Interactions with Essential Medicines & Nirmatrelvir/ritonavir (NMV/r)

Charts revised 15 September 2022

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

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Interaction tables - refer to page 2 for legend, abbreviations and notes

Please note that if a drug is not listed it cannot automatically be assumed it is safe to coadminister.

Drug interaction data for many agents are limited or absent; therefore, risk-benefit assessment for any individual patient rests with prescribers. Management of interactions with nirmatrelvir/ritonavir (Paxlovid) may be complex and full details should be obtained from the website where possible.

A	lassies		
Ana	lgesics Codoino		
	Codeine		
	Diclofenac		
	Fentanyl		
	Hydromorphone Ibuprofen		
	Mefenamic acid		
	Morphine		
<u> </u>	Oxycodone Paracetamol		
	Tramadol		
Ant	tiarrhythmics		
-	Amiodarone		
	Lidocaine		
Ant	ibacterials		
/	Amikacin		
	Amoxicillin		
	Ampicillin		
	Bedaquiline		
	Cefalexin		
	Cefazolin		
	Cefixime		
	Cefotaxime		
	Ceftriaxone		
	Chloramphenicol		
	Ciprofloxacin		
	Clarithromycin (a)		
	Clindamycin		
	Clofazimine		
	Cloxacillin		
	Cycloserine		
	Dapsone		
	Delamanid		
	Doxycycline		
	Erythromycin		
	Ethambutol		
	Ethionamide Gentamicin		
	Imipenem/cilastatin		
	Isoniazid		
	Kanamycin		
	Levofloxacin		
	Linezolid		
	Meropenem		
	Metronidazole		
_	Moxifloxacin		
	Nitrofurantoin		
	Ofloxacin		
	Para-aminosalicylic acid		
	Penicillins		
	Piperacillin		
	Pyrazinamide		
	Pyrazinamide Rifabutin (b)		
□ ×	Pyrazinamide Rifabutin (b) Rifampicin		
	Pyrazinamide Rifabutin (b) Rifampicin Rifapentine		
×	Pyrazinamide Rifabutin (b) Rifampicin Rifapentine Spectinomycin		
×	Pyrazinamide Rifabutin (b) Rifampicin Rifapentine Spectinomycin Streptomycin		
×	Pyrazinamide Rifabutin (b) Rifampicin Rifapentine Spectinomycin Streptomycin Sulfadiazine		
×	Pyrazinamide Rifabutin (b) Rifampicin Rifapentine Spectinomycin Streptomycin Sulfadiazine Tazobactam		
×	Pyrazinamide Rifabutin (b) Rifampicin Rifapentine Spectinomycin Streptomycin Sulfadiazine Tazobactam Tetracyclines		
×	Pyrazinamide Rifabutin (b) Rifampicin Rifapentine Spectinomycin Streptomycin Sulfadiazine Tazobactam Tetracyclines Trimethoprim/		
×	Pyrazinamide Rifabutin (b) Rifampicin Rifapentine Spectinomycin Streptomycin Sulfadiazine Tazobactam Tetracyclines		

	ıtrelvir/ritonavir (Paxlovid) may
	icoagulants/antiplatelets
	Apixaban
	Aspirin (antiplatelet)
	Clopidogrel (stented) (c)
	Dabigatran (d)
	Dalteparin
	Edoxaban <mark>(e)</mark>
	Enoxaparin
	Heparin
	Rivaroxaban
	Streptokinase
	Warfarin
	iconvulsants
×	Carbamazepine
_	Clonazepam
	Ethosuximide
	Lamotrigine
×	Phenobarbital
×	Phenytoin
_	Sodium valproate Valproate semisodium
	(Divalproex sodium)
	Valproic acid
Ant	idepressants
/	Amitriptyline
	Clomipramine
	Fluoxetine
	Lithium
×	St John's Wort
	idiabetics
	Glibenclamide
	Gliclazide
	Insulin
	Metformin
Ant	ifungals
	Amphotericin B
	Fluconazole
	Flucytosine
	Griseofulvin
	Itraconazole (f)
	Ketoconazole (f)
	Nystatin
	Voriconazole
	imalarials
	Amodiaquine
	Artemether
	Artesunate
	Atovaquone
_	Lumefantrine
	Mefloquine
	Piperaquine
	Primaquine
	Proguanil
	Quinine
Ant	ipsychotics Chlororomonia
	Chlorpromazine
	Clozapine
	Fluphenazine
∺	Haloperidol Risperidone
	Nisperiuone

Anxiolytics				
	Diazepam			
	Lorazepam			
	Midazolam			
Beta	a blockers			
	Atenolol			
	Bisoprolol			
	Carvedilol			
	Metoprolol			
	Propranolol			
Bro	nchodilators			
	Aminophylline			
	Ipratropium bromide			
	Salmeterol			
Calc	ium channel blockers			
	Amlodipine			
	Nifedipine			
	Verapamil			
Can	cer drugs			
	Dasatinib <mark>(g)</mark>			
	Erlotinib <mark>(h)</mark>			
	Imatinib <mark>(i)</mark>			
	Methotrexate			
	Vinblastine (j)			
Con	traceptives			
	Ethinylestradiol			
	Etonogestrel (IMP)			
	Etonogestrel (VR)			
	Levonorgestrel (COC)			
	Levonorgestrel (EC)			
	Levonorgestrel (IUD)			
	Levonorgestrel (POP)			
	Medroxyprogesterone			
	(depot injection)			
	Norethisterone (COC)			
	Norethisterone (IM)			
	Norethisterone (POP)			
<u></u>	Norgestrel (COC)			
COV	/ID19 therapies			
	Budesonide (inhaled)			
	Convalescent plasma			
	Dexamethasone			
	Hydrocortisone Infliximab			
	Methylprednisolone COVID19 vaccines			
Gas				
Gas	trointestinal agents Aprepitant			
	Domperidone Lactulose			
	Loperamide Mesalazine			
	Metoclopramide			
	Omeprazole			
	Ondansetron			
	Ranitidine			
	Senna			
HCV	/ antivirals			
	Glecaprevir/pibrentasvir			
	Ledipasvir/sofosbuvir			
	Ombitasvir/paritaprevir/r			

Sofosbuvir/velpatasvir

Her	bals/supplements
	Folic acid
	Magnesium
×	St John's Wort
HIV	antiretrovirals
	Abacavir
	Atazanavir/ritonavir
	Darunavir/ritonavir
	Dolutegravir
	Efavirenz
	Emtricitabine
	Lamivudine
	Lopinavir/ritonavir
	Nevirapine
	Raltegravir
	Tenofovir alafenamide
	Tenofovir-DF
	Zidovudine
Нур	ertension/heart failure
10	Amiloride
	Dopamine
	Enalapril
	Furosemide
	Hydrochlorothiazide
	Isosorbide dinitrate
	Lisinopril
	Losartan
	Methyldopa
	Spironolactone
Immunosuppressants	
	Azathioprine
	Ciclosporin (k)
	Everolimus
Lini	d lowering agents
	Atorvastatin
	Fluvastatin
	Lovastatin
	Simvastatin
Oth	
oui	Allopurinol
	Ergometrine
	Ergotamine
	Levodopa Levothyroxine
Stor	oids
Ster	Beclometasone
	Betamethasone
	Fludrocortisone
	Prednisolone
	Testosterone
	Triamcinolone

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Legend

Colo	our/Symbol	Recommendation for NMV/r use
I	Do not co-administer	Do not use NMV/r \Rightarrow alternative COVID-19 therapy
_		Risk of serious toxicity. Stopping the drug does not mitigate the interaction due to its prolonged half-life.
×	Do not co-administer	Do not use NMV/r \Rightarrow alternative COVID-19 therapy
		Strong inducer can jeopardize NMV/r efficacy due to persisting induction after stopping the drug.
	Do not co-administer	NMV/r use ONLY possible if drug is paused or replaced by a non-interacting drug
		Risk of serious toxicity. Only start NMV/r if the drug can be safely paused or replaced.
		Drug can be resumed at least 3 days (if possible, up to 5 days for narrow therapeutic index drugs) after completing NMV/r therapy.
	Potential interaction	Stop or replace drug if possible or consult specialist for dose adjustment/monitoring to allow use with NMV/r
	Dose adjustment and/or	Ideally, only start NMV/r if the drug can be safely paused or replaced.
	close monitoring required.	Alternatively, dose adjust/monitor. Refer to www.covid19-druginteractions.org for detailed information.
	Potential interaction	Proceed with NMV/r
	Manageable by	Interaction manageable by counselling the patient about potential interaction and advising to temporarily stop
	counselling patient	the drug if feeling unwell.
	Weak interaction	Proceed with NMV/r
	No action needed	Drug metabolized partially by CYP3A4 or with low risk of adverse event from interaction.
	No interaction expected	Proceed with NMV/r

Contraceptive Abbreviations

COC = combined oral contraceptive IUD = intrauterine device EC = emergency contraception IM = intramuscular IMP = implant

POP = progestin only contraceptive pill VR = vaginal ring

Notes

- No dose reduction or monitoring in patients with normal renal function. а
- b Rifabutin dosed 150 mg once daily with NMV/r.
- c Ritonavir decreases clopidogrel efficacy therefore NMV/r cannot be prescribed in high risk situation (i.e. initial period (at least 6 weeks) post coronary stenting). NMV/r is allowed if clopidogrel is used outside this period or if clopidogrel is used as alternative to aspirin (intolerant patients).
- d When used for the treatment of atrial fibrillation, reduce dabigatran to 110 mg twice daily in individuals with normal renal function and to 75 mg twice daily in individuals with moderate renal impairment. Consult www.covid19-druginteractions.org for management in other indications.
- When used for the treatment of atrial fibrillation, reduce edoxaban to 30 mg. Consult www.covid19-druginteractions.org for management in other indications.
- Itraconazole or ketoconazole should not be used at doses >200 mg/day.
- The decision to pause or dose adjust dasatinib should be made in conjunction with the patient's oncologist. g Chronic phase chronic myelogenous leukaemia: pause dasatinib and restart 3 days after completing NMV/r. Alternatively, consider reducing dasatinib dose to 20 mg (in patients receiving 100 mg daily) or 40 mg (in patients receiving 140 mg daily) and monitor for toxicity. Accelerated or blast phase chronic myelogenous leukaemia: do not coadminister, use alternative COVID-19 therapy.
- The decision to pause or dose adjust erlotinib should be made in conjunction with the patient's oncologist. h If it is decided to pause treatment, restart erlotinib 3 days after completing NMV/r treatment. If pausing erlotinib treatment is not feasible, continue full dose erlotinib with patient self-monitoring for rash and diarrhoea. If these do occur, reduce erlotinib dose in 50 mg decrements or re-assess for a short pause.
- The decision to pause imatinib should be made in conjunction with the patient's oncologist. If it is decided to hold treatment, restart imatinib 3 days after completing NMV/r treatment. Alternatively, imatinib may be coadministered with monitoring for adverse effects (fluid retention, nausea and neutropenia). NMV/r is expected to have a modest effect on imatinib exposure. Coadministration with ritonavir (600 mg once daily) for 3 days did not significantly alter imatinib exposure (van Erp NP et al. Clin Cancer Res. 2007;13(24):7394-400).
- The decision to pause or dose adjust vinblastine should be made in conjunction with the patient's oncologist. Vinblastine may be paused in the context of acute infection. Restart vinblastine 3 days after completing NMV/r treatment. Alternatively, vinblastine may be coadministered with close monitoring for haematologic toxicity 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.
- The management of this interaction is challenging and would require dosage adjustment and therapeutic drug monitoring (TDM) of ciclosporin which may not be possible given the short duration of NMV/r treatment. An alternative COVID treatment should be considered. However, if TDM is available, an empiric dose reduction of ciclosporin has been suggested (reduce total daily dose by 80% and administer once daily) during treatment with NMV/r (days 1-5). Ciclosporin concentrations should be assessed on day 6 or 7 and repeated every 2-4 days.

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