COVID-19 and Disease-Modifying Anti-rheumatic Drugs

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Abstract
Purpose of Review Patients on disease-modifying anti-rheumatic drugs (DMARDs) remain concerned about potential risks of severe COVID-19 outcomes. Meanwhile, several DMARDs have been proposed as COVID-19 therapies.

Recent Findings In patients with autoimmune diseases, baseline glucocorticoid use is associated with severe COVID-19. While classes of DMARDs (e.g., conventional synthetic, targeted synthetic, and biologic) do not appear to be associated with higher risk, specific medications such as rituximab and sulfasalazine may be associated. Randomized clinical trials (RCTs) show that glucocorticoids reduce mortality in severe COVID-19. RCTs suggest other agents, such as baricitinib, may improve COVID-19 outcomes in certain populations.

Summary Baseline glucocorticoid use raises the risk of severe COVID-19 in patients with autoimmune diseases, but glucocorticoids are an effective treatment for those with severe COVID-19. Further research is needed to inform DMARD management in autoimmune disease patients during the pandemic and the role of DMARDs in COVID-19 treatment.

Keywords COVID-19 • DMARD • Disease-modifying anti-rheumatic drugs

Introduction
The coronavirus disease 2019 (COVID-19) pandemic, caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), has been an unprecedented global health crisis, with >103 million confirmed cases and >2.2 million deaths worldwide as of February 2021 [1]. Patients with autoimmune diseases and their physicians remain concerned regarding potential heightened risks of severe COVID-19 due to immunosuppressive medications needed to control autoimmune disease activity [2]. Meanwhile, glucocorticoids and several disease-modifying anti-rheumatic drugs (DMARDs), such as hydroxychloroquine and interleukin-6 (IL-6) receptor inhibitors, have been proposed as potential therapies for COVID-19, and observational studies and clinical trials have revealed variable successes with these agents [3, 4–5].

In this review, we examine the literature regarding COVID-19 outcomes in patients with rheumatic diseases on glucocorticoids and/or DMARDs at the time of infection (Table 1). We then review the literature regarding the use of glucocorticoids and DMARDs for the treatment of COVID-19, focusing on data from randomized clinical trials (RCTs) (Table 2).

Risk of Severe COVID-19 in Patients on DMARDs
Early in the COVID-19 pandemic, case series of rheumatic disease patients with COVID-19 reported generally mild clinical courses in patients with inflammatory arthritis, systemic lupus erythematosus, and vasculitis [25, 26, 27]. Early observational studies from longitudinal clinics reported similar incidence of COVID-19 among rheumatic disease patients vs. the general population [28–31]. Although these initial reports were reassuring, a large cohort study of over 17 million patients in the UK (OpenSAFELY) reported higher risk of death from COVID-19 in patients with rheumatoid arthritis, lupus, or psoriasis [32]. Shortly after, comparative cohort studies
Table 1  International physician-reported registry studies examining associations between disease-modifying anti-rheumatic drugs (DMARDs) and severe COVID-19 outcomes in patients with autoimmune diseases

| Reference          | Population                              | Primary outcome | Medication comparator group | Higher odds of primary outcome | Lower odds of primary outcome | No effect on primary outcome |
|--------------------|-----------------------------------------|-----------------|------------------------------|-------------------------------|-------------------------------|------------------------------|
| Gianfrancesco et al. [6••] | Rheumatic disease patients with COVID-19 (n = 600) | Hospitalization | Non-users                    | Prednisone ≥10 mg daily       | Methotrexate monotherapy      | Antimalarials, cs DMARDs     |
| Strangfeld et al. [7••] | Rheumatic disease patients with COVID-19 (n = 3729) | Death           | Methotrexate monotherapy     | Prednisone ≥10 mg daily, rituximab, sulfasalazine, immunosuppressants (azathioprine, cyclophosphamide, cyclosporine, mycophenolate, or tacrolimus), no DMARD | None                          | Leflunomide, antimalarials, TNF inhibitors, abatacept, belimumab, IL-6 inhibitors, IL-17/IL-23/IL-12+23 inhibitors, JAK inhibitors |
| Brenner et al. [8••] | IBD patients with COVID-19 (n = 525) | Severe COVID-19 | Non-users                    | Prednisone, sulfasalazine/5-aminosalicylate | Thiopurine monotherapy; thiopurine, and TNF inhibitor combination therapy; mesalamine or sulfasalazine | None                          | TNF inhibitors³ |
| Ungaro et al. [9••] | IBD patients with COVID-19 (n = 1439) | Severe COVID-19 | TNF inhibitor monotherapy    | None                          | None                          | IL-12/23 antagonists; integrin inhibitors |
| Mahil et al. [10••] | Psoriasis patients with COVID-19 (n = 374) | Hospitalization | Biologics⁴                   | Non-biologics⁵,⁶              | N/A                           | No treatment with biologics or non-biologics |

COVID-19, coronavirus disease 2019; DMARD, disease-modifying anti-rheumatic drug; b/ts DMARD, biologic/targeted synthetic DMARD; cs DMARD, conventional synthetic DMARD; TNF, tumor necrosis factor; IL, interleukin; JAK, janus kinase; IBD, inflammatory bowel disease; vs., versus; N/A, not applicable

¹ Limitations of international physician-reported registry studies include selection bias (i.e., reporting of more severe cases), variable COVID-19 case definitions (i.e., molecular testing vs. suspected cases), unmeasured confounding, and regional variation in COVID-19 testing and treatment

² Severe COVID-19 was defined as intensive care unit admission, mechanical ventilation, or death

³ TNF inhibitors were associated with lower odds of hospitalization but not the primary outcome of severe COVID-19

⁴ Biologics included TNF inhibitors, IL-17 inhibitors, and IL-23/IL-12+23 inhibitors

⁵ Non-biologics included acitretin, apremilast, cyclosporine, methotrexate, fumaric acid esters/dimethylfumarate, and prednisolone

⁶ Accompanying patient-reported data showed less strict social distancing in patients on non-biologic vs. biologic DMARDs
| Reference                  | Location                | Population   | Intervention                                                                 | Primary outcome                                                                 | Key findings                                                                                                                                 |
|----------------------------|-------------------------|--------------|-----------------------------------------------------------------------------|--------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------|
| **Glucocorticoids**        |                         |              |                                                                             |                                                                                |                                                                                                                                             |
| Horby et al. (RECOVERY)    | UK                      | Hospitalized COVID-19 | Dexamethasone (n = 2104) vs. usual care (n = 4321) | 28-day mortality                                                                 | • Dexamethasone reduced 28-day mortality (RR 0.83, 95% CI: 0.74–0.92)  
• Most effective in patients requiring supplemental oxygen and/or intubation  
• Stopped early due to RECOVERY results  
• Fixed-dose and shock-dependent doses of hydrocortisone had 93% and 80% probabilities of superiority to usual care |
| Angus et al. (REMAP-CAP)   | North America, Europe, Australia | COVID-19 in ICU | Fixed-dose hydrocortisone (n = 143) vs. shock-dependent hydrocortisone (n = 152) vs. usual care (n = 108) | Days free of organ support |                                                                                                                                             |
| **Antimalarial therapy**   |                         |              |                                                                             |                                                                                |                                                                                                                                             |
| Self et al. (ORCHID)       | USA                     | Hospitalized COVID-19 | HCQ (n = 242) vs. placebo (n = 237) | Clinical status at 14 days on 7-category ordinal scale | • No significant difference in 14-day ordinal scale or 28-day mortality  
• No significant difference in 15-day ordinal scale  
• Prolonged QT interval and elevated liver enzymes with HCQ |
| Cavalcanti et al.          | Brazil                  | Hospitalized COVID-19 | HCQ (n = 221) vs. HCQ and azithromycin (n = 217) vs. usual care (n = 227) | Clinical status at 15 days on 7-category ordinal scale |                                                                                                                                             |
| Horby et al. (RECOVERY)    | UK                      | Hospitalized COVID-19 | HCQ (n = 1561) vs. usual care (n = 3155) | 28-day mortality | • Patients treated with HCQ were less likely to be alive at 28 days (RR 0.90, 95% CI: 0.83–0.98) and more likely to experience intubation or death (RR 1.14, 95% CI: 1.03–1.27) |
| **IL-6 receptor inhibitors** |                         |              |                                                                             |                                                                                |                                                                                                                                             |
| Stone et al. (BACC Bay)    | Boston, MA, USA         | Hospitalized COVID-19 | TCZ (n = 161) vs. placebo (n = 82) | Intubation or death | • No effect on primary outcome (HR 0.83, 95% CI: 0.38–1.81) or secondary outcomes  
• May have been underpowered due to lower-than-expected event rate  
• No difference in mechanical ventilation/death or clinical status at 14 days  
• Stopped early due to futility on interim analysis |
| Salvanari et al. (RCT-TCZ-COVID-19) | Italy      | Hospitalized COVID-19 | TCZ (n = 60) vs. placebo (n = 66) | Mechanical ventilation or death | • No improvement in clinical status at day 28  
• TCZ arm had shorter time to discharge (20.0 vs. 28.0 days, p = 0.04) and duration of ICU stay (9.8 vs. 15.5 days, p = 0.045) |
| Rosas et al. (COVACTA)     | North America, Europe   | Hospitalized COVID-19 | TCZ (n = 294) vs. placebo (n = 144) | Clinical status at 28 days on 7-category ordinal scale |                                                                                                                                             |
| Hemine et al. (CORIMUNO-TOC-I-1) | France             | Hospitalized COVID-19 | TCZ (n = 64) vs. usual care (n = 67) | Score >5 on WHO-CPS at day 4; intubation or death | • No effect on WHO-CPS scores  
• Trend towards lower risk of intubation or death at 14 days (HR 0.58, 95% posterior probability of efficacy) |
| Salama et al. (EMPACTA)    | North America, South America | Hospitalized COVID-19 | TCZ (n = 249) vs. placebo (n = 128) | Intubation or death | • TCZ arm significantly less likely to experience intubation or death at 28 days (HR 0.56, 95% CI: 0.33–0.97)  
• No improvement in death from any cause |

Days free of organ support
| Reference                      | Location                  | Population     | Intervention | Primary outcome | Key findings                                                                 |
|-------------------------------|---------------------------|----------------|--------------|----------------|------------------------------------------------------------------------------|
| Gordon et al. (REMAP-CAP)     | North America, Europe,   | COVID-19 in    | TCZ (n = 353) or sarilumab (n = 48) vs. usual care (n = 402) | Time to recovery | IL-6 receptor inhibition associated with greater number of organ support free days (10 days TCZ, 11 days sarilumab, 0 days usual care) |
|                               | Australia                 | ICU            |              |                | Odds of survival higher with IL-6 receptor inhibitor (OR 1.66 for TCZ, OR 2.25 for sarilumab, probability of superiority to usual care >99%) |
|                               |                           |                |              |                |                                                                              |
| Kalil et al. (ACTT-2)         | North America, Europe,   | Hospitalized   | Baricitinib (n = 515) vs. placebo (n = 518); both arms received remdesivir | Time to recovery | Baricitinib arm had shorter time to recovery (7.0 vs. 8.0 days, p = 0.03) and higher odds of improved clinical status (OR 1.3, 95% CI: 1.0–1.6) |
|                               | Asia                      | COVID-19       |              |                | Non-significant trend towards lower 28-day mortality (HR 0.65, 95% CI: 0.39–1.09) |
|                               |                           |                |              |                |                                                                              |
| Bureau et al. (CORIMUNO-AN-  | France                    | Hospitalized   | Anakinra (n = 59) vs. usual care (n = 55) | Score >5 on WHO-CPS at day 4; intubation or death | No improvement in WHO-CPS score at 4 days; No improvement in need for intubation or death at 14 days |
| A-1) [21•]                    | Hospitalized COVID-19     |                |              |                |                                                                              |
|                               |                           |                |              |                |                                                                              |
| Colchicine                    | Greece                    | Hospitalized   | Colchicine (n = 55) vs. usual care (n = 50) | Time to deterioration by 2 points on 7-point ordinal scale | Time to deterioration was longer in colchicine arm (20.7 vs. 18.6 days, p = 0.03) |
| Deftereos et al. (GRECCO-19)  | Hospitalized COVID-19     |                |              |                | Clinical deterioration less likely in colchicine arm (OR 0.11, 95% CI: 0.01–0.96) |
| [22•]                         |                           |                |              |                |                                                                              |
| Lopes et al. [23•]            | Brazil                    | Hospitalized   | Colchicine (n = 17) vs. placebo (n = 18) | Days oxygen needed; days hospitalization | Colchicine reduced days of oxygen needed (3.0 vs. 7.0, p = 0.02) and days of hospitalization (6.0 vs. 8.5, p = 0.03) |
| Tardif et al. [24•]           | Non-hospitalized COVID-19 |                | Colchicine (n = 2192) vs. placebo (n = 2189) | Hospitalization or death | No reduction in risk of primary outcome in full cohort (OR 0.79, 95% CI: 0.61–1.03) |
|                               |                           |                |              |                | Slight reduction in risk of primary outcome when limited to patients with PCR-confirmed COVID-19 (OR 0.75, 95% CI: 0.57–0.99) |

**COVID-19**, coronavirus disease 2019; **DMARD**, disease-modifying anti-rheumatic drug; **ICU**, intensive care unit; **MA**, Massachusetts; **USA**, United States of America; **vs.**, versus; **HCQ**, hydroxychloroquine; **TCZ**, tocilizumab; **JAK**, janus kinase; **IL**, interleukin; **WHO-CPS**, World Health Organization 10-point Clinical Progression Scale; **RR**, rate ratio; **CI**, confidence interval; **HR**, hazard ratio; **OR**, odds ratio; **PCR**, polymerase chain reaction
from Wuhan, China, and Boston, MA, reported higher odds of mechanical ventilation in patients with rheumatic diseases vs. comparators without rheumatic diseases [33, 34]. These studies led to heightened concerns that certain risk factors specific to rheumatic disease patients, such as DMARD use, may increase the risk of severe COVID-19 outcomes.

The COVID-19 Global Rheumatology Alliance (GRA) physician-reported registry examined odds of hospitalization among 600 rheumatic disease patients from 40 countries, comparing DMARD users to non-users [6••]. In multivariable models, prednisone doses ≥10 mg daily were associated with higher odds of hospitalization (OR 2.05, 95% CI: 1.06 to 3.96) [6••]. Use of conventional synthetic DMARDs, such as methotrexate, azathioprine, or leflunomide, was not associated with higher odds of hospitalization (OR 1.23, 95% CI: 0.70 to 2.17) [6••]. Interestingly, use of biologic/targeted synthetic DMARDs was associated with lower odds of hospitalization (OR 0.46, 95% CI: 0.22 to 0.93); this latter finding was largely driven by use of tumor necrosis factor (TNF)-alpha inhibitors. Mechanistic studies have suggested up-regulated TNF-alpha may lead to inflammatory cell death, aberrant germline center formation, and less robust humoral immune responses in fatal COVID-19, providing a potential biologic rationale for the protective effect of TNF inhibitors, although further study is needed [35, 36].

The GRA conducted a follow-up study of 3729 patients with rheumatic diseases and COVID-19, with notable differences from the first study including a new primary outcome (i.e., COVID-19-related death) and use of an active medication comparator (i.e., methotrexate monotherapy) [7••]. The follow-up study found that prednisone doses ≥10 mg daily were associated with higher odds of death (OR 1.69, 95% CI: 1.18 to 2.41), consistent with the first study [7••]. Rituximab (OR 4.04, 95% CI: 2.32 to 7.03), sulfasalazine (OR 3.60, 95% CI: 1.66 to 7.78), and other immunosuppressants (including azathioprine, cyclophosphamide, cyclosporine, mycophenolate, and tacrolimus; OR 2.22, 95% CI: 1.43 to 3.46) were associated with higher odds of death compared to methotrexate monotherapy, but there were no higher odds of death with other DMARDs [7••]. Interestingly, patients not on DMARDs also had higher odds of death compared to methotrexate monotherapy (OR 2.11, 95% CI: 1.48 to 3.01) [7••]. Surprisingly, the protective effect associated with TNF inhibitors in the initial study was no longer observed in this follow-up study. Many of the discrepancies between the first and second GRA studies may be related to the change in the comparator group from non-users to methotrexate monotherapy, so it is challenging to compare the two studies. The second GRA study also used a complex hierarchy of disease and medication groupings, which may limit interpretability and generalizability. However, the second GRA study raises important concerns regarding whether patients on rituximab, sulfasalazine, and certain immunosuppressants are at higher risk of poor outcomes.

The inflammatory bowel disease (IBD) community also established a physician-reported registry, called SECURE-IBD, which has been leveraged to examine severe COVID-19 outcomes in medication users vs. non-users [8••]. Among 525 patients from 33 countries, risk factors for severe COVID-19 included systemic corticosteroids (OR 6.9, 95% CI: 2.3 to 20.5) and sulfasalazine/5-aminosalicylate (OR 3.1, 95% CI: 1.3 to 7.7), similar to the GRA findings [8••]. Additionally, the SECURE-IBD registry found that TNF inhibitors were associated with lower risk of COVID-19 hospitalization (OR 0.60, 95% CI: 0.38 to 0.96), similar to the findings of the first GRA study, although there was no effect on the composite outcome of intensive care unit admission, mechanical ventilation, or death (OR 0.90, 95% CI: 0.37 to 2.17) [8••]. The SECURE-IBD registry conducted a follow-up study assessing the odds of severe COVID-19 using TNF inhibitor monotherapy as a comparator group [9••]. This study found higher odds of severe COVID-19 with thiopurine monotherapy (OR 4.08, 95% CI: 1.73 to 9.61), combination therapy with a TNF inhibitor and thiopurine (OR 4.01, 95% CI: 1.65 to 9.78), and mesalamine/sulfasalazine (OR 3.52, 95% CI: 1.93 to 6.45) compared to TNF inhibitor monotherapy, providing more granular detail that thiopurines and mesalamine/sulfasalazine may increase risk of poor outcomes [9••].

Lastly, the psoriasis community also established a physician-reported registry, which found that hospitalization was more frequent in patients using non-biologic DMARDs vs. biologic DMARDs (OR 2.84, 95% CI: 1.31 to 6.18), consistent with the protective effect of biologics observed in the first GRA and SECURE-IBD studies [10••]. However, independent patient-reported data from the psoriasis registry reported stricter social distancing among patients on biologic vs. non-biologic DMARDs, suggesting that adherence to social distancing may be a key mediator in this and other studies [10••]. Although the rheumatology, IBD, and psoriasis registry studies have provided rapid large-scale data on COVID-19 outcomes in patients on DMARDs, these studies are limited by unmeasured confounding and mediators (e.g., adherence to social distancing), selection bias (i.e., physician recall and reporting of more severe cases), variable COVID-19 case definitions, and regional variations in testing and treatment of COVID-19.

In total, evidence from physician-reported registries suggests that patients with autoimmune diseases on glucocorticoids are at higher risk of severe COVID-19 outcomes, while classes of DMARDs (conventional, biologic, and targeted synthetic) do not appear to be associated with higher risk. Some studies suggest that individual DMARDs, such as sulfasalazine, rituximab, and certain immunosuppressants, may be associated with severe COVID-19 outcomes and that TNF inhibitors may protect against severe COVID-19.
although further studies in non-registry data sets are needed to confirm these findings given discrepancies across registry studies. Importantly, in all the aforementioned studies, COVID-19 risk factors observed in the general population, such as older age and cardiovascular and pulmonary comorbidities, were associated with more severe COVID-19 outcomes; these risk factors may be clinically more important than DMARD use in identifying patients at high risk of severe COVID-19.

Based on the available evidence on DMARDs and COVID-19 to date and prior experiences with infections in rheumatic disease patients, the American College of Rheumatology (ACR) has released guidance regarding the management of rheumatic diseases during the ongoing COVID-19 pandemic [37]. For patients with a known SARS-CoV-2 exposure or confirmed COVID-19, the ACR recommends holding conventional synthetic DMARDs, non-IL-6 biologics, and janus kinase (JAK) inhibitors until 7–14 days after symptom resolution [37]. Hydroxychloroquine may be continued during COVID-19, and patients and providers can use shared decision-making regarding IL-6 receptor inhibitors [37]. Regardless of exposure or infection status, glucocorticoids should be kept at the lowest possible dose [37]. These guidelines will be updated as new data emerge [37].

**DMARDs as Therapies for COVID-19**

While the initial stage of COVID-19 is mediated by viral replication, the later stage observed in severe COVID-19 can be associated with a hyperinflammatory syndrome characterized by respiratory and other organ failure, hypercoagulability, and elevated markers of inflammation, including erythrocyte sedimentation rate, C-reactive protein, ferritin, IL-2, IL-6, TNF-alpha, and interferon-gamma (Fig. 1) [38, 39]. While antiviral therapies and monoclonal antibodies to SARS-CoV-2 may be the most effective in the initial viral replication phase, anti-inflammatory therapy may alleviate the hyperinflammation that mediates organ failure and death in severe COVID-19 [38]. Given the urgent need to develop treatments for COVID-19, several DMARDs were proposed as repurposed treatments for the hyperinflammatory phase of COVID-19. To date, several drugs commonly used in autoimmune diseases have been tested in RCTs with variable success.

**Glucocorticoids**

Glucocorticoids were the first anti-inflammatory medication shown to have a mortality benefit in severe COVID-19. In the RECOVERY trial, a randomized open-label adaptive platform trial, dexamethasone 6 mg (oral or intravenous) once daily for up to 10 days was compared to usual care in patients hospitalized with COVID-19 [11••]. Twenty-eight-day mortality was lower in patients receiving dexamethasone vs. usual care (age-adjusted RR 0.83, 95% CI: 0.74 to 0.92) [11••]. Importantly, dexamethasone reduced mortality in patients receiving supplemental oxygen by one-fifth and in patients requiring mechanical ventilation by one-third, but there was no mortality reduction in patients who did not require respiratory support [11••]. This finding suggests that glucocorticoids are effective in patients with severe COVID-19-related respiratory failure but may not have a benefit in patients with early or mild disease.

Further RCTs supported the findings from the RECOVERY trial. In the REMAP-CAP trial, a randomized open-label adaptive platform trial in 121 sites in 8 countries, patients were randomized to a fixed 7-day course of hydrocortisone (50 mg or 100 mg every 6 h), shock-dependent course (50 mg every 6 h when shock was evident), or no hydrocortisone [12]. Although the trial was stopped early due to the release of the RECOVERY results, the fixed-dose and shock-dependent doses of hydrocortisone had 93% and 80% probabilities of superiority for the primary endpoint (days free of cardiovascular and/or respiratory support) respectively compared to no hydrocortisone [12]. Furthermore, a meta-analysis of 7 RCTs (total n = 1703) showed reduced mortality in patients receiving corticosteroids vs. those receiving usual care or placebo (summary OR 0.66, 95% CI: 0.53 to 0.82) [40]. Consistent with this evidence, the Infectious Diseases Society of America (IDSA) guidelines recommend dexamethasone in patients with COVID-19 and hypoxemia and/or critical illness [41]. While RCTs overwhelmingly show glucocorticoids are effective in treating severe COVID-19, the registry studies of patients with autoimmune diseases suggested that glucocorticoid use at baseline was associated with higher risk of severe COVID-19. This may be due to confounding by indication in the registry studies, whereby patients with more severe autoimmune disease may be more likely to be on baseline glucocorticoids, and the more severe autoimmune disease may place patients at higher risk of severe COVID-19 outcomes rather than the use of glucocorticoids itself. Alternatively, the timing of glucocorticoid exposure may impact outcomes such that glucocorticoid use during the viral replication phase of COVID-19 may negatively impact the immune response.

**Antimalarial Therapy**

Early in vitro data and observational studies suggested that antimalarial drugs such as chloroquine and hydroxychloroquine may be effective against SARS-CoV-2 entry and replication, leading to widespread use, stockpiling, and drug shortages prior to any data from RCTs [3]. However, a meta-analysis of three cohort studies (pooled n = 932) showed no association between hydroxychloroquine use and
mortality, and a meta-analysis of two cohort studies (pooled \( n = 1549 \)) showed no association between hydroxychloroquine and a composite outcome of invasive mechanical ventilation and mortality [42].

Data from numerous single-center and multicenter RCTs have since confirmed that hydroxychloroquine does not have efficacy in COVID-19 as a prophylactic or therapeutic. RCTs of healthcare workers and others at high risk of SARS-CoV-2 exposure showed no efficacy for hydroxychloroquine for pre- or post-exposure prophylaxis [43, 44]. In patients with mild to moderate COVID-19, open-label RCTs of hydroxychloroquine vs. standard of care showed no effect of hydroxychloroquine on reducing viral load or duration until viral clearance [45–47]. A randomized, double-blind, placebo-controlled trial of hydroxychloroquine in symptomatic non-hospitalized adults with COVID-19 showed no difference in severity of symptoms at 14 days [48].

Several large RCTs examined hydroxychloroquine efficacy in hospitalized severe COVID-19. The National Institutes of Health (NIH)–sponsored ORCHID trial randomized hospitalized adults with severe COVID-19 to hydroxychloroquine (\( n = 242 \)) or placebo (\( n = 237 \)) [13••]. The trial was stopped early due to futility, and there was no improvement in the 7-category COVID-19 ordinal outcome scale at 14 days or mortality at 28 days [13••]. A multicenter RCT of hydroxychloroquine vs. hydroxychloroquine/azithromycin vs. standard of care in Brazil also showed no benefit in moderate/severe hospitalized COVID-19 [14••]. Additionally, both hydroxychloroquine regimens were associated with QT prolongation on electrocardiogram (which can trigger life-threatening arrhythmias) and elevated liver enzymes [14••].

Lastly, hydroxychloroquine was examined as a treatment arm in large adaptive platform trials. The World Health Organization SOLIDARITY trial reported no clinical benefit from the use of hydroxychloroquine for hospitalized patients with COVID-19 [49]. The aforementioned RECOVERY trial randomized 1561 hospitalized COVID-19 patients to hydroxychloroquine and 3155 to usual care [15••]. Patients treated with hydroxychloroquine were less likely to be alive at 28 days (RR 0.90, 95% CI: 0.83 to 0.98) and more likely to experience invasive mechanical ventilation or death (RR 1.14, 95% CI: 1.03 to 1.27) vs. those receiving usual care, suggesting possible harm from hydroxychloroquine.

There is now abundant evidence from numerous RCTs that hydroxychloroquine does not prevent COVID-19 infection or improve outcomes in mild, moderate, or severe COVID-19, contrary to early observational studies. Hydroxychloroquine has known side effects including QT prolongation, and one of the largest RCTs suggested hydroxychloroquine use may even be associated with lower survival. Therefore, hydroxychloroquine should not be used in the treatment of COVID-19 [41].

**IL-6 Receptor Inhibitors**

Initial observational studies from China reported promising results for IL-6 receptor inhibitors in the treatment of...
COVID-19 [50]. Additionally, a large cohort study of 3924 critically ill patients with COVID-19 in intensive care units (ICUs) in 68 hospitals across the United States (U.S.) showed that treatment with tocilizumab in the first 2 days of ICU admission was associated with a lower risk of death (HR 0.71, 95% CI: 0.56 to 0.92) [51].

Although early observational data were promising, early RCTs had negative or equivocal results. RCTs from the U.S. and Italy failed to reach primary and secondary endpoints [4•, 16]. A global multicenter RCT of tocilizumab vs. placebo (COVACTA) in patients hospitalized for severe COVID-19 also failed to reach its primary endpoint, improvement on a 7-category ordinal scale, although patients treated with tocilizumab had shorter times to discharge (20.0 vs. 28.0 days, \( p = 0.04 \)) and duration of ICU stay (9.8 vs. 15.5 days, \( p = 0.045 \)) [17]. An open-label RCT from France (CORIMUNO-TOCI 1) examined tocilizumab vs. usual care in patients with severe COVID-19 requiring ≥3 L/min of oxygen supplementation [18]. Although this trial showed no effect on the primary outcome (improvement in scores on a clinical progression scale), there was a trend towards lower risk of mechanical ventilation or death at 14 days in the tocilizumab group vs. standard care (median posterior HR 0.58, 95% posterior probability of achieving the predefined efficacy threshold) [18].

Although early trials of tocilizumab had equivocal or negative results, later trials with larger sample sizes and concomitant use of glucocorticoids in the majority (>80%) of patients have shown that tocilizumab may be a promising additive treatment for severe COVID-19. The EMPACTA trial, a placebo-controlled RCT of tocilizumab for patients with severe COVID-19, showed that tocilizumab-treated patients were significantly less likely to progress to mechanical ventilation or death (HR 0.56, 95% CI: 0.33 to 0.97), although there was no improvement in survival alone [5••]. The REMAP-CAP trial of tocilizumab or sarilumab vs. standard care in critically ill COVID-19 patients showed that IL-6 receptor inhibition was associated with significantly greater median organ support-free days and lower mortality compared to standard care [19••]. These two trials suggest that certain patients, especially those who are critically ill in the intensive care unit, may benefit from treatment with tocilizumab in conjunction with glucocorticoids, although the conflicting trial results limit clinical applicability. Currently, the IDSA guidelines recommend against the routine use of tocilizumab in hospitalized COVID-19, although this is a conditional recommendation based on low certainty of evidence and was last updated prior to the availability of the REMAP-CAP results [41].

**JAK Inhibitors**

JAK inhibitors may treat the hyperinflammatory phase of COVID-19 by inhibiting several cytokine pathways including IL-2, IL-6, IL-10, interferon-gamma, and granulocyte-macrophage colony-stimulating factor [20••, 52, 53]. In addition, among JAK inhibitors, baricitinib (a JAK1/JAK2 oral inhibitor) may block viral entry of SARS-CoV-2 by impairing AP2-associated protein kinase 1 and cyclin G–related kinases, which regulate ACE2 receptor–mediated viral endocytosis [53, 54]. A double-blind RCT (ACTT-2) of baricitinib vs. placebo on top of background antiviral therapy with remdesivir was conducted in 1033 patients hospitalized with COVID-19 [20••]. Patients treated with baricitinib had shorter time to recovery (7 vs. 8 days, \( p = 0.03 \)) and higher odds of clinical improvement on an 8-category ordinal scale (OR 1.3, 95% CI: 1.0 to 1.6), and there was a non-significant trend towards lower 28-day mortality (HR 0.65, 95% CI: 0.39 to 1.09) despite the trial being underpowered to detect mortality as an outcome [20••]. Notably, there was no higher incidence of thrombosis or secondary infections in the baricitinib-treated arm [20••].

Further studies are needed to evaluate whether baricitinib provides additive benefit to glucocorticoids in patients with severe COVID-19, especially given the cost difference and availability of baricitinib vs. dexamethasone. There are several ongoing phase 2 and 3 clinical trials of baricitinib (NCT04421027, NCT04321993, NCT04340232, NCT04393051), ruxolitinib (NCT04348071, NCT04377620, NCT04334044, NCT04362137), and tofacitinib (NCT04415151, NCT04469114) that may provide answers regarding whether JAK inhibitors provide additional benefits to glucocorticoids and, if so, which patients are most likely to benefit from combination therapy.

**IL-1 Inhibitors**

Prior to the COVID-19 pandemic, a secondary analysis of a RCT of anakinra vs. placebo showed that anakinra reduced 28-day mortality in patients with sepsis and features of macrophage activation syndrome; no significant safety signals were observed [55]. Given prior studies in sepsis, IL-1 inhibitors were considered promising DMARDs to repurpose for COVID-19. Additionally, transcriptomic profiling of whole blood from patients with COVID-19 showed elevated IL-1 concentrations prior to the nadir of respiratory function, unlike other pro-inflammatory cytokines, which peaked after respiratory deterioration [56]. Early case series and observational studies supported the hypothesis that IL-1 inhibitors may improve COVID-19 outcomes [57–60].

A French multicenter open-label RCT (CORIMUNO-ANA-1) of anakinra vs. usual care was conducted among patients hospitalized for COVID-19 requiring ≥3 L/min of oxygen support [21•]. The trial was stopped early at the recommendation of the data and safety monitoring board; anakinra was not found to improve clinical status at 4 days or need for mechanical ventilation or death at 14 days [21•].
However, this trial was conducted prior to the widespread use of remdesivir and dexamethasone, and further studies are needed to determine whether additive therapy with anakinra may improve outcomes. Ongoing phase 2 and 3 clinical trials of anakinra (NCT0443881, NCT04412291, NCT04362111, NCT04357366) and canakinumab (NCT04348448, NCT04362813) may provide further clarity on the role of IL-1 inhibition in severe COVID-19.

TNF Inhibitors

Anti-TNF therapy has been proposed as a potential treatment for the hyperinflammatory phase of severe COVID-19 because it is overexpressed in patients with severe COVID-19 and associated with poor humoral immune responses in fatal disease [35, 36, 61]. Additionally, patients on baseline TNF inhibitor therapy may have lower odds of severe COVID-19 outcomes in some registry studies, although this may be mediated by other factors such as adherence to social distancing, as previously reviewed [6••, 8•, 10••]. Anti-TNF regimens are currently being investigated in RCTs for COVID-19, including infliximab plus remdesivir (NCT04593940), infliximab (ISRCTN40580903), and adalimumab (ISRCTN33260034, ChiCTR2000030089).

Colchicine

Colchicine has been proposed as a treatment for COVID-19 hyperinflammation due to its ability to inhibit activation of the NLPR3 inflammasome and dampen neutrophil adhesion, migration, and signaling responses [62]. A cohort study from Italy reported that patients treated with colchicine had lower risk of death compared to those treated with usual care (HR 0.15, 95% CI: 0.06 to 0.37) [63]. However, this study used a historical comparator group, which may not be appropriate given the rapidly changing COVID-19 treatment guidelines over time. The Greek Effects of Colchicine in COVID-19 (GRECCO-19) open-label RCT evaluated colchicine vs. usual care in 105 hospitalized COVID-19 patients [22•]. The primary outcome of time to deterioration was longer in the colchicine arm than the usual care arm (20.7 vs. 18.6 days, p = 0.03), and clinical deterioration was less likely in the colchicine arm (OR 0.11, 95% CI: 0.01 to 0.96) [22•]. An interim analysis of a double-blind RCT of colchicine vs. placebo showed that colchicine reduced the duration of hospital stay (6.0 vs. 8.5 days, p = 0.03) and need for supplemental oxygen (53% vs. 83%, p = 0.01) in patients hospitalized with COVID-19 [23•].

Most recently, a double-blind RCT (COLCORONA) of colchicine (0.5 mg twice daily for 3 days followed by once daily for 27 days) vs. placebo was conducted in non-hospitalized patients with COVID-19 diagnosed by molecular testing or clinical presentation [24•]. While there was a trend suggesting that colchicine reduced the occurrence of the primary outcome, a composite of hospitalization or death (OR 0.79, 95% CI: 0.61 to 1.03), this did not achieve statistical significance [24•]. When restricted to patients with COVID-19 confirmed by molecular testing, there was a statistically significant decrease in the odds of the composite outcome (OR 0.75, 95% CI: 0.57 to 0.99), but no statistically significant decrease in the odds of mechanical ventilation (OR 0.50, 95% CI: 0.23 to 1.07) or death (OR 0.56, 95% CI: 0.19 to 1.66) [24•]. Importantly, the trial was terminated early by the investigators due to logistical issues, and therefore it may have been underpowered to detect effects, limiting our ability to apply these findings to non-hospitalized COVID-19 patients in clinical practice [24•].

Conclusions

In patients with autoimmune diseases, baseline glucocorticoid use is associated with severe COVID-19 outcomes. Given that dexamethasone reduces mortality in severe COVID-19, the finding in patients with autoimmune diseases may be related to the timing of glucocorticoid exposure or confounding by indication, as patients with more severe autoimmune disease are more likely to be on glucocorticoids. Overall, classes of DMARDs including biologic, targeted synthetic, and conventional synthetic DMARDs do not appear to be associated with significantly higher risk of severe COVID-19 outcomes in patients with autoimmune diseases. However, registry studies have raised concern regarding specific medications, such as sulfasalazine, rituximab, and certain immunosuppressants; further studies are needed with non-registry data to replicate these observations.

To date, several RCTs have examined glucocorticoids and DMARDs as treatments for COVID-19. Dexamethasone is associated with reduced mortality in severe COVID-19, especially in patients requiring oxygen supplementation and/or mechanical ventilation. Baricitinib and possibly IL-6 receptor inhibitors may improve outcomes in patients with severe COVID-19; however, further studies are needed to determine which patients are most likely to benefit from these medications as additive therapy to glucocorticoids and antivirals and to develop cost-effective strategies for COVID-19 treatment. Additional RCTs are needed to clarify the efficacy of other DMARDs, including IL-1 inhibitors, TNF inhibitors, and colchicine. There are no RCT data supporting the use of antimalarials for the prevention or treatment of COVID-19, and use of these medications for COVID-19 should be discouraged. The landscape of COVID-19 research is
rapidly evolving, and treatment recommendations will continue to change over time based on emerging evidence.

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**Declarations**

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