

Drug Interactions

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Drug Interactions

Drug Interaction:

- *The pharmacologic or clinical response to the administration of a drug combination different from that anticipated from the known effects of the two agents when given alone*
- **May be harmful:** toxicity, reduced efficacy
- **May be beneficial:** synergistic combinations, pharmacokinetic boosting, increased convenience, reduced toxicity, cost reduction

Beneficial Drug Interactions

Saquinavir & ritonavir

- Saquinavir poorly absorbed, TID dosing, high pill burden (18 caps per day!)
- Combination with ritonavir results in 20-fold increase in C_{ps}
- Allows for BID dosing and decreased dose from 1200 mg TID to 1000 mg BID (1600 QD dosing is also possible)

Indinavir, amprenavir, atazanavir + ritonavir

Cyclosporin and ketoconazole

- Difficult to determine doses due to large interpatient variability in CYP3A (and P-gp) activity

PK Interactions between PIs: Pharmacoenhancement

Comparison of “unboosted” and “boosted” medication and resulting plasma drug concentration.

Slide Courtesy of Dr. David Back. 9th Conference on Retroviruses and Opportunistic Infections. February 24-28, 2002.

Beneficial Drug Interactions: RTV + SQV

Graphical illustration of a ritonavir-saquinavir interaction resulting in higher plasma concentrations of saquinavir.

Buss N et al. Br J Clin Pharmacol. 2001 Sep;52(3):255-64.

Epidemiology of Drug-Drug Interactions

True incidence difficult to determine

Data for drug-related hospital admissions do not separate out drug interactions, focus on ADRs

Most data are in the form of case reports

Missing or incomplete information

Patients receiving polypharmacy are at risk

77% of HIV patients on protease inhibitors experience drug interactions

Difficulty in assessing role of OTC and herbal drugs in drug interactions

Questions regarding “active” ingredient in herbal meds

Types of Drug Interactions

Pharmacodynamic

- Related to the drug's effects in the body
- One drug modulates the pharmacologic effect of another: additive, synergistic, or antagonistic

Pharmacokinetic

- What the body does with the drug
- One drug alters the concentration of another
- *Usually* mediated by cytochrome P450 (CYP)

Pharmacokinetic Interactions

Graphic illustration of ADME interactions.

Drug Interactions

Absorption: food, chelation, G.I. motility, pH

Distribution: transport, protein binding

Metabolism: Phase I (CYP450), Phase II (conjugation)

Elimination: Renal (glomerular filtration); transport

Alterations in Absorption

Administration with food

- **Decreased rate of absorption; not extent (\leftrightarrow AUC):**
Common for many drugs; take without regard to meals
- **Decreased extent of absorption (\downarrow AUC):**
Indinavir AUC decreased by 77% with high calorie meal; take on an empty stomach
- **Increased extent of absorption (\uparrow AUC):**
Itraconazole (capsules) AUC increased by 66% with standard meal

Alterations in Absorption: Food Effects

Graphs illustrating that food enhances absorption of indinavir and itraconazole.

Pharm Res. 1999 May;16(5):718-24

Graph

Antimicrob Agents Chemother. 1993 Apr;37(4):778-84.

Alterations in Absorption: Chelation

Chelation

- Irreversible binding of drugs in the GI tract
- Tetracyclines, quinolone antibiotics - ferrous sulfate (Fe^{+2}), antacids (Al^{+3} , Ca^{+2} , Mg^{+2}), dairy products (Ca^{+2})
- Usually separating administration of chelating drugs by 2+ hours decreases interaction effect

Graph of Trovafloxacin +/- Maalox®

J Antimicrob Chemother. 1997 Jun;39 Suppl B:93-7.

Alterations in Absorption: GI Motility

- ↑ GI motility: cisapride, metoclopramide
- ↓ motility: narcotics, antidiarrheals, high calorie meal / viscosity (delayed gastric emptying)

Graph showing that methadone delays absorption of stavudine, ug/L over time, h

Acquir Immune Defic Syndr. 2000 Jul 1;24(3):241-8.

Alterations in Absorption

Graph of Ketoconazole Conc. (mcg/mL) over time.

Sucralfate and ranitidine reduce Ketoconazole absorption.

Piscitelli S et al. Antimicrob Agents Chemother 1991;35:1765-1771

Drug Interactions: Transport Proteins

Graphic illustration of intestinal, hepatic, renal, and brain drug transporters.

Ayrton A, Morgan P. Role of transport proteins in drug absorption, distribution and excretion. *Xenobiotica*. 2001;31:469-97.

Alterations in Absorption: Drug Transport

Efflux proteins

- P-glycoprotein, MRP1, MRP2, OAT3
- Extrude drug from gut back into lumen limiting drug absorption
- Transporter induction may result in ↓ absorption
- Transporter inhibition may result in ↑ absorption
- Effects often difficult to assess (vs. metabolism; vs. anatomic site)
- Inhibition may be of clinical significance for drugs that are large molecules, have low bioavailability, dissolve slowly and/or incompletely (clinical significance may be overstated in medical literature)

Simplified Example of P-gp Function

Graphic illustration of P-glycoprotein function in lymphocytes, brain, g.i. tract, kidney, placenta, and liver.

Drug Interactions: Transport Proteins Concept using the kidney

Graphic illustration with example of clarithromycin blocking P-glycoprotein and digoxin secretion in the kidney.

CLINICAL APPLICATION: HEALTHY HUMAN VOLUNTEERS

	Digoxin + Placebo	Digoxin + Clarithromycin	P
AUC 0-24 hr	14	23	< .05
ClR (mL/min)	57	34	< .05

Rengelshausen et al. Brit J Clin Pharmacol 2003;56:32-8.

Alterations in Absorption: Drug Transport

Uptake proteins

- **OATP: located on the luminal border of enterocytes**
- **Transport drug across lumen and promote absorption**
- **Transporter inhibition may result in ↓ absorption and reduced bioavailability**
- **OATP substrates**
 - Pravastatin, digoxin, fexofenadine, benzylpenicillin
- **OATP inhibitors**
 - Fruit juices, ritonavir, saquinavir, lovastatin, others?
- **In the intestine, OATP functions *OPPOSITE* of P-gp (i.e. P-gp inhibition *INCREASES* drug absorption while OATP *DECREASES* drug absorption for compounds that are substrates of both proteins**

OATP Function: INTESTINE

Graphic illustration of this function.

Alterations in Absorption: Drug Transport

Graph of Fexofenadine conc. over time and changes caused by grapefruit, orange, and apple juice.

Clin Pharmacol Ther. 2002 Jan;71(1):11-20.

Alterations in Absorption: anion exchange resins and Gut Flora Modulation

Anion exchange resins (i.e. cholestyramine)

- Form insoluble complexes with various drugs reducing their absorption
 - Warfarin, digoxin, β -blockers, NSAIDS, others?
- Stagger dose of exchange resin with other meds
 - Difficult due to multiple daily dosing of cholestyramine

Inhibition of drug-metabolizing enteric bacteria

- Antibiotics

Digoxin (*Eubacterium lentum*)

Oral contraceptives (bacteria hydrolyze steroid conjugates)

- √ Reports of unplanned pregnancy: causal relationship with antimicrobial administration is controversial

Distribution: Protein Binding Interactions
Non-restrictively cleared drugs

Eliminating organ removing most of the drug being presented to it, including the fraction bound to plasma proteins

Increase in f_u will not lead to a proportional increase in CL

No examples of clinically significant protein binding interactions have been identified with non-restrictively cleared drugs

Distribution: Protein Binding Interactions

Restrictively cleared drugs

Small fraction of drug extracted during single passage through the eliminating organ ($E \leq f_{ub}$)

Only unbound drug in plasma can be cleared

Increase in f_u leads to proportional increase in total drug CL and decrease in total drug C_{pss}

$C_{pss_{ub}}$ will return to pre-displacement value after transient increase

- Only likely to be clinically significant for drugs with LONG $T_{1/2}$, SMALL V_d , narrow therapeutic range, \uparrow PPB
- Example: warfarin displacement from serum albumin by a metabolite of chloral hydrate (trichloroacetic acid)

Distribution: Protein Binding Interactions

Graphs showing Warfarin Concentration ($\mu\text{g/mL}$) and Prothombin Time (sec) over time.

Warfarin + Tri-CA

Fub : \uparrow

Cub : transient \uparrow (then \leftrightarrow)

Total CL: \uparrow

Ctotal : \downarrow

Cbound : \downarrow

Principles of Clinical Pharmacology, pg 64

Alterations in Distribution: Protein Binding

“...the overall clinical importance of plasma protein binding displacement interactions continues to be overstated...”

“Despite the theoretical and experimental data to the contrary, the concept that plasma protein binding displacement is a common cause of clinically significant interactions may still be widely taught in some medical schools, often appears in textbooks and is accepted by many in the medical community and by drug regulators.”

Sansom LN & Evans AM. Drug Safety 1995;12:227-233.

Rolan PE. Br J Clin Pharmacol 1994;37:125-128.

Distribution: Drug Transport (P-gp)

Graph showing Nelfinavir level, ng/gm over time (hours)

Effect of P-glycoprotein inhibitor.

• **Tissue 14C NFV conc. in brain**
14C NFV + LY-335979 (P-gp inhibitor)

○ *Plasma 14C NFV concentration*
14C NFV + LY-335979 (P-gp inhibitor)

△ **Plasma 14C NFV concentration**
14C NFV + vehicle

Tissue 14C NFV conc. in brain
14C NFV + vehicle

Choo EF et al. Drug Metab Disposit 2000;28:655-660.

Drug Metabolism Interactions

Drug metabolism

- **Chemical modification of a xenobiotic**
- **Phase I (functionalization RX)**
Cytochrome P450 (CYP): i.e. CYP3A4, CYP2D6, CYP1A2 etc.
- **Phase II (synthetic RX)**
Conjugation: i.e. glucuronidation (UGT1A1 etc.)
- **Purpose: detoxification of foreign compounds**
- **Anatomic sites: Liver*, Gut*, kidney, lung, brain etc.**

Drug Metabolism Interactions

Graphic illustration of CYP enzyme induction and inhibition.

CYP 450 Substrates

Drugs may be metabolized by a single isoenzyme

Desipramine/CYP2D6; indinavir/3A4; midazolam/3A, caffeine/CYP1A2;
omeprazole/CYP2C19

Drugs may be metabolized by multiple isoenzymes

Most drugs metabolized by more than one isozyme

Imipramine: CYP2D6, CYP1A2, CYP3A4, CYP2C19

If co-administered with CYP450 inhibitor, some isozymes may “pick up
slack” for inhibited isozyme

Extensive listing + references:

<http://medicine.iupui.edu/flockhart/table.htm>

CYP 450 Enzyme Inhibition

Usually by competitive binding to enzyme site

Onset and offset dependent on the half-life and time to steady-state of the inhibitor

Fluoxetine & CYP2D6; ritonavir and CYP3A4

Time to maximum interaction effect dependent on time required for substrate drug to reach new steady-state

Mechanism-based enzyme inactivation

Grapefruit juice and intestinal CYP3A content

Enzyme Inhibition

Graph showing Drug Conc. over time (days).

A higher steady-state concentration is reached over time with continuous enzyme inhibition.

CYP 450 Inhibitors

The “usual suspects”

Cimetidine (various)

Erythromycin, clarithromycin (3A4)

Ketoconazole, itraconazole (3A4)

HIV protease inhibitors (esp. ritonavir)

Fluoxetine, paroxetine (CYP2D6)

Nefazodone (CYP3A4)

Grapefruit Juice (intestinal CYP3A4 only)

Extensive listing with references:

<http://medicine.iupui.edu/flockhart/table.htm>

Sildenafil (Viagra®) + Grapefruit Juice

Graph showing mean plasma concentration (ng/ml) after dosing (hours)

Jeter A et al. Clin Pharmacol Ther. 2002 Jan;71(1):21-9.

CYP450 Inhibition

Key questions:

- **What is the toxic potential and therapeutic index of the substrate**
Terfenadine or digoxin vs sertraline

- **What are the other pathways involved in the metabolism of the substrate**
Zolpidem vs triazolam

- **Does the substrate have active metabolites?**
Codeine converted to morphine

CYP450 Induction

The “usual suspects”

Rifampin

Rifabutin

Carbamazepine

Phenobarbital

Phenytoin

Nevirapine, efavirenz

St. John’s wort

Troglitazone, pioglitazone

<http://medicine.iupui.edu/flockhart/table.htm>

CYP450 Induction

Gradual onset and offset

(involves increased DNA transcription and synthesis of new CYP enzymes)

Onset and offset

Depends on $T_{1/2}$ of inducer, time to make new CYP proteins, and rate of degradation of CYP proteins

Results in reduction of plasma concentration of substrate drugs

Risk of therapeutic failure

Removal of inducer may lead to toxic concentrations of substrate

Induction may lead to formation of toxic metabolite

Regulation of Drug Metabolism and Transport

Nuclear Receptors (NR)

- Largest known family of transcription factors
- Function as modulators of gene expression
- Ligand (drug, bile acid, hormone etc.) binds to vacant NR in the cytoplasm enters the nucleus & forms homo or heterodimers which complexes to promotor/enhancer regions of target genes
 - Simply put: the gene is “switched on” (or off) causing it to produce (or not produce) mRNA and subsequent proteins

Nuclear receptor activation

Graphic illustration of this activation

Adapted from Urquhart et al J Clin Pharm 2007;47:566-78

Summary of Nuclear Receptor Activators

TARGET GENE	NUCLEAR RECEPTOR	LIGANDS
CYP3A4	PXR, CAR, GR, HNF4, VDR, FXR	PXR: rifampin, dexamethasone + others
CYP2C9	PXR, CAR, GR	CAR: phenobarbital
CYP2C19	CAR, GR	GR: dexamethasone
CYP2B6	PXR, CAR	
MDR1	PXR, CAR	
OATP8	FXR	FXR: chenodeoxycholic acid

Adapted from Urquhart et al J Clin Pharm 2007;47:566-78

Enzyme Induction

Graphic illustration of drug conc. over time (days)

A lower steady-state concentration will be reached over time with continuous enzyme induction.

Induction: Influence of Ritonavir on Olanzapine Disposition in Healthy Volunteers

Chemical structure of olanzapine, a substrate of CYP1A2 and UDPGT.

Graph showing a reduced AUC for olanzapine when given with ritonavir.

Penzak SR et al. J Clin Psychopharm
2002;22:366-70

St. John's wort: CYP3A4 Induction Effects

Graph showing Indinavir Cp ($\mu\text{g/ml}$) over time

- 8 normal volunteers**
- Indinavir AUC determined before and after 14 days
SJW 300 mg TID**
- Indinavir AUC decreased by $57 \pm 19\%$ in presence
of SJW**

Piscitelli SC et al. Lancet 2000;355:547-8

Predicting Drug Interactions: in vitro Screening

Drug development: predicting *in vivo* drug interactions from *in vitro* data Microsomes, hepatocytes, liver slices, purified CYP enzymes etc.

Limitations and caveats

Most systems can only assess inhibition (not induction)

Methadone + ritonavir: discordant *in vivo* / *in vitro* results

Hard to extrapolate data when drugs have multiple CYP pathways

***In vitro* concentrations used may be excessively high**

Ritonavir inhibition of MRP2

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Predicting Drug Interactions: using CYP phenotypes

Probe + putative inhibitor or inducer

Measure probe (+/- metabolite(s) concentration(s)

Ratios of metabolite:parent compound

Examples of CYP probes

- CYP1A2: caffeine
- CYP2C9: tolbutamide; warfarin (+ vitamin K!)
- CYP2C19: S-mephenytoin; omeprazole
- CYP2E1: chlorzoxazone
- CYP2D6: dextromethorphan; debrisoquine; sparteine
- CYP3A4/5: midazolam
- CYP3A4: erythromycin

Herb-Drug Interactions Limitations

Since not regulated by FDA, safety & efficacy not required

- ***Little information available regarding drug interactions***
-

Extrapolation of data to available products difficult

- ***Independent lab tests many products
(<http://www.consumerlabs.com/>)***
- ***6/13 SAMe preparations did not pass testing
no detectable SAMe noted in one product***
- ***8/17 valerian preparations did not pass testing
4 - no detectable levels of valerenic acid
4 - 1/2 the amount claimed on the label***

Evaluation of Specific Drug Interactions

- **What is the time-course of the interaction**
Immediately or over a period of time
- **Is it a drug class effect**
omeprazole vs. lansoprazole
- **Is the interaction clinically significant**
Therapeutic index of drugs
Narrow or wide?
- **How should the interaction be managed?**
DC drug? Switch to another drug? Change dose?

Drug Interactions: General Tools for Evaluation and Management

Familiarity with metabolic pathways

Know where to locate information on interactions

Obtain thorough medication HX at each visit

Maintain high index of suspicion when:

Therapeutic response is less than expected

Toxic effects are present

Choose drugs that are less likely to interact

Consider TDM in certain situations (anti-TB TX)

Anti-TB and anti-HIV therapy

Drug Interactions: Resources

TABLE 5. WEB SITES WITH INFORMATION ABOUT DRUG INTERACTIONS.

www.dml.georgetown.edu/depts/pharmacology (Department of Pharmacology, Georgetown University Medical Center)

www.foodmedinteractions.com (food and drug interactions)

www.hivatis.org (HIV/AIDS Treatment Information Service)

www.hivdent.org (dental information)

hivinsite.ucsf.edu

www.hiv.net (in German)

www.hopkins-aids.edu (Johns Hopkins AIDS Service)

www.iapac.org (International Association of Physicians in AIDS Care)

www.hiv-druginteractions.org (Liverpool HIV Pharmacology Group)

www.medscape.com

Piscitelli SC, Gallicano KD. Interactions Among Drugs for HIV and Opportunistic Infections. N Engl J Med 2001;344:984-96.

THE END