



## South African National Department of Health Rapid Review Report Component: COVID-19

## TITLE: TOCILIZUMAB FOR THE TREATMENT OF COVID-19: EVIDENCE REVIEW OF THE CLINICAL BENEFIT AND HARM

## Date: 26 May 2021 (third update of the initial 15 April 2020 rapid review)

#### **Key findings**

- We updated the rapid review of clinical evidence for the use of tocilizumab with or without other medicines in the management of hospitalised patients with severe COVID-19 requiring oxygen or ventilatory assistance.
- We identified 12 randomised controlled trials (RCTs), and a systematic review and meta-analysis that combined all 12 RCTs.
- Overall tocilizumab reduced all-cause mortality from 29.2% to 25.7% at 28 days: The absolute risk reduction was 3.5% (95% confidence interval (CI) 1.5% to 5.6%), and the relative risk (RR) 0.88 (95% CI 0.81 to 0.95, 11 RCTs, n = 6 937). The number needed to treat to prevent one additional death was 29 (95% CI 18 to 67).
- Tocilizumab was not associated with an increased risk of adverse events (RR 1.23; 95% CI 0.93 to 1.62), or serious adverse events (RR 0.92; 95% CI 0.77 to 1.08).
- We did not identify any reports on the use of tocilizumab in children with COVID-19.

NEMLC THERAPEUTIC GUIDELINES SUB-COMMITTEE RECOMMENDATION:							
	We recommend against	We suggest not to use the	We suggest using either	We suggest	We recommend		
	alternative	to use the alternative	alternative	(conditional)	(strong)		
Type of	(strong)	(conditional)	(conditional)	()	(00003)		
recommendation		X					
Recommendation: The sub-committee suggests not to use tocilizumab. Despite the reduction in death in the							
included trials, tocilizumab is not affordable at the current offered price.							
Rationale: A met	a-analysis of 11 rand	omised controlled tria	lls reporting mortality	showed that toci	lizumab, used in		

combination with corticosteroids, reduced all-cause mortality at day 28 from 29.2% to 25.7% amongst adult patients with COVID-19 with hypoxia and evidence of systemic inflammation (CRP  $\geq$  75mg/L), without an increase in clinically significant adverse events. However, the sub-committee expressed concerns regarding the budget impact and national supply of tocilizumab.

Level of Evidence: I High to moderate certainty evidence *Review indicator:* Reduction in price

**Therapeutic Guidelines Sub-Committee for COVID-19:** Andy Parrish (chair), Gary Reubenson (vice-chair), Marc Blockman, Karen Cohen, Andy Gray, Tamara Kredo, Renee De Waal, Gary Maartens, Jeremy Nel, Helen Rees.

**Note:** Due to the continuous emergence of new evidence, the rapid review will be updated if and when more relevant evidence becomes available. It was noted that, as of 26 May 2021, 46 clinical trials investigating the role of tocilizumab in the management of COVID-19 are registered on the International Clinical Trials Registry Platform (ICTRP), accessed from <a href="https://covid-nma.com/">https://covid-nma.com/</a>

Version	Date	Reviewer(s)	Recommendation and Rationale		
First	15 April 2020 RW, MB		Recommend against using tocilizumab in children or adult patients with COVID-19 as there is current		
			insufficient evidence to recommend routine use - consider in context of clinical trial setting.		
Second	17 November	RW, MB	Recommend against using tocilizumab in children or adult patients with COVID-19 as there is currently		
	2020		insufficient RCT evidence to recommend routine use - consider in context of clinical trial setting.		
Third	5 March 2021	RW, MB,	Suggest the use of tocilizumab in hospitalized hypoxic adult patients with COVID-19 and a CRP $\geq$ 75mg/L,		
		RdW, KC	but with concerns about affordability and possible supply constraints.		
Fourth	20 May 2021	RW, MB	No change		

## BACKGROUND

In patients infected with SARS-CoV-2, disease severity and outcomes are related to the characteristics of the immune response. <sup>1-6</sup> The median time from onset of symptoms of COVID-19 to the development of acute respiratory distress syndrome (ARDS) is as short as 8 days<sup>7</sup>. Interleukin (IL)-6 and other components of the inflammatory cascade play an important role in the inflammatory reaction and immune response<sup>8</sup>. However, excessive cytokine production ('cytokine storm') as part of a hyper-inflammatory response has been suggested as a cause of severe COVID-19.<sup>1-3</sup> There has been some controversy as to whether IL-6 constitutes one of the most important cytokines involved in COVID-19-induced cytokine storms and if there is a correlation between elevated IL-6 levels in patients with COVID-19 and the risks of respiratory failure and the requirement for ventilation.<sup>8, 9, 25</sup>

Retrospective case series and individual case reports from China identified that IL-6 blockade therapy may constitute a novel therapeutic strategy in patients with severe SARS-CoV-2 pneumonia.<sup>8, 10-13</sup>

Tocilizumab (TCZ) is a recombinant humanized monoclonal antibody against human IL-6 receptor of immunoglobulin IgG1 subtype. In South Africa, it is the only commercially available IL-6 inhibitor and is registered for use in the management rheumatoid arthritis. Tocilizumab specifically binds soluble and membrane-bound IL-6 receptors (sIL-6R and mIL-6R) and inhibits the associated signal transduction. As a result, there is biological plausibility associated with tocilizumab's importance in the management COVID-19. <sup>14</sup> However, the World Health Organization guideline provides no guidance on the use of tocilizumab currently, while the Australian and US National Institutes of Health (NIH) guidelines do allow for its use in patients who are within 24 hours of admission to the intensive care unit (ICU) and require invasive or noninvasive mechanical ventilation or high-flow oxygen (Appendix 1).

Existing RECOVERY RCT data was recently published in a peer-reviewed journal, thus warranting an update of this rapid review.

## **RESEARCH QUESTION:**

Should tocilizumab be used for managing severe COVID-19 (with or without elevated IL-6 levels) in patients requiring oxygen or ventilatory assistance?

## **METHODS**

This is the fourth iteration of this rapid review. The initial review was conducted in April 2020, for which we systematically searched four electronic databases (PubMed as well as the Epistemonikos, Cochrane COVID Study Register and Living mapping and living network meta-analysis of COVID-19 studies databases).

The search strategy for the first update focused on randomised controlled trials and systematic reviews as such data had been published subsequent to the initial review. To this end, the Epistemonikos L\*OVE evidence platform (<u>https://app.iloveevidence.com/</u>) was searched for randomised controlled trials and systemic reviews. Relevant records were extracted in a narrative table of results (see update of 26 November 2020).

The second update focused on new studies published after 15 November 2020. The Epistemonikos L\*OVE evidence platform (<u>https://app.iloveevidence.com/</u>) was searched for randomised controlled trials and systemic reviews (see update of 15 April 2021).

This third update was triggered on account of the formal publication of the RECOVERY trial<sup>32</sup> data for tocilizumab in a peer-reviewed journal. The Living Mapping and Living Network Meta-Analysis of COVID-19 studies (https://covid-nma.com/) platform was searched on 18 May 2021 to identify updates to the living meta-analysis for tocilizumab versus standard of care. Relevant records were extracted and summarized in a narrative table of results (see Table 1).

The search strategies for all four reviews are shown in Appendix 2.

## **Eligibility criteria for review**

Population: Patients with confirmed COVID-19 (with or without elevated IL-6 levels), no restriction to age but severe disease requiring oxygen or ventilatory assistance.

Intervention: Tocilizumab in combination with local standard of care at the time. No restriction on dose, frequency, or timing with respect to onset of symptoms/severity of disease.

Comparators: Any (standard of care/placebo or active comparator).

Outcomes: Mortality; duration of ventilatory support including mechanical ventilation; duration of ICU stay; progression to ICU admission, progression to mechanical ventilation, clinical outcome on an ordinal scale at chosen time points, adverse events, adverse reactions.

Study designs: Randomised controlled trials, and systematic reviews of studies in humans.

## RESULTS

The previous version of this review included 9 RCTs found during a search of the Epistemonikos L\*OVE evidence platform (<u>https://app.iloveevidence.com/</u>). Details of the search are provided in Appendix 2. The Cochrane supported living metaanalysis (COVID-NMA)<sup>29</sup> was recently updated to include two additional RCTs<sup>33, 34</sup> as well as the published, peer-reviewed RECOVERY paper in the Lancet.<sup>32</sup> Findings of this living meta-analysis for the outcomes of interest are detailed below in Table 1. The associated forest plots are included in Appendix 3. The main characteristics and outcomes of the 12 included RCTs are summarised in Table 2.

## All-cause mortality at day 28

The COVID-NMA meta-analysis showed that tocilizumab 8mg/kg compared with standard of care/placebo for Mild/Moderate/Severe/Critical COVID-19 (eleven RCTs, 6 937 participants) reduced mortality: RR 0.88 (95% CI 0.81 to 0.95; I<sup>2</sup>=0.0%).

In Horby *et al.* (the Recovery trial)<sup>32</sup> there was some evidence for effect modification by concomitant use of steroids (Chi squared test for interaction p=0.01). In participants that received corticosteroids, day 28 mortality was 29% (482/1644) in the tocilizumab arm and 35% (600/1721) in the usual care arm. Amongst participants that did not receive corticosteroids, day 28 mortality was 39% (139/357) in the tocilizumab arm and 35% (127/367) in the usual care arm.

## WHO ordinal progression score level 7 or above at Day 28

Level 7 and above on the WHO ordinal scale for clinical improvement is defined as "hospitalized patients with severe disease requiring mechanical ventilation ± additional organ support (ECMO, vasopressors or dialysis) OR death"<sup>30</sup>. COVID-NMA meta-analysis showed that tocilizumab 8mg/kg compared with standard of care/placebo for mild/moderate/severe/critical COVID-19 (four RCTs, 1 066 participants) reduced progression to WHO level 7 or above: RR 0.88 (95% CI 0.59 to 1.32; I<sup>2</sup>=65.3%).

## **Adverse events**

Tocilizumab 8mg/kg (except for Wang *et. al.* which allowed for a standard 400 mg dose) compared with standard of care/placebo for mild/moderate/severe/critical COVID-19 (eight RCTs, 1 714 participants): the included studies showed no statistically significant difference in adverse events: RR 1.23 (95% CI 0.93 to 1.62; I<sup>2</sup>=81.3%).

## Serious adverse events

Tocilizumab 8mg/kg (except for Wang *et al.* which allowed for a standard 400 mg dose) compared with standard of care/placebo for mild/moderate/severe/critical COVID-19 (ten RCTs, 2 532 participants): the included studies showed no statistically significant difference in serious adverse events: RR 0.92 (95% CI 0.77 to 1.08; I<sup>2</sup>=0.0%).

## Table 1: Summary of findings of the Cochrane Living Meta-analysis: Tocilizumab compared to standard care/placebo for mild/moderate/severe/critical COVID-19

	Anticipated absolu	ute effects* (95% CI)	Deletive offerst		Containty of the suider of
Outcomes	Risk with Standard care/Placebo	Risk with Tocilizumab	(95% CI)	(studies)	(GRADE)
Clinical improvement D28 b	536 per 1000	<b>569 per 1000</b> (531 to 601)	<b>RR 1.06</b> (0.99 to 1.12)	5625 (8 RCTs) °	<b>⊕⊕⊕</b> ⊖ MODERATE ₫
WHO progression score (level 7 or above) D28	279 per 1000	<b>246 per 1000</b> (165 - 369)	<b>RR: 0.88</b> (0.59 - 1.32)	1066 (4 RCTs) ∘	$\bigoplus_{LOW} \bigcirc_{f,g}$
All-cause mortality D28	292 per 1000	<b>257 per 1000</b> (237 to 278)	<b>RR: 0.88</b> (0.81 - 0.95)	6937 (11 RCTs) ʰ	⊕⊕⊕⊕ HIGH <sup>†</sup>
All-cause mortality D60 or above	133 per 1000	<b>114 per 1000</b> (70 to 186)	<b>RR 0.86</b> (0.53 to 1.40)	519 (2 RCTs) <sup>j</sup>	
Adverse events	429 per 1000	<b>527 per 1000</b> (399 to 695)	<b>RR 1.23</b> (0.93 to 1.62)	1714 (8 RCTs) ™	⊕⊖⊖⊖ VERY LOW n,o,p
Serious adverse events	147 per 1000	<b>136 per 1000</b> (113 to 159)	<b>RR 0.89</b> (0.75 to 1.06)	2532 (10 RCTs) ۹	₩ MODERATE n

\*The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

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

**GRADE Working Group grades of evidence** 

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

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

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

## CONCLUSION

A systematic review of twelve RCTs of the use of tocilizumab in hospitalized patients with COVID-19 demonstrated a reduction in all-cause mortality at day 28. The absolute risk reduction was 3.5% (95% CI 1.5% to 5.6%), while the relative risk was 0.88 (95% CI 0.81 to 0.95, 11 RCTs reporting mortality, n = 6 937). The number needed to treat to prevent one additional death due to COVID-19 was calculated as 29 (95% CI 18 to 67). This result is largely driven by the findings of the RECOVERY trial<sup>32</sup> wherein tocilizumab, when used for hypoxic patients with a CRP of >75mg/L, produced a 4% absolute reduction in 28-day mortality compared with standard of care alone (TCZ = 31% vs. SOC = 35%; RR 0.85; 95% CI, 0.76 to 0.94; p=0.0028). However, there are concerns regarding the national supply of tocilizumab and that the product is unaffordable. On this basis, it is recommended that tocilizumab, used in combination with corticosteroids, not be included in the COVID-19 treatment guidelines for the management of hospitalized hypoxic SARS-CoV-2 infected patients with CRP levels >75mg/L.

**Reviewers:** Roger Wiseman (Liberty Health (Pty) Ltd, South Africa), Marc Blockman (Division of Clinical Pharmacology, Department of Medicine, Groote Schuur Hospital, University of Cape Town).

Additional reviewers: Karen Cohen (Division of Clinical Pharmacology, Department of Medicine, Groote Schuur Hospital, University of Cape Town). Renee De Waal (Centre for Infectious Disease Epidemiology and Research, University of Cape Town).

Declaration of interests: RW, MB, KC and RdW have no interests to declare in respect of tocilizumab.

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## Table 2. Characteristics of 12 randomised controlled trials included in the Cochrane living systematic review

Citation	Study design	Population (n)	Treatment	Main findings	Comments
Published, peer reviewed	Prospective open-label.	TCZ = 350 SRL = 45 Control = 397 Median age: 61.4 years (TCZ = 61.5 years; SRL = 63.4 years; placebo = 61.1 years Setting: Italy, 24 hospitals	The TCZ group received TCZ	Primary endpoint: clinical	Risk of bias assessment:
Salvarani C, Dolci G, Massari M, et al. <sup>21</sup> JAMA internal medicine. 2020. Effect of Tocilizumab vs Standard Care on Clinical Worsening in Patients Hospitalized With COVID-19 Pneumonia: A Randomized Clinical Trial. NCT04346355	<ul> <li>Thospective, open label, randomized clinical trial</li> <li>31 March 2020 to 11 June 2020.</li> <li>The primary aim was to evaluate the efficacy of early administration of tocilizumab vs standard therapy in the first 2 weeks following randomization.</li> </ul>	Participants: patients with COVID-19 confirmed by positive PCR and the presence of acute respiratory failure with a partial pressure of arterial oxygen to fraction of inspired oxygen (PaO2/FIO2) ratio between 200 and 300 mm/Hg, an inflammatory phenotype defined by a temperature greater than 38°C during the last 2 days, and/or serum CRP ≥ 10mg/dL and/or CRP level increased to at least twice the admission measurement. Sample size: 126 (TCZ = 60, Standard Care = 66) Median age = 60.0 years (range 53.0 to 72.0 years).	intravenously within 8 hours from randomization at a dose of 8mg/kg up to a maximum of 800 mg, followed by a second dose after 12 hours. The control arm received supportive care following the treatment protocols of each centre. All drugs were allowed except IL-1 blockers, Jak inhibitors, and tumor necrosis factor inhibitors.	<ul> <li>worsening within 14 days since randomization, defined by the occurrence of 1 of the following events, whichever occurred first:</li> <li>Admission to ICU with mechanical ventilation</li> <li>Death from any cause</li> <li>PaO2/FIO2 ratio less than 150 mm Hg in 1 of the scheduled arterial blood gas measurements or in an emergency measurement, confirmed within 4 hours by a second examination</li> <li>17 of 60 participants (28.3%) in the TCZ group and 17 of 63 (27.0%) in the standard care group showed clinical worsening within 14 days following randomization (rate ratio, 1.05; 95% CI, 0.59-1.86; P = .87)</li> </ul>	MODERATE with some concerns noted. ITT analysis, but an unblinded study with a cross-over design (21.2% of patients in the SOC arm received study treatment due to clinical worsening).
Published, non-peer reviewed Salama C, Han J, Yau L, et. al. <sup>22</sup> medRxiv 2020.	Randomised, double-blind, placebo controlled, Phase III study	Setting: Multi-centre study across 6 countries Participants: hospitalized patients with COVID-19 pneumonia confirmed by positive PCR test and radiographic imaging.	Participants were randomized (2:1) to intravenous tocilizumab (8 mg/kg, maximum 800 mg) or placebo. If participants worsened or did not improve, an additional infusion could be administered 8 to 24 hours after the first.	Primary endpoint: cumulative proportion of participants requiring mechanical ventilation (mechanical invasive ventilation or extracorporeal membrane oxygen) or who had died by Day 28.	Risk of bias assessment: LOW as study was double-blind, placebo- controlled with random allocation sequence and adequate concealment. Data was available for >95% of population and outcomes reported were pre- specified in the protocol.

Tocilizumab in non- ventilated patients hospitalized with Covid-19 pneumonia. DOI: 10.1101/2020.10.21.20210 203 NCT04372186		Sample size: 377 (TCZ = 249, Placebo = 128) Median age (± SD) = TCZ = 56.0 ±14.3 years; placebo = 55.6 ±14.9 years.	Both groups received standard care per local practice which could include antiviral treatment, limited systemic corticosteroids (≤1 mg/kg methylprednisolone or equivalent recommended) and supportive care In the tocilizumab and placebo arms, 55.4% and 67.2% of participants received dexamethasone, respectively, and 52.6% and 58.6% received remdesivir, respectively.	TCZ = 12.0% (95% CI, 8.52% to 16.86%) Placebo = 19.3 % (95% CI, 13.34% to 27.36%) (HR, 0.56 [95% CI, 0.33 to 0.97]; log-rank P=0.036).	
Published, peer reviewed Hermine O, Mariette X, Tharaux PL, et al. for the CORIMUNO-19 Collaborative Group. <sup>19</sup> JAMA internal medicine. 2020. Effect of Tocilizumab vs Usual Care in Adults Hospitalized With COVID-19 and Moderate or Severe Pneumonia: A Randomized Clinical Trial NCT04331808	Randomised, open-labelled, multicenter study. 31 March 2020 to 18 April 2020.	Setting: France, 9 university hospitals Participants: patients with COVID-19 and moderate or severe pneumonia requiring at least 3 L/min of oxygen but without ventilation or ICU admission. Sample size: 130 (TCZ = 63 Usual Care = 67) Median age = TCZ = 64.0 years; UC = 63.3 years.	Participants were randomized on a 1:1 ratio to receive TCZ plus usual care or usual care alone. TCZ was administered at a dose of 8mg/kg IV on Day 1, followed by a fixed dose of 400mg IV on day 3 if the oxygen requirement has not decreased by more than 50%.	Primary outcomes were scores higher than 5 on the World Health Organization 10-point Clinical Progression Scale (WHO-CPS) on day 4 and survival without need of ventilation (including non-invasive ventilation) at day 14. Outcomes amended on 06 April 2020 to include high-flow oxygen in noninvasive ventilation. Primary: 12 participants (19%) had a WHO-CPS score greater than 5 at day 4 vs 19 (28%) in the UC group (median posterior absolute risk difference [ARD] –9.0%; 90% credible interval [Crl], –21.0 to 3.1) At day 14, 12% (95% Cl –28%to 4%) fewer participants needed non- invasive ventilation (NIV) or mechanical ventilation (or died in the TCZ group than in the UC group (24%vs 36%, median posterior hazard ratio [HR] 0.58; 90% Crl, 0.33-1.00).	Risk of bias assessment: MODERATE as this was an unblinded study.

				The number of participants with mechanical ventilation or death at Day 14 was 11 (17%) and 18 (27%) in the TCZ and UC groups respectively.	
Published, peer reviewed Stone JH, Frigault MJ, Serling-Boyd NJ, et al. <sup>20</sup> New England Journal of Medicine. 2020. Efficacy of Tocilizumab in Patients Hospitalized with Covid-19 <u>https://www.nejm.org/doi// full/10.1056/NEJMoa20288</u> <u>36</u> NCT04356937	Randomised, double-blind, placebo controlled. 20 April 2020 to 15 June 2020.	Setting: USA, 7 hospitals in Boston Participants: patients with COVID-19 confirmed either by PCR or serum IgM antibody assay. Participants had to have at least two of the following signs: fever (body temperature >38°C) within 72 hours before enrollment, pulmonary infiltrates, or a need for supplemental oxygen to maintain an oxygen saturation > 92%. At least one of the following laboratory criteria also had to be fulfilled: a CRP > 50 mg/L, ferritin > 500 ng/ml, D-dimer > 1000 ng/ml, LDH > 250 U/L. Sample size: 242 (TCZ = 161, Placebo = 81) Median age = 59.8 years (range 21.7 to 85.4 years).	Participants were randomised on a 2:1 ratio to receive standard care plus a single dose of either tocilizumab (8 mg per kilogram of body weight administered intravenously, not to exceed 800 mg) or placebo Antiviral therapy, hydroxychloroquine, and glucocorticoids were permitted as concomitant treatment. However, some participants received remdesivir as concomitant treatment due to the release of the ACTT-1 trial during this trial. no participants received dexamethasone as the RECOVERY trial results were announced afterwards.	The primary outcome was intubation (or death, for participants who died before intubation) after administration of tocilizumab or placebo. The secondary endpoints were clinical worsening and discontinuation of supplemental oxygen among participants who had been receiving it at baseline. The hazard ratio for intubation or death for TCZ as compared with the placebo group was 0.83 (95% confidence interval [CI], 0.38 to 1.81; P = 0.64). The hazard ratio for disease worsening was 1.11 (95% CI, 0.59 to 2.10; P = 0.73). At 14 days, 18.0% of the participants in the TCZ group and 14.9% of the participants in the placebo group had demonstrated disease worsening. There was no difference in the median time to discontinuation of supplemental oxygen [TCZ = 5.0 days (95% CI, 3.8 to 7.6) vs placebo = 4.9 days (95% CI, 3.8 to 7.8)] in the placebo group (P = 0.69). At 14 days, 24.6% of the participants in the placebo group (P = 0.69). At 14 days, 24.6% of the participants in the placebo group were still receiving supplemental oxygen.	Risk of bias assessment: LOW as the study is a randomized, double- blind, placebo-controlled trial with random allocation sequence and adequate concealment. 242/243 patients analyzed (<5% of total sample size).

Non-peer reviewed Wang D, Fu B, Peng Z, et. al. <sup>24</sup> SSRN. 2020. Tocilizumab Ameliorates the Hypoxia in COVID-19 Moderate Patients with Bilateral Pulmonary Lesions: A Randomized, Controlled, Open-Label, Multicenter Trial. DOI: <u>10.2139/ssrn.3667681</u> ChiCTR2000029765	Randomized, controlled, open-label, multicentre trial 13 February 2020 to 13 March 2020.	Setting: China, 6 hospitals in Anhui and Hubei Participants: Patients PCR confirmed COVID-19 between the ages of 18 and 85 years, had elevated plasma IL-6 levels with moderate or severe disease. Moderate disease was defined as fever or other respiratory symptoms as well as bilateral pulmonary lesions confirmed on chest imaging Severe disease was defined as the presence of any of the following: 1) respiratory rate ≥30 breaths per min; 2) SpO2 ≤ 93% while breathing room air; and/or 3) PaO2/FiO2 ≤ 300 mmHg Sample size: 65 (TCZ = 33, Control = 1460) Median age = 63.0 years (IQR = 55.0 to 71.0 years).	Participants were randomly assigned in a 1:1 ratio to receive either tocilizumab in addition to standard care, or standard care alone. If a participant in the control group progressed to severe disease within 3 days after randomization, they were transferred to the tocilizumab group.	<ul> <li>Primary endpoint: Cure rate. Cure was defined as 1) fever attenuated for continuously for 7 days, 2) two negative COVD-19 PCR tests, 3) CT scan showing absorption of chest effusion by more than 50% percent on discharge.</li> <li>The cure rate for TCZ was 94.12% vs 87.10% for the control group, but the difference was not statistically significant (P = 0.4133).</li> <li>For the secondary endpoints of recovery rate of hypoxia over 14 days and the worsening rate of hypoxia it. TCZ = 91.67% vs 60.00% (p = 0.0328) in the control group. The difference was evident from day 4 and statistically significant from day 12.</li> </ul>	Risk of bias assessment: HIGH, as there were concerns regarding the allocation concealment during randomisation. ITT analysis, but small study (n=65). Study unblinded and possible bias with the measurement of the outcome in particular the measurement of adverse events and serious adverse events. Possible bias regarding selection of the reported results, as adverse events were not mentioned in the registry but reported in the paper. The protocol and statistical plans were not available.
Published non-peer reviewed Rosas I, Bräu N, Waters M, et. al. <sup>23</sup> medRxiv 2020 Tocilizumab in Hospitalized Patients With COVID-19 Pneumonia.	Randomized, double-blind, placebo-controlled trial	Setting: 9 Countries - Canada, Denmark, France, Germany, Italy, Netherlands, Spain, UK and USA. Participants: Patients with PCR confirmed COVID-19 and evidenced by bilateral chest infiltrates on chest x-ray or CT. Participants were also required to have blood oxygen saturation ≤93% or partial pressure of	Participants were randomized (2:1) to receive intravenous tocilizumab (8 mg/kg infusion, maximum 800 mg) or placebo plus standard care. If clinical signs or symptoms did not improve or worsened a second infusion could be administered 8 to 24 hours after the first.	Primary endpoint: Clinical status assessed on a 7-category ordinal scale at day 28. Clinical status was measured at baseline and every day during hospitalization. Clinical status at day 28 was not statistically significantly improved for tocilizumab versus placebo (P=0.36). Median (95% CI) ordinal scale values at day 28: TCZ = 1.0 (1.0 to 1.0) for tocilizumab	Risk of bias assessment: LOW as blinded study with random allocation sequence and adequate concealment. 452/438 patients analysed (>95% of population), and outcomes reported were pre-specified in the registry.

DOI		oxygen/fraction of inspired		Placebo 2.0 (1.0 to 4.0) (odds ratio,	
10.1101/2020.08.27.20183		oxygen <300 mmHg.		1.19 [0.81 to 1.76]).	
442					
		Sample size: 438 (TCZ = 294,		There was no difference in	
NCT04320615		Control = 144)		mortality at day 28 between	
				tocilizumab (19.7%) and placebo	
		Median age = 63.0 years (IOR		(19.4%) (difference, 0.3% [95% Cl, –	
		= 55.0  to  71.0  years		7.6 to 8.2]; nominal P=0.94).	
				Median time to hospital discharge	
				was 8 days shorter with tocilizumab	
				than placebo (20.0 and 28.0,	
				respectively; nominal P=0.037;	
				hazard ratio 1.35 [95% CI 1.02 to	
				1.79]).	
				Median duration of ICU stay was	
				5.8 days shorter with tocilizumab	
				than placebo (9.8 and 15.5,	
				respectively; nominal P=0.045).	
Published, peer-reviewed	Randomised, open-labelled	Setting: , 9 hospitals	Patients were randomised (1:1) to	Primary outcome: clinical status at	Risk of bias assessment:
	trial		receive either TCZ (single intravenous	15 days evaluated on a seven-level	MODERATE as study was
Veiga VC, Prats JAGG, Farias		Participants: Adult patients	infusion of 8 mg/kg) plus standard care	ordinal scale.	unblinded study and the outcomes
DLC et. al. <sup>31</sup>	08 May to 17 July 2020.	with severe PCR-confirmed	or standard care alone.		could have been influenced by the
		SARS-CoV-2 infection.		The trial was prematurely after the	intervention assignment.
BMI 2021		Patients were required to	Standard of care allowed for the	first interim analysis owing to an	
		have experience symptoms	concomitant use of	excess number of deaths at 15 days	
Effect of topilizumah on		for more than 3 days and	hydroxychloroquine, azithromycin,	in the tocilizumab group	
clinical outcomes at 1E days		present with evidence of	corticosteroids, and antibiotics as per		
in patients with severe or		pulmonary infiltrates	local institutional guidelines for	TCZ was not associated with an	
critical coronavirus disease		confirmed by chest CT or	patients with covid-19. Remdesivir	improvement in mechanical	
2019: randomised		radiography and were	was not available in Brazil at the time	ventilation or death at 15 days (18	
controlled trial.		owegen or had been receiving	of the study.	of 65 (28%) patients in the TCZ	
		mechanical ventilation for		group and 13 of 64 (20%) in the	
DOI: 10.1136/bmi.n84		less than 24 hours before		SOC group: odds ratio 1.54, 95% Cl	
DOI: 10.1130/binj.n84		analysis		0.66 - 3.66; P=0.32). Death at 15	
NCTOMODCOF				autoamo accurred in 11 (17%)	
NC104403685		Sample size: 129 (TC7 - 65		nations in the TC7 group	
		SOC = 64		compared with two (3%) in the SOC	
				group (odds ratio 6.42, 1.59 - 43.2).	
		iviean age:			

		TCZ = 57.4 years (SD = 15.7 years) SOC = 57.5 years (SD = 13.5years)			
Published, peer reviewed Horby PW, Pessoa-Amorim G, Peto L et al. for the RECOVERY Collaborative Group <sup>32</sup> Lancet 2021 Tocilizumab in patients admitted to hospital with COVID-19 (RECOVERY): preliminary results of a randomised, controlled, open-label, platform trial DOI: 10.1016/S0140- 6736(21)00676-0 NCT04381936, ISRCTN50189673	Randomised, controlled, open-label, platform trial (Randomised Evaluation of COVID-19 Therapy [RECOVERY] 23 April 2020 and 24 January 2021	Setting: United Kingdom, 131 sites Participants: Adult hospitalized patients with clinically suspected or laboratory confirmed SARS- CoV-2 infection as well as hypoxia (defined as Sp02 of <92% and a CRP level of ≥75 mg/L). Sample size: 4 116 (TCZ = 2 022, SOC = 2 094) At the time of this publication, follow-up was completed for 1602 (79%) of the 2022 TCZ patients and 1664 (79%) of the 2094 patients SOC Mean age: 63.6 years (SD 13.7)	Patients were randomised (1:1) to receive either TCZ plus standard care or standard care alone. TCZ doses were weight-based (800mg if weight >90kg; 600 mg if weight >40 and ≤65 kg; and 8mg/kg if weight ≤40 kg). A second dose could be given 12 to 24 hours later if, at the discretion of the attending clinician, the patient's condition had not improved. Standard of care is not described. At randomisation, 562 (14%) patients were receiving invasive mechanical ventilation, 1686 (41%) were receiving non-invasive respiratory support (including high-flow nasal oxygen, CPAP, and non-invasive ventilation), and 1868 (45%) were receiving no respiratory support other than simple oxygen therapy.	Primary outcome: all-cause mortality at 28 days. Secondary outcomes were time to discharge alive from hospital, and, among patients not receiving invasive mechanical ventilation at randomisation, receipt of invasive mechanical ventilation (including extra-corporeal membrane oxygenation) or death. TCZ was associated with a 4% absolute reduction 28-day mortality compared with SOC alone (621 [31%] of 2022 patients in the tocilizumab group vs. 729 (35%) of 2094 patients in the usual care group; rate ratio 0.85; 95% confidence interval [CI], 0.76 to 0.94; p=0.0028). TCZ was associated with a greater probability of discharge from hospital alive within 28 days (57% vs. 50%; rate ratio 1.22, 95% CI 1.12 to 1.33, p<0.0001). Among those not on invasive mechanical ventilation at baseline, TCZ reduced in the risk of progressing to the pre-specified composite secondary outcome of invasive mechanical ventilation or death when compared to SOC (35% vs. 42%, risk ratio 0.84, 95% CI 0.77 to 0.92,	Risk of bias assessment: MODERATE as study was unblinded and there were some concerns with the measurement of the outcome: Clinical improvement (D28), defined as discharge from hospital. Assessment requires clinical judgement and could be affected by knowledge of intervention receipt.
Published, peer reviewed	Randomised, controlled, open-label phase 3 study	Setting: India, 12 sites	Patients were randomised (1:1) to receive either TCZ plus standard care	Primary outcome: Proportion of patients with progression of	Risk of bias assessment: MODERATE as study was

Soin AS, Kumar K,		Participants: Hospitalised	or standard care alone. TCZ was	COVID-19 from moderate to severe	unblinded and there were some
Choudhary NS, et. al. <sup>33</sup>	30 May 2020 and 31 August	patients aged 18 years or	administered as a single IV infusion at	or from severe to death up to day	concerns with the measurement of
	2020	older with confirmed SARS-	6mg/kg up to a maximum dose of	14.	the outcomes of mortality (D28),
Lancet Respir Med 2021		CoV-2 infection and	480mg. An additional dose of 6mg/kg		adverse events and serious
		moderate to severe disease	(max 480mg) could be administered if	The proportion of patients with	adverse events.
Tocilizumab plus standard		(moderate defined as	clinical symptoms worsened or did not	progressive COVID-19 up to day 14	
care versus standard care in		respiratory rate 15–30 per	show improvement within 12 hours to	was 9% (eight of 91) in the TCZ	
patients in India with		min [revised to 24 per min on	7 days after the first dose.	group and 13% (11 of 88) in the	
moderate to severe COVID-		June 13, 2020] and blood		SOC group. The difference was not	
19- associated cytokine		oxygen saturation [SpO2] 90–	The dosing regimen was selected on	statistically significant (–3.71 [95%	
release syndrome		94%; Severe defined as	the basis of the cost and supply	CI –18.23 to 11.19]; p=0.42).	
(COVINTOC): an open-label,		respiratory rate ≥30 per min	considerations in India and because a		
multicentre, randomised,		or SpO2 <90% in ambient air,	single dose between 4 mg/kg and 8		
controlled, phase 3 trial		or ARDS or septic shock)	mg/kg plus an additional dose to a		
			maximum of 800 mg, if required, had		
https://doi.org/10.1016/		Sample size: 180 (TCZ = 91,	been recommended on the basis of		
S2212 2600(21)00081 2		SOC = 88)	initial reports on the use of tocilizumab		
52215-2000(21)00081-5			in the treatment of COVID-19 in China.		
CTRI/2020/05/025369		Median age: TCZ = 56 years			
CTRI/2020/05/025505		range 47 - 63); SOC = 54 years	Standard of care included		
		(range 43-63)	corticosteroids equivalent to		
			methylprednisolone 1 mg/kg or less if		
			deemed necessary by the treating		
			physician. Supplemental oxygen was		
			recommended to treat hypoxia, and		
			high-flow nasal cannula, non-invasive		
			ventilation, and mechanical ventilation		
			could be considered if hypoxia and		
			respiratory distress progressed.		
Published non-neer	Bandomised controlled	Setting: Tehran Iran	Patients were randomised (1:1) to	Primary outcome: mortality at 28	Small study (n=40) using an
reviewed	double-blind two-arm	Single site	receive either TC7 plus standard care	days	intention to treat analysis.
leviewed	narallel nhase 2 study	Single site	or standard care alone TC7 dose of	duys.	,
	puruler, pruse 2 study		8  mg/kg to a maximum of $800  mg$ was		Overall risk of bias assessment:
Talaschian M, Akhtari M,		Participants: Non-ventilated	administered to the TC7 group within 2	Death was reported in 29.4% of the	HIGH
Mahmoudi M, et. al. 34	10 July 2020 and 10	patients, hospitalised adult	days of hospitalization A second dose	TCZ arm and 21.1% of the SOC arm.	Randomisation: Allocation
	October 2020	patients (18 years and older)	could be given 12 hours later if the	This difference was not statistically	sequence probably random
Research Square 2021		with confirmed SARS-CoV-2	natient's condition was not stable	significant. [Log rank test: P=0.973;	(block randomisation), but
		intection (via positive PCR		Hazard ratio: 1.25; 95% CI: 0.249-	unclear allocation concealment
Tocilizumah failed to		result and atypical CT		4.209].	- MODERATE RISK.
reduce mortality in source		teatures) not responding to			<ul> <li>Deviations from intervention: "In</li> </ul>
COVID-19 patients: Posults		standard COVID-19 treatment			this study, patients, investigators
from a randomized		and have:			and outcome assessors did not
controlled clinical trial					inform which group received an
controlled cliffical trial.	1	1		1	

		1. CRP levels of $\geq 10 \text{ mg/L}$ .			intervention. Besides, a placebo
https://doi.org/10.21202/rs		or IL-6 of 18pg/ml or			was not used in the control
$\frac{1111195.7/1001.01g/10.21203/15}{2 rc}$		lymphopenia			group" Unclear blinding and
.5.15-405921/V1		(<1100/MCL) and			antivirals not distributed
		2  (1100)  (1101)  (1101)			between intervention (TC7) and
IRCT20081027001411N4		2. SpU2 <93% or respiratory			standard of care group (35% vs
		rate higher than 24.			10%) – MODERATE RISK
					Alissing outcome datas 40
		Sample size: 40 patients (TCZ			<ul> <li>Missing outcome data: 40</li> </ul>
		= 20, SOC = 20)			participants randomized; 36
					participants analyzed. 3
		Moon $200 = 61.74 \pm 14.19$			participants in the treatment arm
		weeks ald			and 1 in the standard care arm
		years old			refused to participate before
					start of the intervention –
					MODERATE RISK.
					Measurement of the outcome:
					Outcome assessors were blinded
					– LOW RISK.
					• Selection of the reported results:
					Trial probably not analyzed as
					prespecified. Mortality outcome
					has different time point listed in
					the registry compared to the
					report. Neither are the outcomes
					of clinical improvement and
					serious adverse events not
					prespecified MODERATE RISK.
Published, non-peer	Prospective, randomised,	Setting: Netherlands, 11 sites	Patients were randomised (1:1) to	Primary outcome: mortality at 30	Overall risk of bias assessment:
reviewed	controlled, phase 2 study	_	receive either TCZ plus standard care	days.	MODERATE. Concerns were
		Participante, Upenitalized	or standard care alone. TCZ dose of		noted with the randomization of
	06 April 2020 and 12	Participants: Hospitalised	8mg/kg to a maximum of 800mg was	Death was reparted in 20.4% of the	the patient population.
Rutgers IVI, Westerweel PE,	06 April 2020 and 12	adult patients (18 years and	administered to the TCZ group within 2	TCZ area and 21.1% of the SOC area	
van der Holt B, et.al.	January 2021	Col( 2 infection (via positive	days of hospitalization. A second dose	The difference was not statistically	
		Cov-2 infection (via positive	could be given after 8 hours if hypoxia	inis difference was not statistically	
SSRN 2021		PCR result and have signs	was not resolved.	significant. [Log rank test: P=0.973;	
		compatible with hyper		Hazard ratio: 1.25; 95% CI: 0.249-	
Timely administration of		finiammation, namely a need		4.209].	
tocilizumah improves		for supplemental oxygen	88% of patients received		
survival of hospitalized		(SpO2 < 94% and/or ferritin)	dexametnasone as concomitant	For secondary outcomes measures,	
COVID-19 natients		>2000ug/i or a doubling of	treatment. Remoesivir and	there was no difference in duration	
covid 19 patients.		serum ferritin in 20-48 hours	nydroxychloroquine were both allowed	of hospital stay, percentage of	
			as concomitant therapies.	patients admitted to ICU, number	
http://dx.doi.org/10.2139/s		Sample size: 354 patients		of patients ventilated or duration	
srn.3834311		(TCZ = 174, SOC = 180)		of ventilation. However, the	

Trial NL8504Mean age = 66 years (range 56 - 75 years); SOC = 67 years (60 - 74 years)	median duration of ICU stay was significantly shorter in patients receiving TCZ (9 days, IQR 5-16 days vs. 16 days, IQR 8-30 p=0.025).
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## Appendix 1: GUIDELINE CONSIDERATIONS

## 1. NIH COVID-19 Treatment Guidelines (updated 21 April 2021)<sup>17</sup>

Interleukin-6 Inhibitors Recommendations:

- The Panel recommends using tocilizumab (single intravenous [IV] dose of tocilizumab 8 mg/kg actual body weight up to 800 mg) in combination with dexamethasone (6 mg daily for up to 10 days) in certain hospitalized patients who are exhibiting rapid respiratory decompensation due to COVID-19. These patients are:
  - Recently hospitalized patients (i.e., within first 3 days of admission) who have been admitted to the intensive care unit (ICU) within the prior 24 hours and who require invasive mechanical ventilation, noninvasive ventilation, or high-flow nasal canula (HFNC) oxygen (>0.4 FiO2/30 L/min of oxygen flow) (BIIa); or
  - Recently hospitalized patients (i.e., within first 3 days of admission) not admitted to the ICU who have rapidly increasing oxygen needs and require noninvasive ventilation or HFNC oxygen and who have significantly increased markers of inflammation (CRP ≥75 mg/L) (BIIa).
- For hospitalized patients with hypoxemia who require conventional oxygen therapy, there is insufficient evidence to specify which of these patients would benefit from the addition of tocilizumab. Some Panel members would also give tocilizumab to patients who are exhibiting rapidly increasing oxygen needs while on dexamethasone and have a CRP ≥75 mg/L, but who do not yet require noninvasive ventilation or HFNC oxygen as described above.
- There are insufficient data for the Panel to recommend either for or against the use of sarilumab for hospitalized patients with COVID-19 who are within 24 hours of admission to the ICU and who require invasive mechanical ventilation, noninvasive ventilation, or high-flow oxygen (>0.4 FiO2/30 L/min of oxygen flow).
- The Panel **recommends against** the use of anti-IL-6 monoclonal antibody therapy (i.e., siltuximab) for the treatment of COVID-19, except in a clinical trial (BI).

## 2. World Health Organization: COVID-19 Clinical management: living guidance, 25 January 2021<sup>18</sup>:

It is noted that no guidance has been provided for the use of IL-6 inhibitors in the WHO Therapeutics and COVID-19 living guideline of 31 March 2021.

## 3. Australian guidelines for the clinical care of people with COVID-19. Version 34.1 (updated 13 May 2021)<sup>27</sup> <u>6.3.1 Tocilizumab</u>

**Recommendation:** Consider using tocilizumab for the treatment of COVID-19 in adults who require supplemental oxygen, particularly where there is evidence of systemic inflammation.

In patients hospitalised with COVID-19 who require supplemental oxygen, tocilizumab probably reduces the risk of death. Because of this, the Taskforce gives a conditional recommendation for tocilizumab both within and outside the context of a randomised trial unless contraindicated (e.g. patients with other active, severe infections).

In accordance with the RECOVERY trial, tocilizumab should be administered as a single intravenous infusion over 60 minutes, with the potential for a second dose to be administered either 12 or 24 hours later if the patient's condition has not improved. The suggested dose is dependent on body weight:

- Patients > 90 kg: 800 mg tocilizumab
- $\circ$  Patients > 65 and  $\leq$  90 kg: 600 mg tocilizumab
- Patients > 40 and  $\leq$  65 kg: 400 mg tocilizumab
- Patients ≤ 40 kg: 8 mg/kg tocilizumab

In addition, the RECOVERY and REMAP-CAP trials have demonstrated a significant benefit when using corticosteroids in conjunction with tocilizumab. Use of combined tocilizumab and corticosteroids should be considered in patients hospitalised with COVID-19 who require oxygen, however the optimal sequencing of tocilizumab and corticosteroid use is unclear.

As tocilizumab inhibits the production of C-reactive protein (CRP), a reduction in CRP should not be used as a marker of clinical improvement.

## Search Strategy 1:

Date: 11 April 2020 Period: Prior to 11 April 2020

## Epistemonikos

(title:(coronavirus or covid\* or 2019-ncov or sars-cov-2) or abstract:(coronavirus or covid\* or 2019-ncov or sars-cov-2)) and (title:(tocilizumab or IL-6 inhibitor or interleukin-6 inhibitor) or abstract:(tocilizumab or IL-6 inhibitor or interleukin-6 inhibitor))

Records retrieved: 13 (1 relevant to PICO question)

## PubMed

(((coronavirus[title/abstract] or covid\*[title/abstract] or 2019-ncov[title/abstract] or sars-cov-2[title/abstract])) and (tocilizumab[title/abstract] or IL-6 inhibitor[title/abstract] or interleukin-6 inhibitor[title/abstract]) not ((animals[mh] not humans[mh]))) and ("2019/12/01"[date - publication] : "3000"[date - publication])

Records retrieved: 43 (1 relevant to PICO question)

Living mapping and living network meta-analysis of COVID-19 studies (https://covid-nma.com/)

Tocilizumab Interleukin-6 inhibitor Interleukine-6 inhibitor

**Records retrieved: none** 

Cochrane COVID Study Register (https://covid-19.cochrane.org/)

Tocilizumab AND interleukin-6 inhibitor

Records retrieved: 12 (none relevant to PICO question)

## Search Strategy 2:

Date: 15 November 2020

## **Epistemonikos L\*OVE evidence platform:**

Tocilizumab

**Records retrieved:** 58 (10 RCTs and 48 systematic reviews). 6 RCTs and 1 systematic review were included for review.

## Search Strategy 3:

Date: 13 January 2021

## **Epistemonikos L\*OVE evidence platform:**

Tocilizumab

**Records retrieved:** Randomised controlled trials and systematic reviewed published in December 2020 and January 2021. Three RCTs were eligible, however only one pertained to data not previously reviewed and thus was included in this update. No systematic reviews were found published after 15 November 2020 that included the randomized controlled trial described above.

#### Search Strategy 4:

Date: 18 May 2021

Living mapping and living network meta-analysis of COVID-19 studies (https://covid-nma.com/)

Tocilizumab

**Records retrieved: 12 RCTs** 

# Appendix 3: Forest plots for Cochrane Living Meta-analysis: Tocilizumab compared to Standard of care/Placebo for Mild/Moderate/Severe/Critical COVID-19

Tocilizumab 8 mg/kg Tocilizumab 8 mg/kg	Placebo Placebo	26/259	11/129	4	A	B	C	D	E	Overall	R	isk Ratio (95% CI)
Tocilizumab 8 mg/kg Tocilizumab 8mg/kg	Placebo	26/259	11/129						-		2012/12/2	10001000000000 - 10000000
Tocilizumab 8 mg/kg Tocilizumab 8mg/kg	Placebo Placebo	26/259	11/129			-		-	_			
Tocilizumab 8mg/kg	Placebo						-	-	-	•	1.31%	1.18 [0.60, 2.31]
		9/161	3/82	1	-	•		•	•	-	0.36%	1.53 [0.43, 5.49]
Tocilizumab 8 mg/kg once-off	Standard care	5/20	4/20	· · · ·	-	•			•	-	0.44%	1.25 [0.39, 3.99]
Tocilizumab 8mg/kg	Standard care	7/64	8/67	<u>⊢_</u>		•		•	•	-	0.65%	0.92 [0.35, 2.38]
Tocilizumab 8mg/kg	Placebo	58/301	28/151			•			•	•	3.57%	1.04 [0.69, 1.56]
Tocilizumab 8 mg/kg once-off	Standard care	21/174	34/180	<b></b> 1		•			-	•	2.34%	0.64 [0.39, 1.06]
Tocilizumab 6 mg/kg/day	Standard care	13/90	15/90		•	-		•	-		1.27%	0.87 [0.44, 1.72]
Tocilizumab maximum 800 mg	Standard care	621/2022	729/2094	•		-		-	-	-	76.50%	0.88 [0.81, 0.96]
Tocilizumab 8 mg/kg	Standard care	14/65	6/64						•		0.74%	2.30 [0.94, 5.61]
Tocilizumab 8mg/kg	Standard care	2/60	1/66			•			-		0.10%	2.20 [0.20, 23.65]
Tocilizumab 8 mg/kg	Standard care Total:	98/366 874/3582	142/412 981/3355					-	•		12.72%	0.78 [0.63, 0.96]
	Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg once-off Tocilizumab 6 mg/kg/day Tocilizumab 8 mg/kg Tocilizumab 8 mg/kg Tocilizumab 8 mg/kg Tocilizumab 8 mg/kg	Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg cree-off Tocilizumab Tocilizumab Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab Brigikg Tocilizumab	Tocilizumab Standard care 7/64 Bmg/kg Tocilizumab Placebo 59/301 Bmg/kg once-off Tocilizumab Standard care 21/174 B mg/kg once-off Tocilizumab Standard care 13/90 B mg/kg Tocilizumab Standard care 621/2022 maximum 800 mg Tocilizumab Standard care 14/65 B mg/kg Standard care 2/60 Bmg/kg Tocilizumab Standard care 99/966 B mg/kg Total: 974/3592	Tocilizumab     Standard care     7/64     8/67       Bmg/kg     Placebo     58/301     28/151       Tocilizumab     Placebo     58/301     28/151       Tocilizumab     Standard care     21/174     34/180       Sing/kg once-off     Tocilizumab     Standard care     13/90       Sing/kg order-off     Standard care     13/90     5/90       Tocilizumab     Standard care     621/2022     729/2094       Tocilizumab     Standard care     14/66     6/64       Sing/kg     Standard care     14/66     6/64       Tocilizumab     Standard care     2/60     1/66       Bing/kg     Standard care     9/0366     142/412       Sing/kg     Standard care     9/0355     991/3355	Tocilizumab       Standard care       7/64       8/67         Brig/kg       Placebo       59/301       28/151         Tocilizumab       Standard care       21/174       34/180         Tocilizumab       Standard care       13/90       15/90         Tocilizumab       Standard care       621/2022       729/2094         Tocilizumab       Standard care       621/2022       729/2094         Tocilizumab       Standard care       621/2022       729/2094         Tocilizumab       Standard care       14/65       6/64         8 mg/kg       Tocilizumab       Standard care       2/60       1/66         Brog/kg       Tocilizumab       Standard care       2/60       1/66         Brog/kg       Tocilizumab       Standard care       98/306       142/412         Tocilizumab       Standard care       98/305       98/1/3355       142/412         = 0.00       Total:       874/3582       981/3355       142/412	Tocilizumab Standard care 7/64 8/67 Brig/kg Tocilizumab Placebo 59/301 28/151 Brig/kg orce-off Tocilizumab Standard care 21/174 34/180 Tocilizumab Standard care 13/50 15/90 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 14/66 6/64 Brig/kg Tocilizumab Standard care 2/60 1/66 Brig/kg Tocilizumab Standard care 98/366 142/412 Tocilizumab Standard care 98/366 142/412	Tocilizumab Standard care 7/64 8/67 Brigikg Tocilizumab Placebo 59/301 28/151 Brigikg Tocilizumab Standard care 21/174 34/180 Tocilizumab Standard care 13/90 15/90 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 14/65 6/64 Brigikg Tocilizumab Standard care 14/65 6/64 Brigikg Tocilizumab Standard care 2/60 1/66 Brigikg Tocilizumab Standard care 90/366 142/412 Tocilizumab Standard care 974/3582 991/3355	Tocilizumab Standard care 7/64 8/67 Brigkg Tocilizumab Placebo 58/301 28/151 Brigkg Tocilizumab Standard care 21/174 34/180 Tocilizumab Standard care 13/90 15/90 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 14/66 6/64 Brigkg Tocilizumab Standard care 14/66 6/64 Brigkg Tocilizumab Standard care 2/60 1/66 Brigkg Tocilizumab Standard care 98/386 142/412 Tocilizumab Standard care 98/386 142/412 Brigkg	Tocilizumab Standard care 7/64 8/67 Brig/kg Tocilizumab Placebo 58/301 28/151 Tocilizumab Standard care 21/174 34/180 Tocilizumab Standard care 13/80 15/90 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 14/66 6/64 Tocilizumab Standard care 2/60 1/66 Brig/kg Tocilizumab Standard care 99/366 142/412 Totilizumab Standard care 99/366 142/412 Totilizumab Standard care 99/366 142/412	Tocilizumab Standard care 7/64 8/67 Brigkg Tocilizumab Placebo 58/301 28/151 Tocilizumab Standard care 21/174 34/180 Tocilizumab Standard care 13/50 15/90 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 14/65 6/64 8 mg/kg Tocilizumab Standard care 2/60 1/66 Tocilizumab Standard care 90/366 142/412 Tocilizumab Standard care 90/368 142/412	Tocilizumab Standard care 7/64 8/67 Brigkg Tocilizumab Placebo 58/301 28/151 Tocilizumab Standard care 21/174 34/180 Tocilizumab Standard care 13/50 15/90 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 621/2022 729/2094 Tocilizumab Standard care 14/65 6/64 8 mg/kg Tocilizumab Standard care 2/60 1/66 Tocilizumab Standard care 2/60 1/66 Tocilizumab Standard care 90/366 142/412 Tocilizumab Standard care 90/358 941/3355	Tocilizumab Brigkg         Standard care         7/64         8/67         0.65%           Tocilizumab Brigkg         Placebo         58/301         28/151         0.65%         3.57%           Tocilizumab Brigkg core-off         Standard care         21/174         34/180         0.65%         2.34%           Tocilizumab Brigkg core-off         Standard care         21/174         34/180         0.65%         2.34%           Tocilizumab Brigkg core-off         Standard care         13/80         15/90         0.65%         1.27%           Tocilizumab Brigkg core-off         Standard care         621/2022         729/2094         0.74%         76.50%           Tocilizumab Brigkg         Standard care         14/66         6/64         0.74%         0.10%           Tocilizumab Brigkg         Standard care         2/60         1/66         0.10%         0.10%           Tocilizumab Brigkg         Standard care         98/366         142/412         0.10%         12.72%           = 0.00         Tocilizumab         Standard care         98/365         142/412         0.10%





**Figure 2:** WHO progression score level 7 or above at Day 28 [mechanical ventilation ± additional organ support (ECMO, vasopressors or dialysis) OR death.

					Pharmacol Advers	ogical treatments e events								
Study	Study Dura days	tion Intervention 1	Intervention 2	r1/N1	r2/N2		A	Ri: B	sk of C	Bias D	E	Overail	, e	lisk Ratio (95% Cl
Mild to severe														
Salama C.EMPACTA, 2020 Mild to severe	60	Tocilizumab 8 mg/kg	Placebo	127/259	67/129	•	-	•	-	-			17.30%	0.94 (0.77, 1.16
Stone JH, 2020 Moderate/severe	28	Tocilizumab 8mg/kg	Placebo	37/161	19/82		•	•	•	-	•		12.13%	0.99 (0.61, 1.61
Hermine O.CORIMUNO-19, Moderate/severe	2020 90	Tocilizumab 8mg/kg	Standard care	28/64	36/67	H	•	•	•		•		14 60%	0.81 (0.57, 1.16
Wang D, 2021 Mild to critical	14	Tocilizumab 400 mg	Standard care	20/33	4/32		-	•	-	•			5,90%	4.85 [1.86, 12.63
Rosas I.COVACTA, 2021 Moderate to critical	60	Tocilizumab 8mg/kg	Placebo	237/301	118/151		•	•	-	-	-		18.64%	1.01 (0.91, 1.12
Soin AS,COVINTOC, 2021 Moderate to critical	30	Tocilizumab 6 mg/kg/day	Standard care	33/90	22/90		-	•	-	•	•		12.72%	1.50 (0.95, 2.36
Veiga VC,TOCIBRAS, 2021 Severe	29	Tocilizumab 8 mg/kg	Standard care	29/65	21/64		•	•	•	•	•	•	12.92%	1.36 (0.87, 2.12
Salvarani C, 2020	30	Tocilizumab 8mg/kg	Standard care	13/60	5/66			•		•			5.79%	2,86 [1.08, 7.55
Heterogeneity: $Q = 21.02$ , $p = 0$	00; 1° = 81.3%	( r' = 0.11												
Rsk of bias ratings: Low Risk of Bias Some Concerns High Risk of Bias	A: Bias due B: Bias due C: Bias due D: Bias due E: Bias due	Risk of Bias Domains: to randomization to deviation from intended interve to missing data to outcome measurement to selection of reported result	nton		Intervention 1 bet	ter Interver	ntion 2 t	etter					Forest plot was n	.23 [0.93, 1.62 updated on: 03 22 20

Figure 3: Adverse events: Tocilizumab compared to Standard of care/placebo

					Pharma Seriou	s adverse events									
Study	Study Duration days	Intervention 1	Intervention 2	r1/N1	r2/N2		A	Ris B	k of Bi C	as D	E	Overall	,	Risk Ratio [	95% C
Mild to severe											_				
Salama C, 2020 Mild to severe	60	Tocilizumab 8 mg/kg	Placebo	38/259	25/129	H-		•		•	•	•	13.56%	0.76 [0.48	1.20
Stone JH, 2020 Moderate/severe	28	Tocilizumab 8mg/kg	Placebo	28/161	12/82	<b></b>							7.37%	1.19 [0.64	, 2.2
Talaschian M, 2021 Moderate/severe	28	Tocilizumab 8 mg/kg once-off	Standard care	3/20	0/20			•		-	•	-	0.34%	7.00 [0.38,	127.32
Hermine O, 2020 Moderate/severe	90	Tocilizumab 6mg/kg	Standard care	20/64	29/67	⊢= i							13.76%	0.72 [0.46	, 1.1
Wang D, 2021 Mild to critical	14	Tocilizumab 400 mg	Standard care	0/33	1/32		•	•	•		•	•	0.28%	0.32 [0.01	, 7.8
Rosas I, 2021 Moderate to critical	60	Tocilizumab 8mg/kg	Placebo	113/301	62/151	*	-	•	-	-	•	•	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		-	•		•	•	•	7.42%	1.20 [0.65	. 2.2
Veiga VC, 2021 Severe	29	Tocilizumab 6 mg/kg	Standard care	<mark>11/65</mark>	7/64			•		•	•	•	3.66%	1.55 [0.64	, 3.7
Salvarani C, 2020 Severe/critical	30	Tocilizumab 8mg/kg	Standard care	1/60	2/66			-	-				0.51%	0.55 [0.05	5.9
Gordon AC, 2021	90	Tocilizumab 8 mg/kg	Standard care Total:	9/366 241/1419	11/412 164/1113					•			3.77%	0.92 [0.39	, 2.20
teterogeneity: Q = 0.95, p =	= 0.64; 1" = 0.0%; 1" = 0.00					-									
Risk of bias ratings:	Risk of	Bias Domains:			Intervention 1	hatter i Interver	tion 7 P	ottor					C	0.92 [0.77	, 1.08
Some Concerns High Risk of Blas	B Bas due to faile C Bias due to missi D Bas due to outco E Bias due to selec	tion from intended intervent ng data me measurement tion of reported result	lion		Intervention 1	0.14 1.95	1001 2 1	101104					Forest plot was i	updated on: 0	)5 14 2i



Appendix 4	4:	<b>Evidence</b>	to	decision	framewor	'k
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	JUDGEMENT	EVIDENCE & ADDITIONAL CONSIDERATIONS
EVIDENCE OF BENEFIT	What is the size of the effect for beneficial outcomes?         Large       Moderate       Small       None       Uncertain         X	A meta-analysis of 12 RCTs demonstrated that TCZ is associated with an all-cause mortality benefit at 28 days. ARR 3.5% (95% CI 1.5% to 5.6%); RR 0.88 (95% CI 0.81 to 0.85, eleven RCTs, n = 6 937). NNT 29 (95% CI 18 to 67) to prevent 1 death.
EVIDENCE OF HARMS	What is the size of the effect for harmful outcomes?         Large       Moderate       Small       None       Uncertain         X       X       X       X	No increased risk of infections or adverse events was observed with tocilizumab use in the RCTs <sup>29</sup>
BENEFITS & HARMS	Do the desirable effects outweigh the undesirableharms?FavoursFavoursInterventionInterventioninterventioncontrolControl= Control orUncertainUncertain	Available evidence shows that TCZ is associated with an all- cause mortality benefit at 28 days, with no reports of associated increased risk of infections or adverse events.
QUALITY OF EVIDENCE	What is the certainty/quality of evidence?         High       Moderate       Low       Very low         X       Image: Confident in the evidence         Moderate       quality: confident in the evidence         Moderate quality: mostly confident, but further research may change the effect         Low quality: some confidence, further research likely to change the effect         Very low quality: findings indicate uncertain effect	<ul> <li>The findings are based on the finding of a large RCT and meta- analysis of 12 RCTs.</li> <li>Assessed per outcome – refer to table 1:</li> <li>Mortality at D28: high certainty evidence.</li> <li>WHO progression score (level 7 or above) D28: low certainty evidence</li> <li>Serious adverse events: moderate certainty evidence</li> </ul>
FEASABILITY	Is     implementation     of     this     recommendation       feasible?     Yes     No     Uncertain       Yes     X     X	Tocilizumab is SAHPRA registered and available on the South African market, however price and affordability is a concern.
RESOURCE USE	How large are the resource requirements?         More       Less intensive         Intensive         X       Intensive	Price of medicines/dose (8mg/kg)MedicineOffered price*60 kg patient: Tocilizumab 480mgR 4 435.6475 kg patient: Tocilizumab 600mgR 5 253.92Maximum dose: Tocilizumab 800mgR 7 005.16*Offered price from Roche Products (Pty) Ltd (letter dated 30 March 2021)400mg/20ml = R 3 502.58; 200mg/10ml = R 1 751.34; 80mg/4ml = R 933.06Basic costing analysis:• Clinical inputs:- Public sector hospitalizations in 1 <sup>st</sup> and 2 <sup>nd</sup> waves: ≈ 40000 and60000, respectively (DATCOV survey)(Note: Number of public facilities reporting data improved in the second wave; and modelling data for 3 <sup>rd</sup> wave predictions was not available from South African COVID-19 Modelling Consortium, at the time of this report)- Incidence of elevated CRP>75 mg/l: 30-50% (Western Cape PHDC)> >75 mg/l = 7867/26351 = 30%> >150 mg/l = 4528/26351 = 20%• Assumptions:- DATCOV hospital surveillance data assumed to project total future national COVID-19 hospitalizations. Average for 3 <sup>rd</sup> wave 50000 (lower and upper limit of 40000 and 60000, respectively).Note: Timing of the effect of COVID-19 vaccine on herd immunity needs consideration- Western Cape data regarding the incidence of raised CRP assumed to be generalisable to the whole country. Average of 40% (upper and lower limit of 30% and 50%, respectively).

		<ul> <li>Patient with CRP&gt;75mg/l assumed to be severe, requiring supplemental oxygen.</li> </ul>
		<ul> <li><u>Estimated forecast:</u> <ul> <li>Estimated number of patients who will be eligible for tocilizumab treatment: 20000 (lower and upper limit of 12000 and 30000, respectively).</li> </ul> </li> </ul>
		<ul> <li><u>Estimated budget impact at current SEP:</u></li> <li>Treatment cost (using offered price of 30 March 2021) @R5 250 per patient (75 kg); then total cost estimated as R105 mil (lower and upper limit of R65 mil to R160 mil).</li> </ul>
		<u>References</u> Data on file, Western Cape NHLS     Data on file, DATCOV concerts, NICD
		<ul> <li>Offered price from Roche Products (Pty) Ltd (letter dated 30 March 2021)</li> <li>Rapid review of Tocilizumab for CoVID-19 Update, 3 March 2021</li> </ul>
	Is there important uncertainty or variability about	No specific research surveying patients' or healthcare
. >	how much people value the options?	workers' value of this therapeutic agent is currently available.
ES,	Minor Major Uncertain	
UES, RENC FABI		The Committee was of the opinion that the option would be
FEF EP	Is the option acceptable to key stakeholders?	acceptable to key stakeholders.
VCC	Yes No Uncertain	
- 1	X	
7	Would there be an impact on health inequity?	Tocilizumab currently only available in private sector and is
EQUIT	Yes No Uncertain	expensive.

## Appendix 5: Updating of rapid report

Date	Signal	Rationale
10 January 2021 & 11	Preprints of REMAP-CAP and	The preliminary study results of the tocilizumab arm of the REMAP-CAP and
February 2021	RECOVERY trials	RECOVERY trials have been published in preprint format
6 May 2021	RECOVERY trial published in	The RECOVERY trial data is now published in peer review format.
	Lancet	