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### Key to abbreviations

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<th>Full Name</th>
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### Colour Legend

- **Red**: These drugs should not be coadministered
- **Yellow**: Potential interaction which may require a dose adjustment or close monitoring.
- **Green**: Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
- **White**: No clinically significant interaction expected

---

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Please note that if a drug is not listed it cannot automatically be assumed it is safe to coadminister. No recommendation to use experimental therapy for COVID-19 is made. Drug interaction data for many agents are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers.
**Anaesthetics & Muscle Relaxants**

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**Text Legend**

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
↑↑ Potential increased exposure of COVID drug
↓↓ Potential decreased exposure of COVID drug
↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

This interaction involves drugs identified by [www.crediblemeds.org](http://www.crediblemeds.org) as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined.

Note, please check product labels for any additional cardiac warnings.
Coadministration should be avoided due to the increased risk of haematological toxicity.

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Text Legend

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
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Notes:

Codeine, dihydrocodeine + CLQ or HCLQr
Potential decrease of the analgesic effect due to the reduced conversion to the active metabolite.

Codeine and Tramadol + LPV/r
Potential decrease of the analgesic effect due to the reduced conversion to the active metabolite.

Diamorphine and Morphine + ATv
No effect on systemic exposure but inhibition of P-gp by atazanavir at the blood-brain barrier could potentiate the opiate effect in the CNS.

Diamorphine and Morphine + LPV/r
Ritonavir could reduce systemic exposure of diamorphine and morphine due to induction of glucuronidation. Ritonavir also inhibits P-gp at the blood-brain barrier and could potentiate the opiate effect in the CNS.

Hydrocodone + ATV or LPV/r
Hydrocodone concentrations are increased, but concentrations of the metabolite hydromorphone (which has also analgesic activity) are reduced.

Metamizole + CLQ, HCLQ, IFN-β, RBV, TCZ
Coadministration should be avoided due to the increased risk of haematological toxicity.

Paracetamol + FAVI
The daily dose of paracetamol in adults should be no more than 3000 mg/day (rather than 4000 mg/day).

Key to abbreviations

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<td>Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required</td>
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Key to abbreviations

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<th>ATV</th>
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**Text Legend**

- ↑ Potential increased exposure of the comedication
- ↓ Potential decreased exposure of the comedication
- ↑↑ Potential increased exposure of COVID drug
- ↓↓ Potential decreased exposure of COVID drug
- ↔ No significant effect

**Notes:**

**Amiodarone + LPV/r**

The European product label for LPV/r contraindicates coadministration but the US product label suggests caution and concentration monitoring of amiodarone.

**Colour Legend**

- These drugs should not be coadministered
- Potential interaction which may require a dose adjustment or close monitoring.
- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
- No clinically significant interaction expected

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Antibacterials

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Text Legend

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
⇑ Potential increased exposure of COVID drug
⇓ Potential decreased exposure of COVID drug
→ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

This interaction involves drugs identified by [www.crediblemeds.org](http://www.crediblemeds.org) as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined.

Note, please check product labels for any additional cardiac warnings.

Notes:

No interactions are expected with the COVID-19 therapies listed and the following antibacterials:

- amikacin, amoxicillin, capreomycin, cefalexin, cefazolin, cefotaxime, ceftriaxone, chloramphenicol, clavulanic acid, cloxacillin, cycloserine, dapsone, doxycycline, ethapenem, ethambutol, ethionamide, fluvoxacin, gentamicin, imipenem/cilastatin, isoniazid, kanamycin, meropenem, nitrofurantoin, para-aminosalicylic acid, penicillins, piperaclilin, rifaximin, spectinomycin, streptomycin, tazobactam, tetracyclines, vancomycin.

Clarithromycin + ATV or LPV/r

A dose reduction of clarithromycin may be required for patients with impaired renal function. Refer to product labels for details.

Delamanid + ATV or LPV/r

Coadministration is expected to increase concentrations of DM-6705, a delamanid metabolite which is associated with QT prolongation. Frequent ECG monitoring is recommended.

Linezolid + RBV

Myelosuppression has been reported with both linezolid and ribavirin. Close monitoring of blood counts is recommended.

Linezolid + IFN-β or TCZ

Caution is required due to potential additive haematological toxicity.

Metronidazole and Tindazole + LPV/r

No interaction is expected with lopinavir tablets. Coadministration is not recommended with lopinavir oral solution as it contains alcohol.

Pyrazinamide + FAVI

No effect on pyrazinamide concentrations but coadministration increased blood uric acid concentrations. Monitor uric acid.

Color Legend

- These drugs should not be coadministered
- Potential interaction which may require a dose adjustment or close monitoring
- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required
- No clinically significant interaction expected
Monitor INR with vitamin K antagonists (e.g., acenocoumarol, phenprocoumon, warfarin). However, the US product label recommends no dose modification.

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**Text Legend**
- ↑ Potential increased exposure of the comedication
- ↓ Potential decreased exposure of the comedication
- ↑↑ Potential increased exposure of COVID drug
- ↓↓ Potential decreased exposure of COVID drug
- ↔ No significant effect

**Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.**

- ♥ This interaction involves drugs identified by www.credibledrugs.org as having a known, probable or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. Note, please check product labels for any additional cardiac warnings.

**Notes:**
- **Apixaban + LPV/r**
  The US product label for apixaban suggests to use apixaban at a reduced dose (2.5 mg twice daily) if needed.

- **Betrixaban + ATV or LPV/r**
  The US product label for betrixaban recommends for patients receiving or starting a strong P-gp inhibitor to reduce betrixaban dose and use an initial dose of 80 mg followed by 40 mg once daily.

- **Clopidogrel + ATV or LPV/r**
  Decreased conversion to active metabolite leading to non-responsiveness to clopidogrel. Prasugrel should be preferred to clopidogrel with ATV or LPV/r.

- **Edoxaban + ATV or LPV/r**
  The European product label for edoxaban states to consider a dose reduction of edoxaban from 60 mg to 30 mg with strong P-gp inhibitors, however, the US product label recommends no dose modification.

- **Prasugrel + ATV or LPV/r**
  Concentrations of active metabolite are reduced but without a significant reduction in prasugrel activity.

- **Vitamin K antagonists + ATV or LPV/r**
  Monitor INR with vitamin K antagonists (e.g., acenocoumarol, phenprocoumon, warfarin).
Anticonvulsants

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Text Legend

- ↑ Potential increased exposure of the comedication
- ↓ Potential decreased exposure of the comedication
- ↔ Potential increased exposure of COVID drug
- ↓ Potential decreased exposure of COVID drug
- ↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

This interaction involves drugs identified by www.crediblemeds.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined.

Note, please check product labels for any additional cardiac warnings.

Notes:
- **Valproate + LPV/r**
  - Case report of a 48% decrease in valproate concentration in previously stable patient who developed exacerbated mania on starting lopinavir/ritonavir; dose increase of valproate was required.

Key to abbreviations

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<tr>
<th>ATV</th>
<th>Alazanavir</th>
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Colour Legend

- These drugs should not be coadministered
- Potential interaction which may require a dose adjustment or close monitoring.
- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
- No clinically significant interaction expected
Antidepressants

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**Text Legend**

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
↑↑ Potential increased exposure of COVID drug
↓↓ Potential decreased exposure of COVID drug
↔ No significant effect

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**Key to abbreviations**

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Anti-diabetics

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**Text Legend**

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
↕ Potential increased exposure of COVID drug
↕↓ Potential decreased exposure of COVID drug
↔ No significant effect

**Notes:**

**Antidiabetics + CLQ or HCLQ**

Chloroquine and hydroxychloroquine may enhance the effects of a hypoglycemic treatment and a decrease in doses of insulin or other antidiabetic drugs may be required.

**Canagliflozin + LPV/r**

If coadministration is deemed necessary, increasing canagliflozin to 300 mg once daily may be considered if patients are currently tolerating canagliflozin 100 mg once daily, have an eGFR ≥60 mL/min/1.73m² or CrCl ≥60 mL/min, and require additional glycaemic control. Other glucose-lowering therapies should be considered for patients with an eGFR 45 mL/min/1.73m² to <60 mL/min/1.73m² or CrCl 45 mL/min to <60 mL/min taking canagliflozin 100 mg who are receiving concurrent therapy with a UGT enzyme inducer and who require additional glycaemic control.

**Linagliptin + LPV/r**

The increase in linagliptin exposure is not considered clinically significant as it is mainly eliminated unchanged and has a large safety window.

**Saxagliptin + ATV or LPV/r:**

The US product label for saxagliptin states the recommended dose of saxagliptin to be 2.5 mg once daily when coadministered with strong CYP3A4/5 inhibitors.

**Sitagliptin + ATV or LPV/r**

The increase in sitagliptin exposure is not considered clinically significant as it is mainly eliminated unchanged and has a large safety window.

**Key to abbreviations**

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<th>ATV</th>
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**Colour Legend**

- These drugs should not be coadministered
- Potential interaction which may require a dose adjustment or close monitoring.
- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
- No clinically significant interaction expected
Antifungals

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**Text Legend**

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
↑ Potential increased exposure of COVID drug
↓ Potential decreased exposure of COVID drug
↔ No significant effect

**Notes:**

Griseofulvin + LPV/r

LPV/r oral solution contains alcohol. Consumption of alcohol in association with griseofulvin can result in a ‘disulfiram-like’ type reaction. No such interaction is expected with LPV/r tablets.

Itraconazole or Ketoconazole + ATV or LPV/r

The daily dose of itraconazole or ketoconazole should not exceed 200 mg.

Voriconazole + ATV

The effect of atazanavir on voriconazole exposure is dependent on CYP2C19 metaboliser status. In the majority of patients decreases in both voriconazole and atazanavir exposures may be expected, leading to loss of therapeutic effect and possible development of resistance. The European SmPC for atazanavir recommends a patient's CYP2C19 genotype should be performed if feasible. In patients without a functional CYP2C19 allele, increased voriconazole exposures are expected.

Voriconazole + LPV/r

Coadministration may result in bidirectional interactions leading to increased concentrations of lopinavir/ritonavir and an increase or decrease in voriconazole. Administration of voriconazole with ritonavir (100 mg twice daily) decreased voriconazole AUC by 39%.

**Key to abbreviations**

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<th>Atazanavir</th>
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Antipsychotics/Neuroleptics

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Text Legend

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
★ Potential increased exposure of COVID drug
◆ Potential decreased exposure of COVID drug
→ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

★ This interaction involves drugs identified by www.crediblemeds.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. Note, please check product labels for any additional cardiac warnings.

Notes:

Clozapine + CLQ, HCLQ or RBV
The risk of haematological toxicity may be potentially increased as clozapine, ribavirin, chloroquine and hydroxychloroquine can cause myelosuppression. Closely monitor haematological parameters.

Clozapine + IFN-β or TCZ
Caution is required due to potential additive haematological toxicity.

Quetiapine + ATv or LPV/r
Coadministration contraindicated in the European product label for quetiapine, however, US product label recommends quetiapine should be reduced to one sixth of the original dose if coadministered with a potent CYP3A4 inhibitor.

Key to abbreviations

ATV: Atazanavir
CLQ: Chloroquine
FAVI: Favipiravir
HCLQ: Hydroxychloroquine
IFN-β: Interferon beta
LPV/r: Lopinavir/Ritonavir
RBV: Ribavirin
TCZ: Tocilizumab

Colour Legend

These drugs should not be coadministered
Potential interaction which may require a dose adjustment or close monitoring.
Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
No clinically significant interaction expected
Antivirals – Covid-19 therapies

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**Text Legend**

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
† Potential increased exposure of COVID drug
¶ Potential decreased exposure of COVID drug
↔ No significant effect

**Notes:**

**ATV + LPV/r**
These drugs are not intended to be combined for the treatment of COVID-19.

**ATV + RBV**
A substantial proportion of patients receiving atazanavir experienced significant hyperbilirubinemia and jaundice following initiation of ribavirin and pegylated interferon for the treatment of hepatitis C. Ribavirin-related haemolysis resulted in increased production of bilirubin, the normal clearance of which is by UGT1A1, an enzyme inhibited by atazanavir.

**CLQ or HCLQ**
Chloroquine and hydroxychloroquine should not be coadministered.

**CLQ or HCLQ + LPV/r**
LPV/r may increase concentrations of chloroquine or hydroxychloroquine, but to a moderate extent. Since LPV/r and chloroquine or hydroxychloroquine can cause QT prolongation, ECG monitoring is recommended when coadministering these agents.

**CLQ + IFN-β, RBV or TCZ**
Use with caution due to potential additive toxicity.

**RBV + TCZ**
The risk of haematological toxicity may be potentially increased as ribavirin and tocilizumab can cause myelosuppression. Closely monitor haematological parameters.

**IFN-β + TCZ**
Use with caution due to increased risk of haematological toxicity.
## Antivirals – HCV DDAs

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### Text Legend
- ↑ Potential increased exposure of the comedication
- ↓ Potential decreased exposure of the comedication
- ↑↑ Potential increased exposure of COVID drug
- ↓↓ Potential decreased exposure of COVID drug
- ↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies. This interaction involves drugs identified by www.crediblemeds.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. Note, please check product labels for any additional cardiac warnings.

### Notes:
- * Increased risk of ALT elevations due to an expected or observed significant increase in DAA concentrations.

All DDAs + CLQ and HCLQ

The product labels for chloroquine and hydroxychloroquine recommend caution in impaired hepatic function.

Ledipasvir/sofosbuvir + LPV/r

Case reports of possible interaction between ledipasvir and LPV with patients having drug-induced liver injury manifesting as significant bilirubin rise within two weeks of starting ledipasvir/sofosbuvir while on LPV-containing HIV regimens.

Ombitasvir/Paritaprevir + ATV

Paritaprevir AUC increased by 187%. Exposures of paritaprevir greater than this have been evaluated in phase 2 studies and were not expected to have a clinically meaningful impact on safety.

Ombitasvir/Paritaprevir + Dasabuvir + ATV

Paritaprevir AUC increased by 94%. Exposures of paritaprevir greater than this have been evaluated in phase 2 studies and were not expected to have a clinically meaningful impact on safety. However, the combination carries an increased risk for hyperbilirubinaemia (including oculi icterus), particularly when ribavirin is also prescribed.

Ombitasvir/Paritaprevir + Dasabuvir + FAVI

Coadministration may increase dasabuvir concentrations. However, due to dasabuvir’s large therapeutic index, a clinically relevant effect is not anticipated.

Ombitasvir/Paritaprevir ± Dasabuvir + CLQ

Inhibition of CYP3A4 by ritonavir may decrease exposure to chloroquine active metabolites, but this may not affect overall activity. No a priori dose alteration for chloroquine is recommended.

Ombitasvir/Paritaprevir ± Dasabuvir + HCLQ

Inhibition of CYP3A4 by ritonavir may decrease exposure to hydroxychloroquine, although to a moderate extent due to the multiple elimination pathways. No dose alteration is required.

Sofosbuvir/Velpatasvir + FAVI

Coadministration may increase velpatasvir concentrations. However, due to velpatasvir’s large therapeutic index, a clinically relevant effect is not anticipated.

Sofosbuvir/Velpatasvir/Voxilaprevir + FAVI

Coadministration may increase velpatasvir concentrations. However, due to velpatasvir’s large therapeutic index, a clinically relevant effect is not anticipated.
Antivirals – Others

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Text Legend

- ✦ Potential increased exposure of the comedication
- ▼ Potential decreased exposure of the comedication
- 💚 Potential increased exposure of COVID drug
- 🟡 Potential decreased exposure of COVID drug
- ↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies. ▼ This interaction involves drugs identified by www.crediblemeds.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. Note, please check product labels for any additional cardiac warnings.

Key to abbreviations

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<td>IFN-β</td>
<td>Interferon beta</td>
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Colour Legend

- These drugs should not be coadministered
- Potential interaction which may require a dose adjustment or close monitoring
- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required
- No clinically significant interaction expected
Anxiolytics/Hypnotics/Sedatives

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Text Legend

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
↑↑ Potential increased exposure of COVID drug
↓↓ Potential decreased exposure of COVID drug
↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.
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Note, please check product labels for any additional cardiac warnings.

Key to abbreviations

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Colour Legend

- These drugs should not be coadministered
- Potential interaction which may require a dose adjustment or close monitoring.
- These drugs are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers.
- No clinically significant interaction expected

Please note that if a drug is not listed it cannot automatically be assumed it is safe to coadminister. No recommendation to use experimental therapy for COVID-19 is made. Drug interaction data for many agents are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers.

Charts updated 5 May 2020

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Please check www.covid19-druginteractions.org for updates.
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**Beta Blockers**

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</tbody>
</table>

**Text Legend**

- **↑**: Potential increased exposure of the comedication
- **↓**: Potential decreased exposure of the comedication
- **↑↑**: Potential increased exposure of COVID drug
- **↓↓**: Potential decreased exposure of COVID drug
- **↔**: No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

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Note, please check product labels for any additional cardiac warnings.

**Notes:**

**Beta blockers + CLQ or HCLQ**

Caution and monitoring is advised as beta blockers can prolong the PR interval and chloroquine and hydroxychloroquine have been shown to prolong the QT interval.

---

**Key to abbreviations**

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<th>Abbreviation</th>
<th>Drug</th>
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<td>TCZ</td>
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**Colour Legend**

- **These drugs should not be coadministered**
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- **Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.**
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Bronchodilators

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<td>↔</td>
<td>↓</td>
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</tr>
</tbody>
</table>

**Text Legend**

- Potential increased exposure of the comedication
- Potential decreased exposure of the comedication
- Potential increased exposure of COVID drug
- Potential decreased exposure of COVID drug
  - No significant effect
  - Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.
  - This interaction involves drugs identified by [www.crediblemeds.org](http://www.crediblemeds.org) as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. Note, please check product labels for any additional cardiac warnings.

**Notes:**

**Aminophylline + TCZ**

Aminophylline is a complex of theophylline and ethylenediamine and is given for its theophylline activity. Coadministration may decrease theophylline concentrations.

**Aminophylline or theophylline + IFN-β**

Coadministration may increase theophylline concentrations but this is unlikely to be clinically significant. (Aminophylline is a complex of theophylline and ethylenediamine and is given for its theophylline activity.)

**Indacaterol + ATV or LPV/r**

Exposure can be increased by up to 2-fold with ritonavir (and may be similar with atazanavir), however, this increase does not raise any concerns based on indacaterol’s safety data.
## Calcium Channel Blockers

<table>
<thead>
<tr>
<th></th>
<th>ATV</th>
<th>CLQ</th>
<th>FAVI</th>
<th>HCLQ</th>
<th>IFN-β</th>
<th>LPV/r</th>
<th>RDV</th>
<th>RBV</th>
<th>TCZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amlodipine</td>
<td>↑</td>
<td>←</td>
<td>←</td>
<td>←</td>
<td>←</td>
<td>↑</td>
<td>←</td>
<td>←</td>
<td>←</td>
</tr>
<tr>
<td>Diltiazem</td>
<td>↑125%</td>
<td>←</td>
<td>←</td>
<td>←</td>
<td>←</td>
<td>↑</td>
<td>←</td>
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<td>←</td>
<td>←</td>
<td>←</td>
<td>↑</td>
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<tr>
<td>Nifedipine</td>
<td>↑</td>
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<td>↑</td>
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</tr>
</tbody>
</table>

**Text Legend**

- ↑ Potential increased exposure of the comedication
- ↓ Potential decreased exposure of the comedication
- ↑ Potential increased exposure of COVID drug
- ↓ Potential decreased exposure of COVID drug
- ↔ No significant effect

**Notes:**

- **Calcium channel blockers + CLQ or HCLQ**
  Caution and monitoring is advised as calcium channel blockers can prolong the PR interval and chloroquine and hydroxychloroquine have been shown to prolong the QT interval.

- **Amlodipine + LPV/r**
  If coadministration is indicated, consider a dose reduction for amlodipine of 50%.

- **Diltiazem + ATV**
  If coadministration is indicated, an initial dose reduction of diltiazem by 50% is recommended, with subsequent titration as needed and ECG monitoring.

**Key to abbreviations**

- ATV: Atazanavir
- CLQ: Chloroquine
- FAVI: Favipiravir
- HCLQ: Hydroxychloroquine
- IFN-β: Interferon beta
- LPV/r: Lopinavir/ritonavir
- RDV: Remdesivir
- RBV: Ribavirin
- TCZ: Tocilizumab

**Colour Legend**

- These drugs should not be coadministered
- Potential interaction which may require a dose adjustment or close monitoring.
- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
- No clinically significant interaction expected
Any increase in exposure of levonorgestrel or ulipristal is unlikely to be clinically significant when used as a single dose.

### Contraceptives

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<thead>
<tr>
<th>Drug</th>
<th>ATV</th>
<th>CLQ</th>
<th>FAVI</th>
<th>HCLQ</th>
<th>IFN-β</th>
<th>LPV/r</th>
<th>RDV</th>
<th>RBV</th>
<th>TCZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Desogestrel (COC)</td>
<td>↑</td>
<td>←</td>
<td>↑</td>
<td>←</td>
<td>←</td>
<td>↑</td>
<td>←</td>
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<tr>
<td>Desogestrel (POP)</td>
<td>↑</td>
<td>←</td>
<td>↑</td>
<td>←</td>
<td>←</td>
<td>↑</td>
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<tr>
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<td>←</td>
<td>↑</td>
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<td>←</td>
<td>↓42%</td>
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<td>←</td>
<td>←</td>
<td>↑52%</td>
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<td>Etinogestrel (vaginal ring)</td>
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<td>↑</td>
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<td>←</td>
<td>↑</td>
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<tr>
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<tr>
<td>Levonorgestrel (emergency con.)</td>
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<tr>
<td>Levonorgestrel (implant)</td>
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<tr>
<td>Levonorgestrel (IUD)</td>
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<tr>
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<td>←</td>
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<td>←</td>
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<td>Norelgestromin (patch)</td>
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<td>↑</td>
<td>←</td>
<td>←</td>
<td>↑83%</td>
<td>←</td>
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<td>Norethisterone (COC)</td>
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<td>←</td>
<td>↑47%</td>
<td>←</td>
<td>←</td>
<td>↓17%</td>
<td>←</td>
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<td>Norethisterone (IM depot)</td>
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<tr>
<td>Norethisterone (POP)</td>
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<tr>
<td>Norgestrel (COC)</td>
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### Text Legend

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Effect</th>
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<tbody>
<tr>
<td>↑</td>
<td>Potential increased exposure of the comedication</td>
</tr>
<tr>
<td>↓</td>
<td>Potential decreased exposure of the comedication</td>
</tr>
<tr>
<td>↑↑</td>
<td>Potential increased exposure of COVID drug</td>
</tr>
<tr>
<td>↓↓</td>
<td>Potential decreased exposure of COVID drug</td>
</tr>
<tr>
<td>↔</td>
<td>No significant effect</td>
</tr>
</tbody>
</table>

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

This interaction involves drugs identified by www.crediblemeds.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined.

Note, please check product labels for any additional cardiac warnings.

### Notes:

**COC** – Combined oral contraceptive; **POP** – Progestogen only pill; **IUD** – Intra-uterine device

**Contraceptives + RBV**

Extreme care must be taken to avoid pregnancy in female patients and in female partners of male patients taking ribavirin. The European product labels for ribavirin state that effective contraception must be used during ribavirin treatment and for 4 months after treatment has been concluded in female patients and for 7 months in female partners of male patients. The US product labels for ribavirin state that effective contraception must be used during ribavirin treatment and for 6 months after treatment has been concluded in female patients and female partners of male patients.

**Ethinylestradiol and/or progestins + ATV, LPV/r, FAVI**

Concentrations of ethinylestradiol and progestins may be affected but no action is needed due to the short treatment duration of the COVID-19 therapy.

**Levonorgestrel (emergency contraception) and Ulipristal + ATV or LPV/r**

Any increase in exposure of levonorgestrel or ulipristal is unlikely to be clinically significant when used as a single dose.
Hormone Replacement Therapy

<table>
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<tr>
<th>Drug Interaction</th>
<th>ATV</th>
<th>CLQ</th>
<th>FAVI</th>
<th>HCLQ</th>
<th>IFN-β</th>
<th>LPV/r</th>
<th>RDV</th>
<th>RBV</th>
<th>TCZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drospirenone (HRT)</td>
<td>↑</td>
<td>↔</td>
<td>↑</td>
<td>↔</td>
<td>↔</td>
<td>↑</td>
<td>↔</td>
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<tr>
<td>Dydrogesterone (HRT)</td>
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<tr>
<td>Estradiol</td>
<td>↑</td>
<td>↔</td>
<td>↑</td>
<td>↔</td>
<td>❄</td>
<td>↓</td>
<td>↔</td>
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<tr>
<td>Levonorgestrel (HRT)</td>
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<td>Medroxyprogesterone (oral)</td>
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</tbody>
</table>

**Text Legend**

- **↑** Potential increased exposure of the comedication
- **↓** Potential decreased exposure of the comedication
- **❄** Potential increased exposure of COVID drug
- **↔** No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

This interaction involves drugs identified by www.crediblemeds.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. Note, please check product labels for any additional cardiac warnings.

**Notes:**

**Estradiol and + ATV, FAVI or LPV/r**

Concentrations of estradiol may alter but no action is needed due to the short treatment duration of the COVID-19 therapy.

**Progestins + ATV, FAVI or LPV/r**

Concentrations of progestins may increase but no action is needed due to the short treatment duration of the COVID-19 therapy.
Gastrointestinal Agents

<table>
<thead>
<tr>
<th></th>
<th>ATV</th>
<th>CLQ</th>
<th>FAVI</th>
<th>HCLQ</th>
<th>IFN-β</th>
<th>LPV/r</th>
<th>RDV</th>
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Text Legend

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
† Potential increased exposure of COVID drug
¬ Potential decreased exposure of COVID drug

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.
♥ This interaction involves drugs identified by www.crediblemeds.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined.
Note, please check product labels for any additional cardiac warnings.

Notes:

Antacids + ATV
Antacids can reduce absorption of atazanavir. Atazanavir should be taken at least 2 h before or 1 h after antacids.

Antacids + CLQ or HCLQ
Antacids can reduce absorption of chloroquine and hydroxychloroquine. Antacids should be taken at least 4 h before or 4 h after chloroquine or hydroxychloroquine

Cimetidine, famotidine, ranitidine + ATV
Unboosted atazanavir is not recommended with H2RAs as they can reduce absorption of atazanavir. If coadministration is necessary, atazanavir 400 mg once daily with food should be administered at least 2 hours before and at least 10 hours after a dose of the H2RA.

Esomeprazole, lansoprazole, omeprazole, pantoprazole, rabeprazole + ATV
When possible, discontinue proton pump inhibitor treatment for the duration of atazanavir treatment.

Loperamide + ATV or LPV/r
Caution is advised with high doses of loperamide used for reducing stoma output, particularly as patients may be at increased risk of cardiac events due to electrolytes disturbances.
Please check [www.covid19-druginteractions.org](http://www.covid19-druginteractions.org) for updates.

Please note that if a drug is not listed it cannot automatically be assumed it is safe to coadminister. No recommendation to use experimental therapy for COVID-19 is made.

Drug interaction data for many agents are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers.

Liverpool Drug Interactions Group

Interactions with Experimental COVID-19 Therapies

Gastrointestinal Agents – Anti-emetics

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↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
☞ Potential increased exposure of COVID drug
✂ Potential decreased exposure of COVID drug
↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

This interaction involves drugs identified by [www.crediblemeds.org](http://www.crediblemeds.org) as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined.

Note, please check product labels for any additional cardiac warnings.

Key to abbreviations

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Colour Legend

These drugs should not be coadministered
Potential interaction which may require a dose adjustment or close monitoring.
Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
No clinically significant interaction expected

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Hypertensives – ACE inhibitors

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Text Legend

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
⇑ Potential increased exposure of COVID drug
⇓ Potential decreased exposure of COVID drug
→ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

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Colour Legend

These drugs should not be coadministered
Potential interaction which may require a dose adjustment or close monitoring.
Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
No clinically significant interaction expected

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Please note that if a drug is not listed it cannot automatically be assumed it is safe to coadminister. No recommendation to use experimental therapy for COVID-19 is made. Drug interaction data for many agents are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers.
### Hypertensives – Other agents

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<tr>
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<td>↔</td>
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<td>↔</td>
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**Text Legend**

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
↑↑ Potential increased exposure of COVID drug
↓↓ Potential decreased exposure of COVID drug
↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

• This interaction involves drugs identified by www.credibledrugs.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined.

Note, this interaction involves drugs identified by www.credibledrugs.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined.

**Notes:**

**Doxazosin + ATV or LPV/r**

For patients already taking doxazosin, monitor blood pressure and reduce doxazosin dose as needed if hypotension occurs on starting ATV or LPV/r.

**Isosorbide nitrate + ATV or LPV/r**

Decreased active metabolite.

**Sacubitril + ATV or LPV/r**

Increased active metabolite.

**Terazosin + ATV or LPV/r**

For patients already taking terazosin, monitor blood pressure and reduce terazosin dose as needed if hypotension occurs on starting ATV or LPV/r.

---

**Key to abbreviations**

<table>
<thead>
<tr>
<th>ATV</th>
<th>Alazanavir</th>
<th>LPV/r</th>
</tr>
</thead>
<tbody>
<tr>
<td>CLQ</td>
<td>Chloroquine</td>
<td>RDV</td>
</tr>
<tr>
<td>FAVI</td>
<td>Favipiravir</td>
<td>RBV</td>
</tr>
<tr>
<td>IFN-β</td>
<td>Interferon beta</td>
<td></td>
</tr>
</tbody>
</table>

**Colour Legend**

- These drugs should not be coadministered
- Potential interaction which may require a dose adjustment or close monitoring.
- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
- No clinically significant interaction expected
The European product label for LPV/r does not recommend tadalafil for the treatment of pulmonary arterial hypertension, but the US product label suggests for patients on tadalafil, to avoid the use of tadalafil during the initiation of LPV/r and to stop tadalafil at least 24 hours based upon individual tolerability.

Ambrisentan + ATV or LPV/r
Start ambrisentan at 5 mg and closely monitor the patient for tolerability.

Bosentan + LPV/r
When coadministered patients should be closely observed for bosentan toxicity, especially during the first week of co-administration. For patients on bosentan, the US product label for LPV/r suggests to discontinue bosentan at least 36 hours prior to initiation of LPV/r and after at least 10 days of LPV/r, to resume bosentan at 62.5 mg once daily or every other day based upon individual tolerability.

Riociguat + ATV or LPV/r
The European product label for riociguat does not recommend its use in presence of strong inhibitors of CYPs, P-gp and BCRP; the US product label recommends to start riociguat at a dose of 0.5 mg three times daily and to monitor for signs and symptoms of hypotension.

Tadalafil + ATV
The US product label for ATV suggests for patients receiving atazanavir for at least one week, to start tadalafil at 20 mg once daily and increase to 40 mg once daily based on individual tolerability. For patients on tadalafil, avoid the use of tadalafil when starting atazanavir. Stop tadalafil at least 24 hours before starting atazanavir. At least one week after starting atazanavir, resume tadalafil at 20 mg once daily and increase to 40 mg once daily based on individual tolerability.

Tadalafil + LPV/r
The European product label for LPV/r does not recommend tadalafil for the treatment of pulmonary arterial hypertension, but the US product label suggests for patients on tadalafil, to avoid use of tadalafil during the initiation of LPV/r and to stop tadalafil at least 24 hours prior to starting LPV/r. After at least one week following the initiation of LPV/r, resume tadalafil at 20 mg once daily. Increase to 40 mg once daily based upon individual tolerability.
Interactions with Experimental COVID-19 Therapies

Please check www.covid19-druginteractions.org for updates.

Please note that if a drug is not listed it cannot automatically be assumed it is safe to coadminister. No recommendation to use experimental therapy for COVID-19 is made. Drug interaction data for many agents are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers.

Immunosuppressants

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<th>FAVI</th>
<th>HCLQ</th>
<th>IFN-β</th>
<th>LPV/r</th>
<th>RDV</th>
<th>RBV</th>
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<td>↔</td>
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<tr>
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<tr>
<td>Azathioprine</td>
<td>𝕀殄</td>
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<td>Belatacept</td>
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<tr>
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</tbody>
</table>

Text Legend

↑ Potential increased exposure of the comedication
↓ Potential decreased exposure of the comedication
↑ Potential increased exposure of COVID drug
↓ Potential decreased exposure of COVID drug
↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies. This interaction involves drugs identified by www.crediblemeds.org as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. Note, please check product labels for any additional cardiac warnings.

**Notes:**

**Adalimumab and azathioprine + CLQ or HCLQ**
The risk of haematological toxicity may be potentially increased as adalimumab, azathioprine, chloroquine and hydroxychloroquine can cause myelosuppression. Closely monitor haematological parameters.

Adalimumab + RBV
The risk of haematological toxicity may be potentially increased as adalimumab and ribavirin can cause myelosuppression. Closely monitor haematological parameters.

Adalimumab and basiliximab + TCZ
Avoid coadministration due to the enhanced immunosuppressive effect.

Adalimumab + IFN-β
Caution is required due to potential additive haematological toxicity.

Azathioprine + RBV
Ribavirin may interfere with azathioprine metabolism possibly leading to an accumulation of 6-methylthioinosine monophosphate, which has been associated with myelotoxicity.

Azathioprine + IFN-β or TCZ
Caution is required due to potential additive haematological toxicity.

Pirfenidone + IFN-β
Any increase is pirfenidone is unlikely to be clinically relevant, except in the presence of hepatic impairment as moderate hepatic impairment also increases pirfenidone exposure (by 60%). No a priori dosage adjustment is recommended in patients with hepatic impairment but monitor for increased toxicity.

**Key to abbreviations**

<table>
<thead>
<tr>
<th>ATV</th>
<th>CLQ</th>
<th>FAVI</th>
<th>HCLQ</th>
<th>IFN-β</th>
<th>LPV/r</th>
<th>RDV</th>
<th>RBV</th>
<th>TCZ</th>
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<tbody>
<tr>
<td><em>Alazanavir</em></td>
<td><em>Chloroquine</em></td>
<td><em>Favipiravir</em></td>
<td><em>Hydroxychloroquine</em></td>
<td><em>Interferon beta</em></td>
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</tbody>
</table>

**Colour Legend**

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- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
- No clinically significant interaction expected
### Interactions with Experimental COVID-19 Therapies

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Drug interaction data for many agents are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers.

#### Inotropes & Vasopressors

<table>
<thead>
<tr>
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<th>FAVI</th>
<th>HCLQ</th>
<th>IFN-β</th>
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<tr>
<td>Dobutamine</td>
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</tr>
</tbody>
</table>

**Text Legend**

- **↑**: Potential increased exposure of the comedication
- **↓**: Potential decreased exposure of the comedication
- **†**: Potential increased exposure of COVID drug
- **‡**: Potential decreased exposure of COVID drug

- **↔**: No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

<table>
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<td><strong>♥</strong>: This interaction involves drugs identified by <a href="http://www.crediblemeds.org">www.crediblemeds.org</a> as having a known, possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. Note, please check product labels for any additional cardiac warnings.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**

**Remdesivir**

Pressor requirement to maintain blood pressure is a key exclusion criteria to eligibility for remdesivir use.

See [https://rdvcu.gilead.com/](https://rdvcu.gilead.com/) for further details.
### Lipid Lowering Agents

<table>
<thead>
<tr>
<th></th>
<th>ATV</th>
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<th>HCLQ</th>
<th>IFN-β</th>
<th>LPV/r</th>
<th>RDV</th>
<th>RBV</th>
<th>TCZ</th>
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<td>↑490%</td>
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<tr>
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<tr>
<td>Evolocumab</td>
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<td>↓20%</td>
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<td>↔</td>
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<td>↑33%</td>
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</tr>
</tbody>
</table>

#### Text Legend

- ↑ Potential increased exposure of the comedication
- ↓ Potential decreased exposure of the comedication
- ‾ Power increased exposure of COVID drug
- ‾ Power decreased exposure of COVID drug
- ↔ No significant effect

Numbers refer to increase or decrease in AUC as observed in drug-drug interaction studies.

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Note, please check product labels for any additional cardiac warnings.

#### Notes:

**Atorvastatin + ATV**

Coadministration is not recommended. If the use of atorvastatin is considered necessary, use the lowest possible dose of atorvastatin with careful safety monitoring. The daily atorvastatin dose should not exceed 10 mg.

**Atorvastatin + LPV/r**

Do not exceed a daily dose of 20 mg with careful safety monitoring.

**Evolocumab + TCZ**

Avoid coadministration due to the enhanced immunosuppressive effect.

**Rosuvastatin + ATV or LPV/r**

Do not exceed rosuvastatin 10 mg/day.

---

**Key to abbreviations**

<table>
<thead>
<tr>
<th>ATV</th>
<th>Alazanavir</th>
<th>LPV/r</th>
<th>Lopinavir/ritonavir</th>
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</thead>
<tbody>
<tr>
<td>CLQ</td>
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<td>RDV</td>
<td>Remdesivir</td>
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<td>Remdesivir</td>
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<tr>
<td>IFN-β</td>
<td>Interferon beta</td>
<td>Lopinavir</td>
<td>Ritonavir</td>
</tr>
</tbody>
</table>

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**Colour Legend**

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- Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment unlikely to be required.
- No clinically significant interaction expected

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Please check [www.covid19-druginteractions.org](http://www.covid19-druginteractions.org) for updates.

Please note that if a drug is not listed it cannot automatically be assumed it is safe to coadminister. No recommendation to use experimental therapy for COVID-19 is made.

Drug interaction data for many agents are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers.
A 30% dose reduction of the corticosteroid might be considered during concomitant treatment. Based on DDI study with LPV/r, exposure of prednisolone (obtained also after conversion from prednisone) is increased modestly (+30%). The active metabolite (+30%) of dexamethasone is increased by 108% but no significant effect on adrenal function was seen. Note, there is interaction involves drugs identified by www.crediblemeds.org as having a known possible or conditional risk of QT prolongation and/or TdP. Risk may be related to dose or concentration (due to DDIs) and/or additive if two or more such drugs are combined. No clinically significant interaction expected. These drugs should not be coadministered. No clinically significant interaction expected.

<table>
<thead>
<tr>
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<th>ATV</th>
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</thead>
<tbody>
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<td>Beclometasone</td>
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<td>✯</td>
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**Text Legend**

- Potential increased exposure of the comedication
- Potential decreased exposure of the comedication
- Potential increased exposure of COVID drug
- Potential decreased exposure of COVID drug
- No significant effect

**Notes:**

- Risk of elevated corticosteroid levels, Cushing's syndrome and adrenal suppression.
- This risk is present for oral and injected administration, and also for topical, inhaled or eye drops corticosteroids

**Beclometasone + LPV/r**

Ritonavir (100 mg twice daily) increased the AUC of the active metabolite by 108% but no significant effect on adrenal function was seen. Caution is still warranted, use the lowest possible corticosteroid dose and monitor for corticosteroid side effects.

**Betamethasone or Dexamethasone + ATV, LPV/r or RDV**

Betamethasone and dexamethasone are moderate inducers of CYP3A4 and could decrease exposure and efficacy of ATV, LPV/r or RDV particularly when administered orally or intravenously at high doses or for a long duration.

**Ciclesonide + ATV or LPV/r**

No dose adjustment required but monitor closely, especially for Cushing's syndrome, when using a high dose or prolonged administration.

**Flunisolide + ATV or LPV/r**

Use the lowest possible flunisolide dose with monitoring for corticosteroid side effects.

**Prednisolone or Prednisone + LPV/r**

Based on DDI study with LPV/r, exposure of prednisolone (obtained also after conversion from prednisone) is increased modestly (+30%). A 30% dose reduction of the corticosteroid might be considered during concomitant treatment.