Tocilizumab in COVID-19: a meta-analysis, trial sequential analysis, and meta-regression of randomized-controlled trials

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Abstract

Purpose: Interleukin-6 (IL-6) levels discriminate between patients with mild and severe COVID-19, making IL-6 inhibition an attractive therapeutic strategy. We conducted a systematic review, meta-analysis, trial sequential analysis (TSA), and meta-regression of randomized-controlled trials to ascertain the benefit of IL-6 blockade with tocilizumab for COVID-19.

Methods: We included randomized-controlled trials (RCTs) allocating patients with COVID-19 to tocilizumab. Our control group included standard care or placebo. Trials co-administering other pharmacological interventions for COVID-19 were not excluded. Primary outcome was 28–30 day mortality. Secondary outcomes included progression-to-severe disease defined as need for mechanical ventilation, intensive-care unit (ICU) admission, or a composite.

Results: We identified 10 RCTs using tocilizumab, 9 of which reported primary outcome data (mortality), recruiting 6493 patients with 3358 (52.2%) allocated to tocilizumab. Tocilizumab may be associated with an improvement in mortality (24.4% vs. 29.0%; OR 0.87 [0.74–1.01]; p = 0.07; I² = 10%; TSA adjusted CI 0.66–1.14). Meta-regression suggested a relationship between treatment effect and mortality risk, with benefit at higher levels of risk (logOR vs %risk beta = −0.018 [−0.037 to −0.002]; p = 0.07). Tocilizumab did reduce the need for mechanical ventilation and was associated with a benefit in the composite secondary outcome but did not reduce ICU admission.

Conclusions: For hospitalized COVID-19 patients, there is some evidence that tocilizumab use may be associated with a short-term mortality benefit, but further high-quality data are required. Its benefits may also lie in reducing the need for mechanical ventilation.

Keywords: COVID-19, Immunologic factors, Interleukin-6, Meta-analysis
Introduction

Patients with coronavirus disease 2019 (COVID-19) demonstrate a heterogeneous clinical course ranging from mildly symptomatic disease to acute respiratory distress syndrome (ARDS) and death [1]. Hospital mortality in patients admitted to US hospitals during the first pandemic was 9.6% [2]. Short- and long-term morbidity associated with COVID-19 are also significant [3].

The beneficial effect of dexamethasone on mortality among critically ill patients with COVID-19 highlights the role of an excessive host inflammatory response in the progression of mild disease-to-critical illness and death [4]. In addition to corticosteroids, multiple other immunomodulatory drugs have been proposed as therapeutic candidates [5].

Interleukin-6 (IL-6) is a key regulator of C-reactive protein (CRP) production and fever, biomarkers of the severity of COVID-19 [6]. IL-6 levels also discriminate between patients with mild and severe disease [7], making IL-6 inhibition an attractive therapeutic strategy. However, the absolute levels of IL-6 in patients with COVID-19 are significantly lower than those seen in other systemic inflammatory disorders such as bacterial sepsis [8], raising questions about the potential benefit of IL-6 blockade as a viable therapeutic strategy in COVID-19.

We conducted a systematic review, meta-analysis, and trial sequential analysis (TSA) to ascertain the benefit of tocilizumab, the most commonly used IL-6 antagonist in COVID-19.

Methods

The protocol for this review was registered with the International prospective register of systematic reviews (PROSPERO registration number: CRD42021231300) and is reported according to PRISMA guidelines (Online Resource) [9].

Information sources and search strategy

A systematic search of PubMed, Embase, Cochrane Library, and MedRxiv using a controlled vocabulary (MeSH) and keywords. Date and language restrictions were not applied. The last search update was on 7th March 2021. The Boolean search strategy was as follows: ((Tocilizumab OR Sarilumab OR Interleukin-6 OR IL-6) AND (COVID-19 OR SARS-CoV-2) AND ((Clinical trial OR Randomized OR Trial OR RCT)).

Research papers and review articles were also hand-searched for further relevant trials. Where data on the primary outcome were not available from the manuscript, the corresponding author was contacted for this information.

Take-home message

There is some evidence that the use of tocilizumab may be associated with a short-term mortality benefit in patients with COVID-19. Amongst patients not requiring advanced respiratory support, it may also reduce disease progression to requiring mechanical ventilation. However, most trials are at high risk of bias and further high-quality data is required.

Eligibility criteria

Inclusion and exclusion criteria were determined a priori. All trials comparing patients who received tocilizumab IL-6 blockade in patients with COVID-19 were considered. To avoid potential confounding, where sicker patients were more likely to receive tocilizumab, we only included randomized-controlled trials. We included patients being treated with other COVID-19 therapies (co-interventions), as part of other trials, with the control group defined as those not receiving IL-6 antagonists. Details of co-interventions are provided in the Supplementary Data. Trials enrolling pediatric patients (<18 years were excluded).

Trial selection

Two investigators (NS and TS) independently screened both titles and abstracts to exclude non-relevant trials. Discrepancies were resolved by a third author (NA). Relevant full-text articles were retrieved and analyzed for eligibility using the pre-defined inclusion criteria.

Data collection and analysis

Two investigators (NS and TS) independently extracted information from selected trials using a standardized data collection form. Data were collected on the following: country of trial, total number of participants, dosing of IL-6 receptor antagonist, age and number of patients receiving mechanical ventilation, noninvasive ventilation (NIV), or high-flow nasal oxygen (HFNO) at enrollment.

Primary and secondary outcomes

Primary outcome was 28–30 day mortality. Secondary outcomes included markers of progression-to-severe disease which were defined as either requirement for mechanical ventilation, intensive-care unit (ICU) admission, or a composite of the above.

Subgroup analyses

Our pre-defined sub-group analysis included only patients admitted to intensive care unit (ICU) at enrollment. IL-6
inhibition may be expected to provide the greatest benefit in those at greatest risk of death. Therefore, we performed a meta-regression to investigate the relationship between treatment effect and overall risk. Additionally, as tocilizumab is an IL-6 inhibitor, responsible for regulation of CRP we anticipated, it would provide the greatest benefit in those with a higher baseline CRP. We thus performed a meta-regression to evaluate the interaction between baseline CRP and treatment effect.

**Risk of bias assessment**
Methodological quality of the included randomized-controlled trials was assessed using the Cochrane Collaboration’s tool for assessing risk of bias (RoB2) [10] independently by two authors (NS and TS), with any discrepancies reconciled by a third (NA). The following domains were assessed: randomization process, assignment to intervention, missing outcome data, measurement of outcome, selection of the reported result, other bias, and overall bias. The risk of bias in each domain was judged as either low, high, or some concerns.

**Grading the quality of evidence**
Two authors (NS and TS) assessed the quality of each outcome measure in accordance with the grading of recommendations assessment, development, and evaluation (GRADE) approach (GRADEpro Guideline Development Tool. McMaster University, 2015) [11]. Quality was downgraded on the basis of the following certainty assessment: risk of bias, inconsistency, indirectness, imprecision, and other considerations. Discrepancies were resolved using a third author (NA). Publication bias was assessed using a funnel plot and Harbord’s test [12]. The overall quality of evidence was subsequently rated as “high”, “moderate”, “low”, or “very low”.

**Statistical analysis**
We combined individual trial data for mortality with the reference group taken as the group not randomized to an IL-6 antagonist. The meta-analysis was performed using the review manager (‘Revman’) for Mac (version 5.1, Cochrane Collaboration, Oxford, UK). Statistical heterogeneity was assessed using the $I^2$ methodology. $I^2$ values $>30\%$, $>50\%$, and $>75\%$ were considered to indicate moderate, substantial, and considerable heterogeneity among trials, respectively. A random-effects model was used to analyze data. All $p$ values were two-tailed and considered statistically significant if $<0.05$. Data on dichotomous outcomes are presented as odds ratio (OR), 95% confidence intervals, $p$ values; $I^2$ values. Meta-regression was performed to investigate the effect of overall risk using control group event rate, and average baseline CRP of the treatment group at enrollment, using a random-effects model (Dersimonian-Laird) in Stata (version 16-1, StataCorp, College Station, TX, USA. 2019).

Because type 1 errors may result from meta-analyses with too small sample sizes, we performed Trial Sequential Analysis (TSA) using TSA program version 0.9.5-10 (www.ctu.dk/tsa). TSA tests the credibility of the ascertained results by combining both an estimation of information size (a cumulative sample size of included trials) with an adjusted threshold of statistical significance for the cumulative meta-analysis. Meta-analysis monitoring boundaries (Trial Sequential Monitoring Boundaries) and the required information size (RIS) were quantified, alongside diversity-adjusted information size ($D^2$) and adjusted 95% confidence intervals. Diversity adjustment was performed according to an overall type I error of 5% and power of 80%. Given the novelty of both COVID-19 and the use of IL-6 inhibitors in respiratory disease, RIS was calculated using the relative risk reduction (RRR) obtained from our actual meta-analysis of 15.7%.

**Protocol changes**
The following changes were made to our PROSPERO published protocol. The definition of our control group was extended to include patients receiving standard care or placebo, and other potential COVID-19 treatments either in or out of a clinical trial given the number of platform trials identified. Only one trial reported outcomes for patients stratified by respiratory support, and thus, we were unable to perform this sub-group analysis. We used the random-effect models, rather than a fixed-effects model due to the number of trials identified, but have included the results using both a fixed-effects model and risk ratios as a sensitivity analysis. We performed an additional sensitivity analysis on patients who received sarilumab to investigate a drug versus class treatment effect, and on the trials at low risk of bias.

**Results**
**Search strategy**
Our search strategy identified 2175 results. Following removal of duplicates, 1520 articles remained. Of these, 1504 were excluded on the basis of title/abstract. Of the remaining 16, five were excluded at full review as two were non-randomized [13, 14], two were review articles [15, 16], and one was performed on non-COVID patients [17]. Of the remaining 11 articles [18–28], one trial used sarilumab [22] and one did not report mortality data [18]; the corresponding authors were contacted but did not reply. Thus, nine trials were used for the primary outcome analysis [19–21, 23–28], ten for sensitivity analysis,
[19–28], and ten for secondary analyses [18–21, 23–28]. (Fig. 1). Mortality at day 28–30 was not reported in one trial [19]; we contacted the corresponding author, but the data were not available. In-hospital mortality was therefore used for this trial.

**Trial characteristics**

Only five trials enrolled patients requiring mechanical ventilation [19, 21, 23, 27, 28]. Seven trials enrolled patients receiving NIV [18, 19, 21, 23, 24, 27, 28], while five enrolled patients receiving HFNO [19, 21, 23, 26, 27]. Two trials recruited patients on supplemental oxygen alone [20, 25] (Table 1). Nine trials used tocilizumab [18, 20, 21, 23–28], one trial used sarilumab [22], and one trial used either tocilizumab or sarilumab [19]. Subsequent analyses were performed using data from patients receiving tocilizumab only, with sarilumab used for a sensitivity analysis.

Eight trials used an initial dose of 8 mg/kg, which could be repeated at treating physician discretion within 24 h in seven trials [18, 19, 23–27], or on day 3 in one trial [20]. One trial used a dose of 6 mg/kg, which could be repeated within 7 days if clinical worsening or no improvement [28]. One trial used a weight-based dosing strategy which could be repeated with 24 h at physician discretion [21]. Four trials used a placebo control [22–25], while the control group was defined as standard care in the remaining trials. All trials allowed the use of additional COVID-19 treatments, in particular, glucocorticoids were used as a co-intervention in 72% of enrolled patients. (Online
Table 1  Baseline characteristics for included trials

| Author/group/NCT registration | Country            | Recruitment dates          | Recruitment window | Tocilizumab dosing                                                                 | Control group (n) | Treatment group (n) | Control group (age) | Treatment group (age) | Control group (numbers ventilated) | Treatment group (numbers ventilated) | Control group (numbers on NIV) | Treatment group (numbers on NIV) | Control group (numbers on HFNO) | Treatment group (numbers on HFNO) |
|-------------------------------|--------------------|----------------------------|-------------------|-------------------------------------------------------------------------------------|------------------|---------------------|----------------------|-----------------------|-----------------------------------|-----------------------------------|-------------------------------|-------------------------------|----------------------------------|-----------------------------------|
| Gordon (REMAP-CAP) NCT02735707 | Multi-national     | April 19, 2020–November 19, 2020 | Within 24hrs of ICU admission | 8 mg/kg (maximum 800 mg) repeated at 12-24hrs if needed | 402              | 353                 | 61 ± 13              | 62 ± 13               | 121/402 (30.1%)                   | 104/353 (29.5%)                   | 169/402 (42.0%)                  | 147/353 (41.6%)                  | 110/402 (27.4%)                  | 101/353 (28.6%)                  |
| Horby (RECOVERY) NCT04381936  | United Kingdom     | April 14, 2020–Jan 24, 2021  | Within 21 days of primary randomization | 800 mg if weight > 90 kg, 600 mg if weight > 65 and ≤ 90 kg; 400 mg if weight > 40 and ≤ 65 kg; and 8 mg/kg if weight ≤ 40 kg repeated 12 – 24hrs later if needed | 2094             | 2022                | 64 ± 14              | 63 ± 14               | 294/2094 (14.0%)                   | 268/2022 (13.3%)                   | 867/2094 (41.4%)                  | 819/2022 (40.5%)                  | (included with NIV)              | (included with NIV)              |
| Hermine (CORIMUNO) NCT04331808 | France             | March 31, 2020–April 18, 2020 | Within 72hrs of SAR-CoV-2 diagnosis | 8 mg/kg on day 1 (and 3 if needed) | 67               | 63                  | 64 ± 4               | 65 ± 5               | 0                                 | 0                                 | 0                              | 0                              | 0                              | 0                                 |
| Rosas (COVACTA) NCT04320615   | Multi-national     | NS                          | NS                 | 8 mg/kg (maximum 800 mg) repeated at 8-24hrs if needed | 144              | 294                 | 67 ± 14              | 61 ± 15               | 54/144 (37.5%)                   | 111/294 (37.8%)                   | 40/144 (27.8%)                   | 68/294 (23.1%)                   | (included with NIV)              | (included with NIV)              |
| Salama (EMPACTA) NCT04372186  | Multi-national     | NS                          | Within 48 h of hospital admission | 8 mg/kg (maximum 800 mg) repeated at 12hrs | 128              | 249                 | 56 ± 15              | 56 ± 14               | 0                                 | 0                                 | 0                              | 0                              | 0                              | 0                                 |
| Salvareani (RCT-TCZ-COVID-19) NCT04346355 | Italy               | March 31, 2020–June 11, 2020 | NS                 | 8 mg/kg (maximum 800 mg) repeated at 12hrs | 63               | 60                  | 61 ± 4               | 62 ± 6               | 0                                 | 0                                 | 0                              | 0                              | NS                             | NS                               |
Table 1 (continued)

| Author/group/NCT registration | Country | Recruitment dates | Recruitment window | Tocilizumab dosing | Control group (n) | Treatment group (n) | Control group (age) | Treatment group (age) | Control group (numbers ventilated) | Treatment group (numbers ventilated) | Control group (numbers on NIV) | Treatment group (numbers on NIV) | Control group (numbers on HFNO) | Treatment group (numbers on HFNO) |
|------------------------------|---------|-------------------|--------------------|--------------------|-------------------|--------------------|---------------------|---------------------|-------------------------------|-------------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|
| Soin (COVINTOC) CTRI/2020/05/025369 | India | May 30, 2020–Aug 21, 2020 | NS | 6 mg/kg (maximum 480 mg) repeated up to 7 days later if needed | 88 | 91 | 54±6 | 56±5 | 4/88 (4.5%) | 5/91 (5.5%) | 20/88 (22.7%) | 28/91 (30.8%) | NS | NS |
| Stone (BACC) NCT04356937 | United States | April 20, 2020–June 15, 2020 | Upon hospital admission | 8 mg/kg (maximum 800 mg) | 81 | 161 | 56±6 | 60±7 | 0 | 0 | 5/81 (6.2%) | 5/161 (3.1%) | 0 | 0 |
| Veiga (TOCIBRAS) NCT04403685 | Brazil | May 8, 2020–July 17, 2020 | NS | 8 mg/kg (maximum 800 mg) | 64 | 65 | 57±14 | 57±16 | 10/64 (15.6%) | 11/65 (16.9%) | 26/64 (40.6%) | 15/65 (23.1%) | (included with NIV) | (included with NIV) |
| Zhao NCT04310228 | China | February 2, 2020–March 15, 2020 | NS | 4–8 mg/kg repeated at 24hrs | 7 | 19 | 69±13 | 66±14 | 0 | 0 | 1/7 (14.3%) | 0 | 0 | 0 |

NIV non-invasive ventilation, HFNO high flow nasal oxygen, NS not stated
Table 2 Primary, sub-group, secondary, and sensitivity outcome data for included trials

| Outcome                        | References                  | Intervention group | Control group | Conventional effect estimate (95% CI) | Overall effect | I² (%) |
|--------------------------------|-----------------------------|--------------------|---------------|--------------------------------------|----------------|--------|
| Overall mortality              | [19–21, 23–28]              | 821/3358 (24.4%)   | 909/3135 (29%) | 0.87 (0.74–1.01)                     | Z = 1.82 p = 0.07 | 10     |
| ICU patient mortality          | [19, 21, 23]                | 254/732 (34.7%)    | 297/750 (39.6%)| 0.84 (0.65–1.10)                     | Z = 1.27 p = 0.20 | 24     |
| Disease progression            |                             |                    |               |                                      |                |        |
| Mechanical ventilation         | [20, 21, 23–26, 28]         | 152/1742 (8.7%)    | 152/1454 (10.5%)| 0.70 (0.54–0.89)                     | Z = 2.86 p = 0.004 | 0      |
| ICU admission                  | [20, 23, 26, 28]            | 118/338 (34.9%)    | 117/282 (41.5%)| 0.73 (0.38–1.39)                     | Z = 0.96 p = 0.34 | 60     |
| Composite outcome              | [18–21, 23–27]              | 808/2796 (28.9%)   | 943/2577 (36.6%)| 0.72 (0.59–0.89)                     | Z = 3.14 p = 0.002 | 26     |
| Sensitivity analysis           |                             |                    |               |                                      |                |        |
| Combined IL-6 antagonists mortality |                     | 861/3738 (23%)    | 916/3219 (28.5%)| 0.86 (0.74–1.01)                     | Z = 1.85 p = 0.06 | 10     |
| Sarilumab mortality            | [19, 23]                    | 40/377 (10.6%)     | 149/481 (31%) | 0.72 (0.35–1.51)                     | Z = 0.86 p = 0.39 | 42     |

ICU intensive-care unit

Fig. 2 Effect of tocilizumab on mortality in included trials. a Forest plot of mortality in RCTs listed in descending order of control group mortality. Size of squares for odds ratio reflects weight of trial in pooled analysis. Horizontal bars represent 95% confidence intervals. b Trial sequential analysis of mortality in RCTs. Uppermost and lowermost curves represent trial sequential monitoring boundary lines for benefit and harm, respectively. Horizontal lines represent the traditional boundaries for statistical significance. Triangular lines represent the futility boundary. The cumulative Z curve represents the trial data. A diversity-adjusted required information size (RIS) of 5622 was calculated using α = 0.05 (two sided), β = 0.20 (power 80%). Relative risk of mortality reduction was 15.7%. The cumulative Z curve crosses neither the conventional nor the TSA boundary for benefit or harm, but did cross the boundary for futility having exceed the required information size (RIS). c Meta-regression of log odds ratio for mortality vs. risk (%).
Table 1) Rates of reported adverse events were low, with no differences between the tocilizumab and control arms. (Online Table 2).

Primary outcome
Mortality was defined at 28–30 days in eight trials [20, 21, 23–27], and in-hospital mortality in one trial [19]. A total of 6493 patients with 3358 (51.7%) allocated to the tocilizumab arm and a mean weighted mortality of 26.6% were included. Tocilizumab treatment was not associated with an improvement in mortality compared to standard care (24.4% vs. 29.0%; OR 0.87 [0.74–1.01]; p = 0.07; I² = 10%; TSA adjusted CI 0.66–1.14). The cumulative Z curve crossed neither the conventional nor the TSA boundary for benefit or harm, but did cross the boundary for futility having exceed the required information size (RIS). (Table 2 and Fig. 2a, b) At time of reporting of mortality, 1086 (32.3%) patients in the tocilizumab group, and 1172 (37.4%) patients in the control group remained as inpatients.

Subgroup analyses
Three trials [19, 21, 23] reported mortality for critically ill patients (n=1482) requiring ICU admission at enrollment which did not demonstrate a statistically significant mortality benefit (34.7% vs. 39.6%; OR 0.84 [0.65–1.10]; p = 0.05; I² = 24%) (Online Fig. 1).

Meta-regression suggested a weak relationship between treatment effect and overall risk of mortality (Fig. 2c). There was weak evidence of mortality benefit for higher levels of overall risk (logOR vs %risk beta = −0.018 [−0.037 to −0.002]; p = 0.07). However, there was no evidence of a relationship with baseline CRP (logOR vs. baseline CRP beta = 0.005 [−0.005 to 0.016]; p = 0.32).

Sensitivity analysis
We performed an analysis on the two trials using sarilumab [21, 22]. This included 858 patients with 377 (43.9%) allocated to the sarilumab group and a mean weighted mortality of 22.0%. Sarilumab was not associated with a mortality benefit (10.6% vs. 31.0%; OR 0.72 [0.35–1.51]; p = 0.39; I² = 42%).

An additional analysis was performed incorporating all IL-6 inhibitors. This included 6957 patients of which 3738 (53.7%) were allocated to the treatment arm with a weighted mean mortality of 25.5%. IL6-antagonism was not associated with a mortality benefit (23.0% vs. 28.5%; OR 0.86 [0.74–1.01]; p = 0.06; I² = 10%).

A sensitivity analysis of five trials with low risk of bias [20, 23–26] was performed which included 1314 patients of which 827 (62.9%) were allocated to the treatment arm. Tocilizumab use was not associated with a mortality benefit (12.3% vs. 10.7%; OR 1.09 [0.75–1.57]; p = 0.65; I² = 0%).

An additional sensitivity analysis was performed assessing mortality benefit using a fixed-effects model. Tocilizumab was associated with a mortality benefit on conventional analysis only (OR 0.85 [0.76–0.96]; p = 0.006; I² = 10%; TSA adjusted CI 0.70–1.04). However, analysis using relative risk (RR) with a random-effects model showed a mortality benefit (RR 0.89 [0.82–0.96]; p = 0.005; I² = 10%; TSA adjusted CI 0.80–0.99), as did a fixed-effects model (RR 0.89 [0.83–0.97]; p = 0.006; I² = 0%; TSA adjusted CI 0.81–0.99).

Secondary outcomes
Seven trials including 3196 patients reported progression from a supplemental oxygen requirement to mechanical ventilation [20, 21, 23–26, 28]. Of these, 1742 (54.5%) were allocated to the tocilizumab arm with a mean weighted incidence of 9.5%. Tocilizumab was associated with a reduction in requirement for mechanical ventilation compared to standard care on conventional analysis only (8.7% vs. 10.5%; OR 0.70 [0.54–0.89]; p = 0.004; I² = 0%; TSA adjusted CI 0.43–1.13). The cumulative Z curve crossed the conventional boundary for benefit, but not the TSA boundary with 31.7% of RIS cases accrued (Fig. 3).

Progression to ICU admission was reported in four trials including 620 patients, with 338 (54.5%) allocated to the tocilizumab group and a weighted mean incidence of 37.9% [20, 23, 26, 28]. Tocilizumab was not associated with a reduced rate of ICU admission (34.9% vs. 41.5%; OR 0.86 [0.74–1.01]; p = 0.05; I² = 10%; TSA adjusted CI 0.66–1.14). The cumulative Z curve crossed the conventional boundary for benefit, but not the TSA boundary with 31.7% of RIS cases accrued (Online Fig. 3).

Risk of bias and grade recommendation
The risk of bias was high due to the open label approach taken in six trials [19–21, 26–28]. Ten trials included industry sponsorship [19–27]. Three trials released their results as pre-prints prior to peer review [19, 21, 22] (Online Table 3). Inconsistency among the trials was
low due to low heterogeneity excluding 'ICU admission', and indirectness was adjudicated to be not serious due to the populations and outcomes measured in the trials. Imprecision was judged to be very serious for both 'need for mechanical ventilation' and 'need for ICU admission' due to TSA analysis showing low percentages of RIS cases accrued. While the funnel plot for publication bias was asymmetrical, this was towards the negative trials. Harbord's test suggested a small trial effect ($p = 0.11$), which when adjusted for overall risk effect disappeared ($p = 0.82$). Overall, the quality of evidence by GRADE assessment was marked either 'moderate' or 'very low' (Online Table 4 and Online Fig. 4).

**Discussion**

Among all hospitalized patients with COVID-19, there is some evidence that tocilizumab use may be associated with an overall mortality benefit, although trial sequential analysis suggests futility in continuing trial recruitment. The well-established association between elevated CRP and illness severity in COVID-19 [6] raises the possibility of a mortality benefit in the sickest patients. This finding is supported by meta-regression which suggests a survival benefit for patients at higher mortality risk. This mortality benefit was seen only in the REMAP-CAP and RECOVERY trials where patients in the control arm had the highest mortality compared to other trials. ICU
admission and advanced respiratory support were prerequisites for trial enrollment into REMAP-CAP, in contrast to four of the other trials where these were exclusion criteria.

Among patients with less severe disease, tocilizumab may reduce progression-to-severe disease and reduce the need for mechanical ventilation. However, TSA suggests that further data are required before firm conclusions can be drawn. Caution is required in interpreting the findings given not all patients who receive tocilizumab will be considered appropriate for mechanical ventilation. For example, in the RECOVERY trial, which provides the bulk of the data, almost two-thirds of the patients not mechanically ventilated at enrollment who subsequently died did not receive ventilation. With many ongoing RCTs, the potential benefits of tocilizumab in milder cases of COVID-19 may become clearer.

Following early reports of a cytokine storm associated with severe COVID-19 disease, several immunomodulatory drugs were repurposed with the hope of discovering effective therapeutic strategies [5]. A search of clinicaltrials.gov on 3rd July 2020 identified 1366 registered trials for COVID-19 disease, of which 279 were RCTs assessing immunomodulatory therapies. These include targets against 39 different immune pathways using 90 different drugs or therapies; 47 registered RCTs were evaluating inhibition of IL-6 [5].

While IL-6 values in COVID patients are significantly lower than seen in other inflammatory conditions including non-COVID ARDS, sepsis, and cytokine release syndrome [8], it does discriminate between patients with mild and severe COVID-19 disease [7]. Early observational studies describing the reduction in systemic inflammation biomarkers (CRP, fever) in response to tocilizumab support the biological plausibility of its use in COVID-19 disease, despite the lack of clinical data supporting its use in non-COVID-19 ARDS [29]. The timing of administration of tocilizumab early in the disease remains consistent across trials, although the broad enrollment criteria used may have diluted the effect, as may have the high level of corticosteroid co-prescribing which may explain the lack of correlation seen between treatment effect and baseline CRP value. Early administration of interleukin-6 receptor blockade may interrupt the inflammatory cascade preventing deterioration from mild respiratory failure to into ARDS, multi-organ failure, and eventually death.

There are several limitations to this analysis. It is not possible to evaluate the effect of different dosing strategies on outcome. Seven trials permitted a second dose of tocilizumab, but only one reported outcomes in relation to dose administered [19]. The number of co-interventions (including steroids and anti-viral medication) varied between trials, which we were unable to adjust for. The concurrent use of systemic corticosteroids is of particular relevance given the outcome benefit reported in patients receiving oxygen or advanced respiratory support at randomization [4]. Both the RECOVERY and REMAP-CAP trials demonstrate that estimates of the treatment effect for patients treated either with tocilizumab (or sarilumab) and corticosteroids in combination were greater than with an IL-6 antagonist alone. In both these trials, which account for 75% of the total population, and 88% of the deaths, co-administration of corticosteroids was high. There was no associated mortality benefit seen with tocilizumab in the subset of patients not administered corticosteroids in the RECOVERY trial, suggesting either some interaction between corticosteroids and tocilizumab, or that there is no additional benefit of tocilizumab. Additionally, these data may provide some reassurance surrounding excessive immunosuppression and risk of increased mortality with co-administration of steroids and tocilizumab.

The reported incidence of infectious and other complications varied significantly between trials. This may relate to differences in definitions, screening, and reporting of complications, and variations in patient follow-up. While there is no evidence of increased rates of adverse events with tocilizumab, this finding should be interpreted with caution given the number of reported events is lower than might be expected.

Crucially, the data in this meta-analysis are heavily weighted by two trials [19, 21] with the highest overall risk of mortality. These trials were prone to high risk of bias, having an open label trial design and patients being allocated to treatments based on drug availability at participating sites which may explain why sensitivity analysis of low risk of bias trials failed to show a mortality benefit. While the TSA analysis suggest futility in ongoing recruitment, this should be interpreted in this context and that a smaller, but still significant clinical effect may still exist which would alter the futility boundaries.

It remains difficult to reconcile directly conflicting trial data, where two trials reported a significant improvement in mortality with tocilizumab [19, 21], while another was terminated early due to an excess mortality risk [27]. Further high-quality trial data are required before firm conclusions can be made to guide clinical practice. This includes longer term outcomes as a third of patients remained as inpatients at the data censure cut-point, raising the possibility that tocilizumab may just prolong time to death.

In summary, there is some evidence that tocilizumab use may be associated with a short-term mortality benefit in patients with COVID-19, but further high-quality data are required. Among patients not requiring advanced
respiratory support, tocilizumab may also prevent progression to mechanical ventilation.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1007/s00134-021-06416-z.

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**Author contributions**

Study conception: NA, literature search: NS, TS, and NA; data extraction: NS and TS; assessment of bias: TS and NS; statistics: TS, GA, and NA; drafting manuscript: TS and NA; critical review: EN and MS; finalizing manuscript: all authors.

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**Availability of data**

All authors had access to data.

**Declarations**

**Conflicts of interest**

MS reports grants and advisory board fees from NovoB, grants from the Defence Science and Technology Laboratory, Critical Pressure, Apollo Therapeutics, advisory board and speaker fees (paid to his institution) from Amomred, Biotech, GE, Baxter, Roche, and Bayer, and honorarium for chairing a data monitoring and safety committee from Shionogi.

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