## Resource: ART Drug-Drug Interactions

August 2024

Table 13: Etravirine (ETR) Interactions (also see drug package inserts)			
Class or Drug	Mechanism of Action	Clinical Comments	
Aliskiren	ETR is a minor inhibitor of P-gP and may minimally increase aliskiren concentrations, but this has not been studied.	When using with ETR, monitor for aliskiren-related adverse effects; switch to alternative antihypertensive medicine or ARV if necessary.	
Warfarin	Metabolism of warfarin could potentially increase (or more rarely decrease).	<ul> <li>Use cautiously with warfarin; if use is necessary, increase INR monitoring.</li> <li>If INR increases, decrease warfarin dose.</li> <li>If INR decreases, increase warfarin dose slowly.</li> </ul>	
Antiplatelet medications [Kakuda, et al. 2011; Rathbun and Liedtke 2010]	<ul> <li>Cilostazol: ETR may reduce cilostazol concentrations.</li> <li>Dipyridamole: ETR may induce UGT enzymes, which are responsible for metabolism.</li> <li>Ticagrelor, clopidogrel: ETR reduces ticagrelor concentrations and conversion of clopidogrel to its active metabolite.</li> <li>Vorapaxar: When coadministered with ETR, vorapaxar levels expected to be reduced.</li> </ul>	<ul> <li>Cilostazol: Monitor for antiplatelet effect; may be necessary to use alternative antiplatelet medication or alternative ARV.</li> <li>Dipyridamole: Monitor for antiplatelet effect; use another ARV if necessary.</li> <li>Ticagrelor, clopidogrel: Use with ETR may reduce antiplatelet effect; monitor closely for efficacy and use alternative ARV if possible.</li> <li>Prasugrel: When coadministered with ETR, no dose adjustments are necessary.</li> <li>Vorapaxar: No data available.</li> </ul>	
Statins	<ul> <li>Simvastatin, lovastatin: ETR may decrease concentrations.</li> <li>Atorvastatin, pravastatin, fluvastatin: ETR may modestly reduce concentrations.</li> </ul>	<ul> <li>Simvastatin, lovastatin: Monitor for efficacy. May warrant increases in statin dose. Do not increase dose above maximum recommended statin dose.</li> <li>Atorvastatin, pravastatin, fluvastatin: Monitor for cholesterol-lowering effect of statins. May require increased dose if necessary.</li> </ul>	
Saxagliptin, sitagliptin	ETR may decrease concentrations.	Monitor for efficacy; if necessary, increase dose of DPP-4 inhibitor.	
Inhaled and injected corticosteroids	Coadministration may reduce corticosteroid concentrations.	<b>Dexamethasone (systemic):</b> Consider alternative corticosteroid for long-term use; if benefits of use outweigh risks, monitor for virologic response.	
Trazodone	ETR may decrease trazodone concentrations.	Monitor for antidepressant and/or sedative effects.	
Bupropion	No significant interactions are expected.	Monitor for clinical effect and increase as needed, but do not exceed recommended maximum dose.	



	actions (also see drug package inserts)	
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Alprazolam	ETR may reduce alprazolam concentrations.	Monitor for benzodiazepine withdrawal.
Diazepam	ETR may reduce diazepam concentrations.	No dose adjustments are necessary.
Sleep medications	Zolpidem: ETR may reduce zolpidem concentrations.	<ul> <li>Zolpidem, eszopiclone: Monitor for efficacy; no dose adjustments are recommended.</li> <li>Suvorexant: Monitor for efficacy; do not exceed 20 mg per day.</li> </ul>
Antipsychotics	<ul> <li>Aripiprazole, brexpiprazole: ETR may decrease aripiprazole and brexpiprazole concentrations.</li> <li>Risperidone: ETR may decrease risperidone efficacy.</li> </ul>	<b>Aripiprazole, brexpiprazole, risperidone:</b> Titrate slowly to achieve clinical effect; monitor for efficacy and adverse effects.
Carbamazepine, oxcarbazepine, phenobarbital, phenytoin	Coadministration may significantly reduce concentrations of ARVs through induction of CYP450 system.	<ul> <li>Coadministration is not recommended; use alternative anticonvulsant.</li> <li>If benefit of use outweighs risk, monitor carefully for efficacy and toxicity.</li> <li>Perform TDM if use cannot be avoided.</li> </ul>
Lamotrigine, zonisamide	ETR may reduce lamotrigine or zonisamide efficacy.	Titrate slowly to achieve clinical effect; monitor for efficacy.
Hormonal contraceptives	Information is based on what is known with EFV drug interactions.	<ul> <li>Etonogestrel: No data available; consider alternative or additional contraceptive methods or alternative ARV.</li> <li>Ulipristal: Monitor closely for reduced efficacy.</li> </ul>
Erectile and sexual dysfunction agents	<ul> <li>PDE5 inhibitors: ETR may reduce effectiveness of PDE5 inhibitors (sildenafil, vardenafil, and tadalafil).</li> <li>Flibanserin: ETR may reduce flibanserin concentrations.</li> </ul>	<ul> <li>PDE5 inhibitors: Monitor for clinical effect; if dose increase is needed to achieve desired clinical effect, titrate under medical supervision to lowest effective dose.</li> <li>Flibanserin: Do not coadminister.</li> </ul>
Buprenorphine	No significant interactions are expected.	Titrate opioid or antagonist as required to achieve clinical effect; monitor for signs of withdrawal or opioid toxicity.
Methadone	ETR may slightly increase methadone concentrations.	<ul> <li>Titrate opioid or antagonist as required to achieve clinical effect; monitor for signs of withdrawal or opioid toxicity.</li> <li>Monitor for signs of methadone toxicity; reduce dose if necessary.</li> </ul>
Cyclosporine, tacrolimus	ETR may lower concentrations.	<ul> <li>Adjust cyclosporine and tacrolimus dose based on efficacy and TDM.</li> <li>Conduct TDM more frequently for 2 weeks when starting or stopping NNRTI therapy.</li> </ul>
HCV PIs ("-previr" drugs) [Mak, et al. 2018; Kaur, et al. 2015; Yeh 2015]	ETR may decrease HCV PI levels through CYP3A induction.	Do not coadminister.
Sofosbuvir/velpatasvir (available as coformulated product) [Greig 2016]	ETR may decrease velpatasvir levels through CYP3A induction and (weak) P-gP inhibition.	Do not coadminister.
Daclatasvir [Garrison, et al. 2018]	ETR induces CYP3A, lowering daclatasvir levels.	Increase dose of daclatasvir to 90 mg per day.



Class or Drug	Mechanism of Action	Clinical Comments
Atazanavir (ATV) [Marzolini, et al. 2016; Orrell, et al. 2015]	<ul> <li>ETR is a substrate and inducer of CYP3A4.</li> <li>COBI and ATV are substrates and inhibitors of CYP3A4.</li> </ul>	<ul> <li>Administration with RTV-boosted ATV results in decreased ATV exposure, but decrease is not considered relevant; no dose adjustments are necessary.</li> <li>Due to potential for decreased ARV efficacy, avoid use of ETR with COBI. When these medications are given together, COBI concentrations are decreased.</li> <li>When possible, avoid concomitant use of ETR and unboosted ATV. ETR with unboosted ATV results in significant decreases in ATV exposure.</li> </ul>
Dolutegravir (DTG)	ETR induces UGT1A1 and CYP3A enzymes.	ETR reduces DTG concentrations. Do not use concomitantly
[Green, et al. 2017]	DTG is a substrate of UGT1A1 and CYP3A enzymes.	unless boosted PI is also part of treatment regimen.
Lenacapavir (LEN)	CYP3A4 and P-gP induction associated with concomitant HIV treatment potentially deceases LEN levels.	Do not coadminister.
Rifabutin, rifampin, rifapentine	<ul> <li>Rifabutin: When used concomitantly, increased rifabutin levels are expected and decreased ETR levels may occur.</li> <li>Rifampin, rifapentine: CYP3A induction reduces ETR bioavailability.</li> </ul>	<ul> <li>Rifabutin:         <ul> <li>If ETR and rifabutin are used concomitantly, dose rifabutin at 300 mg daily, with no changes to ETR dose. Continue rifabutin 300 mg daily dosing until at least 2 weeks after rifabutin is stopped.</li> <li>Concomitant use of boosted PI with ETR and rifabutin is contraindicated.</li> </ul> </li> <li>Rifampin, rifapentine: Concomitant use is contraindicated.</li> </ul>
Mpox treatments	<b>Tecovirimat</b> is a weak inducer of CYP3A and a weak inhibitor of CYP2C8 and CYP2C19; use may increase or decrease plasma concentrations of other medications.	<b>Tecovirimat</b> may reduce NNRTI levels, though effects are not likely to be clinically relevant. No dose adjustment in either drug is necessary.
Gender-affirming hormones	<ul> <li>Estradiol: ETR may reduce estradiol levels via CYP3A induction.</li> <li>Finasteride, dutasteride: ETR may reduce finasteride and dutasteride levels via CYP3A induction.</li> <li>Progestins (oral medroxyprogesterone, micronized progesterone): ETR may reduce progestin levels via CYP3A induction.</li> <li>Testosterone: ETR may reduce testosterone levels via CYP3A induction.</li> </ul>	<ul> <li>Estradiol: Monitor serum estradiol levels; may require increased estradiol dose.</li> <li>Finasteride, dutasteride, progestins (oral medroxyprogesterone, micronized progesterone), testosterone: No dose adjustments are necessary.</li> </ul>
ADHD medications	Modafinil: CYP3A4 induction may reduce NNRTI levels.	<b>Modafinil:</b> Avoid concurrent use due to potential loss of virologic response.

**Abbreviations:** ADHD, attention-deficit/hyperactivity disorder; ARV, antiretroviral; COBI, cobicistat; CYP, cytochrome P450; DPP-4, dipeptidyl peptidase-4; EFV, efavirenz; HCV, hepatitis C virus; INR, international normalized ratio; NNRTI, non-nucleoside reverse transcriptase inhibitor; P-gP, P-glycoprotein; PDE5, phosphodiesterase type 5; PI, protease inhibitor; RTV, ritonavir; TDM, therapeutic drug monitoring; UGT, uridine diphosphate glucuronosyltransferase; VIGIV, vaccinia immune globulin intravenous.

No significant interactions/no dose adjustments necessary (see guideline section <u>Drug-Drug Interactions by Common Medication Class</u>): Common oral antibiotics; acid-reducing agents; polyvalent cations; asthma and allergy medications; long-acting beta agonists; nonopioid pain medications; opioid analgesics and tramadol; alpha-adrenergic antagonists for benign prostatic hyperplasia; alcohol, disulfiram, and acamprosate; COVID-19 therapeutics.



## References

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