# STE988734

Generated on January 29, 2017

#### ACE INHIBITORS, PLAIN

ADRB2	adrenoceptor beta 2, surface
	- Clinical Annotations —
hospital utilization when treated with	art failure may have increased emergency department visits and cardiovascular drugs as compared to patients with the AA or all factors may also influence efficacy of cardiovascular drugs.
ADRB1	adrenoceptor beta 1
	- Clinical Annotations —
	art failure may have increased emergency department utilizations as compared to patients with the GG genotype. Other genetic efficacy of cardiovascular drugs.
COMT	catechol-O-methyltransferase
	- Clinical Annotations —
of headache when discontinuing the u compared to patients with the AA ge	abstance withdrawal syndrome may have an increased likelihood use of analgesics (such as opioids, NSAIDs, triptans, ergot) as enotype. Other clinical and genetic factors may also influence withdrawal syndrome who discontinue the use of analgesics.
ANTIINFLAMMATORY AGENTS,	NON-STEROIDS
COMT	catechol-O-methyltransferase
	Clinical Annotations —

Class 3 rs4680 *GA* 

Patients with the AG genotype with substance withdrawal syndrome may have an increased likelihood of headache when discontinuing the use of analgesics (such as opioids, NSAIDs, triptans, ergot) as compared to patients with the AA genotype. Other clinical and genetic factors may also influence likelihood of headache in patients with withdrawal syndrome who discontinue the use of analgesics.

# ANTIVIRALS FOR TREATMENT OF HIV INFECTIONS, COMBINATIONS

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
creased risk for hepato	GG genotype who are co-infected with HIV and tuberculosis (TB) may have a detoxicity when treated with anti-tubercular and antiretroviral drugs as compared A genotype. Other genetic and clinical factors may also influence risk of hepato-
CYP2B6	cytochrome P450, family 2, subfamily B, polypeptide 6
creased risk for hepato	genotype who are co-infected with HIV and tuberculosis (TB) may have a detoxicity when treated with anti-tubercular and antiretroviral drugs as compared T genotype. Other genetic and clinical factors may also influence risk of hepato-
CYP2A7P1	cytochrome P450, family 2, subfamily A, polypeptide 7 pseudogene 1
creased risk for hepato	GG genotype who are co-infected with HIV and tuberculosis (TB) may have a detoxicity when treated with anti-tubercular and antiretroviral drugs as compared T genotype. Other genetic and clinical factors may also influence risk of hepato-
ETA BLOCKING A	GENTS
ADRB2	adrenoceptor beta 2, surface
hospital utilization wh	GG genotype and heart failure may have increased emergency department visits and hen treated with cardiovascular drugs as compared to patients with the AA or genetic and clinical factors may also influence efficacy of cardiovascular drugs.
ADRB1	adrenoceptor beta 1

Patients with the CC genotype and heart failure may have increased emergency department utilization when treated with cardiovascular drugs as compared to patients with the GG genotype. Other genetic and clinical factors may also influence efficacy of cardiovascular drugs.

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ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	Clinical Annotations

#### · Class 4 rs1045642 GG

People with the GG genotype may have decreased exposure to dabigatran compared to patients with the AA and AG genotypes, when also assessed with the rs2032582 allele. Other clinical and genetic factors may affect exposure to dabigatran.

· Class 4 rs2032582 CC

People with the CC genotype may have decreased exposure to dabigatran compared to patients with a variant at this position, including genotypes AA, AC, CT, and TT, when assessed in conjunction with a variant at position rs1045642. Other clinical and genetic factors may affect exposure to dabigatran.

#### DRUGS FOR TREATMENT OF TUBERCULOSIS

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member $1$

#### · Class 3 rs1045642 GG

Patients with the GG genotype and tuberculosis (TB) may have a decreased risk for hepatotoxicity when treated with anti-TB drugs as compared to patients with the AA genotype. Other genetic and clinical factors may also influence hepatotoxicity.

· Class 3 rs1045642 GG

Patients with the GG genotype who are co-infected with HIV and tuberculosis (TB) may have a decreased risk for hepatotoxicity when treated with anti-tubercular and antiretroviral drugs as compared to patients with the AA genotype. Other genetic and clinical factors may also influence risk of hepatotoxicity.

CYP2E1	cytochrome P450, family 2, subfamily E, polypeptide 1

#### · Class 3 rs2031920 CC

Patients with the CC genotype and tuberculosis (TB) may have an increased risk for hepatotoxicity when treated with anti-TB drugs as compared to patients with the CT or TT genotype. However, the majority of studies find no association with hepatotoxicity. Other genetic and clinical factors, such as variations in the NAT2 gene, may also influence risk for hepatotoxicity.

DUX1 double homeobox 1

when treated with anti- majority of studies find	CC genotype and tuberculosis (TB) may have an increased risk for hepatotoxicity TB drugs as compared to patients with the CT or TT genotype. However, the no association with hepatotoxicity. Other genetic and clinical factors, such as gene, may also influence risk for hepatotoxicity.
CYP2B6	cytochrome P450, family 2, subfamily B, polypeptide 6
creased risk for hepatot	GG genotype who are co-infected with HIV and tuberculosis (TB) may have a de-oxicity when treated with anti-tubercular and antiretroviral drugs as compared genotype. Other genetic and clinical factors may also influence risk of hepato-
CYP2A7P1	cytochrome P450, family 2, subfamily A, polypeptide 7 pseudogene 1
	- Clinical Annotations
creased risk for hepatot	genotype who are co-infected with HIV and tuberculosis (TB) may have a de- oxicity when treated with anti-tubercular and antiretroviral drugs as compared genotype. Other genetic and clinical factors may also influence risk of hepato-
NAT2	N-acetyltransferase 2 (arylamine N-acetyltransferase)
	- Clinical Annotations —
when treated with anti is present in a variety of	CC genotype and tuberculosis (TB) may have a decreased risk for hepatotoxicity -TB drugs as compared to patients with the CT or TT genotype. This SNP of NAT2 * alleles resulting in different NAT2 acetylator phenotypes, and is the 2*11. Other genetic and clinical factors may also influence hepatotoxicity.
OPIUM ALKALOIDS	AND DERIVATIVES
CHRNA3	cholinergic receptor, nicotinic, alpha 3 (neuronal)
	- Clinical Annotations
	GG genotype who are in chronic pain and receive opioid medications for treatment sk for addiction as compared to patients with the AA genotype. Other genetic

and clinical factors may also influence risk of opiate addiction.

——- Clinical Annotations ———

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	l Annotations
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#### · Class 3 rs16969968 GG

Patients with the GG genotype who are in chronic pain and receive opioid medications for treatment may be at decreased risk for addiction as compared to patients with the AA genotype. Other genetic and clinical factors may also influence risk of opiate addiction.

#### PLATINUM COMPOUNDS

# ABCB1 ATP-binding cassette, sub-family B (MDR/TAP), member 1 - Clinical Annotations

#### · Class 3 rs1045642 GG

Patients with the GG genotype and non-small-cell lung cancer may have a better response to platinum-based chemotherapy as compared to patients with the AA or AG genotype. This was only seen in those of Asian ethnicity. Other genetic and clinical factors may also influence response to platinum-based chemotherapy.

· Class 3 rs1128503 GG

Patients with the GG genotype and non-small cell lung cancer may have reduced risk of toxicities when treated with platinum-based chemotherapy compared to patients with the AA genotype. Other clinical and genetic factors may affect risk of toxicities in response to platinum-based chemotherapies.

#### PYRIMIDINE ANALOGUES

DPYD	dihydropyrimidine dehydrogenase
	ical Annotations —

#### · Class 1A rs55886062 AA

Patients with the AA genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have a decreased, but not absent, risk for drug toxicity as compared to patients with the AC or CC genotype (DPYD \*1/\*13 or \*13/\*13). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

#### · Class 1A rs3918290 CC

Patients with the CC genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased clearance of fluoropyrimidine drugs and 2) decreased, but not non-existent, risk for drug toxicity as compared to patients with the CT or TT genotype (DPYD \*1/\*2A or \*2A/\*2A). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine based chemotherapy.

#### Class 1A rs67376798 TT

Patients with the TT genotype and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased clearance of the drug and 2) decreased, but not absent, risk and reduced severity of drug toxicity as compared to patients with the AT genotype. Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

· Class 3 rs1801159 TT

Patients with the TT genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) a decreased likelihood of nausea, vomiting, and leukopenia, 2) increased response and 3) increased clearance of fluorouracil as compared to patients with the CT or CC genotype (DPYD \*1/\*5 or \*5/\*5). However, other studies find no associations or contradictory associations with fluoropyrimidine-induced drug toxicity or response. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

#### ACENOCOUMAROL

CYP2C9	cytochrome P450, family 2, subfamily C, polypeptide 9

#### · Class 2A rs1057910 AC

Patients with the AC genotype may require decreased dose of acenocoumarol or closer INR monitoring as compared to patients with the AA genotype. Other genetic and clinical factors may also influence acenocoumarol dose.

· Class 3 rs1799853 CC

Patients with the CC genotype who are taking acenocoumarol may have a decreased risk of a gastrointestinal hemorrhage as compared to patients with the CT or TT genotype. Other genetic and clinical factors may also influence risk of gastrointestinal hemorrhage.

VKORC1	vitamin K epoxide reductase complex, subunit 1

#### $\cdot$ Class 2A rs9934438 GA

Patients with the AG genotype may have decreased dose of acenocoumarol or phenprocoumon as compared to patients with genotype GG. Other genetic and clinical factors may also influence the dose of acenocoumarol or phenprocoumon.

PRSS53	protease, serine, 53

#### Class 2A rs9934438 *GA*

Patients with the AG genotype may have decreased dose of acenocoumarol or phenprocoumon as compared to patients with genotype GG. Other genetic and clinical factors may also influence the dose of acenocoumarol or phenprocoumon.

UGT1A	UDP glucuronosyltransferase 1 family, polypeptide A complex locus

#### · Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

UGT1A10	UDP glucuronosyltransferase 1 family, polypeptide A10

#### $\cdot$ Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

 $\cdot$  Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

UGT1A3	UDP glucuronosyltransferase 1 family, polypeptide A3

### · Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

 $\cdot$  Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

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UDP glucuronosyltransferase 1 family, polypeptide A4

—- Clinical Annotations —

#### · Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

UGT1A5

UDP glucuronosyltransferase 1 family, polypeptide A5

#### · Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

UGT1A6

UDP glucuronosyltransferase 1 family, polypeptide A6

—- Clinical Annotations —

#### · Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

UGT1A7	UDP glucuronosyltransferase 1 family, polypeptide A7
	- Clinical Annotations

 $\cdot$  Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

 $\cdot$  Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

· Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

UGT1A9	UDP glucuronosyltransferase 1 family, polypeptide A9
	nical Annotations

Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

UGT1A1	UDP glucuronosyltransferase 1 family, polypeptide A1

Class 3 rs8330 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

Class 3 rs1042640 GC

Patients with the CG genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk of liver failure due to unintentional acetaminophen overdose.

Class 3 rs10929303 TC

Patients with the CT genotype may have a decreased risk of liver failure due to unintentional acetaminophen overdose as compared to patients with the CC genotype. Other genetic and clinical feature may also influence risk of liver failure due to unintentional acetaminophen avardees

#### Α

factors may also influend	ce risk of liver failure due to unintentional acetaminophen overdose.
AMLODIPINE	
CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
· Class 3 rs2740574 TT  Women with the TT genotype and hypertension may have an increased likelihood of reaching a target mean arterial pressure of ¡= 107 mm Hg when treated with amlodipine as compared to women with the CC genotype. No significant associations were seen when considering a target mean arterial pressure of ¡= 92 mm Hg, or when considering men or men and women together. Other genetic and clinical factors may also influence response to amlodipine.	

CYP3A	cytochrome P450, family 3, subfamily A
	- Clinical Annotations

Class 3 rs2740574

Women with the TT genotype and hypertension may have an increased likelihood of reaching a target mean arterial pressure of i=107 mm Hg when treated with amlodipine as compared to women with the CC genotype. No significant associations were seen when considering a target mean arterial pressure of i=92 mm Hg, or when considering men or men and women together. Other genetic and clinical factors may also influence response to amlodipine.

· Class 3 rs776746 CC

Healthy males with the CC (CYP3A5 \*3/\*3) genotype may have increased metabolism of amlodipine as compared to healthy males with the CT or TT (\*3/\*1 or \*1/\*1) genotype. No significant associations were seen when considering clearance of amlodipine. Other genetic and clinical factors may also influence metabolism of amlodipine.

CYP3A5	cytochrome P450, family 3, subfamily A, polypeptide 5

· Class 3 rs776746 CC

Healthy males with the CC (CYP3A5 \*3/\*3) genotype may have increased metabolism of amlodipine as compared to healthy males with the CT or TT (\*3/\*1 or \*1/\*1) genotype. No significant associations were seen when considering clearance of amlodipine. Other genetic and clinical factors may also influence metabolism of amlodipine.

ZSCAN25	zinc finger and SCAN domain containing 25
	ations —

· Class 3 rs776746 CC

Healthy males with the CC (CYP3A5 \*3/\*3) genotype may have increased metabolism of amlodipine as compared to healthy males with the CT or TT (\*3/\*1 or \*1/\*1) genotype. No significant associations were seen when considering clearance of amlodipine. Other genetic and clinical factors may also influence metabolism of amlodipine.

#### ANASTROZOLE

UGT1A10	UDP glucuronosyltransferase 1 family, polypeptide A10

· Class 4 rs3732219 CC

Patients with the CC genotype may have increased glucuronidation of anastrozole as compared to patients with the CT or TT genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

 $\cdot$  Class 4 rs3732218 GG

Patients with the GG genotype may have increased glucuronidation of anastrozole as compared to patients with the AA or AG genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

otype may have increased glucuronidation of anastrozole as compared to T genotype, as determined by in vitro assays. Glucuronidation allows for tics like anastrozole. Other genetic and clinical factors may also influence to the cole.  G otype may have increased glucuronidation of anastrozole as compared to G genotype, as determined by in vitro assays. Glucuronidation allows for tics like anastrozole. Other genetic and clinical factors may also influence to the cole.
UDP glucuronosyltransferase 1 family, polypeptide A5
- Clinical Annotations
otype may have increased glucuronidation of anastrozole as compared to $T$ genotype, as determined by in vitro assays. Glucuronidation allows for tics like anastrozole. Other genetic and clinical factors may also influence to $G$ otype may have increased glucuronidation of anastrozole as compared to
G genotype, as determined by in vitro assays. Glucuronidation allows for tics like anastrozole. Other genetic and clinical factors may also influence cole.
UDP glucuronosyltransferase 1 family, polypeptide A6

· Class 4 rs3732219 CC

Patients with the CC genotype may have increased glucuronidation of anastrozole as compared to patients with the CT or TT genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

 $\cdot$  Class 4 rs3732218 GG

Patients with the GG genotype may have increased glucuronidation of anastrozole as compared to patients with the AA or AG genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

Patients with the CC genotype may have increased glucuronidation of anastrozole as compared to patients with the CT or TT genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

#### Class 4 rs3732218 *GG*

Patients with the GG genotype may have increased glucuronidation of anastrozole as compared to patients with the AA or AG genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

UGT1A8	UDP glucuronosyltransferase 1 family, polypeptide A8
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#### · Class 4 rs3732219 CC

Patients with the CC genotype may have increased glucuronidation of anastrozole as compared to patients with the CT or TT genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

#### $\cdot$ Class 4 rs3732218 GG

Patients with the GG genotype may have increased glucuronidation of anastrozole as compared to patients with the AA or AG genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

$\mathbf{UGT1A9}$	UDP glucuronosyltransferase 1 family, polypeptide A9

#### · Class 4 rs3732219 CC

Patients with the CC genotype may have increased glucuronidation of anastrozole as compared to patients with the CT or TT genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

#### $\cdot$ Class 4 rs3732218 GG

Patients with the GG genotype may have increased glucuronidation of anastrozole as compared to patients with the AA or AG genotype, as determined by in vitro assays. Glucuronidation allows for the elimination of xenobiotics like anastrozole. Other genetic and clinical factors may also influence glucuronidation of anastrozole.

#### ANTHRACYCLINES AND RELATED SUBSTANCES

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	Clinical Annotations

 $\cdot$  Class 3 rs1045642 GG

Patients with the GG genotype may have 1) decreased exposure to doxorubicin metabolites and 2) decreased response to anthracycline regimens as compared to patients with the AA genotype, however the evidence is highly contradictory. Other genetic and clinical factors may also influence response to anthracycline regimens.

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UDP glucuronosyltransferase 1 family, polypeptide A10

- Clinical Annotations —

#### Class 3 rs6759892 TG

Patients with the GT genotype may have increased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype TT and decreased likelihood as compared to patients with the GG genotype, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

#### · Class 3 rs17863783 GG

Patients with the GG genotype may have decreased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype GT or TT, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

UGT1A6

UDP glucuronosyltransferase 1 family, polypeptide A6

—- Clinical Annotations ————————

#### · Class 3 rs6759892 TG

Patients with the GT genotype may have increased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype TT and decreased likelihood as compared to patients with the GG genotype, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

#### · Class 3 rs17863783 GG

Patients with the GG genotype may have decreased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype GT or TT, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

UGT1A7

UDP glucuronosyltransferase 1 family, polypeptide A7

—- Clinical Annotations

#### $\cdot$ Class 3 rs6759892 TG

Patients with the GT genotype may have increased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype TT and decreased likelihood as compared to patients with the GG genotype, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

#### · Class 3 rs17863783 GG

Patients with the GG genotype may have decreased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype GT or TT, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

#### UGT1A8

UDP glucuronosyltransferase 1 family, polypeptide A8

-- Clinical Annotations -

#### Class 3 rs6759892 TG

Patients with the GT genotype may have increased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype TT and decreased likelihood as compared to patients with the GG genotype, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

#### · Class 3 rs17863783 GG

Patients with the GG genotype may have decreased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype GT or TT, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

UGT1A9

UDP glucuronosyltransferase 1 family, polypeptide A9

-- Clinical Annotations

#### · Class 3 rs6759892 TG

Patients with the GT genotype may have increased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype TT and decreased likelihood as compared to patients with the GG genotype, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

#### · Class 3 rs17863783 GG

Patients with the GG genotype may have decreased likelihood of cardiotoxicity when exposed to anthracyclines and related substances in children with Neoplasms as compared to patients with genotype GT or TT, although this has been contradicted in one study. Other genetic and clinical factors may also influence the risk of toxicity to anthracyclines and related substances.

#### ANTIEPILEPTICS

# ABCB1 ATP-binding cassette, sub-family B (MDR/TAP), member 1 - Clinical Annotations

#### · Class 3 rs1128503 GG

Patients with the GG genotype and specifically localization-related epilepsy syndrome may have a decreased risk for resistance to antiepileptic treatment as compared to patients with the AA genotype. However, all other studies of people with epilepsy have found no association between this variant and antiepileptic resistance. Other genetic and clinical factors may also influence resistance to antiepileptics.

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sodium channel, voltage-gated, type I, alpha subunit

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#### · Class 2B rs3812718 CT

Patients with the CT genotype and epilepsy may be less likely to be resistant to antiepileptic treatment, particularly carbamazepine, as compared to patients with the TT genotype. Other genetic and clinical factors may also influence resistance to antiepileptic drugs.

#### ANTIPSYCHOTICS

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#### · Class 3 rs4680 GA

Patients with the AG genotype may have increased blood pressure when treated with antipsychotics as compared to patients with the GG genotype. Other genetic and clinical factors may also influence blood pressure in patients receiving antipsychotics.

· Class 3 rs4680 GA

Patients with the AG genotype may have increased fasting glucose levels when treated with antipsychotics as compared to patients with the GG genotype. Other genetic and clinical factors may also influence fasting glucose in patients taking antipsychotics.

ABCB1 ATP-binding cassette, sub-family B (MDR/TAP), member 1

- Clinical Annotations

#### · Class 3 rs2032582 CC

Patients with the CC genotype and schizophrenia who responded to treatment with antipsychotics may require an increased dose of antipsychotics as compared to patients with the AA genotype. Other genetic and clinical factors may also influence dose of antipsychotics.

 $\cdot$  Class 3 rs1045642 GG

Patients with the GG genotype and schizophrenia who responded to treatment with antipsychotics may require an increased dose of antipsychotics as compared to patients with the AA genotype. Other genetic and clinical factors may also influence dose of antipsychotics.

#### **ATAZANAVIR**

UGT1A	UDP glucuronosyltransferase 1 family, polypeptide A complex locus

#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

#### Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

#### · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A10

UDP glucuronosyltransferase 1 family, polypeptide A10

— Clinical Annotations	
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#### $\cdot$ Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

#### Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

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UGT1A3	UDP glucuronosyltransferase 1 family, polypeptide A3
	- Clinical Annotations

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Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

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Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

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#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

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Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

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Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A5	UDP glucuronosyltransferase 1 family, polypeptide $A5$

#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

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Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

#### · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A6	UDP glucuronosyltransferase 1 family, polypeptide A6

#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

#### Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

#### · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A7

UDP glucuronosyltransferase 1 family, polypeptide A7

— Clinical Annotations	

#### $\cdot$ Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

#### Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

#### · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGTIA8	UDP glucuronosyltransferase I family, polypeptide A8

#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

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Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

#### · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

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#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

· Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

· Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A1	UDP glucuronosyltransferase 1 family, polypeptide A1

#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

· Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

· Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1
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#### · Class 3 rs1045642 GG

Patients with the GG genotype and HIV may have increased concentrations of atazanavir as compared to patients with the AA genotypes, although this is contradicted in one study. There is no evidence that the GG genotype is associated with hyperbilirubinemia, drug discontinuation, or nephrolithiasis. Other clinical and genetic factors may also influence the concentrations of atazanavir in patients with HIV.

ADRB2	adrenoceptor beta 2, surface
	- Clinical Annotations
triglyceridemia when tr	GG genotype and hypertension may have an increased risk of developing hyperceated with atenolol or metoprolol as compared to patients with the CC or CG c and clinical factors may also influence risk of hypertriglyceridemia.
TORVASTATIN	
ABCC2	ATP-binding cassette, sub-family C (CFTR/MRP), member 2 $$
	- Clinical Annotations —
	CT enotype may have decreased dose of simvastatin and atorvastatin as compared to CC. Other genetic and clinical factors may also influence the dose of simvastatin.
CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
	- Clinical Annotations —
ent drug when treated	TT genotype may be more likely to require a decrease in dose or switch to a differ-with atorvastatin or simvastatin as compared to patients with the CC or CT and clinical factors may also influence dose of simvastatin or atorvastatin, or to a different drug.
CYP3A	cytochrome P450, family 3, subfamily A
	- Clinical Annotations
ent drug when treated	TT genotype may be more likely to require a decrease in dose or switch to a differwith atorvastatin or simvastatin as compared to patients with the CC or CT c and clinical factors may also influence dose of simvastatin or atorvastatin, or to a different drug.
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	- Clinical Annotations

· Class 3 rs2032582 CC

Patients with the CC genotype may have increased risk of drug-induced liver injury compared to patients with the TT genotype. Other factors may affect liver toxicity when treated with atorvastatin.

ADRB2	adrenoceptor beta 2, surface
	nical Annotations ————————————————————————————————————
when treated with benazepril as compared to been seen for systolic blood pressure. Additi	asion may have a greater decrease in diastolic blood pressure o patients with the AA genotype. No significant results have onally, the same study reported no significant differences in genotypes in a different cohort. Other genetic and clinical ic or systolic blood pressure.
BEVACIZUMAB	
VEGFA	vascular endothelial growth factor A
	nical Annotations —
VEGF treatment, as compared to patients may also influence response to anti-VEGF to Class 3 rs699947 AC  Patients with colorectal cancer and the AC capecitabine, fluorouracil, irinotecan, leucon	dal neovascularization may have a better response to anti- with the CC genotype. Other genetic and clinical factors creatment.  C genotype may have a reduced response to bevacizumab, vorin, or oxaliplatin as compared to patients with the CC is may also affect response to chemotherapy in people with
BUPROPION	

#### CYP2B6

cytochrome P450, family 2, subfamily B, polypeptide 6

#### · Class 3 rs3211371 CC

Patients with the CC genotype who are smokers may have a lower chance of smoking cessation when treated with bupropion as compared to patients with the CT or TT genotype, although this is contradicted in one study. Other genetic and clinical factors may also influence likelihood of smoking cessation.

#### · Class 3 rs2279343 AA

Individuals with tobacco use disorder and the AA genotype may have an improved response to bupropion as compared to individuals with the AG and GG genotypes. Other clinical and genetic factors may also affect response to bupropion in individuals with tobacco use disorder.

#### **BUSULFAN**

CYP2C19	cytochrome P450, family 2, subfamily C, polypeptide 19
metabolism of busulfan	genotype (CYP2C19 *1/*1) undergoing transplantation may have decreased as compared to patients with the CT (*1/*17) or TT (*17/*17) genotype. Interpretation exists for this association. Other genetic and clinical factors
CYP2C9	cytochrome P450, family 2, subfamily C, polypeptide 9
	- Clinical Annotations
Patients with the CC (have increased metaboli	CYP2C9 *1/*1) genotype undergoing hemopoietic stem cell transplant may sm of busulfan as compared to patients with the CT (*1/*2) or TT (*2/*2) and clinical factors may also influence metabolism of busulfan.
CAPECITABINE	
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
in people with Colorecta are not associated with of epirubicin and gemcitab genetic and clinical factors.  Class 3 rs2032582  Patients with genotype Colorect	the may have increased risk of hand-foot syndrome when treated with capecitabined Neoplasms as compared to patients with genotype AA. Genotypes AG + GG decreased clinical outcome when treated with capecitabine, cisplatin, docetaxel, ine in people with Pancreatic Neoplasms as compared to genotype AA. Other for may influence the response to capecitabine.  CC  CC may have increased risk of hand-foot syndrome when treated with capecitabine al Neoplasms as compared to patients with genotype AA. Other genetic and
Class 3 rs1128503 Patients with the GG g hand-foot syndrome wh	influence the response to capecitabine. $GG$ genotype and colorectal cancer may have an increased risk of neutropenia or en treated with capecitabine as compared to patients with the AA genotype. al factors may also influence risk of neutropenia or hand-foot syndrome.

### · Class 3 rs602950 AA

CDA

Cancer patients with the AA genotype may have a decreased risk of diarrhea or dehydration when treated with capecitabine-based therapy as compared to patients with the AG or GG genotype. Other genetic and clinical factors may also influence risk of diarrhea and dehydration.

—- Clinical Annotations —

cytidine deaminase

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DPYD:*1/*1 Moderate	
Use label-recommended dosage and administration.	

#### Class 1A rs55886062 AA

Patients with the AA genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have a decreased, but not absent, risk for drug toxicity as compared to patients with the AC or CC genotype (DPYD \*1/\*13 or \*13/\*13). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

#### · Class 1A rs3918290 CC

Patients with the CC genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased clearance of fluoropyrimidine drugs and 2) decreased, but not non-existent, risk for drug toxicity as compared to patients with the CT or TT genotype (DPYD \*1/\*2A or \*2A/\*2A). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine based chemotherapy.

#### $\cdot$ Class 1A rs67376798 TT

Patients with the TT genotype and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased clearance of the drug and 2) decreased, but not absent, risk and reduced severity of drug toxicity as compared to patients with the AT genotype. Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

#### · Class 3 rs1801160 CC

Patients with the CC genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased metabolism of fluorouracil and 2) decreased risk for drug toxicities as compared to patients with the CT or TT genotype (DPYD \*1/\*6 or \*6/\*6). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin) or with other drugs such as bevacizumab, cetuximab, raltitrexed. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

#### $\cdot$ Class 3 rs1801159 TT

Patients with the TT genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) a decreased likelihood of nausea, vomiting, and leukopenia, 2) increased response and 3) increased clearance of fluorouracil as compared to patients with the CT or CC genotype (DPYD \*1/\*5 or \*5/\*5). However, other studies find no associations or contradictory associations with fluoropyrimidine-induced drug toxicity or response. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

Class 3	rs1801158	CC
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Patients with the CC genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) a decreased but not absent risk of toxicity and 2) increased DPYD activity as compared to patients with the CT genotype (DPYD \*1/\*4). However, some studies find no association with drug toxicity or DPYD activity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

VEGFA	vascular endothelial growth factor A
- Clinical Annotations -	

· Class 3 rs699947 AC

Patients with colorectal cancer and the AC genotype may have a reduced response to bevacizumab, capecitabine, fluorouracil, irinotecan, leucovorin, or oxaliplatin as compared to patients with the CC genotype. Other clinical and genetic factors may also affect response to chemotherapy in people with colorectal cancer.

· Class 3 rs2010963 CG

Patients with the CG genotype and colorectal cancer may have a poorer response when treated with capecitabine and oxaliplatin (XELOX) as compared to patients with the CC or GG genotype. Other genetic and clinical factors may also influence response to XELOX treatment.

#### CARBAMAZEPINE

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#### · Class 2B rs3812718 CT

Patients with the CT genotype who are treated with carbamazepine may require a higher dose as compared to patients with the CC genotype. Other genetic and clinical factors may also influence dose of carbamazepine.

 $\cdot$  Class 2B rs3812718 CT

Patients with the CT genotype and epilepsy may be less likely to be resistant to antiepileptic treatment, particularly carbamazepine, as compared to patients with the TT genotype. Other genetic and clinical factors may also influence resistance to antiepileptic drugs.

· Class 3 rs3812718 CT

Patients with epilepsy and the CT genotype may have decreased metabolism of carbamazepine, resulting in increased exposure as compared to patients with the TT genotype.

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1

#### $\cdot$ Class 3 rs1045642 GG

Patient with genotype GG may have decreased likelihood of drug resistance when treated with antiepileptics and carbamazepine in people with Epilepsy as compared to patients with genotype AA. However, contradictory findings have been reported. Other genetic and clinical factors may also influence response to carbamazepine.

 $\cdot$  Class 3 rs1128503 GG

African American and white patients with the GG genotype and epilepsy may have decreased clearance of carbamazepine as compared to patients with the AA or AG genotype. This association was not found in Chinese patients. Other genetic and clinical factors may also influence clearance of carbamazepine.

Class 3 rs1045642 GG

Patients with the GG genotype and epilepsy may have decreased metabolism of carbamazepine and may need a decreased dose as compared to patients with the AG genotype. However, multiple studies have shown no association with dose or concentrations of carbamazepine. Other genetic and clinical factors may also influence concentrations of carbamazepine.

ABCC2	ATP-binding cassette, sub-family C (CFTR/MRP), member 2
	—- Clinical Annotations ————————————————————————————————————

 $\cdot$  Class 3 rs3740066 CT

Patients with the CT genotype may have decreased metabolism of carbamazepine in men with Epilepsy as compared to patients with genotype CC. This association was only significant in male patients. Other genetic and clinical factors may also influence the metabolism of carbamazepine.

CYP3A5	cytochrome P450, family 3, subfamily A, polypeptide 5	

Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) and epilepsy may have decreased clearance and increased concentrations of carbamazepine, and require lower doses of the drug, as compared to patients with the CT (\*1/\*3) or TT (\*1/\*1) genotype. Other genetic and clinical factors may also influence dose or concentrations of carbamazepine.

· Class 3 rs15524 AA

Patients with the AA genotype and epilepsy may have increased concentrations of carbamazepine compared to patients with the AG and GG genotypes when patients were also taking phenytoin or phenobarbital. Other clinical and genetic factors may affect concentrations of carbamazepine.

ZSCAN25	zinc finger and SCAN domain containing $25$
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Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) and epilepsy may have decreased clearance and increased concentrations of carbamazepine, and require lower doses of the drug, as compared to patients with the CT (\*1/\*3) or TT (\*1/\*1) genotype. Other genetic and clinical factors may also influence dose or concentrations of carbamazepine.

· Class 3 rs15524 AA

Patients with the AA genotype and epilepsy may have increased concentrations of carbamazepine compared to patients with the AG and GG genotypes when patients were also taking phenytoin or phenobarbital. Other clinical and genetic factors may affect concentrations of carbamazepine.

	- Clinical Annotations	
	- Clinical Annotations	
	Class 3 rs776746 CC	
	Patients with the CC genotype (CYP3A5 *3/*3) and epilepsy may have decreased clearance and increased concentrations of carbamazepine, and require lower doses of the drug, as compared to patients with the CT (*1/*3) or TT (*1/*1) genotype. Other genetic and clinical factors may also influence dose or concentrations of carbamazepine.	
	Class 3 rs2740574 TT	
	Patients with the TT genotype and epilepsy may have increased clearance of carbamazepine as compared to patients with the CC or CT genotype. Other genetic and clinical factors may also influence clearance of carbamazepine.	
	CYP3A4 cytochrome P450, family 3, subfamily A, polypeptide 4	
	Class 3 rs2242480 <i>CC</i> Patients with the CC genotype (CYP3A4 *1/*1) and epilepsy may have increased concentrations of carbamazepine as compared to patients with the CT (*1/*1G) or TT (*1G/*1G) genotype. However,	
	studies conflict. Other genetic and clinical factors may also influence concentrations of carbamazepine. Class 3 $rs2740574$ $TT$	
	Patients with the TT genotype and epilepsy may have increased clearance of carbamazepine as compated to patients with the CC or CT genotype. Other genetic and clinical factors may also influence clearated of carbamazepine.	
	CYP1A2 cytochrome P450, family 1, subfamily A, polypeptide 2	
•	Class 3 rs762551 AA  Pediatric patients with epilepsy and the AA genotype may have increased clearance of carbamazepine as compared to pediatric patients with epilepsy and the AC or CC genotypes. Other clinical and genetic factors may also influence clearance of carbamazepine in pediatric patients with epilepsy.	

#### **CARVEDILOL**

UGT1A10	UDP glucuronosyltransferase 1 family, polypeptide A10 $$

#### Class 3 rs4148323 GG

Patients with the GG (i.e. UGT1A1 \*1/\*1) genotype and angina or heart failure may have increased glucuronidation of carvedilol as compared to patients with the AA (\*6/\*6) genotype. UGT1A1 is responsible for the glucuronidation of target substrates, rendering them water soluble and allowing for their biliary or renal elimination. Other genetic and clinical factors may also influence metabolism of carvedilol.

responsible for the glucuron	ol as compared to patients with the AA (*6/*6) genotype. UGT1A1 is idation of target substrates, rendering them water soluble and allowing for ation. Other genetic and clinical factors may also influence metabolism of
UGT1A4	UDP glucuronosyltransferase 1 family, polypeptide A4
	- Clinical Annotations
glucuronidation of carvedile responsible for the glucuron	UGT1A1 *1/*1) genotype and angina or heart failure may have increased of as compared to patients with the AA (*6/*6) genotype. UGT1A1 is idation of target substrates, rendering them water soluble and allowing for ation. Other genetic and clinical factors may also influence metabolism of
UGT1A5	UDP glucuronosyltransferase 1 family, polypeptide A5
glucuronidation of carvedile responsible for the glucuron	UGT1A1 *1/*1) genotype and angina or heart failure may have increased of as compared to patients with the AA (*6/*6) genotype. UGT1A1 is idation of target substrates, rendering them water soluble and allowing for ation. Other genetic and clinical factors may also influence metabolism of UDP glucuronosyltransferase 1 family, polypeptide A6
glucuronidation of carvedile responsible for the glucuron	UGT1A1 *1/*1) genotype and angina or heart failure may have increased of as compared to patients with the AA (*6/*6) genotype. UGT1A1 is idation of target substrates, rendering them water soluble and allowing for ation. Other genetic and clinical factors may also influence metabolism of
UGT1A7	UDP glucuronosyltransferase 1 family, polypeptide A7
	- Clinical Annotations —
`	GUGT1A1 *1/*1) genotype and angina or heart failure may have increased of as compared to patients with the AA (*6/*6) genotype. UGT1A1 is

responsible for the glucuronidation of target substrates, rendering them water soluble and allowing for

- Clinical Annotations –

Patients with the GG (i.e. UGT1A1  $^*1/^*1$ ) genotype and angina or heart failure may have increased

**Class 3** rs4148323 *GG* 

their biliary or renal elimination. Other genetic and clinical factors may also influence metabolism of carvedilol. UGT1A8 UDP glucuronosyltransferase 1 family, polypeptide A8 -- Clinical Annotations -Class 3 rs4148323 Patients with the GG (i.e. UGT1A1 \*1/\*1) genotype and angina or heart failure may have increased glucuronidation of carvedilol as compared to patients with the AA (\*6/\*6) genotype. UGT1A1 is responsible for the glucuronidation of target substrates, rendering them water soluble and allowing for their biliary or renal elimination. Other genetic and clinical factors may also influence metabolism of carvedilol. UGT1A9 UDP glucuronosyltransferase 1 family, polypeptide A9 - Clinical Annotations Class 3 rs4148323 GG Patients with the GG (i.e. UGT1A1 \*1/\*1) genotype and angina or heart failure may have increased glucuronidation of carvedilol as compared to patients with the AA (\*6/\*6) genotype. UGT1A1 is responsible for the glucuronidation of target substrates, rendering them water soluble and allowing for their biliary or renal elimination. Other genetic and clinical factors may also influence metabolism of carvedilol. UGT1A1 UDP glucuronosyltransferase 1 family, polypeptide A1 — - Clinical Annotations -Class 3 rs4148323 GG Patients with the GG (i.e. UGT1A1 \*1/\*1) genotype and angina or heart failure may have increased glucuronidation of carvedilol as compared to patients with the AA (\*6/\*6) genotype. UGT1A1 is responsible for the glucuronidation of target substrates, rendering them water soluble and allowing for their biliary or renal elimination. Other genetic and clinical factors may also influence metabolism of carvedilol. **CATECHOLAMINES** ADRB1 adrenoceptor beta 1 - Clinical Annotations -Class 3 rs1801253 GG

Patients with the GG genotype and coronary artery disease may require an increased dose of catecholamines as compared to patients with the CC or CG genotype. Other genetic and clinical factors may also influence required dose of catecholamines.

#### **CELECOXIB**

CYP2C9	cytochrome P450, family 2, subfamily C, polypeptide 9
	- Clinical Annotations —
to patients with the AA ( $CC (*3/*3)$ genotype. Ot	AC P2C9 *1/*3) genotype may have reduced metabolism of celecoxib as compared *1/*1) genotype, and increased metabolism as compared to patients with the ther genetic and clinical factors may also influence metabolism of celecoxib.
LOPIDOGREL	
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
events (MACE such as ca dogrel in people with acu genotypes AA. Contradic	e may have decreased, but not absent, risk of major adverse cardiovascular rdiovascular death, myocardial infarction, or stroke) when treated with clopiate coronary syndrome or myocardial Infarction as compared to people with tory findings have been reported in the literature. Other genetic and clinical the response to clopidogrel.
MED12L	mediator complex subunit 12-like
_	notype may have increased risk of adverse cardiac events when treated with to patients with genotype GG. Other genetic and clinical factors may also
P2RY12	purinergic receptor P2Y, G-protein coupled, 12
_	AA notype may have increased risk of adverse cardiac events when treated with to patients with genotype GG. Other genetic and clinical factors may also
Patients with the AA ger clopidogrel as compared	AA notype may have increased risk of adverse cardiac events when treated with to patients with genotype $GG$ . Other genetic and clinical factors may also
Patients with the AA ger clopidogrel as compared influence the response to	AA notype may have increased risk of adverse cardiac events when treated with to patients with genotype $GG$ . Other genetic and clinical factors may also

Patients with the AG genotype and schizophrenia may have a poorer response when treated with clozapine as compared to patients with the GG genotype. Other genetic and clinical factors may also influence response to clozapine.

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member $1$
	— Clinical Annotations —

#### Class 3 rs1045642 GG

Patients with the GG genotype may have decreased clozapine plasma concentrations, as well as a decreased risk for clozapine-induced agranulocytosis or neutropenia, as compared to patients with the AA genotype. Other genetic and clinical factors may also influence concentrations and risk of clozapine-induced toxicity.

#### CODEINE

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member $1$

#### · Class 3 rs1128503 GG

Breast-feeding infants whose mothers have the GG genotype and are taking codeine may be at decreased risk for CNS depression as compared to those whose mothers have the AA genotype. Other genetic and clinical factors may also influence the risk of CNS depression in breast-feeding infants.

#### **COTININE**

CHRNA3	cholinergic receptor, nicotinic, alpha 3 (neuronal)
	nical Annotations —

#### · Class 3 rs16969968 GG

Individuals with Tobacco Use Disorder and the GG genotype may have decreased concentrations of cotinine, a metabolite of nicotine, as compared to individuals with the AG or AA genotype. Other clinical and genetic factors may also contribute to cotinine concentrations in individuals with Tobacco Use Disorder.

CHRNA5	cholinergic receptor, nicotinic, alpha 5 (neuronal)

#### · Class 3 rs16969968 GG

Individuals with Tobacco Use Disorder and the GG genotype may have decreased concentrations of cotinine, a metabolite of nicotine, as compared to individuals with the AG or AA genotype. Other clinical and genetic factors may also contribute to cotinine concentrations in individuals with Tobacco Use Disorder.

#### **COUMARIN**

CYP2A6	cytochrome P450, family 2, subfamily A, polypeptide 6
Patients with the AT geno	Totype may have increased 7-hydroxylation of coumarin compared to patients ther genetic and clinical factors may also influence metabolism of coumarin.
YCLOPHOSPHAMIDE	
CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
	- Clinical Annotations
Clara 9 9740474 //	
Premenopausal patients w phamide may have a short	with the TT genotype and breast cancer who are treated with cyclophoster period of time before chemotherapy-induced ovarian failure compared to CT genotype. Other genetic and clinical factors may also influence time to trian failure.
CYP3A	cytochrome P450, family 3, subfamily A
Premenopausal patients we phamide may have a short	with the TT genotype and breast cancer who are treated with cyclophoster period of time before chemotherapy-induced ovarian failure compared to CT genotype. Other genetic and clinical factors may also influence time to trian failure.
VEGFA	vascular endothelial growth factor A
Class 3 rs2010963 (	CG
Patients with the CG ger when treated with cyclopl and clinical factors may al	notype and prostate cancer may have longer progression-free survival time nosphamide as compared to patients with the CC genotype. Other genetic so influence length of progression-free survival.
Patients with the AG ger when treated with docetax	action of the control
CYP1B1	cytochrome P450, family 1, subfamily B, polypeptide 1

Class 3	rs1056836	GG
Class 6	10100000	$\alpha \alpha$

Patients with the GG genotype and breast cancer may have a better response when treated with cyclophosphamide, epirubicin and fluorouracil as compared to patients with the CC genotype. Other genetic and clinical factors may also influence response to treatment with cyclophosphamide, epirubicin and fluorouracil. (Note: with a  $\rm C/G$  variant, particularly in a gene on the minus chromosomal strand, and frequencies close to 50

CYP2B6	cytochrome P450, family 2, subfamily B, polypeptide 6
	- Clinical Annotations

#### · Class 3 rs4802101 TT

Patients with the TT genotype may have increased metabolism of cyclophosphamide, resulting in increased concentrations of active cyclophosphamide metabolites, and increased risk of gastrointestinal toxicity, or leukopenia, as compared to patients with the CT or CC genotypes. Other clinical and genetic factors may also influence metabolism of cyclophosphamide, as well as risk of toxicity in patients with lupus.

#### **CYCLOSPORINE**

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$

#### · Class 3 rs1128503 GG

Patients with the GG genotype and myasthenia gravis or organ transplantation may have increased clearance of cyclosporine and therefore may require an increased dose of cyclosporine, compared to patients with the AA genotype. Patients with the GG genotype may also have a decreased risk of infection as compared to those with the AA or AG genotype. Other genetic and clinical factors may also influence clearance and dose of cyclosporine.

#### · Class 3 rs2032582 CC

Patients with the CC genotype may have lower blood trough concentrations of cyclosporine compared to patients with the AA genotype, and may require dose adjustments. Other genetic and clinical factors may also influence cyclosporine blood concentrations.

#### $\cdot$ Class 3 rs1045642 GG

Patients with genotype GG may have decreased intracellular and blood concentrations of cyclosporine in people with Transplantation as compared to patients with genotype AA or AG. However, contradictory findings have been reported. Other genetic and clinical factors may also influence the concentration of cyclosporine.

#### $\cdot$ Class 3 rs2032582 CC

Patients with the CC genotype and cystic fibrosis may have increased clearance of dicloxacillin, when it is coadministered with cyclosporine, as compared to patients with the AA genotype. Other genetic and clinical factors may also influence clearance of dicloxacillin.

CYP3A5	cytochrome P450, family 3, subfamily A, polypeptide 5
Class 2B rs776746	CC

Patients with the CC genotype (CYP3A5 \*3/\*3) may require a lower dose of cyclosporine to reach target blood concentration as compared to patients with the CT (CYP3A5 \*1/\*3) or TT (CYP3A5 \*1/\*1) genotype, although this is contradicted in some studies. Other genetic and clinical factors may also influence dose of cyclosporine.

ZSC	Δ	N	25
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zinc finger and SCAN domain containing 25

-- Clinical Annotations -

#### · Class 2B rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) may require a lower dose of cyclosporine to reach target blood concentration as compared to patients with the CT (CYP3A5 \*1/\*3) or TT (CYP3A5 \*1/\*1) genotype, although this is contradicted in some studies. Other genetic and clinical factors may also influence dose of cyclosporine.

#### CYP3A

cytochrome P450, family 3, subfamily A

—- Clinical Annotations —

#### $\cdot$ Class 2B rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) may require a lower dose of cyclosporine to reach target blood concentration as compared to patients with the CT (CYP3A5 \*1/\*3) or TT (CYP3A5 \*1/\*1) genotype, although this is contradicted in some studies. Other genetic and clinical factors may also influence dose of cyclosporine.

· Class 3 rs35599367 GG

Patients with the GG genotype and organ transplantation administered cyclosporine may have a 1) increased metabolism of cyclosporine 2) increased clearance of cyclosporine and 3) a decreased risk in adverse events (e.g. graft rejection or kidney function) as compared to patients with the AA genotype. Other clinical and genetic factors may also affect metabolism and incidence of adverse events in organ transplant patients administered cyclosporine.

#### CYP3A4

cytochrome P450, family 3, subfamily A, polypeptide 4

—- Clinical Annotations ————

#### · Class 3 rs28371759 AA

Patients with the AA genotype (CYP3A4 \*1/\*1) who underwent kidney transplantation may have decreased metabolism of cyclosporine as compared to patients with the GG genotype (\*18B/\*18B). Other genetic and clinical factors may also influence metabolism of cyclosporine.

 $\cdot$  Class 3 rs35599367 GG

Patients with the GG genotype and organ transplantation administered cyclosporine may have a 1) increased metabolism of cyclosporine 2) increased clearance of cyclosporine and 3) a decreased risk in adverse events (e.g. graft rejection or kidney function) as compared to patients with the AA genotype. Other clinical and genetic factors may also affect metabolism and incidence of adverse events in organ transplant patients administered cyclosporine.

#### CYTARABINE

· Class 3 rs1045642 GC	Y T
decreased response to anthr	type may have 1) decreased exposure to doxorubicin metabolites and 2) acycline regimens as compared to patients with the AA genotype, however adictory. Other genetic and clinical factors may also influence response to
Patients with the GG genot with cytarabine, alone or in	ype and acute myeloid leukemia may have a poorer response when treated combination with daunorubicin, or dexrazoxane as compared to patients e, however some evidence contradicts this. Other genetic and clinical factors
SLCO1B1	solute carrier organic anion transporter family, member 1B1
with de novo acute myeloic mitoxantrone as compared	type may have more favorable event-free and overall survival in children leukemia (AML) treated with cytarabine, daunorubicin, etoposide and to patients with genotype CC. Other genetic and clinical factors may also come in acute myeloid leukemia.
AUNORUBICIN	
SLCO1B1	solute carrier organic anion transporter family, member 1B1
	- Clinical Annotations
· Class 3 rs2291075 CT	י
Patients with the CT geno with de novo acute myeloid	type may have more favorable event-free and overall survival in children d leukemia (AML) treated with cytarabine, daunorubicin, etoposide and to patients with genotype CC. Other genetic and clinical factors may also

# DEFERASIROX

UGT1A10

UDP glucuronosyltransferase 1 family, polypeptide A10

Clinical Annotations

## · Class 3 rs887829 CT

Patients with the CT genotype and beta-thal assemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

· Class 3 rs3806596 TC

Patients with the CT genotype and beta-thalassemia may have worse response to, and decreased concentrations of deferasirox as compared to patients with the CC genotype. Other clinical and genetic factors may also influence concentrations of and response to deferasirox in patients with beta-thalassemia.

#### $\cdot$ Class 4 rs4124874 TG

Pediatric patients with major thalassemia and the GT genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the TT genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype GT, therefore there is no information for patients with the GG or TT genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

#### $\cdot$ Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

#### · Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

UGT1A3	UDP glucuronosyltransferase 1 family, polypeptide A3
	Clinical Annotations

#### · Class 3 rs887829 CT

Patients with the CT genotype and beta-thalassemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

#### · Class 3 rs3806596 TC

Patients with the CT genotype and beta-thalassemia may have worse response to, and decreased concentrations of deferasirox as compared to patients with the CC genotype. Other clinical and genetic factors may also influence concentrations of and response to deferasirox in patients with beta-thalassemia.

#### $\cdot$ Class 4 rs4124874 TG

Pediatric patients with major thalassemia and the GT genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the TT genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype GT, therefore there is no information for patients with the GG or TT genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

#### $\cdot$ Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the

AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

· Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

UGT1A4	UDP glucuronosyltransferase 1 family, polypeptide A4
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· Class 3 rs887829 CT

Patients with the CT genotype and beta-thalassemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

· Class 3 rs3806596 TC

Patients with the CT genotype and beta-thalassemia may have worse response to, and decreased concentrations of deferasirox as compared to patients with the CC genotype. Other clinical and genetic factors may also influence concentrations of and response to deferasirox in patients with beta-thalassemia.

· Class 4 rs4124874 TG

Pediatric patients with major thalassemia and the GT genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the TT genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype GT, therefore there is no information for patients with the GG or TT genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

· Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

· Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

UGT1A5	UDP glucuronosyltransferase 1 family, polypeptide A5
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Patients with the CT genotype and beta-thalassemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

# · Class 3 rs3806596 TC

Patients with the CT genotype and beta-thalassemia may have worse response to, and decreased concentrations of deferasirox as compared to patients with the CC genotype. Other clinical and genetic factors may also influence concentrations of and response to deferasirox in patients with beta-thalassemia.

# · Class 4 rs4124874 TG

Pediatric patients with major thalassemia and the GT genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the TT genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype GT, therefore there is no information for patients with the GG or TT genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

# · Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

## · Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

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UDP glucuronosyltransferase 1 family, polypeptide A6

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#### · Class 3 rs887829 CT

Patients with the CT genotype and beta-thalassemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

# · Class 3 rs3806596 TC

Patients with the CT genotype and beta-thalassemia may have worse response to, and decreased concentrations of deferasirox as compared to patients with the CC genotype. Other clinical and genetic factors may also influence concentrations of and response to deferasirox in patients with beta-thalassemia.

# · Class 4 rs4124874 TG

Pediatric patients with major thalassemia and the GT genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the TT genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype GT, therefore there is no information for patients with the GG or TT genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

#### $\cdot$ Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

# · Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

UGT1A7	UDP glucuronosyltransferase 1 family, polypeptide A7

# · Class 3 rs887829 CT

Patients with the CT genotype and beta-thalassemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

# · Class 3 rs3806596 TC

Patients with the CT genotype and beta-thalassemia may have worse response to, and decreased concentrations of deferasirox as compared to patients with the CC genotype. Other clinical and genetic factors may also influence concentrations of and response to deferasirox in patients with beta-thalassemia.

# $\cdot$ Class 4 rs4124874 TG

Pediatric patients with major thalassemia and the GT genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the TT genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype GT, therefore there is no information for patients with the GG or TT genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

#### · Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

# · Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

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# · Class 3 rs887829 CT

Patients with the CT genotype and beta-thalassemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

# $\cdot$ Class 3 rs3806596 TC

Patients with the CT genotype and beta-thalassemia may have worse response to, and decreased concentrations of deferasirox as compared to patients with the CC genotype. Other clinical and genetic factors may also influence concentrations of and response to deferasirox in patients with beta-thalassemia.

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# $\cdot$ Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

#### · Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

# UGT1A9

UDP glucuronosyltransferase 1 family, polypeptide A9

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# · Class 3 rs887829 CT

Patients with the CT genotype and beta-thalassemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

#### · Class 3 rs3806596 TC

Patients with the CT genotype and beta-thalassemia may have worse response to, and decreased concentrations of deferasirox as compared to patients with the CC genotype. Other clinical and genetic factors may also influence concentrations of and response to deferasirox in patients with beta-thalassemia.

# · Class 4 rs4124874 TG

Pediatric patients with major thalassemia and the GT genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the TT genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype GT, therefore there is no information for patients with the

GG or TT genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

# · Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

# · Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

UGT1A1	UDP glucuronosyltransferase 1 family, polypeptide A1
	- Clinical Annotations

## · Class 3 rs887829 CT

Patients with the CT genotype and beta-thalassemia may have decreased concentrations of deferasirox as compared to patients with the TT genotype. Other genetic and clinical factors may also influence concentrations of deferasirox.

# · Class 4 rs4124874 TG

Pediatric patients with major thalassemia and the GT genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the TT genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype GT, therefore there is no information for patients with the GG or TT genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

# Class 4 rs10929302 GA

Pediatric patients with major thalassemia and the AG genotype may have an increased risk of adverse reactions when administered deferasirox as compared to patients with the GG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the AA or GG genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

#### · Class 4 rs4148323 GG

Pediatric patients with major thalassemia and the GG genotype may have a decreased risk of adverse reactions when administered deferasirox as compared to patients with the AA or AG genotype. Please note, the evidence comes solely from a single case study report of a single individual, a 3 year old female patient with major thalassemia of genotype AG, therefore there is no information for patients with the GG or AA genotypes. Other clinical and genetic factors may also influence risk of adverse reactions in patients with major thalassemia who are administered deferasirox.

Class 3 rs2273697 GG	
	pe and beta-thalassemia may have decreased concentrations of deferasire
	the AG genotype. Other genetic and clinical factors may also influence
concentrations of deferasirox.	
Class 4 rs717620 CT	
Pediatric patients with major reactions when administered the evidence comes solely fro thalassemia of genotype CT,	r thalassemia and the CT genotype may have an increased risk of adver deferasirox as compared to patients with the CC genotype. Please not om a single case study report of a 3 year old female patient with maje therefore there is no information for patients with the CC or TT genotype ctors may also influence risk of adverse reactions in patients with maje tered deferasirox.
CYP1A2	cytochrome P450, family 1, subfamily A, polypeptide 2
Clara 9 700FF1 4.4	
Class 3 rs762551 AA	
	pe and beta-thalassemia may have decreased concentrations of deferasire
Patients with the AA genoty	
Patients with the AA genoty	h the AC or CC genotype. Other genetic and clinical factors may also
Patients with the AA genoty as compared to patients wit influence concentrations of de	h the AC or CC genotype. Other genetic and clinical factors may also
Patients with the AA genoty as compared to patients wit influence concentrations of declass 3 rs2470890 TT Patients with the TT genotypossibly to levels below ther	h the AC or CC genotype. Other genetic and clinical factors may algebrasized and beta-thalassemia may have decreased concentrations of deferasiro
Patients with the AA genoty as compared to patients wit influence concentrations of declass 3 rs2470890 TT Patients with the TT genotypossibly to levels below ther	h the AC or CC genotype. Other genetic and clinical factors may also eferasirox.  Dee and beta-thalassemia may have decreased concentrations of deferasiro expeutic efficacy, as compared to patients with the CC or CT genotype.
Patients with the AA genoty as compared to patients wit influence concentrations of declared as a rs2470890 TT Patients with the TT genotypossibly to levels below ther Other genetic and clinical factors.	pe and beta-thalassemia may have decreased concentrations of deferasiro apeutic efficacy, as compared to patients with the CC or CT genotyp
Patients with the AA genoty as compared to patients wit influence concentrations of declass 3 rs2470890 TT  Patients with the TT genotypossibly to levels below ther Other genetic and clinical factures.  URETICS	h the AC or CC genotype. Other genetic and clinical factors may also eferasirox.  Dee and beta-thalassemia may have decreased concentrations of deferasiro rapeutic efficacy, as compared to patients with the CC or CT genotype ctors may also influence concentrations of deferasirox.
Patients with the AA genoty as compared to patients wit influence concentrations of declass 3 rs2470890 TT  Patients with the TT genotypossibly to levels below ther Other genetic and clinical factures  URETICS  ADRB2	h the AC or CC genotype. Other genetic and clinical factors may all eferasirox.  pe and beta-thalassemia may have decreased concentrations of deferasiro apeutic efficacy, as compared to patients with the CC or CT genotypetors may also influence concentrations of deferasirox.  adrenoceptor beta 2, surface
Patients with the AA genoty as compared to patients wit influence concentrations of declass 3 rs2470890 TT  Patients with the TT genotypossibly to levels below ther Other genetic and clinical factures.  URETICS	h the AC or CC genotype. Other genetic and clinical factors may all eferasirox.  pe and beta-thalassemia may have decreased concentrations of deferasiro apeutic efficacy, as compared to patients with the CC or CT genotypetors may also influence concentrations of deferasirox.  adrenoceptor beta 2, surface
Patients with the AA genoty as compared to patients wit influence concentrations of declass 3 rs2470890 TT  Patients with the TT genotypossibly to levels below ther Other genetic and clinical factorization.  URETICS  ADRB2  Class 3 rs1042713 GG  Patients with the GG genoty hospital utilization when tree	h the AC or CC genotype. Other genetic and clinical factors may algebrasized.  Dee and beta-thalassemia may have decreased concentrations of deferasiron apeutic efficacy, as compared to patients with the CC or CT genotypectors may also influence concentrations of deferasirox.  Adrenoceptor beta 2, surface demands and heart failure may have increased emergency department visits and the concentrations are perfectly and the concentrations.
Patients with the AA genoty as compared to patients wit influence concentrations of declass 3 rs2470890 TT  Patients with the TT genotypossibly to levels below ther Other genetic and clinical factorization.  URETICS  ADRB2  Class 3 rs1042713 GG  Patients with the GG genoty hospital utilization when tree	h the AC or CC genotype. Other genetic and clinical factors may also eferasirox.  pe and beta-thalassemia may have decreased concentrations of deferasiro rapeutic efficacy, as compared to patients with the CC or CT genotypetors may also influence concentrations of deferasirox.  adrenoceptor beta 2, surface  ———————————————————————————————————

· Class 3 rs1801253 GG

Patients with the CC genotype and heart failure may have increased emergency department utilization when treated with cardiovascular drugs as compared to patients with the GG genotype. Other genetic and clinical factors may also influence efficacy of cardiovascular drugs.

# **DOBUTAMINE**

adrenoceptor beta 1
G genotype may have smaller increases in fractional shortening and systolic dobutamine, as compared to healthy males with the CC genotype. No seen for heart rate. Other genetic and clinical factors may also influence systolic blood pressure.
cytochrome P450, family 3, subfamily A, polypeptide 4
type may have decreased clearance of docetaxel and a decreased risk of an accompared to patients with the CC or CT genotype. These patients may a of neurotoxicity with docetaxel treatment, though reports conflict. Other may also influence clearance of and reactions to docetaxel.  cytochrome P450, family 3, subfamily A
cytochrome 1 450, fainny 5, Sublainny A
- Clinical Annotations
type may have decreased clearance of docetaxel and a decreased risk of an accompared to patients with the CC or CT genotype. These patients may a of neurotoxicity with docetaxel treatment, though reports conflict. Other may also influence clearance of and reactions to docetaxel.  cytochrome P450, family 4, subfamily B, polypeptide 1
C
type may have a decreased but not absent risk of toxicity with docetaxel and patients with the CT or TT genotypes. Other genetic and clinical factors are response.
N-acetyltransferase 2 (arylamine N-acetyltransferase)

Class 3 rs1799931 *GG* 

Patients with the GG genotype may have an increased risk of toxicity with docetaxel and thalidomide as compared to patients with the AA genotype. Other genetic and clinical factors may also influence treatment response.

Clinical Approtations	

# · Class 3 rs1570360 AG

Patients with the AG genotype and breast cancer may have a better response to docetaxel treatment as compared to patients with the GG genotype. However, contradictory evidence exists when considering progression-free survival. Other genetic and clinical factors may also influence response to docetaxel.

· Class 3 rs699947 AC

Current literature evidence finds no significant effect of the AC genotype on progression-free survival time in patients taking docetaxel.

· Class 3 rs1570360 AG

Patients with the AG genotype and prostate cancer may have longer progression-free survival time when treated with docetaxel plus oral metronomic cyclophosphamide as compared to patients with the AA genotype. Other genetic and clinical factors may also influence progression-free survival time.

HNF4A	hepatocyte nuclear factor 4, alpha
	ations —

# · Class 3 rs2273618 TC

Patients with nasopharyngeal cancer and the CT genotype may have more severe anemia when treated with docetaxel as compared to patients with the TT genotype. Other clinical and genetic factors may also influence severity of anemia in patients with nasopharyngeal cancer who are treated with docetaxel.

· Class 3 rs3746574 TC

Patients with nasopharyngeal cancer and the CT genotype may have more severe anemia when treated with docetaxel as compared to patients with the TT genotype. Other clinical and genetic factors may also influence severity of anemia in patients with nasopharyngeal cancer who are treated with docetaxel.

#### DOXORUBICIN

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	- Clinical Annotations

# · Class 3 rs1045642 GG

Patients with the GG genotype may have 1) decreased exposure to doxorubicin metabolites and 2) decreased response to anthracycline regimens as compared to patients with the AA genotype, however the evidence is highly contradictory. Other genetic and clinical factors may also influence response to anthracycline regimens.

· Class 3 rs2032582 CC

Patients with the CC genotype may have increased metabolism of doxorubicin in people with Breast Neoplasms as compared to patients with genotype AA. Other genetic and clinical factors may also influence the metabolism of doxorubicin.

#### **EFAVIRENZ**

	$\alpha_1$	A
_ '	Ciinical	Annotations

# $\cdot$ Class 3 rs1045642 GG

Patients with the GG genotype and HIV infection who are treated with efavirenz may have reduced clearance of efavirenz as compared to patients with the AG genotype. Some studies have shown no association between this polymorphism and efavirenz clearance, plasma concentrations or exposure, or PBMC concentrations. Other genetic and clinical factors may also influence efavirenz pharmacokinetics.

# · Class 3 rs2032582 CC

Patients with the CC genotype may have increased likelihood of emerging viral drug resistance when exposed to efavirenz in people with HIV Infections as compared to patients with the AA genotype. This varaint is not associated with plasma exposure of efavirenz. Other genetic and clinical factors may also influence the response to efavirenz

# · Class 4 rs1128503 GG

Patients with GG genotype and HIV may have increased concentrations of efavirenz in plasma compared to patients with AA genotype. However, this association was not significant and was not found in another study of plasma and PBMCs. Other clinical and genetic factors may affect efavirenz concentrations.

CYP2A6	cytochrome P450, family 2, subfamily A, polypeptide 6
	- Cliffical Affilotations

# · Class 3 rs28399433 AA

Patients with the AA genotype and HIV may have decreased plasma concentrations of efavirenz as compared to patients with the AC or CC genotype. Other genetic and clinical factors may also influence plasma concentrations of efavirenz.

	v	D	$^{2}$	P	c
$\mathbf{C}$	1	1	4.	ப	U

cytochrome P450, family 2, subfamily B, polypeptide 6

# · Class 2B rs4803419 CC

Patients with HIV and the CC genotype may have lower plasma concentrations of efavirenz as compared to patients with the TT genotype. Other clinical and genetic factors may also influence plasma concentrations of efavirenz in patients with HIV.

# · Class 3 rs8192709 CC

Patients with genotype CC may have decreased metabolism of efavirenz in people with HIV Infections as compared to patients with genotype CT. Other genetic and clinical factors may also influence the metabolism of efavirenz.

#### · Class 3 rs8192719 CC

Patients with the CC genotype and HIV may have decreased concentrations of efavirenz as compared to patients with the CT or TT genotype. Other genetic and clinical factors, such as rs3745274, may also influence concentrations of efavirenz.

#### **ENALAPRIL**

Patients with the GG ger	cytochrome P450, family 2, subfamily D, polypeptide 6  — Clinical Annotations  GG  notype and depression may have a increased response and remission rate when as compared to patients with the AA genotype. Other genetic and clinical atients response.  N-acetyltransferase 2 (arylamine N-acetyltransferase)
Class 3 rs1065852 Patients with the GG get treated with escitaloprar factors may also effect page	GG notype and depression may have a increased response and remission rate when as compared to patients with the AA genotype. Other genetic and clinical
· Class 3 rs1065852 Patients with the GG generated with escitaloprary	——————————————————————————————————————
CYP2D6	
CYP2D6	cytochrome P450, family 2, subfamily D, polypeptide 6
_	AA notype may have increased concentrations of erlotinib as compared to patient of the genetic and clinical factors may also influence concentrations of erlotinib
	Clinical Annotations
CYP1A2	cytochrome P450, family 1, subfamily A, polypeptide 2
RLOTINIB	
	on and the AC genotype may have an improved response to enalapril as come CC genotype. Other clinical and genetic factors may also influence response
VEGFA	vascular endothelial growth factor A

when treated with anti-TB drugs as compared to patients with the CC genotype. Other genetic and

clinical factors may also influence risk for hepatotoxicity.

-- Clinical Annotations —

# Class 2A rs1799930

Patients with the AA genotype and tuberculosis (TB) may have an increased risk of hepatotoxicity when treated with anti-TB drugs as compared to patients with the GG genotype. They also may have decreased clearance of isoniazid as compared to those with the AG or GG genotype. Other genetic and clinical factors may also influence risk for hepatotoxicity and clearance of isoniazid.

Class 3 rs1799931 GG

Patients with the GG genotype and tuberculosis (TB) may have a decreased risk of hepatotoxicity when treated with anti-TB drugs as compared to patients with the AA or AG genotype. However, some studies find no association with hepatotoxicity. Other genetic and clinical factors may also influence risk of hepatotoxicity.

$\mathbf{F}$	TIT	A N	$\mathbf{n}$	T
$\Gamma_I$	п	$A \cap$	,,,	

ETHANOL	
CHRNA3	cholinergic receptor, nicotinic, alpha 3 (neuronal)
Patients with the GG gene	GG otype may have an increased risk for alcoholism as compared to patients with genetic and clinical factors may also influence risk of alcoholism.
CHRNA5	cholinergic receptor, nicotinic, alpha 5 (neuronal)
	- Clinical Annotations
Patients with the GG gene	GG otype may have an increased risk for alcoholism as compared to patients with genetic and clinical factors may also influence risk of alcoholism.
SLCO1B1	solute carrier organic anion transporter family, member 1B1
Patients with the CT ger with de novo acute myelo mitoxantrone as compared	notype may have more favorable event-free and overall survival in children oid leukemia (AML) treated with cytarabine, daunorubicin, etoposide and d to patients with genotype CC. Other genetic and clinical factors may also atcome in acute myeloid leukemia.
ABCB1	ATP-hinding cassette sub-family R (MDR/TAP) member 1

- Clinical Annotations -

Healthy individuals with t drug levels as compared w ciation with fexofenadine	the GG genotype who are treated with fexofenadine may have higher plasma ith healthy individuals with the AA genotype. Another study found no assoplasma concentrations. Other genetic and clinical factors may also influence exofenadine and dose requirements.
FLUOROURACIL	
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
Patients with GG genotyp	GG e may have decreased risk of diarrhea when treated with fluorouracil in people as compared to patients with genotype AA. Other genetic and clinical factors response to fluorouracil.
ABCC2	ATP-binding cassette, sub-family C (CFTR/MRP), member 2 $$
	- Clinical Annotations
treated with FOLFOX (f genotype. Other genetic a · Class 3 rs717620 C' Patients with the CT gen syndromes when treated v	otype and colon cancer may have a decreased risk of thrombocytopenia when duorouracil, leucovorin, oxaliplatin) as compared to patients with the CO and clinical factors may also influence risk of thrombocytopenia.
CYP1B1	cytochrome P450, family 1, subfamily B, polypeptide 1
	- Clinical Annotations —
	GG notype and breast cancer may have a better response when treated with

Patients with the GG genotype and breast cancer may have a better response when treated with cyclophosphamide, epirubicin and fluorouracil as compared to patients with the CC genotype. Other genetic and clinical factors may also influence response to treatment with cyclophosphamide, epirubicin and fluorouracil. (Note: with a C/G variant, particularly in a gene on the minus chromosomal strand, and frequencies close to 50

DPYD	dihydropyrimidine dehydrogenase
	sing Guideline ————————————————————————————————————

# **DPYD:\*1/\*1** Moderate

# Class 1A rs55886062 AA

Patients with the AA genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have a decreased, but not absent, risk for drug toxicity as compared to patients with the AC or CC genotype (DPYD \*1/\*13 or \*13/\*13). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

# · Class 1A rs3918290 CC

Patients with the CC genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased clearance of fluoropyrimidine drugs and 2) decreased, but not non-existent, risk for drug toxicity as compared to patients with the CT or TT genotype (DPYD \*1/\*2A or \*2A/\*2A). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine based chemotherapy.

# $\cdot$ Class 1A rs67376798 TT

Patients with the TT genotype and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased clearance of the drug and 2) decreased, but not absent, risk and reduced severity of drug toxicity as compared to patients with the AT genotype. Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

# · Class 3 rs1801160 CC

Patients with the CC genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased metabolism of fluorouracil and 2) decreased risk for drug toxicities as compared to patients with the CT or TT genotype (DPYD \*1/\*6 or \*6/\*6). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin) or with other drugs such as bevacizumab, cetuximab, raltitrexed. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

# · Class 3 rs1801159 TT

Patients with the TT genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) a decreased likelihood of nausea, vomiting, and leukopenia, 2) increased response and 3) increased clearance of fluorouracil as compared to patients with the CT or CC genotype (DPYD \*1/\*5 or \*5/\*5). However, other studies find no associations or contradictory associations with fluoropyrimidine-induced drug toxicity or response. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

# · Class 3 rs1801265 GA

Patients with the AG genotype (DPYD \*1/\*9A) and cancer who are treated with fluorouracil may have 1) an increased risk for drug toxicities, 2) decreased response and 3) increased DPYD activity as compared to patients with the AA genotype (DPYD \*1/\*1). Patients with the AG genotype were also found to have increased clearance of fluorouracil as compared to those with the GG genotype (DPYD \*9A/\*9A). However, multiple studies find contradictory or negative evidence for drug toxicities, fluorouracil clearance, and DPYD activity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

Class 3	rs1801158	CC
	IDICOTIO	$\sim$

Patients with the CC genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) a decreased but not absent risk of toxicity and 2) increased DPYD activity as compared to patients with the CT genotype (DPYD \*1/\*4). However, some studies find no association with drug toxicity or DPYD activity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

# · Class 4 rs1801266 GG

Patients with the GG genotype (DPYD \*1/\*1) may have increased DPYD activity as compared to those with the AA or AG genotype (DPYD \*8/\*8 or \*1/\*8). Other genetic and clinical factors may also affect DPYD activity.

# · Class 4 rs1801268 CC

Patients with the CC genotype (DPYD \*1/\*1) may have increased DPYD activity as compared to those with the AC or AA genotype (DPYD \*1/\*10 or \*10/\*10). Other genetic and clinical factors may also affect DPYD activity.

# · Class 4 rs72549306 CC

Patients with the CC genotype (DPYD \*1/\*1) may have increased DPYD activity as compared to those with the AA or AC genotype (DPYD \*11/\*11 or \*1/\*11). Other genetic and clinical factors may also affect DPYD activity.

VEGFA	vascular endothelial growth factor A
	——————————————————————————————————————
capecitabine, fluorouracil, irinotec	ad the AC genotype may have a reduced response to bevacizumab, an, leucovorin, or oxaliplatin as compared to patients with the CC tic factors may also affect response to chemotherapy in people with
IGFBP3	insulin-like growth factor binding protein $3$
	——- Clinical Annotations ————
	d stomach cancer may have a poorer survival outcomes when treated patients with the GT or TT genotype. Other genetic and clinical outcome.

### **FLUVASTATIN**

SLCO1B1	solute carrier organic anion transporter family, member 1B1
	Clinical Annotations —

# Class 3 rs11045819 *CC*

Patients with the CC genotype who are treated with fluvastatin may have a lesser reduction in LDL-C as compared to patients with the AC and AA genotype.

9	otype may have a smaller increase in HDL cholesterol when treated with patients with the AG genotype. Other genetic and clinical factors may also
CYP3A	cytochrome P450, family 3, subfamily A
_	otype may have a smaller increase in HDL cholesterol when treated with patients with the AG genotype. Other genetic and clinical factors may also
GEFITINIB	
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
Cl	- Clinical Annotations
and skin rash when treated	${\cal G}$ type and non-small cell lung cancer may have a decreased risk for diarrhea
Patients with the GG geno and skin rash when treated and clinical factors may als	G type and non-small cell lung cancer may have a decreased risk for diarrhea with gefitinib as compared to patients with the AA genotype. Other genetic
Patients with the GG geno and skin rash when treated and clinical factors may als GEMCITABINE	type and non-small cell lung cancer may have a decreased risk for diarrhea with gefitinib as compared to patients with the AA genotype. Other genetic o influence drug toxicity risk in patients receiving gefitinib.
Patients with the GG geno and skin rash when treated and clinical factors may als  GEMCITABINE  CDA  Class 3 rs1048977 C'  Patients with cancer and the to patients with the TT gen	type and non-small cell lung cancer may have a decreased risk for diarrhea with gefitinib as compared to patients with the AA genotype. Other genetic o influence drug toxicity risk in patients receiving gefitinib.  cytidine deaminase  ———————————————————————————————————
Patients with the GG geno and skin rash when treated and clinical factors may als  GEMCITABINE  CDA  Class 3 rs1048977 C'  Patients with cancer and the to patients with the TT gen	type and non-small cell lung cancer may have a decreased risk for diarrhea with gefitinib as compared to patients with the AA genotype. Other genetic o influence drug toxicity risk in patients receiving gefitinib.  cytidine deaminase  ———————————————————————————————————
Patients with the GG geno and skin rash when treated and clinical factors may als  GEMCITABINE  CDA  Class 3 rs1048977 C  Patients with cancer and th to patients with the TT ger and clinical factors may als	type and non-small cell lung cancer may have a decreased risk for diarrhea with gefitinib as compared to patients with the AA genotype. Other genetic o influence drug toxicity risk in patients receiving gefitinib.  cytidine deaminase  ———————————————————————————————————

Patients with the AG genotype and schizophrenia may have an increased risk for developing extrapyramidal symptoms when treated with haloperidol as compared to patients with the AA or GG genotype. Other genetic and clinical factors may also influence risk for extrapyramidal symptoms when taking haloperidol.

### HMG COA REDUCTASE INHIBITORS

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	Clinical Annotations

# · Class 3 rs1045642 GG

Patients with the GG genotype may have decreased serum creatine kinase levels when treated with hmg CoA reductase inhibitors as compared to patients with the AA genotype. Other genetic and clinical factors may also influence serum creatine kinase levels.

 $\cdot$  Class 3 rs1128503 GG

Patients with the GG genotype may have decreased serum creatine kinase levels when treated with hmg CoA reductase inhibitors as compared to patients with the AA genotypes. Other genetic and clinical factors may also influence serum creatine kinase levels.

#### ILOPERIDONE

CYP2D6	cytochrome P450, family 2, subfamily D, polypeptide 6

# · Class 3 rs1065852 GG

Patients with the GG genotype and schizophrenia may have an increased QTc interval when treated with iloperidone as compared to patients with the AA or AG genotype. Other genetic and clinical factors may also influence QTc interval.

#### **IMATINIB**

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	— Clinical Annotations —

# · Class 3 rs1045642 GG

Patients with the GG genotype and chronic myeloid leukemia may have an increased likelihood of achieving complete molecular response when treated with imatinib, as compared to patients with the AA or AG genotype. However, this was only significant in an exclusively Caucasian population. Additionally, no significant results were seen when considering major molecular response. Other genetic and clinical factors may also influence likelihood of achieving complete molecular response.

# · Class 3 rs1128503 GG

Patients with the GG genotype and chronic myeloid leukemia may have a better response to imatinib treatment as compared to patients with the AA or AG genotype. Other genetic and clinical factors may also influence response to imatinib.

CV	Do	<b>R6</b>
( ; Y	P'	Kh

cytochrome P450, family 3, subfamily A

CYP2B6	cytochrome P450, family 2, subfamily B, polypeptide 6
ment with imatinib a cytogenetic resistance for side effects as con	G genotype and chronic myeloid leukemia may have a 1) a better response to treat as compared to patients with the TT genotype, 2) an increased risk of developing to imatinib as compared to patients with the GT genotype, and 3) a greater risk apared to patients with the GT or TT genotype. Other genetic and clinical factors esponse, resistance and risk of side effects in patients taking imatinib.
CYP2A7P1	cytochrome P450, family 2, subfamily A, polypeptide 7 pseudogene 1
	Clinical Annotations
ment with imatinib a cytogenetic resistance for side effects as con	G genotype and chronic myeloid leukemia may have a 1) a better response to treat as compared to patients with the TT genotype, 2) an increased risk of developing to imatinib as compared to patients with the GT genotype, and 3) a greater risk appared to patients with the GT or TT genotype. Other genetic and clinical factors esponse, resistance and risk of side effects in patients taking imatinib.  cytochrome P450, family 3, subfamily A, polypeptide 5
	Government of the CT and TT genotypes. Other genetic and clinical factors
ZSCAN25	zinc finger and SCAN domain containing 25
	- Clinical Annotations
	genotype and chronic myeloid leukemia have have increased trough concentrations d to patients with the CT and TT genotypes. Other genetic and clinical factors

CC

Class 3 rs776746

CYP3A

Patients with the CC genotype and chronic myeloid leukemia have have increased trough concentrations of imatinib compared to patients with the CT and TT genotypes. Other genetic and clinical factors may affect concentrations of imatinib.

-- Clinical Annotations -

CYP2C9	cytochrome P450, family 2, subfamily C, polypeptide 9
of irbesartan as compared to Other clinical or genetic factor hypertension.	pe and essential hypertension may have decreased metabolism or clearance o patients with the AA genotype, but may have no difference in response. ors may also influence concentrations of irbesartan in patients with essential
may result may in decrease	notype may have increased metabolism and clearance of irbesartan which deposure of irbesartan as compared to patients with the CT genotype. Actors may also influence metabolism of irbesartan.
IRINOTECAN	
SLCO1B1	solute carrier organic anion transporter family, member 1B1
irinotecan or irinotecan-base a different study of similar s	rpe and cancer may have a decreased risk of neutropenia when treated with d regimens, as compared to patients with the CC or CT genotype. However, size found no association between the TT genotype and neutropenia. No a seen for diarrhea. Other genetic and clinical factors may also influence nea.

Patients with the GG genotype and non-small cell lung cancer may have a decreased risk of neutropenia when treated with irinotecan as compared to patients with the AG or GG genotype. No association has been seen for diarrhea. Other genetic and clinical factors may also influence risk of neutropenia.

 $\cdot$  Class 3 rs2306283 AG

Patients with the AG genotype and solid tumors may experience increased risk of neutropenia compared to patients with the AA genotype. However, studies conflict as to this association. Other clinical and genetic factors may affect risk of neutropenia with irinotecan therapy.

UGTIAIU	UDP glucuronosyltransierase 1 family, polypeptide A10
	- Clinical Annotations

# · Class 3 rs3832043 T/del

Patients with the T/del genotype and non-small cell lung cancer may have a decreased risk of diarrhea when treated with irinotecan or irinotecan-based regimens as compared to patients with the del/del genotype. However, a different study of similar size found no association between this genotype and diarrhea. No significant results have been seen when considering neutropenia or tumor response. Other genetic and clinical factors may also influence risk of diarrhea.

· Class 3 rs11692021 TC

Patients with the CT genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the TT genotype, or a decreased risk as compared to patients with the CC genotype. This may be due to decreased enzymatic activity toward SN-38, the active metabolite of irinotecan, found in cells with the C allele as compared to those with the T allele. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan.

· Class 3 rs2070959 AG

Patients with the AG genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the GG genotype, or a decreased risk as compared to patients with the AA genotype. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan.

· Class 3 rs10929302 GA

The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype.

UGT1A8	UDP glucuronosyltransferase 1 family, polypeptide A8

# · Class 3 rs3832043 T/del

Patients with the T/del genotype and non-small cell lung cancer may have a decreased risk of diarrhea when treated with irinotecan or irinotecan-based regimens as compared to patients with the del/del genotype. However, a different study of similar size found no association between this genotype and diarrhea. No significant results have been seen when considering neutropenia or tumor response. Other genetic and clinical factors may also influence risk of diarrhea.

· Class 3 rs11692021 TC

Patients with the CT genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the TT genotype, or a decreased risk as compared to patients with the CC genotype. This may be due to decreased enzymatic activity toward SN-38, the active metabolite of irinotecan, found in cells with the C allele as compared to those with the T allele. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan.

· Class 3 rs2070959 AG

Patients with the AG genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the GG genotype, or a decreased risk as compared to patients with the AA genotype. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan.

 $\cdot$  Class 3 rs10929302 GA

The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype.

UGT1A9	UDP glucuronosyltransferase 1 family, polypeptide A9
	Clinical Annotations ————————————————————————————————————

# · Class 3 rs3832043 T/del

Patients with the T/del genotype and non-small cell lung cancer may have a decreased risk of diarrhea when treated with irinotecan or irinotecan-based regimens as compared to patients with the del/del

genotype. However, a different study of similar size found no association between this genotype and diarrhea. No significant results have been seen when considering neutropenia or tumor response. Other genetic and clinical factors may also influence risk of diarrhea.

# Class 3 rs11692021 TC

Patients with the CT genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the TT genotype, or a decreased risk as compared to patients with the CC genotype. This may be due to decreased enzymatic activity toward SN-38, the active metabolite of irinotecan, found in cells with the C allele as compared to those with the T allele. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan.

# $\cdot$ Class 3 rs2070959 AG

Patients with the AG genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the GG genotype, or a decreased risk as compared to patients with the AA genotype. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan.

#### · Class 3 rs10929302 GA

The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype.

ABCC2
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ATP-binding cassette, sub-family C (CFTR/MRP), member 2

Clinical Annotations	
Cilincai Alliiotations	

# · Class 3 rs2273697 GG

Patients with the GG genotype and colorectal cancer may have decreased metabolism of irinotecan as compared to patients with the AA or AG genotype. Other genetic and clinical factors may also influence metabolism of irinotecan.

# · Class 3 rs3740066 CT

Patients with the CT genotype and non-small cell lung cancer may have a decreased risk of diarrhea when treated with irinotecan as compared to patients with the CC genotype. No association has been seen for neutropenia. Other genetic and clinical factors may also influence risk of diarrhea.

#### UGT1A6

UDP glucuronosyltransferase 1 family, polypeptide A6

# · Class 3 rs11692021 TC

Patients with the CT genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the TT genotype, or a decreased risk as compared to patients with the CC genotype. This may be due to decreased enzymatic activity toward SN-38, the active metabolite of irinotecan, found in cells with the C allele as compared to those with the T allele. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan.

#### $\cdot$ Class 3 rs2070959 AG

Patients with the AG genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the GG genotype, or a decreased risk as compared to patients with the AA genotype. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan.

# $\cdot$ Class 3 rs10929302 GA

The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype. UDP glucuronosyltransferase 1 family, polypeptide A7 UGT1A7 — Clinical Annotations Class 3 rs11692021 TC Patients with the CT genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the TT genotype, or a decreased risk as compared to patients with the CC genotype. This may be due to decreased enzymatic activity toward SN-38, the active metabolite of irinotecan, found in cells with the C allele as compared to those with the T allele. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan. Class 3 rs2070959 Patients with the AG genotype and colorectal cancer may have an increased risk for severe neutropenia when treated with irinotecan as compared to patients with the GG genotype, or a decreased risk as compared to patients with the AA genotype. Other genetic and clinical factors, such as UGT1A1\*28, may also influence the risk for neutropenia in patients taking irinotecan. Class 3 rs10929302 GA The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype. UGT1A3 UDP glucuronosyltransferase 1 family, polypeptide A3 —- Clinical Annotations – Class 3 rs10929302 GA The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype. UGT1A4 UDP glucuronosyltransferase 1 family, polypeptide A4 -- Clinical Annotations Class 3 rs10929302 GA The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype.

UGT1A5

UDP glucuronosyltransferase 1 family, polypeptide A5

—- Clinical Annotations —

· Class 3 rs10929302 GA

The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype.

UGT1A:
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UDP glucuronosyltransferase 1 family, polypeptide A1

- Clinical Annotations

# · Class 3 rs10929302 GA

The AG genotype may be associated with a reduced risk for irinotecan-induced grade 3 or 4 hematological and gastrointestinal toxicities, including neutropenia and diarrhea, as compared to the AA genotype.

#### **VEGFA**

vascular endothelial growth factor A

-- Clinical Annotations

# · Class 3 rs699947 AC

Patients with colorectal cancer and the AC genotype may have a reduced response to bevacizumab, capecitabine, fluorouracil, irinotecan, leucovorin, or oxaliplatin as compared to patients with the CC genotype. Other clinical and genetic factors may also affect response to chemotherapy in people with colorectal cancer.

#### **ISONIAZID**

# NAT2

N-acetyltransferase 2 (arylamine N-acetyltransferase)

-- Clinical Annotations -

# · Class 2A rs1041983 TT

Patients with the TT genotype and tuberculosis (TB) may have an increased risk for hepatotoxicity when treated with anti-TB drugs as compared to patients with the CC genotype. Other genetic and clinical factors may also influence risk for hepatotoxicity.

#### · Class 2A rs1799930 AA

Patients with the AA genotype and tuberculosis (TB) may have an increased risk of hepatotoxicity when treated with anti-TB drugs as compared to patients with the GG genotype. They also may have decreased clearance of isoniazid as compared to those with the AG or GG genotype. Other genetic and clinical factors may also influence risk for hepatotoxicity and clearance of isoniazid.

# · Class 3 rs1799931 GG

Patients with the GG genotype and tuberculosis (TB) may have a decreased risk of hepatotoxicity when treated with anti-TB drugs as compared to patients with the AA or AG genotype. However, some studies find no association with hepatotoxicity. Other genetic and clinical factors may also influence risk of hepatotoxicity.

#### **IVACAFTOR**

**CFTR** cystic fibrosis transmembrane conductance regulator (ATP-binding cassette sub-family C, member 7)

compared to patients wit	notype and cystic fibrosis may not respond when treated with ivacaftor as the AA and AG genotypes. Other genetic and clinical factors may also	
influence the efficacy of ivacaftor.  AMIVUDINE		
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member $1$	
Patients with the GG gene highly active antiretrovira	GG otype and HIV may have an increased risk of virological failure when receiving the therapy (HAART), as compared to patients with the AA genotype. Others may also influence risk of virological failure on HAART.	
AMOTRIGINE		
UGT1A10	UDP glucuronosyltransferase 1 family, polypeptide A10	
	- Clinical Annotations	
serum concentrations of thigher dose as compared to	TT notype and epilepsy who are administered lamotrigine may have increased lamotrigine, as well as improved response to lamotrigine, and may need to patients with the GG genotype. Other clinical and genetic factors may also ponse, and dose of lamotrigine.	
UGT1A4	UDP glucuronosyltransferase 1 family, polypeptide A4	
	- Clinical Annotations	
serum concentrations of thigher dose as compared to	TT notype and epilepsy who are administered lamotrigine may have increased lamotrigine, as well as improved response to lamotrigine, and may need a patients with the GG genotype. Other clinical and genetic factors may also ponse, and dose of lamotrigine.	
UGT1A5	UDP glucuronosyltransferase 1 family, polypeptide A5	

Patients with the TT genotype and epilepsy who are administered lamotrigine may have increased serum concentrations of lamotrigine, as well as improved response to lamotrigine, and may need a higher dose as compared to patients with the GG genotype. Other clinical and genetic factors may also influence metabolism, response, and dose of lamotrigine.

UGT1A6	UDP glucuronosyltransferase 1 family, polypeptide A6

# ullet Class 2B rs2011425 TT

Patients with the TT genotype and epilepsy who are administered lamotrigine may have increased serum concentrations of lamotrigine, as well as improved response to lamotrigine, and may need a higher dose as compared to patients with the GG genotype. Other clinical and genetic factors may also influence metabolism, response, and dose of lamotrigine.

UGT1A7	UDP glucuronosyltransferase 1 family, polypeptide A7		

# $\cdot$ Class 2B rs2011425 TT

Patients with the TT genotype and epilepsy who are administered lamotrigine may have increased serum concentrations of lamotrigine, as well as improved response to lamotrigine, and may need a higher dose as compared to patients with the GG genotype. Other clinical and genetic factors may also influence metabolism, response, and dose of lamotrigine.

UGT1A8	UDP glucuronosyltransferase 1 family, polypeptide A8
	——- Clinical Annotations

# $\cdot$ Class 2B rs2011425 TT

Patients with the TT genotype and epilepsy who are administered lamotrigine may have increased serum concentrations of lamotrigine, as well as improved response to lamotrigine, and may need a higher dose as compared to patients with the GG genotype. Other clinical and genetic factors may also influence metabolism, response, and dose of lamotrigine.

UGT1A9	UDP glucuronosyltransferase 1 family, polypeptide A9

# $\cdot$ Class 2B rs2011425 TT

Patients with the TT genotype and epilepsy who are administered lamotrigine may have increased serum concentrations of lamotrigine, as well as improved response to lamotrigine, and may need a higher dose as compared to patients with the GG genotype. Other clinical and genetic factors may also influence metabolism, response, and dose of lamotrigine.

# **LORAZEPAM**

	- Clinical Annotations
subjects with the CC genot	AC type may have decreased clearance of oxazepam or lorazepam as compared to type, or increased clearance as compared to subjects with the AA genotype. factors may also influence the oral clearance of oxazepam or lorazepam.
LOSARTAN	
CYP2C9	cytochrome P450, family 2, subfamily C, polypeptide 9
· ·	notype who are treated with losartan may have decreased metabolism of abjects with the AA genotype. Other genetic and clinical factors may also
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	- Clinical Annotations
9	G otype may have poorer response to losartan in people with hypertension as the AA or AG genotype. Other genetic and clinical factors may also influence
ТРМТ	thiopurine S-methyltransferase
	- Clinical Annotations
experience decreased GI to as compared to patients w	T TT genotype and Precursor Cell Lymphoblastic Leukemia-Lymphoma may xicity when treated with mercaptopurine and may require an increased dose ith the CT or CC genotypes. Other genetic and clinical factors may also GI toxicity and dose of mercaptopurine in pediatric patients with Precursor

Cell Lymphoblastic Leukemia-Lymphoma.

Class 4 rs3931660 AA

Patients with the AA genotype may have increased TPMT activity toward mercaptopurine as compared to patients with the AT genotype. Other genetic and clinical factors may also influence TPMT activity.

# **METHADONE**

an increased dose of the drug as factors may also influence dose  Class 3 rs2279343 AA  Patients with the AA genotype an increased dose of the drug	who are being treated with methadone for heroin addiction may require as compared to patients with the GG genotype. However, one study his variant and methadone dose. Other genetic and clinical factors may
CYP2A7P1	cytochrome P450, family 2, subfamily A, polypeptide 7 pseudogene 1
	who are being treated with methadone for heroin addiction may require compared to patients with the TT genotype. Other genetic and clinical of methadone.
CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
when treated with methadone in TT or CT. Other genetic and colors are safety as a second color of the treatment of the treatm	who are heroin dependent may have less severe side effects and opioid eated with methadone as compared to patients with genotype eated with methadone as compared to patients with genotype eated with methadone as compared to patients with the CC genotype.

# METHOTREXATE

SLCO1B1 solute carrier organic anion transporter family, member 1B1

— Clinical Annotations

# · Class 3 rs4149056 TT

Pediatric patients with the TT genotype and acute lymphoblastic leukemia may have increased clearance of methotrexate as compared to patients with the CC or CT genotype. Other genetic and clinical factors may also influence clearance of methotrexate.

# $\cdot$ Class 3 rs2306283 AG

Pediatric patients with the AG genotype and acute lymphoblastic leukemia may have increased clearance of methotrexate as compared to patients with the GG genotype. Other genetic and clinical factors may also influence clearance of methotrexate.

ABCC2	ATP-binding cassette, sub-family C (CFTR/MRP), member 2 $$
	- Clinical Annotations
as compared to patients study done in an Asian por decreased clearance of me distribution of volume of  Class 3 rs717620 C  Patients with lymphoblas drug toxicity and decreas to patients with the CC a French population with	otype may have increased clearance and distribution of volume of methotrexate with the CC genotype. Please note: the opposite effect was observed in a oppulation with various types of lymphomas. The T allele was associated with ethotrexate. Other clinical and genetic factors may also affect clearance and methotrexate.  The ticleukemia-lymphoma and the CT genotype may have an increased risk of sed clearance of methotrexate when treated with methotrexate as compared genotype. Please note: the opposite effect was observed in a study done in a various types of lymphomas. The C allele was associated with decreased et. Other clinical and genetic factors may also influence risk of drug toxicity
METOPROLOL	
ADRB2	adrenoceptor beta 2, surface
	- Clinical Annotations
triglyceridemia when trea	enotype and hypertension may have an increased risk of developing hyperted with atenolol or metoprolol as compared to patients with the CC or CG and clinical factors may also influence risk of hypertriglyceridemia.
MIDAZOLAM	
CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
to patients with the AG g midazolam.  • Class 4 rs12721627	otype and tumors may have increased metabolism of midazolam as compared genotype. Other genetic and clinical factors may also influence metabolism of $GG$ ruct caring the G variant is not associated with decreased clearance of mida-

-- Clinical Annotations

cytochrome P450, family 3, subfamily A

CYP3A

Patients with the GG genotype and tumors may have increased metabolism of midazolam as compared to patients with the AG genotype. Other genetic and clinical factors may also influence metabolism of midazolam.

Class 4 rs12721627 GG

The expression of a construct caring the G variant is not associated with decreased clearance of midazolam in transfected cells.

т.	T (	$\mathbf{T}$	<b>P</b>	TTI	гт т	$\mathbf{T}$
I۷	41	JК	P	н	IIV	H

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1
	— Clinical Annotations —

· Class 3 rs1045642 GG

Patients with the GG genotype may have decreased pain reduction when treated with morphine in cancer patients as compared to patients with genotype AA. Other genetic and clinical factors may also influence response to morphine.

#### **NEVIRAPINE**

CYP2B6	cytochrome P450, family 2, subfamily B, polypeptide 6
	- Clinical Annotations —

· Class 2A rs3745274 GG

Patients with the GG genotype and HIV infection may have increased clearance of and decreased exposure to nevirapine as compared to patients with the TT or GT genotype. Other genetic and clinical factors may also influence clearance of nevirapine and exposure to drug.

· Class 3 rs28399499 TT

Patients with the TT genotype and HIV may have a decreased risk for Stevens-Johnson Syndrome/toxic epidermal necrolysis (SJS/TEN) when treated with nevirapine as compared to patients with the CC or CT genotype. Other genetic and clinical factors may also influence risk for developing SJS/TEN when receiving nevirapine.

CYP2A7P1	cytochrome P450, family 2, subfamily A, polypeptide 7 pseudogene 1
	- Clinical Annotations

# Class 2A rs3745274 GG

Class 3 rs776746

CC

Patients with the GG genotype and HIV infection may have increased clearance of and decreased exposure to nevirapine as compared to patients with the TT or GT genotype. Other genetic and clinical factors may also influence clearance of nevirapine and exposure to drug.

clinical factors may also influence	clearance of nevirapine and exposure to drug.
CYP3A5	cytochrome P450, family 3, subfamily A, polypeptide 5
	——- Clinical Annotations ————————————————————————————————————

Patients with the CC genotype and HIV infection who are treated with nevirapine may have increased clearance of the drug as compared to patients with the CT and TT genotype. Association with clearance was not found in a larger cohort in a separate study. Patients may also have differences in alanine aminotransferase levels, but association with toxicity has not been reported. Other genetic and clinical factors may also influence clearance of nevirapine.

ZSCAN25	zinc finger and SCAN domain containing 25
	linical Annotations —
Class 3 rs776746 CC	

Patients with the CC genotype and HIV infection who are treated with nevirapine may have increased clearance of the drug as compared to patients with the CT and TT genotype. Association with clearance was not found in a larger cohort in a separate study. Patients may also have differences in alanine aminotransferase levels, but association with toxicity has not been reported. Other genetic and clinical factors may also influence clearance of nevirapine.

CYP3A cytochrome P450, family 3, subfamily A -- Clinical Annotations -

# Class 3 rs776746 CC

Patients with the CC genotype and HIV infection who are treated with nevirapine may have increased clearance of the drug as compared to patients with the CT and TT genotype. Association with clearance was not found in a larger cohort in a separate study. Patients may also have differences in alanine aminotransferase levels, but association with toxicity has not been reported. Other genetic and clinical factors may also influence clearance of nevirapine.

ABCC10 ATP-binding cassette, sub-family C (CFTR/MRP), member 10 -- Clinical Annotations

# Class 3 rs2125739 TT

Patients with the TT genotype and HIV may have increased concentrations of nevirapine as compared to patients with the CC genotype. Other genetic and clinical factors may also influence concentrations of nevirapine.

CYP2D6 cytochrome P450, family 2, subfamily D, polypeptide 6 -- Clinical Annotations

# Class 3 rs28371706 GG

Pediatric patients with the GG genotype and HIV may have increased clearance of nevirapine as compared to pediatric patients with the AA or AG genotype. No significant association was seen in adults. Other genetic and clinical factors may also influence clearance of nevirapine.

#### **NICOTINE**

Patients with with the GG of this SNP	the AA genotype may have increased metabolism of nicotine as compared to patients or AG genotype. Other variants within the CYP2A6 gene should be considered - allele is part of the *7, *10, *19, *36, *37 CYP2A6 alleles. Other genetic and clinical factors ence metabolism of nicotine.
DDC	dopa decarboxylase (aromatic L-amino acid decarboxylase)
	- Clinical Annotations —
	2718541 $$ $$ $$ $$ $$ $$ $$ $$ $$ $$
NIFEDIPINE	
CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
· Class 4 rs4 In vitro, the c	A987161 $AA$ onstruct expressing the wild type allelic protein has average nifedipine metabolism.
CYP3A	cytochrome P450, family 3, subfamily A
	- Clinical Annotations —
women with to f nifedipine.  Class 4 rs4	nen with the CC genotype may have decreased clearance of nifedipine as compared to he CT or TT genotype. Other genetic and clinical factors may also influence clearance
CYP3A5	cytochrome P450, family 3, subfamily A, polypeptide 5
	- Clinical Annotations
_	$^{7}$ 76746 $^{\prime}$ $^{$

——- Clinical Annotations —

_	e CC genotype may have decreased clearance of nifedipine as compared to $\Gamma$ genotype. Other genetic and clinical factors may also influence clearance
ONDANSETRON	
CYP3A5	cytochrome P450, family 3, subfamily A, polypeptide 5
with the TT genotype. Oth	type may have increased metabolism of ondansetron as compared to patients ner genetic and clinical factors may also influence metabolism of ondansetron.
ZSCAN25	zinc finger and SCAN domain containing 25
_	type may have increased metabolism of ondansetron as compared to patients ner genetic and clinical factors may also influence metabolism of ondansetron.
CYP3A	cytochrome P450, family 3, subfamily A
with the TT genotype. Oth	type may have increased metabolism of ondansetron as compared to patients her genetic and clinical factors may also influence metabolism of ondansetron.
OPIOIDS	
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	- Clinical Annotations —
Patients with the GG geno	type may have a decreased risk of opioid dependence when exposed to opioids with the AG genotype. Other clinical and genetic factors may also influence upon exposure to opioids.

cate chol-O-methyl transfer as e

 $\mathbf{COMT}$ 

—- Clinical Annotations ————

Class 3 rs4680 GA  Patients with the AG genotype with substance withdrawal syndrome may have an increased likelihood of headache when discontinuing the use of analgesics (such as opioids, NSAIDs, triptans, ergot) as compared to patients with the AA genotype. Other clinical and genetic factors may also influence likelihood of headache in patients with withdrawal syndrome who discontinue the use of analgesics.	
PACLITAXEL	
CYP2C8	cytochrome P450, family 2, subfamily C, polypeptide 8
	- Clinical Annotations
Patients with the TT genoty the AA or AT genotypes, h may also influence clearance  Class 4 rs11572080 C  Patients with the CC genoty the CT or TT genotypes, h may also influence clearance Class 4 rs10509681 T  Patients with the TT genotypes	where this has not been shown in vivo. Other genetic and clinical factors are of paclitaxel.  The type may have increased metabolism of paclitaxel as compared to patients to patients per however this has not been shown in vivo. Other genetic and clinical pers, however this has not been shown in vivo. Other genetic and clinical
CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
	- Clinical Annotations
Patients with the GG general	otype may have increased metabolism of paclitaxel as compared to pagenotypes. Other genetic and clinical factors may also influence paclitaxel

# · Class 3 rs12721627 GG

CYP3A

Patients with the GG genotype may have increased metabolism of paclitaxel as compared to patients with the CC or CG genotypes. Other genetic and clinical factors may also influence paclitaxel metabolism.

——- Clinical Annotations —

cytochrome P450, family 3, subfamily A

# Class 3 rs776746 CC

Patients with the CC genotype may have decreased but not absent risk of neurotoxicity when treated with paclitaxel as compared to patients with the TT genotype. Other genetic and clinical factors may also influence risk of toxicity with paclitaxel.

CYP3A5	cytochrome P450, family 3, subfamily A, polypeptide 5
	- Clinical Annotations —
Class 3 rs776746 CC	
Patients with the CC genot	ype may have decreased but not absent risk of neurotoxicity when treated to patients with the TT genotype. Other genetic and clinical factors may with paclitaxel.
ZSCAN25	zinc finger and SCAN domain containing 25
	- Clinical Annotations —
Class 3 rs776746 CC	
Patients with the CC genot	type may have decreased but not absent risk of neurotoxicity when treated to patients with the TT genotype. Other genetic and clinical factors may with paclitaxel.
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
rate when treated with pac	ociated with increased disease control rate and increased overall survival elitaxel in Asians with metastatic breast cancer as compared to genotype
Genotype GG may be assorate when treated with pactors AG. However, contradictors Caucasians. Other genetic a Class 3 rs1045642 GG Patients with the GG genot when treated with paclitax genetic and clinical factors in	ciated with increased disease control rate and increased overall survival clitaxel in Asians with metastatic breast cancer as compared to genotype by findings have been reported and no association have been reported for and clinical factors may influence the response to paclitaxel.  Graph may have decreased risk of Neutropenia and Neurotoxicity Syndromes are in cancer patients as compared to patients with genotype AA. Other may influence the risk of adverse events to paclitaxel.
Genotype GG may be assorate when treated with pactors AG. However, contradictors Caucasians. Other genetic a Class 3 rs1045642 GG Patients with the GG genot when treated with paclitax	ciated with increased disease control rate and increased overall survival elitaxel in Asians with metastatic breast cancer as compared to genotype by findings have been reported and no association have been reported for and clinical factors may influence the response to paclitaxel.  Graph may have decreased risk of Neutropenia and Neurotoxicity Syndromes are in cancer patients as compared to patients with genotype AA. Other may influence the risk of adverse events to paclitaxel.
Genotype GG may be assorate when treated with pactors AG. However, contradictors Caucasians. Other genetic a Class 3 rs1045642 GG Patients with the GG genot when treated with paclitax genetic and clinical factors in	ciated with increased disease control rate and increased overall survival clitaxel in Asians with metastatic breast cancer as compared to genotype by findings have been reported and no association have been reported for and clinical factors may influence the response to paclitaxel.  Graph may have decreased risk of Neutropenia and Neurotoxicity Syndromes are in cancer patients as compared to patients with genotype AA. Other may influence the risk of adverse events to paclitaxel.
Genotype GG may be assorate when treated with pactors. AG. However, contradictory. Caucasians. Other genetic at Class 3 rs1045642 GG. Patients with the GG genot when treated with paclitax genetic and clinical factors of GINTERFERON ALFA.	ciated with increased disease control rate and increased overall survival elitaxel in Asians with metastatic breast cancer as compared to genotype by findings have been reported and no association have been reported for and clinical factors may influence the response to paclitaxel.  Graph may have decreased risk of Neutropenia and Neurotoxicity Syndromes tel in cancer patients as compared to patients with genotype AA. Other may influence the risk of adverse events to paclitaxel.
Genotype GG may be assorate when treated with pactors. AG. However, contradictory. Caucasians. Other genetic at Class 3 rs1045642 GG. Patients with the GG genot when treated with paclitax genetic and clinical factors of GINTERFERON ALFA.	ciated with increased disease control rate and increased overall survival clitaxel in Asians with metastatic breast cancer as compared to genotype by findings have been reported and no association have been reported for and clinical factors may influence the response to paclitaxel.  Grappe may have decreased risk of Neutropenia and Neurotoxicity Syndromes are in cancer patients as compared to patients with genotype AA. Other may influence the risk of adverse events to paclitaxel. 2A  — Dosing Guideline  — Dosing Guideline

Patients with genotype CT may have decreased response to daclatasvir, peginterferon alfa-2a, peginterferon alfa-2b and ribavirin in people with Hepatitis C, Chronic as compared to genotypes CC. SVR24 rates are higher in patients treated with the combination of daclatasvir and pegIFN-alfa/RBV than those receiving pegIFN-alfa/RBV alone across all IL28B genotypes (CC, CT, or TT) regardless of viral subtypes. Other genetic and clinical factors may also influence the response to daclatasvir therapy.

ay have decreased response to daclatasvir, peginterferon alfa-2a, peginter people with Hepatitis C, Chronic as compared to genotypes CC. SVR24 created with the combination of daclatasvir and pegIFN-alfa/RBV that RBV alone across all IL28B genotypes (CC, CT, or TT) regardless of viral clinical factors may also influence the response to daclatasvir therapy.
low density lipoprotein receptor
likelihood of sustained virological response when treated with pegylated ompared to patients with the AA or AG genotype. Other genetic and ence likelihood of sustained virological response.  2B
interferon, lambda 3
——————————————————————————————————————
2979860C Strong

feron alfa-2b and ribavirin in people with Hepatitis C, Chronic as compared to genotypes CC. SVR24 rates are higher in patients treated with the combination of daclatasvir and pegIFN-alfa/RBV than those receiving pegIFN-alfa/RBV alone across all IL28B genotypes (CC, CT, or TT) regardless of viral subtypes. Other genetic and clinical factors may also influence the response to daclatasvir therapy.

IFNL4

IFNL4

interferon, lambda 4 (gene/pseudogene)

Class 3 rs12979860 CT Patients with genotype CT may	y have decreased response to daclatasvir, peginterferon alfa-2a, peginter-
feron alfa-2b and ribavirin in p rates are higher in patients tre those receiving pegIFN-alfa/RE	eople with Hepatitis C, Chronic as compared to genotypes CC. SVR24 eated with the combination of daclatasvir and pegIFN-alfa/RBV than BV alone across all IL28B genotypes (CC, CT, or TT) regardless of viral inical factors may also influence the response to daclatasvir therapy.
LDLR	low density lipoprotein receptor
	Clinical Annotations —
· Class 3 rs14158 GG	
HIV may have an increased like interferon and ribavirin as con-	be who are co-infected with chronic hepatitis C, genotype 1 or 4, and selihood of sustained virological response when treated with pegylated inpared to patients with the AA or AG genotype. Other genetic and ince likelihood of sustained virological response.
PHENPROCOUMON	
VKORC1	vitamin K epoxide reductase complex, subunit 1
· Class 2A rs9934438 GA	
Patients with the AG genotype	be may have decreased dose of acenocoumarol or phenprocoumon as otype GG. Other genetic and clinical factors may also influence the dose oumon.
PRSS53	protease, serine, 53
· Class 2A rs9934438 GA	
Patients with the AG genotyp	be may have decreased dose of acenocoumarol or phenprocoumon as otype GG. Other genetic and clinical factors may also influence the dose bumon.
PHENYTOIN	
SCN1A	sodium channel, voltage-gated, type I, alpha subunit
• Class 2B rs3812718 CT	who are treated with phenytoin may require a higher dose as compared to
	Other genetic and clinical factors may also influence dose of phenytoin.

LDLR	low density lipoprotein receptor
	- Clinical Annotations —
ment as compared to patien	Type and vascular diseases may have a poorer response to pravastatin treatts with the TT genotype, or a better response as compared to patients with netic and clinical factors may also influence pravastatin response.
ROPOFOL	
UGT1A10	UDP glucuronosyltransferase 1 family, polypeptide A10
9	type may have decreased but not non-existent risk of adverse effects when npared to patients with the AA or AG genotype. Other genetic and clinical
UGT1A8	UDP glucuronosyltransferase 1 family, polypeptide A8
	type may have decreased but not non-existent risk of adverse effects when npared to patients with the AA or AG genotype. Other genetic and clinical
UGT1A9	UDP glucuronosyltransferase 1 family, polypeptide A9
Patients with the GG geno	type may have decreased but not non-existent risk of adverse effects when mpared to patients with the AA or AG genotype. Other genetic and clinical esponse to propofol.
CYP2B6	cytochrome P450, family 2, subfamily B, polypeptide 6

# Class 3 rs3745274 GG

Patients under general anaesthesia with genotypes GG may need increased dose of propofol as compared to patients with genotype TT or GT. Other genetic and clinical factors may also influence the dose of propofol.

CYP2A7P1	cytochrome P450, family 2, subfamily A, polypeptide 7 pseudogene 1
	Clinical Annotations
· Class 3 rs3745274 (	GG
_	esthesia with genotypes GG may need increased dose of propofol as compared TT or GT. Other genetic and clinical factors may also influence the dose of
QUETIAPINE	
COMT	catechol-O-methyltransferase
CI	
quetiapine as compared to	notype and schizophrenia may have a poorer response to treatment with patients with the GG genotype, or a better response as compared to patients ther genetic and clinical factors may also influence quetiapine response.
Patients with the CG generated to	notype and schizophrenia may have a poorer response to treatment with patients with the GG genotype, or a better response as compared to patients ther genetic and clinical factors may also influence quetiapine response.
RALOXIFENE	
UGT1A8	UDP glucuronosyltransferase 1 family, polypeptide A8
· Class 3 rs1042597	CC
	with the CC genotype and schizophrenia may have increased response to

Post menopausal women with the CC genotype and schizophrenia may have increased response to raloxifene compared to patients with the CG genotype. Other genetic and clinical factors may affect response to raloxifene.

#### **RANIBIZUMAB**

VEGFA	vascular endothelial growth factor A
	annotations —

· Class 3 rs2010963 CG

Patients with the CG genotype and choroidal neovascularization may have a better response to anti-VEGF treatment, as compared to patients with the CC genotype. Other genetic and clinical factors may also influence response to anti-VEGF treatment.

# REPAGLINIDE

SLCO1B1	solute carrier organic anion transporter family, member 1B1
· Class 3 rs2306283 A While the GG genotype is shown for the GA genotype	associated with reduced plasma concentrations of repaglinide, no results are
CYP2C8	cytochrome P450, family 2, subfamily C, polypeptide 8
compared to patients with in blood glucose lowering $\epsilon$	(CYP2C8*1/*1) genotype may have decreased metabolism of repaglinide the CT genotype (CYP2C8*3/*1). No association was found with differences efficacy. Please note, the study supporting this annotation was carried out in genetic and clinical factors may also influence metabolism of repaglinide.
NAT2	N-acetyltransferase 2 (arylamine N-acetyltransferase)
Class 2A rs1041983	TT
when treated with anti-TI clinical factors may also in	otype and tuberculosis (TB) may have an increased risk for hepatotoxicity drugs as compared to patients with the CC genotype. Other genetic and fluence risk for hepatotoxicity.
Patients with the AA gen when treated with anti-TE decreased clearance of ison clinical factors may also in	AA otype and tuberculosis (TB) may have an increased risk of hepatotoxicity drugs as compared to patients with the GG genotype. They also may have liazed as compared to those with the AG or GG genotype. Other genetic and affluence risk for hepatotoxicity and clearance of isoniazid.
Patients with the GG geno treated with anti-TB drug	type and tuberculosis (TB) may have a decreased risk of hepatotoxicity when gs as compared to patients with the AA or AG genotype. However, some with hepatotoxicity. Other genetic and clinical factors may also influence
SLCO1B1	solute carrier organic anion transporter family, member 1B1

· Class 3 rs11045819 CC

Patients with the CC genotype may have decreased clearance of rifampin as compared to patients with the AC genotype. Other genetic and clinical factors may also influence rifampin clearance.

	~1 -		. ~
•	Class 3	rs2306283	AG

Patients with the AG genotype may have increased clearance of rifampin as compared to patients with the GG genotype. Other genetic and clinical factors may also influence rifampin clearance.

#### RISPERIDONE

# ABCB1 ATP-binding cassette, sub-family B (MDR/TAP), member 1 — Clinical Annotations

# · Class 3 rs1045642 GG

Patients with the GG genotype and schizophrenia may have a shorter QTc interval when treated with risperidone as compared to patients with the AA or AG genotype. Other genetic and clinical factors may also influence QTc interval in patients taking risperidone.

CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4

# $\cdot$ Class 3 rs35599367 GG

Patients with the GG genotype and psychiatric disorders may have increased clearance of risperidone compared to patients with the AG genotype. Other clinical and genetic factors likely affect risperidone pharmacokinetics.

CYP3A	cytochrome P450, family 3, subtamily A

# · Class 3 rs35599367 GG

Patients with the GG genotype and psychiatric disorders may have increased clearance of risperidone compared to patients with the AG genotype. Other clinical and genetic factors likely affect risperidone pharmacokinetics.

#### RITONAVIR.

UGT1A	UDP glucuronosyltransferase 1 family, polypeptide A complex locus

# · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

# · Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

Class 3 rs8330 *GC* 

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

# Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

· Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

· Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A3

UDP glucuronosyltransferase 1 family, polypeptide A3

— Clinical Annotations

#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

· Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

· Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

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# · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

# · Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

# · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A5	UDP glucuronosyltransferase 1 family, polypeptide $A5$

# · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

#### · Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

#### · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A6	UDP glucuronosyltransferase 1 family, polypeptide A6

#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

# Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

# · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A7

UDP glucuronosyltransferase 1 family, polypeptide A7

— Clinical Annotations	
- Chinca Annotations	

# · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

# Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

# · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGTIA8	UDP glucuronosyltransferase I family, polypeptide A8

#### · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

# · Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

#### · Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.


# · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

· Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

 $\cdot$  Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

UGT1A1	UDP glucuronosyltransferase 1 family, polypeptide A1

# · Class 3 rs10929303 TC

Patients with the CT genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the TT genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

· Class 3 rs1042640 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to patients with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in people with HIV who are taking atazanavir and ritonavir.

 $\cdot$  Class 3 rs8330 GC

Patients with the CG genotype and HIV may have a decreased risk of nephrolithiasis when treated with atazanavir and ritonavir as compared to patients with the GG genotype and an increased risk of nephrolithiasis as compared to people with the CC genotype. Other genetic and clinical factors may also affect risk of nephrolithiasis in patients with HIV who are taking atazanavir and ritonavir.

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1

#### · Class 3 rs1045642 GG

Patients with the GG genotype and HIV may have an increased risk of virological failure when receiving highly active antiretroviral therapy (HAART), as compared to patients with the AA genotype. Other genetic and clinical factors may also influence risk of virological failure on HAART.

 $\cdot$  Class 3 rs1045642 GG

Patients with the GG genotype and HIV may have increased concentrations of atazanavir as compared to patients with the AA genotypes, although this is contradicted in one study. There is no evidence that the GG genotype is associated with hyperbilirubinemia, drug discontinuation, or nephrolithiasis. Other clinical and genetic factors may also influence the concentrations of atazanavir in patients with HIV.

ROSIGLITAZONE

CYP2C8	cytochrome P450, family 2, subfamily C, polypeptide 8
a larger change in HbA1c, an (CYP2C8*3/*3) or CT (CYP2c	C8*1/*1) genotype may have decreased metabolism of rosiglitazone, d an increased risk of edema as compared to patients with the CC C8*3/*1) genotype. One study found no association with blood glucose al factors may also influence metabolism of rosiglitazone, risk of edema
SLCO1B1	solute carrier organic anion transporter family, member 1B1
0 11	e may have decreased response to rosiglitazone in people with type II to patients with genotype CC or CT. Other genetic and clinical factors e to rosiglitazone.
VEGFA	vascular endothelial growth factor A
	may have decreased response to sildenafil in men with Erectile Dysfuncith genotype CC. Other genetic and clinical factors may also influence
SIMVASTATIN	
HMGCR	3-hydroxy-3-methylglutaryl-CoA reductase
	- Clinical Annotations
Class 4 mg2946669 CC	

The GG genotype may be associated with decreased induction of full-length transcripts and increased expression of spliced HMGCRv1 transcript as compared to AA genotype. ABCC2 ATP-binding cassette, sub-family C (CFTR/MRP), member 2 -- Clinical Annotations --Class 3 rs717620 Patients with the CT genotype may have decreased dose of simvastatin and atorvastatin as compared to patients with genotype CC. Other genetic and clinical factors may also influence the dose of simvastatin. ABCB1 ATP-binding cassette, sub-family B (MDR/TAP), member 1 -- Clinical Annotations --Class 3 rs1128503 GG Patients with the GG genotype and hypercholesterolemia may lesser reduction in LDL and total cholesterol when treated with simvastatin as compared to patients with the AA or AG genotype. Other genetic and clinical factors may also influence cholesterol levels. Class 3 rs1128503 GG Patients with the GG genotype and hypercholesterolemia may have an increased risk for myalgia when treated with simvastatin as compared to patients with the AA genotype. Other genetic and clinical factors may also influence risk for myalgia. CYP3A4 cytochrome P450, family 3, subfamily A, polypeptide 4 -- Clinical Annotations Class 3 rs2740574 TTPatients with the TT genotype may be more likely to require a decrease in dose or switch to a different drug when treated with atorvastatin or simvastatin as compared to patients with the CC or CT genotype. Other genetic and clinical factors may also influence dose of simvastatin or atorvastatin, or likelihood of switching to a different drug. CYP3A cytochrome P450, family 3, subfamily A —- Clinical Annotations -Class 3 rs2740574

Patients with the TT genotype may be more likely to require a decrease in dose or switch to a different drug when treated with atorvastatin or simvastatin as compared to patients with the CC or CT genotype. Other genetic and clinical factors may also influence dose of simvastatin or atorvastatin, or likelihood of switching to a different drug.

#### **SORAFENIB**

		Clinical Annotations –	
Class 3 rs2010963	CG		

Patients with the CG genotype may have increased risk of hand-foot syndrome when treated with sorafenib in people with Carcinoma, Renal Cell as compared to patients with genotype CC. Other genetic and clinical factors may also influence the toxicity to sorafenib.

· Class 3 rs1570360 AG

Patients with the AG genotype may have unfavorable progression-free survival when treated with sorafenib in people with Carcinoma, Renal Cell as compared to patients with genotype GG. Other genetic and clinical factors may also influence the response to sorafenib.

· Class 3 rs2010963 CG

Patients with the CG genotype may have increased progression-free survival and increased overall survival when treated with sorafenib in people with Hepatocellular Carcinoma as compared to patients with genotype GG. Other genetic and clinical factors may also influence the response to sorafenib.

SLCO1B1	solute carrier organic anion transporter family, member 1B1
	- Clinical Annotations —

# $\cdot$ Class 3 rs4149056 TT

Patients with the TT genotype may have increased likelihood of developing Thrombocytopenia when treated with sorafenib as compared to patients with genotype CC. Other genetic and clinical factors may also influence the response to sorafenib.

 $\cdot$  Class 3 rs2306283 AG

Patients with the AG genotype may have decreased likelihood of developing Diarrhea when treated with sorafenib as compared to patients with genotype AA. Other genetic and clinical factors may also influence the response to sorafenib.

ABCC2	ATP-binding cassette, sub-family C (CFTR/MRP), member 2
	—- Clinical Annotations ————

# $\cdot$ Class 3 rs717620 CT

Patients with the CT genotype may have decreased risk of skin rash when treated with sorafenib in people with Carcinoma, Renal Cell as compared to patients with genotype CC. Other genetic and clinical factors may also influence the toxicity to sorafenib.

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1
	- Clinical Annotations —

#### · Class 3 rs1045642 GG

Patients with the GG genotype may have decreased risk of hypertension when treated with sorafenib in people with Carcinoma, Renal Cell as compared to patients with genotype AA or AG. Other genetic and clinical factors may also influence the toxicity to sorafenib.

# SULFONAMIDES, UREA DERIVATIVES

	AC rith the AC genotype were not statistically significant.
SUNITINIB	
VEGFA	vascular endothelial growth factor A
of developing grade 3 hy	AC enotype may have higher increase in systolic blood pressure and increased risk ypertension when treated with sunitinib as compared to patients with genotype clinical factors may also influence the response to sunitinib.
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member $1$
	- Clinical Annotations —
· Class 3 rs1045642	GG
when treated with suni no association between Other genetic and clinic	enotype and renal cell carcinoma may have an increased risk for adverse effects tinib as compared to patients with the AA or AG genotype. One study found this SNP and thrombocytopenia, neutropenia, anemia or hand-food syndrome. cal factors may also influence risk for sunitinib toxicities.  CC
when treated with sunit	carcinoma and the CC genotypes may have an increased risk of neutropenia inib as compared to patients with any of the following genotypes: AA, AC, AT netic factors may also influence risk of neutropenia in patients with renal cell ted with sunitinib.  GG
Patients with renal cell increased risk of neutronalthough this has been	carcinoma and the GG genotype who are treated with sunitinib may have an penia, leukopenia, and diarrhea as compared to patients with the AA genotypes, a contradicted by some studies. Other clinical and genetic factors may also a patients with renal cell carcinoma who are administered sunitinib.  CC

Patients with renal cell carcinoma and the CC genotype may have an incressed response to sunitinib as compared to patients with the AA genotypes. There is no association between this SNP and overall or progression free survival. Response here refers to stable disease or partial response and non-response to progressive disease. Other clinical and genetic factors may also influence response to sunitinib in patients with renal cell carcinoma.

#### **TACROLIMUS**

# · Class 3 rs1045642 GG

Patients with the GG genotype who are undergoing organ transplantation may have increased clearance and dose requirements of tacrolimus, as compared to patients with the AA or AG genotype. However, the vast majority of studies find no association between this SNP and clearance or dose of tacrolimus. Other genetic and clinical factors, such as CYP3A5\*3, may also influence clearance and dose of tacrolimus.

#### · Class 3 rs1045642 GG

Patients with the GG genotype who are CYP2C19 extensive metabolizers and are receiving tacrolimus after renal transplantation may have increased plasma concentrations of (R)-lansoprazole but no significant differences in the frequency of gastroesophageal symptoms as compared to patients with the AA genotype. Other genetic and clinical factors may also influence lansoprazole clearance.

# · Class 3 rs1045642 GG

Pediatric patients with the GG genotype who are treated with prednisone and tacrolimus may have an increased risk of remaining on steroids 1 year after heart transplantation compared to patients with the AA or AG genotype. Other genetic and clinical factors may also influence risk of remaining on steroids 1 year after transplantation.

# $\cdot$ Class 3 rs1045642 GG

Patients who receive a kidney with the GG genotype may have increased estimated glomerular filtration rate (eGFR) when treated with tacrolimus as compared to patients with the AA or AG genotype. No significant results were seen when recipient genotype was considered. Other genetic and clinical factors may also influence eGFR.

# · Class 3 rs2032582 CC

Patients with CC genotype may have lower success rate in achieving short-term remission when treated with tacrolimus in people with Colitis, Ulcerative as compared to patients with the AA genotype. The majority of studies find no association with dose of tacrolimus in people with transplantations as compared and genotypes of this SNP. Other genetic or clinical factors may influence response and dose of tacrolimus.

#### $\cdot$ Class 3 rs1128503 GG

Patients with the GG genotype who are undergoing organ transplantation may have decreased concentrations of tacrolimus as compared to patients with the AA or AG genotype. However, the majority of the literature evidence shows no association between this variant and tacrolimus concentrations, clearance or dose. Other genetic and clinical factors may also influence concentrations of tacrolimus.

#### · Class 3 rs2032582 CC

Patients with the CC genotype who are undergoing organ transplantation may have increased metabolism and dose requirements of tacrolimus, as compared to patients with the AA, AC, CT or TT genotypes. However, the majority of studies have found no association between this polymorphism and metabolism or dose of tacrolimus. Other genetic and clinical factors, such as CYP3A5\*3, may also influence metabolism and dose of tacrolimus.

#### $\cdot$ Class 3 rs1045642 GG

Patients with the GG genotype and ulcerative colitis may have a poorer chance at achieving remission when treated with tacrolimus as compared to patients with the AA genotype. Other genetic and clinical factors may also influence likelihood of ulcerative colitis remission.

#### $\cdot$ Class 3 rs1045642 GG

Patients with the GG genotype who are undergoing kidney transplantation and are treated with tacrolimus may have decreased risk of experiencing transplant rejection as compared to patients with the AG genotype. However, the majority of studies find no association between this polymorphism and risk for transplant rejection. Other genetic and clinical factors may also influence risk of transplant rejection.

Patients with the GG genotype who are undergoing kidney transplantation may have a decreased risk of hypokalemia when treated with tacrolimus as compared to patients with the AG genotype. Other genetic and clinical factors may also influence risk of hypokalemia.

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cytochrome P450, family 3, subfamily A, polypeptide 5

— Dosing Guidel	ino		
Dosing Guider	.1116		

# CYP3A5:\*1A/\*1A Strong

Increase starting dose 1.5 to 2 times recommended starting dose. Total starting dose should not exceed 0.3mg/kg/day. Use therapeutic drug monitoring to guide dose adjustments.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing liver transplantation may have an increased risk for renal dysfunction when treated with tacrolimus as compared to patients with the CT or TT genotype (\*1/\*3 or \*1/\*1). Other genetic and clinical factors may also influence risk for renal dysfunction.

#### · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing organ transplantation may have a decreased risk for neurotoxicity when treated with tacrolimus as compared to patients with the CT (\*1/\*3) genotype. Other genetic and clinical factors may also influence risk for neurotoxicity in patients receiving tacrolimus.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing kidney transplantation may have decreased systolic and diastolic blood pressure when treated with tacrolimus as compared to patients with the CT or TT (\*1/\*3 or \*1/\*1) genotype. However, the majority of studies show no association between the CC genotype and blood pressure. Other genetic and clinical factors may also influence changes in blood pressure in patients receiving tacrolimus.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing organ transplantation may have a decreased risk for infections when treated with tacrolimus as compared to patients with the CT or TT (\*1/\*3 or \*1/\*1) genotype. Other genetic and clinical factors may also influence risk for infections in patients receiving tacrolimus.

#### · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) and ulcerative colitis may have an increased chance of achieving remission when treated with tacrolimus as compared to patients with the CT (\*1/\*3) genotype. Other genetic and clinical factors may also influence chance of remission from ulcerative colitis.

ZSCAN25	zinc finger and SCAN domain containing $25$
	nnotations —

# Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing liver transplantation may have an increased risk for renal dysfunction when treated with tacrolimus as compared to patients with the CT

or TT genotype ( $^*1/^*3$  or  $^*1/^*1$ ). Other genetic and clinical factors may also influence risk for renal dysfunction.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing organ transplantation may have a decreased risk for neurotoxicity when treated with tacrolimus as compared to patients with the CT (\*1/\*3) genotype. Other genetic and clinical factors may also influence risk for neurotoxicity in patients receiving tacrolimus.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing kidney transplantation may have decreased systolic and diastolic blood pressure when treated with tacrolimus as compared to patients with the CT or TT (\*1/\*3 or \*1/\*1) genotype. However, the majority of studies show no association between the CC genotype and blood pressure. Other genetic and clinical factors may also influence changes in blood pressure in patients receiving tacrolimus.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing organ transplantation may have a decreased risk for infections when treated with tacrolimus as compared to patients with the CT or TT (\*1/\*3 or \*1/\*1) genotype. Other genetic and clinical factors may also influence risk for infections in patients receiving tacrolimus.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) and ulcerative colitis may have an increased chance of achieving remission when treated with tacrolimus as compared to patients with the CT (\*1/\*3) genotype. Other genetic and clinical factors may also influence chance of remission from ulcerative colitis.

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$\mathbf{\mathbf{\mathcal{C}}}$	1	Г	O.	А

cytochrome P450, family 3, subfamily A

— Clinical Ann	notations	
—- Unincai Ani	notations —	

#### · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing liver transplantation may have an increased risk for renal dysfunction when treated with tacrolimus as compared to patients with the CT or TT genotype (\*1/\*3 or \*1/\*1). Other genetic and clinical factors may also influence risk for renal dysfunction.

#### · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing organ transplantation may have a decreased risk for neurotoxicity when treated with tacrolimus as compared to patients with the CT (\*1/\*3) genotype. Other genetic and clinical factors may also influence risk for neurotoxicity in patients receiving tacrolimus.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing kidney transplantation may have decreased systolic and diastolic blood pressure when treated with tacrolimus as compared to patients with the CT or TT (\*1/\*3 or \*1/\*1) genotype. However, the majority of studies show no association between the CC genotype and blood pressure. Other genetic and clinical factors may also influence changes in blood pressure in patients receiving tacrolimus.

# · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) undergoing organ transplantation may have a decreased risk for infections when treated with tacrolimus as compared to patients with the CT or TT (\*1/\*3 or \*1/\*1) genotype. Other genetic and clinical factors may also influence risk for infections in patients receiving tacrolimus.

#### · Class 3 rs776746 CC

Patients with the CC genotype (CYP3A5 \*3/\*3) and ulcerative colitis may have an increased chance of achieving remission when treated with tacrolimus as compared to patients with the CT (\*1/\*3) genotype. Other genetic and clinical factors may also influence chance of remission from ulcerative colitis.

# · Class 3 rs35599367 GG

Transplant recipients with the GG genotype (also known as CYP3A4 \*1/\*1) may have increased metabolism of tacrolimus, resulting in decreased exposure and a higher dose requirement (among other pharmacokinetic parameters), as compared to patients with the AG or AA genotype (CYP3A4 \*22/\*1 and \*22/\*22). Around half of studies report an association, and half report no association. One study found an association with liver transplant donor genotype but not recipient genotype. Other genetic and clinical factors, such as CYP3A5 \*3 (rs776746), may also influence metabolism of tacrolimus.

#### · Class 4 rs4986910 AA

Patients with the AA genotype who are undergoing kidney transplantation may require an increased dose of tacrolimus as compared to patients with the GG genotype. Other genetic and clinical factors, such as CYP3A5\*3, may also influence dose of tacrolimus.

CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
	- Clinical Annotations
	- Chincal Affilotations

# · Class 3 rs28371759 AA

Patients with the AA genotype (CYP3A4 \*1/\*1) who underwent kidney transplantation may have decreased metabolism of tacrolimus as compared to patients with the AG genotype (\*1/\*18B). Other genetic and clinical factors, such as rs776746 (CYP3A5\*3), may also influence metabolism of tacrolimus.

· Class 3 rs35599367 GG

Transplant recipients with the GG genotype (also known as CYP3A4 \*1/\*1) may have increased metabolism of tacrolimus, resulting in decreased exposure and a higher dose requirement (among other pharmacokinetic parameters), as compared to patients with the AG or AA genotype (CYP3A4 \*22/\*1 and \*22/\*22). Around half of studies report an association, and half report no association. One study found an association with liver transplant donor genotype but not recipient genotype. Other genetic and clinical factors, such as CYP3A5 \*3 (rs776746), may also influence metabolism of tacrolimus.

#### · Class 4 rs4986910 AA

Patients with the AA genotype who are undergoing kidney transplantation may require an increased dose of tacrolimus as compared to patients with the GG genotype. Other genetic and clinical factors, such as CYP3A5\*3, may also influence dose of tacrolimus.

#### **TAMOXIFEN**

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	- Clinical Annotations

# $\cdot$ Class 3 rs1045642 GG

Women with the GG genotype and breast cancer may have a decreased chance of disease recurrence when treated with tamoxifen as compared to patients with the AG genotype. Other genetic and clinical factors may also influence breast cancer recurrence.

	also influence risk of endometrial cancer.
CYP3A	cytochrome P450, family 3, subfamily A
Class 3 rs2740574	TT
cancer following tamoxid	notype and breast cancer may have a decreased risk of developing endometria fen treatment as compared to patients with the CT genotype. Other geneticalso influence risk of endometrial cancer.
ABCC2	ATP-binding cassette, sub-family C (CFTR/MRP), member 2 $$
	- Clinical Annotations —
Class 4 rs717620 (	CT
treated with tamoxifen factors may also influence	enotype and breast neoplasms may have increased disease-free survival whereast compared to patients with the CC genotype. Other genetic and clinicate disease-free survival with tamoxifen treatment.
UGT1A10	UDP glucuronosyltransferase 1 family, polypeptide A10
	obi graculonosymianolerase i lammy, polypepulae ilio
	- Clinical Annotations
Class 3 rs2011425 Patients with the TT g n-glucuronide when taki	——————————————————————————————————————
Class 3 rs2011425 Patients with the TT g n-glucuronide when taki clinical and genetic factor	——————————————————————————————————————
Class 3 rs2011425 Patients with the TT g n-glucuronide when taki	——————————————————————————————————————
Class 3 rs2011425 Patients with the TT g n-glucuronide when taki clinical and genetic factor	——————————————————————————————————————
Class 3 rs2011425 Patients with the TT g n-glucuronide when taki clinical and genetic facto  UGT1A4  Class 3 rs2011425 Patients with the TT g n-glucuronide when taki	——————————————————————————————————————
Class 3 rs2011425 Patients with the TT g n-glucuronide when taki clinical and genetic facto  UGT1A4  Class 3 rs2011425 Patients with the TT g n-glucuronide when taki	— Clinical Annotations  TT enotype and breast cancer may have decreased concentrations of tamoxifering tamoxifen compared to patients with the GG and GT genotypes. Other ors may affect the metabolism of tamoxifen.  UDP glucuronosyltransferase 1 family, polypeptide A4  — Clinical Annotations  TT enotype and breast cancer may have decreased concentrations of tamoxifering tamoxifen compared to patients with the GG and GT genotypes. Other

	UDP glucuronosyltransferase 1 family, polypeptide A6
	- Clinical Annotations —
n-glucuronide when taking tak	pe and breast cancer may have decreased concentrations of tamoxifen moxifen compared to patients with the GG and GT genotypes. Other by affect the metabolism of tamoxifen.
UGT1A7	UDP glucuronosyltransferase 1 family, polypeptide A7
	- Clinical Annotations —
n-glucuronide when taking tak	pe and breast cancer may have decreased concentrations of tamoxifen moxifen compared to patients with the GG and GT genotypes. Other by affect the metabolism of tamoxifen.
UGT1A8	UDP glucuronosyltransferase 1 family, polypeptide A8
	pe and breast cancer may have decreased concentrations of tamoxifen moxifen compared to patients with the GG and GT genotypes. Other
_	y affect the metabolism of tamoxifen.
_	
clinical and genetic factors ma	y affect the metabolism of tamoxifen.
Class 3 rs2011425 TT Patients with the TT genotypen-glucuronide when taking tak	UDP glucuronosyltransferase 1 family, polypeptide A9  ———————————————————————————————————
Class 3 rs2011425 TT Patients with the TT genotypen-glucuronide when taking tak	UDP glucuronosyltransferase 1 family, polypeptide A9  — Clinical Annotations  — ce and breast cancer may have decreased concentrations of tamoxifen moxifen compared to patients with the GG and GT genotypes. Other

Patients with the TT genotype and breast cancer may have decreased concentrations of tamoxifenn-glucuronide when taking tamoxifen compared to patients with the GG and GT genotypes. Other

	Clin	ical	Anı	nota	tions
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# · Class 1A rs55886062 AA

Patients with the AA genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have a decreased, but not absent, risk for drug toxicity as compared to patients with the AC or CC genotype (DPYD \*1/\*13 or \*13/\*13). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

# · Class 1A rs3918290 *CC*

Patients with the CC genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased clearance of fluoropyrimidine drugs and 2) decreased, but not non-existent, risk for drug toxicity as compared to patients with the CT or TT genotype (DPYD \*1/\*2A or \*2A/\*2A). Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine based chemotherapy.

#### $\cdot$ Class 1A rs67376798 TT

Patients with the TT genotype and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) increased clearance of the drug and 2) decreased, but not absent, risk and reduced severity of drug toxicity as compared to patients with the AT genotype. Fluoropyrimidines are often used in combination chemotherapy such as FOLFOX (fluorouracil, leucovorin and oxaliplatin), FOLFIRI (fluorouracil, leucovorin and irinotecan) or FEC (fluorouracil, epirubicin and cyclophosphamide) or with other drugs such as bevacizumab, cetuximab, raltitrexed. The combination and delivery of the drug may influence risk for toxicity. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

#### $\cdot$ Class 3 rs1801159 TT

Patients with the TT genotype (DPYD \*1/\*1) and cancer who are treated with fluoropyrimidine-based chemotherapy may have 1) a decreased likelihood of nausea, vomiting, and leukopenia, 2) increased response and 3) increased clearance of fluorouracil as compared to patients with the CT or CC genotype (DPYD \*1/\*5 or \*5/\*5). However, other studies find no associations or contradictory associations with fluoropyrimidine-induced drug toxicity or response. Other genetic and clinical factors may also influence response to fluoropyrimidine-based chemotherapy.

CYP2A6	cytochrome P450, family 2, subfamily A, polypeptide 6

#### · Class 4 rs28399433 AA

Hepatic cells with the AA genotype may have increased expression of the CYP2A6 gene, resulting in increased metabolism of tegafur, as compared to those with the AC or CC genotype. Other genetic and clinical factors may also influence CYP2A6 expression and tegafur metabolism.

# **THALIDOMIDE**

	- Clinical Annotations —
0 0.	e may have a decreased but not absent risk of toxicity with docetaxel and atients with the CT or TT genotypes. Other genetic and clinical factors response.
NAT2	N-acetyltransferase 2 (arylamine $N$ -acetyltransferase)
	- Clinical Annotations
	e may have an increased risk of toxicity with docetaxel and thalidomide the AA genotype. Other genetic and clinical factors may also influence
ΓICAGRELOR	
CYP3A4	cytochrome P450, family 3, subfamily A, polypeptide 4
	be and acute coronary syndrome may have decreased concentrations of ts with the CT genotype. Other factors may affect concentrations of
SLCO1B1	solute carrier organic anion transporter family, member 1B1
	be and acute coronary syndrome may have decreased concentrations of swith the CC and CT genotypes. Other factors may affect concentrations
FOLBUTAMIDE	
CYP2C9	cytochrome P450, family 2, subfamily C, polypeptide 9
	e may have increased metabolism of tolbutamide as compared to patients.  Other genetic and clinical factors may also influence tolbutamide

metabolism.

 $\cdot$  Class 3 rs2070959 AG

SLC22A1	solute carrier family $22$ (organic cation transporter), member $1$
	- Clinical Annotations
exposed to tramadol in healt genetic or clinical factors may Class 3 rs34130495 GG Patients with the GG genoty when exposed to tramadol in	pe may have lower plasma concentrations of O-desmethyltramadol when thy individuals as compared to patients with the TT genotype. Other y influence the response to tramadol.
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
response to tramadol.  VALPROIC ACID	
UGT1A10	UDP glucuronosyltransferase 1 family, polypeptide A10
	e may require an increased dose of valproic acid compared to patients with etic and clinical factors may also influence a patients dose requirements.
UGT1A6	UDP glucuronosyltransferase 1 family, polypeptide A6
0 0.	e may require an increased dose of valproic acid compared to patients with etic and clinical factors may also influence a patients dose requirements.
UGT1A7	UDP glucuronosyltransferase 1 family, polypeptide A7

the AA genotype. Other	er genetic and clinical factors may also influence a patients dose requirements.
UGT1A8	UDP glucuronosyltransferase 1 family, polypeptide A8
_	AG enotype may require an increased dose of valproic acid compared to patients with er genetic and clinical factors may also influence a patients dose requirements.
UGT1A9	UDP glucuronosyltransferase 1 family, polypeptide A9
	- Clinical Annotations
Patients with the AG ge	$AG$ enotype may require an increased dose of valproic acid compared to patients with $\Delta G$ genetic and clinical factors may also influence a patients dose requirements.
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member $1$
Patients with genotype pared to patients with g to have different respons	GG and depressive disorder may have increased response to venlafaxine com- genotype AA or AG. Patients with GG genotype and narcolepsy were not found se to venlafaxine compared to patients with other genotypes. Other clinical and a affect response to venlafaxine.
CYP2D6	cytochrome P450, family 2, subfamily D, polypeptide 6
	- Clinical Annotations —
· Class 4 rs36754300 Patients with the GG g	GG senotype were not studied.
/ERAPAMIL	
ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member $1$
· Class 3 rs2032582	CC

Patients with the AG genotype may require an increased dose of valproic acid compared to patients with

Patients with the CC genotype may have decreased metabolism of verapamil as compared to patients with the AA or AC genotype. Other genetic and clinical factors may also impact the metabolism of verapamil.

# · Class 3 rs1045642 GG

Patients with the GG genotype may have decreased metabolism of verapamil as compared to patients with the AA or AG genotype. Other genetic and clinical factors may also impact the metabolism of verapamil.

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V	11	ľΑ	ιN	/	IIN	H)

CYP4F2	cytochrome P450, family 4, subfamily F, polypeptide 2

# · Class 3 rs2108622 CC

Patients with the CC genotype may have decreased steady-state levels of vitamin E when taking vitamin E supplements as compared to patients with the CT and TT genotypes. Other clinical and genetic factors may also influence steady-state levels of vitamin E in patients taking vitamin E supplements.

 $\cdot$  Class 4 rs3093105 AA

The AA genotype may be associated with decreased CYP4F2 activity and decreased vitamin e metabolism as compared to the AC or CC genotype. This is based solely on an in vitro study in a haploid heterologous cell system. Other clinical and genetic factors may also influence metabolism of vitamin e.

#### WARFARIN

CYP2C9	cytochrome P450, family 2, subfamily C, polypeptide 9
	— Dosing Guideline ————————————————————————————————————

#### CYP2C9:\*3/\*1 N/A

Estimate the anticipated stable dose of warfarin using the algorithms available on http://www.warfarindosing.org, the IWPC Pharmacogenetic Dosing Algorithm or the FDA-approved drug label

#### · Class 1A rs1057910 AC

Patients with the AC genotype: 1) may require a decreased dose of warfarin as compared to patients with the AA genotype 2) may have an increased risk for adverse events as compared to patients with the AA genotype.

 $\cdot$  Class 2A rs7900194 GG

Patients with the GG genotype who are treated with warfarin may require a higher maintenance dose as compared to patients with the AG or GG genotype. Other clinical or genetic factors may also influence warfarin dose.

 $\cdot$  Class 2A rs56165452 TT

Patients with the TT genotype may required higher dose of warfarin as compared to patients with the CT or CC genotype. Other clinical or genetic factors may also influence warfarin dose. This variant rs56165452 defines CYP2C9\*4.

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# CYP2C9:\*3/\*1 N/A

Estimate the anticipated stable dose of warfarin using the algorithms available on http://www.warfarindosing.org, the IWPC Pharmacogenetic Dosing Algorithm or the FDA-approved drug label

-- Clinical Annotations ----

# · Class 1B rs9934438 GA

Patients with the AG genotype who are treated with warfarin may require a lower dose as compared to patients with the GG genotype, and a higher dose as compared to patients with the AA genotype. Other clinical and genetic factors may also influence a patients required dose of warfarin.

 $\cdot$  Class 2A rs9923231 CT

Patients with genotype CT may require shorter time to therapeutic INR when treated with warfarin as compared with patients with genotype CC. Other genetic and clinical factors may also influence the response to warfarin.

 $\cdot$  Class 2A rs9923231 CT

Patients with the CT genotype may have increased risk of over-anticoagulation when treated with warfarin as compared with patients with genotype CC. Other genetic and clinical factors may also influence the toxicity to warfarin.

 $\cdot$  Class 2B rs7196161 GA

Patients with the AG genotype may require an increased dose of warfarin as compared to patients with the GG genotype and a decreased dose of warfarin as compared to patients with the AA genotype. Other clinical and genetic factors may also influence the dose of warfarin.

· Class 3 rs17880887 GG

Patients with the GG genotype may require lower dose of warfarin as compared to patients with the GT or TT genotype. Other genetic and clinical factors may also influence warfarin dose. This variant rs17880887 is part of VKORC1 H8 and H9 haplotypes.

 $\cdot$  Class 3 rs9923231 CT

Patients with the CT genotype may spent less time in INR therapeutic range (TTR) when treated with warfarin as compared with patients with genotype CC. Contradictory findings have also been reported. Other genetic and clinical factors may also influence the response to warfarin.

· Class 3 rs9934438 GA

Patients with the AG genotype may have decreased time in the rapeutic range of INR (TTR) when treated with warfarin as compared to genotype GG. Other genetic and clinical factors may also influence the response to warfarin.

PRSS53 protease, serine, 53

- Clinical Annotations

#### · Class 1B rs9934438 GA

Patients with the AG genotype who are treated with warfarin may require a lower dose as compared to patients with the GG genotype, and a higher dose as compared to patients with the AA genotype. Other clinical and genetic factors may also influence a patients required dose of warfarin.

 $\cdot$  Class 2A rs9923231 CT

Patients with genotype CT may require shorter time to the rapeutic INR when treated with warfarin as compared with patients with genotype CC. Other genetic and clinical factors may also influence the response to warfarin.

# Class 2A rs9923231 CT

Patients with the CT genotype may have increased risk of over-anticoagulation when treated with warfarin as compared with patients with genotype CC. Other genetic and clinical factors may also influence the toxicity to warfarin.

· Class 3 rs9923231 CT

Patients with the CT genotype may spent less time in INR therapeutic range (TTR) when treated with warfarin as compared with patients with genotype CC. Contradictory findings have also been reported. Other genetic and clinical factors may also influence the response to warfarin.

· Class 3 rs9934438 GA

Patients with the AG genotype may have decreased time in the rapeutic range of INR (TTR) when treated with warfarin as compared to genotype GG. Other genetic and clinical factors may also influence the response to warfarin.

BCKDK	branched chain ketoacid dehydrogenase kinase
	- Clinical Annotations

# Class 2A rs9923231 CT

Patients with genotype CT may require shorter time to the rapeutic INR when treated with warfarin as compared with patients with genotype CC. Other genetic and clinical factors may also influence the response to warfarin.

 $\cdot$  Class 2A rs9923231 CT

Patients with the CT genotype may have increased risk of over-anticoagulation when treated with warfarin as compared with patients with genotype CC. Other genetic and clinical factors may also influence the toxicity to warfarin.

· Class 3 rs9923231 CT

Patients with the CT genotype may spent less time in INR therapeutic range (TTR) when treated with warfarin as compared with patients with genotype CC. Contradictory findings have also been reported. Other genetic and clinical factors may also influence the response to warfarin.

CYP4F2	cytochrome P450, family 4, subfamily F, polypeptide 2		

#### · Class 3 rs2108622 CC

Patients with the CC genotype may have increased international normalized ratio variability (INR-var) when treated with warfarin as compared to patients with genotype TT or CT in European-Americans after the warfarin dose-titration phase. Other genetic and clinical factors may also influence the response to warfarin.

UGT1A10	UDP glucuronosyltransferase 1 family, polypeptide A10
	- Clinical Annotations
	- Chinical Affilotations

#### · Class 3 rs887829 CT

Patients with the CT genotype and heart valve replacement may require a larger stable dose of warfarin compared to patients with the CC genotypes. Other clinical and genetic factors affect stable dose of warfarin.

	rpe and heart valve replacement may require a larger stable dose of warfaring the CC genotypes. Other clinical and genetic factors affect stable dose of
UGT1A4	UDP glucuronosyltransferase 1 family, polypeptide A4
	pe and heart valve replacement may require a larger stable dose of warfaring the CC genotypes. Other clinical and genetic factors affect stable dose of
UGT1A5	UDP glucuronosyltransferase 1 family, polypeptide A5
UGT1A6	UDP glucuronosyltransferase 1 family, polypeptide A6  ———————————————————————————————————
	ope and heart valve replacement may require a larger stable dose of warfaring the CC genotypes. Other clinical and genetic factors affect stable dose of
UGT1A7	UDP glucuronosyltransferase 1 family, polypeptide A7
· · ·	the CC genotypes. Other clinical and genetic factors affect stable dose of
UGT1A8	UDP glucuronosyltransferase 1 family, polypeptide A8

Class 3 rs887829 CT

Patients with the CT genotype and heart valve replacement may require a larger stable dose of warfarin compared to patients with the CC genotypes. Other clinical and genetic factors affect stable dose of warfarin.

UGT1A9	UDP glucuronosyltransferase 1 family, polypeptide A9
	- Clinical Annotations
Class 3 rs887829	CT

Patients with the CT genotype and heart valve replacement may require a larger stable dose of warfarin compared to patients with the CC genotypes. Other clinical and genetic factors affect stable dose of warfarin.

UGT1A1 UDP glucuronosyltransferase 1 family, polypeptide A1

— Clinical Annotations —

· Class 3 rs887829 CT

Patients with the CT genotype and heart valve replacement may require a larger stable dose of warfarin compared to patients with the CC genotypes. Other clinical and genetic factors affect stable dose of warfarin.

# ZIDOVUDINE

ABCB1	ATP-binding cassette, sub-family B (MDR/TAP), member 1 $$
	- Clinical Annotations

· Class 3 rs1045642 GG

Patients with the GG genotype and HIV may have an increased risk of virological failure when receiving highly active antiretroviral therapy (HAART), as compared to patients with the AA genotype. Other genetic and clinical factors may also influence risk of virological failure on HAART.