As of Apr. 30, 2020, coronavirus disease 2019 (COVID-19) has resulted in more than 3 million cases and more than 210,000 deaths worldwide. The World Health Organization (WHO) has declared the COVID-19 outbreak a pandemic. The pathogen, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), is a novel Betacoronavirus that shares a phylogenetic similarity to SARS-CoV (about 79%) and Middle East respiratory syndrome (MERS)-CoV (about 50%).

Many clinical trials of potential COVID-19 treatments are underway, but current strategies for treatment are based to a considerable extent on preclinical studies and previous experiences from SARS and MERS. Clinicians have administered a number of antiviral treatments to patients with COVID-19. Optimal decision-making regarding these agents requires systematic summaries of the best available evidence about antiviral agents.

**RESEARCH**

**Efficacy and safety of antiviral treatment for COVID-19 from evidence in studies of SARS-CoV-2 and other acute viral infections: a systematic review and meta-analysis**

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**ABSTRACT**

**BACKGROUND:** Antiviral medications are being given empirically to some patients with coronavirus disease 2019 (COVID-19). To support the development of a COVID-19 management guideline, we conducted a systematic review that addressed the benefits and harms of 7 antiviral treatments for COVID-19.

**METHODS:** We searched MEDLINE, Embase, Cochrane Central Register of Controlled Trials (CENTRAL), PubMed and 3 Chinese databases (CNKI, WANFANG and SinoMed) through Apr. 19, medRxiv and Chinaxiv through Apr. 27, and Chongqing VIP through Apr. 30, 2020. We included studies of ribavirin, chloroquine, hydroxychloroquine, umifenovir (arbidol), favipiravir, interferon and lopinavir/ritonavir. If direct evidence from COVID-19 studies was not available, we included indirect evidence from studies of severe acute respiratory syndrome (SARS) and Middle East respiratory syndrome (MERS) for efficacy outcomes and other acute respiratory viral infections for safety outcomes.

**RESULTS:** In patients with nonsevere COVID-19 illness, the death rate was extremely low, precluding an important effect on mortality. We found only very low-quality evidence with little or no suggestion of benefit for most treatments and outcomes in both nonsevere and severe COVID-19. An exception was treatment with lopinavir/ritonavir, for which we found low-quality evidence for a decrease in length of stay in the intensive care unit (risk difference 5 d shorter, 95% confidence interval [CI] 0 to 9 d) and hospital stay (risk difference 1 d shorter, 95% CI 0 to 2 d). For safety outcomes, evidence was of low or very low quality, with the exception of treatment with lopinavir/ritonavir for which moderate-quality evidence suggested likely increases in diarrhea, nausea and vomiting.

**INTERPRETATION:** To date, persuasive evidence of important benefit in COVID-19 does not exist for any antiviral treatments, although for each treatment evidence has not excluded important benefit. Additional randomized controlled trials involving patients with COVID-19 will be needed before such treatments can be administered with confidence.
We provide a systematic review conducted to support a clinical practice guideline that offers recommendations to address currently used antiviral treatments (i.e., ribavirin, chloroquine, hydroxychloroquine, umifenovir, favipiravir, interferon and lopinavir/ritonavir) for COVID-19. Because remdesivir was unavailable at the time the panel determined the scope of the guideline, we did not include it in our review; however, results for the first randomized controlled trials (RCTs) of remdesivir are now available. The review includes RCTs and observational studies in patients with COVID-19, in patients with SARS and MERS, and in patients with influenza.

**Methods**

**Study design and data sources**

We adhered to the Preferred Reporting Items for Systematic Reviews and Meta-Analysis (PRISMA) Statement. Appendix 1 (available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.200647/-/DC1) presents the protocol of this systematic review. Appendix 2 (available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.200647/-/DC1) presents our detailed search strategy for 2 independent searches. We searched MEDLINE, Embase, the Cochrane Central Register of Controlled Trials (CENTRAL), PubMed and 3 Chinese databases (China National Knowledge Infrastructure [CNKI], Wanfang and SinoMed) through Apr. 19, 2020, and medRxiv and Chinaxiv preprints through Apr. 27, 2020. We also searched another Chinese database (Chongqing VIP Information) through Apr. 30, 2020. We used search terms that comprised a combination of SARS, MERS, COVID-19 and drugs of interest, as developed by an experienced medical librarian (R.C.).

Owing to concerns about inadequate evidence for safety, we performed an expanded search for safety outcomes in other acute respiratory infections that included PubMed, Embase and CENTRAL through Mar. 19, 2020. The search terms included combinations of drugs of interest, respiratory infectious diseases and terms specific for identifying RCTs. We also sought other eligible studies from reference lists of eligible published articles for both of our searches.

We considered a study eligible if it met the following criteria: patients enrolled in the trial had a diagnosis of COVID-19, SARS, MERS or other acute respiratory infectious diseases (Appendix 1); the trial involved antiviral treatments of interest (i.e., ribavirin, chloroquine, hydroxychloroquine, umifenovir, favipiravir, interferon and lopinavir/ritonavir); the trial evaluated efficacy (i.e., mortality, mechanical ventilation rate, length of stay in the intensive care unit [ICU], length of hospital stay, virologic outcomes, disease progression rate or relief of symptoms) or safety (symptomatic and serious adverse events) outcomes; and the study type was RCT, cohort study or case-control study. We did not exclude by language.

We included the best available evidence using the following hierarchy: COVID-19 RCTs; COVID-19 observational studies with adjusted analysis; RCTs involving SARS and MERS; observational studies in SARS and MERS with adjusted analysis; for safety outcomes only, RCTs addressing acute respiratory infectious diseases other than SARS and MERS (Appendix 1); observational studies without adjusted analysis; and studies comparing the drugs of interest with another antiviral agent. For each outcome, when studies from the higher categories provided evidence of higher quality than studies in lower categories, we included only the higher-quality evidence. Because of communication and time costs of informing the guideline panel of evidence summary changes, we no longer included evidence identified after Apr. 19, 2020, with very low quality that were unlikely to change the recommendations.

Pairs of reviewers independently screened titles and abstracts and reviewed the full texts of potentially eligible studies to determine eligibility. They resolved any disagreements by discussion. Two reviewers independently extracted data, including names of authors, publication year, study country or region, study design, patient population, sample size for each group, age, sex, percentage of patients who were critically ill, interventions and comparison regimen, and outcomes. Reviewers resolved any disagreement by discussion.

Two reviewers independently assessed risk of bias for each study using a modification of the Cochrane criteria for RCTs, a modification of the Newcastle–Ottawa instrument for cohort studies and an instrument developed specifically for case-control studies.

The Grading of Recommendations, Assessment, Development, and Evaluation (GRADE) approach informed the assessment of quality of evidence for each of our outcomes (Table 1). Mortality, mechanical ventilation and length of stay in the ICU were assessed only for the population of patients with severe illness, whereas we assessed rate of disease progression and symptom-based outcomes for only the nonsevere population. For efficacy outcomes, we rated down 1 level for indirectness if evidence came from studies involving patients with SARS or MERS. For safety outcomes, we did not rate down for indirectness for patients with SARS or MERS; however, we rated down 1 level for other acute respiratory infections. We present evidence using the GRADE Summary of Findings tables.

<table>
<thead>
<tr>
<th>Quality</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>We are very confident that the true effect lies close to that of the estimate of the effect.</td>
</tr>
<tr>
<td>Moderate</td>
<td>We are moderately confident in the effect estimate. The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.</td>
</tr>
<tr>
<td>Low</td>
<td>Our confidence in the effect estimate is limited. The true effect may be substantially different from the estimate of the effect.</td>
</tr>
<tr>
<td>Very low</td>
<td>We have very little confidence in the effect estimate. The true effect is likely to be substantially different from the estimate of the effect.</td>
</tr>
</tbody>
</table>
Statistical analysis
Using numbers of events and patients in RCTs and cohort studies, and adjusted relative estimates reported by original observational studies, we conducted our meta-analysis with Review Manager (version 5.3). Mantel-Haenszel random-effect models provided methods to calculate risk ratios (RRs) for RCTs and for cohort studies of dichotomous outcomes. We used DerSimonian and Laird inverse variance random-effect models to pool adjusted RRs and odds ratios (ORs). For studies that made their original data set available, we conducted a multiple regression analysis to calculate relative estimates.

Our target populations were patients with nonsevere and severe COVID-19 illness. We adhered to the WHO definition of pneumonia for severe COVID-19: fever or suspected respiratory infection, plus 1 of the following: respiratory rate > 30 breaths/min, severe respiratory distress, or peripheral oxygen saturation (SpO₂) ≤ 93% on room air.11 We identified the baseline risk of each outcome in patients with severe and nonsevere COVID-19 from published studies, choosing the most representative populations. We applied relative effects to baseline risks to estimate risk differences. When no patient in the control group experienced an event, the incidence and associated 95% confidence interval (CI) in the intervention group provided the estimated risk difference. Where hazard ratios (HRs) were provided, we estimated risk of an event in the intervention group from HRs and risk in the control group.12

Ethics approval
Ethics approval was not required for this systematic review.

Results
Figure 1 and Appendix 3 (available at www.cmaj.ca/lookup/ suppl/doi:10.1503/cmaj.200647/-/DC1) show flow charts for the study. We included 19 studies that were conducted in China (n = 11); 7 of the studies are preprints15,17,18,24,27,28,32, Saudi Arabia (n = 2), Singapore (n = 1), United States (n = 1 [preprint]19), France (n = 1) and Canada (n = 1).14-31 Two of these studies were multiregional.16,29 Seven studies used an RCT design, 11 used a cohort design and 1 used a case-control design. The studies enrolled patients with COVID-19 (n = 12), MERS (n = 2), SARS (n = 4) and influenza (n = 1). The interventions used in the studies included ribavirin (n = 3), hydroxychloroquine (n = 5), favipiravir (n = 3), interferon (n = 3), lopinavir/ritonavir (n = 2), umifenovir (n = 1), ribavirin and interferon (n = 1), and umifenovir and lopinavir/ritonavir (n = 1). Table 2 presents characteristics of the included studies. We did not find any eligible studies that addressed the use of chloroquine.

Among 7 RCTs, 418,21,26 were open label and rated down for lack of blinding (2 of these studies were preprints15,18). One RCT did not blind patients or physicians (preprint).17 We determined that the other 2 RCTs16,29 were low risk of bias (1 of these RCTs is a preprint29). Figure 2 presents the risk-of-bias assessment for RCTs. Table 3 and Table 4 present the risk-of-bias assessment for observational studies.

A large epidemiologic study in China that involved 173 patients with severe COVID-19 provided our baseline mortality estimate of 10.4%.33 Table 5 presents baseline risk data for all outcomes.

Ribavirin
Efficacy
Two retrospective cohort studies29,30 that enrolled 1334 patients with SARS (mixed severity of illness) provided mortality estimates. One of these studies was conducted in Hong Kong and Canada, which was analyzed separately,29 and the other was in Singapore.30 Pooled results suggested uncertain effects of treatment using ribavirin on mortality (OR 0.83, 95% CI 0.49 to 1.41) (Appendix 4, Supplementary Figure 1, available at www.cmaj.ca/lookup/suppl/ doi:10.1503/cmaj.200647/-/DC1). One case-control study that involved 51 patients with MERS21 (mixed severity of illness) provided similar findings (OR 0.66, 95% CI 0.04 to 12.36). Both SARS and MERS studies provided very low-quality evidence for effects of treatment using ribavirin on mortality in patients with severe COVID-19 illness (Appendix 5, Supplementary Table 2, available at www.cmaj.ca/lookup/suppl/ doi:10.1503/cmaj.200647/-/DC1).

Safety
One retrospective cohort study that involved 306 patients31 with SARS and mixed severity of illness reported that ribavirin increased the incidence of anemia (defined as a decrease in hemoglobin level of 20 g/L; OR 3.00, 95% CI 1.77 to 5.16) and bradycardia (defined as a heart rate < 55 beats/min; OR 2.30, 95% CI 1.21 to 4.20). Because both outcomes were surrogates (i.e., anemia for symptomatic anemia and bradycardia for symptomatic bradycardia), we rated down for indirectness and judged quality of evidence as very low (Appendix 5, Supplementary Tables 1 and 2).

Hydroxychloroquine
Efficacy
Three RCTs17,18,21 (2 of these RCTs are preprints17,18) that involved 240 patients with nonsevere and 2 patients with severe COVID-19 illness compared treatment with hydroxychloroquine and treatment without hydroxychloroquine, providing very low-quality evidence of minimal effects on viral clearance at day 14 (RR 0.98, 95% CI 0.89 to 1.07; Appendix 4, Supplementary Figure 2), progression from nonsevere to severe illness (RR 0.96, 95% CI 0.10 to 9.66; Appendix 4, Supplementary Figure 3) or clinical recovery at day 7 (RR 1.10, 95% CI 0.44 to 2.77).17 Hydroxychloroquine might result in a shorter duration of fever (mean difference [MD] 1 d shorter, 95% CI 0.36 to 1.64 d shorter; very low-quality evidence; Appendix 5, Supplementary Table 3).

In addition, 2 observational studies (preprints19,20) that enrolled patients with COVID-19 (181 with severe and 255 with mixed-severity illness) provided very low-quality evidence for effects of hydroxychloroquine on mortality (RR 1.48, 95% CI 0.42 to 5.24; Appendix 4, Supplementary Figure 4, and Appendix 5, Supplementary Table 4). One of these studies also reported inconclusive results of the use of hydroxychloroquine while patients were receiving mechanical ventilation (Appendix 5, Supplementary Table 4).19
Safety

Two RCTs\(^{18,21}\) (1 of these studies is a preprint\(^{19}\) that enrolled 178 patients with nonsevere and 2 patients with severe COVID-19 illness reported that no patient had diarrhea in the treatment group without hydroxychloroquine; however, 10.6% (95% CI 4.0% to 17.1%) of patients in the hydroxychloroquine treatment group had diarrhea (low-quality evidence; Appendix 5, Supplementary Tables 3 and 4). An RCT that involved 62 patients with

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**Figure 1:** Flow chart for the determination of included studies. CNKI = China National Knowledge Infrastructure.
nonsevere COVID-19 illness (preprint)\textsuperscript{17} reported an incidence of headache or rash in the intervention group of 3.2% (95% CI 0% to 9.4%), with none of these events in the control group. An RCT (preprint)\textsuperscript{18} that enrolled 148 patients with nonsevere and 2 with severe COVID-19 reported an incidence of both nausea and blurred vision in 1.4% (95% CI 0% to 4.2%) of patients and an incidence of vomiting in 2.9% (95% CI 0% to 6.8%); none of these events occurred in the control group. The quality of evidence for headache, rash, nausea, vomiting and blurred vision was very low (Appendix 5, Supplementary Tables 3 and 4).

### Table 2 (part 1 of 2): Characteristics of the included studies

<table>
<thead>
<tr>
<th>Reference</th>
<th>Dosage and administration</th>
<th>Antiviral agent comparison</th>
<th>Study design</th>
<th>Country</th>
<th>Participant population</th>
<th>No. of participants</th>
<th>Age, mean ± SD*</th>
<th>Percentage of population who were male</th>
<th>Percentage of population with severe disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cai et al., 2020\textsuperscript{14}</td>
<td>Favipiravir 1600 mg po b.i.d. on day 1 and 600 mg po b.i.d. on days 2–14 plus interferon-(\alpha) (60 (\mu)g b.i.d.) by aerosol inhalation</td>
<td>Lopinavir/ritonavir (200 mg/50 mg) 500 mg po b.i.d. on days 1–14 plus interferon-(\alpha) 60 (\mu)g b.i.d. by aerosol inhalation</td>
<td>Cohort China</td>
<td>Nonsevere COVID-19</td>
<td>80</td>
<td>47.0 (35.8–61.0)†</td>
<td>43.8</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Chen et al., 2020\textsuperscript{15}‡</td>
<td>Favipiravir 1600 mg po b.i.d. on day 1 and 600 mg po b.i.d. for 7–10 d</td>
<td>Umifenovir (200 mg) po t.i.d. for 7–10 d</td>
<td>RCT China</td>
<td>COVID-19 with mixed severity</td>
<td>236</td>
<td>NR</td>
<td>46.6</td>
<td>11.4</td>
<td></td>
</tr>
<tr>
<td>MDVI\textsuperscript{16}</td>
<td>Favipiravir 1200 mg po b.i.d. for 1 d, followed by 800 mg po b.i.d. for 4 d</td>
<td>Placebo</td>
<td>RCT Multiple countries</td>
<td>Influenza with unspecified severity</td>
<td>386</td>
<td>42.7 (20.0–80.0)¶</td>
<td>45.3</td>
<td>NR</td>
<td></td>
</tr>
<tr>
<td>Chen et al., 2020\textsuperscript{17}‡</td>
<td>Hydroxychloroquine (200 mg) po b.i.d. for 5 d</td>
<td>No hydroxychloroquine</td>
<td>RCT China</td>
<td>Nonsevere COVID-19</td>
<td>62</td>
<td>44.7 ± 15.3</td>
<td>46.8</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Tang et al., 2020\textsuperscript{18}‡</td>
<td>Hydroxychloroquine: loading dose of 1200 mg daily for 3 d followed by a maintenance dose of 800 mg daily for remaining treatment days (total treatment duration: 2 wk for patients with mild/moderate disease or 3 wk for patients with severe disease)</td>
<td>No hydroxychloroquine</td>
<td>RCT China</td>
<td>COVID-19 with mixed severity</td>
<td>150</td>
<td>46.1 ± 14.7</td>
<td>54.7</td>
<td>1.3</td>
<td></td>
</tr>
<tr>
<td>Magagnoli et al., 2020\textsuperscript{19}‡</td>
<td>Hydroxychloroquine dose not mentioned</td>
<td>No hydroxychloroquine</td>
<td>Cohort US</td>
<td>COVID-19 with mixed severity</td>
<td>255</td>
<td>NR</td>
<td>100.0</td>
<td>NR</td>
<td></td>
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<tr>
<td>Mahervas et al., 2020\textsuperscript{20}‡</td>
<td>Hydroxychloroquine 600 mg/d</td>
<td>No hydroxychloroquine</td>
<td>Cohort France</td>
<td>COVID-19 with mixed severity</td>
<td>181</td>
<td>60.0 (52.0–68.0)†</td>
<td>71.1</td>
<td>NR</td>
<td></td>
</tr>
<tr>
<td>Chen et al., 2020\textsuperscript{17}‡</td>
<td>Hydroxychloroquine (400 mg) po daily for 5 d plus interferon-(\alpha) by aerosol inhalation (80.0% of patients used umifenovir)</td>
<td>Interferon-(\alpha) by aerosol inhalation (66.7% of patients used umifenovir and 13.3% used lopinavir/ritonavir)</td>
<td>RCT China</td>
<td>Nonsevere COVID-19</td>
<td>30</td>
<td>48.6 ± 4.1</td>
<td>70.0</td>
<td>0</td>
<td></td>
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<tr>
<td>Interferon versus no interferon, ribavirin versus no ribavirin</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Al Ghandi et al., 2016\textsuperscript{21}</td>
<td>A: interferon-(\alpha) B: interferon-(\beta) C: ribavirin</td>
<td>A: no interferon-(\alpha) B: no interferon-(\beta) C: no ribavirin</td>
<td>Case–control Saudi Arabia</td>
<td>MERS with mixed severity</td>
<td>51</td>
<td>54.0 (36.5–58.0)†</td>
<td>78.4</td>
<td>37.3</td>
<td></td>
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<tr>
<td>Interferon plus ribavirin versus ribavirin alone</td>
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<tr>
<td>Shalhoub et al., 2015\textsuperscript{22}</td>
<td>Interferon-x2a (180 (\mu)g) by sc injection weekly; interferon-(\beta)1a (44 (\mu)g) by sc injection 3 times per week plus ribavirin (2 g loading dose) po followed by 600 mg q.12h</td>
<td>Ribavirin (2 g loading dose) po followed by 600 mg q.12h</td>
<td>Cohort Saudi Arabia</td>
<td>MERS with mixed severity</td>
<td>32</td>
<td>60.0 (42.0–73.0)†</td>
<td>56.0</td>
<td>NR</td>
<td></td>
</tr>
</tbody>
</table>
One RCT that enrolled 23 patients with nonsevere COVID-19 illness (preprint)\(^2^8\) provided limited evidence of uncertain effects of treatment using umifenovir on viral clearance at day 14, cough alleviation at day 7, fever at day 7 and progression to severe illness (Appendix 5, Supplementary Table 5). With additional indirectness, this trial reported even lower-quality evidence for delayed viral clearance in patients with severe COVID-19 illness (Appendix 5, Supplementary Table 6). An observational study in Wuhan, China, that enrolled 504 patients with mixed severities of COVID-19 illness (preprint)\(^3^2\) reported a large decrease in mortality among those who received umifenovir (OR 0.18, 95% CI 0.08 to 0.45). However, we found the quality of evidence to be very low because of the observational study design and suboptimal adjustment for disease severity (Appendix 5, Supplementary Table 6).

### Table 2 (part 2 of 2): Characteristics of the included studies

<table>
<thead>
<tr>
<th>Reference</th>
<th>Study intervention</th>
<th>Antiviral agent comparison</th>
<th>Study design</th>
<th>Country</th>
<th>Participant population</th>
<th>No. of participants</th>
<th>Age, mean ± SD*</th>
<th>Percentage of population who were male</th>
<th>Percentage of population with severe disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zhou et al., 2020(^‡)‡</td>
<td>Interferon-α2b (5 mIU) by aerosol inhalation twice daily plus umifenovir (200 mg) po 3 times daily</td>
<td>Umifenovir 200 mg po 3 times daily</td>
<td>Cohort</td>
<td>China</td>
<td>COVID-19 with mixed severity</td>
<td>70</td>
<td>48.7 ± 18.1</td>
<td>44.3</td>
<td>NR</td>
</tr>
<tr>
<td>Li et al., 2005(^‡)</td>
<td>Interferon-α (1 mIU/d) by im or sc injection for 6–10 d</td>
<td>No interferon</td>
<td>Cohort</td>
<td>China</td>
<td>SARS with mixed severity</td>
<td>87</td>
<td>28.1 ± 9.5</td>
<td>18.4</td>
<td>71.3</td>
</tr>
<tr>
<td>Cao et al., 2020(^‡)</td>
<td>Lopinavir/ritonavir (400/100 mg) po b.i.d. for 14 d</td>
<td>No lopinavir/ritonavir</td>
<td>RCT</td>
<td>China</td>
<td>Severe COVID-19</td>
<td>199</td>
<td>58.0 (49.0–68.0) †</td>
<td>60.3</td>
<td>100.0</td>
</tr>
<tr>
<td>Yan et al., 2020(^‡)</td>
<td>Lopinavir/ritonavir (400/100 mg) po b.i.d. for 10 d or longer</td>
<td>No lopinavir/ritonavir</td>
<td>Cohort</td>
<td>China</td>
<td>COVID-19 with mixed severity</td>
<td>120</td>
<td>52.0 (35.0–63.0) †</td>
<td>45</td>
<td>25.8</td>
</tr>
<tr>
<td>Li et al., 2020(^‡)‡</td>
<td>A: lopinavir/ritonavir (200 mg/50 mg) 500 mg po q.12h for 7–14 d or umifenovir (200 mg) po t.i.d. for 7–14 d</td>
<td>No lopinavir/ritonavir or umifenovir</td>
<td>RCT</td>
<td>China</td>
<td>Nonsevere COVID-19</td>
<td>44</td>
<td>49.4 ± 14.9</td>
<td>47.7</td>
<td>0</td>
</tr>
<tr>
<td>Lau et al., 2009(^‡)</td>
<td>Ribavirin</td>
<td>No ribavirin</td>
<td>Cohort</td>
<td>Canada</td>
<td>SARS with mixed severity</td>
<td>953</td>
<td>48.7, 36.8 NR</td>
<td>48.7, 36.8 NR</td>
<td>NR</td>
</tr>
<tr>
<td>Leong et al., 2004(^‡)</td>
<td>Ribavirin (1.2 g) po t.i.d. or 400 mg by intravenous injection q.8h for sicker patients who are severely ill and those who could not take it by mouth; mean treatment duration was 5.6 d</td>
<td>No ribavirin</td>
<td>Cohort</td>
<td>Singapore</td>
<td>SARS with mixed severity</td>
<td>229</td>
<td>39.1 ± 16.8</td>
<td>31.9</td>
<td>20.1 (as outcome)</td>
</tr>
<tr>
<td>Muller et al., 2007(^‡)</td>
<td>Ribavirin (2 g loading dose) by intravenous injection, followed by 1 g q.8h for 4 d, followed by 500 mg q.6h for 3 d**</td>
<td>No ribavirin</td>
<td>Cohort</td>
<td>Canada</td>
<td>SARS with mixed severity</td>
<td>306</td>
<td>NR</td>
<td>37.3</td>
<td>41.6†</td>
</tr>
<tr>
<td>Liu et al., 2020(^‡)‡</td>
<td>Umifenovir (dose not mentioned)</td>
<td>No umifenovir</td>
<td>Cohort</td>
<td>China</td>
<td>COVID-19 with mixed severity</td>
<td>504</td>
<td>59.5 ± 14.9</td>
<td>51.4</td>
<td>NR</td>
</tr>
</tbody>
</table>

Note: b.i.d. = twice a day, COVID-19 = coronavirus disease 2019, im = intramuscular, IQR = interquartile range, MERS = Middle East respiratory syndrome, NR = not reported, po = by mouth, q 6h = every 6 hours, q 8h = every 8 hours, q 12h = every 12 hours, RCT = randomized controlled trial, SARS = severe acute respiratory syndrome, sc = subcutaneous, SD = standard deviation, t.i.d. = 3 times per day.

*Unless stated otherwise.
†Median (IQR).
‡Preprint.
§The course of treatment in both groups was 7–10 days. If necessary, the treatment time could have been extended to 10 days according to the judgment of researchers.
¶Mean (range).
**Only 155 of 183 participants received this treatment regimen; the other 28 patients received several lower-dose treatment regimens.
††Calculated from the baseline characteristic, admission oxygen saturation < 95%.

### Umifenovir

**Efficacy**

One RCT that enrolled 23 patients with nonsevere COVID-19 illness (preprint)\(^2^8\) provided limited evidence of uncertain effects of treatment using umifenovir on viral clearance at day 14, cough alleviation at day 7, fever at day 7 and progression to severe illness (Appendix 5, Supplementary Table 5). With additional indirectness, this trial reported even lower-quality evidence for delayed viral clearance in patients with severe COVID-19 illness (Appendix 5, Supplementary Table 6). An observational study in Wuhan, China, that enrolled 504 patients with mixed severities of COVID-19 illness (preprint)\(^3^2\) reported a large decrease in mortality among those who received umifenovir (OR 0.18, 95% CI 0.08 to 0.45). However, we found the quality of evidence to be very low because of the observational study design and suboptimal adjustment for disease severity (Appendix 5, Supplementary Table 6).
Safety
The RCT that enrolled 23 patients with nonsevere COVID-19 illness (preprint) reported that no patients in either the treatment or control groups had diarrhea or decreased appetite (very low-quality evidence; Appendix 5, Supplementary Tables 5 and 6).

Favipiravir

Efficacy
One RCT that enrolled 236 patients (preprint) of mixed severity compared interferon-α plus umifenovir with umifenovir alone and reported on time to viral clearance and length of hospital stay. We performed a multiple linear regression analysis based on the original data set (Appendix 6, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.200647/-/DC1). We found that patients receiving interferon-α tended to have a shorter time to viral clearance (MD 4.6 d, 95% CI –0.5 to 9.6 d) and and a shorter length of hospital stay (MD 4.4 d, 95% CI –1.5 to 10.3 d) (Appendix 5, Supplementary Tables 11 and 12). However, the quality of evidence was very low because of concurrent use of umifenovir and imprecision of the results.

Safety
One retrospective cohort study (preprint) and 1 case–control study that involved a total of 83 patients with MERS of mixed severity provided very low-quality evidence and raised the possibility of decreased mortality with interferon-α (OR 0.23, 95% CI 0.04 to 1.32) (Appendix 4, Supplementary Figure 5, and Appendix 5, Supplementary Table 14) and interferon-β (OR 0.37, 95% CI 0.07 to 2.05) (Appendix 4, Supplementary Figure 6, and Appendix 5, Supplementary Table 15).

Lopinavir/ritonavir

Efficacy
One RCT that enrolled 199 patients with severe COVID-19 (preprint) compared treatment with lopinavir/ritonavir with no lopinavir/ritonavir treatment and reported on mortality, viral clearance at day 14, mechanical ventilation and length of stay in ICU and hospital. Another RCT compared treatment with lopinavir/ritonavir with no lopinavir/ritonavir treatment in 28 patients with nonsevere COVID-19 (preprint) and reported on mortality, viral clearance at day 14, cough alleviation at day 7, progression from nonsevere to severe illness and fever at day 7. Because no patients died in the latter RCT, we included only mortality data from the RCT involving patients with severe illness. Another observational study enrolling 120 patients with mixed severity of COVID-19 reported on viral clearance at day 23 (preprint). We conducted a meta-analysis on viral clearance at day 14 (Appendix 4, Supplementary Figure 7).
For nonsevere COVID-19 patients, lopinavir/ritonavir may provide little or no reduction in viral clearance at day 14 (RD –0.7%, 95% CI –17.1% to 20.7%, low-quality evidence; 26 [preprint]). The observational study raised the possibility of increased viral clearance at day 23 with lopinavir/ritonavir treatment, but the study failed to adjust for disease severity, making this result of very low quality (preprint).27 Also, there was very low-quality evidence of uncertain effects of lopinavir/ritonavir on cough alleviation at day 7, progression to severe illness, fever at day 7 and length of hospital stay14 (preprint) 19 (Appendix 5, Supplementary Table 16).

For severe COVID-19 patients, lopinavir/ritonavir may result in a small decrease in mortality (RD 2.4% fewer deaths, 95% CI 5.7% decrease to 3.1% increase, low-quality evidence), and reductions in length of ICU stay (RD 5 d shorter, 95% CI 0 to 9 d, low-quality evidence) and hospital stay (RD 1 d shorter, 95% CI 0 to 2 d, low-quality evidence)26 (Appendix 5, Supplementary Table 17).

<table>
<thead>
<tr>
<th>Study</th>
<th>From the same population</th>
<th>Assessment of exposure</th>
<th>Outcome not present at start</th>
<th>Adjustment</th>
<th>Assessment of prognostic factors</th>
<th>Assessment of outcome</th>
<th>Adequate follow-up</th>
<th>Similar co-interventions</th>
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Table 3: Risk-of-bias assessment for included cohort studies using the modified Newcastle–Ottawa scale

<table>
<thead>
<tr>
<th>Study</th>
<th>Assessment of exposure</th>
<th>Cases had developed the outcome and controls had not</th>
<th>Selection of cases</th>
<th>Selection of controls</th>
<th>Matching or adjustment</th>
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Table 4: Risk-of-bias assessment in the case–control study using the modified Newcastle–Ottawa scale

Table 5: Baseline risk for patients with severe and nonsevere coronavirus disease 2019

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Patients with nonsevere COVID-19</th>
<th>Patients with severe COVID-19</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mortality</td>
<td>NA</td>
<td>10.4%33</td>
</tr>
<tr>
<td>Length of hospital stay</td>
<td>11 d23</td>
<td>13 d23</td>
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<tr>
<td>Length of stay in the ICU</td>
<td>NA</td>
<td>11 d26</td>
</tr>
<tr>
<td>Mechanical ventilation</td>
<td>NA</td>
<td>38.7%23</td>
</tr>
<tr>
<td>Viral clearance at day 14</td>
<td>71.4% (preprint)28</td>
<td>56.3%28</td>
</tr>
<tr>
<td>Viral clearance at day 7</td>
<td>71.4% (preprint)28</td>
<td>32.4%28</td>
</tr>
<tr>
<td>Progressing from nonsevere to severe disease</td>
<td>14.3% (preprint)28</td>
<td>NA</td>
</tr>
</tbody>
</table>

Note: COVID-19 = coronavirus disease 2019, ICU = intensive care unit, NA = not applicable.
*Used day 5 instead of day 7.
Safety
One RCT that involved 194 patients with severe COVID-19\textsuperscript{26} and another RCT that involved 28 patients with nonsevere COVID-19 (preprint)\textsuperscript{28} reported no diarrhea in their control groups. The incidence of diarrhea in the intervention group was 6.0\% (95\% CI 1.7\% to 10.4\%\textsuperscript{26} (preprint))\textsuperscript{28} moderate-quality evidence; Appendix 5, Supplementary Tables 16 and 17). The RCT with 194 patients\textsuperscript{28} reported that lopinavir/ritonavir probably increased nausea (MD 9.5\%, 95\% CI 3.6\% to 15.4\%) and vomiting (MD 6.3\%, 95\% CI 1.4\% to 11.2\%) (both moderate-quality evidence; Appendix 5, Supplementary Tables 16 and 17). This study also reported very low-quality evidence that raised the possibility of an increase in stomach ache (Appendix 5, Supplementary Tables 16 and 17).

**Interpretation**

Our systematic review did not find persuasive evidence of benefit for any antiviral agent in patients with COVID-19. We found no direct evidence for treatment using ribavirin in a population of patients with COVID-19, and results from studies evaluating SARS or MERS provided no support for a reduction in mortality with ribavirin treatment.\textsuperscript{22,29,30} We found that interferon did not show a benefit on viral clearance or length of hospital stay in patients with COVID-19.

Hydroxychloroquine and umifenovir failed to show benefit in viral clearance, disease progression or symptom relief in patients with nonsevere COVID-19. In patients with severe COVID-19, treatment using hydroxychloroquine did not show reductions in mortality or mechanical ventilation. Umifenovir appeared to reduce mortality based on 1 observational study with very low-quality evidence. Very low-quality evidence raised the possibility that favipiravir may accelerate clinical recovery relative to umifenovir and accelerate viral clearance relative to lopinavir/ritonavir.

Included RCTs addressed lopinavir/ritonavir in patients with both nonsevere and severe COVID-19. However, the sample size in the RCT that evaluated nonsevere COVID-19 was only 28 patients, resulting in very wide CIs for all outcomes. Based on the RCT evaluating patients with severe COVID-19, it is possible that lopinavir/ritonavir reduced 28-day mortality, length of ICU stay and length of hospital stay, but the evidence was of low quality. Moderate-quality evidence showed substantial increases in gastrointestinal adverse effects with lopinavir/ritonavir.

Adverse effects remain a concern with each of these drugs. Gastrointestinal upset and potential drug–drug interactions are the primary concerns with lopinavir/ritonavir. Hydroxychloroquine and chloroquine widen the QT-interval and based on case reports from the FDA Adverse Event Reporting System database, the US Food and Drug Administration issued a warning about the risk of drug-induced sudden cardiac death associated with use of chloroquine or hydroxychloroquine with or without azithromycin in patients with COVID-19.\textsuperscript{34}

Strengths of our review include a study team with methodologic, pharmacologic and clinical expertise from working directly to treat patients with COVID-19, consideration of both direct and indirect evidence, a comprehensive and current literature search, and review of eligibility, risk of bias and data abstraction in duplicate. By using the GRADE approach we focused on the highest-quality evidence available and carefully considered indirectness. It also directed us to focus on absolute effects and to produce succinct, informative evidence summaries using table formats.

Since the COVID-19 pandemic began, other efforts have been made to summarize the available evidence about antiviral treatments. Although previous reviews of studies of antiviral treatments in MERS and SARS have been published, they have not been brought together or put in the context of COVID-19 using a rigorous methodologic perspective.\textsuperscript{35–41} For example, the Public Health Agency of Canada (PHAC) published a rapid review on the efficacy and safety of antiviral or antibody treatments for coronavirus.\textsuperscript{35} They included all the known antiviral treatments and antibodies for their potential treatment in coronavirus and searched for all types of studies including preclinical (animal) studies. The methodology of such efforts, including that of PHAC, is limited in 2 aspects. First, they did not use a formal system such as GRADE for rating the certainty of the evidence. In the context of applying evidence from one patient group to another, rating systems need a formal approach to evaluating indirectness — how skeptical we should be when we apply, for instance, evidence from studies evaluating SARS and MERS to COVID-19. Such ratings are crucial for decision-makers to understand evidence credibility. Second, they did not calculate the absolute risk of these agents based on a baseline risk of patients with COVID-19, which limited their application to real-world management of COVID-19. Our review addressed these issues.

Results from ongoing RCTs (preprint)\textsuperscript{42} will certainly increase the quality of the evidence and may provide convincing evidence of benefit that our review did not. Nevertheless, clinicians need guidance at present, and our review serves that purpose.

In vitro and animal studies that evaluated treatment of COVID-19 using remdesivir, chloroquine and hydroxychloroquine have shown inhibitory effects against SARS-CoV-2, which may be promising for antiviral treatment.\textsuperscript{43–45} Investigators have also reported that umifenovir suppresses reproduction of SARS-CoV-2 in cell cultures.\textsuperscript{46} Cell culture and animal studies have provided evidence of activity of ribavirin, high-dose interferon\textsuperscript{47,48} and lopinavir/ritonavir\textsuperscript{49–51} against coronaviruses. Given that each drug has adverse effects, such studies do not provide sufficient rationale for use in humans with COVID-19.\textsuperscript{52}

**Limitations**

The primary limitation of our review is the very low-quality evidence that is currently available to inform the benefit and harms of available antiviral agents, which suggests uncertainty about their effects. This uncertainty comes primarily from estimates of relative effects but also from estimates of baseline risk in COVID-19 that came from single studies with limited sample sizes.

In addition, we restricted our review to interventions in which there was some published evidence. However, given the uncertainty about individual agents, our conclusions of uncertainty about benefit also apply to combinations of these agents. Moreover, combinations of agents are likely to have greater harms than single agents.
Conclusion
This review provides evidence to support COVID-19 guideline recommendations. To date, persuasive evidence of important benefit does not exist for any antiviral treatment, although important benefit has not been excluded for each agent. Owing to the very low risk of death in patients with nonsevere COVID-19, antiviral treatment will not result in important reductions to mortality in these patients. Confident administration of any antiviral treatment requires the conduct of RCTs showing patient-relevant benefits.

References

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**Contributors:** Wei Liu, Pengxiang Zhou, Ken Chen, Zhikang Ye, Fang Liu, Xiaotong Li, Na He, Ning Shen, Suodi Zhai and Gordon Guyatt contributed to the conception or design of the study. Wei Liu, Pengxiang Zhou and Ken Chen drafted the manuscript. Wei Liu, Pengxiang Zhou, Ken Chen, Zhikang Ye, Fang Liu, Xiaotong Li, Na He, Ziyang Wu, Qi Zhang, Xuepeng Gong, Qiyu Tang, Xin Du, Yingqiu Ying, Xiaohan Xu, Yahui Zhang, Jinyu Liu, Yun Li, Ning Shen, Rachel Couban, Quazi Ibrahim, Gordon Guyatt and Suodi Zhai acquired, analyzed or interpreted the data. All of the authors revised the manuscript critically for important intellectual content, gave final approval of the version to be published and agreed to be accountable for all aspects of the work. Wei Liu, Pengxiang Zhou and Ken Chen contributed equally to this work.

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**Data sharing:** All of our data are available as tables and figures in the article or in the appendices.

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