

DRUG INTERACTIONS WITH TOBACCO SMOKE

Many interactions between tobacco smoke and medications have been identified. Note that in most cases it is the tobacco smoke—not the nicotine—that causes these drug interactions. Tobacco smoke interacts with medications through pharmacokinetic (PK) and pharmacodynamic (PD) mechanisms. PK interactions affect the absorption, distribution, metabolism, or elimination of other drugs, potentially causing an altered pharmacologic response. The majority of PK interactions with smoking are the result of induction of hepatic cytochrome P450 enzymes (primarily CYP1A2). PD interactions alter the expected response or actions of other drugs. The amount of tobacco smoking needed to have an effect has not been established, and the assumption is that any smoker is susceptible to the same degree of interaction. The most clinically significant interactions are depicted in the shaded rows.

•	ction. The most clinically significant interactions are depicted in the shaded rows.
Drug/Class	MECHANISM OF INTERACTION AND EFFECTS
Pharmacokinetic Interactions	
Alprazolam (Xanax)	 Conflicting data on significance, but possible
Bendamustine (Treanda)	■ Metabolized by CYP1A2. Manufacturer recommends using with caution in smokers due to likely ↓
	bendamustine concentrations, with ↑ concentrations of its two active metabolites.
Caffeine	■ ↑ Metabolism (induction of CYP1A2); ↑ clearance (56%). Caffeine levels likely ↑ after cessation.
Chlorpromazine (Thorazine)	■ ↓ Area under the curve (AUC) (36%) and serum concentrations (24%).
	
Clopidogrel (Plavix)	 Metabolism (induction of CYP1A2) of clopidogrel to its active metabolite.
	 Clopidogrel's effects are enhanced in smokers (≥10 cigarettes/day): significant ↑ platelet inhibition, ↓ platelet
	aggregation; while improved clinical outcomes have been shown, may also ↑ risk of bleeding.
Clozapine (Clozaril)	↑ Metabolism (induction of CYP1A2); ↓ plasma concentrations (18%).
	■ ↑ Levels upon cessation may occur; closely monitor drug levels and reduce dose as required to avoid toxicity.
Erlotinib (Tarceva)	↑ Clearance (24%); ↓ trough serum concentrations (2-fold).
Flecainide (Tambocor)	↑ Clearance (61%); ↓ trough serum concentrations (25%). Smokers may need ↑ dosages.
Fluvoxamine (Luvox)	■ ↑ Metabolism (induction of CYP1A2); ↑ clearance (24%); ↓ AUC (31%); ↓ plasma concentrations (32%).
	■ Dosage modifications not routinely recommended but smokers may need ↑ dosages.
Haloperidol (Haldol)	■ ↑ Clearance (44%); ✓ serum concentrations (70%).
Heparin	■ Mechanism unknown but ↑ clearance and ↓ half-life are observed. Smoking has prothrombotic effects.
	■ Smokers may need ↑ dosages due to PK and PD interactions.
Insulin, subcutaneous	Possible
	endogenous substances that cause insulin resistance.
	PK & PD interactions likely not clinically significant; smokers may need ↑ dosages.
Irinotecan (Camptosar)	 ↑ Clearance (18%); serum concentrations of active metabolite, SN-38 (~40%; via induction of glucuronidation); systemic exposure resulting in lower hematologic toxicity and may reduce efficacy.
	■ Smokers may need ↑ dosages.
Mexiletine (Mexitil)	↑ Clearance (25%; via oxidation and glucuronidation); ↓ half-life (36%).
	 ↑ Metabolism (induction of CYP1A2); ↑ clearance (98%); ↓ serum concentrations (12%).
Olanzapine (Zyprexa)	 Dosage modifications not routinely recommended but smokers may need ↑ dosages.
Propranolol (Inderal)	■ ↑ Clearance (77%; via side-chain oxidation and glucuronidation).
Ropinirole (Requip)	■ V Cmax (30%) and AUC (38%) in study with patients with restless legs syndrome.
	■ Smokers may need ↑ dosages.
Tacrine (Cognex)	↑ Metabolism (induction of CYP1A2); ↓ half-life (50%); serum concentrations 3-fold lower.
	■ Smokers may need ↑ dosages.
Theophylline	↑ Metabolism (induction of CYP1A2); ↑ clearance (58–100%); ↓ half-life (63%).
(Theo Dur, etc.)	 Levels should be monitored if smoking is initiated, discontinued, or changed. Maintenance doses are
	considerably higher in smokers; ↑ clearance with second-hand smoke exposure.
Tricyclic antidepressants (e.g.,	 Possible interaction with tricyclic antidepressants in the direction of ↓ blood levels, but the clinical significance is
imipramine, nortriptyline)	not established.
Tizanidine (Zanaflex)	 ↓ AUC (30-40%) and ↓ half-life (10%) observed in male smokers.
Warfarin	■ ↑ Metabolism (induction of CYP1A2) of R-enantiomer; however, S-enantiomer is more potent and effect on INR
	is inconclusive. Consider monitoring INR upon smoking cessation.
Pharmacodynamic Interaction	
Benzodiazepines (diazepam, chlordiazepoxide)	■ ↓ Sedation and drowsiness, possibly caused by nicotine stimulation of central nervous system.
Beta-blockers	 Less effective antihypertensive and heart rate control effects; possibly caused by nicotine-mediated sympathetic
	activation.
	■ Smokers may need ↑ dosages.
Corticosteroids, inhaled	Smokers with asthma may have less of a response to inhaled corticosteroids.
Hormonal contraceptives	■ ↑ Risk of cardiovascular adverse effects (e.g., stroke, myocardial infarction, thromboembolism) in women who
	smoke and use oral contraceptives. Ortho Evra patch users shown to have 2-fold \uparrow risk of venous
	thromboembolism compared to oral contraceptive users, likely due to ↑ estrogen exposure (60% higher levels). ↑ Risk with age and with heavy smoking (≥15 cigarettes per day) and is quite marked in women ≥35 years old.
Onicide (numerous trans	 ↑ Risk with age and with neavy smoking (≥15 cigarettes per day) and is quite marked in women ≥35 years old. ↓ Analgesic effect; smoking might ↑ the metabolism of propoxyphene (15–20%) and pentazocine (40%).
Opioids (propoxyphene, pentazocine)	■ Analgesic effect; smoking might 1 the metabolism of propoxyphene (15–20%) and pentazocine (40%). Mechanism unknown.
	■ Smokers may need ↑ opioid dosages for pain relief.
	1