%	High for mortality Notes: Non-randomized	



	received NSAID and 1924095 received alternative treatment schemes	diabetes 9.6%, chronic lung disease 2.4%, asthma %, coronary heart disease 0.5%, chronic kidney disease 2.8%, cancer 5.2%,		study with retrospective design. Regression was implemented to adjust for potential confounders (age, sex, relevant comorbidities, use of selected prescription drugs, vaccination, and deprivation).
<u>Imam et al</u> ; ⁴⁹⁸ peer- reviewed; 2020	Patients with moderate to critical COVID-19 infection. 466 received NSAID and 839 received alternative treatment schemes	Mean age 61 ± 16.3, male 53.8%, hypertension 56.2%, diabetes 30.1%, chronic lung disease 8.2%, asthma 8.8%, coronary heart disease 15.9%, chronic kidney disease 17.5%, immunosuppression 1%, cancer 6.4%,	NR	High for mortality Notes: Non-randomized study with retrospective design. Regression was implemented to adjust for potential confounders (not specified).
<u>Esba et al;</u> ⁴⁹⁹ preprint; 2020	Patients with mild to severe COVID-19 infection. 146 received NSAID and 357 received alternative treatment schemes	Median age 41.7 ± 30, male 57.2%, hypertension 20.4%, diabetes 22.5%, chronic lung disease 5.2%, chronic kidney disease 3.2%, cancer 1.4%	NR	High for mortality Notes: Non-randomized study with retrospective design. Regression was implemented to adjust for potential confounders (age; sex; comorbidities: hypertension, diabetes mellitus (DM), dyslipidemia, asthma, or chronic obstructive pulmonary disease (COPD), cardiovascular disease (CVD), renal or liver impairment, and malignancy).

Norelgestromin and Ethinylestradiol Uncertainty in potential benefits and harms. Further research is needed.



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Cortés-Algara et</u> <u>al</u> ; ⁵⁰⁰ peer reviewed; 2021	Patients with moderate COVID-19 infection. 30 assigned to norelgestromin and ethinylestradiol 6 mg/ 0.6 mg and 14 assigned to SOC	Mean age 58.6 , male 38.6%, hypertension 29.5%, diabetes 34.1%, obesity 6.8%	Corticosteroids 65.9%, hydroxychloroquine 65.9%, azithromycin 93.2%, vaccinated 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncerta	Nov inty in potential benefits a	7 aferon and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Zheng et al</u> ; ⁴⁰⁴ preprint; 2020	Patients with moderate to severe COVID-19 infection. 30 assigned to	Median age 44.5 ± NR, male 47.1%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution,	Mortality: No information Invasive mechanical



	novaferon 40 microg			infection, and adverse	ventilation: No
	twice a day (inh), 30 assigned to novaferon			events	information
	plus lopinavir-			Notes: Non-blinded	Symptom
	ritonavir 40 microg			study. Concealment of	resolution or
	twice a day (inh) +			allocation is probably	improvement: No
	400/100 mg a day and 29 assigned to			inappropriate.	information
	lopinavir-ritonavir				Symptomatic
	1				infection
					(prophylaxis
					studies): No
					information
					Adverse events: No
					information
					Hospitalization: No
					information
		Nutritio	nal support		
	Uncertai	inty in potential benefits a	and harms. Further resea	arch is needed.	
Study; publication	Patients and interventions analyzed	Comorbidities	Additional	Risk of bias and study limitations	Interventions effects
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE
		Comorbidities			vs standard of care and GRADE certainty of the
status		Comorbidities			vs standard of care and GRADE
		Comorbidities			vs standard of care and GRADE certainty of the
status		Comorbidities Mean age 52.7 \pm 10.8,		limitations High for mortality and	vs standard of care and GRADE certainty of the
status	Patients with severe COVID-19 infection.	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation;	vs standard of care and GRADE certainty of the evidence
status RCT Leal et al; ⁵⁰¹	Patients with severe COVID-19 infection. 40 assigned to	Mean age 52.7 ± 10.8,	interventions	limitations High for mortality and mechanical ventilation; high for symptom	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○
status RCT Leal et al; ⁵⁰¹	Patients with severe COVID-19 infection. 40 assigned to nutritional support	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical
status RCT Leal et al; ⁵⁰¹	Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very
status RCT Leal et al; ⁵⁰¹	Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic acid, glutamine,	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty
status RCT Leal et al; ⁵⁰¹	interventions analyzed Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic acid, glutamine, vegetable protein,	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very
status RCT Leal et al; ⁵⁰¹	interventions analyzed Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic acid, glutamine, vegetable protein, vitamin C, zinc,	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○
status RCT Leal et al; ⁵⁰¹	interventions analyzed Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic acid, glutamine, vegetable protein, vitamin C, zinc, selenium, vitamin D,	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty
status RCT Leal et al; ⁵⁰¹	interventions analyzed Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic acid, glutamine, vegetable protein, vitamin C, zinc,	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom
status RCT Leal et al; ⁵⁰¹	interventions analyzed Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic acid, glutamine, vegetable protein, vitamin C, zinc, selenium, vitamin D, resveratrol, omega-3,	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or
status RCT Leal et al; ⁵⁰¹	interventions analyzed Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic acid, glutamine, vegetable protein, vitamin C, zinc, selenium, vitamin D, resveratrol, omega-3, L-arginine, magnesium	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information
status RCT Leal et al; ⁵⁰¹	interventions analyzed Patients with severe COVID-19 infection. 40 assigned to nutritional support with spirulin, folic acid, glutamine, vegetable protein, vitamin C, zinc, selenium, vitamin D, resveratrol, omega-3, L-arginine, magnesium and probiotics and 40	Mean age 52.7 ± 10.8, male 65%, CHD 33.7%,	interventions	limitations High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably	vs standard of care and GRADE certainty of the evidence Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No



					(prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncertai	Omega-3 inty in potential benefits a	B fatty acids nd harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	•				
<u>Sedighiyan et al</u> ; ⁵⁰² Preprint; 2020	Patients with mild to moderate COVID-19. 15 assigned to omega-3 670 mg three times a day for 2 weeks and 15 assigned to SOC	Mean age 66.7 ± 2.5, male 60%	Hydroxychloroquine 100%,	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No
<u>Doaei et al</u> ; ⁵⁰³ peer reviewed; 2021	Patients with critical COVID-19 infection. 28 assigned to omega-3 1000 mg a day and 73 assigned to SOC	Mean age 64 ± 14, male 59.4%	NR	Some concerns for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Blinding is probably inappropriate. Significant loss to follow-up.	information Symptomatic infection (prophylaxis studies): No information Adverse events: No information
<u>COVID-Omega-F</u> <u>trial;</u> ⁵⁰⁴ Arnardottir	Patients with moderate to severe	Mean age 81.1 ± 6.1, male 45%, hypertension	NR	Low for mortality and mechanical ventilation;	Hospitalization: No information





et al; preprint; 2021	COVID-19 infection. 10 assigned to omega-3 10 g a day for 5 days and 12 assigned to SOC	64%, diabetes 41%, COPD 13%, CHD 64%, CKD 23%, cancer 18%		High for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
	Uncertai	inty in potential benefits a	P-101 nd harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>PRANA trial</u> ; ⁵⁰⁵ Gusdon et al; peer reviewed; 2022	Patients with severe to critical COVID-19 infection. 17 assigned to OP-101 2 to 8 mg/kg once and 7 assigned to SOC	Median age 61, male 70.8%, hypertension 45.8%, diabetes 58.3%	Corticosteroids 100%, remdesivir 75%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: Very low certainty ⊕○○○Symptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information

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Opaganib may not r	Opaganib Dpaganib may not reduce mortality or mechanical ventilation; it may not increase severe adverse events but it may increase symptom resolutio or improvement. Further research is needed.							
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT	l							
<u>ABC-110 trial</u> ; ⁵⁰⁶ Winthrop et al; peer-reviewed; 2021	Patients with moderate to severe COVID-19 infection. 22 assigned to opaganib 1000 mg a day for 14 days and 18 assigned to SOC	Median age 58 ± 29.8, male 64.3%	Corticosteroids 92.8%, remdesivir 45.2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: RR 0.94 (95%CI 0.66 to 1.34) RD -0.9% (95%CI - 5.5% to -5.4%); Low certainty ⊕⊕○○ Invasive mechanical			
<u>Carvalho</u> <u>Neuenschwander et</u> al; ⁵⁰⁷ preprint; 2022	Patients with severe COVID-19 infection. 230 assigned to opaganib 500 mg a day for 14 days and 233 assigned to SOC	Mean age 56.5, male 65.4%, diabetes 35%	Corticosteroids 94.2%, remdesivir 17.3%, convalescent plasma 1.7%; Vaccinated 0.3%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	ventilation: RR 0.94 (95%CI 0.68 to 1.24); RD -1% (95%CI - 5.5% to -4.1%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Symptom resolution or improvement: RR 1.1 (95%CI 0.95 to 1.27); RD 6% (95%CI -3% to -16.4%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No information Adverse events: RR 0.96 (95%CI 0.69 to 1.34); RD -0.4% (95%CI -3.2% to - 3.5%); Low certainty $\oplus \oplus \bigcirc \bigcirc$			
					Hospitalization: No			





					information			
Otilimab Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT				·	•			
OSCAR trial; ⁵⁰⁸ Patel et al; preprint; 2021	Patients with severe to critical COVID-19 infection. 386 assigned to otilimab 90 mg once and 393 assigned to SOC	Mean age 59.6 ± 12, male 71.6%, hypertension 49.7%, diabetes 36.7%, CHD 11.9%	Corticosteroids 83%, remdesivir 34%, tocilizumab 1.2%, convalescent plasma 6%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationGroup by laxis studies): No informationAdverse events: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Hospitalization: No information			
	Uncerta	C inty in potential benefits :	ZONE and harms. Further rese	arch is needed.				
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			



RCT					
PROBIOZOVID trial; ⁵⁰⁹ Araimo et al; peer-reviewed; 2020	Patients with moderate to severe COVID-19. 14 assigned to ozone 250 ml ozonized blood and 14 assigned to standard of care	Mean age 61.7 ± 13.2, male 50%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very
SEOT trial; ⁵¹⁰ Shah et al; Peer reviewed; 2020	Patients with mild to moderate COVID-19. 30 assigned to ozone 150 ml rectal insufflation plus 5 ml with venous blood once a day for 10 days and 30 assigned to SOC	Mean age 43.8 ± 9, male 80%, diabetes 10%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
P2Y12 in combinatio				may not improve time to s is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			· ·		
<u>ACTIV-4a trial</u> ; ⁵¹¹ Berger et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 293 assigned to P2Y12 inhibitors (ticagrelor 120 mg a day or	Mean age 52.7, male 58.5%, hypertension 48.4%, diabetes 25.8%, COPD 5.4%, asthma 11.2%, CKD 3.9%, cerebrovascular disease	Corticosteroids 64.1%, remdesivir 52%, tocilizumab 2.8%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: RR 1.02 (95%CI 0.64 to 1.62); RD 0.3% (95%CI - 5.7% to 9.9%); Low certainty ⊕⊕○○





REMAP-CAP - P2Y12 trial; ⁶⁶ Bradbury et al; peer reviewed; 2021	prasugrel 5 to 10 mg a day or clopidogrel 75 mg a day) in combination with full dose anticoagulants and 269 assigned to SOC in combination with full dose anticoagulants Patients with severe to critical COVID-19 infection. 455 assigned to P2Y12 inhibitors clopidogrel 75 mg a day or ticagrelor 120 mg a day or prsugrel 60 mg once followed by 5 to 10 mg a day for 14 days and 529 assigned to SOC	0.7% Median age 57, male 67.2%, hypertension %, diabetes 39.3%, CHD 5.1%, CKD 3.9%	Corticosteroids 97.4%, remdesivir 22%, tocilizumab 43.7%	Low for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Invasive mechanical ventilation: Very low certainty ⊕○○○Symptom resolution or improvement: RR 0.97 (95%CI 0.94 to 1.02); RD -1.8% (95%CI -3.6% to 1.2%); Low certainty ⊕⊕○○Symptomatic infection (prophylaxis studies): No informationAdverse events: RR 3.1 (95%CI 1.32 to 7.29); RD 21.4% (95%CI -3.3% to 64.2%); Low certainty ⊕⊕○○Hospitalization: No information
	Uncerta	Palmitoyle	e thanolamide and harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT				·	
Fessler et al; ⁵¹² peer reviewed; 2022	Patients with mild COVID-19 infection. 30 assigned to Palmitoylethanolamid e 230 to 300 mg twice	Mean age 25.5, male %, hypertension 3.3%, asthma 6.6%	Vaccinated 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	Mortality: No information Invasive mechanical ventilation: No





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	a day for 4 weeks and 30 assigned to SOC	Peg-interfe	ron (IFN) alfa	Notes: Concealment of allocation probably inappropriate.	information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncerta	inty in potential benefits a		arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT		•	•	•	
	Patients with mild to moderate COVID-19 infection. 20 assigned to pegylated interferon alfa 1 µg/kg once and 19 assigned to SOC	Mean age 49.2 ± 13.5, male 75%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very
Bushan et al; ⁵¹⁴ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 119 assigned to Peg Interferon Alfa 1 μg/kg subcutaneous	Mean age 49.9 ± 15.3, male 70.8%	Corticosteroids 59.9%, remdesivir 21.5%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	low certainty ⊕○○○ Symptomatic infection (prophylaxis



	[SC] injection once and 123 assigned to SOC	Peg-interfer	on (IFN) lamda	Notes: Non-blinded study. Concealment of allocation probably inappropriate.	 studies): No information Adverse events: No information Hospitalization: No information
	Uncertai	inty in potential benefits a		urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
ILIAD trial; ⁵¹⁵ Feld et al; preprint; 2020	Patients with mild to severe COVID-19. 30 assigned to peg-IFN lambda 180 µg subcutaneous injection once and 30 assigned to standard of care	Median age 46 ± 22, male 58%, comorbidities 15%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: No information Invasive mechanical ventilation: No information Symptom
<u>COVID-Lambda</u> <u>tria</u> l; ⁵¹⁶ Jagannathan et al; preprint; 2020	Patients with mild COVID-19. 60 assigned to peg-IFN lambda 180 mcg subcutaneous injection once and 60 assigned to standard of care	Median age 36 ± 53, male 68.3%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events:
Chung et al; <u>NCT04343976;</u> other; 2022	Patients with moderate to severe COVID-19 infection. 7 assigned to Peg-IFN lambda 180 µg once	Mean age 54.5, male 78.6%,	NR	NA	Very low certainty ⊕○○○ Hospitalization: Very low certainty ⊕○○○

PROTECT trial; <u>NCT04344600;</u> Sulkowski et al;	and 7 assigned to SOC Patients with exposed to COVID-19 infection. 2 assigned to	Age >65 50, male 16.7%	NR	NA	
other; 2022	Peg-IFN lambda 180 μg once and 4 assigned to SOC				
	Uncertai	Pembr inty in potential benefits a	'Olizumab nd harms. Further resea	nrch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
COPERNICO trial; ⁵¹⁷ Sanchez- Conde et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 7 assigned to pembrolizumab 200 mg on days 1 and 21 and 5 assigned to SOC	Mean age 68, male 75%	Corticosteroids 100%, remdesivir 33%	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty @OOOInvasive mechanical ventilation: Very low certainty @OOOSymptom resolution or improvement: Very low certainty @OOOSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty @OOOHospitalization: No



					information				
	Pentoxifylline Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence				
RCT									
<u>Maldonado et al</u> ; ⁵¹⁸ peer-reviewed; 2020	Patients with severe to critical COVID-19. 26 assigned to pentoxifylline 400 mg three times a day while hospitalized and 12 assigned to standard of care	Mean age 57.5 ± 11.7, male 55.2%, hypertension 39.4%, diabetes 50%, obesity 55.2%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement:No				
<u>Azizi et al</u> ; ⁵¹⁹ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 40 assigned to pentoxifylline 1200 mg a day for 10 days and 32 assigned to SOC	Mean age 59, male 35%, hypertension 18%, diabetes 32%, CHD 12.5%, cerebrovascular disease 5.5%	Corticosteroids 55.5%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate. Significant loss to follow-up.	information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information				
	Uncorta	Pirf einty in potential benefits a	enidone	urch is needed					
	Uncertai	incy in potential benefits a	ing harms, purtiler resea						
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence				



RCT					
Zhang et al; ⁵²⁰ peer reviewed; 2022	Patients with severe COVID-19 infection. 73 assigned to pirfenidone 1200 mg a day for 28 days and 73 assigned to SOC	Mean age 62, male 64.4%, hypertension 34.3%, diabetes 12.3%, COPD 6.2%, CHD 5.5%, CKD 1.4%, cerebrovascular disease 3.4%, cancer 2.7%,	Corticosteroids 84.3%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncerta	Plit inty in potential benefits :	idepsin and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>APLICOV-PC</u> <u>trial</u> ; ⁵²¹ Varona et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 45 assigned to plitidepsin three doses of 1.5 to 2.5 mg	Mean age 51, male 66.6%, hypertension 20%, diabetes 17.8%, COPD 6.7%, asthma 11.1%, CHD 4.4%, CKD 2.2%, obesity	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events Notes:	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty





		22.2%			⊕○○○ Symptom resolution or
					<pre>improvement:No information Symptomatic infection (prophylaxis studies): No information Adverse events:</pre>
					Very low certainty ⊕○○○ Hospitalization: No information
	Uncertai	PNB001 (CC inty in potential benefits a	K-A antagonist)		
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
BCR-PNB-001 <u>trial</u> ; ⁵²² Lattaman et al; preprint; 2021	Patients with moderate COVID-19 infection. 20 assigned to PNB001 200 mg a day for 14 days and 20 assigned to SOC	Mean age 52, 65% male	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: No information Symptom resolution or
					<pre>improvement: Very low certainty ⊕○○○ Symptomatic</pre>





		Polymerized typ			infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Mendez-Flores et al; ⁵²³ preprint; 2021	moderate COVID-19 infection. 44 assigned to PT1C 25 mg	Mean age 48.5 ± 14.1, male 41.6%, hypertension 20.2%, diabetes 16.9%, COPD 2.3%, asthma 4.5%, CHD 0%, cancer 0%, obesity 28.1%	Corticosteroids 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: Very low certainty

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					$\oplus \bigcirc \bigcirc \bigcirc \bigcirc$			
Potassium canrenoate Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT								
SpiroCOVID19 trial; ⁵²⁴ Karolak et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 24 assigned to potassium canrenoate 400 mg a day for 7 days and 25 assigned to SOC	Mean age 62, male 53.1%, hypertension 63.2%, diabetes 28.6%, COPD %, asthma %, CHD 14.2%, cerebrovascular disease 2%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationinfection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information			
	Uncerta	Povidone inty in potential benefits :	iodine spray	arch is needed.				
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			



RCT					
<u>Seet et al</u> , ²⁹⁴ peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 735 assigned to povidone iodine spray 3 times a day for 42 days and 619 assigned to SOC (vitamin C)	Mean age 33, male 100%, hypertension 1%, diabetes 0.3%	NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty \oplus \(\circ>\)Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationGymptomatic informationSymptomatic informationSymptomatic informationSymptomatic infection (prophylaxis studies): Very low certainty \oplus \(\circ>\)Adverse events: Very low certainty \oplus \(\circ>\)Hospitalization: Very low certainty \oplus \(\circ>\)
Probiotics ma	y increase symptom reso		biotics The effect on other of	utcomes is uncertain. Further r	esearch is needed.
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Wang et al</u> ; ⁵²⁵ peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 98 assigned to probiotics 2 lozenges a day for 30 days and 95 assigned to SOC	Mean age 36 ± 8, male 29%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$



PROCOV-19-2020 trial; ⁵²⁶ Ivashkin et al; peer reviewed; 2021	Patients with moderate to critical COVID-19 infection. 99 assigned to probiotics three times a day for 14 days and 101 assigned to SOC	Mean age 64 ± , male 46%	NR	study. Concealment of allocation is probably inappropriate. High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): RR 1.89 (95%CI 1.4 to 2.56); RD 53.9.8% (95%CI 24.2% to 94.5%); Low certainty ⊕⊕○○
	Individuals exposed to SARS-CoV-2 infection. 91 assigned to probiotics 1 capsule a day for 28 days and 91 assigned to SOC	Age 18-64 62%, male 36.8%, hypertension 12.1%, diabetes 3.8%, COPD 1.1%, cancer 2.7%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Adverse events: No information Hospitalization: No information
<u>ABB-COVID19</u> <u>trial</u> ; ⁵²⁸ Gutiérrez- Castrellón et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 147 assigned to probiotics 1 capsule a day for 30 days and 146 assigned to SOC	Median age 37 ± , male 46.3%, hypertension 19.6%, diabetes 10.3%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>Saviano et al</u> ; ⁵²⁹ peer reviewed; 2022	Patients with severe COVID-19 infection. 40 assigned to probiotics (<i>Bifidobacterium lactis</i> LA 304, <i>Lactobacillus</i> <i>salivarius</i> LA 302)and <i>Lactobacillus</i> <i>acidophilus</i> LA 201) twice a day for 10 days and 40 assigned to SOC	Mean age 59.6, male 55%, hypertension 38.7%, diabetes 17.5%, COPD 8.7%	Corticosteroids 100%; vaccinated 18.7%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
		0	esterone		

Uncertainty in potential benefits and harms. Further research is needed.



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			·		
Ghandehari et al; ⁵³⁰ preprint; 2020	Patients with severe COVID-19. 18 assigned to progesterone 100 mg twice a day for 5 days and 22 assigned to standard of care	Mean age 55.3 ± 16.4, male 100%, hypertension 48%, diabetes 25%, obesity 45%	Corticosteroids 60%, remdesivir 60%, hydroxychloroquine 2.5%, tocilizumab 12.5%, azithromycin 50%, convalescent plasma 5%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: No information
	Uncertai	Prol inty in potential benefits a	ectin-M and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Prolectin-M trial</u> ; ⁵³¹ Sigamani et al;	Patients with mild COVID-19. 5 assigned	Mean age 28.5 ± 3.85, male 20%	NR	High for mortality and mechanical ventilation;	Mortality: No information





preprint; 2020	to prolectin-M 40 g a day and 5 assigned to standard of care	Pr inty in potential benefits a	opolis	high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Bee-Covid trial</u> ; ⁵³² Duarte Silveira et al; Preprint; 2020	Patients with moderate to critical COVID-19. 82 assigned to propolis 400–800 mg a day for 7 days and 42 assigned to SOC	Mean age 50 ± 12.8, male 69.4%, hypertension 45.2%, diabetes 21%, COPD 7.3%, asthma %, obesity 51.6%	Corticosteroids 80.6%, hydroxychloroquine 3.2%, azithromycin 95.2%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty \bigoplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \bigoplus \bigcirc \bigcirc Symptom resolution or improvement: Very low certainty \bigoplus \bigcirc \bigcirc

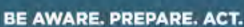




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	Uncertai	Pros inty in potential benefits a	tacyclin Ind harms. Further resea	rch is needed.	Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
COMBAT- COVID trial; ⁵³³ Johansson et al; peer reviewed; 2021	Patients with critical COVID-19 infection. 41 assigned to prostacyclin 1 ng/kg/min for 3 days and 39 assigned to SOC	Mean age 67, male 66.2%, hypertension 61.2%, COPD 12.5%, CKD 2.5%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○





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					Hospitalization: No information				
Prostacyclin (inhaled) Uncertainty in potential benefits and harms. Further research is needed.									
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence				
RCT	•	-			<u>.</u>				
ThIlo trial; ⁵³⁴ Haeberle et al; preprint; 2021	Patients with critical COVID-19 infection. 72 assigned to prostacyclin (inhaled) 3 times a day for 5 days and 72 assigned to SOC	Mean age 60, male 75%, hypertension 58.6%, diabetes 28.5%, COPD 7.6%, asthma 4.9%, CKD 6.9%, cancer 2.8%	Corticosteroids 51.4%, remdesivir 42.4%, tocilizumab 16%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: RR 1.05 (95%CI 0.64 to 1.7); RD 0.8% (95%CI - 5.7% to 11.2%); Low certainty ⊕⊕○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No				
		Prove	lutamide	<u> </u>					
	Uncertai	inty in potential benefits a		urch is needed.					
Study; publication status	Patients and interventions	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care				



	analyzed				and GRADE certainty of the evidence
RCT					
<u>Cadegiani et al</u> ; ⁵³⁵ Preprint; 2020	Patients with mild COVID-19. 114 assigned to proxalutamide 200 mg a day for 15 days and 100 assigned to SOC	NR	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Randomization and concealment methods probably not appropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very
<u>004 trial</u> ; ⁵³⁶	Patients with mild to moderate COVID-19 infection. 171 assigned to proxalutamide 200 mg a day for 15 days and 65 assigned to SOC	Mean age 45.3 ± 13, male 54.2%, hypertension 22.5%, diabetes 8.9%, COPD 0%, asthma 5%, CKD 0.4%, cancer 17%, obesity 15.7%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Concealment of allocation and blinding probably inappropriate.	low certainty ⊕○○○ Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic
<u>KP-DRUG-SARS-</u> <u>003 trial;⁵³⁷</u> Cadegiani et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 423 assigned to proxalutide 300 mg a day for 14 days and 355 assigned to SOC	Median age 51 ± , male 59.6%, hypertension 27.6%, diabetes 12.5%, COPD 2.3%, asthma %, CHD %, CKD 0%	Steroids 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Randomization scheme was modified during the study.	infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: RR
AB-DRUG-SARS- 005 trial, ⁵³⁸ Cadegiani et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 75 assigned to proxalutamide 200 mg a day for 7 days and 102 assigned to SOC	Mean age 44.2 ± 12.1, male 0%, hypertension 31.1%, diabetes 8.5%, COPD 0.6%, obesity 18.1%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Randomization process presented as "Blocked" but described	Hospitalization: RR 0.07 (95%CI 0.01 to 0.52); RD -4.5% (95%CI -4.7% to - 2.3%); Very low certainty ⊕○○○

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				as a cluster randomization.					
Pyridostigmine Uncertainty in potential benefits and harms. Further research is needed.									
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence				
RCT		-							
PISCO trial; ⁵³⁹ Fragoso-Saavedra et al; preprint; 2021	Patients with moderate to severe COVID-19 infection. 94 assigned to pyridostigmine 60 mg a day for 14 days and 94 assigned to SOC	Median age 52 ± 20, male 59.6%, hypertension 35.1%, diabetes 36.2%, COPD 4.3%, asthma %, CHD 2.1%, obesity 43.1%	Corticosteroids 74.5%, tocilizumab 5.3%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation and blinding probably inappropriate.	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or improvement: Very low certainty \oplus \bigcirc \bigcirc Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: No information				
	Uncerta	QU inty in potential benefits a	ercetin and harms. Further resea	urch is needed.					
Study; publication status	Patients and interventions	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care				



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	analyzed				and GRADE certainty of the evidence
RCT				•	
<u>Onal et al</u> , ⁵⁴⁰ peer review; 2020	Patients with moderate to severe COVID-19. 49 assigned to quercetin 1000 mg and 380 assigned to SOC	Age > 50 65.7%, male 56.6%, hypertension 38.7%, diabetes 28.2%, COPD 6%, asthma 13.9%, CHD 22.6%, CKD 0.2%, cancer 3.6%, obesity 0.9%	Hydroxychloroquine 97.5%, favipiravir 13.2%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Randomization and concealment process probably inappropriate. Non-blinded study.	Mortality: Very low certainty ⊕○○○
<u>Di Pierro et al</u> ; ⁵⁴¹ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 21 assigned to quercetin 400- 600 mg a day for 14days and 21 assigned to SOC	Mean age 49.3 ± 19.5, male 47.6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic
Shohan et al; ⁵⁴² peer reviewed; 2022	Patients with severe to critical COVID-19 infection. 30 assigned to quercetin 1000 mg a day for 7 days and 30 assigned to SOC	Mean age 51.8, male 56.6%, hypertension 20%, asthma 6.6%, CHD 15%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes:	infection (prophylaxis studies): Very low certainty ⊕○○○ Adverse events: No information
<u>Rondanelli et al</u> ; ⁵⁴³ peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 60 assigned to quercetin 500 mg a day and 60 assigned to SOC	Mean age 49.3 ± 12.9, male 52.5%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Hospitalization: Very low certainty ⊕○○○

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Raloxifene Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT								
Nicastri et al; ⁵⁴⁴ peer reviewed; 2021	Patients with moderate COVID-19 infection. 42 assigned to raloxifene 60 to 120 mg for 14 days and 19 assigned to SOC	Mean age 56.7 ± 10.1, male 54.1%, hypertension 26.2%, diabetes 0.66%, COPD %, asthma 1.6%	Corticosteroids 14.7%, remdesivir 1.6%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: No information			
			mipril					
	Uncerta	inty in potential benefits a	ind harms. Further resea	irch is needed.				
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			





RASTAVI trial; ⁵⁴⁵ Amat-Santos et al; preprint; 2020	Individuals exposed to SARS-CoV-2 infection. 50 assigned to ramipril 2.5 mg a day progressively increased to 10 mg a day and 52 assigned to standard of care	Mean age 82.3 ± 6.1, male 56.9%, hypertension 54.15%, diabetes 20.65%, chronic lung disease 7.35%, coronary heart disease 22.45%, chronic kidney disease 34.15%, cerebrovascular disease 11.15%	NR	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationGymptomatic infection (prophylaxis studies): Very low certainty ⊕○○○Adverse events: No informationHospitalization: No information
	Uncerta	RD-X19 (l inty in potential benefits a	ight therapy) and harms. Further resea	urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			l	L	
<u>EB-P12-01 trial;</u> ⁵⁴⁶ Stasko et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 20 assigned to RD-X19 light dose of 16 J/cm2 twice a day and 11 assigned to SOC	Median age 40 ± 20.6, male 52%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very



		combinant super			low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT		L		L	
Li et al; ⁵⁴⁷ peer- reviewed; 2020	Patients with moderate to severe COVID-19 infection. 46 assigned to recombinant super- compound interferon 12 million IU twice daily (nebulization) and 48 assigned to interferon alfa	Median age 54 ± 23.5, male 46.8%, hypertension 19.1%, diabetes 9.6%, chronic lung disease 1.1%, coronary heart disease 7.4%, cerebrovascular disease 5.3%, liver disease 6.4%	Corticosteroids 9.6%, ATB 22.3%, intravenous immunoglobulin 3.2%, lopinavir-ritonavir 44.7%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No



Regdabivimab may			n mortality and mech	ibody) anical ventilation are uncerta	information Hospitalization: No information in. Further research is
Study; publication status	Patients and interventions analyzed	Comorbidities	eeded. Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Streinu-Cercel et</u> <u>al;⁵⁴⁸ Peer reviewed;</u> 2021	Patients with mild to moderate COVID-19 infection. 204 assigned to regdanvimab 40- 80 mg/kg once and 103 assigned to SOC	Mean age 51 ± 20, male 44.6%, comorbidities 73%	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty
<u>CT-P59 1.2 trial</u> ; ⁵⁴⁹ Kim et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 15 assigned to regdanvimab 20 to 80 mg once and 3 assigned to SOC	Median age 52 ± 8, male 100%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	 ⊕○○○ Symptom resolution or improvement: RR 1.24 (95%CI 1.05 to 1.46); RD 4.2% (95%CI 9% to 80%); Low certainty ⊕⊕○○ Symptomatic infection (prophylaxis studies): No information
					Adverse events: Very low certainty ⊕○○○ Hospitalization: Very low certainty



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REGEN-COV (casirivimab and imdevimab) REGEN-COV probably reduces mortality and mechanical ventilation in seronegative severe to critical patients. In mild patients REGEN-COV probably reduces hospitalizations and in exposed individuals it reduces symptomatic infections.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT								
Weinreich et al; ⁵⁵⁰ preprint; 2020	Patients with recent onset mild disease with risk factors for severe COVID-19 infection. 2091 assigned to REGEN-COV (casirivimab and imdevimab) 1.2 to 2.4 g single infusion and 2089 assigned to SOC	Median age 50 ± 21, male 48.7%, obesity 58%, comorbidities 100%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: RR 0.83 (95%CI 0.63 to 1.09); RD -2.7% (95%CI - 5.9% to 1.4%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Mortality (seronegative): RR 0.79 (95%CI 0.71 to 0.89); RD -3.2% (95%CI -4.6% to -			
<u>RECOVERY -</u> <u>REGEN-COV</u> <u>trial;</u> ⁵⁵¹ Horby et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 4839 assigned to REGEN- COV (Regeneron) 8 g once and 4946 assigned to SOC	Mean age 61.9 ± 14.4, male 63%, diabetes 26.5%, COPD %, CHD 21%, CKD 5%	Corticosteroids 94%, azithromycin 3%, baricitinib 9%; vaccinated 8%	Low for mortality and mechanical ventilation; some Concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	 1.8%); Moderate certainty ⊕⊕⊕○ Invasive mechanical ventilation: RR 0.79 (95%CI 0.54 to 1.14); RD -3.6% (95%CI -8% to 2.4%); Low certainty ⊕⊕○○ Invasive mechanical ventilation (seronegative): RR 			
<u>O'Brien et al</u> ; ⁵⁵² peer reviwed; 2021	Patients with early asymptomatic COVID-19 infection. 100 assigned to REGEN-COV (Regeneron) 1.2 g once and 104 assigned	Mean age 40.9 ± 18, male 45.4%, diabetes 7.8%, CKD 2.5%, immunosuppressive therapy 1.5%, obesity 13.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	0.82 (95%CI 0.74 to 0.9); RD -3.1% (95%CI -4.5% to - 1.7%); Moderate certainty ⊕⊕⊕⊖ Symptom			

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O'Brien et al; ⁵⁵³ peer reviewed; 2021	to SOC Individuals exposed to SARS-CoV-2 infection. 841 assigned to REGN-COV2 (Regeneron) 1200 mg once and 842 assigned to SOC	Median age 43 ± 25, male 45.9%, 6.8%, CKD 1.9%, immunosuppresive therapy 1%, obesity 34.1%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	resolution or improvement: RR 1.06 (95%CI 1 to 1.12); RD 3.6% (95%CI 0% to 7.2%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Symptom resolution or
OPTIMISE-C19 <u>trial;</u> ⁹³ McCreary et al; peer reviewed; 2022	Patients with mild COVID-19 infection disease and risk factors for severity. 922 assigned to REGN- CoV2 (Regeneron) and 1013 assigned to bamlanivimab +/- etesevimab	Mean age 56 ± 16, male 46%, hypertension 53%, diabetes 25%, COPD 19%, asthma %, CHD 18%, CKD 6.5%, immunosuppresive therapy 27%, obesity 48%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	improvement (seronegative): RR 1.1 (95%CI 1.06 to 1.14); RD 6% (95%CI 3.6% to 8.5%); Moderate certainty ⊕⊕⊕⊖ Symptomatic infection
Somersan-Karakaya <u>et al</u> ; ⁵⁵⁴ peer- reviewed; 2021	Patients with moderate to severe COVID-19 infection. 804 assigned to REGN-COV2 (Regeneron) 2.4 to 8 gr once and 393 assigned to SOC	Median age 62 ± , male 54.1%	Corticosteroids 74.8%, remdesivir 54.9%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	(prophylaxis studies): RR 0.24 (95%CI 0.08 to 0.76); RD -13.2% (95%CI - 16% to -4.2%); High certainty ⊕⊕⊕⊕ Adverse events: RR 0.51 (95%CI 0.38 to
<u>R10933-10987-</u> <u>COV-20145 trial;</u> ⁵⁵⁵ Portal Celhay et al; preprint; 2021	Patients with mild COVID-19 infection. 584 assigned to REGN-COV2 (Regeneron) 300 - 2400 mg once and 77 assigned to SOC	Mean age 34.6, male 44.3%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	0.67); RD -5% (95%CI -6.3% to - 3.4%); Moderate certainty ⊕⊕⊕○ Hospitalization: RR 0.28 (95%CI 0.19 to 0.42); RD -3.5%
<u>Isa et al</u> ; ⁵⁵⁶ preprint; 2021	Patients with COVID- 19 infection. assigned to REGN-COV2 (Regeneron) and assigned to	Median age 48 ± 22, male 55.1%, hypertension 14.7%, asthma 5.2%, CHD 0.8%, CKD 0.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	(95%CI -3.9% to - 2.8%); Moderate certainty ⊕⊕⊕⊖





<u>Weinreich et al</u> , ⁵⁵⁷ preprint; 2021	Patients with mild to moderate COVID-19 infection. 434 assigned to REGN-COV2 (Regeneron) 2400 TO 8000 mg once and 231 assigned to SOC	Median age 42 ± 21, male 47.1%, obesity 37.3%, Risk factor for hospitalization 60.5%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events				
<u>OPTIMISE-C19</u> <u>trial</u> ; ⁵⁵⁸ Huang et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 2454 assigned to REGN- COV2 (Regeneron) one infusion and 1104 assigned to sotrovimab one infusion	Mean age 54 ± 18, male %, hypertension 30%, diabetes 12%, CHD 16%, CKD 4.7%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events				
MANTICO trial; ⁹⁶ Mazzaferri et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 107 assigned to sotrovomab 500 mg once and 106 assigned to bamlanivimab + etesevimab 700/1400 mg once and 106 assigned to REGEN- COV2 600/600 mg once		Vaccinated 28.6%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.				
PLATCOV - Regen trial; ³⁸¹ Schilling et al; peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 10 assigned to REGEN-COV 1200 mg once and 41 assigned to SOC	Mean age 27 , male 39%	Corticosteroids %, remdesivir %, hydroxychloroquine %, lopinavir-ritonavir %, tocilizumab %, azithromycin %, convalescent plasma %; Vaccinated %	Low for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.				
	Remdesivir n hospitalized patients with moderate to critical disease, remdesivir probably reduces mortality and mechanical ventilation, and it may improv time to symptom resolution without increasing severe adverse events. In patients with recent onset mild COVID-19, it may reduce hospitalizations. However, the certainty is low because of risk of bias and imprecision.							
Study; publication	Patients and	Comorbidities	Additional	Risk of bias and study	Interventions effects			



status	interventions analyzed		interventions	limitations	vs standard of care and GRADE certainty of the evidence
RCT					
ACTT-1 trial; Beigel et al; ⁵⁵⁹ peer- reviewed; 2020	Patients with mild to critical COVID-19 infection. 541 assigned to remdesivir intravenously 200 mg loading dose on day 1 followed by a 100 mg maintenance dose administered daily on days 2 through 10 or until hospital discharge or death and 522 assigned to standard of care	Mean age 58.9 ± 15, male 64.3%, hypertension 49.6%, diabetes 29.7%, chronic lung disease 7.6%, coronary heart disease 11.6%,	NR	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: RR 0.93 (95%CI 0.89 to 1.03); RD -1.1% (95%CI - 1.8% to 0.5%); Moderate certainty ⊕⊕⊕○ Invasive mechanical ventilation: RR 0.76 (95%CI 0.56 to 1.04); RD -4.2% (95%CI - 7.6% to 0.7%); Moderate certainty ⊕⊕⊕○
SIMPLE trial; Goldman et al; ⁵⁶⁰ peer-reviewed; 2020	Patients with severe COVID-19 infection. 200 assigned to remdesivir (5 days) 200 mg once followed 100 mg for 5 days and 197 assigned to remdesivir (10 days)	Median age 61.5 ± 20, male 63.7%, hypertension 49.8%, diabetes 22.6%, asthma 12.3%	NR	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Symptom resolution or improvement: RR 1.1 (95%CI 0.96 to 1.28); RD 6% (95%CI -2.4% to 17%); Low certainty ⊕⊕⊖⊖ Symptomatic infection (prophylaxis studies): No
<u>CAP-China</u> <u>remdesivir 2 trial;⁵⁶¹</u> Wang et al; peer- reviewed; 2020	critical COVID-19	Median age 65 ± 7.5, male 60.5%, hypertension 43%, diabetes 23.7%, coronary heart disease 7.2%	Corticosteroids 65.6%, lopinavir-ritonavir 28.4%, IFN 32.2%, ATB 91.1%	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	information Severe Adverse events: RR 0.77 (95%CI 0.46 to 1.29); RD -2.3% (95%CI - 5.5% to 3%); Low certainty ⊕⊕○○ Hospitalization: RR 0.28 (95%CI 0.11 to



SIMPLE 2 trial; Spinner et al; ⁵⁶² peer-reviewed; 2020	Patients with moderate COVID-19 infection. 384 assigned to remdesivir 200 mg on day 1 followed by 100 mg a day for 5 to 10 days and 200 assigned to standard of care	Median age 57 ± 9, male 61.3%, hypertension 42%, diabetes 40%, asthma 14%, coronary heart disease 56%	Corticosteroids 17%, hydroxychloroquine 21.33%, lopinavir- ritonavir 11%, tocilizumab 4%	Some concerns for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Additional treatments unbalanced between arms which suggests that patients might have been treated differently.	0.75); RD -3.4% (95%CI -4.3% to - 1.2%); Low certainty ⊕⊕⊖⊖
WHO SOLIDARITY; ²⁸¹ Pan et al; peer reviewed; 2020	Patients with moderate to critical COVID-19 infection. 4146 assigned to remdesivir 200 mg once followed by 100 mg a day for 10 days and 4129 assigned to SOC	Age range 50 – 69 years old 46.2%, male 63.4%, diabetes 27.2%, COPD 6.8%, asthma 5.9%, CHD 22.5%	Steroids 67.7%, convalescent plasma 3.3%, Anti IL6 4.5%	Low for mortality and mechanical ventilation; some Concerns for symptom resolution, infection and adverse events Notes: Non-blinded study wich might have introduced bias to symptoms and adverse events outomes results.	
<u>Mahajan et al</u> ; ⁵⁶³ peer reviewed; 2021	Patients with mild to severe COVID-19 infection. 34 assigned to remdesivir 200 mg once followed by 100 mg once a day for 5 days and 36 assigned to SOC	Mean age 57.7 ± 13.1, male 65.5%, hypertension 45.7%, diabetes 60%, asthma 1.4%, CHD 12.9%, CKD 4.3%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Abd-Elsalam et al</u> ; ⁵⁶⁴ peer reviewed; 2021	moderate COVID-19	Mean age 53 ± 15, male 59.5%, hypertension 33%, diabetes 34%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	





	days and 100 assigned to SOC			Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>Sarhan et al</u> ; ⁵⁶⁵ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 52 assigned to remdesivir 200 mg once followed by 100 mg a day for 5 days plus tocilizumab and 56 assigned to HCQ 400 mg once followed by 200 mg a day for 5 days plus tocilizumab	Mean age 57, male 72%, hypertension 61.7%, diabetes 47.6%, COPD 2.8%, asthma 13.1%, CHD 21.5%, CKD 4.7%,	Hydroxychloroquine 52.3%, tocilizumab 100%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>PINETREE trial</u> ; ⁵⁶⁶ Gottlieb et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 279 assigned to remdesivir 200 mg once followed by 100 mg on days two and three and 283 assigned to SOC	Mean age 50 ± 15, male 53.1%, hypertension 47.7%, diabetes 61.6%, COPD 24%, CKD 3.2%, immunosuppresion 4.1%, cancer 5.3%, obesity 55.2%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>CATCO trial</u> ; ⁵⁶⁷ Ali et al; peer reviewed; 2021	Patients with moderate to critical COVID-19 infection. 170 assigned to remdesivir 200 mg once followed by 100 mg a day for 10 days and 153 assigned to SOC	NR	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
	Uncerta	Remdesiv inty in potential benefits a	v ir (inhaled) Ind harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence





RCT					
Gilead et al; <u>NCT04539262;</u> other; 2021	Patients with mild to moderate COVID-19 infection. 109 assigned to remdesivir (inh) 31 to 62 mg a day for 3 to 5 days and 45 assigned to SOC	Age > 60 years old 12.9%, male 50%	NR	NA	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationSevere Adverse events: No informationHospitalization: Very low certainty ⊕○○○
	Uncertai	Rep inty in potential benefits a	Darixin Ind harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>REPAVID-19</u> <u>trial</u> ; ⁵⁶⁸ Landoni et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 36 assigned to reparixin 3600 mg a day for 7 days and 19 assigned to SOC	Mean age 61.7, male 76.4%, hypertension 43.6%, diabetes 23.6%, COPD %, CHD 12.7%, CKD 7.3%, obesity 20%	Corticosteroids 92.7%, remdesivir 23.6%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○ Invasive mechanical ventilation: Very low certainty





				Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	 ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Severe Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncerta	Rese inty in potential benefits a	veratrol and harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	<u>I</u>				
<u>McCreary et al</u> ; ⁵⁶⁹ peer-reviewed; 2021	Patients with mild COVID-19 infection. 50 assigned to resveratrol 4 g a day for 7 days and 50 assigned to SOC	Mean age 56 ± 9, male 43%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty
<u>Reszinate trial</u> ; ⁵⁷⁰ Kaplan et al; preprint; 2021	Patients with mild COVID-19 infection. 14 assigned to resveratrol + zinc 4000/150 mg once a day for five days and 16 assigned to SOC	Mean age 42.4, male 40%	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	 OO Symptom resolution or improvement: No information Symptomatic

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					<pre>infection (prophylaxis studies): No information Severe Adverse events: Very low certainty ⊕○○○ Hospitalization: Very low certainty ⊕○○○</pre>
		G-CSF (in patien inty in potential benefits a			
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			•		
Cheng et al, ⁵⁷¹ peer- reviewed; 2020	Patients with moderate to severe COVID-19 and lymphopenia. 100 assigned to rhG-CSF six doses and 100 assigned to standard of care	Mean age 45 ± 15, male 56%	Lopinavir-ritonavir 15.5%, IFN 9%, umifenovir 18%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Severe Adverse events: Very low



					certainty ⊕○○○ Hospitalization: No information					
	rhG-CSF (inhaled) Uncertainty in potential benefits and harms. Further research is needed.									
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence					
RCT	-			-						
SARPAC trial; ⁵⁷² Lambrecht et al; preprint; 2021	Patients with severe COVID-19 infection. 40 assigned to rhG- CSF (inhaled) 125 µg twice daily for 5 days and 41 assigned to SOC	Mean age 60 ± 20, male 61%, hypertension 17.1%, diabetes 17.1%, CHD 2.4%, CKD 2.4%, cancer 4.9%	Corticosteroids 22%, hydroxychloroquine 63.4%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanica ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events:					



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
BTI-202 trial; ⁵⁷³ DiNubile et al; peer reviewed; 2022	Patients with severe COVID-19 infection. 31 assigned to rhu- pGSN 12 mg/kg three times and 30 assigned to SOC	Mean age 62.1 ± 11.6, male 57.4%, hypertension 41%, diabetes 32.8%	Corticosteroids 100%, remdesivir 98.4%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: Very low certainty ⊕○○○Symptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
	Uncerta	Rib inty in potential benefits a	avirin and harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Chen et al</u> ; ⁴⁰⁵ preprint; 2020	Patients with mild to moderate COVID-19	Mean age 42.5 ± 11.5, male 45.5%	NR	High for mortality and invasive mechanical	Mortality: No information



	infection. 33 assigned to ribavirin 2 g IV loading dose followed by orally 400-600 mg every 8 h for 14 days,			ventilation; high for symptom resolution, infection, and adverse events	Invasive mechanical ventilation: No information
	36 assigned to lopinavir-ritonavir and 32 assigned to ribavirin plus lopinavir-			Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Symptom resolution or improvement: No information
	ritonavir				Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
	Uncertai	Ribavirin plus i inty in potential benefits a	interferon beta- nd harms. Further resea		
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					





Hung et al; ⁵⁷⁴ peer- reviewed; 2020	Patients with mild to moderate COVID-19 infection. 86 assigned to ribavirin plus interferon beta-1b 400 mg every 12 hours (ribavirin), and subcutaneous injection of one to three doses of interferon beta-1b 1 mL (8 million international units [IU]) on alternate days, for 14 days and 41 assigned to standard of care	Median age 52 ± 15, male 54%, hypertension 18.3%, diabetes 13.3%, coronary heart disease 7.9% cerebrovascular disease 1.5%, cancer 1.5%	Corticosteroids 6.2%, ATB 53.3%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic informationAdverse events: No informationAdverse events: No informationHospitalization: No information
Ru	xolitinib may reduce mo		olitinib ainty of the evidence was	low. Further research is n	eeded.
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Cao et al</u> ; ⁵⁷⁵ peer- reviewed; 2020	Patients with severe COVID-19 infection. 22 assigned to ruxolitinib 5 mg twice a day and 21 assigned to standard of care	Mean age 63 ± 10, male 58.5%, hypertension 39%, diabetes 19.5%, coronary heart disease 7.3%,	Corticosteroids 70.7%, IVIG 43.9%, umifenovir 73%, oseltamivir 27%	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: RR 0.72 (95%CI 0.59 to 0.89); RD -4.5% (95%CI - 6.5% to -1.7%); Low certainty ⊕⊕○○ Invasive mechanical
<u>RUXCOVID</u> <u>trial; 576</u> Han et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection.	Mean age 56.5 ± 13.3, male 54%, diabetes 21.9%, obesity 47%	NR	Low for mortality and mechanical ventilation; low for symptom	ventilation: Very low certainty ⊕○○○



RUXCOVID- DEVENT trial; NCT04377620; other; 2021	287 assigned to ruxolitinib 10 mg a day for 14 to 28 days and 145 assigned to SOC Patients with critical COVID-19 infection. 164 assigned to ruxolitinib 10 to 30 mg a day and 47 assigned to SOC	Mean age 63.4 ± 12.7, male 64.9% Sabi inty in potential benefits a	NR Zabulin and harms. Further resea	resolution, infection and adverse events Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Symptom resolution or improvement: RR 1.05 (95% CI 0.89 to) 1.24); RD 3% (95% CI -6.7% to 14.5%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	-	<u> </u>			
Barnette et al; ⁵⁷⁷ peer reviewed; 2022	Patients with severe COVID-19 infection. 98 assigned to sabizabulin 9 mg for up to 21 days and 52 assigned to SOC	Mean age 59.7 ± 14.7, male 68%, hypertension 60%, diabetes 37.3%, COPD %, CHD 4.7%, CKD 10%, cancer 5.3%, obesity 32.4%	Corticosteroids 82.7%, remdesivir 32.7%, tocilizumab 10%, baricitinib 12%; vaccinated 44.7%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information



					infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
Sarilumab may re	duce mortality and mecl		ilumab ements; however, the cer	tainty of the evidence is lov	w. Further research is
·			eeded.		
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>REMAP-CAP -</u> <u>tocilizumab trial;</u> ⁵⁷⁸ Gordon et al; peer- reviewed; 2020	Patients with severe to critical COVID-19 infection. 353 assigned to TCZ 8 mg/kg once or twice, 48 assigned to sarilumab 400 mg once and 402 assigned to SOC	CHD 10.2%,	Corticosteroids 75.6%, remdesivir 32.8%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have	Mortality: RR 0.97 (95%CI 0.81 to 1.16); RD -0.5% (95%CI - 3% to 2.6%); Low certainty ⊕⊕⊖⊖ Invasive mechanical ventilation: RR 0.98
				introduced bias to symptoms and adverse events outcomes results.	(95%CI 0.68 to 1.42); RD -0.3% (95%CI - 5.5% to 7.3%); Low
<u>Lescure et al</u> ; ⁵⁷⁹ peer-reviewed; 2020	Patients with severe to critical COVID-19. 332 assigned to sarilumab 200-400 mg once and 84 assigned to SOC	Mean age 59 ± 18, male 62.7%, hypertension 42.5%, diabetes 26.4%, COPD 4.3%, asthma 4.1%, CHD 5.3%, CKD 4.3%, cancer 10.1%, obesity 20.7%	Corticosteroids 46.4%, hydroxychloroquine 34.5%, azithromycin 46.4%,	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	certainty ⊕⊕⊖⊖ Symptom resolution or improvement: RR 1.02 (95%CI 0.97 to 1.06); RD 1.2% (95%CI -1.8% to
<u>Sarilumab-</u> COVID19 Study	Patients with severe to critical COVID-19	Critical patient population: mean age 61	Corticosteroids 34.3%,	Low for mortality and mechanical ventilation;	3.6%); Moderate certainty ⊕⊕⊕⊖



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<u>trial</u> ; ⁵⁸⁰ Sivapalasingam, et al; preprint; 2021 (two studies reported) <u>CORIMUNO-</u>	infection. 1148 assigned to sarilumab 200-400 mg once and 376 assigned to SOC Patients with	± 20, male 68.4%, hypertension 52.1%, diabetes 18.7%, obesity 46.5% Median age 62, male %,	Steroids 20.1%,	Low for symptom resolution, infection, and adverse events Low for mortality and	Symptomatic infection (prophylaxis studies): No information
<u>SARI trial</u> ; ⁵⁸¹ Mariette, et al, peer reviewed; 2021	moderate to severe COVID-19 infection. 68 assigned to sarilumab 400 mg once and 76 assigned to SOC	hypertension 25.1%, diabetes 30.5%, COPD 6.3%, asthma 8%, CKD 11.8%, cancer 3%,	remdesivir 0%, hydroxychloroquine 14.6%, azithromycin 39.6%	mechanical ventilation; high for symptom resolution, infection, and adverse events	Severe adverse events: RR 1.03 (95%CI 0.91 to 1.17); RD 0.3% (95%CI - 0.9% to 1.7%); Moderate certainty
<u>CORIMUNO-</u> <u>SARI ICU trial;⁵⁸²</u> Hermine et al; peer reviewed; 2021	Patients with critical COVID-19 infection. 48 assigned to sarilumab 400 mg once and 33 assigned to SOC	Median age 61, male 76.5%, diabetes 31.2%, COPD 3.7%, asthma 4.9%, CKD 13.5%, cancer 1.2%,	Steroids 19.7%, remdesivir 0%, hydroxychloroquine 4.9%, lopinavir- ritonavir 1.2%, azithromycin 2.5%, convalescent plasma 0%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	⊕⊕⊕○ Hospitalization: No information
SARCOVID trial; ⁵⁸³ García Vicuña et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 20 assigned to sarilumab 400 mg once and 10 assigned to SOC	Median age 61.5, male 67%, hypertension 43%, diabetes 17%, COPD 7%, CHD 10%, CKD 13%, obesity 10%	Steroids 83%, remdesivir 0%, hydroxychloroquine 20%, lopinavir- ritonavir 17%, tocilizumab %, azithromycin 60%, convalescent plasma 0%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
SARICOR trial; ⁵⁸⁴ Merchante et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 76 assigned to sarilumab 200-400 mg once and 39 assigned to SOC	Median age 59, male 68%, hypertension 41%, diabetes 15%, COPD 13%, CHD 4%, CKD 2%,	Steroids 90%, remdesivir 12%, convalescent plasma 0%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded	





<u>SARTRE trial</u> ; ⁵⁸⁵ Sancho-Lopez et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 99 assigned to sarilumab 200-400 mg once and 102 assigned to SOC	Median age 60, male 70.2%, hypertension 40.8%, diabetes 16.4%, COPD 9.5%, CHD 12.4%, CKD 3%, cancer 3%, obesity 3.5%	Steroids 100%, remdesivir 1%, convalescent plasma 0%	study which might have introduced bias to symptoms and adverse events outcomes results. Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
IRB 3305 trial; ⁵⁸⁶ Branch-Elliman et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 20 assigned to sarilumab 200 to 400 mg (subcutaneous) once and 30 assigned to SOC	Mean age 72.3 ± 12.7, male 92%, hypertension 86%, diabetes 50%, COPD 32%, asthma 16%, CHD 70%, CKD 18%, cancer 48%, obesity 62%	Corticosteroids 86%, remdesivir 80%, hydroxychloroquine 4%, tocilizumab 2%, convalescent plasma 2%;	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
	Uncerta	Secul inty in potential benefits a	K inumab Ind harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	•				
BISHOP trial; ⁵⁸⁷ Gomes Resende et al; peer reviewed; 2022	Patients with severe COVID-19 infection. 25 assigned to secukinumab 300 mg once and 23 assigned to SOC	Mean age 54 ± 21.5, male 52%, hypertension 48%, diabetes 34%, CHD 8%, obesity 48%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Symptom





				symptoms and adverse events outcomes results.	resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Severe adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncerta	nty in potential benefits a	nicapoc and harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>COVIPOC trial</u> ; ⁵⁸⁸ Granfeldt et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 20 assigned to senicapoc 50 mg twice and 26 assigned to SOC	Median age 66, male 65.2%, hypertension 34.8%, diabetes 28.3%, COPD 26%, CKD 4.5%, cancer 15.2%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information





status inter RCT Panatto et al; ⁵⁸⁹ peer Patie		Sen nty in potential benefits a Comorbidities	ntinox and harms. Further re Additional interventions	esearch is needed. Risk of bias and study limitations	Interventions effects
status inter RCT Panatto et al; ⁵⁸⁹ peer Patie		Comorbidities			
<u>Panatto et al</u> ; ⁵⁸⁹ peer Patie	L. L				vs standard of care and GRADE certainty of the evidence
-					
36 as 0.005	VID-19 infection. ssigned to sentinox 95% 3 to 5 times a and 18 assigned to	Mean age 40.1 ± 13.7, male 81%, any commorbidities 4%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationSevere adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Hospitalization: $\forall e \bigcirc \bigcirc \bigcirc$



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Tian et al</u> ; ⁵⁹⁰ peer reviewed; 2021	Patients with moderate COVID-19 infection. 27 assigned to short-wave diathermy and 13 assigned to SOC	Median age 65 ± 18, male 62.5%, hypertension 30%, diabetes %, COPD 45%, CHD 30%, CKD 7.5%, cerebrovascular disease 27.5%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Concealment of allocation and blinding probably inappropriate.	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No information Severe adverse events: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Hospitalization: No information
	Uncerta	Silc inty in potential benefits a	lenafil nd harms. Further resea	arch is needed.	
Study; publication tatus	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence



UNAB-003 trial; ⁵⁹¹ Santamarina et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 20 assigned to sildenafil 75 mg a day for 7 days and 20 assigned to SOC	Median age 57, male 82.5%, diabetes 20%, COPD 0%, asthma 5%	Corticosteroids 82.5%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Blinding and concealment of allocation probably inappropriate.	Mortality: Very low certainty @OOOInvasive mechanical ventilation: Very low certainty @OOOSymptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationSevere adverse events: Very low certainty @OOOHospitalization: No information
	Uncerta	Siltu inty in potential benefits a	uximab and harms. Further resea	urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	•		•	•	
COV-AID-2 trial, ⁵⁹² other; 2021	Patients with severe to critical COVID-19 infection. 77 assigned to siltuximab 11 mg/kg once and 72 assigned to SOC	Median age 64	Corticosteroids 59%, remdesivir 3.4%, convalescent plasma 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes: Risk of bias assessment extracted from a systematic review.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom



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	Uncertai	Sily inty in potential benefits a	/marin Ind harms. Further resea	urch is needed.	resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Severe adverse events: No information Hospitalization: No information
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Aryan et al</u> , ⁵⁹³ peer reviewed; 2022	Patients with severe COVID-19 infection. 25 assigned to silymarin 210 mg a day for 14 days and 25 assigned to SOC	Mean age 49 ± 11.1, male 48%	Corticosteroids 100%, remdesivir 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information





					Severe adverse events: Very low certainty ⊕○○○ Hospitalization: No			
information Sitagliptin Uncertainty in potential benefits and harms. Further research is needed.								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT								
Asadipooya et al; ⁵⁹⁴ preprint; 2021	Patients with moderate to severe COVID-19 infection. 66 assigned to sitagliptin 100 mg a day and 87 assigned to SOC	Mean age 57.5 ±, male 51.2%, hypertension 29%, diabetes 27.1%, COPD 8.4%, asthma %, CHD 21.2%, CKD 6.4%, cancer 5.9%, obesity 18.7%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Severe adverse events: No information Hospitalization: No information			





	and probably does not improve time to symptom resolution.							
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT								
<u>Kasgari et al</u> ; ⁴⁰⁸ peer- reviewed; 2020	moderate COVID-19	Median age 52.5 ± NR, male 37.5%, hypertension 35.4%, diabetes 37.5%, chronic lung disease 2%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded	Mortality: RR 1.14 (95%CI 0.83 to 1.56); RD 2.2% (95%CI - 2.7% to 9%); Low certainty ⊕⊕○○ Invasive mechanical			
	plus lopinavir- ritonavir			study. Concealment of allocation is probably inappropriate.	ventilation: RR 1.02 (95%CI 0.59 to 1.76); RD 0.3% (95%CI - 7.1% to 13.1.7%);			
<u>Sadeghi et al</u> ; ⁵⁹⁵ peer-reviewed; 2020	33 assigned to sofosbuvir/daclatasvir 400/60 mg once a day for 14 days and 33	Median age 58 ± 13, male 20.21%, hypertension 34.8%, diabetes 42.4%, chronic lung disease 22.7%, asthma 3%, coronary heart disease 15.1%, cancer 4.5%, obesity 25.7%	Corticosteroids 30.2%, lopinavir-ritonavir 48.4%, antibiotics 89.4%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Only outcome assessors and data analysts were blinded. Concealment of	7.1% to 13.1.7%); Low certainty ⊕⊕○○ Symptom resolution or improvement: RR 1.01 (95%CI 0.95 to 1.08); RD 0.6% (95%CI -3% to 4.8%); Moderate certainty ⊕⊕⊕○			
				allocation is probably inappropriate.	Symptomatic infection			
<u>Yakoot et al</u> , ⁵⁹⁶ preprint; 2020	Patients with mild to severe COVID-19. 44 assigned to sofosbuvir/daclatasvir 400/60 mg once a day for 10 days and 45 assigned to standard of care	Median age 49 ± 27, male 42.7%, hypertension 26%, diabetes 19%, COPD %, asthma 1%, coronary heart disease 8%	Hydroxychloroquine 100% azithromycin 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of	(prophylaxis studies): Very low certainty ⊕○○○ Adverse events: No information			
				allocation is probably inappropriate.	Hospitalization: Very low certainty ⊕○○○			



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<u>Roozbeh et al</u> ; ⁵⁹⁷ Peer reviewed; 2020	Patients with moderate COVID-19. 27 assigned to sofosbuvir/daclatasvir 400/60 mg once a day for 7 days and 28 assigned to SOC	Median age 53 ± 16, male 47%, comorbidities 38%	Azithromycin 100%, hydroxychloroquine 100%	High for symptom resolution, infection, and adverse events Notes: Blinding method possibly inappropriate which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Sali et al</u> ; ⁴⁰⁶ Peer reviewed; 2020	Patients with moderate to severe COVID-19. 22 assigned to sofosbuvir 400 mg a day and 32 assigned to lopinavir- ritonavir 400/100 mg every 12 hours	Mean age 56.5 ± 14, male 53.7%, diabetes 33%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
DISCOVER trial; ⁵⁹⁸ Mobarak et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 541 assigned to sofosbuvir/daclatasvir 400/60 mg a day for 10 days and 542 assigned to SOC	Median age 58, male 54%, hypertension 34%, diabetes 26%, COPD 2.1%, asthma 4.8%, CHD 9.1%	Steroids 69.9%, remdesivir 15.6%, hydroxychloroquine 12.8%, lopinavir- ritonavir 33.1%, azithromycin 22.1%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>Alavi-moghaddam</u> <u>et al</u> ; ⁵⁹⁹ Preprint; 2021	Patients with severe to critical COVID-19 infection. 27 assigned to sofosbuvir 400 mg a day and 30 assigned to SOC	Mean age 57.2 ±, male 49.1%, hypertension 21%, diabetes 29.8%, COPD 7%, CHD 19.3%, CKD 1.7%, obesity 1.7%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Yadollahzadeh et</u> <u>al;</u> ⁴⁰⁹ Preprint; 2021	Patients with mild to moderate COVID-19 infection. 58 assigned	Mean age 57.4 ± 15, male 44.6%, hypertension 25%,	Hydroxychloroquine 100%	High for mortality and mechanical ventilation; High for symptom	





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	to sofosbuvir/daclatasvir 400/60 mg a day for 10 days and 54 assigned to lopinavir- ritonavir 400/100 mg twice a day for 7 days	diabetes 21.4%, COPD 3.6%, CHD 15.2%, CKD 6.2%, immunosuppression 3.6%, cancer 10.7%		resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
<u>Khalili et al;</u> 600 Peer reviewed; 2020	Patients with mild to moderate COVID-19. 42 assigned to sofosbuvir/ledipasvir 400/90 mg a day for 10 days and 40 assigned to SOC	Median age 62.2 ± 23.1, hypertension 45.1%, diabetes 45.1%, COPD 4.9%, CHD 31.7%, cancer 3.6%	Corticosteroids 8.5%, hydroxychloroquine 10.9%,	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
Elgohary et al; ⁶⁰¹ preprint; 2021	Patients with moderate COVID-19 infection. 125 assigned to sofosbuvir/ledipasvir 400/90 mg once a day for 15 days and 125 assigned to SOC	Mean age 43 ±, male 0.4%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.
SOVECOD trial; ⁶⁰² Sayad et al; peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 40 assigned to sofosbuvir/velpatasvir 400/100 mg once a day for 10 days and 40 assigned to SOC	Mean age 54.1 ± 17.8, male 55%, hypertension 30%, diabetes 20%, COPD 10%, CHD 17.5%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>El-Bendari et al</u> ; ⁶⁰³ peer reviewed; 2021	Patients with moderate to severe	Mean age 53 ± 15, male 54.6%, hypertension	NR	High for mortality and mechanical ventilation;







	COVID-19 infection. 96 assigned to sofosbuvir/daclatasvir 400/60 mg a day for 14 days and 78 assigned to SOC	21.3%, diabetes 37.3%, asthma 1.7%, CHD 10.9%		high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
Abbass et al; ⁶⁰⁴ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 80 assigned to sofosbuvir/daclatasvir 400/60 a day or sofosbuvir/ravidasvir 400/200 mg a day for 10 days and 40 assigned to SOC	Mean age 44.6 ± 4.7, male 53.3%, diabetes 18.3%, asthma 1.6%, CHD 75.8%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Table 1 shows more severe patients in SOC (68% vs 59%).	
<u>Medhat et al</u> ; ⁶⁰⁵ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 70 assigned to sofosbuvir/ledipasvir 400/90 mg a day for 14 days and 73 assigned to SOC	Mean age 45, male 51%, hypertension 20.9%, diabetes 20.3%	Corticosteroids 49%, hydroxychloroquine 8.4%,	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
	Patients with severe COVID-19 infection. 50 assigned to sofosbuvir 400 mg a day for 7 days and 50 assigned to SOC	Mean age 53.8 ± , male 44%, diabetes 7%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>COVER HCW</u> trial; ⁴⁸⁷ Sokhela et al;	Patients with exposed to COVID-19	Median age 24, male 51.9%, hypertension	Vaccinated 0%	Low for mortality and mechanical ventilation;	





peer reviewed; 2022	infection. 265 assigned to sofosbuvir/daclatasvir 400/60 mg a day for 24 weeks and 283 assigned to SOC	8.2%, diabetes 1.1%, COPD 2.2%		high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Significant loss to follow-up.	
Sotrovimab	probably reduces hospi		ovimab h mild recent onset COV	ID-19 with risk factors for	severe disease.
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>COMET-ICE</u> <u>trial;</u> ⁶⁰⁷ Gupta et al; peer reviewed; 2021	Patients with mild to moderate recent onset with risk factors COVID-19 infection. 528 assigned to sotrovimab 500 mg once and 529 assigned to SOC	Median age 53, male 45.9%, hypertension %, diabetes 21.6%, COPD 5.6%, asthma 16.8%, CHD 0.7%, CKD 1.2%, obesity 63.4%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes: Stopped early for benefit	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$
<u>OPTIMISE-C19</u> <u>trial</u> ; ⁵⁵⁸ Huang et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 2454 assigned to REGN- COV2 (Regeneron) one infusion and 1104 assigned to sotrovimab one infusion	Mean age 54 ± 18, male %, hypertension 30%, diabetes 12%, CHD 16%, CKD 4.7%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: RR 0.34 (95%CI 0.16 to 0.68); RD -6.7% (95%CI -8.6% to -
<u>MANTICO trial</u> ; [%] Mazzaferri et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 107 assigned to sotrovomab 500 mg once and 106 assigned		Vaccinated 28.6%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	



	to bamlanivimab + etesevimab 700/1400 mg once and 106 assigned to REGEN- COV2 600/600 mg once	immunosuppression19. 6%, obesity 25.4%		Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	3.3%); Moderate certainty ⊕⊕⊕○ Hospitalization: RR 0.20 (95%CI 0.08 to 0.48); RD -3.8% (95%CI -4.6% to - 2.5%); Moderate certainty ⊕⊕⊕○
	Uncertai	Spiroi inty in potential benefits a	10lactone and harms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT				•	
<u>Asadipooya et al</u> , ⁵⁹⁴ preprint; 2021	Patients with moderate to severe COVID-19 infection. 50 assigned to spironolactone 100 mg a day and 87 assigned to SOC	Mean age 57.5 ±, male 51.2%, hypertension 29%, diabetes 27.1%, COPD 8.4%, asthma %, CHD 21.2%, CKD 6.4%, cancer 5.9%, obesity 18.7%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Symptom resolution or
<u>Bharti et al</u> ; ⁶⁰⁸ preprint; 2022	Patients with severe COVID-19 infection. 74 assigned to spironolactone 50 mg once followed by 25 mg a day for 21 days and 46 assigned to SOC	Mean age 48.8 ± 14.3, male 61.7%, hypertension 28.3%, diabetes 34.2%, COPD 1.7%, asthma 3.3%, CHD 5.8%, CKD 0.8%, cancer 0.8%	Corticosteroids 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Significant loss to follow up. Selective reporting: Patients with symptom progression were excluded.	 improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Severe adverse events: No information Hospitalization: No





Study; publication status Patinte inte ana RCT RESIST trial;63 Patie mod preprint; 2021 COV 221 COV 221	tients and terventions halyzed tients with oderate to severe DVID-19 infection. 1 assigned to		Additional interventions	. Further research is neede Risk of bias and study limitations	ed. Interventions effects vs standard of care and GRADE certainty of the evidence
status inte ana RCT RESIST trial; ⁶³ Patie Ghati et al; mod preprint; 2021 COV 221 atory once and	terventions halyzed tients with oderate to severe DVID-19 infection. 1 assigned to	Mean age 53.1 ± 9.2,	interventions		vs standard of care and GRADE certainty of the
RESIST trial, ⁶³ Patie Ghati et al; mod preprint; 2021 COV 221 ator once and	oderate to severe DVID-19 infection. 1 assigned to				
Ghati et al; mod preprint; 2021 COV 221 atory once and	oderate to severe DVID-19 infection. 1 assigned to				
	orvastatin 40 mg ce a day for 10 days d 219 assigned to OC	hypertension 28.6%, diabetes 27.7%, CHD 1.1%, CKD 2.4%	Corticosteroids 27.3%, remdesivir 20.6%, hydroxychloroquine 9.9%, tocilizumab 0.6%, convalescent plasma 0.2%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Blinding and concealment probably inappropriate.	Mortality: RR 0.92 (95%CI 0.73 to 1.15); RD -1.3% (95%CI - 4.3% to 2.4%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty
NSPIRATION-S trial; ⁶⁰⁹ Bikdeli et al; peer reviewed; 2022 to at day f	tical COVID-19 ection. 290 assigned	Median age 57 ± , male 56.4%, hypertension 31.5%, diabetes 16.7%, COPD 8%	Corticosteroids 93.4%, remdesivir 66.3%, hydroxychloroquine 7.5%, lopinavir- ritonavir 0.7%, tocilizumab 14.5%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	⊕○○○ Symptom resolution or improvement: Very low certainty ⊕○○○
peer reviewed; 2021 COV 76 as atory 7 to		Mean age 51.8 ± 17.4, male 50.6%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to	Symptomatic infection (prophylaxis studies): No information Adverse events: No information
				symptoms and adverse events outcomes results.	Hospitalization: No information



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
SENTAD-COVID trial; ⁶¹¹ Carmenate et al; preprint; 2021	moderate to critical COVID-19 infection. 69 assigned to stem- cell nebulization twice,	Mean age 45.1 ± 10.4, male 46.5%, hypertension 26.6%, diabetes 22.3%, COPD %, asthma 10.7%, CHD 9.3%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: Very low certainty ⊕○○○Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information





Steroids (corticosteroids) Corticosteroids reduce mortality and probably reduce invasive mechanical ventilation requirements in patients with severe COVID-19 infection with moderate certainty. Corticosteroids may not significantly increase the risk of severe adverse events. Higher doses (i.e., dexamethasone 12 mg a day) may not be more effective than standard doses (i.e., dexamethasone 6 mg a day).								
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence			
RCT				-				
<u>GLUCOCOVID</u> <u>trial</u> ; ⁶¹² Corral- Gudino et al; preprint; 2020	Patients with moderate to severe COVID-19 infection. 56 assigned to methylprednisolone 40 mg twice daily for 3 days followed by 20 mg twice daily for 3 days and 29 assigned to standard of care	Mean age 69.5 ± 11.5, male 61.9%, hypertension 47.6%, diabetes 17.5%, chronic lung disease 7.9%, cerebrovascular disease 12.7%	Hydroxychloroquine 96.8%, lopinavir- ritonavir 84.1%, azithromycin 92%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably	Mortality: RR 0.90 (95%CI 0.80 to 1.01); RD -1.6% (95%CI - 3.2% to 0.2%); Moderate certainty ⊕⊕⊕○ Invasive mechanical ventilation: RR 0.87 (95%CI 0.73 to 1.04);			
				inappropriate.	RD -2.2% (95%CI - 4.7% to 0.7%);			
Metcovid trial; ⁶¹³ Prado Jeronimo et al; peer-reviewed; 2020	Patients with severe COVID-19 infection. 194 assigned to methylprednisolone 0.5 mg/kg twice a day for 5 days and 199 assigned to standard of care	Mean age 55 ± 15 , male 64.6% , hypertension 48.9% , diabetes 29.1% , chronic lung disease 0.5% , asthma 2.5% , coronary heart disease 6.9% , alcohol use disorder 27% , liver disease 5.5%	Remdesivir 0%, tocilizumab 0%, convalescent plasma 0%	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	4.7% to 0.7%); Moderate certainty ⊕⊕⊕○ Symptom resolution or improvement: RR 1.19 (95%CI 0.95 to 1.5); RD 11.5% (95%CI -3% to 30%);			
<u>RECOVERY -</u> <u>Dexamethasone</u> <u>trial</u> ; ⁶¹⁴ Horby et al; peer-reviewed; 2020	Patients with moderate to critical COVID-19 infection. 2104 assigned to dexamethasone 6 mg once daily for 10 days and 4321 assigned to standard of care	Mean age 66.1 ± 15.7, male 64%, diabetes 24%, chronic lung disease 21%, asthma NR%, coronary heart disease 27%, chronic kidney disease 8%, liver disease 2%, any comorbidities 56%	Corticosteroids NA%, remdesivir 0.08%, hydroxychloroquine 1%, lopinavir-ritonavir 0.5%, tocilizumab 3%, azithromycin 25%	Low for mortality and invasive mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse	 (95%CI -3% to 30%); Low certainty ⊕⊕○○ Symptomatic infection (prophylaxis studies): No information Severe adverse events: RR 0.89 			



				events outcomes results.	(95%CI 0.68 to 1.17); RD -1.1% (95%CI -
DEXA-COVID19 trial; ⁶¹⁵ Villar et al; unpublished; 2020	Patients with severe to critical COVID-19. Seven assigned to dexamethasone 20 mg a day for 5 days followed by 10 mg a day for 5 days and 12 assigned to standard of care	NR	NR	Low for mortality and invasive mechanical ventilation Notes: RoB judgment from published SR.	3.3% to 1.7%); Low certainty ⊕⊕⊖⊖ Hospitalization: No information
<u>CoDEX trial</u> ; ⁶¹⁶ Tomazini et al; peer-reviewed; 2020	Patients with critical COVID-19. 151 assigned to dexamethasone 20 mg a day for 5 days followed by 10 mg a day for 5 days and 148 assigned to standard of care	male 62.5%, hypertension 66.2%, diabetes 42.1%, coronary heart disease 7.7%, chronic kidney disease 5.3%, obesity	hydroxychloroquine 21.4%, azithromycin 71.2%, ATB 87%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>REMAP-CAP</u> <u>trial;</u> ⁶¹⁷ Arabi et al; peer-reviewed; 2020	Patients with severe to critical COVID-19. 278 assigned to hydrocortisone 50 mg every 6 hours for 7 days and 99 assigned to standard of care	Mean age 59.9 ± 13, male 71%, diabetes 32%, chronic lung disease 20.3%, coronary heart disease 7.5%, chronic kidney disease 9.2%, immunosuppression 4.9%	NR	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
COVID STEROID trial; ⁶¹⁸ Munch et al; PEER- REVIEWED; 2022	Patients with severe to critical COVID-19. 16 assigned to hydrocortisone 200 mg a day for 7 days and 14 assigned to	NR	NR	Low for mortality and invasive mechanical ventilation Notes: Risk of bias judgment from	



			•		
	standard of care			published SR.	
trial; ⁶¹⁹ Dequin et al; peer-reviewed; 2020	Patients with severe to critical COVID-19. 76 assigned to hydrocortisone 200 mg a day progressively reduced to 50 mg a day for 7 to 14 days and 73 assigned to standard of care	Median age 64.7 ± 19.3, male 69.8%, hypertension %, diabetes 18.1%, chronic lung disease 7.4%, immunosuppression 6%	Remdesivir 3.4%, hydroxychloroquine 46.9%, lopinavir- ritonavir 14.1%, tocilizumab 2%, azithromycin 34.2%	Low for mortality and invasive mechanical ventilation; Low for symptom resolution, infection, and adverse events	
<u>Corticosteroids-</u> <u>SARI trial</u> ; ⁶¹⁵ Unpublished; 2020	Patients with severe to critical COVID-19. 24 assigned to methylprednisolone 40 mg twice a day for 5 days and 23 assigned to standard of care	NR	NR	Low for mortality and invasive mechanical ventilation Notes: Risk of bias judgment from published SR.	
	Patients with severe to critical COVID-19. 14 assigned to methylprednisolone 1000 mg/day for three days followed by prednisolone 1 mg/kg for 10 days, and 15 assigned to standard of care	Mean age 64 ± 13.5	Hydroxychloroquine 100%, lopinavir- ritonavir 100%, azithromycin 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Edalatifard et al</u> ; ⁶²¹ peer-reviewed; 2020	Patients with severe COVID-19. 34 assigned to methylprednisolone 250 mg/day for 3 days and 28 assigned to standard of care	Mean age 58.5 ± 16.6, male 62.9%, hypertension 32.3%, diabetes 35.5%, chronic lung disease 9.7%, coronary heart disease 17.7%, chronic kidney disease 11.3%, cancer 4.8%	Hydroxychloroquine 100%, lopinavir- ritonavir 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	





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<u>Tang et al</u> ; ⁶²² Peer reviewed; 2020	Patients with moderate to severe COVID-19. 43 assigned to methylprednisolone 1 mg/kg for 7 days and 43 assigned to SOC	Median age 56 ± 27, male 47.7%, hypertension 36%, diabetes 9.3%, COPD 3.5%, asthma 2.4%, CHD 7%, CKD 1.2%	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	
<u>Jamaati et al</u> ; ⁶²³ Peer-reviewed; 2020	Patients with moderate to severe COVID-19. 25 assigned to dexamethasone 20 mg a day for 5 days followed by 10 mg a day until day 10 and 25 assigned to SOC	Median age 62 ± 16.5, male 72%, hypertension 50%, diabetes 54%, COPD 20%, CHD 14%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>Rashad et al</u> ; ⁶²⁴ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 75 assigned to dexamethasone 4 mg/kg a day for 3 days followed by 8 mg a day for 10 days and 74 assigned to TCZ	Mean age 62, male 56.9%, hypertension 47.7%, diabetes 28.4%, COPD 1.8%, asthma 2.7%, CHD 12.8%, CKD 8.2%, cancer 0.9%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate. Significant loss to follow-up as patients who died in the first 3 days after randomization were excluded.	
<u>Ghanei et al</u> ; ⁸³ peer reviewed; 2021	Patients with severe COVID-19 infection. 116 assigned to predninoslone 25 mg a day for 5 days and 110 assigned to SOC	Mean age 58.1 ± 16.3, male 51.5%, hypertension 24.7%, diabetes 12.2%, asthma 4.5%, CHD 8.9%, CKD 1.2%	Convalescent plasma 1.8%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	



CORTIVID trial; ⁶²⁵ Les et al; peer reviewed; 2021	Patients with moderate COVID-19 infection. 34 assigned to methylprednisolone and 37 assigned to SOC	Mean age 58.4, male 69%, hypertension 32.4%, diabetes 18.3%, COPD 1.4%, asthma 2.8%, CKD 7%	Remdesivir 8.5%, tocilizumab 28.2%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	
<u>Ranjbar et al</u> ; ⁶²⁶ Preprint; 2020	Patients with severe to critical COVID-19 infection. 44 assigned to methylprednisolone 2 mg/kg daily for 5 days followed by tapering using same scheme at half dose every 5 days, 42 assigned to dexamethasone 6 mg a day for 10 days	Mean age 58.7 ± 17.4, male 56.9%, hypertension 45.3%, diabetes 32.5%, CHD 30.2%, CKD 2.3%	NR	Some concerns for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Unbalanced prognostic factors (age and gender).	Mortality: RR 0.97 (95%CI 0.78 to 1.21); RD -0.5% (95%CI - 3.5% to $3.4%$); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
COVID STEROID 2 trial; ⁶²⁷ Munch et al; preprint; 2021	Patients with severe to critical COVID-19 infection. 497 assigned to dexamethasone 12 mg a day for 10 days and 485 assigned to dexamethasone 6 mg a day for 10 days	Median age 64.5 ± 18, male 69%, diabetes 30.3%, COPD 12%, CHD 14%	Remdesivir 62.8%, tocilizumab 10.1%, convalescent plasma 2.8%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Symptom resolution or improvement: RR 0.99 (95%CI 0.9 to 1.08); RD -0.6% (95%CI -5.5% to 4.8%); Low certainty ⊕⊕⊖⊖
Maskin et al; ⁶²⁸ preprint; 2021	Patients with critical COVID-19 infection. 49 assigned to dexamethasone 16 mg a day for 5 days followed by 8 mg a day for 5 days and 49 assigned to dexamethasone 6 mg a day for 10 days	Mean age 61.8 ± 13.4, male 70%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Symptomatic infection (prophylaxis studies): No information Adverse events: RR 0.82 (95%CI 0.6 to 1.11); RD -1.8% (95%CI -4.1% to 1.1%); Low certainty
<u>Toroghi et al;</u> ⁶²⁹ peer reviewed; 2021	Patients with severe COVID-19 infection. 86 assigned to dexamethasone 16 to 24 mg a day and 47	Mean age 58, male 60.2%, hypertension 36%, diabetes 22.5%, COPD 6%, CHD 17.3%, CKD 1.5%,	Remdesivir 75.2%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	 1.1%); Low certainty ⊕⊕○○ Hospitalization: No information





	assigned to dexamethasone 8 mg a day for up to 10 days	cerebrovascular disease 6%, cancer 2.3%		Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
HIGHLOWDEXA trial; ⁶³⁰ Taboada et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 98 assigned to dexamethasone 20 mg once a day for 5 days dexamethasone and 102 assigned to dexamethasone 6 mg once a day for 10 days	Mean age 64.3 ± 14.3, male 61.8%, hypertension 48%, diabetes 19%, COPD 7%, asthma 5%, CHD 13.5%, CKD 3.5%, obesity 53%	Remdesivir 10%, tocilizumab 12%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>Naik et al</u> ; ⁶³¹ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 21 assigned to dexamethasone 20 mg a day for 3 days and 21 assigned to TCZ 6 mg/kg once	Median age 50.5, male 57.1%, hypertension 57.1%, diabetes 35.7%, COPD 4.8%, asthma 2.4%, CHD %, CKD 0%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
	Patients with severe COVID-19 infection. 151 assigned to three boluses of 1 g of methylprednisolone intravenously and 150 assigned to SOC	Median age 64 , male 72.1%, hypertension 52.2%, diabetes 14.9%, COPD 4.4%, obesity 22.9%	Corticosteroids 88.4%, remdesivir 15.3%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
COVIDICUS trial; ⁶³³ Bouadma et al; peer reviewed; 2022	Patients with severe to critical COVID-19 infection. 270 assigned to dexamethasone 14 mg a day for 5 days followed by dexamethasone 4 mg a day for 5 days and 276	Median age 67, male 75.8%, hypertension 55.4%, diabetes 37%, cancer 11.2%,	Corticosteroids %, remdesivir 17%, hydroxychloroquine 1.1%, lopinavir- ritonavir 2.2%, tocilizumab 1%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	



Dastenae et al; ⁶³⁴ peer reviewed; 2022	critical COVID-19 infection. 73 assigned to methylprednisolone 60 mg a day for 10 days and 71 assigned to dexamethasone 8 mg a	%, CHD 11.9%, CKD	Remdesivir 88.1%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded	
	day for 10 days		ed corticosteroio		
Inhaled corticostero		symptom resolution but important outcomes are		important effect on hospit rch is needed.	alizations. Their effects
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
	Patients with mild to moderate COVID-19. 71 assigned to inhlaed budesonide 800 µg twice a day and 69 assigned to SOC	Mean age 45 ± 56, male 42.4%	NR	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Symptom resolution or improvement: RR 1.09 (95%CI 0.99 to 1.2); RD 5.5% (95%CI -0.6% to 12.1%); Low certainty
<u>PRINCIPLE</u> <u>trial</u> ; ⁶³⁶ Yu et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 787 assigned to inhaled budesonide	Mean age 64.2 ± 7.6, male 48%, hypertension 44.3%, diabetes 21.4%, COPD 12.6%, CHD	NR	Some concerns for mortality and mechanical ventilation; Some concerns for	⊕⊕○○ Symptomatic infection





	800µg twice daily for 14 days and 1069 assigned to SOC	15.8%, cerebrovascular disease 5.6%		symptom resolution, infection, and adverse events Notes: Non-blinded study. Significant loss to follow-up.	(prophylaxis studies): No information Hospitalization: RR 0.9 (95%CI 0.7 to 1.15); RD -0.5%
Song et al; ⁶³⁷ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 35 assigned to inhaled ciclesonide 320 µg twice per day for 14 days and 26 assigned to SOC	Median age 53 ± 26, male 47%, hypertension 27.8%, diabetes 14.7%, cerebrovascular disease 3.3%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	(95%CI -1.4% to 0.7%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ Adverse events: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
ALV-020-001 trial; ⁶³⁸ Clemency et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 197 assigned to inhaled ciclesonide 640 μg a day for 30 days and 203 assigned to SOC	Mean age 43.3 ± 16.9, male 44.8%, hypertension 22.3%, diabetes 7.5%, asthma 6.5%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
CONTAIN trial; ⁶³⁹ Ezer et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 105 assigned to inhaled ciclesonide 1200 μg + 200 μg intranasal a day and 98 assigned to SOC	Median age 35 ± 19, male 46.3%, hypertension 5.9%, diabetes 2.5%, asthma 5%, CHD 0.5%, cancer 1%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	



<u>Alsultan et al</u> ; ¹⁴² peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 14 assigned to inhaled steroids budesonide 200 mcg twice a day for 5 days and 21 assigned to SOC	age 60 to 80 65.3, male 38.8%, diabetes 53.1%, CKD 8.2%, cerebrovascular disease 4.1%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>COVERAGE</u> <u>trial</u> ; ⁶⁴⁰ Duvignaud et al; peer reviewed; 2021	Patients with mild COVID-19 infection. 110 assigned to inhaled ciclesonide 640 µg of ciclesonide per day for 10 days and 107 assigned to SOC	Median age 63, male 48.9%, hypertension 41%, diabetes 15.2%, COPD 3.2%, CHD 5%, cerebrovascular disease 8.7%, cancer 5.9%, obesity 29.4%	Vaccinated13.8%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
TACTIC-COVID <u>trial</u> ; ⁶⁴¹ Agusti et al; other; 2021	Patients with moderate to severe COVID-19 infection. 58 assigned to budesonide (inh) 400 μg/12 h and 62 assigned to SOC	Mean age 51.1 ± 13.7, male 47.1%,	Corticosteroids 17.8%, remdesivir 8.5%, hydroxychloroquine 8.5%, lopinavir- ritonavir 5.9%, tocilizumab 0.8%, azithromycin 9.3%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
<u>Terada et al</u> ; ¹²⁰ peer reviewed; 2022	Patients with mild to severe COVID-19 infection. 56 assigned to camostat 600 mg + ciclesonide (inhaled) 1200 µg a day and 61 assigned to SOC	Mean age 58.3, male 64.9%, diabetes 24.8%, COPD 9.4%, CHD 2.6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.



ACTIV-6 - Fluticazone trial, ⁶⁴² Naggie et al; preprint; 2022	Patients with mild to moderate COVID-19 infection. 656 assigned to fluticazone 200 µg once a day for 14 days and 621 assigned to SOC	Median age 45, male 36.8%, hypertension 26.1%, diabetes 9.7%, COPD 1.4%, asthma 13%, CHD 4.7%, CKD 0.8%, cancer 3.4%,	Corticosteroids %, remdesivir 0.1%, monoclonar antibodies 2.7%, paxlovid 0.1%; Vaccinated 65.2%,	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
	Uncertai	Steroids (nasa inty in potential benefits a	l corticosteroids		
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Yildiz et al; ⁴⁷¹ peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 50 assigned to nasal steroids and 50 assigned to SOC	asthma %, CHD 14%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information
		Sulo	odexide		



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
ERSul trial; ⁶⁴³ Gonzalez Ochoa et al; preprint; 2020	Patients with mild (early within 3 days of onset) COVID-19. 124 assigned to sulodexide 500 RLU twice a day for 3 weeks and 119 assigned to standard of care	Median age 52 ± 10.6, male 47.4%, hypertension 34.2%, diabetes 22.2%, COPD 23%, coronary heart disease 21%	Corticosteroids 62.5%, hydroxychloroquine 33.7%, ivermectin 43%	Some concerns for mortality and mechanical ventilation; some concerns for symptom resolution, infection, and adverse events Notes: Significant loss to follow-up.	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: Very low certainty \oplus \bigcirc \bigcirc
	Uncerta	Tafe inty in potential benefits a	noquine and harms. Further resea	urch is needed.	
Study; publication tatus	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence





Dow et al. ⁶⁴⁴ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 45 assigned to tafenoquine 200 mg a day for 3 days followed by 200 mg once next week and 41 assigned to SOC	Mean age 43 ± 15, male 47.7%	Vaccinated 32.6%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: No informationInvasive mechanical ventilation: No informationSymptom resolution or improvement: : Very low certainty $\oplus \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty $\oplus \bigcirc \bigcirc$ Hospitalization: : Very low certainty $\oplus \bigcirc \bigcirc$
	Uncertai	TD-0903 (inhal inty in potential benefits a	ed JAK-inhibit		
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Singh et al</u> ; ⁶⁴⁵ Preprint; 2021	Patients with severe to critical COVID-19 infection. 19 assigned to TD-0903 1-10 mg once a day for 7 days and 6 assigned to SOC	Mean age 57.1 ± 12.3, male 68%, hypertension 68%, diabetes 40%	Corticosteroids 92%, remdesivir 12%,	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom



				allocation is probably inappropriate.	resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
Tenofovir + emtricit	abine may not reduce m	ortality but may reduce m	- emtricitabine hechanical ventilation. H h is needed.	owever, certainty of the ev	idence was low. Further
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
AR0-CORONA trial; ⁶⁴⁶ Parientti et al; peer reviewed; 2021	Patients with mild to moderate COVID-19 infection. 30 assigned to tenofovir + emtricitabine 245/200 mg twice a day on day one followed by 245/200 mg a day for 7 days and 30 assigned to SOC	Mean age 42 ± 15, male 43%, hypertension 5%, diabetes 3.3%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: RR 0.97 (95%CI 0.49 to 1.92); RD -0.5% (95%CI - 8.2% to 14.7%); Low certainty ⊕⊕○○ Invasive mechanical ventilation: RR 0.76 (95%CI 0.49 to 1.18); RD -4.2% (95%CI - 8.8% to 3.1%); Low
<u>ARTAN-C19</u> <u>trial</u> , ⁶⁴⁷ Lima et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 81 assigned to tenofovir +/- emtricitabine 300/200 mg once a	Mean age 38 ± 14.9, male 35%, hypertension 17%, diabetes 10%, asthma 6%, CHD 3%, cancer 1%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events	8.8% to 3.1%); Low certainty ⊕⊕⊖⊖ Symptom resolution or improvement: Very low certainty



EPICOS trial; ³⁰⁸ Polo et al; preprint; 2021	day and 41 assigned to SOC Individuals exposed to SARS-CoV-2 infection. 233 assigned to tenofovir +/- emtricitabine 245/200 mg a day and 223 assigned to SOC	Mean age 38.5, male 38%, hypertension 7.4%, diabetes 1.3%, COPD 0%, asthma 3.7%, CHD 0.4%, cancer 1.1%,	NR	Notes: Concealment of allocation probably inappropriate. Significant loss to follow-up. High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	$ \begin{array}{c} \bigoplus \bigcirc \bigcirc \bigcirc \\ \hline \\ Symptomatic \\ infection \\ (prophylaxis \\ studies): Very low \\ certainty \bigoplus \bigcirc \bigcirc \bigcirc \\ \hline \\ Adverse events: \\ Very low certainty \\ \bigoplus \bigcirc \bigcirc \bigcirc \\ \hline \\ Hospitalization: \\ Very low certainty \\ \bigoplus \bigcirc \bigcirc \bigcirc \\ \hline \end{array} $
<u>Gaitan-Duarte et</u> <u>al</u> ; ¹⁴⁸ preprint; 2021	Patients with moderate to severe COVID-19 infection. 160 assigned to emtricitabine/ tenofovir 200/300 mg once a day for 10 days and 161 assigned to SOC	Mean age 55.4 ± 12.8, male 68%, hypertension 28%, diabetes 12%, COPD 4%	Corticosteroids 98%,	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
PanCOVID19 trial; ¹⁰⁴ Montejano et al; peer reviewed; 2022	Patients with moderate COVID-19 infection. 177 assigned to tenofovir +/- emtricitabine 400/490 mg once followed by 200/245 mg once a day for 14 days and 178 assigned to SOC	Median age 67, male 64.5%, hypertension 61.1%, diabetes 27.3%, obesity 16.1%	Corticosteroids 100%, remdesivir 12.7%, baricitinib 50.5%; Vaccinated 91%	Low for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
	Uncerta	Thal inty in potential benefits a	idomide and harms. Further rese	arch is needed	
Study; publication status	Patients and interventions	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care





	analyzed				and GRADE certainty of the evidence
RCT					
<u>Amra et al</u> ; ⁶⁴⁸ preprint; 2021	Patients with severe COVID-19 infection. 28 assigned to thalidomide 100 mg a day for 14 days and 23 assigned to SOC	Mean age 62 ± 10, male 54.9%, hypertension 33.3%, diabetes 37.2%, COPD 5.9%, CHD 9.8%	Corticosteroids 100%, hydroxychloroquine 100%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty \oplus \bigcirc \bigcirc Invasive mechanical ventilation: Very low certainty \oplus \bigcirc \bigcirc Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty \oplus \bigcirc \bigcirc Hospitalization: No
<u>Haghighi et al</u> ; ⁶⁴⁹ preprint; 2021	Patients with moderate to severe COVID-19 infection. 25 assigned to thalidomide 100 mg a day for 14 days and 25 assigned to SOC	Median age 51 ± 18, male 68%, hypertension 24%, diabetes 16%, CHD 8%, cancer 14%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
	Uncerta	Thym inty in potential benefits a	oquinone and harms. Further rese	arch is needed	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Bencheqroun et al; ⁶⁵⁰ peer reviewed; 2022	Patients with mild to moderate COVID-19 infection. 23 assigned to thymoquinone	Age >55 29.1%, male 43.6%, hypertension 40%, diabetes 18.2%, obesity 38.2%	Vaccinated 16.4%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and	Mortality: No information Invasive mechanical



				1	i
	3000 mg a day and 19 assigned to SOC			adverse events Notes:	ventilation: No information
					Symptom resolution or improvement: No information
					Symptomatic infection (prophylaxis studies): No information
					Adverse events: Very low certainty ⊕○○○
					Hospitalization: No information
	Uncerta	Tissue plasmino			
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	•		•		
<u>STARS trial;</u> ⁶⁵¹ Barret et al; peer reviewed; 2021	Patients with critical COVID-19 infection. 25 assigned to tPa	Mean age 61, male 74%, hypertension 36%, diabetes 34%, COPD	Corticosteroids 52%, remdesivir 40%,	High for mortality and mechanical ventilation; high for symptom	Mortality: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
	50 mg bolus with or without drip and heparin and 25 assigned to SOC	62%, asthma %, CHD 66%, immunosuppressive therapy 66%		resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Invasive mechanical ventilation: No information Symptom resolution or improvement: No information
TACOVID trial; ⁵² Rashidi et al; peer reviewed; 2022	Patients with severe to critical COVID-19 infection. 5 assigned to	Mean age 56.5, male 80%, hypertension 40%, diabetes 10%, CHD	NR	High for mortality and mechanical ventilation; high for symptom	Symptomatic infection (prophylaxis



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	tPa 50 mg in 24 hs and 5 assigned to UFH 15000 IU a day	20%, CKD 0%, cancer 0%, obesity 20%		resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	<pre>studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information</pre>
Tixagevimab-cilga	vimab probably reduc	es mortality, hospitaliza	ab–cilgavimab ations, and SARS-CO vere adverse events.	V-2 infections in exposed	individuals, and may
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
PROVENT trial; ⁶⁵² Levin et al; peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 3441 assigned to tixagevimab- cilgavimab 300 mg once and 1731 assigned to SOC	Mean age 53.5 ± 15, male 53.9%, hypertension 35.9%, diabetes 14.1%, COPD 5.3%, asthma 11.1%, CHD 8.1%, CKD 5.2%, immunosuppresive therapy 3.3%, cancer 7.4%, obesity 41.7%	Vaccinated 0%	Low for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Most patients were not blinded which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: RR 0.72 (95%CI 0.54 to 0.96); RD -4.5% (95%CI - 7.4% to -0.6%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ Invasive mechanical ventilation No information
<u>TACKLE trial</u> ; ⁶⁵³ Montgomery et al; peer reviewed; 2022	moderate COVID-19 infection. 452 assigned to tixagevimab- cilgavimab 600 mg	Mean age 46.1 ± 15.2, male 50%, hypertension 28%, diabetes 12%, immunosuppression therapy 5%, cancer 4%, obesity 43%	Corticosteroids 2.8%; vaccinated 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Symptom resolution or improvement: RR 1.03 (95%CI 0.99 to 1.08); RD 2% (95%CI -0.6% to 4.7%); Moderate certainty ⊕⊕⊕○
<u>TICO trial</u> ; ⁶⁵⁴ Lane et al; peer reviewed; 2022	Patients with moderate COVID-19 infection. 710 assigned to tixagevimab-	Mean age 46.1 ± 15.2, male 50%, hypertension 28%, diabetes 12%, CHD 9%, CKD 2%,	Corticosteroids 73%, remdesivir 63.3%; vaccinated 26.5%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and	Symptomatic infection (prophylaxis studies): RR 0.18 (95%CI 0.09 to 0.35);

	cilgavimab 600 mg once and 707 assigned to SOC	immunosuppression 5%, cancer 4%, obesity 43%	izumah	adverse events	RD -14.2% (95%CI - 15.8% to -11.2%); Moderate certainty $\oplus \oplus \oplus \bigcirc$ Adverse events: RR 0.95 (95%CI 0.69 to 1.31); RD -0.5% (95%CI -3.2% to 3.2%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Hospitalization: RR 0.42 (95%CI 0.24 to 0.74); RD -2.8% (95%CI -3.6% to 1.3%); Moderate certainty $\oplus \oplus \oplus \bigcirc$
Tocil	lizumab reduces mortali		lizumab tion requirements withou	it increasing severe advers	se events.
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	· · · · · · · · · · · · · · · · · · ·		· · · · · · · · · · · · · · · · · · ·		·
<u>COVACTA trial;</u> Rosas et al; ⁶⁵⁵ peer- reviewed; 2020	Patients with severe COVID-19. 294 assigned to tocilizumab 8 mg/kg once and 144 assigned to standard of care	Mean age 60.8 ± 14, male 70%, hypertension 62.1%, diabetes 38.1%, chronic lung disease 16.2%, coronary heart disease 28%, obesity 20.5%	Corticosteroids 42.2%, convalescent plasma 3.6%, Antivirals 31.5%	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: RR 0.86 (95%CI 0.79 to 93); RD -2.2% (95%CI - 3.4% to -1.1%); High certainty ⊕⊕⊕⊕ Invasive mechanical
<u>Wang et al</u> ; ⁶⁵⁶ preprint; 2020	Patients with moderate to severe COVID-19. 34 assigned to tocilizumab 400 mg once or twice and 31 assigned to standard of care	Median age 63 ± 16, male 50.8%, hypertension 30.8%, diabetes 15.4%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded	ventilation: RR 0.84 (95%CI 0.79 to 0.91); RD -2.8% (95%CI - 3.6% to -1.6%); High certainty ⊕⊕⊕ Symptom resolution or



<u>Zhao et al</u> ; ²³¹ peer- reviewed; 2020	Patients with moderate to critical COVID-19 infection. 13 assigned to favipiravir 3200 mg once followed by 600 mg twice a day for 7 days, 7 assigned to tocilizumab 400 mg once or twice and 5 assigned to favipiravir plus tocilizumab	Mean age 72 ± 40, male 54%, hypertension 42.3%, diabetes 11.5%, coronary heart disease 23.1%	NR	study. Concealment of allocation is probably inappropriate. High for mortality and invasive mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	improvement: RR 1.08 (95%CI 1.02 to 1.14); RD 4.8% (95%CI 1.2% to 8.5%); Low certainty ⊕⊕○○ Symptomatic infection (prophylaxis studies): No information Adverse events: RR 0.95 (95%CI 0.87 to 1.04); RD -0.5%
<u>RCT-TCZ-</u> <u>COVID-19 trial</u> ; ⁶⁵⁷ Salvarani et al; peer- reviewed; 2020	Patients with severe COVID-19. 60 assigned to tocilizumab 8 mg/kg twice on day 1 and 66 assigned to standard of care	Median age 60 ± 19, male 61.1%, hypertension 44.4%, diabetes 15.1%, COPD 3.2%, obesity 32.2%	Hydroxychloroquine 91.3%, azithromycin 20.6%, antivirals 41.3%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	1.04); RD -0.5% (95%CI -1.3% to 0.4%); Moderate certainty ⊕⊕⊕⊖ Hospitalization: No information
BACC Bay Tocilizumab Trial trial; ⁶⁵⁸ Stone et al; peer-reviewed; 2020	Patients with severe COVID-19. 161 assigned to tocilizumab 8 mg/kg once and 81 assigned to standard of care	Median age 59.8 ± 15.1, male 58%, hypertension 49%, diabetes 31%, COPD 9%, asthma 9%, coronary heart disease 10%, chronic kidney disease 17%, cancer 12%	Corticosteroids 9.5%, remdesivir 33.9%, hydroxychloroquine 3.7%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
CORIMUNO- <u>TOCI 1 trial</u> ; ⁶⁵⁹ Hermine et al; peer- reviewed; 2020	Patients with moderate to severe COVID-19. 63 assigned to tocilizumab 8 mg/kg once followed by an optional 400 mg dose on day 3 and 67	Median age 63.6 ± 16.2, male 67.7%, diabetes 33.6%, COPD 4.7%, asthma 6.3%, coronary heart disease 31.2%, chronic kidney disease 14%, cancer 7%	Corticosteroids 43%, remdesivir 0.7%, hydroxychloroquine 6.2%, lopinavir- ritonavir 3%, azithromycin 15.4%	Low for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have	





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	assigned to standard of care			introduced bias to symptoms and adverse events outcomes results.	
<u>EMPACTA trial</u> ; ⁶⁶⁰ Salama et al; preprint; 2020	Patients with moderate to severe COVID-19. 249 assigned to tocilizumab 8 mg/kg once and 128 assigned to standard of care	Mean age 55.9 ± 14.4, male 59.2%, hypertension 48.3%, diabetes 40.6%, COPD 4.5%, asthma 11.4%, coronary heart disease 1.9%, cerebrovascular disease 3.4%, obesity 24.4%	Corticosteroids 59.4%, remdesivir 54.6%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
<u>REMAP-CAP -</u> <u>tocilizumab trial;</u> ⁵⁷⁸ Gordon et al; peer- reviewed; 2020		CHD 10.2%,	Corticosteroids 75.6%, remdesivir 32.8%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Veiga et al</u> ; ⁶⁶¹ peer reviewed; 2020		Mean age 57.4 ± 14.6, male 68%, hypertension 49.6%, diabetes 32.6%, COPD 3%, CHD 5.5%, cancer 7%,	Corticosteroids 71.3%	Low for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
RECOVERY-TCZ <u>trial</u> ; ⁶⁶² Horby et al; peer reviewed; 2020	critical COVID-19.	Mean age 63.6 ± 13.6, male 67.3%, diabetes 28.5%, COPD 23%, asthma %, CHD 23%, CKD 5.5%	Corticosteroids 82%, hydroxychloroquine 2%, lopinavir-ritonavir 3%, azithromycin 9%	Low for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events	



				Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>PreToVid trial;</u> 663 Rutgers et al; preprint; 2021	Patients with severe COVID-19 infection. 174 assigned to TCZ 8 mg/kg once or twice and 180 assigned to SOC	Median age 66.5 ± 16.5, male 67%, comorbidities 74.3%		Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events	
				Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Talaschian et al</u> ; ⁶⁶⁴ preprint; 2021	Patients with severe COVID-19 infection. 17 assigned to TCZ 8 mg/kg once or twice and 19 assigned to SOC	Mean age 61.7 ± 14.2, male 52.7%, hypertension 50%, diabetes 36.1%, COPD 8.3%, asthma %, CHD 44.4%, CKD 2.8%, cancer 0%	Corticosteroids 33.3%, hydroxychloroquine 63.9%, lopinavir- ritonavir 8.3%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Concealment of allocation and blinding	
				probably inappropriate.	
<u>Hamed et al</u> ; ⁶⁶⁵ peer reviewed; 2021	Patients with severe COVID-19 infection. 23 assigned to TCZ 400 mg once and 26 assigned to SOC	Mean age 48 ±, male 85.5%, hypertension 36.8%	Corticosteroids 100%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events	
				Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>ARCHITECTS</u> <u>trial</u> ; ⁵⁹² other; 2021	Patients with severe to critical COVID-19 infection. 10 assigned to TCZ 8 mg/kg once or twice and 11	Median age 61 ±	Corticosteroids 95.2%, remdesivir 90.4%, convalescent plasma 100%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	



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	assigned to SOC			Notes: Risk of bias assessment extracted from a systematic review.
<u>CORIMUNO-</u> <u>TOCI ICU trial</u> ; ⁵⁸² Hermine et al; Peer reviewed; 2021	Patients with critcal COVID-19 infection. 49 assigned to TCZ 8 mg/kg once or twice and 43 assigned to SOC	Mean age 64.2 ± , male 71.7%, diabetes 35.5%, COPD 7.8%, asthma 5.5%, CHD %, CKD 6.6%, cancer 2.2%,	Steroids 33.6%, remdesivir 0%, hydroxychloroquine 0%, lopinavir-ritonavir 4.3%, azithromycin 4.3%, convalescent plasma 0%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.
<u>COV-AID trial; et</u> <u>al</u> ; ⁵⁹² other; 2021	Patients with severe to critical COVID-19 infection. 81 assigned to TCZ 8 mg/kg once and 72 assigned to SOC	Median age 63	Corticosteroids 52.6%, remdesivir 5.8%, convalescent plasma 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes: Risk of bias assessment extracted from a systematic review.
<u>COVIDOSE-2 trial;</u> <u>et al</u> ; ⁵⁹² other; 2021	Patients with moderate to severe COVID-19 infection. 20 assigned to TCZ 40-120 mg once and 8 assigned to SOC	Median age 65	Corticosteroids 30%, remdesivir 75%, convalescent plasma 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes: Risk of bias assessment extracted from a systematic review.
COVIDSTORM trial; ⁶⁶⁶ Broman et al; peer reviewed; 2021	critical COVID-19 infection. 57 assigned to TCZ 400 to 800 mg	Median age 58.5 ± 13.9, male 55.8%, hypertension 37.2%, diabetes 24.4%, COPD 3.5%, asthma 14%, CHD 5.81%, cancer 11.6%, obesity 63.5%	Steroids 77%, remdesivir 0%, convalescent plasma 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of





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				allocation probably inappropriate.	
<u>COVITOZ-01 trial;</u> <u>et al</u> ; ⁵⁹² other; 2021	Patients with moderate to severe COVID-19 infection. 17 assigned to TCZ 8 mg/kg once or twice and 9 assigned to SOC	Median age 57	Corticosteroids 100%, remdesivir 52.9%, convalescent plasma 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes: Risk of bias assessment extracted from a systematic review.	
HMO-0224-20 trial; ⁵⁹² other; 2021	Patients with severe to critical COVID-19 infection. 37 assigned to TCZ 8 mg/kg once and 17 assigned to SOC	Median age 63	Corticosteroids 85.2%, remdesivir 22.2%, convalescent plasma 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Concealment of allocation probably inappropriate.	
<u>et al;</u> ⁶⁶⁷ Rosas et al;	Patients with severe to critical COVID-19 infection. 430 assigned to TCZ 8 mg/kg once or twice and 210 assigned to SOC	Median age 6, male 63.2%, hypertension 61.7%, diabetes 39.5%, CHD 23.4%	Corticosteroids 88.1%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
ImmCoVA trial; ⁵⁹² other; 2021	Patients with severe to critical COVID-19 infection. 22 assigned to TCZ 8 mg/kg once and 27 assigned to SOC	Median age 24	Corticosteroids 96%, remdesivir 14.5%, convalescent plasma 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes: Risk of bias assessment extracted from a systematic review.	





TOCOVID trial; ⁵⁹² other; 2021	Patients with moderate to severe COVID-19 infection. 136 assigned to TCZ 400 to 600 mg once and 134 assigned to SOC	Median age 53	Corticosteroids 35%, remdesivir 0.5%, convalescent plasma 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events Notes: Risk of bias	
				assessment extracted from a systematic review.	
<u>COVINTOC trial;</u> <u>et al;</u> 668 Soin et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 91 assigned to TCZ 6 mg/kg once or twice and 88 assigned to SOC	Median age 55 , male 85.5%, hypertension 39.4%, diabetes 41.1%, COPD 2.2%, CHD 15%, CKD 4.4%	Corticosteroids 91%, remdesivir 41.6%, convalescent plasma 0%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to	
				symptoms and adverse events outcomes results.	
TOCIDEX trial; ⁶⁶⁹ Hermine et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 224 assigned to TCZ 400 mg once and 226 assigned to SOC	Median age 63 ± 21, male 68%, hypertension 37.1%, diabetes 23.8%, COPD %, asthma 8.4%, CHD 13.5%, CKD 7.2%	Corticosteroids 100%, convalescent plasma 1.3%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events	
				Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Karampitsakos et</u> <u>al;</u> ⁶⁷⁰ preprint; 2022	Patients with severe COVID-19 infection. 125 assigned to baricitinib 4 mg a day for 14 days and 126 assigned to TCZ 8	Mean age 72.5, male 59.4%, hypertension 53.8%, cancer 9.2%, obesity 8%	Corticosteroids 100%, remdesivir 100%; vaccinated 20.3%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events	
	mg/kg once			Notes: Non-blinded study. Concealment of allocation probably inappropriate.	





MARIPOSA trial; ⁶⁷¹ Kumar et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 49 assigned to TCZ 4 mg/kg and 48 assigned to TCZ 8 mg/kg	Mean age 56.8 ± 14.3, male 58.7%	Corticosteroids 22.7%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Mortality: Very low certainty \oplus ()Invasive mechanical ventilation: Very low certainty (\oplus ()Symptom resolution or improvement: Very low certainty (\oplus ()Symptom resolution or improvement: Very low certainty (\oplus ()Symptom resolution or improvement: Very low certainty (\oplus ()Symptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty (\oplus ()Hospitalization: No information
	I	Tofa	acitinib	1	
	Tofacitinib may increas	se symptom resolution or i	improvement and may ir	ncrease severe adverse ever	ıts.
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
STOP-COVID trial; ⁶⁷² Guimaraes et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 144 assigned to tofacitinib 10 mg twice a day for 14 days and 145 assigned to	Mean age 56 ± 14, male 65.1%, hypertension 50.2%, diabetes 23.5%	Corticosteroids 78.5%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty $\bigoplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: No information



Murugesan et al; ⁶⁷³ peer reviewed; 2021	SOC Patients with moderate to severe COVID-19 infection. 50 assigned to tofacitinib 20 mg a day for 14 days and 50 assigned to SOC	Mean age 46.5, male 74%, diabetes 36%, COPD 1%, CHD 5%	Corticosteroids 100%, remdesivir 98%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Symptom resolution or improvement: RR 1.1 (95%CI 0.98 to 1.23); RD 6.1% (95%CI 1.2% to 13.9%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Symptomatic infection (prophylaxis studies): No information Adverse events: RR 3.22 (95%CI 1.12 to 8.56); RD 22.6% (95%CI 1.2% to 77.1%); Low certainty $\oplus \oplus \bigcirc \bigcirc$ Hospitalization: No
	Unconto		anilast	wah is woodod	information
	Uncerta	inty in potential benefits a	and narms. Further resea	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Saeedi-Boroujeni et <u>al</u> , ⁶⁷⁴ peer reviewed; 2021	Patients with severe COVID-19 infection. 30 assigned to tranilast 300 mg a day for 7 days and 30 assigned to SOC	Mean age 59.5, male 63.3%, hypertension 36.7%, diabetes 26.7%, COPD 16.6%, CKD 6.6%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or





	Uncerta	Tria	Zavirin nd harms. Further resea	inappropriate.	<pre>improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information Adverse events: No information Hospitalization: No information</pre>
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Wu et al, ⁶⁷⁵ peer- reviewed; 2020	Patients with mild to critical COVID-19. 26 assigned to triazavirin 250 mg orally three or four times a day for 7 days and 26 assigned to standard of care		Corticosteroids 44.2%, hydroxychloroquine 26.9%, lopinavir- ritonavir 9.6%, antibiotics 69.2%, interferon 48.1%, umifenovir 61.5%, ribavirin 28.9%	Low for mortality and invasive mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): No information



					Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncerta	TX inty in potential benefits a	A-127 and harms. Further res	search is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	1			- !	
AAAT0535 trial; ⁶⁷⁶ Wagener et al; peer reviewed; 2022	Patients with severe COVID-19 infection. 11 assigned to TXA- 127 0.5 mg/kg a day for 10 days and 9 assigned to SOC	Mean age 56, male 65%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: No information



Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
Lau et al; ⁶⁷⁷ peer reviewed; 2022	Patients with severe COVID-19 infection. 15 assigned to UVB escalating protocol for 8 days and 15 assigned to SOC	Mean age 66.9, male 60%, hypertension 50%, diabetes 16.7%	Corticosteroids 93.3%, remdesivir 76.7%, tocilizumab 30%, vaccinated 33.3%, Regeneron 3.3%	Low for mortality and mechanical ventilation; Low for symptom resolution, infection and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
1		Umi	fenovir	l	
	Uncerta	inty in potential benefits a	nd harms. Further resea	irch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Chen et al</u> ; ²²¹ preprint; 2020	Patients with moderate to critical COVID-19 infection.	Mean age NR ± NR, male 46.6%, hypertension 27.9%,	NR	High for mortality and invasive mechanical ventilation; high for	Mortality: Very low certainty ⊕○○○



	116 assigned to favipiravir 1600 mg twice the first day followed by 600 mg twice daily for 7 days and 120 assigned to umifenovir 200 mg three times daily for 7 days	diabetes 11.4%		symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: No information
ELACOI trial; ⁴⁰² Li et al; peer-reviewed; 2020		Mean age 49.4 ± 14.7, male 41.7%	Corticosteroids 12.5%, IVIG 6.3%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
<u>Nojomi et al</u> ; ⁶⁷⁸ preprint; 2020	Patients with severe COVID-19. 50 assigned to umifenovir 100 mg two twice a day for 7 to 14 days and 50 assigned to lopinavir-ritonavir 400 mg a day for 7 to 14 days	Mean age 56.4 ± 16.3, male 60%, hypertension 39%, diabetes 28%, asthma 2%, coronary heart disease 9%, chronic kidney disease 2%	Hydroxychloroquine 100%	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Yethindra et al</u> ; ⁶⁷⁹ peer-reviewed; 2020	Patients with mild COVID-19. 15 assigned to umifenovir 200 mg three times a day for 1 to 5 days and 15 assigned to standard of care	Mean age 35.5 ± 12.1, male 60%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of	



				allocation is probably inappropriate.	
<u>Ghaderkhani S et al</u> (<u>Tehran University</u> <u>of Medical Sciences</u>) <u>trial;</u> ⁶⁸⁰ Ghaderkhani et al; preprint; 2020	Patients with mild to moderate COVID-19. 28 assigned to umifenovir 200 mg three times a day for 10 days and 25 assigned to standard of care	Mean age 44.2 ± 19, male 39.6%,	Hydroxychloroquine 100%	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	
<u>UAIIC trial</u> ; ⁶⁸¹ Darazam et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 51 assigned to umifenovir 600 mg a day for 10 days and 50 assigned to SOC	Mean age 61.2 ± 15.8, male 56.4%, hypertension 46.4%, diabetes 31.6%, COPD 10%, asthma 6.1%, CHD 11.2%, CKD 7.1%, cancer 1%	Corticosteroids 3%	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Ramachandran et</u> <u>al;</u> ⁶⁸² preprint; 2021	Patients with mild to moderate COVID-19 infection. 60 assigned to umifenovir 800 mg twice a day for 14 days and 63 assigned to SOC	Mean age 46.7 ± 1.9, male 74.8%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	
	Uncerta	Ver inty in potential benefits a	apamil and harms. Further rese	arch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					





ReCOVery-SIRIO trial; ¹⁸ Navarese et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 72 assigned to verapamil 120 to 480 mg a day and 72 assigned to SOC	Median age 61.3 , male 62.3%, diabetes 23.7%, COPD 6.5%, cancer 7%	Remdesivir 1.9%, hydroxychloroquine 2.3%, azithromycin 6%, convalescent plasma 1.9%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: Very low certainty ⊕○○○Symptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No
					information
	Vilobelimab probal		belimab probably does not increa	ase severe adverse events.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT	·				
<u>Vlaar et al;</u> ⁶⁸³ peer- reviewed; 2020	Patients with severe COVID-19 infection. 15 assigned to vilobelimab 800 mg IV with a maximum of seven doses and 15 assigned to standard of care	Mean age 60 ± 9, male 73%, hypertension 30%, diabetes 27%, obesity 20%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably	Mortality: RR 0.76 (95%CI 0.6 to 0.98); RD -3.8% (95%CI - 6.4% to -0.3%); Moderate certainty ⊕⊕⊕○ Invasive mechanical ventilation: No information

PANAMO trial (phase 3); ⁶⁸⁴ Vlaar et al; peer reviewed; 2022	Patients with critical COVID-19 infection. 177 assigned to vilobelimab 800 mg (six infusions) and 191 assigned to SOC	Mean age 56.3, male 68.5%, hypertension 46.2%, diabetes 29.6%, COPD 2%, CHD 7%, CKD 6.2%, cancer 1.1%, obesity 40.7%	NR	inappropriate. Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: RR 0.94 (95%CI 0.8 to 1.11); RD -0.6% (95%CI -2% to 1.1%); Moderate certainty ⊕⊕⊕○ Hospitalization: No information
	Uncertai	Vita inty in potential benefits a	amin B nd harms. Further resea	rch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Majidi et al;</u> 685 peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 40 assigned to Vit B IM thiamine (10 mg), riboflavin (4 mg), nicotinamide (40 mg), and dexpanthenol (6 mg) once a day for 14 days and 45 assigned to SOC	Mean age 61.2	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	Mortality: No information Invasive mechanical ventilation: No information Symptom resolution or improvement: No information Symptomatic infection

					(prophylaxis studies): No information Adverse events: No information Hospitalization: No information
Vitamin C may incr	ease symptom resolution	ı or improvement. Vitami	amin C n C effects on other imp eeded.	ortant outcomes are uncer	tain. Further research is
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT			•	-	•
<u>Zhang et al</u> ; ⁶⁸⁶ preprint; 2020	Patients with severe COVID-19 infection. 26 assigned to vitamin C 12 g twice a day for 7 days and 28 assigned to standard of care	Mean age 67.4 ± 12.4, male 66.7%, hypertension 44.4%, diabetes 29.6%, chronic lung disease 5.6%, coronary heart disease 22.2%, chronic kidney disease 1.85%, cancer 5.6%, nervous system disease 20.4%	NR	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: Very low certainty ⊕○○○ Symptom resolution or improvement: RR
Kumari et al; ⁶⁸⁷ Peer reviewed; 2020	Patients with severe COVID-19. 75 assigned to Vit C 50 mg/kg a day and 75 assigned to SOC	Mean age 52.5 ± 11.5	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	1.16 (95%CI 1.01 to 1.33); RD 9.7% (95%CI 0.6% to 20%); Low certainty ⊕⊕○○ Symptomatic infection (prophylaxis studies): No information
<u>Jamali Moghadam</u> <u>Siahkali et a</u> l; ⁶⁸⁸		Mean age 59.2 ± 17, male 50%, hypertension	Hydroxychloroquine 100%, lopinavir-	High for mortality and mechanical ventilation;	Adverse events: No



Preprint; 2020	assigned to Vit C 5 g a day for 5 days and 30 assigned to SOC	41.6%, diabetes 38.3%, COPD 10%,	ritonavir 100%	High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	information Hospitalization: Very low certainty ⊕○○○
COVIDAtoZ - Vit C trial; ⁶⁸⁹ Thomas et al; peer reviewed; 2020	Patients with mild COVID-19. 48 assigned to Vit C 8000 mg a day and 50 assigned to SOC	Mean age 45.2 ± 14.6, male 38.3%, hypertension 32.7%, diabetes 13.6%, COPD %, asthma 15.4%	Corticosteroids 8.4%,	Low for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
VCACS trial; ⁶⁹⁰ Tehrani et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 18 assigned to Vit C 8 gr a day for 5 days and 26 assigned to SOC	Mean age 59.5, male 59%, hypertension 40.9%, diabetes 34%, COPD 7%, CHD 22.7%, CKD 9.1%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
Beigmohammadi et al; ⁶⁹¹ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 30 assigned to multivitamin vitamin D 600000 UI once, vitamin A 25000 UI a day, vitamin E 300 UI a day, vitamin C 2000 mg a day in addition to others for 7 days. and 30 assigned to SOC	51.6%, hypertension 33.3%, diabetes 18.3%, asthma 13.3%, cancer 5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	



Study; publication status	Patients and interventions analyzed		Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
Vitamin D does not	t reduce SARS-COV-2 in		amin D duals and probably does	not reduce hospitalization	s. Vitamin D effects on
<u>Fogleman C et al</u> <u>trial;⁴²² peer</u> reviewed; 2022	Patients with mild to moderate COVID-19 infection. 32 assigned to vitamin C 1000 mg a day for 14 days and 34 assigned to SOC	Median age 52, male 44.9%, hypertension 26.5%, diabetes 16.3%	Vaccinated 2%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	
<u>Coppock et al</u> ; ⁶⁹⁴ peer reviewed; 2021		Mean age 60, male 50%, hypertension 62.1%, diabetes 34.8%, COPD 19.7%	Corticosteroids 77.3%, remdesivir 92.4%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
ALLIANCE trial; ⁶⁹³ Ried et al; peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 162 assigned to vitamin C 400 mg/kg a day for 7 days and 75 assigned to SOC	Mean age 62.3 ± 15.7, male 50%, diabetes 35%, COPD 34%, CHD 36%, cancer 4%,	Hydroxychloroquine 100%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>Majidi et al</u> ; ⁶⁹² peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 31 assigned to vitamin C 500 mg a day and 69 assigned to SOC	Mean age 62.4 ± , male 60%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	



RCT					
COVIDIOL trial; Entrenas Castillo et al; ⁶⁹⁵ peer-reviewed; 2020	Patients with moderate to severe COVID-19. 50 assigned to vitamin D 0.532 once followed by 0.266 twice and 26 assigned to standard of care	Mean age 52.95 ± 10, male 59.2%, hypertension 34.2%, diabetes 10.5%, chronic lung disease 7.9%, coronary heart disease 3.9%, immunosuppression 9.2%	Hydroxychloroquine 100%, azithromycin 100%	High for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Mortality: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$ Invasive mechanical ventilation: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
<u>SHADE trial;</u> ⁶⁹⁶ Rastogi et al; peer- reviewed; 2020	Patients with mild to moderate COVID-19. 16 assigned to vitamin D 60000 IU a day for 7 days and 24 assigned to standard of care	Mean age 48.7 ± 12.4, male 50%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	Symptom resolution or improvement: Very low certainty ⊕○○○ Symptomatic infection (prophylaxis studies): RR 1.06 (95%CI 0.91 to 1.24)
<u>Murai et al</u> ; ⁶⁹⁷ peer- reviewed; 2020	Patients with severe COVID-19. 117 assigned to vitamin D 200,000 IU once and 120 assigned to standard of care	Mean age 56.3 ± 14.6, male 56.3%, hypertension 52.5%, diabetes 35%, COPD %, asthma 6.3%, coronary heart disease 13.3%, chronic kidney disease 1%,	NR	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	RD 1% (95%CI -1.6% to 4.2%); High certainty ⊕⊕⊕⊕ Adverse events: RR 1.03 (95%CI 0.84 to 1.26); RD 0.3% (95%CI -1.6% to 2.7%); Low certainty
<u>Lakkireddy et al;</u> ⁶⁹⁸ preprint; 2021	Patients with mild to moderate with low plasmatic vitamin D COVID-19 infection. 44 assigned to vitamin D 60000 IU a day for 8 to 10 days and 43 assigned to SOC	Mean age 45.5 ± 13.3, male 75%	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	 ⊕⊕○○ Hospitalization: RR 1.2 (95%CI 0.83 to 1.74); RD 1% (95%CI -0.8% to 3.6%); Moderate certainty ⊕⊕⊕○

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<u>Sabico et al</u> ; ⁶⁹⁹ peer reviewed; 2021	Patients with moderate to critical COVID-19 infection. 36 assigned to vitamin D 5000 IU for 14 days and 33 assigned to vitamin D 1000 IU for 14 days	Mean age 49.8 ± 14.3, male 49.3%, hypertension 55%, diabetes 51%, COPD %, asthma 4%, CHD 6%, CKD 7%, obesity 33%	NR	Low for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
<u>Maghbooli et al</u> ; ⁷⁰⁰ peer reviewed; 2021	Patients with moderate to severe COVID-19 infection. 53 assigned to vitamin D3 25 µg a day for 30 days and 53 assigned to SOC	Mean age 49.1 ± 14.1, male 60.4%, hypertension 31.1%, diabetes 23.6%, COPD 10.3%, CHD 12.3%, CKD 2.8%	Corticosteroids 46.2%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.	
Beigmohammadi et al; ⁷⁰¹ peer reviewed; 2021	Patients with severe to critical COVID-19 infection. 30 assigned to multivitamin vitamin D 600000 UI once, vitamin A 25000 UI a day, vitamin E 300 UI a day, vitamin C 2000 mg a day in addition to others for 7 days, and 30 assigned to SOC	Mean age 52 ± 9, male 51.6%, hypertension 33.3%, diabetes 18.3%, asthma 13.3%, cancer 5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
REsCue trial; ⁷⁰² Bishop et al; preprint; 2021	Patients with mild to moderate COVID-19 infection. 65 assigned to vitamin D calcifediol 300 mcg a day for three days followed by 60 mcg a day for 27 days and 69 assigned to SOC	Mean age 43, male 41%, hypertension 21.6%, diabetes 6%, asthma 2.2%, CKD 3%, obesity 40%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events	



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<u>Karonova et al</u> ; ⁷⁰³ peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 45 assigned to cholecalciferol 50,000 IU/week for 2 weeks followed by 500 UI/day for 3 months and 46 assigned to cholecalciferol 5000 IU/day for 3 months	Mean age 35 ± 2, male 15.3%, obesity 16.5%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
<u>COVID-VIT-D</u> <u>trial</u> ; ⁷⁰⁴ Cannata- Andía et al; peer reviewed; 2021	Patients with severe COVID-19 infection. 274 assigned to vitamin D Cholecalciferol 100.000UI once and 269 assigned to SOC	Median age 58, male 65%, hypertension 43.8%, diabetes 24.7%, COPD 4.2%, asthma 5.5%, CHD 21.2%	Corticosteroids 29.9%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	
CORONAVIT trial; ⁷⁰⁵ Jolliffe et al; preprint; 2021	Individuals exposed to SARS-CoV-2 infection. 3030 assigned to vitamin D 800 to 3200 UI a day and 2949 assigned to SOC	Median age 60.2, male 67%, hypertension 3.7%, diabetes 4.2%, COPD 1.8%, asthma 15.3%, CHD 19.5%, obesity 20.1%	NR; Vaccinated 1.3%	Low for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
Villasis-Keever et <u>al;</u> ⁷⁰⁶ peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 150 assigned to vitamin D 4,000 IU cholecalciferol a day for 30 days and 152 assigned to SOC	male 30%, hypertension	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate. Significant loss to follow up.	



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CARED-TRIAL trial; ⁷⁰⁷ Mariani et al; peer reviewed; 2021	Patients with moderate COVID-19 infection. 115 assigned to vitamin D 500 000 IU of vitamin D3 once and 103 assigned to SOC	Mean age 59.1 ± 10.6, male 52.8%, hypertension 43.1%, diabetes 26.6%, COPD 11.9%, CHD 4.6%, cancer 0.9%, obesity 39.9%	NR	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events
COVIT-TRIAL <u>trial</u> ; ⁷⁰⁸ Annweiler et al; peer reviewed; 2022	Patients with mild to severe COVID-19 infection. 127 assigned to vitamin D cholecalciferol 400.000 UI once and 127 assigned to vitamin D 50.000 UI	Median age 88, male 46%, hypertension 70%, diabetes 21%, COPD 7%, CHD 43%, CKD 17%, cerebrovascular disease 19%, cancer 7%, obesity 22%	Corticosteroids 15%, hydroxychloroquine 0.4%,azithromycin 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>Karonova et al</u> ; ⁷⁰⁹ peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 65 assigned to vitamin D cholecalciferol 100,000 IU and 64 assigned to SOC	Mean age 60.5, male 59.2%, hypertension 73.6%, diabetes 31.8%, COPD %, CHD 23.3%, obesity 38.8%	Vaccinated 0%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events
<u>Romero-</u> <u>Ibarguengoitia et</u> <u>al</u> ; ⁷¹⁰ preprint; 2022	Individuals exposed to SARS-CoV-2 infection. 43 assigned to vitamin D 52,000 IU a month for 6 months and 42 assigned to SOC	Mean age 44.4 ± 11.1, male 58.8%, hypertension 10%, diabetes 7%, asthma 4.7%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.
<u>Cervero et al</u> ; ⁷¹¹ peer reviewed; 2022	Patients with severe COVID-19 infection. 41 assigned to vitamin D cholecalciferol 10000 IU a day for 14 days and 44 assigned to Vit D 2000 IU a day	Median age 65 ± , male 71%, hypertension 48%, diabetes 22%	Corticosteroids 87%, remdesivir 15%, tocilizumab 25%, azithromycin 44%,	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded





	for 14 days			study. Concealment of allocation probably inappropriate.
Abroug et al; ⁷¹² preprint; 2022	Patients with mild with persistently positive PCR test at 14 days COVID-19 infection. 57 assigned to vitamin D cholecalciferol 200,000 IU once and 60 assigned to SOC	Mean age 42.7 ± 14, male 55.6%, hypertension 6.8%, diabetes 12%, asthma 6.8%	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.
D-COVID trial; ⁷¹³ De Niet et al; peer reviewed; 2022	Patients with moderate to severe COVID-19 infection. 21 assigned to cholecalciferol 25.000 UI a day for 4 days followed by 25.000 UI a week for 6 weeks and 22 assigned to SOC	Mean age 66, male 53.5%, hypertension 55.8%, diabetes 37.2%, COPD 32.6%, CKD 18.6%	Corticosteroids 100%, remdesivir 100%; Vaccinated 14%	High for mortality and mechanical ventilation; High for symptom resolution, infection and adverse events Notes: Concealment of allocation probably inappropriate.
Brunvoll et al; ⁷¹⁴ peer reviewed; 2022	Patients with exposed to COVID-19 infection. 17278 assigned to Vit D 400 IU a day in the form of cod liver oil for 164 days (median) and 17323 assigned to SOC	Mean age 44.9 ± 13.4, male 35.4%, comorbidities 22.2%	Vaccinated 35.6%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:
<u>Van Helmond et</u> <u>al:</u> ⁷¹⁵ preprint; 2022	Patients with exposed COVID-19 infection. 299 assigned to cholecalciferol 5000 IU a day and 578 assigned to SOC	Mean age 49, male 21.2%, diabetes 6.6%, cancer 5.5%,	NR	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.





Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
POLYCOR trial; ⁷¹⁶ Gaborit et al; preprint; 2021	Patients with severe COVID-19 infection. 12 assigned to XAV-19 0.5 to 2 mg/kg on days 1 and 5 and 5 assigned to SOC	Mean age 71 ± 24, male 64.7%, hypertension 47.1%, diabetes 11.8%, COPD %, asthma 17.6%, CHD 29.4%, CKD 5.9%, cancer 11.8%, obesity 17.6%	Corticosteroids 100%, remdesivir 47.1%	Low for mortality and mechanical ventilation; low for symptom resolution, infection, and adverse events	Mortality: Very low certainty ⊕○○○ Invasive mechanicativentilation: No information Symptom resolution or improvement: No information Symptomatic infection (prophylaxis studies): No information Adverse events: Very low certainty ⊕○○○ Hospitalization: No information
	Uncertai	Zilu inty in potential benefits a	ICOPlan and harms. Further rese	arch is needed.	
Study; publication tatus	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence



ZILU-COV trial; ⁷¹⁷ Leeuw et al; peer- reviewed; 2021	Patients with severe COVID-19 infection. 54 assigned to zilucoplan 32.4 mg a day, subcutaneously, for 14 days and 24 assigned to SOC	Median age 63, male 87%, hypertension 46%, diabetes 23%, asthma %, CHD 24%, CKD 5%	Corticosteroids 86%, remdesivir 12%	High for mortality and mechanical ventilation; high for symptom resolution, infection and adverse events Notes: Non-blinded study. Concealment of allocation probably inappropriate.	Mortality: Very low certainty ⊕○○○Invasive mechanical ventilation: No informationSymptom resolution or improvement: No informationSymptomatic infection (prophylaxis studies): No informationAdverse events: Very low certainty ⊕○○○Hospitalization: No information
Zinc may not impr	ove symptom resolution.		Zinc of the evidence was low b	ecause of imprecision. Its e	effects on other clinical
Study; publication	imp Patients and	oortant outcomes are unce Comorbidities	ertain. Further research Additional	is needed. Risk of bias and study	Interventions effects
status	interventions analyzed		interventions	limitations	vs standard of care and GRADE certainty of the evidence
RCT	•		•	-	
<u>Hassan et al</u> ; ⁷¹⁸ preprint; 2020	Patients with mild to critical COVID-19. 49 assigned to zinc 220	Mean age 45.9 ± 17.5, male 58.2%, hypertension 10.4%,	NR	High for mortality and mechanical ventilation; high for symptom	Mortality: Very low certainty $\oplus \bigcirc \bigcirc \bigcirc$
	mg twice a day and 56 assigned to standard of care	diabetes 11.2%, coronary heart disease 3%		resolution, infection, and adverse events Notes: Concealment of	Invasive mechanical ventilation: Very low certainty ⊕○○○
				allocation probably inappropriate.	Symptom



Abd-Elsalam et al; ⁷¹⁹ peer-reviewed; 2020	Patients with mild to critical COVID-19. 96 assigned to zinc 220 mg twice a day for 15 days and 95 assigned to standard of care	57.7%, hypertension 18.4%, diabetes 12.9%	Hydroxychloroquine 100%,	High for mortality and mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	resolution or improvement: RR 1.01 (95%CI 0.91 to 1.12); RD 0.6% (95%CI -5.4% to 7.3%); Low certainty ⊕⊕○○ Symptomatic infection
<u>Abdelmaksoud et</u> <u>al;</u> ⁷²⁰ Peer reviewed; 2020	Patients with mild to critical COVID-19. 49 assigned to Zinc 220 mg twice a day and 56 assigned to SOC	NR	NR	High for mortality and mechanical ventilation; High for symptom resolution, infection, and adverse events Notes: Non-blinded study. Concealment of allocation is probably inappropriate.	 (prophylaxis studies): Very low certainty ⊕○○○ Adverse events: No information Hospitalization: Very low certainty ⊕○○○
<u>COVIDAtoZ -Zinc</u> <u>trial;</u> ⁶⁸⁹ Thomas et al; ; 2020	Patients with mild COVID-19. 58 assigned to Zinc 50 mg a day and 50 assigned to SOC	Mean age 45.2 ± 14.6, male 38.3%, hypertension 32.7%, diabetes 13.6%, COPD %, asthma 15.4%	Corticosteroids 8.4%,	Low for mortality and mechanical ventilation; Some concerns for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results.	
ZINC COVID <u>trial;</u> ⁷²¹ Patel et al; Peer reviewed; 2020	Patients with severe to critical COVID-19. 15 assigned to Zinc 0.24 mg/kg a day for 7 days and 18 assigned to SOC	Mean age 61.8 ± 16.9, male 63.6%, hypertension 48.4%, diabetes 18.2%, COPD 6%, CHD 21.2%,	Corticosteroids 75.8%, remdesivir 30.3%,	Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events	
<u>Seet et al</u> ; ²⁹⁴ peer reviewed; 2021	Individuals exposed to SARS-CoV-2 infection. 634 assigned	Mean age 33 , male 100%, hypertension 1%, diabetes 0.3%	NR	Low for mortality and mechanical ventilation; High for symptom	



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Reszinate trial; ⁵⁷⁰ Kaplan et al; preprint; 2021	to zinc 80 mg and 500 mg a day for 42 days and 619 assigned to SOC (vitamin C) Patients with mild COVID-19 infection. 14 assigned to resveratrol + zinc 4000/150 mg once a day for five days and	Mean age 42.4, male 40%	NR	resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse events outcomes results. Low for mortality and mechanical ventilation; Low for symptom resolution, infection, and adverse events Notes:	
Stambouli et al; ²⁰⁵ peer reviewed; 2022	16 assigned to SOC Individuals exposed to SARS-CoV-2 infection. 59 assigned to zinc 15 mg a day for 6 weeks and 56 assigned to SOC	Mean age 38.4 ± 10.7, male 61%, hypertension 4.1%, diabetes 2.3%, COPD 0.6%, asthma 1.2%	Vaccinated 0%	Low for mortality and mechanical ventilation; low for symptom resolution, infection and adverse events Notes:	
	Uncerta	α-lip inty in potential benefits a	oic acid and harms. Further resea	urch is needed.	
Study; publication status	Patients and interventions analyzed	Comorbidities	Additional interventions	Risk of bias and study limitations	Interventions effects vs standard of care and GRADE certainty of the evidence
RCT					
<u>Zhong et al</u> ; ⁷²² preprint; 2020	Patients with critical COVID-19 infection. 8 assigned to α-lipoic acid 1200 mg infusion once daily for 7 days and 9 assigned to standard of care	Median age 63 ± 7, male 76.5%, hypertension 47%, diabetes 23.5%, coronary heart disease 5.9%	NR	Low for mortality and invasive mechanical ventilation; high for symptom resolution, infection, and adverse events Notes: Non-blinded study which might have introduced bias to symptoms and adverse	Mortality: Very low certainty ⊕○○○ Invasive mechanical ventilation: No information Symptom resolution or improvement: No information





		events outcomes results.	Symptomatic infection (prophylaxis studies): No information
			Adverse events: No information
			Hospitalization: No information





Appendix 1. Summary of findings tables

Summary of findings Table 1. (Interactive online version)

Population: Patients with severe COVID-19 disease Intervention: Corticosteroids Comparator: Standard of care

Outcome Timeframe	Study results and measurements	Absolute effec		Certainty of the Evidence (Quality of evidence)	Plain language summary	
		Standard of care	Steroids	(Quality of evidence)		
Mortality 28 days	Relative risk: 0.9 (CI 95% 0.8 - 1.01) Based on data from 8000 participants in 12 studies	160 per 1000 Difference: 16 fe (Cl 95% 32 few		Moderate Due to serious imprecision ¹	Steroids probably decreases mortality	
Mechanical ventilation	Relative risk: 0.87 (Cl 95% 0.72 - 1.05) Based on data from 5942	172 per 1000	150 per 1000	Moderate Due to serious	Steroids probably decreases mechanical	
28 days	participants in 6 studies Follow up 28	Difference: 22 fe v (CI 95% 48 few		imprecision ²	ventilation	
Symptom resolution	Relative risk: 1.27 (Cl 95% 0.98 - 1.65)	606 per 1000	770 per 1000	Moderate	Steroids probably	
or improvement Based on data from 646	Based on data from 646 participants in 5 studies	Difference: 164 m (Cl 95% 12 fewe		Due to serious risk of bias ³	increases symptom resolution or improvement	
Severe adverse	Relative risk: 0.89 (Cl 95% 0.68 - 1.17)	102 per 1000	91 per 1000	Low	Steroids may have little o	
events 28 days	events Based on data from 833	Difference: 11 fer (CI 95% 33 fewe		Due to serious risk of bias, Due to serious imprecision ⁴	no difference on severe adverse events	
Mortality (High vs standard dose) (Low	Relative risk: 0.97 (Cl 95% 0.78 - 1.21)	160 per 1000	155 per 1000	Low	High dose steroids (i.e dexamethasone 12 mg a day) may not decreases	
	Based on data from 2060 participants in 5 studies	Difference: 5 fev (Cl 95% 35 fewe	ver per 1000 er – 34 more)	Due to very serious imprecision ⁵	mortality in comparison to standard dose steroids (i.e dexamethasone 6 mg a day)	
Severe adverse events (High vs. standard dose) 28 days	Relative risk: 0.82 (Cl 95% 0.6 – 1.11)	102 per 1000	84 per 1000	Low	High dose steroids (i.e dexamethasone 12 mg a day) may not increase	
	Based on data from 1280 participants in 2 studies	Difference: 18 fe v (CI 95% 41 fewe		Due to very serious imprecision ⁶	severe adverse events in comparison to standard dose steroids (i.e dexamethasone 6 mg a day)	

1. Imprecision: serious. 95%CI includes no mortality reduction;

2. Imprecision: serious. 95%CI include no IVM reduction;





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- 3. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias;
- 4. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** Low number of patients;
- 5. Imprecision: very serious. 95%CI includes no mortality decrease;
- 6. Imprecision: very serious. Low number of patients, Wide confidence intervals.



Summary of findings Table 2. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Remdesivir Comparator: Standard of care

Outcome	Study results and	Absolute eff	ect estimates	Certainty of the Evidence	Blain language our	
Timeframe	measurements	SOC	Remdesivir	(Quality of evidence)	Plain language summar	
Mechanical ventilation	Relative risk: 0.76 (Cl 95% 0.56 - 1.04) Based on data from 9730	173 per 1000	131 per 1000	Moderate Due to serious	Remdesivir probably decrease mechanical	
28 days	participants in 7 studies Follow up Median 28 days		fewer per 1000 ewer - 7 more)	imprecision ¹	ventilation requirements	
Mortality	Relative risk: 0.93 (Cl 95% 0.89 - 1.03) Based on data from 10855	160 per 1000	149 per 1000	Moderate Due to serious	Remdesivir probably	
28 days	participants in 8 studies Follow up Median 28 days		fewer per 1000 ewer - 5 more)	imprecision ²	reduces mortality	
Symptom resolution	Relative risk: 1.1 (Cl 95% 0.96 - 1.28)	606 per 1000	667 per 1000	Low Due to serious risk of	Remdesivir may improve	
or improvement 28 days	Based on data from 1981 participants in 4 studies Follow up 28 days		more per 1000 wer - 170 more)	bias, Due to serious imprecision ³	symptom resolution or improvement	
Severe adverse	Relative risk: 0.77 (CI 95% 0.46 - 1.29)	102 per 1000	79 per 1000	Low Due to serious risk of	Remdesivir may have littl	
events	events Based on data from 2430 participants in 4 studies Difference: 23 fewer per 1000 (Cl 95% 55 fewer - 30 more)	bias, Due to serious	or no difference on sever adverse events			
Hospitalization (in patients with non-	Relative risk: 0.28 (Cl 95% 0.11 - 0.75)	48 per 1000	13 per 1000	Low	Remdesivir may decreas	
patients with non- severe disease) 28 days	Based on data from 562 participants in 1 study Follow up Median 28 days	Difference: 35 fewer per 1000 (Cl 95% 43 fewer - 12 fewer)		Due to very serious imprecision ⁵	hospitalizations (in patients with non-severe disease)	

1. **Imprecision: serious.** Wide confidence intervals;

2. Imprecision: serious. Wide confidence intervals;

 Risk of Bias: serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Imprecision: serious. 95%CI includes significant benefits and absence of benefits;

4. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** 95%ci included significant severe adverse events increase;

5. Imprecision: very serious.





Summary of findings Table 3. (Interactive online version)

Population: Patients with COVID-19 infection or exposed to COVID-19 Intervention: Hydroxychloroquine (HCQ) Comparator: Standard of care

Outcome	Study results and	Absolute effect es	stimates	Certainty of the	Plain language	
Timeframe	measurements	SOC	HCQ	Evidence (Quality of evidence)	summary	
Mortality 15 days	Relative risk: 1.09 (Cl 95% 1 - 1.19) Based on data from 10904 participants in 16 studies	I.	171 per 1000	Moderate Due to serious risk of bias ¹	HCQ probably increases mortality	
	Parto-parto 11 10 01000	Difference: 14 more (Cl 95% 0 fewer - 3				
Mechanical ventilation	Relative risk: 1.08 (Cl 95% 0.93 - 1.25) Based on data from 8667	173 per 1000	187 per 1000	Moderate Due to serious risk of	Hcq probably has little or no difference on	
15 days	participants in 10 studies		Difference: 14 more per 1000 (CI 95% 12 fewer - 43 more)		mechanical ventilation	
	Relative risk: 1.01 (Cl 95% 0.93 - 1.1)	606 per 1000	612 per 1000	Moderate	Hcq probably has little or	
	Based on data from 6601 participants in 10 studies Follow up 28 days	Difference: 6 more (Cl 95% 42 fewer -		Due to serious inconsistency ³	no difference on symptor resolution or improvement	
COVID-19 infection (in exposed individuals) (Low risk	Relative risk: 0.88 (CI 95% 0.72 - 1.11) Based on data from 4523	174 per 1000	153 per 1000	Low Due to serious	Hcq may have little or no difference on covid-19	
of bias studies)	participants in 6 studies	Difference: 21 fewer (Cl 95% 49 fewer -		imprecision, Due to serious inconsistency ⁴	infections (in exposed individuals)	
Hospitalizations (in patients with non- severe disease) (CI 95% Based on d	Relative risk: 0.82 (CI 95% 0.61 - 1.1) Based on data from 4255	48 per 1000	39 per 1000	Low Due to very serious	Hcq may have little or no difference on hospitalizations in	
	participants in 9 studies	Difference: 9 fewer (Cl 95% 19 fewer -		imprecision ⁵	patients with non-severe disease	
Severe adverse events	Relative risk: 0.9 (Cl 95% 0.66 - 1.22) Based on data from 10381	102 per 1000	92 per 1000	Low Due to serious risk of	Hcq may have little or no difference on severe	
	participants in 20 studies	Difference: 10 fewer (Cl 95% 35 fewer -		bias, Due to serious imprecision ⁶	difference on severe adverse events	

1. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias;

2. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias;

 Risk of Bias: no serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Inconsistency: serious. I2 82%; Imprecision: no serious. Secondary to inconsistency;

4. **Inconsistency: serious.** The direction of the effect is not consistent between the included studies; **Imprecision: serious.** 95%CI includes no infection reduction;





- 5. Imprecision: very serious. 95%CI includes significant benefits and harms;
- 6. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** Low number of patients.





Summary of findings Table 4. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Lopinavir-ritonavir (LPV) Comparator: Standard of care

Outcome Time frameStudy results and measurementsAbsolute		Absolute eff	ect estimates	Certainty of the evidence (quality of evidence)	Plain text summary
		SOC	LPV	(quality of ordeneo)	
Mortality 28 days	Relative risk: 1.01 (CI 95% 0.92 - 1.11) Based on data from 8059	160 per 1000	162 per 1000	Moderate Due to serious imprecision ¹	LPV probably has little or no difference on mortality
	patients in 4 studies Follow-up median 28 days	Difference: 2 more per 1000 (CI 95% 13 fewer - 18 more)			
Mechanical ventilation 28 days	Relative risk: 1.07 (CI 95% 0.98 - 1.17) Based on data from 7622	173 per 1000	185 per 1000	High	LPV does not reduce mechanical ventilation
	patients in 4 studies Follow-up median 28 days	Difference: 12 more per 1000 (CI 95% 3 fewer - 29 more)			
Symptom resolution or improvement	Relative risk: 1.03 (CI 95% 0.92 - 1.15) Based on data from 5239	606 per 1000) 624 Moderate per 1000 Due to serious risk of bias	Moderate Due to serious risk of bias ²	LPV probably has little or no difference on symptom resolution
28 days	patients in 2 studies Follow-up 28 days	Difference: 18 more per 1000 (CI 95% 48 fewer - 91 more)			or improvement
Symptomatic infection (exposed individuals)	Relative risk: 1.4 (CI 95% 0.78 - 2.54) Based on data from 318	174 per 1000	244 per 1000	Very low Due to serious risk of bias, Due to very serious	We are uncertain whether LPV increases or decreases
	patients in 1 study	10	70 more per 00 wer - 268 more)	imprecision ³	symptomatic infection in exposed individuals
	Relative risk: 0.6 (CI 95% 0.37 - 0.98) Based on data from 199	102 per 1000	61 per 1000	Low Due to serious risk of bias, Due to serious imprecision ⁴	LPV may have little or no difference on severe adverse events
	patients in 1 study	Difference: 41 fewer per 1000 (CI 95% 64 fewer - 2 fewer)			
Hospitalization	Relative risk: 1.22 (CI 95% 0.61 - 2.47)	48 per 1000	59 per 1000	Very low	We are uncertain whether LPV



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1. Imprecision: Serious. 95%CI includes significant mortality reduction and increase;

 Risk of bias: Serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Imprecision: No serious. Secondary to inconsistency;

3. Risk of bias: Serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; Imprecision: Very serious. 95%CI includes significant benefits and harms;

 Risk of bias: Serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Imprecision: Serious. Low number of patients;

5. Imprecision: Very serious. 95%CI includes significant benefits and harms.





Summary of findings Table 5. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Convalescent plasma Comparator: Standard of care

Outcome	Study results and	Absolute eff	bsolute effect estimates Certainty of the Evidence Plain language s		Plain language summary
Timeframe	tromo moscuromonte	(Quality of evidence)	Fiam language summary		
Mechanical ventilation	Relative risk: 1.03 (Cl 95% 0.94 - 1.11) Based on data from 14363	173 per 1000	176 per 1000	High	Convalescent plasma has little or no difference on
28 days	participants in 22 studies Follow up Median 28 days		more per 1000 wer - 19 more)		mechanical ventilation
Mortality 28 days	Relative risk: 0.98 (Cl 95% 0.93 - 1.03) Based on data from 24156	160 per 1000	157 per 1000	High	Convalescent plasma has little or no difference on
28 days	participants in 50 studies Follow up Median 28 days		fewer per 1000 ewer - 5 more)		mortality
Symptom resolution or improvement	Relative risk: 0.99 (Cl 95% 0.95 - 1.02) Based on data from 14487	606 per 1000	600 per 1000	High	Cp has little or no difference on symptom resolution or
28 days	participants in 13 studies Follow up 28 days		fewer per 1000 wer - 12 more)		improvement
Hospitalizations	Relative risk: 0.77 (Cl 95% 0.57 - 1.03) Based on data from 2642	48 per 1000	37 per 1000	Moderate Due to serious	Coucalescent plasma probably has little or no
	participants in 4 studies		fewer per 1000 ewer - 1 more)	imprecision ²	difference on hospitalizations
Severe adverse	Relative risk: 1.05 (Cl 95% 0.9 - 1.22)	102 per 1000	104 per 1000	Low	Convalescent may have
events	Based on data from 7451 participants in 17 studies		more per 1000 wer - 22 more)	Due to serious imprecision, Due to serious risk of bias ³	little or no difference on severe adverse events
Symptomatic	Relative risk: 0.92 (CI 95% 0.32 - 2.62)	174 per 1000	160 per 1000	Very low	We are uncertain whether
infection	Based on data from 168 participants in 1 study Difference: 14 fewer per 1000 (Cl 95% 118 fewer - 282 more) Due to extremely serious imprecision ⁴	Due to extremely serious imprecision ⁴	cp increases or decreases symptomatic infection		
Specific severe adverse events	Based on data from 20000 participants in 1 study	events were: TR	f severe adverse ALI 0.1%, TACO gic reactions 0.1%	Very low Due to very serious risk of bias ⁵	We are uncertain whether lpv increases or decreases severe adverse events

1. Inconsistency: no serious. Point estimates vary widely;

2. Imprecision: serious. Wide confidence intervals;

3. Risk of Bias: serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; Imprecision: serious. Wide confidence intervals;

4. Imprecision: ~extreme_serious. Wide confidence intervals;





5. **Risk of Bias: very serious.** Although adverse events were rare, we assume that some might have been missed and assumed as related to disease progression. RCT are needed to determine interventions safety.



Summary of findings Table 6. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Tocilizumab (TCZ) Comparator: Standard of care

Outcome	Study results and	Absolute effect estimates		Certainty of the Evidence	Plain language
Timeframe	measurements	SOC	TCZ	(Quality of evidence)	summary
Mortality	ortality Based on data from 8541	High	TCZ decreases mortality		
28 days	participants in 21 studies Follow up Median 28 days		fewer per 1000 wer - 11 fewer)		
Mechanical	Relative risk: 0.84 (CI 95% 0.79 - 0.91) Based on data from 7655	173 per 1000	145 per 1000	High	TCZ decreases
28 days p	participants in 21 studies Follow up Median 28 days	Difference: 28 fewer per 1000 (Cl 95% 36 fewer - 16 fewer)		1	mechanical ventilation
Symptom resolution	Relative risk: 1.08 (CI 95% 1.02 - 1.14) Based on data from 7077	606 per 1000	648 per 1000	Low Due to serious	TCZ may increase
28 days participant	participants in 11 studies Follow up 28 days	Difference: 48 more per 1000 (Cl 95% 12 more - 85 more)		imprecision, Due to serious risk of bias ²	symptom resolution or improvement
Severe adverse	Relative risk: 0.95 (CI 95% 0.86 - 1.04)	102 per 1000	97 per 1000	Moderate	Tcz probably has little o
events	Based on data from 5412 participants in 17 studies	Difference: 5 fewer per 1000 (Cl 95% 14 fewer - 4 more)		Due to serious risk of bias ³	no difference on severe adverse events

1. Imprecision: no serious. 95% included significant and trivial reduction mechanical ventilation requirement reduction ;

2. Risk of Bias: serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; Imprecision: serious. 95%CI includes significant benefits and absence of benefits ;

3. Risk of Bias: serious. Imprecision: no serious. 95%ci included significant severe adverse events increase.



Summary of findings Table 7. (Interactive online version)

Population: Patients with COVID-19 infection

Intervention & comparator: Anticoagulants in intermediate (i.e., enoxaparin 1 mg/kg a day); anticoagulants in full dose (i.e., enoxaparin 1 mg/kg twice a day); anticoagulants in prophylactic dose (i.e., enoxaparin 40 mg a day); no anticoagulants

Outcome	Study results and	Absolute effect estimates	Certainty of the Evidence	Plain language summary	
Timeframe	measurements	SOC ACO	(Quality of evidence)		
Mortality (full or intermediate dose vs. prophylactic dose in hospitalized patients) (excluding high risk of bias studies)	Relative risk: 0.99 (Cl 95% 0.83 - 1.19) Based on data from 5874 participants in 10 studies	160 158 per 1000 per 1000 Difference: 2 fewer per 1000 (Cl 95% 27 fewer - 30 more)	Moderate Due to serious imprecision ¹	Anticoagulants in intermediate or full dose probably have little or no difference on mortality in comparison with prophylactic dose	
studies)				propriylactic dose	
Venous thromboembolic events (intermediate dose vs. prophylactic	Relative risk: 0.82 (Cl 95% 0.43 - 1.59) Based on data from 1115	70 57 per 1000 per 1000	Low	Anticoagulants in intermediate dose may	
dose vs. propriyactic dose in hospitalized patients)	participants in 4 studies	Difference: 13 fewer per 1000 (Cl 95% 40 fewer - 41 more)	Due to very serious imprecision ²	slightly reduce venous thromboembolic events	
Clinically important bleeding (prophylactic dose	Relative risk: 2.5 (Cl 95% 0.49 - 12.8)	9 23 per 1000 per 1000	Very low	It is uncertain if anticoagulants in prophylactic dose increase or decrease clinically important bleeding	
vs. no anticoagulants in mild ambulatory patients)	Based on data from 444 participants in 1 study	Difference: 14 more per 1000 (Cl 95% 5 fewer - 106 more)	Due to very serious imprecision ³		
Venous thromboembolic events (full dose vs. prophylactic dose in	Relative risk: 0.56 (Cl 95% 0.44 - 0.71) Based on data from 5235	70 39 per 1000 per 1000	High	Anticoagulants in intermediate or full dose probably decreases venous thromboembolic events (ful dose)	
hospitalized patients)	participants in 8 studies	Difference: 31 fewer per 1000 (CI 95% 39 fewer - 20 fewer)			
Major bleeding (full or intermediate dose vs. prophylactic dose	Relative risk: 1.56 (Cl 95% 1.08 - 2.25) Based on data from 6343	19 30 per 1000	Moderate Due to serious	Anticoagulants in intermediate or full dose	
in hospitalized patients)	participants in 11 studies	Difference: 11 more per 1000 (Cl 95% 2 more - 24 more)	imprecision ⁴	probably increases major bleeding	
Hospitalization (prophylactic dose vs. no anticoagulants	Relative risk: 0.94 (Cl 95% 0.55 - 1.59) Based on data from 1549	48 45 per 1000 per 1000	Low Due to serious risk of	Anticoagulants may have	
in mild ambulatory patients)	participants in 4 studies	Difference: 3 fewer per 1000 (Cl 95% 22 fewer - 28 more)	bias, Due to serious imprecision ⁵	little or no difference on hospitalization	
Symptom resolution or improvement (prophylactic dose	Relative risk: 1.08 (Cl 95% 0.92 - 1.27)	606654per 1000per 1000	Low	Anticoagulants may have little or no difference on	
vs. no anticoagulants in mild ambulatory patients)	Based on data from 444 participants in 1 study	Difference: 48 more per 1000 (Cl 95% 48 fewer - 164 more)	Due to very serious imprecision ⁶	little or no difference on symptom resolution or improvement	





Mortality: Intermediate dose (i.e enoxaparin 40 mg every 12 hs) vs. prophylactic dose (i.e enoxaparin 40 mg a day) ⁷ 28 days	Relative risk: 0.29 (Cl 95% 0.13 - 0.64) Based on data from 843 participants in 2 studies	160 46 per 1000 per 1000 Difference: 114 fewer per 1000 (Cl 95% 139 fewer - 58 fewer)	Very low Due to very serious risk of bias ⁸	We are uncertain whether ACO intermediate dose increases or decreases mortality in comparison to ACO prophylactic dose
Mortality: Therapeutic dose (i.e enoxaparin 1 mg/kg every 12 hs) vs. prophylactic dose (i.e enoxaparin 40 mg a day) ⁹ 28 days	Relative risk: 2.02 (Cl 95% 0.7 - 5.8) Based on data from 2409 participants in 5 studies	160 323 per 1000 per 1000 Difference: 163 more per 1000 (Cl 95% 48 fewer - 768 more)	Very low Due to very serious risk of bias, Due to very serious imprecision ¹⁰	We are uncertain whether ACO in therapeutic dose increases or decreases mortality in comparison to ACO in prophylactic dose

1. Imprecision: serious. Low number of patients;

2. Imprecision: very serious. 95%CI includes significant benefits and harms;

3. Imprecision: very serious. 95%CI includes harms and absence of harms;

4. Imprecision: serious. 95%CI includes harms and absence of harms;

5. Risk of Bias: serious. Imprecision: serious. 95%CI includes harms and absence of harms;

6. Imprecision: very serious. 95%CI includes harms and absence of harms;

7. Therapeutic dose (i.e enoxaparin 40 mg every 12 hs) vs. prophylactic dose (i.e enoxaparin 40 mg a day)

8. Risk of Bias: very serious.

9. Therapeutic dose (i.e enoxaparin 1 mg/kg every 12 hs) vs. prophylactic dose (i.e enoxaparin 40 mg a day)

10. Risk of Bias: very serious. Imprecision: very serious. 95%CI includes significant mortality reduction and increase.



Summary of findings Table 8. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Non-corticosteroids anti-inflammatory drugs (NSAID) Comparator: Standard of care

Outcome Time frame	Study results and measurements	Absolute effect estimates		Certainty of the evidence (quality of evidence)	Plain text summary
		SOC	NSAID		
Mortality 28 days	Odds Ratio: 0.83 (CI 95% 0.66 - 1.05) Based on data from	160 per 1000	137 per 1000	Very low Due to very serious risk of bias ¹	We are uncertain whether NSAID increases or decreases
	2465490 patients in 6 studies	10	23 fewer per 000 Wewer - 7 more)		mortality

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1. Risk of bias: Very serious.





Summary of findings Table 9. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Interferon beta-1a (IFN-B-1a) Comparator: Standard of care

Outcome Timeframe	Study results and	Absolute effect estimates		Certainty of the Evidence	Plain language	
	measurements	SOC	IFN	(Quality of evidence)	summary	
Mortality	Relative risk: 0.99 (Cl 95% 0.75 - 1.31) Based on data from 6869	160 per 1000	171 per 1000	Moderate Due to serious	IFN probably has little or no difference on	
28 days	patients in 6 studies Follow up Median 28 days		fewer per 1000 wer - 50 more)	imprecision ¹	mortality	
Mechanical ventilation	Relative risk: 1.01 (Cl 95% 0.87 - 1.18) Based on data from 5052	173 per 1000	168 per 1000	Moderate Due to serious	IFN probably has little or no difference on mechanical ventilation	
28 days	patients in 4 studies Follow up 28 days		more per 1000 ewer - 31 more)	imprecision ²		
Symptom resolution or improvement	Relative risk: 0.96 (Cl 95% 0.92 - 0.99) Based on data from 969	606 per 1000	582 per 1000	Moderate	IFN probably has little or no difference on	
28 days	patients in 1 study Follow up 28 days		fewer per 1000 ewer - 6 fewer)	Due to serious imprecision ³	symptom resolution or improvement	
Severe adverse events	Relative risk: 0.94 (Cl 95% 0.65 - 1.37) Based on data from 877	102 per 1000	96 per 1000	Low Due to very serious	IFN may have little or	
28 days	patients in 1 study Follow up 28 days		fewer per 1000 wer - 38 more)	imprecision ⁴	adverse events	
Symptom resolution or improvement	Hazard Ratio: 2.19 (CI 95% 1.03 - 4.69)	606 per 1000	870 per 1000	Low Due to very serious imprecision ⁶	IFN (inhaled) may	
(inhaled) ⁵ 30 days	Based on data from 81 patients in 1 study Follow up 28 days		4 more per 1000 ore - 381 more)		increase symptom resolution or improvement	

1. **Imprecision: serious.** 95%CI includes significant mortality reduction and increase;

2. **Risk of Bias: no serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Imprecision: serious.** 95% included significant mechanical ventilation requirement reduction and increase;

3. Imprecision: serious. 95%CI includes significant benefits and absence of benefits;

4. Imprecision: very serious. 95%CI includes significant benefits and absence of benefits;

5. Nebulizations;

6. Imprecision: very serious. 95%CI includes significant benefits and absence of benefits.





Summary of findings Table 10. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Bamlanivimab +/- etesevimab Comparator: Standard of care

Outcome Timeframe	Study results and	Absolute et	ffect estimates	Certainty of the		
	measurements	SOC	Bamlanivimab +/- etesevimab	Evidence (Quality of evidence)	Plain language summary	
Mortality	Relative risk: 0.68 (Cl 95% 0.17 - 2.8) Based on data from 2315	160 per 1000	109 per 1000	Very low Due to serious	We are uncertain whether bamlanivimab increases or decreases mortality	
	patients in 3 studies		1 fewer per 1000 fewer - 288 more)	imprecision, Due to very serious imprecision ¹		
Symptom resolution	Relative risk: 1.02 (CI 95% 0.99 - 1.06)	606 per 1000	618 per 1000	Moderate	Bamlanivimab probably has little or no difference on	
or improvement ²	Based on data from 1750 patients in 3 studies	Difference: 12 more per 1000 (Cl 95% 6 fewer - 36 more)		Due to serious imprecision ³	symptom resolution or improvement	
Symptomatic	Relative risk: 0.56 (Cl 95% 0.39 - 0.81) Based on data from 961 patients in 1 study Follow up 28 days	174 per 1000	97 per 1000	Moderate	Bamlanivimab probably	
infection		Difference: 77 fewer per 1000 (Cl 95% 106 fewer - 33 fewer)		Due to serious imprecision ⁴	decreases symptomatic infection	
Severe adverse	Hazard Ratio: 1.12 (CI 95% 0.75 - 1.66)	102 per 1000	114 per 1000	Low	Bamlanivimab may not	
events⁵	Based on data from 3661 patients in 6 studies		2 more per 1000 fewer - 62 more)	Due to very serious imprecision ⁶	increase severe adverse events	
Hospitalization ⁷	Hazard Ratio: 0.37 (Cl 95% 0.21 - 0.65)	48 per 1000	18 per 1000	Moderate	Bamlanivimab +/-	
	Based on data from 1804 patients in 3 studies		0 fewer per 1000 Fewer - 17 fewer)	Due to serious imprecision ⁸	etesevimab probably decreases hospitalization	

1. **Imprecision: very serious.** 95%CI includes significant benefits and harms;

2. Symptomatic infection in persons at risk or exposed to SARS-COV2;

3. Imprecision: serious. 95%CI includes benefits and absence of benefits;

4. Imprecision: serious. OIS not met;

5. Symptomatic infection in persons at risk or exposed to SARS-COV2;

6. Imprecision: very serious. 95%CI includes significant benefits and harms;

7. Symptomatic infection in persons at risk or exposed to SARS-COV2;

8. **Imprecision: serious.** Low number of patients



Summary of findings Table 11. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Favipiravir Comparator: Standard of care

Outcome Timeframe	Study results and	Absolute effe	ect estimates	Certainty of the Evidence	Plain language summary	
	measurements	SOC	Favipravir	(Quality of evidence)		
Mortality	Relative risk: 1.08 (Cl 95% 0.77 - 1.5) Based on data from 3247	160 per 1000	173 per 1000	Low Due to very serious	Favipiravir may increase	
28 days	participants in 12 studies Follow up Median 28 days	Difference: 13 (CI 95% 37 fev	more per 1000 wer - 80 more)	imprecision ¹	mortality	
Mechanical	Relative risk: 1.27 (Cl 95% 0.91 - 1.76)	173 per 1000	220 per 1000	Low	Favipravir may increase	
ventilation 28 days	Based on data from 1632 participants in 6 studies Follow up Median 28 days		Difference: 47 more per 1000 (Cl 95% 16 fewer - 131 more)		mechanical ventilation	
Symptom resolution or improvement (Low	Relative risk: 1.01 (CI 95% 0.97 - 1.05) Based on data from 2029	606 per 1000	612 per 1000	High	Favipiravir has little or no	
RoB studies) 28 days	participants in 4 studies Follow up 28 days	Difference: 6 I (CI 95% 18 fee	nore per 1000 wer - 30 more)		resolution or improvement	
Hospitalization (in patients with non-	Relative risk: 1.33 (Cl 95% 0.64 - 1.78)	48 per 1000	64 per 1000	Low	Favipravir may have little	
severe disease)	Based on data from 824 participants in 5 studies Follow up 28 days	Difference: 16 (CI 95% 17 fev	more per 1000 wer - 37 more)	Due to very serious imprecision ³	hospitalization (in patient with non-severe disease	
Severe adverse	Relative risk: 0.92 (CI 95% 0.56 - 1.52)	606 per 1000	558 per 1000	Very low Due to very serious	We are uncertain whethe favipiravir increases or	
events 30 days	Based on data from 2557 participants in 9 studies	Difference: 48 (CI 95% 267 fe	fewer per 1000 wer - 315 more)	imprecision, Due to serious risk of bias ⁴	adverse events	

1. Imprecision: very serious. 95%CI includes significant mortality reduction and increase;

2. Imprecision: very serious. 95%CI includes significant benefits and harms;

3. Imprecision: very serious. 95%CI includes significant benefits and absence of benefits ;

4. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: very serious.** 95%CI includes significant benefits and absence of benefits ;





Summary of findings Table 12. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Ivermectin Comparator: Standard of care

Outcome	Study results and measurements	Absolute effect estimates	Certainty of the	
Timeframe		SOC Ivermectin	Evidence (Quality of evidence)	Plain language summary
Mortality (Low risk of bias studies)	Relative risk: 1 (Cl 95% 0.8 - 1.24) Based on data from 6522 participants in 13 studies	160 158 per 1000 per 1000 Difference: 0 fewer per 1000 (Cl 95% 32 fewer - 38 more)	Moderate Due to serious imprecision ¹	lvermectin probably has little or no difference on mortality
Mechanical ventilation (Low risk of bias studies)	Relative risk: 0.82 (Cl 95% 0.58 - 1.17) Based on data from 3288 participants in 9 studies	173 142 per 1000 per 1000 Difference: 31 fewer per 1000 (Cl 95% 73 fewer - 29 more)	Very low Due to very serious imprecision ²	We are uncertain whether ivermectin increases or decreases mechanical ventilation (low risk of bias studies)
Symptom resolution or improvement (Low risk of bias studies)	Relative risk: 1.04 (CI 95% 0.98 - 1.1) Based on data from 3238 participants in 7 studies	606 630 per 1000 per 1000 Difference: 24 more per 1000 (Cl 95% 12 fewer - 61 more)	Moderate Due to serious imprecision ³	Ivermectin probably has little or no difference on symptom resolution or improvement
Symptomatic infection (Low risk of bias studies) ⁴	Relative risk: 1.01 (CI 95% 0.54 - 1.89) Based on data from 536 participants in 1 study	174 176 per 1000 per 1000 Difference: 2 more per 1000 (Cl 95% 80 fewer - 155 more)	Very low Due to very serious imprecision ⁵	We are uncertain whether ivermectin increases or decreases symptomatic infection
Severe adverse events	Relative risk: 1.05 (CI 95% 0.69 - 1.62) Based on data from 2831 participants in 8 studies Follow up 28 days	102 107 per 1000 per 1000 Difference: 5 more per 1000 (Cl 95% 32 fewer - 63 more)	Very low Due to very serious imprecision ⁶	Ivermectin may have little or no difference on severe adverse events
Hospitalization (in non-severe patients)	Relative risk: 0.9 (CI 95% 0.74 - 1.1) Based on data from 6315 participants in 11 studies Follow up 28 days	48 43 per 1000 per 1000 Difference: 5 fewer per 1000 (Cl 95% 12 fewer - 5 more)	Moderate Due to serious imprecision ⁷	Ivermectin probably has little or no difference on hospitalization

1. Imprecision: serious. 95%CI includes significant benefits and harms;

2. Imprecision: very serious. Wide confidence intervals;

- 3. Imprecision: serious. Wide confidence intervals;
- 4. Symptomatic infection in persons at risk or exposed to SARS-COV2;

5. Imprecision: very serious. Low number of patients;

6. Imprecision: very serious. 95%CI includes significant benefits and absence of benefits;

7. Imprecision: serious. Less than 200 events.





Summary of findings Table 13. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Baricitinib Comparator: Standard of care

Outcome	Study results and	Absolute effect estimates		Certainty of the Evidence	Plain language summary	
Timeframe	measurements		(Quality of evidence)	Flain language summary		
Mortality	Relative risk: 0.73 (CI 95% 0.57 - 0.92) Based on data from 11102	160 per 1000	117 per 1000	High	Baricitinib decreases	
	participants in 5 studies		fewer per 1000 wer - 13 fewer)		mortality	
Invasive mechanical	Relative risk: 0.83 (Cl 95% 0.66 - 1.04)	173 per 1000	144 per 1000	Moderate	Baricitinib probably	
	participants in 3 studies Follow up 30 days	Difference: 29 fewer per 1000 (Cl 95% 59 fewer - 7 more)		Due to serious imprecision ¹	decreases invasive mechanical ventilation	
Symptom resolution or improvement	Relative risk: 1.27 (CI 95% 1.13 - 1.42) Based on data from 2659	606 per 1000	770 per 1000	Moderate Due to serious risk of	Baricitinib probably improves symptom	
or improvement	participants in 3 studies Follow up 30 days		4 more per 1000 ore - 255 more)	bias ²	resolution or improvemen	
Severe adverse	Relative risk: 0.78 (CI 95% 0.64 - 0.95)	102 per 1000	80 per 1000	Moderate Due to serious risk of bias ³	Baricitinib probably has litt	
events	Based on data from 2659 participants in 3 studies Follow up 30 days		fewer per 1000 ewer - 5 fewer)		or no difference on severe adverse events	

1. Imprecision: serious. Wide confidence intervals;

2. Risk of Bias: serious. Incomplete data and/or large loss to follow up;

3. Risk of Bias: serious. Incomplete data and/or large loss to follow up.



Summary of findings Table 14. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Azithromycin Comparator: Standard of care

Outcome Timeframe	Study results and measurements	Absolute effect estimates SOC Azythromicin	Certainty of the Evidence (Quality of evidence)	Plain language summary	
Mortality	Relative risk: 1.01 (Cl 95% 0.92 - 1.1) Based on data from 8967 participants in 6 studies	160 162 per 1000 per 1000 Difference: 2 more per 1000 (Cl 95% 13 fewer - 16 more)	Moderate Due to serious imprecision ¹	Azythromicin probably has little or no difference on mortality	
Invasive mechanical ventilation	Relative risk: 0.92 (Cl 95% 0.77 - 1.1) Based on data from 8947 participants in 5 studies	173 159 per 1000 per 1000 Difference: 14 fewer per 1000 (Cl 95% 40 fewer - 17 more)	Moderate Due to serious imprecision ²	Azythromicin probably has little or no difference on invasive mechanical ventilation	
Symptom resolution or improvement ³	Relative risk: 1.02 (Cl 95% 0.99 - 1.04) Based on data from 9690 participants in 6 studies	606 618 per 1000 per 1000 Difference: 12 more per 1000 (Cl 95% 6 fewer - 24 more)	High	Azythromicin has little or no difference on symptom resolution or improvement	
Severe adverse events	Relative risk: 1.23 (CI 95% 0.51 - 2.96) Based on data from 439 participants in 1 study Follow up 28 days	102 125 per 1000 per 1000 Difference: 23 more per 1000 (Cl 95% 50 fewer - 200 more)	Very low Due to very serious imprecision, Due to very serious risk of bias ⁴	We are uncertain whether azythromicin increases or decreases severe adverse events	
Hospitalizations	Relative risk: 0.98 (Cl 95% 0.52 - 1.86) Based on data from 493 participants in 2 studies Follow up 21 days	48 47 per 1000 per 1000 Difference: 1 fewer per 1000 (Cl 95% 23 fewer - 41 more)	Low Due to serious risk of bias, Due to serious imprecision ⁵	Azythromicin may have litt or no difference on hospitalizations	

1. Imprecision: serious. 95%CI includes significant benefits and harms;

2. Imprecision: serious. 95%CI includes significant benefits and harms;

3. Symptomatic infection in persons at risk or exposed to SARS-COV2;

4. Risk of Bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Imprecision: very serious. 95%CI includes significant benefits and absence of benefits;

5. Risk of Bias: serious. Inadequate concealment of allocation during randomization process, resulting in potential for selection bias, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias, Incomplete data and/or large loss to follow up; Imprecision: serious. 95%CI includes significant benefits and absence of benefits.





Summary of findings Table 15. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Colchicine Comparator: Standard of care

Outcome	Study results and	Absolute effect estimates		Certainty of the		
Timeframe	measurements	SOC	Colchicine	Evidence (Quality of evidence)	Plain language summary	
Mortality	Relative risk: 0.99 (Cl 95% 0.92 - 1.05) Based on data from 18353 patients in 13 studies	160 per 1000	158 per 1000 fewer per 1000	Moderate Due to serious imprecision ¹	Colchicine probably has little or no difference on mortality	
			ewer - 8 more)			
Invasive mechanical ventilation	Relative risk: 0.98 (Cl 95% 0.89 - 1.07) Based on data from 17053	173 per 1000	170 per 1000	Moderate Due to serious	Colchicine probably has little or no difference on invasive mechanical	
	patients in 7 studies Follow up 30 days		fewer per 1000 ewer - 12 more)	imprecision ²	ventilation	
Symptom resolution or improvement	Relative risk: 1 (CI 95% 0.98 - 1.02) Based on data from 11784	173 per 1000	175 per 1000	High	Colchicine has little or no difference on symptom	
	patients in 5 studies Follow up 30 days		more per 1000 ewer - 3 more)		resolution or improvement	
Severe adverse events	Relative risk: 0.78 (Cl 95% 0.61 - 0.99) Based on data from 4880	102 per 1000	80 per 1000	High	Colchicine has little or no difference on severe	
	patients in 3 studies Follow up 30 days		fewer per 1000 ewer - 1 fewer)		adverse events	
Pulmonary embolism	Relative risk: 5.55 (Cl 95% 1.23 - 25.0) Based on data from 4399	0.9 per 1000	5.0 per 1000	Low Due to very serious	Colchicine may have little o no difference on pulmonary	
	patients in 1 study Follow up 30 days		more per 1000 nore - 21.6 more)	imprecision ³	embolism	
Hospitalization (in patients with non- severe disease)	Relative risk: 0.81 (Cl 95% 0.63 - 1.04) Based on data from 4777	48 per 1000	39 per 1000	Moderate Due to serious	Colchicine probably has little or no difference on	
severe uisease)	patients in 2 studies Follow up 30 days		fewer per 1000 ewer - 2 more)	imprecision ⁴	hospitalization (in patients with non-severe disease)	

1. Imprecision: serious. 95%CI includes significant benefits and harms;

2. Imprecision: serious. 95%CI includes benefits and harms;

3. **Imprecision: very serious.** 95%CI includes significant benefits and absence of benefits , low number of patients, wide confidence intervals;

4. Imprecision: serious. Low number of patients.





Summary of findings Table 16. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Sofosbuvir +/- daclatasvir, ledipasvir, or velpatasvir Comparator: Standard of care

	Absolute ef	fect estimates		
Study results and measurements	SOC	Sofosbuvir +/- daclatasvir, ledipasvir or velpatasvir	Certainty of the Evidence (Quality of evidence)	Plain language summary
Relative risk: 1.02 (Cl 95% 0.59 - 1.76) Based on data from 1163 participants in 2 studies Follow up 30 days			Low Due to very serious imprecision ¹	Sofosbuvir +/- daclatasvir, ledipasvir or velpatasvir may have little or no difference on invasive mechanical ventilation
Relative risk: 1.14 (Cl 95% 0.83 - 1.56) Based on data from 1163 participants in 2 studies			Low Due to very serious imprecision ²	Sofosbuvir alone or in combination may have little or no difference on mortalit
Relative risk: 0.35 (Cl 95% 0.06 - 2.18) Based on data from 628 participants in 2 studies			Very low Due to serious risk of bias, Due to very serious imprecision ³	We are uncertain whether sofosbuvir +/- daclatasvir, ledipasvir or velpatasvir increases or decreases severe adverse events
Relative risk: 1.01 (Cl 95% 0.95 - 1.08) Based on data from 1163 participants in 2 studies Follow up 7 days			Moderate Due to serious imprecision ⁴	Sofosbuvir alone or in combination probably has little or no difference on symptom resolution or improvement
Relative risk: 0.52 (Cl 95% 0.3 - 0.89) Based on data from 548 participants in 1 study			Very low Due to serious risk of bias, Due to very serious imprecision ⁵	We are uncertain whether sofosbuvir +/- daclatasvir, ledipasvir or velpatasvir increases or decreases symptomatic infection
	measurementsRelative risk: 1.02 (CI 95% 0.59 - 1.76)Based on data from 1163 participants in 2 studies Follow up 30 daysRelative risk: 1.14 (CI 95% 0.83 - 1.56)Based on data from 1163 participants in 2 studiesRelative risk: 0.35 (CI 95% 0.06 - 2.18)Based on data from 628 participants in 2 studiesRelative risk: 1.01 (CI 95% 0.95 - 1.08)Based on data from 1163 participants in 2 studiesRelative risk: 0.52 (CI 95% 0.3 - 0.89) Based on data from 548	Study results and measurementsSOCRelative risk: 1.02 (CI 95% 0.59 - 1.76)173 per 1000Based on data from 1163 participants in 2 studies Follow up 30 daysDifference: 3 (CI 95% 71 fetRelative risk: 1.14 (CI 95% 0.83 - 1.56)160 per 1000Based on data from 1163 participants in 2 studiesDifference: 22 (CI 95% 27 fetRelative risk: 0.35 (CI 95% 0.06 - 2.18)102 per 1000Based on data from 628 participants in 2 studiesDifference: 66 (CI 95% 96 fetRelative risk: 1.01 (CI 95% 0.95 - 1.08)606 per 1000Based on data from 1163 participants in 2 studiesDifference: 66 (CI 95% 96 fetRelative risk: 0.52 (CI 95% 0.3 - 0.89)Difference: 67 per 1000Based on data from 548 participants in 1 study174 per 1000	measurementsSOCdaclatasvir, ledipasvir or velpatasvirRelative risk: 1.02 (CI 95% 0.59 - 1.76)173176 per 1000Based on data from 1163 participants in 2 studies Follow up 30 days173176 per 1000Relative risk: 1.14 (CI 95% 0.83 - 1.56)Difference: 3 more per 1000 (CI 95% 71 fewer - 131 more)Relative risk: 1.14 (CI 95% 0.83 - 1.56)160182 per 1000Based on data from 1163 participants in 2 studiesDifference: 22 more per 1000 (CI 95% 27 fewer - 90 more)Relative risk: 0.35 (CI 95% 0.06 - 2.18)102 per 100036 per 1000Based on data from 628 participants in 2 studies102 per 100036 per 1000Relative risk: 1.01 (CI 95% 0.95 - 1.08)606 per 1000612 per 1000Relative risk: 1.01 (CI 95% 0.95 - 1.08)606 per 1000612 per 1000Based on data from 1163 participants in 2 studiesDifference: 6 more per 1000 per 1000Relative risk: 0.52 (CI 95% 0.3 - 0.89)174 per 100090 per 1000Relative risk: 0.52 (CI 95% 0.3 - 0.89)174 per 100090 per 1000	Study results and measurementsSofosition for the contraction of the contraction

1. Imprecision: very serious. 95%CI includes significant benefits and harms;

2. Imprecision: very serious. 95%CI includes significant benefits and harms;

 Risk of Bias: serious. Incomplete data and/or large loss to follow up, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; Inconsistency: serious. Imprecision: very serious. Wide confidence intervals;

4. Inconsistency: serious. Imprecision: serious. Wide confidence intervals;

5. **Risk of Bias: serious.** Incomplete data and/or large loss to follow up, Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias; **Inconsistency: serious. Imprecision: very serious.** Wide confidence intervals.



Summary of findings Table 17. (Interactive online version)

Patients with COVID-19 infection Intervention: REGEN-COV (casirivimab and imdevimab) Comparator: Standard of care

		Absolute effect estimates		Certainty of the		
Outcome Timeframe	Study results and measurements	SOC	REGEN-COV (casirivimab and imdevimab)	Evidence (Quality of evidence)	Plain language summary	
Mortality	Relative risk: 0.83 (Cl 95% 0.63 - 1.09) Based on data from 16845	160 per 1000	133 per 1000	Low Due to serious inconsistency, Due to	Regen-cov (casirivimab and imdevimab) may	
	participants in 4 studies		7 fewer per 1000 fewer - 14 more)	serious imprecision ¹	decrease mortality	
Mortality (seronegative)	Relative risk: 0.79 (CI 95% 0.71 - 0.89) Based on data from 3673	160 per 1000	126 per 1000	Moderate Due to serious	Regen-cov (casirivimab and imdevimab) probabl	
(seronegauve)	participants in 2 studies		4 fewer per 1000 ewer - 18 fewer)	indirectness ²	decreases mortality in seronegative patients	
Invasive mechanical ventilation	Relative risk: 0.79 (Cl 95% 0.54 - 1.14) Based on data from 14575	173 per 1000	137 per 1000	Low Due to very serious	Regen-cov (casirivimab and imdevimab) may	
ventilation	participants in 3 studies Follow up 30 days		6 fewer per 1000 fewer - 24 more)	imprecision ³	decrease invasive mechanical ventilation	
Invasive mechanical ventilation	Relative risk: 0.82 (CI 95% 0.74 - 0.9) Based on data from 3603	173 per 1000	142 per 1000	Moderate Due to serious	Regen-cov (casirivimab and imdevimab) probably decreases invasive	
(seronegative)	participants in 2 studies		1 fewer per 1000 ewer - 17 fewer)	indirectness, Due to serious imprecision ⁴	mechanical ventilation in seronegative patients	
Symptom resolution or improvement	Relative risk: 1.06 (CI 95% 1.0 - 1.12) Based on data from 14746	606 per 1000	642 per 1000	Low Due to serious	Regen-cov (casirivimab and imdevimab) may increase symptom	
	participants in 3 studies	Difference: 36 more per 1000 (Cl 95% 0 fewer - 73 more)		imprecision, Due to serious inconsistency ⁵	resolution or improvement	
Symptom resolution or improvement	Relative risk: 1.1 (Cl 95% 1.06 - 1.14) Based on data from 6277	606 per 1000	667 per 1000	Moderate Due to serious	Regen-cov (casirivimab and imdevimab) probably increases symptom	
(seronegative)	participants in 3 studies Follow up 30 days		1 more per 1000 more - 85 more)	indirectness ⁶	resolution or improvement in seronegative patients	
Hospitalization (in patients with non- severe disease)	Relative risk: 0.28 (Cl 95% 0.19 - 0.42)	48 per 1000	13 per 1000	Moderate Due to serious imprecision ⁷	Regen-cov (casirivimab and imdevimab) probably reduces hospitalization ir	





	Based on data from 6732 participants in 4 studies Follow up 30 days	Difference: 35 fewer per 1000 (Cl 95% 39 fewer - 28 fewer)			patients with recent onset non-severe disease	
Symptomatic infection (in exposed		174 per 1000	42 per 1000	High	Regen-cov (casirivimab and imdevimab) decreases symptomatic	
individuals)	participants in 3 studies Follow up 30 days		2 fewer per 1000 ewer - 42 fewer)	8	infection in exposed individuals	
Severe adverse events	Relative risk: 0.51 (Cl 95% 0.38 - 0.67) Based on data from 12360	102 per 1000	52 per 1000	Moderate	Regen-cov (casirivimab and imdevimab) probably	
	participants in 6 studies Diff			imprecision ⁹	has little or no difference on severe adverse events	

1. **Risk of Bias: no serious.** Incomplete data and/or large loss to follow up; **Inconsistency: serious.** The confidence interval of some of the studies do not overlap with those of most included studies/ the point estimate of some of the included studies.; **Imprecision: serious.** Wide confidence intervals;

2. Risk of Bias: no serious. Incomplete data and/or large loss to follow up; Indirectness: serious. Subgroup analysis; Imprecision: very serious.

3. Risk of Bias: no serious. Incomplete data and/or large loss to follow up; Imprecision: very serious. Wide confidence intervals;

4. Risk of Bias: no serious. Incomplete data and/or large loss to follow up; Indirectness: serious. Subgroup analysis;

 Inconsistency: serious. The confidence interval of some of the studies do not overlap with those of most included studies/ the point estimate of some of the included studies; Imprecision: serious. Wide confidence intervals;

6. Indirectness: serious. Subgroup analysis;

7. Risk of Bias: no serious. Incomplete data and/or large loss to follow up; Imprecision: serious. Low number of events;

8. Risk of Bias: no serious. Incomplete data and/or large loss to follow up;

9. Imprecision: serious. Wide confidence intervals.



Summary of findings Table 18. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Sotrovimab Comparator: Standard of care

Outcome	Study results and	Absolute effe	ct estimates	Certainty of the Evidence	Plain language	
Timeframe	measurements	Standard of care	Sotrovimab	(Quality of evidence)	summary	
Mortality	Relative risk: 0.2 (CI 95% 0.01 - 4.16) Based on data from 1057	160 per 1000	32 per 1000	Very low Due to extremely serious	We are uncertain wheth	
	participants in 1 study	Difference: 128 f (CI 95% 158 few		imprecision ¹	decreases mortality	
Mechanical	Relative risk: 0.11 (CI 95% 0.01 - 2.06)	174 per 1000	19 per 1000	Very low	We are uncertain whether sotrovimab increases or decreases mechanical ventilation	
ventilation	Based on data from 1057 participants in 1 study	Difference: 155 f (CI 95% 172 few		Due to extremely serious imprecision ²		
Hospitalization	Relative risk: 0.2 (Cl 95% 0.08 - 0.48)	48 per 1000	10 per 1000	Moderate	Sotrovimab probably decreases hospitalizatio	
	Based on data from 1057 participants in 1 study	Difference: 38 fo (CI 95% 44 few		Due to serious imprecision ³		
Hospitalization (sotrovimab vs.	Relative risk: 1.07 (Cl 95% 0.88 - 1.3)	48 per 1000	51 per 1000	High	Sotrovimab has little o	
REGEN-COV)	Based on data from 3558 participants in 1 study	Difference: 3 m (Cl 95% 6 fewe		, ingri	hospitalization compar- to REGEN-COV	
Severe adverse	Relative risk: 0.34 (Cl 95% 0.18 - 0.68)	102 per 1000	35 per 1000	Moderate	Sotrovimab probably h	
events	Based on data from 1057 participants in 1 study	Difference: 67 fo (Cl 95% 84 few		Due to serious imprecision ⁴	little or no difference o severe adverse event	

1. Imprecision: ~extremely_serious. Very low number of events;

2. Imprecision: ~extremely_serious. Very low number of events;

3. Imprecision: serious;

4. Imprecision: serious. Low number of patients.



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Summary of findings Table 19. (Interactive online version)

Patients with COVID-19 infection Intervention: Inhaled corticosteroids Comparator: Standard of care

Outcome	Study results and	Absolute ef	fect estimates	Certainty of the		
Timeframe	measurements	SOC	Inhaled coticosteroids	Evidence (Quality of evidence)	Plain language summar	
Relative risk: 1.09 Symptom resolution or improvement ¹ Based on data from 3919	(CI 95% 0.99 - 1.2) Based on data from 3919	606 per 1000	661 per 1000	Low Due to serious risk of bias, Due to serious	Inhaled coticosteroids ma increase symptom	
	participants in 8 studies		5 more per 1000 wer - 121 more)	imprecision ²	resolution or improvemer	
Invasive mechanical ventilation	Relative risk: 0.94 (Cl 95% 0.44 - 1.98) Based on data from 1560	173 per 1000	163 per 1000	Very low Due to serious risk of bias,	We are uncertain whethe inhaled corticosteroids increases or decreases	
	participants in 1 study		fewer per 1000 wer - 170 more)	Due to very serious imprecision ³	invasive mechanical ventilation	
	Relative risk: 0.82 (Cl 95% 0.44 - 1.53)	160 per 1000	131 per 1000	Very low Due to serious risk of bias.	We are uncertain whethe	
Mortality	Based on data from 2345 participants in 5 studies		fewer per 1000 ewer - 85 more)	Due to very serious imprecision ⁴	increases or decreases mortality	
Severe adverse	Relative risk: 0.5 (Cl 95% 0.23 - 1.12)	102 per 1000	51 per 1000	Very low Due to serious risk of bias,	We are uncertain whethe inhaled coticosteroids	
	Based on data from 2014 participants in 4 studies		fewer per 1000 ewer - 12 more)	Due to very serious imprecision ⁵	increases or decreases severe adverse events	
Hospitalizations (C Based	Relative risk: 0.9 (CI 95% 0.7 - 1.15) Based on data from 3953	48 per 1000	43 per 1000	Moderate	Inhaled coticosteroids probably has little or no	
	participants in 5 studies	Difference: 5 fewer per 1000 (Cl 95% 14 fewer - 7 more)		bias ⁶	difference on hospitalizations	

1. Symptomatic infection in persons at risk or exposed to SARS-COV2

2. Risk of Bias: serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; Imprecision: serious. Wide confidence intervals;

3. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: very serious.** 95%CI includes significant benefits and harms;

4. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: very serious.** 95%CI includes significant benefits and harms;

5. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias; **Imprecision: very serious.** 95%CI includes significant benefits and absence of benefits , Wide confidence intervals;

6. Risk of Bias: serious. Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias.



Summary of findings Table 20. (Interactive online version)

Patients with COVID-19 infection Intervention: Fluvoxamine Comparator: Standard of care

Outcome	Study results and	Absolute eff	ect estimates	Certainty of the Evidence	Plain language	
Timeframe	measurements	SOC	Fluvoxamine	(Quality of evidence)	summary	
Mortality	Relative risk: 0.69 (Cl 95% 0.36 - 1.27) Based on data from 1497 patients in 1 study		110 per 1000 fewer per 1000 ewer - 43 more)	Very low Due to very serious imprecision ¹	There were too few who experienced the mortality in order to determine whether fluvoxamine made a difference	
Mechanical ventilation	Relative risk: 0.77 (Cl 95% 0.45 - 1.3) Based on data from 1497 patients in 1 study		133 per 1000 fewer per 1000 ewer - 52 more)	Very low Due to very serious imprecision ²	There were too few who experienced the mortality in order to determine whether fluvoxamine made a difference	
Hospitalizations	Relative risk: 0.79 (Cl 95% 0.6 - 1.03) Based on data from 2302 patients in 3 studies		38 per 1000 fewer per 1000 ewer - 1 more)	Moderate Due to serious imprecision ³	Fluvoxamine probably ha little or no difference or hospitalizations	
Severe adverse events ⁴	Relative risk: 0.81 (CI 95% 0.54 - 1.22) Based on data from 1649 patients in 2 studies		83 per 1000 fewer per 1000 ewer - 22 more)	Low Due to very serious imprecision ⁵	Fluvoxamine may not increase severe adverse events	

1. Imprecision: very serious. 95%CI includes significant benefits and harms;

2. Imprecision: very serious. 95%CI includes significant benefits and harms;

3. Imprecision: serious. 95%CI includes significant benefits and absence of benefits;

4. Symptomatic infection in persons at risk or exposed to SARS-COV2;

5. **Imprecision: very serious.** Wide confidence intervals.





Summary of findings Table 21. (Interactive online version)

Patients with COVID-19 infection Intervention: Molnupiravir Comparator: Standard of care

Outcome	Study results and	Absolute effect estimates		Certainty of the Evidence	Plain language	
Timeframe	measurements	Standard of care	Molnupiravir	(Quality of evidence)	summary	
Symptom resolution	Relative risk: 1.17 (Cl 95% 1.1 - 1.3) Based on data from 1513	606 per 1000	1000 per 1000	Low Due to very serious risk of	Molnupiravir may	
	participants in 2 studies Follow up 5	Difference: 394 (CI 95% 394 mo		bias ¹	resolution	
Mortality	Relative risk: 0.35 (CI 95% 0.06 - 2.19)	160 per 1000	56 per 1000	Very low	We are uncertain whether molnupiravir increases or decreases mortality	
	Based on data from 2202 participants in 4 studies	Difference: 104 f (CI 95% 150 few		Due to very serious imprecision ²		
Mechanical	Relative risk: 0.36 (Cl 95% 0.11 - 1.12)	173 per 1000	62 per 1000	Very low	We are uncertain whethe	
ventilation	Based on data from 1610 participants in 1 studies	Difference: 111 f (Cl 95% 154 fev		Due to very serious imprecision ³	molnupiravir increases or decreases mortality	
Hospitalization	Relative risk: 0.6 (Cl 95% 0.44 - 0.81)	48 per 1000	29 per 1000	High	Molnupiravir decreases	
	Based on data from 4050 participants in 6 studies (CI 95% 27 fewer - 9 fewer)		hospitalization			
	Relative risk: 0.75 (CI 95% 0.48 - 1.19)	102 per 1000	77 per 1000	Low	Molnupiravir may have	
	Based on data from 2219 participants in 4 studies Follow up 29	Difference: 25 f (CI 95% 53 few		Due to very serious imprecision ⁴	little or no difference on severe adverse events	

1. **Risk of Bias: very serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias;

2. Imprecision: very serious. 95%CI includes significant benefits and harms, Low number of patients;

3. Imprecision: very serious. 95%CI includes significant benefits and harms, Low number of patients;

4. Imprecision: very serious. 95%CI includes significant benefits and absence of benefits.



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Summary of findings Table 22. (Interactive online version)

Patients with COVID-19 infection Intervention: Nirmatrelvir-ritonavir Comparator: Standard of care

Outcome	Study results and	Absolute effe	ct estimates	Certainty of the		
Timeframe	measurements	Standard of care	Nirmatrelvir- ritonavir	Evidence (Quality of evidence)	Plain language summary	
Mortality	Relative risk: 0.04 (CI 95% 0.0 - 0.68) Based on data from 2085	160 per 1000	6 per 1000	Very low Due to very serious	We are uncertain whether nirmatrelvir-ritonavir	
	participants in 1 study	Difference: 154 f (CI 95% 160 fev		imprecision ¹	increases or decreases mortality	
Hospitalization	Relative risk: 0.12 (CI 95% 0.06 - 0.25)	48 per 1000	6 per 1000	Moderate	Nirmatrelvir-ritonavir probably decreases hospitalizations	
	Based on data from 2085 participants in 1 study	Difference: 42 fe (Cl 95% 45 few		Due to serious imprecision ²		
Relative risk: 0.49 Severe adverse (CI 95% 0.3 - 0.8)	102 per 1000	50 per 1000	Moderate	Nirmatrelvir-ritonavir probably has little or no		
events	Based on data from 2224 participants in 1 study Follow up 29	Difference: 52 fe (CI 95% 71 few		er per 1000	difference on severe adverse events	

1. Imprecision: very serious. 95%CI includes significant benefits and harms, low number of patients;

2. Imprecision: serious. 95%CI includes significant benefits and absence of benefits;

3. **Imprecision: serious.** Low number of events.



Summary of findings Table 23. (Interactive online version)

Patients with COVID-19 infection Intervention: Ruxolitinib Comparator: Standard of care

Outcome	Study results and	Absolute effect estimates		Certainty of the Evidence	Plain language	
Timeframe	measurements	Standard of care	Molnupiravir	(Quality of evidence)	summary	
Mortality	Relative risk: 0.72 (Cl 95% 0.59 - 0.89) Based on data from 686	160 per 1000	21 per 1000	Low Due to serious imprecision	Ruxolitinib may reduce	
	participants in 3 studies	Difference: 45 fe (Cl 95% 66 few		and incosistency ¹	mortality	
	Relative risk: 0.99 (CI 95% 0.49 - 1.99)	173 per 1000	171 per 1000	Very low Due to very serious	It is uncertain if ruxolitinib increases or	
ventilation	Based on data from 474 patients in 2 study	Difference: 2 fe (Cl 95% 32 fewe		imprecision ²	decreases mechanical ventilation	
Severe adverse	Relative risk: 1.12 (Cl 95% 0.69 - 1.82) Based on data from 679	102 per 1000	114 per 1000	Very low	It is uncertain if ruxolitinib increases or	
events	participants in 3 studies	Difference: 12 n (Cl 95% 79 fewe	nore per 1000 er - 100 more)	Due to very serious imprecision ²	decreases mechanical ventilation	
Symptom resolution	Relative risk: 1.05 (Cl 95% 0.89 – 1.24)	606 per 1000	606 per 1000	Low	Ruxolitinb may no	
- ,	Based on data from 685 participants in 3 studies	Difference: 30 more per 1000 (CI 95% 66 fewer - 145 more)		Due to very serious imprecision ²	increase symptom resolution	

1. Imprecision: serious. Low number of patients; Inconsistency: serious. Significant not explained heterogeneity;

2. Imprecision: very serious. 95%CI including important benefits and harms.



Summary of findings Table 24. (Interactive online version)

Patients with COVID-19 infection Intervention: CD24Fc Comparator: Standard of care

Outcome	Study results and	Absolute effect estimates		Certainty of the Evidence	Plain language	
Timeframe	measurements	SOC	CD24Fc	(Quality of evidence)	summary	
Mortality	Relative risk: 0.9 (CI 95% 0.49 - 1.69) Based on data from 234	160 per 1000	144 per 1000	Very low Due to extremely	We are uncertain whethe CD24Fc increases or	
	participants in 1 study Follow up 29 days		fewer per 1000 ver - 110 more)	serious imprecision ¹	decreases mortality	
	Relative risk: 0.57 (CI 95% 0.34 - 0.96)	173 per 1000	99 per 1000	Low Due to serious	CD24Fc may decrease	
ventilation	Based on data from 234 participants in 1 study Follow up 29 days		fewer per 1000 ewer - 7 fewer)	imprecision, Due to very serious imprecision ²	invasive mechanical ventilation	
Symptom resolution	Relative risk: 1.18 (Cl 95% 1.0 - 1.39)	606 per 1000	715 per 1000	Low	CD24Fc may increase	
or improvement	Based on data from 234 participants in 1 study Follow up 29 days		more per 1000 ver - 236 more)	Due to very serious imprecision ³	symptom resolution or improvement	
Severe adverse (CI 95%) events Based on o participan	Relative risk: 0.98 (Cl 95% 0.61 - 1.57)	102 per 1000	100 per 1000	Very low	We are uncertain whethe CD24Fc increases or	
	Based on data from 234 participants in 1 study Follow up 29 days		fewer per 1000 wer - 58 more)	Due to extremely serious imprecision ⁴	decreases severe adverse events	

1. Imprecision: ~extreme_serious. Low number of patients, Wide confidence intervals;

2. Imprecision: very serious. Wide confidence intervals, Low number of patients;

3. Imprecision: very serious;

4. Imprecision: ~extreme_serious. Wide confidence intervals, Low number of patients.



Summary of findings Table 25. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Vitamin D Comparator: Standard of care

Outcome	Study results and	Absolute effect estimates		Certainty of the	Plain language	
Timeframe	measurements	SOC	Vitamin D	Evidence (Quality of evidence)	summary	
Symptom resolution or improvement	Relative risk: 1.78 (CI 95% 1.1 - 2.94) Based on data from 43 participants in 1 studies	606 per 1000 Difference: 473 (CI 95% 61 mc	1079 per 1000 3 more per 1000 ore - 1176 more)	Very low Due to very serious imprecision, Due to serious risk of bias ¹	We are uncertain whether vitamin d increases or decrease invasive mechanical ventilation	
Mortality	Relative risk: 1.24 (CI 95% 0.8 - 1.91) Based on data from 1234 participants in 6 studies		198 per 1000 more per 1000 wer - 146 more)	Very low Due to very serious imprecision, Due to serious risk of bias ²	We are uncertain whether vitamin D increases or decrease mortality	
Invasive mechanical ventilation	Relative risk: 0.55 (Cl 95% 0.31 - 1.0) Based on data from 561 participants in 3 studies		95 per 1000 fewer per 1000 fewer - 0 fewer)	Very low Due to very serious imprecision, Due to serious risk of bias ³	We are uncertain whether vitamin d increases or decrease invasive mechanical ventilation	
Symptomatic infection (Excluding high RoB studies)	Relative risk: 1.06 (Cl 95% 0.91 - 1.24) Based on data from 40580 participants in 2 studies		184 per 1000 more per 1000 ewer - 42 more)	High	Vitamin D has little or no difference on symptomatic infectior (excluding high rob studies)	
Hospitalization	Relative risk: 1.2 (CI 95% 0.83 - 1.74) Based on data from 40882 participants in 3 studies		58 per 1000 more per 1000 wer - 36 more)	Moderate Due to serious imprecision ⁴	Vitamin D probably does not reduce hospitalizations	
Severe adverse events	Relative risk: 1.03 (Cl 95% 0.84 - 1.89) Based on data from 6197 participants in 2 studies Follow up 29 days		105 per 1000 more per 1000 ewer - 91 more)	Low Due to serious risk of bias, Due to serious imprecision ⁵	Vitamin D may not increase severe adverse events	

1. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; **Imprecision: very serious.** Wide confidence intervals, Low number of patients;

2. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; **Imprecision: very serious.** Low number of patients, Wide confidence intervals;

3. **Risk of Bias: serious.** Inadequate concealment of allocation during randomization process, resulting in potential for selection bias; **Imprecision: very serious.** Wide confidence intervals, Low number of patients;

4. Imprecision: serious. Low number of patients;

5. Risk of Bias: serious. Imprecision: serious. Wide confidence intervals, Low number of patients;





Summary of findings Table 26. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Tixagevimab–Cilgavimab Comparator: Standard of care

Outcome Timeframe	Study results and measurements	Absolute effect estimates		Certainty of the	Plain language
		SOC	Tixagevimab– Cilgavimab	Evidence (Quality of evidence)	summary
Symptom resolution or improvement	Relative risk: 1.03 (Cl 95% 0.99 - 1.08) Based on data from 1417 participants in 1 study	606 per 1000	624 per 1000	Moderate Due to serious	Tixagevimab– cilgavima probably has little or no difference on symptom
		Difference: 18 more per 1000 (Cl 95% 6 fewer - 48 more)		imprecision ¹	resolution or improvement
Mortality	Relative risk: 0.72 (Cl 95% 0.54 - 0.96) Based on data from 7492 participants in 3 studies	160 per 1000	115 per 1000	Moderate Due to serious	Tixagevimab– cilgavima probably decreases
		Difference: 45 fewer per 1000 (Cl 95% 74 fewer - 6 fewer)		imprecision ²	mortality
Symptomatic infection	Relative risk: 0.18 (CI 95% 0.09 - 0.35) Based on data from 5172 participants in 1 study Follow up 29 days	174 per 1000	31 per 1000	Moderate	Tixagevimab– cilgavima
		Difference: 143 fewer per 1000 (CI 95% 158 fewer - 113 fewer)		Due to serious risk of bias ³	probably decreases symptomatic infection
Severe adverse events	Relative risk: 0.95 (CI 95% 0.69 - 1.31) Based on data from 7492 participants in 3 studies	102 per 1000	97 per 1000	Low	Tixagevimab– cilgavima may have little or no
		Difference: 5 fewer per 1000 (Cl 95% 32 fewer - 32 more)		Due to very serious imprecision ⁴	difference on severe adverse events
Hospitalization	Relative risk: 0.42 (Cl 95% 0.24 - 0.74) Based on data from 903 participants in 1 study	102 per 1000	43 per 1000	Moderate	Tixagevimab– cilgavima
		Difference: 59 fewer per 1000 (Cl 95% 78 fewer - 27 fewer)		Due to serious imprecision ⁵	probably decreases hospitalization

1. Imprecision: serious. Low number of patients;

2. Imprecision: serious. Low number of patients;

3. **Risk of Bias: serious.** Inadequate/lack of blinding of participants and personnel, resulting in potential for performance bias, Inadequate/lack of blinding of outcome assessors, resulting in potential for detection bias;

4. Risk of Bias: serious. Imprecision: very serious. Wide confidence intervals;

5. Imprecision: serious. Low number of patients.





Summary of findings Table 27. (Interactive online version)

Population: Patients with COVID-19 infection Intervention: Vilobelimab Comparator: Standard of care

Outcome Timeframe	Study results and measurements	Absolute effect estimates		Certainty of the Evidence	Plain language
		SOC	Vilobelomab	(Quality of evidence)	summary
Mortality	Relative risk: 0.76 (Cl 95% 0.6 - 0.98) Based on data from 398 participants in 2 studies	160 per 1000	122 per 1000	Moderate	Vilobelimab probably
		Difference: 38 fewer per 1000 (Cl 95% 64 fewer - 3 fewer)		Due to serious imprecision ¹	decreases mortality
Severe adverse events	Relative risk: 0.94 (Cl 95% 0.8 - 1.11) Based on data from 298 participants in 2 studies	102 per 1000	96 per 1000		Vilobemilab probably
		Difference: 6 fewer per 1000 (Cl 95% 20 fewer - 11 more)		Moderate Due to serious imprecision ²	makes little or no difference on severe adverse events

7. **Imprecision: serious.** Low number of patients;

8. Imprecision: serious. Wide confidence intervals;



References

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- World Health Organization. Commentaries: Off-label use of medicines for COVID-19 (Scientific brief, 31 March 2020) [Internet]. Geneva: World Health Organization; 2020 [cited 7 December 2020]. Available from: https://www.who.int/newsroom/commentaries/detail/off-label-use-of-medicines-for-covid-19
- The L·OVE Platform. Methods for the special L·OVE of coronavirus infection [Internet] Santiago: Epistemonikos Foundation; 2020 [cited 7 December 2020]. Available from: https://app.iloveevidence.com/covid-19
- World Health Organization. WHO R&D Blueprint novel Coronavirus: outline of trial designs for experimental therapeutics. WHO reference number WHO/HEO/R&D Blueprint (nCoV)/2020.4. Geneva: World Health Organization; 2020. Available at: https://apps.who.int/iris/bitstream/handle/10665/330694/WHO-HEO-RDBlueprintnCoV-2020.4-eng.pdf?ua=1
- 4. Schünemann HJ, Cuello C, Akl EA, Mustafa RA, Meerpohl JJ, Thayer K, et al. GRADE Guidelines: 18. How ROBINS-I and other tools to assess risk of bias in nonrandomized studies should be used to rate the certainty of a body of evidence. J Clin Epidemiol 2019;111(July):105–14. Available from: https://doi.org/10.1016/j.jclinepi.2018.01.012.
- 5. Docherty AB, Mulholland RH, Lone NI, Cheyne CP, De Angelis D, Diaz-Ordaz K, et al. Changes in UK hospital mortality in the first wave of COVID-19: the ISARIC WHO Clinical Characterisation Protocol prospective multicentre observational cohort study. MedRxiv 2020. Available from:

http://medrxiv.org/lookup/doi/10.1101/2020.12.19.20248559

- 6. International Severe Acute Respiratory and emerging Infections Consortium, Hall M, Pritchard M, Dankwa EA, Baillie JK, Carson G, et al. ISARIC Clinical Data Report 20 November 2020 [Internet]. MedRxiv 2020. Available from: http://medrxiv.org/lookup/doi/10.1101/2020.07.17.20155218
- Chu DK, Akl EA, Duda S, Solo K, Yaacoub S, Schünemann HJ, et al. Physical distancing, face masks, and eye protection to prevent person-to-person transmission of SARS-CoV-2 and COVID-19: a systematic review and meta-analysis. Lancet 2020;395:1973-1987. Available from: https://doi.org/10.1016/S0140-6736(20)31142-9.





- 8. Hultcrantz M, Rind D, Akl EA, et al. The GRADE Working Group clarifies the construct of certainty of evidence. J Clin Epidemiol 2017; 87: 4–13.
- Zeng L, Brignardello-Petersen R, Hultcrantz M, et al. GRADE guidelines 32: GRADE offers guidance on choosing targets of GRADE certainty of evidence ratings. Journal of Clinical Epidemiology 2021; 137: 163–75.
- Sterne JAC, Savović J, Page MJ, Elbers RG, Blencowe NS, Boutron I, et al. RoB 2: A revised tool for assessing risk of bias in randomised trials. BMJ 2019;366:14898. Available from: https://doi.org/10.1136/bmj.14898.
- 11. Guyatt GH, Oxman AD, Vist GE, et al. GRADE: an emerging consensus on rating quality of evidence and strength of recommendations. BMJ 2008; 336: 924–26.
- 12. Axfors C, Schmitt AM, Janiaud P, van 't Hooft J, Abd-Elsalam S, Abdo EF, et al.. Mortality outcomes with hydroxychloroquine and chloroquine in COVID-19: an international collaborative meta-analysis of randomized trials [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.09.16.20194571.
- Fontana P, Casini A, Robert-Ebadi H, Glauser F, Righini M, Blondon M. Venous thromboembolism in COVID-19: systematic review of reported risks and current guidelines. Swiss Med Wkly 2020;150:w20301. Available from: https://doi.org/10.4414/smw.2020.20301.
- 14. Pan-American Health Organization. Guidelines for critical care of seriously ill adult patients with coronavirus (COVID-19) in the Americas: short version v-1. Washington DC: PAHO;2020. Available from: https://iris.paho.org/handle/10665.2/52184
- 15. Yuan X, Yi W, Liu B, Tian S, Cao F, Wang R, et al. Pulmonary radiological change of COVID-19 patients with 99mTc-MDP treatment [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.04.07.20054767.
- 16. Fakharian A, Barati S, Mirenayat M, Rezaei M, Haseli S, Torkaman P, et al. Evaluation of adalimumab effects in managing severe cases of COVID-19: A randomized controlled trial. International Immunopharmacology. 2021 Oct;99:107961.
- McElvaney OJ, McEvoy NL, Boland F, McElvaney OF, Hogan G, Donnelly K, et al. A randomized, double-blind, placebo-controlled trial of intravenous alpha-1 antitrypsin for acute respiratory distress syndrome secondary to COVID-19. Med. 2022 Mar;S2666634022001295.



- 18. Navarese EP, Podhajski P, Andreotti F, La Torre G, Gajda R, Radziwanowski A, et al. Ion channel inhibition with amiodarone or verapamil in symptomatic hospitalized nonintensive-care COVID-19 patients: The ReCOVery-SIRIO randomized trial. Cardiol J. 2022 Jul 29;VM/OJS/J/88627.
- 19. Siami Z, Aghajanian S, Mansouri S, Mokhames Z, Pakzad R, Kabir K, et al. Effect of Ammonium Chloride in addition to standard of care in outpatients and hospitalized COVID-19 patients: a randomized clinical trial. International Journal of Infectious Diseases. 2021 Apr;S1201971221003544.
- 20. Roshon M, Lemos-Filho L, Cherevka H, Goldberg L, Salottolo K, Bar-Or D. A Randomized Controlled Trial to Evaluate the Safety and Efficacy of a Novel Inhaled Biologic Therapeutic in Adults with Respiratory Distress Secondary to COVID-19 Infection. Infect Dis Ther [Internet]. 2021 Nov 14 [cited 2021 Dec 6]; Available from: https://link.springer.com/10.1007/s40121-021-00562-z
- 21. Bureau S, Dougados M, Tibi A, Azoulay E, Cadranel J, Emmerich J, et al. Effect of anakinra versus usual care in adults in hospital with COVID-19 and mild-to-moderate pneumonia (CORIMUNO-ANA-1): a randomised controlled trial. The Lancet Respiratory Medicine. 2021 Jan;S2213260020305567.
- 22. Kyriazopoulou E, Poulakou G, Milionis H, Metallidis S, Adamis G, Tsiakos K, et al. Early Anakinra Treatment for COVID-19 Guided by Urokinase Plasminogen Receptor [Internet]. Infectious Diseases (except HIV/AIDS); 2021 May [cited 2021 May 24]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.05.16.21257283
- 23. Declercq J, Van Damme KFA, De Leeuw E, Maes B, Bosteels C, Tavernier SJ, et al. Effect of anti-interleukin drugs in patients with COVID-19 and signs of cytokine release syndrome (COV-AID): a factorial, randomised, controlled trial. The Lancet Respiratory Medicine. 2021 Oct;S2213260021003775.
- 24. Kharazmi AB, Moradi O, Haghighi M, Kouchek M, Manafi-Rasi A, Raoufi M, et al. A randomized controlled clinical trial on efficacy and safety of anakinra in patients with severe COVID-19. Immun Inflamm Dis. 2021 Nov 11;iid3.563.
- 25. Elmekaty EZI, Maklad A, Abouelhassan R, Munir W, Mohamed Ibrahim MI, Nair A, et al. Efficacy of Anakinra in the Management of Patients with COVID-19 Infection: A Randomized Clinical Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jul



BE AWARE, PREPARE, ACT.

www.paho.org/coronavirus

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[cited 2022 Jul 18]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2022.07.04.22277207

- 26. Audemard-Verger A, Le Gouge A, Pestre V, Courjon J, Langlois V, Vareil MO, et al. Efficacy and safety of anakinra in adults presenting deteriorating respiratory symptoms from COVID-19: A randomized controlled trial. Plavec D, editor. PLoS ONE. 2022 Aug 4;17(8):e0269065.
- 27. Cohen JB, Hanff TC, William P, Sweitzer N, Rosado-Santander NR, Medina C, et al. Continuation versus discontinuation of renin-angiotensin system inhibitors in patients admitted to hospital with COVID-19: a prospective, randomised, open-label trial. Lancet Respir Med. 2021 Jan 7.
- 28. Lopes RD, Macedo AVS, de Barros E Silva PGM, Moll-Bernardes RJ, dos Santos TM, Mazza L, et al. Effect of Discontinuing vs Continuing Angiotensin-Converting Enzyme Inhibitors and Angiotensin II Receptor Blockers on Days Alive and Out of the Hospital in Patients Admitted With COVID-19: A Randomized Clinical Trial. JAMA. 2021 Jan 19;325(3):254.
- 29. Bauer A, Schreinlechner M, Sappler N, Dolejsi T, Tilg H, Aulinger BA, et al. Discontinuation versus continuation of renin-angiotensin-system inhibitors in COVID-19 (ACEI-COVID): a prospective, parallel group, randomised, controlled, open-label trial. The Lancet Respiratory Medicine. 2021 Jun;S2213260021002149.
- 30. Tornling G, Batta R, Porter JC, Williams B, Bengtsson T, Parmar K, et al. Seven days treatment with the angiotensin II type 2 receptor agonist C21 in hospitalized COVID-19 patients; a placebo-controlled randomised multi-centre double-blind phase 2 trial. EClinicalMedicine. 2021 Nov;41:101152.
- 31. Comparison of Losartan and Amlodipine Effects on the Outcomes of Patient with COVID-19 and Primary Hypertension: A Randomized Clinical Trial. International Journal of Clinical Practice [Internet]. 2021 Mar [cited 2021 Mar 4]; Available from: https://onlinelibrary.wiley.com/doi/10.1111/ijcp.14124
- 32. Puskarich M, Cummins NW, Ingraham N, Wacker DA, Reilkoff R, Driver BE, et al. Effect of Losartan on Symptomatic Outpatients with COVID-19: A Randomized Clinical Trial. SSRN Journal [Internet]. 2021 [cited 2021 Mar 24]; Available from: https://www.ssrn.com/abstract=378746



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- 33. Geriak M, Haddad F, Kullar R, Greenwood KL, Habib M, Habib C, et al. Randomized Prospective Open Label Study Shows No Impact on Clinical Outcome of Adding Losartan to Hospitalized COVID-19 Patients with Mild Hypoxemia. Infect Dis Ther [Internet]. 2021 May 11 [cited 2021 May 18]; Available from: https://link.springer.com/10.1007/s40121-021-00453-3
- 34. Duarte M, Pelorosso F, Nicolosi LN, Victoria Salgado M, Vetulli H, Aquieri A, et al. Telmisartan for treatment of Covid-19 patients: An open multicenter randomized clinical trial. EClinicalMedicine. 2021 Jul;37:100962.
- 35. Najmeddin F, Solhjoo M, Ashraf H, Salehi M, Rasooli F, Ghoghaei M, et al. Effects of Renin-Angiotensin-Aldosterone Inhibitors on Early Outcomes of Hypertensive COVID-19 Patients: A Randomized Triple-Blind Clinical Trial. American Journal of Hypertension. 2021 Jul 15;hpab111.
- 36. Puskarich MA, Ingraham NE, Merck LH, Driver BE, Wacker DA, Black LP, et al. Effect of losartan on hospitalized patients with COVID-19-induced lung injury: A randomized clinical trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Aug [cited 2021 Nov 24]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.08.25.21262623
- Freilich D, Victory J, Jenkins P, Gadomski A. COVIDMED An early pandemic randomized clinical trial of losartan treatment for hospitalized COVID-19 patients. Contemporary Clinical Trials Communications. 2022 Oct;29:100968.
- 38. Sharma A, Elharram M, Afilalo J, Flannery A, Afilalo M, Tselios C, et al. A Randomized Controlled Trial of Renin-Angiotensin-Aldosterone System Inhibitor Management in Patients Admitted in Hospital with COVID-19. American Heart Journal. 2022 Feb;S0002870322000242.
- 39. Bertoldi Lemos AC, do Espírito Santo DA, Salvetti MC, Gilio RN, Agra LB, Pazin-Filho A, Miranda CH. Therapeutic versus prophylactic anticoagulation for severe COVID-19: a randomized phase II clinical trial (HESACOVID). Thromb Res 2020;196:359-366. Available from: https://doi.org/10.1016/j.thromres.2020.09.026.
- 40. The REMAP-CAP, ACTIV-4a, and ATTACC Investigators. Therapeutic Anticoagulation with Heparin in Critically III Patients with Covid-19. N Engl J Med. 2021 Aug 4;NEJMoa2103417.
- 41. INSPIRATION Investigators, Sadeghipour P, Talasaz AH, Rashidi F, Sharif-Kashani B, Beigmohammadi MT, et al. Effect of Intermediate-Dose vs Standard-Dose Prophylactic

PAHOS



Anticoagulation on Thrombotic Events, Extracorporeal Membrane Oxygenation Treatment, or Mortality Among Patients With COVID-19 Admitted to the Intensive Care Unit: The INSPIRATION Randomized Clinical Trial. JAMA [Internet]. 2021 Mar 18 [cited 2021 Mar 22]; Available from: https://jamanetwork.com/journals/jama/fullarticle/2777829

42. Perepu U, Chambers I, Wahab A, Ten Eyck P, Wu C, Dayal S, et al. Standard Prophylactic Versus Intermediate Dose Enoxaparin in Adults with Severe COVID-19: A Multi-Center, Open-Label, Randomised Controlled Trial. SSRN Journal [Internet]. 2021 [cited 2021 May 18]; Available from: https://www.ssrn.com/abstract=3840099

- 43. The ATTACC, ACTIV-4a, and REMAP-CAP Investigators, Lawler PR, Goligher EC, Berger JS, Neal MD, McVerry BJ, et al. Therapeutic Anticoagulation in Non-Critically Ill Patients with Covid-19 [Internet]. Intensive Care and Critical Care Medicine; 2021 May [cited 2021 May 27]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.05.13.21256846
- 44. Lopes RD, de Barros e Silva PGM, Furtado RHM, Macedo AVS, Bronhara B, Damiani LP, et al. Therapeutic versus prophylactic anticoagulation for patients admitted to hospital with COVID-19 and elevated D-dimer concentration (ACTION): an open-label, multicentre, randomised, controlled trial. The Lancet. 2021 Jun;S0140673621012034.
- 45. Sholzberg M, Tang GH, Rahhal H, AlHamzah M, Kreuziger LB, Ainle FN, et al. Effectiveness of therapeutic heparin versus prophylactic heparin on death, mechanical ventilation, or intensive care unit admission in moderately ill patients with covid-19 admitted to hospital: RAPID randomised clinical trial. BMJ. 2021 Oct 14;n2400.
- 46. Spyropoulos AC, Goldin M, Giannis D, Diab W, Wang J, Khanijo S, et al. Efficacy and Safety of Therapeutic-Dose Heparin vs Standard Prophylactic or Intermediate-Dose Heparins for Thromboprophylaxis in High-risk Hospitalized Patients With COVID-19: The HEP-COVID Randomized Clinical Trial. JAMA Intern Med [Internet]. 2021 Oct 7 [cited 2021 Oct 15]; Available from:

https://jamanetwork.com/journals/jamainternalmedicine/fullarticle/2785004

PAHO®

 Marcos M, Carmona-Torre F, Vidal Laso R, Ruiz-Artacho P, Filella D, Carbonell C, et al. Therapeutic vs. prophylactic bemiparin in hospitalized patients with non-severe COVID-19 (BEMICOP): an open-label, multicenter, randomized trial. Thromb Haemost. 2021 Oct 12;a-1667-7534.



- 48. Oliynyk O, Barg W, Slifirczyk A, Oliynyk Y, Dubrov S, Gurianov V, et al. Comparison of the Effect of Unfractionated Heparin and Enoxaparin Sodium at Different Doses on the Course of COVID-19-Associated Coagulopathy. Life. 2021 Sep 30;11(10):1032.
- 49. Morici N, Podda G, Birocchi S, Bonacchini L, Merli M, Trezzi M, et al. Enoxaparin for thromboprophylaxis in hospitalized COVID-19 patients: The X-COVID-19 Randomized Trial. Eur J Clin Investigation [Internet]. 2021 Dec 26 [cited 2022 Jan 7]; Available from: https://onlinelibrary.wiley.com/doi/10.1111/eci.13735
- 50. Muñoz-Rivas N, Aibar J, Gabara-Xancó C, Trueba-Vicente Á, Urbelz-Pérez A, Gómez-Del Olmo V, et al. Optimal thromboprophylaxis strategies in non-critically ill patients with COVID-19 pneumonia. The PROTHROMCOVID Randomized Controlled Trial [Internet]. Cardiovascular Medicine; 2022 May [cited 2022 Jun 2]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.05.03.22274594</u>
- 51. Blondon M, Cereghetti S, Pugin J, Marti C, Darbellay Farhoumand P, Reny J, et al. Therapeutic anticoagulation to prevent thrombosis, coagulopathy, and mortality in severe COVID-19: The Swiss COVID-HEP randomized clinical trial. Res Pract Thromb Haemost [Internet]. 2022 May [cited 2022 Jun 2];6(4). Available from: <u>https://onlinelibrary.wiley.com/doi/10.1002/rth2.12712</u>
- 52. Rashidi F, Barco S, Rezaeifar P, Sadeghipour P, Ghodrati S, Bakhshandeh H, et al. Tissue plasminogen activator for the treatment of adults with critical COVID-19: A pilot randomized clinical trial. Thrombosis Research. 2022 Aug;216:125–8.
- 53. Kumar D, Kaimaparambil V, Chandralekha S, Lalchandani J. Oral Rivaroxaban in the Prophylaxis of COVID-19 Induced Coagulopathy. J Assoc Physicians India. 2022 Feb;70(2):11–2.
- 54. Connors JM, Brooks MM, Sciurba FC, Krishnan JA, Bledsoe JR, Kindzelski A, et al. Effect of Antithrombotic Therapy on Clinical Outcomes in Outpatients With Clinically Stable Symptomatic COVID-19: The ACTIV-4B Randomized Clinical Trial. JAMA. 2021 Oct 11;
- 55. Ananworanich J, Mogg R, Dunne MW, Bassyouni M, David CV, Gonzalez E, et al. Randomized study of rivaroxaban vs. placebo on disease progression and symptoms resolution in high-risk adults with mild COVID-19. Clinical Infectious Diseases. 2021 Sep 15;ciab813.



- 56. Barco S, Voci D, Held U, Sebastian T, Bingisser R, Colucci G, et al. Enoxaparin for primary thromboprophylaxis in symptomatic outpatients with COVID-19 (OVID): a randomised, open-label, parallel-group, multicentre, phase 3 trial. The Lancet Haematology. 2022 Jun;S2352302622001752.
- 57. Cools F, Virdone S, Sawhney J, Lopes RD, Jacobson B, Arcelus JI, et al. Thromboprophylactic low-molecular-weight heparin versus standard of care in unvaccinated, at-risk outpatients with COVID-19 (ETHIC): an open-label, multicentre, randomised, controlled, phase 3b trial. The Lancet Haematology. 2022 Aug;9(8):e594– 604.
- 58. Kumar DrGS, Vadgaonkar DrA, Purunaik DrS, Shelatkar R, Vaidya VG, Ganu DrG, et al. Efficacy and Safety of Aspirin, Promethazine, and Micronutrients for Rapid Clinical Recovery in Mild to Moderate COVID-19 Patients: A Randomized Controlled Clinical Trial. Cureus [Internet]. 2022 May 30 [cited 2022 Jul 18]; Available from: <u>https://www.cureus.com/articles/96829-efficacy-and-safety-of-aspirin-promethazine-andmicronutrients-for-rapid-clinical-recovery-in-mild-to-moderate-covid-19-patients-arandomized-controlled-clinical-trial</u>
- 59. Mehboob R, Ahmad F, Qayyum A, Rana MA, Tariq MA, Akram J. Aprepitant as a combinant with dexamethasone reduces the inflammation via neurokinin 1 receptor antagonism in severe to critical COVID-19 patients and potentiates respiratory recovery: a novel therapeutic approach [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.08.01.20166678.
- 60. Redondo-Calvo FJ, Padín JF, Muñoz-Rodríguez JR, Serrano-Oviedo L, López-Juárez P, Porras Leal ML, et al. Aprotinin treatment against SARS-CoV-2: A randomized phase III study to evaluate the safety and efficacy of a pan-protease inhibitor for moderate COVID-19. Eur J Clin Investigation [Internet]. 2022 Apr 5 [cited 2022 Apr 27]; Available from: <u>https://onlinelibrary.wiley.com/doi/10.1111/eci.13776</u>
- 61. Khodashahi R, Naderi H, Bojdy A, Heydari AA, Sani AT, Ghabouli MJ, et al. Comparison the Effect of Arbidol Plus Hydroxychloroquine vs Hydroxychloroquine Alone in Treatment of COVID-19 Disease: A Randomized Clinical Trial. CRMR. 2021 Mar 1;16(4):252–62.



- 62. Hellou E, Mohsin J, Elemy A, Hakim F, Mustafa-Hellou M, Hamoud S. Effect of ArtemiC in patients with COVID-19: A Phase II prospective study. J Cellular Molecular Medi. 2022 May 19;jcmm.17337.
- 63. Trieu V, Saund S, Rahate PV, Barge VB, Nalk KS, Windlass H, et al. Targeting TGF-β pathway with COVID-19 Drug Candidate ARTIVeda/PulmoHeal Accelerates Recovery from Mild-Moderate COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Feb [cited 2021 Feb 16]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.01.24.21250418
- 64. Ghati N, Bhatnagar S, Mahendran M, Thakur A, Prasad K, Kumar D, et al. Statin and aspirin as adjuvant therapy in hospitalised patients with SARS-CoV-2 infection: a randomised clinical trial (RESIST trial). BMC Infect Dis. 2022 Dec;22(1):606.
- 65. Abani O, Abbas A, Abbas F, Abbas M, Abbasi S, Abbass H, et al. Aspirin in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, openlabel, platform trial. The Lancet. 2021 Nov;S0140673621018250.
- 66. REMAP-CAP Writing Committee for the REMAP-CAP Investigators, Florescu S, Stanciu D, Zaharia M, Kosa A, Codreanu D, et al. Effect of Antiplatelet Therapy on Survival and Organ Support–Free Days in Critically Ill Patients With COVID-19: A Randomized Clinical Trial. JAMA [Internet]. 2022 Mar 22 [cited 2022 Apr 4]; Available from: <u>https://jamanetwork.com/journals/jama/fullarticle/2790488</u>
- Nekoukar Z, Ala S, Moradi S, Hill A, Davoudi Badabi AR, Alikhani A, et al. Comparison of the Efficacy and Safety of Atazanavir/Ritonavir Plus Hydroxychloroquine with Lopinavir/Ritonavir Plus Hydroxychloroquine in Patients with Moderate COVID-19, A Randomized, Double-blind Clinical Trial. Iran J Pharm Res. 2021;20(4):278–88.
- Jain MK, Lemos JA de, McGuire DK, Ayers C, Eiston JL, Sanchez CL, et al. Atovaquone for Treatment of COVID-19: A Prospective Randomized, Double-Blind, Placebo-Controlled Clinical Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 May [cited 2022 Jun 2]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.05.24.22275411
- 69. Bruen C, Al-Saadi M, Michelson E, Tanios M, Mendoza-Ayala R, Miller J, et al. Auxora Improves Outcomes in Patients With Severe COVID-19 Pneumonia: A Randomized Clinical Trial. SSRN Journal [Internet]. 2021 [cited 2021 Dec 20]; Available from: https://www.ssrn.com/abstract=3976177



- 70. Carvelli J, Meziani F, Dellamonica J, Cordier P-Y, Allardet-Servent J, Fraisse M, et al. Avdoralimab (anti-C5aR1 mAb) Versus Placebo in Patients With Severe COVID-19: Results From a Randomized Controlled Trial (FORCE). SSRN Journal [Internet]. 2022
 [cited 2022 Feb 21]; Available from: <u>https://www.ssrn.com/abstract=4028533</u>
- 71. Youssef JG, Lee R, Javitt J, Lavin P, Jayaweera D. Effectiveness of ZYESAMITM (Aviptadil) in Accelerating Recovery and Shortening Hospitalization in Critically-III Patients with COVID-19 Respiratory Failure: Interim Report from a Phase 2B/3 Multicenter Trial. SSRN Journal [Internet]. 2021 [cited 2021 Apr 8]; Available from: https://www.ssrn.com/abstract=3794262
- 72. Singh H, Srivastava S, Yadav B, Rai AK, Jameela S, Muralidharan S, et al. AYUSH-64 as an adjunct to standard care in mild to moderate COVID-19: An open-label randomized controlled trial in Chandigarh, India. Complementary Therapies in Medicine. 2022 Jun;66:102814.
- 73. Chorlton J, Hollowood Z, Dyer C, Lockhart D, Boekman P, McCafferty K, et al. A randomised, double-blind, placebo-controlled, multicentre clinical trial of AZD1656 in diabetic patients hospitalised with COVID-19: The ARCADIA Trial - implications for therapeutic immune modulation. eClinicalMedicine. 2022 Sep;51:101604.
- 74. Klussmann JP, Lehmann C, Grosheva M, Sahin K, Nagy E, Szijártó V, et al. COVID-19: Azelastine nasal spray Reduces Virus-load In Nasal swabs (CARVIN). Early intervention with azelastine nasal sprays reduces viral load in SARS-CoV-2 infected patients. First report on a double-blind placebo-controlled phase II clinical trial. [Internet]. In Review; 2021 Sep [cited 2021 Sep 21]. Available from:

https://www.researchsquare.com/article/rs-864566/v1

PAHO®

- 75. Sekhavati E, Jafari F, SeyedAlinaghi S, Jamali Moghadam Siahkali S, Sadr S, Tabarestani M, et al. Safety and effectiveness of azithromycin in patients with COVID-19: an open-label randomized trial. Int Journal Antimicrob Ag 2020;56(4):106143. Available from: https://doi.org/10.1016/j.ijantimicag.2020.106143.
- 76. Guvenmez O, Keskin H, Ay B, Birinci S, Kanca MF. The comparison of the effectiveness of lincocin® and azitro® in the treatment of COVID-19-associated pneumonia: a prospective study. J Popul Ther Clin Pharmacol 2020;27(S Pt1):e5–10. Available from : https://doi.org/10.15586/jptcp.v27iSP1.684.



- 77. Furtado RHM, Berwanger O, Fonseca HA, Corrêa TD, Ferraz LR, Lapa MG, et al. Azithromycin in addition to standard of care versus standard of care alone in the treatment of patients admitted to the hospital with severe COVID-19 in Brazil (COALITION II): a randomised clinical trial. Lancet 2020;396:959-67. Available from: https://doi.org/10.1016/S0140-6736(20)31862-6.
- 78. Horby PW, Roddick A, Spata E, Staplin N, Emberson JR, Pessoa-Amorim G, Peto L, et al. 2020. Azithromycin in Hospitalised Patients with COVID-19 (RECOVERY): A Randomised, Controlled, Open-Label, Platform Trial. Preprint. Infectious Diseases (except HIV/AIDS). https://doi.org/10.1101/2020.12.10.20245944.
- 79. Rashad A, Nafady A, Hassan M, Mansour H, Taya U, Bazeed S, et al. Therapeutic efficacy of macrolides in management of patients with mild COVID-19. ResearchSquare [Internet]. 2021
- 80. Butler CC, Dorward J, Yu L-M, Gbinigie O, Hayward G, Saville BR, et al. Azithromycin for community treatment of suspected COVID-19 in people at increased risk of an adverse clinical course in the UK (PRINCIPLE): a randomised, controlled, open-label, adaptive platform trial. The Lancet. 2021 Mar;S014067362100461X.
- 81. Hinks TS, Cureton L, Knight R, Wang A, Cane JL, Barber VS, et al. A randomised clinical trial of azithromycin versus standard care in ambulatory COVID-19 – the ATOMIC2 trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Apr [cited 2021 May 3]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.04.21.21255807
- 82. Oldenburg CE, Pinsky BA, Brogdon J, Chen C, Ruder K, Zhong L, et al. Effect of Oral Azithromycin vs Placebo on COVID-19 Symptoms in Outpatients With SARS-CoV-2 Infection: A Randomized Clinical Trial. JAMA [Internet]. 2021 Jul 16 [cited 2021 Aug 2]; Available from: https://jamanetwork.com/journals/jama/fullarticle/2782166
- 83. Ghanei M, Solaymani-Dodaran M, Qazvini A, Ghazale AH, Setarehdan SA, Saadat SH, et al. The efficacy of corticosteroids therapy in patients with moderate to severe SARS-CoV-2 infection: a multicenter, randomized, open-label trial. Respir Res. 2021 Dec;22(1):245.
- 84. Gyselinck I, Liesenborghs L, Belmans A, Engelen MM, Betrains A, Van Thillo Q, et al. Azithromycin for treatment of hospitalised COVID-19 patients: a randomised, multicentre, open-label clinical trial (DAWn-AZITHRO). ERJ Open Res. 2022 Jan;8(1):00610–2021.

PAHOS





- 85. Ren Z, Luo H, Yu Z, Song J, Liang L, Wang L, et al. A randomized, open-label, controlled clinical trial of azvudine tablets in the treatment of mild and common COVID-19, a pilot study. Adv Sci 2020;7:2001435. Available from: https://doi.org/10.1002/advs.202001435.
- 86. Lou Y, Liu L, Qiu Y. Clinical outcomes and plasma concentrations of baloxavir marboxil and favipiravir in COVID-19 patients: an exploratory randomized, controlled trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.04.29.20085761.
- 87. Chen P, Nirula A, Heller B, Gottlieb RL, Boscia J, Morris J, et al. SARS-CoV-2 neutralizing antibody LY-CoV555 in outpatients with COVID-19. N Engl J Med 2020; NEJMoa2029849. Available from: https://doi.org/10.1056/NEJMoa2029849.
- ACTIV-3/TICO LY-CoV555 Study Group. A Neutralizing Monoclonal Antibody for Hospitalized Patients with Covid-19. N Engl J Med. 2020 Dec 22;NEJMoa2033130.
- 89. Gottlieb RL, Nirula A, Chen P, Boscia J, Heller B, Morris J, et al. Effect of Bamlanivimab as Monotherapy or in Combination With Etesevimab on Viral Load in Patients With Mild to Moderate COVID-19: A Randomized Clinical Trial. JAMA [Internet]. 2021
- 90. Cohen MS, Nirula A, Mulligan MJ, Novak RM, Marovich M, Yen C, et al. Effect of Bamlanivimab vs Placebo on Incidence of COVID-19 Among Residents and Staff of Skilled Nursing and Assisted Living Facilities: A Randomized Clinical Trial. JAMA [Internet]. 2021 Jun 3 [cited 2021 Jun 15]; Available from: https://jamanetwork.com/journals/jama/fullarticle/2780870
- 91. Dougan M, Nirula A, Azizad M, Mocherla B, Gottlieb RL, Chen P, et al. Bamlanivimab plus Etesevimab in Mild or Moderate Covid-19. N Engl J Med. 2021 Jul 14;NEJMoa2102685.
- 92. Chen P, Datta G, Li YG, Chien J, Price K, Chigutsa E, et al. First in Human Study of Bamlanivimab in a Randomized Trial of Hospitalized Patients with COVID-19. Clinical Pharmacology & Therapeutics. 2021 Aug 28;cpt.2405.
- 93. Huang DT, McCreary EK, Bariola JR, Minnier TE, Wadas RJ, Shovel JA, et al. Effectiveness of Casirivimab-Imdevimab and Sotrovimab During a SARS-CoV-2 Delta Variant Surge: A Cohort Study and Randomized Comparative Effectiveness Trial. JAMA Netw Open. 2022 Jul 14;5(7):e2220957.



- 94. nasopharyngeal SARS-CoV-2 RNA levels but not symptom duration in non-hospitalized adults with COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Dec [cited 2021 Dec 30]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.12.17.21268009
- 95. Huang DT, McCreary EK, Bariola JR, Minnier TE, Wadas RJ, Shovel JA, et al. Effectiveness of casirivimab and imdevimab, and sotrovimab during Delta variant surge: a prospective cohort study and comparative effectiveness randomized trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Dec [cited 2022 Jan 10]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.12.23.21268244
- 96. Mazzaferri F, Mirandola M, Savoldi A, De Nardo P, Morra M, Tebon M, et al. Exploratory data on the clinical efficacy of monoclonal antibodies against SARS-CoV-2 Omicron Variant of Concern [Internet]. Infectious Diseases (except HIV/AIDS); 2022 May [cited 2022 Jun 2]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2022.05.06.22274613

- 97. Dougan M, Azizad M, Chen P, Feldman B, Frieman M, Igbinadolor A, et al. Bebtelovimab, alone or together with bamlanivimab and etesevimab, as a broadly neutralizing monoclonal antibody treatment for mild to moderate, ambulatory COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Mar [cited 2022 Jul 25]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.03.10.22272100</u>
- 98. Kalil AC., Patterson TF, Mehta AK, Tomashek KM, Wolfe CR, Ghazaryan V, Marconi VC, et al. 2020. Baricitinib plus Remdesivir for Hospitalized Adults with Covid-19. New England Journal of Medicine, December, NEJMoa2031994. https://doi.org/10.1056/NEJMoa2031994.
- 99. Marconi VC, Ramanan AV, de Bono S, Kartman CE, Krishnan V, Liao R, 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. The Lancet Respiratory Medicine. 2021 Sep;S2213260021003313.
- 100. Ely EW, Ramanan AV, Kartman CE, de Bono S, Liao R, Piruzeli MLB, et al. Baricitinib plus Standard of Care for Hospitalised Adults with COVID-19 on Invasive Mechanical Ventilation or Extracorporeal Membrane Oxygenation: Results of a Randomised, Placebo-Controlled Trial [Internet]. Infectious Diseases (except



HIV/AIDS); 2021 Oct [cited 2021 Oct 18]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.10.11.21263897

- 101. Abani O, Abbas A, Abbas F, Abbas J, Abbas K, Abbas M, et al. Baricitinib in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial and updated meta-analysis. The Lancet. 2022 Jul;400(10349):359–68.
- 102. Wolfe CR, Tomashek KM, Patterson TF, Gomez CA, Marconi VC, Jain MK, et al. Baricitinib versus dexamethasone for adults hospitalised with COVID-19 (ACTT-4): a randomised, double-blind, double placebo-controlled trial. The Lancet Respiratory Medicine. 2022 May;S2213260022000881.
- 103. Karampitsakos T, Papaioannou O, Tsiri P, Katsaras M, Katsimpris A, Kalogeropoulos AP, et al. Tocilizumab versus baricitinib in hospitalized patients with severe COVID-19: an open label, randomized controlled trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jun [cited 2022 Jul 6]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.06.13.22276211</u>
- 104. Montejano R, de la Calle-Prieto F, Velasco M, Guijarro C, Queiruga-Parada J, Jiménez-González M, et al. Tenofovir Disoproxil Fumarate/Emtricitabine and Baricitinib for Patients at High Risk of Severe COVID-19: The PANCOVID Randomized Clinical Trial. Clinical Infectious Diseases. 2022 Jul 30;ciac628.
- 105. Padmanabhan U, Mukherjee S, Borse R, Joshi S, Deshmukh R. Phase II clinical trial for evaluation of BCG as potential therapy for COVID-19 [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.10.28.20221630.
- 106. Raghavan K, Dedeepiya VD, Suryaprakash V, Rao K-S, Ikewaki N, Sonoda T, et al. Beneficial effects of novel aureobasidium pullulans strains produced beta-1,3-1,6 glucans on interleukin-6 and D-dimer levels in COVID-19 patients; results of a randomized multiple-arm pilot clinical study. Biomedicine & Pharmacotherapy. 2021 Sep;112243.
- 107. Pushkala S, Seshayyan S, Theranirajan E, Sudhakar D, Raghavan K, Dedeepiya VD, et al. Efficient control of IL-6, CRP and Ferritin in Covid-19 patients with two variants of Beta-1,3-1,6 glucans in combination, within 15 days in an open-label prospective clinical trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Dec

PAHO®





[cited 2021 Dec 30]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2021.12.14.21267778

- 108. Delić N, Matetic A, Domjanović J, Kljaković-Gašpić T, Šarić L, Ilić D, et al. Effects of Different Inhalation Therapy on Ventilator-Associated Pneumonia in Ventilated COVID-19 Patients: A Randomized Controlled Trial. Microorganisms. 2022 May 28;10(6):1118.
- 109. Rybakov A.R., Zhebelenko Y.G., Dubrov S.O., Vdovenko D.V., Kavardakova N.V., Matsibokh S.V., et al. The Results of the Clinical Study: An Open-label Multicenter Randomized Trial to Evaluate the Efficacy of Bioven, Manufactured by Biopharma Plasma, LLC, in Complex Therapy of Patients with Pneumonia Induced by COVID-19/SARS-COV-2 / PE3YJbTATU KJIHIЧHOFO ДОСЛІДЖЕННЯ «ВІДКРИТЕ БАГАТОЦЕНТРОВЕ РАНДОМІЗОВАНЕ ДОСЛІДЖЕННЯ З ОЦІНКИ ЕФЕКТИВНОСТІ ПРЕПАРАТУ БІОВЕН, ВИРОБНИЦТВА ТОВ «БІОФАРМА ПЛАЗМА», В КОМПЛЕКСНІЙ ТЕРАПІЇ ПАЦІЄНТІВ З ПНЕВМОНІЄЮ, ЩО ВИКЛИКАНА КОРОНАВІРУСНОЮ ІНФЕКЦІЄЮ COVID-19. Pain, Anaesthesia and Intensive Care. 2020;4(93):9–21.
- 110. Barzin Tond S, Balenci L, Khajavirad N, Salehi M, Tafakhori A, Shahmohammadi MR, et al. Inflawell® improves neutrophil-to-lymphocyte ratio and shortens hospitalization in patients with moderate COVID-19, in a randomized doubleblind placebo-controlled clinical trial. Inflammopharmacol [Internet]. 2022 Feb 24 [cited 2022 Mar 11]; Available from: <u>https://link.springer.com/10.1007/s10787-022-00928-w</u>
- 111. Li T, Sun L, Zhang W, Zheng C, Jiang C, Chen M, et al. Bromhexine hydrochloride tablets for the treatment of moderate COVID-19: an open-label randomized controlled pilot study. Clin Transl Sci 2020;13(6):1096-1102. Available from: https://doi.org/10.1111/cts.12881.
- 112. Ansarin K, Tolouian R, Ardalan M, Taghizadieh A, Varshochi M, Teimouri S, et al. 2020. Effect of bromhexine on clinical outcomes and mortality in COVID-19 patients: a randomized clinical trial. Bioimpacts 2020;10(4):209–15. Available from: https://doi.org/10.34172/bi.2020.27.
- 113. Mikhaylov EN, Lyubimtseva TA, Vakhrushev AD, Stepanov D, Lebedev DS, Vasilieva EYu, et al. Bromhexine Hydrochloride Prophylaxis of COVID-19 for Medical



Personnel: A Randomized Open-Label Study. Tharmalingam J, editor. Interdisciplinary Perspectives on Infectious Diseases. 2022 Jan 29;2022:1–7.

- 114. Tolouian R, Mulla ZD, Jamaati H, Babamahmoodi A, Marjani M, Eskandari R, et al. Effect of bromhexine in hospitalized patients with COVID-19. J Investig Med. 2021 Mar 15; jim-2020-001747.
- 115. Tolouian R, Moradi O, Mulla ZD, Ziaie S, Haghighi M, Esmaily H, et al. Bromhexine, for Post Exposure COVID-19 Prophylaxis: A Randomized, Double-Blind, Placebo Control Trial. SSRN Journal [Internet]. 2021 [cited 2022 Jan 11]; Available from: https://www.ssrn.com/abstract=3989849
- 116. Elamir YM, Amir H, Lim S, Rana YP, Lopez CG, Feliciano NV, et al. A randomized pilot study using calcitriol in hospitalized COVID-19 patients. Bone. 2022 Jan;154:116175.
- 117. Gunst JD, Staerke NB, Pahus MH, Kristensen LH, Bodilsen J, Lohse N, et al. Efficacy of the TMPRSS2 inhibitor camostat mesilate in patients hospitalized with Covid-19-a double-blind randomized controlled trial. EClinicalMedicine. 2021 Apr;100849.
- 118. Chupp G, Spichler-Moffarah A, Søgaard OS, Esserman D, Dziura J, Danzig L, et al. A Phase 2 Randomized, Double-Blind, Placebo-controlled Trial of Oral Camostat Mesylate for Early Treatment of COVID-19 Outpatients Showed Shorter Illness Course and Attenuation of Loss of Smell and Taste [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jan [cited 2022 Feb 16]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.01.28.22270035
- 119. Kinoshita T., Masahiro Shinoda, Yasuhiro Nisizaki, Katsuya Shiraki, Yuji Hirai, Yoshiko Kichikawa, et al. Phase 3, multicentre, double-blind, randomised, parallel-group, placebo-controlled study of camostat mesilate (FOY-305) for the treatment of COVID-19 (CANDLE study). medRxiv [Internet]. 2022; Available from: http://www.epistemonikos.org/documents/17575b0440c65ac971614982e92008e43db429 7f
- 120. Terada J, Fujita R, Kawahara T, Hirasawa Y, Kinoshita T, Takeshita Y, et al. Favipiravir, camostat, and ciclesonide combination therapy in patients with moderate COVID-19 pneumonia with/without oxygen therapy: An open-label, single-center phase 3 randomized clinical trial. eClinicalMedicine. 2022 Jul;49:101484.

PAHO®

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- 121. Tobback E, Degroote S, Buysse S, Delesie L, Van Dooren L, Vanherrewege S, et al. Efficacy and safety of camostat mesylate in early Covid-19 disease in an ambulatory setting: A randomized placebo-controlled phase II trial. International Journal of Infectious Diseases. 2022 Jul;S1201971222003885.
- 122. Caricchio R, Abbate A, Gordeev I, Meng J, Hsue PY, Neogi T, et al. Effect of Canakinumab vs Placebo on Survival Without Invasive Mechanical Ventilation in Patients Hospitalized With Severe COVID-19: A Randomized Clinical Trial. JAMA. 2021 Jul 20;326(3):230–9.
- 123. Cremer PC, Sheng CC, Sahoo D, Dugar S, Prada RA, Wang TKM, et al. Double-Blind Randomised Proof-of-Concept Trial of C anakinumab in Patients with C OVID-19 Associated C ardiac Injury and Heightened Inflammation. European Heart Journal Open. 2021 Jul 29;0eab002.
- 124. Crippa JAS, Pacheco JC, Zuardi AW, Guimarães FS, Campos AC, Osório F de L, et al. Cannabidiol for COVID-19 Patients with Mild to Moderate Symptoms (CANDIDATE Study): A Randomized, Double-Blind, Placebo-Controlled Clinical Trial. Cannabis and Cannabinoid Research. 2021 Oct 7;can.2021.0093.
- 125. Welker J, Pulido JD, Catanzaro AT, Malvestutto CD, Li Z, Cohen JB, et al. Efficacy and safety of CD24Fc in hospitalised patients with COVID-19: a randomised, double-blind, placebo-controlled, phase 3 study. The Lancet Infectious Diseases. 2022 Mar;S1473309922000585.
- 126. Perlin DS, Neil GA, Anderson C, Zafir-Lavie I, Roadcap L, Raines S, et al. CERC-002, a human anti-LIGHT mAb reduces respiratory failure and death in hospitalized COVID-19 ARDS patients [Internet]. Pharmacology and Therapeutics; 2021 Apr [cited 2021 Apr 12]. Available from: http://www.dwiw.eu/dwiwi10.1101/2021.04.02.21254748

http://medrxiv.org/lookup/doi/10.1101/2021.04.03.21254748

PAHO®

- 127. Thakar A, Panda S, Sakthivel P, Brijwal M, Dhakad S, Choudekar A, et al. Chloroquine nasal drops in asymptomatic & mild COVID-19: An exploratory randomized clinical trial. Indian J Med Res. 2021;0(0):0.
- 128. Cruz LR, Baladron I, Rittoles A, Diaz PA, Valenzuela C, Santana R, et al. Treatment with an anti-CK2 synthetic peptide improves clinical response in COVID-19 patients with pneumonia: a randomized and controlled clinical trial [Preprint]. MedRxiv 2020. Available from: <u>https://doi.org/10.1101/2020.09.03.20187112</u>.



- 129. Lonze BE, Spiegler P, Wesson RN, Alachkar N, Petkova E, Weldon EP, et al. A Randomized Double-Blinded Placebo Controlled Trial of Clazakizumab for the Treatment of COVID-19 Pneumonia With Hyperinflammation. Critical Care Medicine [Internet]. 2022 May 18 [cited 2022 Jun 7];Publish Ahead of Print. Available from: https://journals.lww.com/10.1097/CCM.00000000005591
- 130. Song J-Y, Kim Y-S, Eom J-S, Kim J-Y, Lee J-S, Lee J, et al. Oral antiviral clevudine compared with placebo in Korean COVID-19 patients with moderate severity [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Dec [cited 2021 Dec 29]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.12.09.21267566
- 131. Altay O, Yang H, Aydin M, Alkurt G, Altunal N, Kim W, et al. Combined metabolic cofactor supplementation accelerates recovery in mild-to-moderate COVID-19 [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.10.02.20202614.
- 132. Altay O, Arif M, Li X, Yang H, Aydın M, Alkurt G, et al. Combined Metabolic Activators Accelerates Recovery in Mild-to-Moderate COVID-19. Adv Sci. 2021 Sep;8(17):2101222.
- 133. Hu Q, Zhang Q-Y, Peng C-F, Ma Z, Han Y-L. Efficiency of Nicotinamide-based Supportive Therapy in Lymphopenia for Patients With COVID-19: A Randomized Controlled Trial [Internet]. In Review; 2022 Jan [cited 2022 Jan 18]. Available from: <u>https://www.researchsquare.com/article/rs-1173313/v1</u>
- 134. Deftereos SG, Giannopoulos G, Vrachatis DA, Siasos GD, Giotaki SG,
 Gargalianos P, 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.
 Available from: https://doi.org/10.1001/jamanetworkopen.2020.13136.
- 135. Lopes MI, Bonjorno LP, Giannini MC, Amaral NB, Menezes PI, Dib SM, et al. Beneficial effects of colchicine for moderate to severe COVID-19: a randomised, doubleblinded, placebo-controlled clinical trial. RMD Open. 2021 Feb;7(1):e001455.
- 136. Farhad S, Pourfarzi F, Ataei S. The impact of colchicine on the COVID-19 patients: a clinical trial study [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-69374/v1.

PAHO®

137. Tardif J-C, Bouabdallaoui N, L'Allier PL, Gaudet D, Shah B, Pillinger MH, et al. Colchicine for community-treated patients with COVID-19 (COLCORONA): a phase 3,



randomised, double-blinded, adaptive, placebo-controlled, multicentre trial. The Lancet Respiratory Medicine. 2021 May;S2213260021002228.

- Colchicine in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. The Lancet Respiratory Medicine. 2021 Oct;S2213260021004355.
- 139. Pascual-Figal DA, Roura-Piloto AE, Moral-Escudero E, Bernal E, Albendin-Iglesias H, Pérez-Martínez MT, et al. Colchicine in Recently Hospitalized Patients with COVID-19: A Randomized Controlled Trial (COL-COVID). IJGM. 2021 Sep;Volume 14:5517–26.
- 140. Dorward J, Yu LM, Hayward G, Saville BR, Gbinigie O, Van Hecke O, et al. Colchicine for COVID-19 in the community (PRINCIPLE): a randomised, controlled, adaptive platform trial. Br J Gen Pract. 2022 Mar 23;BJGP.2022.0083.
- 141. Diaz R, Orlandini A, Castellana N, Caccavo A, Corral P, Corral G, 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 Dec 29;4(12):e2141328.
- 142. Alsultan M, Obeid A, Alsamarrai O, Anan MT, Bakr A, Soliman N, et al. Efficacy of Colchicine and Budesonide in Improvement Outcomes of Patients with Coronavirus Infection 2019 in Damascus, Syria: A Randomized Control Trial. Lanzafame M, editor. Interdisciplinary Perspectives on Infectious Diseases. 2021 Dec 31;2021:1–7.
- 143. Pourdowlat G, Saghafi F, Mozafari A, Sahebnasagh A, Abedini A, Nabi Meybodi M, et al. Efficacy and safety of colchicine treatment in patients with COVID -19: A prospective, multicenter, randomized clinical trial. Phytotherapy Research. 2022 Feb 2;ptr.7319.
- 144. 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. Annals of Medicine and Surgery. 2022 Apr;103593.
- 145. Pimenta Bonifácio L, Ramacciotti E, Agati LB, Vilar FC, Tojal da Silva AC, Louzada-Junior P, et al. Efficacy and Safety of Ixekizumab vs. Low-Dose IL-2 vs. Colchicine vs. Standard of Care on the Treatment of Patients Hospitalized with Moderate



to Critical Covid-19: A Pilot Randomized Clinical Trial (STRUCK: Survival Trial Using Cyto kine Inhibitors). SSRN Journal [Internet]. 2022 [cited 2022 Jun 8]; Available from: https://www.ssrn.com/abstract=4095747

- 146. Cecconi A, Martinez-Vives P, Vera A, Lavilla Olleros C, Barrios A, Fonseca Aizpuru E, et al. Efficacy of short-course colchicine treatment in hospitalized patients with moderate to severe COVID-19 pneumonia and hyperinflammation: a randomized clinical trial. Sci Rep. 2022 Dec;12(1):9208.
- 147. Rabbani A, Rafique A, Wang X, Campbell D, Wang D, Brownell N, et al. Colchicine for the Treatment of Cardiac Injury in Hospitalized Patients With Coronavirus Disease-19. Front Cardiovasc Med. 2022 Jun 17;9:876718.
- 148. Gaitán-Duarte HG, Álvarez-Moreno C, Rincón-Rodríguez CJ, Yomayusa-González N, Cortés JA, Villar JC, et al. Effectiveness of Rosuvastatin plus Colchicine, Emtricitabine/Tenofovir and a combination of them in Hospitalized Patients with SARS Covid-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Jul [cited 2021 Aug 2]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.07.06.21260085
- 149. Li L, Zhang W, Hu Y, Tong X, Zheng S, Yang J, et al. Effect of convalescent plasma therapy on time to clinical improvement in patients with severe and lifethreatening COVID-19: a randomized clinical trial. JAMA 2020;324(5):460-70. Available from: https://doi.org/10.1001/jama.2020.10044.
- 150. Gharbharan A, Jordans CCE, GeurtsvanKessel C, den Hollander JG, Karim F, Mollema PN, et al. Convalescent plasma for COVID-19: a randomized clinical trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.07.01.20139857.
- 151. Avendano-Sola C, Ramos-Martinez A, Munez-Rubio E, Ruiz-Antoran B, de Molina RM, Torres F, et al. Convalescent plasma for COVID-19: a multicenter, randomized clinical trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.08.26.20182444.
- 152. Agarwal A, Mukherjee A, Kumar G, Chatterjee P, Bhatnagar T, Malhotra P, et al. Convalescent plasma in the management of moderate COVID-19 in India: an open-label parallel-arm phase II multicentre randomized controlled trial (PLACID Trial) [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.09.03.20187252.
- 153. Simonovich VA, Burgos Pratx LD, Scibona P, Beruto MV, Vallone MG, VázquezC, et al. A randomized trial of convalescent plasma in COVID-19 severe pneumonia. N

PAHO®





Engl J Med 2020; NEJMoa2031304. Available from: https://doi.org/10.1056/NEJMoa2031304.

- 154. Bajpai M, Kumar S, Maheshwari A, Chabra K, Kale P, Gupta A, et al. Efficacy of convalescent plasma therapy compared to fresh frozen plasma in severely ill COVID-19 patients: a pilot randomized controlled trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.10.25.20219337.
- 155. AlQahtani M, Abdulrahman A, AlMadani A, Yousif AlAli S, Al Zamrooni AM, Hejab A, et al. Randomized controlled trial of convalescent plasma therapy against standard therapy in patients with severe COVID-19 disease [Preprint]. 2020 MedRxiv 2020. Available from: https://doi.org/10.1101/2020.11.02.20224303.
- 156. Libster R, Pérez Marc G, Wappner D, Coviello S, Bianchi A, Braem V, et al. Early High-Titer Plasma Therapy to Prevent Severe Covid-19 in Older Adults. N Engl J Med. 2021 Jan 6;NEJMoa2033700.
- 157. Ray, Yogiraj, Shekhar Ranjan Paul, Purbita Bandopadhyay, Ranit D'Rozario, Jafar Sarif, Deblina Raychaudhuri, Debaleena Bhowmik, et al. 2022. "A Phase 2 Single Center Open Label Randomised Control Trial for Convalescent Plasma Therapy in Patients with Severe COVID-19." *Nature Communications* 13 (1): 383. <u>https://doi.org/10.1038/s41467-022-28064-7</u>.
- 158. Horby PW, Estcourt L, Peto L, Emberson JR, Staplin N, Spata E, et al. Convalescent plasma in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Mar [cited 2021 Mar 11]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.03.09.21252736
- 159. Baklaushev V, Averyanov AV, Sotnikova AG, Perkina AS, Ivanov A, Yusubalieva GM, et al. Safety and Efficacy of Convalescent Plasma for COVID-19: The First Results of a Clinical Study. Journal of Clinical Practice [Internet]. 2020 Jul 17 [cited 2021 Feb 14]; Available from: https://journals.ecovector.com/clinpractice/article/view/35168
- 160. O'Donnell MR, Grinsztejn B, Cummings MJ, Justman JE, Lamb MR, Eckhardt CM, et al. A randomized double-blind controlled trial of convalescent plasma in adults with severe COVID-19. Journal of Clinical Investigation [Internet]. 2021 May 11 [cited 2021 May 17]; Available from: http://www.jci.org/articles/view/150646

PAHO®

161. Gonzalez JLB, González Gámez M, Mendoza Enciso EA, Esparza Maldonado RJ, Palacios DH, Campos SD, et al. Efficacy and safety of convalescent plasma and intravenous immunoglobulin in critically ill COVID-19 patients. A controlled clinical trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Mar [cited 2021 Apr 5]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.03.28.21254507

Pouladzadeh M, Safdarian M, Eshghi P, Abolghasemi H, Bavani AG, Sheibani B, et al. A randomized clinical trial evaluating the immunomodulatory effect of convalescent plasma on COVID-19-related cytokine storm. Internal and emergency medicine [Internet]. 2021; Available from: http://www.epistemonikos.org/documents/1996674ceda1dbb24d8246a2f7b3b4f65135369 3

- 163. Bennett-Guerrero E, Romeiser JL, Talbot LR, Ahmed T, Mamone LJ, Singh SM, et al. Severe Acute Respiratory Syndrome Coronavirus 2 Convalescent Plasma Versus Standard Plasma in Coronavirus Disease 2019 Infected Hospitalized Patients in New York: A Double-Blind Randomized Trial. Critical Care Medicine [Internet]. 2021 Apr 16 [cited 2021 Apr 27];Publish Ahead of Print. Available from: https://journals.lww.com/10.1097/CCM.00000000005066
- 164. Hamdy Salman O, Ail Mohamed HS. Efficacy and safety of transfusing plasma from COVID-19 survivors to COVID-19 victims with severe illness. A double-blinded controlled preliminary study. Egyptian Journal of Anaesthesia. 2020 Jan 1;36(1):264–72.
- 165. Körper S, Weiss M, Zickler D, Wiesmann T, Zacharowski K, M.Corman V, et al. High Dose Convalescent Plasma in COVID-19: Results from the Randomized Trial CAPSID [Internet]. Infectious Diseases (except HIV/AIDS); 2021 May [cited 2021 May 20]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.05.10.21256192
- 166. Writing Committee for the REMAP-CAP Investigators, Estcourt LJ, Turgeon AF, McQuilten ZK, McVerry BJ, Al-Beidh F, et al. Effect of Convalescent Plasma on Organ Support-Free Days in Critically Ill Patients With COVID-19: A Randomized Clinical Trial. JAMA. 2021 Oct 4;
- 167. The CONCOR-1 Study Group, CONCOR-1 writing committee, Bégin P, Callum J, Jamula E, Cook R, et al. Convalescent plasma for hospitalized patients with COVID-19 and the effect of plasma antibodies: a randomized controlled, open-label trial [Internet].



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Infectious Diseases (except HIV/AIDS); 2021 Jul [cited 2021 Jul 6]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.06.29.21259427

- 168. Sekine L, Arns B, Fabro BR, Cipolatt MM, Machado RRG, Durigon EL, et al. Convalescent plasma for COVID-19 in hospitalised patients: an open-label, randomised clinical trial. Eur Respir J. 2021 Jul 8;2101471.
- 169. Kirenga B, Byakika-Kibwika P, Muttamba W, Kayongo A, Loryndah NO, Mugenyi L, et al. Efficacy of convalescent plasma for treatment of COVID-19 in Uganda. BMJ Open Resp Res. 2021 Aug;8(1):e001017.
- Korley FK, Durkalski-Mauldin V, Yeatts SD, Schulman K, Davenport RD,
 Dumont LJ, et al. Early Convalescent Plasma for High-Risk Outpatients with Covid-19.
 N Engl J Med. 2021 Aug 18;NEJMoa2103784.
- 171. Devos T, Van Thillo Q, Compernolle V, Najdovski T, Romano M, Dauby N, et al. Early high antibody-titre convalescent plasma for hospitalised COVID-19 patients: DAWn-plasma. Eur Respir J. 2021 Aug 26;2101724.
- 172. Bar KJ, Shaw PA, Choi GH, Aqui N, Fesnak A, Yang JB, et al. A randomized controlled study of convalescent plasma for individuals hospitalized with COVID-19 pneumonia. Journal of Clinical Investigation [Internet]. 2021 Nov 17 [cited 2021 Dec 13]; Available from: http://www.jci.org/articles/view/155114
- 173. Menichetti F, Popoli P, Puopolo M, Spila Alegiani S, Tiseo G, Bartoloni A, et al. Effect of High-Titer Convalescent Plasma on Progression to Severe Respiratory Failure or Death in Hospitalized Patients With COVID-19 Pneumonia: A Randomized Clinical Trial. JAMA Netw Open. 2021 Nov 29;4(11):e2136246.
- Millat-Martinez P, Gharbharan A, Alemany A, Rokx C, Geurtsvankessel C, Papageourgiou G, et al. Convalescent plasma for outpatients with early COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Dec [cited 2021 Dec 8]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.11.30.21266810
- 175. Sullivan DJ, Gebo KA, Shoham S, Bloch EM, Lau B, Shenoy AG, et al. Early Outpatient Treatment for Covid-19 with Convalescent Plasma. N Engl J Med. 2022 Mar 30;NEJMoa2119657.
- 176. Holm K, Lundgren MN, Kjeldsen-Kragh J, Ljungquist O, Böttiger B, Wikén C, et al. Convalescence plasma treatment of COVID-19: results from a prematurely terminated



467

randomized controlled open-label study in Southern Sweden. BMC Res Notes. 2021 Dec;14(1):440.

- 177. Ortigoza MB, Yoon H, Goldfeld KS, Troxel AB, Daily JP, Wu Y, et al. Efficacy and Safety of COVID-19 Convalescent Plasma in Hospitalized Patients: A Randomized Clinical Trial. JAMA Intern Med [Internet]. 2021 Dec 13 [cited 2022 Jan 6]; Available from: <u>https://jamanetwork.com/journals/jamainternalmedicine/fullarticle/2787090</u>
- Baldeón ME, Maldonado A, Ochoa-Andrade M, Largo C, Pesantez M, Herdoiza M, et al. Effect of convalescent plasma as complementary treatment in patients with moderate COVID -19 infection. Transfusion Medicine. 2022 Jan 9;tme.12851.
- De Santis GC, Oliveira LC, Garibaldi PMM, Almado CEL, Croda J, Arcanjo
 GGA, et al. High-Dose Convalescent Plasma for Treatment of Severe COVID-19. Emerg
 Infect Dis [Internet]. 2022 Mar [cited 2022 Feb 14];28(3). Available from:
 https://wwwnc.cdc.gov/eid/article/28/3/21-2299_article.htm
- 180. van den Berg K, Glatt TN, Vermeulen M, Little F, Swanevelder R, Barrett C, et al. Convalescent plasma in the treatment of moderate to severe COVID-19 pneumonia: a randomized controlled trial (PROTECT-Patient Trial). Sci Rep. 2022 Dec;12(1):2552.
- 181. Axfors C, Janiaud P, Schmitt AM, van't Hooft J, Smith ER, Haber NA, et al. Association between convalescent plasma treatment and mortality in COVID-19: a collaborative systematic review and meta-analysis of randomized clinical trials. BMC Infect Dis. 2021 Dec;21(1):1170.
- 182. Fernández-Sánchez V, Ventura-Enríquez Y, Cabello-Gutiérrez C, Pérez-Calatayud ÁA, Rosa ECD la, Fareli-González CJ, et al. Convalescent Plasma to Treat Covid-19: a Randomized Double Blind 2 Centers Trial [Internet]. In Review; 2022 Apr [cited 2022 Apr 25]. Available from: <u>https://www.researchsquare.com/article/rs-1277990/v1</u>
- 183. Thorlacius-Ussing L, Brooks PT, Nielsen H, Jensen BA, Wiese L, Sækmose SG, et al. A randomized placebo-controlled trial of convalescent plasma for adults hospitalized with COVID-19 pneumonia. Sci Rep. 2022 Sep 30;12(1):16385.
- 184. Song ATW, Rocha V, Mendrone-Júnior A, Calado RT, De Santis GC, Benites BD, et al. Treatment of severe COVID-19 patients with either low- or high-volume of convalescent plasma versus standard of care: A multicenter Bayesian randomized open-

PAHOS



label clinical trial (COOP-COVID-19-MCTI). The Lancet Regional Health - Americas. 2022 Jun;10:100216.

- 185. Bajpai M, Maheshwari A, Dogra V, Kumar S, Gupta E, Kale P, et al. Efficacy of convalescent plasma therapy in the patient with COVID-19: a randomised control trial (COPLA-II trial). BMJ Open. 2022 Apr 6;12(4):e055189.
- 186. Bartelt LA, Markmann AJ, Nelson B, Keys J, Root H, Henderson HI, et al. Outcomes of convalescent plasma with defined high- versus lower-neutralizing antibody titers against SARS-CoV-2 among hospitalized patients: CoronaVirus Inactivating Plasma (CoVIP), double-blind phase 2 study [Internet]. Infectious Diseases (except HIV/AIDS); 2022 May [cited 2022 May 31]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.04.29.22274387</u>
- 187. Shoham S, Bloch EM, Casadevall A, Hanley D, Lau B, Gebo K, et al. Transfusing convalescent plasma as post-exposure prophylaxis against SARS-CoV-2 infection: a double-blinded, phase 2 randomized, controlled trial. Clinical Infectious Diseases. 2022 May 17;ciac372.
- 188. Rojas M, Rodríguez Y, Hernández JC, Díaz-Coronado JC, Vergara JAD, Vélez VP, et al. Safety and efficacy of convalescent plasma for severe COVID-19: a randomized, single blinded, parallel, controlled clinical study. BMC Infect Dis. 2022 Dec;22(1):575.
- 189. Bargay-Lleonart J, Sarubbo F, Arrizabalaga M, Guerra JM, Borràs J, El Haji K, et al. Reinforcement of the Standard Therapy with Two Infusions of Convalescent Plasma for Patients with COVID-19: A Randomized Clinical Trial. JCM. 2022 May 27;11(11):3039.
- 190. Self WH, Wheeler AP, Stewart TG, Schrager H, Mallada J, Thomas CB, et al. Neutralizing COVID-19 Convalescent Plasma in Adults Hospitalized with COVID-19: A Blinded Randomized Placebo-Controlled Trial. Chest. 2022 Jul;S0012369222012016.
- Balcells ME, Rojas L, Le Corre N, Martínez-Valdebenito C, Ceballos ME, Ferrés M, et al. Early versus deferred anti-SARS-CoV-2 convalescent plasma in patients admitted for COVID-19: A randomized phase II clinical trial. PLoS Med. 2021 Mar;18(3):e1003415.
- 192. Joyner MJ, Bruno KA, Klassen SA, Kunze KL, Johnson PW, Lesser ER, et al. Safety update: COVID-19 convalescent plasma in 20,000 hospitalized patients. Mayo

PAHO®





Clin Proc 2020;95(9):1888–97. Available from: https://doi.org/10.1016/j.mayocp.2020.06.028

- 193. Leucker TM, Osburn WO, Reventun P, Smith K, Claggett B, Kirwan B-A, et al. Effect of Crizanlizumab, a P-Selectin Inhibitor, in COVID-19. JACC: Basic to Translational Science. 2021 Dec;S2452302X21003156.
- 194. Askari G, Sahebkar A, Soleimani D, Mahdavi A, Rafiee S, Majeed M, et al. The efficacy of curcumin-piperine co-supplementation on clinical symptoms, duration, severity, and inflammatory factors in COVID-19 outpatients: a randomized double-blind, placebo-controlled trial. Trials. 2022 Dec;23(1):472.
- 195. Khan A, Iqtadar S, Mumtaz SU, Heinrich M, Pascual-Figal DA, Livingstone S, et al. Oral Co-Supplementation of Curcumin, Quercetin, and Vitamin D3 as an Adjuvant Therapy for Mild to Moderate Symptoms of COVID-19—Results From a Pilot Open-Label, Randomized Controlled Trial. Front Pharmacol. 2022 Jun 7;13:898062.
- 196. Kosiborod MN, Esterline R, Furtado RHM, Oscarsson J, Gasparyan SB, Koch GG, et al. Dapagliflozin in patients with cardiometabolic risk factors hospitalised with COVID-19 (DARE-19): a randomised, double-blind, placebo-controlled, phase 3 trial. The Lancet Diabetes & Endocrinology. 2021 Jul;S2213858721001807.
- 197. Chen J, Xia L, Liu L, Xu Q, Ling Y, Huang D, et al. Antiviral activity and safety of darunavir/cobicistat for the treatment of COVID-19. Open Forum Infect Dis 2020;7(7):ofaa241. Available from: <u>https://doi.org/10.1093/ofid/ofaa241</u>.
- 198. Nickols NG, Mi Z, DeMatt E, Biswas K, Clise CE, Huggins JT, et al. Effect of Androgen Suppression on Clinical Outcomes in Hospitalized Men With COVID-19: The HITCH Randomized Clinical Trial. JAMA Netw Open. 2022 Apr 1;5(4):e227852.
- 199. Madurka I, Vishnevsky A, Soriano JB, Gans SJ, Ore DJS, Rendon A, et al. DFV890: a new oral NLRP3 inhibitor—tested in an early phase 2a randomised clinical trial in patients with COVID-19 pneumonia and impaired respiratory function. Infection [Internet]. 2022 Sep 14 [cited 2022 Sep 28]; Available from: <u>https://link.springer.com/10.1007/s15010-022-01904-w</u>
- 200. Hosseinzadeh A, Emamian MH, Tavakolian A, Kia V, Ebrahimi H, Sheibani H, et al. Application of nasal spray containing dimethyl sulfoxide (DSMO) and ethanol during the COVID-19 pandemic may protect healthcare workers: A randomized controlled trials



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[Internet]. Infectious Diseases (except HIV/AIDS); 2021 Jul [cited 2021 Jul 14]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.07.06.21259749</u>

- 201. Porter JC, Inshaw J, Solis VJ, Denneny E, Evans R, Temkin MI, et al. Antiinflammatory therapy with nebulised dornase alfa in patients with severe COVID-19 pneumonia [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Apr [cited 2022 Apr 28]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.04.14.22272888
- 202. Sobngwi E, Zemsi S, Guewo-Fokeng M, Katte J-C, Kounfack C, Mfeukeu-Kuate L, et al. Doxycycline is a safe alternative to Hydroxychloroquine + Azithromycin to prevent clinical worsening and hospitalization in mild COVID-19 patients: An open label randomized clinical trial (DOXYCOV) [Internet]. Pharmacology and Therapeutics; 2021 Jul [cited 2021 Aug 3]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2021.07.25.21260838

- 203. Butler CC, Yu L-M, Dorward J, Gbinigie O, Hayward G, Saville BR, et al. Doxycycline for community treatment of suspected COVID-19 in people at high risk of adverse outcomes in the UK (PRINCIPLE): a randomised, controlled, open-label, adaptive platform trial. The Lancet Respiratory Medicine. 2021 Jul;S2213260021003106.
- 204. Dhar R, Kirkpatrick J, Gilbert L, Khanna A, Modi MM, Chawla RK, et al. Doxycycline for the prevention of progression of COVID-19 to severe disease requiring intensive care unit (ICU) admission: a randomized, controlled, open-label, parallel group trial (DOXPREVENT.ICU) [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Feb [cited 2022 Feb 15]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2022.01.30.22269685

- 205. Stambouli N, Driss A, Gargouri F, Bahrini K, Arfaoui B, Abid R, et al. COVID-19 prophylaxis with Doxycycline and Zinc in Health Care Workers: A prospective randomized double-blind clinical trial. International Journal of Infectious Diseases. 2022 Jun;S1201971222003496.
- 206. Sasson J, Donlan AN, Ma JZ, Haughey H, Coleman R, Nayak U, et al. Safety and Efficacy of Dupilumab for the Treatment of Hospitalized Patients with Moderate to Severe COVID 19: A Phase IIa Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Apr [cited 2022 Apr 27]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.03.30.22273194



- 207. Cadegiani FA, McCoy J, Wambier CG, Goren A. 5-alpha-reductase inhibitors reduce remission time of COVID-19: results from a randomized double blind placebo controlled interventional trial in 130 SARS-CoV-2 positive men [Preprint]. MedRxiv 2020. Available from: <u>https://doi.org/10.1101/2020.11.16.20232512</u>.
- 208. Cadegiani FA, McCoy J, Gustavo Wambier C, Goren A. Early Antiandrogen Therapy With Dutasteride Reduces Viral Shedding, Inflammatory Responses, and Timeto-Remission in Males With COVID-19: A Randomized, Double-Blind, Placebo-Controlled Interventional Trial (EAT-DUTA AndroCoV Trial – Biochemical). Cureus [Internet]. 2021 Feb 1 [cited 2021 Feb 14]
- 209. Moslemi M, Hejazian SM, Shaddelan M, Javanali F, Mirghaffari A, Sadeghi A, et al. Evaluating the effect of Edaravone on clinical outcome of patients with severe COVID-19 admitted to ICU: a randomized clinical trial. Inflammopharmacol [Internet].
 2022 Jun 20 [cited 2022 Jul 5]; Available from: https://link.springer.com/10.1007/s10787-022-01001-2
- 210. Delgado-Enciso I, Paz-Garcia J, Barajas-Saucedo CE, Mokay-Ramírez KA, Meza-Robles C, Lopez-Flores R, et al. Patient-reported health outcomes after treatment of COVID-19 with nebulized and/or intravenous neutral electrolyzed saline combined with usual medical care versus usual medical care alone: a randomized, open-label, controlled trial [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-68403/v1.
- 211. Gutiérrez-García R, De La Cerda-Angeles JC, Cabrera-Licona A, Delgado-Enciso I, Mervitch-Sigal N, Paz-michel B. Nasopharyngeal and oropharyngeal rinses with neutral electrolyzed water prevents COVID-19 in front-line health professionals: A randomized, open-label, controlled trial in a general hospital in Mexico City. Biomed Rep. 2021 Dec 15;16(2):11.
- 212. Matli K, Al Kotob A, Jamaleddine W, Al Osta S, Salameh P, Tabbikha R, et al. Managing Endothelial Dysfunction in COVID -19 with Statins, Beta Blockers, Nicorandil and Oral Supplements: A Pilot, DOUBLE-BLIND, PLACEBO-CONTROLLED, Randomized Clinical Trial. Clinical Translational Sci. 2022 Jul 8;cts.13369.
- 213. Olha Holubovska, Denisa Bojkova, Stefano Elli, Marco bechtel, David Boltz, Miguel Muzzio, et al. Enisamium is an inhibitor of the SARS-CoV-2 RNA polymerase



and shows improvement of recovery in COVID-19 patients in an interim analysis of a clinical trial. medRxiv [Internet]. 2021.

- 214. Mukae H, Yotsuyanagi H, Ohmagari N, Doi Y, Sakaguchi H, Sonoyama T, et al. Efficacy and safety of ensitrelvir in patients with mild-to-moderate COVID-19: the phase 2b part of a randomized, placebo-controlled, phase 2/3 study [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jun [cited 2022 Jul 5]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.06.22.22276792.
- 215. ACTIV-3/TICO Study Group*. Efficacy and Safety of Ensovibep for Adults Hospitalized With COVID-19: A Randomized Controlled Trial. Ann Intern Med. 2022 Aug 9;M22-1503.
- 216. Welén K, Rosendal E, Gisslén M, Lenman A, Freyhult E, Fonseca-Rodríguez O, et al. A Phase 2 Trial of the Effect of Antiandrogen Therapy on COVID-19 Outcome: No Evidence of Benefit, Supported by Epidemiology and In Vitro Data. European Urology. 2021 Dec;S0302283821022247.
- 217. Amoushahi A, Moazam E, Reza Tabatabaei A, Ghasimi G, Salvatori P, Grant-Whyte I, et al. Efficacy and Safety of Nebulized Ethanol Inhalation in COVID-19 Treatment. A Randomized, Clinical Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jun [cited 2022 Jul 8]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.06.15.22276427
- 218. Samimagham H, Azad M, Haddad M, Arabi M, Hooshyar D, KazemiJahromi M. The Efficacy of Famotidine in improvement of outcomes in Hospitalized COVID-19 Patients: A phase III randomised clinical trial. ResearchSquare [Internet]. 2021; Available from:

http://www.epistemonikos.org/documents/a38a60b031b058f125e2d5572d2bc7678b6764 98

- 219. Brennan CM, Nadella S, Zhao X, Dima RJ, Jordan-Martin N, Demestichas BR, 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 Feb 10;gutjnl-2022-326952.
- 220. Pahwani S, Jadwani M, Dhanwani A, Gul M, Lal D, Rakesh F, et al. Efficacy of Oral Famotidine in Patients Hospitalized With Severe Acute Respiratory Syndrome Coronavirus 2. Cureus [Internet]. 2022 Feb 20 [cited 2022 May 2]; Available from:



https://www.cureus.com/articles/78980-efficacy-of-oral-famotidine-in-patientshospitalized-with-severe-acute-respiratory-syndrome-coronavirus-2

- 221. Chen C, Huang J, Cheng Z, Wu J, Chen S, Zhang Y, et al. Favipiravir versus arbidol for COVID-19: a randomized clinical trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.03.17.20037432.
- 222. Ivashchenko AA, Dmitriev KA, Vostokova NV, Azarova VN, Blinow AA, Egorova AN, et al. Interim results of a phase II/III multicenter randomized clinical trial of AVIFAVIR in hospitalized patients with COVID-19. MedRxiv 202. Available from: https://doi.org/10.1101/2020.07.26.20154724.
- 223. Doi Y, Hibino M, Hase R, Yamamoto M, Kasamatsu Y, Hirose M, et al. A prospective, randomized, open-label trial of early versus late favipiravir in hospitalized patients with COVID-19. Antimicrob Agents Chemother 2020; 64:e01897-20. Available from: https://doi.org/10.1128/AAC.01897-20.
- 224. Dabbous HM, El-Sayed MH, El Assal G, Elghazaly H, Ebeid FFS, Sherief AF, et al. A randomized controlled study of favipiravir vs hydroxychloroquine in COVID-19 management: what have we learned so far? [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-83677/v1.
- 225. Zhao H, Zhu Q, Zhang C, Li J, Wei M, Qin Y, et al. Tocilizumab combined with favipiravir in the treatment of COVID-19: a multicenter trial in a small sample size. Biomed Pharmacother 2021; 133:110825. Available from: https://doi.org/10.1016/j.biopha.2020.110825.
- 226. Khamis F, Al Naabi H, Al Lawati A, Ambusaidi Z, Al Sharji M, Al Barwani U, et al. Randomized controlled open label trial on the use of favipiravir combined with inhaled interferon beta-1b in hospitalized patients with moderate to severe COVID-19 pneumonia. Int J Infect Dis 2020; 102:538-43. Available from: https://doi.org/10.1016/j.ijid.2020.11.008.
- 227. Ruzhentsova TA, Oseshnyuk RA, Soluyanova TN, Dmitrikova EP, Mustafaev DM, Pokrovskiy KA, et al. Phase 3 trial of coronavir (favipiravir) in patients with mild to moderate COVID-19. Am J Transl Res. 2021;13(11):12575–87.
- 228. Udwadia ZF, Singh P, Barkate H, Patil S, Rangwala S, Pendse A, et al. Efficacy and safety of favipiravir, an oral RNA-dependent RNA polymerase inhibitor, in mild-tomoderate COVID-19: a randomized, comparative, open-label, multicenter, phase 3

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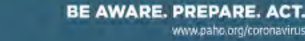
clinical trial [Preprint]. Int J Infect Dis 2020. Available from: https://doi.org/10.1016/j.ijid.2020.11.142.

- 229. Ogarev Mordovia State University, Saransk, Russian Federation, Balykova LA, Govorov AV, A.I.Evdokimov Moscow State University of Medicine and Dentistry, Moscow, Russian Federation, Vasilyev AO, A.I.Evdokimov Moscow State University of Medicine and Dentistry, Moscow, Russian Federation, et al. Characteristics of COVID-19 and possibilities of early causal therapy. Results of favipiravir use in clinical practice. Infekc bolezni. 2020;18(3):30–40.
- 230. Solaymani-Dodaran M, Ghanei M, Bagheri M, Qazvini A, Vahedi E, Hassan Saadat S, et al. Safety and efficacy of Favipiravir in moderate to severe SARS-CoV-2 pneumonia. International Immunopharmacology. 2021 Jun;95:107522.
- 231. Zhao H, Zhang C, Zhu Q, Chen X, Chen G, Sun W, et al. Favipiravir in the treatment of patients with SARS-CoV-2 RNA recurrent positive after discharge: A multicenter, open-label, randomized trial. International Immunopharmacology. 2021 Aug;97:107702.
- 232. Bosaeed M, Mahmoud E, Alharbi A, Altayeib H, Albayat H, Alharbi F, et al. Favipiravir and Hydroxychloroquine Combination Therapy in Patients with Moderate to Severe COVID-19 (FACCT): An Open-Label, Multicentre, Randomised, Controlled Trial. SSRN Journal [Internet]. 2021 [cited 2021 May 5]; Available from: <u>https://www.ssrn.com/abstract=3829663</u>
- 233. Shinkai M, Tsushima K, Tanaka S, Hagiwara E, Tarumoto N, Kawada I, et al. Efficacy and Safety of Favipiravir in Moderate COVID-19 Pneumonia Patients without Oxygen Therapy: A Randomized, Phase III Clinical Trial. Infect Dis Ther [Internet].
 2021 Aug 27 [cited 2021 Sep 6]; Available from: https://link.springer.com/10.1007/s40121-021-00517-4
- 234. Atipornwanich K, Kongsaengdao S, Harnsomburana P, Nanna R, Chtuparisute C, Saengsayan P, et al. Various Combinations of Favipiravir, Lopinavir-Ritonavir, Darunavir-Ritonavir, High-Dose Oseltamivir, and Hydroxychloroquine for the Treatment of COVID-19: A Randomized Controlled Trial (FIGHT-COVID-19 Study). SSRN Journal [Internet]. 2021 [cited 2021 Oct 13]; Available from: https://www.ssrn.com/abstract=3936499



- 235. Shenoy S, Munjal S, Youha SA, Alghounaim M, Almazeedi S, Alshamali Y, et al. Favipiravir In Adults with Moderate to Severe COVID-19: A Phase 3 Multicentre, Randomized, Double-Blinded, Placebo-Controlled Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Nov [cited 2021 Nov 26]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.11.08.21265884
- 236. Holubar M, Subramanian A, Purington N, Hedlin H, Bunning B, Walter KS, et al. Favipiravir for treatment of outpatients with asymptomatic or uncomplicated COVID-19: a double-blind randomized, placebo-controlled, phase 2 trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Nov [cited 2021 Dec 8]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.11.22.21266690</u>
- 237. Chuah CH, Chow TS, Hor CP, Cheng JT, Ker HB, Lee HG, et al. Efficacy of Early Treatment with Favipiravir on Disease Progression among High Risk COVID-19 Patients: A Randomized, Open-Label Clinical Trial. Clinical Infectious Diseases. 2021 Nov 19;ciab962.
- 238. Finberg RW, Ashraf M, Julg B, Ayoade F, Marathe JG, Issa NC, et al. US201 Study: A Phase 2, Randomized Proof-of-Concept Trial of Favipiravir for the Treatment of COVID-19. Open Forum Infectious Diseases. 2021 Dec 1;8(12):ofab563.
- 239. Bosaeed M, Alharbi A, Mahmoud E, Alrehily S, Bahlaq M, Gaifer Z, et al. Efficacy of favipiravir in adults with mild COVID-19: a randomized, double-blind, multicenter, placebo-controlled trial clinical trial. Clinical Microbiology and Infection. 2022 Jan;S1198743X21007345.
- 240. Hassaniazad M, Farshidi H, Gharibzadeh A, Bazram A, Khalili E, Noormandi A, et al. Efficacy and safety of favipiravir plus interferon-beta versus lopinavir/ritonavir plus interferon-beta in moderately ill patients with COVID-19: A randomized clinical trial. Journal of Medical Virology. 2022 Mar 24;jmv.27724.
- 241. Lowe DM, Brown L-AK, Chowdhury K, Davey S, Yee P, Ikeji F, et al. Favipiravir, lopinavir-ritonavir or combination therapy (FLARE): a randomised, double blind, 2x2 factorial placebo-controlled trial of early antiviral therapy in COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Feb [cited 2022 Mar 31]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.02.11.22270775
- 242. Tabarsi P, Vahidi H, Saffaei A, Hashemian SMR, Jammati H, Daraei B, et al. Favipiravir Effects on the Control of Clinical Symptoms of Hospitalized COVID-19

PAHO®



Cases: An Experience with Iranian Formulated Dosage Form. Iran J Pharm Res. 2021;20(4):1–8.

- 243. AlQahtani M, Kumar N, Aljawder D, Abdulrahman A, Alnashaba F, Fayyad MA, et al. Randomized controlled trial of favipiravir, hydroxychloroquine, and standard care in patients with mild/moderate COVID-19 disease. Sci Rep. 2022 Mar 23;12(1):4925.
- 244. Rahman SMA, Kabir A, Abdullah ABM, Alam MB, Azad KAK, Miah MT, et al. Safety and efficacy of favipiravir for the management of COVID-19 patients: A preliminary randomized control trial. Clinical Infection in Practice. 2022 Jul;15:100145.
- 245. McMahon JH, Lau JSY, Coldham A, Roney J, Hagenauer M, Price S, et al. Favipiravir in Early Symptomatic COVID-19, A Randomised Placebo-Controlled Trial. SSRN Journal [Internet]. 2022 [cited 2022 Jul 6]; Available from: <u>https://www.ssrn.com/abstract=4135325</u>
- 246. Golan Y, Campos JAS, Woolson R, Cilla D, Hanabergh R, Gonzales-Rojas Y, et al. Favipiravir in patients with early mild-to-moderate COVID-19: a randomized controlled trial. Clinical Infectious Diseases. 2022 Sep 6;ciac712.
- 247. Sirijatuphat R, Manosuthi W, Niyomnaitham S, Owen A, Copeland KK, Charoenpong L, et al. Early Treatment of Favipiravir in COVID-19 Patients Without Pneumonia: A Multicentre, Open-Labelled, Randomized Control Study [Internet]. Pharmacology and Therapeutics; 2022 Jun [cited 2022 Sep 19]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.06.06.22275902</u>
- 248. Davoodi L, Abedi SM, Salehifar E, Alizadeh-Navai R, Rouhanizadeh H, Khorasani G, Hosseinimehr SJ. Febuxostat therapy in outpatients with suspected COVID-19: a clinical trial. Int J Clin Pract 2020; 74:e13600. Available from: <u>https://doi.org/10.1111/ijcp.13600</u>.
- Chirinos J, Lopez-Jaramillo P, Giamarellos-Bourboulis E, Dávila-del-Carpio G,
 Bizri A, Andrade-Villanueva J, et al. A Randomized Trial of Lipid Metabolism
 Modulation with Fenofibrate for Acute Coronavirus Disease 2019 [Internet]. In Review;
 2022 Aug [cited 2022 Aug 20]. Available from:

https://www.researchsquare.com/article/rs-1933913/v1

250. E. Zarehoseinzade, A. Allami, M. Ahmadi, B. Bijani, N. Mohammadi. Finasteride in hospitalized adult males with Covid-19: A risk factor for severity of the disease or an



adjunct treatment: A randomized controlled clinical trial. The Medical Journal of The Islamic Republic of Iran [Internet]. 2021;35(1). Available from: http://www.epistemonikos.org/documents/f3b23e45ed8faff34c8ba4b500fc9bfc82d32f81

- 251. Lenze EJ, Mattar C, Zorumski CF, Stevens A, Schweiger J, Nicol GE, et al. Fluvoxamine vs placebo and clinical deterioration in outpatients with symptomatic COVID-19: a randomized clinical trial. JAMA 2020 Published online November 12, 2020. Available from: <u>https://doi.org/10.1001/jama.2020.22760</u>.
- 252. Reis G, dos Santos Moreira-Silva EA, Silva DCM, Thabane L, Milagres AC, Ferreira TS, 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. The Lancet Global Health. 2021 Oct;S2214109X21004484.
- 253. Seo H, Kim H, Bae S, Park S, Chung H, Sung H sup, et al. Fluvoxamine Treatment of Patients with Symptomatic COVID-19 in a Community Treatment Center: A Preliminary Result of Randomized Controlled Trial. Infect Chemother. 2022;54(1):102.
- 254. Bramante CT, Huling JD, Tignanelli CJ, Buse JB, Liebovitz DM, Nicklas JM, et al. Randomized Trial of Metformin, Ivermectin, and Fluvoxamine for Covid-19. N Engl J Med. 2022 Aug 18;387(7):599–610.
- 255. Strich JR, Tian X, Samour M, King CS, Shlobin O, Reger R, et al. Fostamatinib for the treatment of hospitalized adults with COVID-19 A randomized trial. Clinical Infectious Diseases. 2021 Sep 1;ciab732.
- 256. Soltani R, Nasirharandi S, Khorvash F, Nasirian M, Dolatshahi K, Hakamifard A. The effectiveness of gabapentin and gabapentin/montelukast combination compared with dextromethorphan in the improvement of COVID-19- related cough: A randomized, controlled clinical trial. Clinical Respiratory J. 2022 Jul 31;crj.13529.
- 257. Gaughan E, Sethi T, Quinn T, Hirani N, Mills A, Bruce AM, et al. GB0139, an inhaled small molecule inhibitor of galectin-3, in COVID-19 pneumonitis: a randomised, controlled, open-label, phase 2a experimental medicine trial of safety, pharmacokinetics, and potential therapeutic value [Internet]. Respiratory Medicine; 2021 Dec [cited 2021 Dec 30]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.12.21.21267983
- 258. Criner GJ, Lang FM, Gottlieb RL, Mathews KS, Wang TS, Rice TW, et al. Anti-GM-CSF Monoclonal Antibody Gimsilumab for COVID-19 Pneumonia: A Randomized,

PAHOS

Double-Blind, Placebo-Controlled Trial. Am J Respir Crit Care Med. 2022 Mar 15;rccm.202108-1859OC.

- 259. Shogenova LV, Petrikov SS, Zhuravel SV, Gavrilov PV, Utkina II, Varfolomeev SD, et al. Thermal Helium-Oxygen Mixture as Part of a Treatment Protocol for Patients with COVID-19. Annals RAMS. 2020 Dec 4;75(5S):353–62.
- 260. Dupuis J, Laurin P, Tardif J-C, Hausermann L, Rosa C, Guertin M-C, et al. Fourteen-days Evolution of COVID-19 Symptoms During the Third Wave in Nonvaccinated Subjects and Effects of Hesperidin Therapy: A randomized, double-blinded, placebo-controlled study [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Oct [cited 2021 Oct 13]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2021.10.04.21264483

- 261. Jarczak D, Roedl K, Fischer M, de Heer G, Burdelski C, Frings DP, et al. Effect of Hemadsorption in Critically Ill Patients with COVID-19 (CYTOCOV-19): A Prospective Randomized Controlled Pilot Trial [Internet]. In Review; 2021 Jul [cited 2021 Nov 23]. Available from: <u>https://www.researchsquare.com/article/rs-704552/v1</u>
- 262. Borba MGS, Val FFA, Sampaio VS, Alexandre MAA, Melo GC, Brito M, et al. Effect of high vs low doses of chloroquine diphosphate as adjunctive therapy for patients hospitalized with severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection: a randomized clinical trial. JAMA Netw Open 2020;3(4):e208857. Available from: https://doi.org/10.1001/jamanetworkopen.2020.8857.
- 263. Huang M, Tang T, Pang P, Li M, Ma R, Lu J, et al. Treating COVID-19 with chloroquine. J Mol Cell Biol 2020;12(4):322–25. Available from: https://doi.org/10.1093/jmcb/mjaa014.
- 264. The RECOVERY Collaborative Group. Effect of hydroxychloroquine in hospitalized patients with COVID-19. N Engl J Med 2020;383:2030-40. Available from: https://doi.org/10.1056/NEJMoa2022926.
- 265. Mitja O, Ubals M, Corbacho M, Alemany A, Suner C, Tebe C, et al. A clusterrandomized trial of hydroxychloroquine as prevention of COVID-19 transmission and disease [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.07.20.20157651.
- 266. Boulware DR, Pullen MF, Bangdiwala AS, Pastick KA, Lofgren SM, Okafor EC, et al. A randomized trial of hydroxychloroquine as postexposure prophylaxis for COVID-



19. N Engl J Med 2020;383:517-25. Available from: https://doi.org/10.1056/NEJMoa2016638.

- 267. Cavalcanti AB, Zampieri FG, Rosa RG, Azevedo LCP, Veiga VC, Avezum A, et al. Hydroxychloroquine with or without azithromycin in mild-to-moderate COVID-19. N Engl J Med 2020;383:2041-52. Available from: https://doi.org/10.1056/NEJMoa2019014.
- 268. Kamran SM, Mirza ZH, Naseem A, Saeed F, Azam R, Ullah N, et al. Clearing the fog: is HCQ effective in reducing COVID-19 progression: a randomized controlled trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.07.30.20165365.
- 269. Skipper CP, Pastick KA, Engen NW, Bangdiwala AS, Abassi M, Lofgren SM, et al. Hydroxychloroquine in nonhospitalized adults with early COVID-19: a randomized trial. Ann Int Med 2020;173(8):623-31. Available from: https://doi.org/10.7326/M20-4207.
- 270. Mitjà O, Corbacho-Monné M, Ubals M, Tebe C, Peñafiel J, Tobias A, et al. Hydroxychloroquine for early treatment of adults with mild COVID-19: a randomizedcontrolled trial. Clin Infect Dis 2020; ciaa1009. Available from: https://doi.org/10.1093/cid/ciaa1009.
- 271. Tang W, Cao Z, Han M, Wang Z, Chen J, Sun W, et al. Hydroxychloroquine in patients with mainly mild to moderate coronavirus disease 2019: open label, randomised controlled trial. BMJ 2020;369:m1849. Available from: https://doi.org/10.1136/bmj.m1849.
- 272. Chen Z, Hu J, Zhang Z, Jiang SS, Han S, Yan D, et al. Efficacy of hydroxychloroquine in patients with COVID-19: results of a randomized clinical trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.03.22.20040758.
- 273. Chen L, Zhang Z-y, Fu J-g, Feng Z-p, Zhang S-z, Han Q-y, et al. Efficacy and safety of chloroquine or hydroxychloroquine in moderate type of COVID-19: a prospective open-label randomized controlled study [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.06.19.20136093.
- 274. Chen C-P, Lin Y-C, Chen T-C, Tseng T-Y, Wong H-L, Kuo C-Y, et al. A multicenter, randomized, open-label, controlled trial to evaluate the efficacy and tolerability of hydroxychloroquine and a retrospective study in adult patients with mild to



moderate coronavirus disease 2019 (COVID-19) [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.07.08.20148841.

- 275. Chen J, Liu D, Liu L, Liu P, Xu Q, Xia L, et al. A pilot study of hydroxychloroquine in treatment of patients with moderate COVID-19. 浙江大学学报 (医学版) (Journal of Zhejiang University. Medical Sciences) 2020; 49(2):215–19. Available from: https://doi.org/10.3785/j.issn.1008-9292.2020.03.03.
- 276. Abd-Elsalam S, Esmail ES, Khalaf M, Abdo EF, Medhat MA, Abd El Ghafar MS, et al. Hydroxychloroquine in the treatment of COVID-19: a multicenter randomized controlled study. Am J Trop Med Hyg 2020; 13(4):635-39. Available from: https://doi.org/10.4269/ajtmh.20-0873.
- 277. Rajasingham R, Bangdiwala AS, Nicol MR, Skipper CP, Pastick KA, Axelrod ML, et al. Hydroxychloroquine as pre-exposure prophylaxis for COVID-19 in healthcare workers: a randomized trial. Clin Infect Dis 2020; ciaa1571. Available from: https://doi.org/10.1093/cid/ciaa1571.
- 278. Ulrich RJ, Troxel AB, Carmody E, Eapen J, Bäcker M, DeHovitz JA, 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. Available from: https://doi.org/10.1093/ofid/ofaa446.
- 279. Grau-Pujol B, Camprubí D, Marti-Soler H, Fernández-Pardos M, Carreras-Abad C, et al. Pre-exposure prophylaxis with hydroxychloroquine for COVID-19: initial results of a double-blind, placebo-controlled randomized clinical trial [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-72132/v1.
- 280. Abella BS, Jolkovsky EL, Biney BT, Uspal JE, Hyman MC, Frank I, et al. Efficacy and safety of hydroxychloroquine vs placebo for pre-exposure SARS-CoV-2 prophylaxis among health care workers: a randomized clinical trial. JAMA Int Med 2020 published online September 30. Available from:

https://doi.org/10.1001/jamainternmed.2020.6319.

- 281. Remdesivir and three other drugs for hospitalised patients with COVID-19: final results of the WHO Solidarity randomised trial and updated meta-analyses. The Lancet. 2022 May;S0140673622005190.
- 282. Barnabas RV, Brown ER, Bershteyn A, Stankiewicz Karita HC, Johnston C, Thorpe LE, Kottkamp A, et al. Hydroxychloroquine as Postexposure Prophylaxis to



Prevent Severe Acute Respiratory Syndrome Coronavirus 2 Infection : A Randomized Trial. Annals of Internal Medicine 2020. <u>https://doi.org/10.7326/M20-6519</u>.

- 283. Self WH, Semler MW, Leither LM, Casey JD, Angus DC, Brower RG, 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. Available from: https://doi.org/10.1001/jama.2020.22240.
- 284. Brown SM, Peltan I, Kumar N, Leither L, Webb BJ, Starr N, et al. Hydroxychloroquine vs. azithromycin for hospitalized patients with COVID-19 (HAHPS): results of a randomized, active comparator trial. Ann Am Thor Soc 2020; published online 9 November 2020. Available from: https://doi.org/10.1513/AnnalsATS.202008-940OC.
- 285. Dubée V, Roy P-M, Vielle B, Parot-Schinkel E, Blanchet O, Darsonval A, et al. Hydroxychloroquine in mild-to-moderate COVID-19: a placebo-controlled double blind trial. Clinical Microbiology and Infection. 2021 Apr;S1198743X21001403.
- 286. Omrani AS, Pathan SA, Thomas SA, Harris TRE, Coyle PV, Thomas CE, et al. Randomized double-blinded placebo-controlled trial of hydroxychloroquine with or without azithromycin for virologic cure of non-severe COVID-19. EClinicalMedicine 2020;29: 100645. Available from: https://doi.org/10.1016/j.eclinm.2020.100645.
- 287. Dabbous HM, El-Sayed MH, Assal GE, Elghazaly H, Ebeid FF, Sherief AF, et al. A Randomized Controlled Study Of Favipiravir Vs Hydroxychloroquine In COVID-19 Management: What Have We Learned So Far? [Internet]. In Review; 2020 Sep [cited 2020 Oct 1]. Available from: https://www.researchsquare.com/article/rs-83677/v1
- 288. Hernandez-Cardenas C, Thirion-Romero I, Rivera-Martinez NE, Meza-Meneses P, Remigio-Luna A, Perez-Padilla R. Hydroxychloroquine for the Treatment of Severe Respiratory Infection by Covid-19: A Randomized Controlled Trial. medRxiv [Internet]. 2021; Available from:

http://www.epistemonikos.org/documents/0881ad73607247595bdf210de533bbd94651b0 b4

289. Johnston C, Brown ER, Stewart J, Karita HCS, Kissinger PJ, Dwyer J, et al. Hydroxychloroquine with or without azithromycin for treatment of early SARS-CoV-2 infection among high-risk outpatient adults: A randomized clinical trial. EClinicalMedicine. 2021 Feb;100773.

PAHOS



- 290. Purwati, Budiono, Rachman BE, Yulistiani, Miatmoko A, Nasronudin, et al. A Randomized, Double-Blind, Multicenter Clinical Study Comparing the Efficacy and Safety of a Drug Combination of Lopinavir/Ritonavir-Azithromycin, Lopinavir/Ritonavir-Doxycycline, and Azithromycin-Hydroxychloroquine for Patients Diagnosed with Mild to Moderate COVID-19 Infections. Huyut Z, editor. Biochemistry Research International. 2021 Feb 9;2021:1–12.
- 291. Beltran Gonzalez JL, González Gámez M, Mendoza Enciso EA, Esparza Maldonado RJ, Hernández Palacios D, Dueñas Campos S, et al. Efficacy and Safety of Ivermectin and Hydroxychloroquine in Patients with Severe COVID-19: A Randomized Controlled Trial. Infectious Disease Reports. 2022 Mar 3;14(2):160–8.
- 292. Amaravadi RK, Giles L, Carberry M, Hyman MC, Frank I, Nasta SD, et al. Hydroxychloroquine for SARS-CoV-2 positive patients quarantined at home: The first interim analysis of a remotely conducted randomized clinical trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Feb [cited 2021 Mar 4]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.02.22.21252228
- 293. Galan LEB, Santos NM dos, Asato MS, Araújo JV, de Lima Moreira A, Araújo AMM, et al. Phase 2 randomized study on chloroquine, hydroxychloroquine or ivermectin in hospitalized patients with severe manifestations of SARS-CoV-2 infection. Pathogens and Global Health. 2021 Mar 8;1–8.
- 294. Seet RCS, Quek AML, Ooi DSQ, Sengupta S, Lakshminarasappa SR, Koo CY, et al. Positive impact of oral hydroxychloroquine and povidone-iodine throat spray for COVID-19 prophylaxis: an open-label randomized trial. International journal of infectious diseases : IJID : official publication of the International Society for Infectious Diseases [Internet]. 2021; Available from:

http://www.epistemonikos.org/documents/f0a6f1dede7897794397549169853a5d5c7c6c0 e

- 295. Reis G, Moreira Silva EADS, Medeiros Silva DC, Thabane L, Singh G, Park JJH, 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 network open. 2021;4(4):e216468.
- 296. Réa-Neto Á, Bernardelli RS, Câmara BMD, Reese FB, Queiroga MVO, Oliveira MC. An open-label randomized controlled trial evaluating the efficacy of

PAHOS



chloroquine/hydroxychloroquine in severe COVID-19 patients. Sci Rep. 2021 Dec;11(1):9023.

- 297. Syed F, Hassan M, Arif MA, Batool S, Niazi R, Laila U e, et al. Pre-exposure Prophylaxis With Various Doses of Hydroxychloroquine Among Healthcare Personnel With High-Risk Exposure to COVID-19: A Randomized Controlled Trial. Cureus [Internet]. 2021 Dec 21 [cited 2022 Feb 16]; Available from: <u>https://www.cureus.com/articles/77806-pre-exposure-prophylaxis-with-various-doses-ofhydroxychloroquine-among-healthcare-personnel-with-high-risk-exposure-to-covid-19-arandomized-controlled-trial</u>
- 298. Sivapalan P, Suppli Ulrik C, Sophie Lapperre T, Dahlin Bojesen R, Eklöf J, Browatzki A, et al. Azithromycin and hydroxychloroquine in hospitalised patients with confirmed COVID-19–a randomised double-blinded placebo-controlled trial. Eur Respir J. 2021 Jun 3;2100752.
- 299. Byakika-Kibwika P, Sekaggya-Wiltshire C, Semakula JR, Nakibuuka J, Musaazi J, Kayima J, et al. Safety and efficacy of hydroxychloroquine for treatment of non-severe COVID-19 among adults in Uganda: a randomized open label phase II clinical trial. BMC Infect Dis. 2021 Dec;21(1):1218.
- 300. Schwartz I, Boesen ME, Cerchiaro G, Doram C, Edwards BD, Ganesh A, et al. Assessing the efficacy and safety of hydroxychloroquine as outpatient treatment of COVID-19: a randomized controlled trial. cmajo. 2021 Apr;9(2):E693–702.
- 301. Naggie S, Milstone A, Castro M, Collins SP, Seetha L, Anderson DJ, et al. Hydroxychloroquine for pre-exposure prophylaxis of COVID-19 in health care workers: a randomized, multicenter, placebo-controlled trial (HERO-HCQ) [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Aug [cited 2021 Aug 30]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.08.19.21262275
- 302. Rodrigues C, Freitas-Santos RS, Levi JE, Senerchia AA, Lopes ATA, Santos SR, et al. Hydroxychloroquine plus azithromycin early treatment of mild COVID-19 in outpatient setting: a randomized, double-blinded, placebo-controlled clinical trial evaluating viral clearance. International Journal of Antimicrobial Agents. 2021 Aug;106428.
- 303. Babalola OE, Yahaya N, Ajayi AA, Ogedengbe JO, Thairu Y, Omede O. A Randomized Controlled Trial of Ivermectin Monotherapy Versus Hydroxychloroquine,



Ivermectin, and Azithromycin Combination Therapy in Covid-19 Patients in Nigeria [Internet]. In Review; 2021 Oct [cited 2021 Oct 12]. Available from: https://www.researchsquare.com/article/rs-950352/v1

- 304. Panda PK, Singh BO, Moirangthem B, Bahurupi YA, Saha S, Saini G, et al. Antiviral Combination Clinically Better Than Standard Therapy in Severe but Not in Non-Severe COVID-19. CPAA. 2021 Sep;Volume 13:185–95.
- 305. Ahmad B, ul Hassan N, Sehar B, Zeb F, e Nayab D, Siddiqui FA. Effect of Chloroquine and Hydroxychloroquine on Cytokine Release Syndrome in Patients with COVID-19. Clin Med Res. 2021 Dec;19(4):179–82.
- 306. McKinnon J, Wang D, Zervos M, Saval M, Marshall-Nightengale L, Kilgore P, et al. Safety and Tolerability of Hydroxychloroquine in healthcare workers and first responders for the prevention of COVID-19: WHIP COVID-19 Study. International Journal of Infectious Diseases. 2021 Dec;S1201971221012431.
- 307. Rojas-Serrano J, Portillo-Vásquez AM, Thirion-Romero I, Vázquez-Pérez J, Mejía-Nepomuceno F, Ramírez-Venegas A, et al. Hydroxychloroquine for prophylaxis of COVID-19 in health workers: A randomized clinical trial. Triche EW, editor. PLoS ONE. 2022 Feb 9;17(2):e0261980.
- 308. Polo R, García-Albéniz X, Terán C, Morales M, Rial-Crestelo D, Garcinuño MA, et al. Daily tenofovir disoproxil fumarate/emtricitabine and hydroxychloroquine for preexposure prophylaxis of COVID-19: a double-blind placebo controlled randomized trial in healthcare workers. Clinical Microbiology and Infection. 2022 Aug;S1198743X22003706.
- 309. Avezum Á, Oliveira GBF, Oliveira H, Lucchetta RC, Pereira VFA, Dabarian AL, et al. Hydroxychloroquine versus placebo in the treatment of non-hospitalised patients with COVID-19 (COPE Coalition V): A double-blind, multicentre, randomised, controlled trial. The Lancet Regional Health Americas. 2022 Jul;11:100243.
- 310. Roy-García IA, Moreno-Noguez M, Rivas-Ruiz R, Zapata-Tarres M, Perez-Rodriguez M, Ortiz-Zamora MA, et al. "Efficacy and Safety of Fixed Combination of Hydroxychloroquine with Azithromycin Versus Hydroxychloroquine and Placebo in Patients with Mild COVID-19: Randomized, double blind, Placebo controlled trial" [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Apr [cited 2022 Apr 25]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.04.06.22273531</u>



- 311. Tirupakuzhi Vijayaraghavan BK, Jha V, Rajbhandari D, Myatra SN, Ghosh A, Bhattacharya A, et al. Hydroxychloroquine plus personal protective equipment versus personal protective equipment alone for the prevention of laboratory-confirmed COVID-19 infections among healthcare workers: a multicentre, parallel-group randomised controlled trial from India. BMJ Open. 2022 Jun;12(6):e059540.
- 312. Elshafie AH, Elsawah HK, Hammad M, Sweed EM, Seif AS, Abdel Ghaffar MM, et al. Ivermectin role in COVID-19 treatment (IRICT): single-center, adaptive, randomized, double-blind, placebo-controlled, clinical trial. Expert Review of Antiinfective Therapy. 2022 Jul 12;1–10.
- 313. Choudhary R, Ali O, Singh BK. Study on Hydroxychloroquinine Sulfate Being Given to the Admitted COVID -19 Positive Patients at Institute of JLNMCH, Bhagalpur, Bihar, India. Cureus [Internet]. 2022 Jun 28 [cited 2022 Aug 16]; Available from: <u>https://www.cureus.com/articles/100191-study-on-hydroxychloroquinine-sulfate-beinggiven-to-the-admitted-covid--19-positive-patients-at-institute-of-jlnmch-bhagalpur-biharindia</u>
- 314. Hadanny A, Finci S, Catalogna M, Abu Hamed R, Korin C, Gabriella L, et al. Hyperbaric Oxygen Therapy for COVID-19 Patients: A Prospective, Randomized Controlled Trial. SSRN Journal [Internet]. 2020 [cited 2021 Apr 19]; Available from: https://www.ssrn.com/abstract=3745115
- 315. Cannellotto M, Duarte M, Keller G, Larrea R, Cunto E, Chediack V, et al. Hyperbaric oxygen as an adjuvant treatment for patients with COVID-19 severe hypoxaemia: a randomised controlled trial. Emerg Med J. 2021 Dec 14;emermed-2021-211253.
- 316. Kjellberg A, Douglas J, Hassler A, Al-Ezerjawi S, Boström E, Abdel-Halim L, et al. COVID-19 induced acute respiratory distress syndrome treated with Hyperbaric Oxygen: Interim safety report from a multicenter, randomised, open-label phase II clinical trial (COVID-19-HBO). ResearchSquare [Internet]. 2022; Available from: http://www.epistemonikos.org/documents/db28a97c703a59e6c2ea3d021c759a1dc577c11 f
- Ali S, Uddin SM, Shalim E, Sayeed MA, Anjum F, Saleem F, et al.
 Hyperimmune anti-COVID-19 IVIG (C-IVIG) treatment in severe and critical COVID-19 patients: A phase I/II randomized control trial. EClinicalMedicine. 2021 Jun;100926.

PAHO®

- 318. Parikh D, Chaturvedi A, Shah N, Patel P, Patel R, Ray S. Safety and efficacy of COVID-19 hyperimmune globulin (HIG) solution in the treatment of active COVID-19 infection- Findings from a Prospective, Randomized, Controlled, Multi-Centric Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Jul [cited 2021 Aug 17]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.07.26.21261119
- 319. Polizzotto MN, Nordwall J, Babiker AG, Phillips A, Vock DM, Eriobu N, et al. Hyperimmune immunoglobulin for hospitalised patients with COVID-19 (ITAC): a double-blind, placebo-controlled, phase 3, randomised trial. The Lancet. 2022 Feb;399(10324):530–40.
- 320. Huygens S, Hofsink Q, Nijhof IS, Goorhuis A, Kater AP, te Boekhorst PA, et al. SARS-CoV-2 hyperimmune globulin for severely immunocompromised patients with COVID-19: a randomised, controlled, double-blind, phase 3 trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Apr [cited 2022 Apr 27]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.04.04.22273314</u>
- 321. Prasenohadi P, Burhan E, Dhunny S, Suharno W, Wabnitz P, Kim YW, et al. Double-Blind, Randomized, Placebo-Controlled Study on hzVSF-v13, a Novel Anti-Vimentin Monoclonal Antibody Drug as Add-on Standard of Care in the Management of Patients with Moderate to Severe COVID-19. JCM. 2022 May 24;11(11):2961.
- 322. Coutre SE, Barnett C, Osiyemi O, Hoda D, Ramgopal M, Fort AC, et al. Ibrutinib for Hospitalized Adults With Severe Coronavirus Disease 2019 Infection: Results of the Randomized, Double-Blind, Placebo-Controlled iNSPIRE Study. Open Forum Infectious Diseases. 2022 May 1;9(5):ofac104.
- 323. Mansour E, Palma AC, Ulaf RG, Ribeiro LC, Bernardes AF, Nunes TA, et al. Pharmacological inhibition of the kinin-kallikrein system in severe COVID-19: a proofof-concept study [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.08.11.20167353.
- 324. Kosmopoulos A, Bhatt DL, Meglis G, Verma R, Pan Y, Quan A, et al. A Randomized Trial of Icosapent Ethyl in Ambulatory Patients with COVID-19. iScience. 2021 Aug;103040.
- 325. Aman J, Duijvelaar E, Botros L, Kianzad A, Schippers JR, Smeele PJ, et al. Imatinib in patients with severe COVID-19: a randomised, double-blind, placebo-



on World Health Organization controlled, clinical trial. The Lancet Respiratory Medicine. 2021 Jun;S221326002100237X.

- 326. Ravichandran R, Mohan SK, Sukumaran SK, Kamaraj D, Daivasuga SS, Ravi SOAS, et al. An open label randomized clinical trial of Indomethacin for mild and moderate hospitalised Covid-19 patients. Sci Rep. 2022 Dec;12(1):6413.
- 327. Fisher BA, Veenith T, Slade D, Gaskell C, Rowland M, Whitehouse T, et al. Namilumab or infliximab compared with standard of care in hospitalised patients with COVID-19 (CATALYST): a randomised, multicentre, multi-arm, multistage, open-label, adaptive, phase 2, proof-of-concept trial. The Lancet Respiratory Medicine. 2021 Dec;S2213260021004604.
- 328. Lopardo G, Belloso WH, Nannini E, Colonna M, Sanguineti S, Zylberman V, et al. RBD-specific polyclonal F(ab')2 fragments of equine antibodies in patients with moderate to severe COVID-19 disease: A randomized, multicenter, double-blind, placebo-controlled, adaptive phase 2/3 clinical trial. EClinicalMedicine. 2021 Apr;100843.
- 329. Esquivel-Moynelo I, Perez-Escribano J, Duncan-Robert Y, Vazque-Blonquist D, Bequet-Romero M, Baez-Rodriguez L, et al. Effect and safety of combination of interferon alpha-2b and gamma or interferon alpha-2b for negativization of SARS-CoV-2 viral RNA: preliminary results of a randomized controlled clinical trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.07.29.20164251
- 330. Davoudi-Monfared E, Rahmani H, Khalili H, Hajiabdolbaghi M, Salehi M, Abbasian L, et al. Efficacy and safety of interferon beta-1a in treatment of severe COVID-19: a randomized clinical trial [Preprint] MedRxiv 2020. Available from: https://doi.org/10.1101/2020.05.28.20116467.
- 331. Darazam I, Pourhoseingholi M, Shokouhi S, Irvani S, Mokhtari M, Shabani M, et al. Role of Interferon Therapy in Severe COVID-19: The COVIFERON Randomized Controlled Trial. ResearchSquare [Internet]. 2021.
- 332. Darazam I, Hatami F, Rabiei M, Pourhoseingholi M, Shabani M, Shokouhi S, et al. An Investigation Into the Beneficial Effects of High-Dose Interferon beta 1-a, Compared to Low-Dose Interferon Beta 1-a (the base therapeutic regimen) in moderate to severe COVID-19. ResearchSquare [Internet]. 2021.



- 333. Kalil AC, Mehta AK, Patterson TF, Erdmann N, Gomez CA, Jain MK, et al. Efficacy of interferon beta-1a plus remdesivir compared with remdesivir alone in hospitalised adults with COVID-19: a double-bind, randomised, placebo-controlled, phase 3 trial. The Lancet Respiratory Medicine. 2021 Oct;S2213260021003842.
- 334. Ranieri VM, Pettilä V, Karvonen MK, Jalkanen J, Nightingale P, Brealey D, et al. Effect of Intravenous Interferon β-1a on Death and Days Free From Mechanical Ventilation Among Patients With Moderate to Severe Acute Respiratory Distress Syndrome: A Randomized Clinical Trial. JAMA. 2020 Feb 25;323(8):725.
- 335. Castro-Rodriguez JA, Fish EN, Kollmann T, Iturriaga C, Karpievitch Y, Shannon C, et al. Interferon Beta-1α ring prophylaxis to reduce household transmission of SARS-CoV-2: the Containing Coronavirus Disease-19 randomized clinical trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jun [cited 2022 Jul 8]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.06.13.22276369
- 336. Monk PD, Marsden RJ, Tear VJ, Brookes J, Batten TN, Mankowski M, et al. Safety and efficacy of inhaled nebulised interferon beta-1a (SNG001) for treatment of SARS-CoV-2 infection: a randomised, double-blind, placebo-controlled, phase 2 trial. Lancet Respir Med 2020; published online 12 November 2020. Available from: https://doi.org/10.1016/S2213-2600(20)30511-7.
- 337. Rahmani H, Davoudi-Monfared E, Nourian A, Khalili H, Hajizadeh N, Jalalabadi NZ, et al. Interferon β-1b in treatment of severe COVID-19: a randomized clinical trial. Int Immunopharmacol 2020;88:106903. Available from: https://doi.org/10.1016/j.intimp.2020.106903.
- 338. Tam AR, Zhang RR, Lung KC, Liu R, Leung KY, Liu D, et al. Early treatment of high-risk hospitalized COVID-19 patients with a combination of interferon beta-1b and remdesivir: a phase 2 open-label randomized controlled trial. Clinical Infectious Diseases. 2022 Jun 28;ciac523.
- 339. Myasnikov AL, Berns SA, Talyzin PA, Ershov FI. Interferon gamma in the treatment of patients with moderate COVID-19. Voprosy virusologii. 2021 Mar 7;66(1):47–54.
- 340. Fu W, Yan L, Liu L, Hu H, Cheng X, Liu P, et al. An open-label, randomized trial of the combination of IFN-κ plus TFF2 with standard care in the treatment of patients



World Health Organization

with moderate COVID-19. EclinicalMedicine 2020;27:100547. Available from: https://doi.org/10.1016/j.eclinm.2020.100547.

- 341. Chahla RE, Medina Ruiz L, Ortega ES, Morales MF, Barreiro F, George A, et al. A Randomized Trial - Intensive Treatment Based in Ivermectin and Iota-carrageenan as Pre-exposure Prophylaxis for COVID-19 in Healthcare Agents [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Mar [cited 2021 Apr 2]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.03.26.21254398
- 342. Figueroa JM, Lombardo M, Dogliotti A, Flynn LP, Giugliano RP, Simonelli G, et al. Efficacy of a nasal spray containing Iota-Carrageenan in the prophylaxis of COVID-19 in hospital personnel dedicated to patients care with COVID-19 disease A pragmatic multicenter, randomized, double-blind, placebo-controlled trial (CARR-COV-02) [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Apr [cited 2021 Apr 20]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.04.13.21255409
- 343. Ojeda RA. Clinical study to verify the effectiveness and safety of the modified isothymol or carvacrol compound against SARS-CoV-2 in COVID-19 patients [Internet]. In Review; 2022 Jul [cited 2022 Aug 30]. Available from: <u>https://www.researchsquare.com/article/rs-1809364/v1</u>
- 344. Kumar S, de Souza R, Nadkar M, Guleria R, Trikha A, Joshi SR, Loganathan S, Vaidyanathan S, Marwah A, and Athalye S. A Two-Arm, Randomized, Controlled, Multi-Centric, Open-Label Phase-2 Study to Evaluate the Efficacy and Safety of Itolizumab in Moderate to Severe ARDS Patients Due to COVID-19. [Preprint]. Allergy and Immunology 2020. https://doi.org/10.1101/2020.12.01.20239574.
- 345. Shouman W., Nafae M., Awad Hegazy A., et al. Use of Ivermectin as a potential chemoprophylaxis for COVID-19 in Egypt : A Randomised clinical trial Journal of Clinical and Diagnostic Research, doi:10.7860/JCDR/2020/46795.0000
- 346. Chowdhury ATMM, Shahbaz M, Karim MR, Islam J, Guo D, He S. A randomized trial of ivermectin-doxycycline and hydroxychloroquine-azithromycin therapy on COVID19 patients [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-38896/v1.
- 347. Podder C, Chowdhury N, Sina M, Haque W. Outcome of ivermectin treated mild to moderate COVID-19 cases: a single-centre, open-label, randomised controlled study



[Internet]. IMC J Med Sci 2020;14(2):002. Available from:

http://www.imcjms.com/registration/journal_abstract/353

- 348. Hashim HA, Maulood MF, Rasheed AM, Fatak DF, Kabah KK, Abdulamir AS. Controlled randomized clinical trial on using ivermectin with doxycycline for treating COVID-19 patients in Baghdad, Iraq [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.10.26.20219345.
- 349. Mahmud R, Rahman MdM, Alam I, Ahmed KGU, Kabir AKMH, Sayeed SKJB, et al. Ivermectin in combination with doxycycline for treating COVID-19 symptoms: a randomized trial. J Int Med Res. 2021 May;49(5):030006052110135.
- 350. Elgazzar A, Hany B, Youssef SA, Hafez M, Moussa H. Efficacy and safety of ivermectin for treatment and prophylaxis of COVID-19 pandemic [Preprint]. ResearchSquare 2020. Available from: <u>https://doi.org/10.21203/rs.3.rs-100956/v1</u>.
- 351. Krolewiecki A, Lifschitz A, Moragas M, Travacio M, Valentini R, Alonso DF, et al. Antiviral effect of high-dose ivermectin in adults with COVID-19: A proof-of-concept randomized trial. EClinicalMedicine. 2021 Jul;37:100959.
- 352. Niaee MS, Gheibi N, Namdar P, Allami A, Zolghadr L, Javadi A, Amin Karampour, et al. 2020. Ivermectin as an adjunct treatment for hospitalized adult COVID-19 patients: a randomized multi-center clinical trial [Preprint]. ResearchSquare 2020. https://doi.org/10.21203/rs.3.rs-109670/v1.
- 353. Sabeena A, Karim MM, Ross ag, Hossain ms, Clemens jd, Sumiya MK, Phru CS, et al. A Five Day Course of Ivermectin for the Treatment of COVID-19 May Reduce the Duration of Illness. International Journal of Infectious Diseases 2020. S1201971220325066. https://doi.org/10.1016/j.ijid.2020.11.191.
- 354. Chaccour C, Casellas A, Blanco-Di Matteo A, Pineda I, Fernandez-Montero A, Ruiz-Castillo P, et al. The effect of early treatment with ivermectin on viral load, symptoms and humoral response in patients with non-severe COVID-19: A pilot, doubleblind, placebo-controlled, randomized clinical trial. EClinicalMedicine. 2021 Jan;100720.
- Zeeshan Khan Chachar A, Ahmad Khan K, Asif M, Tanveer K, Khaqan A, Basri R. Effectiveness of Ivermectin in SARS-CoV-2/COVID-19 Patients. ijSciences. 2020;9(09):31–5.
- 356. Babalola OE, Bode CO, Ajayi AA, Alakaloko FM, Akase IE, Otrofanowei E, et al. Ivermectin shows clinical benefits in mild to moderate COVID19: a randomized

PAHOS



controlled double-blind, dose-response study in Lagos. QJM: An International Journal of Medicine. 2021 Feb 18;hcab035.

- 357. Kirti R, Roy R, Pattadar C, Raj R, Agarwal N, Biswas B, et al. Ivermectin as a potential treatment for mild to moderate COVID-19: A double blind randomized placebo-controlled trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Jan [cited 2021 Jan 11]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.01.05.21249310</u>
- 358. Mohan A, Tiwari P, Suri T, Mittal S, Patel A, Jain A, et al. Ivermectin in mild and moderate COVID-19 (RIVET-COV): a randomized, placebo-controlled trial [Internet]. In Review; 2021 Feb [cited 2021 Jun 5]. Available from: https://www.researchsquare.com/article/rs-191648/v1
- 359. Shahbaznejad L, Davoudi A, Eslami G, Markowitz JS, Navaeifar MR, Hosseinzadeh F, et al. Effect of ivermectin on COVID-19: A multicenter double-blind randomized controlled clinical trial. Clinical Therapeutics. 2021 May;S0149291821002010.
- 360. Hill A, Abdulamir A, Ahmed S, Asghar A, Babalola OE, Basri R, et al. Metaanalysis of randomized trials of ivermectin to treat SARS-CoV-2 infection [Internet]. In Review; 2021 Jan [cited 2021 Jan 29]. Available from: <u>https://www.researchsquare.com/article/rs-148845/v1</u>
- 361. Samaha AA, Mouawia H, Fawaz M, Hassan H, Salami A, Bazzal AA, et al. Effects of a Single Dose of Ivermectin on Viral and Clinical Outcomes in Asymptomatic SARS-CoV-2 Infected Subjects: A Pilot Clinical Trial in Lebanon. Viruses. 2021 May 26;13(6):989.
- 362. Efficacy of Ivermectin in COVID-19 Patients with Mild to Moderate Disease
 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Feb [cited 2021 Mar 9].
 Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.02.02.21250840</u>
- 363. Okumuş N, Demirtürk N, Çetinkaya RA, Güner R, Avcı İY, Orhan S, et al. Evaluation of the effectiveness and safety of adding ivermectin to treatment in severe COVID-19 patients. BMC Infect Dis. 2021 Dec;21(1):411.
- 364. López-Medina E, López P, Hurtado IC, Dávalos DM, Ramirez O, Martínez E, et al. Effect of Ivermectin on Time to Resolution of Symptoms Among Adults With Mild COVID-19: A Randomized Clinical Trial. JAMA [Internet]. 2021 Mar 4 [cited 2021 Mar 9]; Available from: <u>https://jamanetwork.com/journals/jama/fullarticle/2777389</u>



ww.paho.org/coronavirus

- 365. Pott-Junior H, Bastos Paoliello MM, de Queiroz Constantino Miguel A, da Cunha AF, de Melo Freire CC, Neves FF, et al. Use of ivermectin in the treatment of Covid-19: a pilot trial. Toxicology Reports. 2021 Mar;S2214750021000445.
- 366. Kishoria N, Mathur SL, Parmar V, Kaur RJ, Agarwal H, Parihar BS, et al. Ivermectin as Adjuvant to Hydroxycholoroquine in Patients Resistant to Standard Treatment for SARS-CoV-2: Results of an Open-label Randomized Clinical Study. PIJR. 2020 Aug 15;1–4.
- 367. Abd-Elsalam S, Noor RA, Badawi R, Khalaf M, Esmail ES, Soliman S, et al. Clinical Study Evaluating the Efficacy of Ivermectin in COVID-19 Treatment: A Randomized Controlled Study. J Med Virol. 2021 Jun 2;jmv.27122.
- 368. Biber A, Harmelin G, Lev D, Ram L, Shaham A, Nemet I, et al. The effect of ivermectin on the viral load and culture viability in early treatment of nonhospitalized patients with mild COVID-19 – a double-blind, randomized placebo-controlled trial. International Journal of Infectious Diseases. 2022 Sep;122:733–40.
- Faisal R, Shah SFA, Hussain M. Potential use of azithromycin alone and in combination with ivermectin in fighting against the symptoms of COVID-19. TPMJ. 2021 May 10;28(05):737–41.
- 370. Vallejos J, Zoni R, Bangher M, Villamandos S, Bobadilla A, Plano F, et al. Ivermectin to prevent hospitalizations in patients with COVID-19 (IVERCOR-COVID19) a randomized, double-blind, placebo-controlled trial. BMC Infect Dis. 2021 Dec;21(1):635.
- 371. Buonfrate D, Chesini F, Martini D, Roncaglioni MC, Fernandez MLO, Alvisi MF, et al. High dose ivermectin for the early treatment of COVID-19 (COVER study): a randomised, double-blind, multicentre, phase II, dose-finding, proof of concept clinical trial. International Journal of Antimicrobial Agents. 2022 Jan;106516.
- 372. Manomaipiboon A, Pholtawornkulchai K, Pupipatpab S, Suraamornkul S, Maneerit J, Ruksakul W, et al. Efficacy and safety of ivermectin in the treatment of mild-to-moderate COVID-19 infection: A randomized, double blind, placebo, controlled trial [Internet]. In Review; 2022 Feb [cited 2022 Feb 15]. Available from: <u>https://www.researchsquare.com/article/rs-1290999/v1</u>
- 373. Lim SCL, Hor CP, Tay KH, Mat Jelani A, Tan WH, Ker HB, et al. Efficacy of Ivermectin Treatment on Disease Progression Among Adults With Mild to Moderate

PAHO®





COVID-19 and Comorbidities: The I-TECH Randomized Clinical Trial. JAMA Intern Med [Internet]. 2022 Feb 18 [cited 2022 Feb 22]; Available from: https://jamanetwork.com/journals/jamainternalmedicine/fullarticle/2789362

- 374. Reis G, Silva EASM, Silva DCM, Thabane L, Milagres AC, Ferreira TS, et al. Effect of Early Treatment with Ivermectin among Patients with Covid-19. N Engl J Med. 2022 Mar 30;NEJMoa2115869.
- 375. Rocha C de la, Cid-Lopez MA, Venegas-Lopez BI, Gómez-Mendez SC, Sánchez-Ortiz A, Pérez-Ríos AM, et al. Ivermectin compared with placebo in the clinical evolution of Mexican patients with asymptomatic and mild COVID-19: a randomized clinical trial [Internet]. In Review; 2022 May [cited 2022 Jun 8]. Available from: https://www.researchsquare.com/article/rs-1640339/v1
- 376. Accelerating COVID-19 Therapeutic Interventions and Vaccines (ACTIV)-6 Study Group, Naggie S. Ivermectin for Treatment of Mild-to-Moderate COVID-19 in the Outpatient Setting: A Decentralized, Placebo-controlled, Randomized, Platform Clinical Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jun [cited 2022 Jul 26]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.06.10.22276252</u>
- 377. Rezai MS, Ahangarkani F, Hill A, Ellis L, Mirchandani M, Davoudi A, et al. Non-effectiveness of Ivermectin on Inpatients and Outpatients With COVID-19; Results of Two Randomized, Double-Blinded, Placebo-Controlled Clinical Trials. Front Med. 2022 Jun 16;9:919708.
- 378. Angkasekwinai N, Rattanaumpawan P, Chayakulkeeree M, Phoompoung P, Koomanachai P, Chantarasut S, et al. Safety and Efficacy of Ivermectin for the Prevention and Treatment of COVID-19: A Double-Blinded Randomized Placebo-Controlled Study. Antibiotics. 2022 Jun 12;11(6):796.
- 379. Mirahmadizadeh A, Semati A, Heiran A, Ebrahimi M, Hemmati A, Karimi M, et al. Efficacy of single-dose and double-dose ivermectin early treatment in preventing progression to hospitalization in mild COVID -19: A multi-arm, parallel-group randomized, double-blind, placebo-controlled trial. Respirology. 2022 Jun 23;resp.14318.
- 380. George B, Moorthy M, Kulkarni U, Selvarajan S, Rupali P, Christopher DJ, et al. Single Dose of Ivermectin is not Useful in Patients with Hematological Disorders and COVID-19 Illness: A Phase II B Open Labelled Randomized Controlled Trial. Indian J

PAHOS



Hematol Blood Transfus [Internet]. 2022 May 27 [cited 2022 Jul 1]; Available from: https://link.springer.com/10.1007/s12288-022-01546-w

- 381. Schilling WH, Jittamala P, Watson JA, Ekkapongpisit M, Siripoon T, Ngamprasertchai T, et al. Pharmacometric assessment of the *in vivo* antiviral activity of ivermectin in early symptomatic COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jul [cited 2022 Jul 26]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.07.15.22277570
- 382. Nimitvilai S, Suputtamongkol Y, Poolvivatchaikarn U, Rassamekulthana D, Rongkiettechakorn N, Mungaomklang A, et al. A randomized controlled trial of combined ivermectin and zinc sulfate versus combined hydroxychloroquine, darunavir/ritonavir, and zinc sulfate among adult patients with asymptomatic or mild coronavirus-19 infection. J Global Infect Dis. 2022;14(2):69.
- 383. Aref ZF, Bazeed SEES, Hassan MH, Hassan AS, Rashad A, Hassan RG, et al. Clinical, Biochemical and Molecular Evaluations of Ivermectin Mucoadhesive Nanosuspension Nasal Spray in Reducing Upper Respiratory Symptoms of Mild COVID-19. Int J Nanomedicine. 2021;16:4063–72.
- 384. Sakoulas G, Geriak M, Kullar R, Greenwood K, Habib M, Vyas A, et al. Intravenous immunoglobulin (IVIG) significantly reduces respiratory morbidity in COVID-19 pneumonia: a prospective randomized trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.07.20.20157891.
- 385. Gharebaghi N, Nejadrahim R, Mousavi SJ, Sadat-Ebrahimi S-R, Hajizadeh R. The use of intravenous immunoglobulin gamma for the treatment of severe coronavirus disease 2019: a randomised placebo-controlled double-blind clinical trial [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-40899/v2.
- 386. Tabarsi P, Barati S, Jamaati H, Haseli S, Marjani M, Moniri A, et al. Evaluating the effects of intravenous immunoglobulin (IVIG) on the management of severe COVID-19 cases: a randomized controlled trial [Internet]. Int Immunopharmacol 2020:107205. Available from: <u>https://doi.org/10.1016/j.intimp.2020.107205</u>.
- 387. R S R, Barge VB, Darivenula AK, Dandu H, Kartha RR, Bafna V, et al. A Phase II Safety and Efficacy Study on Prognosis of Moderate Pneumonia in COVID-19 patients with Regular Intravenous Immunoglobulin Therapy. The Journal of Infectious Diseases. 2021 Feb 15;jiab098.

PAHO®



495

388. Haran JP, Zheng Y, Knobil K, Palma NA, Lawrence JF, Wingertzahn MA. Targeting the Microbiome With KB109 in Outpatients with Mild to Moderate COVID-19 Reduced Medically Attended Acute Care Visits and Improved Symptom Duration in Patients With Comorbidities [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Mar [cited 2021 Apr 5]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2021.03.26.21254422

- 389. Fiorentino G, Coppola A, Izzo R, Annunziata A, Bernardo M, Lombardi A, et al. Effects of adding L-arginine orally to standard therapy in patients with COVID-19: A randomized, double-blind, placebo-controlled, parallel-group trial. Results of the first interim analysis. EClinicalMedicine. 2021 Sep;101125.
- 390. Endam LM, Tremblay C, Filali A, Desrosiers MY. Intranasal Application of Lactococcus Lactis W 136 Bacteria Early in SARS-Cov-2 Infection May Have a Beneficial Immunomodulatory Effect: A Proof-of-concept Study [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Apr [cited 2021 May 3]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.04.18.21255699</u>
- 391. Algahtani FD, Elabbasy MT, Samak MA, Adeboye AA, Yusuf RA, Ghoniem ME. The Prospect of Lactoferrin Use as Adjunctive Agent in Management of SARS-CoV-2 Patients: A Randomized Pilot Study. Medicina. 2021 Aug 19;57(8):842.
- 392. Hu K, Wang M, Zhao Y, Zhang Y, Wang T, Zheng Z, et al. A small-scale medication of leflunomide as a treatment of COVID-19 in an open-label blank-controlled clinical trial [Internet]. Virol Sin 2020. Available from: https://doi.org/10.1007/s12250-020-00258-7.
- 393. Wang M, Zhao Y, Hu W, Zhao D, Zhang Y, Wang T, et al. Treatment of COVID-19 patients with prolonged post-symptomatic viral shedding with leflunomide -- a singlecenter, randomized, controlled clinical trial [Internet]. Clin Infect Dis 2020; ciaa1417. Available from: <u>https://doi.org/10.1093/cid/ciaa1417</u>.
- 394. Temesgen Z, Burger CD, Baker J, Polk C, Libertin CR, Kelley CF, et al. Lenzilumab in hospitalised patients with COVID-19 pneumonia (LIVE-AIR): a phase 3, randomised, placebo-controlled trial. The Lancet Respiratory Medicine. 2021 Dec;S221326002100494X.
- 395. Roostaei A, Meybodi Z, Mosavinasab S, Karimzadeh I, Sahebnasagh A, Gholinataj M, et al. Efficacy and Safety of Levamisole Treatment in Clinical

PAHO®



496

Presentations of Patients With COVID-19: A Double-Blind, Randomized, Controlled Trial. ResearchSquare [Internet]. 2021.

- 396. Asgardoon MH, koochak HE, Kazemi-Galougahi MH, Dehnavi AZ, Khodaei B, Behkar A, et al. Efficacy of Levamisole with Standard Care Treatment vs Standard Care in Clinical Presentations of Non-Hospitalized Patients with COVID-19: A Randomized Clinical Trial [Internet]. In Review; 2021 Nov [cited 2021 Dec 6]. Available from: <u>https://www.researchsquare.com/article/rs-964097/v1</u>
- 397. Lomakin NV, Bakirov BA, Protsenko DN, Mazurov VI, Musaev GH, Moiseeva OM, et al. The efficacy and safety of levilimab in severely ill COVID-19 patients not requiring mechanical ventilation: results of a multicenter randomized double-blind placebo-controlled phase III CORONA clinical study. Inflamm Res [Internet]. 2021 Sep 29 [cited 2021 Oct 12]; Available from: https://link.springer.com/10.1007/s00011-021-01507-5
- 398. Abuhasira R, Ayalon-Dangur I, Zaslavsky N, Koren R, Keller M, Dicker D, et al. A Randomized Clinical Trial of Linagliptin vs. Standard of Care in Patients Hospitalized With Diabetes and COVID-19. Front Endocrinol. 2021 Dec 22;12:794382.
- 399. Guardado-Mendoza R, Garcia-Magaña MA, Martínez-Navarro LJ, Macías-Cervantes HE, Aguilar-Guerrero R, Suárez-Pérez EL, et al. Effect of linagliptin plus insulin in comparison to insulin alone on metabolic control and prognosis in hospitalized patients with SARS-CoV-2 infection. Sci Rep. 2022 Dec;12(1):536.
- 400. Spuch C, López-García M, Rivera-Baltanás T, Cabrera-Alvargonzález JJ, Gadh S, Rodrigues-Amorim D, et al. Efficacy and Safety of Lithium Treatment in SARS-CoV-2 Infected Patients. Front Pharmacol. 2022 Apr 14;13:850583.
- 401. Cao B, Wang Y, Wen D, Liu W, Wang J, Fan G, et al. A trial of lopinavirritonavir in adults hospitalized with severe Covid-19. N Engl J Med 2020; 382(19): 1787–99. Available from: https://doi.org/10.1056/NEJMoa2001282.
- 402. Li Y, Xie Z, Lin W, Cai W, Wen C, Guan Y, et al. Efficacy and safety of lopinavir/ritonavir or arbidol in adult patients with mild/moderate COVID-19: an exploratory randomized controlled trial [Internet]. Clin Advance 2020, published online 4 May 2020. Available from: https://doi.org/10.1016/j.medj.2020.04.001.
- 403. RECOVERY Collaborative Group. Lopinavir–ritonavir in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform

PAHO®



trial. Lancet 2020; 396 (10259): 1345-52. Available from: https://doi.org/10.1016/S0140-6736(20)32013-4.

- Zheng F, Zhou Y, Zhou Z, Ye F, Huang B, Huang Y, et al. A novel protein drug, novaferon, as the potential antiviral drug for COVID-19 [Preprint]. MedRxiv 2020.
 Available from: https://doi.org/10.1101/2020.04.24.20077735.
- 405. Chen Y-K, Huang Y-Q, Tang S-Q, Xu X-L, Zeng Y-M, He X-Q, et al. Comparative effectiveness and safety of ribavirin plus interferon-alpha, lopinavir/ritonavir plus interferon-alpha and ribavirin plus lopinavir/ritonavir plus interferon-alpha in patients with mild to moderate novel coronavirus pneumonia: results of a randomized, open-labeled prospective study [Preprint]. 2020. Available from SSRN: <u>https://doi.org/10.2139/ssrn.3576905</u>.
- 406. Shahnaz Sali, Davood Yadegarinia, Sara Abolghasemi, Shabnam Tehrani, Babak Gharaei, Neda Khabiri, et al. Comparison of the Efficacy of Sofosbuvir and Kaletra on Outcome of Covid-19. Is Sofosbuvir A Potential Treatment For COVID-19? Novelty in Biomedicine [Internet]. 2021
- 407. Purwati, Budiono, Rachman BE, Yulistiani, Miatmoko A, Nasronudin, et al. A Randomized, Double-Blind, Multicenter Clinical Study Comparing the Efficacy and Safety of a Drug Combination of Lopinavir/Ritonavir-Azithromycin, Lopinavir/Ritonavir-Doxycycline, and Azithromycin-Hydroxychloroquine for Patients Diagnosed with Mild to Moderate COVID-19 Infections. Huyut Z, editor. Biochemistry Research International. 2021 Feb 9;2021:1–12.
- 408. Kasgari HA, Moradi S, Shabani AM, Babamahmoodi F, Badabi ARD, Davoudi L, et al. Evaluation of the efficacy of sofosbuvir plus daclatasvir in combination with ribavirin for hospitalized COVID-19 patients with moderate disease compared with standard care: a single-centre, randomized controlled trial. J Antimicrob Chemother 2020; 75(11):3373-78. Available from: <u>https://doi.org/10.1093/jac/dkaa332</u>.
- 409. Yadollahzadeh M, Eskandari M, Roham M, Zamani F, Laali A, Kalantari S, et al. Evaluation of Sovodak (Sofosbuvir/Daclatasvir) Treatment Outcome in COVID-19 Patient's Compared with Kaletra (Lopinavir/ritonavir): a Randomized Clinical Trial [Internet]. In Review; 2021 Mar [cited 2021 Mar 25]. Available from: <u>https://www.researchsquare.com/article/rs-257762/v1</u>



498

- 410. Labhardt ND, Smit M, Petignat I, Perneger T, Marinosci A, Ustero P, et al. Efficacy of Lopinavir-Ritonavir Prophylaxis for Individuals Exposed to SARS-CoV-2: The COPEP Pragmatic Open-Label, Cluster Randomized Trial. SSRN Journal [Internet]. 2021 [cited 2021 Jul 14]; Available from: <u>https://www.ssrn.com/abstract=3878828</u>
- 411. Papachristofilou A, Finazzi T, Blum A, Zehnder T, Zellweger N, Lustenberger J, et al. Low-Dose Radiation Therapy for Severe COVID-19 Pneumonia: A Randomized Double-Blind Study. International Journal of Radiation Oncology*Biology*Physics. 2021 Mar;S036030162100239X.
- 412. Ganesan G, Ponniah S, Sundaram V, Kumar Marimuthu P, Pitchaikannu V, Chandrasekaran M, et al. Whole lung Irradiation as a Novel treatment for COVID-19: Final Results of the Prospective Randomized trial (WINCOVID trial). Radiotherapy and Oncology. 2021 Dec;S0167814021090721.
- 413. Singh P, Mandal A, Singh D, Kumar S, Kumar A, Rakesh A, et al. Interim Analysis of Impact of Adding Low Dose Pulmonary Radiotherapy to Moderate COVID-19 Pneumonia Patients: IMpaCt-RT Study. Front Oncol. 2022 Mar 29;12:822902.
- 414. Cremer PC, Abbate A, Hudock K, McWilliams C, Mehta J, Chang SY, et al. Mavrilimumab in patients with severe COVID-19 pneumonia and systemic hyperinflammation (MASH-COVID): an investigator initiated, multicentre, double-blind, randomised, placebo-controlled trial. The Lancet Rheumatology. 2021 Mar;S2665991321000709.
- 415. Farnoosh G, Akbariqomi M, Badri T, Bagheri M, Izadi M, Saeedi-Boroujeni A, et al. Efficacy of a Low Dose of Melatonin as an Adjunctive Therapy in Hospitalized Patients with COVID-19: A Randomized, Double-blind Clinical Trial. Archives of Medical Research. 2021 Jun;S0188440921001417.
- Davoodian N, Sharifimood F, Salarbashi D, Elyasi S, Baniasad A, Bejestani FS. The Effect of Melatonin as an Adjuvant Therapy on COVID-19: A Randomized Clinical Trial. SSRN Journal [Internet]. 2021 [cited 2021 Jul 14]; Available from: <u>https://www.ssrn.com/abstract=3878090</u>
- Alizadeh Z, Keyhanian N, Ghaderkhani S, Dashti-Khavidaki S, Shokouhi
 Shoormasti R, Pourpak Z. A Pilot Study on Controlling Coronavirus Disease 2019
 (COVID-19) Inflammation Using Melatonin Supplement. IJAAI [Internet]. 2021 Aug 11

PAHOS



[cited 2021 Aug 30]; Available from: https://publish.knepublishing.com/index.php/IJAAI/article/view/6959

- 418. Mousavi SA, Heydari K, Mehravaran H, Saeedi M, Alizadeh-Navaei R, Hedayatizadeh-Omran A, et al. Melatonin effects on sleep quality and outcomes of COVID-19 patients: An open-label, randomized, controlled trial. J Med Virol. 2021 Sep 8;jmv.27312.
- 419. Hasan ZT, Atrakji DrMQYMAA, Mehuaiden DrAK. The Effect of Melatonin on Thrombosis, Sepsis and Mortality Rate in COVID-19 Patients. International Journal of Infectious Diseases. 2021 Oct;S1201971221007980.
- 420. García-García I, Seco-Meseguer E, Ruiz-Seco P, Navarro-Jimenez G, Martínez-Porqueras R, Espinosa-Díaz M, et al. Melatonin in the Prophylaxis of SARS-CoV-2 Infection in Healthcare Workers (MeCOVID): A Randomised Clinical Trial. JCM. 2022 Feb 21;11(4):1139.
- 421. Alizadeh Z, Keyhanian N, Ghaderkhani S, Dashti-Khavidaki S, Shokouhi Shoormasti R, Pourpak Z. A Pilot Study on Controlling Coronavirus Disease 2019 (COVID-19) Inflammation Using Melatonin Supplement. IJAAI [Internet]. 2021 Aug 11 [cited 2021 Aug 30]; Available from: <u>https://publish.knepublishing.com/index.php/IJAAI/article/view/6959</u>
- Fogleman C, Cohen D, Mercier A, Farrell D, Rutz J, Bresz K, et al. A Pilot of a Randomized Control Trial of Melatonin and Vitamin C for Mild-to-Moderate COVID-19. J Am Board Fam Med. 2022 Jul;35(4):695–707.
- 423. Guzman-Esquivel J, Galvan-Salazar HR, Guzman-Solorzano HP, Cuevas-Velazquez AC, Guzman-Solorzano JA, Mokay-Ramirez KA, et al. Efficacy of the use of mefenamic acid combined with standard medical care vs. standard medical care alone for the treatment of COVID-19: A randomized double-blind placebo-controlled trial. Int J Mol Med. 2022 Mar;49(3):29.
- 424. Shu L, Niu C, Li R, Huang T, Wang Y, Huang M, et al. Treatment of severe COVID-19 with human umbilical cord mesenchymal stem cells. Stem Cell Res Ther 2020;11(1):361. Available from: https://doi.org/10.1186/s13287-020-01875-5.
- 425. Shi L, Huang H, Lu X, Yan X, Jiang X, Xu R, et al. Treatment with human umbilical cord-derived mesenchymal stem cells for COVID-19 patients with lung



damage: a randomised, double-blind, placebo controlled phase 2 trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.10.15.20213553.

- 426. Lanzoni G, Linetsky E, Correa D, Cayetano SM, Marttos AC, Alvarez RA, et al. Umbilical cord mesenchymal stem cells for COVID-19 ARDS: a double blind, phase 1/2a, randomized controlled trial [Preprint]. 2020. Available from SSRN: <u>https://doi.org/10.2139/ssrn.3696875</u>.
- 427. Dilogo IH, Aditianingsih D, Sugiarto A, Burhan E, Damayanti T, Sitompul PA, et al. Umbilical cord mesenchymal stromal cells as critical COVID -19 adjuvant therapy: A randomized controlled trial. STEM CELLS Transl Med. 2021 Jun 8;sctm.21-0046.
- 428. Zhu R, Yan T, Feng Y, Liu Y, Cao H, Peng G, et al. Mesenchymal stem cell treatment improves outcome of COVID-19 patients via multiple immunomodulatory mechanisms. Cell Res [Internet]. 2021 Oct 26 [cited 2021 Nov 4]; Available from: https://www.nature.com/articles/s41422-021-00573-y
- 429. Fathi-Kazerooni M, Fattah-Ghazi S, Darzi M, Makarem J, Nasiri R, Salahshour F, et al. Safety and efficacy study of allogeneic human menstrual blood stromal cells secretome to treat severe COVID-19 patients: clinical trial phase I & II. Stem Cell Res Ther. 2022 Dec;13(1):96.
- 430. Rebelatto CLK, Senegaglia AC, Franck CL, Daga DR, Shigunov P, Stimamiglio MA, et al. Safety and long-term improvement of mesenchymal stromal cell infusion in critically COVID-19 patients: a randomized clinical trial. Stem Cell Res Ther. 2022 Dec;13(1):122.
- 431. Karyana M, Djaharuddin I, Rif'ati L, Arif M, Choi MK, Angginy N, et al. Safety of DW-MSC infusion in patients with low clinical risk COVID-19 infection: a randomized, double-blind, placebo-controlled trial. Stem Cell Res Ther. 2022 Dec;13(1):134.
- 432. Farkhad NK, Sedaghat A, Reihani H, Moghadam AA, Moghadam AB, Ghaebi NK, et al. Mesenchymal Stem Cell therapy for COVID-19-induced ARDS patients. A successful phase1, randomized, control-placebo group, clinical trial [Internet]. In Review; 2022 Jan [cited 2022 Sep 7]. Available from: <u>https://www.researchsquare.com/article/rs-1240880/v1</u>
- 433. Reis G, dos Santos Moreira Silva EA, Medeiros Silva DC, Thabane L, Cruz Milagres A, Ferreira TS, et al. Effect of early treatment with metformin on risk of



501

emergency care and hospitalization among patients with COVID-19: The TOGETHER randomized platform clinical trial. The Lancet Regional Health - Americas. 2022 Feb;6:100142.

- 434. Ventura-López C, Cervantes-Luevano K, Aguirre-Sánchez JS, Flores-Caballero JC, Alvarez-Delgado C, Bernaldez-Sarabia J, et al. Treatment with metformin glycinate reduces SARS-CoV-2 viral load: An in vitro model and randomized, double-blind, Phase IIb clinical trial. Biomedicine & Pharmacotherapy. 2022 Aug;152:113223.
- 435. Hamidi-Alamdari D, Hafizi-Lotfabadi S, Bagheri-Moghaddam A, Safari H, Mozdourian M, Javidarabshahi Z, et al. Methylene Blue for Treatment of Hospitalized COVID-19 Patients: A Randomized, Controlled, Open-label Clinical Trial, Phase 2. Rev Invest Clin. 2021;73(3):190–8.
- 436. Borges M, Borges M, Borges J, Bastidas R. Estudio Experimental: Manejo del Metisoprinol en Pacientes con COVID-19. uct. 2020 Aug 10;24(103):41–50.
- Clemente-Moragón A, Martínez-Milla J, Oliver E, Santos A, Flandes J,
 Fernández I, et al. Metoprolol in Critically Ill Patients With COVID-19. Journal of the
 American College of Cardiology. 2021 Sep;78(10):1001–11.
- 438. Kazempour M, Izadi H, Chouhdari A, Rezaeifard M. Anti-inflammatory Effect of Metronidazole in Hospitalized Patients with Pneumonia due to COVID-19. Iran J Pharm Res. 2021;20(3):532–40.
- Painter WP, Holman W, Bush JA, Almazedi F, Malik H, Eraut NCJE, et al. Human Safety, Tolerability, and Pharmacokinetics of a Novel Broad-Spectrum Oral Antiviral Compound, Molnupiravir, with Activity Against SARS-CoV-2 [Internet]. Infectious Diseases (except HIV/AIDS); 2020 Dec [cited 2020 Dec 30]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2020.12.10.20235747</u>
- Khoo SH, FitzGerald R, Fletcher T, Ewings S, Jaki T, Lyon R, et al. Optimal dose and safety of molnupiravir in patients with early SARS-CoV-2: a phase 1, dose-escalating, randomised controlled study [Internet]. Pharmacology and Therapeutics; 2021 May [cited 2021 May 14]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.05.03.21256309
- 441. Fischer WA, Eron JJ, Holman W, Cohen MS, Fang L, Szewczyk LJ, 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 Dec 23;eabl7430.

- 442. Jayk Bernal A, Gomes da Silva MM, Musungaie DB, Kovalchuk E, Gonzalez A, Delos Reyes V, et al. Molnupiravir for Oral Treatment of Covid-19 in Nonhospitalized Patients. N Engl J Med. 2021 Dec 16;NEJMoa2116044.
- 443. Tippabhotla SK, Lahiri DrS, D RR, Kandi C, V NP. Efficacy and Safety of Molnupiravir for the Treatment of Non-Hospitalized Adults With Mild COVID-19: A Randomized, Open-Label, Parallel-Group Phase 3 Trial. SSRN Journal [Internet]. 2022 [cited 2022 Mar 7]; Available from: <u>https://www.ssrn.com/abstract=4042673</u>
- 444. Zou R, Peng L, Shu D, Zhao L, Lan J, Tan G, et al. Antiviral Efficacy and Safety of Molnupiravir Against Omicron Variant Infection: A Randomized Controlled Clinical Trial. Front Pharmacol. 2022 Jun 15;13:939573.
- Khoo SH, FitzGerald R, Saunders G, Middleton C, Ahmad S, Edwards CJ, et al. A Randomised -Controlled Phase 2 trial of Molnupiravir in Unvaccinated and Vaccinated Individuals with Early SARS-CoV-2 [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jul [cited 2022 Aug 11]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2022.07.20.22277797

- Arribas JR, Bhagani S, Lobo SM, Khaertynova I, Mateu L, Fishchuk R, et al.
 Randomized Trial of Molnupiravir or Placebo in Patients Hospitalized with Covid-19.
 NEJM Evidence [Internet]. 2022 Jan 25 [cited 2022 Aug 19];1(2). Available from: https://evidence.nejm.org/doi/10.1056/EVIDoa2100044
- Caraco Y, Crofoot GE, Moncada PA, Galustyan AN, Musungaie DB, Payne B, et al. Phase 2/3 Trial of Molnupiravir for Treatment of Covid-19 in Nonhospitalized Adults. NEJM Evidence [Internet]. 2022 Jan 25 [cited 2022 Aug 19];1(2). Available from: https://evidence.nejm.org/doi/10.1056/EVIDoa2100043
- 448. Kerget B, Kerget F, Aydın M, Karaşahin Ö. Effect of montelukast therapy on clinical course, pulmonary function, and mortality in patients with COVID-19. Journal of Medical Virology. 2021 Dec 27;jmv.27552.
- Mukhtar K, Qassim S, DanJuma MI, Mohamedali M, Al Farhan H, Khudair MF, El Tayeh AR, et al. On the Possible Beneficial Role for the Regular Use of Potent Mouthwash Solutions as a Preventive Measure for COVID19 Transmission; Invoking the



503

Evolutionary Biology and Game Theory. [Preprint] 2020. https://doi.org/10.1101/2020.11.27.20234997.

- 450. Azmawati MN, Baharom N, Wan Sulaiman W, Rashid ZZ, Wong KK, Ali UK, Othman SN, et al. Early viral clearance among COVID-19 patients when gargling with povidone-iodine and essential oils: A pilot clinical trial. [Preprint] 2020. <u>https://doi.org/10.1101/2020.09.07.20180448</u>.
- 451. Guenezan J, Garcia M, Strasters D, Jousselin C, Lévêque N, Frasca D, et al. Povidone Iodine Mouthwash, Gargle, and Nasal Spray to Reduce Nasopharyngeal Viral Load in Patients With COVID-19: A Randomized Clinical Trial. JAMA Otolaryngol Head Neck Surg [Internet]. 2021 Feb 4 [cited 2021 Feb 14]; Available from: <u>https://jamanetwork.com/journals/jamaotolaryngology/fullarticle/2775984</u>
- 452. Elzein R, Abdel-Sater F, Fakhreddine S, Hanna PA, Feghali R, Hamad H, et al. In vivo evaluation of the virucidal efficacy of Chlorhexidine and Povidone-iodine mouthwashes against salivary SARS-CoV-2 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Mar [cited 2021 Mar 22]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.03.07.21252302
- 453. Santos PS da S, Orcina B da F, Machado RRG, Vilhena FV, Alves LM da C, Zangrando MSR, et al. Beneficial effects of a mouthwash containing an antiviral phthalocyanine derivative on the length of hospital stay for COVID-19 [Internet]. In Review; 2021 Mar [cited 2021 Mar 23]. Available from: https://www.researchsquare.com/article/rs-330173/v1
- 454. Carrouel, Valette, Gadea, Esparcieux, Illes, Langlois, et al. Use of an antiviral mouthwash as an additional barrier measure in the SARS-CoV-2 transmission in adults with asymptomatic to mild COVID-19: A multicenter, randomized, double-blind controlled trial [Internet]. In Review; 2021 Mar [cited 2021 Mar 25]. Available from: https://www.researchsquare.com/article/rs-315468/v1
- 455. Huang YH, Huang JT. Use of chlorhexidine to eradicate oropharyngeal SARS-CoV-2 in COVID-19 patients. J Med Virol. 2021 Apr;jmv.26954.
- 456. Eduardo F de P, Corrêa L, Heller D, Daep CA, Benitez C, Malheiros Z, et al. Salivary SARS-CoV-2 load reduction with mouthwash use: A randomized pilot clinical trial. Heliyon. 2021 Jun;7(6):e07346.



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- 457. Di Domênico MB, Collares K, dos Santos RB, Lenz U, Antunes VP, Godinho V, et al. Hydrogen peroxide as auxiliary treatment for COVID-19: A randomized doubleblind clinical trial. Epidemiol Health. 2021 Aug 3;e2021051.
- 458. Damião Costa D, Brites C, Nunes Vaz S, Souza de Santana D, Dos Santos JN, Cury PR. Chlorhexidine mouthwash reduces the salivary viral load of SARS-CoV-2: a randomized clinical trial. Oral Dis. 2021 Nov 26
- 459. Ferrer MD, Barrueco ÁS, Martinez-Beneyto Y, Mateos-Moreno MV, Ausina-Márquez V, García-Vázquez E, et al. Clinical evaluation of antiseptic mouth rinses to reduce salivary load of SARS-CoV-2. Sci Rep. 2021 Dec;11(1):24392.
- 460. Poleti ML, Gregório D, Bistaffa AGI, Fernandes KBP, Vilhena FV, Santos PS da S, et al. The use of a mouthwash and a dentifrice containing antimicrobial phthalocyanine derivative on the reduction of clinical symptoms of COVID-19: A randomized tripleblinded clinical trial [Internet]. In Review; 2021 Dec [cited 2022 Jan 5]. Available from: <u>https://www.researchsquare.com/article/rs-1139111/v1</u>
- 461. Alemany A, Perez-Zsolt D, Raïch-Regué D, Muñoz-Basagoiti J, Ouchi D, Laporte-Villar C, et al. Cetylpyridinium Chloride Mouthwash to Reduce Shedding of Infectious SARS-CoV-2: A Double-Blind Randomized Clinical Trial. J Dent Res. 2022 Jun 21;002203452211023.
- 462. Barrueco ÁS, Mateos-Moreno MV, Martínez-Beneyto Y, García-Vázquez E, González AC, Ferrero JZ, et al. Effect of Oral Antiseptics in Reducing SARS-CoV-2 Infectivity: Evidence from a Randomized Double-blind Clinical Trial. Emerging Microbes & Infections. 2022 Jul 7;1–23.
- Miller RA, Guru P, Bauer P, Robles J, Tomaszewski C, Overcash JS, et al.
 Clinical Results with a B Cell Activating Anti-CD73 Antibody for the Immunotherapy of COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Sep [cited 2021 Sep 29]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.09.13.21263406
- 464. Sehgal IS, Guleria R, Singh S, Siddiqui MS, Agarwal R. A randomised trial of *Mycobacterium w* in critically ill patients with COVID-19: ARMY-1. ERJ Open Res. 2021 Apr;7(2):00059–2021.
- 465. Alencar JCG de, Moreira CdL, Müller AD, Chaves CE, Fukuhara MA, Silva EA da, Miyamoto MdFS, et al. Double-blind, randomized, placebo-controlled trial with N-



acetylcysteine for treatment of Severe Acute Respiratory Syndrome caused by COVID-19. Clin Infect Dis 2020: ciaa1443. Available from: https://doi.org/10.1093/cid/ciaa1443.

- 466. Gaynitdinova VV, Avdeev SN, Merzhoeva ZM, Berikkhanov ZG-M, Medvedeva IV, Gorbacheva TL. N-acetylcysteine as a part of complex treatment of moderate COVID-associated pneumonia. Pul'monologiâ (Mosk). 2021 Feb 19;31(1):21–9.
- 467. Taher A, Lashgari M, Sedighi L, Rahimi-bashar F, Poorolajal J, Mehrpooya M. A pilot study on intravenous N-Acetylcysteine treatment in patients with mild-to-moderate COVID19-associated acute respiratory distress syndrome. Pharmacol Rep [Internet].
 2021 Jun 10 [cited 2021 Jun 21]; Available from: https://link.springer.com/10.1007/s43440-021-00296-2
- 468. Quinn TM, Gaughan EE, Bruce A, Antonelli J, O'Connor R, Li F, et al. Randomised Controlled Trial of Intravenous Nafamostat Mesylate in COVID pneumonitis: Phase 1b/2a Experimental Study to Investigate Safety, Pharmacokinetics and Pharmacodynamics [Internet]. Respiratory Medicine; 2021 Oct [cited 2021 Oct 18]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.10.06.21264648
- 469. Hassaniazad M, Eftekhar E, Inchehsablagh BR, Kamali H, Tousi A, Jaafari MR, et al. A triple-blind, placebo-controlled, randomized clinical trial to evaluate the effect of curcumin-containing nanomicelles on cellular immune responses subtypes and clinical outcome in COVID -19 patients. Phytotherapy Research. 2021 Sep 19;ptr.7294.
- 470. Kimura KS, Freeman MH, Wessinger BC, Gupta V, Sheng Q, Huang LC, et al. Interim analysis of an open-label randomized controlled trial evaluating nasal irrigations in non-hospitalized patients with COVID-19. Int Forum Allergy Rhinol 2020;10(12):1325-28. Available from: https://doi.org/10.1002/alr.22703.
- 471. Yildiz E, Koca Yildiz S, Kuzu S, Günebakan Ç, Bucak A, Kahveci OK.
 Comparison of the Healing Effect of Nasal Saline Irrigation with Triamcinolone
 Acetonide Versus Nasal Saline Irrigation alone in COVID-19 Related Olfactory
 Dysfunction: A Randomized Controlled Study. Indian J Otolaryngol Head Neck Surg
 [Internet]. 2021 Jul 10 [cited 2021 Nov 23]; Available from:
 https://link.springer.com/10.1007/s12070-021-02749-9
- George CE, Scheuch G, Seifart U, Inbaraj LR, Chandrasingh S, Nair IK, et al.
 COVID-19 symptoms are reduced by targeted hydration of the nose, larynx and trachea.
 Sci Rep. 2022 Dec;12(1):4599.

PAHOS



473. Baxter AL, Schwartz KR, Johnson RW, Kuchinski A-M, Swartout KM, Srinivasa Rao ASR, et al. Rapid initiation of nasal saline irrigation to reduce severity in high-risk COVID+ outpatients: a randomized clinical trial compared to a national dataset observational arm [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Aug [cited 2022 Jan 3]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2021.08.16.21262044

- 474. Nesari TM, Bhardwaj A, ShriKrishna R, Ruknuddin G, Ghildiyal S, Das A, et al. Neem (Azadirachta Indica A. Juss) Capsules for Prophylaxis of COVID-19 Infection: A Pilot, Double-Blind, Randomized Controlled Trial. Altern Ther Health Med. 2021 Apr 23;
- 475. Abdulamir AS, Gorial FI, Saadi SJ, Maulood MF, Hashim HA, abdulrrazaq MK. Effectiveness and Safety of Niclosamaide as Add-on Therapy to the Standard of Care Measures in COVID-19 Management: Randomized controlled clinical trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Jun [cited 2021 Jul 9]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.06.10.21258709
- 476. Cairns DM, Dulko D, Griffiths JK, Golan Y, Cohen T, Trinquart L, et al. Efficacy of Niclosamide vs Placebo in SARS-CoV-2 Respiratory Viral Clearance, Viral Shedding, and Duration of Symptoms Among Patients With Mild to Moderate COVID-19: A Phase 2 Randomized Clinical Trial. JAMA Netw Open. 2022 Feb 9;5(2):e2144942.
- 477. Labro G, Tubach F, Belin L, Dubost JL, Osman D, Muller G, et al. Nicotine patches in patients on mechanical ventilation for severe COVID-19: a randomized, double-blind, placebo-controlled, multicentre trial. Intensive Care Med. 2022 Jul;48(7):876–87.
- 478. Ashraf S, Ashraf S, Ashraf M, Imran MA, Kalsoom L, Siddiqui UN, et al. Honey and Nigella sativa against COVID-19 in Pakistan (HNS-COVID-PK): A multi-center placebo-controlled randomized clinical trial [Internet]. Infectious Diseases (except HIV/AIDS); 2020 Nov [cited 2021 May 4]. Available from: http://medrxiv.org/lookup/doi/10.1101/2020.10.30.20217364
- 479. Koshak AE, Koshak EA, Mobeireek AF, Badawi MA, Wali SO, Malibary HM, et al. Nigella sativa for the treatment of COVID-19: An open-label randomized controlled clinical trial. Complementary Therapies in Medicine. 2021 Sep;61:102769.



- Hammond J, Leister-Tebbe H, Gardner A, Abreu P, Bao W, Wisemandle W, et al.
 Oral Nirmatrelvir for High-Risk, Nonhospitalized Adults with Covid-19. N Engl J Med.
 2022 Feb 16;NEJMoa2118542.
- 481. Rocco PRM, Silva PL, Cruz FF, Junior MACM, Tierno PFGMM, Moura MA, et al. Early use of nitazoxanide in mild COVID-19 disease: randomized, placebo-controlled trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.10.21.20217208.
- 482. Vinicius Fontanesi Blum, Sérgio Cimerman, James R. Hunter, Paulo Tierno, Acioly Lacerda, Alexandre Soeiro, et al. Nitazoxanide In Vitro Efficacy Against SARS CoV-2 and In Vivo Superiority to Placebo to Treat Moderate COVID-19 – A Phase 2 Randomized Double-Blind Clinical Trial. SSRN [Internet]. 2021
- 483. Silva M, Espejo A, L Pereyra M, Lynch M, Thompson M, Taconelli H, et al. Efficacy of Nitazoxanide in reducing the viral load in COVID-19 patients. Randomized, placebo-controlled, single-blinded, parallel group, pilot study. [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Mar [cited 2021 Mar 8]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.03.03.21252509
- 484. Rossignol J-F, Bardin MC, Oaks JB, Bostick BG, Vora KN, Fulgencio J, et al. Early treatment with nitazoxanide prevents worsening of mild and moderate COVID-19 and subsequent hospitalization [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Apr [cited 2021 Apr 29]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2021.04.19.21255441

- 485. Fowotade A, Bamidele F, Egbetola B, Fagbamigbe AF, Adeagbo BA, Adefuye BO, et al. Efficacy and safety of nitazoxanide combined with ritonavir-boosted atazanavir for the treatment of mild to moderate COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Feb [cited 2022 Feb 16]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.02.03.22270152
- 486. Medhat MA, El-Kassas M, Karam-Allah H, Al Shafie A, Abd-Elsalam S, Moustafa E, et al. Sofosbuvir/ledipasvir in combination or nitazoxanide alone are safe and efficient treatments for COVID-19 infection: A randomized controlled trial for repurposing antivirals. Arab Journal of Gastroenterology. 2022 May;S1687197922000326.



- 487. Sokhela S, Bosch B, Hill A, Simmons B, Woods J, Johnstone H, et al. Randomized clinical trial of nitazoxanide or sofosbuvir/daclatasvir for the prevention of SARS-CoV-2 infection. Journal of Antimicrobial Chemotherapy. 2022 Aug 12;dkac266.
- 488. Moni M, Madathil T, Sathyapalan DT, Menon V, Gutjahr G, Edathadathil F, et al. A Feasibility Trial to Evaluate the Composite Efficacy of Inhaled Nitric Oxide in the Treatment of Covid 19 Pneumonia : Impact on Viral Load and Clinical Outcomes [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Apr [cited 2021 May 5]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.04.15.21255300
- 489. Winchester S, John S, Jabbar K, John I. Clinical Efficacy of Nitric Oxide Nasal Spray (NONS) for the Treatment of Mild COVID-19 Infection. Journal of Infection. 2021 May;S0163445321002516.
- 490. Strickland B, Albala L, Coffey EC, Carroll RW, Zapol WM, Ichinose F, et al. Safety and practicality of high dose inhaled nitric oxide in emergency department COVID-19 patients. The American Journal of Emergency Medicine. 2022 Aug;58:5–8.
- 491. Tandon M, Wu W, Moore K, Winchester S, Tu YP, Miller C, et al. SARS-CoV-2 accelerated clearance using a novel nitric oxide nasal spray (NONS) treatment: A randomized trial. The Lancet Regional Health - Southeast Asia. 2022 Jun;100036.
- Mobarak S, Salasi M, Hormati A, Khodadadi J, Ziaee M, Abedi F, et al. Evaluation of the Effect of Sofosbuvir and Daclatasvir in Hospitalised COVID-19 Patients: A Randomized Double-Blind Clinical Trial (DISCOVER). SSRN Journal [Internet]. 2021 [cited 2021 Mar 24]; Available from: https://www.ssrn.com/abstract=3792895
- 493. Eilidh B, Barlow-Pay F, Short R, Vilches-Moraga A, Price A, McGovern A, et al. Prior routine use of non-steroidal anti-inflammatory drugs (NSAIDs) and important outcomes in hospitalised patients with COVID-19. J Clin Med 2020;9(8):2586. Available from: https://doi.org/10.3390/jcm9082586.
- 494. Jeong HE, Lee H, Shin HJ, Choe YJ, Filion KB, Shin J-Y. Association between NSAIDs use and adverse clinical outcomes among adults hospitalised with COVID-19 in South Korea: a nationwide study [Preprint] MedRxiv 2020. Available from: https://doi.org/10.1101/2020.06.01.20119768.
- 495. Lund LC, Kristensen KB, Reilev M, Christensen S, Thomsen RW, Christiansen CF, et al. Adverse outcomes and mortality in users of non-steroidal anti-inflammatory

PAHOS



drugs who tested positive for SARS-CoV-2: a Danish nationwide cohort study. PLOS Med 2020;17(9):e1003308. Available from: https://doi.org/10.1371/journal.pmed.1003308.

- 496. Rinott E, Kozer E, Shapira Y, Bar-Haim A, Youngster I. Ibuprofen use and clinical outcomes in COVID-19 patients. Clin Microbiol Infect 2020;26(9):1259.e5-1259.e7. Available from: https://doi.org/10.1016/j.cmi.2020.06.003.
- 497. Wong AYS, MacKenna B, Morton C, Schultze A, Walker AJ, Bhaskaran K, et al. OpenSAFELY: do adults prescribed non-steroidal anti-inflammatory drugs have an increased risk of death from COVID-19? [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.08.12.20171405.
- 498. Imam Z, Odish F, Gill I, O'Connor D, Armstrong J, Vanood A, et al. Older age and comorbidity are independent mortality predictors in a large cohort of 1305 COVID-19 patients in Michigan, United States. J Intern Med 2020;288(4):469–76. Available from: https://doi.org/10.1111/joim.13119.
- 499. Esba LCA, Alqahtani RA, Thomas A, Shamas N, Alswaidan L, Mardawi G.
 Ibuprofen and NSAIDs use in COVID-19 infected patients is not associated with worse outcomes [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-85148/v1.
- 500. Alfredo CA, Noemí CR, Samuel RL, Daniel OC, Rodrigo RB, Paul MT, et al. Effect of Norelgestromin and Ethinylestradiol in Transdermal Patches on the Clinical Outcomes and Biochemical Parameters of COVID-19 Patients: A Clinical Trial Pilot Study. Pharmaceuticals. 2022 Jun 17;15(6):757.
- 501. Leal F, Garcia A, Abarca L del C, Gonzalez D, Cruz G, Montell M, et al. Effect of a Nutritional Support System to Increase Survival and Reduce Mortality in Patients with COVID-19 in Stage III and Comorbidities: A Blinded Randomized Controlled Clinical Trial. SSRN Journal [Internet]. 2021 [cited 2021 Nov 4]; Available from: https://www.ssrn.com/abstract=3949424
- 502. Mohsen Sedighiyan, Hamed Abdollahi, Elmira Karimi, Mostafa Badeli, Reza Erfanian, Shima Raeesi, et al. Omega-3 polyunsaturated fatty acids supplementation improve clinical symptoms in patients with covid-19: A randomized clinical trial. Authorea [Internet]. 2021.



- 503. Doaei S, Gholami S, Rastgoo S, Gholamalizadeh M, Bourbour F, Bagheri SE, et al. The effect of omega-3 fatty acid supplementation on clinical and biochemical parameters of critically ill patients with COVID-19: a randomized clinical trial. J Transl Med. 2021 Dec;19(1):128.
- 504. Arnardottir H, Pawelzik S-C, Sarajlic P, Quaranta A, Kolmert J, Religa D, et al. Immunomodulation by intravenous omega-3 fatty acid treatment in older subjects hospitalized for COVID-19: a single-blind randomized controlled trial [Internet]. Respiratory Medicine; 2021 Dec [cited 2022 Jan 10]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.12.27.21268264</u>
- 505. Gusdon AM, Faraday N, Aita JS, Kumar S, Mehta I, Choi HA, et al. Dendrimer nanotherapy for severe COVID-19 attenuates inflammation and neurological injury markers and improves outcomes in a phase2a clinical trial. Sci Transl Med. 2022 Jul 20;14(654):eabo2652.
- 506. Winthrop KL, Skolnick AW, Rafiq AM, Beegle SH, Suszanski J, Koehne G, et al. Opaganib in Coronavirus Disease 2019 Pneumonia: Results of a Randomized, Placebo-Controlled Phase 2a Trial. Open Forum Infectious Diseases. 2022 Jul 4;9(7):ofac232.
- 507. Fernando Carvalho Neuenschwander, Ofra Barnett-Griness, Stefania Piconi, Yasmin Maor, Eduardo Sprinz, Nimer Assy, et al. Effect of Opaganib on Supplemental Oxygen and Mortality in Patients with Severe SARS-CoV-2 Pneumonia. medRxiv [Internet]. 2022; Available from:

http://www.epistemonikos.org/documents/f770cb51808746db8a984f4669b7473f0b8d70e 3

- 508. Patel J, Beishuizen A, Ruiz XB, Boughanmi H, Cahn A, Criner GJ, et al. A Randomized Trial of Otilimab in Severe COVID-19 Pneumonia (OSCAR) [Internet]. Intensive Care and Critical Care Medicine; 2021 Apr [cited 2021 Apr 28]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.04.14.21255475
- 509. Araimo F, Imperiale C, Tordiglione P, Ceccarelli G, Borrazzo C, Alessandri F, et al. Ozone as adjuvant support in the treatment of COVID-19: a preliminary report of probiozovid trial [Preprint] J Med Virol 2020: jmv.26636. Available from: https://doi.org/10.1002/jmv.26636.
- 510. Shah M, Captain J, Vaidya V, Kulkarni A, Valsangkar K, Nair PMK, et al. Safety and efficacy of ozone therapy in mild to moderate COVID-19 patients: A phase 1/11

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randomized control trial (SEOT study). International Immunopharmacology. 2021 Feb;91:107301.

- 511. Berger JS, Kornblith LZ, Gong MN, Reynolds HR, Cushman M, Cheng Y, et al. Effect of P2Y12 Inhibitors on Survival Free of Organ Support Among Non–Critically Ill Hospitalized Patients With COVID-19: A Randomized Clinical Trial. JAMA. 2022 Jan 18;327(3):227.
- 512. Fessler SN, Liu L, Chang Y, Yip T, Johnston CS. Palmitoylethanolamide Reduces Proinflammatory Markers in Unvaccinated Adults Recently Diagnosed with COVID-19: A Randomized Controlled Trial. The Journal of Nutrition. 2022 Sep 9;nxac154.
- 513. Pandit A, Bhalani N, Bhushan BLS, Koradia P, Gargiya S, Bhomia V, et al. Efficacy and Safety of Pegylated Interferon alfa-2b in Moderate COVID-19: A phase II, randomized, controlled, open-label study. International Journal of Infectious Diseases. 2021 Mar;S1201971221002320.
- 514. Bushan S, Wanve S, Koradia P, Bhomia V, Soni P, Chakraborty S, et al. Efficacy and Safety of Pegylated Interferon-α2b in Moderate COVID-19: A phase 3, randomized, comparator-controlled, open-label study. International Journal of Infectious Diseases. 2021 Aug;S1201971221006779.
- 515. Feld JJ, Kandel C, Biondi MJ, Kozak RA, Zahoor MA, Lemieux C, et al. Peginterferon-lambda for the treatment of COVID-19 in outpatients [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.11.09.20228098.
- Jagannathan P, Andrews J, Bonilla H, Hedlin H, Jacobson K, Balasubramanian V, et al. Peginterferon lambda-1a for treatment of outpatients with uncomplicated COVID-19: a randomized placebo-controlled trial [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.11.18.20234161.
- 517. Sánchez-Conde M, Vizcarra P, Pérez-García JM, Gion M, Martialay MP, Taboada J, et al. Pembrolizumab in combination with tocilizumab in high-risk hospitalized patients with COVID-19 (COPERNICO): A randomized proof-of-concept phase II study. Int J Infect Dis. 2022 Aug 17;123:97–103.
- 518. Maldonado V, Hernandez-Ramírez C, Oliva-Pérez EA, Sánchez-Martínez CO, Pimentel-González JF, Molina-Sánchez JR, Jiménez-Villalba YZ, Chávez-Alderete J, and Loza-Mejía MA. Pentoxifylline Decreases Serum LDH Levels and Increases Lymphocyte Count in COVID-19 Patients: Results from an External Pilot Study.

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International Immunopharmacology 2020. 90 (January): 107209. https://doi.org/10.1016/j.intimp.2020.107209.

- 519. Azizi H, Rouhani N, Shaki F, Karimpour-razkenari E, Ghazaeian M, Salehifar E, et al. Pentoxifylline effects on hospitalized patients with COVID19: A randomized, double-blind clinical trial. International Immunopharmacology. 2021 Oct;108227.
- 520. Zhang F, Wei Y, He L, Zhang H, Hu Q, Yue H, et al. A trial of pirfenidone in hospitalized adult patients with severe coronavirus disease 2019. Chinese Medical Journal. 2022 Feb 5;135(3):368–70.
- 521. Varona JF, Landete P, Lopez-Martin JA, Estrada V, Paredes R, Guisado-Vasco P, et al. Preclinical and randomized phase I studies of plitidepsin in adults hospitalized with COVID-19. Life Sci Alliance. 2022 Apr;5(4):e202101200.
- 522. Lattmann E, Bhalerao P, ShashiBhushan B, Nargundkar N, Lattmann P, Pillai KS, et al. Randomized, Comparative, Clinical Trial to Evaluate Efficacy and Safety of PNB001 in Moderate COVID-19 Patients [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Apr [cited 2021 May 3]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.04.16.21255256
- 523. Méndez-Flores S, Priego-Ranero Á, Azamar-Llamas D, Olvera-Prado H, Rivas-Redondo KI, Ochoa-Hein E, et al. Effect of polymerized type I collagen in hyperinflammation of adult outpatients with symptomatic COVID-19: a double blind, randomised, placebo-controlled clinical trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 May [cited 2021 May 21]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.05.12.21257133
- 524. Kotfis K, Karolak I, Lechowicz K, Zegan-Barańska M, Pikulska A, Niedźwiedzka-Rystwej P, et al. Mineralocorticoid Receptor Antagonist (Potassium Canrenoate) Does Not Influence Outcome in the Treatment of COVID-19-Associated Pneumonia and Fibrosis—A Randomized Placebo Controlled Clinical Trial. Pharmaceuticals. 2022 Feb 5;15(2):200.
- 525. Wang Q, Lin X, Xiang X, Liu W, Fang Y, Chen H, et al. Oropharyngeal Probiotic ENT-K12 Prevents Respiratory Tract Infections Among Frontline Medical Staff Fighting Against COVID-19: A Pilot Study. Front Bioeng Biotechnol. 2021 Jun 24;9:646184.
- 526. Ivashkin V, Fomin V, Moiseev S, Brovko M, Maslennikov R, Ulyanin A, et al. Efficacy of a Probiotic Consisting of Lacticaseibacillus rhamnosus PDV 1705,

PAHO®



Bifidobacterium bifidum PDV 0903, Bifidobacterium longum subsp. infantis PDV 1911, and Bifidobacterium longum subsp. longum PDV 2301 in the Treatment of Hospitalized Patients with COVID-19: a Randomized Controlled Trial. Probiotics & Antimicro Prot [Internet]. 2021 Oct 13 [cited 2021 Oct 20]; Available from: https://link.springer.com/10.1007/s12602-021-09858-5

- 527. Wischmeyer PE, Tang H, Ren Y, Bohannon L, Ramirez ZE, Andermann TM, et al. Daily Lactobacillus Probiotic versus Placebo in COVID-19-Exposed Household Contacts (PROTECT-EHC): A Randomized Clinical Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jan [cited 2022 Jan 11]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.01.04.21268275</u>
- 528. Gutiérrez-Castrellón P, Gandara-Martí T, Abreu Y Abreu AT, Nieto-Rufino CD, López-Orduña E, Jiménez-Escobar I, et al. Probiotic improves symptomatic and viral clearance in Covid19 outpatients: a randomized, quadruple-blinded, placebo-controlled trial. Gut Microbes. 2022 Dec 31;14(1):2018899.
- 529. Saviano A, Potenza A, Siciliano V, Petruzziello C, Tarli C, Migneco A, et al. COVID-19 Pneumonia and Gut Inflammation: The Role of a Mix of Three Probiotic Strains in Reducing Inflammatory Markers and Need for Oxygen Support. JCM. 2022 Jun 28;11(13):3758.
- 530. Ghandehari S, Matusov Y, Pepkowitz S, Stein D, Kaderi T, Narayanan D, et al. Progesterone in addition to standard of care versus standard of care alone in the treatment of men admitted to the hospital with moderate to severe COVID-19: a randomised control phase 1 trial [Preprint]. 2020. Available from SSRN: https://doi.org/10.2139/ssrn.3709835.
- 531. Sigamani A, Shetty Madhavi S, Sudhishma RM, Chugani A, Chen-Walden H, Kutty T, and Platt D. Galectin Antagonist Use in Mild Cases of SARS-CoV-2 Cases; Pilot Feasibility Randomised, Open Label, Controlled Trial. [Preprint] 2020. https://doi.org/10.1101/2020.12.03.20238840.
- 532. Marcelo Augusto Duarte Silveira, David De Jong, Erica Batista dos Santos Galvao, Juliana Caldas Ribeiro, Thiago Cerqueira Silva, Andresa Aparecida Berretta, et al. Efficacy of propolis as an adjunct treatment for hospitalized COVID-19 patients: a randomized, controlled clinical trial. medRxiv [Internet]. 2021.



- 533. Johansson PI, Søe-Jensen P, Bestle MH, Clausen NE, Kristiansen KT, Lange T, et al. Prostacyclin in Mechanically Ventilated Patients with COVID-19 and Severe Endotheliopathy: A Multicenter, Randomized, Clinical Trial. Am J Respir Crit Care Med. 2021 Nov 23
- 534. Haeberle HA, Calov S, Martus P, Higuita LMS, Koeppen M, Goll A, et al. Inhaled Prostacyclin Improves Oxygenation in Patients with COVID-19-induced Acute Respiratory Distress Syndrome – a randomized controlled multicenter trial [Internet]. In Review; 2022 May [cited 2022 May 31]. Available from: https://www.researchsquare.com/article/rs-1652838/v1
- 535. Cadegiani F, McCoy J, Wambier C, Kovacevic M, Shapiro J, Sinclair R, et al. Proxalutamide (GT0918) Reduces the Rate of Hospitalization and Death in COVID-19 Male Patients: A Randomized Double-Blinded Placebo-Controlled Trial. ResearchSquare [Internet]. 2020.
- 536. Cadegiani FA, McCoy J, Gustavo Wambier C, Vaño-Galván S, Shapiro J, Tosti A, et al. Proxalutamide Significantly Accelerates Viral Clearance and Reduces Time to Clinical Remission in Patients with Mild to Moderate COVID-19: Results from a Randomized, Double-Blinded, Placebo-Controlled Trial. Cureus [Internet]. 2021 Feb 22 [cited 2021 Mar 4]
- 537. Cadegiani FA, Zimerman RA, Fonseca DN, Correia MN, Muller MP, Bet DL, et al. Final Results of a Randomized, Placebo-Controlled, Two-Arm, Parallel Clinical Trial of Proxalutamide for Hospitalized COVID-19 Patients: A Multiregional, Joint Analysis of the Proxa-Rescue AndroCoV Trial. Cureus [Internet]. 2021 Dec 25 [cited 2022 Jan 12]; Available from: <u>https://www.cureus.com/articles/80171-final-results-of-arandomized-placebo-controlled-two-arm-parallel-clinical-trial-of-proxalutamide-forhospitalized-covid-19-patients-a-multiregional-joint-analysis-of-the-proxa-rescueandrocov-trial</u>
- 538. Cadegiani FA, Zimerman RA, do Nascimento Fonseca D, do Nascimento Correia M, McCoy J, Wambier CG, et al. Proxalutamide (GT0918) Reduces the Rate of Hospitalization in mild-to-moderate COVID-19 Female Patients: A Randomized Double-Blinded Placebo-Controlled Two-Arm Parallel Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Jul [cited 2021 Jul 29]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.07.06.21260086



539. Fragoso-Saavedra S, Núñez I, Audelo-Cruz BM, Arias-Martínez S, Manzur-Sandoval D, Quintero-Villegas A, et al. Pyridostigmine in adults with severe SARS-CoV-2 infection: the PISCO trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Apr [cited 2021 May 4]. Available from:

http://medrxiv.org/lookup/doi/10.1101/2021.04.28.21255834

- 540. Önal H, Arslan B, Üçüncü Ergun N, Topuz Ş, Yilmaz Semerci S, Kurnaz ME, et al. Treatment of COVID-19 patients with quercetin: a prospective, single center, randomized, controlled trial. Turk J Biol. 2021;45(4):518–29.
- 541. Di Pierro F, Iqtadar S, Khan A, Ullah Mumtaz S, Masud Chaudhry M, Bertuccioli A, et al. Potential Clinical Benefits of Quercetin in the Early Stage of COVID-19: Results of a Second, Pilot, Randomized, Controlled and Open-Label Clinical Trial. Int J Gen Med. 2021;14:2807–16.
- 542. Shohan M, Nashibi R, Mahmoudian-Sani M-R, Abolnezhadian F, Ghafourian M, Alavi SM, et al. The therapeutic efficacy of quercetin in combination with antiviral drugs in hospitalized COVID-19 patients: A randomized controlled trial. European Journal of Pharmacology. 2022 Jan;914:174615.
- 543. Rondanelli M, Perna S, Gasparri C, Petrangolini G, Allegrini P, Cavioni A, et al. Promising Effects of 3-Month Period of Quercetin Phytosome® Supplementation in the Prevention of Symptomatic COVID-19 Disease in Healthcare Workers: A Pilot Study. Life. 2022 Jan 4;12(1):66.
- 544. Nicastri E, Marinangeli F, Pivetta E, Torri E, Reggiani F, Fiorentino G, et al. A phase 2 randomized, double-blinded, placebo-controlled, multicenter trial evaluating the efficacy and safety of raloxifene for patients with mild to moderate COVID-19. eClinicalMedicine. 2022 Jun;48:101450.
- 545. Amat-Santos IJ, Santos-Martinez S, López-Otero D, Nombela-Franco L, Gutiérrez-Ibanes E, Del Valle R, et al. Ramipril in high risk patients with COVID-19. J Am Coll Cardiol 2020;76(3):268–76. Available from: https://doi.org/10.1016/j.jacc.2020.05.040.
- 546. Stasko N, Cockrell AS, Kocher JF, Henson I, Emerson D, Wang Y, et al. A randomized, controlled, feasibility study of RD-X19 in subjects with mild-to-moderate COVID-19 in the outpatient setting. Clin Transl Sci. 2022 Feb 8;

PAHO®

- 547. Li C, Luo F, Liu C, Xiong N, Xu Z, Zhang W, et al. Effect of a genetically engineered interferon-alpha versus traditional interferon-alpha in the treatment of moderate-to-severe COVID-19: a randomised clinical trial. Annals of Medicine. 2021 Jan 1;53(1):391–401.
- 548. Streinu-Cercel A, Săndulescu O, Preotescu L-L, Kim JY, Kim Y-S, Cheon S, et al. Efficacy and Safety of Regdanvimab (CT-P59): A Phase 2/3 Randomized, Double-Blind, Placebo-Controlled Trial in Outpatients With Mild-to-Moderate Coronavirus Disease 2019. Open Forum Infectious Diseases. 2022 Apr 1;9(4):ofac053.
- 549. Kim JY, Jang YR, Hong JH, Jung JG, Park J-H, Streinu-Cercel A, et al. Safety, Virologic Efficacy, and Pharmacokinetics of CT-P59, a Neutralizing Monoclonal Antibody Against SARS-CoV-2 Spike Receptor-Binding Protein: Two Randomized, Placebo-Controlled, Phase I Studies in Healthy Individuals and Patients With Mild SARS-CoV-2 Infection. Clinical Therapeutics. 2021 Aug;S0149291821003088.
- 550. Weinreich DM, Sivapalasingam S, Norton T, Ali S, Gao H, Bhore R, et al. REGN-COV2, a Neutralizing Antibody Cocktail, in Outpatients with Covid-19. N Engl J Med. 2020 Dec 17;NEJMoa2035002.
- 551. Abani O, Abbas A, Abbas F, Abbas M, Abbasi S, Abbass H, et al. Casirivimab and imdevimab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. The Lancet. 2022 Feb;399(10325):665–76.
- 552. O'Brien MP, Forleo-Neto E, Sarkar N, Isa F, Hou P, Chan K-C, et al. Effect of Subcutaneous Casirivimab and Imdevimab Antibody Combination vs Placebo on Development of Symptomatic COVID-19 in Early Asymptomatic SARS-CoV-2 Infection: A Randomized Clinical Trial. JAMA [Internet]. 2022 Jan 14 [cited 2022 Jan 18]; Available from: https://jamanetwork.com/journals/jama/fullarticle/2788256
- 553. Herman GA, O'Brien MP, Forleo-Neto E, Sarkar N, Isa F, Hou P, et al. Efficacy and safety of a single dose of casirivimab and imdevimab for the prevention of COVID-19 over an 8-month period: a randomised, double-blind, placebo-controlled trial. The Lancet Infectious Diseases. 2022 Jul;S1473309922004169.
- 554. Somersan-Karakaya S, Mylonakis E, Menon VP, Wells JC, Ali S, Sivapalasingam S, et al. Casirivimab and Imdevimab for the Treatment of Hospitalized Patients With COVID-19. The Journal of Infectious Diseases. 2022 Jul 27;jiac320.

PAHOO

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- 555. Portal-Celhay C, Forleo-Neto E, Eagan W, Musser BJ, Davis JD, Turner KC, et al. Phase 2 dose-ranging study of the virologic efficacy and safety of the combination COVID-19 antibodies casirivimab and imdevimab in the outpatient setting [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Nov [cited 2021 Dec 13]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.11.09.21265912</u>
- 556. Isa F, Forleo-Neto E, Meyer J, Zheng W, Rasmussen S, Armas D, et al. Repeat Subcutaneous Administration of REGEN-COV[®] in Adults is Well-Tolerated and Prevents the Occurrence of COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Nov [cited 2021 Dec 1]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.11.10.21265889</u>
- 557. Weinreich DM, Sivapalasingam S, Norton T, Ali S, Gao H, Bhore R, et al. REGEN-COV Antibody Cocktail Clinical Outcomes Study in Covid-19 Outpatients [Internet]. Infectious Diseases (except HIV/AIDS); 2021 May [cited 2021 May 24]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.05.19.21257469</u>
- 558. Huang DT, McCreary EK, Bariola JR, Minnier TE, Wadas RJ, Shovel JA, et al. Effectiveness of casirivimab and imdevimab, and sotrovimab during Delta variant surge: a prospective cohort study and comparative effectiveness randomized trial [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Dec [cited 2022 Jan 10]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.12.23.21268244
- 559. Beigel JH, Tomashek KM, Dodd LE, Mehta AK, Zingman BS, Kalil AC, et al. Remdesivir for the treatment of COVID-19 — final report. N Engl J Med 2020;383:1813-26. Available from: https://doi.org/10.1056/NEJMoa2007764.
- 560. Goldman JD, Lye DCB, Hui DS, Marks KM, Bruno R, Montejano R, et al. Remdesivir for 5 or 10 days in patients with severe COVID-19. N Engl J Med 2020;383:1827-37. Available from: https://doi.org/10.1056/NEJMoa2015301.
- 561. Wang Y, Zhang D, Du G, Du R, Zhao J, Jin Y, et al. Remdesivir in adults with severe COVID-19: a randomised, double-blind, placebo-controlled, multicentre trial. Lancet 2020;395(10236):1569–78. Available from: https://doi.org/10.1016/S0140-6736(20)31022-9.
- 562. Spinner CD, Gottlieb RL, Criner GJ, Arribas López JR, Cattelan AM, Viladomiu AS, et al. Effect of remdesivir vs standard care on clinical status at 11 days in patients



with moderate COVID-19: a randomized clinical trial. JAMA 2020;324(11):1048-57. Available from: https://doi.org/10.1001/jama.2020.16349.

- 563. Mahajan L, Singh A, Gifty. Clinical outcomes of using remdesivir in patients with moderate to severe COVID-19: A prospective randomised study. Indian J Anaesth. 2021;65(13):41.
- 564. Abd-Elsalam S, Ahmed OA, Mansour NO, Abdelaziz DH, Salama M, Fouad MHA, et al. Remdesivir Efficacy in COVID-19 Treatment: A Randomized Controlled Trial. The American Journal of Tropical Medicine and Hygiene [Internet]. 2021 Sep 10 [cited 2021 Oct 19]; Available from: https://www.ajtmh.org/view/journals/tpmd/aop/article-10.4269-ajtmh.21-0606/article-

10.4269-ajtmh.21-0606.xml

PAHO®

- 565. Sarhan RM, Harb HS, Abou Warda AE, Salem-Bekhit MM, Shakeel F, Alzahrani SA, et al. Efficacy of the early treatment with tocilizumab-hydroxychloroquine and tocilizumab-remdesivir in severe COVID-19 Patients. Journal of Infection and Public Health. 2021 Nov;S1876034121003452.
- 566. Gottlieb RL, Vaca CE, Paredes R, Mera J, Webb BJ, Perez G, et al. Early Remdesivir to Prevent Progression to Severe Covid-19 in Outpatients. N Engl J Med. 2021 Dec 22;NEJMoa2116846.
- 567. Ali K, Azher T, Baqi M, Binnie A, Borgia S, Carrier FM, et al. Remdesivir for the treatment of patients in hospital with COVID-19 in Canada: a randomized controlled trial. CMAJ. 2022 Jan 19;cmaj.211698.
- 568. Landoni G, Piemonti L, Monforte A d'Arminio, Grossi P, Zangrillo A, Bucci E, et al. A Multicenter Phase 2 Randomized Controlled Study on the Efficacy and Safety of Reparixin in the Treatment of Hospitalized Patients with COVID-19 Pneumonia. Infect Dis Ther [Internet]. 2022 May 26 [cited 2022 Jun 6]; Available from: https://link.springer.com/10.1007/s40121-022-00644-6
- 569. McCreary MR, Schnell PM, Rhoda DA. Randomized double-blind placebocontrolled proof-of-concept trial of resveratrol for outpatient treatment of mild coronavirus disease (COVID-19). Sci Rep. 2022 Dec;12(1):10978.
- 570. Kaplan HG, Wang K, Reeves KM, Scanlan JM, Nunn CC, Kieper DA, et al. Resveratrol and Zinc in the Treatment of Outpatients With COVID-19 – The Reszinate Study - A Phase 1/2 Randomized Clinical Trial Utilizing Home Patient-Obtained Nasal



ww.paho.org/coronavirus

and Saliva Viral Sampling. SSRN Journal [Internet]. 2021 [cited 2021 Oct 13]; Available from: https://www.ssrn.com/abstract=3934228

- 571. Cheng L-l, Guan W-j, Duan C-y, Zhang N-f, Lei C-l, Hu Y, et al. Effect of recombinant human granulocyte colony–stimulating factor for patients with coronavirus disease 2019 (COVID-19) and lymphopenia: a randomized clinical trial. JAMA Intern Med 2020; published online 10 September 2020. Available from: https://doi.org/10.1001/jamainternmed.2020.5503.
- 572. Bosteels C, Damme KV, De Leeuw E, Declercq J, Maes B, Bosteels V, et al. Early treatment with inhaled GM-CSF improves oxygenation and anti-viral immunity in COVID-19 induced lung injury – a randomized clinical trial [Internet]. In Review; 2021 Oct [cited 2021 Oct 21]. Available from: https://www.researchsquare.com/article/rs-959220/v1
- 573. DiNubile MJ, Parra S, Salomó AC, Levinson SL. Adjunctive Recombinant Human Plasma Gelsolin for Severe Coronavirus Disease 2019 Pneumonia. Open Forum Infectious Diseases. 2022 Aug 2;9(8):ofac357.
- 574. Hung IF, Lung KC, Tso EY, Liu R, Chung TW, Chu MY, et al. Triple combination of interferon beta-1b, lopinavir-ritonavir, and ribavirin in the treatment of patients admitted to hospital with COVID-19: an open-label, randomised, phase 2 trial. Lancet 2020;395(10238):1695–1704. Available from: https://doi.org/10.1016/S0140-6736(20)31042-4.
- 575. Cao Y, Wei J, Zou L, Jiang T, Wang G, Chen L, et al. Ruxolitinib in treatment of severe coronavirus disease 2019 (COVID-19): a multicenter, single-blind, randomized controlled trial. J Allergy Clin Immunol 2020;146(1):137-46.E3. Available from: https://doi.org/10.1016/j.jaci.2020.05.019.
- 576. Han MK, Antila M, Ficker JH, Gordeev I, Guerreros A, Bernus AL, et al. Ruxolitinib in addition to standard of care for the treatment of patients admitted to hospital with COVID-19 (RUXCOVID): a randomised, double-blind, placebo-controlled, phase 3 trial. The Lancet Rheumatology. 2022 Mar;S2665991322000443.
- 577. Barnette KG, Gordon MS, Rodriguez D, Bird TG, Skolnick A, Schnaus M, et al. Oral Sabizabulin for High-Risk, Hospitalized Adults with Covid-19: Interim Analysis. NEJM Evidence [Internet]. 2022 Jul 6 [cited 2022 Jul 13]; Available from: <u>https://evidence.nejm.org/doi/10.1056/EVIDoa2200145</u>



- 578. The REMAP-CAP Investigators. Interleukin-6 Receptor Antagonists in Critically Ill Patients with Covid-19. N Engl J Med. 2021 Feb 25;NEJMoa2100433.
- 579. Lescure F-X, Honda H, Fowler RA, Lazar JS, Shi G, Wung P, et al. Sarilumab in patients admitted to hospital with severe or critical COVID-19: a randomised, doubleblind, placebo-controlled, phase 3 trial. The Lancet Respiratory Medicine. 2021 Mar;S2213260021000990.
- 580. Sivapalasingam S, Lederer DJ, Bhore R, Hajizadeh N, Criner G, Hossain R, et al. A Randomized Placebo-Controlled Trial of Sarilumab in Hospitalized Patients with Covid-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2021 May [cited 2021 May 20]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.05.13.21256973
- 581. Mariette X, Hermine O, Tharaux P-L, Resche-Rigon M, Porcher R, Ravaud P, et al. Sarilumab in adults hospitalised with moderate-to-severe COVID-19 pneumonia (CORIMUNO-SARI-1): An open-label randomised controlled trial. The Lancet Rheumatology. 2022 Jan;4(1):e24–32.
- 582. Hermine O, Mariette X, Porcher R, Resche-Rigon M, Tharaux P-L, Ravaud P. Effect of Interleukin-6 Receptor Antagonists in Critically Ill Adult Patients with COVID-19 Pneumonia: two Randomised Controlled Trials of the CORIMUNO-19 Collaborative Group. Eur Respir J. 2022 Feb 3;2102523.
- 583. García-Vicuña R, Rodriguez-García SC, Abad-Santos F, Bautista Hernández A, García-Fraile L, Barrios Blandino A, et al. Subcutaneous IL-6 Inhibitor Sarilumab vs. Standard Care in Hospitalized Patients With Moderate-To-Severe COVID-19: An Open Label Randomized Clinical Trial. Front Med. 2022 Feb 23;9:819621.
- 584. Merchante N, Cárcel S, Garrido-Gracia JC, Trigo-Rodríguez M, Esteban Moreno MÁ, León-López R, et al. Early Use of Sarilumab in Patients Hospitalised with COVID-19 Pneumonia and Features of Systemic Inflammation. Antimicrob Agents Chemother. 2021 Dec 13;AAC.02107-21.
- 585. Sancho-López A, Caballero-Bermejo AF, Ruiz-Antorán B, Múñez Rubio E, García Gasalla M, Buades J, et al. Efficacy and Safety of Sarilumab in patients with COVID19 Pneumonia: A Randomized, Phase III Clinical Trial (SARTRE Study). Infect Dis Ther [Internet]. 2021 Oct 17 [cited 2021 Nov 2]; Available from: https://link.springer.com/10.1007/s40121-021-00543-2

PAHOS





- 586. Branch-Elliman W, Ferguson R, Doros G, Woods P, Leatherman S, Strymish J, et al. Subcutaneous sarilumab for the treatment of hospitalized patients with moderate to severe COVID19 disease: A pragmatic, embedded randomized clinical trial. De Socio GV, editor. PLoS ONE. 2022 Feb 25;17(2):e0263591.
- 587. Resende GG, da Cruz Lage R, Lobê SQ, Medeiros AF, Costa e Silva AD, Nogueira Sá AT, et al. Blockade of interleukin seventeen (IL-17A) with secukinumab in hospitalized COVID-19 patients – the BISHOP study. Infectious Diseases. 2022 Aug 3;54(8):591–9.
- 588. Granfeldt A, Andersen LW, Vallentin MF, Hilberg O, Hasselstrøm JB, Sørensen LK, et al. Senicapoc treatment in COVID -19 Patients with Severe Respiratory Insufficiency A Randomized, OPEN-LABEL, Phase II Trial. Acta Anaesthesiol Scand. 2022 Apr 11;aas.14072.
- 589. Panatto D, Orsi A, Bruzzone B, Ricucci V, Fedele G, Reiner G, et al. Efficacy of the Sentinox Spray in Reducing Viral Load in Mild COVID-19 and Its Virucidal Activity against Other Respiratory Viruses: Results of a Randomized Controlled Trial and an In Vitro Study. Viruses. 2022 May 12;14(5):1033.
- 590. Tian F, Wang J, Xi X, He M, Zhao C, Feng F, et al. Efficacy and safety of shortwave diathermy treatment for moderate COVID-19 patients: a prospective, double-blind, randomized controlled clinical study. European journal of physical and rehabilitation medicine [Internet]. 2021; Available from:

http://www.epistemonikos.org/documents/356ba654e07f6231b50fd2a20e44ae587685ad9

- 591. Santamarina MG, Beddings I, Lomakin FM, Boisier Riscal D, Gutiérrez Claveria M, Vidal Marambio J, et al. Sildenafil for treating patients with COVID-19 and perfusion mismatch: a pilot randomized trial. Crit Care. 2022 Dec;26(1):1.
- 592. The WHO Rapid Evidence Appraisal for COVID-19 Therapies (REACT) Working Group, Domingo P, Mur I, Mateo GM, Gutierrez M del M, Pomar V, et al. Association Between Administration of IL-6 Antagonists and Mortality Among Patients Hospitalized for COVID-19: A Meta-analysis. JAMA [Internet]. 2021 Jul 6 [cited 2021 Jul 13]; Available from: https://jamanetwork.com/journals/jama/fullarticle/2781880
- 593. Aryan H, Farahani RH, Chamanara M, Elyasi S, Jaafari MR, Haddad M, et al. Evaluation of the efficacy of oral nano-silymarin formulation in hospitalized patients

PAHO®

with COVID-19: A double-blind placebo-controlled clinical trial. Phytotherapy Research. 2022 Jul 20;ptr.7537.

- 594. Asadipooya K, Abbasi F, Adatorwovor R, Davarpanah MA, Mansoori Y, Hajiani M, et al. A Randomized Single Blind Controlled Trial of Combination Therapy (Spironolactone and Sitagliptin) in Hospitalized Adult Patients with Covid-19. SSRN Journal [Internet]. 2021 [cited 2021 Aug 3]; Available from: https://www.ssrn.com/abstract=3889411
- 595. Sadeghi A, Asgari AA, Norouzi A, Kheiri Z, Anushirvani A, Montazeri M, et al. Sofosbuvir and daclatasvir compared with standard of care in the treatment of patients admitted to hospital with moderate or severe coronavirus infection (COVID-19): a randomized controlled trial. J Antimicrob Chemother 2020;75(11):3379-85. Available from: https://doi.org/10.1093/jac/dkaa334.
- 596. Yakoot M, Eysa B, Gouda E, Hill A, Helmy SA, Elsayed MR, et al. Efficacy and safety of sofosbuvir/daclatasvir in the treatment of COVID-19: a randomized, controlled study [Preprint]. 2020. Available from SSRN: https://doi.org/10.2139/ssrn.3705289.
- 597. Roozbeh F, Saeedi M, Alizadeh-Navaei R, Hedayatizadeh-Omran A, Merat S, Wentzel H, et al. Sofosbuvir and daclatasvir for the treatment of COVID-19 outpatients: a double-blind, randomized controlled trial. Journal of Antimicrobial Chemotherapy. 2020 Dec 18;dkaa501.
- 598. Mobarak S, Salasi M, Hormati A, Khodadadi J, Ziaee M, Abedi F, et al. Evaluation of the Effect of Sofosbuvir and Daclatasvir in Hospitalised COVID-19 Patients: A Randomized Double-Blind Clinical Trial (DISCOVER). SSRN Journal [Internet]. 2021 [cited 2021 Mar 24]; Available from: https://www.ssrn.com/abstract=3792895
- 599. Alavi-moghaddam M, Haghighi M, Sabaghian T, Soroureddin Z, Chaboki BG. Safety and Efficacy of Sofosbuvir in Hospitalized Adult Patients with SARS-CoV-2: A Preliminary Report. SSRN Journal [Internet]. 2021 [cited 2021 Mar 24]; Available from: https://www.ssrn.com/abstract=3790463
- 600. Khalili H, Nourian A, Ahmadinejad Z, Emadi Kouchak H, Jafari S, Dehghan Manshadi SA, et al. Efficacy and safety of sofosbuvir/ ledipasvir in treatment of patients with COVID-19; A randomized clinical trial. Acta Biomed. 2020 Nov 10;91(4):e2020102.



- 601. Elgohary MA-S, Hasan EM, Ibrahim AA, Ahmed Abdelsalam MF, Abdel-Rahman RZ, Zaki AI, et al. Efficacy of Sofosbuvir plus Ledipasvir in Egyptian patients with COVID-19 compared to standard treatment: Randomized controlled trial [Internet]. Epidemiology; 2021 May [cited 2021 May 26]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.05.19.21257429
- 602. Sayad B, Khodarahmi R, Najafi F, Miladi R, Mohseni Afshar Z, Mansouri F, et al. Efficacy and safety of sofosbuvir/velpatasvir versus the standard of care in adults hospitalized with COVID-19: a single-centre, randomized controlled trial. Journal of Antimicrobial Chemotherapy. 2021 May 25;dkab152.
- 603. El-Bendary M, Abd-Elsalam S, Elbaz T, El-Akel W, Cordie A, Elhadidy T, et al. Efficacy of combined Sofosbuvir and Daclatasvir in the treatment of COVID-19 patients with pneumonia: a multicenter Egyptian study. Expert Review of Anti-infective Therapy. 2021 Jul 15;1–5.
- 604. Abbass S, Salama M, Salman T, Sabry A, Abdel-Razek W, Kamal E, et al. fficacy and safety of Sofosbuvir plus Daclatasvir or Ravidasvir in patients with COVID-19, A Randomized Controlled Trial. J Med Virol. 2021 Aug 11;jmv.27264.
- 605. Medhat MA, El-Kassas M, Karam-Allah H, Al Shafie A, Abd-Elsalam S, Moustafa E, et al. Sofosbuvir/ledipasvir in combination or nitazoxanide alone are safe and efficient treatments for COVID-19 infection: A randomized controlled trial for repurposing antivirals. Arab Journal of Gastroenterology. 2022 May;S1687197922000326.
- 606. Bozorgmehr R, Amiri F, Hosein Zadeh M, Ghorbani F, Khameneh Bagheri A, Yazdi E, et al. Effect of Sofosbuvir on Length of Hospital Stay in Moderate COVID-19 Cases; a Randomized Controlled Trial. Arch Acad Emerg Med. 2022;10(1):e46.
- 607. Gupta A, Gonzalez-Rojas Y, Juarez E, Crespo Casal M, Moya J, Rodrigues Falci D, et al. Effect of Sotrovimab on Hospitalization or Death Among High-risk Patients With Mild to Moderate COVID-19: A Randomized Clinical Trial. JAMA [Internet]. 2022 Mar 14 [cited 2022 Mar 28]; Available from:

https://jamanetwork.com/journals/jama/fullarticle/2790246

PAHO®

608. Wadhwa B, Malhotra V, Kerai S, Husain F, Pandey NB, Saxena KN, et al. Phase
2 randomised placebo-controlled trial of spironolactone and dexamethasone versus
dexamethasone in COVID-19 hospitalised patients in Delhi [Internet]. Respiratory

BE AWARE. PREPARE. ACT. www.paho.org/coronavirus

Medicine; 2022 Jul [cited 2022 Jul 21]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.07.01.22277163

- 609. INSPIRATION-S Investigators. Atorvastatin versus placebo in patients with covid-19 in intensive care: randomized controlled trial. BMJ. 2022 Jan 7;376:e068407.
- 610. Ghafoori M, Saadati H, Taghavi M, Azimian A, Alesheikh P, Mohajerzadeh MS, et al. Survival of the hospitalized patients with COVID-19 receiving atorvastatin: A randomized clinical trial. J Med Virol. 2022 Mar 10;
- 611. Carmenate YV, Alkaabi FM, Aleman YMC, Valverde CAV, Ahmed YM, Sanna P, et al. Safety and Efficacy of Autologous Non-Hematopoietic Enriched Stem Cell Nebulization in Covid-19 Patients. A Randomized Clinical Trial, Abu Dhabi 2020. [Internet]. In Review; 2021 Jun [cited 2021 Jun 18]. Available from: https://www.researchsquare.com/article/rs-558653/v1
- 612. GLUCOCOVID investigators, Corral-Gudino L, Bahamonde A, Arnaiz-Revillas F, Gómez-Barquero J, Abadía-Otero J, et al. Methylprednisolone in adults hospitalized with COVID-19 pneumonia: An open-label randomized trial (GLUCOCOVID). Wien Klin Wochenschr [Internet]. 2021 Feb 3 [cited 2021 Feb 11]; Available from: http://link.springer.com/10.1007/s00508-020-01805-8
- 613. Jeronimo CMP, Farias MEL, Almeida Val FF, Sampaio VS, Alexandre MAA, Melo GC, et al. Methylprednisolone as adjunctive therapy for patients hospitalized with COVID-19 (metcovid): a randomised, double-blind, phase IIb, placebo-controlled trial. Clin Infect Dis 2020: ciaa1177. Available from: https://doi.org/10.1093/cid/ciaa1177.
- Horby P, Lim WS, Emberson J, Mafham M, Bell J, Linsell L, et al. Effect of dexamethasone in hospitalized patients with COVID-19: preliminary report [Preprint] MedRxiv 2020. Available from: https://doi.org/10.1101/2020.06.22.20137273.
- 615. The WHO Rapid Evidence Appraisal for COVID-19 Therapies (REACT) Working Group. Association between administration of systemic Corticosteroids and mortality among critically ill patients with COVID-19: a meta-analysis. JAMA 2020;324:1330-41. Available from: https://doi.org/10.1001/jama.2020.17023.
- 616. Tomazini BM, Maia IS, Cavalcanti AB, Berwanger O, Rosa RG, Veiga VC, et al. Effect of dexamethasone on days alive and ventilator-free in patients with moderate or severe acute respiratory distress syndrome and COVID-19: the CoDEX randomized



clinical trial. JAMA 2020; 324(13):1307-16. Available from: https://doi.org/10.1001/jama.2020.17021.

- 617. The Writing Committee for the REMAP-CAP Investigators, et al. Effect of hydrocortisone on mortality and organ support in patients with severe COVID-19: the REMAP-CAP COVID-19 corticosteroid domain randomized clinical trial. JAMA 2020; 324(13):1317-29. <u>https://doi.org/10.1001/jama.2020.17022</u>.
- 618. Munch MW, Granholm A, Kjær MN, Aksnes TS, Sølling CG, Christensen S, et al. Long-term mortality and health-related quality of life in the COVID STEROID trial. Acta Anaesthesiol Scand. 2022 Apr;66(4):543–5.
- 619. Dequin P-F, Heming N, Meziani F, Plantefève G, Voiriot G, Badié J, et al. Effect of hydrocortisone on 21-day mortality or respiratory support among critically ill patients with COVID-19: a randomized clinical trial. JAMA 2020;324(13):1298-1306. Available from: https://doi.org/10.1001/jama.2020.16761.
- 620. Farahani RH, Mosaed R, Nezami-Asl A, Chamanara N, Soleiman-Meigooni S, Kalantar S, et al. Evaluation of the efficacy of methylprednisolone pulse therapy in treatment of Covid-19 adult patients with severe respiratory failure: randomized, clinical trial [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-66909/v1.
- 621. Edalatifard M, Akhtari M, Salehi M, Naderi Z, Jamshidi A, Mostafaei S, et al. Intravenous methylprednisolone pulse as a treatment for hospitalised severe COVID-19 patients: results from a randomised controlled clinical trial [Preprint]. Eur Respir J 2020; published online 17 September 2020. Available from: https://doi.org/10.1183/13993003.02808-2020.
- 622. Tang X, Feng Y-M, Ni J-X, Zhang J-Y, Liu L-M, Hu K, et al. Early Use of Corticosteroid May Prolong SARS-CoV-2 Shedding in Non-Intensive Care Unit Patients with COVID-19 Pneumonia: A Multicenter, Single-Blind, Randomized Control Trial. Respiration. 2021 Jan 22;1–11.
- 623. Jamaati H, Hashemian SM, Farzanegan B, Malekmohammad M, Tabarsi P, Marjani M, et al. No clinical benefit of high dose corticosteroid administration in patients with COVID-19: A preliminary report of a randomized clinical trial. European Journal of Pharmacology. 2021 Apr;897:173947.



- 624. Rashad A, Mousa S, Nafady-Hego H, Nafady A, Elgendy H. Short term survival of critically ill COVID-19 Egyptian patients on assisted ventilation treated by either Dexamethasone or Tocilizumab. Sci Rep. 2021 Dec;11(1):8816.
- 625. Les I, Loureiro-Amigo J, Capdevila F, Oriol I, Elejalde I, Aranda-Lobo J, et al. Methylprednisolone Pulses in Hospitalized COVID-19 Patients Without Respiratory Failure: A Randomized Controlled Trial. Front Med (Lausanne). 2022;9:807981.
- 626. Ranjbar K, Shahriarirad R, Erfani A, Khodamoradi Z, Saadi MHG, Mirahmadizadeh A, et al. Methylprednisolone or Dexamethasone, Which One Is the Superior Corticosteroid in the Treatment of Hospitalized COVID-19 Patients: A Triple-Blinded Randomized Controlled Trial [Internet]. In Review; 2021 Feb [cited 2021 Feb 14]. Available from: https://www.researchsquare.com/article/rs-148529/v1
- 627. Munch MW, Myatra SN, Tirupakuzhi Vijayaraghavan BK, Saseedharan S, Benfield T, Wahlin RR, et al. Dexamethasone 12 mg versus 6 mg for patients with COVID-19 and severe hypoxia: an international, randomized, blinded trial [Internet]. Intensive Care and Critical Care Medicine; 2021 Jul [cited 2021 Jul 30]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.07.22.21260755
- 628. Maskin LP, Bonelli I, Olarte GL, Palizas F, Velo AE, Lurbet MF, et al. High-Versus Low-Dose Dexamethasone for the Treatment of COVID-19-related Acute Respiratory Distress Syndrome: A Multicenter and Randomized Open-label Clinical Trial [Internet]. Intensive Care and Critical Care Medicine; 2021 Sep [cited 2021 Sep 24]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.09.15.21263597
- 629. Toroghi N, Abbasian L, Nourian A, Davoudi-Monfared E, Khalili H, Hasannezhad M, et al. Comparing efficacy and safety of different doses of dexamethasone in the treatment of COVID-19: a three-arm randomized clinical trial. Pharmacol Rep [Internet]. 2021 Nov 27 [cited 2021 Dec 1]; Available from: <u>https://link.springer.com/10.1007/s43440-021-00341-0</u>
- 630. Taboada M, Rodríguez N, Varela PM, Rodríguez MT, Abelleira R, González A, et al. Effect of high *versus* low dose of dexamethasone on clinical worsening in patients hospitalised with moderate or severe COVID-19 Pneumonia: an open-label, randomised clinical trial. Eur Respir J. 2021 Dec 16;2102518.
- 631. Naik NB, Puri GD, Kajal K, Mahajan V, Bhalla A, Kataria S, et al. High-Dose Dexamethasone Versus Tocilizumab in Moderate to Severe COVID-19 Pneumonia: A

PAHOS





Randomized Controlled Trial. Cureus [Internet]. 2021 Dec 11 [cited 2022 Jan 24]; Available from: <u>https://www.cureus.com/articles/78251-high-dose-dexamethasone-versus-tocilizumab-in-moderate-to-severe-covid-19-pneumonia-a-randomized-controlled-trial</u>

- 632. Salvarani C, Massari M, Costantini M, Franco Merlo D, Lucia Mariani G, Viale P, et al. Intravenous methylprednisolone pulses in hospitalised patients with severe COVID-19 pneumonia, A double-blind, randomised, placebo-controlled trial. Eur Respir J. 2022 Mar 31;2200025.
- 633. Bouadma L, Mekontso-Dessap A, Burdet C, Merdji H, Poissy J, Dupuis C, et al. High-Dose Dexamethasone and Oxygen Support Strategies in Intensive Care Unit Patients With Severe COVID-19 Acute Hypoxemic Respiratory Failure: The COVIDICUS Randomized Clinical Trial. JAMA Intern Med. 2022 Jul 5;
- 634. Dastenae ZH, Bahadori A, Dehghani M, Asadi-Samani M, Izadi I, Shahraki HR. Comparison of the effect of intravenous dexamethasone and methylprednisolone on the treatment of hospitalized patients with COVID-19: A randomized clinical trial. International Journal of Infectious Diseases. 2022 Jul;S1201971222004131.
- 635. Ramakrishnan S, Nicolau DV, Langford B, Mahdi M, Jeffers H, Mwasuku C, et al. Inhaled budesonide in the treatment of early COVID-19 (STOIC): a phase 2, openlabel, randomised controlled trial. The Lancet Respiratory Medicine. 2021 Apr;S2213260021001600.
- 636. Yu L-M, Bafadhel M, Dorward J, Hayward G, Saville BR, Gbinigie O, 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. The Lancet. 2021 Aug;S014067362101744X.
- 637. Song J-Y, Yoon J-G, Seo Y-B, Lee J, Eom J-S, Lee J-S, et al. Ciclesonide Inhaler Treatment for Mild-to-Moderate COVID-19: A Randomized, Open-Label, Phase 2 Trial. JCM. 2021 Aug 12;10(16):3545.
- 638. Clemency BM, Varughese R, Gonzalez-Rojas Y, Morse CG, Phipatanakul W, Koster DJ, et al. A randomized controlled trial of inhaled ciclesonide for outpatient treatment of symptomatic COVID-19 infections [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Sep [cited 2021 Sep 13]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.09.07.21261811

рано©

- 639. Ezer N, Belga S, Daneman N, Chan A, Smith BM, Daniels S-A, et al. Inhaled and intranasal ciclesonide for the treatment of covid-19 in adult outpatients: CONTAIN phase II randomised controlled trial. BMJ. 2021 Nov 2;e068060.
- 640. Duvignaud A, Lhomme E, Onaisi R, Sitta R, Gelley A, Chastang J, 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 Mar 15;S1198-743X(22)00108-2.
- 641. Agustí A, De Stefano G, Levi A, Muñoz X, Romero-Mesones C, Sibila O, et al. Add-on inhaled budesonide in the treatment of hospitalised patients with COVID-19: a randomised clinical trial. Eur Respir J. 2022 Mar;59(3):2103036.
- 642. Accelerating Covid-19 Therapeutic Interventions and Vaccines (ACTIV)-6
 Study Group, Naggie S. Inhaled Fluticasone for Outpatient Treatment of Covid-19: A
 Decentralized, Placebo-controlled, Randomized, Platform Clinical Trial [Internet].
 Infectious Diseases (except HIV/AIDS); 2022 Jul [cited 2022 Jul 25]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.07.12.22277548
- 643. Gonzalez Ochoa AJ, Raffetto JD, Hernandez AG, Zavala NA, Gutierrez O, Vargas A, and Loustaunau J. Sulodexide in the Treatment of Patients with Early Stages of COVID-19: A Randomised Controlled Trial. MedRxiv 2020. https://doi.org/10.1101/2020.12.04.20242073.
- 644. Dow GS, Smith BL. A phase II, double blind, placebo-controlled, randomized evaluation of the safety and efficacy of tafenoquine in patients with mild-moderate COVID-19 disease. New Microbes and New Infections. 2022 Apr;47:100986.
- 645. Singh D, Bogus M, Moskalenko V, Lord R, Moran EJ, Crater GD, et al. A phase
 2 study of the inhaled pan-JAK inhibitor TD-0903 in severe COVID-19: Part 1 [Internet].
 Respiratory Medicine; 2021 Mar [cited 2021 Mar 24]. Available from:
 http://medrxiv.org/lookup/doi/10.1101/2021.03.09.21252944
- 646. Parienti J-J, Prazuck T, Peyro-Saint-Paul L, Fournier A, Valentin C, Brucato S, et al. Effect of Tenofovir Disoproxil Fumarate and Emtricitabine on nasopharyngeal SARS-CoV-2 viral load burden amongst outpatients with COVID-19: A pilot, randomized, open-label phase 2 trial. EClinicalMedicine. 2021 Jun;100993.
- 647. Arruda EAG, Pires-Neto RJ, Medeiros MS, Quirino-Filho J, Clementino M, Gondim RNDG, et al. Clinical Trial of Efficacy and Toxicity of Disoproxil Tenofovir

PAHO®

Fumarate and Emtricitabine for Mild to Moderate SARS-CoV-2 Infections [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Sep [cited 2021 Oct 12]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.09.28.21264242

- 648. Amra B, Ashrafi F, Soltaninejad F, Feizi A, Salmasi M. Thalidomide for the treatment of severe Covid-19: A randomized clinical trial [Internet]. In Review; 2021 Apr [cited 2021 Apr 8]. Available from: https://www.researchsquare.com/article/rs-379635/v1
- 649. Shirin Haghighi, Soodeh Ramezaninejad, Atousa Hakamifard, et al. The Effects of Thalidomide as an Adjuvant Treatment Besides of Dexamethasone and Remdesivir on Patients with Moderate COVID-19. Available online at: https://ssrn.com/abstract=3941711
- 650. Bencheqroun H, Ahmed Y, Kocak M, Villa E, Barrera C, Mohiuddin M, et al. A Randomized, Double-Blind, Placebo-Controlled, Multicenter Study to Evaluate the Safety and Efficacy of ThymoQuinone Formula (TQF) for Treating Outpatient SARS-CoV-2. Pathogens. 2022 May 7;11(5):551.
- 651. Barrett CD, Moore HB, Moore EE, Wang DJ, Hajizadeh N, Biffl WL, et al. STudy of Alteplase for Respiratory failure in SARS-Cov2 COVID-19 (STARS): A Vanguard Multicenter, Rapidly Adaptive, Pragmatic, Randomized, Controlled Trial. Chest. 2021 Sep;S0012369221040630.
- 652. Levin MJ, Ustianowski A, De Wit S, Launay O, Avila M, Templeton A, et al. Intramuscular AZD7442 (Tixagevimab–Cilgavimab) for Prevention of Covid-19. N Engl J Med. 2022 Apr 20;NEJMoa2116620.
- 653. Montgomery H, Hobbs FDR, Padilla F, Arbetter D, Templeton A, Seegobin S, et al. Efficacy and safety of intramuscular administration of tixagevimab–cilgavimab for early outpatient treatment of COVID-19 (TACKLE): a phase 3, randomised, doubleblind, placebo-controlled trial. The Lancet Respiratory Medicine. 2022 Jun;S2213260022001801.
- 654. Tixagevimab–cilgavimab for treatment of patients hospitalised with COVID-19: a randomised, double-blind, phase 3 trial. The Lancet Respiratory Medicine. 2022 Jul;S2213260022002156.
- 655. Rosas IO, Bräu N, Waters M, Go RC, Malhotra A, Hunter BD, et al. Tocilizumab in patients hospitalised with COVID-19 pneumonia: Efficacy, safety, viral clearance, and



antibody response from a randomised controlled trial (COVACTA). eClinicalMedicine. 2022 May;47:101409.

- 656. Wang D, Fu B, Peng Z, Yang D, Han M, Li M, et al. Tocilizumab ameliorates the hypoxia in COVID-19 moderate patients with bilateral pulmonary lesions: a randomized, controlled, open-label, multicenter trial [Preprint]. 2020. Available from SSRN: https://doi.org/10.2139/ssrn.3667681.
- 657. Salvarani C, Dolci G, Massari M, Merlo DF, Cavuto S, Savoldi L, et al. Effect of tocilizumab vs standard care on clinical worsening in patients hospitalized with COVID-19 pneumonia: a randomized clinical trial [Preprint]. JAMA Int Med 2020; published online 20 October 2020. Available from:

https://doi.org/10.1001/jamainternmed.2020.6615.

- 658. Stone JH, Frigault MJ, Serling-Boyd NJ, Fernandes AD, Harvey L, Foulkes AS, et al. Efficacy of tocilizumab in patients hospitalized with COVID-19 [Preprint]. N Engl J Med 2020; published online 21 October 2020. Available from: https://doi.org/10.1056/NEJMoa2028836.
- 659. Hermine O, Mariette X, Tharaux P-L, Resche-Rigon M, Porcher R, Ravaud P, and the CORIMUNO-19 Collaborative Group. Effect of tocilizumab vs usual care in adults hospitalized with COVID-19 and moderate or severe pneumonia: a randomized clinical trial [Preprint]. JAMA Int Med 2020; published online 20 October 2020. Available from: https://doi.org/10.1001/jamainternmed.2020.6820.
- 660. Salama C, Han J, Yau L, Reiss WG, Kramer B, Neidhart JD, et al. Tocilizumab in Patients Hospitalized with Covid-19 Pneumonia. N Engl J Med. 2020 Dec 17;NEJMoa2030340.
- 661. Veiga VC, Prats JAGG, Farias DLC, Rosa RG, Dourado LK, Zampieri FG, et al. Effect of tocilizumab on clinical outcomes at 15 days in patients with severe or critical coronavirus disease 2019: randomised controlled trial. BMJ. 2021 Jan 20;n84.
- 662. Horby PW, Campbell M, Staplin M, et al. Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): a randomised, controlled, open-label, platform trial. The Lancet. 2021 May;397(10285):1637–45.
- 663. Rutgers A, Westerweel PE, van der Holt B, Postma S, van Vonderen MGA, Piersma DP, et al. Timely Administration of Tocilizumab Improves Survival of



Hospitalized COVID-19 Patients. SSRN Journal [Internet]. 2021 [cited 2021 May 12]; Available from: https://www.ssrn.com/abstract=3834311

- 664. Talaschian M, Akhtari M, Mahmoudi M, Mostafaei S, Jafary M, Husseini AS, et al. Tocilizumab Failed to Reduce Mortality in Severe COVID-19 Patients: Results From a Randomized Controlled Clinical Trial [Internet]. In Review; 2021 May [cited 2021 May 14]. Available from: https://www.researchsquare.com/article/rs-463921/v1
- 665. Hamed DM, Belhoul KM, Al Maazmi NA, Ghayoor F, Moin M, Al Suwaidi M, et al. Intravenous methylprednisolone with or without tocilizumab in patients with severe COVID-19 pneumonia requiring oxygen support: A prospective comparison. Journal of Infection and Public Health. 2021 Aug;14(8):985–9.
- 666. Broman N, Feuth T, Vuorinen T, Valtonen M, Hohenthal U, Löyttyniemi E, et al. Early administration of tocilizumab in hospitalized COVID-19 patients with elevated inflammatory markers; COVIDSTORM – A prospective, randomized, single center, open label study. Clinical Microbiology and Infection. 2022 Mar;S1198743X22001045.
- 667. Rosas IO, Diaz G, Gottlieb RL, Lobo SM, Robinson P, Hunter BD, et al. Tocilizumab and remdesivir in hospitalized patients with severe COVID-19 pneumonia: a randomized clinical trial. Intensive Care Med [Internet]. 2021 Oct 5 [cited 2021 Oct 12]; Available from: https://link.springer.com/10.1007/s00134-021-06507-x
- 668. Soin AS, Kumar K, Choudhary NS, Sharma P, Mehta Y, Kataria S, et al. Tocilizumab plus standard care versus standard care in patients in India with moderate to severe COVID-19-associated cytokine release syndrome (COVINTOC): an open-label, multicentre, randomised, controlled, phase 3 trial. The Lancet Respiratory Medicine. 2021 May;9(5):511–21.
- 669. Hermine O, Mariette X, Porcher R, Djossou F, Nguyen Y, Arlet JB, et al. Tocilizumab plus dexamethasone versus dexamethasone in patients with moderate-tosevere COVID-19 pneumonia: A randomised clinical trial from the CORIMUNO-19 study group. eClinicalMedicine. 2022 Apr;46:101362.
- 670. Karampitsakos T, Papaioannou O, Tsiri P, Katsaras M, Katsimpris A, Kalogeropoulos AP, et al. Tocilizumab versus baricitinib in hospitalized patients with severe COVID-19: an open label, randomized controlled trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jun [cited 2022 Jul 6]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.06.13.22276211</u>



- 671. Kumar PN, Hernández-Sánchez J, Nagel S, Feng Y, Cai F, Rabin J, et al. Safety and Efficacy of Tocilizumab 4 or 8 mg/kg in Hospitalized Patients With Moderate to Severe Coronavirus Disease 2019 Pneumonia: A Randomized Clinical Trial. Open Forum Infectious Diseases. 2022 Jan 1;9(1):ofab608.
- 672. Guimarães PO, Quirk D, Furtado RH, Maia LN, Saraiva JF, Antunes MO, et al. Tofacitinib in Patients Hospitalized with Covid-19 Pneumonia. N Engl J Med. 2021 Jun 16;NEJMoa2101643.
- 673. Murugesan H, Cs G, Nasreen HS, Santhanam S, M G, Ravi S, et al. An Evaluation of Efficacy and Safety of Tofacitinib, A JAK Inhibitor in the Management of Hospitalized Patients with Mild to Moderate COVID-19 - An Open-Label Randomized Controlled Study. J Assoc Physicians India. 2022 Dec;69(12):11–2.
- 674. Saeedi-Boroujeni A, Nashibi R, Ghadiri AA, Nakajima M, Salmanzadeh S, Mahmoudian-Sani MR, et al. Tranilast as an Adjunctive Therapy in Hospitalized Patients with Severe COVID- 19: A Randomized Controlled Trial. Archives of Medical Research. 2022 Mar;S0188440922000248.
- 675. Wu X, Yu K, Wang Y, Xu W, Ma H, Hou Y, et al. Efficacy and safety of triazavirin therapy for coronavirus disease 2019: a pilot randomized controlled trial. Engineering 2020;6(10):1185-91. Available from: https://doi.org/10.1016/j.eng.2020.08.011.
- 676. Wagener G, Goldklang MP, Gerber A, Elisman K, Eiseman KA, Fonseca LD, et al. A randomized, placebo-controlled, double-blinded pilot study of angiotensin 1–7 (TXA-127) for the treatment of severe COVID-19. Crit Care. 2022 Dec;26(1):229.
- 677. Lau FH, Powell CE, Adonecchi G, Danos DM, DiNardo AR, Chugden RJ, et al.
 Pilot phase results of a prospective, randomized controlled trial of narrowband ultraviolet
 B phototherapy in hospitalized COVID -19 patients. Experimental Dermatology. 2022
 Jun 13;exd.14617.
- 678. Nojomi M, Yasin Z, Keyvani H, Makiani MJ, Roham M, Laali A, et al. Effect of arbidol on COVID-19: a randomized controlled trial [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-78316/v1.
- 679. Yethindra V, Tagaev T, Uulu MS, Parihar Y. Efficacy of umifenovir in the treatment of mild and moderate COVID-19 patients. Int J Res Pharm Sci 2020;11(SPL1):506–09. Available from: https://doi.org/10.26452/ijrps.v11iSPL1.2839.

PAHOS



- 680. Ghaderkhani S, Khaneshan AS, Salami A, Alavijeh PE, Kouchak HE, Khalili H, et al. Efficacy and safety of arbidol in treatment of patients with COVID-19 infection: a randomized clinical trial [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-91430/v1.
- 681. Alavi Darazam I, Shokouhi S, Mardani M, Pourhoseingholi MA, Rabiei MM, Hatami F, et al. Umifenovir in hospitalized moderate to severe COVID-19 patients: A randomized clinical trial. International Immunopharmacology. 2021 Oct;99:107969.
- 682. Ramachandran R, Bhosale V, Reddy H, Atam V, Faridi M, Fatima J, et al. Phase III, Randomized, Double-Blind, Placebo Controlled Trial of Efficacy, Safety and Tolerability of Antiviral Drug Umifenovir vs Standard Care of Therapy in Non-Severe Covid-19 Patients. SSRN Journal [Internet]. 2021 [cited 2021 Sep 29]; Available from: https://www.ssrn.com/abstract=3919585
- 683. Vlaar APJ, de Bruin S, Busch M, Timmermans SAMEG, van Zeggeren IE, Koning R, et al. Anti-C5a antibody IFX-1 (vilobelimab) treatment versus best supportive care for patients with severe COVID-19 (PANAMO): an exploratory, open-label, phase 2 randomised controlled trial. The Lancet Rheumatology [Internet]. [cited 2020 Sep 29]; Available from: <u>https://doi.org/10.1016/S2665-9913(20)30341-6</u>
- 684. Vlaar APJ, Witzenrath M, van Paassen P, Heunks LMA, Mourvillier B, de Bruin S, et al. Anti-C5a antibody (vilobelimab) therapy for critically ill, invasively mechanically ventilated patients with COVID-19 (PANAMO): a multicentre, double-blind, randomised, placebo-controlled, phase 3 trial. The Lancet Respiratory Medicine. 2022 Sep;S2213260022002971.
- 685. Majidi N, Bahadori E, Shekari S, Gholamalizadeh M, Tajadod S, Ajami M, et al. Effects of supplementation with low-dose group B vitamins on clinical and biochemical parameters in critically ill patients with COVID-19: a randomized clinical trial. Expert Review of Anti-infective Therapy. 2022 Sep 28;1–7.
- 686. Zhang J, Rao X, Li Y, Zhu Y, Liu F, Guo G, et al. High-dose vitamin C infusion for the treatment of critically ill COVID-19 [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-52778/v1.
- 687. Kumari P, Dembra S, Dembra P, Bhawna F, Gul A, Ali B, et al. The Role of Vitamin C as Adjuvant Therapy in COVID-19. Cureus [Internet]. 2020 Nov 30 [cited



2021 Jan 11]; Available from: https://www.cureus.com/articles/45284-the-role-of-vitamin-c-as-adjuvant-therapy-in-covid-19

- 688. Jamali Moghadam Siahkali S, Zarezade B, Koolaji S, Alinaghi S, Zendehdel A, Tabarestani M, et al. Safety and Effectiveness of High-Dose Vitamin C in Patients with COVID-19; A Randomized Controlled open-label Clinical Trial . ResearchSquare [Internet]. 2021.
- 689. Thomas S, Patel D, Bittel B, Wolski K, Wang Q, Kumar A, et al. Effect of High-Dose Zinc and Ascorbic Acid Supplementation vs Usual Care on Symptom Length and Reduction Among Ambulatory Patients With SARS-CoV-2 Infection: The COVID A to Z Randomized Clinical Trial. JAMA Netw Open. 2021 Feb 12;4(2):e210369.
- 690. Tehrani S, Yadegarynia D, Abrishami A, Moradi H, Gharaei B, Rauofi M, et al. An investigation into the Effects of Intravenous Vitamin C on Pulmonary CT Findings and Clinical Outcomes of Patients with COVID 19 Pneumonia A Randomized Clinical Trial. Urology Journal. 2021 Nov 8;(Instant 2021):6863.
- 691. Beigmohammadi MT, Bitarafan S, Hoseindokht A, Abdollahi A, Amoozadeh L, Soltani D. The effect of supplementation with vitamins A, B, C, D, and E on disease severity and inflammatory responses in patients with COVID-19: a randomized clinical trial. Trials. 2021 Dec;22(1):802.
- 692. Majidi N, Rabbani F, Gholami S, Gholamalizadeh M, BourBour F, Rastgoo S, et al. The Effect of Vitamin C on Pathological Parameters and Survival Duration of Critically Ill Coronavirus Disease 2019 Patients: A Randomized Clinical Trial. Front Immunol. 2021 Dec 15;12:717816.
- 693. Ried K, BinJemain T, Sali A. Therapies to Prevent Progression of COVID-19, Including Hydroxychloroquine, Azithromycin, Zinc, and Vitamin D3 With or Without Intravenous Vitamin C: An International, Multicenter, Randomized Trial. Cureus [Internet]. 2021 Nov 25 [cited 2022 Jan 10]; Available from: <u>https://www.cureus.com/articles/76496-therapies-to-prevent-progression-of-covid-19-including-hydroxychloroquine-azithromycin-zinc-and-vitamin-d3-with-or-without-intravenous-vitamin-c-an-international-multicenter-randomized-trial</u>
- 694. Coppock D, Violet PC, Vasquez G, Belden K, Foster M, Mullin B, et al.
 Pharmacologic Ascorbic Acid as Early Therapy for Hospitalized Patients with COVID-19: A Randomized Clinical Trial. Life. 2022 Mar 19;12(3):453.

PAHOS



- 695. Castillo ME, Costa LME, Barrios JMV, Díaz JFA, Miranda JL, Bouillon R, Gomez JMQ. Effect of calcifediol treatment and best available therapy versus best available therapy on intensive care unit admission and mortality among patients hospitalized for COVID-19: a pilot randomized clinical study [Preprint]. J Steroid Biochem Mol Biol 2020;203:105751. Available from: https://doi.org/10.1016/j.jsbmb.2020.105751.
- 696. Rastogi A, Bhansali A, Khare N, Suri V, Yaddanapudi N, Sachdeva N, et al. Short term, high-dose vitamin D supplementation for COVID-19 disease: a randomised, placebo-controlled, study (SHADE Study) [Preprint]. Postgrad Med J 2020; published online 12 November 2020. Available from: https://doi.org/10.1136/postgradmedj-2020-139065.
- 697. Murai IH, Fernandes AL, Sales LP, Pinto AJ, Goessler KF, Duran CSC, et al. Effect of a Single High Dose of Vitamin D3 on Hospital Length of Stay in Patients With Moderate to Severe COVID-19: A Randomized Clinical Trial. JAMA. 2021 Feb 17
- 698. Lakkireddy M, Gadiga SG, Malathi RD, Karra ML, Raju ISSVPM, Ragini, et al. Impact of Pulse D Therapy on The Inflammatory Markers in Patients With COVID-19. [Internet]. In Review; 2021 Feb [cited 2021 Mar 8]. Available from: https://www.researchsquare.com/article/rs-152494/v1
- 699. Sabico S, Enani MA, Sheshah E, Aljohani NJ, Aldisi DA, Alotaibi NH, et al. Effects of a 2-Week 5000 IU versus 1000 IU Vitamin D3 Supplementation on Recovery of Symptoms in Patients with Mild to Moderate Covid-19: A Randomized Clinical Trial. Nutrients. 2021 Jun 24;13(7):2170.
- 700. Maghbooli Z, Sahraian MA, Jamalimoghadamsiahkali S, Asadi A, Zarei A, Zendehdel A, et al. Treatment With 25-Hydroxyvitamin D3 (Calcifediol) Is Associated With a Reduction in the Blood Neutrophil-to-Lymphocyte Ratio Marker of Disease Severity in Hospitalized Patients With COVID-19: A Pilot Multicenter, Randomized, Placebo-Controlled, Double-Blinded Clinical Trial. Endocrine Practice. 2021 Oct;S1530891X21012593.
- 701. Gaborit B, Dailly E, Vanhove B, Josien R, Lacombe K, Dubee V, et al. Pharmacokinetics and safety of XAV-19, a swine glyco-humanized polyclonal anti-SARS-CoV-2 antibody, for COVID-19-related moderate pneumonia: a randomized, double-blind, placebo-controlled, phase IIa study [Internet]. Infectious Diseases (except

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HIV/AIDS); 2021 Apr [cited 2021 Apr 28]. Available from: http://medrxiv.org/lookup/doi/10.1101/2021.04.15.21255549

- 702. Bishop CW, Ashfaq A, Melnick JZ, Vazquez-Escarpanter E, Fialkow JA, Strugnell SA, et al. Results From the REsCue Trial: A Randomized Controlled Trial with Extended-Release Calcifediol in Symptomatic Outpatients with COVID-19 [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Feb [cited 2022 Feb 17]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.01.31.22270036</u>
- 703. Karonova TL, Chernikova AT, Golovatyuk KA, Bykova ES, Grant WB, Kalinina OV, et al. Vitamin D Intake May Reduce SARS-CoV-2 Infection Morbidity in Health Care Workers. Nutrients. 2022 Jan 24;14(3):505.
- 704. Cannata-Andía JB, Díaz-Sottolano A, Fernández P, Palomo-Antequera C, Herrero-Puente P, Mouzo R, et al. A single-oral bolus of 100,000 IU of cholecalciferol at hospital admission did not improve outcomes in the COVID-19 disease: the COVID-VIT-D—a randomised multicentre international clinical trial. BMC Med. 2022 Dec;20(1):83.
- 705. Jolliffe DA, Holt H, Greenig M, Talaei M, Perdek N, Pfeffer P, et al. Vitamin D Supplements for Prevention of Covid-19 or other Acute Respiratory Infections: a Phase 3 Randomized Controlled Trial (CORONAVIT) [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Mar [cited 2022 Apr 25]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.03.22.22271707
- 706. Villasis-Keever MA, López-Alarcón MG, Miranda-Novales G, Zurita-Cruz JN, Barrada-Vázquez AS, González-Ibarra J, et al. Efficacy and Safety of Vitamin D Supplementation to Prevent COVID-19 in Frontline Healthcare Workers. A Randomized Clinical Trial. Archives of Medical Research. 2022 Jun;53(4):423–30.
- Mariani J, Antonietti L, Tajer C, Ferder L, Inserra F, Sanchez Cunto M, et al. High-dose vitamin D versus placebo to prevent complications in COVID-19 patients: Multicentre randomized controlled clinical trial. Putzu A, editor. PLoS ONE. 2022 May 27;17(5):e0267918.
- Annweiler C, Beaudenon M, Gautier J, Gonsard J, Boucher S, Chapelet G, et al. High-dose versus standard-dose vitamin D supplementation in older adults with COVID-19 (COVIT-TRIAL): A multicenter, open-label, randomized controlled superiority trial. Cannegieter SC, editor. PLoS Med. 2022 May 31;19(5):e1003999.

PAHOS





- 709. Karonova TL, Golovatyuk KA, Kudryavtsev IV, Chernikova AT, Mikhaylova AA, Aquino AD, et al. Effect of Cholecalciferol Supplementation on the Clinical Features and Inflammatory Markers in Hospitalized COVID-19 Patients: A Randomized, Open-Label, Single-Center Study. Nutrients. 2022 Jun 23;14(13):2602.
- 710. Romero-Ibarguengoitia ME, Gutiérrez-González D, Cantú-López C, Sanz-Sánchez MA, González-Cantú A. Effect of Vitamin D 3 supplementation vs. dietary-hygienic measures on SARS-COV-2 infection rates in hospital workers with 25-hydroxyvitamin D3 [25(OH)D3] levels ≥20 ng/mL [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Jul [cited 2022 Jul 22]. Available from: http://medrxiv.org/lookup/doi/10.1101/2022.07.12.22277450
- 711. Cervero M, López-Wolf D, Casado G, Novella-Mena M, Ryan-Murua P, Taboada-Martínez ML, et al. Beneficial Effect of Short-Term Supplementation of High Dose of Vitamin D3 in Hospitalized Patients With COVID-19: A Multicenter, Single-Blinded, Prospective Randomized Pilot Clinical Trial. Front Pharmacol. 2022 Jul 4;13:863587.
- 712. Abroug H, Maatouk A, Bennasrallah C, Dhouib W, Fredj MB, Zemni I, et al. Effect of vitamin D supplementation versus placebo on recovery delay among COVID-19 patients: A randomized-controlled clinical trial [Internet]. In Review; 2022 Jul [cited 2022 Jul 26]. Available from: <u>https://www.researchsquare.com/article/rs-1276203/v1</u>
- De Niet S, Trémège M, Coffiner M, Rousseau AF, Calmes D, Frix AN, et al.
 Positive Effects of Vitamin D Supplementation in Patients Hospitalized for COVID-19: A Randomized, Double-Blind, Placebo-Controlled Trial. Nutrients. 2022 Jul 26;14(15):3048.
- 714. Brunvoll SH, Nygaard AB, Ellingjord-Dale M, Holland P, Istre MS, Kalleberg KT, et al. Prevention of covid-19 and other acute respiratory infections with cod liver oil supplementation, a low dose vitamin D supplement: quadruple blinded, randomised placebo controlled trial. BMJ. 2022 Sep 7;e071245.
- 715. van Helmond N, Brobyn TL, LaRiccia PJ, Cafaro T, Hunter K, Roy S, et al. Vitamin D3 Supplementation at 5000 IU Daily for the Prevention of Influenza-Like Illness in Healthcare Workers: A Randomized Clinical Trial [Internet]. Infectious Diseases (except HIV/AIDS); 2022 Sep [cited 2022 Sep 30]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2022.09.16.22280047</u>



- 716. Gaborit B, Dailly E, Vanhove B, Josien R, Lacombe K, Dubee V, et al. Pharmacokinetics and safety of XAV-19, a swine glyco-humanized polyclonal anti-SARS-CoV-2 antibody, for COVID-19-related moderate pneumonia: a randomized, double-blind, placebo-controlled, phase IIa study [Internet]. Infectious Diseases (except HIV/AIDS); 2021 Apr [cited 2021 Apr 28]. Available from: <u>http://medrxiv.org/lookup/doi/10.1101/2021.04.15.21255549</u>
- 717. De Leeuw E, Van Damme KFA, Declercq J, Bosteels C, Maes B, Tavernier SJ, et al. Efficacy and safety of the investigational complement C5 inhibitor zilucoplan in patients hospitalized with COVID-19: an open-label randomized controlled trial. Respir Res. 2022 Aug 9;23(1):202.
- 718. Hassan M, Abdelmaksoud A, Ghweil A, Rashad A, Aref Z, Khodeary A, et al. Olfactory disturbances as presenting manifestation among Egyptian patients with COVID-19: possible role of zinc [Preprint]. ResearchSquare 2020. Available from: https://doi.org/10.21203/rs.3.rs-107577/v1.
- 719. Abd-Elsalam S, Soliman S, Esmail ES, Khalaf M, Mostafa EF, Medhat MA, Ahmed OA, El Ghafar MSA, Alboraie M, and Hassany SM. Do Zinc Supplements Enhance the Clinical Efficacy of Hydroxychloroquine?: A Randomized, Multicenter Trial. Biological Trace Element Research 2020. https://doi.org/10.1007/s12011-020-02512-1.
- 720. Abdelmaksoud AA, Ghweil AA, Hassan MH, Rashad A, Khodeary A, Aref ZF, et al. Olfactory Disturbances as Presenting Manifestation Among Egyptian Patients with COVID-19: Possible Role of Zinc. Biol Trace Elem Res [Internet]. 2021 Jan 7 [cited 2021 Jan 11]; Available from: http://link.springer.com/10.1007/s12011-020-02546-5
- 721. Patel O, Chinni V, El-Khoury J, Perera M, Neto AS, McDonald C, et al. A pilot double-blind safety and feasibility randomised controlled trial of high-dose intravenous zinc in hospitalised COVID-19 patients. J Med Virol. 2021 Feb 25;jmv.26895.
- 722. Zhong M, Sun A, Xiao T, Yao G, Sang L, Zheng X, Zhang J, et al. A randomized, single-blind, group sequential, active-controlled study to evaluate the clinical efficacy and safety of α-lipoic acid for critically ill patients with coronavirus disease 2019 (COVID-19) [Preprint]. MedRxiv 2020. Available from: https://doi.org/10.1101/2020.04.15.20066266.

