Nirmatrelvir/Ritonavir (Paxlovid) Drug Interactions:

This is not an exhaustive list. Consultation with a pharmacist who can obtain a complete medication, recreational, and natural health product history from the patient is recommended prior to prescribing nirmatrelvir/ritonavir.

Severity	Recommendation	Rationale
Contraindicated	Use alternative COVID agent.	Stopping the drug will not mitigate the interaction (e.g., prolonged half-life,
Contraindicated (use within past 14 days)	Do not use nirmatreivir/ritonavir.	narrow therapeutic index, prolonged enzyme-inducing effects which may decrease effectiveness of nirmatrelvir/ritonavir). Do not coadminister due to risk of serious toxicity.
Do not coadminister	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Significant \uparrow in drug concentrations expected. Do not coadminister due to risk of serious toxicity.
Caution	Therapy modification required (see Appendix).	Significant \uparrow/\downarrow in drug concentrations expected, which may lead to serious toxicity or impaired efficacy. Only coadminister if the interacting drug can be safely held or dose-adjusted and closely monitored (see Appendix). Expert consultation may be useful.
Drug interaction not likely to be clinically relevant	Continue with standard dosing.	Although mentioned in the monograph, clinically relevant interaction is not anticipated (e.g., minimal impact on certain metabolic pathways, wide therapeutic index, and short course of nirmatrelvir/ritonavir).
	Contraindicated Contraindicated (use within past 14 days) Do not coadminister Caution Drug interaction not likely to be clinically	Contraindicated (use within past 14 days) Do not coadminister Hold and restart 2 days after completing nirmatrelvir/ritonavir. Caution Therapy modification required (see Appendix). Continue with standard dosing.

- Abemaciclib (Verzenio)
- Alfuzosin (*Xatral*)
- Alprazolam (*Xanax*)
- ▲ Amiodarone
- ✓ Amitriptyline
- Amlodipine (*Norvasc*)
- ▲ Apalutamide (*Erleada*)
- Apixaban (*Eliquis*)
- Aripiprazole (Abilify), oral
- Atorvastatin (*Lipitor*)
- ✓ Atovaquone
- ▲ Bosentan (*Tracleer*)
- Bosutinib (*Bosulif*)
- Brexpiprazole (Rexulti)
- ✓ Budesonide
- Bupropion
- Buspirone (Buspar)
- ▲ Carbamazepine (Tegretol)
- Ceritinib (Zykadia)
- Cisapride
- Citalopram
- Clarithromycin
- Clomipramine
- Clonazepam
- Clopidogrel (*Plavix*)
- Clorazepate
- ▲ Clozapine (*Clozaril*)
- Cobimetinib (*Cotellic*)
- Colchicine in renal/hepatic impairment
- Cyclosporine (*Neoral*)
- Dabigatran
- ▲ Dabrafenib (*Tafinlar*)
- Dasatinib (Sprycel)
- Dexamethasone, high dose
- Diazepam (Valium)
- Digoxin
- Diltiazem (Tiazac, Cardizem)

†ED = erectile dysfunction ‡PAH = pulmonary arterial hypertension

- ✓ Divalproex
- Dofetilide
- ✓ Dronabinol
- ▲ Dronedarone (*Multag*)
- Edoxaban (*Lixiana*)
- Elagolix (Orilissa)
- Encorafenib (*Braftovi*)
- ▲ Enzalutamide
- Ergot alkaloids (e.g., dihydroergotamine, ergonovine)
- **▲** Eslicarbazepine
- Ethinyl estradiol
- Everolimus (Certican)
- Felodipine
- ▲ Fentanyl (*Duragesic*)
- ▲ Flecainide
- ✓ Fluoxetine
- Flurazepam
- ✓ Fluvoxamine
- Fostamatinib (Tavalisse)
- ✓ Fusidic acid, topical
- Glecaprevir/Pibrentasvir (Maviret)
- Hydrocodone
- Ibrutinib (*Imbruvica*)
- ✓ Imipramine
- ✓ Itraconazole
- Ketoconazole
- ✓ Lamotrigine
- Lomitapide (*Juxtapid*)
- **▲** Lorlatinib (*Lorbrena*)
- Lovastatin
- ▲ Lurasidone (*Latuda*)
- ✓ Maprotiline
- ✓ Maraviroc
- Meperidine (*Demerol*)
- Methamphetamine

- Metoprolol
- Midazolam, oral
- ▲ Mitotane (Lysodren)
- Modafinil
- Neratinib (Nerlynx)
- Nifedipine
- Nilotinib (Tasigna)
- Nitrazepam (Mogadon)
- ✓ Nortriptyline
- ▲ Oxcarbazepine
- Oxycodone (*Percocet*, OxyNEO)
- ✓ Paroxetine
- ♠ Phenobarbital
- ♠ Phenytoin (*Dilantin*)
- Pimozide
- Primidone
- ▲ Propafenone
- Quetiapine (Seroquel)
- ▲ Quinidine
- Quinine
- ✓ Raltegravir
- ▲ Ranolazine (*Corzyna*)
- Rifabutin
- A Rifampin
- ▲ Rifapentine
- Risperidone (Risperdal), oral
- ▲ Risperidone, long-acting injection (Risperdal Consta)
- Rivaroxaban (Xarelto)
- Rosuvastatin (Crestor)
- Salmeterol (Serevent, Advair)
- ✓ Sertraline
- ♦ Sildenafil for ED[†] (Viagra)
- ▲ Sildenafil for PAH[‡] (*Revatio*)

- Silodosin (Rapaflo)
- Simvastatin
- Sirolimus (Rapamune)
- ▲ Sonidegib (*Odomzo*)
- ▲ St. John's wort (Hypericum perforatum)
- Tacrolimus (Prograf, Advagraf, Envarsus)
- ◆ Tadalafil for ED[†] (*Cialis*)
- ▲ Tadalafil for PAH[‡] (*Adcirca*)
- Tamsulosin (*Flomax*)
- ▲ Tepotinib (*Tepmetko*)

✓ Theophylline

- Ticagrelor (Brilinta)
- ✓ Timolol
- Tramadol
- Triazolam (Halcion)
- ✓ Trimipramine
- Vardenafil (Levitra) for ED[†]
- ▲ Vardenafil (*Levitra*) for PAH[‡]
- ▲ Venetoclax (Venclexta)
- ✓ Venlafaxine
- Verapamil
- Vinblastine
- Vincristine
- ✓ Voriconazole
- Warfarin
- Ziprasidone (Zeldox)
- Zolpidem (Sublinox, Ambien)
- Zopiclone (*Imovane*)

Liverpool's COVID-19 Interaction Checker https://www.covid19-druginteractions.org/ University Health Network &

Kingston Health Sciences Centre Paxlovid-Oncology DDI https://www.antimicrobialstewardship.com/paxlovid-ddi-oncology

Appendix: Nirmatrelvir/ritonavir (Paxlovid) Drug Interactions

June 6, 2022. This document will be updated as more information becomes available.

Guiding principles for managing drug interactions categorized as • and •.

There is limited drug interaction data for nirmatrelvir/ritonavir (which is a potent CYP3A4/P-glycoprotein inhibitor). Most potential interactions listed below are based on known/anticipated effects with ritonavir alone or with other protease inhibitors. In some instances, pharmacokinetic interaction data for other potent CYP3A4 inhibitors (e.g., clarithromycin, ketoconazole) are included in this table to help predict the potential extent of an interaction effect with nirmatrelvir/ritonavir.

General recommendation: • •

Hold the interacting drug for one week (i.e., beginning on the first day of nirmatrelvir/ritonavir and resuming two days after completing nirmatrelvir/ritonavir).

> Ritonavir inhibition is not immediately reversible.

If holding a drug for one week is not a safe option:

- Use an alternative COVID-19 agent for orgument

Caution:

Some drugs may need to be held longer due to a greater sensitivity to ritonavir inhibition (e.g., calcineurin inhibitors).

In many instances, replacing a drug is not feasible, and may introduce more risk of harm or error (e.g., patient takes both the held and new drug, forgets to restart original drug, etc).

> Recommendations in this appendix are based on Canadian product monographs, the <u>Liverpool COVID-19 Drug Interactions Database</u> (University of Liverpool, 2022), <u>Lexi-Interact Online Database</u> (Hudson OH, Wolters Kluwer, 2022), and additional references as noted.

Disclaimer

This document is intended for use by experienced clinicians, including prescribers and pharmacists. The information is not intended to replace sound professional judgment in individual situations, and should be used in conjunction with other reliable sources of information. Clinicians should always consider the risk/benefit profile for their individual patient, discuss these risks with the patient or caregiver before initiating therapy, and closely monitor for treatment benefit and adverse effects.

Neither the Ontario COVID-19 Science Advisory Table, the University of Waterloo, nor the authors and their respective institutions are responsible for deletions or inaccuracies in information or for claims of injury resulting from any such deletions or inaccuracies. Mention of specific drugs, drug doses, or drug combinations within this document does not constitute endorsement by the Ontario COVID-19 Science Advisory Table, the University of Waterloo, or the authors and their respective institutions.

This document is intended to complement (but is separate from) the Ontario COVID-19 Science Advisory Table Drugs and Biologics Clinical Practice Guidelines.

Drug	Recommendation	Comments
Abemaciclib (Verzenio)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. Alternatively, for patients who have not previously had dose reduction for toxicity, consider a dose reduction to 50 mg once daily with close monitoring for toxicity.	Decisions to hold or dose-adjust should be made in conjunction with the patient's oncologist. Cyclin-dependent kinase inhibitors are generally held for acute infection. Abemaciclib AUC increased over 3-fold when coadministered with clarithromycin.
Alfuzosin (<i>Xatral</i>)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. Alternatively, may consider giving every other day in patients with heightened risk of urinary retention. Monitor for hypotension.	Alfuzosin AUC increased 3-fold when coadministered with ketoconazole 400 mg.
◆ Alprazolam (<i>Xanax</i>)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. Alternatively, reduce alprazolam dose by at least 50% and monitor for increased effects.	Alprazolam AUC increased 148% and half-life increased from 13 to 30 hours when coadministered with ritonavir 200 mg x 4 doses.
Amlodipine (Norvasc)	Reduce amlodipine dose by 50% or take dose every other day. Restart usual dose 2 days after completing nirmatrelvir/ritonavir. Monitor blood pressure. May consider continuing with usual dosing in patients at low risk of bradycardia or hypotension.	Amlodipine AUC increased 2-fold when coadministered with indinavir/ritonavir or paritaprevir/ritonavir.

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Recommendation **Comments** Drug Apixaban (Eliquis) If possible, use alternative COVID-19 agent. If not possible, Canadian monograph states that ensure stable renal function, then: coadministration with ritonavir is contraindicated. However, US product A) If already on low dose (2.5 mg BID) apixaban, continue. monograph suggests to decrease 5 mg twice daily dose to 2.5 mg twice daily when combined B) If acute venous thromboembolism (VTE): with strong inhibitors of CYP3A4 and **!** Low risk of clot: P-glycoprotein. Hold apixaban. 12 hours after the last dose of apixaban, Eliquis (U.S.) Prescribing Information. Accessed February 8, start nirmatrelvir/ritonavir AND aspirin 81 mg daily. Finish 2022. https://www.accessdata.fda.gov/drugsatfda_docs/label/2012/2 aspirin 1 day after completing nirmaltrevir/ritonavir. 02155s000lbl.pdf Restart apixaban 2 days after completing nirmatrelvir/ritonavir. Observational data from Italy found a 70 to 490% increase in apixaban levels in combination High risk of clot: with antivirals containing ritonavir in hospitalized Hold apixaban. 12 hours after the last dose of apixaban, patients. start nirmatrelvir/ritonavir AND therapeutic dosing of a Testa S, Prandoni P, Paoletti O et al. Direct oral anticoagulant subcutaneous low molecular weight heparin (LMWH) plasma levels' striking increase in severe COVID-19 respiratory such as: syndrome patients treated with antiviral agents: The Cremona experience. J Thromb Haemost. 2020;18:1320–1323. Dalteparin 200 units/kg daily <u>or</u> 100 units/kg https://doi.org/10.1111/jth.14871 every 12 hours if >90 kg; Enoxaparin 1 mg/kg every 12 hours (preferred) OR 1.5 mg/kg once every 24 hours; **High risk of clot includes:** Tinzaparin 175 anti-Xa units/kg once daily. Clot within past 6 months Finish LMWH 1 day after completing nirmatrelvir/ritonavir. Clot at any time in past when Restart apixaban 2 days after completing anticoagulation interrupted nirmatrelvir/ritonavir. Active cancer with clot at any point in cancer journey **C)** If atrial fibrillation: Diagnosis of antiphospholipid Decrease apixaban to 2.5 mg BID. Resume usual dose 2 days antibody syndrome after completing nirmatrelvir/ritonavir. See Paxlovid for a Patient on a DOAC for more details. https://uwaterloo.ca/pharmacy/sites/ca.pharmacy/files/ uploads/files/paxlovid_for_a_patient_on_a_doac.pdf Aripiprazole AUC increased almost 2-fold when Aripiprazole Reduce aripiprazole oral dose by 50% and resume usual dose 2 days after completing nirmatrelvir/ritonavir. (Abilify), oral coadministered with ketoconazole. No clinically relevant interaction expected with Monitor for confusion, restlessness, and sedation. long-acting injection. Hold and restart 2 days after completing nirmatrelvir/ritonavir. Atorvastatin AUC increased almost 6-fold when Atorvastatin coadministered with lopinavir/ritonavir 400/100 Alternatively, reduce atorvastatin to 10 mg daily. Resume mg twice daily. usual dose 2 days after completing nirmatrelvir/ritonavir. Decisions to hold or dose-adjust should be made Bosutinib (*Bosulif*) Hold bosutinib and start nirmatrelvir/ritonavir 24 hours after the last bosutinib dose. Restart bosutinib 2 days after in conjunction with the patient's oncologist. completing nirmatrelvir/ritonavir. Bosutinib AUC increased almost 9-fold when coadministered with ketoconazole. Reduce brexpiprazole dose by 50% and resume usual dose 2 Brexpiprazole AUC increased 97% when Brexpiprazole days after completing nirmatrelvir/ritonavir. coadministered with ketoconazole. (Rexulti) Monitor for confusion, restlessness, sedation. Buspirone (*Buspar*) Hold and restart 2 days after completing nirmatrelvir/ritonavir. Buspirone AUC increased 19-fold when coadministered with itraconazole 200 mg/day Alternatively, reduce buspirone dose to 2.5 mg daily if the for 4 days. usual dose is 20 to 30 mg/day.

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Drug	Recommendation	Comments
Ceritinib (Zykadia)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. Alternatively, consider reducing ceritinib dose by 33% and monitor for toxicity.	Canadian monograph recommends to avoid concomitant use. However, US monograph suggests reducing dose by 33%, rounded to nearest 150 mg dosage strength. Zykadia (U.S.) Prescribing Information. Accessed February 8, 2022. https://www.accessdata.fda.gov/drugsatfda_docs/label/2019/205755s016lbl.pdf
		Decision to hold or dose-adjust ceritinib should be made in conjunction with the patient's oncologist.
		Ceritinib AUC increased 3-fold when single dose coadministered with ketoconazole.
Cisapride	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Potential for serious and/or life-threatening adverse effects, including cardiac arrhythmias.
Clonazepam	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Due to prolonged benzodiazepine half-life, coadministration is not recommended.
	If an anxiolytic is needed, use lorazepam, oxazepam, or temazepam at usual doses.	coadministration is not recommended.
Clopidogrel (<i>Plavix</i>)	Acute coronary syndrome (ACS)/percutaneous coronary intervention (PCI):	Coadministration will decrease the antiplatelet effect of clopidogrel.
	 If <1 month since ACS: Use alternative COVID-19 agent. 	Clopidogrel active metabolite AUC decreased by
	 If <3 months since ACS or <1 month since PCI (no ACS): <p>Consider switching clopidogrel to prasugrel (if age <75, weight >60 kg, and no history of stroke/TIA) and resume clopidogrel 2 days after completing nirmatrelvir/ritonavir; </p> 	51 to 69% when coadministered with ritonavir.
	 If >3 months since ACS or >1 month since PCI (no ACS): Continue clopidogrel with acetylsalicylic acid (ASA) during nirmatrelvir/ritonavir therapy. If not taking ASA, consider switching to prasugrel (if age <75, weight >60 kg, and no history of stroke/TIA) and resume clopidogrel 2 days after completing nirmatrelvir/ritonavir. 	
Clorazepate	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Due to prolonged benzodiazepine half-life, coadministration is not recommended.
	If an anxiolytic is needed, use lorazepam, oxazepam, or temazepam at usual doses.	Coadministration is not recommended.
Cobimetinib(Cotellic)	Hold cobimetinib and start nirmatrelvir/ritonavir 24 hours after the last cobimetinib dose. Restart cobimetinib 2 days after completing nirmatrelvir/ritonavir.	Decisions to hold or dose-adjust should be made in conjunction with the patient's oncologist.
		Cobimetinib AUC increased almost 7-fold when coadministered with ketoconazole.
Colchicine in renal/	Coadministration is contraindicated in patients with renal and/or hepatic impairment.	Drug interaction could lead to potentially life-threatening/fatal adverse events.
hepatic Impairment	In patients with <u>normal renal/hepatic function</u> , colchicine may be administered at a lowered dose if practical: • Treatment of gout flares: 0.6 mg x 1 dose, then 0.3 mg (½ tablet) 1 hour later. Repeat dose no earlier than 3 days.	
	 Prevention of gout flares: a) If on 0.6 mg twice daily: decrease to 0.3 mg once daily; b) If on 0.3 mg twice daily: decrease to 0.3 mg once every 2 days. 	
	 Treatment of Familial Mediterranean fever: maximum 0.6 mg (or 0.3 mg twice daily). 	
	In all cases, resume usual colchicine dose 2 days after completing nirmatrelvir/ritonavir.	

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Drug Recommendation Comments Cyclosporine Reduce cyclosporine total daily dose by 80% and start Check cyclosporine concentrations 2 days after the last dose of nirmatrelvir/ritonavir. (Neoral) nirmatrelvir/ritronavir 12 hours after the last cyclosporine dose. Continue at reduced dose throughout • If subtherapeutic: increase cyclosporine dose. nirmatrelvir/ritonavir therapy. Consider resumption of twice daily dosing. • If therapeutic: continue with current Resuming transplant immunotherapy after the last dose of cyclosporine dose. nirmatrelvir/ritonavir should be guided by therapeutic drug • If supratherapeutic: reduce or hold current monitoring and in conjunction with the patient's transplant cyclosporine dose. provider. In all cases, repeat cyclosporine level in 2 to 4 days and continue to dose-adjust accordingly. Dabigatran AUC increased almost 2-fold when Dabigatran If possible, use alternative COVID-19 agent. If not possible, then: coadministered with nirmatrelyir/ritonavir. A) If already on low dose (110 mg BID) dabigatran, continue. B) If acute venous thromboembolism (VTE): ! Low risk of clot: Hold dabigatran. 12 hours after the last dose of dabigatran, start nirmatrelvir/ritonavir AND aspirin 81 mg daily. Finish aspirin 1 day after completing nirmatrelvir/ritonavir. Restart dabigatran 2 days after completing nirmaltrevir/ritonavir. A High risk of clot: Hold dabigatran. 12 hours after the last dose of High risk of clot includes: dabigatran, start nirmatrelvir/ritonavir AND therapeutic Clot within past 6 months dosing of a subcutaneous low molecular weight heparin Clot at any time in past when (LMWH) such as: anticoagulation interrupted Dalteparin 200 units/kg daily OR 100 units/kg Active cancer with clot at any every 12 hours if >90 kg; point in cancer journey Enoxaparin 1 mg/kg every 12 hours (preferred) Diagnosis of antiphospholipid OR 1.5 mg/kg once every 24 hours; antibody syndrome • Tinzaparin 175 anti-Xa units/kg once daily. Finish LMWH 1 day after completing nirmatrelvir/ritonavir. Restart dabigatran 2 days after completing See Paxlovid for a Patient on a DOAC for nirmatrelvir/ritonavir. more details. https://uwaterloo.ca/pharmacy/sites/ca.ph C) If atrial fibrillation: armacy/files/uploads/files/paxlovid_for_a Decrease dabigatran to 110 mg BID (if eGFR>50 mL/minute) _patient_on_a_doac.pdf OR decrease to 75 mg BID (if eGFR 30-50 mL/minute). Resume usual dose 2 days after completing nirmatrelvir/ritonavir. Chronic phase chronic myelogenous leukemia (CML): Decisions to hold or dose-adjust dasatinib should Dasatinib Hold and restart 2 days after completing nirmatrelvir/ritonavir. (Sprycel) be made in conjunction with the patient's Alternatively, consider reducing dasatinib dose to 20 to 40 mg oncologist. and monitor for toxicity. Dasatinib AUC increased 5-fold when **Accelerated or blast phase CML:** coadministered with ketoconazole. Do not coadminister; use alternate COVID-19 therapy. Dexamethasone, **High dose** (≥**20 mg daily):** Reduce dexamethasone dose by Dexamethasone AUC increased almost 3-fold 50% and resume usual dose 2 days after completing when coadministered with voriconazole. high dose nirmatrelvir/ritonavir. Li M, Zhu L, Chen L et al. Assessment of drug-drug interactions between voriconazole and glucocorticoids. J Chemother. Low dose (<20 mg daily): Continue with usual dose during 2018;30(5):296-303. doi: 10.1080/1120009X.2018.1506693. nirmatrelvir/ritonavir. Potential for risk of dexamethasone toxicity with high doses (≥20 mg daily). Clinically significant interaction is not expected with dexamethasone at low doses, including when used for COVID-19 treatment.

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Drug	Recommendation	Comments
Diazepam (<i>Valium</i>)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. If an anxiolytic is needed, use lorazepam, oxazepam, or temazepam at usual doses.	Due to prolonged benzodiazepine half-life, coadministration is not recommended.
Digoxin	Reduce digoxin dose by 50% <u>OR</u> hold and restart 2 days after completing nirmatrelvir/ritonavir.	
Diltiazem (<i>Tiazac,</i> <i>Cardizem</i>)	Reduce diltiazem dose by 50% or take dose every other day. Restart usual dose 2 days after completing nirmatrelvir/ritonavir. Monitor heart rate and blood pressure. May consider continuing with usual dosing in patients at low risk of bradycardia or hypotension.	Concentrations of calcium channel blockers are expected to increase when coadministered with nirmatrelvir/ritonavir.
Dofetilide	If possible, use alternative COVID-19 agent. Alternatively, hold dofetilide and restart 2 days after completing nirmatrelvir/ritonavir.	Dofetilide is metabolized to a small extent through CYP3A4.
Edoxaban (<i>Lixiana</i>)	If possible, use alternative COVID-19 agent. If not possible, ensure stable renal function, then: A) If already on low dose (30 mg once daily) edoxaban, continue. B) If acute venous thromboembolism (VTE): Low risk of clot: Hold edoxaban. 24 hours after the last dose of edoxaban, start nirmatrelvir/ritonavir AND aspirin 81 mg daily. Finish aspirin 1 day after completing nirmaltrevir/ritonavir. Restart edoxaban 2 days after completing nirmatrelvir/ritonavir.	No drug interaction data available with protease inhibitors but up to a 2-fold increase in exposure is anticipated. Canadian product monograph recommends caution when using with ritonavir; 30 mg daily dose is recommended with P-glycoprotein inhibitors.
	 High risk of clot: Hold edoxaban. 24 hours after the last dose of edoxaban, start nirmatrelvir/ritonavir AND therapeutic dosing of a subcutaneous low molecular weight heparin (LMWH) such as: Dalteparin 200 units/kg daily OR 100 units/kg every 12 hours if >90 kg; Enoxaparin 1 mg/kg every 12 hours (preferred)	 High risk of clot includes: Clot within past 6 months Clot at any time in past when anticoagulation interrupted Active cancer with clot at any point in cancer journey Diagnosis of antiphospholipid antibody syndrome
	Finish LMWH 1 day after completing nirmatrelvir/ritonavir. Restart edoxaban 2 days after completing nirmatrelvir/ritonavir. C) If atrial fibrillation: Decrease edoxaban to 30 mg daily. Resume usual dose 2 days after completing nirmatrelvir/ritonavir.	See Paxlovid for a Patient on a DOAC for more details. https://uwaterloo.ca/pharmacy/sites/ca.pharmacy/files/uploads/files/paxlovid_for_a_patient_on_a_doac.pdf
Elagolix (<i>Orili</i> ssa)	Potential for increased elagolix concentrations and possibly decreased nirmatrelvir concentrations. Continue with usual elagolix dose during nirmatrelvir/ritonavir therapy and monitor for elagolix toxicity.	Potential for serious adverse effects, including suicidal ideation and elevation of hepatic transaminases. Elagolix AUC increased over 2-fold when coadministered with ketoconazole 400 mg daily.

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Drug	Recommendation	Comments
Encorafenib (<i>Braftovi</i>)	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Decisions to hold or dose-adjust encorafenib should be made in conjunction with the
_	Alternatively, consider reducing encorafenib dose as follows and monitoring for toxicity:	patient's oncologist.
	 If taking 450 mg per day: reduce to 150 mg daily. If taking 150 to 300 mg per day: reduce dose to 75 mg daily. 	Encorafenib AUC increased 3-fold when coadministered with posaconazole.
	Resume usual encorafenib dose 2 days after completing nirmatrelvir/ritonavir.	
Ergot alkaloids (e.g., dihydroergotamine, ergonovine)	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Potential for serious and/or life threatening adverse effects, including acute ergot toxicity
Everolimus (<i>Certican</i>)	Hold everolimus and start nirmatrelvir/ritonavir 12 hours after last everolimus dose.	Check everolimus concentrations 2 days afte last dose of nirmatrelvir/ritonavir.
	Resuming transplant immunotherapy after the last dose of nirmatrelvir/ritonavir should be guided by therapeutic drug monitoring and in conjunction with the patient's transplant provider.	 If therapeutic/sub-therapeutic: resume everolimus at 25 to 50% baseline dose. Repeat level every 2 to 4 days and adjust dose accordingly. If supratherapeutic: continue to hold everolimus; repeat level in 2 to 4 days to assess resumption.
Felodipine	Reduce felodipine dose by 50% or take dose every other day. Restart usual dose 2 days after completing nirmatrelvir/ritonavir.	Concentrations of calcium channel blocke are expected to increase when coadminist with nirmatrelvir/ritonavir.
	Monitor blood pressure. May consider continuing with usual dosing in patients at low risk of bradycardia or hypotension.	
Flurazepam	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Due to prolonged benzodiazepine half-life,
	If an anxiolytic is needed, use lorazepam, oxazepam, or temazepam at usual doses.	coadministration is not recommended.
Fostamatinib (<i>Tavali</i> sse)	Monitor for toxicity including diarrhea, hypertension, hepatotoxicity, and neutropenia. If significant toxicity occurs, consider interruption of fostamatinib with reintroduction 2 days after completing nirmatrelvir/ritonavir.	Fostamatinib active metabolite AUC increase 102% when coadministered with ketoconazole.
Glecaprevir/ Pibrentasvir (<i>Maviret</i>)	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Glecaprevir exposure is increased over 4-fold with ritonavir and is associated with increase risk of alanine aminotransferase (ALT) elevation.
		In patients who are planning to start Hepatitis C (HCV) treatment, glecaprevir/pibrentasvir treatment should be deferred.
Hydrocodone	Reduce dose by about 50% or switch to equivalent dose of hydromorphone: • Multiply hydrocodone dose by 0.25 to get equivalent	Hydrocodone is metabolized to active metabolites: hydromorphone and norhydrocodone.
	 hydromorphone dose. Consider further reducing hydromorphone dose by 25 to 50% to account for cross tolerance. 	Hydrocodone AUC increased by 90% when coadministered with ritonavir/ombitasvir/paritaprevir combination.
	Monitor for signs of opioid toxicity. Resume usual hydrocodone dose 2 days after completing nirmatrelvir/ritonavir.	

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Drug	Recommendation	Comments
Ibrutinib (<i>Imbruvica</i>)	Consider alternate COVID-19 therapy. Alternatively, consider holding ibrutinib and starting nirmatrelvir/ritonavir 12 hours after the last ibrutinib dose. Restart ibrutinib 2 days after completing nirmatrelvir/ritonavir.	Decisions to hold or dose-adjust ibrutinib should be made in conjunction with the patient's oncologist. It may be dangerous to interrupt therapy in patients with high volume chronic lymphocytic leukemia or mantle cell lymphoma due to disease flare and/or cytokine release.
		Ibrutinib AUC increased 26-fold when coadministered with ketoconazole.
Lomitapide (<i>Juxtapid</i>)	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Lomitapide AUC increased 27-fold when coadministered with ketoconazole.
Lovastatin	Stop lovastatin at least 12 hours before starting nirmatrelvir/ritonavir. Restart 5 days after completing nirmatrelvir/ritonavir.	Contraindicated due to potential for severe toxicity including rhabdomyolysis and elevated liver function tests.
Meperidine (<i>Demerol</i>)	 Do not coadminister. Switch meperidine to an equivalent dose of hydromorphone: Multiply meperidine dose by 0.02 to get equivalent hydromorphone dose. Consider further reducing hydromorphone dose by 25 to 50% to account for cross tolerance. Monitor for signs of opioid toxicity. Resume usual meperidine 	Normeperidine AUC increased 50% when coadministered with ritonavir. Higher levels of normeperidine can cause central nervous system excitation and seizures.
Midazolam, oral	dose 2 days after completing nirmatrelvir/ritonavir. Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Coadministration may result in large increases in or midazolam concentrations with the potential for serious events such as prolonged or increased sedation or respiratory depression.
Modafinil	No dose adjustment required. Monitor for anxiety and agitation.	Coadministration could potentially increase modafi exposure due to CYP3A4 inhibition. Modafinil is a moderate inducer of CYP3A4, but a clinically significant effect on nirmetrelvir/ritonavir exposure is unlikely.
Neratinib (<i>Nerlynx</i>)	Hold and start nirmatrelvir/ritonavir 24 hours after the last neratinib dose. Restart neratinib 2 days after completing nirmatrelvir/ritonavir.	Decisions to hold or dose-adjust should be made in conjunction with the patient's oncologist. Neratinib AUC increased almost 5-fold when coadministered with ketoconazole.
Nifedipine	Reduce nifedipine dose by 50% or take dose every other day. Restart usual dose 2 days after completing nirmatrelvir/ritonavir. Monitor blood pressure. May consider continuing with usual dosing in patients at low risk of bradycardia or hypotension.	Concentrations of calcium channel blockers are expected to increase when coadministered with nirmatrelvir/ritonavir.
Nilotinib (Tasigna)	Chronic phase chronic myelogenous leukemia (CML): Hold nilotinib if possible, restart 2 days after completing nirmatrelvir/ritonavir. Alternatively, consider dose reduction to 400 mg PO daily and monitor for toxicity. Accelerated or blast phase CML: Do not coadminister. Consider an alternate COVID-19 therapy.	Decisions to hold or dose-adjust nilotinib should be made in conjunction with the patient's oncologist. Canadian monograph recommends holding if using CYP3A4 inhibitors, or monitoring for QTc if treatme interruption is not possible. A 50% dose reduction i recommended based on expected effect on nilotini exposures. Deeken JF, Pantanowitz I, Dezube BJ. Targeted therapies to treat non-AIDS-defining cancers in patients with HIV on HAART therapy treatment considerations. <i>Curr Opin Oncol</i> 2009; 21(5): 445-54. doi: 10.1097/CC.0b013e32832f3e04 Nilotinib AUC increased 3-fold when coadministere with ketoconazole.

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Drug	Recommendation	Comments
Nitrazepam (Mogadon)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. If an anxiolytic is needed, use lorazepam, oxazepam, or temazepam at usual doses.	Due to prolonged benzodiazepine half-life, coadministration is not recommended.
Oxycodone (Percocet, OxyNEO)	 Reduce dose of oxycodone by 66% or switch to equivalent dose of hydromorphone: Multiply oxycodone dose by 0.3 to get equivalent hydromorphone dose. Consider further reducing hydromorphone dose by 25 to 50% to account for cross tolerance. Monitor for signs of opioid toxicity. Resume usual oxycodone dose 2 days after completing nirmatrelvir/ritonavir. 	Oxycodone half-life increased 2-fold and AUC increased between 3 and 4-fold when coadministered with other potent 3A4 inhibitors (i.e., voriconazole).
Quetiapine (Seroquel)	Reduce to one-sixth of original dose and resume usual dose 2 days after completing nirmatrelvir/ritonavir. Monitor for confusion, dizziness, and sedation.	Quetiapine AUC increased 5 to 8-fold when coadministered with ketoconazole.
Quinine	For treatment of leg cramps: Hold and restart 2 days after completing nirmatrelvir/ritonavir. For treatment of malaria: Use an alternative COVID-19 agent.	Quinine AUC increased 4-fold and conversion to active metabolite was markedly inhibited when coadministered with ritonavir 200 mg twice daily.
Rifabutin	Reduce rifabutin to 150 mg once daily; return to 300 mg once daily 2 days after completing nirmatrelvir/ritonavir.	Canadian monograph recommends 150 mg three times a week, but the dose been found to be too low and contributes to resistance. The Department of Health and Human Services Panel on Antiretroviral Guidelines for Adults and Adolescents recommends using rifabutin 150 mg daily when used with a ritonavir-boosted protease inhibitor. https://clinicalinfo.hiv.gov/en/guidelines/adult-and-adolescent-arv/drug-interactions-between-protease-inhibitors-and-other-drugs?view=full Significant increases in exposures of rifabutin (>3-fold) and metabolite (>40-fold) observed when coadministered with lopinavir/ritonavir 400/100 mg twice daily.
Risperidone (<i>Risperdal</i>),	Reduce risperidone dose by 25 to 50% and resume usual dose 2 days after completing nirmatrelvir/ritonavir.	Risperidone AUC increased up to 2-fold when coadministered with ketoconazole.
oral	Monitor for confusion, extrapyramidal symptoms, and sedation.	Avoid coadministration in patients stabilized on risperidone long-acting injection.
Rivaroxaban (Xarelto)	Next page	Next page

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Drug Recommendation **Comments** If possible, use alternative COVID-19 agent. If not possible, then: Rivaroxaban AUC and Cmax increased by 153% and Rivaroxaban 55%, respectively, when coadministered with ritonavir (Xarelto) A) If acute venous thromboembolism (VTE): 600 mg twice daily in healthy volunteers. **Low risk of clot:** Observational data from Italy found a 600 to 3000% Hold rivaroxaban. 24 hours after the last dose of increase in rivaroxaban levels in combination with rivaroxaban, start nirmatrelvir/ritonavir AND aspirin 81 mg antivirals containing ritonavir in hospitalized patients. daily. Finish aspirin 1 day after completing Testa S, Prandoni P, Paoletti O et al. Direct oral anticoagulant plasma nirmaltrevir/ritonavir. Restart rivaroxaban 2 days after levels' striking increase in severe COVID-19 respiratory syndrome completing nirmatrelvir/ritonavir. patients treated with antiviral agents: The Cremona experience. J Thromb Haemost. 2020;18:1320-1323. A High risk of clot: https://doi.org/10.1111/jth.14871 Hold rivaroxaban. 24 hours after the last dose of rivaroxaban, start nirmatrelvir/ritonavir AND therapeutic dosing of a subcutaneous low molecular weight heparin High risk of clot includes: (LMWH) such as: Clot within past 6 months Dalteparin 200 units/kg daily OR 100 units/kg Clot at any time in past when every 12 hours if >90 kg; anticoagulation interrupted Enoxaparin 1 mg/kg every 12 hours (preferred) Active cancer with clot at any OR 1.5 mg/kg once every 24 hours; point in cancer journey • Tinzaparin 175 anti-Xa units/kg once daily. Diagnosis of antiphospholipid antibody syndrome Finish LMWH 1 day after completing nirmatrelvir/ritonavir. Restart rivaroxaban 2 days after completing nirmatrelvir/ritonavir. See Paxlovid for a Patient on a DOAC for B) If atrial fibrillation: more details. Hold rivaroxaban. 24 hours after the last dose of rivaroxaban, https://uwaterloo.ca/pharmacy/sites/ca.ph start nirmatrelvir/ritonavir AND edoxaban 30 mg daily. Finish armacy/files/uploads/files/paxlovid_for_a edoxaban 1 day after completing nirmatrelvir/ritonavir. Restart _patient_on_a_doac.pdf rivaroxaban 2 days after completing nirmatrelvir/ritonavir. Rosuvastatin AUC increased 2-fold and Cmax Rosuvastatin Hold and restart 2 days after completing nirmatrelvir/ritonavir. increased almost 5-fold when coadministered with Alternatively, reduce to 10 mg daily. Resume usual dose 2 lopinavir/ritonavir 400/100 mg twice daily. days after completing nirmatrelvir/ritonavir. Hold and restart 2 days after completing nirmatrelvir/ritonavir. Potential for serious and/or life-threatening Salmeterol adverse effects, including cardiac arrhythmias (Serevent, Advair) (prolonged QTc). Hold and restart 2 days after completing nirmatrelvir/ritonavir. Sildenafil AUC increased 2 to 11-fold when Sildenafil for coadministered with protease inhibitors. erectile Alternatively, reduce dose to 25 mg once every 48 hours. dysfunction Resume usual dose 2 days after completing nirmatrelvir/ritonavir. (Viagra) Silodosin AUC increased over 3-fold when Silodosin Hold and restart 2 days after completing nirmatrelvir/ritonavir. (Rapaflo) coadministered with ketoconazole. Stop simvastatin at least 12 hours before starting nirmatrelvir/ Simvastatin Contraindicated due to potential for severe ritonavir. Restart 5 days after completing nirmatrelvir/ritonavir. toxicity including rhabdomyolysis and elevated liver function tests. Sirolimus Hold sirolimus and start nirmatrelvir/ritonavir 24 to 48 hours Check sirolimus concentration 2 days after last (Rapamune) after the last sirolimus dose. dose of nirmatrelvir/ritonavir. • If therapeutic/subtherapeutic: resume Resuming transplant immunotherapy after the last dose of sirolimus at 50% of baseline dose. Repeat nirmatrelvir/ritonavir should be done in conjunction with the level every 7 days and dose-adjust patient's transplant provider. Use therapeutic drug monitoring accordingly. to guide sirolimus dose re-adjustment after completion of • If supratherapeutic: continue to hold nirmatrelvir/ritonavir. sirolimus and repeat level in 5 to 7 days to assess resumption.

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Drug	Recommendation	Comments
 Tacrolimus (Prograf, Advagraf, Envarsus) 	Immediate release (<i>Prograf</i> , generics): hold tacrolimus and start nirmatrelvir/ritonavir 12 hours after the last tacrolimus dose. Extended (<i>Advagraf</i>) or prolonged (<i>Envarsus</i>) release: hold the long acting tacrolimus and start nirmatrelvir/ritonavir 24 hours after the last tacrolimus dose. Resuming transplant immunotherapy after the last dose of nirmatrelvir/ritonavir should be guided by therapeutic drug monitoring and in conjunction with the patient's transplant provider.	 For all forms of tacrolimus: check tacrolimus concentrations 2 days after the last dose of nirmatrelvir/ritonavir. If therapeutic/subtherapeutic: resume tacrolimus at 25 to 75% of baseline dose; repeat level every 2 to 4 days and adjust dose accordingly. If supratherapeutic: continue to hold tacrolimus; repeat level in 2 to 4 days to assess resumption.
Tadalafil for erectile dysfunction (Cialis)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. Alternatively, reduce the dose to 10 mg once every 72 hours. Resume usual dose 2 days after completing nirmatrelvir/ritonavir.	Tadalafil AUC increased 124% when coadministered with ritonavir 200 mg twice daily.
Tamsulosin (Flomax)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. Alternatively, may consider using 0.4 mg daily or giving every other day in patients with heightened risk of urinary retention. Monitor for hypotension.	Tamsulosin AUC increased almost 3-fold when coadministered with ketoconazole.
• Ticagrelor (Brilinta)	 Acute coronary syndrome (ACS)/percutaneous coronary intervention (PCI): If <1 month since ACS: Suggest alternative COVID-19 agent. If <3 months since ACS or <1 month since PCI (no ACS): Switch to prasugrel (if age <75, weight >60 kg, and no history of stroke/TIA) during nirmatrelvir/ritonavir therapy. If >3 months since ACS or >1 month since PCI (no ACS): Consider temporarily holding ticagrelor (i.e., no switching) during nirmatrelvir/ritonavir therapy and resuming after. If not taking acetylsalicylic acid (ASA), consider switching to prasugrel (if age <70, weight >60 kg, and no history of stroke/TIA) or half-dose of ticagrelor (45 mg twice daily). 	Ticagrelor AUC increased 36% when coadministered with a single dose of ritonavir 100 mg.
◆ Tramadol	Reduce tramadol dose by 50% and monitor for pain relief and opioid toxicity. Resume usual dose 2 days after completing nirmatrelvir/ritonavir.	Inhibition of CYP3A4 may increase tramadol concentrations. Inhibition of CYP2D6 can decrease conversion of tramadol to a more active metabolite, but this is not expected to be significant when coadministered with nirmatrelvir/ritonavir.
• Triazolam (<i>Halcion</i>)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. If an anxiolytic is needed, use lorazepam, oxazepam, or temazepam at usual doses.	Due to prolonged benzodiazepine half-life, coadministration is not recommended. Triazolam half-life increased from 4 to 50 hours when coadministered with ritonavir 200 mg x 4 doses.
 Vardenafil (<i>Levitra</i>) for erectile dysfunction 	Hold and restart 2 days after completing nirmatrelvir/ritonavir.	Vardenafil AUC increased 49-fold when coadministered with ritonavir 600 mg twice daily.
Verapamil	Reduce verapamil dose by 50% or take dose every other day. Restart usual dose 2 days after completing nirmatrelvir/ritonavir. Monitor blood pressure. May consider continuing with usual	Concentrations of calcium channel blockers are expected to increase when coadministered with nirmatrelvir/ritonavir.
	dosing in patients at low risk of bradycardia or hypotension.	

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Drug	Recommendation	Comments
Vinblastine	Vinblastine may be held in the context of acute infection. Restart vinblastine at least 2 days after completing nirmatrelvir/ritonavir.	Decisions to hold or dose-adjust should be made in conjunction with the patient's oncologist.
	Alternatively, vinblastine may be coadministered with close monitoring for hematologic and neurotoxicity. Some providers may wish to empirically reduce vinblastine dose, especially in patients who have previously experienced or are at high risk for toxicity.	Vinblastine AUC increased almost 2-fold when coadministered with ritonavir. Increased risk of autonomic and peripheral neurotoxicity and neutropenia have been reported with coadministration of ritonavir and vinblastine.
Vincristine	Vincristine may be held in the context of acute infection. Restart vincristine 2 days after completing nirmatrelvir/ritonavir.	Decisions to hold or dose-adjust should be made in conjunction with the patient's oncologist.
	Alternatively, vincristine may be coadministered with close monitoring for hematologic and neurotoxicity. Some providers may wish to empirically reduce vincristine dose, especially in patients who have previously experienced or are at high risk for toxicity.	Increased rates of hematologic toxicity and neuropathy (including autonomic neuropathy) have been reported with coadministration of ritonavir and vincristine.
Warfarin	Monitor for signs of increased bleeding and bruising. Check international normalized ratio (INR) if clinically indicated.	Potential for increased warfarin concentrations when coadministered with nirmatrelvir/ritonavir.
Ziprasidone (<i>Zeldox</i>)	No dose adjustment required. Monitor for dizziness, extrapyramidal symptoms, and sedation.	Only one-third of ziprasidone dose is metabolized by CYP450. Ziprasidone AUC increased 35 to 40% when coadministered with ketoconazole.
Zolpidem (Sublinox, Ambien)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. If coadministration required, reduce zolpidem dose by 50%.	Zolpidem AUC increased 70% when coadministered with ketoconazole.
Zopiclone (<i>Imovane</i>)	Hold and restart 2 days after completing nirmatrelvir/ritonavir. If coadministration required, reduce zopiclone dose by 50%.	Potential for increased zopiclone exposures when coadministered with nirmatrelvir/ritonavir.