

NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX:

SPECIMEN DETAILS

ORDERED BY

SPECIMEN TYPE: COLLECTION DATE: **RECEIVED DATE:** REPORT DATE: 11/11/2022

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Complete Panel

Risk Management

Type III Hyperlipoproteinemia

Not Associated with Type III Hyperlipoproteinemia

The patient is positive for both the APOE c.388 T>C (Cys130Arg) and the APOE c.526 C>T (Arg176Cys) mutations. The patient's genotype is ϵ^2/ϵ^4 (frequency: 0.73-2.9%).

The APOE E2 form is associated with a slower conversion of IDL to LDL, lower plasma cholesterol, and higher triglycerides. The APOE E4 confers a limitation of HDL binding to its receptor, and is associated with increased plasma cholesterol, LDL, and triglycerides. Individuals with the APOE ε2/ε4 genotype may have higher lipid levels.

Consider dietary adjustments (very low fat diet) and lipid-lowering therapy based on lipid profiles and other risk factors.

Hyperhomocysteinemia - Depression

No Increased Risk of Hyperhomocysteinemia

The patient carries one copy of the MTHFR c.665C>T variant (heterozygous). MTHFR enzyme activity is reduced (60% of normal activity). Patients diagnosed with depression often have low folate levels and homocysteine is a highly sensitive marker of folate status. Functional folate deficiency is indicated by elevated homocysteine. The patient's small reduction in MTHFR activity is not a risk factor for hyperhomocysteinemia. Patients diagnosed with depression: as lower folate levels are associated with poorer antidepressant response, and baseline levels of folate within the normal range predict antidepressant response, testing for homocysteine levels and serum folate levels may be informative for this patient before prescribing methylfolate as an antidepressant-augmenting agent.

Thrombophilia

Increased Risk of Thrombosis

The patient carries one copy (heterozygous) of the F5 c.1601G>A variant (also known as Factor V Leiden) and one copy (heterozygous) of the F2 c.*97G>A variant (also known as Factor II 20210G>A).

The patient's risk of thrombosis is 20 times higher than average (average risk of clotting is about 1 in 1000 for anyone in a year). Other risk factors may have additive effects on thrombotic risk, increasing it further.

Anticoagulation:

Post-VTE patients with a low or moderate bleeding risk: long-term anticoagulation may be considered with periodic reevaluation to assess risks versus benefits.

Asymptomatic individual without a history of thrombosis: a short course of prophylactic anticoagulation may be considered in high-risk settings such as surgery, pregnancy, or prolonged immobilization. Decisions regarding prophylactic anticoagulation should be based on a risk/benefit assessment.

Estrogen-containing contraceptive and hormone replacement therapy: women with or without prior history of thrombotic events should avoid estrogen containing contraception and hormone replacement therapy.

Hyperhomocysteinemia - Thrombosis

No Increased Risk of Hyperhomocysteinemia

The patient carries one copy of the MTHFR c.665C>T variant (heterozygous) and does not carry the MTHFR c.1286A>C variant. MTHFR enzyme activity is reduced (60% of normal activity).

Based on results for the MTHFR c.665C>T variant, the patient has a small reduction in MTHFR activity, which is not a risk factor for

hyperhomocysteinemia. Hyperhomocysteinemia is associated with a risk for venous thromboembolism (VTE). Unless other risk factors are present, the patient is not expected to have an increased risk for VTE.

The patient's MTHFR activity is slightly reduced.



Manchester		PATIENT INFORMATION		SPECIMEN DETAILS		ORDERED BY
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A medication has potentially reduced efficacy, increased toxicity or the patient has an increased risk for the indicated condition.		ACTIONABLE	pharr (CPIC imple	macogenetic exper , DPWG, FDA. EMA ementation in a clir	t groups, cons (). Recommend	ations by international ortia or regulatory bodies lations are suitable for uidelines may change as
Guidelines exist for adjusting dosage, increased vigil the patient has a moderate risk for the indicated cor The medication can be prescribed according to stan	ndition. Idard	INFORMATIVE	There		,	findings documenting the sm or drug interaction.
regimens or the patient's risk for the indicated cond not increased.	ition is		Reco	mmendations are i ng is optional.	nformative and	d implementation in a clinical





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Potentially Impacted Medications

CATEGORY	DRUG CLASS	STANDARD PRECAUTIONS	USE WITH CAUTION	CONSIDER ALTERNATIVES
Anticancer Agents	Antifolates		Methotrexate (Trexall®)	
	Angiotensin II Receptor Antagonists	Azilsartan (Edarbi®, Edarbyclor®) Candesartan (Atacand®) Eprosartan (Teveten®) Irbesartan (Avapro®) Losartan (Cozaar®, Hyzaar®) Olmesartan (Benicar®) Telmisartan (Micardis®) Valsartan (Diovan®, Entresto®)		
	Antiarrhythmics	Amiodarone (Nexterone®, Pacerone®) Disopyramide (Norpace®) Quinidine (Quinidine®) Sotalol (Betapace®, Sorine®, Sotylize®)		
Cardiovascular	Anticoagulants	Apixaban (Eliquis®) Betrixaban (Bevyxxa®) Dabigatran Etexilate (Pradaxa®) Edoxaban (Savaysa®) Fondaparinux (Arixtra®) Rivaroxaban (Xarelto®) Warfarin (Coumadin®)		
	Antiplatelets	Prasugrel (Effient®) Ticagrelor (Brilinta®) Vorapaxar (Zontivity®)		Clopidogrel (Plavix®)
	Beta Blockers	Atenolol (Tenormin®) Bisoprolol (Zebeta®) Labetalol (Normodyne®, Trandate®)		
	Diuretics	Torsemide (Demadex®)		
	Statins		Fluvastatin (Lescol®) Pravastatin (Pravachol®) Rosuvastatin (Crestor®)	Atorvastatin (Lipitor®) Lovastatin (Mevacor®, Altoprev®, Advicor®) Pitavastatin (Livalo®) Simvastatin (Zocor®)
	Meglitinides	Nateglinide (Starlix®) Repaglinide (Prandin®, Prandimet®)		
Diabetes	Sulfonylureas	Chlorpropamide (Diabinese®) Glimepiride (Amaryl®) Glipizide (Glucotrol®) Glyburide (Micronase®) Tolbutamide (Orinase®)		
	Antiemetics	Aprepitant (Emend-oral®) Dronabinol (Marinol®) Fosaprepitant (Emend-IV®) Rolapitant (Varubi®)		



Genetic Test Results For Patient s7akg3i

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V Ur	anchest	NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX:	SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: 11/11/2022	,
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CATEGORY	DRUG CLASS	STANDARD PRECAUTIONS	USE WITH CAUTION	CONSIDER ALTERNATIVES
Gastrointestinal	Proton Pump Inhibitors	Dexlansoprazole (Dexilant ®, Kapidex®) Esomeprazole (Nexium ®) Lansoprazole (Prevacid ®) Omeprazole (Prilosec®) Pantoprazole (Protonix®) Rabeprazole (Aciphex®)		
Infections	Antifungals	Amphotericin B (AmBisome®, Abelcet®) Anidulafungin (Eraxis®) Caspofungin (Cancidas®) Fluconazole (Diflucan®) Isavuconazonium (Cresemba®) Itraconazole (Sporanox®) Micafungin (Mycamine®) Posaconazole (Noxafil®) Voriconazole (Vfend®)		
	Anti-HIV Agents	Dolutegravir (Tivicay®, Triumeq®) Doravirine (Pifeltro®) Efavirenz (Sustiva®) Etravirine (Edurant®) Raltegravir (Isentress®, Dutrebis®) Rilpivirine (Intelence®)		
	Antimalarials	Proguanil (Malarone®)		
	Fibromyalgia Agents	Milnacipran (Savella®)		
	Muscle Relaxants	Carisoprodol (Soma®) Cyclobenzaprine (Flexeril®, Amrix®) Metaxalone (Skelaxin®) Methocarbamol (Robaxin®)	Tizanidine (Zanaflex®)	
Pain	NSAIDs	Celecoxib (Celebrex®) Diclofenac (Voltaren®) Flurbiprofen (Ansaid®) Ibuprofen (Advil®, Motrin®) Indomethacin (Indocin®) Ketoprofen (Orudis®) Ketorolac (Toradol®) Meloxicam (Mobic®) Nabumetone (Relafen®) Naproxen (Aleve®) Piroxicam (Feldene®) Sulindac (Clinoril®)		



Antiender Ruber Der Statistikken d	$\langle \nabla \rangle \mathbf{M}$	anchoct	PATIENT INFORMATION	SPECIMEN DETAILS	ORDERED BY
CATEGORY DRUG CLASS STANDARD PRECAUTIONS USE WITH CAUTION CONSIDER ALTERNATIVES Affentani (Affenta®) Bupencorphine (Butrans®), Bupencorphine (Butrans®), Bupencorphine (Butrans®), Dipidids Affentani (Affenta®) Bupencorphine (Disudd®, Dipidids Fentaryl (Actiq ®) Hydrocondrew (Vicdin ®) Methadone (Dolophine®)) Suffentani (Sufenta®) Suffentani (Sufenta®) Suffentani (Sufenta®) Suffentani (Sufenta®) Suffentani (Sufenta®) Suffentani (Sufenta®) Fentaryl (Actiq ®) Hydrocondrew (Vicdin ®) Morphine (MS Contin ®) Antiaddictives Buppropion (Wellbutrin®, Zyban®, Aptenzia P(Sufenta®)) Suffentani (Sufenta®) Suffentani (Sufenta®) Destroamphetamine (Adderal®, Victor®) Clonidine (Apaya %) Destroamphetamine (Moderal®, Clonidine (Apaya) Destroamphetamine (Moderal®, Clonidine (Apaya %) Destroamphe	V Un	iversity	ACC #: s7akg3i DOB: 1/1/1900	COLLECTION DATE: RECEIVED DATE:	
Altentanii (Altenta®) Buprenorphine (Butran®), Buprenorphine (Dilaudid®, Ebago®) Fentanyl (Actiq®) Hydrocodone (Vicodin®), Merphine (Opana®), Numorphan®), Suferanal (Sufenta®), Tapentadi (Nukynna®) Antiaddictives Bupropion (Wellburin®, Zyban®, Altexane (Vivitol®, Contrave®) Pentanyl (Actiq ®) Hydrocodone (Vicodin®), Morphine (MS Contin®) Antiaddictives Bupropion (Wellburin®, Zyban®, Altexane (Vivitol®, Contrave®) Desmethylphenidate (Focalin®), Matrexane (Vivitol®, Contrave®) Anti-ADHD Agents Burapoin (Riviart®), Destroamphetamine (Desertine®), Clonidine (Kapaya®), Destroamphetamine (Vivanse®) Desmethylphenidate (Focalin®), Methylphenidate (Focalin®, Aptensin %, Concera®, Ganabache (Intuniv®), Destroamphetamine (Vivanse®) Anti-ADHD Agents Birviarcetan (Riviart®), Contare(Egretol®), Canabache (Intuniv®), Esticataregine (Riviart®), Esticataregine (Riviart®), Prenobarbital (Luminal®), Esticataregine (Riviart®), Primidane (Mysoline®), Esticataregine (Riviart®), Primidane (Mysoline®), Esticataregine (Riviart®), Prenobarbital (Luminal®), Esticataregine (Riviart®), Primidane (Riviart®), Primidane (Riviart®), Primidane (Riviart®), Primidane (Zonegine), Zonisamide (Zonegane), Xipriare (Conegine), Zonisamide (Zonegane)	FOR ACADEMIC PUR	RPOSES ONLY - NOT FOR CLINICAL U	JSE		
Peychetropi Burgenorphine (Butrans®, Burgenex®) Pertanyl (Actiq®) Burgenex®) Exalgo®) Pertanyl (Actiq®) Hydromorphone (Dilaudi®, Exalgo®) Hydrocodone (Vicodin®) Methadoue (Dolophine®)) Suffertanil (Sufenta®) Numorphan®) Suffertanil (Sufenta®) Numorphan®) Suffertanil (Sufenta®) Numorphan®) Suffertanil (Sufenta®) Antiaddictives Burgepion (Wellburtin®, Zyban %, Aplecizin®, Contraxe®) Numerophite (Marcane) Desmethylphenidate (Ritalin®, Aplecizin®, Contraxe®) Anti-ADHD Agenta Burgepion (Wellburtin®, Zyban %, Aplecizin®, Contraxe®) Evekce®) Desmethylphenidate (Ritalin®, Aplecizin®, Contraxe®) Canabadio (Intuniv%) Burgepion (Wellburtin®, Zyban %, Aplecizin®, Contraxe®) Evekce®) Contrate (Ritalin®, Contraxe®) Canabadio (Intuniv%) Burgepion (Wellburtin®, Zyban %, Aplecizin®, Contraxe®) Evekce®) Contrate (Riverave) Canabadio (Intuniv%) Burgepion (Wellburtin®, Zyban %, Aplecizin®, Contraxe®) Evekce®) Contrate (Riverave) Canabadio (Intuniv%) Burgepion (Wellburtin®, Zyban %, Aplecizin®, Canabadio (Intuniv®) Evekce®) Canabadio (Intuniv%) Laccosmic (Wot	CATEGORY	DRUG CLASS	STANDARD PRECAUTIONS	USE WITH CAUTION	CONSIDER ALTERNATIVES
Antiaddictives Aplenzin®, Contrave®) Naltrexone (Vivitol®, Contrave®) Ampletamine (Adderal®, Evekeo®) Dexmethylphenidate (Focalin®) Methylphenidate (Rialin®, Aptensio XR®, Concerta®, Guanfacine (Intuniv®) Lisdexamfetamine (Vyvanse®) Brivaracetam (Briviact®) Carbanabidol (Epidolex®) Carbanabidol (Epidolex®) Carbanabidol (Epidolex®) Carbanazepine (Fegretol®, Carbarazepine (Fegretol®, Carbarazepine (Portga®) Felbarate (Felbatol®) Esticarbazepine (Aptensio XR®, Concerta®, Metadate ER®, Quillivant ER®) Anticonvulsants Brivaracetam (Briviact®) Carbarazepine (Fegretol®, Carbarazepine (Portga®) Felbarate (Felbatol®)) Ethosuximide (Zarontin®) Etosumide (Zarontin®) Etacosmide (Vimpa®) Prempontin (Cerebyx®) Gabapentin (Neurontin®) Lacosmide (Vimpa®) Prempontin (Cerebyx®) Gabapentin (Neurontin®) Lacosmide (Vimpa®) Prempontin (Cinetols Netwy) Prempontin (Dilantin®) Prempontin (Dilantin®) Prempontin (Dilantin®) Pregabalin (Lyrica®) Rufinamide (Banzel®) Tigabatin (Gabitril®) Topiramate (Topamax®) Vigabatrin (Sabril®) Psychotropic Vigabatrin (Sabril®)		Opioids	Buprenorphine (Butrans®, Buprenex®) Hydromorphone (Dilaudid®, Exalgo®) Levorphanol (Levo Dromoran®) Meperidine (Demerol®) Methadone (Dolophine®) Oxymorphone (Opana®, Numorphan®) Sufentanil (Sufenta®)	Hydrocodone (Vicodin®)	
Anti-ADHD Agents Evekeo®) Clonidine (Kapvay®) Dextromphetamine (Dexedrate) Guanfacine (Intuniv®) Lisdexamfetamine (Vyvanse®) Dexmethylphenidate (Focalin®) Methylphenidate (Ritalin®, Aptensio X8*, Concerta®, Metadate ER®, Quillivant ER®) Brivaracetam (Briviact®) Carbamazepine (Tegretol®, Carbamazepine (Tegretol®, Carbamazepine (Tegretol®, Carbatrol®, Epitol®) Esticarbazepine (Potiga®) Febbanete (Fotiga®) Febbanete (Fotiga®) Febbanete (Fotiga®) Eascarbazepine (Trigretol®, Carbatrol®, Epitol®) Estosuximide (Zarontin®) Exogabine (Potiga®) Febbanete (Fotiga®) Febbanete (Fotiga®) Febbanete (Fotiga®) Eascarbazepine (Trigretal) Carobarbite (Lamotrigine (Lamotri		Antiaddictives	Aplenzin [®] , Contrave [®])		
Psychotropic Canabidiol (Epidiolex®) Vigabatrin (Sabril®) Carbamazepine (Tegretol®, Carbamazepine (Aption®) Eslicarbazepine (Aption®) Eslicarbazepine (Aption®) Ethosuximide (Zarontin®) Etogabine (Potiga®) Felbamate (Felbatol®) Felbamate (Felbatol®) Gabapentin (Neurontin®) Phenobarbital (Luminal®) Lacosamide (Vimpat®) Primidone (Mysoline®) Levetiracetam (Keppra®) Zonisamide (Zonegra®) Oxcarbazepine (Trileptal®, Oxtellar XR®) XR®) Perampanel (Fycompa®) Phenytoin (Dilantin®) Phenytoin (Dilantin®) Pregabalin (Lyrica®) Rufinamide (Banzel®) Topiramate (Topamax®) Valproic Acid (Depakene®) Valproic Acid (Depakene®)		Anti-ADHD Agents	Evekeo®) Clonidine (Kapvay®) Dextroamphetamine (Dexedrine®) Guanfacine (Intuniv®)	Methylphenidate (Ritalin®, Aptensio XR®, Concerta®,	
Antidementia Agents Memantine (Namenda®)	Psychotropic	Anticonvulsants	Cannabidiol (Epidiolex®) Carbamazepine (Tegretol®, Carbatrol®, Epitol®) Eslicarbazepine (Aptiom®) Ethosuximide (Zarontin®) Ezogabine (Potiga®) Felbamate (Felbatol®) Fosphenytoin (Cerebyx®) Gabapentin (Neurontin®) Lacosamide (Vimpat®) Lamotrigine (Lamictal®) Levetiracetam (Keppra®) Oxcarbazepine (Trileptal®, Oxtellar XR®) Perampanel (Fycompa®) Phenytoin (Dilantin®) Pregabalin (Lyrica®) Rufinamide (Banzel®) Tiagabine (Gabitril®) Topiramate (Topamax®) Valproic Acid (Depakene®)	Primidone (Mysoline®)	
		Antidementia Agents			



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CATEGORY	DRUG CLASS	STANDARD PRECAUTIONS	USE WITH CAUTION	CONSIDER ALTERNATIVES
	Antidepressants	Amitriptyline (Elavil®) Citalopram (Celexa®) Clomipramine (Anafranil®) Doxepin (Silenor®) Duloxetine (Cymbalta®) Escitalopram (Lexapro®) Imipramine (Tofranil®) Levomilnacipran (Fetzima®) Mirtazapine (Remeron®) Sertraline (Zoloft®) Trazodone (Oleptro®) Trimipramine (Surmontil®) Vilazodone (Viibryd®)		
	Antipsychotics	Asenapine (Saphris®) Cariprazine (Vraylar®) Fluphenazine (Prolixin®) Loxapine (Loxitane®, Adasuve®) Lurasidone (Latuda®) Pimavanserin (Nuplazid®) Quetiapine (Seroquel®) Thiothixene (Navane®) Trifluoperazine (Stelazine®) Ziprasidone (Geodon®)	Clozapine (Clozaril®) Olanzapine (Zyprexa®)	
	Benzodiazepines	Alprazolam (Xanax®) Clonazepam (Klonopin®) Diazepam (Valium®)	Clobazam (Onfi®)	
	Other Neurological Agents	Flibanserin (Addyi®)		
Rheumatology	Anti-Hyperuricemics and Anti-Gout Agents	Colchicine (Mitigare®) Febuxostat (Uloric®)		
	Immunomodulators	Apremilast (Otezla®) Tofacitinib (Xeljanz®)	Leflunomide (Arava®)	
Transplantation	Immunosuppressants	Tacrolimus (Prograf®)		
	5-Alpha Reductase Inhibitors for Benign Prostatic Hyperplasia	Dutasteride (Avodart®) Finasteride (Proscar®)		
Urologicals	Alpha-Blockers for Benign Prostatic Hyperplasia	Alfuzosin (UroXatral®) Doxazosin (Cardura®) Silodosin (Rapaflo®) Terazosin (Hytrin®)		
Unorogicals	Antispasmodics for Overactive Bladder	Oxybutynin (Ditropan®) Solifenacin (Vesicare®) Trospium (Sanctura®)		
	Phosphodiesterase Inhibitors for Erectile Dysfunction	Avanafil (Stendra®) Sildenafil (Viagra®) Tadalafil (Cialis®) Vardenafil (Levitra®)		





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Dosing Guidance

-		
t orvastatin pitor®	Increased Atorvastatin Exposure (SLCO1B1: Decreased Function) The patient's genotype is associated with possible increased atorvastatin exposure. Patients may be at ar	ACTIONABLE increased
	myopathy risk.	
	Consider starting atorvastatin at doses ≤40 mg. If doses >40 mg are needed, consider combination thera atorvastatin plus a non-statin guideline directed therapy).	apy (e.g.,
lopidogrel	Reduced Exposure to Clopidogrel Active Metabolite (CYP2C19: Intermediate Metabolizer)	ACTIONABLE
lavix®	The patient's genotype is associated with possible decreased clopidogrel exposure. Patients may be at a for adverse cardiac and cerebrovascular events.	n increased risk
	ACS, PCI, and Neurovascular Indications: Consider an alternative such as prasugrel (contraindicated in TIA/stroke) or ticagrelor. In patients with AC clopidogrel cannot be avoided, alternative dosing strategies exist and may be considered.	CS or PCI, if
ovastatin	Increased Lovastatin Exposure (SLCO1B1: Decreased Function)	ACTIONABLE
levacor®, Altoprev®,	The patient's genotype is associated with possible increased lovastatin exposure. Patients may be at an in	ncreased
dvicor®	myopathy risk. Consider an alternative statin based on disease-specific guidelines. If lovastatin use is warranted, conside ≤20 mg per day.	r limiting dose to
itavastatin	Increased Pitavastatin Exposure (SLCO1B1: Decreased Function)	ACTIONABLE
valo®	The patient's genotype is associated with possible increased pitavastatin exposure. Patients may be at ar myopathy risk with doses >1 mg per day.	increased
	Consider starting pitavastatin at doses ≤ 2 mg. If doses > 2 mg are needed, consider an alternative statin therapy (e.g., pitavastatin plus a non-statin guideline directed medical therapy).	or combination
imvastatin	Increased Simvastatin Exposure (SLCO1B1: Decreased Function)	ACTIONABLE
ocor®	The patient's genotype is associated with possible increased simvastatin exposure. Patients may be at an myopathy risk with doses >20 mg.	increased
	Consider an alternative statin. If simvastatin use is warranted, consider limiting dose to <20 mg.	
lobazam	Possible Sensitivity to Clobazam (CYP2C19: Intermediate Metabolizer)	ACTIONABLE
nfi®	In CYP2C19 intermediate metabolizers, plasma levels of the active metabolite N-desmethylclobazam wer than those found in CYP2C19 normal metabolizers. The dose adjustment for intermediate metabolizers is established, and therefore the recommendation for poor metabolizers is proposed. The starting dose sho mg/day, and dose titration should proceed slowly according to weight. Patients should be titrated initiall (≤30 kg body weight) or 20 mg/day (>30 kg body weight). If necessary and based upon clinical response titration to the maximum doses 20 mg/day (≤30 kg body weight) or 40 mg/day (>30 kg body weight) m day 21.	s not well ould be 5 y to 10 mg /day y, an additional
lozapine	Non-Response to Clozapine (CYP1A2: Normal Metabolizer - Higher Inducibility)	INFORMATIVE
lozaril ®	between high clozapine doses and the risk of seizures, and therefore careful monitoring is recommended	during dosing
-		Dine Non-Response to Clozapine (CYP1A2: Normal Metabolizer - Higher Inducibility) ® Smokers have a high risk for non-response at standard doses and may require higher doses. There is an between high clozapine doses and the risk of seizures, and therefore careful monitoring is recommended adjustment. Smoking cessation will increase plasma drug levels, leading to adverse events. Therefore, the



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V	Univers	sity	NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX:	SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: 11/11/20)22
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<u> </u>	Dexmethylphenid ate	Decreased Resp	onse to Dexmethylphenidat	e (COMT: Intermediate COM	T Activity) INFORMATIVE
	Focalin ®			al response to dexmethylphenida t. Therapy should be initiated in s	te. Dosage should be individualized mall doses, with gradual weekly
	Fentanyl	Altered Respons	e to Fentanyl (OPRM1: Alter	ed OPRM1 Function)	INFORMATIVE
	Actiq®	the patient's genot the patient may re	type has been shown to be assoc quire higher doses of this drug. I	iated with reduced analgesia at s	Acute postoperative and cancer pain: standard fentanyl doses. Therefore, erapeutic window, it is advised to minimal side effects.
<u>^</u>	Fluvastatin	Increased Fluvas Metabolizer)	statin Exposure (SLCO1B1: De	ecreased Function; CYP2C9: N	Normal ACTIONABLE
	Lescol®	The patient's geno	ommended dosage and adminis	increased fluvastatin exposure. Fl tration, but patients may be at an	uvastatin can be prescribed at i increased risk for myopathy with
	Hydrocodone	Altered Respons	e to Hydrocodone (OPRM1:	Altered OPRM1 Function)	INFORMATIVE
	Vicodin®	genotype has beer	n shown to be associated with re	>G variant. Acute postoperative a duced analgesia and increased o to increased hydrocodone doses,	pioid side effects at standard or high
<u>^</u>	Leflunomide	•		9: Intermediate Metabolizer)	
	Arava®	that patients with on hepatotoxicity. The	decreased CYP2C19 activity have	a higher risk of developing gasti e dose adjustment. If leflunomide	nomide. Preliminary studies indicate rointestinal side effects and e is prescribed at standard dosing,
		treatment, and eve		neters should be checked no mor s of therapy. Blood pressure shou	e than 6 months before beginning Ild be checked before beginning
	Methotrexate	Increased Risk fo	or Methotrexate Toxicity (M	THFR: Reduced MTHFR Activ	vity) INFORMATIVE
	Trexall®	Leukemia or lymp likelihood of treatr and adjust the dos response to metho between individual patients. However, effects and adjust t	homa patients who are treated we nent interruptions due to metho e accordingly. Other genetic and trexate treatment. Nonmaligna Is carrying the MTHFR c.665C>T there is insufficient data to calcu	l clinical factors may also influence nt conditions: a limited number variant and methotrexate-induce ulate dose adjustment. Monitor p	nens might have an increased ent closely for increased side effects the patient's risk for toxicity and of studies found an association ed toxicity in rheumatoid arthritis
	Methylphenidate	Decreased Resp	onse to Methylphenidate (C	OMT: Intermediate COMT Ad	ctivity) INFORMATIVE
	Ritalin [®] , Aptensio XR [®] , Concerta [®] , Metadate ER [®] , Quillivant ER [®]			al response to methylphenidate. I t. Therapy should be initiated in s	
<u>^</u>	Morphine	Altered Respons	e to Morphine (OPRM1: Alte	ered OPRM1 Function)	INFORMATIVE
	owered By				
	ranslational		Genetic Test Results For Patie	nt s7akg3i	Page 8 of 58

		hostor	PATIENT INFORMATION	SPECIMEN DETAILS	ORDERED BY
V	Manc Univer	rsity	NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX:	SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: 11/11/20)22
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	MS Contin®	genotype has be nausea and vom	iting during the first 24-hour posto sing regimen needs to be individu	duced analgesia at standard mor operative period. Therefore, the p	phine doses and decreased risk for atient may require higher doses of
	Olanzapine	Non-Response	e to Olanzapine (CYP1A2: Norr	nal Metabolizer - Higher Ind	ucibility) INFORMATIV
<u> </u>	Zyprexa ®	There is little evid for non-response may increase pla	dence regarding the impact of CYF	P1A2 genetic variants on olanzapi oring is recommended during do e events. Therefore, therapeutic d	ne response. Smokers may be at risk sing adjustment. Smoking cessation
Ŷ	Phenobarbital	Possible Sensit	tivity to Phenobarbital (CYP2C	19: Intermediate Metabolize	r) INFORMATIV
_	Luminal®	lower clearance of with this antiepil		abolizers, no significant changes i al can be prescribed at standard	19 intermediate metabolizers have a n clinical outcome has been reported label-recommended dosage and
<u>^</u>	Pravastatin	Increased Prav	astatin Exposure (SLCO1B1: De	ecreased Function)	ACTIONABL
	Pravachol®		-		ravastatin can be prescribed at increased myopathy risk with doses
Ŷ	Primidone	Possible Sensit	tivity to Primidone (CYP2C19:	Intermediate Metabolizer)	INFORMATIV
	Mysoline ®	lower clearance of has been reported	•	e) than normal metabolizers, no s refore, primidone can be prescrib	intermediate metabolizers have a ignificant changes in clinical outcome ed at standard label-recommended
Ŷ	Rosuvastatin	Increased Rosu	uvastatin Exposure (SLCO1B1:	Decreased Function)	ACTIONABL
	Crestor [®]		notype is associated with possible ecommended dosage and administ		Rosuvastatin can be prescribed at increased myopathy risk with doses
<u>î</u>	Tizanidine	Possible Non-I Inducibility)	Response to Tizanidine (CYP1/	A2: Normal Metabolizer - Hig	jher INFORMATIV
	Zanaflex®	There is little evid for non-response and the risk of hy adjustment. Smo	e and may require higher doses. The provide the provided	nere is an association between hig . Therefore, careful monitoring is na drug levels, leading to excessiv	e hypotension and sedation. Careful
<u>^</u>	Zonisamide	Possible Sensit	tivity to Zonisamide (CYP2C19	: Intermediate Metabolizer)	INFORMATIV
	Zonegran®	intermediate me change in the cli	y involved in the metabolism of zo tabolizers have a slightly lower (15 nical outcome has been reported v commended dosage and administ	%) zonisamide clearance than no vith this antiepileptic drug. There	rmal metabolizers, no significant fore, zonisamide can be prescribed at
	Alfentanil Alfenta®	Normal Respo	nse to Alfentanil		INFORMATIV
P	rowered By		Genetic Test Results For Patie	nt s7akg3i	

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NAME: Patient s7akg3i

SPECIMEN DETAILS

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S S	oftware	FOR ACADEMIC PURPOSES ONLY - DO NOT DISTRIBUTE - NOT FOR CLINICAL USE	Page 10 of 5
	Powered By	Genetic Test Results For Patient s7akg3i	
√	Anidulafungin Eraxis®	Normal Response to Anidulafungin	ACTIONABL
	AmBisome®, Abelcet®	Pharmacogenetic guidance: Amphotericin B is excreted very slowly (over weeks to months) by the of a given dose being excreted in the biologically active form. Details of possible metabolic pathway genetically guided drug selection or dosing recommendations are available. Polypharmacy guidan medications such as aminoglycosides, cyclosporine, and pentamidine may enhance the potential for induced renal toxicity, and should be used concomitantly only with great caution. Intensive monitori is recommended in patients requiring any combination of nephrotoxic medications.	vs are unknown. No nce: Nephrotoxic r amphotericin B-
√	Amphotericin B	Normal Response to Amphotericin B	ACTIONABL
√	Amphetamine Adderall®, Evekeo®	Good Response to Amphetamine salts (COMT: Intermediate COMT Activity) The patient's genotype result predicts a favorable response to amphetamine stimulants. Amphetami administered at the lowest effective dose, and dosage should be individually adjusted.	INFORMATIV
		Neuropathic Pain: Amitriptyline therapy can be prescribed according to standard recommended de administration.	osage and
		Psychiatric Conditions: Amitriptyline therapy can be prescribed according to standard recommend administration. Consider therapeutic drug monitoring to guide dose adjustments.	led dosage and
\checkmark	Amitriptyline Elavil®	Normal Amitriptyline Exposure (CYP2C19: Intermediate Metabolizer) The patient's reduced CYP2C19 activity is unlikely to result in increased amitriptyline exposure.	ACTIONABL
V	Amiodarone Nexterone®, Pacerone®	Normal Exposure to Amiodarone Pharmacogenetic guidance: Amiodarone is metabolized to N-desethylamiodarone. This process is by CYP3A. No genetically guided drug selection or dosing adjustments are recommended. Polypha administration of amiodarone with drugs that are, a strong inducer or inhibitor of CYP3A may affect In addition, co-administration of amiodarone with drugs known to prolong QT interval can precipita QT syndrome.	rmacy guidance: Co drug plasma levels.
		which results in a loss of efficacy.	
		guidance: The concomitant use of alprazolam with CYP3A4 inhibitors may result in increased alpraz prolonged sedation. Impairment of motor skills are also observed with some combinations. Monitor exaggerated sedative effects. If possible, alprazolam should be avoided in patients receiving strong such as ketoconazole, itraconazole and ritonavir. Drugs that induce CYP3A enzymes may decrease a	patients for patients of CYP3A4
V	Alprazolam Xanax®	Normal Response to Alprazolam Pharmacogenetic guidance: Alprazolam is primarily eliminated by metabolism via CYP3A4 and CYP polymorphisms of these genes are not expected to affect the efficacy or safety profiles of this drug.	P3A5. Genetic Polypharmacy
		drug levels may increase.	INFORMATIV
	UroXatral®	Pharmacogenetic guidance: No genetically-guided drug selection or dosing recommendations are Polypharmacy guidance: Alfuzosin is extensively metabolized by CYP3A4 into pharmacologically ir Alfuzosin is contraindicated with strong CYP3A4 inhibitors, as the risk for QTc prolongation in increased at higher concentrations. Take caution when this drug is prescribed with CYP3A4 mode	nactive metabolites. duced by this drug i
	Alfuzosin	Normal Response to Alfuzosin	INFORMATIV
		Pharmacogenetic guidance : alfentanil is primarily metabolized by CYP3A4 and CYP3A5. Studies in showed that CYP3A5 genotype had no effect on the systemic or apparent oral clearances, or pharma alfentanil. Polypharmacy guidance: Alfentanil should be used with caution when prescribed to pat inhibitors or inducers.	acodynamics of
	FOR ACADEMIC PURPOSES ONLY - NOT	SEX: REPORT DATE: 11/11/2022 FOR CLINICAL USE 11/11/2022 11/11/2022	
	Univer	ACC #: s7akg3i COLLECTION DATE: DOB: 1/1/1900 RECEIVED DATE:	



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NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX:

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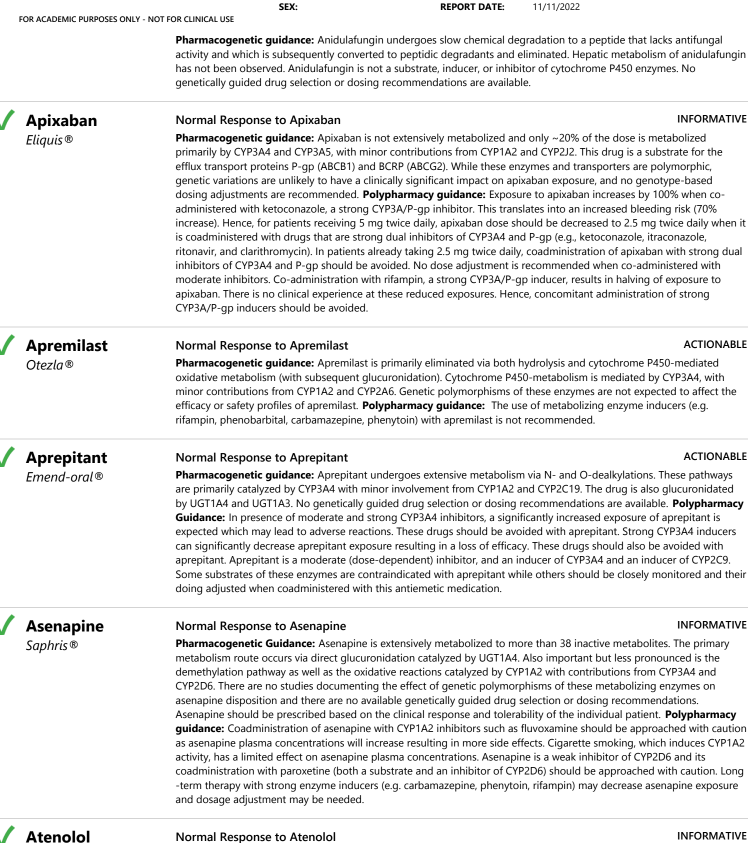
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Translational

Tenormin[®]

Genetic Test Results For Patient s7akg3i

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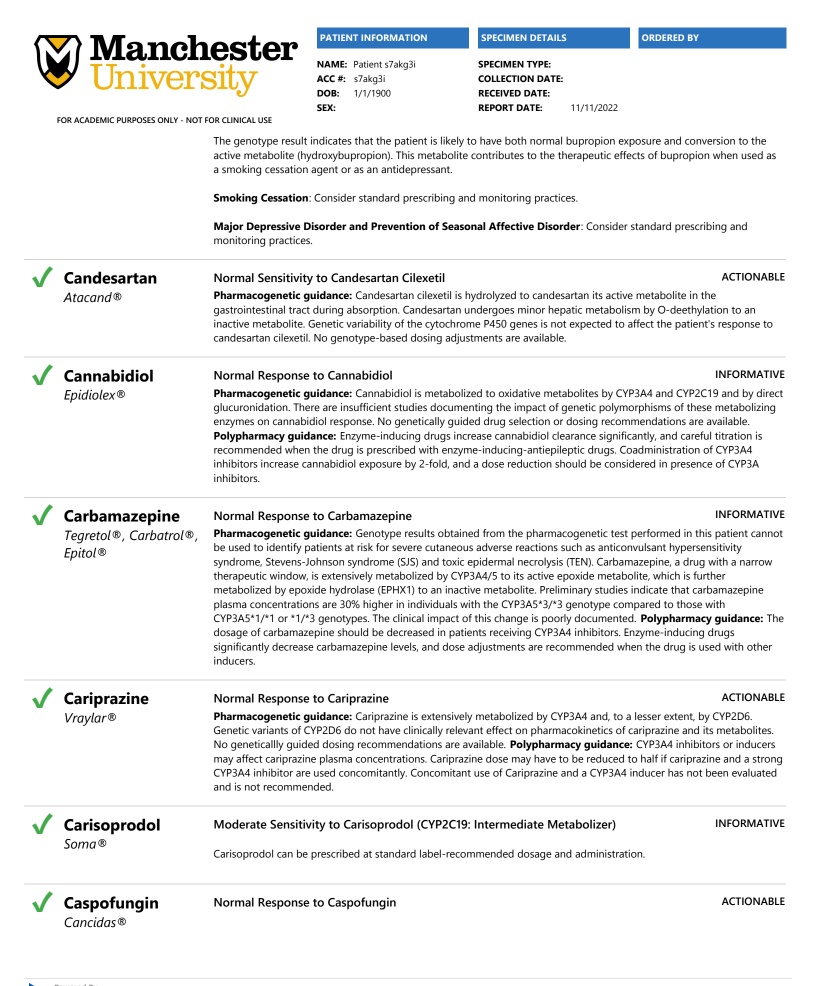
NAME: Patient s7akg3i 1/1/1900

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SPECIMEN TYPE: COLLECTION DATE: ACC #: s7akg3i DOB: **RECEIVED DATE:** SEX: REPORT DATE: 11/11/2022 FOR ACADEMIC PURPOSES ONLY - NOT FOR CLINICAL USE Pharmacogenetic guidance: The bioavailability of atenolol is approximately 40-50% and renal excretion eliminates approximately 90% of the absorbed drug in its unchanged form. A negligible amount of the drug is metabolized. Atenolol is a substrate of several organic anion and cation transporters including SLC22A1, SLC22A2, SLC47A1, and SLC47A2. No genetically-guided drug selection or dosing recommendations are available. Avanafil INFORMATIVE Normal Response to Avanafil Stendra® Pharmacogenetic guidance: no genetically guided drug selection or dosing recommendations are available. Polypharmacy guidance: Avanafil is extensively metabolized by CYP3A4, therefore Avanafil should not be used with strong CYP3A4 inhibitors such as ketoconazole, itraconazole, voriconazole, ritonavir, atazanavir, clarithromycin, indinavir, itraconazole, nefazodone, nelfinavir, saquinavir, and telithromycin. If taking a moderate CYP3A4 inhibitor, such as erythromycin, amprenavir, aprepitant, diltiazem, fluconazole, fosamprenavir, or verapamil, the dose should be no more than 50 mg in a 24-hour period. Inducers of CYP3A4 may decrease the concentrations of avanafil. Azilsartan INFORMATIVE Normal Azilsartan Exposure (CYP2C9: Normal Metabolizer) Edarbi[®], Edarbyclor[®] Azilsartan medoxomil is hydrolyzed to azilsartan, its active metabolite, in the gastrointestinal tract during absorption. Azilsartan is further metabolized to inactive metabolites by CYP2C9. Consider standard label-recommended dosage and administration ACTIONABLE **Betrixaban** Normal Response to Betrixaban Pharmacogenetic guidance: The predominant metabolic pathway of betrixaban is amide hydrolysis with minor Bevyxxa® cytochrome P450 enzymes-based metabolism (less than 1% of the drug is metabolized by CYP1A1, CYP1A2, CYP2B6, CYP2C9, CYP2C19, CYP2D6 and CYP3A4). The main elimination pathway of the drugs is biliary excretion followed by urinary excretion. Betrixaban is a substrate for the efflux transport protein P-gp (ABCB1) and while this transporter is polymorphic, genetic variations are unlikely to have a clinically significant impact on betrixaban exposure, and no genotype-based dosing adjustments are available. Polypharmacy guidance: Concomitant use with P-gp inhibitors such as amiodarone, azithromycin, verapamil, ketoconazole, clarithromycin results in increased plasma levels of betrixaban and increased risk of bleeding. Dosing reduction and close monitoring are recommended in presence of P-gp inhibitors. **Bisoprolol** INFORMATIVE Normal Response to Bisoprolol Zebeta® Pharmacogenetic guidance: Bisoprolol is eliminated by renal and non-renal pathways with 50% of the total dose being metabolized in the liver and 50% being excreted via the kidneys unchanged. Bisoprolol is predominantly metabolized by CYP3A4 with smaller contribution from CYP2D6. Limited studies suggest that bisoprolol plasma concentrations and its beta-adrenergic inhibition are not affected by CYP2D6 genetic variability. No genetically-guided drug selection or dosing recommendations are available. Brivaracetam ACTIONABLE Normal Sensitivity to Brivaracetam (CYP2C19: Intermediate Metabolizer) **Briviact**® Brivaracetam is primarily metabolized by hydrolysis and to a minor extent by hydroxylation, which is mediated by CYP2C19. In CYP2C19 intermediate metabolizers, the plasma concentration of brivaracetam is increased by 22%, but this change is not clinically significant. Brivaracetam can be prescribed at the standard label recommended dosage. INFORMATIVE **Buprenorphine** Normal Response to Buprenorphine Pharmacogenetic guidance: no genetically guided drug selection or dosing recommendations are available. Butrans[®], Buprenex[®] Buprenorphine is primarily metabolized by CYP3A4 to norbuprenorphine and by UGT enzymes (mainly UGT1A1 and 2B7). The effects of genetic variants in these enzymes on its response have not been studied. Polypharmacy guidance: The concomitant use of buprenorphine with all CYP3A4 inhibitors may result in an increase in the drug levels, which could increase or prolong adverse drug effects. Monitor patients receiving buprenorphine with a CYP3A4 inhibitor. CYP and UGT inducers may decrease buprenorphine levels. **Bupropion** INFORMATIVE Normal Bupropion Exposure (CYP2B6: Normal Metabolizer) Wellbutrin[®], Zyban[®], Aplenzin[®], Contrave[®]







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		Pharmacogenetic guidance: Caspofungin i undergoes also spontaneous chemical degra dominant mechanism influencing plasma cle are available. Polypharmacy guidance: Co- rifampin, efavirenz, nevirapine, phenytoin, or caspofungin concentrations which may requ	adation. Distribution, rather th arance. No genetically guide administration of caspofungi carbamazepine) may result i	han excretion or biotransforma d drug selection or dosing rec in with metabolizing enzyme ii	ation, is the commendations nducers (e.g.,
\checkmark	Celecoxib	Normal Celecoxib Exposure (CYP2C9:	Normal Metabolizer)		ACTIONABLE
	Celebrex®	Celecoxib therapy can be initiated at standar	d label-recommended dosag	ge and administration.	
		Consider initiating treatment at the lowest e warranted when celecoxib is administered w			stment may be
		Osteoarthritis, Rheumatoid Arthritis, Ank the lowest effective dosage for the shortest			: Consider using
		Acute Migraine: Consider using for the few	est number of days per mont	th, as needed.	
		Osteoarthritis and Hypertension (co-form the shortest duration consistent with the pat	-	Consider using the lowest effe	ctive dosage for
	Chlorpropamide	Normal Exposure to Chlorpropamide			INFORMATIVE
	Diabinese [®]	Pharmacogenetic guidance: Chlorpropami While this clearance pathway is diminished in to be clinically significant. No genetically gui guidance: Co-administration of chlorpropar chlorpropamide concentrations possibly lead CYP2C19 inducers may result in lower chlorp	n subjects with reduced CYP2 ded drug selection or dosing nide with a strong CYP2C9 ar ding to hypoglycemia. Co-adu	2C9 activity, such a change has g recommendations are availab nd/or CYP2C19 inhibitors may ministration with a strong CYP	not been shown ble. Polypharmacy result in higher
\checkmark	Citalopram Celexa®	Normal sensitivity to Citalopram (CYP2	2C19: Intermediate Metab	oolizer)	ACTIONABLE
	Celexa	Citalopram can be prescribed at standard lal	pel-recommended dosage an	nd administration.	
	Clomipramine	Normal Clomipramine Exposure (CYP2	C19: Intermediate Metab	olizer)	ACTIONABLE
	Anafranil®	The patient's reduced CYP2C19 activity is un	likely to result in increased cl	lomipramine exposure.	
		Psychiatric Conditions: Clomipramine thera administration. Consider therapeutic drug m			d dosage and
	Clonazepam	Normal Response to Clonazepam			INFORMATIVE
	Klonopin®	Pharmacogenetic guidance: No genetically Polypharmacy guidance: clonazepam is ex acetylated by N-acetyltransferases. This drug inducers.	tensively metabolized by CYP	P3A4 to an amino metabolite t	hat is further
✓	Clonidine Kapvay®	Normal Exposure to Clonidine			INFORMATIVE



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I DATE: 11/11/2022

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Pharmacogenetic guidance: Clonidine is metabolized by CYP2D6 along with CYP3A4 and CYP1A2. About 40-60% of the dose is excreted in urine as unchanged drug. Preliminary studies indicate that individuals lacking CYP2D6 activity, have increased clonidine exposure compared to subjects with normal CYP2D6 activity. The clinical relevance of this changed is not well understood and there is insufficient data to calculate dose adjustments. Other preliminary studies indicate that individuals with high CYP2D6 activity (pregnant women), have decreased clonidine exposure and may require higher doses to reach target therapeutic plasma concentrations and respond to therapy. No genetically guided drug selection or dosing adjustments are recommended. **Polypharmacy guidance**: Co-administration of clonidine with inhibitors of CYP2D6 or CYP3A4 may cause an increase in clonidine plasma concentrations while the co-administration with CYP3A4 inducers may cause a decrease in clonidine plasma concentrations. Caution should be used when co-administering drugs that can affect renal function.

Normal Response to Colchicine

Colchicine *Mitigare*®

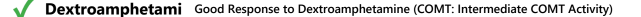
Pharmacogenetic guidance: Colchicine in eliminated both by renal excretion and metabolism. While 50% of the absorbed dose is eliminated unchanged in urine, less than 20% is metabolized by CYP3A4. Glucuronidation is also a metabolic pathway for colchicine. Colchicine is a substrate of P-glycoprotein (encoded by ABCB1 gene) and its efflux by this transporter is important in its disposition. Colchicine has a narrow therapeutic index. Preliminary and limited studies indicate a lack of an effect of CYP3A4 or ABCB1 genetic polymorphisms on clinical response to colchicine in individuals with familial Mediterranean fever (FMF). There are no available genetically-guided drug selection or dosing recommendations. **Polypharmacy guidance:** Because colchicine is a substrate for both the CYP3A4 metabolizing enzyme and the P-glycoprotein efflux transporter, inhibition of either of these pathways may lead to colchicine-related toxicity. Inhibition of both CYP3A4 and P-gp by dual inhibitors such as clarithromycin has been reported to produce life-threatening or fatal colchicine toxicity due to significant increases in systemic colchicine levels. Therefore, concomitant use of colchicine and inhibitors of CYP3A4 or P-glycoprotein should be avoided.

INFORMATIVE Cyclobenzaprine Normal Response to Cyclobenzaprine Pharmacogenetic guidance: No genetically guided drug selection or dosing recommendations are available. Flexeril®, Amrix® Cyclobenzaprine is excreted primarily as a glucuronide via the kidneys, and as an N-demethylated metabolite by CYP3A4, CYP1A2, and to a lesser extent CYP2D6. Due to the minor involvement of CYP2D6 in the metabolism of cyclobenzaprine, the polymorphism of this enzyme is not of concern in its the clinical use. Dabigatran INFORMATIVE Normal Response to Dabigatran Etexilate Pradaxa® Pharmacogenetic guidance: Dabigatran is eliminated primarily unchanged by the kidneys. After oral administration, dabigatran etexilate is converted to its active form dabigatran by esterases. A small portion (20%) of dabigatran dose is also conjugated to form pharmacologically active acyl glucuronides. Dabigatran is not a substrate, inhibitor, or inducer of

CYP450 enzymes. Dabigatran etexilate is a substrate of the efflux transporter P-gp (ABCB1). Common genetic polymorphism of the ABCB1 gene (2677G>T/A and 3435 C>T) do not appear to affect dabigatran exposure. **Polypharmacy guidance:** <u>1-Reduction in Risk of Stroke and Systemic Embolism in Non-valvular AF</u>: In patients with moderate renal impairment (CrCl 30-50 mL/min), concomitant use of the P-gp inhibitor dronedarone or systemic ketoconazole can be expected to produce dabigatran exposure similar to that observed in severe renal impairment. Consider reducing the dose of dabigatran to 75 mg twice daily. Dose adjustment is not necessary when coadministered with other P-gp inhibitors. In patients with CrCl<30 mL/min, avoid use of concomitant P-gp inhibitors with dabigatran. <u>2-Treatment of DVT and PE Reduction in the Risk of Recurrence of DVT and PE</u>: Avoid use of concomitant P-gp inhibitors with CrCl <50 mL/min.

Dexlansoprazole Dexilant[®], Kapidex[®] Increased Exposure to Dexlansoprazole (CYP2C19: Intermediate Metabolizer)

The patient's genotype may be associated with a slightly increased dexlansoprazole exposure following standard dosing. Consider prescribing dexlansoprazole at standard label-recommended dosage and administration. Once efficacy is achieved, in the setting of chronic PPI therapy (beyond 12 weeks), consider a 50% reduction in the daily dose to minimize the risk of adverse events from prolonged acid suppression.



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	Dexedrine®		type result predicts a favorable re e lowest effective dose, and dosa		-	netamine should be
√	Diazepam Valium®		tivity to Diazepam (CYP2C19:			INFORMATIV
		Diazepam can be p	prescribed at standard label-reco	mmended dosage and	administration.	
\checkmark	Diclofenac	Normal Diclofen	ac Exposure			INFORMATIV
	Voltaren ®	50% of diclofenaci CYP2C8, CYP2C19 drug is also directly affect the response Polypharmacy gu toxicity of whereas	guidance: Diclofenac is extensive is eliminated as a 4-hydroxymeta and CYP3A4 are also involved in y glucuronidated by UGT2B7 and to diclofenac. No dosing recom idance: Co-administration of dic co-administration with CYP2C9 is warranted when diclofenac is ac	bolite, a reaction media the formation of a 5-hy UGT2B4. Genetic poly mendations or genetica lofenac with CYP2C9 in nducers may lead to co	ated by CYP2C9. Other CYI ydroxymetabolite. A substa morphisms of CYP2C9 hav ally guided drug selection hibitors may enhance the ompromised efficacy of dic	P enzymes including antial portion of the e not been found to are recommended. drug exposure and
	Disopyramide	Normal Exposur	e to Disopyramide			INFORMATIV
		CYP2D6 have not b adjustments are re Polypharmacy gu disopyramide plasi	excreted in urine as unchanged been found to affect patient resp commended. No genetically guid idance : Co-administration of dis ma concentrations, which could r ase in disopyramide plasma conc nction.	onse to disopyramide. led drug selection or d opyramide with inhibito esult in a fatal interacti	No genetically guided drug osing adjustments are recor- ors of CYP3A4 may cause a on. Co-administration with	g selection or dosing ommended. an increase in o CYP3A4 inducers
	Deluterreit	Normal Respons	se to Dolutegravir			
\checkmark	Dolutegravir					ACTIONABL
\checkmark	Dolutegravir Tivicay®, Triumeq®	contribution from (have increased pla required for dolute	guidance: Dolutegravir is elimir CYP3A. Although UGT1A1 poor n sma levels of dolutegravir, these egravir due to genetic variations i Irugs that are strong enzyme ind	netabolizers or patients changes are not clinica n UGT1A1. Polypharm	s taking inhibitors of UGT1 Ily significant. No dosing a a cy guidance : Coadminis	l a minor A1 activity adjustments are tration of
✓ ✓	Tivicay®, Triumeq®	contribution from (have increased pla required for dolute dolutegravir with d	CYP3A. Although UGT1A1 poor r sma levels of dolutegravir, these egravir due to genetic variations i lrugs that are strong enzyme ind	netabolizers or patients changes are not clinica n UGT1A1. Polypharm	s taking inhibitors of UGT1 Ily significant. No dosing a a cy guidance : Coadminis	l a minor A1 activity Idjustments are tration of sma concentrations
✓ ✓	-	contribution from 0 have increased pla required for dolute dolutegravir with d of this drug. Normal Exposur Pharmacogenetic dosing recomment with drugs that are occur, which may c	CYP3A. Although UGT1A1 poor r sma levels of dolutegravir, these egravir due to genetic variations i lrugs that are strong enzyme ind	netabolizers or patients changes are not clinica n UGT1A1. Polypharm ucers, such as rifampin, y metabolized by CYP3. (acy guidance: Doravir as significant decreases avirine. Co-administrati	a taking inhibitors of UGT1 Ily significant. No dosing a acy guidance: Coadminis may result in reduced pla A. No genetically guided d ine is contraindicated whe s in doravirine plasma cond	a minor A1 activity adjustments are tration of sma concentrations ACTIONABL Irug selection or n co-administered centrations may
√ √ √	Tivicay®, Triumeq® Doravirine	contribution from 0 have increased pla required for dolute dolutegravir with d of this drug. Normal Exposur Pharmacogenetic dosing recomment with drugs that are occur, which may c	CYP3A. Although UGT1A1 poor r sma levels of dolutegravir, these egravir due to genetic variations i lrugs that are strong enzyme ind e to Doravirine guidance : Doravirine is primarily dations are available. Polypharm e strong CYP3A enzyme inducers decrease the effectiveness of dora ilt in increased plasma concentra	netabolizers or patients changes are not clinica n UGT1A1. Polypharm ucers, such as rifampin, y metabolized by CYP3. (acy guidance: Doravir as significant decreases avirine. Co-administrati	a taking inhibitors of UGT1 Ily significant. No dosing a acy guidance: Coadminis may result in reduced pla A. No genetically guided d ine is contraindicated whe s in doravirine plasma cond	a minor A1 activity adjustments are tration of sma concentrations ACTIONABL Irug selection or n co-administered centrations may
√ √ √	Tivicay®, Triumeq® Doravirine Pifeltro®	contribution from 0 have increased pla required for dolute dolutegravir with d of this drug. Normal Exposur Pharmacogenetic dosing recommend with drugs that are occur, which may c of CYP3A may resu Normal Respons Pharmacogenetic Polypharmacy gu	CYP3A. Although UGT1A1 poor r sma levels of dolutegravir, these egravir due to genetic variations i lrugs that are strong enzyme ind e to Doravirine guidance : Doravirine is primarily dations are available. Polypharm e strong CYP3A enzyme inducers decrease the effectiveness of dora ilt in increased plasma concentra	netabolizers or patients changes are not clinica n UGT1A1. Polypharm ucers, such as rifampin, y metabolized by CYP3. Tacy guidance : Doravir as significant decreases avirine. Co-administrati tions of doravirine.	a taking inhibitors of UGT1 Ily significant. No dosing a acy guidance: Coadminis may result in reduced place A. No genetically guided d ine is contraindicated whe is in doravirine plasma com- on of doravirine with drug	a minor A1 activity adjustments are tration of sma concentrations ACTIONABL Irug selection or n co-administered centrations may s that are inhibitors INFORMATIV available.

	7 Manal	hactor	PATIENT INFORMATION	SPECIMEN DETAILS	ORDERED BY
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		The patient's redu	ced CYP2C19 activity is unlikely t	o result in increased doxepin exp	osure.
		-	tions: Doxepin therapy can be p nsider therapeutic drug monitor	rescribed according to standard r ing to guide dose adjustments.	ecommended dosage and
		Insomnia: Doxepi	n can be prescribed according to	the standard recommended dos	age and administration.
	Dronabinol	Normal Dronab	nol Exposure (CYP2C9: Nor	nal Metabolizer)	ACTIONABI
-	Marinol®		type predicts a normal CYP2C9 age and administration.	netabolic activity. Dronabinol can	be prescribed at standard label-
	Duloxetine	Normal Exposur	e to Duloxetine		ACTIONABI
	Cymbalta®	these clearance pa to be clinically sigr Polypharmacy gu	thways are diminished in subjec iificant. No genetically guided d idance : Co-administration of du	ts with reduced enzyme activity, thrug selection or dosing recommer loxetine with a CYP1A2 inhibitor s	a lesser extent by CYP2D6. While hese changes have not been shown ndations are recommended. should be avoided. Co-administratio . Duloxetine is a moderate inhibitor o
	Dutasteride	Normal Respons	se to Dutasteride		INFORMATI
	Avodart®	Polypharmacy gu CYP3A4 inhibitors	idance: Dutasteride is extensive on dutasteride has not been stu		3A4 and CYP3A5. The effect of poter drug-drug interactions, use caution
	Edoxaban	Normal Respons	se to Edoxaban		INFORMATI
	Savaysa ®	via hydrolysis (mea the efflux transpor Studies indicate th edoxaban or its ac	Jiated by carboxylesterase 1; CES ter P-gp and its active metabolit at the two common variants SLC tive metabolite. There are no ge	51), conjugation, and oxidation by e (formed by CES1) is a substrate :O1B1 rs4149056 and ABCB1 rs10	n urine. There is minimal metabolisn CYP3A4. Edoxaban is a substrate of of the uptake transporter SLCO1B1. 45642 do not affect the exposure to lations. Polypharmacy guidance : nended for concomitant P-gp
	Efavirenz	Normal Efaviren	z Exposure (CYP2B6: Norma	l Metabolizer)	ACTIONAB
	Sustiva®			ely to have a normal efavirenz exp mmended dosage and administra	bosure following standard dosing. ation (600 mg/day).
/	Eprosartan	Normal Sensitiv	ity to Eprosartan		ACTIONABI
	Teveten ®	Pharmacogenetic Eprosartan is not r	guidance: Eprosartan is eliminanetabolized by the cytochrome l	, , , , , , , , , , , , , , , , , , ,	, primarily as unchanged compound of the cytochrome P450 genes is no adjustments are available.
/	Escitalopram	Normal Sensitiv	ity to Escitalopram (CYP2C1	9: Intermediate Metabolizer)	ACTIONAB
	Lexapro ®	Escitalopram can b	e prescribed at standard label-r	ecommended dosage and admini	stration.
	Eslicarbazepine Aptiom®	Normal Respons	se to Eslicarbazepine		INFORMATI
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	software		EMIC PURPOSES ONLY - DO NOT DISTRIE		Page 17 of

	7) Manch	lactor	PATIENT INFORMATION	SPECIMEN DETAILS	ORDERED BY
V	Manch Univer	sity	NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX:	SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: 11/11/20	022
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		be used to identify syndrome, Stevens- converted by a redu excretion unchange are available. Polyp	patients at risk for severe cutan Johnson syndrome (SJS) and to ictase to its active metabolite, e d and as a glucuronide conjuga	eous adverse reactions such as a exic epidermal necrolysis (TEN). Es eslicarbazepine. Eslicarbazepine is ate. No genetically guided drug s esence of enzyme-inducing drugs	slicarbazepine acetate (prodrug) is
\checkmark	Esomeprazole	Slightly Increased	Exposure to Esomeprazol	e (CYP2C19: Intermediate Me	etabolizer) INFORMATIV
	Nexium [®]			lightly increased esomeprazole ex el-recommended dosage and ad	xposure following standard dosing. Iministration.
\checkmark	Ethosuximide	Normal Response	e to Ethosuximide		INFORMATIV
	Zarontin [®]	Polypharmacy guid with caution when p	dance: ethosuximide is extensivorescribed with CYP3A4 inhibito		therefore this drug should be used ethosuximide clearance, and higher
	Etravirine	Normal Exposure	to Etravirine		ACTIONABL
	Edurant®	metabolites are sub etravirine is negligik guidance : Co-admi	sequently glucuronidated by ur ole. No genetically guided drug nistration of etravirine with dru ct or adverse reaction profile of	ridine diphosphate glucuronosylt selection or dosing recommenda gs that inhibit or induce CYP3A4,	YP3A4, CYP2C9 and CYP2C19. The ransferase. Renal elimination of ations are available. Polypharmacy , CYP2C9, and/or CYP2C19 may alter er of CYP3A and a weak inhibitor of
	Ezogabine	Normal Response	e to Ezogabine		INFORMATIV
	Potiga®	metabolite, no dose metabolized primar oxidative metabolisi are not expected to	adjustment is necessary in the ily via glucuronidation (by UGT m of ezogabine by cytochrome affect its efficacy or toxicity pro clearance by 30%, and dose inc	se individuals. Polypharmacy gu 1A4 and UGT1A1) and acetylation P450 enzymes, and genetic varia ofiles. Enzyme-inducing drugs suc	e in the exposure of ezogabine active Jidance: Ezogabine is extensively n (by NAT2). There is no evidence of ations in these metabolizing enzymes ch as carbamazepine and phenytoin n this drug is coadministered with
	Febuxostat	Normal Response	e to Febuxostat		INFORMATIV
-	Uloric®	metabolized both b cytochrome P450 er glucuronidated prin subjects with UGT1/ of these changes is febuxostat, there are available. Polyphar	y glucuronidation (40%) and ox nzymes (CYPs): CYP1A2, CYP2C harily by UGT1A1 and UGT1A3. A1*28 allele-UGT1A3*2a allele a not known. Although serious sk e no genetic biomarkers for pre macy guidance: Concomitant a h as theophylline, azathioprine	8 and CYP2C9 as well as other no Preliminary studies indicate that and decreased in those with the L kin and hypersensitivity reactions edicting such reactions; no genot administration of febuxostat, a xa	lative metabolism involves several on-CYP enzymes. Febuxostat is also febuxostat clearance is increased in JGT1A1*6 allele. The clinical relevance have been reported in patients taking ype-based recommendations are
	Felbamate	Normal Response			INFORMATIV

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	University

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PATIENT INFORMATION

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SPECIMEN DETAILS

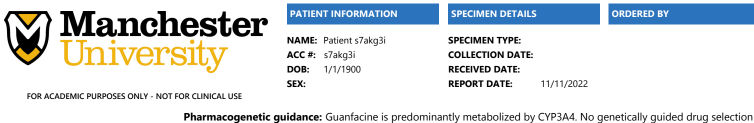
SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE

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X	Univer	sity	ACC #: DOB:	s7akg3i 1/1/1900	COLLECTION DATE: RECEIVED DATE:		
	FOR ACADEMIC PURPOSES ONLY - NO	FOR CLINICAL USE	SEX:		REPORT DATE:	11/11/2022	
		Polypharmacy gu 50% is present as minor for drug eli enzyme-inducing	uidance: A metabolite mination w antiepilept	bout 40-50% of a s and conjugates hen the drug is g ic drugs, which r	guided drug selection or dosi absorbed felbamate dose app 5. Felbamate is a substrate of 6 given as a monotherapy. This esults in a 30-50% decrease ir nt must be considered in pres	ears unchanged in urine CYP3A4 and CYP2E1, bu pathway is enhanced by n felbamate plasma conc	e, and an additional t these pathways are concomitant use of
\checkmark	Finasteride Proscar®		c guidance	: no genetically	guided drug selection or dosin nsively metabolized in human		
					ive not been studied. Because ients taking CYP3A4 enzyme		g-drug interactions,
\checkmark	Flibanserin	Normal Exposu	re to Flib	anserin (CYP2C	19: Intermediate Metabol	izer)	ACTIONABLE
	Addyi®	Flibanserin is prim	arily metal d to have a	bolized by CYP3A normal clearanc	cquired, generalized hypoa 4 and, to a lesser extent, by C e and a typical exposure to fli	YP2C19. The genotype	results predict that the
\checkmark	Fluconazole	Normal Respon	se to Flue	conazole			ACTIONABLE
	Diflucan®	approximately 80 pharmacokinetics or dosing recomm CYP2C9 and CYP2 therapeutic windo	% of the ac of flucona nendations C19 enzyn w metabo	Iministered dose zole is markedly are available. Po nes. Fluconazole 1 lized by CYP2C9,	t extensively metabolized and appearing in the urine as unc affected by reduction in renal lypharmacy guidance: Flucco reated patients who are conc CYP2C19 or CYP3A4 should b ation of the drug due to its lo	hanged drug and 11% a function. No genetically mazole is a moderate in omitantly treated with c re monitored. The enzym	s metabolites. The guided drug selection hibitor of CYP3A4, rugs with a narrow
\checkmark	Fluphenazine	Normal Exposu	re to Flup	henazine			INFORMATIVE
	Prolixin®	polymorphisms of selection or dosin inhibitors of CYP3 CYP3A4 inducers	^E CYP2D6 h g adjustme A4 may ca may cause	ave not been fou ents are recomme use an increase in a decrease in flu	metabolized by CYP2D6, CYF and to affect patient response ended. Polypharmacy guidar n fluphenazine plasma concerr ohenazine plasma concentrati tine) did not increase fluphen	to fluphenazine. No gen nce: Co-administration of trations while the co-actions. The co-administrations	netically guided drug of fluphenazine with Iministration with ion of fluphenazine
1	Flurbiprofen	Normal Flurbip	rofen Exp	osure (CYP2C9	: Normal Metabolizer)		ACTIONABLE
	Ansaid®	Rheumatoid Arth	nritis and	Osteoarthritis : F	urbiprofen therapy can be ini t effective dosage for the shor		
					d of the dosing range in geria with CYP2C9 inhibitors or inde		adjustment may be
\checkmark	Fondaparinux	Normal Respon	se to Fon	daparinux			INFORMATIVE
-	Arixtra®	CYPs, and therefo profiles. No genet concomitant use o	re genetic ically guid of fondapa risk of hem	variations in thes ed drug selection rinux with aspirin norrhage prior to	s eliminated unchanged throu e metabolizing enzymes are r or dosing recommendations or NSAIDS may enhance the initiation of therapy with fond orrhage.	not expected to affect its are available. Polyphar risk of hemorrhage. Disc	efficacy or toxicity macy guidance: The continue agents that
\checkmark	Fosaprepitant	Normal Respon	se to Fos	aprepitant			ACTIONABLE
	Powered By Translational oftware		Genet	ic Test Results For	Patient s7akg3i		Page 19 of 58

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	7) Manch	octor	PATIENT INFORMATION	SPECIMEN DETAILS	ORDERED BY
Ų	Manch Univers	sity	NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX:	SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: 11/11/2	022
	FOR ACADEMIC PURPOSES ONLY - NOT	FOR CLINICAL USE			
	Emend-IV®	intravenous adminis metabolism via N-a CYP1A2 and CYP2C dosing recommend inhibitors, a significa should be avoided v a loss of efficacy. Th inhibitor, and an inc	stration. Its antiemetic effects ar and O-dealkylations. These path 19. The drug is also glucuronida ations are available. Polypharn antly increased exposure of apr with fosaprepitant. Strong CYP3 nese drugs should also be avoid ducer of CYP3A4 and an inducer while others should be closely m	re attributable to aprepitant. Apro- toways are primarily catalyzed by 0 ated by UGT1A4 and UGT1A3. Not nacy Guidance: In presence of m epitant is expected which may le A4 inducers can significantly dec led with fosaprepitant. Aprepitan r of CYP2C9. Some substrates of	CYP3A4 with minor involvement from o genetically guided drug selection or
\checkmark	Fosphenytoin	Normal Phenytoi Metabolizer)	n (Fosphenytoin Active Me	tabolite) Exposure (CYP2C9:	Normal ACTIONABLE
	Cerebyx [®]	Fosphenytoin is a p CYP2C9 enzyme act	ivity. Fosphenytoin can be pres		ent is expected to have a normal e and a standard maintenance dose. ize the maintenance dosage.
\checkmark	Gabapentin	Normal Response	e to Gabapentin		INFORMATIVE
	Neurontin®	Polypharmacy gui Genetic variations in	dance: Gabapentin is eliminate	are not expected to affect its effic	nmendations are available. on and is not metabolized by CYPs. acy or toxicity profiles. Gabapentin
\checkmark	Glimepiride	Normal Exposure	e to Glimepiride		ACTIONABLE
	Amaryl®	subjects with reduce guided drug selection glimepiride with a s	ed CYP2C9 activity, such a chan on or dosing adjustments are re trong CYP2C9 inhibitor may res	olized by CYP2C9. While this clea ge has not been shown to be clir ecommended. Polypharmacy gu sult in higher glimepiride concent P2C9 inducer may result in lower	nically significant. No genetically Jidance : Co-administration of
\	Glipizide	Normal Exposure	e to Glipizide		INFORMATIVE
	Glucotrol ®	with reduced CYP2C selection or dosing strong CYP2C9 inhil	9 activity, such a change has no recommendations are available bitor may result in higher glipizi	ot been shown to be clinically sig	
\	Glyburide	Normal Exposure	e to Glyburide		ACTIONABLE
-	Micronase ®	clearance pathways clinically significant. guidance: Co-admi concentrations, leac	are diminished in subjects with No genetically guided drug sel nistration of glyburide with strc	reduced enzyme activity, these lection or dosing recommendation ong CYP2C9 and/or CYP3A4 inhib Co-administration with strong C	a lesser extent by CYP3A4. While these changes have not been shown to be ons are recommended. Polypharmacy oitors may result in higher glyburide YP2C9 and/or CYP3A4 inducers may
\checkmark	Guanfacine Intuniv®	Normal Response	e to Guanfacine		INFORMATIVE



		Pharmacogenetic guidance: Guanfacine is predominantly metabolized by CYP3A4. No genetically g or dosing recommendations are available and guanfacine extended-release should be titrated based response and tolerability of the individual patient. Polypharmacy guidance : The dose of guanfacine should be reduced to one half of the standard dose when co-medicated with a strong CYP3A4 inhibitor is dis should be increased to the standard recommended dose. Guanfacine dose should be increased up to recommended dose when used in combination with a strong CYP3A4 inducer (e.g., phenytoin, carbar St. John's wort etc.). When the CYP3A4 inducer is discontinued, the dose should be reduced to the sta- recommended dose within 7-14 days.	on the clinical extended-release bitor (e.g., continued, the dose double the mazepine, rifampin,
	Hydromorphone	Normal Response to Hydromorphone	INFORMATIVE
	Dilaudid®, Exalgo®	No genetically guided drug selection or dosing recommendations are available. Hydromorphone is n CYPs, and genetic variations in these metabolizing enzymes are not expected to affect its efficacy or the Hydromorphone can be prescribed at standard label-recommended dosage and administration.	
\checkmark	Ibuprofen	Normal Ibuprofen Exposure (CYP2C9: Normal Metabolizer)	ACTIONABLE
	Advil®, Motrin®	Pain, Dysmenorrhea, Rheumatoid Arthritis, Osteoarthritis, Fever and Other Anti-Inflammatory therapy can be initiated at standard label-recommended dosage and administration. Consider using dosage for the shortest duration consistent with the patient treatment goals.	
		Consider initiating treatment at the lowest end of the dosing range in geriatric patients. A dosage ad warranted when ibuprofen is administered with CYP2C9 inhibitors or inducers.	justment may be
\checkmark	Imipramine	Normal Imipramine Exposure (CYP2C19: Intermediate Metabolizer)	INFORMATIVE
	Tofranil®	The patient's reduced CYP2C19 activity is unlikely to result in increased imipramine exposure.	
		Psychiatric Conditions: Imipramine therapy can be prescribed according to standard recommended administration. Consider therapeutic drug monitoring to guide dose adjustments.	dosage and
	Indomethacin	Normal Indomethacin Exposure	INFORMATIVE
	Indocin®	Pharmacogenetic guidance : Indomethacin is metabolized mainly by O-demethylation to its inactive desmethyl indomethacin, a reaction catalyzed by CYP2C9. Genetic polymorphisms of CYP2C9 have no affect the response to indomethacin. No genetically guided drug selection or dosing recommendation	ot been found to
\checkmark	Irbesartan	Normal Irbesartan Exposure (CYP2C9: Normal Metabolizer)	INFORMATIVE
	Avapro®	Irbesartan can be prescribed at standard label-recommended dosage and administration.	
\checkmark	Isavuconazonium	Normal Response to Isavuconazonium	ACTIONABLE
	Cresemba®	Pharmacogenetic guidance: Isavuconazonium sulfate is a prodrug that is rapidly hydrolyzed in plas butylcholinesterase into its active moiety isavuconazole. Isavuconazole is extensively metabolized CYI and Common genetic polymorphism of these metabolizing enzymes gene are not expected to affect exposure. No genetically guided drug selection or dosing recommendations are available. Polyphare Isavuconazole is a sensitive CYP3A4 substrate and its use with strong CYP3A4 inhibitors or inducers of	P3A4 and CYP3A5 isavuconazole macy guidance:
\checkmark	Itraconazole Sporanox®	Normal Response to Itraconazole	ACTIONABLE



PATIENT	INFOR	MATION

NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX: SPECIMEN DETAILS

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\checkmark	Lansoprazole Prevacid®	Increased Exposure to Lansoprazole (CYP2C19: Intermediate Metabolizer)	INFORMATIVE
	Lamictal®	Pharmacogenetic guidance: Genotype results obtained from the pharmacogenetic test performed i be used to identify patients at risk for severe cutaneous adverse reactions such as anticonvulsant hyp syndrome, Stevens-Johnson syndrome (SJS) and toxic epidermal necrolysis (TEN). Lamotrigine is meta glucuronidation, which is mediated primarily by UGT1A4 with some contribution from UGT1A1 and U insufficient studies documenting the impact of genetic polymorphisms of these metabolizing enzyme response. No genetically guided drug selection or dosing recommendations are available. Polypharr Enzyme-inducing drugs increase lamotrigine clearance significantly, and higher doses of this drug are maintain therapeutic concentrations. Coadministration of valproic acid, an inhibitor of UGT enzymes, lamotrigine levels and may result in serious lamotrigine adverse effects (neurological and cutaneous), with a slow titration schedule is recommended when lamotrigine is added to existing valproic acid tree.	ersensitivity abolized by IGBT2B7. There are es on lamotrigine macy guidance: e required to increases . A low starting dose
\checkmark	Lamotrigine	Normal Response to Lamotrigine	INFORMATIVE
✓	Lacosamide Vimpat®	Normal Exposure to Lacosamide Pharmacogenetic guidance: Lacosamide is primarily cleared by renal excretion and metabolized by and CYP2C19. While these clearance pathways are diminished in subjects with reduced enzyme activi have not been shown to be clinically significant. No genetically guided drug selection or dosing adjus recommended. Polypharmacy guidance: Co-administration of lacosamide, in patients with reduced strong CYP2C9 and/or CYP3A4 inhibitors may result in higher lacosamide concentrations.	ty, these changes stments are
✓ 	Labetalol Normodyne®, Trandate®	Normal Response to Labetalol Pharmacogenetic guidance: Labetalol is extensively metabolized by UGT2B7, UGT1A1, and CYP2C19 metabolites. Preliminary studies indicate that following a single 200-mg oral dose, labetalol plasma co -fold higher in Chinese individuals with the CYP2C19 *2/*2 genotype than those with the CYP2C19 *1 clinical impact of this change is unknown. Polypharmacy guidance: Cimetidine increases the bioava and clinical monitoring is advised when both drugs are coadministered.	oncentrations are 2.9 /*1 genotype. The
✓	Ketorolac Toradol®	Normal Response to Ketorolac Pharmacogenetic guidance: Ketorolac is metabolized by glucuronidation (UGT enzymes) and oxidat catalyzing the oxidation are not well characterized. No genetically guided drug selection or dosing re available.	
✓	Ketoprofen Orudis®	Normal Response to Ketoprofen Pharmacogenetic guidance: Ketoprofen is primarily eliminated by glucuronidation (by UGT1A3, UG and no major implication of CYP2C9 in the metabolism of this drug has been demonstrated. No gene selection or dosing recommendations are available.	
		Pharmacogenetic guidance: Itraconazole is extensively metabolized to several metabolites by CYP3. metabolite is hydroxy-itraconazole, which has in vitro antifungal activity comparable to itraconazole; concentrations of this metabolite are about twice those of itraconazole. No genetically guided drug s recommendations are available. Polypharmacy guidance: Coadministration of itraconazole with pot may decrease the bioavailability of itraconazole and hydroxy-itraconazole to such an extent that effic. Therefore, administration of potent CYP3A4 inducers with itraconazole is not recommended and the should be avoided 2 weeks before and during treatment with itraconazole. Potent CYP3A4 inhibitors bioavailability of itraconazole and these drugs should be used with caution when coadministered with Itraconazole inhibit the metabolism of drugs metabolized by CYP3A4 or transported by P-glycoprote in increased plasma concentrations of these drugs and/or their active metabolite(s) when they are coelevated plasma concentrations may increase or prolong both therapeutic and adverse effects of these using concomitant medication, it is recommended that the corresponding label be consulted for info contraindications or need for dose adjustments.	trough plasma election or dosing tent CYP3A4 inducers acy may be reduced. use of these drugs may increase the h this antifungal. in, which may result administered. These se drugs. When



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		The patient's genotype may be associated with a slightly increased lansoprazole exposure following stan Consider prescribing lansoprazole at standard label-recommended dosage and administration. Once effi in the setting of chronic PPI therapy (beyond 12 weeks), consider a 50% reduction in the daily dose to m adverse events from prolonged acid suppression.	icacy is achieved,
	Levetiracetam	Normal Response to Levetiracetam	INFORMATIVE
	Keppra®	Pharmacogenetic guidance: No genetically guided drug selection or dosing recommendations are ava Polypharmacy guidance: Levetiracetam is minimally metabolized by non-CYP enzymes (esterases) and excreted unchanged in urine. Coadministration of enzyme-inducing antiepileptic drugs produce modest levetiracetam plasma levels.	is primarily
	Levomilnacipran	Normal Response to Levomilnacipran	INFORMATIVE
	• Fetzima®	Pharmacogenetic guidance: Levomilnacipran is moderately metabolized by desethylation, which is cata by CYP3A4, with minor contributions by CYP2C8, CYP2C19, CYP2D6, and CYP2J2. More than 58% of the cin urine as unchanged levomilnacipran, and 18% as N-desethyl levomilnacipran. Genetic polymorphisms expected to have a significant impact on levomilnacipran exposure. no genetically guided drug selection recommendations are available. Polypharmacy guidance : the daily levomilnacipran dose should not excoadministered with strong CYP3A4 inhibitors, such as ketoconazole, itrazonazole, and ritonavir.	dose is excreted of CYPs are not or dosing
	Levorphanol	Normal Response to Levorphanol	INFORMATIVE
	Levo Dromoran®	Pharmacogenetic guidance: Levorphanol is metabolized by glucuronidation which is mediated by UGT studies documenting the impact of genetic polymorphisms of this metabolizing enzyme on levorphanol no genetically guided drug selection or dosing recommendations are available. Polypharmacy guidance inducing drugs are expected to increase levorphanol clearance significantly.	response. And
	Lisdexamfetamine	Good Response to Lisdexamfetamine (COMT: Intermediate COMT Activity)	INFORMATIVE
_	Vyvanse ®	The patient's genotype result predicts a favorable response to amphetamine stimulants. Lisdexamfetami administered at the lowest effective dose, and dosage should be individually adjusted.	ne should be
	Losartan	Normal Response to Losartan (CYP2C9: Normal Metabolizer)	INFORMATIVE
	Cozaar®, Hyzaar®	Losartan is metabolized to its active metabolite by CYP2C9 and CYP3A4. The patient's genotype predicts exposure to losartan and its active metabolite. Losartan can be prescribed at label-recommended dosag administration.	
	Loxapine	Normal Response to Loxapine	INFORMATIVE
-	Loxitane®, Adasuve®	Pharmacogenetic guidance: Loxapine is metabolized extensively in the liver following oral administration metabolites formed. Loxapine metabolism occurs via hydroxylation and oxidation catalyzed by CYP1A2 accontributions from CYP3A4, CYP2D6 and FMO. There are no studies documenting the effect of genetic p these metabolizing enzymes on Loxapine disposition and there are no available genetically-guided drug dosing recommendations. Polypharmacy guidance: Loxapine is a central nervous system (CNS) depress concurrent use of Loxapine with other CNS depressants (<i>e.g.</i> , alcohol, opioid analgesics, benzodiazepine antidepressants, general anesthetics, phenothiazines, sedative/hypnotics, muscle relaxants, and/or illicit c can increase the risk of respiratory depression, hypotension, profound sedation, and syncope. Therefore, reduction/modification of CNS depressants if used concomitantly with Loxapine. Loxapine has anticholine concomitant use with other anticholinergic drugs can increase the risk of adverse reactions, including exercise and urinary retention.	along with olymorphisms of selection or sant. The s, tricyclic CNS depressants) consider dose ergic activity and
\checkmark	Lurasidone Latuda ®	Normal Response to Lurasidone	ACTIONABLE



Translational

PATIENT INFORMATION	
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SPECIMEN DETAILS

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		Pharmacogenetic gui available. Polypharma increase in lurasidone not be administered with moderate CYP3A4 strong inducers of CV moderate CYP3A4 inducer.	acy guid plasma with str 4 inhibit YP3A sh	lance: The cor concentrations ong CYP3A4 i ors. Monitor p ould not be a	ncomitant use o s, which could i inhibitors . Lur atients receivir dministered v	of lurasidone v ncrease or pro asidone dose s g lurasidone a /ith lurasidon	vith all CYP3A4 inh plong adverse drug should not exceed and any CYP3A4 in e. If lurasidone is	hibitors may re g effects. Lura 40 mg when hibitor. Rifam used concomi	sult in an sidone should administered apin or other tantly with a
	Meloxicam	Normal Meloxicam	Exposi	ure (CYP2C9:	: Normal Met	abolizer)			ACTIONABL
	Mobic®	Pain, Rheumatoid Ar dosage and administra patient treatment goal	thritis a ation. Co	nd Osteoarth	ritis: Meloxica	m therapy can			
		Consider initiating treat warranted when melow					-	losage adjustr	nent may be
	Memantine	Normal Response t	o Mem	antine					INFORMATIVE
	Namenda ®	Pharmacogenetic Gu hepatic metabolism to metabolite). CYP450 er documenting the effect response. No genetica Memantine is predom not expected to intera of drugs that use the s ranitidine, quinidine, a	o three ir nzymes cts of ge illy guide inantly r oct with r same rer	nactive metabo do not play a s netic variabilit ed drug selecti enally eliminat nemantine. Be nal cationic sys	blites (N-glucur significant role y in metabolizi ion or dosing m ted, and drugs ecause memant item, including	onide, 6hydi in the metabo ng enzymes or ecommendatio that are substr ine is elimination hydrochloroth	roxy metabolite, a lism of memantin organic cationic ons are available. I rates and/or inhib ed in part by tubu iiazide, triamteren	nd 1-nitroso-d e. There are no transporters of Polypharmacy itors of the CY lar secretion, c e, metformin,	leaminated o studies n memantine / Guidance: P450 system are coadministration
./	Meperidine	Normal Response t	o Mepe	eridine					INFORMATIVE
•	Demerol®	Pharmacogenetic gui is metabolized to norm variants in these enzyr meperidine metabolisu ritonavir, meperidine's these findings, the risk increased concentratic This combination shou	idance: meperidi mes have m is incr s exposu c of narco ons of no	no genetically ne by multiple e not been stur eased resulting re is significan otic-related ad ormeperidine s	CYPs, includin died. Polypha g in higher leve tly reduced wh lverse effects fi suggest a poten	g CYP2B6, CYF rmacy guidan els of its neuro ile normeperic rom this comb	23A4, and CYP2C1 ce: In patients tak toxic metabolite n line concentration ination appears to	9. The effects ing strong CY ormeperidine. is are increase be minimal. H	of genetic 'P inducers , In presence of d. Based on However,
./	Metaxalone	Normal Response t	o Meta	xalone					INFORMATIVE
	Skelaxin®	Pharmacogenetic gui CYP2D6, CYP2E1, and extent. no genetically	idance: CYP3A4	Metaxalone is . Genetic polyr	morphisms of t	hese enzymes	are unlikely to aff		
	Methadone	Normal Methadone	e Expos	ure (CYP2B6	i: Normal Me	tabolizer)			INFORMATIVE
	Dolophine [®]	The patient's genotype	e is asso	ciated with a n	normal methad	one exposure	following standard	d dosing.	
		For Addiction Treatm	nent: Co	nsider standar	rd prescribing a	nd monitoring	g practices.		
		For Pain Managemen exposure when this dr							
✓	Methocarbamol Robaxin®	Normal Response t	o Meth	locarbamol					INFORMATIVE
	Powered By		Genetic	Test Results Fo	r Patient s7akg	13i			

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\checkmark	Universi	ty

PATIENT INFORMATION

NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX: SPECIMEN DETAILS

ORDERED BY

SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: 11/11/2022

FOR ACADEMIC PURPOSES ONLY - NOT FOR CLINICAL USE

Pharmacogenetic guidance: Methocarbamol is metabolized via dealkylation and hydroxylation. The enzymes responsible for the metabolism of this drug have not been characterized. No genetically guided drug selection or dosing recommendations are available.

\checkmark	Micafungin	Normal Response to Micafungin	ACTIONABLE
	Mycamine [®]	Pharmacogenetic guidance: Micafungin is metabolized by arylsulfatase, catechol-O-methyltransf P450 enzymes. Even though micafungin is a substrate for and a weak inhibitor of CYP3A in vitro, hy is not a major pathway for micafungin metabolism in vivo. No genetically guided drug selection or recommendations are available.	/droxylation by CYP3A
	Milnacipran	Normal Response to Milnacipran	INFORMATIVE
	Savella®	Pharmacogenetic guidance: milnacipran is minimally metabolized by UGT enzymes and primarily in urine. No genetically guided drug selection or dosing recommendations are available. Polyphar coadministration of drugs that inhibit or induce CYP or UGT enzymes are unlikely to affect the exp	macy guidance:
√	Mirtazapine	Normal Exposure to Mirtazapine	ACTIONABLE
	Remeron ®	Pharmacogenetic guidance : Mirtazapine is metabolized by CYP2D6 as well as CYP1A2 and CYP3A clearance pathways are diminished in subjects with reduced enzyme activity, these changes have n clinically significant. No genetically guided drug selection or dosing recommendations are recomm guidance : Co-administration of mirtazapine with CYP inhibitors did not result in clinically relevant changes. While co-administration with strong CYP inducers (ex. phenytoin, carbamazepine, rifampi mirtazapine concentrations and a lack of efficacy.	ot been shown to be hended. Polypharmacy pharmacokinetics
	Nabumetone	Normal Response to Nabumetone	INFORMATIVE
	Relafen ®	Pharmacogenetic guidance: Nabumetone is a prodrug, which is converted by CYP1A2 to an active that is further metabolized by CYP2C9 to an inactive metabolite. Theoretically, individuals with reduce (i.e. CYP2C9 poor metabolizers) may have higher levels of the active metabolite, but it is unknown waltered drug response. No genetically guided drug selection or dosing recommendations are availa Guidance: CYP1A2 inhibitors may inhibit the activation of nabumetone to its active metabolite rest the therapeutic effects of this drug. On the other hand, CYP1A2 inducers (i.e. smoking) may result in nabumetone active metabolite, which may affect the response to this drug.	uced CYP2C9 activity whether this results in able. Polypharmacy ulting in a reduction in
\checkmark	Naltrexone	Good Response to Naltrexone (OPRM1: Altered OPRM1 Function)	INFORMATIVE
	Vivitrol®, Contrave®	<u>Treatment of alcohol dependence</u> : the patient has the OPRM1 118GG homozygous genotype that good clinical outcome with naltrexone therapy. Naltrexone-treated patients carrying two copies of allele are more likely to respond to this drug. They have a higher percentage of days abstinent and heavy drinking days than those who are not carriers of this allele. This association has not been replaced studies.	the OPRM1 118A>G G a lower percentage of
./	Naproxen	Normal Sensitivity to Naproxen	INFORMATIVE
	Aleve ®	Pharmacogenetic guidance: UGT2B7 is responsible for hepatic naproxen acyl glucuronidation, wl elimination pathway for this drug (60% of total clearance). CYP2C9 and CYP1A2 are responsible for desmethylnaproxen but this pathway is not the primary pathway for the elimination for naproxen. of CYP2C9 has not been found to affect the response to naproxen. No genetically guided drug sele recommendations are available.	the formation of O- Genetic polymorphism
√	Nateglinide	Normal Sensitivity to Nateglinide (SLCO1B1: Decreased Function)	INFORMATIVE
-	Starlix®	The patient carries one copy of the SLCO1B1 521T>C variant, which is associated with intermediate Nateglinide can be prescribed at label-recommended standard dosage and administration.	e transporter function.
\checkmark	Nateglinide	Normal Nateglinide Exposure (CYP2C9: Normal Metabolizer)	INFORMATIVE
	Powered By	Genetic Test Results For Patient s7akg3i	
	software	FOR ACADEMIC PURPOSES ONLY - DO NOT DISTRIBUTE - NOT FOR CLINICAL USE	Page 25 of 58

🚫 Mana	hostor	PATIENT INFORMATION	SPECIMEN DETAILS	ORDERED BY
Unive	ehester rsity	NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900	SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE:	
FOR ACADEMIC PURPOSES ONLY -	NOT FOR CLINICAL USE	SEX:	REPORT DATE: 11/11/2	2022
Starlix®	The patient's gene dosage and admin	•••••	to nateglinide, and this drug ca	n be prescribed at label-recommended
Olmesartan Benicar®	Pharmacogenetic gastrointestinal tr	genes is not expected to affect the	mil is hydrolyzed to olmesartan rtually no further metabolism of	ACTIONABLE its active metabolite in the olmesartan. Genetic variability of the tan medoxomil. No genotype-based
Omeprazole Prilosec®	The patient's gene Consider prescrib in the setting of c	ing omeprazole at standard label	ightly increased omeprazole exp -recommended dosage and adn	bosure following standard dosing. ninistration. Once efficacy is achieved, in the daily dose to minimize the risk of
O xcarbazepine	Normal Respor	se to Oxcarbazepine		INFORMATIVE
Trileptal®, Oxtellar XR®	be used to identif syndrome, Steven by a reductase to eliminated by dire or dosing recomn	y patients at risk for severe cutan is-Johnson syndrome (SJS) and to its active monohydroxylated active trenal excretion, glucuronidation	eous adverse reactions such as a xic epidermal necrolysis (TEN). C ve metabolite: 10-hydroxycarbaz on, and hydroxylation (minimal). armacy guidance: In the presen	ic test performed in this patient cannot anticonvulsant hypersensitivity Dxcarbazepine (prodrug) in converted zepine (MHD). This active metabolite is No genetically guided drug selection ace of enzyme-inducing drugs, the
🗸 Oxybutynin	Normal Respor	nse to Oxybutynin		INFORMATIVE
Ditropan®	Pharmacogenetic Polypharmacy go CYP3A4 strong inl	c guidance: no genetically guide uidance: Oxybutynin is extensive hibitor (itraconazole) increases ox rug to patients taking CYP3A4 en	ly metabolized in humans by CY sybutynin serum concentrations.	P3A4, and coadministration of a
Vxvmorphone	Normal Respon	nse to Oxymorphone		INFORMATIVE
Oxymorphone Opana®, Numorpha	n® No genetically gu CYPs, and genetic		enzymes are not expected to af	ymorphone is not metabolized by fect its efficacy or toxicity profiles.
Opana®, Numorpha	n® No genetically gu CYPs, and genetic Oxymorphone car	ided drug selection or dosing rec variations in these metabolizing n be prescribed at standard label-	enzymes are not expected to aft recommended dosage and adm	cymorphone is not metabolized by fect its efficacy or toxicity profiles. ninistration.
	n No genetically gu CYPs, and genetic Oxymorphone car Increased Export The patient's gene Consider prescribt in the setting of c	ided drug selection or dosing rec variations in these metabolizing n be prescribed at standard label- sure to Pantoprazole (CYP2C otype may be associated with a sl ing pantoprazole at standard labe	enzymes are not expected to aff recommended dosage and adm 19: Intermediate Metabolize ightly increased pantoprazole ex el-recommended dosage and ad	symorphone is not metabolized by fect its efficacy or toxicity profiles. inistration. INFORMATIVE sposure following standard dosing. Iministration. Once efficacy is achieved,
Opana®, Numorpha Pantoprazole Protonix®	n No genetically gu CYPs, and genetic Oxymorphone car Increased Export The patient's gene Consider prescribi in the setting of c adverse events from	ided drug selection or dosing rec variations in these metabolizing n be prescribed at standard label- sure to Pantoprazole (CYP2C otype may be associated with a sl ing pantoprazole at standard labe hronic PPI therapy (beyond 12 we	enzymes are not expected to aff recommended dosage and adm 19: Intermediate Metabolize ightly increased pantoprazole ex el-recommended dosage and ad	rymorphone is not metabolized by fect its efficacy or toxicity profiles. ninistration. (er) INFORMATIVI xposure following standard dosing. Iministration. Once efficacy is achieved, in the daily dose to minimize the risk o
Opana®, Numorpha	 No genetically gu CYPs, and genetic Oxymorphone car Increased Export The patient's genetic Consider prescribinin the setting of cardioverse events from Normal Respont Pharmacogenetic and CYP3A5. No genetic Enzyme-inducing should be increase Coadministration 	ided drug selection or dosing rec variations in these metabolizing n be prescribed at standard label- sure to Pantoprazole (CYP2C otype may be associated with a sl ing pantoprazole at standard labe hronic PPI therapy (beyond 12 we om prolonged acid suppression. nse to Perampanel c guidance: Perampanel is elimin genetically guided drug selection of drugs decrease perampanel plas ed when it is added to a stable th with strong enzyme-inducers oth	enzymes are not expected to aff recommended dosage and adm 19: Intermediate Metabolize ightly increased pantoprazole ex el-recommended dosage and ad eeks), consider a 50% reduction mated either unchanged or follow or dosing recommendations are ima concentrations by 50-60%, a erapy regimen containing enzyr iers than antiepileptic drugs (e.g	rect its efficacy or toxicity profiles. inistration. inistration. inistration. inistration. Once efficacy is achieved, in the daily dose to minimize the risk of INFORMATIVE ving oxidative metabolism by CYP3A4 e available. Polypharmacy guidance: and the initial dosage of the drug ne-inducing antiepileptic drugs.
 Opana ®, Numorpha Pantoprazole Protonix ® Perampanel 	 No genetically gu CYPs, and genetic Oxymorphone car Increased Export The patient's genetic Consider prescribinin the setting of car adverse events from Normal Response Pharmacogenetic and CYP3A5. No genetic Enzyme-inducing should be increase Coadministration Coadministration by 20%. 	ided drug selection or dosing rec variations in these metabolizing n be prescribed at standard label- sure to Pantoprazole (CYP2C otype may be associated with a sl ing pantoprazole at standard labe hronic PPI therapy (beyond 12 we om prolonged acid suppression. nse to Perampanel c guidance: Perampanel is elimin genetically guided drug selection of drugs decrease perampanel plas ed when it is added to a stable th with strong enzyme-inducers oth	enzymes are not expected to aff recommended dosage and adm 19: Intermediate Metabolize ightly increased pantoprazole ex el-recommended dosage and ad eeks), consider a 50% reduction atted either unchanged or follow or dosing recommendations are sma concentrations by 50-60%, a erapy regimen containing enzyr lers than antiepileptic drugs (e.g P3A4 inhibitors such as ketocona	symorphone is not metabolized by fect its efficacy or toxicity profiles. inistration. Pr) INFORMATIVE syposure following standard dosing. Iministration. Once efficacy is achieved, in the daily dose to minimize the risk of INFORMATIVE ving oxidative metabolism by CYP3A4 e available. Polypharmacy guidance: and the initial dosage of the drug ne-inducing antiepileptic drugs. ., rifampin) should be avoided.
 Opana ®, Numorpha Pantoprazole Protonix ® Perampanel Fycompa ® 	 No genetically gu CYPs, and genetic Oxymorphone car Increased Export The patient's genetic Consider prescribinin the setting of car adverse events from Normal Response Pharmacogenetic and CYP3A5. No genetic Enzyme-inducing should be increase Coadministration Coadministration by 20%. 	ided drug selection or dosing rec variations in these metabolizing n be prescribed at standard label- sure to Pantoprazole (CYP2C otype may be associated with a sl ing pantoprazole at standard labe hronic PPI therapy (beyond 12 we om prolonged acid suppression. nse to Perampanel c guidance: Perampanel is elimin genetically guided drug selection of drugs decrease perampanel plas ed when it is added to a stable th with strong enzyme-inducers oth with perampanel with strong CYF	enzymes are not expected to aff recommended dosage and adm 19: Intermediate Metabolize ightly increased pantoprazole ex- el-recommended dosage and ad eeks), consider a 50% reduction mated either unchanged or follow or dosing recommendations are sma concentrations by 50-60%, a rerapy regimen containing enzyr ivers than antiepileptic drugs (e.g. 23A4 inhibitors such as ketocona mal Metabolizer)	tymorphone is not metabolized by fect its efficacy or toxicity profiles. ininistration. Pr) INFORMATIVE typosure following standard dosing. Iministration. Once efficacy is achieved, in the daily dose to minimize the risk or INFORMATIVE ving oxidative metabolism by CYP3A4 e available. Polypharmacy guidance: and the initial dosage of the drug me-inducing antiepileptic drugs. ., rifampin) should be avoided. azole increases perampanel exposure

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	V I In ivor	eitzz		Patient s7akg3i s7akg3i	SPECIMEN TYPE: COLLECTION DATE:		
		SILY	DOB:	1/1/1900	RECEIVED DATE:		
			SEX:		REPORT DATE:	11/11/2022	
	FOR ACADEMIC PURPOSES ONLY - NC		in diant	a that the national is an	nested to have a nermal		una activity. Dhanytain can ha
	Dilantin®	prescribed at a star	dard load		ard maintenance dose. C		me activity. Phenytoin can be peutic drug monitoring and
/	Pimavanserin	Normal Respons	e to Pim	avanserin			INFORMATIV
	Nuplazid®	by CYP2J2, CYP2D6 major active metab Polypharmacy gui QT prolongation or (e.g., quinidine, pro (e.g., ziprasidone, cl of pimavanserin wit drug is coadminister	and oth olite (AC- dance: P in combi cainamid hlorprom h CYP3A red with	er CYP and FMO enzyr 279). There are no ava imavanserin prolongs ination with other drug e) or Class 3 antiarrhyt azine, thioridazine), an 4 inhibitor increases pi	nes. CYP3A4 is the majo ilable genetically-guided the QT interval and its us gs known to prolong QT hmics (e.g., amiodarone, d certain antibiotics (e.g mavanserin exposure an rs. Coadministration of p	r enzyme resp d drug selectio se should be a interval incluo sotalol), certa ., gatifloxacin, d a dose redu	d CYP3A5 and to a lesser extent consible for the formation of its on or dosing recommendations. avoided in patients with known ding Class 1A antiarrhythmics ain antipsychotic medications moxifloxacin). Concomitant use loction of 50% is needed when th with strong CYP3A inducers may
/	Piroxicam	Normal Piroxicar	n Expos	ure (CYP2C9: Norm	al Metabolizer)		ACTIONABL
	Feldene ®						d label-recommended dosage consistent with the patient
					he dosing range in geria P2C9 inhibitors or induc		A dosage adjustment may be
/	Posaconazole	Normal Respons	e to Pos	aconazole			ACTIONABL
	Noxafil®	and feces account f direct glucuronidat glycoprotein are en drug selection or de inducers may affect	or approx on, mino zymes ar osing reco posacon	ximately 17% of the ac r oxidation and dealky id transporters that pla ommendations are ava	Iministered dose. The me lation. CYP3A4 (and pos ay a role in the eliminatic ilable. Polypharmacy g rations. Concomitant use	etabolic pathv sibly CYP1A1 on of this antif uidance: UGT	excreted metabolites in urine vays for posaconazole include and CYP3A5), UGT1A4, and P- fungal. No genetically guided and P-glycoprotein inhibitors o zole and these agents should be
/	Prasugrel	Normal Respons	e to Pra	sugrel			ACTIONABL
-	Effient®	converted to the ac Prasugrel active me efficacy or safety pr drug selection or de	tive meta tabolite e ofile are osing rece	bolite primarily by CY exposure and platelet i also unaffected by CYF	P3A4 and CYP2B6, and to reactivity are not affected P2B6, CYP3A5, and CYP2 illable. Polypharmacy g	o a lesser exte d by CYP2C19 C9 genetic va	a thiolactone, which is then ent by CYP2C9 and CYP2C19. genetic variants. Prasugrel riants. No genetically-guided sugrel can be administered with
/	Pregabalin	Normal Respons	e to Pre	gabalin			INFORMATIV
	Lyrica®	Polypharmacy gui Genetic variations in	dance: P n these m	regabalin is eliminated netabolizing enzymes a		excretion and	ndations are available. l is not metabolized by CYPs. or toxicity profiles. Pregabalin can
	Proguanil	Normal Exposure	to Pro	guanil			INFORMATIV

(\mathbf{X})	Manchester
\checkmark	University

PATIENT INFORMATION						
NAME:	Patient s7akg3i					

SPECIMEN DETAILS

SPECIMEN TYPE:

Y	Univer	-	CC #: s7	atient s7akg3i 'akg3i '1/1900	SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE:		
	FOR ACADEMIC PURPOSES ONLY - NOT		:X :		REPORT DATE:	11/11/2022	
		cycloguanil. Preliminary exposure compared to proguanil metabolic rat and there is insufficient	studies subjects ios acros data to vailable.	indicate that ind with normal CYF ss CYP2C19 meta calculate dose ac Polypharmacy	drug that is primarily met ividuals with reduced CYP 2C19 function, but there is bolizer status. The clinical djustments. No genetically guidance : Co-administrat proguanil) exposure.	2C19 function, have reduce s considerable overlap of relevance of this change guided drug selection or	ed cycloguanil cycloguanil and is not well understood dosing
	Quetiapine	Normal Response to	Quetia	pine			INFORMATIV
	Seroquel®	Pharmacogenetic guid CYP2D6 are also respon compared to CYP3A4. N effect) is further metabol CYP3A4, CYP2D6 and C metabolite N-desalkylq genetically guided drug the clinical response an reduced to one sixth o itraconazole, indinavir, n by 6 fold. Quetiapine do treatment (e.g. > 7-14 of	lance: C sible for I-desalk Ilized by YP3A5 e uetiapin selectio d tolerat forigin itonavir, ose shou lays) of a	vetiapine is prece quetiapine is prece quetiapine meta ylquetiapine, a p c CYP2D6 and CY nzymes may be r e. However, the c on or dosing reco bility of the indivi al dose when co nefazodone). W Ild be increased to a potent CYP3A4	lominantly metabolized to abolism but their role in th narmacologically active me P3A4. Preliminary studies responsible in variable exp dinical significance of thes mmendations are availabl dual patient. Polypharma medicated with a potent hen the CYP3A4 inhibitor up to 5 fold of the original inducer (e.g., phenytoin, c ose should be reduced to	e overall metabolism of t etabolite (responsible of t have shown that genetic osures to quetiapine and e changes is not establish e. Quetiapine dose should cy guidance : Quetiapine CYP3A4 inhibitor (e.g., ke is discontinued, the dose dose when used in comb arbamazepine, rifampin,	his drug is minor he antidepressant polymorphisms of to its active ed yet and no d be titrated based or dose should be toconazole, should be increased ination with a chronic St. John's wort etc.).
	Quinidine Quinidine®	metabolizing enzyme for Polypharmacy guidan	l ance : Ir or quinid ce : Co-a of quinic	n vitro studies usi line. No genetica dministration of line. This may res	ng human liver microsom Ily guided drug selection o drugs/herbs that are know sult in adverse events or su	or dosing adjustments are on to induce or inhibit CY	recommended. P3A can change
	Rabeprazole	Slightly Increased Ex	posure	to Rabeprazo	e (CYP2C19: Intermedi	ate Metabolizer)	INFORMATIV
	Aciphex [®]		-		a slightly increased rabepra pel-recommended dosage		standard dosing.
/	Raltegravir	Normal Response to	Ralteg	ravir			ACTIONABL
	Isentress®, Dutrebis®	metabolizers or patient are not clinically signific	s taking ant. No y guida i	inhibitors of UGT dosing adjustme nce: Coadministi	inated mainly through me 1A1 activity have increase nts are required for ralteg ation of raltegravir with du ntrations of this drug.	d plasma levels of raltegr ravir in patients who carry	avir, these changes genetic variants of
/	Repaglinide	Normal Sensitivity to	o Repag	glinide (SLCO1I	31: Decreased Function)	INFORMATIV
	Prandin®, Prandimet®	-			IT>C variant. This genotyp recommended standard d		
/	Rilpivirine	Normal Exposure to	Rilpivir	ine			ACTIONABL
-	Intelence ®	Pharmacogenetic guid selection or dosing reco	l ance : R ommend	ilpivirine is prima ations are availa	rily eliminated by metabo ole. Polypharmacy guida sma concentrations of rilp	nce: Co-administration or	
	Rivaroxaban	Normal Response to	Rivaro	xaban			INFORMATIV
	Powered By Translational	(ienetic Te	est Results For Pa	tient s7akg3i		
S s	oftware	FOR ACADEMIC F	URPOSES	ONLY - DO NOT DIST	RIBUTE - NOT FOR CLINICAL USE		Page 28 of 5

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V	Manch Univer	sity	NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX:	SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE:	11/11/2022	
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	Xarelto®	(ABCB1) and BCRP (safety profiles of riv. strong CYP3A4 inhit concomitant use of phenytoin, rifampin, as combined P-gp a increased exposure	ABCG2) transporters. Genetic polaroxaban. Polypharmacy guida pitors (e.g., ketoconazole, itracon rivaroxaban with drugs that are o	lymorphisms of these (nce: Avoid concomitar azole, lopinavir/ritonav combined P-gp and str th renal impairment co (e.g., diltiazem, verapa	genes are not nt use of rivarc vir, ritonavir, in rong CYP3A4 i badministered mil, dronedarc	ndinavir, and conivaptan). Avoid nducers (e.g., carbamazepine, rivaroxaban with drugs classified one, and erythromycin) have
\checkmark	Rolapitant	Normal Response	e to Rolapitant			ACTIONABLE
	Varubi®	hydroxylated rolapit selection or dosing decrease rolapitant moderate CYP2D6 in while others should medication. Rolapit glycoprotein (P-gp).	ant). Rolapitant is eliminated prin recommendations are available. exposure resulting in a loss of ef	marily through the hep Polypharmacy Guida ficacy. These drugs sho rates (e.g. thioridazine loing adjusted when co g efflux transporters: b	batic/biliary roo nce: Strong C buld be avoide , pimozide) are badministered breast-cancer-r	YP3A4 inducers can significantly ed with rolapitant. Rolapitant is a e contraindicated with rolapitant with this antiemetic resistance protein (BCRP) and P-
\	Rufinamide	Normal Response	e to Rufinamide			INFORMATIVE
	Banzel®	Polypharmacy guid not involved in its m efficacy or toxicity p rufinamide plasma l Patients stabilized o	guidance: No genetically guided dance: Rufinamide is extensively hetabolism. Therefore, genetic va rofiles. Coadministration of enzy evels, while coadministration of v n rufinamide should begin valpro n valproate should begin rufinam	metabolized by carbo riations in these metab me-inducing antiepile valproate increases the pate therapy at a low c	xylesterases. C polizing enzym ptic drugs proc e drug levels ar	ytochrome P450 enzymes are nes are not expected to affect its duce modest decreases in nd requires dose adjustment.
\checkmark	Sertraline Zoloft®	Normal Sensitivit	y to Sertraline (CYP2C19: Int	ermediate Metabol	izer)	ACTIONABLE
	2010/1 9	Sertraline can be pre	escribed at standard label-recom	mended dosage and a	administration.	
√	Sildenafil	Normal Response	e to Sildenafil			INFORMATIVE
-	Viagra®	CYP3A5*3/*3 genot unknown. Polyphar patients taking stre	guidance: Preliminary findings in ype compared to those with CYP macy guidance: Sildenafil is me ong CYP3A inhibitors, sildenafi um single dose of 25 mg in a 4	3A5*1/*1 genotype. Th tabolized by CYP3A4 (i l exposure is signific a	ne clinical sign major route) a antly increase	ificance of this change is and CYP2C9 (minor route). In ad, and it is recommended not
\checkmark	Silodosin	Normal Response	e to Silodosin			INFORMATIVE
-	Rapaflo®	Pharmacogenetic g metabolites. no gen silodosin is contrair	guidance: silodosin is extensively etically guided drug selection or ndicated with potent CYP3A4 inh caution when this drug is prescri	dosing recommendati ibitors, as the risk for s	ions are availal erious adverse	ble. Polypharmacy guidance: e events is increased at higher
\checkmark	Solifenacin Vesicare®	Normal Response	e to Solifenacin			INFORMATIVE

(\mathbf{N})	Manchester
	University

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		Pharmacogenetic guidance: no genetically guided drug selection or dosing recommendations are ava Polypharmacy guidance: Coadministration of a CYP3A4 strong inhibitor increases solifenacin serum concentrations significantly. Therefore, it is recommended not to exceed a 5 mg daily dose of solife coadministered with strong CYP3A4 inhibitors, as the risk for QTc prolongation induced by this d at higher concentrations. Although the effects of moderate CYP3A4 inhibitors were not examined, use this drug is administered with moderate CYP3A4 inhibitors.	enacin when rug is increased
\checkmark	Sotalol	Normal Exposure to Sotalol	INFORMATIVE
-	Betapace®, Sorine®, Sotylize®	Pharmacogenetic guidance : Excretion of sotalol is predominantly via the kidney in the unchanged forr lower doses are necessary in conditions of renal impairment. No genetically guided drug selection or do are recommended. Polypharmacy guidance : Co-administration of sotalol with drugs that can prolong can increase the patient's risk for developing drug induced long QT syndrome.	sing adjustments
	Sufentanil	Normal Response to Sufentanil	INFORMATIVE
	Sufenta®	Pharmacogenetic guidance: No genetically guided drug selection or dosing recommendations are ava Polypharmacy guidance: Sufentanil is primarily metabolized by CYP3A4 and so should be used with ca prescribed with CYP3A4 inhibitors or inducers.	
	Sulindac	Normal Response to Sulindac	INFORMATIVE
	Clinoril®	Pharmacogenetic guidance: Sulindac is primarily eliminated by glucuronidation which is catalyzed by sincluding UGT1A3, UGT1A9 and UGT2B7. The role of CYP2C9 in sulindac metabolism is of minor relevan guided drug selection or dosing recommendations are available.	
	Tacrolimus	Typical response to Tacrolimus (CYP3A5: Poor Metabolizer)	ACTIONABLE
	Prograf®	The genotype result predicts that the patient does not express the CYP3A5 protein. Therefore, there is n patient may metabolize tacrolimus more rapidly. Careful titration of tacrolimus in response to therapeut monitoring is recommended until a favorable response is achieved.	
	Tadalafil	Normal Response to Tadalafil	INFORMATIVE
-	Cialis®	Pharmacogenetic guidance: no genetically guided drug selection or dosing recommendations are ava Polypharmacy guidance: Tadalafil is extensively metabolized by CYP3A4. Tadalafil for Use as Needed taking concomitant potent inhibitors of CYP3A4, such as ketoconazole or ritonavir, the maximum recom vardenafil is 10 mg, not to exceed once every 72 hours. Tadalafil for Once Daily Use — For patients ta strong inhibitors of CYP3A4, the maximum recommended dose is 2.5 mg. Although specific interactions studied, other CYP3A4 moderate inhibitors would likely increase tadalafil exposure. The exposure of tad when coadministered with rifampin or other CYP3A4 inducers. This can be anticipated to decrease the e for once-daily use, though the magnitude of decreased efficacy is unknown.	I — For patients mended dose of iking concomitant have not been alafil is reduced
	Tapentadol	Normal Response to Tapentadol	INFORMATIVE
	Nucynta ®	No genetically guided drug selection or dosing recommendations are available. Tapentadol is not metal and genetic variations in these metabolizing enzymes are not expected to affect its efficacy or toxicity p Tapentadol can be prescribed at standard label-recommended dosage and administration.	
	Telmisartan	Normal Sensitivity to Telmisartan	ACTIONABLE
-	Micardis®	Pharmacogenetic guidance: Telmisartan is metabolized by conjugation to form a pharmacologically in glucuronide. Telmisartan is not metabolized by the cytochrome P450 isoenzymes. Genetic variability of t P450 genes is not expected to affect the patient's response to telmisartan. No genotype-based dosing a available.	the cytochrome
\checkmark	Terazosin Hytrin®	Normal Response to Terazosin	INFORMATIVE



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Pharmacogenetic guidance: no genetically guided drug selection or dosing recommendations are available. **Polypharmacy guidance:** The enzymes involved in metabolizing terazosin have not been characterized.

√	Thiothixene Navane®	Normal Response to Thiothixene Pharmacogenetic guidance: Thiothixene is metabolized by UGTs and by cytochrome P450 enzymes of CYP3A4). No genetically guided drug selection or dosing recommendations are available. Polypharma likely that strong enzyme inducers may lead to substantial decreases in thiothixene plasma concentrat potential for reduced effectiveness. Consider increasing the dose of thiothixene when concomitantly u CYP3A4 inducers (e.g., carbamazepine).	acy guidance: It is ions with the
✓	Tiagabine Gabitril®	Normal Response to Tiagabine Pharmacogenetic guidance: no genetically guided drug selection or dosing recommendations are as Polypharmacy guidance: Tiagabine is extensively metabolized by CYP3A4, and therefore this drug sh caution when prescribed with CYP3A4 inhibitors. Inducers of CYP3A4 increase tiagabine clearance by 2 initial dosage of the drug should be considered carefully when added to a stable therapy regimen con inducing antiepileptic drugs.	ould be used with 2-fold, and the
✓	Ticagrelor Brilinta®	Normal Response to Ticagrelor Pharmacogenetic guidance: Ticagrelor is extensively metabolized by CYP3A4 and CYP3A5 to both ac metabolites, and this drug does not require bioactivation to achieve its antiplatelet effect. The drug is P-glycoprotein, encoded by the ABCB1 gene. Studies have shown that the efficacy and safety profile of depend on CYP2C19 or CYP3A5 metabolizer statuses. Moreover, preliminary studies indicate that releve variants within the ABCB1, SLCO1B1, CYP3A4 and UGT2B7 genes do not affect ticagrelor exposure, eff profiles. No genetically-guided drug selection or dosing recommendations are available. Polypharma presence of strong CYP3A4 inhibitors, significantly increased exposure to ticagrelor is expected which adverse reactions such as dyspnea or bleeding. These drugs should be avoided with ticagrelor. Strong can significantly decrease ticagrelor exposure (resulting in a loss of efficacy) and these drugs should al Ticagrelor is a weak inhibitor of CYP3A4 and P-glycoprotein and some substrates of these proteins sho monitored and their dosing adjusted when coadministered with this medication.	also a substrate of f ticagrelor do not vant genetic icacy or safety cy guidance: In may lead to CYP3A4 inducers so be avoided.
√	Tofacitinib Xeljanz®	Normal Exposure to Tofacitinib Pharmacogenetic guidance: Tofacitinib is metabolized primarily by CYP3A4 with some contribution f Genetic variations in the CYP2C19 gene do not significantly influence tofacitinib exposure. Tofacitinib at standard dosing, but consider a dose reduction if a CYP2C19 poor metabolizer is also prescribed a G such as ketoconazole, erythromycin, diltiazem, troleandomycin, nefazodone, fluconazole, verapamil or inhibitors. Polypharmacy guidance : Tofacitinib dose should be reduced if a patient is taking strong C (e.g., ketoconazole), or if a patient is taking a moderate CYP3A4 inhibitor (e.g., alprazolam) with a stror inhibitor (e.g., fluconazole).	may be prescribed CYP3A4 inhibitor HIV protease CYP3A4 inhibitors
✓	Tolbutamide Orinase®	Normal Exposure to Tolbutamide Pharmacogenetic guidance: Tolbutamide is extensively metabolized by CYP2C9. While this clearance diminished in subjects with reduced CYP2C9 activity, such a change has not been shown to be clinicall genetically guided drug selection or dosing adjustments are recommended. Polypharmacy guidance of tolbutamide with a strong CYP2C9 inhibitor may result in higher tolbutamide concentrations possib hypoglycemia. Co-administration with a strong CYP2C9 inducer may result in lower tolbutamide conce lack of efficacy.	y significant. No : Co-administration ly leading to
✓	Topiramate Topamax®	Normal Response to Topiramate	INFORMATIVE



PATIEN	PATIENT INFORMATION						
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ACC COLLECTION DATE: DO **RECEIVED DATE:** SEX REPORT DATE: 11/11/2022 FOR ACADEMIC PURPOSES ONLY - NOT FOR CLINICAL USE Pharmacogenetic guidance: no genetically guided drug selection or dosing recommendations are available. Polypharmacy guidance: About 50% of absorbed topiramate dose appears unchanged in urine, and an additional 50% is present as metabolites and conjugates. Topiramate metabolism by cytochrome P450 enzymes is minor for its elimination when the drug is given as a monotherapy. However, this pathway is enhanced by concomitant use of enzymeinducing antiepileptic drugs, and may result in reduced topiramate plasma concentrations. Thus, this drug should be titrated slowly, and dose adjustment must be considered in presence of inducers. Concomitant administration of valproic acid and topiramate has been associated with hyperammonemia with and without encephalopathy. INFORMATIVE Torsemide Normal Torsemide Exposure (CYP2C9: Normal Metabolizer) Demadex[®] The patient's genotype predicts a normal exposure to torsemide and this drug can be prescribed at label-recommended dosage and administration. INFORMATIVE Trazodone Normal Response to Trazodone Oleptro® Pharmacogenetic guidance: Trazodone is metabolized to its active metabolite m-chlorophenylpiperazine by CYP3A4. This metabolite which may contribute to adverse events, is further metabolized by CYP2D6. The impact of genetic polymorphisms of this enzyme on the clinical response to trazodone is not well documented. No genetically guided drug selection or dosing recommendations are available. Polypharmacy guidance: It is likely that CYP3A4 inhibitors may lead to substantial increases in trazodone plasma concentrations with the potential for adverse effects. If trazodone is used with a potent CYP3A4 inhibitor, the risk of cardiac arrhythmia may be increased. Therefore coadministration of trazodone with drugs that are inhibit CYP3A4 should be approached with caution. **Trifluoperazine** INFORMATIVE Normal Response to Trifluoperazine Pharmacogenetic guidance: Thrifluoperazine extensively metabolized by oxidation, sulfoxidation, hydroxylation and Stelazine[®] direct glucuronidation catalyzed by UGT1A4. No genetically guided drug selection or dosing recommendations are available. Polypharmacy guidance: It is likely that strong enzyme inducers may lead to substantial decreases in trifluoperazine plasma concentrations with the potential for reduced effectiveness. Trimipramine INFORMATIVE Normal Trimipramine Exposure (CYP2C19: Intermediate Metabolizer) The patient's reduced CYP2C19 activity is unlikely to result in increased trimipramine exposure. Surmontil[®] Psychiatric Conditions: Trimipramine therapy can be prescribed according to standard recommended dosage and administration. Consider therapeutic drug monitoring to guide dose adjustments. INFORMATIVE Trospium Normal Response to Trospium Sanctura® Pharmacogenetic guidance: no genetically guided drug selection or dosing recommendations are available. Polypharmacy guidance: CYP enzymes do not contribute significantly to the elimination of trospium. No major drugdrug interactions are expected with CYP inhibitors or inducers. Valproic Acid INFORMATIVE Normal Response to Valproic acid Pharmacogenetic guidance: Genotype results obtained from the pharmacogenetic test performed in this patient cannot Depakene[®] be used to identify patients carrying mutations in mitochondrial DNA polymerase y (POLG). Valproic acid is contraindicated in patients known to have mitochondrial disorders caused by mutations in mitochondrial DNA polymerase y (POLG; e.g., Alpers-Huttenlocher Syndrome) and children under two years of age who are suspected of having a POLG-related disorder. Valproic acid is extensively metabolized in the liver, which occurs primarily by glucuronidation with probable contributions of UGT1A6, UGT1A9, and UGT2B7. This drug is also metabolized by a minor CYP-dependent oxidation pathway, which includes multiple enzymes such as CYP2A6, CYP2C9, and CYP2C19. There are insufficient studies documenting the impact of genetic polymorphisms of these metabolizing enzymes on valproic acid response, and no genetically guided drug selection or dosing recommendations are available. Polypharmacy guidance: enzyme-inducing drugs increase valproic acid clearance 2-fold, and higher doses of this drug are required to maintain therapeutic concentrations when added to a therapy regimen containing enzyme-inducing antiepileptic drugs.



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V	Valsartan Diovan®, Entresto®	formation of a min contribution of CY	guidance: Valsartan is excreted or metabolite, valeryl 4-hydroxy P2C9 in the overall disposition o response to valsartan. No geno	valsartan, which accounts f valsartan, genetic variabi	for about 9% of a dose. O lity of the CYP2C9 gene is	Given the limited
	Vardenafil Levitra®	CYP3A5*3/*3 genc Polypharmacy gu inhibitors such as l patients receiving should not be exc For itraconazole: 24-hour period. F	se to Vardenafil guidance: Preliminary findings itype compared to those with CY idance: The dosage of vardenaf ketoconazole, itraconazole, riton moderate CYP3A4 inhibitors suc ceeded in a 72-hour period. Fo 400 mg daily. For clarithromyc or ketoconazole: 200 mg daily hould not be exceeded in a 24	'P3A5*1/*1 genotype. The il may require adjustment avir, indinavir, saquinavir, a h as erythromycin. For rite r indinavir, saquinavir, at cin: a single dose of 2.5 n r. For itraconazole: 200 m	clinical impact of this cha in patients receiving stror atazanavir, or clarithromyc onavir, a single dose of 2 tazanavir, or ketoconazc ng vardenafil should not ng daily. For erythromyc	nge is unknown. ng CYP3A4 cin, as well as in 2.5 mg vardenafil ole: 400 mg daily. t be exceeded in a in: a single dose o
√	Vigabatrin Sabril®	Polypharmacy gu Therefore, genetic	se to Vigabatrin guidance: no genetically guide idance: Vigabatrin is eliminated variations in these metabolizing prescribed at standard label-reco	primarily through renal exercises are not expected	cretion and is not metabo d to affect its efficacy or to	olized by CYPs.
✓	Vilazodone Viibryd®	Pharmacogenetic a minor role in the available. Polypha plasma concentrat with a strong inhib erythromycin), the readjusted to the c to 2-fold when cor	guidance: Vilazodone is predor biotransformation of this drug. rmacy guidance: It is likely that ions with the potential for adver- itor of CYP3A4 (e.g., ketoconazo dose should be reduced to 20 n original level when the CYP3A4 ir acomitantly used with strong CYI If CYP3A4 inducers are discontin	No genetically guided dru CYP3A4 inhibitors may le- se effects. Vilazodone shor le). During coadministration of for patients with intoler nhibitor is discontinued. Co P3A4 inducers (e.g., carbar	g selection or dosing reco ad to substantial increase uld be reduced to 20 mg i on with moderate inhibito rable adverse events. The onsider increasing the dos mazepine). The maximum	ommendations are s in vilazodone if co-administered rs of CYP3A4 (e.g., dose can be se of vilazodone up
✓	Vorapaxar Zontivity®	polymorphisms of contraindicated in because of the inco CYP3A4 inhibitors increases in vorapa	Se to Vorapaxar guidance: vorapaxar is metabo these genes are not expected to people who have had a stroke, t reased bleeding risk. Polypharm (e.g., ketoconazole, itraconazole, axar exposure may increase blee amazepine, phenytoin, rifampin,	affect the efficacy or safe ransient ischemic attack (T hacy guidance: Avoid con , lopinavir/ritonavir, ritona ding risk. Avoid concomita	ty profiles of this drug. Vo FIA), or intracranial hemor comitant use of vorapaxa vir, indinavir, and conivap	orapaxar is rhage, (ICH) r with strong tan). Significant
✓	Voriconazole Vfend®		ity to Voriconazole (CYP2C1			ACTIONABL
\	Warfarin Coumadin®	Average Dosing	Requirements are Expected	(CYP2C9 *1/*1; VKORC	1 -1639G>A G/A)	ACTIONABL



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When initiating warfarin treatment for indications with a target INR of 2-3, consider one of the following methods to estimate dosing requirements:

FDA Label: CYP2C9 and VKORC1 genotype results indicate an expected therapeutic dose of 5-7 mg/day.

Pharmacogenomics algorithms/calculators available at www.warfarindosing.org:

Caucasians and Asians: Use the patient's demographics and other clinical factors along with CYP2C9 and VKORC1 genotypes to calculate the expected therapeutic dose.

Africans and African Americans: Use the patient's demographics and other clinical factors along with CYP2C9 and VKORC1 genotypes to calculate the expected therapeutic dose.

The provided recommendations in Africans and African Americans apply only when all the following CYP2C9 alleles are tested: *5, *6, *8, *11.



Normal Response to Ziprasidone

Pharmacogenetic guidance: Ziprasidone is primarily cleared following extensive metabolism. CYP3A4 is the major CYP contributing to the oxidative metabolism of ziprasidone with minor involvement from CYP1A2. Less than one-third of ziprasidone metabolic clearance is mediated by cytochrome P450 catalyzed oxidation and approximately two-thirds via reduction involving glutathione as well as aldehyde oxidase. No genetically guided drug selection or dosing recommendations are available. Individualization of ziprasidone dose with careful weekly titration is required. Dosage adjustments should generally occur at intervals of no less than 2 days, as steady-state plasma concentrations are achieved within 1 to 3 days. In order to ensure use of the lowest effective dose, patients should ordinarily be observed for improvement for several weeks before upward dosage adjustment. When deciding among the alternative treatments available, the prescriber should consider the finding of **ziprasidone's greater capacity to prolong the QT/QTc interval** compared to several other antipsychotic drugs. **Polypharmacy guidance:** Although coadministration of strong CYP3A4 inhibitors are expected to result in modest increases in ziprasidone plasma concentrations, a closer monitoring of the patient's response and a dose reduction may be considered. Ziprasidone dose may need to be increased when used in combination with a chronic treatment of a potent CYP3A4 inducer (e.g., phenytoin, carbamazepine, rifampin, St. John's wort etc.).



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Test Details

Gene	Genotype	Phenotype	Alleles Tested
CYP2C9	*1/*1	Normal Metabolizer	*2, *3, *4, *5, *6, *8, *11, *12, *13, *15, *25
CYP2C19	*1/*2	Intermediate Metabolizer	*2, *3, *4A, *4B, *5, *6, *7, *8, *17
CYP3A5	*3/*3	Poor Metabolizer	*3, *6, *7
VKORC1	-1639G>A G/A	Intermediate Warfarin Sensitivity	-1639G>A
APOE	ε2/ε4	Altered APOE function	ε2, ε4, (ε3 is reference)
CYP2D6	Indeterminate	Unknown Phenotype	*2, *3, *4, *4M, *6, *7, *8, *9, *10, *11, *12, *14, *15, *17, *18, *19, *20, *29, *41, *114
CYP2B6	*1/*1	Normal Metabolizer	*6, *9, *18, *18.002
CYP1A2	*1F/*1F	Normal Metabolizer - Higher Inducibility	*1C, *1F, *1K, *1L, *7, *11
COMT	Val158Met A/G	Intermediate COMT Activity	Val158Met
OPRM1	A118G G/G	Altered OPRM1 Function	A118G
CYP3A4	*1/*22	Intermediate Metabolizer	*2, *17, *22
SLCO1B1	*1/*5	Decreased Function	*5
F2 F5	rs1799963 GA rs6025 CT	Increased Risk of Thrombosis	rs1799963, rs6025
MTHFR	c.1286A>C AA c.665C>T CT	No Increased Risk of Hyperhomocysteinemia	c.1286A>C, c.665C>T
MTHFR	c.665C>T CT	Reduced MTHFR Activity	c.1286A>C, c.665C>T

Limitation: This test will not detect all the known alleles that result in altered or inactive tested genes. This test does not account for all individual variations in the individual tested. Absence of a detectable gene mutation does not rule out the possibility that a patient has different phenotypes due to the presence of an undetected polymorphism or due to other factors such as drug-drug interactions, comorbidities and lifestyle habits.

Methodology: Array based assays detect listed alleles, including all common and most rare variants with known clinical significance at analytical sensitivity and specificity >99%.

Disclaimer: Manchester University developed the Genotype test. The performance characteristics of this test were determined by Manchester University. It has not been cleared or approved by the U.S. Food and Drug Administration.

Translational Software Disclaimer: The information presented on this report is provided as general educational health information. The content is not intended to be a substitute for professional medical advice, diagnosis, or treatment. Only a physician, pharmacist or other healthcare professional should advise a patient on the use of the medications prescribed.

The pharmacogenetic assay involves use of reporting software and genotype-phenotype associations performed by Translational Software (www.translationalsoftware.com). The software has not been evaluated by the Food and Drug Administration. The software, and the report generated by the software, is not intended to diagnose, treat, cure, or prevent any disease. A qualified designee within the lab uses Translational Software to generate and subsequently review the report. The pharmacogenetic report is one of multiple pieces of information that clinicians should consider in guiding their therapeutic choice for each patient. It remains the responsibility of the health-care provider to determine the best course of treatment for a patient. Adherence to dose guidelines does not necessarily assure a successful medical outcome.

APOE Monograph

Clinical Utility





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Apolipoproteins (APO) are structural constituents of lipoprotein particles that have critical roles in blood lipid metabolism and transport. Apolipoprotein E (APOE) is a major constituent of triglyceride-rich chylomicrons, very low-density lipoproteins (VLDL), and some subclasses of high-density lipoproteins (HDL). APOE acts as a lipid carrier and transports cholesterol from the cells in the blood vessel wall to the liver for excretion. It also binds to several cell surface receptors to support membrane homeostasis and injury repair in the brain. Defects in APOE can result in hyperlipidemia, which is an important risk factor in the genesis of atherosclerosis and subsequent development of cardiovascular disease (CVD).

Assay Interpretation

There are three common APOE alleles designated ϵ_2 , ϵ_3 , and ϵ_4 , resulting from combinations of the two common variants c.388T>C (rs429358, p.Cys130Arg) and c.526 C>T (rs7412, p.Arg176Cys). These alleles result in E2, E3, and E4 protein isoforms, respectively. The approximate allele frequencies for most populations are 8-12% for ϵ_2 , 74-78% for ϵ_3 , and 14-15% for ϵ_4 .

The reference ranges for both mutations of APOE are c.388T/T and c.526C/C. This is consistent with a ɛ3/ɛ3 genotype and a normal APOE function.

Clinical Implications





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The apolipoprotein E3 is considered the normal isoform and is associated with normal lipid metabolism. The apolipoprotein E2 isoform shows defective binding of remnants to hepatic lipoprotein and delayed clearance from plasma. It is associated with a slower conversion of intermediate-density lipoproteins (IDL) to low-density lipoproteins (LDL), lower cholesterol, and higher triglycerides, compared to the normal apolipoprotein E3 isoform. The apolipoprotein E4 isoform results in the down regulation of the LDL receptor and is associated with increased total cholesterol, triglycerides, and LDL.

1 - Type III Hyperlipoproteinemia

<u>Result Interpretation</u>: APOE genotyping can be used to confirm the diagnosis of suspected type III hyperlipoproteinemia. Patients with symptoms (xanthomas) and with a lipid profile consistent with type III hyperlipidemia are candidates for APOE genotype analysis. Homozygosity for the APOE ϵ^2 allele is strongly associated with type III hyperlipoproteinemia. Over 90% of individuals presenting the clinical type III hyperlipoproteinemia have the rare ϵ^2/ϵ^2 genotype. These individuals have an increased risk of premature vascular disease.

Only 1-5% of individuals with the APOE ϵ^2/ϵ^2 genotype develop type III hyperlipoproteinemia, suggesting that other genetic, hormonal, or environmental factors must contribute the expression of this disease. Despite the accumulation of remnants in the circulation, most (>95%) APOE ϵ^2 homozygous subjects are normolipidemic or even hypolipidemic because of low LDL cholesterol levels. Other non- ϵ^2/ϵ^2 APOE genotypes (ϵ^3/ϵ^3 , ϵ^2/ϵ^3 , ϵ^2/ϵ^4 , ϵ^3/ϵ^4 , ϵ^4/ϵ^4) are not associated with type III hyperlipoproteinemia.

Summary: patients with a lipid profile (tendinous/tuberous xanthomas, elevated cholesterol, triglycerides, very low density lipoprotein (VLDL)) consistent with type III hyperlipoproteinemia are candidates for APOE genotyping. Most patients with type III hyperlipoproteinemia are homozygous for the APOE ϵ^2 allele and homozygosity for ϵ^2 allele is the only genotype associated with type III hyperlipoproteinemia. Factors in addition to the defective receptor binding activity of the apolipoprotein E2 isoform are necessary for manifestation of this disorder and only 1-5% of APOE ϵ^2 homozygous develop type III hyperlipoproteinemia.

2 - Atherosclerotic Cardiovascular Disease

<u>Result Interpretation</u>: genetic testing of APOE genotyping has been proposed for individual cardiovascular risk assessment and/or to predict the response to statin therapy. APOE genotyping can be informative in individuals with premature coronary heart disease or a family history of premature coronary heart disease. The APOE ϵ 4 allele has been linked to pure elevations of low-density lipoproteins (LDL), and the ϵ 4/ ϵ 4 and ϵ 3/ ϵ 4 genotypes are associated with increased serum cholesterol levels.

Although the evidence is not fully consistent, it has been estimated that having the $\epsilon 3/\epsilon 4$ or $\epsilon 4/\epsilon 4$ genotype is associated with on average a 30 -40% increased risk of cardiovascular disease relative to the common $\epsilon 3/\epsilon 3$ genotype.

There is some evidence that having an ϵ^2/ϵ^2 or ϵ^2/ϵ^3 genotype may be associated with a lower CVD risk in Caucasians normolipidemic patients. A proposed explanation of such observation is the association between APOE2 allele and lower Lp(a) concentrations.

Summary: the APOE ϵ 4 allele results in the down regulation of the LDL receptor and is associated with increased total cholesterol, triglycerides, and LDL. Several studies indicate that the ϵ 3/ ϵ 4, ϵ 2/ ϵ 4 or ϵ 4/ ϵ 4 genotypes are associated with increased plasma cholesterol levels. The presence of the ϵ 3/ ϵ 4 or ϵ 4/ ϵ 4 genotype has been suggested as a risk factor for atherosclerosis and coronary heart disease. Thus, the APOE genotype may be useful and informative in individuals with a family history of coronary heart disease as an additional tool for assessing their risk in conjunction with other clinical or laboratory findings. No clinical practice guidelines or position statements from U.S. professional associations were identified that recommended the use of APOE genotyping in cardiovascular risk assessment, including but not limited to the following: the 2013 American College of Cardiology/American Heart Association guidelines for the assessment of cardiovascular risk in asymptomatic patients; the 2009 U.S. Preventive Services Task Force (USPSTF) recommendations on the use of nontraditional risk factors for the assessment of coronary heart disease; the American Diabetes Association and the American College of Cardiology Foundation consensus conference publication.





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Apolipoprotein E Gene Variants and Risk of Coronary Heart Disease: A Meta-Analysis. Biomed Res Int. 2016;2016:3912175. 28: Moriarty PM et al. Lipoprotein(a) Mass Levels Increase Significantly According to APOE Genotype: An Analysis of 431 239 Patients. Arterioscler Thromb Vasc Biol. 2017 Mar;37(3):580-588. 29: Mack S et al. A genome-wide association meta-analysis on lipoprotein(a) concentrations adjusted for apolipoprotein(a) isoforms. J Lipid Res. 2017 May 16.





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COMT Monograph

Clinical Utility

Catechol-O-Methyltransferase (COMT) is an enzyme responsible for the metabolism of catecholamines and catechol-estrogens in the central nervous system and other organs. Dopamine is cleared mainly by COMT in the frontal cortex, and a reduced activity of this enzyme results in higher synaptic levels of dopamine, which affects prefrontal cortex cognitive response to certain drugs. A single nucleotide polymorphism of the COMT gene produces an amino acid change from valine to methionine (Val158Met) and reduces the enzyme activity by 3- to 4-fold.

Assay Interpretation

The most well studied COMT genetic variant (rs4680; 472G>A) is the one resulting in a valine to methionine change at codon 158. The variant allele called the Met allele is found in 30-47% of Caucasians, 23% of Africans, and 27-32% of Asians. The phenotype is defined by the presence of the reduced activity Met allele (A variant). The wild-type genotype (Val/Val; GG) predicts a high/normal COMT activity, the heterozygous genotype (Val/Met; GA) predicts an intermediate COMT activity, and the homozygous (Met/Met; AA) results in a low COMT activity.

The reference range for COMT metabolic status is COMT Val158Met GG (Val/Val) (wild-type), which is consistent with a high/normal COMT activity.

Clinical Implications

Several complex associations with the Val158Met variant as a risk factor for numerous diseases have been found, but seem to have limited predictive value. The response to some psychotropic medications seems to be dependent to some extent upon the COMT status. In general, the wild-type genotype result predicts a good response to methylphenidate and amphetamines in the treatment of attention deficit hyperactivity disorder. In the treatment of pain, patients who are homozygous for the Met allele require lower doses of morphine to achieve analgesia.

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PATIENT INFORMATION

 NAME:
 Patient s7akg3i

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PECIMEN DETAILS

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CYP1A2 Monograph

Clinical Utility

The cytochrome P450 1A2 (CYP1A2) accounts for 13% of total CYP in the human liver, and is responsible for metabolizing 8-10% of commonly used drugs as well as natural compounds such as caffeine. A large inter-individual variability in the elimination of drugs that are metabolized by CYP1A2 has been observed, which has been ascribed to genetic variations and environmental factors. CYP1A2 activity is highly inducible (increased) by environmental factors including smoking (tobacco), some drugs, and several dietary compounds (cruciferous vegetables).

Assay Interpretation

More than 20 different alleles have been characterized for the CYP1A2 gene, and some have been shown to affect enzyme activity and its sensitivity towards inducers (inducibility). The CYP1A2*1F is the most studied allele and results in a rapid metabolizer phenotype in the presence of inducers, while CYP1A2*1K and *1C alleles result in enzymes that are less active and less sensitive to induction. The CYP1A2*1F allele is found in 25-50% of Caucasians, 30% of Asians, and 50% of Ethiopians. The genotype-phenotype relationship for CYP1A2 is not well established, and can be expressed in terms of metabolic capacity as well as sensitivity towards induction (inducibility). Individuals are predicted to have CYP1A2 normal, intermediate, or poor metabolic capacity, with high, possible, or low inducibility depending on their genotype.

The reference range for CYP1A2 metabolic status is CYP1A2 *1A/ *1A, which is consistent with a normal metabolizer that is possibly inducible.

Clinical Implications

CYP1A2 genotype can help identify patients with high or low sensitivity to inducing agents, especially those released during smoking. The clinical relevance of this sensitivity becomes important in patients who are smokers or who quit smoking. Patients with the highly inducible genotype CYP1A2*1F/*1F can experience loss of response to drug substrates while they are exposed to dietary or environmental inducers, and therefore may require higher doses.

The following drugs used in the management of pain and various psychiatric conditions are metabolized extensively by CYP1A2 and are sensitive to its function: clozapine (Clozaril), duloxetine (Cymbalta), olanzapine (Zyprexa), and tizanidine (Zanaflex). CYP1A2 also metabolizes other important drugs such as melatonin, ondansetron (Zofran), pirfenidone (Esbriet), pomalidomide (Pomalyst), ramelteon (Rozerem), ropivacaine (Naropin), and tasimelteon (Hetlioz).

CYP1A2 metabolism is highly sensitive to inhibition and induction, and the occurrence of drug-drug interactions can have profound effects on the pharmacokinetics, response, and safety profiles of many CYP1A2 drug substrates.

Inhibitors

Some known strong CYP1A2 inhibitors include: ciprofloxacin (Cipro), enoxacin (Penetrex), fluvoxamine (Luvox) and zafirlukast (Accolate).

Some known **moderate to weak** CYP1A2 inhibitors include: allopurinol (Zyloprim), mexiletine (Mexitil), norfloxacin (Norflox), peginterferon alfa-2a (Pegasys), obeticholic acid (Ocaliva), oral contraceptives, ticlopidine (Ticlid), vemurafenib (Zelboraf) and zileuton (Zyflo).

Inducers

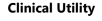
Known CYP1A2 inducers include: carbamazepine (Tegretol), montelukast (Singulair), moricizine (Ethmozine), omeprazole (Prilosec), phenobarbital, phenytoin (Dilantin), primidone (Mysoline) and rifampin (Rifadin).

Some dietary and environmental compounds found in charcoal-grilled food, cigarette smoke and cruciferous vegetables can also increase CYP1A2 activity.

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CYP2B6 Monograph





NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX: SPECIMEN DETAILS

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The cytochrome P450 2B6 (CYP2B6) enzyme is responsible for the metabolism of 4% of frequently used medications. It is expressed primarily in the liver and also in the brain. This enzyme is highly polymorphic: to date, a large number of variants have been identified. The CYP2B6 assay identifies some common variants that are associated with variability in enzyme activity, which has important clinical implications for efavirenz dosing.

Assay Interpretation

Genetic polymorphism in CYP2B6 result in different enzyme isoforms that are fully active (normal function), partially active (decreased function), inactive (no function), or have increased activity (increased function). The CYP2B6*1 allele is considered wild-type and encodes a functionally active enzyme (normal). Common decreased function alleles include the *6, *7, and *9 alleles. The *4 and *22 alleles are increased function alleles while the *18 allele is a no-function allele. Of note, common functional polymorphisms in CYP2B6 alter enzyme activity in a substrate-dependent manner. For most substrates, activity of the *9 variant is exceptionally low, activity of the *4 variant is similar or greater than that of the *1, while the activity of the *6 variant lies between *9 and *4, depending on substrate.

The most clinically relevant decreased function allele is the CYP2B6*6 allele. It is found in 7-18%, 10-17%, 23% and 33% of Caucasians, Asians, Mexican-Americans and African-Americans, respectively. The CYP2B6*18 allele is found only in individuals of African descent, with a frequency of 4-7%.

An individual carrying two normal function alleles is considered a normal metabolizer. An individual carrying one normal function allele and one decreased function allele OR one normal function allele and one no function allele OR one increased function allele and one decreased function allele OR one increased function allele and one no function allele is considered an intermediate metabolizer. An individual carrying two decreased function alleles OR two no function alleles OR one decreased function allele and one no function allele is considered a poor metabolizer. An individual carrying one normal function allele and one increased function allele is considered a rapid metabolizer. An individual carrying two increased function alleles is considered an ultra-rapid metabolizer.

The reference range for CYP2B6 metabolic status is CYP2B6 *1/ *1, which is consistent with a normal metabolizer.

Clinical Implications

CYP2B6 plays a role in the metabolism of the following drugs: artemisinin, bupropion (Wellbutrin), cyclophosphamide (Cytoxan), efavirenz (Sustiva), ketamine (Ketalar), meperidine (Demerol), methadone (Dolophine), nevirapine (Viramune), propofol (Diprivan), and selegiline (Eldepryl).

The impact of CYP2B6 polymorphism on pharmacokinetics and clinical response have been documented in adult and pediatric patients taking efavirenz. Patients who are intermediate or poor metabolizers are likely to have higher dose-adjusted trough concentrations of efavirenz following standard treatment when compared with normal metabolizers. Therefore, these patients are subject to increased risk of CNS adverse events. Decreased initial dosage and therapeutic monitoring are recommended for intermediate and poor metabolizers. Patients who are rapid or ultra-rapid metabolizers are likely to have slightly higher efavirenz clearance with no clinically significant effect on efficacy or toxicity. Efavirenz can be initiated with standard dosing for normal, rapid and ultra-rapid metabolizers. Specific dosing strategies for pediatric patients have been published by a panel of experts (Department of Human and Health Services-Panel on Antiretroviral Therapy and Medical Management of Children Living with HIV). These are based and on CYP2B6 genotype, weight, age and comorbidities.

Inhibitors or inducers of the CYP2B6 enzyme may modify its activity and change the patient's metabolizer status. This can result in drug-drug interactions when a drug substrate is prescribed with known CYP2B6 inhibitors or inducers.

Inhibitors

Some known CYP2B6 inhibitors include: clopidogrel (Plavix), darunavir (Prezista), prasugrel (Effient), ritonavir (Norvir), thiotepa (Tepadina), ticlopidine (Ticlid) and voriconazole (Vfend).

Inducers

Some CYP2B6 inducers include: artemether (Coartem), carbamazepine (Tegretol), dabrafenib (Tafinlar), efavirenz (Sustiva), metamizole, nevirapine (Viramune), phenobarbital, phenytoin (Dilantin), rifampin (Rimactane), ritonavir (Norvir) and St. John's wort.





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CYP2C19 Monograph

Clinical Utility

The cytochrome P450 2C19 (CYP2C19) is involved in the metabolism of 10% of clinically important medications. This enzyme is highly polymorphic and more than 30 different alleles have been identified in various ethnicities. The CYP2C19 assay identifies common and rare variants that are associated with variability in CYP2C19 enzyme, which has important pharmacological and toxicological implications for antidepressants, benzodiazepines, antiplatelets, and proton-pump inhibitors.

Assay Interpretation

CYP2C19 enzyme activity defines a normal or abnormal (intermediate, poor, rapid or ultra-rapid) metabolizer status for a given individual. Several variant alleles have been identified and result in different isoforms of the CYP2C19 enzyme that functionally exhibit normal function, reduction function, no function or increased function. The CYP2C19*1 allele is considered wild-type/reference allele and CYP2C19 *11, *13 and *18 encodes a functionally active enzyme (normal function allele). The alleles *2, *3 *4-*8, *22, *24, and *35-*37 encode an inactive enzyme and are referred to as no function alleles while the *9, *10, *16,*19, *25 and *26 alleles are classified as reduced function alleles. The CYP2C19*17 is an increased function allele.

A new joint statement by the Association for Molecular Pathology (AMP) and the College of American Pathologists (CAP), provides a twotiered classification of CYP2C19 alleles for inclusion during clinical testing. Tier 1 alleles that should be included in all clinical tests, include CYP2C19 *2, *3 and *17. Tier 2 alleles which may also be included in clinical tests include CYP2C19 *4A, *4B, *5, *6, *7, *8, *9, *10 and *35.

The CYP2C19 genotype-phenotype relationship is established based on the allele's function. An Individual with two normal function alleles is considered a normal metabolizer while an individual carrying two no function alleles is considered a poor metabolizer. An individual carrying one normal function allele with one no function allele or one no function allele with one increased function allele is classified as an intermediate metabolizer. Individuals with two increased function alleles are considered ultra-rapid metabolizers and carriers of one increased function allele with one normal function allele are referred to as rapid metabolizers. Because of limited evidence, individuals carrying two decreased function alleles OR those carrying one normal or increased function allele with one decreased function allele are provisionally categorized as intermediate metabolizers.

The reference range for CYP2C19 metabolic status is CYP2C19 *1/*1 genotype, which is consistent with a normal metabolizer.

Clinical Implications





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There is substantial evidence linking the CYP2C19 polymorphisms to variability in the pharmacological and safety profiles of the following therapies used in psychiatric conditions and pain management: amitriptyline (Elavil), sertraline (Zoloft), clobazam (Onfi), citalopram (Celexa), escitalopram (Lexapro), diazepam (Valium), imipramine (Tofranil), and carisoprodol (Soma). CYP2C19 plays a minor role in the elimination of methadone (Dolophine).

Cardiovascular medications that are metabolized by CYP2C19 include the prodrug clopidogrel (Plavix), propranolol (Inderal), and cilostazol (Pletal). Proton-pump inhibitors such as omeprazole (Prilosec), esomeprazole (Nexium), lansoprazole (Prevacid), dexlansoprazole (Dexilant), and pantoprazole (Protonix) are major substrates of CYP2C19.

Brivaracetam (Briviact) is primarily metabolized by hydrolysis and by hydroxylation. The hydrolysis reaction is mediated by amidase and the hydroxylation is mediated by CYP2C19. Brivaracetam exposure is increased by 22% and 42%, in CYP2C19 intermediate and poor metabolizers, respectively. CYP2C19 poor metabolizers and patients using inhibitors of CYP2C19 may therefore require dose reduction to avoid concentration-dependent toxicity.

CYP2C19 is significantly involved in the metabolism of voriconazole (Vfend) and the disposition of this antifungal is affected by CYP2C19 genetic polymorphisms. The CPIC dosing guideline for voriconazole recommends selecting an alternative agent that is not dependent on CYP2C19 metabolism in adults who are CYP2C19 ultrarapid metabolizers, rapid metabolizers or poor metabolizers. In pediatric patients, an alternative agent should be used in patients who are ultrarapid metabolizers or poor metabolizers.

Inhibitors or inducers of CYP2C19 enzyme may modify its activity and change the patient's metabolizer status. This can result in drug-drug interactions when a drug substrate is prescribed with known CYP2C19 inhibitors or inducers.

Inhibitors

Some known CYP2C19 inhibitors include: armodafinil (Nuvigil), delavirdine (Rescriptor), esomeprazole (Nexium), etravirine (Intelence), felbamate (Felbatol), fluconazole (Diflucan), fluoxetine (Prozac), fluvoxamine (Luvox), moclobemide (Manerix), modafinil (Provigil), omeprazole (Prilosec), oxcarbazepine (Trileptal), ticlopidine (Ticlid), topiramate (Topamax) and voriconazole (Vfend).

Inducers

Some known CYP2C19 inducers include: apalutamide (Erleada), artemether (Coartem), carbamazepine (Tegretol), efavirenz (Sustiva), enzalutamide (Xtandi), fosphenytoin (Cerebyx), phenobarbital, phenytoin (Dilantin), primidone (Mysoline), rifampin (Rifadin) and St. John's wort.

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CYP2C9 Monograph

Clinical Utility

The cytochrome P450 2C9 (CYP2C9) is involved in the metabolism of 15% of clinically important medications. This enzyme is highly polymorphic: to date, 60 alleles have been identified. The CYP2C9 assay identifies some common variants that are associated with variability in CYP2C9 enzyme activity, which has important pharmacological and toxicological implications for anticonvulsants, anticoagulants, and certain antidiabetics.

Assay Interpretation





NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX: SPECIMEN DETAILS

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CYP2C9 enzyme activity defines a normal or abnormal (intermediate and poor) metabolizer status for a given individual. Several variant alleles have been identified and result in different CYP2C9 isoforms that functionally are fully active, partially active, or inactive. The CYP2C9 *1 (wildtype) and CYP2C9*9 alleles encode functionally active enzymes. A number of CYP2C9 alleles such as *2, *4, *5, *8, *11, *12 and *31 encode partially active enzymes and are classified as decreased function alleles. Other CYP2C9 alleles such as *3, *6, *13, *15 and *25 are considered no -function alleles encoding inactive enzymes. Many other alleles with an unknown/uncertain functional effect have also been identified and can be revealed from testing.

A new joint statement by the Association for Molecular Pathology (AMP) and the College of American Pathologists (CAP), provides a twotiered classification of CYP2C9 alleles for clinical testing. Tier 1 alleles that should be included in all tests, these include CYP2C9 *2, *3, 5. *6, *8 and *11. Tier 2 alleles which may also be included in clinical tests include CYP2C9 *12, *13 and *15.

The genotype-phenotype relationship is established based on the allele's function. Individuals with two normal function alleles are considered normal (extensive) metabolizers (AS = 2.0). Individuals with one normal function allele with one decreased function allele are considered intermediate metabolizers (AS = 1.5). Individuals with one normal function allele and one no-function allele or two decreased function alleles are considered intermediate metabolizers (AS = 1.5). Individuals with one normal function allele and one no-function allele or two decreased function alleles are considered intermediate metabolizers (AS = 1.5). Individuals with one decreased function allele and one no function allele are considered poor metabolizers (AS = 0.5). Individuals with two no-function alleles are considered poor metabolizers (AS = 0.5). Individuals with two no-function alleles are considered poor metabolizers (AS = 0.5). Individuals with two no-function alleles are considered poor metabolizers (AS = 0.5). Individuals with two no-function alleles are considered poor metabolizers (AS = 0.5). Individuals with two no-function alleles are considered poor metabolizers (AS = 0.5). Individuals with two no-function alleles are considered poor metabolizers (AS = 0.5). Individuals with two no-function alleles are considered poor metabolizers (AS = 0.5). Individuals with two no-function alleles are considered poor metabolizers (AS = 0.5). The phenotype of carriers of one or two CYP2C9 unknown/uncertain function alleles cannot be predicted accurately (unknown phenotype).

The reference range for CYP2C9 metabolic status is CYP2C9 *1/*1 genotype, which is consistent with an AS of 2.0 and a normal metabolizer.

Clinical Implications

Abnormal CYP2C9 activity affects the therapeutic outcome of a variety of drugs used to treat cardiovascular and other conditions. Following the administration of drug substrates, the clinical manifestation in a poor or an intermediate metabolizer depends on the characteristics of the drug (i.e., the amount of drug/metabolites that is cleared by the enzyme), and the safety and pharmacological profiles of the drug and its metabolites. Within the medications used to treat cardiovascular conditions, there is compelling evidence that the response to certain angiotensin II inhibitors, statins, and anticoagulants is altered in individuals exhibiting abnormal CYP2C9 activity.

CYP2C9 plays a role in the metabolism of the following psychotropic drugs: fluoxetine (Prozac), phenytoin (Dilantin), and primidone (Mysoline). Several NSAIDs and Cox-2 inhibitors are substrates of CYP2C9, and patients with reduced CYP2C9 activity may have higher plasma levels of celecoxib (Celebrex), flurbiprofen (Ocufen), piroxicam (Feldene), or meloxicam (Mobic). CYP2C9 plays a minor role in the elimination of diclofenac (Voltaren), sulindac (Clinoril), and naproxen (Aleve).

Cardiovascular medications that are metabolized by CYP2C9 include warfarin (Coumadin), fluvastatin (Lescol), losartan (Cozaar), and irbesartan (Avapro).

The FDA has added a contraindication on the use of siponimod (Mayzent) in patients with a CYP2C9 *3/*3 genotype.

Inhibitors or inducers of the CYP2C9 enzyme may modify its activity and change the patient's metabolizer status. This can result in drug-drug interactions when a drug substrate is prescribed with known CYP2C9 inhibitors or inducers.

Inhibitors

Some known CYP2C9 inhibitors include: amiodarone (Cordarone), capecitabine (Xeloda), chloramphenicol, cimetidine (Tagamet), co-trimoxazole (Septra), danazol (Danocrine), delavirdine (Rescriptor), disulfiram (Antabuse), efavirenz (Sustiva), etravirine (Intelence), fluconazole (Diflucan), 5-fluorouracil (Adrucil), fluoxetine (Prozac), fluvastatin (Lescol), fluvoxamine (Luvox), gemfibrozil (Lopid), lomitapide (Juxtapid), metronidazole (Flagyl), miconazole (Oravig), oxandrolone (Oxandrin), phenytoin (Dilantin), sulfamethoxazole (Bactrim), sulfinpyrazone (Anturane), tamoxifen (Nolvadex), tigecycline (Tygacil), toremifene (Fareston), voriconazole (Vfend) and zafirlukast (Accolate).

Inducers

Some known CYP2C9 inducers include: alpelisib (Piqray), apalutamide (Erleada), aprepitant (Emend), bosentan (Tracleer), carbamazepine (Tegretol), dabrafenib (Tafinlar), elvitegravir (Genvoya, Stribild), enzalutamide (Xtandi), phenobarbital, phenytoin (Dilantin), primidone (Mysoline), rifampin (Rifadin, Rimactane), rifapentine (Priftin), ritonavir (Norvir) and St. John's wort.





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CYP2D6 Monograph

Clinical Utility

The cytochrome P450 2D6 (CYP2D6) is involved in the metabolism of 20% of clinically important medications. This enzyme is highly polymorphic: more than 100 different variants have been identified. The CYP2D6 assay identifies common variants that are associated with variability in CYP2D6 enzyme activity, which has important pharmacological and toxicological implications for antidepressants, antipsychotics, opioids, beta-blockers, and antiarrhythmics.

Assay Interpretation





PATIENT	INFORMATION

ACC #: s7akg3i DOB: 1/1/1900 SEX:

SPECIMEN DETAILS

ORDERED BY

NAME: Patient s7akg3i

SPECIMEN TYPE: COLLECTION DATE: **RECEIVED DATE:** REPORT DATE: 11/11/2022

FOR ACADEMIC PURPOSES ONLY - NOT FOR CLINICAL USE

CYP2D6 enzyme activity defines a normal or abnormal (intermediate, poor, or ultra-rapid) metabolizer status for a given individual. Many alleles have been identified and result in different CYP2D6 isoforms that functionally have normal activity, decreased activity, no activity or in some cases unknown activity. The CYP2D6*1 (reference or wild-type allele) is inferred when no variants tested by the assay are detected. Genotyping tests usually interrogate for both sequence variants and structural variants and results are used to infer CYP2D6 alleles or haplotypes as defined by PharmVar. Sequence variants include single nucleotide polymorphisms (SNPs) and small insertions and deletions. Structural variations include entire gene deletions (CYP2D6*5), gene duplication/multiplication (CYP2D6*1xN, *2xN and *4xN), and hybrids with the CYP2D7 gene. Moreover, alleles that contain hybrids can be found on their own (as single entities) or in combination with other gene copies (tandems).

Commonly detected alleles include those with normal function (e.g. CYP2D6 *1, *2 and *35), increased function (e.g. CYP2D6*1xN, *2xN), reduced function (e.g. CYP2D6*9, *10, *10-*36, *17, *29, and *41) and no-function (e.g. CYP2D6 *3, *4, *4N, *5, *6, *7, *8, *11, *12, *36, *4xN).

The genotype-phenotype relationship is established using a scoring system that assigns an activity value to every CYP2D6 allele, in order to assign a predicted phenotype. For a given genotype, the activity values of the constituent alleles are added together to calculate the CYP2D6 activity score. This activity score (AS) is then used to assign a predicted phenotype. The AS system was first published in 2008 and was recently refined by an expert group that included both the Clinical Pharmacogenetics Implementation Consortium (CPIC) and the Dutch Pharmacogenetics Working Group (DPWG) members. The group reviewed recent findings and heterogeneities in phenotype assignment.

The consensus method jointly recommended by CPIC and the DPWG redefined the translation of CYP2D6 diplotypes to phenotype as follows:

- CYP2D6 ultra-rapid metabolizers (UMs) when the calculated AS is > 2.25
- CYP2D6 normal metabolizers (NMs) when the calculated AS = x is within the range of $1.25 \le x \le 2.25$
- CYP2D6 intermediate metabolizers (IMs) when the calculated AS = x is within the range of 0 < x < 1.25
- CYP2D6 poor metabolizers (PMs) when the calculated AS = 0

After literature review, the standardization expert group also obtained consensus to 'downgrade' the activity value for the CYP2D6*10 allele from 0.5 to 0.25 to better reflect the level of enzyme reduction that is observed with this allele. The group proposed the following activity value when the functional effect of the CYP2D6 allele is known:

- fully functional CYP2D6 alleles are assigned an activity value of 1.0 (e.g. CYP2D6 *2, *35).
- reduced function CYP2D6 alleles (except CYP2D6*10) have an activity value of 0.5 (Note that the value of 0.5 does not indicate a 50%) reduction in activity, but signals decreased function, i.e., has functional activity somewhere between no-function and full function).
- CYP2D6*10 reduced function allele is assigned an activity value of 0.25.
- non-functional CYP2D6 alleles are assigned an activity value of 0 (e.g. CYP2D6 *4, *5, *36, *36xN, *4x2N).

For alleles with two or more gene copies, the value of the allele is multiplied by the number of gene copies (e.g. AS of CYP2D6*1x3N = 3 calculated as the AS of *1 which is 1 multiplied by 3). For a tandem, the sum of each allele value is used to assign the score (e.g. AS of CYP2D6*36-*10 = 0.25 calculated as the sum of AS of CYP2D6*36 which is 0 and AS of CYP2D6*10 which is 0.25).

The reference range for CYP2D6 metabolic status is a CYP2D6 *1/ *1 genotype, which is consistent with an AS of 2.0 and a normal metabolizer status.

Clinical Implications





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There is substantial evidence linking the CYP2D6 polymorphisms to variability in the pharmacological and safety profiles of the following psychotropics: desipramine (Norpramin), imipramine (Tofranil), amitriptyline (Elavil), nortriptyline (Pamelor), haloperidol (Haldol), trimipramine (Surmontil), venlafaxine (Effexor), doxepin (Silenor), aripiprazole (Abilify), atomoxetine (Strattera), risperidone (Risperdal), clomipramine (Anafranil), and pimozide (Orap).

CYP2D6 polymorphisms have been shown to affect the pharmacological and safety profiles of the following analgesics: codeine, tramadol (Ultram), and hydrocodone (Vicodin). Codeine and tramadol are prodrugs that need to be activated by CYP2D6. Poor metabolizers are at high risk of therapy failure when given codeine or tramadol. On the other hand, ultra-rapid metabolizers may experience increased toxicity when given standard dosage of codeine or tramadol. Because CYP3A4 is also involved in the metabolism of oxycodone, patients with abnormal CYP2D6 activity may still experience adequate analgesia when taking this drug. CYP2D6 polymorphism has been shown to affect dihydrocodeine (Synalgos-DC) pharmacokinetics and can potentially alter the response to this drug.

Morphine, oxymorphone (Opana), hydromorphone (Dilaudid), butorphanol (Stadol), fentanyl (Duragesic), buprenorphine (Butrans), methadone (Dolophine), morphine (Avinza), and tapentadol (Nucynta) are not substrates of CYP2D6, and the patient's response to these drugs is not expected to be affected by polymorphisms in this enzyme.

Several important cardiovascular medications are metabolized by CYP2D6, and include: metoprolol (Lopressor), flecainide (Tambocor), and propafenone (Rythmol).

Eliglustat (Cerdelga) is a glucosylceramide synthase inhibitor used for the long-term treatment of adult patients with Gaucher disease type 1. Both the FDA-approved drug label and the EMA Summary of Product Characteristics for this drug state that CYP2D6 ultra-rapid metabolizers may not achieve adequate concentrations of eliglustat to achieve a therapeutic effect. Therefore this drug should not be used in these individuals. Patients who are CYP2D6 intermediate and normal metabolizers have a recommended dose of 84 mg twice daily, while poor metabolizers have a recommended dose of 84 mg once daily.

Cevimeline (Evoxac) is a muscarinic agonist indicated for the treatment of symptoms of dry mouth in patients with Sjögrens Syndrome. Both CYP2D6 and CYP3A are responsible for the metabolism of cevimeline, and this drug should be used with caution in individuals who are CYP2D6 poor metabolizers, as they may be at a higher risk of adverse events.

Inhibitors of the CYP2D6 enzyme may modify its activity and change the patient's metabolizer status. This can result in drug-drug interactions when a drug substrate is prescribed with known CYP2D6 inhibitors. Although there are no known clinical inducers of CYP2D6, the pharmacokinetics of a drug substrate can be affected by inducers of other enzymes (such as CYP3A) that are involved in the metabolism of that drug.

Inhibitors

Some known **strong and moderate** CYP2D6 inhibitors include: abiraterone (Zytiga), bupropion (wellbutrin), cinacalcet (Sensipar), cobicistat (Stribild), dacomitinib (Vizimpro), duloxetine (Cymbalta), ecstasy, fluoxetine (Prozac), paroxetine (Paxil), quinidine (Quinidex), rolapitant (Varubi), terbinafine (Lamisil), mirabegron (Myrbetriq), panobinostat (Farydak), peginterferon alfa-2b (Sylatron) and tipranavir/ritonavir (Aptivus).

Some known **weak** CYP2D6 inhibitors include: amiodarone (Cordarone), celecoxib (Celebrex), clobazam (Onfi), desvenlafaxine (Pristiq), diltiazem (Cardiazem), diphenhydramine (Benadryl), Echinacea, escitalopram (Lexapro), febuxostat (Uloric), gefitinib (Iressa), hydralazine (Apresoline), hydroxychloroquine (Plaquenil), imatinib (Gleevec), lorcaserin (Belviq), methadone (Dolophine), perphenazine (Trilafon), propafenone (Rythmol), ranitidine (Zantac), ritonavir (Norvir), sertraline (Zoloft), telithromycin (Ketek), venlafaxine (Effexor) and verapamil (Isoptin, Covera-HS).





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1: Swen et al. Pharmacogenetics: from bench to byte--an update of guidelines. Clin Pharmacol Ther. 2011 May;89(5):662-73. 2: Zhou SF. Polymorphism of human cytochrome P450 2D6 and its clinical significance: part II. Clin Pharmacokinet. 2009;48(12):761-804. 3: Zhou SF. Polymorphism of human cytochrome P450 2D6 and its clinical significance: Part I. Clin Pharmacokinet. 2009;48(11):689-723. 4: Preskorn SH. Clinically important differences in the pharmacokinetics of the ten newer "atypical" antipsychotics: Part 2. Metabolism and elimination. J Psychiatr Pract. 2012 Sep;18(5):361-8. 5: Preskorn SH. Clinically important differences in the pharmacokinetics of the ten newer "atypical" antipsychotics: part 1. J Psychiatr Pract. 2012 May;18(3):199-204. 6: D'Empaire et al. Antidepressant treatment and altered CYP2D6 activity: are pharmacokinetic variations clinically relevant? J Psychiatr Pract. 2011 Sep;17(5):330-9. 7: Hicks et al. Clinical Pharmacogenetics Implementation Consortium Guideline for CYP2D6 and CYP2C19 Genotypes and Dosing of Tricyclic Antidepressants. Clin Pharmacol Ther. 2013 Jan 16. 8: Gaedigk et al. The CYP2D6 activity score: translating genotype information into a gualitative measure of phenotype. Clin Pharmacol Ther. 2008 Feb;83(2):234-42. 9: Crews et al. Clinical Pharmacogenetics Implementation Consortium (CPIC) guidelines for codeine therapy in the context of cytochrome P450 2D6 (CYP2D6) genotype. Clin Pharmacol Ther. 2012 Feb;91(2):321-6. 10: Meyer et al. Absorption, distribution, metabolism and excretion pharmacogenomics of drugs of abuse. Pharmacogenomics. 2011 Feb;12(2):215-3. 11: Evoxac FDA Prescribing Label. 12: Cerdelga FDA Prescribing Label. 13: Brown JT, Bishop JR, Sangkuhl K, Nurmi EL, Mueller DJ, Dinh JC, Gaedigk A, Klein TE, Caudle KE, McCracken JT, de Leon J, Leeder JS. Clinical Pharmacogenetics Implementation Consortium Guideline for Cytochrome P450 (CYP)2D6 Genotype and Atomoxetine Therapy. Clin Pharmacol Ther. 2019 Feb 22. 14: Pharmacogene Variation (PharmVar) Consortium. PharmVar CYP2D6. https://www.pharmvar.org/gene/CYP2D6. Published 2019. Accessed January 6, 2020. 15: Nofziger C, Turner AJ, Sangkuhl K, et al. PharmVar GeneFocus: CYP2D6. Clin Pharmacol Ther. 2020;107(1):154-170. 16: Gaedigk A. Complexities of CYP2D6 gene analysis and interpretation. Int Rev Psychiatry. 2013;25(5):534-553. 17: Pharmacogene Variation (PharmVar) Consortium. Structural Variation CYP2D6. 2019:9. https://a.storyblok.com/f/70677/x/2de9d1f5e1/cyp2d6_structural-variation_v1-7.pdf. Accessed January 6, 2020. 18: Caudle KE, Sangkuhl K, Whirl-Carrillo M, et al. Standardizing CYP2D6 Genotype to Phenotype Translation: Consensus Recommendations from the Clinical Pharmacogenetics Implementation Consortium and Dutch Pharmacogenetics Working Group. Clin Transl Sci. October 2019.

CYP3A4 Monograph

Clinical Utility

The cytochrome P450 3A4 and 3A5 (CYP3A4 and CYP3A5) account for 40-80% of total CYP in human liver and intestine, respectively. Most importantly, CYP3A enzymes metabolize 50% of commonly used drugs. CYP3A4 and CYP3A5 enzymes have overlapping substrate specificity, and the contribution of CYP3A5 in the overall metabolism is smaller than the one for CYP3A4. The overall CYP3A metabolism status is expected to affect drugs that have a narrow therapeutic index.

Assay Interpretation

A limited number of variants identified within the CYP3A4 and CYP3A5 genes have been associated with significant alterations in enzyme activity and subsequent variability in therapeutic response. For CYP3A5, individuals with the less prevalent "normal metabolizer phenotype" may metabolize drugs faster than those with the more common "poor metabolizer phenotype". This may result in increased toxicity or loss of efficacy.

The CYP3A4*1B variant is the most studied, and results in an enzyme with a moderately decreased activity. It occurs in 50% of African-Americans, 3-5% of Caucasians, and <1% of Asians. The CYP3A4*2, *3, *12, and *17 are also considered decreased activity alleles. Recently, the CYP3A4 *22 allele has been characterized as a decreased function allele that can be clinically relevant (associated with a decreased clearance of certain substrates). The genotype-phenotype relationship for CYP3A4 is not well established, and individuals are predicted to have a CYP3A4 normal or intermediate metabolic capacity.

The reference range for CYP3A4 metabolic status is CYP3A4 *1/ *1, which is consistent with a normal metabolizer.

The variant results in an enzyme with no activity, and is the most common variant in the general population. This variant is found on all the CYP3A5*3 alleles. The CYP3A5 *6 and *7 are also no function alleles. The functional effects of the CYP3A5 alleles *2, *4, *5 *8, and *9 are not well established. The CYP3A5 *1 functional allele produces an active enzyme, and is found in 5% of Caucasians, 20% of Asians, and 15-50% of Africans. Individuals with two CYP3A5 no function alleles are classified as poor metabolizers while those carrying one copy of a functional CYP3A5*1 allele are considered intermediate metabolizers. A subject carrying two copies of the functional CYP3A5*1 allele is considered a normal metabolizer. **CYP3A5 poor metabolizers represent 50% of Asians and 90% of Caucasians.**

The reference range for CYP3A5 metabolic status is CYP3A5 *1/*1, which is consistent with a normal metabolizer. This genotype is the least prevalent in Caucasians and Asians.

Clinical Implications



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CYP3A4 and CYP3A5 genotypes can help identify patients with high or low overall CYP3A activity. Although these two enzymes metabolize many drugs, the response of only a few (such as narrow therapeutic index drugs) is expected to change significantly by genetic polymorphisms. Fentanyl (Duragesic) is a narrow therapeutic drug that is mainly metabolized by CYP3A. There is limited evidence suggesting that the response to this drug is altered in individuals with abnormal CYP3A activity.

The following drugs used in pain management and various psychiatric conditions are metabolized extensively by CYP3A: fentanyl (Duragesic), oxycodone (Oxycontin), buprenorphine (Suboxone), carbamazepine (Tegretol), quetiapine (Seroquel), ziprasidone (Geodon), alprazolam (Xanax), midazolam (Versed), triazolam (Halcion), nefazodone (Serzone), trazodone (Oleptro), vilazodone (Vibryd), zaleplon (Sonata), and zolpidem (Ambien). CYP3A contributes to a small extent in the elimination of methadone (Dolophine).

Within the major therapeutic classes used in cardiovascular conditions, the following drugs are substantially metabolized by CYP3A: atorvastatin (Lipitor), simvastatin (Zocor), lovastatin (Mevacor), nifedipine (Procardia), verapamil (Verelan), nicardipine (Cardene), felodipine (Plendil), nisoldipine (Sular), clopidogrel (Plavix), prasugrel (Effient), ticagrelor (Brilinta), cilostazol (Pletal), amiodarone (Cordarone), quinidine (Qualaquin), disopyramide (Norpace), losartan (Cozaar), rivaroxaban (Xarelto), and apixaban (Eliquis).

CYP3A metabolism is highly sensitive to inhibition and induction when a patient is taking multiple drugs. In this case, occurrence of drug-drug interactions can have profound effects on the pharmacokinetics, as well as the responses and safety profiles of many CYP3A drug substrates.

Inhibitors

Some known **strong** CYP3A inhibitors include: boceprevir (Victrelis), clarithromycin (Biaxin), clofazimine (Lamprene), conivaptan (Vaprisol), grapefruit juice (high dose), idelalisib (Zydelig), itraconazole (Sporanox), ketoconazole (Nizoral), lopinavir, (Kaletra), nefazodone (Serzone), nelfinavir (Viracept), paritaprevir (Technivie), posaconazole (Noxafil), ribociclib (Kisqali), ritonavir (Norvir), saquinavir (Invirase), telaprevir (Incivek), telithromycin (Ketek), tipranavir (aptivus), troleandomycin (TAO) and voriconazole (Vfend).

Some known **moderate** CYP3A inhibitors include: amprenavir (agenerase), aprepitant (Emend), atazanavir (Reyataz), ciprofloxacin (Cipro), darunavir (Prezista), diltiazem (cardizem), erythromycin (Eryc), fosamprenavir (Lexiva), fluconazole (Diflucan), grapefruit juice (low dose), imatinib (Gleevec), quinupristin/dalfopristin (Synercid) and verapamil (Isoptin, Covera-HS).

Some known weak CYP3A inhibitors include: amiodarone (Cordarone), amlodipine (Norvasc), atorvastatin (Lipitor), bicalutamide (Casodex), cilostazol (Pletal), cimetidine (Tagamet), fluoxetine (Prozac), fluoxamine (Luvox), ranitidine (Zantac), ranolazine (Ranexa), sertraline (Zoloft) and ticagrelor (Brilinta).

Inducers

Some known **strong** CYP3A inducers include: apalutamide (Erleada), carbamazepine (Tegretol), enzalutamide (Xtandi), fosphenytoin (Cerebyx), lumacaftor (Orkambi), phenobarbital, phenytoin (Dilantin), primidone (Mysoline), rifampin (Rifadin), rifapentine (Priftin) and St. John's wort.

Some known **moderate** CYP3A inducers include: artemether (Coartem), bosentan (Tracleer), dabrafenib (Tafinlar), efavirenz (Sustiva), etravirine (Intelence), ivosidenib (tibsovo), modafinil (Provigil), nafcillin (Unipen), nevirapine (Viramune) and rifabutin (Mycobutin).

Some known **weak** CYP3A inducers include: aprepitant (Emend), clobazam (Onfi), dexamethasone (Decadron), Echinacea, fosamprenavir (Lexiva), lesinurad (Zurampic), methylprednisolone (Medrol), midostaurin (Rydapt), oxcarbazepine (Trileptal), pioglitazone (Actos) and rufinamide (Banzel).

References

1- Metabolic Drug Interactions. RH Levy, KE Thummel, WF Trager. Publisher: Lippincott Williams & Wilkins (March 15, 2000). 2: Zhou et al. Polymorphism of human cytochrome P450 enzymes and its clinical impact. Drug Metab Rev. 2009;41(2):89-295 3- Isoherranen et al. The influence of CYP3A5 expression on the extent of hepatic CYP3A inhibition is substrate-dependent: an in vitro-in vivo evaluation. Drug Metab Dispos. 2008 Jan;36(1):146-54. 4- Williams et al. A significant drug-metabolizing role for CYP3A5? Drug Metab Dispos. 2003 Dec;31(12):1526-30. 5- Elens et al. A new functional CYP3A4 intron 6 polymorphism significantly affects tacrolimus pharmacokinetics in kidney transplant recipients. Clin Chem. 2011 Nov;57(11):1574-83. 6- Elens et al. Effect of a new functional CYP3A4 polymorphism on calcineurin inhibitors' dose requirements and trough blood levels in stable renal transplant patients. Pharmacogenomics. 2011 Oct;12 (10):1383-96. 7-Lamba et al. Genetic contribution to variable human CYP3A-mediated metabolism. Adv Drug Deliv Rev. 2002 Nov 18;54(10):1271-94.

CYP3A5 Monograph

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The CYP3A4*1B variant is the most studied, and results in an enzyme with a moderately decreased activity. It occurs in 50% of African-Americans, 3-5% of Caucasians, and <1% of Asians. The CYP3A4*2, *3, *12, and *17 are also considered decreased activity alleles. Recently, the CYP3A4 *22 allele has been characterized as a decreased function allele that can be clinically relevant (associated with a decreased clearance of certain substrates). The genotype-phenotype relationship for CYP3A4 is not well established, and individuals are predicted to have a CYP3A4 normal or intermediate metabolic capacity.

The reference range for CYP3A4 metabolic status is CYP3A4 *1/ *1, which is consistent with a normal metabolizer.

The variant results in an enzyme with no activity, and is the most common variant in the general population. This variant is found on all the CYP3A5*3 alleles. The CYP3A5 *6 and *7 are also no function alleles. The functional effects of the CYP3A5 alleles *2, *4, *5 *8, and *9 are not well established. The CYP3A5 *1 functional allele produces an active enzyme, and is found in 5% of Caucasians, 20% of Asians, and 15-50% of Africans. Individuals with two CYP3A5 no function alleles are classified as poor metabolizers while those carrying one copy of a functional CYP3A5*1 allele are considered intermediate metabolizers. A subject carrying two copies of the functional CYP3A5*1 allele is considered a normal metabolizer. **CYP3A5 poor metabolizers represent 50% of Asians and 90% of Caucasians.**

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Some known **moderate** CYP3A inhibitors include: amprenavir (agenerase), aprepitant (Emend), atazanavir (Reyataz), ciprofloxacin (Cipro), darunavir (Prezista), diltiazem (cardizem), erythromycin (Eryc), fosamprenavir (Lexiva), fluconazole (Diflucan), grapefruit juice (low dose), imatinib (Gleevec), quinupristin/dalfopristin (Synercid) and verapamil (Isoptin, Covera-HS).

Some known **weak** CYP3A inhibitors include: amiodarone (Cordarone), amlodipine (Norvasc), atorvastatin (Lipitor), bicalutamide (Casodex), cilostazol (Pletal), cimetidine (Tagamet), fluoxetine (Prozac), fluoxamine (Luvox), ranitidine (Zantac), ranolazine (Ranexa), sertraline (Zoloft) and ticagrelor (Brilinta).

Inducers

Some known **strong** CYP3A inducers include: apalutamide (Erleada), carbamazepine (Tegretol), enzalutamide (Xtandi), fosphenytoin (Cerebyx), lumacaftor (Orkambi), phenobarbital, phenytoin (Dilantin), primidone (Mysoline), rifampin (Rifadin), rifapentine (Priftin) and St. John's wort.

Some known **moderate** CYP3A inducers include: artemether (Coartem), bosentan (Tracleer), dabrafenib (Tafinlar), efavirenz (Sustiva), etravirine (Intelence), ivosidenib (tibsovo), modafinil (Provigil), nafcillin (Unipen), nevirapine (Viramune) and rifabutin (Mycobutin).

Some known **weak** CYP3A inducers include: aprepitant (Emend), clobazam (Onfi), dexamethasone (Decadron), Echinacea, fosamprenavir (Lexiva), lesinurad (Zurampic), methylprednisolone (Medrol), midostaurin (Rydapt), oxcarbazepine (Trileptal), pioglitazone (Actos) and rufinamide (Banzel).

References

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Factor II Monograph

Clinical Utility

The F2 gene encodes the coagulation factor II, or prothrombin. It is a vitamin K–dependent proenzyme that functions in the blood coagulation cascade. It is a precursor to thrombin, which converts fibrinogen into fibrin, which in turn strengthens a protective clot.

The F2 c.*97G>A variant (also known as Factor II 20210G>A) in the F2 gene results in increased levels of plasma prothrombin and a concurrent increased risk for thrombosis. Prothrombin-related thrombophilia is characterized by venous thromboembolism (VTE). This risk of thrombosis is also increased when variants exist for other coagulation factors such as F5 c.1601G>A variant (also known as Factor V Leiden), or in presence of non-genetic risk factors such as obesity, injury, surgery, smoking, pregnancy, use of estrogen-containing contraceptives, or replacement therapy. The clinical expression of prothrombin-related thrombophilia is variable, and many individuals may never develop thrombosis, while others may experience venous thrombotic events or pregnancy complications.

Assay Interpretation

The Factor II thrombophilia is the second most common inherited risk factor for thrombosis. The F2 c.*97G>A variant (also known as Factor II 20210G>A) is associated with a hypercoagulable state. In the United States, the prevalence of this variant is 1.1% in Caucasians and Hispanics and 0.3% in African-Americans. The prevalence of heterozygosity is 2%-5% in whites and 0%-0.3% in African-Americans. The prevalence of homozygosity is approximately one in 10,000.

The reference range for F2 c.*97G>A variant is F2 c.*97G>A G/G.

Clinical Implications

The F2 c.*97G>A variant is associated with an elevation of plasma prothrombin levels to about 30% above normal in heterozygotes and to 70% above normal in homozygotes. Heterozygotes are at a 2- to 5-fold increased risk of an initial VTE. The risk for VTE in F2 c.*97G>A homozygotes is not well defined but is presumed to be higher than in F2 c.*97G>A heterozygotes. F2 c.*97G>A homozygotes tend to develop thrombosis more frequently and at a younger age. Individuals who are doubly heterozygotes for F5 c.1601G>A variant and F2 c.*97G>A variant have an estimated 20-fold increased risk when compared to individuals without either variant suggesting a multiplicative elevation in risk. Certain circumstantial factors can increase the risk of thrombosis, and include: travel, central venous catheter use, pregnancy, oral contraceptive use, hormone replacement therapy (HRT), selective estrogen receptor modulators (SERMs), organ transplantation, injury, age, and surgery.

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PATIENT INFORMATION

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Factor V Leiden Monograph

Clinical Utility

The F5 gene encodes the coagulation factor V. In normal conditions, Factor V is inactivated during the clotting process by the activated protein C (APC). In subjects with Factor V Leiden thrombophilia, a variant in the gene produces a Factor V that cannot be inactivated normally by APC. As a result, the clotting process remains active longer than usual, leading to more thrombin generation. This hypercoagulable state is also increased when other variants exist in other coagulation factors such as F2 c.*97G>A variant (also known as Factor II 20210G>A), or in the presence of non-genetic risk factors such as obesity, injury, surgery, smoking, pregnancy, or use of estrogen-containing contraceptive or estrogen containing replacement therapy. The clinical expression of Factor V Leiden thrombophilia is variable. Many individuals may never develop thrombosis, while others may experience venous thrombotic events or pregnancy complications. Certain circumstantial factors can increase the risk of thrombosis, and include: travel, central venous catheter use, pregnancy, oral contraceptive use, hormone replacement therapy (HRT), selective estrogen receptor modulators (SERMs), organ transplantation, injury, age, and surgery. These factors are associated with the first thrombotic episode in at least 50% of individuals with a F5 c.1601G>A variant.

Assay Interpretation

The F5 c.1601G>A variant (also known as Factor V Leiden) is the most common known inherited risk factor for thrombosis. The F5 c.1601G>A variant refers to a base change (from G to A at position 1691) in the gene. As a result, Factor V is inactivated to a lesser extent and persists for longer in the circulation, leading to hypercoagulability. In the US, the frequency of the F5 c.1601G>A variant varies by ethnicity, with about 5% of Caucasians, 2% of Hispanics, and 1% of African-Americans having one variant. Only 1 in 5000 individuals have two F5 c.1601G>A variants.

The reference range for F5 c.1601G>A variant is F5 c.1601G>A G/G.

Clinical Implications

About 1 in 1000 people in the U.S. experience a first venous thromboembolism (VTE) each year. VTE is caused by inherited and environmental factors, and while the F5 c.1601G>A variant is present in only 15-20% of individuals with a first VTE, it is found in 50% of individuals with recurrent VTE or estrogen-related thrombosis. The risk for VTE is increased 3- to 8-fold in F5 c.1601G>A heterozygotes and 9- to 80-fold in homozygotes. This risk is increased further if other genetic or circumstantial factors are present. A heterozygote individual for both the F5 c.1601G>A variant and the F2 c.*97G>A variant (compound heterozygote) has an even greater risk of VTE (20-fold) than an individual with a variant in only one factor. This illustrates the multiplicative effect of these two factors on overall thrombotic risk.

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MTHFR Monograph

Clinical Utility

Methylenetetrahydrofolate reductase (MTHFR) is involved in folate metabolism and is essential for the remethylation of homocysteine. Two common variants in the MTHFR gene: c.665C>T (legacy name 677C>T) and c.1286A>C (legacy name 1298A>C), result in an enzyme with decreased activity, which is linked to increased plasma homocysteine levels (i.e. hyperhomocysteinemia). Mild to moderate hyperhomocysteinemia has been identified as a risk factor for venous thromboembolism and other cardiovascular diseases such as coronary heart disease and stroke. Other conditions in which hyperhomocysteinemia is found, include recurrent pregnancy loss, placental infarction and birth defects. However, the causal role of MTHFR variants in these conditions is not well established.





SPECIMEN DETAILS

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Assay Interpretation

The approximate minor allele frequencies for most populations are 30-50% for the MTHFR c.1286A>C variant and 18-30% for the MTHFR c.665C>T variant. Heterozygotes and homozygotes for the MTHFR c.665C>T variant have 60% and 30% of normal MTHFR activity, respectively. Heterozygotes and homozygotes for the MTHFR c.1286A>C variant have 80% and 60% of normal MTHFR activity, respectively.

The reference ranges for both variants of MTHFR are c.665C>T C/C and c.1286A>C A/A. This is consistent with a normal MTHFR activity.

Clinical Implications

The MTHFR assay provides information about potential causes of elevated homocysteine, and approaches for addressing it.

- Homozygosity for the MTHFR c.665C>T variant (individual with MTHFR c.665C>T T/T genotype) predisposes for hyperhomocysteinemia (especially during times of folate insufficiency) and an increase in premature cardiovascular disease. Measurement of total plasma homocysteine is informative in this case.
- Compound heterozygosity (individual with MTHFR c.665C>T C/T and MTHFR c.1286A>C A/C genotypes) is not associated with an increase in plasma homocysteine level. Measurement of total plasma homocysteine is informative in this case.
- Heterozygosity or homozygosity for the MTHFR c.1286A>C A>C variant alone (individual with MTHFR c.1286A>C A/C or MTHFR c.1286A>C C/C genotypes) does not increase homocysteine levels. Similarly, heterozygosity for the MTHFR c.665C>T variant alone (individuals with MTHFR c.665C>T C/T genotype) does not increase homocysteine levels.
- Hyperhomocysteinemia related to MTHFR genetic variants has been associated with neural tube defects, stillbirths and recurrent pregnancy loss. However, because hyperhomocysteinemia is multifactorial, involving a combination of other genetic, physiologic and environmental factors, the presence of MTHFR variants in an individual should not be used alone to predict the risk of these conditions.
- MTHFR in depression: Low MTHFR activity may exacerbate folate deficiency in patients with depression and may increase the risk of depressive relapse or delay the response to antidepressants. Testing for homocysteine levels and serum folate levels are recommended in patients with depression. Methylfolate may substantially benefit patients with hyperhomocysteinemia and depression when used as an adjuvant to antidepressant medication.
- The response to methotrexate, a drug used in cancer and autoimmune diseases, is affected by the presence of MTHFR genetic variants. Methotrexate intolerance is observed in individuals that are heterozygous or homozygous for the MTHFR c.665C>T variant.

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OPRM1 Monograph

Clinical Utility

"Mu" opioid receptors are the most important site of action of opioid drugs. Single polymorphisms in the human mu-opioid receptor (OPRM1) have been investigated for their role in human nociception, opiate efficacy, and addiction.

Assay Interpretation

The variant mostly studied is a single substitution at position 118, from an adenine to a quanine (A118G). This variant reduces the OPRM1 receptor signaling efficiency induced by exogenous opioids. Reduced OPRM1 mRNA expression levels were observed in carriers of the G variant. The variant allele (G) is present in 7-15% of Caucasians, 1.5% of African-Americans, and up to 48.5% of Asians. The major interest of this particular SNP is due to its pharmacological and physiological consequences; however the exact mechanism by which the altered receptor influences opioid analgesia is still unresolved. The presence of the G allele seems to reduce the effect of exogenous agonists but increase the effects of exogenous antagonists.

The reference range for the A118G SNP is A118G AA, and is associated with a normal OPRM1 receptor signaling efficiency.

Clinical Implications

The presence of the G allele (A118G) seems to be associated with pain sensitivity as well as opioid dosage requirements. But only weak evidence of these associations is available to date. It is suggested that patients carrying the G allele report higher intensity pain. In terms of drug response, patients with the G allele have a favorable response to the anti-addictive drug naltrexone. Several studies conducted in postsurgical settings or in cancer analgesia showed that G allele carriers require slightly higher doses of morphine or fentanyl. This association still needs to be confirmed in larger studies and does not hold in other situations such as labor pain.

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SLCO1B1 Monograph

Clinical Utility

The SLCO1B1 gene encodes a liver-specific transporter involved in the removal of endogenous compounds (bile acids, bilirubin) and drugs such as statins from the blood to the liver. Some variants of the SLCO1B1 gene result in a low-functioning protein, which impairs statin clearance, and may lead to an increased risk of muscle pain, tenderness, or weakness, called myopathy. Certain medications can potently inhibit SLCO1B1, causing clinically significant drug interactions.

Assay Interpretation





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The functional SLCO1B1 activity (phenotype) is estimated from the genotype at 521T>C. Non-carriers of the 521T>C variant are predicted to have a SLCO1B1 normal function, those with one copy of the 521T>C variant have a SLCO1B1 decreased function, and those with two copies of the variant have poor SLCO1B1 function.

There are several variants of the SLCO1B1 that define over 15 alleles. One relatively common variant 521T>C (rs4149056) results in SLCO1B1 decreased function, which affects the transport of drug substrates such as statins. This variant is present alone on the *5 allele and in presence with another variant (388A>G; rs2306283) on the *15 allele. Both alleles are low-activity alleles (reduced hepatic uptake), and have a combined frequency of 15-20% in Caucasians, 10-15% in Asians, and 2% in sub-Saharan Africans and African-Americans.

The reference range for the 521T>C mutation of SLCO1B1 is 521 TT. This is consistent with normal SLCO1B1 function.

Clinical Implications

All statins are substrates of SLCO1B1, but the effects of SLCO1B1 genetic polymorphism differ between individual statins. The effect is the largest on simvastatin, and individuals with the 521T>C variant have increased levels of the active simvastatin form. The variant is strongly associated with simvastatin-induced myopathy (with or without CK elevation), especially with high-dose simvastatin therapy. More than 60% of the myopathy cases could be attributed to its presence. The clinical spectrum of statin-induced myopathy ranges from a mild and common myalgia to a life-threatening and rare rhabdomyolysis. Other known risk factors for statin-induced myopathy include a high-statin dose, interacting drugs that raise statin levels, age, hypothyroidism, and certain inherited muscle disorders.

At therapeutic doses, the apparent sensitivity levels of the five statins to the presence of the 521T>C variant are

simvastatin>pitavastatin>atorvastatin>pravastatin>rosuvastatin. Carriers of the 521 T>C variant should avoid high-dose simvastatin therapy. These patients can take other statins, such as atorvastatin, pitavastatin, rosuvastatin, or pravastatin, but at reduced doses. Fluvastatin is not affected by the 521 T>C variant and could therefore be considered a suitable alternative.

Other drugs that are substrates of SLCO1B1 transporter include enalapril, olmesartan, valsartan, atrasentan, repaglinide, nateglinide, methotrexate, and bosentan. However, there is insufficient evidence documenting the impact of the 521 T>C variant on the systemic exposure and safety profile of these drugs.

Inhibitors

Inhibitors of SLCO1B1 transporter may alter its activity and result in increased levels of drug substrates. These include boceprevir (Victrelis), clarithromycin (Biaxin), cyclosporine (Gengraf, Neoral, Restasis, Sandimmune), eltrombopag (Promacta), gemfibrozil (Lopid), paritaprevir (Technivie), protease inhibitors, simeprevir (Olysio), telaprevir (Incivek) and teriflunomide (Aubagio).

Inducers

Inducers of SLCO1B1 transporter include: apalutamide (Erleada).

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VKORC1 Monograph

Clinical Utility

The Vitamin K epoxide reductase complex, subunit 1 (VKORC1) is the target of anticoagulants such as warfarin, acenocoumarol and phenprocoumon. VKORC1 catalyzes the conversion of oxidized Vitamin K to reduced Vitamin K. By inhibiting this conversion, warfarin leads to decreased availability of the reduced Vitamin K that serves as a cofactor for gamma-glutamyl carboxylase and blocks the formation of functionally-active clotting factors, resulting in reduced coagulation. Polymorphisms within the VKORC1 gene result in variable expression levels of the VKORC1 enzyme and altered response towards coumarin anticoagulants. VKORC1 genetic testing defines three clinical phenotypes: high, moderate, and low sensitivity to warfarin. VKORC1 genetic testing is usually used in conjunction with CYP2C9 and CYP4F2 genetic testing to optimize warfarin dosing and minimize the risks of bleeding or thrombotic complications.

Assay Interpretation

Several relevant variants in the VKORC1 gene have been identified and correlate well with warfarin dose requirements. In Europeans, the genotype for a single variant (-1694G>A; rs9923231) accounts for 25% of variability in warfarin dose requirements, with the minor A allele associated with lower doses. This association is also observed in East Asians, where the minor allele occurs more frequently than in Europeans (88% vs 41%). In Africans, the minor allele frequency for VKORC1 -1639G>A is 10% which is considerably lower than in Europeans. Several studies in Africans and African-Americans identified common variants that are associated with higher warfarin dose requirements in patients of African descent.

The reference range for VKORC1 -1639G>A is G/G and is associated with a normal VKORC1 enzyme expression phenotype.

Clinical Implications

The common VKORC1 -1639G>A variant is associated with low-dose warfarin requirements. When CYP2C9 and VKORC1 genotypes are combined with other patient-specific parameters such as demographic (age, weight, height), clinical (disease, co-medications), and environmental factors (smoking), they account for 50% of warfarin dose variation between individuals. Several validated warfarin pharmacogenetic dosing algorithms have been developed and are available to calculate initial and maintenance warfarin doses in both adults and children. Moreover, the FDA prescribing label for warfarin includes genotype-specific dose ranges based on CYP2C9 and VKORC1 genotypes.

References

1: Food and Drug Administration: Coumadin® Label accessed on Jan 2013. 2: Gage et al. Use of pharmacogenetic and clinical factors to predict the therapeutic dose of warfarin. Clin Pharmacol Ther. 2008 Sep;84(3):326-31 3: Schelleman et al. New genetic variant that might improve warfarin dose prediction in African Americans. Br J Clin Pharmacol. 2010 Sep;70(3):393-9 4: Johnson et al. Clinical Pharmacogenetics Implementation Consortium. Clinical Pharmacogenetics Implementation Consortium Guidelines for CYP2C9 and VKORC1 genotypes and warfarin dosing. Clin Pharmacol Ther. 2011 Oct;90(4):625-9. 5: Klein et al. Estimation of the warfarin dose with clinical and pharmacogenetic data. International Warfarin Pharmacogenetics Consortium. N Engl J Med. 2009 Feb 19;360 (8):753-64 6: Rieder et al. Effect of VKORC1 haplotypes on transcriptional regulation and warfarin dose. N Engl J Med. 2005 Jun 2;352(22):2285-93. 7: Johnson JA, Caudle KE, Gong L, Whirl-Carrillo M, Stein CM, Scott SA, Lee MT, Gage BF, Kimmel SE, Perera MA, Anderson JL, Pirmohamed M, Klein TE, Limdi NA, Cavallari LH, Wadelius M. Clinical Pharmacogenetics Implementation Consortium (CPIC) Guideline for Pharmacogenetics-Guided Warfarin Dosing: 2017 Update. Clin Pharmacol Ther. 2017 Sep;102(3):397-404.





DAT	ENIT	INFO	DM		\mathbf{n}
PAL		INFU	KW	АП	UI

NAME: Patient s7akg3i ACC #: s7akg3i DOB: 1/1/1900 SEX: SPECIMEN DETAILS

SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: 11/11/2022

FOR ACADEMIC PURPOSES ONLY - NOT FOR CLINICAL USE

Patient Information Card

This is a summary genetic report for your patient to share with other healthcare providers. The card can be cut out along the dashed line and carried with the patient.

Manchester REPORT DETAILS Diversity Patient: Patient s7akg3i DOB: 1/1/1900 ACC #: s7akg3i Pharmacogenetic Test Summary		REPORT DETAILS					
		Patient: Patient s7akg3i	VKORC1	-1639G>A G/A Intermediate Warfarin Ser			
			MTHFR		ased Risk of mocysteinemia		
		MTHFR	c.665C>T CT Reduced	MTHFR Activity			
CYP2C19	*1/*2	Intermediate Metabolizer		plete report contact Manchester University Master of in Pharmacogenomics Program			
CYP2C9	*1/*1	Normal Metabolizer	For a compl				
CYP2D6	Indeterminate	Unknown Phenotype		www.manchester.edu/pgx			
CYP3A4	*1/*22	Intermediate Metabolizer					
CYP3A5	*3/*3	Poor Metabolizer		softwar			

