REVIEW

Immunopathology of Hyperinflammation in COVID-19

Joshua N. Gustine and Dennis Jones

From the Department of Pathology and Laboratory Medicine, Boston University School of Medicine, Boston, Massachusetts

The rapid spread of coronavirus disease 2019 (COVID-19), caused by severe acute respiratory syndrome—coronavirus 2 (SARS-CoV-2), has resulted in an unprecedented public health crisis worldwide. Recent studies indicate that a hyperinflammatory syndrome induced by SARS-CoV-2 contributes to disease severity and mortality in COVID-19. In this review, an overview of the pathophysiology underlying the hyperinflammatory syndrome in severe COVID-19 is provided. The current evidence suggests that the hyperinflammatory syndrome results from a dysregulated host innate immune response. The gross and microscopic pathologic findings as well as the alterations in the cytokine milieu, macrophages/monocytes, natural killer cells, T cells, and neutrophils in severe COVID-19 are summarized. The data highlighted include the potential therapeutic approaches undergoing investigation to modulate the immune response and abrogate lung injury in severe COVID-19. (Am J Pathol 2021, 191: 4e17; https://doi.org/10.1016/j.ajpath.2020.08.009)
marrow and reticuloendothelial organs. Collectively, this hyperinflammatory response shares many biological and clinical characteristics with the macrophage-activation syndrome seen in virus-induced hemophagocytic lymphohistiocytosis, suggesting a significant role of the host innate immune system in the immunopathology of COVID-19. Indeed, the efficacy of several immunomodulatory agents in attenuating the immune response to SARS-CoV-2 and in abrogating lung injury is being investigated. In this review, we discuss the inflammatory response in patients with COVID-19; in particular, we highlight the data that suggest that a dysregulated innate immune response drives the hyperinflammatory syndrome in severe COVID-19.

Initial Pathologic Findings

The pathologic examination of decedents from COVID-19 has provided important insights into the pathogenesis of the disease. The predominant pattern of lung injury associated with COVID-19 has been identified as diffuse alveolar damage accompanied by platelet–fibrin microthrombi in the pulmonary vessels. On postmortem analysis, gross examination has revealed heavy, congested, and diffusely edematous lung parenchyma, consistent with a clinical diagnosis of ARDS. Microscopy has identified diffuse alveolar hemorrhage in different phases. Most cases have been in the early or intermediate proliferative phase with edema, while features consistent with the fibrotic phase were rare. Histologic features accompanying the intra-alveolar and interstitial exudate have included capillary congestion, dilated alveolar ducts and collapsed alveoli, hyaline membrane formation, and desquamation of pneumocytes. Desquamated pneumocytes have been reported to have an apparent viral cytopathic effect and were present in alveolar spaces. Electron microscopy has identified viral particles within type 1 and type 2 pneumocytes, and immunofluorescence has localized SARS-CoV-2 antigen to the ACE2 bronchiolar epithelium.

It has been reported that in the majority of cases, microthrombi were present in the small- and medium-sized pulmonary arterial vessels. Increased levels of CD61 megakaryocytes were also found in alveolar capillaries accompanying the microthrombi. Microthrombi have been identified in nearly all major organs, including the lung, heart, brain, and liver; the location of microthrombi appears to correlate with endothelial ACE2 expression. These microthrombi likely contribute to organ dysfunction and mortality in COVID-19. Ackermann et al. reported on a study comparing postmortem findings of lung tissue from patients infected with COVID-19, influenza A (hemagglutinin type 1 and neuraminidase type 1: H1N1), and age-matched, uninfected controls. Alveolar capillary microthrombi were ninefold as prevalent in the lungs of COVID-19 patients compared to those in patients with influenza A. In addition, COVID-19 patients had increased numbers of ACE2 endothelial cells and significant histologic changes to endothelial cell morphology, including disruption of intercellular junctions, cell swelling, and loss of contact with basement membrane. Consistent with the findings from a previous study evaluating endothelial cells from glomerular capillary loops, electron microscopy identified SARS-CoV-2 viral particles within alveolar endothelial cells. In vitro studies have also shown that SARS-CoV-2 can directly infect an engineered human blood vessel organoid via the ACE2 receptor. These findings suggest that direct viral invasion by SARS-CoV-2 may trigger endothelitis and contribute to endothelial injury in COVID-19. Further work is needed for confirming these data as well as for elucidating the mechanism underlying the prothrombotic state in COVID-19.

In studies to date, all COVID-19 patients had a mononuclear inflammatory infiltrate in the lung parenchyma, composed primarily of lymphocytes and macrophages. The lymphocytic infiltrate included both CD4 and CD8 T cells in the bronchiolar and alveolar interstitium, with a CD4 T-cell predominance. CD4 T cells appeared to aggregate around small vessels that often contained microthrombi. Large numbers of CD68 macrophages and multinucleated cells were also identified and localized to the alveolar lumen. The presence of macrophages appeared to correlate with disease advancement. These patient findings have been recapitulated in phenotypic studies using transgenic mice bearing human ACE2 infected with SARS-CoV-2, which showed the characteristic interstitial pneumonia with infiltration of lymphocytes and macrophages in the alveolar interstitium, as well as macrophages in the alveolar lumen.

A recent study identified the presence of hemophagocytosis in the bone marrow and reticuloendothelial organs of patients with COVID-19. Bryce et al. reported a comprehensive autopsy series of 67 patients who succumbed to COVID-19. In that study, hemophagocytic macrophages were identified in the lymph nodes (9/11), spleen (9/22), bone marrow (4/6), heart, and liver. Abnormally high mean peak levels of C-reactive protein, ferritin, IL-6, IL-8, and tumor necrosis factor (TNF)-α accompanied the histopathologic findings in these patients. The presence of hemophagocytosis with concomitant elevations in inflammatory cytokines suggests that alveolar macrophage activation can induce a hemophagocytic lymphohistiocytosis phenotype in patients with severe COVID-19. Severe cases of SARS-CoV-1 infection shared similar pathologic findings.

Cytokine Storm

Early studies reporting outcomes in COVID-19 identified that elevated clinical inflammatory markers were prognostic of disease severity and mortality. Two multicenter, retrospective studies in hospitalized patients in China evaluated predictors of mortality in COVID-19. Non-survivors had elevated C-reactive protein, lactate
dehydrogenase, serum ferritin, and serum IL-6 levels at the
time of hospital admission compared to those in surviv-
ors. In addition, nonsurvivors had elevated inflam-
mmatory markers throughout the entire clinical course, and the
clinical deterioration that preceded death tracked with
increasing levels of inflammation. The prevalence of
ARDS was also higher in the group with elevated inflam-
matory markers.

Consistent with these initial clinical findings, plasma
sampling from patients with severe COVID-19 revealed a
proinflammatory cytokine profile. The initial report included
41 patients admitted to the hospital with COVID-19-related
pneumonia in Wuhan, China. Compared to healthy adults,
patients with COVID-19 in that study had higher plasma
concentrations of IL-1β, IL-1Rz, IL-7, IL-8, IL-10, basic
fibroblast growth factor, granulocyte colony-stimulating
factor (G-CSF), granulocyte macrophage (GM-CSF), interferon
(IFN)-γ, induced protein (IP)-10/CXCL10, monocyte
chemotactic protein (MCP)-1/C-C motif chemokine ligand
(CCL)-2, macrophage inflammatory protein (MIP)-1α/CCL3,
MIP-1β/CCL4, platelet-derived growth factor, TNF-α, and
vascular endothelial growth factor. The mean plasma con-
centrations of IL-2, IL-7, IL-10, G-CSF, IP-10/CXCL10,
MCP-1/CCL2, MIP-1α/CCL3, and TNF-α were higher in the
subgroup in the intensive care unit versus those in
non-intensive care unit patients. A study in 21 patients with
COVID-19 admitted with pneumonia reported similar find-
ings; patients with moderate or severe disease had abnor-
mally elevated levels of IL-1β, IL-2 receptor (R), IL-6, IL-8,
IL-10, and TNF-α. In a large-scale, retrospective study in
1484 patients at Mount Sinai Hospital (New York, NY),
plasma levels of IL-6, IL-8, and TNF-α were higher in those
with severe COVID-19. Moreover, IL-6 and TNF-α levels
at the time of hospitalization were independently associ-
ated with disease severity and mortality in that multivariate
analysis. Elevated plasma levels of IL-6, IL-8, and TNF-α also
peaked before death in an autopsy series of COVID-19
patients, and correlated with pathologic evidence of hemo-
phagocytosis. These data suggest that a cytokine storm
underpins the immunopathology of severe COVID-19.

TY et al proposed a possible mechanism underlying the
cytokine storm in severe COVID-19. SARS-CoV-2 is a
cytopathic virus that induces the death of infected cells during
viral replication. Viral replication in epithelial cells may
also cause high levels of pyroptosis, which is an inflamma-
tory form of programmed cell death observed in infection with
cytopathic viruses. Pyroptosis could represent an inciting
event for the hyperinflammatory response to SARS-CoV-2
infection. IL-1β is released from cells undergoing
pyroptosis, and high levels are present in both serum and
bronchoalveolar fluid (BALF) in patients with severe COVID-
19. Pathogen-associated molecular patterns, such as viral
RNA, are also released by infected epithelial cells. Neigh-
boring lung epithelial cells and resident alveolar macrophages
detect the pathogen-associated molecular patterns using
dxample pattern-recognition receptors and IL-1R stimulates the secre-
tion of proinflammatory cytokines, including IL-6, IFN-γ,
MCP-1/CCL2, MIP-1α/CCL3, MIP-1β/CCL4, and IP-10/
CXCL3. The cytokines then recruit activated macrophages
and T cells to the site of infection, whereby these immune
effector cells potentiate inflammation with additional cytokine
secretion and destruction of lung parenchyma. Similar to
SARS-CoV-1 and MERS-CoV, the cytokine profile observed
in patients with COVID-19 is consistent with a predominant
Th1 cell— and classic M1 macrophage—polarized
response. Consequently, a proinflammatory feedback
loop is established, triggering a cytokine storm that circulates
and causes local tissue damage as well as systemic effects, such
as septic shock, multiorgan failure, and hemophagocytosis in
reticuloendothelial organs.

Notably, the tropism of SARS-CoV-2 for ACE2+ type 2
pneumocytes may contribute to the development of cytokine
storm. Viral binding by SARS-CoV-2 to the ACE2 receptor is
primarily restricted to type 2 pneumocytes on the lung
epithelium. Although typically associated with surfactant
production and alveolar repair, type 2 pneumocytes have a
specialized role in the innate immune response. Type 2
pneumocytes express toll-like receptors (TLRs) that activate
inflammatory NF-κB signaling in response to binding of viral
RNA. The activation of NF-κB triggers the production of
cytokines that can induce an inflammatory program in resident
macrophages and recruit activated monocytes and T cells to the
lung. Accordingly, increased IL-6 levels in hyperplastic type 2
pneumocytes infected with SARS-CoV-2 have been
measured. ACE2+ infected pneumocytes also have been
reported to express high levels of proinflammatory cytokines
(i.e., MCP-1/CCL2, TNF-α, IL-1β, and IL-6) in an autopsy
series of patients with SARS-CoV-1 infection; cytokine
expression was unchanged in uninfected cells. Direct inva-
sion and stimulation of TLR-containing ACE2+ type 2
pneumocytes may therefore exacerbate cytokine production in
COVID-19.

Dysregulated IFN signaling also likely has an impact on the
immunopathology of severe COVID-19. The findings
from initial studies have suggested that SARS-CoV-2 can
suppress IFN signaling and impair viral clearance from
infected cells. SARS-CoV-2 was reported to be sensitive to
pretreatment with types I and III IFNs. However, a study using
infected cell lines, primary bronchiolar cells, and a ferret model with SARS-CoV-2 infection demonstrated
decreased type I and III IFN signatures. Patients with
severe COVID-19 also appeared to have an impaired type I
IFN signature compared to that in mild or moderate cases.
Whole-blood transcriptome analysis revealed that the
expression of type I IFN was inversely correlated with the
viral load and NF-κB–driven inflammatory response (i.e.,
IL-6 and TNF-α levels) in COVID-19 patients. Moreover,
a multicenter observational study reported that the admin-
istration of IFN-α2b early in the disease course of COVID-
19 improved in-hospital mortality in infected patients.
These findings suggest that a weak IFN response during
early infection with SARS-CoV-2 may allow for the development of severe COVID-19.

Additional work is needed for ascertaining the mechanism that SARS-CoV-2 employs to impair IFN signaling. However, the findings from functional studies in other pathogenic coronaviruses, such as SARS-CoV-1 and MERS-CoV, have elucidated the viral proteins that antagonize IFN signaling and release. Example virulence mechanisms include the inhibition of TNF receptor–associated factor family member–associated NF-κB activator–binding kinase (TBK)-1–dependent phosphorylation, IFN regulatory factor 3 activation, and IFN production.\(^{32–44}\) A murine model of SARS-CoV-1 infection also showed that delayed type I IFN signaling and rapid viral replication induced a hyperinflammatory state with lung pathology.\(^{45}\) These mice had an excessive accumulation of activated macrophages in the lung, as well as elevated cytokine levels and an inadequate virus-specific T-cell response.\(^{45}\) As such, impaired viral clearance of SARS-CoV-2 due to the antagonism of IFN signaling might enable continuous TLR stimulation and pyroptosis of infected type 2 pneumocytes, thereby promoting the hyperinflammatory state observed in severe COVID-19.

### Dysregulated Macrophage Activation

A proinflammatory macrophage microenvironment defines the landscape of bronchoalveolar immune cells in patients with severe COVID-19. In line with initial pathologic findings, single-cell RNA sequencing (scRNA-seq) of BALF collected from COVID-19 patients identified an abundance of macrophages in severe disease.\(^{31,46}\) Further analysis revealed that the macrophages are primarily inflammatory monocyte derived, with a relative paucity of resident alveolar macrophages.\(^{31}\) The macrophages in severe disease highly express the genes ficolin-1 (FCNI) and SPP1, while in moderate disease they preferentially express FABP4.\(^{31}\) Consistent with the cytokine pattern in peripheral blood,\(^{4}\) macrophages expressing FCNI and SPP1 in BALF have gene-expression signatures characteristic of classic M1 macrophages.\(^{31}\) These macrophages express genes that encode peripheral monocyte-like markers (ie, S100A8, FCN1, and CD14), chemokines (ie, MCP1/CCL2, MIP1A/CCL3, and INP10/CXCL10), and inflammatory transcription factors (ie, NFκB, STAT1, STAT2, and IFN regulatory factors).

A proinflammmatory subset of alternative M2 macrophages that express both profibrotic genes (ie, TREM2, TGFBI1, and SPP1) and immunoregulatory genes (ie, A2M and GPR3) was also identified in patients with severe COVID-19.\(^{31}\) These findings suggest that the pathogenic role of macrophages in severe COVID-19 may extend beyond acute inflammation to include pulmonary fibrosis. Early reports indicate fibrotic lung patterns with reduced diffusion and total lung capacity in survivors of severe COVID-19, as well as persistent ground-glass opacities on imaging at the time of hospital discharge.\(^{47,48}\)

Gene expression of cytokines and chemokines has been reported to be markedly increased in lung macrophages. Based on scRNA-seq data, lung macrophages from patients with severe COVID-19 had increased expression of IL1B, IL6, TNF, and genes encoding several chemokines, including MCP1/CCL2, MIP1A/CCL3, MIP1B/CCL4, and MCP3/CCL7.\(^{31,46,49}\) These chemokines are potent recruiters of inflammatory monocytes to the site of infection. Conversely, patients with moderate disease have been reported to express higher levels of CXCL16, which can bind to the CXCR6 receptor and attract CD8\(^+\) T cells that may be specific for SARS-CoV-2.\(^{31}\) In both severe and moderate cases, the gene expression levels of the CXCR3 ligands CXCL9, CXCL10, and CXCL11 were higher compared to those in healthy controls.\(^{31}\) These transcriptional changes in lung macrophages also correlated with BALF cytokine levels. Patients with severe disease had significantly higher levels of proinflammatory cytokines in BALF, especially IL-1β, IL-6, IL-8, TNF-α, and IFN-γ.\(^{31,45}\) Collectively, these data suggest that lung macrophages recruit inflammatory monocytes and produce cytokines that contribute to hyperinflammation in severe COVID-19.

A significant expansion of inflammatory monocytes is also present in the peripheral blood of COVID-19 patients. Flow cytometric analysis have been used for identifying increased populations of CD14\(^+\)CD16\(^-\), GM-CSF/CD14\(^+\), and IL-6/CD14\(^-\) subsets of inflammatory monocytes.\(^{50,51}\) The percentage of CD14\(^+\)CD16\(^-\) monocytes producing IL-6 was correlated with disease severity.\(^{50}\) Data from scRNA-seq of peripheral blood mononuclear cells demonstrated similar findings, wherein CD14\(^+\)IL-1β\(^+\) monocytes, IFN-activated monocytes, and IL1B-associated inflammasome signatures were observed.\(^{52–54}\) Akin to lung macrophages, inflammatory monocytes in the peripheral blood showed an enrichment of genes encoding cytokine signaling and inflammation characteristic of classic M1 macrophages.\(^{55}\) The scRNA-seq analysis additionally suggested that cytokine activation drives the systemic expansion of monocyte populations in the peripheral blood of COVID-19 patients.\(^{52,54}\) Hence, the cytokine environment likely generated by type 2 pneumocytes and resident alveolar macrophages after SARS-CoV-2 infection triggered the expansion and recruitment of inflammatory monocytes to the lung.

Pathologic stimulation of TLR/IL-1R signaling represents a significant mechanism activating inflammatory monocytes and macrophages in COVID-19. It has been shown that after TLR/IL-1R stimulation, Bruton tyrosine kinase (BTK) drives the signaling cascade that activates both NF-κB and nucleotide-binding oligomerization domain-containing protein–like receptor protein (NLRP)-3 inflammasome secretion of IL-1β.\(^{56–57}\) Accordingly, recent data showed increased phosphorylation of BTK in IL-6/CD14\(^+\) monocytes derived from patients with severe COVID-19.\(^{51}\) The entire population of monocytes had phosphorylated BTK,\(^{31}\) consistent with the notion of systemic monocytic activation. Transcriptomic profiling of whole blood and single monocytes revealed similar
findings. Patients with severe COVID-19 had increased expression of TLR and IL1R and downstream signaling molecules (ie, MYD88, IRAK4, IRAK1, TRAF6, and RELA/p65).\textsuperscript{52,53} The expression of these signaling molecules also tracked with disease severity in one patient with longitudinal samples.\textsuperscript{53} Myeloid differentiation primary response protein 8 (MyD88) is an adaptor protein that triggers BTK activation and mediates signaling for IL-1, IL-18, and all TLRs except TLR3.\textsuperscript{58} MyD88\textsuperscript{WT/WT} mice infected with SARS-CoV-1 exhibited decreased lung pathology and cytokine levels as well as impaired monocyte recruitment to the lung versus MyD88\textsuperscript{WT/KO} mice.\textsuperscript{59} Likewise, in a small cohort of severe COVID-19 patients, treatment with a BTK inhibitor normalized C-reactive protein and IL-6 levels and improved oxygenation.\textsuperscript{51} The findings from these studies indicate that targeting the TLR/IL-1R signaling cascade may attenuate the hyperinflammatory response in severe COVID-19.

**Impaired Natural Killer Cell Response**

Studies have shown that peripheral natural killer (NK) cells are depleted and exhibit an exhausted phenotype in patients with severe COVID-19. Peripheral NK cell counts were significantly lower in COVID-19 patients with severe disease versus those in patients with mild disease or in healthy controls.\textsuperscript{52,60–63} Whereas antiviral cytotoxic CD56\textsuperscript{dim} NK cells were primarily depleted in ventilator-dependent patients, all COVID-19 patients had decreased cytokine-producing CD56\textsuperscript{high} NK cells peripherally.\textsuperscript{60} The uniform reduction of CD56\textsuperscript{high} NK cells suggests that peripheral NK cells do not directly contribute cytokines toward the cytokine storm in COVID-19. The percentage of peripheral NK cells expressing the activation markers CD16, CD107a, IFN-γ, IL-2, and TNF-α was also significantly lower in COVID-19 patients compared to healthy controls, as was the mean fluorescence intensity of cytotoxic granzyme B.\textsuperscript{61,62} scRNA-seq revealed evidence of exhausted NK cells with increased transcription of the inhibitory receptor genes LAG3, PDCD1, and HAVCR2.\textsuperscript{60} Both NK and CD8\textsuperscript{+} T cells also had increased expression of the immune checkpoint natural killer group 2 member A (NKG2A), which inhibits cell cytotoxicity.\textsuperscript{61} The successful treatment of COVID-19 is associated with the reversal of functional NK cell exhaustion, as evidenced by the down-regulation of NKG2A in the convalescent period.\textsuperscript{51}

It remains unclear whether the peripheral NK cell depletion is due to the trafficking of cells to infected lung tissue or cell death. A study employing bulk RNA-seq analyzed the immune composition of BALF from COVID-19 patients collected a median of 8 days after symptom onset.\textsuperscript{46} The results demonstrated no change in the proportion of activated NK cells, while COVID-19 patients had a decreased proportion of resting NK cells in BALF versus healthy controls.\textsuperscript{46} Conversely, a study using scRNA-seq of BALF collected a median of 12 days after symptom onset reported increased NK cells in COVID-19 patients.\textsuperscript{31} The discrepant findings between studies could have been attributed to the timing differences in BALF sample collection, and suggest that NK cells may be recruited to the lung later in the disease course of COVID-19.\textsuperscript{64} Trafficking of NK cells is further supported by the expansion of lung macrophages that produce CXCR3 ligands (CXCL9, CXCL10, and CXCL11) in COVID-19.\textsuperscript{31} CXCR3 has been previously shown to promote NK cell recruitment to the lung in influenza infection.\textsuperscript{65} It is possible that cell death may also contribute to peripheral NK-cell depletion, given that the transcription of TP53 and apoptosis pathways in peripheral lymphocytes are up-regulated in COVID-19 patients.\textsuperscript{69} While the absence of ACE2 receptors makes SARS-CoV-2 infection of NK cells unlikely,\textsuperscript{34} circulating cytokines could play a role, given that peripheral NK-cell counts are inversely correlated with serum IL-6 levels.\textsuperscript{52} Additional studies are nonetheless needed for delineating the processes involved in peripheral NK-cell depletion, including whether the cytokine storm leads to NK-cell apoptosis.

The cytokine milieu also may confer the exhausted phenotype of peripheral NK cells in COVID-19. In particular, studies have reported that IL-6 and TNF-α impaired the cytolytic function of NK cells.\textsuperscript{54,61,62,66–68} In healthy donor NK cells, in vitro stimulation of the soluble IL-6R by IL-6 reduced the production of perforin and granzyme, which can be reversed by IL-6R antagonism with tocilizumab.\textsuperscript{66} These findings suggest that IL-6 might hasten the down-regulation of granzyme expression on NK cells reported in COVID-19.\textsuperscript{61} However, additional experiments are needed for confirming these findings in NK cells derived from COVID-19 patients. The decreased expression of CD16 on NK cells indicates that reduced NK-cell licensing is also likely occurring; CD16 expression was reported to be inversely correlated with serum IL-6 levels in COVID-19 patients.\textsuperscript{62} In addition, ligand-receptor analysis with scRNA-seq of peripheral blood revealed interactions between inflammatory monocytes and NK cells in severe COVID-19.\textsuperscript{54} The data showed that TNF-α may interact with its receptor, TNFR1, on NK cells.\textsuperscript{54} The impact of this cytokine–receptor interaction in COVID-19 needs investigation, but in the tumor microenvironment it results in the down-regulation of the major NK cell–activating receptor Nkp46, especially on CD56\textsuperscript{high} NK cells.\textsuperscript{57} Taken together, these findings suggest that crosstalk with monocytes and macrophages via cytokines might mediate NK-cell dysfunction and impair the clearance of SARS-CoV-2.\textsuperscript{68} Persistent infection with SARS-CoV-2 could subsequently exacerbate the proinflammatory feedback loop underlying the cytokine storm in severe COVID-19.

**Role of T Cells in Hyperinflammation**

The occurrence of a profound lymphopenia in patients with COVID-19 is well established. The absolute lymphocyte count is prognostic of disease severity and mortality, with significant reductions in peripheral CD4\textsuperscript{+} and CD8\textsuperscript{+} T cells reported in both moderate and severe cases of
COVID-19. Serum cytokine levels are inversely correlated with CD4+ and CD8+ T-cell counts, and improved cytokine levels have been associated with the reversal of lymphopenia in the convalescent period of recovered patients. In decedents who had lymphopenia, autopsy evaluations have revealed the extensive cell death of lymphocytes in the lymph nodes and spleen; SARS-CoV-2—infected spleens and lymph nodes also were found to have overexpressed the apoptotic protein Fas. Likewise, transcriptomic analysis identified up-regulation of apoptotic pathways in lymphocytes from the peripheral blood of COVID-19 patients. Hemophagocytic macrophages present in the spleen and lymph nodes accompanied the lymphocyte depletion, suggesting a role for aberrantly activated macrophages in lymphopenia. These macrophages have been found to express high levels of IL-6, which may promote lymphocyte necrosis and apoptosis. Indeed, antagonism of the IL-6R with tocilizumab resulted in increased peripheral lymphocyte counts in COVID-19. Trafficking of peripheral T cells to the lung also likely contributes to the lymphopenia. As previously discussed, autopsy studies have reported a pulmonary lymphocytic interstitial infiltrate with CD4+ and CD8+ T cells, and scRNA-seq of BALF identified a heterogeneous expansion of CD8+ T cells. Additional studies are needed for clarifying the mechanism and functional impact of lymphopenia, such as on the clearance of SARS-CoV-2.

The T-cell composition of surviving lymphocytes may exacerbate the hyperinflammatory state in COVID-19. In conjunction with IL-6– and GM-CSF—producing monocytes, patients with severe COVID-19 have been found to have increased proportions of pathogenic Th1 CD4+ T cells that express IL-6, GM-CSF, and IFN-γ. In particular, patients requiring intensive care unit admission had significantly higher proportions of Th1 cells co-expressing GM-CSF and IFN-γ. These Th1 cells can produce proinflammatory cytokines that contribute to the cytokine storm and migrate to the lung to potentiate tissue damage. Zheng et al postulated that GM-CSF links the activation of inflammatory monocytes and Th1 cells in severe cases. Moreover, flow cytometry has revealed decreased levels of immunosuppressive T-regulatory cells in patients with severe COVID-19. scRNA-seq of peripheral blood similarly identified depleted T-regulatory cells, as well as up-regulated inflammatory and cytokine pathways in CD4+ T cells. These findings collectively suggest that the reduction in T-regulatory cells disrupts immune homeostasis and promotes a central role for CD4+ Th1 cells in the pathogenesis of COVID-19—related hyperinflammation.

**Neutrophils and Neutrophil Extracellular Traps**

Increasing evidence indicates that the dysregulated myeloid response to SARS-CoV-2 extends to neutrophils in severe COVID-19. An elevated absolute neutrophil count and neutrophil/lymphocyte ratio have been reported to be prognostic of ARDS and death in COVID-19. Patients with severe COVID-19 also had increased levels of neutrophil extracellular traps (NETs), which are webs of DNA material with antimicrobials and oxidant enzymes extruded by neutrophils to control infections. Serum samples collected from patients hospitalized with COVID-19 had higher levels of cell-free DNA, myeloperoxidase (MPO)–DNA complexes, and citrullinated histone H3 (Cit-H3) compared to those from patients with mild/moderate disease and healthy controls; the latter two markers are considered specific for NET remnants. Elevated plasma MPO–DNA complexes have been associated with a decreased PaO2/FiO2 ratio, a need for mechanical ventilation, and death in COVID-19. Increasing levels of MPO–DNA complexes also tracked with oxygen deterioration in patients with longitudinal samples. The milieu of patients with COVID-19 appears to promote the formation of NETs, as the exposure of healthy neutrophils to serum collected from hospitalized COVID-19 patients has been reported to trigger significant NETosis. Further work is needed for delineating the mechanism of NET formation in COVID-19, but the activation of neutrophils by viral nucleic acids and cytokines might have a role.

Importantly, NETs may contribute to the development of lung injury and microthrombi in COVID-19. Dysregulated NETosis has been previously shown to cause excessive lung injury and immunothrombosis in response to other respiratory viruses and bacteria. A recent study that evaluated autopsy lung specimens for NETs reported pathologic evidence suggestive of dysregulated NETosis in COVID-19. Immunofluorescence revealed a neutrophilic infiltrate in the lungs, with numerous Cit-H3 and MPO neutrophils as well as lattices of extracellular DNA with Cit-H3 and MPO. In addition, the study reported the co-localization of Cit-H3+ neutrophils likely undergoing NETosis with platelet factor 4–positive platelets in pulmonary blood vessels, suggesting a possible role of NET and platelet interaction in microthrombi formation in COVID-19. Whether excessive immunothrombosis by NETs triggers platelet aggregation and the prothrombotic state in COVID-19 warrants investigation.

**Multisystem Inflammatory Syndrome in Children**

Recent case series of children have identified a hyperinflammatory vasculopathy associated with COVID-19, deemed the *multisystem inflammatory syndrome in children* (MIS-C). This syndrome has been reported to have a wide range of clinical presentations that overlap with Kawasaki disease, toxic shock syndrome, and macrophage-activation syndrome/hemophagocytic lymphohistiocytosis. Nearly all patients with MIS-C have presented with fever. Multiple organ systems have been reported to be
involved in most patients, with the gastrointestinal (92%), cardiovascular (80%), hematologic (76%), mucocutaneous (74%), and respiratory (70%) systems most commonly involved. The majority of patients have required admission to the intensive care unit for treatment. Invasive and noninvasive mechanical ventilation for respiratory insufficiency or failure has been needed in 20% and 17% of patients, respectively. The mortality rate from MIS-C has been estimated as approximately 0% to 2%. An important finding in MIS-C has been the high incidence of cardiovascular involvement. In reports from the United States and Europe, the majority of patients had elevated brain natriuretic peptide and troponin levels. Nearly half of the patients had hypotension or cardiogenic shock, which prompted vasopressor support in one-quarter of patients. On echocardiography, ventricular dysfunction (ejection fraction <55%) was identified in approximately half of patients. Approximately 8% to 21% of patients also had coronary artery aneurysm identified in either the left descending artery or the right coronary artery, consistent with a Kawasaki-like vasculitis presentation of MIS-C.

The immunologic findings in MIS-C have been similar to those described in adults with severe COVID-19. Nearly all MIS-C patients had multiple laboratory markers of inflammation, including an elevated erythrocyte sedimentation rate, C-reactive protein, and ferritin. Serum cytokine analysis showed elevated IL-6, IL-10, and soluble IL-2R levels. Patients with MIS-C also displayed lymphopenia, including reduced levels of CD4+ and CD8+ T cells as well as NK cells. Notably, the majority of MIS-C patients had positive SARS-CoV-2 serologies, and the median time interval elapsed between COVID-19 symptom onset and MIS-C symptom onset was 25 days. These findings support the current hypothesis that MIS-C represents an immune-mediated SARS-CoV-2 postinfectious process. Although MIS-C is a rare complication of COVID-19, additional studies are needed for understanding the pathophysiology, therapeutic management, and long-term sequelae of this complication.

Therapeutic Targeting of Hyperinflammation

The evidence of a dysregulated host immune response has prompted the investigation of immunomodulatory therapies for abrogating lung injury in COVID-19. There are >2500 studies of COVID-19 currently registered with clinicaltrials.gov, including many evaluating immunomodulators alone or in combination with other agents. These clinical trials will need to determine the optimal disease stage of COVID-19 at which immunomodulators are most effective and safe in patients with hyperinflammation. The published clinical trials of immunomodulators in COVID-19 are reviewed in the subsequent paragraph and the developmental stage of each clinical approach as well as ongoing randomized studies are summarized in Table 1.

Corticosteroids have widespread inhibitory effects on the immune system and are efficacious in managing acute inflammatory processes. The randomized, controlled, open-label RECOVERY trial (Dexamethasone in Hospitalized Patients with Covid-19—preliminary report) evaluated the efficacy of the corticosteroid dexamethasone (n = 2104) versus usual care (n = 4321) in hospitalized COVID-19 patients. Dexamethasone 6 mg once daily was given for up to 10 days. The study demonstrated a significantly lower 28-day mortality in patients randomized to receive dexamethasone versus usual care (22.9% versus 25.7%; P < 0.001). In a prespecified subgroup analysis, dexamethasone reduced 28-day mortality by 36% in patients on mechanical ventilation (HR = 0.64; 95% CI, 0.51 to 0.81) and by 18% in patients receiving oxygen without mechanical ventilation (HR = 0.82; 95% CI, 0.72 to 0.94), whereas dexamethasone had no benefit in patients receiving no respiratory support (HR = 1.19; 95% CI, 0.91 to 1.55). Dexamethasone was also more effective in patients randomized to receive treatment for >7 versus ≤7 days after the onset of COVID-19 symptoms (P < 0.001). Dexamethasone was the first therapy reported to improve survival in patients with COVID-19.

The identification of respiratory support as a predictor of dexamethasone efficacy has provided insights into the disease stages of COVID-19. Previous studies have shown that inflammatory markers track with disease progression and clinical deterioration in COVID-19. Consistent with these findings, the benefit of immune suppression with dexamethasone was apparent in only patients requiring respiratory support, as well as those randomized to receive treatment for >7 days after symptom onset. These data suggest that the patients were in a later disease stage dominated by immunopathology. Conversely, patients not requiring respiratory support, who were likely without significant inflammation, had no benefit, and a trend of increasing death rate from dexamethasone treatment. It is possible that dexamethasone may have suppressed the viral clearance of SARS-CoV-2 in these patients. The majority of patients without respiratory support had symptoms for <1 week at the time of randomization, which corresponds to the timing of peak viral replication of SARS-CoV-2. The use of systemic corticosteroids have been shown to impair viral clearance in patients with SARS-MERS or SARS-CoV-1 infection. Longitudinal evaluation of inflammatory markers and viral loads with dexamethasone use will further clarify these findings in COVID-19.

The cytokine storm in patients with severe COVID-19 has prompted the evaluation of selective cytokine inhibitors. Tocilizumab, a recombinant humanized monoclonal antibody directed against both the soluble and membrane-bound forms of IL-6R, has been reported to be effective in the treatment of the cytokine-release syndrome associated with chimeric antigen receptor T-cell therapy. Case series have suggested that tocilizumab is effective in the treatment of severe COVID-19, wherein patients had rapid decreases...
inflammatory markers (ie, C-reactive protein, ferritin, D-dimer) and improved oxygenation (ie, PaO\textsubscript{2}/FiO\textsubscript{2} ratio) and lymphocyte count.\textsuperscript{84–86} Two recent nonrandomized, observational studies recapitulated these findings and demonstrated that tocilizumab might improve survival in severe COVID-19.\textsuperscript{87,88} Somers et al\textsuperscript{87} reported that tocilizumab use was associated with a 45% reduced risk for death versus supportive care in patients with COVID-19 on mechanical ventilation (HR = 0.55; 95% CI, 0.33 to 0.90). Tocilizumab use was also associated with an increased risk for superinfections in this population of mechanically ventilated patients (54% versus 26%; P < 0.001), although there was no difference in the 28-day case fatality rate among patients receiving tocilizumab with and without

### Table 1  
**Selected Candidate Immunomodulator Therapies for Severe COVID-19**

| Drug class     | Rationale                                                                 | Clinical evidence                                      | Current status                                                                 |
|---------------|---------------------------------------------------------------------------|--------------------------------------------------------|-------------------------------------------------------------------------------|
| Corticosteroids | Broad immunosuppression                                                   | Randomized, controlled RECOVERY trial showed dexamethasone decreased 28-day mortality vs supportive care (22.9% vs 25.7%) in hospitalized COVID-19 patients; subgroup analysis revealed improved survival for patients on mechanical ventilation (HR = 0.64; 95% CI, 0.51–0.81) and patients requiring supplemental oxygenation but not on mechanical ventilation (HR = 0.82; 95% CI, 0.72–0.94), but no survival benefit for patients not requiring supplemental oxygenation (HR = 1.19; 95% CI, 0.91–1.55) | Treatment with dexamethasone has been incorporated into NIH COVID-19 Treatment Guidelines for hospitalized patients who are on mechanical ventilation and patients who require supplemental oxygenation* |
| IL-6 antagonists | Activity in cytokine release syndrome associated with CAR T cell therapy | Limited; retrospective, nonrandomized cohort studies suggest that benefit in severe COVID-19, but confirmation required | Randomized, controlled trials in progress (NCT04306705, NCT04366355, NCT04320615, NCT04331808, NCT0435071, NCT04322773, NCT0433914, NCT04330638) |
| IL-1 antagonists | Activity in macrophage activation syndrome and hemophagocytic lymphohistiocytosis | Limited; retrospective, nonrandomized cohort studies suggest that benefit in severe COVID-19, but confirmation required | Randomized, controlled trials in progress (NCT04341584, NCT04324021, NCT04330638) |
| BTK inhibitors | Inhibits TLR signaling and cytokine production in activated macrophages; prevents lethal lung injury in mouse influenza model | Limited; possible benefit suggested from small case series and uncontrolled pilot study that showed normalization of inflammatory markers (ie, CRP, IL-6) and improved oxygenation with treatment, but confirmation required | Randomized, controlled trials in progress (NCT04375397, NCT04380688, NCT04382586) |
| Hydroxychloroquine | In vitro activity against SARS-CoV-2; inhibits TLR signaling and cytokine production | Inconsistent results from retrospective, nonrandomized cohort studies; recent randomized, controlled trials demonstrated no benefit in COVID-19 patients with mild/moderate or severe disease as well as when used for post-exposure prophylaxis | Emergency Use Authorization revoked by US FDA on June 15, 2020\textsuperscript{1} |

Detailed information for each clinical trial can be accessed by searching the unique NCT identifier number on www.clinicaltrials.gov.

*https://www.covid19treatmentguidelines.nih.gov/immune-based-therapy/immunomodulators/corticosteroids, last accessed August 19, 2020.

1https://www.fda.gov/news-events/press-announcements/coronavirus-covid-19-update-fda-revokes-emergency-use-authorization-chloroquine-and, last accessed August 19, 2020.  
BTK, Bruton tyrosine kinase; CAR, chimeric antigen receptor; CRP, C-reactive protein; FDA, US Food and Drug Administration; HR, hazard ratio; NIH, National Institutes of Health; RECOVERY, Randomized Evaluation of Covid-19 Therapy\textsuperscript{78}; TLR, toll-like receptor.
superinfection (22% versus 15%; \( P = 0.42 \)). Likewise, in patients admitted with COVID-19–related pneumonia, Guaraldi et al reported that tocilizumab use was associated with a 39% reduced risk for invasive mechanical ventilation or death versus supportive care (HR = 0.61; 95% CI, 0.40 to 0.91; \( P = 0.02 \)).

Two nonrandomized, observational studies have also evaluated the efficacy of the IL-1R antagonist anakinra in blocking the activity of IL-1\( \beta \) and inflammasome signaling in patients with severe COVID-19. Anakinra has been reported to be effective in patients with secondary hemophagocytic lymphohistiocytosis, including when triggered by viral infections, as well as sepsis in those with hyperinflammation. In severe COVID-19, anakinra use was associated with a rapid reduction in C-reactive protein and a progressive improvement in respiratory function (ie, \( \text{PaO}_2/\text{FiO}_2 \) ratio). Cavalli et al reported a higher 21-day survival with anakinra use versus supportive care (90% versus 56%; \( P = 0.009 \)), and Huet et al showed that anakinra use was associated with a 78% decreased risk for mortality versus supportive care (HR = 0.22; 95% CI, 0.10 to 0.49; \( P = 0.0002 \)). Similar to IL-6R antagonism with tocilizumab, anakinra use also was associated with an increased risk for infection. Randomized, controlled studies of the clinical benefits of tocilizumab, anakinra, and additional selective cytokine inhibitors in severe COVID-19 are ongoing.

The dependence on BTK to transmit proinflammatory TLR/IL-1R signaling in monocytes has prompted the
consideration of BTK inhibitors in the abrogation of the cytokine storm in COVID-19. BTK inhibitors are approved and widely used in patients with B-cell malignancies and chronic graft-versus-host disease. In these patients, the use of the BTK inhibitor ibrutinib was associated with markedly reduced levels of inflammatory cytokines and chemokines (ie, IL-6, TNF-α, GM-CSF, IP-10/CXCL10, MCP-1/CCL2, MIP-1α/CCL3, and MIP-1β/CCL4) that overlap with those elevated in the plasma and BALF in severe COVID-19 patients.93–95 Ibrutinib was reported to block pulmonary cytokine production and fatal lung injury in a murine model of influenza.96 Treon et al97 first reported a possible clinical benefit of ibrutinib in a case series of Waldenström macroglobulinemia patients infected with COVID-19. Results from a pilot study of another BTK inhibitor, acalabrutinib, were subsequently reported in patients hospitalized with severe COVID-19 (n = 19).51 After the initiation of acalabrutinib treatment, most patients had normalization of C-reactive protein, IL-6, and lymphocyte levels.51 This decrease in inflammation was correlated with improved respiratory function and oxygenation; 73% of patients requiring supplemental oxygenation were discharged on room air, and 50% of patients on mechanical ventilation were extubated.51 Randomized, controlled studies evaluating the BTK inhibitors ibrutinib, acalabrutinib, and zanubrutinib in severe COVID-19 are ongoing and results are awaited.

Hydroxychloroquine has been proposed as a treatment of COVID-19 based on in vitro activity against SARS-CoV-2. Although the mechanism of antiviral activity is poorly defined, hydroxychloroquine has been reported to increase the pH of endosomes used by the virus for cell entry and to interfere with viral binding to cellular receptors.58 Anti-inflammatory activity with hydroxychloroquine has also been demonstrated, including the inhibition of TLR signaling and cytokine production.59 Nonrandomized, observational studies have reported conflicting results on the clinical benefit of hydroxychloroquine in severe COVID-19.100–103 However, recent results from randomized, controlled clinical trials have indicated that hydroxychloroquine is not effective in hospitalized patients with severe COVID-19.104 In addition to dexamethasone, the RECOVERY trial also evaluated the use of hydroxychloroquine (n = 1561) versus usual care (n = 3155) in hospitalized COVID-19 patients.104 The study found no difference in 28-day mortality between patients randomized to receive hydroxychloroquine versus usual care (26.8% versus 25.0%; P = 0.18).104 Moreover, hydroxychloroquine was associated with a longer time to discharge (median, 16 versus 13 days) and with an increased risk for mechanical ventilation or death among patients not on invasive mechanical ventilation at baseline compared to usual care (HR = 1.12; 95% CI, 1.01 to 1.25).104 The World Health Organization recently discontinued the SOLIDARITY study, a randomized, controlled trial of hydroxychloroquine versus supportive care in hospitalized COVID-19 patients, as hydroxychloroquine did not reduce mortality (WHO, https://www.who.int/news-room/detail/04-07-2020-who-discontinues-hydroxychloroquine-and-lopinavir-ritonavir-treatment-arms-for-covid-19, last accessed July 18, 2020). Randomized, controlled studies of hydroxychloroquine in patients with mild or moderate COVID-19 disease as well as postexposure prophylaxis similarly showed no clinical benefit.105–107

Conclusion

The rapid spread of SARS-CoV-2 infection has resulted in an unprecedented public health crisis worldwide. Although the majority of cases of COVID-19 are asymptomatic or mild, COVID-19 can manifest as severe viral pneumonia that leads to the need for mechanical ventilation, or death. The natural history of COVID-19 includes an initial stage of viral replication that can be followed by a second stage of immunopathology driven by a hyperinflammatory response to SARS-CoV-2 infection. This review presented the current evidence indicating that a dysregulated host innate immune response underlies the hyperinflammatory syndrome in COVID-19; the disease mechanism is summarized in Figure 1. The syndrome shares overlapping features with a viral-induced hemophagocytic lymphohistiocytosis, including evidence of macrophage activation with a cytokine storm, NK-cell and CD8+ T-cell impairment, and hemophagocytic macrophages. Therapies modulating the immune response may be crucial for treating and preventing immunopathology in patients who progress to severe disease. Given the rapidly evolving nature of the COVID-19 pandemic, studies in which the peer-review process has not yet been completed were discussed herein to present the most recent data available, but the findings should be interpreted with caution as the conclusions may change. Future studies are needed for identifying the exact drivers of the pathologic inflammation and biomarkers that predict a hyperinflammatory response to SARS-CoV-2 infection. The results of these studies will enable the identification of the most appropriate immunomodulating agents as well as the optimal timing of such interventions to achieve maximal therapeutic benefit in patients with COVID-19.

Acknowledgment

We thank Tyler Kalajian for generating the figure showing the disease mechanism in severe COVID-19.

References

1. Zhu N, Zhang D, Wang W, Li X, Yang B, Song J, Zhao X, Huang B, Shi W, Lu R, Niu P, Zhan F, Ma X, Wang D, Xu W, Wu G, Gao GF, Tan W: A novel coronavirus from patients with pneumonia in China, 2019. N Engl J Med 2020, 382:727–733
2. Lu R, Zhao X, Li J, Niu P, Yang B, Wu H, Wang W, Song H, Huang B, Zhu N, Bi Y, Ma X, Zhan F, Wang L, Hu T, Zhou H, Hu Z,
14

Bryce C, Grimes Z, Pujadas E, Ahuja S, Beasleys MB, Albrecht R, et al: Pathophysiology of SARS-CoV-2: targeting of endothelial cells renders a complex disease with thrombotic microangiopathy and aberrant immune response. The Mount Sinai COVID-19 autopsy experience. medRxiv 2020. [Epub] doi:10.1016/2020.05.18.20099960

Ramos-Casals M, Brito-Zerón P, López-Guillermo A, Khamashia MA, Bosch X: Adult haemophagocytic syndrome. Lancet 2014, 383:1503–1516

Ackermann M, Verleden SE, Kuehnel M, Haverich A, Welte T, Laenger F, Vanstapel A, Werlein C, Stark H, Tzankov A, Li WW, Li VW, Mentzer SJ, Jonigk D: Pulmonary vascular endotheliitis, thrombosis, and angiogenesis in COVID-19. N Engl J Med 2020, 383:120–128

Varga Z, Flammer AJ, Steiger P, Haberecker M, Andermatt R, Zinkenagel AS, Mehra MR, Schuepbach RA, Ruschitzka F, Moch H: Endothelial cell infection and endothelitis in COVID-19. Lancet 2020, 395:1417–1418

Monteil V, Kwon H, Prado P, Hagleitner A, Wimmer RA, Stahl M, Leopoldi A, Garreta E, Hurtado del Pozo C, Prosper F, Romero JPM, Wirnsberger G, Zhang H, Slutsky AS, Conder R, Montserrat N, Mirazimi A, Penninger JM: Inhibition of SARS-CoV-2 infections in engineered human tissues using clinical-grade soluble human ACE2. Cell 2020, 181:905–913.e7

Bao L, Deng W, Huang B, Gao H, Liu J, Ren L, et al: The pathogenicity of SARS-CoV-2 in hACE2 transgenic mice. Nature 2020, 583:830–833

Nicholls JM, Poon LLM, Lee KC, Ng WF, Lai ST, Leung CY, Chu CM, Hui PK, Mak KL, Lim W, Yan KW, Chan KH, Tsang NC, Guan Y, Yuen KY, Malik Peiris JS: Lung pathology of fatal severe acute respiratory syndrome. Lancet 2003, 361:1773–1778

Chen G, Wu D, Guo W, Cao Y, Huang D, Wang H, Wang T, Zhang X, Chen H, Yu H, Zhang M, Wu S, Song J, Chen T, Han M, Li J, Xiao Y, Zhao J, Ning Q: Clinical and immunological features of severe and moderate Coronavirus disease 2019. J Clin Invest 2020, 130:2620–2629

Tay MZ, Poh CM, Rénia L, MacAry PA, Ng LFP: The trinity of COVID-19: immunity, inflammation and intervention. Nat Rev Immunol 2020, 20:363–374

Schifanella L, Anderson JL, Galli M, Corbellino M, Lai A, Wicking G, Grzywacz B, Klatt NR, Haase AT, Schacker TW: Massive viral replication and cytotoxic effects in early COVID-19 pneumonia. arXiv 2020. [Epub] doi:10.1016/2020.05.18.20099960

Zhang H, Zhou P, Wei Y, Yue H, Wang Y, Hu M, Zhang S, Cao T, Chu Y, Yang C, Guo G, Chen X, Chen Y, Lei M, Liu H, Zhao J, Peng P, Wang C-Y, Du R: Histopathologic changes and SARS-CoV-2 immunostaining in the lung of a patient with COVID-19. Ann Intern Med 2020, 172:629–632

Fink SL, Cookson BT: Apoptosis, pyroptosis, and necroinflammation: a new paradigm of necroinflammatory injury in systemic inflammation and intervention. Nat Med 2020, 26:1907–1916

Yang M: Cell pyroptosis, a potential pathogenic mechanism of 2019-nCoV infection. SSRN 2020. [Epub] doi:10.2139/ssrn.3527420

Yap JKY, Moriyama M, Iwasaki A: Inflammotrypsins and pyroptosis as therapeutic targets for COVID-19. J Immunol 2020, 205:307–312

St John AL, Rathore APS: Early insights into immune responses during COVID-19. J Immunol 2020, 205:555–564

Liao M, Liu Y, Yuan J, Wen Y, Xu G, Zhao J, Cheng L, Li J, Wang X, Wang F, Liu L, Amit I, Zhang S, Zhang H: Single-cell landscape of bronchoalveolar immune cells in patients with COVID-19. Nat Med 2020, 26:842–844

Mills CD, Kincaid K, Alt JM, Heilmann ML, Hill AM: M-1/M-2 macrophages and the Th1/Th2 paradigm. J Immunol 2000, 164:6166–6173

Zhang B, Zhou X, Qiu Y, Feng F, Feng J, Jia Y, Zhu H, Hu K, Liu J, Liu Z, Wang S, Gong Y, Zhou C, Zhu T, Cheng Y, Liu Z, Deng H, Tao F, Ren Y, Cheng B, Gao L, Wu X, Yu L, Huang Z, Mao Z, Song Q, Zhu B: Clinical characteristics of 82 death cases with COVID-19. PLoS One 2020, 15(7):e0235458

Gustine and Jones

Zhou W, Zhao L, Chen J, Meng Y, Wang J, Lin Y, Yuan J, Xie Z, Ma J, Liu WJ, Wang D, Xu W, Holmes EC, Gao GF, Wu G, Chen W, Shi W, Tan W: Genomic characterisation and epidemiology of 2019 novel coronavirus: implications for virus origins and receptor binding. Lancet 2020, 395:565–574

Li Q, Guan X, Wu P, Wang X, Zhou L, Tong Y, et al: Early transmission dynamics in Wuhan, China, of novel coronavirus-infected pneumonia. N Engl J Med 2020, 382:1199–1207

Huang C, Wang Y, Li X, Ren L, Zhao J, Hu Y, Zhang L, Fan G, Xu J, Gu X, Cheng Z, Yu T, Xie J, Wei Y, Wu W, Xie Y, Yin W, Li H, Liu M, Xiao Y, Gao H, Guo L, Xie J, Wang G, Jiang R, Gao Z, Jin Q, Wang J, Cao B: Clinical features of patients with infected with novel coronavirus in Wuhan, China. Lancet 2020, 395:497–506

Guan W-J, Ni Z-Y, Hu Y, Liang W-H, Ou C-Q, He J-X, et al: China Medical Treatment Expert Group for Covid-19: Clinical characteristics of Coronavirus disease 2019 in China. N Engl J Med 2020, 382:1708–1720

Richardson S, Hirsch JS, Narasimhan M, Crawford JM, McGinn T, Davidson KW: Northwell COVID-19 Research Consortium: Presenting characteristics, comorbidities, and outcomes among 5700 patients hospitalized with COVID-19 in the New York City area. JAMA 2020, 323:2052–2059

Chen N, Zhou M, Dong X, Qiu J, Gong F, Han Y, Qiu Y, Wang J, Liu Y, Wei Y, Xia J, Yu T, Zhang X, Zhang L: Epidemiological and clinical characteristics of 99 cases of 2019 novel coronavirus pneumonia in Wuhan, China: a descriptive study. Lancet 2020, 395:507–513

Zhou F, Yu T, Du R, Fan G, Liu Y, Liu Z, Xiang J, Wang Y, Song B, Gu X, Guan L, Wei Y, Li H, Wu X, Xu J, Tu S, Zhang Y, Chen H, Cao B: Clinical course and risk factors for mortality of adult inpatients with COVID-19 in Wuhan, China: a retrospective cohort study. Lancet 2020, 395:1054–1062

Wu C, Chen X, Cai Y, Xia J, Zhou X, Xu S, Huang H, Zhang L, Zhou X, Du C, Zhang Y, Song J, Wang S, Chao Y, Yang Z, Xu J, Zhou X, Chen D, Xiong W, Xu L, Zhou F, Jiang J, Bai C, Zheng J, Song Y: Risk factors associated with acute respiratory distress syndrome and death in patients with Coronavirus disease 2019 pneumonia in Wuhan, China. JAMA Intern Med 2020, 180:934–943

Mehta P, McAuley DF, Brown M, Sanchez E, Tattersall RS, Manson JJ: COVID-19: consider cytokine storm syndromes and immunosuppression. Lancet 2020, 395:1033–1034

Ruan Q, Yang K, Wang W, Jiang L, Song J: Clinical predictors of mortality due to COVID-19 based on an analysis of data of 150 patients from Wuhan, China. Intensive Care Med 2020, 46:846–848

Del Valle DM, Kim-schulze S, Hsin-hui H, Beckmann ND, Nirenberg S, Wang B, Lavin Y, Swartz T, Madduri D, Stock A, Marron T, Xie H, Patel MK, van Oekelen O, Rahman A, Kovatch P, Aberg J, Schade T, Jagannath S, Mazumdar M, Channay Y, FitzBetancourt A, Mendu DR, Jiang J, Reich D, Sigel K, Cordon-Carlo C, Feldmann M, Parekh S, Mead M, Gujatic S: An inflammatory cytokine signature helps predict COVID-19 severity and survival. Nature Medicine 2020, 26:1636–1643

Xu Z, Shi L, Wang Y, Zhang J, Huang L, Zhang C, Liu S, Zhao P, Liu H, Zhu L, Tai Y, Bai C, Gao T, Song J, Xia P, Dong J, Zhao J, Wang F-S: Pathological findings of COVID-19 associated with acute respiratory distress syndrome. Lancet Respir Med 2020, 8:420–422

Fox SE, Akmatbekov A, Harbert JL, Li G, Quincy Brown J, Vander Heide RS: Pulmonary and cardiac pathology in African American patients with COVID-19: an autopsy series from New Orleans. Lancet Respir Med 2020, 8:681–686

Carsana L, Sonzogni A, Nasr A, Rossi RS, Pellegrinelli A, Zerbi P, Rech R, Colombo R, Antonini S, Corbellino M, Galli M, Catena E, Tosoni A, Gianatti A, Nebuloni M: Pulmonary post-mortem findings in a series of COVID-19 cases from northern Italy: a two-centre descriptive study. Lancet Infect Dis 2020, 20:1135–1140
34. Ziegler CGK, Allon SJ, Nyquist SK, Mbano IM, Miao VN, Tzouanas CN, et al; HCA Lung Biological Network: SARS-CoV-2 receptor ACE2 is an interferon-stimulated gene in human airway epithelial cells and is detected in specific cell subsets across tissues. Cell 2020, 181:1016–1035.e19

35. Pechkovsky DV, Goldmann T, Ludvig C, Prasse A, Vollmer E, Müller-Quernheim J, Zissel G: CCR2 and CXCR3 agonistic chemokines are differently expressed and regulated in human alveolar epithelial cells type II. Respir Res 2005, 6:75

36. Thorley AJ, Grandolfo D, Lim E, Goldstraw P, Young A, Tettley TD: Innate immune responses to bacterial ligands in the peripheral human lung–role of alveolar epithelium, TLR expression and signalling. PLoS One 2011, 6:e21827

38. He L, Ding Y, Zhang Q, Che X, He Y, Shen H, Wang H, Li Z, Zhao L, Geng J, Deng Y, Yang L, Li J, Cai J, Qiu L, Wen K, Xu X, Jiang S: Expression of elevated levels of pro-inflammatory cytokines in SARS-CoV-infected ACE2+ cells in SARS patients: relation to the acute lung injury and pathogenesis of SARS. J Pathol 2006, 210: 288–297

39. Blanco-Melo D, Nilsson-Payant BE, Liu W-C, Uhl S, Hoagland D, Jiang S: Expression of elevated levels of pro-inflammatory cytokines in SARS-CoV-infected ACE2+ cells in SARS patients: relation to the acute lung injury and pathogenesis of SARS. J Pathol 2006, 210: 288–297

40. Adjadji J, Yatim N, Barnabei L, Corneau A, Boussier J, Smith N, Pérè H, Charbit B, Bondet V, Chevenier-Gobeaux C, Braillat P, Carlier N, Gauzit R, Morbec C, Péne F, Marin N, Roche N, Srwebel T-A, Merkling SH, Treluyer J-M, Mangan M, Düwell P, Mouthon L, Blanc C, Tharaux P-L, Rozenberg F, Fischer A, Duffy D, Rieux-Laucat F, Kernéis S, Terrier B: Impaired type I interferon activity and inflammatory responses in severe COVID-19 patients. Science 2020, 369:718–724

41. Wang N, Zhan Y, Zhu L, Hou Z, Liu F, Song P, Qiu F, Wang X, Zou X, Wan D, Qian X, Wang S, Guo Y, Yu H, Cui M, Tong G, Xu Y, Zheng Z, Lu Y, Hong P: Retrospective multicenter cohort study shows early interferon therapy is associated with favorable response to SARS-CoV-2 drives development of COVID-19. Cell 2020, 181:1016–1035.e19

42. Chen X, Yang X, Zheng Y, Yang Y, Xing Y, Chen Z: SARS coronavirus papain-like protease inhibits the type I interferon signaling pathway through interaction with the STING-TRAF3-TBK1 complex. Protein Cell 2014, 5:369–381

43. Yang Y, Ye F, Zhu N, Wang W, Deng Y, Zhao Z, Tan W: Middle East respiratory syndrome coronavirus ORF4b protein inhibits type I interferon production through both cytoplasmic and nuclear targets. Sci Rep 2015, 5:17554

44. Lui PY, Wong LY, Fung CL, Siu KL, Yeung ML, Yuen KS, Chan CP, Woo PC, Yuen KY, Jin DY: Middle East respiratory syndrome coronavirus M protein suppresses type I interferon expression through the inhibition of TBK1-dependent phosphorylation of IRF3. Emerg Microbes Infect 2016, 5:e39

45. Channappanavar R, Fehr Anthony R, Vijay R, Mack M, Zhao J, Meyerholz David K, Perlman S: Dysregulated type I interferon and inflammatory monocyte-macrophage responses cause lethal pneumonia in SARS-CoV-infected mice. Cell Host Microbe 2016, 19: 181–193

46. Zhou Z, Ren L, Zhang L, Zhong J, Xiao Y, Jia Z, Guo L, Yang J, Wang C, Jiang S, Yang D, Zhang G, Li H, Chen F, Xu Y, Chen M, Gao Z, Yang J, Dong J, Liu B, Zhang X, Wang W, He K, Jin Q, Li M, Wang J: Heightened innate immune responses in the respiratory tract of COVID-19 patients. Cell Host Microbe 2020, 27:883–890.e2

47. Mo X, Jian W, Su Z, Chen M, Peng H, Peng P, Lei C, Li S, Chen R, Zhong N: Abnormal pulmonary function in COVID-19 patients at time of hospital discharge. Eur Respir J 2020, 55:2001217

48. Wang Y, Dong C, Hu Y, Li C, Ren Q, Zhang X, Shi H, Zhou M: Temporal changes of CT findings in 90 patients with COVID-19 pneumonia: a longitudinal study. Radiology 2020, 296:E55–E64

49. Xiong Y, Liu Y, Cao L, Wang D, Guo M, Jiang A, Guo D, Hu W, Yang J, Tang Z, Wu H, Lin Y, Zhang M, Zhang Q, Shi M, Liu Y, Zhou Y, Lan K, Chen Y: Transcriptomic characteristics of bronchoalveolar lavage fluid and peripheral blood mononuclear cells in COVID-19 patients. Emerg Microbes Infect 2020, 9:761–770

50. Zhou Y, Fu B, Zhang X, Wang D, Zhao C, Qi Y, Sun R, Tian Z, Xu X, Wei H: Pathogenic T-cells and inflammatory monocytes incite inflammatory storms in severe COVID-19 patients. Nat Sci Rev 2020, 7:998–1002

51. Roschewski M, Lounakis MS, Sharman JP, Roswarski J, Goy A, Monticelli MA, Roshon M, Wrzesinski SH, Desai JV, Zaraka M, Collen J, Rose KM, Hamdy A, Izumi R, Wright GW, Chung KK, Baselga J, Staudt LM, Wilson WH: Inhibition of Bruton tyrosine kinase in patients with severe COVID-19. Sci Immunol 2020, 5:eabd0110

52. Wen W, Wu S, Wang H, Le W, Zhang Y, Zheng Y, Liu X, Xie L, Li J, Ye J, Dong L, Cui X, Miao Y, Wang D, Dong J, Xiao C, Chen W, Wang H: Immune cell profiling of COVID-19 patients in the recovery stage by single-cell sequencing. Cell Discov 2020, 2:631

53. Ong EZ, Chan YFZ, Leong WY, Lee NMY, Balabin RM: Mokximmun A, Haji Mohideen SM, Chan KS, Tan AT, Bertolotti A, Ooi EE, Low JGH: A dynamic immune response shapes COVID-19 progression. Cell Host Microbe 2020, 27:879–882.e2

54. Guo C, Li B, Ma H, Wang X, Cai P, Yu Q, Zhu L, Jin L, Jiang C, Fang J, Liu Q, Zong D, Zhang W, Lu Y, Li K, Gao X, Fu B, Liu L, Ma X, Gao J, Wei H, Ji T, Lin J, Qi K: Single-cell analysis of severe COVID-19 patients reveals a monocyte-driven inflammatory storm attenuated by tocilizumab. Nat Commun 2020, 11:3924

55. Page TH, Urbania AM, Espirito Santo A, Danks L, Smallie T, Williams LM, Horwood NJ: Bruton’s tyrosine kinase regulates TLR7/8-induced TNF transcription via nuclear factor-[kappa]B recruitment. Biochem Biophys Res Commun 2018, 499:260–266

56. Ito M, Shicita T, Okada M, Komine R, Noguchi Y, Yoshimura A, Morita R: Bruton’s tyrosine kinase is essential for NLRP3 inflammasome activation and contributes to ischaemic brain injury. Nat Commun 2015, 6:7360

57. Bittner ZA, Liu X, Dickhöfer S, Kalbacher H, Bock K, Andreeva L, Marcu A, Stevanovic S, Mangan M, Düwell P, Lovotti M, Herster F, Löfler MW, Shankar S, Tapia-Abellán A, Wolz O-O, Schilling NA, Kümmerle-Deschner J, Wagner S, Delor A, Grimbacher B, Wu H, Latz E, Weber ANR: BTK operates a phospho-tyrosine switch to regulate NLRP3 inflammasome activity. bioRxiv 2019:864702

58. Akira S, Uematsu S. Takeuchi O. Pathogen recognition and innate immunity. Cell 2006, 124:783–801

59. Sheahan T, Morrison TE, Funkhouser W, Uematsu S, Akira S, Baric RS, Heise MT: MyD88 is required for protection from lethal infection with a mouse-adapted strain of SARS-CoV. PLoS Pathog 2008, 4:e1000240

60. Wilk AJ, Rustagi A, Zhao NQ, Roque J, Martinez-Colón GJ, McKechnie JL, Ivison GT, Rangathan T, Vergara R, Hollis T, Simpson LJ, Grant P, Subramanian A, Rogers AJ, Blish CA: A single-cell atlas of the peripheral immune response in patients with severe COVID-19. Nat Med 2020, 26:1070–1076

61. Zheng M, Gao Y, Wang G, Song G, Liu S, Sun D, Xu Y, Tian Z: Functional exhaustion of antiviral lymphocytes in COVID-19 patients. Cell Mol Immunol 2020, 17:536–546

62. Wang F, Nie J, Wang H, Zhao Q, Xiong Y, Deng L, Song S, Ma Z, Mo P, Zhang Y: Characteristics of peripheral lymphocyte subset alteration in COVID-19 pneumonia. J Infect Dis 2020, 221:1764–1774

63. Giamarellos-Bourboulis EJ, Netea MG, Rovina N, Akinosoglou K, Antoniadou A, Antonakos N, Damoraki G, Gkavogianni T,
Gustine and Jones

Adami M-E, Katsaounou P, Nagano M, Kyriakopoulou M, Dimopoulos G, Koutsodimitropoulos I, Velissaris D, Koufarypiss P, Karageorgos A, Katrini K, Lekaikis V, Lupsa M, Kotsaki A, Reniers G, Theodoulou D, Panou V, Koukaki E, Koulouris N, Gogos C, Koutsoukou A: Complex immune dysregulation in COVID-19 patients with severe respiratory failure. Cell Host Microbe 2020, 27:992–1000.e3

McKechnie JL, Blish CA: The innate immune system: fighting on the front lines or fanning the flames of COVID-19? Cell Host Microbe 2020, 27:863–869

Carlin LE, Hemann EA, Zacharias ZR, Heusel JW, Legge KL: Natural killer cell recruitment to the lung during influenza A virus infection is dependent on CXCR3, CCR5, and virus exposure dose. Front Immunol 2018, 9:781

Cifaldi L, Prencipe G, Caiello I, Bracaglia C, Locatelli F, De Benedetti F, Strippoli R: Inhibition of natural killer cell cytotoxicity by IL-6: implications for the pathogenesis of macrophage activation syndrome. Arthritis Rheumatol 2015, 67:3037–3046

Vagnats A, Massoudende M, Stoll G, Routy B, Fluckiger A, Yamazaki T, Iribarren K, Duong CPM, Fend L, Caignard A, Cremer I, LeCesne A, Adam J, Honoré C, Mir O, Chaigneau L, Berger A, Validire P, Christidis C, Brun-Ly VL, Smyth MJ, Mariette X, Salomon BL, Kroemer G, Rusakiewicz S, Zitvogel L: TNFR2/BIRC3-TRAF1 signaling pathway as a novel NK cell immune checkpoint in cancer. Oncoimmunology 2018, 7:e1386826

Vabret N, Britton GJ, Gruber C, Hegde S, Kim J, Kuksin M, et al; Sinai Immunology Review Project: Immunology of COVID-19: current state of the science. Immunity 2020, 52:910–941

Chen Y, Feng Z, Diao B, Wang R, Wang G, Wang C, Tan Y, Liu L, Wang C, Liu Y, Liu Y, Yuan Z, Ren L, Wu Y: The novel severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) directly decimates human spleens and lymph nodes. medRxiv 2020. [Epub ahead of print] doi:10.1101/2020.03.27.20045427

Egeblad M, Schiffman JD, Yost CC: Neutrophil extracellular traps in pulmonary infection is dependent on CXCR3, CCR5, and virus exposure dose. Front Immunol 2020, 27:992

Feldstein LR, Rose EB, Horwitz SM, Collins JP, Newhams MM, Porto BN, Stein RT: Neutrophil extracellular traps in pulmonary diseases: too much of a good thing? Front Immunol 2016, 7:311

Merson L, Hayden FG, Fowler RA: Corticosteroid therapy for critically ill patients with Middle East respiratory syndrome. Am J Respir Crit Care Med 2016, 193:1507–1517

Vuk et al; RECOVERY Collaborative Group: Dexamethasone in hospitalized patients with Covid-19–preliminary report. N Engl J Med 2020. [Epub ahead of print] doi:10.1056/NEJMoa2021436

Teachey DT, Rheingold SR, Maude SL, Seif AE, Nichols KE, Suppa EK, Kals B, Bergh RA, Fitzgerald JC, Aplenc R, Gore L, Grupp SA: Cytokine release syndrome after blinatumomab treatment related to abnormal macrophage activation and ameliorated with cytokine-directed therapy. Blood 2013, 121:5154–5157

Maude SL, Frey N, Shaw PA, Aplenic R, Barrett DM, Seif AE, Nichols KE, Suppa EK, Kals B, Bergh RA, Fitzgerald JC, Aplenic R, Gore L, Grupp SA: Cytokine release syndrome related to abnormal macrophage activation and ameliorated with cytokine-directed therapy. Blood 2013, 121:5154–5157

Hanauer DA, Martin E, Sharma P, Fung CM, Pogue JM: Tocilizumab for COVID-19: a single center experience. J Med Virol 2020, 92:1517–1521

Xu X, Han M, Li T, Sun W, Wang D, Fu B, Zhou Y, Zheng X, Yang Y, Li X, Zhang X, Pan A, Wei H: Effective treatment of severe COVID-19 patients with tocilizumab. Proc Natl Acad Sci U S A 2020, 117:10970–10975

Sciascia S, Aprà F, Baffa A, Baldovino S, Boaro D, Boero R, Boffa R, Calcagno A, Cecchi I, Cininnella G, Converso M, Cozzi M, Crosioso P, De Iaco F, Di Perri G, Eandi M, Fenoglio R, Giusti M, Imperiale D, Imperiale G, Livigni S, Manno E, Massara C, Milone V, Natale G, Navarra M, Oddone V, Osella S, Piccioni P, Rasin M, Roccatello D, Rossi D: Pilot prospective open, single-arm multicentre study on off-label use of tocilizumab in patients with severe COVID-19. Clin Exp Rheumatol 2020, 38:529–532

Luo P, Liu Y, Qiu L, Liu X, Liu D, Li J: Tocilizumab treatment in COVID-19: a single center experience. J Med Virol 2020, 92:814–818

Somers EC, Eschenauer GA, Troost JP, Golob JL, Gandhi TN, Wang L, Zhou N, Petty LA, Baang JH, Dillman NO, Frame D, Gregg KS, Kaur DL, Nagel J, Patel TS, Zhou S, Lauring AS, Hanauer DA, Martin E, Sharma P, Fung CM, Pogue JM: Tocilizumab for treatment of mechanically ventilated patients with COVID-19. Clin Infect Dis 2020. [Epub ahead of print] doi:10.1093/cid/ciaa954

Guandalini G, Meschiari M, Cozzi-Lepri A, Milic J, Tonelli R, Menozzi M, Franceschini E, Cuomo G, Orlando G, Borghi V, Santoro A, Di Gaetano M, Puzzolante C, Carli F, Bedini A, Corradi L, Fantini R, Castaniere I, Tabbi L, Girardi M, Teledeschi S, Giannella M, Bartoletti M, Pascale R, Dolci G, Brugioni L,
90. Huet T, Beausser H, Voisin O, Jouveshomme S, Dauriat G, Lazareth I, Sacco E, Naccache J-M, Bézie Y, Laplanche S, Le Berre A, Le Pavec J, Salmeron S, Emmerich J, Mourad J-I, Chatellier G, Gayem G. Anakinra for severe forms of COVID-19: a cohort study. Lancet Rheumatol 2020, 2:e325–e331.

91. Mehta P, Cron RQ, Hartwell J, Manson JJ, Tattersall RS. Silencing the cytokine storm: the use of intravenous anakinra in haemophagocytic lymphohistiocytosis or macrophage activation syndrome. Lancet Rheumatol 2020, 2:e358–e367.

92. Shakoory B, Carcillo JA, Chatham WW, Amdur RL, Zhao H, Di Mare B, Grifoni A, Piazza P, De Cornaro S, Dauriat G, et al. Hydroxychloroquine with or without azithromycin in combination against SARS-CoV-2: what molecular dynamics studies of virus-host interactions reveal. Int J Antimicrob Agents 2020, 56:106020.

93. Niemann CU, Herman SEM, Maric I, Gomez-Rodriguez J, Biancott A, Chang BY, Martyr S, Stetler-Stevenson M, Yang G, Treon SP, Hunter ZR. CXCL13 levels are elevated in patients with Waldenström macroglobulinemia, and are predictive of major response to ibrutinib. Cancer Res 2016, 22:1572–1580.

94. Gero MC, Pals ST, Kersten MJ, Xu L, Yang G, Treon SP, Hunter ZR. Interleukin-1 blockade is associated with reduced mortality in sepsis patients with features of macrophage activation syndrome: reanalysis of a prior phase III trial. Crit Care Med 2016, 44:275–281.

95. Niemann CU, Herman SEM, Marie I, Gomez-Rodriguez J, Biancott A, Chang BY, Martyr S, Stetler-Stevenson M, Yuan CM, Calvo KR, Braylan RC, Valdez J, Lee YS, Wong DH, Jones J, Sun C, Marti GE, Farooqui MZH, Wiestner A. Disruption of in vivo chronic lymphocytic leukemia tumor-microenvironment interactions by ibrutinib—findings from an investigator-initiated phase II study. J Clin Cancer Res 2016, 22:1572–1580.

96. Florence JM, Krupa A, Booshieh LM, Davis SA, Matthay MA, Kurdowska AK. Inhibiting Bruton’s tyrosine kinase rescues mice from lethal influenza-induced acute lung injury. Am J Physiol Lung Cell Mol Physiol 2018, 315:L52–L58.

97. Teun SP, Castillo JJ, Skarbiñik AP, Soumerai JD, Ghobrial IM, Guerrera ML, Meid K, Yang G. The BTK inhibitor ibrutinib may protect against pulmonary injury in COVID-19-infected patients. Blood 2020, 135:1912–1915.

98. Fantini J, Chahinian H, Yahi N. Synergistic antiviral effect of hydroxychloroquine and azithromycin in combination against SARS-CoV-2: what molecular dynamics studies of virus-host interactions reveal. Int J Antimicrob Agents 2020, 56:106020.

99. Schrenzeiner E, Dörner T. Mechanisms of action of hydroxychloroquine and chloroquine: implications for rheumatology. Nat Rev Rheumatol 2020, 16:155–166.

100. Gautret P, Lagier J-C, Parola P, Hoang VT, Meddeb L, Mailhe M, Doudier B, Courjon J, Giordanengo V, Vieira VE, Tissot Dupont H, Honoré S, Colson P, Chabrière E, La Scola B, Rolain J-M, Brouqui P, Raoult D. Hydroxychloroquine and azithromycin as a treatment of COVID-19: results of an open-label non-randomized clinical trial. Int J Antimicrob Agents 2020, 56:105949.

101. Arshad S, Kilgore P, Chaudhry ZS, Jacobsen G, Wang DD, Huisiting K, et al. Hydroxychloroquine and azithromycin, and combination in patients hospitalized with COVID-19. Int J Infect Dis 2020, 97:396–403.

102. Geleris J, Sun Y, Platt J, Zucker J, Baldwin M, Hripsicas G, Labelia A, Manson DK, Kubin C, Barr RG, Sobieszczyn ME, Schluger NW. Observational study of hydroxychloroquine in hospitalized patients with Covid-19. N Engl J Med 2020, 382:2411–2418.

103. Mahéva M, Tran V-T, Roumier M, Chabról A, Paule R, Guillaud C, Fois E, Lepeule R, Szwebel T-A, Lescure F-X, Schlemmer F, Matignon M, Khellaf M, Cricks E, Terrier B, Morbieu C, Legendre D, Sang J, Schoindre Y, Pawlotsky J-M, Michel M, Perrodeau E, Carlier N, Roche N, de Lastours V, Ourghanlian C, Kernen S, Ménager P, Moutoun L, Audureau E, Ravaud P, Godeau B, Gallien S, Cuesta-Blumes N. Clinical efficacy of hydroxychloroquine in patients with COVID-19 pneumonia who require oxygen: observational comparative study using routine care data. BMJ 2020, 369:m1844.

104. Horby P, Mathias M, Linsell L, Bell JL, Staplin N, Emerson JR, Wiselka M, Ustianowski A, Elmahi E, Prudon B, Whitehouse A, Felton T, Williams J, Faccenda J, Underwood J, Baillie JK, Chappell L, Faust SN, Jaki T, Jeffery K, Lim WS, Montgomery A, Rowan K, Tarning J, Watson JA, White NJ, Juszkiewicz E, Haynes R, Landray MJ; RECOVERY Collaborative Group. Effect of hydroxychloroquine in hospitalized patients with COVID-19: preliminary results from a multi-centre, randomized, controlled trial. medRxiv 2020 [Epub] doi:10.1101/2020.07.15.20151852.

105. Cavalcanti AB, Zampieri FG, Rosa RG, Azevedo LCP, Veiga VC, Aveuzem A, Damiani LP, Marcadenti A, Kawano-Dourado L, Lisboa T, Junqueira DLM, de Barros e Silva PGM, Tramujas L, Abreu-Silva EO, Laranjeira LN, Soares AT, Echenique LS, Pereira AJ, Freitas FGR, Gebara OCE, Dantas VCS, Furtado RHM, Milan EP, Golani NA, Cardoso FF, Maia IS, Hoffmann Filho CR, Kormann APM, Amazonas RB, Bocchi de Oliveira MF, Serpa-Neto A, Falavigna M, Lopes RD, Machado FR, Berwanger O. Hydroxychloroquine with or without azithromycin in moderate Covid-19: a multi-centres controlled trial. medRxiv 2020 [Epub] doi:10.1101/2020.07.15.20151852.

106. Mitjá O, Corbacho-Monné M, Ulbas M, Tebe C, Petafiel J, Tobias A, et al; BCN PEP-CoV-2 Research Group. Hydroxychloroquine for early treatment of adults with mild Covid-19: a randomized-controlled trial. Clin Infect Dis 2020. [Epub ahead of print] doi:10.1093/cid/ciaa009.

107. Boluware DR, Pullen MF, Bangdiwala AS, Pastick KA, Lofgren SM, Okafor EC, Skipper CP, Nascene AA, Nicol MR, Abassi M, Engen NW, Cheng MP, LaBar D, Lother SA, MacKenzie LJ, Huitsing K, et al; Henry Ford COVID-19 Task Force. Treatment with hydroxychloroquine, azithromycin, and combination in patients hospitalized with COVID-19. Int J Antimicrob Agents 2020, 56:105949.