**Assessment of Evidence for COVID-19-Related Treatments: Updated 5/15/2020**

The information contained in this evidence table is emerging and rapidly evolving because of ongoing research and is subject to the professional judgment and interpretation of the practitioner due to the uniqueness of each medical facility’s approach to the care of patients with COVID-19 and the needs of individual patients. ASHP provides this evidence table to help practitioners better understand current approaches related to treatment and care. ASHP has made reasonable efforts to ensure the accuracy and appropriateness of the information presented. However, any reader of this information is advised ASHP is not responsible for the continued currency of the information, for any errors or omissions, and/or for any consequences arising from the use of the information in the evidence table in any and all practice settings. Any reader of this document is cautioned that ASHP makes no representation, guarantee, or warranty, express or implied, as to the accuracy and appropriateness of the information contained in this evidence table and will bear no responsibility or liability for the results or consequences of its use. Public access to AHFS Drug Information® (https://www.ahfscdi.com/login) is currently available with the username "ahfs@ashp.org" and password "covid-19." ASHP's patient medication information is available at http://www.safemedication.com/.

Select entries were updated on 5/15/2020; these can be identified by the date that appears in the Drug(s) column.

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**Updated 5-15-20. The current version of this document can be found on the ASHP COVID-19 Resource Center.**

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**ANTIVIRAL AGENTS**

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<th>Rationale</th>
<th>Trials or Clinical Experience</th>
<th>Dosage*</th>
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<tr>
<td>Baloxavir</td>
<td>8:18.92</td>
<td>Antiviral</td>
<td>Only limited clinical trial data available to date to evaluate use of baloxavir for treatment of COVID-19</td>
<td>Protocol for two registered Chinese trials (ChiCTR2000029544, ChiCTR2000029548) specifies an oral baloxavir marboxil dosage of 80 mg on day 1 and on day 4, and another dose of 80 mg on day 7 (as needed); not to exceed 3 total doses.</td>
<td>No data to date support use in the treatment of COVID-19</td>
</tr>
<tr>
<td><strong>Updated 5/13/20</strong></td>
<td></td>
<td></td>
<td>Exploratory, open-label, randomized controlled study at a single center in China (ChiCTR2000029544): 29 adults hospitalized with COVID-19 receiving antiviral treatment with lopinavir/ritonavir, darunavir/copegicst, or umifenovir (Arbidol®), in combination with inhaled interferon-α, were randomized to treatment with baloxavir marboxil (80 mg orally on day 1 and on day 4, and 80 mg orally on day 7 as needed) (n=10), favipiravir (1600 or 2200 mg orally on day 1, followed by 600 mg three times daily for up to 14 days) (n=9), or control (standard antiviral treatment) (n=10). Percentage of pts with viral conversion (2 consecutive tests with undetectable viral RNA results) after 14 days of treatment was 70, 77, and 100% in the baloxavir, favipiravir, and control groups, respectively.</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Another randomized controlled trial registered in China: 1 CHICCTR2000029548 Protocol for two registered Chinese trials (ChiCTR2000029544, ChiCTR2000029548) specifies an oral baloxavir marboxil dosage of 80 mg on day 1 and on day 4, and another dose of 80 mg on day 7 (as needed); not to exceed 3 total doses.</td>
<td></td>
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</tr>
<tr>
<td>Chloroquine</td>
<td>8:30.08</td>
<td>Antimalarial</td>
<td>Only limited clinical trial data available to date to evaluate use of chloroquine for treatment or prevention of COVID-19</td>
<td>Optimal dosage and duration of treatment not known</td>
<td>Efficacy and safety of chloroquine for treatment or prevention of COVID-19 not established</td>
</tr>
<tr>
<td>Phosphate</td>
<td></td>
<td></td>
<td>Clinical experience in treating pts with COVID-19 accumulating; reports of possible clinical benefits, including decrease in viral load and duration of illness; only limited data available to date to support efficacy and identify possible safety concerns in pts with COVID-19.</td>
<td>Consider: 500 mg of chloroquine phosphate is equivalent to 300 mg of chloroquine base.</td>
<td>Additional data needed to determine whether in vitro activity against SARS-CoV-2 corresponds with clinical efficacy for treatment or prevention of COVID-19</td>
</tr>
<tr>
<td><strong>Updated 4/29/20</strong></td>
<td></td>
<td></td>
<td>Double-blind randomized phase 2b study in Brazil (Borba et al) to evaluate two different chloroquine dosages as adjunctive therapy in hospitalized adults with</td>
<td>Various dosages recommended or being investigated for treatment of COVID-19</td>
<td>Additional data needed to substantiate initial reports of efficacy for treatment and identify optimal dose and duration</td>
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<td></td>
<td></td>
<td></td>
<td>Viral activity against SARS-CoV-1 and MERS-CoV</td>
<td>Oral chloroquine phosphate dosage suggested in the EUA: For treatment of hospitalized adults and adolescents weighing 50 kg or more when</td>
<td>Additional data needed regarding</td>
</tr>
</tbody>
</table>

**Drug(s)**: Baloxavir, Chloroquine Phosphate

**AHFS Class**: Antiviral, Antimalarial

**Rationale**: Antiviral active against influenza viruses, In vitro activity against various viruses, including coronaviruses

**Trials or Clinical Experience**: Only limited clinical trial data available to date to evaluate use of baloxavir for treatment of COVID-19, Exploratory, open-label, randomized controlled study at a single center in China (ChiCTR2000029544): 29 adults hospitalized with COVID-19 receiving antiviral treatment with lopinavir/ritonavir, darunavir/copegicst, or umifenovir (Arbidol®), in combination with inhaled interferon-α, were randomized to treatment with baloxavir marboxil (80 mg orally on day 1 and on day 4, and 80 mg orally on day 7 as needed) (n=10), favipiravir (1600 or 2200 mg orally on day 1, followed by 600 mg three times daily for up to 14 days) (n=9), or control (standard antiviral treatment) (n=10). Percentage of pts with viral conversion (2 consecutive tests with undetectable viral RNA results) after 14 days of treatment was 70, 77, and 100% in the baloxavir, favipiravir, and control groups, respectively. Another randomized controlled trial registered in China: 1 CHICCTR2000029548 Protocol for two registered Chinese trials (ChiCTR2000029544, ChiCTR2000029548) specifies an oral baloxavir marboxil dosage of 80 mg on day 1 and on day 4, and another dose of 80 mg on day 7 (as needed); not to exceed 3 total doses.  

**Dosage**: Protocol for two registered Chinese trials (ChiCTR2000029544, ChiCTR2000029548) specifies an oral baloxavir marboxil dosage of 80 mg on day 1 and on day 4, and another dose of 80 mg on day 7 (as needed); not to exceed 3 total doses.  

**Comments**: No data to date support use in the treatment of COVID-19, Efficacy and safety of chloroquine for treatment or prevention of COVID-19 not established, Additional data needed to determine whether in vitro activity against SARS-CoV-2 corresponds with clinical efficacy for treatment or prevention of COVID-19, Additional data needed to substantiate initial reports of efficacy for treatment and identify optimal dose and duration
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<td>Chloroquine</td>
<td>ASHP COVID-19</td>
<td>Has immunomodulatory activity that theoretically could contribute to an anti-inflammatory response in patients with viral infections 1-3, 13, 15-16</td>
<td>Known pharmacokinetics and toxicity profile</td>
<td>a clinical trial is not available or participation not feasible, 1 g on day 1, then 500 mg daily for 4-7 days of total treatment based on clinical evaluation 25</td>
<td>toxicity profile when used in patients with COVID-19</td>
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**Rationale**

**Known pharmacokinetics and toxicity profile**

severe COVID-19 (NCT04323527): The first 81 enrolled pts were randomized 1:1 to receive high-dose chloroquine (600 mg twice daily for 10 days) or lower-dose chloroquine (450 mg twice daily on day 1, then 450 mg once daily on days 2-5); all pts also received azithromycin and ceftriaxone and some also received oseltamivir. An unplanned interim analysis was performed and the high-dose arm of the study was halted because of toxicity concerns, particularly QTc prolongation and ventricular tachycardia, and because more deaths were reported in this arm. By day 13, 16/41 pts (39%) treated with the high-dose regimen had died vs 6/40 (15%) treated with the lower-dose regimen. QTc >500 msec occurred more frequently in the high-dose group (18.9%) than in the lower-dose group (11.1%). The high-dose arm included more pts prone to cardiac complications than the lower-dose arm. Data were insufficient to evaluate efficacy. Study continuing using only the lower dosage. 37

Multiple clinical trials to evaluate chloroquine for the treatment of COVID-19 are registered at clinicaltrials.gov (some listed below): 10

NCT04323527
NCT04328493
NCT04331600
NCT04333628
NCT04353336
NCT04360759
NCT04362332

Several clinical trials to evaluate chloroquine for prevention of COVID-19 in the healthcare setting are registered at clinicaltrials.gov: 10

NCT04303507
NCT04333732
NCT04349371

**Dosage**

- **Oral chloroquine phosphate:** 500 mg twice daily for 7 days (adults 18-65 years weighing >50 kg); 500 mg twice daily on days 1 and 2, then 500 mg once daily on days 3-7 (adults weighing <50 kg) 31

- **Oral chloroquine phosphate:** Initial dose of 600 mg (of chloroquine) followed by 300 mg (of chloroquine) 12 hours later on day 1, then 300 mg (of chloroquine) twice daily on days 2-5 4

**Notes**

- Chloroquine suggested as possible option and included in Chinese guidelines for treatment of COVID-19. 11
- NIH COVID-19 Treatment Guidelines Panel states that clinical data are insufficient to recommend either for or against the use of chloroquine for the treatment of COVID-19. 33
- IDSA recommends that chloroquine be used for the treatment of COVID-19 in the context of a clinical trial. 38
- NIH COVID-19 Treatment Guidelines Panel does not recommend the use of any agents, including chloroquine, for preexposure prophylaxis (PrEP) or post-exposure prophylaxis (PEP) for prevention of SARS-CoV-2 infection outside of clinical trials. 35
- Because chloroquine is associated with QT prolongation, caution is advised in pts at risk for QT prolongation or receiving other drugs associated with arrhythmias; 36, 39 diagnostic testing and monitoring recommended to minimize risk of adverse effects, including drug-induced cardiac effects. 4, 15, 36, 39
- FDA issued a safety alert regarding adverse cardiac effects (e.g., prolonged QT interval, ventricular tachycardia, ventricular fibrillation) reported with use of chloroquine or hydroxychloroquine (either alone or in conjunction with azithromycin or other drugs known to prolong QT interval) in hospital and outpatient settings; FDA cautions against use of chloroquine or hydroxychloroquine outside of a clinical trial or hospital setting and urges healthcare professionals and pts to report adverse effects involving these drugs to FDA MedWatch. 39

**Comments**

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Emergency use authorization (EUA) for chloroquine: FDA issued an EUA that permits distribution of the drug from the strategic national stockpile (SNS) for use only in adults and adolescents weighing 50 kg or more hospitalized with COVID-19 for whom a clinical trial is not available or participation not feasible. \[^{24, 25}\]

To request the drug, healthcare providers should contact local or state health departments; \[^{25}\] distribution to states will be managed by the Office of the Assistant Secretary for Preparedness and Response (ASPR) and FEMA. \[^{29}\]

To mitigate risks of this unapproved use, the EUA includes certain mandatory requirements (including adverse event reporting to FDA MedWatch). \[^{24, 25}\]

FDA states that, based on the totality of scientific evidence available, it is reasonable to believe that the drug may be effective in treating COVID-19 and that, when used under the EUA conditions, known and potential benefits outweigh known and potential risks. \[^{24}\]

Consult the EUA, \[^{24}\] EUA fact sheet for healthcare providers, \[^{25}\] and EUA fact sheet for patients and parent/caregivers \[^{27}\] for additional information.

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<thead>
<tr>
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<tbody>
<tr>
<td>Favipiravir (Avigan®, Favilavir)</td>
<td>8:18.32 Antiviral</td>
<td>Broad-spectrum antiviral with in vitro activity against various viruses, including coronaviruses [^{1-5}] In vitro evidence of activity against SARS-CoV-2 in infected Vero E6 cells reported with high concentrations of the drug [^{1, 2, 16}] Licensed in Japan and China for treatment of influenza [^{2, 4, 6}]</td>
<td>Only very limited clinical trial data available to date to evaluate use of favipiravir in the treatment of COVID-19 Open-label, prospective, randomized, multicenter study in 236 adults with COVID-19 pneumonia in China (ChiCTR2000030254): Favipiravir (1600 mg orally twice daily on day 1, then 600 mg orally twice daily thereafter for 7–10 days) was associated with greater clinical recovery rate at 7 days (61 vs 52%) compared with the control group treated with umifenovir (Arbidol®; 200 mg 3 times daily for 7–10 days). Stratified by disease severity, clinical recovery rate at day 7 in pts with moderate COVID-19 pneumonia was 71% in the favipiravir group vs 56% in the umifenovir group; clinical recovery rate in those with severe to critical COVID-19</td>
<td>A favipiravir dosage of 1600 mg twice daily on day 1, then 600 mg twice daily thereafter for 7–10 days was used in one open-label COVID-19 study [^{6}] A favipiravir dosage of 1600 mg twice daily on day 1, then 600 mg twice daily thereafter for 14 days was used in one open-label COVID-19 study [^{15}] Protocol in one ongoing trial (NCT04336904) for treatment of moderate COVID-19 specifies a favipiravir dosage of 1800 mg twice daily on day 1, then 600 mg three times daily thereafter for up to 14 days [^{7}] Protocol in one ongoing trial (NCT04346628) for treatment of</td>
<td>Emergency use authorization (EUA) for chloroquine: FDA issued an EUA that permits distribution of the drug from the strategic national stockpile (SNS) for use only in adults and adolescents weighing 50 kg or more hospitalized with COVID-19 for whom a clinical trial is not available or participation not feasible. [^{24, 25}] To request the drug, healthcare providers should contact local or state health departments; [^{25}] distribution to states will be managed by the Office of the Assistant Secretary for Preparedness and Response (ASPR) and FEMA. [^{29}] To mitigate risks of this unapproved use, the EUA includes certain mandatory requirements (including adverse event reporting to FDA MedWatch). [^{24, 25}] FDA states that, based on the totality of scientific evidence available, it is reasonable to believe that the drug may be effective in treating COVID-19 and that, when used under the EUA conditions, known and potential benefits outweigh known and potential risks. [^{24}] Consult the EUA, [^{24}] EUA fact sheet for healthcare providers, [^{25}] and EUA fact sheet for patients and parent/caregivers [^{27}] for additional information.</td>
</tr>
</tbody>
</table>

\[^{a}\] FAVIPIRAVIR DOSAGE a

\[^{b}\] Not commercially available in the US

\[^{c}\] Efficacy and safety of favipiravir for treatment of COVID-19 not established

\[^{d}\] Additional data needed to substantiate initial reports of efficacy for treatment of COVID-19 and identify optimal dosage and treatment duration

\[^{e}\] Given the lack of pharmacokinetic and safety data for the high favipiravir dosages proposed for treatment of COVID-19, the drug should be used with caution at such dosages. \[^{19, 20}\] Favipiravir is associated with QT prolongation. \[^{21}\] Some have suggested close cardiac and hepatic monitoring during treatment, as well as monitoring of plasma and tissue concentrations of the drug and, if
Drug(s) | AHFS Class | Rationale | Trials or Clinical Experience | Dosage | Comments
--- | --- | --- | --- | --- | ---

pneumonia was 6% vs 0%, respectively. Twice as many pts in the favipiravir group had severe to critical disease compared with the group receiving umifenovir. 6

In a small, open-label, nonrandomized study in patients with non-severe COVID-19 in China (ChiCTR2000029600), favipiravir (1600 mg orally twice daily on day 1, then 600 mg orally twice daily on days 2–14) (n=35) was associated with decreased median time to viral clearance (4 vs 11 days) and higher improvement rate on chest CT imaging on day 14 (91 vs 62%) compared with the control group receiving lopinavir/ritonavir (n=45); both groups also received aerosolized interferon α-1b. 15

Italy: Randomized, placebo-controlled multicenter trial (NCT04336904) to evaluate efficacy and safety of favipiravir in pts with moderate COVID-19 7

US: Randomized, controlled open-label proof-of-concept trial (NCT04358549) of favipiravir for the treatment of COVID-19 7 10

US: Randomized, open-label trial (NCT04346628) to evaluate efficacy of favipiravir in pts with mild, uncomplicated COVID-19 7

Multiple clinical trials initiated in pts with COVID-19 in China, Japan, and other countries to evaluate favipiravir alone or in combination with other antivirals or other agents (some listed below): 7 9

NCT04310228
NCT04319900
NCT04303299
NCT04333589
NCT04336904
NCT04345419
NCT04351295
NCT04349241
NCT04356495
NCT04358549
NCT04359615

mild COVID-19 specifies a favipiravir dosage of 1800 mg on day 1, then 800 mg twice daily on days 2–10 7

Protocol in one ongoing trial (NCT04349241) for treatment of non-severe COVID-19 specifies a favipiravir dosage of 1600 mg every 12 hours on day 1, then 600 mg every 12 hours on days 2–10 7

Protocol in one ongoing trial (NCT04358549) for treatment of COVID-19 specifies a favipiravir dosage of 1800 mg twice daily on day 1, then 1000 mg twice daily on days 2–14 7

Protocol in one ongoing trial (NCT04373733; PIONEER) for early treatment of suspected or confirmed COVID-19 specified a favipiravir dosage of 1800 mg twice daily on day 1, followed by 800 mg twice daily on days 2–10 7

Because high favipiravir concentrations are required for in vitro activity against SARS-CoV-2, 1, 5, 13 it has been suggested that high favipiravir dosages, like those used in the treatment of Ebola virus disease, should be considered for the treatment of COVID-19. 11, 19, 20 One such favipiravir regimen used in the treatment of Ebola virus disease includes a loading dosage of 6000 mg (doses of 2400 mg, 2400 mg, and 1200 mg given 8 hours apart on day 1), then a maintenance dosage of 1200 mg every 12 hours on days 2–10. 12, 13 One pharmacokinetic simulation model suggested that a dosage of 2400 mg twice daily on day 1, followed by 1600 mg twice daily on days 2–10 should achieve adequate favipiravir trough plasma concentrations and may be more pharmacologically relevant. 19

possible, the active metabolite. 19, 20, 21

Early embryonic deaths and teratogenicity observed in animal studies. Favipiravir is contraindicated in women with known or suspected pregnancy and precautions should be taken to avoid pregnancy during treatment with the drug. 14

If favipiravir is used in pts receiving acetaminophen, the maximum recommended daily dosage of acetaminophen is 3 g 17, 18
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</thead>
<tbody>
<tr>
<td>HIV Protease Inhibitors</td>
<td>8:18.08.08</td>
<td>HIV Protease Inhibitors</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Updated 4/24/20</td>
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<tr>
<td><strong>Lopinavir (LPV):</strong></td>
<td>8</td>
<td>In vitro activity against SARS-CoV-2 in Vero E6 cells; also has in vitro activity against SARS-CoV-1 and MERS-CoV; some evidence of benefit in animal studies for treatment of MERS-CoV Atazanavir (ATV): ATV alone or with ritonavir (ATV/RTV) has in vitro activity against SARS-CoV-2 in Vero E6 cells, human epithelial pulmonary cells (A549) and human monocytes Darunavir (DRV): In one study, DRV with cobicistat had no in vitro activity against SARS-CoV-2 at clinically relevant concentrations in Caco-2 cells; in another study, high DRV concentrations were required for in vitro inhibition of SARS-CoV-2 in Vero E6 cells Nelfinavir (NFV), Saquinavir (SQV), and Tipranavir (TPV): In vitro activity against SARS-CoV-2 in Vero E6 cells</td>
<td><strong>Lopinavir and Ritonavir (LPV/RTV; Kaletra®)</strong> randomized, open-label trial in China in hospitalized adults with severe COVID-19 compared LPV/RTV in conjunction with standard care (99 pts vs standard care alone (100 pts). Primary end point was time to clinical improvement (time from randomization to improvement of two points on a seven-category ordinal scale or hospital discharge, whichever came first). In ITT population, <strong>time to clinical improvement was not shorter with LPV/RTV compared with standard care</strong> (median time to clinical improvement 16 days in both groups); in modified ITT population, median time to clinical improvement 15 days in LPV/RTV group and 16 days in standard care only group. The 28-day mortality rate was numerically lower in LPV/RTV group (19.2% vs 25% in ITT population; 16.7% vs 25% in modified ITT population). Some evidence that LPV/RTV initiation within 12 days after symptom onset is associated with shorter time to clinical improvement. <strong>No significant differences in reduction of viral RNA load, duration of viral RNA detectability, duration of oxygen therapy, duration of hospitalization, or time from randomization to death.</strong> LPV/RTV stopped early in 13 pts because of adverse effects.</td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days</td>
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<tr>
<td><strong>LPV/RTV (COVID-19):</strong></td>
<td></td>
<td>Efficacy for the treatment of COVID-19, with or without other antivirals, not definitely established</td>
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<tr>
<td>Darunavir: DRV:</td>
<td></td>
<td>No data to date to support use in the treatment of COVID-19. Manufacturer states they have no clinical or pharmacologic evidence to support use of DRV/cobicistat for treatment of COVID-19 and initial unpublished results from a study in China indicated that a 5-day regimen of DRV/cobicistat was not effective for treatment of COVID-19</td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for no longer than 10 days with or without interferon (5 million units of interferon-α or equivalent twice daily given in 2 mL of sterile water by nebulization) and with or without ribavirin for up to 10 days</td>
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<tr>
<td>Atazanavir, Nelfinavir, Saquinavir, Tipranavir:</td>
<td></td>
<td>No data to date to support use in the treatment of COVID-19</td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 14 days with ribavirin (4-g oral loading dose, then 1.2 g orally every 8 hours)</td>
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<tr>
<td>Lopinavir (LPV):</td>
<td>9</td>
<td>In vitro activity against SARS-CoV-2 in Vero E6 cells</td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 14 days with ribavirin (various regimens) and/or interferon-α; LPV 400 mg/RTV 100 mg orally twice daily with interferon β1b (0.25 mg/mL sub-Q on alternate days) for 14 days</td>
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<td>Lopinavir and Ritonavir (LPV/RTV; Kaletra®):  Open-label trial in China in hospitalized adults with severe COVID-19 compared LPV/RTV in conjunction with standard care (99 pts vs standard care alone (100 pts). Primary end point was time to clinical improvement (time from randomization to improvement of two points on a seven-category ordinal scale or hospital discharge, whichever came first). In ITT population, time to clinical improvement was not shorter with LPV/RTV compared with standard care (median time to clinical improvement 16 days in both groups); in modified ITT population, median time to clinical improvement 15 days in LPV/RTV group and 16 days in standard care only group. The 28-day mortality rate was numerically lower in LPV/RTV group (19.2% vs 25% in ITT population; 16.7% vs 25% in modified ITT population). Some evidence that LPV/RTV initiation within 12 days after symptom onset is associated with shorter time to clinical improvement. No significant differences in reduction of viral RNA load, duration of viral RNA detectability, duration of oxygen therapy, duration of hospitalization, or time from randomization to death. LPV/RTV stopped early in 13 pts because of adverse effects.</td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days</td>
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<tr>
<td>Lopinavir (LPV):</td>
<td>2</td>
<td>In vitro activity against SARS-CoV-2 in Vero E6 cells</td>
<td></td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days</td>
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<td>Lopinavir (LPV):</td>
<td>7</td>
<td>In vitro activity against SARS-CoV-2 in Vero E6 cells</td>
<td></td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days</td>
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<td>Lopinavir (LPV):</td>
<td>9</td>
<td>In vitro activity against SARS-CoV-2 in Vero E6 cells</td>
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<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days</td>
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<td>Darunavir: DRV:</td>
<td>17</td>
<td>In one study, DRV with cobicistat had no in vitro activity against SARS-CoV-2 at clinically relevant concentrations in Caco-2 cells; in another study, high DRV concentrations were required for in vitro inhibition of SARS-CoV-2 in Vero E6 cells</td>
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<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days</td>
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<tr>
<td>Lopinavir (LPV):</td>
<td>18</td>
<td>In vitro activity against SARS-CoV-2 in Vero E6 cells</td>
<td></td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days</td>
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<tr>
<td>Lopinavir (LPV):</td>
<td>19</td>
<td>In vitro activity against SARS-CoV-2 in Vero E6 cells</td>
<td></td>
<td><strong>LPV/RTV (COVID-19):</strong> LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days</td>
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**Dosagea:**
- **LPV/RTV (COVID-19):** LPV 400 mg/RTV 100 mg orally twice daily for 10-14 days
- **LPV/RTV (COVID-19):** LPV 400 mg/RTV 100 mg orally twice daily for no longer than 10 days with or without interferon (5 million units of interferon-α or equivalent twice daily given in 2 mL of sterile water by nebulization) and with or without ribavirin for up to 10 days
- **LPV/RTV (COVID-19):** LPV 400 mg/RTV 100 mg orally twice daily for 14 days with ribavirin (4-g oral loading dose, then 1.2 g orally every 8 hours)
- **LPV/RTV (COVID-19):** LPV 400 mg/RTV 100 mg orally twice daily for 14 days with ribavirin (various regimens) and/or interferon-α; LPV 400 mg/RTV 100 mg orally twice daily with interferon β1b (0.25 mg/mL sub-Q on alternate days) for 14 days

**Comments:***

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<tr>
<th>Drug(s)</th>
<th>AHFS Class</th>
<th>Rationale</th>
<th>Trials or Clinical Experience</th>
<th>Dosage*</th>
<th>Comments</th>
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<tr>
<td>Hydroxychloroquine</td>
<td>8:30.08</td>
<td>Antimalarial</td>
<td>In vitro activity against various viruses, including coronaviruses. In vitro activity against SARS-CoV-2 in infected Vero E6 cells reported; may be more potent than chloroquine in vitro, but some data are conflicting and additional study needed.</td>
<td>Only limited clinical trial data available to date to evaluate use of hydroxychloroquine for treatment or prevention of COVID-19</td>
<td>Optimal dosage and duration of treatment not known. Various dosages recommended or being investigated for treatment of COVID-19. Oral hydroxychloroquine sulfate dosage suggested in the EUA: For treatment of hospitalized adults and adolescents weighing 50 kg or more when a clinical trial is not available or Efficacy and safety of hydroxychloroquine for treatment or prevention of COVID-19 not established. Additional data needed to determine whether in vitro activity against SARS-CoV-2 corresponds with clinical efficacy for treatment or prevention of COVID-19. Additional data needed to substantiate initial reports of efficacy for treatment.</td>
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<td>Hydroxychloroquine</td>
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<td>Has immunomodulatory activity that theoretically could contribute to an anti-inflammatory response in patients with viral infections. Known pharmacokinetics and toxicity profile. Hydroxyl analog of chloroquine with similar mechanisms of action and adverse effects, may have more favorable dose-related toxicity profile than chloroquine, but cardiotoxicity (e.g., prolonged QT interval) is a concern with both drugs. Received hydroxychloroquine sulfate (400 mg daily for 5 days) with conventional treatments and 15 pts received conventional treatments alone, both groups received interferon and most pts also received umifenovir (Arbidol®) or LPV/RTV. Primary end point was conversion to negative PCR in pharyngeal swabs on day 7. Negative PCR reported at day 7 in 13 pts (86.7%) treated with hydroxychloroquine and 14 pts (93.3%) not treated with the drug (data unclear for 3 pts); median duration from hospitalization to negative conversion and to temperature normalization were similar in both groups; evidence of radiologic progression on CT in 5 pts treated with the drug and 7 pts not treated with the drug (all pts showed improvement at follow-up). Hydroxychloroquine randomized, parallel-group study in adults in China (ChiCTR2000029559): 31 pts with COVID-19 and pneumonia received hydroxychloroquine sulfate (200 mg twice daily for 5 days) and standard treatment (O2, antiviral agents, antibacterial agents, immunoglobulin, with or without corticosteroids) and 31 other pts received standard treatment alone (control group). Exclusion criteria included severe and critical illness. Pts assessed at baseline and 5 days after treatment initiation for time to clinical recovery (TTCR, defined as normalization of fever and cough relief maintained for &gt;72 hours), clinical characteristics, and changes on chest CT. It was concluded that hydroxychloroquine was associated with symptom relief since time to fever normalization was shorter in hydroxychloroquine group (2.2 days) vs control group (3.2 days), time to cough remission was shorter in hydroxychloroquine group, and pneumonia improved in 25/31 pts (80.6%) in hydroxychloroquine group vs 17/31 pts (54.8%) in control group. Total of 4 pts progressed to severe illness (all in the participation not feasible, 800 mg on day 1, then 400 mg daily for 4-7 days of total treatment based on clinical evaluation. Oral hydroxychloroquine sulfate: 400 mg twice daily on day 1, then 200 mg twice daily on days 2-5. Oral hydroxychloroquine sulfate: 400 mg once or twice daily for 5-10 days. Oral hydroxychloroquine sulfate: 600 mg twice daily on day 1, then 400 mg daily on days 2-5. Oral hydroxychloroquine sulfate: 100-200 mg twice daily for 5-14 days. Oral hydroxychloroquine sulfate: 200 mg 3 times daily for 10 days. and identify optimal dose and duration. Additional data needed before any conclusions can be made regarding possible benefits and safety of using hydroxychloroquine with azithromycin. (See Azithromycin in this Evidence Table.) Additional data needed regarding toxicity profile when used in patients with COVID-19. Hydroxychloroquine suggested as possible option and included in Chinese guidelines for treatment of COVID-19. NIH COVID-19 Treatment Guidelines Panel states that clinical data are insufficient to recommend either for or against use of hydroxychloroquine for the treatment of COVID-19. IDSA recommends that hydroxychloroquine be used for the treatment of COVID-19 in the context of a clinical trial. NIH COVID-19 Treatment Guidelines Panel recommends against the use of a combined regimen of hydroxychloroquine and azithromycin for the treatment of COVID-19, except in the context of a clinical trial. IDSA recommends that a combined regimen of hydroxychloroquine and azithromycin be used for the treatment of COVID-19 only in the context of a clinical trial. NIH COVID-19 Treatment Guidelines Panel does not recommend the use of any agents, including hydroxychloroquine, for preexposure prophylaxis (PrEP) or postexposure prophylaxis (PEP) for prevention of SARS-CoV-2 infection outside of clinical trials. Because hydroxychloroquine is associated with QT prolongation, caution is</td>
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control group). Note: This study did not include pts with severe disease and pts received other anti-infectives in addition to hydroxychloroquine. At study entry, 9 pts without fever and 9 pts without cough were included in hydroxychloroquine group and 14 pts without fever and 16 pts without cough were included in control group; unclear how these pts were addressed in TTCR calculations. Although initial registered study protocol specified 2 different hydroxychloroquine treatment groups and a placebo group (each with 100 pts) and primary end points of time to negative nucleic acid and T-cell recovery, data provided only for certain clinical symptoms in 62 pts without severe disease and PCR results not reported.

**Hydroxychloroquine with azithromycin, open-label, nonrandomized study in France (Gautret et al):** Preliminary data from an ongoing study in hospitalized pts with confirmed COVID-19 was used to assess efficacy of hydroxychloroquine used alone or with azithromycin; untreated pts were used as a negative control. The primary end point was negative PCR results in nasopharyngeal samples at day 6. Data from 14 pts treated with hydroxychloroquine (200 mg 3 times daily for 10 days), 6 pts treated with hydroxychloroquine and azithromycin (500 mg on day 1, then 250 mg daily on days 2-5), and 16 pts in the control group were analyzed. At day 6, 8/14 (57%) in the hydroxychloroquine group, 6/6 (100%) in the hydroxychloroquine and azithromycin group, and 2/16 (12.5%) in the control group had negative PCR results. At day 8, a positive PCR was reported in a pt treated with both drugs who had tested negative at day 6. Note: This was a small nonrandomized study that didn’t appear to be designed to compare hydroxychloroquine vs hydroxychloroquine and azithromycin (pts received antibiotics advised in pts at risk for QT prolongation or receiving other drugs associated with arrhythmias). 

FDA issued a safety alert regarding adverse cardiac effects (e.g., prolonged QT interval, ventricular tachycardia, ventricular fibrillation) reported with use of chloroquine or hydroxychloroquine (either alone or in conjunction with azithromycin or other drugs known to prolong QT interval) in hospital and outpatient settings; FDA cautions against use of chloroquine or hydroxychloroquine outside of a clinical trial or hospital setting and urges healthcare professionals and pts to report adverse effects involving these drugs to FDA MedWatch.

**Emergency use authorization (EUA) for hydroxychloroquine:** FDA issued an EUA that permits distribution of the drug from the strategic national stockpile (SNS) for use only in adults and adolescents weighing 50 kg or more hospitalized with COVID-19 for whom a clinical trial is not available or participation not feasible. To request the drug, healthcare providers should contact local or state health departments; distribution to states will be managed by the Office of the Assistant Secretary for Preparedness and Response (ASPR) and FEMA. To mitigate risks of this unapproved use, the EUA includes certain mandatory requirements (including adverse event reporting to FDA MedWatch). FDA states that, based on the totality of scientific evidence available, it is reasonable to believe that the drug may be effective in treating COVID-19 and that, when used under the EUA conditions, known and potential benefits outweigh known and potential risks.
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<th>Drug(s)</th>
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<td>to prevent bacterial superinfection based on clinical judgment. Data on disease severity was unclear (some asymptomatic pts were included when study initiated) and information on disease progression and clinical outcomes was not presented. Hydroxychloroquine with azithromycin open-label, uncontrolled study in France (Molina et al): 11 adults hospitalized with COVID-19 received hydroxychloroquine (600 mg daily for 10 days) and azithromycin (500 mg on day 1, then 250 mg daily on days 2-5). At time of treatment initiation, 8/11 pts had significant comorbidities associated with poor outcomes and 10/11 had fever and received O₂. Within 5 days, 1 pt died and 2 transferred to ICU; the regimen discontinued in 1 pt after 4 days because of prolonged QT interval. Nasopharyngeal samples were still PCR positive at days 5 and 6 in 8/10 pts tested. <strong>Note:</strong> In this small uncontrolled study, hydroxychloroquine and azithromycin regimen did not result in rapid viral clearance or provide clinical benefit. Hydroxychloroquine with azithromycin uncontrolled, retrospective, observational study in France (Gautret et al): 80 adults with confirmed COVID-19 (including 6 pts included in a previous study by the same group) were treated with hydroxychloroquine (200 mg 3 times daily for 10 days) and azithromycin (500 mg on day 1, then 250 mg daily on days 2-5). Majority (92%) were considered low risk for clinical deterioration (low national early warning score for COVID-19 based on age, respiratory rate, O₂ saturation, temperature, BP, pulse, level of consciousness); only 15% had fever; 4 pts were asymptomatic carriers; mean time from onset of symptoms to treatment initiation was 4.9 days. Clinical outcome, contagiousness as assessed by nasopharyngeal PCR assay and culture, and length of stay in infectious disease (ID) unit were...</td>
<td>**</td>
<td>healthcare providers, ** and EUA fact sheet for patients and parent/caregivers ** for additional information.</td>
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<td>Drug(s)</td>
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<td>hydroxychloroquine (with or without azithromycin)</td>
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<td>evaluated in pts who were treated for at least 3 days and followed for at least 6 days. Favorable outcome was reported for 81.3%; 15% required O₂; 3 pts transferred to ICU; 1 pt died; mean time to discharge from ID unit was 4.1 days. At day 8, PCR results were negative in 93% of those tested; at day 5, viral cultures were negative in 97.5% of those tested. <strong>Note:</strong> Almost all pts were considered low risk for clinical deterioration (including 4 pts described as asymptomatic carriers) and it is unclear how many would have had spontaneous conversion to negative nasopharyngeal samples during same time frame. Although 80 pts were enrolled, PCR results available for fewer pts beginning on day 3 and only 60 pts represented in day 6 data. This was an uncontrolled study and data presented cannot be used to determine whether a regimen of hydroxychloroquine with azithromycin provides benefits in terms of disease progression or decreased infectiousness, especially for pts with more severe disease.</td>
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Azithromycin did not reduce the risk of mechanical ventilation. 40  **Note:** The pt population included only elderly males 59-75 years of age, many with significant comorbidities. This analysis did not look at efficacy measures.

**Efficacy measures:** Initial studies evaluating hydroxychloroquine based efficacy of the drug on negative conversion in nasopharyngeal samples at day 6 or 7. 7, 18 RT-PCR tests using upper and lower respiratory specimens (including nasopharyngeal and oropharyngeal swabs) are recommended for diagnosis of COVID-19; 10, 21 however, dynamics of SARS-CoV-2 in infected patients (untreated or treated) and presence of the virus at various body sites over the course of infection have not been fully determined. 22, 23

Multiple clinical trials to evaluate hydroxychloroquine for treatment of COVID-19 are registered at clinicaltrials.gov (some listed below): 10
- NCT04329923
- NCT04332991
- NCT04334967
- NCT04335552
- NCT04341727
- NCT04345692
- NCT04350450
- NCT04351620
- NCT04353037
- NCT04362332

Multiple clinical trials to evaluate hydroxychloroquine for prevention of COVID-19 in the healthcare setting or in household contacts of pts with the disease are registered at clinicaltrials.gov (some listed below): 10
- NCT04303507
- NCT04318015
- NCT04318444
- NCT04328961
- NCT04330144
<table>
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<tr>
<td>Neuraminidase inhibitors (e.g., oseltamivir)</td>
<td>8:18.28</td>
<td>Antivirals active against influenza viruses</td>
<td>In a retrospective case series of 99 patients with COVID-19 at single center in Wuhan from 1/1/20 to 1/20/20, 76% of patients received antiviral treatment, including oseltamivir (75 mg orally every 12 hours). At the time of evaluation, 58% of patients remained hospitalized, 31% had been discharged, and 11% had died.</td>
<td>Dosage of oseltamivir in the case series of 99 patients was 75 mg orally every 12 hours. 1</td>
<td>No data to date support use in the treatment of COVID-19</td>
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<td>Remdesivir</td>
<td>8:18.32</td>
<td>Antiviral</td>
<td>Various clinical trials initiated in US, China, and other countries</td>
<td>Optimal dosage and duration of treatment not known 25, 26</td>
<td>Not commercially available; most promising direct-acting antiviral (DAA) currently being investigated for COVID-19 Efficacy and safety of remdesivir for treatment of COVID-19 not established NIH COVID-19 Treatment Guidelines Panel states that clinical data are insufficient to recommend either for or against the use of remdesivir for the treatment of COVID-19 20</td>
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Updated 5/15/20

Updated 5/8/20
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<th>Drug(s)</th>
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| 12 hours after virus inoculation was associated with some benefits (lower disease severity scores, fewer pulmonary infiltrates, lower virus titers in bronchoalveolar lavage samples) compared with vehicle control; remdesivir treatment did not reduce viral loads or infectious virus titers in nose, throat, or rectal swabs compared with vehicle control.  
In vitro activity against SARS-CoV and MERS-CoV; active in animal models of SARS and MERS; prevented MERS in Rhesus macaques when given before infection and provided benefits when given after animal already infected.  
Pharmacokinetic data available from evaluations for Ebola.  
158 pts treated with remdesivir and 78 pts treated with placebo; 32% of pts also received interferon α-2b, 28% also received LPV/RTV, and 66% also received corticosteroids during hospitalization. Median time to clinical improvement was not significantly different in remdesivir group (21 days) vs placebo group (23 days); 28-day mortality rate was similar in both groups (14 vs 13%). When remdesivir was initiated within 10 days of symptom onset, median time to clinical improvement was numerically shorter (but not statistically significant) compared with placebo group (18 vs 23 days). Duration of invasive mechanical ventilation was numerically shorter (but not statistically significant) in remdesivir group; only a small percentage of pts (0.4%) were on invasive mechanical ventilation at time of enrollment. Remdesivir did not result in significant reduction in SARS-CoV-2 viral load in nasopharyngeal, oropharyngeal, and sputum samples. Remdesivir was discontinued in 18 pts (12%) because of adverse effects. Note: Enrollment was terminated before the pre-specified number of pts was attained (lack of available pts); trial was insufficiently powered to detect assumed differences in clinical outcome.  
Phase 3 randomized, open-label trial in hospitalized adults with severe COVID-19 (NCT04292899) sponsored by the manufacturer (Gilead): Initial study protocol designed to evaluate safety and antiviral activity of 5- and 10-day regimens of remdesivir (200 mg IV on day 1, followed by 100 mg once daily for total of 5 or 10 days) in conjunction with standard of care in pts not receiving mechanical ventilation; protocol subsequently modified to add extension arms to evaluate safety and efficacy of 10-day regimen of remdesivir in conjunction with standard of care in pts who are or are not receiving mechanical ventilation.  
Manufacturer announced that data available for the initial 397 pts not requiring mechanical ventilation at 200 mg IV on day 1, then 100 mg IV daily on days 2-10 (arm 1).  
NIH adaptive study protocol: 200 mg IV on day 1, then 100 mg IV for duration of hospitalization up to 10 days total.  
Compassionate use access protocol: 200 mg IV on day 1, then 100 mg IV on days 2-10.  
Emergency use authorization (EUA) for adults and children weighing 40 kg or more: Loading dose of 200 mg by IV infusion on day 1, followed by 100 mg by IV infusion once daily on days 2-10 (for pts requiring invasive mechanical ventilation and/or ECMO) or followed by 100 mg by IV infusion once daily on days 2-5 with option to extend treatment up to day 10 if needed (for pts not requiring mechanical ventilation and/or ECMO).  
Emergency use authorization (EUA) for children weighing 3.5 to less than 40 kg: 5 mg/kg by IV infusion on day 1, followed by 2.5 mg/kg by IV infusion once daily on days 2-10 (for pts requiring invasive mechanical ventilation and/or ECMO) or followed by 2.5 mg/kg by IV infusion once daily on days 2-5 with option to extend treatment up to day 10 if needed (for pts not requiring mechanical ventilation and/or ECMO).  
Emergency use authorization (EUA) for remdesivir: FDA issued an EUA on May 1, 2020 that permits use of the drug for the treatment of COVID-19 only in hospitalized adults and children with suspected or laboratory-confirmed COVID-19 who have severe disease (defined as oxygen saturation [SpO₂] 94% or lower on room air or requiring supplemental oxygen, mechanical ventilation, or ECMO) and requires that the drug be administered by a healthcare provider in an inpatient hospital setting via IV infusion at dosages recommended in the EUA. Distribution of remdesivir under this EUA is controlled by the US government for use consistent with the terms and conditions of the EUA. The manufacturer (Gilead) donated remdesivir for use under the EUA; distribution to hospitals and other healthcare facilities is being directed by the HHS Office of the Assistant Secretary for Preparedness and Response (ASPR) in collaboration with state health departments. To request remdesivir for use under the EUA, healthcare providers should contact their state health departments. The EUA requires that healthcare facilities and healthcare providers administering remdesivir comply with certain mandatory record keeping and reporting requirements (including adverse event reporting to FDA MedWatch). Consult the EUA fact sheet for healthcare providers and EUA fact sheet for patients and parent/caregivers for additional information.
study entry (200 received a 5-day regimen and 197 received a 10-day regimen) indicate similar clinical improvement with both treatment durations. **Time to clinical improvement for 50% of pts was 10 days in the 5-day treatment group vs 11 days in the 10-day treatment group.** At day 14, 129/200 pts (64.5%) in the 5-day group and 106/197 pts (53.8%) in the 10-day group achieved clinical recovery. Pts who received remdesivir within 10 days of symptom onset had improved outcomes compared with those treated after more than 10 days of symptoms. **Note:** Data regarding this initial pt population (e.g., disease severity and comorbidities at study enrollment, additional supportive treatment received) not provided to date.

**Phase 3 randomized, open-label trial in pts with moderate COVID-19 (NCT04292730)** sponsored by the manufacturer (Gilead) is evaluating safety and antiviral activity of 5- and 10-day regimens of remdesivir in conjunction with standard of care compared with standard of care alone. **Note:** Data regarding this initial pt population (e.g., disease severity and comorbidities at study enrollment, additional supportive treatment received) not provided to date.

**Phase 3 adaptive, randomized, placebo-controlled trial (NCT04280705; ACTT) in hospitalized adults sponsored by NIAID:** Pts received remdesivir (200 mg IV on day 1, then 100 mg once daily for duration of hospitalization up to 10 days total) or placebo. **Sponsor announced** that preliminary data analysis (total of 1063 pts) indicated **shorter median time to recovery in remdesivir group (11 days) vs placebo group (15 days)** and suggested that remdesivir treatment may have provided a survival benefit (mortality rate 8% in remdesivir group vs 11.6% in placebo group). **Note:** Data regarding the pt population (e.g., disease severity and comorbidities at study enrollment, time to initiation of treatment after symptom onset, additional supportive treatment received) not provided to date.

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<td>study entry (200 received a 5-day regimen and 197 received a 10-day regimen) indicate similar clinical improvement with both treatment durations. <strong>Time to clinical improvement for 50% of pts was 10 days in the 5-day treatment group vs 11 days in the 10-day treatment group.</strong> At day 14, 129/200 pts (64.5%) in the 5-day group and 106/197 pts (53.8%) in the 10-day group achieved clinical recovery. Pts who received remdesivir within 10 days of symptom onset had improved outcomes compared with those treated after more than 10 days of symptoms. <strong>Note:</strong> Data regarding this initial pt population (e.g., disease severity and comorbidities at study enrollment, additional supportive treatment received) not provided to date.</td>
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<td><strong>Expanded access IND protocol</strong> (NCT04323761): The manufacturer (Gilead) established a protocol for emergency access to remdesivir for the treatment of severe acute COVID-19 in hospitalized adults and children 12 years of age or older.</td>
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<td><strong>Compassionate use access</strong>: The manufacturer (Gilead) is transitioning from individual compassionate use requests to an expanded access program for emergency access to the drug for severely ill pts with confirmed COVID-19. New individual compassionate use requests cannot be accepted, with the possible exception of requests for pregnant women and children &lt;18 years of age with confirmed infections and severe manifestations of the disease. <a href="https://rdvcu.gilead.com/">17</a></td>
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<td>Compassionate use access (NCT04302766): May be available for DoD personnel through treatment IND protocol sponsored by US Army Medical Research and Development Command.</td>
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<td><strong>Data from the manufacturer’s compassionate use program</strong>: Preliminary data are available for a cohort of 53 adults from multiple sites in the US, Italy, Japan, and other countries who were hospitalized with severe COVID-19 and received treatment with remdesivir; 40 pts received the full 10-day regimen (200 mg IV on day 1, then 100 mg IV on days 2-10), 10 pts received 5-9 days and 3 pts received less than 5 days of treatment with the drug. At baseline, 30 pts (57%) were receiving mechanical ventilation and 4 (18%) were receiving extracorporeal membrane oxygenation (ECMO). Over a median follow-up of 18 days after first dose, 36 pts (68%) showed clinical improvement based on oxygen-support status and 8 pts (15%) worsened. There were 7 deaths (13%), including 6 pts receiving invasive ventilation. Adverse effects (e.g., increased hepatic enzymes, diarrhea, rash, renal impairment, hypotension) were</td>
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reported in 32 pts (60%); 12 pts (23%) had serious adverse effects (e.g., multiple organ dysfunction syndrome, septic shock, acute kidney injury, hypotension); 4 pts (8%) discontinued the drug because of adverse effects. Note: Data presented for this small cohort of pts offers only limited information regarding efficacy and safety of remdesivir for treatment of COVID-19. There was no control group and, although supportive therapy could be provided at the discretion of the clinician, it is unclear whether pts at any of the various study sites also received other therapeutic agents being used for treatment of COVID-19. In addition, data were not presented regarding the effects of remdesivir on viral load.

Adaptive, randomized, double-blind trial to compare a regimen of remdesivir alone vs a regimen of remdesivir with baricitinib (ACTT2): This next iteration of NIAID’s Adaptive COVID-19 Treatment Trial (ACTT) will evaluate possible benefits of using baricitinib (a Janus kinase [JAK] inhibitor) in conjunction with remdesivir in hospitalized adults with laboratory-confirmed SARS-CoV-2 infection and evidence of lung involvement (abnormal chest x-rays, need for supplemental oxygen or mechanical ventilation). Pts will be randomized 1:1 to receive remdesivir (200 mg IV on day 1, then 100 mg IV once daily for the duration of hospitalization or up to 10 days total) or the same remdesivir dosage given with oral baricitinib (4 mg once daily for the duration of hospitalization or up to 14 days total).
<table>
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<tr>
<th>Drug(s)</th>
<th>AHFS Class</th>
<th>Rationale</th>
<th>Trials or Clinical Experience</th>
<th>Dosage</th>
<th>Comments</th>
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<tbody>
<tr>
<td>Umifenovir (Arbidol®)</td>
<td>8:18.92 Antiviral</td>
<td>Broad-spectrum antiviral with in vitro activity against various viruses, including coronaviruses and SARS-CoV-2. Licensed in China, Russia, Ukraine, and possibly other countries for prophylaxis and treatment of influenza.</td>
<td>Retrospective cohort study in 50 adults with COVID-19 in China suggests better viral suppression with umifenovir vs LPV/RTV. All pts received conventional therapy, including interferon-α-2b. At 7 days after hospital admission, SARS-CoV-2 was undetectable in 50% of pts treated with umifenovir vs 23.5% treated with LPV/RTV; at 14 days, viral load undetectable in all pts treated with umifenovir vs 44.1% treated with LPV/RTV. Duration of positive SARS-CoV-2 RNA positive test was shorter with umifenovir vs LPV/RTV.</td>
<td>Dosage recommended for treatment of COVID-19 in China: Adults, 200 mg orally 3 times daily for no more than 10 days.</td>
<td>Not commercially available in the US</td>
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<td></td>
<td>Updated 5/8/20</td>
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<td><em>Retrospective cohort study</em> in 33 adults with COVID-19 in China suggests more favorable outcome with LPV/RTV plus umifenovir vs LPV/RTV alone: Primary end point was negative conversion in nasopharyngeal samples and progression or improvement of pneumonia. At 7 days, SARS-CoV-2 undetectable in nasopharyngeal specimens in 12/16 pts (75%) treated with LPV/RTV plus umifenovir vs 6/17 pts (35%) treated with LPV/RTV alone; at 14 days, undetectable in 15/16 pts (94%) treated with both drugs vs 9/17 pts (53%) treated with LPV/RTV alone. At 7 days, chest CT scans were improving in 11/16 pts (69%) treated with both drugs vs 5/17 pts (29%) treated with LPV/RTV alone.</td>
<td><em>Dosage used or being investigated in COVID-19 clinical trials</em>: 200 mg orally 3 times daily for duration of 7-10 days or longer.</td>
<td>Included in some guidelines for treatment of COVID-19. Efficacy for the treatment of COVID-19 not established.</td>
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</table>
When favipiravir was compared with umifenovir, clinical recovery rate was greater in those treated with favipiravir than in those treated with umifenovir.  

(See Favipiravir in this Evidence Table.)

**Randomized, single-center, partially blind-ed trial in China (NCT0425885)** evaluated efficacy of umifenovir in conjunction with standard care vs LPV/RTV in conjunction with standard care vs standard care without an antiviral in hospitalized adults with mild/moderate COVID-19. Data for the 86 enrolled pts suggest no difference in mean time for positive-to-negative conversion of SARS-CoV-2 nucleic acid in respiratory specimens and no difference in clinical outcomes between pts treated with umifenovir or LPV/RTV compared with no antiviral therapy

**NCT04260594 (not yet recruiting):** Randomized, open-label trial evaluating efficacy and safety of umifenovir in conjunction with standard of care in adults with COVID-19

## SUPPORTING AGENTS

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<tr>
<th>Drug(s)</th>
<th>AHFS Class</th>
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<th>Trials or Clinical Experience</th>
<th>Dosage</th>
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<tbody>
<tr>
<td>Anakinra</td>
<td>92:36 Disease-modifying Anti-rheumatic Drug</td>
<td>Recombinant human interleukin-1 (IL-1) receptor antagonist; may potentially combat cytokine release syndrome (CRS) symptoms in severely ill patients</td>
<td>Currently no known published clinical trial evidence supporting efficacy or safety of anakinra in treating COVID-19</td>
<td>Various dosage regimens are being studied</td>
<td>Insufficient clinical data to recommend either for or against use in the treatment of COVID-19</td>
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<td>Updated 4/24/20</td>
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<td>Encouraging preliminary results reported in China with another disease-modifying antirheumatic drug, tocilizumab</td>
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<td>Safety profile well established in patients with sepsis and has been studied extensively in pediatric patients with rheumatologic conditions</td>
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<td><strong>Italy:</strong> Phase 3 randomized, open-label, multicenter trial (NCT04324021) initiated by the manufacturer (Swedish Orphan Biovitrum) to evaluate efficacy and safety of anakinra or emapalumab with standard of care in reducing hyperinflammation and respiratory distress in patients with COVID-19 is recruiting</td>
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<td><strong>(Note:</strong> Anakinra is approved only for subcutaneous administration in the U.S.)</td>
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</table>
### Drug(s) | AHFS Class | Rationale | Trials or Clinical Experience | Dosage* | Comments
--- | --- | --- | --- | --- | ---
Ascorbic acid | 88:12 Vitamin C | Antioxidant and cofactor for numerous physiologic reactions; may support host defenses against infection and protect host cells against infection-induced oxidative stress | Other noncomparative, open-label trials are recruiting in Greece (NCT04356366, NCT04339712) and Belgium (NCT04330638) | Various dosages of IV ascorbic acid used in COVID-19 studies; 50 mg/kg IV every 6 hours for 4 days used in NCT03680274 | Current data not specific to COVID-19; additional study needed

**Updated 5/6/20**

**IV ascorbic acid:**
- Phase 3 randomized, blinded, placebo-controlled trial (NCT03680274; LOVIT) evaluating effect of high-dose IV ascorbic acid on mortality and persistent organ dysfunction in septic ICU patients (including COVID-19 patients); other clinical trials of high-dose IV ascorbic acid for treatment of COVID-19 also registered: NCT04264533, NCT04323514, NCT04363216, NCT04357782, NCT04344184

**Oral ascorbic acid:**
- Randomized, open-label study (NCT04324728; COVIDAtoZ) initiated to evaluate oral ascorbic acid (8 g daily), zinc, or both in combination in symptomatic outpatients receiving a positive COVID-19 test result
- Included at lower dosages as an active or placebo-equivalent comparator (control) in other COVID-19 prevention or treatment studies
- Included as a component of some hydroxychloroquine-based combination regimens being studied for prevention or treatment of COVID-19

**Other infections:**
- Sepsis: Meta-analysis of several small studies suggested beneficial effects from IV ascorbic acid; however, primary end points not improved in CITRIS-ALI study (NCT02106975) in patients with sepsis and ARDS or in VITAMINS study (NCT03333278) in patients with septic shock; additional studies under way

**Note:** May interfere with laboratory tests based on oxidation-reduction reactions (e.g., blood and urine glucose testing, nitrite and bilirubin concentrations, leukocyte counts). Manufacturer states to delay oxidation-reduction reaction-based tests until 24 hours after infusion, if possible.
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<th>Drug(s)</th>
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<tr>
<td>Azithromycin</td>
<td>8:12.12 Macrolides</td>
<td>Antibacterial with some in vitro activity against some viruses (e.g., influenza A H1N1, Zika)</td>
<td>Pneumonia: Limited study data available regarding ascorbic acid (oral) in hospitalized patients with pneumonia. Common cold: Effect of oral supplementation studied extensively; decreases duration of symptoms, may decrease incidence of common cold in individuals under heavy physical stress but not in overall population.</td>
<td>Adjunctive therapy in certain respiratory viral infections: Although contradictory results reported, some evidence of beneficial immunomodulatory or anti-inflammatory effects when used in pts with some viral infections (e.g., influenza). However, in a retrospective cohort study in critically ill pts with laboratory-confirmed MERS, there was no statistically significant difference in 90-day mortality rates or clearance of MERS-CoV RNA between those who received macrolide therapy and those who did not.</td>
<td>Adjunctive treatment in certain viral infections: 500 mg once daily has been used.</td>
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Updated 4/24/20

Updated 5-15-20. The current version of this document can be found on the ASHP COVID-19 Resource Center. This work is licensed under a Creative Commons Attribution-NonCommercial 4.0 International License.
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<tr>
<th>Drug(s)</th>
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<th>Trials or Clinical Experience</th>
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<tr>
<td>Baricitinib</td>
<td>92:36 Disease-modifying Anti-rheumatic Drug</td>
<td>Janus kinase (JAK) 1 and 2 inhibitor; disrupts regulators of endocytosis (AP2-associated protein kinase 1 [AAK1] and cyclin G-associated kinase [GAK]), which may help reduce viral entry and inflammation; also may interfere with intracellular virus particle assembly (^1, 2)</td>
<td>Currently no known published controlled clinical trial evidence supporting efficacy or safety in patients with COVID-19</td>
<td>Therapeutic dosages of baricitinib (2 or 4 mg orally once daily) are sufficient to inhibit AAK1 (^1, 2)</td>
<td>Minimal interaction with CYP enzymes and drug transporters and low protein binding of baricitinib allow for combined use with antiviral agents and other drugs (^4, 14)</td>
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<td>Updated 5/15/20</td>
<td>Ability to inhibit a variety of proinflammatory cytokines, including interferon, has been raised as a possible concern with the use of JAK inhibitors in the management of hyperinflammation resulting from viral infections such as COVID-19 (^5)</td>
<td></td>
<td>Dosage information not yet available (see Trials or Clinical Experience)</td>
<td>NIH COVID-19 Treatment Guidelines Panel recommends against use of JAK inhibitors for the treatment of COVID-19 except in the context of a clinical trial; the panel states that at present the broad immunosuppressive effect of JAK inhibitors outweighs the potential for benefit (^11)</td>
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\(^a\) NIH COVID-19 Treatment Guidelines Panel recommends against use of JAK inhibitors for the treatment of COVID-19 except in the context of a clinical trial; the panel states that at present the broad immunosuppressive effect of JAK inhibitors outweighs the potential for benefit.
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<tr>
<td>Colchicine</td>
<td>92:16 Antigout Agents</td>
<td>Exerts broad anti-inflammatory and immunomodulatory effects through multiple mechanisms, including inhibition of NOD-like receptor protein 3 (NLRP3) inflammasome assembly and disruption of cytoskeletal functions through inhibition of microtubule polymerization. May combat the hyper-inflammatory state of COVID-19 (e.g., cytokine storm) by suppressing proinflammatory cytokines and chemokines. NLRP3 inflammasome activation results in release of interleukins, including IL-1β. In experimental models of acute respiratory distress syndrome/acute lung injury (ARDS/ALI), the NLRP3 inflammasome had a major role in the development of lung injury.</td>
<td>Adaptive phase 2/3 clinical trial: Open-label study planned to evaluate safety and efficacy of baricitinib in hospitalized patients with COVID-19 (NCT04340232). Other planned clinical trials will evaluate baricitinib in combination with or without an antiviral agent for the treatment of COVID-19 (NCT04346147, NCT04320277, NCT04345289, NCT04321993). Minimal anecdotal experience and no clinical trial data reported to date in COVID-19. Phase 3, randomized, double-blind, placebo-controlled study (NCT04322682; COL-CORONA) initiated in adults with COVID-19 and at least one high-risk criterion to evaluate effect of colchicine on mortality, hospitalization rate, and need for mechanical ventilation; study excludes enrollment of currently hospitalized patients; enrollment target is approximately 6000 pts. Other registered randomized, open-label, parallel-group studies (not yet recruiting) will evaluate effects of colchicine plus standard treatment vs standard treatment alone on various outcome measures (e.g., mortality, markers of myocardial damage, clinical status, need for mechanical ventilation, duration of hospitalization) in adults with COVID-19: NCT04326790, NCT04322565, NCT04328480, NCT04350320, NCT04355143.</td>
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<td>Safety and efficacy for treatment of COVID-19 not established.</td>
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<td>Drug(s)</td>
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<td>Corticosteroids (general)</td>
<td>68:04 Adrenals</td>
<td>Potential to limit COVID-19-related myocardial damage also has been hypothesized based on the drug’s mechanisms of action and promising results of ongoing research on colchicine in various cardiac conditions. SARS-CoV-1 envelope (E) protein, a viroporin involved in replication and virulence, activates the NLRP3 inflammasome in vitro in Vero E6 cells by forming calcium-permeable ion channels, leading to increased IL-1β production. Evidence suggests that cytokine storm, a hyperinflammatory state resembling secondary hemophagocytic lymphohistiocytosis (HLH), is a contributing factor in COVID-19-associated mortality. Immunosuppression from corticosteroids has been proposed as a treatment option for such hyperinflammation. May improve dysregulated immune response caused by sepsis (possible complication of infection with the drugs can inhibit immune response, potentially limiting COVID-19-related myocardial damage).</td>
<td>Observational studies: Evidence suggests that corticosteroid use in patients with SARS, MERS, and influenza was associated with no survival benefit and possible harm (e.g., delayed viral clearance, avascular necrosis, psychosis, diabetes). Uncontrolled observational data from the recent COVID-19 outbreak in China suggest a possible treatment benefit of methylprednisolone in COVID-19 patients with acute respiratory distress syndrome (ARDS). (See Methylprednisolone in this Evidence Table.) Pending results of randomized controlled clinical studies specifically evaluating corticosteroids for COVID-19, indirect evidence from studies in patients with community-acquired pneumonia, ARDS, and other viral infections has been used to inform treatment decisions for COVID-19 patients. Systemic corticosteroid therapy has been studied in several randomized controlled studies for the treatment of ARDS; overall evidence is low to moderate in quality and</td>
<td>In general, low to moderate dosages of corticosteroids are recommended in intubated patients with ARDS. Regimens used in China were typically methylprednisolone 40-80 mg IV daily for a course of 3-6 days. Some experts suggest that equivalent dosages of dexamethasone (i.e., 7-15 mg daily, typically 10 mg daily) may have an advantage of producing less fluid retention, since dexamethasone has less mineralocorticoid activity. This dosage of dexamethasone is consistent with those used in the DEXA-ARDS trial. Higher dosages have been suggested for cytokine storm. (See Comments column.)</td>
<td>Data on the use of corticosteroids in COVID-19 are limited. The benefits and risks of corticosteroid therapy should be carefully weighed before use in patients with COVID-19. NIH, CDC, WHO, IDSA, and other experts have issued guidelines for the use of corticosteroids in patients with COVID-19 based on the currently available information. Recommendations are made according to the severity of illness, indications, and underlying medical conditions and should be considered on a case-by-case basis. General recommendations: WHO, CDC, NIH, and IDSA generally recommend against the routine use of corticosteroids for the treatment of COVID-19 unless indicated for another reason (e.g., asthma or COPD exacerbation, refractory septic shock). Non-critical patients: Corticosteroids generally should not be used in the treatment of early or mild disease since the drugs can inhibit immune response,</td>
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<td>Drug(s)</td>
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<td>COVID-19</td>
<td>5, 8, 9, 14, 17</td>
<td>and increase BP when low</td>
<td>most studies were performed prior to the prelung protection strategy era. In a recent multicenter, unblinded, randomized controlled study (DEXA-ARDS trial), the effects of dexamethasone in conjunction with conventional care were evaluated in hospitalized patients with moderate-to-severe ARDS receiving lung-protective mechanical ventilation. Treatment with IV dexamethasone at a dosage of 20 mg once daily on days 1-5, followed by 10 mg once daily on days 6-10 resulted in reduced duration of mechanical ventilation and reduced overall mortality (i.e., 15% increase in 60-day survival) compared with conventional treatment alone. Based on results of this study, a clinical trial (NCT04325061) has been initiated to specifically evaluate the use of dexamethasone in patients with ARDS due to COVID-19. Other clinical trials have been initiated in various countries to evaluate use of IV corticosteroids (e.g., dexamethasone, hydrocortisone), oral corticosteroids (e.g., prednisone), or inhaled corticosteroids (e.g., budesonide) for treatment of COVID-19 pneumonia or ARDS, including the following trials registered at clinicaltrials.gov: NCT04327401 NCT04344288 NCT04344730 NCT04348305 NCT04355637 NCT04359511 NCT04360876 (For registered clinical trials evaluating use of methylprednisolone, see Methylprednisolone in this Evidence Table.) Randomized controlled studies evaluating use of corticosteroids (e.g., hydrocortisone, dexamethasone, methylprednisolone, prednisolone) in septic shock suggest a small, but uncertain mortality reduction. NIH recommends against the routine use of systemic corticosteroids for the treatment of COVID-19 in hospitalized patients unless they are in the intensive care unit. Critically ill patients: The Surviving Sepsis Campaign COVID-19 subcommittee (a joint initiative of the Society of Critical Care Medicine and the European Society of Intensive Care Medicine) recommends against the routine use of systemic corticosteroids in mechanically ventilated adults with COVID-19 and respiratory failure (without ARDS). However, these experts generally support a weak recommendation to use low-dose, short-duration systemic corticosteroids in the sickest patients with COVID-19 and ARDS. NIH also recommends against the routine use of systemic corticosteroids for the treatment of mechanically ventilated COVID-19 patients without ARDS. However, the NIH panel states that there is insufficient evidence for or against the use of systemic corticosteroids in mechanically ventilated patients with COVID-19 and ARDS. IDSA suggests against using corticosteroids in hospitalized patients with COVID-19 pneumonia; however, in those with ARDS due to COVID-19, systemic corticosteroids may be used in the context of a clinical trial. Cytokine storm: There is no well-established or evidence-based treatment for cytokine storm in patients with COVID-19. However, some experts suggest that use of more potent immunosuppression with corticosteroids may be beneficial in such patients.</td>
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experts suggest higher dosages of corticosteroids (e.g., IV methylprednisolone 60–125 mg every 6 hours for up to 3 days) followed by tapering of the dose when inflammatory markers (e.g., C-reactive protein levels) begin to decrease. The decision to use corticosteroids in patients with early signs of cytokine storm should be balanced with the known adverse effects.

**Septic shock:** The effect of corticosteroids in COVID-19 patients with sepsis or septic shock may be different than the effects seen in those with ARDS. The Surviving Sepsis Campaign and NIH suggest the use of low-dose corticosteroid therapy (e.g., hydrocortisone 200 mg daily as an IV infusion or intermittent doses) over no corticosteroid therapy in adults with COVID-19 and refractory shock. Clinicians considering corticosteroids for such patients with COVID-19 should balance the potential small reduction in mortality with potential effects of prolonged coronavirus shedding. If corticosteroids are prescribed, monitor and treat adverse effects including hyperglycemia, hypernatremia, and hypokalemia.

**Patients receiving corticosteroid therapy for chronic conditions:** NIH states that oral corticosteroids used for the treatment of an underlying condition prior to COVID-19 infection (e.g., primary or secondary adrenal insufficiency, rheumatologic diseases) should not be discontinued. Supplemental or stress dosages of corticosteroids may be indicated on an individual basis in patients with such conditions. The guidelines also recommend that inhaled corticosteroids used daily for the management of asthma and COPD to control airway inflammation should not be discontinued in patients with COVID-19.
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Endocrinology experts state that patients with primary or secondary adrenal insufficiency who are receiving prolonged corticosteroid therapy should follow usual steroid “sick day rules” since these individuals may not be able to mount a normal stress response in the event of COVID-19 infection. If such individuals develop symptoms such as fever and a dry continuous cough, they should immediately double their daily oral corticosteroid dosage and continue with this regimen until the fever subsides. These guidelines also apply to patients who are receiving prolonged therapy (> 3 months) with corticosteroids for underlying inflammatory conditions, including asthma, allergy, and rheumatoid arthritis. In such patients whose condition worsens or in those experiencing vomiting or diarrhea, treatment with parenteral corticosteroids may be necessary. Administration of physiologic stress doses of corticosteroids (e.g., IV hydrocortisone 50-100 mg 3 times daily) and not pharmacologic doses should be considered in all cases to avoid potentially fatal adrenal failure. Additional study is needed to determine the optimum corticosteroid stress dosage regimens in patients with COVID-19. There is some evidence suggesting that continuous IV infusion of hydrocortisone (following an initial IV bolus dose) may provide more stable circulating cortisol concentrations in patients with adrenal insufficiency and reduce the potentially harmful effects of peak and trough concentrations of cortisol on the immune system.

**Pregnancy considerations:** For pregnant women with COVID-19, NIH guidelines state that the antenatal use of corticosteroids that cross the placenta (i.e., betamethasone, dexamethasone) is generally reserved for when administration is required for fetal benefit.
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<tr>
<td>COVID-19 Convalescent Plasma</td>
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<td>Plasma obtained from patients who have recovered from COVID-19 (i.e., COVID-19 convalescent plasma) that contains antibodies against SARS-CoV-2, including neutralizing antibodies, may provide short-term passive immunity to the virus; theoretically, such immunity may prevent or contribute to recovery from the infection, possibly as the result of viral neutralization and/or other mechanisms.</td>
<td>Uncontrolled pilot study of COVID-19 convalescent plasma in China: 10 adults with severe COVID-19 received a single transfusion of COVID-19 convalescent plasma (containing SARS-CoV-2 neutralizing antibody titers of 1:640 or greater) with standard care; all patients received antiviral therapy (e.g., umifenovir [Arbidol®], ribavirin, oseltamivir, peramivir, interferon α) and 6 patients also received methylprednisolone. The median time from onset of symptoms to transfusion of convalescent plasma was 16.5 days. COVID-19 symptoms (fever, cough, shortness of breath, chest pain) improved in all patients within 1-3 days after the transfusion and all patients showed radiological improvement in pulmonary lesions. Titors of neutralizing antibody increased in 5 patients after the transfusion, but remained the same in 4 patients. Prior to the transfusion, RT-PCR tests for SARS-CoV-2 RNA were positive in 7 patients and negative in 3 patients; after transfusion, SARS-CoV-2 RNA was undetectable in 3 patients on day 2, 3 patients on day 3, and 1 patient on day 6.</td>
<td>Efficacy and safety of COVID-19 convalescent plasma for the treatment of COVID-19 not established.</td>
<td>Other systemic corticosteroids do not cross the placenta, and pregnancy is not a reason to restrict their use if otherwise indicated. ACOG recommends against administration of antenatal corticosteroids for fetal benefit in the late preterm period (i.e., 34 weeks and 0 days through 36 weeks and 6 days) in patients with suspected or confirmed COVID-19 because the benefits of such therapy in late preterm are less well established. Treatment should be individualized, weighing the neonatal benefits of antenatal corticosteroid therapy with the risks of potential harm to the pregnant patient.</td>
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<td>Drug(s)</td>
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<td>received convalescent plasma less than 14 days after onset of symptoms had better outcomes than those who received such plasma later in the course of the disease.</td>
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<td>2 transfusions of COVID-19 convalescent plasma (containing SARS-CoV-2 neutralizing antibody end point dilution titers of 80-480 depending on the donor); patients continued to receive antiviral treatments (e.g., LPV/RTV, favipiravir, umifenovir [Arbidol®], darunavir, interferon α-1b) and methylprednisolone. Patients received the convalescent plasma transfusions 10-22 days after hospital admission. Following the transfusions, body temperature normalized within 3 days in 4/5 patients, sequential organ failure assessment (SOFA) scores improved in all patients (decreased from initial scores of 2-10 to 1-4 on day 12), titers of SARS-CoV-2 IgG, IgM, and neutralizing antibody increased in all patients, and viral loads decreased and became negative within 12 days. ¹⁰</td>
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<td>Retrospective observational study in China: 6 critically ill adults with COVID-19 were treated with convalescent plasma at a median of 21.5 days after first detection of viral shedding. Although viral clearance was observed in all patients following transfusion, death occurred in 5 out of 6 patients. ¹⁶</td>
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<td>Uncontrolled descriptive study in China: 6 adults with COVID-19 received convalescent plasma initiated at a relatively late stage of the disease (most patients received 2 or 3 plasma transfusions); various laboratory, radiologic, and clinical improvements were reported. ¹⁸</td>
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<td>Although there is some evidence that suggests possible benefits from use of convalescent plasma in patients with COVID-19, available data to date are largely from case reports or series; confirmation from randomized controlled studies is required. ¹, ¹⁰, ¹³</td>
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<td>Multiple clinical trials have been initiated in the US and other countries to evaluate use of COVID-19 convalescent plasma in</td>
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<td>Potential risks associated with COVID-19 convalescent plasma therapy (e.g., inadvertent transmission of other infectious agents, allergic reactions, thrombotic complications, transfusion-associated circulatory overload, transfusion-related acute lung injury [TRALI], antibody-dependent enhancement of infection) and steps to mitigate such risks not fully determined and require further evaluation. ¹, ⁵, ⁹, ²³, ²⁴</td>
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<td>FDA issued a guidance for industry to provide recommendations to healthcare providers and investigators regarding administration and study of investigational COVID-19 convalescent plasma. This guidance document includes recommendations regarding pathways for access to COVID-19 convalescent plasma, patient eligibility to receive such plasma, collection of such plasma (including donor eligibility and qualifications), product labeling, and recordkeeping. ¹¹</td>
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<td>FDA states that COVID-19 convalescent plasma is regulated as an investigational product and there currently are 3 available pathways for administering or studying use of such plasma:</td>
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<td>1). Clinical Trials: Requests to study use of COVID-19 convalescent plasma should be submitted to FDA under the traditional investigational new drug (IND) regulatory pathway. ¹¹</td>
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<td>2). Expanded Access IND: For patients with serious or immediately life-threatening COVID-19 who are not eligible or are unable to participate in randomized clinical trials, an expanded access IND can be used. A National Expanded Access Treatment Protocol has been established to facilitate access through participation of acute care facilities under an IND that is already in place. ¹¹ Information on a protocol that is currently in place is available at</td>
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<td>Drug(s)</td>
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<td>various settings (e.g., postexposure prophylaxis, treatment of different stages of the disease). Some of the trials that are currently recruiting are listed below. For additional trials, see clinicaltrials.gov:</td>
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<td>NCT04374370 (Expanded Access)</td>
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<td><a href="https://www.uscovidplasma.org">https://www.uscovidplasma.org</a></td>
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<td>3. Single Patient Emergency IND (eIND): Licensed physicians seeking to administer COVID-19 convalescent plasma to individual patients with serious or life-threatening disease may request an eIND from the FDA. Consult the FDA guidance document for specific information on applying for an eIND.</td>
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<td>Donor eligibility: FDA guidance suggests that COVID-19 convalescent plasma be collected from individuals with laboratory-confirmed evidence of COVID-19 infection and complete resolution of symptoms for at least 14 days before donation (a negative result for COVID-19 by a diagnostic test is not necessary to qualify the donor).</td>
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<td>Antibody titers in donor plasma: If measurement of antibody titers is available, FDA recommends a neutralizing antibody titer of at least 1:160 (a titer of 1:80 may be considered acceptable if an alternative matched unit of plasma is not available).</td>
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<td>Patient eligibility: For healthcare providers seeking an eIND for the treatment of patients with severe or life-threatening disease, consideration should be given to following the patient eligibility criteria used in the National Expanded Access Treatment Protocol <a href="https://www.uscovidplasma.org">https://www.uscovidplasma.org</a>. According to the protocol, severe disease is defined as one or more of the following: shortness of breath, respiratory frequency 30/minute or greater, blood oxygen saturation 93% or lower, PaO2/FiO2 ratio less than 300, lung infiltrates greater than 50% within 24-48 hours, and life-threatening disease is defined as one or more of the following: respiratory failure, septic shock, multiple organ dysfunction or failure.</td>
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<td>Drug(s)</td>
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<td>Epoprostenol (inhaled)</td>
<td>48:48</td>
<td>Vasodilating Agent; may be useful in the adjunctive treatment of acute respiratory distress syndrome (ARDS), a potential complication of COVID-19 1-9</td>
<td>No studies evaluating use specifically in COVID-19 patients 10</td>
<td>Various dosages of inhaled epoprostenol have been used in ARDS studies 2-9</td>
<td>Additional studies are needed to evaluate the potential role of inhaled epoprostenol in the treatment of ARDS 6,9. The Surviving Sepsis Campaign states that due to the lack of adequately powered randomized controlled studies, a recommendation cannot be made for or against use of inhaled prostacyclins in COVID-19 patients with severe ARDS 10</td>
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<td>Methylprednisolone (DEPO-Medrol®, SOLU-Medrol®)</td>
<td>68:04</td>
<td>Adrenal</td>
<td>Retrospective, observational, single-center study: In 201 patients with confirmed COVID-19 pneumonia who developed ARDS, methylprednisolone appeared to reduce the risk of death. 6 Among patients with ARDS, of those who received methylprednisolone treatment, 23 of 50 (46%) patients died, while of those who did not receive methylprednisolone, 21 of 34 (61.8%) died. 6</td>
<td>Dosage used in the retrospective study (Wu et al) not provided. 6</td>
<td>Findings from observational studies suggest that for patients with COVID-19 pneumonia who progress to ARDS, methylprednisolone treatment may be beneficial. However, results should be interpreted with caution because of potential bias (drug used in sickest patients) and small sample size. Confirmation from randomized controlled studies is needed. 6,13 (See Corticosteroids in this Evidence Table.)</td>
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Methylprednisolone was suggested as an alternative to inhaled nitric oxide due to its similar efficacy, lower potential for systemic adverse effects, lower cost, and ease of delivery 1-2,9.

Inhaled epoprostenol has been suggested as an alternative to inhaled nitric oxide due to its similar efficacy, lower potential for systemic adverse effects, lower cost, and ease of delivery 1, 2, 9.

Experience in patients with ARDS indicates that inhaled epoprostenol can substantially reduce mean pulmonary artery pressure and improve oxygenation in such patients; however, data demonstrating clinical benefit are lacking 3, 6-9.

Various dosages of inhaled epoprostenol have been used in ARDS studies 2-9.

Dosages up to 50 ng/kg per minute have been used (titrated to response). 1, 4, 6, 9.

To provide a clinically important increase in PaO₂ and reduction in pulmonary artery pressure, data from these studies suggest that the most effective and safe dosage appears to be 20-30 ng/kg per minute in adults and 30 ng/kg per minute in pediatric patients 9.

Additional studies are needed to evaluate the potential role of inhaled epoprostenol in the treatment of ARDS 6,9.

The Surviving Sepsis Campaign states that due to the lack of adequately powered randomized controlled studies, a recommendation cannot be made for or against use of inhaled prostacyclins in COVID-19 patients with severe ARDS 10.

Dosage used in the retrospective study (Wu et al) not provided. 6

Dosage used in the retrospective study (Wang et al) was 1-2 mg/kg daily IV for 5-7 days. 13

Dosage used in the randomized, controlled study (NCT04244591) was 40 mg IV every 12 hours for 5 days. 23

Findings from observational studies suggest that for patients with COVID-19 pneumonia who progress to ARDS, methylprednisolone treatment may be beneficial. However, results should be interpreted with caution because of potential bias (drug used in sickest patients) and small sample size. Confirmation from randomized controlled studies is needed. 6,13 (See Corticosteroids in this Evidence Table.)

Dosage used in the randomized, controlled study (NCT04244591) was recently completed in China that compared use of methylprednisolone in conjunction with standard care in patients with confirmed COVID-19 infection that progressed to acute respiratory failure; results have not yet been posted. 23

(See Corticosteroids in this Evidence Table.)

Open-label, multicenter, randomized controlled study (NCT04244591) was recently completed in China that compared use of methylprednisolone in conjunction with standard care in patients with confirmed COVID-19 infection that progressed to acute respiratory failure; results have not yet been posted. 23
<table>
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<tr>
<th>Drug(s)</th>
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<th>Trials or Clinical Experience</th>
<th>Dosage*</th>
<th>Comments</th>
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<tr>
<td>Nitric oxide (inhaled)</td>
<td>48:48 Vasodilating Agent</td>
<td>Selective pulmonary vasodilator; may be useful in the adjunctive treatment of acute respiratory distress syndrome (ARDS), a potential complication of COVID-19</td>
<td>Multiple clinical trials have been initiated in various countries to evaluate use of methylprednisolone for treatment of COVID-19 pneumonia or severe acute respiratory syndrome, including the following trials registered at clinicaltrials.gov: NCT03852537, NCT04263402, NCT04323592, NCT04329650, NCT04343729. A non-randomized pilot study registered at clinicaltrials.gov (NCT04355247) has been initiated to evaluate use of methylprednisolone for the prevention of COVID-19 cytokine storm and progression to respiratory failure.</td>
<td>In the Chen et al. study in severe SARS patients, inhaled nitric oxide therapy was given for ≥3 days (30 ppm on day 1, followed by 20 and 10 ppm on days 2 and 3, respectively, then weaned on day 4; therapy was resumed at 10 ppm if deteriorating oxygenation occurred). Phase 2 clinical trial protocol (NCT04306393) for treatment of mechanically ventilated COVID-19 patients: 80 ppm for the first 48 hours, followed by 40 ppm and then subsequently wean.</td>
<td>Therapeutic guidelines for the treatment of ARDS state that inhaled nitric oxide may be considered in patients with severe hypoxemia not responsive to conventional ventilation strategies; however, routine use not recommended. The Surviving Sepsis Campaign suggests a trial of inhaled pulmonary vasodilator therapy as rescue therapy in mechanically ventilated adults with COVID-19, severe ARDS, and hypoxemia despite optimized ventilation and other rescue strategies; if rapid improvement in oxygenation is not observed, treatment should be tapered off. Clinical trials evaluating inhaled nitric oxide for the treatment or prevention of COVID-19 are planned or underway (NCT04338828, NCT04305457, NCT04306393, NCT04312243). On March 20th, 2020, Bellerophon Therapeutics announced that the FDA granted emergency expanded access allowing its inhaled nitric oxide delivery system (INOpulse®) to be immediately used for the treatment of COVID-19.</td>
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Updated: 4/22/2020
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<th>Drug(s)</th>
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<td><strong>Ruxolitinib</strong></td>
<td>10:00</td>
<td>Janus kinase (JAK) 1 and 2 inhibitor; 7 may potentially combat cytokine release syndrome (CRS) in severely ill patients 6,5. Ability to inhibit a variety of proinflammatory cytokines, including interferon, has been raised as a possible concern with the use of JAK inhibitors in the management of hyperinflammation resulting from viral infections such as COVID-19 5,7.</td>
<td>Currently no known published clinical trial evidence supporting efficacy or safety in patients with COVID-19. Phase 3 randomized, double-blind, placebo-controlled clinical trial (NCT04362137; RUXCOVID) evaluating ruxolitinib plus standard of care vs standard of care alone is being initiated in patients ≥12 years of age with COVID-19-associated cytokine storm (sponsored by Incyte in U.S. and Novartis outside of U.S.) 1,10. <strong>Expanded-access (managed-access, compassionate use) program</strong> (NCT04337359) available for eligible adults and children ≥6 years of age with severe or very severe COVID-19 illness; address inquiries to Incyte (855-463-3463 or <a href="mailto:medinfo@incyte.com">medinfo@incyte.com</a>) 1,2. <strong>Expanded-access program</strong> (NCT04355793) available for emergency treatment of cytokine storm from COVID-19 infection in adults and pediatric patients ≥12 years of age; address inquiries to Incyte (855-463-3463 or <a href="mailto:medinfo@incyte.com">medinfo@incyte.com</a>) 9. Other earlier-phase, smaller, and/or open-label clinical trials registered: NCT04331665 NCT04334044 NCT04338958 NCT04348071 NCT04359290 NCT04354714 NCT04348695 ChiCTR2000029580 (in Chinese Clinical Trial Registry) 5,6.</td>
<td>Various dosages are being evaluated 3,6,10. Phase 3 study (NCT04362137): Ruxolitinib 5 mg twice daily for 14 days with possible extension to 28 days 10.</td>
<td>NIH COVID-19 Treatment Guidelines Panel recommends against use of JAK inhibitors for the treatment of COVID-19 except in the context of a clinical trial; the panel states that at present the broad immunosuppressive effect of JAK inhibitors outweighs the potential for benefit 8.</td>
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<td><strong>Sarilumab</strong></td>
<td>92:36</td>
<td>Recombinant humanized monoclonal antibody specific for the interleukin-6 (IL-6) receptor; may potentially combat cytokine release syndrome (CRS) and pulmonary symptoms in severely ill patients 5,2,5.</td>
<td>Currently no known published clinical trial evidence supporting efficacy or safety in treatment of patients with COVID-19. However, based on encouraging results in China with a similar drug, tocilizumab, a U.S.-based, phase 2/3, randomized, double-blind, placebo-controlled study evaluating efficacy and safety of sarilumab in patients hospitalized with severe COVID-19 is currently under way 1,4.</td>
<td>Not available (see Trials or Clinical Experience)</td>
<td>NIH COVID-19 Treatment Guidelines Panel states that there are insufficient clinical data to recommend either for or against use of sarilumab in the treatment of COVID-19 7.</td>
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**Updated 5/13/20. The current version of this document can be found on the ASHP COVID-19 Resource Center.**  
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<th>Drug(s)</th>
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<th>Trials or Clinical Experience</th>
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<td>Siltuximab</td>
<td>10:00</td>
<td>Antineoplastic agents</td>
<td>Recombinant chimeric monoclonal antibody specific for the interleukin-6 (IL-6) receptor; may potentially combat cytokine release syndrome (CRS) symptoms (e.g., fever, organ failure, death) in severely ill patients.</td>
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<td>For compassionate use access or investigator-sponsored clinical studies, contact the manufacturer (Sanofi Genzyme) for further information (1-800-633-1610).</td>
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<td>(Sylvant®)</td>
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<td><strong>Italy</strong>: Early (non-peer-reviewed) findings from an observational case-control study of the first 21 patients with COVID-19 and pneumonia/acute respiratory distress syndrome (ARDS) who participated in a compassionate use program (SISCO study; NCT04322188) in one hospital and were followed for up to 7 days showed reduced and normalized C-reactive protein (CRP) levels (a marker of systemic inflammation) by day 5 in all 16 siltuximab-treated patients with sufficient available data. An interim analysis revealed that the condition of 33% of the siltuximab-treated patients improved and no clinically relevant change in condition was reported in 43% of patients while 24% of patients worsened, including one patient who died and another with cerebral vascular event. This cohort study with patients treated with standard therapy is ongoing.</td>
<td>In the SISCO study in Italy, patients received an initial dose of siltuximab 11 mg/kg by IV infusion over 1 hour; a second dose could be administered at the physician’s discretion (5 of the first 21 patients received a second dose after 2-3 days).</td>
<td>Other clinical trials evaluating siltuximab in the treatment of COVID-19 currently are recruiting in Belgium (NCT04330638) and Spain (NCT04329650).</td>
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<td>Other clinical trials evaluating siltuximab in the treatment of COVID-19 currently are recruiting in Belgium (NCT04330638) and Spain (NCT04329650).</td>
<td>Yazaki et al. (2020).</td>
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<td>Sirolimus</td>
<td>92:44</td>
<td>Immunosuppressive agent (mTOR inhibitor)</td>
<td>mTOR complex 1 (mTORC1) is involved in the replication of various viruses, including coronavirus.</td>
<td>In vitro studies demonstrated inhibitory activity against MERS-CoV infection.</td>
<td>Dosage being investigated in NCT04341675 trial: 6 mg orally on day 1 followed by 2 mg daily for a maximum treatment duration of 14 days or until hospital discharge.</td>
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Updated 5-15-20. The current version of this document can be found on the ASHP COVID-19 Resource Center.
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<td>Tocilizumab (Actemra&lt;sup&gt;®&lt;/sup&gt;)</td>
<td>92:36 Disease-modifying Anti-rheumatic Drug</td>
<td>Recombinant humanized monoclonal antibody specific for the interleukin-6 (IL-6) receptor; may potentially combat cytokine release syndrome (CRS) symptoms in severely ill COVID-19 patients&lt;sup&gt;1&lt;/sup&gt;,&lt;sup&gt;4&lt;/sup&gt;,&lt;sup&gt;6&lt;/sup&gt;,&lt;sup&gt;10&lt;/sup&gt;,&lt;sup&gt;12&lt;/sup&gt;</td>
<td>Case reports and observational studies describing use of tocilizumab in patients with COVID-19 reported from various areas of the world&lt;sup&gt;1&lt;/sup&gt;,&lt;sup&gt;6&lt;/sup&gt;,&lt;sup&gt;10&lt;/sup&gt;,&lt;sup&gt;12&lt;/sup&gt; In preliminary data from a non-peer-reviewed, single-arm, observational Chinese trial (Xu et al.) involving 21 patients with severe or critical COVID-19 infection, patients demonstrated rapid fever reduction and a reduced need for supplemental oxygen within several days after receiving tocilizumab (initially given as a single 400-mg dose by IV infusion; this dose was repeated within 12 hours in 3 patients because of continued fever)&lt;sup&gt;3&lt;/sup&gt; In a retrospective, observational study in China (Luo et al.) involving 15 patients moderately to critically ill with COVID-19, tocilizumab (80-600 mg per dose) was given, and was used in conjunction with methylprednisolone in 8 of the patients. About one-third of the patients received 2 or more doses of tocilizumab. Elevated C-reactive protein (CRP) levels rapidly decreased in most patients following treatment, and a gradual decrease in IL-6 levels was noted in patients who stabilized following tocilizumab administration. Clinical outcomes were equivocal.&lt;sup&gt;10&lt;/sup&gt; A single-center, retrospective observational study of 20 kidney transplant recipients in Italy with COVID-19 hospitalized for pneumonia included 6 patients who received tocilizumab. Half of the patients experienced reduced oxygen requirements and 2 (33%) showed improved radiologic findings following administration; 2 (33%) of the 6 tocilizumab-treated patients died.&lt;sup&gt;12&lt;/sup&gt; Zhang et al. from China reported on a patient with COVID-19 and multiple myeloma who appeared to be successfully treated with tocilizumab&lt;sup&gt;13&lt;/sup&gt;</td>
<td>IV infusion: China recommends an initial dose of 4–8 mg/kg infused over more than 60 minutes. If initial dose not effective, may administer second dose (in same dosage as initial dose) after 12 hours. No more than 2 doses should be given; maximum single dose is 800 mg&lt;sup&gt;2&lt;/sup&gt; US/Global randomized, placebo-controlled trial (manufacturer sponsored; COVACTA): Will evaluate an initial IV infusion of 8 mg/kg (up to a maximum dose of 800 mg); one additional dose may be given if symptoms worsen or show no improvement&lt;sup&gt;8&lt;/sup&gt;</td>
<td>In China, tocilizumab can be used to treat severely or critically ill COVID-19 patients with extensive lung lesions and high IL-6 levels&lt;sup&gt;2&lt;/sup&gt; NIH COVID-19 Treatment Guidelines Panel states that there are insufficient clinical data to recommend either for or against use of tocilizumab in the treatment of COVID-19&lt;sup&gt;9&lt;/sup&gt; The role of routine cytokine measurements (e.g., IL-6, CRP) in determining the severity of and treating COVID-19 requires further study&lt;sup&gt;14&lt;/sup&gt;</td>
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<td><strong>Currently, there are no well-controlled published studies on the efficacy and safety of tocilizumab for the treatment of COVID-19; however, numerous clinical trials are planned or underway globally.</strong>&lt;sup&gt;1, 5, 7, 8&lt;/sup&gt; China: Randomized, multicenter, controlled clinical trial evaluating efficacy &amp; safety in 188 patients with COVID-19 under way through 5/10/20. <strong>Results not yet available.</strong> Chinese Clinical Trial Registry link: <a href="http://www.chictr.org.cn/showprof.aspx?proj=49409">http://www.chictr.org.cn/showprof.aspx?proj=49409</a> US/Global randomized, placebo-controlled trial: Manufacturer (Roche) conducting a randomized, double-blind, placebo-controlled phase 3 trial (COVACTA; NCT04320615) in collaboration with the US Health and Human Services’ Biomedical Advanced Research and Development Authority (BARDA); the study will evaluate safety and efficacy of tocilizumab in combination with standard of care compared with placebo. Expected to enroll about 330 patients globally, including in the U.S., beginning in April 2020&lt;sup&gt;7, 8&lt;/sup&gt;</td>
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<table>
<thead>
<tr>
<th>Drug(s)</th>
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<th>Rationale</th>
<th>Trials or Clinical Experience</th>
<th>Dosage</th>
<th>Comments</th>
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</thead>
<tbody>
<tr>
<td>ACE Inhibitors, Angiotensin II Receptor Blockers (ARBs)</td>
<td>24:32</td>
<td>Renin-Angiotensin-Aldosterone System Inhibitor</td>
<td>Hypothetical harm: Human pathogenic coronaviruses bind to their target cells through angiotensin-converting enzyme 2 (ACE2). Expression of ACE2 may be increased in patients treated with ACE inhibitors or ARBs. Increased expression of ACE2 may potentially facilitate COVID-19 infections.</td>
<td>Data are lacking; no evidence of harm or benefit with regards to COVID-19 infection. Clinical trial underway: Initiation of losartan in adult patients with COVID-19 requiring hospitalization; primary outcome measure: sequential organ failure assessment (SOFA) respiratory score. (NCT04312009)</td>
<td>American Heart Association (AHA), American College of Cardiology (ACC), Heart Failure Society of America (HFSA), European Society of Cardiology (ESC) recommend to continue treatment with renin-angiotensin-alderosterone system (RAAS) antagonists in those patients who are currently prescribed such agents. NIH COVID-19 Treatment Guidelines Panel states patients who are receiving an ACE inhibitor or ARB for cardiovascular disease (or other indications) should continue receiving these drugs; recommends against use of ACE inhibitors or ARBs for the treatment of COVID-19 except in the context of a clinical trial. Patients with cardiovascular disease are at an increased risk of serious COVID-19 infections. Abrupt withdrawal of RAAS inhibitors in high-risk patients (e.g., heart failure patients, patients with prior myocardial infarction) may lead to clinical instability and adverse health outcomes.</td>
</tr>
</tbody>
</table>

<p>| Anticoagulants (low molecular weight heparin [LMWH], unfractionated heparin [UFH]) | 20:12.04.16 | Heparins | There is increasing evidence that patients with severe COVID-19 develop a hypercoagulable state, which has been associated with poor outcomes (e.g., progressive respiratory failure, acute respiratory distress syndrome [ARDS], death). Coagulation abnormalities observed in these patients include prothrombotic disseminated intravascular coagulation (DIC), venous thromboembolism, elevated D-dimer levels, high fibrinogen levels, and microvascular and macrovascular. | Limited data from a retrospective study in China suggest that patients with severe COVID-19 infection or markedly elevated levels of D-dimer (&gt;6 x ULN) may have decreased mortality when given prophylactic doses of LMWH or UFH. However, prospective studies are needed to confirm these findings. A randomized open-label clinical trial (NCT04345848) is currently being conducted to evaluate prophylactic- and therapeutic-dose anticoagulation in hospitalized adults with severe COVID-19 infection. | Additional study is needed to understand the anticoagulant needs of COVID-19 patients. Several organizations have published interim guidance for the management of COVID-19-associated coagulopathy. The International Society for Thrombosis and Haemostasis (ISTH) and the American Society of Hematology (ASH) recommend that all hospitalized COVID-19 patients, including non-ICU patients, receive prophylactic-dose LMWH unless contraindicated (e.g., active bleeding, platelet count &lt;25x10^9/L, fibrinogen less than 0.5 g/L). Abnormal PT or aPTT is not a contraindication for prophylaxis. |</p>
<table>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Thrombosis. 1-6, 8, 11, 14, 18</td>
<td>Early anticoagulation in patients with severe COVID-19 infection may reduce the risk of thrombotic complications and improve clinical outcomes. 2, 4, 5, 14, 25, 27</td>
<td>WHO recommends pharmacologic prophylaxis with LMWH (preferred) or UFH (5000 units sub-Q twice daily) in adults and adolescents without contraindications. 25</td>
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<td>An additional benefit of heparins is their anti-inflammatory effects. 5, 7, 8, 17</td>
<td>Although LMWH is generally preferred, 4, 5, 25 UFH also has been used for thrombo prophylaxis; practical concerns (e.g., need for frequent monitoring, convenience of administration, risk of medical staff exposure) may influence institutional choice of anticoagulant. 8, 9, 14, 20, 27</td>
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<td>Because of the severity of coagulopathy in critically ill COVID-19 patients and reports of thrombotic complications that have continued to occur despite standard prophylaxis with LMWH or UFH, some clinicians suggest that higher prophylactic doses or even therapeutic doses be considered; however, high-quality randomized controlled studies are needed to evaluate these approaches. 11, 14, 17, 20-24, 26, 27</td>
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<td>The American Society of Hematology (ASH) states that therapeutic anticoagulation is not required in COVID-19 patients unless there is documented VTE or atrial fibrillation. 4 The efficacy of intermediate or full therapeutic anticoagulation for critically ill COVID-19 patients without documented VTE is being evaluated. 4 In patients already on anticoagulation for VTE or atrial fibrillation, therapeutic doses of anticoagulant therapy should continue but may need to be held if the platelet count is less than 30-50 x 10^9/L or if fibrinogen is less than 1 g/L. 4</td>
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<td>The risk of venous thromboembolism and anticoagulation requirements should be assessed in all patients on an individual basis. 4, 5, 10, 17, 18</td>
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<td>Bleeding appears to be infrequent in COVID-19 patients. 5 However, standard risk factors for bleeding should be considered and patients should be individually assessed to balance risk of thrombosis with risk of bleeding. 4</td>
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<tr>
<td>Famotidine</td>
<td>56:28.12</td>
<td>Histamine H₂ Antagonists</td>
<td>Computer-aided, structure-based, virtual screening of libraries of compounds against SARS-CoV-2 proteins suggested potential for famotidine to interact with viral proteases involved in coronavirus replication. <strong>1–4</strong> Anecdotal observations: Observations based on retrospective medical record review indicated that many Chinese COVID-19 survivors had received famotidine for chronic heartburn; mortality rate appeared to be lower in hospitalized COVID-19 patients receiving famotidine than in patients not receiving the drug (14 vs 27%); observations did not control for possible confounding (e.g., socioeconomic) factors. <strong>3</strong> Retrospective matched cohort study of COVID-19 patients initially hospitalized in non-ICU setting at a single New York medical center indicated that the risk for the composite outcome of death or intubation was reduced (mainly due to difference in mortality) in patients who received famotidine within 24 hours of hospital admission vs those who did not receive the drug. <strong>7</strong> Currently no known published prospective clinical trial evidence supporting efficacy or safety for treatment of COVID-19. Randomized, double-blind, historical-controlled, comparative trial (NCT04370262) initiated in New York in hospitalized adults with moderate to severe COVID-19; trial includes 2 active treatment groups (high-dose IV famotidine with oral hydroxychloroquine, IV placebo with oral hydroxychloroquine) and a historical control group receiving neither of these drugs (patients treated during early stages of the COVID-19 pandemic in New York); targeted enrollment is 600 patients in each active treatment group; 2 interim analyses planned. <strong>5</strong> Dosage in NCT04370262: Famotidine is being given IV in 120-mg doses (proposed total daily dosage of 360 mg) for maximum of 14 days or until hospital discharge, whichever comes first. <strong>5</strong> Proposed daily dosage in NCT04370262 is 9 times the usual manufacturer-recommended IV adult dosage; the study excludes patients with creatinine clearance (CrCl) ≤50 mL/minute, including dialysis patients; renally impaired patients may be at increased risk of adverse CNS effects since drug half-life is closely related to CrCl. <strong>6</strong> Safety and efficacy for treatment of COVID-19 not established.</td>
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<td>HMG-CoA Reductase Inhibitors (statins)</td>
<td>24:06</td>
<td>Antilipemic Agents</td>
<td>In addition to lipid-lowering effects, statins have anti-inflammatory and immunomodulatory effects which may prevent acute lung injury. <strong>1</strong> Statins affect ACE2 as part of their function in reducing endothelial dysfunction. <strong>2, 8</strong> Data are lacking on the use of statins in patients with COVID-19. Preliminary findings have shown mixed results with other respiratory illnesses; some observational studies suggest statin therapy is associated with a reduction in various cardiovascular outcomes and possibly mortality in patients hospitalized with COVID-19. NIH COVID-19 Treatment Guidelines Panel states patients who are receiving a statin for the treatment or prevention of cardiovascular disease should continue statin therapy; <strong>2</strong> recommends against use of statins for the treatment of COVID-19 except in the context of a clinical trial. <strong>2</strong></td>
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<tr>
<td>Immune Globulin (IGIV, IVIG, γ-globulin)</td>
<td>80:04 Immune Globulin</td>
<td>Commercially available immune globulin (IGIV, IVIG, γ-globulin) is derived from pooled plasma; contains many antibodies normally present in adult human blood; used for replacement therapy in pts with primary humoral immunodeficiency unable to produce sufficient IgG antibodies and also used to provide passive immunity to certain viral infections in other individuals.</td>
<td>SARS Experience: IGIV has been used in some pts for the treatment of SARS. Benefits in such pts were unclear because of comorbidities, differences in stage of illness, and effect of other treatments; IGIV may have contributed to hypercoagulable state and thrombotic complications in some pts. COVID-19 case reports in China (Cao et al): Treatment with IGIV at the early stage of clinical deterioration was reported to provide some clinical benefit in 3 adults with severe COVID-19; 2 pts also received antivirals and 1 pt also received short-term steroid treatment. Patients were afebrile within 1-2 days and breathing difficulties gradually improved within 3-5 days of IGIV administration.</td>
<td>IGIV dosage of 0.3-0.5 g/kg daily for 5 days has been used in some pts with COVID-19; IGIV dosage of 0.5 g/kg daily for 5 days being investigated in a clinical trial in China.</td>
<td>Role of commercially available immune globulin (IGIV, IVIG, γ-globulin) in the treatment of COVID-19 unclear. The Surviving Sepsis Campaign COVID-19 subcommittee suggests that IGIV not be used routinely in critically ill adults with COVID-19 because efficacy data not available, currently available IGIV preparations may not contain antibodies against SARS-CoV-2, and IGIV can be associated with increased risk of severe adverse effects (e.g., anaphylaxis, aseptic meningitis, renal failure, thromboembolism, hemolytic reactions, transfusion-related lung injury). IGIV mentioned in Chinese guidelines as other therapeutic measure for treatment of severe and critical cases of COVID-19 in children.</td>
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<td>Drug(s)</td>
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<tr>
<td>Ivermectin</td>
<td>8:08</td>
<td>Anthelmintic</td>
<td>In vitro activity against some human and animal viruses&lt;sup&gt;1,6&lt;/sup&gt;</td>
<td></td>
<td>No data to date to support use in the treatment of COVID-19</td>
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<tr>
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<td></td>
<td>In vitro evidence of activity against SARS-CoV-2 in infected Vero-hSLAM cells reported with high concentrations of the drug&lt;sup&gt;1&lt;/sup&gt;</td>
<td>Currently no known published data regarding efficacy or safety in the treatment of COVID-19</td>
<td></td>
<td>Ivermectin plasma concentrations attained with dosages recommended for treatment of parasitic infections are substantially lower than concentrations associated with in vitro inhibition of SARS-CoV-2; pharmacokinetic modeling predicts that plasma concentrations attained with dosages up to 10 times higher than usual dosage also are substantially lower than concentrations associated with in vitro inhibition of the virus&lt;sup&gt;9&lt;/sup&gt;</td>
</tr>
<tr>
<td>Nebulized drugs</td>
<td>8:08</td>
<td>Anthelmintic</td>
<td>Potential harm: Concern that use of nebulized drugs (e.g., albuterol) for the management of respiratory conditions in patients with COVID-19 infection may distribute the virus into the air and expose close contacts&lt;sup&gt;1,2&lt;/sup&gt;</td>
<td></td>
<td>American College of Allergy, Asthma &amp; Immunology (ACAAI) recommends that nebulized albuterol should be administered in a location that minimizes exposure to close contacts who do not have COVID-19 infection. In the home, choose a location where air is not recirculated (e.g., porch, patio, or garage) or areas where surfaces can be cleaned easily or may not need cleaning.&lt;sup&gt;1&lt;/sup&gt; In hospitals, clinicians typically use nebulizers to deliver medications such as albuterol, but are being encouraged to switch to use of metered-dose inhalers because of the risk of the virus becoming airborne when treating patients infected with COVID-19. &lt;sup&gt;2&lt;/sup&gt;</td>
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<td>Nebulizer treatment used in clinical practice to treat influenza and other respiratory infections is thought to generate droplets or aerosols. In one study, nebulized saline delivered droplets in the small- and medium-size aerosol/droplet range. These results may have infection control implications for airborne infections, including severe acute respiratory syndrome and pandemic influenza infection.&lt;sup&gt;3&lt;/sup&gt;</td>
<td></td>
<td>No data to date inappropriate use of ivermectin products intended for animals as an attempt to self-medicate for the treatment of COVID-19&lt;sup&gt;8&lt;/sup&gt;</td>
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<tr>
<td>Niclosamide</td>
<td>8:08</td>
<td>Anthelmintic</td>
<td>Broad antiviral activity</td>
<td></td>
<td>Protocol in one ongoing trial (&lt;a&gt;NCT04372082&lt;/a&gt;) for treatment of COVID-19 specifies a niclosamide dosage of 2 g on day 1, then 500 mg twice daily for 10 days&lt;sup&gt;3&lt;/sup&gt;</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Currently no known published clinical trial data regarding efficacy or safety in the treatment of COVID-19</td>
<td></td>
<td>Not commercially available in the US</td>
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<td></td>
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<td></td>
<td>In vitro evidence of activity against SARS-CoV and MERS-CoV&lt;sup&gt;1,2&lt;/sup&gt;</td>
<td></td>
<td>No data to date support use in treatment of COVID-19</td>
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<tbody>
<tr>
<td>Nitazoxanide</td>
<td>8:30.92 Antiprotozoal</td>
<td>In vitro activity against various viruses, including coronaviruses (^4,5)</td>
<td>Randomized, open-label, controlled trial in France  (NCT04372082; HydiLIC) to evaluate niclosamide in adults with SARS-CoV-2 infection (asymptomatic or onset of symptoms less than 8 days previously) and comorbidities (^3)</td>
<td>Dosages investigated for treatment of influenza and influenza-like illness or being investigated for other viral infections: Adults and adolescents (≥12 years of age): 500 or 600 mg orally twice daily for 5 days (^6,7,8)</td>
<td>Current data not specific to COVID-19; additional study needed (^1)</td>
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**Updated 5/1/20**

**Experience in treating influenza:** In a randomized, placebo-controlled study in 624 otherwise healthy adult and adolescent patients with acute uncomplicated influenza, treatment with nitazoxanide reduced duration of symptoms by approximately 1 day \(^6\).

**Experience in treating influenza-like illness:** In two studies for the treatment of influenza-like illness symptoms associated with viral respiratory infection in 186 adults and pediatric pts, treatment with nitazoxanide reduced duration of symptoms (4 days versus ≥7 days with placebo). \(^7\) In another study in 260 adults and pediatric pts hospitalized with influenza-like illness (≥50% with pneumonia at presentation), treatment with nitazoxanide did not reduce the duration of hospital stay (primary end point) or duration of symptoms \(^7\).

**COVID-19:** Randomized, double-blind, placebo-controlled proof-of-concept trial (NCT04348409) initiated to evaluate nitazoxanide for treatment of moderate COVID-19 \(^8\).

**Two randomized, double-blind, placebo-controlled clinical trials** have been initiated by the manufacturer (Romark) to evaluate efficacy and safety for pre- or post-exposure prophylaxis of COVID-19 and other viral respiratory illnesses in healthcare workers (NCT04359680) and post-exposure prophylaxis of COVID-19 and other viral respiratory illnesses in elderly residents of long-term care facilities (NCT04343248) \(^8\).
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</thead>
<tbody>
<tr>
<td>Nonsteroidal Anti-inflammatory Agents (NSAIAs)</td>
<td>28:08.04 Nonsteroidal Anti-inflammatory Agent (NSAIA)</td>
<td><strong>Ibuprofen:</strong> Speculative link between ibuprofen and increased ACE2 expression leading to worse outcomes in COVID-19 patients, and should NOT be used in patients with COVID-19.</td>
<td>Multiple other clinical trials planned or initiated to evaluate nitazoxanide in combination with other drugs (chloroquine, hydroxychloroquine, or ivermectin) or alone for treatment of COVID-19.</td>
<td>Ibuprofen: None; anecdotal.</td>
<td>Ibuprofen: A letter published in The Lancet Respir Med stated that increased expression of ACE2 could facilitate infection with COVID-19. The letter states that thiazolidinediones and ibuprofen can increase ACE2; however, this appears to be based on animal studies. 1,4 A statement attributed to WHO spokesperson Christian Lindmeier recommending paracetamol and avoiding ibuprofen as a self-medicating was widely circulated in the media; however, such a position could not be found on the WHO website or other official sources. WHO has stated “after a rapid review of the literature, is not aware of published clinical or population-based data on this topic.” As of 3/18/20 (via Twitter) “WHO does not recommend against the use of ibuprofen.” <a href="https://twitter.com/WHO/status/1240409217997189128">https://twitter.com/WHO/status/1240409217997189128</a></td>
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<td>Indomethacin: Possible antiviral activity against other coronaviruses SARS-CoV &amp; CanineCoV (interferes with viral RNA synthesis).</td>
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<td>Indomethacin: Speculative; one in vitro &amp; animal model study with other coronaviruses SARS-CoV &amp; CanineCoV.</td>
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### Tissue Plasminogen Activator (t-PA; alteplase)

**Updated 4/29/20**

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<tr>
<td><strong>20:12.20</strong></td>
<td>20:12.20 Thrombolytic agents</td>
<td>A consistent finding in patients with severe COVID-19 is a hypercoagulable state, which may contribute to their risk of poor outcomes (e.g., progressive respiratory failure, acute respiratory distress syndrome [ARDS], death).</td>
<td>Results of a small phase 1 study conducted in 2001 suggest possible benefit of plasminogen activators for the treatment of ARDS. In this study, 20 patients with ARDS secondary to trauma and/or sepsis who failed to respond to standard ventilator therapy and were not expected to survive were treated with urokinase or streptokinase; such therapy improved PaO₂ and also appeared to improve survival. A registered open-label randomized trial (NCT04357730) will evaluate systemic fibrinolytic therapy with t-PA versus standard care in mechanically ventilated COVID-19 patients with severe respiratory failure. A registered open-label nonrandomized pilot study (NCT04356833) will evaluate an open-label systemic fibrinolytic therapy trial (NCT04357730) will evaluate t-PA (alteplase) dosages of 50 mg (administered as a 10-mg IV bolus followed by IV administration of the remaining 40 mg over a total time of 2 hours) and 100 mg (administered as a 10-mg IV bolus dose followed by IV administration of the remaining 90 mg over a total time of 2 hours); a heparin infusion will be initiated immediately following completion of the alteplase infusion. Other dosage regimens have been evaluated in patients with ARDS associated with COVID-19, including t-PA has been proposed as a salvage treatment for COVID-19 patients (e.g., those with decompensating respiratory function who do not have access to mechanical ventilation or extracorporeal membrane oxygenation [ECMO]). However, there are currently no clinical trial data to inform this practice and a lack of clinical experience with the use of fibrinolytic agents in ARDS patients in general. Several institutions (Beth Israel Deaconess, University of Colorado Anschultz Medical Campus, Denver Health) are currently testing the use of t-PA as salvage therapy in patients with severe COVID-19.</td>
<td>utility of diagnostic signs in detecting infections. <a href="https://www.fda.gov/drugs/drug-safety-and-availability/fda-advises-patients-use-non-steroidal-anti-inflammatory-drugs-nsaids-covid-19">https://www.fda.gov/drugs/drug-safety-and-availability/fda-advises-patients-use-non-steroidal-anti-inflammatory-drugs-nsaids-covid-19</a>. Therefore, currently no compelling evidence to support an association between ibuprofen and negative outcomes in patients with COVID-19. However, some experts have recommended preferentially using acetaminophen for treatment of fever. NIH COVID-19 Treatment Guidelines Panel states that patients who are receiving NSAIA for other conditions should continue receiving the drugs; states antipyretic strategy (e.g., use of acetaminophen or NSAIA) should be no different between patients with or without COVID-19. The Surviving Sepsis Campaign COVID-19 guidelines state that until more evidence is available, use of acetaminophen over no treatment for fever control is suggested (weak recommendation).</td>
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<td>D-dimer levels, high fibrinogen levels, and microvascular and macrovascular thrombosis. 1, 2, 5-10, 13, 14, 16</td>
<td>inhaled formulation of t-PA (via nebulization) in patients with ARDS due to COVID-19; the inhaled formulation of t-PA is investigational at this time</td>
<td>an initial t-PA (alteplase) dose of 25 mg administered IV over 2 hours, followed by an IV infusion of 25 mg of t-PA over the subsequent 22 hours, with a dose not to exceed 0.9 mg/kg (Beth Israel Deaconess et al study); however, the optimum dose, route of administration, and duration of treatment remain to be determined. 1, 9, 14</td>
<td>COVID-19 under the FDA compassionate use program. Preliminary findings from the first few cases reported an initial, but transient improvement in PaO₂/FiO₂ (P/F) ratio. 9 The American Society of Hematology states that treatment of the underlying pathology is paramount in COVID-19 patients with coagulopathies; supportive care should be individualized and standard risk factors for bleeding should be considered. 8</td>
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</table>

<sup>a</sup> See US prescribing information for additional information on dosage and administration of drugs commercially available in the US for other labeled indications.
REFERENCES

ACE Inhibitors and Angiotensin II Receptor Blockers (ARBs)


Anakinra:


Anticoagulants


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Ascorbic acid:


Azithromycin:


Baloxavir:

Baricitinib:

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Chloroquine and Hydroxychloroquine:

**Corticosteroids, including methylprednisolone:**

COVID-19 Convalescent Plasma:


Epoprostenol:


Favipiravir:


Favipiravir:


HIV Protease Inhibitors:


HMG-CoA Reductase Inhibitors (statins)


Immunoglobulin (IGIV, IVIG, y-globulin):

Ivermectin:
Nebulized drugs:

Neuraminidase Inhibitors (e.g., oseltamivir):

Niclosamide:

Nitazoxanide:

Nitric Oxide (inhaled):

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NSAIDs, including ibuprofen:

Remdesivir:
Ruxolitinib

Sarilumab:

Siltuximab:

Sirolimus:

Tissue Plasminogen Activator (t-PA; alteplase):
1. Moore HB, Barrett CD, Moore EE et al. Is there a role for tissue plasminogen activator (tPA) as a novel treatment for refractory COVID-19 associated acute respiratory distress syndrome (ARDS)? J Trauma Acute Care Surg. DOI: 10.1097/TA.0000000000002694

Tocilizumab:
7. PMID 32192578 DOI: 10.1016/S0140-6736(20)30628-0.

Umifenovir:
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