

Institute of Clinical Epidemiology, National Institutes of Health, UP Manila In cooperation with the Philippine Society for Microbiology and Infectious Diseases Funded by the Department of Health

EVIDENCE SUMMARY

Among patients with COVID-19, should tocilizumab be used for treatment?

Updated by: Katherine Ruth O. Relato, MD, Carol Stephanie C. Tan-Lim, MD, MSc (Clinical Epidemiology), Natasha Ann R. Esteban-Ipac, MD

Initial review by: Ian Theodore Cabaluna, RPh, MD, GDip (Epi), Howell Henrian G Bayona, MSc, CSP-PASP

RECOMMENDATIONS

We recommend the addition of tocilizumab to systemic steroids in patients showing rapid respiratory deterioration and/or requiring high doses of oxygen (high-flow nasal cannula, noninvasive or invasive mechanical ventilation) and with elevated biomarkers of inflammation (CRP). (Moderate certainty of evidence, Strong recommendation)

We recommend against the use of tocilizumab among patients with COVID-19 infection who do not require oxygen. (Very low certainty of evidence, Strong recommendation)

Consensus issues

Report on adverse events, although an important outcome, was not rated as a critical outcome to be included in the decision making. The over-all assessment of the certainty of evidence is based only on critical outcomes identified by the consensus panel, hence, the quality of evidence was retained as moderate.

There is no new evidence on the use of tocilizumab among patients with COVID-19 infection who do not require oxygen. Given the lack of evidence, potential adverse effects and risks, this recommendation also considers indiscriminate use to avoid misuse or overuse of tocilizumab among patients who do not require oxygen.

PREVIOUS RECOMMENDATIONS

We recommend the addition of tocilizumab to systemic steroids in patients showing rapid respiratory deterioration and/or requiring high doses of oxygen (high-flow nasal cannula, noninvasive or invasive mechanical ventilation) and with elevated biomarkers of inflammation (CRP). (Moderate certainty of evidence; Strong recommendation)

We recommend against the use of tocilizumab among patients with COVID-19 infection who do not require oxygen. (Very low certainty of evidence; Strong recommendation)

Previous consensus issues

The high cost and limited availability of tocilizumab should be considered in our local setting. The indiscriminate use, potential adverse effects (i.e., leukemia, TB reactivation), risks, and lack of evidence of tocilizumab on COVID-19 patients who do not require oxygenation should be taken into consideration as well. Tocilizumab may also be administered in the earlier part of the therapy in order to prevent the occurrence of cytokine storm.



What's new in this version?

This version includes three (3) RCTs (Derde 2021, Hamed 2021, and Hermine 2021) as well as an update on previous trials (Hermine 2020).

Key Findings

Fifteen (15) randomized controlled clinical trials (RCTs) (N = 8,937) that investigated the effectiveness of tocilizumab among confirmed hospitalized COVID-19 patients compared to placebo and/or standard of care was found. Tocilizumab significantly reduced all-cause mortality, time to clinical improvement, and the need for mechanical ventilation, with no significant increase in adverse events among hospitalized patients.

Introduction

Tocilizumab is a humanized anti-IL-6Ra monoclonal antibody globally approved for management of various types of arthritis. IL-6 is an important cytokine in both acute and chronic inflammation. Tocilizumab binds to the IL-6 receptors, thus inhibiting inflammation by reducing circulating neutrophils, neutrophil infiltration, circulation of dendritic cells, and serum macrophage migration. Safety of tocilizumab intravenous and subcutaneous administration was reported with the most common adverse events being infections such as pneumonia and cellulitis. Elevated liver enzyme levels, neutropenia, and changes in lipid levels were reported. Serious adverse effects include myocardial infarction and stroke.[1]

Hyperinflammation contributes to the severity of COVID-19.[2] IL-6 is an important prognostic marker for survival of the disease. It is also independently associated with the severity and predictive outcome of ventilation and organ damage.[3] Recent WHO Therapeutic and COVID-19 Living Guideline strongly recommends the use of tocilizumab in severe or critical patients.

Review Methods

A systematic search was done from the date of last search May 18, 2021 until September 15, 2021 to check for new trials in the COVID-NMA living data. Trials found in the COVID-NMA were included. Search was done in Medline, Cochrane Library, Google scholar using free text, MeSH terms and advance search using the terms coronavirus infections, COVID-19, severe acute respiratory syndrome coronavirus 2, and tocilizumab. Screening was done in various trial registries for ongoing trials. Preprint search was also done in medrxiv, chinaxiv and biorxiv. Randomized controlled trials on tocilizumab compared to placebo or standard of care on COVID-19 patients, regardless of severity were included.

Results

Characteristics of included studies

Fifteen (15) randomized controlled clinical trials (RCTs) (N = 8,937) that evaluated the effectiveness of tocilizumab among confirmed hospitalized COVID-19 patients compared to placebo or standard of care were found. All of the trials reviewed were also included in the COVID-NMA Living Data which was last updated on September 15, 2021. Three (3) of the 15 trials were preprints. The summary on the characteristics of included studies can be found in Appendix 2.

Four (4) studies were multinational trials [5-8], while the rest were conducted in the United Kingdom [9], France [10,11], Italy [12], USA [13], China [14], Brazil [15], India [16], the Netherlands [17], Iran [18], and Dubai.[19] Study participants in all trials were suspected/confirmed COVID-19 patients, 18 years old and above, and with either presence of pulmonary infiltrates or requiring supplemental oxygen. Six (6) trials [7,10-13,18] excluded



patients on mechanical ventilation at the start of the trial, while two (2) trials [10,11] enrolled critical patients admitted in the intensive care unit that were receiving respiratory or cardiovascular organ support. Four (4) trials included elevated laboratory markers such as C-reactive protein (CRP), d-dimer, and ferritin in their inclusion criteria.[12-14,18] Standard of care varied per study but usually involved the administration of anticoagulants, steroids, or anti-viral drugs (e.g., darunavir/cobicistat, darunavir/ritonavir, lopinavir/ritonavir, or remdesivir).[5,8,12] Two (2) of the trials included steroids with tocilizumab as intervention.[11,19] All studies evaluated tocilizumab given via the intravenous route. There were no studies that evaluated tocilizumab given subcutaneously. There were only few moderate cases (not requiring oxygen) (121/8,937, 1.35%) that were included in these trials.

Overall Certainty of Evidence

Eleven (11) out of 15 trials were open label trials and there was an imbalance in the administration of steroids and anti-viral drugs as part of the standard of care among the participants of the studies. There were serious risk of bias, inconsistency, and imprecision in one important outcome (adverse events), however, given that the adverse events were not rated as critical outcomes to be included in the decision making by the panel, the overall quality of evidence is retained to be moderate. The risk of bias summary is shown in Appendix 4. The GRADE evidence summary is presented in Appendix 5.

Effectiveness Outcomes

The COVID-NMA project provided pooled analysis for clinical improvement, time to clinical improvement, time to death, adverse events, and serious adverse events (last updated July, 7, 2021).[20] Results on mortality and the initiation of mechanical ventilation were also pooled showing significant benefit in reducing these outcomes.

Mortality

Based on 14 RCTs (N = 8,489), tocilizumab reduced all-cause mortality at day 14 to day 90 followup compared to standard of care (RR 0.88, 95% CI 0.82-0.94, $l^2=0\%$). Tocilizumab significantly reduced all-cause mortality at day 28 (RR 0.88, 95% CI 0.82-0.95, $l^2=7\%$) but had no significant effect on all-cause mortality at day 90 (RR 0.88, 95% CI 0.76-1.02, $l^2=0\%$). Pooled effect of tocilizumab based on co-administration of steroids also significantly reduced mortality however, it was noted to have borderline heterogeneity with RR 0.86, 95% CI 0.79-0.93, $l^2=43\%$, P=0.07.

Subgroup analyses on the effect of tocilizumab on mortality stratified according to oxygen requirement and co-administration of steroids were done (Appendix 5). Subgroup according to oxygen requirement did not show significant benefit across groups, including those requiring oxygen supplementation (RR 0.89, 95% CI 0.76-1.04, $l^2=0\%$); those requiring non-invasive ventilation (RR 0.89, 95% CI 0.79-1.00, $l^2=0\%$); and those requiring invasive mechanical ventilation (RR 0.98, 95% CI 0.82-1.16, $l^2=0\%$). A subgroup analysis by co-administration of steroids demonstrated significant reduction in mortality among patients who were also given steroids (RR 0.80, 95% CI 0.73-0.88, $l^2=17\%$) however, there was no significant benefit among patients who were not given steroids (RR 1.08, 95% CI 0.91-1.29, $l^2=0\%$).

Clinical improvement

Eight (8) RCTs (N = 5,585) showed no significant difference in clinical improvement at day 28 among patients given tocilizumab compared to standard of care with borderline heterogeneity (RR 1.06, 95% CI 0.99-1.12, I²=39.6%, p=0.07). There was a significantly shorter time to clinical improvement among those given tocilizumab (HR 1.25, 95% CI 1.14-1.38, I²=9.9%).

Need for mechanical ventilation



There was significant reduction in the need for mechanical ventilation among patients given tocilizumab compared to standard of care (RR 0.78, 95% CI 0.68-0.89, I²=0%; 9 RCTs).

Safety Outcomes

There was no significant difference in the risk for adverse events (RR 1.23, 95% CI 0.93-1.62, I^2 =81.3%) and serious adverse events between the tocilizumab group and the control group (RR 0.92, 95% CI 0.77-1.08, I^2 =0%), however, there is significant heterogeneity with the risk for adverse events. Common adverse events noted with tocilizumab were abnormal liver function tests, leukopenia, and neutropenia.[12-15] There was no evidence for increased risk for serious secondary infection.[5-7,10,13]

Recommendations from Other Groups

Regulatory Agency	Recommendation
Infectious Diseases Society of America (IDSA) (as of September 14, 2021)	Suggests the use of tocilizumab in adult hospitalized patients with progressive severe or critical COVID-19 who have elevated markers in addition to standard of care (i.e., steroids). Severe COVID-19 cases are patients with SpO2 ≤ 94% on room air, while critical cases are those on mechanical ventilation and ECMO (Low certainty of evidence, Conditional recommendation).[21]
US National Institutes of Health (NIH) (as of September 15, 2021)	Recommends the use of tocilizumab in combination with dexamethasone in hospitalized patients exhibiting rapid respiratory decompensation. These are COVID-19 patients that are: (1) admitted to an ICU unit within 24 hours and who require mechanical ventilation, non-invasive ventilator (NIV) or high-flow nasal cannula (HFNC) and (2) non-ICU patients but requiring NIV or HFNC AND have significantly increased markers of inflammation (Moderate certainty of evidence, other randomized trials or subgroup analyses of randomized trials).[22]
World Health Organization (WHO) (as of September 24, 2021)	Recommends treatment with the use of IL-6 receptor blocker (tocilizumab or sarilumab) in severe to critical COVID-19 patients, with recommendation to give both corticosteroids and IL-6 receptor blocker in the said patients. This is based on a high certainty of evidence for mortality and mechanical ventilation.[4]

Table 1. Summary of Recommendations from Other Groups

Research Gaps

As of September 16, 2021, there are 18 ongoing clinical trials on tocilizumab for COVID-19 patients registered on *clinicaltrials.gov* and EU Clinical Trials Register. Four (4) of these trials have been completed and are awaiting results.



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Appendix 1. Evidence to Decision

Table 1. Summary of initial judgements prior to the panel discussion (N = 9)

FACTORS			RESEARCH EVIDENCE/ADDITIONAL CONSIDERATIONS				
Problem	No	Yes (9)					
Benefits	Large (2)	Moderate (7)	Small	Uncertain			 Tocilizumab reduces all-cause mortality with significant reduction in mortality on subgroup analysis by co-administration of steroids There was significant reduction in the need for mechanical ventilation
Harm	Large	Small (7)	Uncertain (2)				• Tocilizumab showed no significant difference in the risk for adverse events and serious adverse events between the 2 groups
Certainty of Evidence	High (1)	Moderate	Low (3)	Very low (5)			• The overall quality of evidence is very low due serious risk of bias, inconsistency and imprecision in one important outcome.
Balance of effects	Favors drug (9)	Does not favor drug	Uncertain				• Patients given tocilizumab had significant benefit in all-cause mortality and need for mechanical ventilation, with no significant increase in adverse events.
Values	Important uncertainty or variability (1)	Possibly important uncertainty or variability (4)	Possibly NO important uncertainty or variability (4)	No important uncertainty or variability			
Resources Required	Uncertain	Large cost (9)	Moderate Cost	Negligible cost	Moderate savings	Large savings	 Cost of Php 28,830.84 for retail price of 400mg/20ml, 20ml vial; total drug regimen cost per patient per treatment course would be Php 57,661.68



						 Issues on drug shortage and high demand
Certainty of evidence of required resources	No included studies (1)	Very low (1)	Low	Moderate (4)	High (3)	 The cost is regulated by Presidential Executive Order 104
Cost effectiveness	No included studies (5)	Favors the comparison (1)	Does not favor either the intervention or the comparison	Favors the intervention (3)		
Equity	Uncertain (2)	Reduced (3)	Probably no impact	Increased (4)		
Acceptability	Uncertain (3)	No	Yes (6)			
Feasibility	Uncertain (2)	No	Yes (7)			



Appendix 2. Search Yield and Results

		DATE AND TIME	RES	ULTS	
DATABASE	SEARCH STRATEGY / SEARCH TERMS	OF SEARCH	Yield	Eligible	
Medline	{"Coronavirus Infections"[Mesh] OR "Coronavirus"[Mesh] OR coronavirus OR novel coronavirus OR NCOV OR "COVID-19" [Supplementary Concept] OR covid19 OR covid 19 OR covid-19 OR "severe acute respiratory syndrome coronavirus 2" [Supplementary Concept] OR severe acute respiratory syndrome coronavirus 2 OR SARS2 OR SARS 2 OR SARS COV2 OR SARS COV 2 OR SARS-COV-2} AND Tocilizumab	9/7/2021 5:09PM	16	13	
CENTRAL	MeSH descriptor: [Coronaviridae Infections] explode all trees OR MeSH descriptor: [Coronavirus] explode all trees OR coronavirus OR novel coronavirus OR NCOV OR covid19 OR covid 19 OR covid-19 OR severe acute respiratory syndrome coronavirus 2 OR SARS2 OR SARS 2 OR SARS COV2 OR SARS COV 2 OR SARS-COV-2} AND Tocilizumab	9/7/2021 9:05PM	24	12	
COVID-NMA Initiative	Tocilizumab	9/7/2021 4:46PM	17	15	
Google Scholar	Tocilizumab AND COVID AND randomized trial	9/7/2021 920PM	13	10	
		T	1	I	
ClinicalTrials.gov	COVID-19 COVID-19 Pneumonia, Investigational Trials, Tocilizumab	9/15/2021 9:30PM	21	14	
Chinese Clinical Trial Registry	COVID Tocilizumab	9/15/2021 9:40PM	3	0	
EU Clinical Trials Register	COVID Tocilizumab	9/15/2021 9:50PM	6	4	
Republic of Korea - Clinical Research Information Service	COVID Tocilizumab	9/15/2021 9:52PM	0	0	
Japan Primary Registries Network/ NIPH Clinical Trials Search	COVID Tocilizumab	9/15/2021 9:53 M	0	0	
CenterWatch	COVID Tocilizumab	9/15/2021 9:56PM	0	0	
			1		
chinaxiv.org	COVID Tocilizumab	9/15/2021 10:01PM	1	0	
Medrxiv.org	COVID Tocilizumab Filter: May 18-Sept 7	9/15/2021 10:23PM	5	5	
Biorxiv.org	COVID Tocilizumab Filter: May 18-Sept 7	9/15/2021	0	0	



Title/Author	Study Design	Country	Population	Intervention Group(s)	Control	Outcomes
Horby 2021 RECOVERY	Open label RCT	United Kingdom	Suspected or confirmed COVID-19 patients N = 4,116	Tocilizumab 8 mg/kg	Standard care	 All-cause mortality at day 28 Time to discharge Receipt of invasive mechanical ventilation Use of non-invasive respiratory support Time to successful cessation of invasive mechanical ventilation Use of renal dialysis/hemofiltration Major cardiac arrhythmia Serious adverse events
Gordon 2020 REMAP-CAP	Adaptive RCT	United Kingdom, France, the Netherlands, Australia	ICU admitted critical COVID-19 patients AND receiving respiratory or cardiovascular organ support N = 755	Group 1: Tocilizumab 8 mg/kg Group 2: Sarilumab	Standard care	 Respiratory and cardiovascular organ support- free days Survival Time to ICU discharge Time to hospital discharge WHO scale at day 14 Progression to invasive mechanical ventilation, ECMO or death Serious adverse events
Hermine 2020	Open label RCT	France	Moderate, severe or critical COVID-19 patients with oxygen levels of 3 L/min or higher but without non-invasive ventilation (NIV) or mechanical ventilation (MV) N = 131	Tocilizumab 8 mg/kg	Usual care	 Mortality on day 4 and day 14 Mechanical ventilation on day 4 and day 14 Clinical status (WHO CPS) at day 7 and day 14 Overall survival Time to discharge Time to oxygen supply independency C-reactive protein levels Adverse events
Rosas 2020 COVACTA	Double-blind, placebo controlled RCT	Canada, Denmark, France, Germany, Italy, Netherlands, Spain, UK, USA	Severe COVID-19 patients N = 452	Tocilizumab 8 mg/kg	Placebo	 Clinical status at day 28 Mortality Ventilator free days Time to improvement Time to hospital discharge Adverse events
Salama 2020	Double-blind, placebo controlled RCT	USA, Mexico, Kenya, South Africa, Peru, Brazil	Hospitalized COVID-19 pneumonia patients not on continuous	Tocilizumab 8mg/kg	Placebo	 Invasive mechanical ventilation or ECMO Mortality Time to hospital discharge or readiness for discharge

Appendix 3: Characteristics of Included Studies



Salvarani 2020	Open-label	Italy	positive airway pressure, bilevel positive airway pressure, or mechanical ventilation. N = 388 Non-ICU COVID-19	Tocilizumab	Standard of care	 Time to at least a two-category improvement in clinical status Time to clinical failure Clinical worsening at day 14:
	RCI		patients. N = 126	8 mg/kg		 Admission to ICU with mechanical ventilation Death from any cause PaO2/FIO2 ratio less than 150 mmHg
Stone 2020	Double-blind, placebo controlled RCT	USA	Confirmed COVID-19 patients not on oxygen above 10 L/minute N = 243	Tocilizumab 8 mg/kg	Placebo	 Mortality Mechanical ventilation Clinical worsening Time to improvement Time to death Duration of supplemental O2 Admission to ICU
Wang 2020	Open-label RCT	China	Moderate or severe COVID-19 patients with elevated IL-6. N = 65	Tocilizumab	Standard care	 Cure rate Recovery rate of hypoxia over 14 days, Worsening rate of hypoxia during hospitalization, Duration of hospital stay, Time to negative virus load.
Veiga 2021	Open-label RCT	Brazil	Severe or critical COVID- 19 patients N = 129	Tocilizumab 8 mg/kg	Standard care	 Clinical status at Day 15 All-cause mortality In-hospital mortality Sequential organ failure assessment score Clinical status at day 8 and day 29 Ventilator-free days within 29 days Time to independence from supplemental oxygen Duration of hospital stay
Soin 2021 (COVINTOC)	Open-label RCT	India	Moderate to severe Covid-19 patients Moderate – RR 15-30 AND SpO2 90-94% Severe- RR>30 OR SpO2 <90% OR ARDS OR septic shock	Tocilizumab 6 mg/ kg	Standard Care	 Clinical progression Mortality Clinical improvement Time to clinical improvement Ventilator free days Organ failure free days ICU admission Time to hospital discharge Time to negative result on RT-PCR Adverse events



			N = 183			Serious adverse events
Rutgers 2021 (<i>Pre-print</i>)	Open label RCT	The Netherlands	Hospitalized COVID 19 patients with the following conditions: Need for supplemental Oxygen Ferritin >2000 ug/l or doubling serum ferritin in 20-48 hours N = 354	Tocilizumab 8 mg/kg	Standard of care	 30-day mortality Duration of hospital stay ICU admission Duration of ICU stay Duration of mechanical ventilation Time to mechanical ventilation Time to death
Talaschian 2021	Double blind RCT	Iran	 COVID-19 patients with the following conditions: Elevated CRP (>10 mg/L)/ IL 6 (> 18 pg/ml) / Lymphopenia (WBC< 1100/MCL) O2 sat <93% or RR >24 Not connected to mechanical ventilator Not responding to standard COVID-19 treatment N = 40 	Tocilizumab 8 mg/kg	Standard of care	 Clinical improvement 28-day mortality Time to improvement
New Studies Add	led (as of Sept	15, 2021)	1	1	1	1
Derde 2021 (Pre-print)	Open label, Adaptive RCT	US, Australia, Belgium, Canada, Croatia, Germany, Hungary, Ireland	ICU admitted critical COVID-19 patient AND receiving respiratory or cardiovascular organ	Group 1: Tocilizumab 8 mg/kg Group 2: Sarilumab Group 3: Anakinra	Standard care	 Respiratory and cardiovascular organ support- free days Survival In-bospital mortality 90 days
		Netherlands, New Zealand, Portugal, Romania, Spain, UK	support N = 2,274	Group 4: Interferon B1a		 Time to ICU discharge Time to hospital discharge
Hamed 2021	Open label RCT	Dubai	Hospitalized COVID-19 patients AND lung infiltrates >50% of lung	Group 1: Methylprednisolone	Historical control group	 All-cause mortality day 45 Admission to ICU Length of ICU stay



			fields within 48 hrs admission, O2 saturation <93% at rest on room air N = 76	Group 2: Methylprednisolone and Tocilizumab		 Invasive ventilation Days on ventilation Length of hospital stay
Hermine 2021 (CORIMUNO- TOCI-DEX) (Pre-print)	Open label RCT	France	Moderate to severe COVID-19 patients requiring oxygen but without ventilation support, high flow or mech vent, WHO class 5 N = 453	Tocilizumab 8 mg/kg at Day 1 PLUS Dexamethasone 10 mg/d for 5 days and tapering up to 10 days	Dexamethasone 10mg/d for 5 days and tapering up to 10 days	 Survival without mechanical ventilation at day 14 WHO-CPS progression Time to oxygen supply independency Time to hospital discharge Adverse events



Appendix 4. Study Appraisal





Appendix 5. GRADE Evidence Profile Author(s): K. Relato; I. Cabaluna; A. Garcia Question: Tocilizumab compared to standard of care in Covid-19

Bibliography: https://covid-nma.com/

			Certainty assess			№ of patients		Effect				
Nº of studies	Study design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Tocilizumab	Standard of care	Relative (95% Cl)	Absolute (95% Cl)	Certainty	Importance
All-cause mortality (follow up: range 14 days to 90 days)												
14	randomised trials	serious ^a	not serious	not serious	not serious	none	1214/4475 (27.1%)	1165/4014 (29.0%)	RR 0.88 (0.82 to 0.94)	35 fewer per 1,000 (from 52 fewer to 17 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
Clinical Improvement at Day 28												
8	randomised trials	serious ^a	not serious	not serious	not serious	none	1778/2952 (60.2%)	1434/2673 (53.6%)	RR 1.06 (0.99 to 1.12)	32 more per 1,000 (from 0 fewer to 70 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Time to Clinical Improvement												
7	randomised trials	serious ^a	not serious	not serious	not serious	none	2183 participants	1325 participants	-	HR 1.25 higher (1.14 higher to 1.38 higher)	⊕⊕⊕⊖ MODERATE	CRITICAL
Need for mechanical ventilation												
9	randomised trials	serious ^a	not serious	not serious	not serious	none	342/2741 (12.5%)	408/2624 (15.8%)	RR 0.78 (0.68 to 0.89)	34 fewer per 1,000 (from 50 fewer to 17 fewer)	⊕⊕⊕⊖ MODERATE	CRITICAL
Adverse	Events											
8	randomised trials	serious ^a	serious ^b	not serious	serious °	none	524/1033 (50.7%)	292/681 (42.9%)	RR 1.23 (0.93 to 1.62)	99 more per 1,000 (from 30 fewer to 266 more)	⊕○○○ VERY LOW	IMPORTANT
Serious	Adverse Even	ts										
10	randomised trials	serious ^a	not serious	not serious	serious °	none	241/1419 (17.0%)	164/1113 (14.7%)	RR 0.92 (0.77 to 1.08)	12 fewer per 1,000 (from 34 fewer to 12 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
CI: Confide	ence interval; R	RR: Risk ratio; HR:	Hazard Ratio									
Explana a. some con b. Heteroger c. Wide conf	ntions cerns due to imb neity: I2=81.3% idence interval w	alance in the adminis	stration of steroids ar efit and harm.	nd antivirals								



Appendix 6. Forest Plots

	Teeilinumeh		Stondard of Core		Diale Datia		Diels Detie	
	rocilizu	map	Standard of G	Jare		RISK RAUO	RISK RAUO	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl	
Derde 2021	317	943	150	406	17.2%	0.91 [0.78, 1.06]	-	
Gordon 2021	98	366	142	412	11.0%	0.78 [0.63, 0.96]	+	
Hamed 2021	2	26	5	27	0.4%	0.42 [0.09, 1.95]		
Hermine 2020	7	64	11	67	0.9%	0.67 [0.28, 1.61]		
Hermine 2021	18	224	24	226	2.0%	0.76 [0.42, 1.35]		
Horby 2021	621	2022	729	2094	58.8%	0.88 [0.81, 0.96]		
Rosas 2021	58	301	28	151	3.1%	1.04 [0.69, 1.56]		
Rutgers 2021	21	174	34	180	2.7%	0.64 [0.39, 1.06]		
Salama 2020	29	259	15	129	1.6%	0.96 [0.54, 1.73]		
Salvarani 2020	2	60	1	66	0.1%	2.20 [0.20, 23.65]		
Soin 2021	13	90	15	90	1.2%	0.87 [0.44, 1.72]		
Stone 2020	9	161	3	82	0.3%	1.53 [0.43, 5.49]		
Talaschian 2021	5	20	2	20	0.2%	2.50 [0.55, 11.41]		
Veiga 2021	14	65	6	64	0.5%	2.30 [0.94, 5.61]		
Total (95% CI)		4775		4014	100.0%	0.88 [0.82, 0.94]	*	
Total events	1214		1165					
Heterogeneity: Chi ² =	12.85, df:	= 13 (P :	= 0.46); I ² = 0%	, ,				7
Test for overall effect:	Z = 3.62 (P = 0.00)03)				U.U1 U.1 1 1U 1U Equation of the second	U
	,						Favours (experimental) Favours (control)	

Figure 1. Pooled effect of tocilizumab on all-cause mortality (day 14 to day 90)

	Experim	ental	Contr	ol		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Gordon 2021	98	366	142	412	13.5%	0.78 [0.63, 0.96]	-=-
Hermine 2020	7	64	8	67	0.8%	0.92 [0.35, 2.38]	
Hermine 2021	15	224	19	226	1.9%	0.80 [0.42, 1.53]	
Horby 2021	621	2022	729	2094	72.4%	0.88 [0.81, 0.96]	
Rosas 2021	58	301	28	151	3.8%	1.04 [0.69, 1.56]	+
Rutgers 2021	21	174	34	180	3.4%	0.64 [0.39, 1.06]	_
Salama 2020	26	259	11	129	1.5%	1.18 [0.60, 2.31]	_
Salvarani 2020	2	60	1	66	0.1%	2.20 [0.20, 23.65]	
Soin 2021	13	90	15	90	1.5%	0.87 [0.44, 1.72]	
Stone 2020	9	161	3	82	0.4%	1.53 [0.43, 5.49]	
Talaschian 2021	5	20	2	20	0.2%	2.50 [0.55, 11.41]	
Veiga 2021	14	65	6	64	0.6%	2.30 [0.94, 5.61]	
Total (95% CI)		3806		3581	100.0%	0.88 [0.82, 0.95]	*
Total events	889		998				
Heterogeneity: Chi ² =	11.87, df=	= 11 (P =	= 0.37); I ²				
Test for overall effect:	Z = 3.15 (P = 0.00	2)				Eavours (experimental) Eavours (control)
							ratears [experimental] ratears [control]

Figure 2. Pooled effect of tocilizumab on all-cause mortality day 28

	Tocilizumab		Standard of Care			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Derde 2021	317	943	150	406	77.9%	0.91 [0.78, 1.06]	
Hamed 2021	2	26	5	27	1.8%	0.42 [0.09, 1.95]	
Hermine 2020	7	64	11	67	4.0%	0.67 [0.28, 1.61]	
Hermine 2021	18	224	24	226	8.9%	0.76 [0.42, 1.35]	-+-
Salama 2020	29	259	15	129	7.4%	0.96 [0.54, 1.73]	-+-
Total (95% CI)		1516		855	100.0%	0.88 [0.76, 1.02]	•
Total events	373		205				
Heterogeneity: Chi ² =	1.80, df=	4 (P = 0).77); I² = 0%				
Test for overall effect	Z = 1.71 (P = 0.09	3)				Favours [experimental] Favours [control]

Figure 3. Pooled effect of tocilizumab on all-cause mortality day 90



					Clinical	improvement D28								
Study	Follow up days	Intervention 1	Intervention 2	r1/N1	r2/N2		A	в	Risk o C	of Bias D	E	Overall	Ris	k Ratio [95% Cl]
Mild to severe Salama C, 2020	28	Tocilizumab 8 mg/kg	Placebo	218/259	107/129	•	_	_			_		19.30%	1.01 [0.92, 1.12]
Mild to severe Stone JH, 2020 Moderate/severe	28	Tocilizumab 8mg/kg	Placebo	147/161	72/82		2	Ę	ļ,	2	2	2.1	19.44%	1.04 [0.95, 1.14]
Talaschian M, 2021 Moderate/severe	28	Tocilizumab 8 mg/kg once-off	Standard care	12/20	15/20	 4							1.78%	0.80 [0.52, 1.24]
Hermine O, 2020 Mild to critical	28	Tocilizumab 8mg/kg	Standard care	52/64	49/67							-	8.05%	1.11 [0.92, 1.34]
Rosas I, 2021 Moderate to critical	28	Tocilizumab 8mg/kg	Placebo	103/301	41/151	F=1					-		3.51%	1.26 [0.93, 1.71]
Horby P, 2021 Moderate to critical	28	Tocilizumab maximum 800 mg	Standard care	1150/2022	1044/2094	•					-		27.67%	1.14 [1.08, 1.21]
Veiga VC, 2021 Severe	29	Tocilizumab 8 mg/kg	Standard care	42/65	48/64	H#9							5.79%	0.86 [0.69, 1.08]
Salvarani C, 2020	30	Tocilizumab 8mg/kg	Standard care	54/60	58/66	•							14.46%	1.02 [0.91, 1.16]
Heterogeneity: Q = 13.13, p	$= 0.07; 1^2 = 39.6\%; \tau^2 = 0$.00												
Risk of bias ratings: Low Risk of Bias Some Concerns High Risk of Bias	Risk of A: Bias due to rand B: Bias due to devic C: Bias due to outc D: Bias due to outc E: Bias due to selec	Bias Domains: omization iton from intended interven ing data me measurement tion of reported result	Total: Total events:	2952 1778	2673 1434 ntervention 2 b	etter Interven 0.14 1.95 Risk Ratio	tion 1 t	petter		Data s	ource:	the COVID-NMA	1. A initiative (http	06 [0.99, 1.12] ps://covid-nma.com/)



	Tocilizumab		zumab Standard of Care			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Fixed, 95% Cl	IV, Fixed, 95% CI
Hamed 2021	6	26	9	27	2.3%	0.69 [0.29, 1.67]	
Hermine 2020	5	63	14	67	1.9%	0.38 [0.15, 0.99]	
Hermine 2021	15	224	20	226	4.3%	0.76 [0.40, 1.44]	
Horby 2021	215	1754	273	1800	63.9%	0.81 [0.68, 0.95]	
Rosas 2021	51	183	33	90	13.8%	0.76 [0.53, 1.09]	
Rutgers 2021	18	174	27	180	5.7%	0.69 [0.39, 1.21]	
Soin 2021	14	91	13	88	3.6%	1.04 [0.52, 2.09]	
Stone 2020	11	161	8	82	2.3%	0.70 [0.29, 1.67]	
Veiga 2021	7	65	11	64	2.3%	0.63 [0.26, 1.51]	
Total (95% CI)		2741		2624	100.0%	0.78 [0.68, 0.89]	•
Total events Heterogeneity: Chi² = Test for overall effect:	342 3.57, df = Z = 3.70 (i	8 (P = 0 P = 0.00	408).89); I² = 0%)02)				0.01 0.1 1 10 100 Favours [experimental] Favours [control]

Figure 5. Pooled effect of tocilizumab on initiation of mechanical ventilation



				Tir	me to clini	cal improvement							
Study	Study Duration days	Intervention 1	Intervention 2	N1	N2		A	Ri B	sk of C	Bias D	E	Overall	Estimate [95% CI]
Mild to severe													
Salama C, 2020 Mild to severe	60	Tocilizumab 8 mg/kg	Placebo	259	129	11	-		-				17.37% 1.14 [0.92, 1.41]
Stone JH, 2020	28	Tocilizumab 8mg/kg	Placebo	161	82	H H H							10.41% 1.06 [0.80, 1.41]
Hermine O, 2020	90	Tocilizumab 8mg/kg	Standard care	64	67								5.42% 1.52 [1.02, 2.27]
Rosas I, 2021	60	Tocilizumab 8mg/kg	Placebo	301	151	H=1							11.52% 1.31 [1.00, 1.71]
Salvarani C, 2020	30	Tocilizumab 8mg/kg	Standard care	60	66	H-1				-			6.18% 0.92 [0.63, 1.34]
Derde L, 2021	90	Tocilizumab 8 mg/kg once-off	Standard care	972	418	-					-		26.36% 1.31 [1.11, 1.55]
Gordon AC, 2021	90	Tocilizumab 8 mg/kg	Standard care	366	412	-					2		22.73% 1.41 [1.17, 1.69]
Heterogeneity: Q = 7.59, p =	0.27; $l^2 = 9.9\%$; $\tau^2 = 0.00$						-			_			
Risk of bias ratings: Low Risk of Bias Some Concerns High Risk of Bias	Risk of B A: Bias due to randon B: Bias due to deviati C: Bias due to deviati D: Bias due to outcom E: Bias due to selectic	as Domains: iization on from intended intervention j data re measurement on of reported result	Tota	al: 2183 Interv	1325 ention 2 better	Intervent	tion 1 be	etter		Data se	ource: 1	ihe COVID-NMA	1.25 [1.14, 1.38] initiative (https://covid-nma.com/)
						Hazard Ratio							

Figure 6. Pooled effect of tocilizumab on time to clinical improvement (hazard ratio) (Source: covid-nma.com)

Study	Study Duration days	Intervention 1	Intervention	2 r1/N1	r2/N2				A	Б В	Risk of C	Bias D	E	Overall	R	isk Ratio [95% Cl]
Mild to severe																
Salama C, 2020	60	Tocilizumab	Placebo	127/259	67/129		÷								17.30%	0.94 [0.77, 1.16]
Mild to severe		8 mg/kg					1				•					
Stone JH, 2020	28	Tocilizumab	Placebo	37/161	19/82		н і н								12.13%	0.99 [0.61, 1.61]
Moderate/severe		8mg/kg					1		-	-	-	-				
Hermine O, 2020	90	Tocilizumab	Standard care	28/64	36/67		H								14.60%	0.81 [0.57, 1.16]
Moderate/severe		8mg/kg					1									
Wang D, 2021	14	Tocilizumab	Standard care	20/33	4/32			-							5.90%	4.85 [1.86, 12.63]
Mild to critical		400 mg									-					
Rosas I, 2021	60	Tocilizumab	Placebo	237/301	118/151		÷.								18.64%	1.01 [0.91, 1.12]
Moderate to critical		8mg/kg														
Soin AS, 2021	30	Tocilizumab	Standard care	33/90	22/90		÷								12.72%	1.50 [0.95, 2.36]
Moderate to critical		6 mg/kg/day							-							
Veiga VC, 2021	29	Tocilizumab	Standard care	29/65	21/64		÷.								12.92%	1.36 [0.87, 2.12]
Severe		8 mg/kg														
Salvarani C, 2020	30	Tocilizumab 8mg/kg	Standard care	13/60	5/66		-	+	-	-	-				5.79%	2.86 [1.08, 7.55]
Heterogeneity: Q = 21.02, p	$p = 0.00; ^2 = 81.3\%; \tau^2 = 0$.11					1					-				
Risk of bias ratings: Low Risk of Bias Some Concerns High Risk of Bias	Risk of A: Bias due to randt B: Bias due to devia C: Bias due to outer D: Bias due to outer E: Bias due to selec	Blas Domains: mization tion from intended interve ng data me measurement tion of reported result	Total: Total events: ntion	1033 524	8 681 292 Intervention	n 1 better 0.14	1.95 sk Ratio	Interventi	on 2 b	etter		Data so	ource:	the COVID-NM	1 A initiative (h	.23 [0.93, 1.62] ttps://covid-nma.com/)

Adverse events





					Serio	ous adverse	events						
Study	Study Duration	Intervention 1	Intervention 2	r1/N1	r2/N2				Ri	isk of	Bias		Bisk Batio 195% Cll
	days							A	в	C	DE	Overall	
Mild to severe													
Salama C, 2020 Mild to severe	60	Tocilizumab 8 mg/kg	Placebo	38/259	25/129		н						13.56% 0.76 [0.48, 1.20]
Stone JH, 2020	28	Tocilizumab 8mg/kg	Placebo	28/161	12/82	H	•	- 21	Ξ.	2.	22		7.37% 1.19 [0.64, 2.21]
Talaschian M, 2021	28	Tocilizumab 8 mg/kg once-off	Standard care	3/20	0/20	, <u> </u>	-	- 21	Ξ.	Ξ.	2.2	2	0.34% 7.00 [0.38, 127.32]
Hermine O, 2020	90	Tocilizumab 8mg/kg	Standard care	20/64	29/67	H B	4		2	2.	22		13.75% 0.72 [0.46, 1.14]
Wang D, 2021 Mild to critical	14	Tocilizumab 400 mg	Standard care	0/33	1/32	-	-	- 21	с.	20	22	2.1	0.28% 0.32 [0.01, 7.66]
Rosas I, 2021 Moderate to critical	60	Tocilizumab 8mg/kg	Placebo	113/301	62/151	+	H	- 21	Ξ.	Ξ.	22		49.34% 0.91 [0.72, 1.16]
Soin AS, 2021 Moderate to critical	30	Tocilizumab 6 mg/kg/day	Standard care	18/90	15/90	H	•	- 21	с.	2.	22		7.42% 1.20 [0.65, 2.23]
Veiga VC, 2021	29	Tocilizumab 8 mg/kg	Standard care	11/65	7/64	H		- 2	с.	24	22		3.66% 1.55 [0.64, 3.74]
Salvarani C, 2020 Severe/critical	30	Tocilizumab 8mg/kg	Standard care	1/60	2/66			- 01	Э.	21	22		0.51% 0.55 [0.05, 5.91]
Gordon AC, 2021	90	Tocilizumab 8 mg/kg	Standard care	9/366	11/412	H-4	-	- 21	Ξ.	Ξ.	22		3.77% 0.92 [0.39, 2.20]
Heterogeneity: Q = 6.95, p = 0	0.64; $l^2 = 0.0\%$; $\tau^2 = 0.00$									۰.		-	
Fisk of bias ratings: Low Risk of Bias Some Concerns High Risk of Bias	Risk of Bia A: Bias due to random B: Bias due to deviatio C: Bias due to missing D: Bias due to outcom E: Bias due to selectio	as Domains: ization n from intended interven data e measurement n of reported result	Total:	1419 241	1113 164 Intervention	0.14	Interv 1.95	rention 2 b	etter	C	Data source	e: the COVID-NM	0.92 [0.77, 1.08] A initiative (https://covid-nma.com/)
						Risk F	Ratio						



	Experim	ental	Contr	ol		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
1.1.1 Requiring Oxyg	jen Supple	mentati	on				
Hamed 2021	2	26	5	27	0.6%	0.42 [0.09, 1.95]	
Hermine 2020	7	64	8	67	1.0%	0.92 [0.35, 2.38]	
Hermine 2021	18	224	24	226	3.1%	0.76 [0.42, 1.35]	—
Horby 2021	175	935	202	933	26.6%	0.86 [0.72, 1.04]	-
Rosas 2021	6	78	2	44	0.3%	1.69 [0.36, 8.03]	
Salama 2020	26	259	11	129	1.9%	1.18 [0.60, 2.31]	_
Salvarani 2020	2	60	1	66	0.1%	2.20 [0.20, 23.65]	
Stone 2020	9	161	3	82	0.5%	1.53 [0.43, 5.49]	
Subtotal (95% CI)		1807		1574	34.3%	0.89 [0.76, 1.04]	•
Total events	245		256				
Heterogeneity: Chi² =	= 3.89, df = 1	7 (P = 0	.79); I ^z = I	0%			
Test for overall effect	: Z = 1.44 (F	P = 0.15)				
1.1.2 Requiring non-	invasive ve	entilatio	n				
Horby 2021	296	819	350	867	44.7%	0.90 (0.79, 1.01)	
Rosas 2021	13	94	8	39	1.5%	0.67 (0.30, 1.50)	
Subtotal (95% CI)		913		906	46.2%	0.89 [0.79, 1.00]	•
Total events	309		358				
Heterogeneity: Chi ^z =	= 0.48, df = 1	1 (P = 0	49); I ^z = I	0%			
Test for overall effect	: Z = 1.93 (F	P = 0.05)				
1.1.3 Requiring invas	sive mecha	nical ve	entilation	I			
Horby 2021	125	268	142	294	17.8%	0.97 [0.81, 1.15]	+
Rosas 2021	20	113	9	55	1.6%	1.08 [0.53, 2.22]	
Subtotal (95% CI)		381		349	19.4%	0.98 [0.82, 1.16]	♦
Total events	145		151				
Heterogeneity: Chi ² =	= 0.09, df = 1	1 (P = 0	.76); I ^z = I	0%			
Test for overall effect	: Z = 0.29 (F	P = 0.77)				
Total (95% CI)		3101		2829	100.0%	0.91 [0.83, 0.99]	•
Total events	699		765				
Heterogeneity: Chi ² =	= 5.29, df = 1	11 (P = I	0.92); I^z =	:0%			
Test for overall effect	: Z = 2.30 (F	P = 0.02)				U.U1 U.1 1 1U 1UU
Test for subgroup dif	ferences: C	Chi ² = 0.	87. df = 2	(P = 0.	.65), I ^z = 0	1%	ravours (experimental) ravours (control)

Figure 9. Pooled effect of tocilizumab on mortality according to oxygen requirement



	Experimental		al Standard of Care			Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl	
4.1.1 Steroids								-
Hamed 2021	2	26	5	27	0.6%	0.42 [0.09, 1.95]		
Hermine 2020	1	10	6	12	0.7%	0.20 [0.03, 1.40]		
Hermine 2021	18	224	24	226	2.9%	0.76 [0.42, 1.35]		
Horby 2021	457	1664	585	1721	70.2%	0.81 [0.73, 0.89]		
Rosas 2021	29	106	18	79	2.5%	1.20 [0.72, 2.00]		
Salama 2020	27	200	24	112	3.8%	0.63 [0.38, 1.04]		
Subtotal (95% CI)		2230		2177	80.7%	0.80 [0.73, 0.88]	•	
Total events	534		662					
Heterogeneity: Chi ² =	6.01, df=	5 (P = 0.	31); I ^z = 17%					
Test for overall effect:	Z=4.48 (F	• < 0.00	001)					
4.1.2 Without Steroid	1							
Hermine 2020	14	53	18	55	2.2%	0.81 [0.45, 1.45]		
Horby 2021	139	357	127	367	15.3%	1.13 [0.93, 1.36]		
Rosas 2021	29	188	10	65	1.8%	1.00 [0.52, 1.94]		
Salama 2020	2	49	0	16	0.1%	1.70 [0.09, 33.67]		
Subtotal (95% CI)		647		503	19.3%	1.08 [0.91, 1.29]	•	
Total events	184		155					
Heterogeneity: Chi ² =	1.26, df = 3	3 (P = 0.	.74); I² = 0%					
Test for overall effect:	Z = 0.87 (F	P = 0.39)					
Total (95% CI)		2877		2680	100.0%	0.86 [0.79, 0.93]	•	
Total events	718		817					
Heterogeneity: Chi ² =	15.82, df=	= 9 (P = I	0.07); I² = 43%					
Test for overall effect:	Z=3.60(F	^o = 0.00	03)				Favours [experimental] Favours [control]	
Test for subgroup diff	ferences: C	≥hi ² = 8.9	50, df = 1 (P =	0.004),	I ^z = 88.29	%	r arears (experimental) i aroars (control)	

Figure 10. Pooled effect of tocilizumab on mortality with and without co-administration of steroids



Appendix 7. Pooled Results of Trials

Outcomes	Pooled Relative Risk	95% CI	Certainty of evidence (GRADE)
All-cause mortality (14 RCTs, N = 8,489)	0.88	0.82 to 0.94	Moderate
Clinical improvement D28 (8 RCTs, N = 5,625)	1.06	0.99 to 1.12	Moderate
Need for mechanical ventilation (9 RCTs, N = 5,365)	0.78	0.68 to 0.89	Moderate
Time to clinical improvement (Hazard ratio) (7 RCTs, N = 3,508)	1.25	1.14 to 1.38	Moderate
Adverse events (8 RCTs, N = 1,714)	1.23	0.93 to 1.62	Very Low
Serious adverse events (10 RCTs, N = 2,532)	0.92	0.77 to 1.08	Moderate



Appendix 8. Subgroup Analysis

	Pooled Relative Risk	95% CI	Certainty of Evidence
By oxygen requirement			
Requiring O2 supplementation (8 RCTs, N = 3,381)	0.89	0.76 to 1.04	Low
Requiring non-invasive ventilation (2 RCTs, N = 1,819)	0.89	0.79 to 1.00	Low
Requiring mechanical ventilation (2 RCTs, N = 730)	0.98	0.82 to 1.16	Low
By co-administration with steroids			
With steroids (6 RCTs, N = 4,407)	0.80	0.73 to 0.88	Moderate
Without steroids (4 RCTs, N = 1,150)	1.08	0.91 to 1.29	Low



Appendix 9. Characteristics of Ongoing Studies

Study Title	Patients (n)	Interventions	Outcomes	Method
1. Tocilizumab in COVID-19 Pneumonia (TOCIVID-19)	No age limit, COVID-19 confirmed by RT-PCR, hospitalized due to clinical/instrumental diagnosis of pneumonia, oxygen saturation ≤93% or requiring oxygen therapy or mechanical ventilation either non-invasive or invasive (intubated) (N = 402)	Experimental: Tocilizumab 8 mg/kg (up to a maximum of 800mg per dose). A second administration (same dose) can be given after 12 hours if respiratory function has not recovered, at discretion of the Investigator.	Primary: Lethality rate two weeks after registration	Single-arm, open-label, parallel cohort
2. Efficacy of Early Administration of Tocilizumab in COVID-19 Patients <i>Terminated</i>	>18 years old, RT-PCR confirmed COVID-19 patients, hospitalized due to clinical or instrumental diagnosis, presence of ARDS, with at least one of the ff: temperature of > 38C in the past two days, CRP >/= 10mg/dl, CRP increase of at least 2x the basal value (N = 126)	Experimental: Tocilizumab within 8 hours from entering the study plus standard of care; 8 mg/kg IV up to a maximum of 800mg with repetition of the same dosage after 12 hours Control: Standard of care	Primary: Entry into intensive care with invasive mechanical ventilation or death from any cause or clinical aggravation	Randomized parallel, open label
3. A Study to Investigate Intravenous Tocilizumab in Participants with Moderate to Severe COVID-19 Pneumonia (MARIPOSA) Completed	>18 years old, confirmed COVID-19 by RT-PCR Severe patients: SpO2 = 93%<br or PaO2/FiO2 < 300 mmHg Moderate patients: CRP > 2 x upper limit of normal (ULN) is required	Experimental: Tocilizumab 4 mg/kg IV plus standard of care Active Comparator: Tocilizumab 8 mg/kg plus standard of care	 Primary:Area Under the Curve from Day 0-28 (AUC0- d28) of Tocilizumab, Secondary: Maximum Serum Concentration (Cmax) of Tocilizumab Clearance (CL) of Tocilizumab Volume of the Central Compartment (Vc) of Tocilizumab Serum Concentration of C-reactive Protein (CRP) Serum Concentration of Ferritin Serum Concentration of Soluble Interleukin-6 Receptor Serum Concentration of Interleukin-6 (IL-6) 	Randomized, parallel, open label
4. A Study to Evaluate the Efficacy and Safety of Tocilizumab in Hospitalized Participants With COVID-19 Pneumonia (EMPACTA)	>18 years old, hospitalized COVID-19 pneumonia confirmed by RT-PCR and radiographic imaging, spO2 <94% while on ambient air (N = 388)	Experimental: Tocilizumab 8 mg/kg maximum dose 800mg plus standard of care Control: Placebo plus standard of care	Primary: Cumulative proportion of participants requiring mechanical ventilation by day 28	Randomized Parallel, Double-blind, Placebo- controlled
5. A Study to Evaluate the Efficacy and Safety of Remdesivir Plus Tocilizumab Compared with Remdesivir Plus Placebo in Hospitalized Participants with Severe COVID-19 Pneumonia (REMDACTA)	>/= 12 years old, hospitalized with COVID-19 pneumonia confirmed by RT-PCR and evidenced by chest X-ray or CT scan Requiring > 6 L/min supplemental oxygen to maintain SpO2 > 93%	Experimental: Remdesivir plus Tocillizumab Control: Remdesivir plus Placebo	Primary: Time from randomization to hospital discharge or ready for discharge day 28	Randomized, parallel, doiuble-blind



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Completed	(N = 649)			
6. The Use of Tocilizumab in the Management of Patients Who Have Severe COVID- 19 With Suspected Pulmonary Hyperinflammation	>/= 18 years old, RT-PCR positive, acute respiratory failure radiologic pneumonia, breathing spontaneously >50% Oxygen and MEWS score >7, if intubated PaO2/Fio2 = 200 and PEEP 5 cm H2O (N = 500)	Experimental: Tocilizumab 8 mg/kg max 800mg Control: Placebo	Survival	Randomized, parallel, open label
7. Randomized, Unicentric, Open, Controlled Clinical Trial, in Phase III to Demonstrate the Effectiveness of Tocilizumab	18 -to 99 years old, bilateral pneumonia by SARS-COV-2 without response in 48 to 72 hours after starting the treatment according to local protocol, persistent elevated inflammatory markers (N = 60)	Experimental: Tocilizumab >75 kg = 600mg, <75kg = 400mg Control: Methylprednisolone 250mg IV for 3 days	Primary: Respiratory situation at 24 hours Secondary: Immune hyperactivation situation Mechanical ventilation In-hospital mortality	Randomized, parallel, open
8. Checkpoint Blockade in COVID-19 Pandemic (COPERNICO)	18 to 80 years old, confirmed COVID-19 with RT-PCR, or IgM, IgG, diagnostic confirmation of pneumonia on x-ray and CT scan, ARDS, SOFA score = 3,<br total lymphocyte <0.8x106/mL, SpO2 =92%, life expectancy of 10 years	Experimental: Tocilizumab 8 mg/kg plus Pembrolizumab 200mg Control: Standard of care	Primary: Percentage of patients with normalization of SpO2 ≥ 96% on room air	Randomized parallel, open- label
9. Low-dose Tocilizumab Versus Standard of Care in Hospitalized Patients With COVID-19 (COVIDOSE-2)	≥18 years old hospitalized, T ≥ 38 degrees C, positive test for active SARS-CoV-2 infection, radiographic evidence of infiltrates on chest radiograph (CXR) or computed tomography (CT) (N = 332)	Experimental sub study A: Tocilizumab 40mg Experimental sub study A: Tocilizumab 120mg Control A: Tocilizumab free standard of care Experimental sub study B: Tocilizumab 40mg Experimental sub study B: Tocilizumab 120mg Control B: Tocilizumab 400mg or 8mg/kg plus standard of care	Primary: Time to recovery	Randomized parallel, open label
10. COVID 19: Experimental use of tocilizumab (Roactemra®) in severe SARS-CoV-2 related pneumonia.	Positive COVID status as defined by positive RT-PCR, hospitalized patients aged ≥ 18 and ≤ 75 years old Signs of severe COVID-19 pneumonia (3 of the following): patient wheezing or unable to speak in full sentences while at rest/with minimal effort, respiratory rate >22, PaO2 <65 mmHg or SpO2 <90%, repeated chest imaging is significantly worsening despite being on standard of care, which may	Experimental: Tocilizumab Control: Standard of care	Clinical status assessed using a 7-category ordinal scale at Day 28	Randomized open label



	include anti-viral treatment, low dose steroids and antibiotics.			
11. A randomized clinical trial (IIIb) of eficacy of a single dose of Tocilizumab or a combination of Tocilizumab plus Vitamin D (single IM dose) for the treatment of the COVID-19 hyperimmune complication. Assessment of IL-6.	Moderate to severe COVID-19 patients (WHO severity scale 4- 7) needing oxygen therapy At least 2 of the following: Dímer D> 1.500, CRP> 60 or Ferritin >800 Alternatively, IL-6 >40	Experimental: Tocilizumab plus Vitamin D3 2000 IU Control: Tocilizumab	Global survival rate	Randomized, open label
12. A prospective, randomized, double blinded placebo-controlled trial to evaluate the efficacy and safety of tocilizumab in patients with severe COVID- 19 pneumonia (TOC-COVID)	≥ 18 years old, proof of SARS- CoV-2, severe respiratory failure: ambient air SpO2 ≤ 92% or need of ≥ 6L O2/min or NIV (non- invasive ventilation) or IMV (invasive mechanical ventilation)	Experimental: Tocilizumab Control: Placebo	Ventilator free days (d) (VFD) in the first 28 days after randomization	Randomized double-blind placebo- controlled
13. A multi-center, randomized, open-label study in patients with COVID-19 and respiratory distress not requiring mechanical ventilation, to compare standard-of-care with anakinra and tocilizumab treatment. The Immunomodulation-CoV Assessment (ImmCoVA) study	Age ≥18 years old, confirmed SARS-CoV-2 infection as determined by polymerase chain reaction (PCR) or other commercial or public health assay < 3 days prior to screening SARS-CoV-2 infection with duration at least 7 days (as determined by onset of symptoms), 5 liters/minute of oxygen for at least 8 hours to maintain SpO2 at ≥93%, CRP > 70 mg/L with no non-SARS- Cov2 infections, Ferritin > 500 µg/L, at least two points on a scale of 0-3 where 1 point is awarded for each value of: lymphocytes < 1x 10(9)/L; D- dimer ≥ 0.5 mg/L and; Lactate Dehydrogenase ≥ 8 microkatal/L	Experimental: Anakinra and Tocilizumab Control: standard of care	Time to recovery [Time Frame: Day 1 through Day 29]	Randomized, open label
14. Trial of Tocilizumab for Treatment of Severe COVID- 19: ARCHITECTS	Hospitalized with COVID-19 pneumonia, based on chest X- ray or CT scan AND Evidence of hyperinflammation: IL-6>40pg/mL (if available) OR CRP >2 mg/dL OR ferritin >2000 ng/mL AND iv. One or more of the following: impending need for requiring invasive or non-invasive mechanical ventilation OR shock requiring vasopressor (without evidence of bacterial / fungal infection) OR need for extracorporeal membrane oxygenation (ECMO) OR severe, refractor ARDS (PaO2/FiO2<200 mmHg)	Experimental: Tocilizumab Control: Placebo	Clinical status (on a 7-point ordinal scale) at day 28	Randomized, double blind Placebo control
15. CORIMUNO-19 - Tocilizumab Trial - TOCI (CORIMUNO-TOCI) 2	Patients included in the CORIMUNO-19 cohort Patients belonging to one of the 2 following groups:	Experimental: Tocilizumab 8mg/kg day 1, if no response second injection Day 3 Control:	Survival without needs of ventilator utilization WHO progression scale <=5	Randomized, parallel open label



	Group 1: Cases meeting all of the following criteria: Requiring more than 3L/min of oxygen, OMS/WHO progression scale = 5, No NIV or High flow Group 2: Cases meeting all of the following criteria Respiratory failure AND (requiring mechanical ventilation OR NIV OR High flow), WHO progression scale >=6, No do- not-resuscitate order (DNR order)	Standard of care	Cumulative incidence of successful tracheal extubation WHO progression scale <=7	
16. Treatment of COVID-19 Patients with Anti-interleukin Drugs (COV-AID)	Recent (≥ 6 days of flu-like symptoms or malaise yet ≤16 days of flu-like symptoms or malaise prior to randomization) infection with COVID-19. Confident COVID-19 diagnosis Presence of hypoxia signs of cytokine release syndrome Chest X-ray or CT scan showing bilateral infiltrates within last 2 days Admitted to specialized COVID- 19 ward or an ICU ward taking care of COVID-19 patients Age ≥ 18 years old	Experimental: Anakinra, Siltuximab, Tocilizumab Control: Placebo and standard of care	Time to Clinical Improvement	Randomized open label
17. COVID-19: Salvage Tocilizumab as a Rescue Measures (COVISTORM) Completed	Hospitalized with COVID-19 disease Age >/= 18 years old SARS-CoV-2 NhO positive SpO2 93% on ambient air or<br respiratory rate >30 /min Any 2 of the 4: P-IL-6 > 2 x ULN / P-ferritin > 2 x ULN / P-FIDD >1.5 mg/l / P- CRP >40 mg/l without obvious presence of bacterial infection (normal values: P -IL-6 <5.9 ng/l; P- ferritin, men 30-400 mikrog/l, women 13-150 mikrog/l ; P-FIDD (Fibrin degradation products, D- dimer) <0.5 mg/l; P-CRP <10 mg/l)	Experimental: Tocilizumab Control: Standard of care	Clinical status at day 28	Randomized Parallel Open Label
18. Efficacy of Tocilizumab in Modifying the Inflammatory Parameters of Patients With COVID-19 (COVITOZ-01) (COVITOZ- 01) <i>Terminated</i>	 > 18 years of age, mild-moderate SARS-CoV-2 pneumonia confirmed microbiologically ≤7 days before randomization, and presents: a. Basal oxygen saturation> 90% b. CURB-65 ≤1 c. PaO2 / FiO2≥300 or SatO2 / FiO2≥315 The patient is hospitalized or meets hospital admission criteria; the patient is not expected to enter the ICU or die in the next 24 hours. 	Experimental 1: Tocilizumab single dose and standard of care Experimental 2: Tocilizumab two doses and standard of care Control: Standard of care	Change in IL-12 values in the 3 study groups from the start of treatment (D0) and on days D + 1 and D + 3	Randomized single group open label