

PATIENT INFORMATION

NAME: 656677961 ACC #: 656677961 DOB: 1/1/1900 SEX:

SPECIMEN DETAILS

ORDERED BY

SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: 9/1/2019

FOR ACADEMIC PURPOSES ONLY - NOT FOR CLINICAL USE

Complete Panel

Risk Management

Type III Hyperlipoproteinemia

Unknown Association with Type III Hyperlipoproteinemia

The patient's results for the APOE c.388 T>C (Cys130Arg) mutation and/or APOE c.526 C>T (Arg176Cys) mutations are indeterminate.

The patient's risk for type III hyperlipoproteinemia cannot be estimated accurately.

Consult with a genetic counselor for more information.

\checkmark

Hyperhomocysteinemia - Depression

No Increased Risk of Hyperhomocysteinemia

The patient does not carry the MTHFR C677T variant. MTHFR enzyme activity is normal.

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. With a normal MTHFR activity, this patient can process folate normally and is unlikely to have elevated plasma levels of homocysteine.

<u>Patients diagnosed with depression</u>: 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

No Increased Risk of Thrombosis

The patient does not carry the Factor V Leiden G1691A mutation or Factor II G20210A mutation (wild-type).

The patient's risk of thrombosis is not increased (average risk of clotting is about 1 in 1000 for anyone in a year). However, because this test cannot find all of the inherited reasons for abnormal clotting, other factors may affect this risk assessment.

Unless other genetic and/or circumstantial risk factors are present (e.g., smoking or obesity...), estrogen-containing contraceptive and hormone replacement therapy can be used by the patient.

Hyperhomocysteinemia - Thrombosis

No Increased Risk of Hyperhomocysteinemia

The patient carries one MTHFR A1298C mutation (heterozygous). MTHFR enzyme activity is reduced (80% of normal activity). The patient's slightly reduced MTHFR activity is not a risk factor for hyperhomocysteinemia. Unless other risk factors are present, the patient is not

expected to have an increased risk for venous thromboembolism (VTE). The patient's MTHFR activity is slightly reduced.

| A medication has potentially reduced efficacy, increased toxicity or the patient has an increased risk for the indicated condition. Guidelines exist for adjusting dosage, increased vigilance or the patient has a moderate risk for the indicated condition. | ACTIONABLE | Recommendations based upon publications by international pharmacogenetic expert groups, consortia or regulatory bodies (CPIC, DPWG, FDA. EMA). Recommendations are suitable for implementation in a clinical setting. Guidelines may change as knowledge arises. |
|---|-------------|--|
| The medication can be prescribed according to standard regimens or the patient's risk for the indicated condition is not increased. | INFORMATIVE | There are insufficient or contradictory findings documenting the impact of a given genetic polymorphism or drug interaction. Recommendations are informative and implementation in a clinical setting is optional. |





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SPECIMEN TYPE: **COLLECTION DATE:** 1/1/1900 RECEIVED DATE:

1/1/1900

REPORT DATE: 2/8/2018

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) | | |
| | Antianginal Agents | Ranolazine (Ranexa) | | |
| | Antiarrhythmics | | Flecainide (Tambocor) Mexiletine (Mexitil) Propafenone (Rythmol) | |
| 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) Carvedilol (Coreg) Labetalol (Normodyne, Trandate) Propranolol (Inderal) | Metoprolol (Lopressor) Nebivolol (Bystolic) Timolol (Timoptic) | |
| | Diuretics | Torsemide (Demadex) | | |
| | Statins | Atorvastatin (Lipitor) Fluvastatin (Lescol) Lovastatin (Mevacor, Altoprev, Advicor) Pitavastatin (Livalo) Pravastatin (Pravachol) Rosuvastatin (Crestor) Simvastatin (Zocor) | | |
| | Meglitinides | Nateglinide (Starlix) Repaglinide (Prandin, Prandimet) | | |
| Diabetes | Sulfonylureas | Chlorpropamide (Diabenese) Glimepiride (Amaryl) Glipizide (Glucotrol) Glyburide (Micronase) Tolbutamide (Orinase) | | |



| V Un | anchest iversity | ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN DETAILSSPECIMEN TYPE:COLLECTION DATE:1/1/1900RECEIVED DATE:1/1/1900REPORT DATE:2/8/2018 | ORDERED BY |
|------------------|------------------------|--|--|-----------------------|
| CATEGORY | DRUG CLASS | STANDARD PRECAUTIONS | USE WITH CAUTION | CONSIDER ALTERNATIVES |
| Gastrointestinal | Antiemetics | Aprepitant (Emend-oral) Dolasetron (Anzemet) Dronabinol (Marinol) Fosaprepitant (Emend-i.v) Granisetron (Sancuso, Sustol) Netupitant-Palonosetron (Akynzeo) Ondansetron (Zofran, Zuplenz) Palonosetron (Aloxi) Rolapitant (Varubi) | Metoclopramide (Reglan) | |
| | Proton Pump Inhibitors | Rabeprazole (Aciphex) | Dexlansoprazole (Dexilant, Kapidex) Esomeprazole (Nexium) Lansoprazole (Prevacid) Omeprazole (Prilosec) Pantoprazole (Protonix) | |
| 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) Raltegravir (Isentress, Dutrebis) | | |
| | Antimalarials | Proguanil (Malarone) | | |





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|----------|---------------------|--|--|-----------------------|
| | Fibromyalgia Agents | Milnacipran (Savella) | | |
| | Muscle Relaxants | Cyclobenzaprine (Flexeril, Amrix) Metaxalone (Skelaxin) Methocarbamol (Robaxin) | Carisoprodol (Soma) 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) | | |
| | Opioids | Alfentanil (Alfenta) Buprenorphine (Butrans, Buprenex) Dihydrocodeine (Synalgos-DC) Fentanyl (Actiq) Hydromorphone (Dilaudid, Exalgo) Levorphanol (Levo Dromoran) Meperidine (Demerol) Methadone (Dolophine) Morphine (MS Contin) Oxymorphone (Opana, Numorphan) Sufentanil (Sufenta) Tapentadol (Nucynta) | Codeine (Codeine; Fioricet with Codeine) Hydrocodone (Vicodin) Oxycodone (Percocet, Oxycontin) Tramadol (Ultram) | |
| | Antiaddictives | Bupropion (Wellbutrin, Zyban, Aplenzin, Contrave) | Naltrexone (Vivitrol, Contrave) | |
| | Anti-ADHD Agents | Amphetamine (Adderall, Evekeo) Atomoxetine (Strattera) Clonidine (Kapvay) Dextroamphetamine (Dexedrine) Guanfacine (Intuniv) Lisdexamfetamine (Vyvanse) | Dexmethylphenidate (Focalin) Methylphenidate (Ritalin, Aptensio XR, Concerta, Metadate ER, Quillivant ER) | |



| (X) Manchester | | | SPECIMEN DETAILS | ORDERED BY |
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| CATEGORY | DRUG CLASS | STANDARD PRECAUTIONS | USE WITH CAUTION | CONSIDER ALTERNATIVES |
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| | Anticonvulsants | Brivaracetam (Briviact) 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) Phenobarbital (Luminal) Phenytoin (Dilantin) Pregabalin (Lyrica) Primidone (Mysoline) Rufinamide (Banzel) Tiagabine (Gabitril) Topiramate (Topamax) Valproic Acid (Depakote, Depakene) Vigabatrin (Sabril) Zonisamide (Zonegran) | | |
|--------------|---------------------|---|--|--|
| Psychotropic | Antidementia Agents | Donepezil (Aricept) Galantamine (Razadyne) Memantine (Namenda) | | |
| | Antidepressants | Desvenlafaxine (Pristiq) Duloxetine (Cymbalta) Fluoxetine (Prozac, Sarafem) Fluvoxamine (Luvox) Levomilnacipran (Fetzima) Mirtazapine (Remeron) Nefazodone (Serzone) Paroxetine (Paxil, Brisdelle) Trazodone (Oleptro) Vilazodone (Viibryd) Vortioxetine (Trintellix) | Amoxapine (Amoxapine) Desipramine (Norpramin) Maprotiline (Ludiomil) Nortriptyline (Pamelor) Protriptyline (Vivactil) Sertraline (Zoloft) | Amitriptyline (Elavil) Citalopram (Celexa) Clomipramine (Anafranil) Doxepin (Silenor) Escitalopram (Lexapro) Imipramine (Tofranil) Trimipramine (Surmontil) Venlafaxine (Effexor) |
| | Antipsychotics | Aripiprazole (Abilify, Aristada) Asenapine (Saphris) Brexpiprazole (Rexulti) Cariprazine (Vraylar) Chlorpromazine (Thorazine) Haloperidol (Haldol) Loxapine (Loxitane, Adasuve) Lurasidone (Latuda) Paliperidone (Invega) Pimavanserin (Nuplazid) Pimozide (Orap) Quetiapine (Seroquel) Thiothixene (Navane) Trifluoperazine (Stelazine) Ziprasidone (Geodon) | Clozapine (Clozaril) Fluphenazine (Prolixin) Iloperidone (Fanapt) Olanzapine (Zyprexa) Perphenazine (Trilafon) | Risperidone (Risperdal) Thioridazine (Mellaril) |



Genetic Test Results For Patient 33169



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|-----------------|---|--|--------------------------|------------------------------|
| | Benzodiazepines | Alprazolam (Xanax) Clobazam (Onfi) Clonazepam (Klonopin) | Diazepam (Valium) | |
| | Other Neurological Agents | Deutetrabenazine (Austedo) Dextromethorphan / Quinidine (Nuedexta) Flibanserin (Addyi) Valbenazine (Ingrezza) | Tetrabenazine (Xenazine) | |
| Rheumatology | Anti-Hyperuricemics and Anti-Gout Agents | Colchicine (Mitigare) Febuxostat (Uloric) Lesinurad (Zurampic) | | |
| Kileumatology | Immunomodulators | Apremilast (Otezla) Leflunomide (Arava) Tofacitinib (Xeljanz) | | |
| Transplantation | Immunosuppressants | | Tacrolimus (Prograf) | |
| | 5-Alpha Reductase Inhibitors for Benign Prostatic Hyperplasia | Dutasteride (Avodart) Finasteride (Proscar) | | |
| | Alpha-Blockers for Benign Prostatic Hyperplasia | Alfuzosin (UroXatral) Doxazosin (Cardura) Silodosin (Rapaflo) Tamsulosin (Flomax) Terazosin (Hytrin) | | |
| Urologicals | Antispasmodics for Overactive Bladder | Darifenacin (Enablex) Fesoterodine (Toviaz) Mirabegron (Myrbetriq) Oxybutynin (Ditropan) Solifenacin (Vesicare) Tolterodine (Detrol) Trospium (Sanctura) | | |
| | Phosphodiesterase Inhibitors for Erectile Dysfunction | Avanafil (Stendra) Sildenafil (Viagra) Tadalafil (Cialis) Vardenafil (Levitra) | | |



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Dosing Guidance INFORMATIVE Amitriptyline Increased Sensitivity to Amitriptyline (CYP2C19: Rapid Metabolizer) Elavil Consider an alternative drug, or consider prescribing amitriptyline at standard dose and monitor the plasma concentrations of amitriptyline and nortriptyline to guide dose adjustments. ACTIONABLE Citalopram Insufficient Response to Citalopram (CYP2C19: Rapid Metabolizer) At standard label-recommended dosage, citalopram plasma concentrations levels are expected to be low which may Celexa result in a loss of efficacy. Consider an alternative medication. If citalopram is warranted, consider increasing the dose to a maximum of 150% and titrate based on the clinical response and tolerability. INFORMATIVE **Clomipramine** Increased Sensitivity to Clomipramine (CYP2C19: Rapid Metabolizer) Anafranil Consider an alternative drug, or consider prescribing clomipramine at standard dose and monitor the plasma concentrations of clomipramine and desmethyl-clomipramine to guide dose adjustments. INFORMATIVE Doxepin Increased Sensitivity to Doxepin (CYP2C19: Rapid Metabolizer) Silenor Consider an alternative drug, or consider prescribing doxepin at standard dose and monitor the plasma concentrations of doxepin and desmethyl-doxepin to guide dose adjustments. ACTIONABLE Escitalopram Insufficient Reponse to Escitalopram (CYP2C19: Rapid Metabolizer) At standard label-recommended dosage, escitalopram plasma concentrations levels are expected to be low which may Lexapro result in a loss of efficacy. Consider an alternative medication. If escitalopram is warranted, consider increasing the dose to a maximum of 150% and titrate based on the clinical response and tolerability. Imipramine Increased Sensitivity to Imipramine (CYP2C19: Rapid Metabolizer) INFORMATIVE Tofranil Consider an alternative drug, or consider prescribing imipramine at the standard dose and monitor the plasma concentrations of imipramine and desipramine to guide dose adjustments. ACTIONABLE Risperidone Increased Sensitivity to Risperidone (CYP2D6: Intermediate Metabolizer) Risperdal Consider an alternative drug, OR prescribe risperidone, be extra alert of adverse events, and adjust dosage in response to clinical response and tolerability. There is insufficient data to allow calculation of dose adjustment. ACTIONABLE Thioridazine Increased Sensitivity to Thioridazine (CYP2D6: Intermediate Metabolizer) Reduced cytochrome CYP2D6 activity results in elevated plasma levels of thioridazine, would be expected to augment the Mellaril prolongation of the QTc interval associated with thioridazine, and may increase the risk of serious, potentially fatal, cardiac arrhythmias, such as Torsades de pointes-type arrhythmias. Such an increased risk may result also from the additive effect of coadministering thioridazine with other agents that prolong the QTc interval. Therefore, thioridazine is contraindicated in patients with reduced levels of CYP2D6 activity. Trimipramine INFORMATIVE Increased Sensitivity to Trimipramine (CYP2C19: Rapid Metabolizer) Surmontil Consider an alternative drug, or consider prescribing trimipramine at standard dose and monitor the plasma concentrations of trimipramine and desmethyl-trimipramine to guide dose adjustments.



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| V | Manch Univer | | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: 1/1/1900 RECEIVED DATE: 1/1/1900 REPORT DATE: 2/8/2018 | |
| \sim | FOR ACADEMIC PURPOSES ONLY - NOT | FOR CLINICAL USE | | | |
| (\times) | Venlafaxine | Increased Sensitiv | ivity to Venlafaxine (CYP2D | 6: Intermediate Metabolizer) | ACTIONABLE |
| | Effexor | | ÷ . | xine, be extra alert of adverse events ethylvenlafaxine plasma concentratio | · · · |
| (\mathbf{X}) | Voriconazole | Non-Response to | o Voriconazole (CYP2C19: R | Rapid Metabolizer) | ACTIONABLE |
| Ŭ | Vfend | Voriconazole plasma response and effect | na concentrations are expected tiveness and subsequent diseas | to be low if a standard dose is used, se progression. Consider an alternativ iconazole, liposomal amphotericin B | ve medication that is not |
| | Amoxapine | Possible Sensitivi | vity to Amoxapine (CYP2D6 | : Intermediate Metabolizer) | INFORMATIVE |
| | Amoxapine | contribution of this in higher amoxapine | s enzyme in the metabolism of t ne concentrations potentially lea tients with decreased CYP2D6 f | amoxapine is metabolized by CYP2E this drug is not well documented. De ading to higher adverse events. There unction; therapy must be initiated ca | creased CYP2D6 activity may result e are no established dosing |
| <u>^</u> | Carisoprodol | Altered Sensitivit | ty to Carisoprodol (CYP2C1 | 9: Rapid Metabolizer) | INFORMATIVE |
| | Soma | | t data to allow calculation of do carefully monitor the patient fo | ise adjustment. If carisoprodol is pres or side effects. | scribed, it is recommended to use a |
| <u>^</u> | Clopidogrel | Increased Respon | nse to Clopidogrel (CYP2C1 | l9: Rapid Metabolizer) | ACTIONABLE |
| | Plavix | | prescribed at standard label-re eeding while taking clopidogre | commended dosage. Individuals wit I. | h the *17 allele may have an |
| | Clozapine | Unknown Respor | nse to Clozapine (CYP1A2: | Unknown Phenotype) | INFORMATIVE |
| | Clozaril | response to standar careful monitoring i | rd doses. There is an associatio is recommended during dosing events. Therefore, therapeutic o | cannot be predicted accurately, smo n between high clozapine doses and adjustment. Smoking cessation may drug monitoring accompanied by do | the risk of seizures, and therefore y increase plasma drug levels, |
| <u>^</u> | Codeine | Possible Non-Res | esponse to Codeine (CYP2D | 6: Intermediate Metabolizer) | ACTIONABLE |
| | Codeine; Fioricet with Codeine | Codeine can be pres insufficient pain relie | escribed at standard label-recor | patient may or may not experience a nmended dosage and administratior zed by CYP2D6 may also be consider orphone). | n, with monitoring for symptoms of |
| | Desipramine | Moderate Sensiti | ivity to Desipramine (CYP2 | D6: Intermediate Metabolizer) | ACTIONABLE |
| | Norpramin | | 5 | nmended standard starting dose. Mo gly until a favorable response is achie | • |
| <u>^!</u> | Dexlansoprazole | Insufficient Respo | oonse to Dexlansoprazole ((| CYP2C19: Rapid Metabolizer) | INFORMATIVE |
| | Dexilant, Kapidex | | | lose by 200% and be alert to insuffic use and consider dose increase of 200 | - |
| | | | | | |

| V | Manch Univers | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | | 1/1/1900 1/1/1900 2/8/2018 | |
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| | FOR ACADEMIC PURPOSES ONLY - NOT | FOR CLINICAL USE | | | | |
| <u>î</u> | Dexmethylphenid ate | Decreased Respo | onse to Dexmethylphenidat | te (COMT: Intermediate | e COMT Activity) | INFORMATIV |
| | Focalin | | ype result predicts a less optim eds and response of the patier | | | |
| | Diazepam | Possible Altered | Sensitivity to Diazepam (C | YP2C19: Rapid Metabo | lizer) | INFORMATIVI |
| | Valium | metabolizers. Howe | ultra-rapid metabolizers metal ever, there is insufficient data to 's response and adjust the dos | allow calculation of dose | | |
| Ŷ | Esomeprazole | Insufficient Resp | onse to Esomeprazole (CYF | 2C19: Rapid Metaboliz | er) | INFORMATIV |
| | Nexium | | er pylori eradication: increase d extra alert to insufficient respon | | | e. |
| | Flecainide | Increased Sensiti | vity to Flecainide (CYP2D6 | : Intermediate Metabo | izer) | ACTIONABL |
| _ | Tambocor | metabolizer may re | g a lower flecainide dose. When quire a 25% dose reduction. Ca recommended until a favorable | areful titration with ECG red | cording and monitoring of | |
| Â | Fluphenazine | Possible Sensitiv | ity to Fluphenazine (CYP2D | 06: Intermediate Metab | olizer) | INFORMATIV |
| | Prolixin | fluphenazine conc are no established of cautiously with oral dosage are apparent | tabolized by CYP2D6, CYP1A2 a entrations potentially leading dosing adjustments for patients or parenteral fluphenazine hyd nt, an equivalent dose of fluphe s may be necessary. | g to higher adverse even s with decreased CYP2D6 f drochloride. When the pha | ts such as extrapyramida unction therefore, therapy rmacological effects and a | al symptoms. There win must be initiated an appropriate |
| Ŵ | Hydrocodone | Possible Altered | Response to Hydrocodone | (CYP2D6: Intermediate | e Metabolizer) | INFORMATIV |
| | Vicodin | Decreased conversi intermediate metab taking hydrocodon | on of hydrocodone to the more polizers. However, there is insuf e. Adequate pain relief can be a plized by CYP2D6 may also be c | e active metabolite hydror ficient evidence whether tl achieved by increasing the | norphone is expected in C nese patients have decreas dose in response to pain | sed analgesia when symptoms. Other |
| | lloperidone | Moderate Sensit | ivity to lloperidone (CYP2D | 6: Intermediate Metab | olizer) | ACTIONABLE |
| | Fanapt | reduced CYP2D6 ac patients taking ilop | e is associated with QTc prolon tivity. Iloperidone must be titra eridone experience symptoms ins, or syncope), the prescriber | ited slowly from a low star that could indicate the occ | ting dose to avoid orthost currence of cardiac arrhyth | atic hypotension. If imias (e.g., |
| Δ | Lansoprazole | Insufficient Resp | onse to Lansoprazole (CYP | 2C19: Rapid Metabolize | er) | INFORMATIVE |
| <u>/!</u> \ | Prevacid | | | lose by 200% and be alert | | |

| | Manch Univer | lester | PATIENT INFORMATION NAME: Patient 33169 NAME: 23160 | SPECIMEN DETAILS | | ORDERED BY |
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| \wedge | Maprotiline | Possible Sensitiv | ity to Maprotiline (CYP2D6: | Intermediate Metab | olizer) | INFORMATIV |
| <u>·</u> · · · | Ludiomil | Like other tricyclic a CYP2D6 activity res established dosing dosage and gradua | and tetracyclic antidepressants, ults in higher maprotiline conce adjustments for patients with d illy adjusted according to the pa maintenance therapy. | maprotiline is metabolize ntrations potentially lead ecreased CYP2D6 functic | ed by CYP2D6 a ding to higher a on therefore, the | as well as CYP1A2. Decreased adverse events. There are no erapy must be initiated at a lov |
| <u>^</u> | Methylphenidate Ritalin, Aptensio XR, Concerta, Metadate ER, Quillivant ER | The patient's genot | onse to Methylphenidate (C type result predicts a less optima eeds and response of the patien | al response to methylphe | enidate. Dosage | e should be individualized |
| <u>^</u> | Metoclopramide Reglan | There is no data do metabolizers. Meto | ity to Metoclopramide (CYF ocumenting the changes in plasm iclopramide can be prescribed a sible increase of side effects. | na concentrations of me | toclopramide ir | |
| <u>^</u> | Metoprolol Lopressor | Based on the genot dosage. <u>Heart Failu</u> lower dose. When c <u>Other indication</u> s: C dose. When compa | ivity to Metoprolol (CYP2De type result, this patient may be a tre: Consider alternative beta-blu compared to a normal metaboliz Consider alternative beta-blocke ired to a normal metabolizer, an ribed, be alert to adverse events | at risk of excessive beta- ockers such as bisoprolo zer, an intermediate met rs such as bisoprolol or a intermediate metabolize | blockade when l or carvedilol, o abolizer may re atenolol, or pre er may require a | or prescribe metoprolol at a equire a 50% dose reduction. scribe metoprolol at a lower |
| Ŷ | Mexiletine | Increased Sensiti | ivity to Mexiletine (CYP2D6 | Intermediate Metab | olizer) | ACTIONABL |
| | Mexitil | | g a lower mexiletine dose. A slo recommended until a favorable | | - | nitoring of mexiletine plasma |
| Ŵ | Naltrexone | Altered Response | e to Naltrexone (OPRM1: No | ormal OPRM1 Functio | n) | INFORMATIV |
| | Vivitrol, Contrave | outcome with naltro respond to this dru | ol dependence: the patient has exone therapy. Naltrexone-treat g, and may have higher relapse sistently across studies. | ed patients not carrying | the OPRM1 11 | 8A>G G allele are less likely to |
| | Nebivolol | Normal Sensitivi | ty to Nebivolol (CYP2D6: In | termediate Metaboliz | zer) | ACTIONABL |
| Â | | Nebivolol can be p | | mmended dosage and a | dministration. (| Caution is recommended during |
| <u>^</u> | Bystolic | • | favorable response is achieved. | | | |
| <u>^</u> | Nortriptyline | up-titration until a | | D6: Intermediate Met | abolizer) | ACTIONABL |

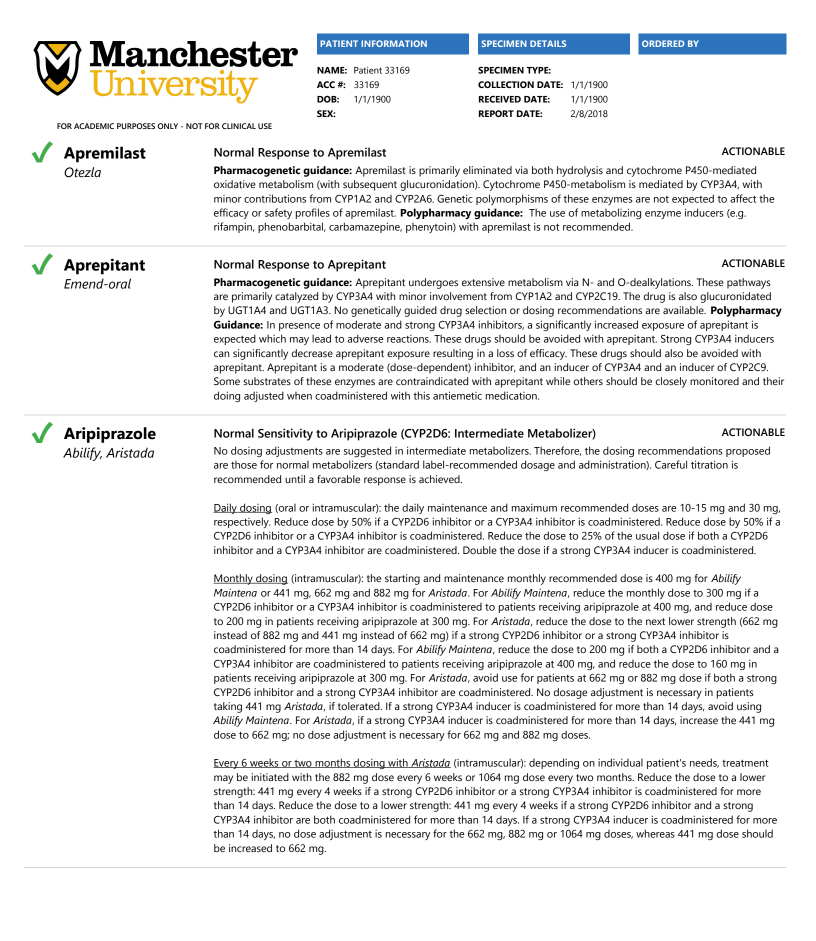


| | Mancl | hactor | PATIENT INFORMATION | SPECIMEN DETAILS | ORDERED BY | | |
|--------------------|---|---|---|---|--|--|--|
| | FOR ACADEMIC PURPOSES ONLY - NO | rsity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: Comparison | | /1900 /1900 /2018 | | |
| $\mathbf{\Lambda}$ | Olananina | | nse to Olanzapine (CYP1A2: | Linknown Dhonotyno) | INFORMATIV | | |
| <u>··</u> > | Olanzapine <i>Zyprexa</i> | There is little evider CYP1A2 metabolism doses, and careful r | nce regarding the impact of CYI n status cannot be predicted ac nonitoring is recommended du lverse events. Therefore, therap | 1A2 genetic variants on olan curately, smoking may increat ring dosing adjustment. Smol | zapine response. Although the patient's se the risk of non-response to standard king cessation may increase plasma drug panied by dose reduction may be neede | | |
| Ŷ | Omeprazole | Insufficient Resp | onse to Omeprazole (CYP2 | C19: Rapid Metabolizer) | ACTIONABI | | |
| | Prilosec | | er pylori eradication: increase d extra alert to insufficient respon | | | | |
| <u>^</u> | Oxycodone Percocet, Oxycontin | Decreased conversi metabolizers. Howe oxycodone. Adequa | e Altered Response to Oxycodone (CYP2D6: Intermediate Metabolizer) A ed conversion of oxycodone to the more active metabolite oxymorphone is expected in CYP2D6 intern zers. However, there is insufficient evidence whether these patients have decreased analgesia when ta ne. Adequate pain relief can be achieved by increasing the dose in response to pain symptoms. Other abolized by CYP2D6 may also be considered (i.e., morphine, oxymorphone, buprenorphine, fentanyl, n romorphone). | | | | |
| Ŷ | Pantoprazole | Insufficient Resp | onse to Pantoprazole (CYP2 | 2C19: Rapid Metabolizer) | ACTIONAB | | |
| | Protonix | Helicobacter pylori eradication: increase dose by 400% and be alert to insufficient response. Other: be extra alert to insufficient response and consider dose increase of 400%. | | | | | |
| Ŷ | Perphenazine Trilafon | Patients with a decr concentrations and | essible Sensitivity to Perphenazine (CYP2D6: Intermediate Metabolizer) tients with a decreased CYP2D6 function will eliminate perphenazine more slowly, which can resu ncentrations and possibly more adverse events (extrapyramidal symptoms). Consider close monit duction to avoid toxicity. | | | | |
| <u>^</u> | Propafenone | Moderate Sensiti | vity to Propafenone (CYP2 | D6: Intermediate Metabo | lizer) ACTIONABI | | |
| | Rythmol | | | | y and adjust the dose in response to h as sotalol, disopyramide, quinidine, or | | |
| | | inhibitors may signi | ficantly increase the plasma con other adverse events. Therefore | ncentration of propafenone a | y with CYP3A4 inhibitors and CYP2D6 nd thereby increase the risk of propafenone with both a CYP2D6 inhibito | | |
| <u>^</u> | Protriptyline | Possible Sensitivi | ity to Protriptyline (CYP2D6 | : Intermediate Metaboliz | er) INFORMATI\ | | |
| | Vivactil | | g protriptyline at 25% of recom etabolites and titrate according | | ese. Monitor plasma concentrations of is achieved. | | |
| <u>^</u> | Sertraline | Possible Reduced | l Response to Sertraline (C | /P2C19: Rapid Metabolize | er) INFORMATIN | | |
| | Zoloft | Sertraline can be prescribed at standard label-recommended dosage and administration. If patient does not recommended maintenance dosing, consider an alternative medication. | | | | | |

| | Manc Univer | hester sity | PATIENT INFORMATION NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 | SPECIMEN DETAILS SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: | | ORDERED BY |
|----------|----------------------------------|--|---|---|---|--|
| F | FOR ACADEMIC PURPOSES ONLY - N | OT FOR CLINICAL USE | SEX: | REPORT DATE: | 2/8/2018 | |
| <u>^</u> | Tacrolimus Prograf | The genotype resul tacrolimus more ra at increased risk for dose 1.5 to 2 times | onse to Tacrolimus (CYP3A t predicts that the patient expre- bidly, resulting in low tacrolimus acute transplant rejection while recommended starting dose wi dose should not exceed 0.3mg | sses the CYP3A5 protein s trough levels. Studies h e taking a standard dose th close monitoring is sta | . Therefore, th ave shown pa of tacrolimus | tients with this genotype may b . Therefore, increasing starting |
| <u>^</u> | Tetrabenazine Xenazine | For treating chore required. The first w weekly intervals by CYP2D6 is 100 mg | ty to Tetrabenazine (CYP2D a associated with Huntington veek's starting dose is 12.5 mg of 12.5 mg to a tolerated dose. The with a maximum single dose ose of tetrabenazine should be r | 's disease: Individualizat daily; second week, 25 m the maximum daily dose of 37.5 mg. If serious ac | ion of dose w g (12.5 mg tw in CYP2D6 i dverse events | ice daily); then slowly titrate at ntermediate metabolizers of |
| <u>î</u> | Timolol | Possible Sensitiv | ity to Timolol (CYP2D6: Inte | ermediate Metabolize | r) | ACTIONABI |
| _ | Timoptic | | ic beta-blockade (e.g., bradycar activity. Monitor patient for trea | | | treatment by patients with |
| <u>î</u> | Tizanidine | Unknown Respo | nse to Tizanidine (CYP1A2: I | Jnknown Phenotype) | | INFORMATI |
| | Zanaflex | CYP1A2 metabolisn higher doses. There excessive sedation. increase plasma dru | nce regarding the impact of CYF n status cannot be predicted acc is an association between high Therefore, careful monitoring is ug levels, leading to excessive hy eeded in patients who have qui | curately, smokers may be tizanidine plasma conce recommended during d ypotension and sedation | e at risk for no intrations and losing adjustn | n-response and may require the risk of hypotension and nent. Smoking cessation may |
| <u>î</u> | Tramadol | Possible Non-Re | sponder to Tramadol (CYP2 | D6: Intermediate Me | tabolizer) | ACTIONABI |
| _ | Ultram | needs to be individ than codeine, or a r | ed higher doses or may not exp ualized and careful weekly titrat non-opioid analgesic such as a I not sensitive to CYP2D6 functio | ion is recommended. If r NSAID or a COX-2 inhibit | no response, c or. Unless cor | consider alternative opioids othe ntraindicated, available |
| / | Alfentanil | Normal Respons | e to Alfentanil | | | INFORMATIV |
| - | Alfenta | Pharmacogenetic showed that CYP3A | guidance : alfentanil is primarily 5 genotype had no effect on th rmacy guidance: Alfentanil sho | e systemic or apparent o | ral clearances | s, or pharmacodynamics of |
| / | Alfuzosin | Normal Respons | e to Alfuzosin | | | INFORMATI |
| - | UroXatral | Polypharmacy gui Alfuzosin is contrai | r concentrations. Take caution | metabolized by CYP3A4 i inhibitors, as the risk f | into pharmaco or QTc prolo | ologically inactive metabolites. ngation induced by this drug |



| $\mathbf{\nabla}$ | 7) Manel | nactor | PATIENT INFORMATION | SPECIMEN DETAILS | 5 | ORDERED BY |
|-------------------|-------------------------------------|--|--|---|--|--|
| V | Manch Univer | J | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | FOR ACADEMIC PURPOSES ONLY - NO | T FOR CLINICAL USE | | | | |
| | Alprazolam Xanax | Pharmacogenetic polymorphisms of guidance: The cor prolonged sedatio exaggerated sedat | se to Alprazolam guidance: Alprazolam is primar these genes are not expected to ncomitant use of alprazolam with in. Impairment of motor skills are tive effects. If possible, alprazolar ole, itraconazole and ritonavir. D loss of efficacy. | affect the efficacy or said CYP3A4 inhibitors may also observed with som n should be avoided in p | fety profiles of result in increa e combination patients receiv | this drug. Polypharmacy ased alprazolam levels and ns. Monitor patients for ing strong inhibitors of CYP3A4 |
| | Amphetamine | Normal Exposu | re to Amphetamine (CYP2D6 | : Intermediate Metak | oolizer) | INFORMATIV |
| | Adderall, Evekeo | Amphetamine can | be prescribed at standard label- herapeutic needs and response c | recommended dosage a | | tion. Individualize the dosage |
| | Amphetamine | Good Response | to Amphetamine salts (CON | 1T: Intermediate CON | IT Activity) | INFORMATIV |
| | Adderall, Evekeo | | type result predicts a favorable r e lowest effective dose, and dose | | | Amphetamines should be |
| | Amphotericin B AmBisome, Abelcet | Pharmacogenetic of a given dose be genetically guided medications such induced renal toxic | se to Amphotericin B guidance: Amphotericin B is ex ing excreted in the biologically a drug selection or dosing recom as aminoglycosides, cyclosporine city, and should be used concom n patients requiring any combina | ctive form. Details of po mendations are available , and pentamidine may itantly only with great ca | ssible metabo e. Polypharma enhance the p aution. Intensiv | lic pathways are unknown. No acy guidance: Nephrotoxic otential for amphotericin B- |
| | Anidulafungin | Normal Respon | se to Anidulafungin | | | ACTIONABL |
| | Eraxis | activity and which has not been obse | : guidance: Anidulafungin under is subsequently converted to pe rved. Anidulafungin is not a subs drug selection or dosing recom | otidic degradants and el strate, inducer, or inhibite | iminated. Hep or of cytochro | atic metabolism of anidulafungi |
| | Apixaban | Normal Respon | se to Apixaban | | | INFORMATIV |
| | Eliquis | Pharmacogenetic primarily by CYP3/ efflux transport pri- genetic variations dosing adjustment administered with increase). Hence, f is coadministered ritonavir, and clarit inhibitors of CYP3/ | : guidance: Apixaban is not exter A4 and CYP3A5, with minor contri- oteins P-gp (ABCB1) and BCRP (A are unlikely to have a clinically si ts are recommended. Polypharm ketoconazole, a strong CYP3A/P or patients receiving 5 mg twice with drugs that are strong dual in thromycin). In patients already ta A4 and P-gp should be avoided. | ibutions from CYP1A2 a ABCG2). While these enzy gnificant impact on apix hacy guidance: Exposur -gp inhibitor. This transle daily, apixaban dose sho nhibitors of CYP3A4 and king 2.5 mg twice daily, | nd CYP2J2. Th ymes and tran aban exposure e to apixaban ates into an in ould be decrea P-gp (e.g., kei coadministrati ecommended | is drug is a substrate for the sporters are polymorphic, e, and no genotype-based increases by 100% when co- creased bleeding risk (70% sed to 2.5 mg twice daily when toconazole, itraconazole, ion of apixaban with strong dua when co-administered with |



| $\overline{\mathbf{N}}$ | 🕻 Manch | lester | PATIENT INFORMATION | SPECIMEN DETAILS | | ORDERED BY |
|-------------------------|----------------------------------|--|--|---|---|--|
| | FOR ACADEMIC PURPOSES ONLY - NOT | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: Image: Set the set | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| / | Asenapine | Normal Respons | e to Asenapine | | | INFORMATIV |
| | Saphris | Pharmacogenetic metabolism route of demethylation path CYP2D6. There are asenapine dispositi Asenapine should b guidance: Coadmi as asenapine plasm activity, has a limite coadministration w -term therapy with | Guidance: Asenapine is extension occurs via direct glucuronidation mway as well as the oxidative rea no studies documenting the effec- on and there are no available ge- pe prescribed based on the clinic nistration of asenapine with CYP na concentrations will increase re- ed effect on asenapine plasma co | catalyzed by UGT1A4. A ctions catalyzed by CYP1 ect of genetic polymorple netically guided drug se cal response and tolerab 1A2 inhibitors such as fl sulting in more side effe oncentrations. Asenapine and an inhibitor of CYP2 | Iso importan A2 with cont hisms of these election or do ility of the inc uvoxamine sh ects. Cigarette e is a weak inl 2D6) should b | t but less pronounced is the ributions from CYP3A4 and e metabolizing enzymes on sing recommendations. lividual patient. Polypharmacy rould be approached with cautio smoking, which induces CYP1A2 hibitor of CYP2D6 and its e approached with caution. Long |
| / | Atenolol | Normal Respons | e to Atenolol | | | INFORMATIV |
| | Tenormin | Pharmacogenetic approximately 90% Atenolol is a substr | guidance: The bioavailability of of the absorbed drug in its unc rate of several organic anion and tically-guided drug selection or o | hanged form. A negligib cation transporters incl | le amount of uding SLC22A | the drug is metabolized. 1, SLC22A2, SLC47A1, and |
| / | Atomoxetine | Normal Sensitivi | ty to Atomoxetine (CYP2D6 | : Intermediate Metab | olizer) | ACTIONABL |
| | Strattera | recommended unti | e prescribed at standard label-re il a favorable response is achieve : up to 70 kg, and 100 mg for pa | d. The maximum recom | mended daily | dose is 1.4 mg/kg for patients |
| / | Atorvastatin | Normal Myopatl | hy Risk (SLCO1B1: Normal Fu | inction) | | INFORMATIV |
| - | Lipitor | are present, atorvas -specific guidelines | a concentrations are not expecte statin can be prescribed at stanc . (Other myopathy predisposing nigh statin dose, comedications, | ard FDA-recommended factors include advance | starting dose | s and adjusted based on disease |
| / | Atorvastatin | Normal Respons | e to Atorvastatin (CYP3A4: | Normal Metabolizer) | | INFORMATIV |
| - | Lipitor | | It indicates that the patient does enzyme activity). The patient is equirements. | | | |
| / | Avanafil | Normal Respons | e to Avanafil | | | INFORMATIV |
| - | Stendra | Polypharmacy gui strong CYP3A4 in indinavir, itraconaz as erythromycin, ar | guidance: no genetically guided idance: Avanafil is extensively m hibitors such as ketoconazole, ir ole, nefazodone, nelfinavir, saqu nprenavir, aprepitant, diltiazem, -hour period. Inducers of CYP3A | etabolized by CYP3A4, t rraconazole, voriconazol inavir, and telithromycin fluconazole, fosamprena | herefore Ava e, ritonavir, at . If taking a m avir, or verapa | nafil should not be used with azanavir, clarithromycin, ioderate CYP3A4 inhibitor, such mil, the dose should be no more |
| | Azilsartan | Normal Sensitivi | ty to Azilsartan Medoxomil | (CYP2C9: Normal Me | tabolizer) | INFORMATIV |
| | Edarbi, Edarbyclor | | - | | | inal tract during absorption. |

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|-------------|--|---|---|---|--|---|
| | Univer | rsity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: Image: Compare the second secon | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | FOR ACADEMIC PURPOSES ONLY - NO | | | | | |
| | Betrixaban Bevyxxa | cytochrome P450 e CYP2C9, CYP2C19, d urinary excretion. B polymorphic, genet genotype-based do as amiodarone, azit | guidance: The predominant mon nzymes-based metabolism (less CYP2D6 and CYP3A4). The mair etrixaban is a substrate for the ic variations are unlikely to hav using adjustments are available. | s than 1% of the drug is r n elimination pathway of efflux transport protein P e a clinically significant ir Polypharmacy guidanc zole, clarithromycin result | netabolized b the drugs is b -gp (ABCB1) a npact on betri e: Concomita ts in increased | y CYP1A1, CYP1A2, CYP2B6, iliary excretion followed by ind while this transporter is xaban exposure, and no nt use with P-gp inhibitors such plasma levels of betrixaban and |
| | Bisoprolol | Normal Response | e to Bisoprolol | | | INFORMATIV |
| | Zebeta | Pharmacogenetic metabolized in the CYP3A4 with smalle | guidance: Bisoprolol is elimina liver and 50% being excreted vi er contribution from CYP2D6. Li libition are not affected by CYP | a the kidneys unchanged mited studies suggest the | l. Bisoprolol is at bisoprolol p | vith 50% of the total dose being predominantly metabolized by plasma concentrations and its guided drug selection or dosing |
| | Brexpiprazole | Normal Sensitivit | ty to Brexpiprazole (CYP2D | 6: Intermediate Metal | bolizer) | ACTIONABL |
| - | Rexulti | | ents are needed in CYP2D6 inte d dosage and administration. C | | | can be prescribed at standard favorable response is achieved. |
| | | daily maintenance of | doses and maximum recommer ing dose is 1 mg once daily. Th | nded dose are 1-2 mg an | d 3 mg, respe | e 0.5 mg or 1 mg once daily. The ctively. <u>Schizophrenia:</u> the um recommended dose are 2-4 |
| | | coadministered. Ad | vith comedications: reduce dos minister a quarter of the usual YP3A4 inhibitor are coadministe | dose if both a strong/mo | derate CYP2D | - |
| | Brivaracetam | Normal Sensitivit | ty to Brivaracetam (CYP2C1 | 9: Rapid Metabolizer) | | ACTIONABL |
| | Briviact | | narily metabolized by hydrolysi tam can be prescribed at the st | | | on, which is mediated by |
| | Buprenorphine | Normal Response | e to Buprenorphine | | | INFORMATIV |
| - | Butrans, Buprenex | Buprenorphine is pr The effects of gene concomitant use of increase or prolong | guidance: no genetically guide rimarily metabolized by CYP3A4 tic variants in these enzymes or buprenorphine with all CYP3A4 adverse drug effects. Monitor decrease buprenorphine levels. | 4 to norbuprenorphine ar n its response have not be 4 inhibitors may result in | nd by UGT enz een studied. P an increase in | rymes (mainly UGT1A1 and 2B7) Polypharmacy guidance: The the drug levels, which could |
| | Bupropion | Normal Response | e to Bupropion (CYP2B6: N | ormal Metabolizer) | | INFORMATIV |
| - | Wellbutrin, Zyban, Aplenzin, Contrave | Bupropion is metab therapeutic effects | olized to its active metabolite h of bupropion when used as a si ors are present, individuals who | nydroxybupropion by CYI moking cessation agent o | or as an antide | pressant. Unless other genetic |

| | 7 Manal | loctor | PATIEN | IT INFORMATION | SPECIMEN DETAILS | | ORDERED BY |
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| V | FOR ACADEMIC PURPOSES ONLY - NO | | NAME: ACC #: DOB: SEX: | Patient 33169 33169 1/1/1900 | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | | | | | | | |
| | Candesartan Atacand | gastrointestinal tract inactive metabolite. | uidance : during a Genetic v | Candesartan cilexetil bsorption. Candesarta ariability of the cytoch | 5 | atic metabolis expected to a | ACTIONABLI e metabolite in the m by O-deethylation to an affect the patient's response to |
| | Carbamazepine | Normal Response | to Carb | amazepine | | | INFORMATIVE |
| | Tegretol, Carbatrol, Epitol | be used to identify p syndrome, Stevens-Jo therapeutic window, metabolized by epox plasma concentration CYP3A5*1/*1 or *1/* dosage of carbamaze | atients a ohnson s is extens kide hydr ns are 30 3 genoty epine sho | t risk for severe cutane yndrome (SJS) and to ively metabolized by (olase (EPHX1) to an in % higher in individual pes. The clinical impac ould be decreased in p | ous adverse reactions s kic epidermal necrolysis CYP3A4/5 to its active ep active metabolite. Prelin s with the CYP3A5*3/*3 t of this change is poorl atients receiving CYP3A | uch as anticor (TEN). Carbam poxide metabo ninary studies genotype com y documented 4 inhibitors. En | indicate that carbamazepine npared to those with d. Polypharmacy guidance: The |
| | Cariprazine | Normal Response | to Cari | orazine | | | ACTIONABLE |
| | Vraylar | Genetic variants of C No geneticallly guide may affect cariprazin | YP2D6 d ed dosing e plasma used co | o not have clinically re g recommendations ar o concentrations. Carip | levant effect on pharma e available. Polypharm a razine dose may have to | cokinetics of c acy guidance b be reduced t | lesser extent, by CYP2D6. cariprazine and its metabolites. cCYP3A4 inhibitors or inducers to half if cariprazine and a strong inducer has not been evaluated |
| | Carvedilol | Normal Sensitivity | / to Car | vedilol (CYP2D6: In | termediate Metaboli | zer) | ACTIONABLE |
| - | Coreg | • | | at standard label-recon ng until a favorable res | mmended dosage and a ponse is achieved. | dministration. | Careful titration is |
| | Caspofungin | Normal Response | to Casp | oofungin | | | ACTIONABLE |
| | Cancidas | Pharmacogenetic g undergoes also spon dominant mechanism are available. Polyph rifampin, efavirenz, n | uidance itaneous n influen harmacy evirapine | Caspofungin is cleare chemical degradation cing plasma clearance guidance: Co-admini | Distribution, rather tha No genetically guided stration of caspofungin nazepine) may result in | n excretion or drug selection with metaboliz | lysis and N-acetylation. The drug biotransformation, is the or dosing recommendations zing enzyme inducers (e.g., hingful reductions in |
| \ | Celecoxib Celebrex | Normal Sensitivity | / to Cele | ecoxib (CYP2C9: No | rmal Metabolizer) | | ACTIONABLE |
| | CEIEDIEX | Celecoxib can be pre | scribed a | at standard label-recor | nmended dosage and a | dministration. | |
| | Chlorpromazine | Normal Response | to Chlo | rpromazine (CYP2I | 06: Intermediate Met | abolizer) | INFORMATIVE |
| - | ■ Thorazine | Chlorpromazine is m at standard label rec | etabolize | ed by CYP2D6, CYP3A4 | and flavin-containing r | nonooxygenas | ses. This drug can be prescribed |

| | Manch | noctor | PATIENT INFORMATION | SPECIMEN DETAILS | ORDERED BY |
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| V | Univer | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: 1/1/1 RECEIVED DATE: 1/1/1 REPORT DATE: 2/8/2 | 900 |
| | FOR ACADEMIC PURPOSES ONLY - NO | T FOR CLINICAL USE | | | |
| | Chlorpropamide <i>Diabenese</i> | The patient's genot | ty to Chlorpropamide (CYP) ype predicts a normal exposure age and administration (dose til | to chlorpropamide, and this c | INFORMATIVI drug can be prescribed at label- evels of glucose/glycosylated |
| | Clobazam | Normal Sensitivit | to Clobazam (CYP2C19: F | apid Metabolizer) | ACTIONABL |
| | Onfi | function. Rapid and metabolite of cloba prescribed. Therefo standard label-reco clinical efficacy and concentrations of cl Recommended dail | ultra-rapid metabolizers have zam. However, there is insuffici re, the dosing recommendation mmended dosage and adminis tolerability. Do not proceed wi obazam and its active metabol | a higher capacity to metabolize ent data to allow calculation o for normal metabolizers is pre- tration. Individualize dosing w th dose escalation more rapidl ite require 5 and 9 days, respe- starting dose 5 mg; day 7: 10 r | |
| \ | Clonazepam Klonopin | Polypharmacy gui | guidance: No genetically guide dance: clonazepam is extensive | ely metabolized by CYP3A4 to | INFORMATIVE commendations are available. an amino metabolite that is further prescribed with CYP3A4 inhibitors or |
| \checkmark | Clonidine | Normal Sensitivit | ty to Clonidine (CYP2D6: In | termediate Metabolizer) | INFORMATIV |
| | Карvау | remainder undergo CYP3A and CYP1A2 | ing hepatic metabolism. CYP2D | 6 plays a major role in clonidir t standard label recommendec | unchanged by the kidneys, with the ne oxidative metabolism, followed by I-dosage and administration. The dose patient. |
| | Colchicine | Normal Response | e to Colchicine | | INFORMATIV |
| - | Mitigare | Pharmacogenetic absorbed dose in el metabolic pathway this transporter is ir indicate a lack of ar with familial Medite recommendations. enzyme and the P-o toxicity. Inhibition of threatening or fatal | guidance: Colchicine in elimina iminated unchanged in urine, le for colchicine. Colchicine is a su nportant in its disposition. Colc effect of CYP3A4 or ABCB1 ge erranean fever (FMF). There are Polypharmacy guidance: Beca glycoprotein efflux transporter, f both CYP3A4 and P-gp by du | ess than 20% is metabolized by ubstrate of P-glycoprotein (enc hicine has a narrow therapeuti netic polymorphisms on clinica no available genetically-guide ause colchicine is a substrate for inhibition of either of these pa al inhibitors such as clarithrom icant increases in systemic col | nd metabolism. While 50% of the y CYP3A4. Glucuronidation is also a coded by ABCB1 gene) and its efflux by ic index. Preliminary and limited studies al response to colchicine in individuals d drug selection or dosing or both the CYP3A4 metabolizing thways may lead to colchicine-related hycin has been reported to produce life- chicine levels. Therefore, concomitant |
| | Cyclobenzaprine | Normal Response | e to Cyclobenzaprine | | INFORMATIVE |
| - | Flexeril, Amrix | Pharmacogenetic Cyclobenzaprine is CYP1A2, and to a le | guidance: No genetically guide excreted primarily as a glucuror | nide via the kidneys, and as an minor involvement of CYP2D | commendations are available. N-demethylated metabolite by CYP3A4, 6 in the metabolism of cyclobenzaprine, |



| Manch | octor | PATIENT INFORMATION | SPECIMEN DETAILS | ; | ORDERED BY |
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| FOR ACADEMIC PURPOSES ONLY - NOT | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: Image: Compare the second secon | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| Dabigatran Etexilate | Normal Response | e to Dabigatran | | | INFORMATIV |
| Pradaxa | dabigatran etexilate also conjugated to fi CYP450 enzymes. Da polymorphism of the Polypharmacy guid moderate renal impa ketoconazole can be Consider reducing th with other P-gp inhi <u>2-Treatment of DVT</u> | orm pharmacologically active a abigatran etexilate is a substrate e ABCB1 gene (2677G>T/A and lance: <u>1-Reduction in Risk of St</u> airment (CrCl 30-50 mL/min), co e expected to produce dabigatr he dose of dabigatran to 75 mg bitors. In patients with CrCl<30 | dabigatran by esterases. cyl glucuronides. Dabiga e of the efflux transporte 3435 C>T) do not appe roke and Systemic Embo oncomitant use of the P- an exposure similar to th twice daily. Dose adjus mL/min, avoid use of co | A small porti- atran is not a s er P-gp (ABCB ear to affect da <i>lism in Non-va</i> -gp inhibitor of nat observed i tment is not no poncomitant P- | on (20%) of dabigatran dose is substrate, inhibitor, or inducer of (1). Common genetic abigatran exposure. <u>alvular AF</u> : In patients with dronedarone or systemic in severe renal impairment. necessary when coadministered |
| Darifenacin Enablex | | e to Darifenacin (CYP2D6: Ir | | | ACTIONABLI |
| Desvenlafaxine Pristiq | | y to Desvenlafaxine (CYP2E | | | ACTIONABL |
| Deutetrabenazine Austedo | For treating chorea required. The first we | y to Deutetrabenazine (CYF a associated with Huntington eek's starting dose is 6 mg onco and up to a maximum recomm | s disease: Individualizate e daily followed by a slo | tion of dose w w titration at v | weekly intervals by 6 mg per day |
| Dextroamphetami | Normal Exposure | to Dextroamphetamine (C | YP2D6: Intermediate | Metabolize | r) INFORMATIV |
| ne Dexedrine | | e can be prescribed at standard the therapeutic needs and res | | sage and adn | ninistration. Individualize the |
| • | Good Response to | o Dextroamphetamine (CO | MT: Intermediate CO | MT Activity |) INFORMATIV |
| ne Dexedrine | | rpe result predicts a favorable re lowest effective dose, and dosa | | | Dextroamphetamine should be |
| Dextromethorpha n / Quinidine | Normal Sensitivit Metabolizer) | y to Dextromethorphan-Qเ | iinidine (CYP2D6: Int | ermediate | ACTIONABL |
| Nuedexta | the dextromethorph | dobulbar Affect : quinidine is a an-quinidine combination to in quinidine can be prescribed acc | crease the systemic bio | availability of | |

| N . N | /> Manol | nester | PATIENT INFORMATION | SPECIMEN DETAILS | | ORDERED BY |
|--------------------------------|--|--|---|--|--|--|
| | | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | | | | | | |
| | Diclofenac Voltaren | Individuals with a n | ty to Diclofenac (CYP2C9: N ormal CYP2C9 activity (i.e norm d-dosage and administration. | | prescribed dicl | INFORMATIVI ofenac according to standard |
| | Dihydrocodeine | Normal Respons | e to Dihydrocodeine (CYP2 | D6: Intermediate Met | abolizer) | INFORMATIV |
| | Synalgos-DC | intermediate metal | on of dihydrocodeine to the m polizers. However, there is insuf eine. Adequate pain relief can b | icient evidence whether | these patients | have decreased analgesia when |
| | Dolasetron Anzemet | Normal Respons | e to Dolasetron (CYP2D6: I | ntermediate Metaboli | zer) | INFORMATIV |
| | | Dolasetron can be | prescribed at standard label-rec | ommended dosage and | administratior | ۱. |
| | Dolutegravir | Normal Respons | e to Dolutegravir | | | ACTIONABL |
| | Tivicay, Triumeq | have increased plas required for dolute | TYP3A. Although UGT1A1 poor sma levels of dolutegravir, these gravir due to genetic variations rugs that are strong enzyme inc | changes are not clinicall in UGT1A1. Polypharma | y significant. N I cy guidance : | No dosing adjustments are Coadministration of |
| | | | | | | |
| | Donepezil | Normal Respons | e to Donepezil (CYP2D6: In | termediate Metaboliz | er) | INFORMATIV |
| \ | Donepezil Aricept | Donepezil can be p | e to Donepezil (CYP2D6: In rescribed at standard label-reco l a favorable response is achiev | mmended dosage and a | | |
| ✓ ✓ | - | Donepezil can be p recommended unti | rescribed at standard label-reco l a favorable response is achiev e to Doxazosin | ommended dosage and a ed. | dministration. | Careful titration is INFORMATIV |
| | Aricept | Donepezil can be p recommended unti Normal Respons Pharmacogenetic Polypharmacy gui | rescribed at standard label-reco l a favorable response is achiev | ommended dosage and a ed. d drug selection or dosir | dministration. ng recommend | Careful titration is INFORMATIV lations are available. |
| く く く | Aricept Doxazosin Cardura Dronabinol | Donepezil can be p recommended unti Normal Respons Pharmacogenetic Polypharmacy gui known to influence | rescribed at standard label-reco l a favorable response is achiev e to Doxazosin guidance: no genetically guide dance: doxazosin is metabolize | ommended dosage and a ed. d drug selection or dosir d by multiple enzymes. T | dministration. ng recommend | Careful titration is INFORMATIV dations are available. d data on the effects of drugs |
| ✓ ✓ ✓ | Aricept Doxazosin Cardura | Donepezil can be p recommended unti Normal Respons Pharmacogenetic Polypharmacy gui known to influence Normal Sensitivi The patient's genot | rescribed at standard label-reco l a favorable response is achiev e to Doxazosin guidance: no genetically guide dance: doxazosin is metabolize the metabolism of doxazosin. | ommended dosage and a ed. d drug selection or dosir d by multiple enzymes. T Normal Metabolizer) | dministration. ng recommeno 'here is limiteo | Careful titration is INFORMATIV lations are available. I data on the effects of drugs INFORMATIV |
| ✓ ✓ ✓ ✓ | Aricept Doxazosin Cardura Dronabinol Marinol Duloxetine | Donepezil can be p recommended unti Normal Respons Pharmacogenetic Polypharmacy gui known to influence Normal Sensitivi The patient's genot recommended dos | rescribed at standard label-reco l a favorable response is achiev e to Doxazosin guidance: no genetically guide dance: doxazosin is metabolize the metabolism of doxazosin. ty to Dronabinol (CYP2C9: type predicts a normal CYP2C9 | ommended dosage and a ed. d drug selection or dosir d by multiple enzymes. T Normal Metabolizer) netabolic activity. Dronal | dministration. ng recommeno 'here is limiteo binol can be p | Careful titration is INFORMATIVI dations are available. d data on the effects of drugs INFORMATIVI rescribed at standard label- |
| ✓ ✓ ✓ | Aricept Doxazosin Cardura Dronabinol Marinol | Donepezil can be p recommended unti Normal Respons Pharmacogenetic Polypharmacy gui known to influence Normal Sensitivi The patient's genot recommended dos Normal Sensitivi | rescribed at standard label-reco l a favorable response is achiev e to Doxazosin guidance: no genetically guide dance: doxazosin is metabolize the metabolism of doxazosin. ty to Dronabinol (CYP2C9: sype predicts a normal CYP2C9 age and administration. | ommended dosage and a ed. d drug selection or dosir d by multiple enzymes. T Normal Metabolizer) netabolic activity. Dronal | dministration. ng recommenc There is limited binol can be p izer) | INFORMATIVE dations are available. d data on the effects of drugs INFORMATIVE rescribed at standard label- INFORMATIVE |
| ✓ ✓ ✓ ✓ | Aricept Doxazosin Cardura Dronabinol Marinol Duloxetine | Donepezil can be p recommended unti Normal Respons Pharmacogenetic Polypharmacy gui known to influence Normal Sensitivi The patient's genot recommended dos Normal Sensitivi Duloxetine can be Normal Respons | rescribed at standard label-reco l a favorable response is achiev e to Doxazosin guidance: no genetically guide dance: doxazosin is metabolize the metabolism of doxazosin. ty to Dronabinol (CYP2C9: type predicts a normal CYP2C9 age and administration. ty to Duloxetine (CYP2D6: I prescribed at standard label-reco | ommended dosage and a ed. d drug selection or dosir d by multiple enzymes. T Normal Metabolizer) netabolic activity. Dronal ntermediate Metabol ommended dosage and | dministration. ng recommend There is limited binol can be p izer) administratior | Careful titration is INFORMATIVI dations are available. d data on the effects of drugs INFORMATIVI rescribed at standard label- INFORMATIVI n. INFORMATIVI |

| | A Manch | noctor | PATIENT INFORMATION | SPECIMEN DETAILS | | ORDERED BY |
|---|---------------------------------|--|---|---|---|--|
| | FOR ACADEMIC PURPOSES ONLY - NO | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: Image: Comparison of the second | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | Edoxaban | Normal Response | e to Edoxaban | | | INFORMATIVE |
| · | Savaysa | Pharmacogenetic of via hydrolysis (medi efflux transporter P- SLCO1B1. Prelimina does not affect edo: | | jugation, and oxidation rmed by carboxylesteras single nucleotide polyr harmacy guidance: Avo | by CYP3A4. I se 1) is a subs norphism (rse pid the conco | strate of the uptake transporter 4149056) of the SLCO1B1 gene |
| | Eprosartan | Normal Sensitivit | y to Eprosartan | | | ACTIONABLE |
| | Teveten | Pharmacogenetic g Eprosartan is not me | juidance: Eprosartan is elimina | 450 enzymes. Genetic va | ariability of th | marily as unchanged compound. ne cytochrome P450 genes is not istments are available. |
| | Eslicarbazepine | Normal Response | e to Eslicarbazepine | | | INFORMATIVE |
| | Aptiom | syndrome, Stevens converted by a redu excretion unchange are available. Polyp | batients at risk for severe cutane Johnson syndrome (SJS) and to: Ictase to its active metabolite, en d and as a glucuronide conjuga harmacy guidance: In the pre- sed, and higher doses of the dru | kic epidermal necrolysis slicarbazepine. Eslicarba te. No genetically guide sence of enzyme-inducii | (TEN). Eslicar zepine is elim d drug select | bazepine acetate (prodrug) is ninated primarily by renal ion or dosing recommendations |
| | Ethosuximide | Normal Response | e to Ethosuximide | | | INFORMATIVE |
| - | Zarontin | Polypharmacy guid with caution when p | guidance: No genetically guide dance: ethosuximide is extensiv prescribed with CYP3A4 inhibito ed when the drug is coadministe | ely metabolized by CYP3 rs. Inducers of CYP3A4 i | BA4, and ther ncrease ethos | efore this drug should be used |
| | Ezogabine | Normal Response | e to Ezogabine | | | INFORMATIVE |
| - | Potiga | metabolite, no dose metabolized primari oxidative metabolisi are not expected to | adjustment is necessary in thes ily via glucuronidation (by UGT1 n of ezogabine by cytochrome affect its efficacy or toxicity pro clearance by 30%, and dose inc | e individuals. Polyphar A4 and UGT1A1) and ac P450 enzymes, and gene files. Enzyme-inducing c | macy guidar etylation (by etic variations lrugs such as | NAT2). There is no evidence of s in these metabolizing enzymes carbamazepine and phenytoin |
| | Febuxostat | Normal Response | e to Febuxostat | | | INFORMATIVE |
| - | Uloric | Pharmacogenetic of metabolized both bo cytochrome P450 er metabolized to an a are no available gen administration of pr | guidance: Febuxostat is elimina y glucuronidation and oxidative nzymes (CYPs): CYP1A2, CYP2C8 cyl glucuronide, primarily by UC | pathways. The oxidative and CYP2C9 as well as GT1A1 with contribution or dosing recommendati hibitor, with substrate dr | e metabolism other non-CY s from UGT1/ ons. Polyph a ugs such as t | of this drug involves several /P enzymes. Febuxostat is also A3, UGT1A9 and UGT2B7. There armacy guidance: Concomitant theophylline, azathioprine or |



| | /) Mane | hester | PATIENT INFORMATION | SPECIMEN DETAILS | | ORDERED BY |
|-------------|--|---|--|--|--|--|
| V | Univer | rsity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | FOR ACADEMIC PURPOSES ONLY - N | NOT FOR CLINICAL USE | | | | |
| | Felbamate | Normal Respons | | | | |
| | Felbatol | Polypharmacy gui 50% is present as n minor for drug elim enzyme-inducing a | guidance: No genetically guide dance: About 40-50% of absord netabolites and conjugates. Felb nination when the drug is given ntiepileptic drugs, which results slowly, and dose adjustment mu | ped felbamate dose appe namate is a substrate of C as a monotherapy. This p in a 30-50% decrease in | ears unchange YP3A4 and CY pathway is enh felbamate pla | d in urine, and an additional (P2E1, but these pathways are lanced by concomitant use of Isma concentrations. Felbamate |
| | Fentanyl | Good Response | to Fentanyl (OPRM1: Norma | al OPRM1 Function) | | INFORMATIV |
| | Actiq | experience good ar | | ses. Because fentanyl has | a narrow the | r pain: the patient is expected to rapeutic window, it is advised to nal side effects. |
| √ | Fesoterodine Toviaz | | ty to Fesoterodine (CYP2D6 | | | ACTIONABL |
| | | Fesoterodine can b | e prescribed at standard label-r | ecommended dosage an | d administrati | on. |
| | Finasteride | Normal Respons | e to Finasteride | | | INFORMATIV |
| V | i masteriae | i torinar nespons | e to i masteriae | | | |
| V | Proscar | Pharmacogenetic Polypharmacy gui moderate CYP3A4 | guidance: no genetically guide dance: Finasteride is extensively inhibitors on finasteride have no prescribing this drug to patients | y metabolized in humans ot been studied. Because | by CYP3A4. T of the potenti | he effects of potent or |
| v V | | Pharmacogenetic Polypharmacy gui moderate CYP3A4 use caution when p | guidance: no genetically guide dance: Finasteride is extensively inhibitors on finasteride have no | y metabolized in humans ot been studied. Because taking CYP3A4 enzyme in | by CYP3A4. T of the potenti | he effects of potent or |
| ✓ ✓ | Proscar | Pharmacogenetic Polypharmacy gui moderate CYP3A4 i use caution when p Normal Exposure For treating prem Flibanserin is prima | guidance: no genetically guide dance: Finasteride is extensively inhibitors on finasteride have no prescribing this drug to patients to Flibanserin (CYP2C19: R enopausal women with acquir rily metabolized by CYP3A4 and to have a normal clearance and | y metabolized in humans ot been studied. Because taking CYP3A4 enzyme in apid Metabolizer) red, generalized hypoac d, to a lesser extent, by C | by CYP3A4. T of the potenti nhibitors. tive sexual do YP2C19. The g | The effects of potent or al for drug-drug interactions, ACTIONABL |
| ✓ ✓ | Proscar Flibanserin | Pharmacogenetic Polypharmacy gui moderate CYP3A4 i use caution when p Normal Exposure For treating prem Flibanserin is prima patient is expected | guidance: no genetically guide dance: Finasteride is extensively inhibitors on finasteride have no prescribing this drug to patients to Flibanserin (CYP2C19: R enopausal women with acquir rily metabolized by CYP3A4 and to have a normal clearance and ecautions. | y metabolized in humans ot been studied. Because taking CYP3A4 enzyme in apid Metabolizer) red, generalized hypoac d, to a lesser extent, by C | by CYP3A4. T of the potenti nhibitors. tive sexual do YP2C19. The g | The effects of potent or al for drug-drug interactions, ACTIONABL esire disorder (HSDD): enotype results predict that the |
| ✓ ✓ | Proscar Flibanserin Addyi | Pharmacogenetic Polypharmacy gui moderate CYP3A4 i use caution when p Normal Exposure For treating preme Flibanserin is prima patient is expected follow standard pre Normal Respons Pharmacogenetic approximately 80% pharmacokinetics c or dosing recomme CYP2C9 and CYP2C therapeutic window | guidance: no genetically guide dance: Finasteride is extensively inhibitors on finasteride have no prescribing this drug to patients e to Flibanserin (CYP2C19: R enopausal women with acquir rily metabolized by CYP3A4 and to have a normal clearance and ecautions. e to Fluconazole guidance: Fluconazole not exten- of the administered dose appe | y metabolized in humans of been studied. Because taking CYP3A4 enzyme in capid Metabolizer) red, generalized hypoac d, to a lesser extent, by C ¹ a typical exposure to flik ensively metabolized and aring in the urine as unch ed by reduction in renal f armacy guidance: Flucor d patients who are conco C19 or CYP3A4 should be | by CYP3A4. T of the potenti nhibitors. tive sexual de YP2C19. The g panserin. Use I is eliminated function. No g nazole is a mo pmitantly treat e monitored. T | The effects of potent or al for drug-drug interactions, ACTIONABL esire disorder (HSDD): enotype results predict that the abel-recommended dosage and ACTIONABL primarily by renal excretion, wit and 11% as metabolites. The enetically guided drug selectior derate inhibitor of CYP3A4, red with drugs with a narrow |
| ✓ ✓ ✓ | Proscar Flibanserin Addyi Fluconazole | Pharmacogenetic Polypharmacy gui moderate CYP3A4 is use caution when p Normal Exposure For treating preme Filibanserin is prima patient is expected follow standard pres Normal Respons Pharmacogenetic approximately 80% pharmacokinetics co or dosing recomme CYP2C9 and CYP2C therapeutic window fluconazole persister | guidance: no genetically guide dance: Finasteride is extensively inhibitors on finasteride have no prescribing this drug to patients e to Flibanserin (CYP2C19: R enopausal women with acquir rily metabolized by CYP3A4 and to have a normal clearance and ecautions. e to Fluconazole guidance: Fluconazole not extension of the administered dose append of fluconazole is markedly affect endations are available. Polypha (19 enzymes. Fluconazole treate of metabolized by CYP2C9, CYP2 | y metabolized in humans ot been studied. Because taking CYP3A4 enzyme in apid Metabolizer) red, generalized hypoac d, to a lesser extent, by CV a typical exposure to flik ensively metabolized and aring in the urine as unch ed by reduction in renal f armacy guidance: Flucor d patients who are conco C19 or CYP3A4 should be of the drug due to its lor | by CYP3A4. T of the potenti nhibitors. tive sexual do YP2C19. The g banserin. Use I hanged drug a function. No g nazole is a mo pmitantly treat e monitored. T ng half-life. | The effects of potent or al for drug-drug interactions, ACTIONABL esire disorder (HSDD): enotype results predict that the abel-recommended dosage and ACTIONABL primarily by renal excretion, wit and 11% as metabolites. The enetically guided drug selectior derate inhibitor of CYP3A4, red with drugs with a narrow |
| ✓ ✓ ✓ | Proscar Flibanserin <i>Addyi</i> Fluconazole <i>Diflucan</i> | Pharmacogenetic Polypharmacy gui moderate CYP3A4 i use caution when p Normal Exposure For treating preme Flibanserin is prima patient is expected follow standard pre Normal Respons Pharmacogenetic approximately 80% pharmacokinetics c or dosing recomme CYP2C9 and CYP2C therapeutic window fluconazole persists Normal Sensitivi Fluoxetine is metab | guidance: no genetically guide dance: Finasteride is extensively inhibitors on finasteride have no prescribing this drug to patients the to Flibanserin (CYP2C19: R enopausal women with acquir rily metabolized by CYP3A4 and to have a normal clearance and ecautions. to fucconazole guidance: Fluconazole not exten- of the administered dose appe- of fluconazole is markedly affect endations are available. Polypha (19 enzymes. Fluconazole treate of metabolized by CYP2C9, CYP2 s 4-5 days after discontinuation | y metabolized in humans of been studied. Because taking CYP3A4 enzyme in apid Metabolizer) red, generalized hypoac d, to a lesser extent, by C a typical exposure to flit ensively metabolized and aring in the urine as unch ed by reduction in renal f armacy guidance: Flucor d patients who are conce C19 or CYP3A4 should be of the drug due to its lor ntermediate Metaboli orfluoxetine and to othe | by CYP3A4. T of the potenti nhibitors. tive sexual de YP2C19. The g banserin. Use I is eliminated hanged drug a function. No g nazole is a mo omitantly treat e monitored. T ng half-life. zer) r metabolites I | The effects of potent or al for drug-drug interactions, ACTIONABL esire disorder (HSDD): enotype results predict that the abel-recommended dosage and ACTIONABL primarily by renal excretion, wit and 11% as metabolites. The renetically guided drug selection derate inhibitor of CYP3A4, ed with drugs with a narrow The enzyme inhibiting effect of INFORMATIV by multiple enzymes including |
| ✓ ✓ ✓ | Proscar Flibanserin Addyi Fluconazole Diflucan Fluoxetine | Pharmacogenetic Polypharmacy gui moderate CYP3A4 is use caution when p Normal Exposure For treating preme Filibanserin is prima patient is expected follow standard pression Normal Respons Pharmacogenetic approximately 80% pharmacokinetics of or dosing recomme CYP2C9 and CYP2C therapeutic window fluconazole persists Normal Sensitivi Fluoxetine is metab CYP2D6, CYP2C19, administration. | guidance: no genetically guide dance: Finasteride is extensively inhibitors on finasteride have no orescribing this drug to patients e to Flibanserin (CYP2C19: R enopausal women with acquir rily metabolized by CYP3A4 and to have a normal clearance and ecautions. e to Fluconazole guidance: Fluconazole not extended of the administered dose append of the administered dose append of fluconazole is markedly affect endations are available. Polypha 19 enzymes. Fluconazole treate w metabolized by CYP2C9, CYP2 5 4-5 days after discontinuation ty to Fluoxetine (CYP2D6: In polized to its active metabolite n | y metabolized in humans ot been studied. Because taking CYP3A4 enzyme in apid Metabolizer) red, generalized hypoac d, to a lesser extent, by CV a typical exposure to flik ensively metabolized and aring in the urine as unch ed by reduction in renal farmacy guidance: Flucor d patients who are conco C19 or CYP3A4 should be of the drug due to its lor ntermediate Metaboli orfluoxetine and to othe e can be prescribed at sta | by CYP3A4. T of the potenti nhibitors. tive sexual do YP2C19. The g banserin. Use I hanged drug a function. No g nazole is a mo omitantly treat e monitored. T ng half-life. zer) r metabolites I andard label-r | The effects of potent or al for drug-drug interactions, ACTIONABL esire disorder (HSDD): enotype results predict that the abel-recommended dosage and ACTIONABL primarily by renal excretion, wit and 11% as metabolites. The renetically guided drug selection derate inhibitor of CYP3A4, ed with drugs with a narrow The enzyme inhibiting effect of INFORMATIV by multiple enzymes including |

| | A IVIANCI | hester | PATIENT INFORMATION | SPECIMEN DETAILS | | ORDERED BY |
|----------------------|-----------------------------------|---|--|---|---|---|
| V | Mancl Univer | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | FOR ACADEMIC PURPOSES ONLY - NO | DT FOR CLINICAL USE | | | | |
| | Fluvastatin | Normal Myopath | y Risk (SLCO1B1: Normal F | unction) | | INFORMATIVE |
| | Lescol | present, fluvastatin specific guidelines. (| concentrations are not expecte can be prescribed at standard Other myopathy predisposing gh statin dose, comedications | FDA-recommended starti factors include advanced | ing doses and | - |
| | Fluvastatin | Normal Sensitivit | y to Fluvastatin (CYP2C9: I | Normal Metabolizer) | | ACTIONABLE |
| | Lescol | present, fluvastatin specific guidelines. (| can be prescribed at standard | FDA-recommended starti isposing factors include a | ing doses and advanced age | (265), diabetes, hypothyroidism, |
| | Fluvoxamine | Normal Sensitivit | y to Fluvoxamine (CYP2D6 | : Intermediate Metab | olizer) | ACTIONABLE |
| | Luvox | | prescribed at standard label re a favorable response is achiev | - | d administratio | on. Careful titration is |
| | Fondaparinux | Normal Response | e to Fondaparinux | | | INFORMATIVE |
| | | | and a set a construction of the set and a | • - !! | | |
| | | profiles. no genetica concomitant use of may enhance the ris | fondaparinux with aspirin or N | osing recommendations a SAIDS may enhance the tion of therapy with fonce | are available. I risk of hemorr | o affect its efficacy or toxicity Polypharmacy guidance: The hage. Discontinue agents that ess essential. If co-administration |
| | Fosaprepitant | profiles. no genetica concomitant use of may enhance the ris is necessary, monito Normal Response | Ily guided drug selection or de fondaparinux with aspirin or N k of hemorrhage prior to initia pratients closely for hemorrha to Fosaprepitant | osing recommendations a SAIDS may enhance the tion of therapy with fonc age. | are available. I risk of hemorr laparinux unle | Polypharmacy guidance: The hage. Discontinue agents that ess essential. If co-administration ACTIONABLE |
| √ | Fosaprepitant Emend-i.v | profiles. no genetica concomitant use of may enhance the ris is necessary, monito Normal Response Pharmacogenetic g intravenous adminis metabolism via N- a CYP1A2 and CYP2C dosing recommenda inhibitors, a significa should be avoided v a loss of efficacy. Th inhibitor, and an inc | ally guided drug selection or de fondaparinux with aspirin or N k of hemorrhage prior to initia or patients closely for hemorrha e to Fosaprepitant guidance: Fosaprepitant is a p stration. Its antiemetic effects a and O-dealkylations. These pat 19. The drug is also glucuronid ations are available. Polyphari antly increased exposure of ap with fosaprepitant. Strong CYP ese drugs should also be avoid lucer of CYP3A4 and an induce while others should be closely r | osing recommendations a SAIDS may enhance the tion of therapy with fond age. rodrug of aprepitant which re attributable to aprepit hways are primarily cataly ated by UGT1A4 and UG macy Guidance: In prese repitant is expected which BA4 inducers can significat led with fosaprepitant. A r of CYP2C9. Some subst | are available. I risk of hemorr laparinux unle ch is rapidly cc ant. Aprepitan yzed by CYP3A T1A3. No gene cnce of modera h may lead to antly decrease prepitant is a r rates of these | Polypharmacy guidance: The hage. Discontinue agents that iss essential. If co-administration ACTIONABLE onverted to aprepitant following at undergoes extensive A4 with minor involvement from etically guided drug selection or ate and strong CYP3A4 adverse reactions. These drugs aprepitant exposure resulting in moderate (dose-dependent) enzymes are contraindicated |
| | | profiles. no genetica concomitant use of may enhance the ris is necessary, monito Normal Response Pharmacogenetic Q intravenous adminis metabolism via N- a CYP1A2 and CYP2C dosing recommenda inhibitors, a significa should be avoided v a loss of efficacy. Th inhibitor, and an inc with fosaprepitant v antiemetic medicati | ally guided drug selection or de fondaparinux with aspirin or N k of hemorrhage prior to initia or patients closely for hemorrha e to Fosaprepitant guidance: Fosaprepitant is a p stration. Its antiemetic effects a and O-dealkylations. These pat 19. The drug is also glucuronid ations are available. Polyphari antly increased exposure of ap with fosaprepitant. Strong CYP ese drugs should also be avoid lucer of CYP3A4 and an induce while others should be closely r | osing recommendations a SAIDS may enhance the tion of therapy with fonc- age. Todrug of aprepitant which re attributable to aprepit hways are primarily cataly ated by UGT1A4 and UG nacy Guidance: In prese repitant is expected which 8A4 inducers can significat ded with fosaprepitant. A r of CYP2C9. Some subst monitored and their doing | are available. I risk of hemorr laparinux unle ch is rapidly co ant. Aprepitan yzed by CYP3A T1A3. No gene ence of modera h may lead to antly decrease prepitant is a i rates of these g adjusted wh | Polypharmacy guidance: The hage. Discontinue agents that iss essential. If co-administration ACTIONABLE onverted to aprepitant following at undergoes extensive A4 with minor involvement from etically guided drug selection or ate and strong CYP3A4 adverse reactions. These drugs aprepitant exposure resulting in moderate (dose-dependent) enzymes are contraindicated |
| ✓ ✓ | Emend-i.v | profiles. no genetica concomitant use of may enhance the ris is necessary, monito Normal Response Pharmacogenetic g intravenous adminis metabolism via N- a CYP1A2 and CYP2C dosing recommenda inhibitors, a significa should be avoided v a loss of efficacy. Th inhibitor, and an ind with fosaprepitant v antiemetic medicati Normal Sensitivit The genotype result | Illy guided drug selection or de fondaparinux with aspirin or N k of hemorrhage prior to initia or patients closely for hemorrha e to Fosaprepitant guidance: Fosaprepitant is a p stration. Its antiemetic effects a and O-dealkylations. These pat 19. The drug is also glucuronid ations are available. Polyphari antly increased exposure of ap with fosaprepitant. Strong CYP ese drugs should also be avoid lucer of CYP3A4 and an induce while others should be closely r on. y to Fosphenytoin (CYP2C s indicate that the patient is a g dose and a standard mainte | osing recommendations a SAIDS may enhance the tion of therapy with fond age. rodrug of aprepitant which re attributable to aprepit hways are primarily cataly ated by UGT1A4 and UG nacy Guidance: In prese repitant is expected which BA4 inducers can significat ded with fosaprepitant. A r of CYP2C9. Some subst nonitored and their doing 9: Normal Metabolize CYP2C9 substrate norma | are available. I risk of hemorr laparinux unle ch is rapidly cc ant. Aprepitan yzed by CYP3A T1A3. No gene ince of modera h may lead to antly decrease prepitant is a r rates of these g adjusted wh r) I metabolizer. | Polypharmacy guidance: The hage. Discontinue agents that iss essential. If co-administration ACTIONABLE onverted to aprepitant following at undergoes extensive A4 with minor involvement from etically guided drug selection or ate and strong CYP3A4 adverse reactions. These drugs aprepitant exposure resulting in moderate (dose-dependent) enzymes are contraindicated en coadministered with this ACTIONABLE Fosphenytoin can be prescribed |
| | Emend-i.v Fosphenytoin | profiles. no genetica concomitant use of may enhance the ris is necessary, monito Normal Response Pharmacogenetic g intravenous adminis metabolism via N- a CYP1A2 and CYP2C dosing recommenda inhibitors, a significa should be avoided v a loss of efficacy. Th inhibitor, and an inc with fosaprepitant v antiemetic medicati Normal Sensitivit The genotype result at a standard loadin | Ally guided drug selection or de fondaparinux with aspirin or N ik of hemorrhage prior to initia or patients closely for hemorrha e to Fosaprepitant guidance: Fosaprepitant is a p tration. Its antiemetic effects a and O-dealkylations. These pat 19. The drug is also glucuronid ations are available. Polyphar antly increased exposure of ap with fosaprepitant. Strong CYP ese drugs should also be avoid lucer of CYP3A4 and an induce while others should be closely r on. y to Fosphenytoin (CYP2C s indicate that the patient is a g dose and a standard mainte y. | osing recommendations a SAIDS may enhance the tion of therapy with fond age. rodrug of aprepitant which re attributable to aprepit hways are primarily cataly ated by UGT1A4 and UG nacy Guidance: In prese repitant is expected which BA4 inducers can significat ded with fosaprepitant. A r of CYP2C9. Some subst nonitored and their doing 9: Normal Metabolize CYP2C9 substrate norma | are available. I risk of hemorr laparinux unle ch is rapidly cc ant. Aprepitan yzed by CYP3A T1A3. No gene ince of modera h may lead to antly decrease prepitant is a r rates of these g adjusted wh r) I metabolizer. | Polypharmacy guidance: The hage. Discontinue agents that iss essential. If co-administration ACTIONABLE onverted to aprepitant following at undergoes extensive A4 with minor involvement from etically guided drug selection or ate and strong CYP3A4 adverse reactions. These drugs aprepitant exposure resulting in moderate (dose-dependent) enzymes are contraindicated en coadministered with this ACTIONABLE Fosphenytoin can be prescribed |

| | Manch Univer | sity | PATIENT INFORMATION NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 | SPECIMEN DETAILS SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: | | БҮ |
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| | FOR ACADEMIC PURPOSES ONLY - NO | T FOR CLINICAL USE | SEX: | REPORT DATE: | 2/8/2018 | |
| | Galantamine | Normal Sensitivit | ty to Galantamine (CYP2I | 06: Intermediate Metabo | olizer) | INFORMATIV |
| | Razadyne | Galantamine can be with weekly titratior | e prescribed at standard labe n is recommended. | -recommended dosage and | d administration. Individu | alization of dose |
| √ | Glimepiride | Normal Sensitivit | ty to Glimepiride (CYP2C | 9: Normal Metabolizer) | | ACTIONABL |
| | Amaryl | • | prescribed according to stan levels of glucose/glycosylate | | osage and administration | (dose titration in |
| √ | Glipizide | Normal Sensitivit | ty to Glipizide (CYP2C9: N | lormal Metabolizer) | | INFORMATIV |
| | Glucotrol | | escribed according to standar I levels of glucose/glycosylate | | age and administration (d | ose titration in |
| \checkmark | Glyburide | Normal Sensitivit | ty to Glyburide (CYP2C9: | Normal Metabolizer) | | ACTIONABL |
| | Micronase | • | rescribed according to standa l levels of glucose/glycosylate | | sage and administration (| dose titration in |
| | Sancuso, Sustol | desmethylgranisetro women reported an clearance of the dru within the CYP3A4 of an association with is unclear and no ge Inducers or inhibito an in vivo pharmaco of granisetron with | guidance: Granisetron is extern on by CYP3A4, CYP3A5 and C in increased granisetron cleara ug in subjects with the CYP3A or ABCB1 genes, had no effec granisetron efficacy and ABC enetically guided drug selections of CYP1A1 and CYP3A4 en okinetic interaction with stror metabolizing enzyme induce change is not known. | YP1A1. A preliminary pharm ince in carriers of the CYP1A 5*3/*3 genotype. The same at on granisetron clearance B1 genetic polymorphisms. on or dosing recommendat zymes may affect the clear on CYP3A4 inhibitors such a | nacokinetic study conduct A1*2A increased function a study showed that gene while other reports in car The significance of these tions are available. Polyp ance of granisetron. Howe s ketoconazole is not kno | ted in pregnant allele and a lower tic polymorphisms cer patients found preliminary findings harmacy guidance: ever, the potential fo own. Administration |
| | Guanfacine | Normal Response | e to Guanfacine | | | INFORMATIV |
| - | Intuniv | or dosing recomme response and tolera should be reduced t ketoconazole, itracc should be increased recommended dose | guidance: Guanfacine is predendations are available and grability of the individual patien to one half of the standard onazole, indinavir, ritonavir, n d to the standard recommende when used in combination v . When the CYP3A4 inducer is e within 7-14 days. | uanfacine extended-release t. Polypharmacy guidance dose when co-medicated w efazodone). When the stror led dose. Guanfacine dose s with a strong CYP3A4 induc | should be titrated based a: The dose of guanfacine with a strong CYP3A4 inhi ng CYP3A4 inhibitor is dis should be increased up to er (e.g., phenytoin, carbar | on the clinical extended-release bitor (e.g., continued, the dose o double the mazepine, rifampin, |
| \checkmark | Haloperidol | Normal Sensitivit | ty to Haloperidol (CYP2D | 6: Intermediate Metabo | lizer) | ACTIONABL |
| | Haldol | | prescribed at standard label- l a favorable response is achie | | administration. Careful ti | tration is |
| | Hydromorphone | Normal Response | e to Hydromorphone | | | INFORMATIV |
| - | Dilaudid, Exalgo | No genetically guid CYPs, and genetic v | led drug selection or dosing i variations in these metabolizin in be prescribed at standard l | ng enzymes are not expecte | d to affect its efficacy or | |
| | | | | | | |

| V | Manch Univer | ester sity | PATIENT INFORMATION NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: 1/1/1900 | | ORDERED BY //1/1900 //1/1900 2/8/2018 | |
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| | FOR ACADEMIC PURPOSES ONLY - NOT | FOR CLINICAL USE | | | , | |
| | Ibuprofen | Normal Sensitivit | y to Ibuprofen (CYP2C9: No | rmal Metabolizer) | IN | FORMATIVI |
| - | Advil, Motrin | | ormal CYP2C9 activity (i.e norma -dosage and administration. | l metabolizers) can be pr | escribed ibuprofen according to | standard |
| | Indomethacin Indocin | Normal Sensitivit | y to Indomethacin (CYP2C9 | : Normal Metabolizer) | IN | FORMATIVE |
| | maocin | Indomethacin can be | e prescribed at standard label re | commended-dosage and | administration. | |
| | Irbesartan | Normal Sensitivit | y to Irbesartan (CYP2C9: No | rmal Metabolizer) | IN | FORMATIVE |
| | Avapro | Irbesartan can be pr | escribed at standard label-recor | nmended dosage and ad | ministration. | |
| \ | Isavuconazonium Cresemba | • | to Isavuconazonium uidance: Isavuconazonium sulf | ate is a prodrug that is ra | | CTIONABLE |
| | Cresemba | butylcholinesterase | nto its active moiety isavucona | cole. Isavuconazole is exte | ensively metabolized CYP3A4 and | |
| | | exposure. No geneti | cally guided drug selection or d | osing recommendations | e not expected to affect isavucor are available. Polypharmacy gu i I inhibitors or inducers contraind | dance: |
| | Itraconazole | exposure. No geneti Isavuconazole is a se Normal Response | cally guided drug selection or d ensitive CYP3A4 substrate and it to Itraconazole | osing recommendations s use with strong CYP3A4 | are available. Polypharmacy gu i I inhibitors or inducers contraind A | dance: cated. CTIONABLI |
| | Itraconazole Sporanox | exposure. No geneti Isavuconazole is a se Normal Response Pharmacogenetic g metabolite is hydrox concentrations of th recommendations a may decrease the bi Therefore, administr should be avoided 2 bioavailability of itra Itraconazole inhibit in increased plasma elevated plasma con using concomitant n | cally guided drug selection or d ensitive CYP3A4 substrate and it to Itraconazole guidance: Itraconazole is extens y-itraconazole, which has in vitr is metabolite are about twice th re available. Polypharmacy gui oavailability of itraconazole and ation of potent CYP3A4 inducer weeks before and during treatr conazole and these drugs shou the metabolism of drugs metab concentrations of these drugs a centrations may increase or pro- | osing recommendations s use with strong CYP3A4 ively metabolized to seve o antifungal activity com ose of itraconazole. No g dance: Coadministration hydroxy-itraconazole to s with itraconazole is not ment with itraconazole. P d be used with caution w olized by CYP3A4 or trans nd/or their active metabo- long both therapeutic ar | are available. Polypharmacy gu i I inhibitors or inducers contraind | dance: cated. .CTIONABLE nain lasma or dosing &A4 inducers be reduced. ese drugs ease the ifungal. may result ered. These When |
| ✓ ✓ | | exposure. No geneti Isavuconazole is a se Normal Response Pharmacogenetic g metabolite is hydrox concentrations of th recommendations a may decrease the bi Therefore, administr should be avoided 2 bioavailability of itra Itraconazole inhibit in increased plasma elevated plasma con using concomitant n | cally guided drug selection or d ensitive CYP3A4 substrate and it to ltraconazole guidance: Itraconazole is extens y-itraconazole, which has in vitr is metabolite are about twice th re available. Polypharmacy gui oavailability of itraconazole and ation of potent CYP3A4 inducer weeks before and during treatr conazole and these drugs shou the metabolism of drugs metab concentrations of these drugs a centrations may increase or pro- nedication, it is recommended to need for dose adjustments. | osing recommendations s use with strong CYP3A4 ively metabolized to seve o antifungal activity com ose of itraconazole. No g dance: Coadministration hydroxy-itraconazole to s with itraconazole is not ment with itraconazole. P d be used with caution w olized by CYP3A4 or trans nd/or their active metabo- long both therapeutic ar | are available. Polypharmacy gui I inhibitors or inducers contraind aral metabolites by CYP3A4. The r parable to itraconazole; trough p enetically guided drug selection of itraconazole with potent CYP3 such an extent that efficacy may recommended and the use of th botent CYP3A4 inhibitors may incr then coadministered with this and sported by P-glycoprotein, which blite(s) when they are coadminist d adverse effects of these drugs. bel be consulted for information of | dance: cated. CTIONABLE nain lasma or dosing BA4 inducers be reduced. ese drugs ease the ifungal. may result ered. These When on possible |
| ✓ ✓ | Sporanox | exposure. No geneti Isavuconazole is a se Normal Response Pharmacogenetic g metabolite is hydrox concentrations of th recommendations at may decrease the bi Therefore, administr should be avoided 2 bioavailability of itra Itraconazole inhibit f in increased plasma elevated plasma con using concomitant n contraindications or Normal Response Pharmacogenetic g and no major implic | cally guided drug selection or d ensitive CYP3A4 substrate and it to Itraconazole guidance: Itraconazole is extens y-itraconazole, which has in vitr is metabolite are about twice th re available. Polypharmacy gui oavailability of itraconazole and ation of potent CYP3A4 inducer weeks before and during treati conazole and these drugs shou the metabolism of drugs metab concentrations of these drugs a centrations may increase or pro- nedication, it is recommended t need for dose adjustments. to Ketoprofen guidance: Ketoprofen is primari | osing recommendations s use with strong CYP3A4 ively metabolized to seve o antifungal activity com ose of itraconazole. No g dance: Coadministration hydroxy-itraconazole to s with itraconazole is not nent with itraconazole. P d be used with caution w olized by CYP3A4 or trans nd/or their active metabo long both therapeutic ar hat the corresponding lal | are available. Polypharmacy gui I inhibitors or inducers contraind aral metabolites by CYP3A4. The r parable to itraconazole; trough p enetically guided drug selection of itraconazole with potent CYP3 such an extent that efficacy may recommended and the use of th botent CYP3A4 inhibitors may incr then coadministered with this and sported by P-glycoprotein, which blite(s) when they are coadminist d adverse effects of these drugs. bel be consulted for information of | dance: cated. CTIONABLE nain lasma or dosing SA4 inducers be reduced. ese drugs ease the ifungal. may result ered. These When on possible FORMATIVE I UGT2B7) |
| ✓ ✓ ✓ | Sporanox Ketoprofen | exposure. No geneti Isavuconazole is a se Normal Response Pharmacogenetic g metabolite is hydrox concentrations of th recommendations at may decrease the bi Therefore, administr should be avoided 2 bioavailability of itra Itraconazole inhibit f in increased plasma elevated plasma con using concomitant n contraindications or Normal Response Pharmacogenetic g and no major implic | cally guided drug selection or d ensitive CYP3A4 substrate and it to Itraconazole guidance: Itraconazole is extens y-itraconazole, which has in vitr is metabolite are about twice th re available. Polypharmacy gui oavailability of itraconazole and ation of potent CYP3A4 inducer weeks before and during treatr conazole and these drugs shou the metabolism of drugs metab concentrations of these drugs a centrations may increase or pro- nedication, it is recommended to need for dose adjustments. to Ketoprofen guidance: Ketoprofen is primari ation of CYP2C9 in the metabol recommendations are available. | osing recommendations s use with strong CYP3A4 ively metabolized to seve o antifungal activity com ose of itraconazole. No g dance: Coadministration hydroxy-itraconazole to s with itraconazole is not nent with itraconazole. P d be used with caution w olized by CYP3A4 or trans nd/or their active metabo long both therapeutic ar hat the corresponding lal | are available. Polypharmacy gui I inhibitors or inducers contraind aral metabolites by CYP3A4. The r parable to itraconazole; trough p enetically guided drug selection of itraconazole with potent CYP3 such an extent that efficacy may recommended and the use of th botent CYP3A4 inhibitors may incr then coadministered with this and sported by P-glycoprotein, which olite(s) when they are coadminist d adverse effects of these drugs. bel be consulted for information of IN idation (by UGT1A3, UGT1A9 and demonstrated. No genetically gui | dance: cated. CTIONABLE nain lasma or dosing SA4 inducers be reduced. ese drugs ease the ifungal. may result ered. These When on possible FORMATIVE I UGT2B7) |

| | Manch | octor | PATIENT INFORMATION | SPECIMEN DETAILS | ; | ORDERED BY |
|--------------|---|---|---|---|---|---|
| | FOR ACADEMIC PURPOSES ONLY - NOT | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: Image: Second Sec | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | | | | | | INFORMATIV |
| V | Labetalol Normodyne, Trandate | metabolites. Prelimi -fold higher in Chin clinical impact of th | guidance: Labetalol is extensive inary studies indicate that follow ese individuals with the CYP2C1 | /ing a single 200-mg ora 9 *2/*2 genotype than t rmacy guidance: Cimet | al dose, labeta hose with the | and CYP2C19 to inactive alol plasma concentrations are 2. |
| \checkmark | Lacosamide | Normal Sensitivit | y to Lacosamide (CYP2C19: | Rapid Metabolizer) | | INFORMATIV |
| | Vimpat | | wolved in the metabolism of lac ard label-recommended dosage | 5 | P2C9 and CYF | P3A, and this drug can be |
| √ | Lamotrigine | Normal Response | e to Lamotrigine | | | INFORMATIV |
| | | syndrome, Stevens- glucuronidation, wh insufficient studies of response. No genet Enzyme-inducing di maintain therapeuti lamotrigine levels a | documenting the impact of gen ically guided drug selection or c rugs increase lamotrigine cleara c concentrations. Coadministrat | xic epidermal necrolysis T1A4 with some contrib etic polymorphisms of t dosing recommendation nce significantly, and hig ion of valproic acid, an i gine adverse effects (new | (TEN). Lamot ution from U(hese metabol s are available gher doses of nhibitor of U(urological and | rigine is metabolized by GT1A1 and UGBT2B7. There are izing enzymes on lamotrigine e. Polypharmacy guidance: this drug are required to GT enzymes, increases d cutaneous). A low starting dose |
| \checkmark | Leflunomide | Normal Sensitivit | ty to Leflunomide (CYP2C19 | : Rapid Metabolizer) | | INFORMATIV |
| | Arava | count (CBC) and live | prescribed according to standa er function parameters should b e initial 6 months of therapy. Blo ter. | e checked no more thar | n 6 months be | efore beginning treatment, and |
| | Lesinurad | Normal Sensitivit | ty to Lesinurad (CYP2C9: No | ormal Metabolizer) | | ACTIONABL |
| | Zurampic | | ype predicts a normal CYP2C9 n age and administration. | netabolic activity. Lesinu | rad can be pr | escribed at standard label- |
| | Levetiracetam | Normal Response | e to Levetiracetam | | | INFORMATIV |
| | Keppra | Polypharmacy guid | guidance: No genetically guide dance: Levetiracetam is minima d in urine. Coadministration of e na levels. | lly metabolized by non- | CYP enzymes | (esterases) and is primarily |
| \checkmark | Levomilnacipran | Normal Response | e to Levomilnacipran | | | INFORMATIV |
| - | Fetzima | by CYP3A4, with mi in urine as unchang expected to have a | | YP2C19, CYP2D6, and C N-desethyl levomilnaci cipran exposure. no gen | (P2J2. More t pran. Genetic etically guide | han 58% of the dose is excreted polymorphisms of CYPs are not d drug selection or dosing |

| | Manch | actor | PATIENT INFORMATION | SPECIMEN DETAILS | | ORDERED BY |
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| | FOR ACADEMIC PURPOSES ONLY - NOT | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: Image: Comparison of the second | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| / | Levorphanol | Normal Response | e to Levornhanol | | | INFORMATIVI |
| | Levo Dromoran | Pharmacogenetic g studies documenting no genetically guide | • | phisms of this metaboli mmendations are availa | zing enzyme ble. Polypha | ediated by UGT2B7. There are no on levorphanol response. And |
| / | Lisdexamfetamine | Normal Exposure | to Lisdexamfetamine (CYP2 | 2D6: Intermediate M | etabolizer) | INFORMATIVE |
| _ | Vyvanse | | an be prescribed at standard lab the therapeutic needs and resp | | ge and admin | istration. Individualize the |
| | Lisdexamfetamine | Good Response to | o Lisdexamfetamine (COMT | : Intermediate COM | Г Activity) | INFORMATIVE |
| - | Vyvanse | | pe result predicts a favorable re lowest effective dose, and dosa | | | Lisdexamfetamine should be |
| / | Losartan | Normal Response | e to Losartan (CYP2C9: Norn | nal Metabolizer) | | INFORMATIV |
| - | Cozaar, Hyzaar | | zed to its active metabolite by C and its active metabolite. Losa | | | |
| / | Lovastatin | Normal Myopath | y Risk (SLCO1B1: Normal Fu | nction) | | INFORMATIVE |
| - | Mevacor, Altoprev, Advicor | are present, lovastat specific guidelines. C | na concentration is not expecte in can be prescribed at standarc Other myopathy predisposing fa atin dose, comedications, and fe | d FDA-recommended st ctors include advanced | arting doses a | |
| / | Lovastatin | Normal Response | e to Lovastatin (CYP3A4: No | rmal Metabolizer) | | INFORMATIVE |
| - | Mevacor, Altoprev, Advicor | | indicates that the patient does enzyme activity). The patient is e irements. | - | | |
| | Loxapine | Normal Response | e to Loxapine | | | INFORMATIVE |
| _ | Loxitane, Adasuve | metabolites formed. contributions from C these metabolizing e dosing recommenda concurrent use of Lc antidepressants, ger can increase the risk reduction/modificati | Loxapine metabolism occurs vi CYP3A4, CYP2D6 and FMO. Ther enzymes on Loxapine dispositio ations. Polypharmacy guidance oxapine with other CNS depressi- neral anesthetics, phenothiazine of respiratory depression, hypo ion of CNS depressants if used on h other anticholinergic drugs ca | a hydroxylation and oxio e are no studies docum n and there are no avail e: Loxapine is a central r ants (<i>e.g.</i> , alcohol, opioi s, sedative/hypnotics, m tension, profound seda concomitantly with Loxa | dation catalyz enting the eff able genetica nervous syste d analgesics, uscle relaxan tion, and syno pine. Loxapin | fect of genetic polymorphisms of Ily-guided drug selection or m (CNS) depressant. The benzodiazepines, tricyclic ts, and/or illicit CNS depressants) cope. Therefore, consider dose e has anticholinergic activity and |



| \mathbf{N} | 🖌 Mancl | iester | PATIENT INFORMATION | SPECIMEN DETAILS | , | ORDERED BY |
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| V | Univer | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| • | FOR ACADEMIC PURPOSES ONLY - NC | OT FOR CLINICAL USE | | | | |
| \checkmark | Lurasidone | - | se to Lurasidone | | | ACTIONABL |
| | Latuda | available. Polypha increase in luraside not be administer with moderate CYI strong inducers o | red with strong CYP3A4 inhib P3A4 inhibitors. Monitor patient of CYP3A should not be admin inducer, it may be necessary to | tant use of lurasidone wit ch could increase or prolo itors. Lurasidone dose sh ts receiving lurasidone an istered with lurasidone. | h all CYP3A4 ong adverse d ould not exce d any CYP3A4 . If lurasidone | inhibitors may result in an rug effects. Lurasidone should eed 40 mg when administered i inhibitor. Rifampin or other |
| | Meloxicam | Normal Sensitiv | ity to Meloxicam (CYP2C9: | Normal Metabolizer) | | INFORMATIV |
| | Mobic | | a concentrations are not expecte sage and administration. | ed to be altered. Meloxica | m can be pres | scribed at standard label- |
| | Memantine | Normal Respons | se to Memantine | | | INFORMATIV |
| | | | m to three inactive metabolites 50 enzymes do not play a signif | | | |
| | | metabolite). CYP45 documenting the e response. No gene Memantine is prec not expected to in of drugs that use t | 50 enzymes do not play a signif effects of genetic variability in n etically guided drug selection or dominantly renally eliminated, a | icant role in the metabolis netabolizing enzymes or c dosing recommendation nd drugs that are substrate memantine is eliminated including hydrochlorothia | sm of meman organic cation is are available tes and/or inh l in part by tul azide, triamter | tine. There are no studies ic transporters on memantine e. Polypharmacy Guidance: nibitors of the CYP450 system are bular secretion, coadministration rene, metformin, cimetidine, |
| ✓ | Meperidine | metabolite). CYP45 documenting the e response. No gene Memantine is prec not expected to in of drugs that use t ranitidine, quinidir | 50 enzymes do not play a signif effects of genetic variability in n etically guided drug selection or dominantly renally eliminated, a teract with memantine. Because the same renal cationic system, | icant role in the metabolis netabolizing enzymes or c dosing recommendation nd drugs that are substrate memantine is eliminated including hydrochlorothia | sm of meman organic cation is are available tes and/or inh l in part by tul azide, triamter | tine. There are no studies ic transporters on memantine e. Polypharmacy Guidance: nibitors of the CYP450 system are bular secretion, coadministration rene, metformin, cimetidine, |
| √ | Meperidine Demerol | metabolite). CYP45 documenting the e response. No gene Memantine is prec not expected to in of drugs that use t ranitidine, quinidir Normal Respons Pharmacogenetic is metabolized to r variants in these ei meperidine metab ritonavir, meperidi these findings, the increased concent | 50 enzymes do not play a signif effects of genetic variability in n etically guided drug selection or dominantly renally eliminated, a teract with memantine. Because the same renal cationic system, ne, and nicotine, could potential se to Meperidine c guidance: no genetically guid normeperidine by multiple CYPs nzymes have not been studied. polism is increased resulting in h ine's exposure is significantly re- erisk of narcotic-related adverse | icant role in the metabolis netabolizing enzymes or con- dosing recommendation and drugs that are substrate ememantine is eliminated including hydrochlorothia ly result in altered plasma ed drug selection or dosir s, including CYP2B6, CYP3 Polypharmacy guidance igher levels of its neuroto duced while normeperidir e effects from this combin | sm of memani- organic cation is are available tes and/or inh d in part by tul azide, triamter a levels of both ng recommen- A4, and CYP2: In patients to pic metabolite ne concentrati ation appears | tine. There are no studies ic transporters on memantine e. Polypharmacy Guidance: hibitors of the CYP450 system are bular secretion, coadministration rene, metformin, cimetidine, h agents. INFORMATIV dations are available. Meperidine C19. The effects of genetic taking strong CYP inducers , e normeperidine. In presence of ions are increased. Based on |
| √ √ | Demerol | metabolite). CYP45 documenting the e response. No gene Memantine is prec not expected to in of drugs that use t ranitidine, quinidir Normal Respons Pharmacogenetic is metabolized to r variants in these en meperidine metab ritonavir, meperidi these findings, the increased concent This combination s | 50 enzymes do not play a signif effects of genetic variability in n etically guided drug selection or dominantly renally eliminated, a iteract with memantine. Because the same renal cationic system, he, and nicotine, could potential se to Meperidine c guidance: no genetically guid normeperidine by multiple CYPs nzymes have not been studied. bolism is increased resulting in h ine's exposure is significantly re- erisk of narcotic-related adverse rations of normeperidine sugge should be avoided is possible. | icant role in the metabolis netabolizing enzymes or con- dosing recommendation and drugs that are substrate ememantine is eliminated including hydrochlorothia ly result in altered plasma ed drug selection or dosir s, including CYP2B6, CYP3 Polypharmacy guidance igher levels of its neuroto duced while normeperidir e effects from this combin | sm of memani- organic cation is are available tes and/or inh d in part by tul azide, triamter a levels of both ng recommen- A4, and CYP2: In patients to pic metabolite ne concentrati ation appears | tine. There are no studies ic transporters on memantine e. Polypharmacy Guidance: hibitors of the CYP450 system are bular secretion, coadministration rene, metformin, cimetidine, h agents. INFORMATIV dations are available. Meperidine C19. The effects of genetic taking strong CYP inducers , e normeperidine. In presence of ions are increased. Based on is to be minimal. However, |
| ✓ ✓ | - | metabolite). CYP45 documenting the e response. No gene Memantine is prec not expected to in of drugs that use t ranitidine, quinidir Normal Respons Pharmacogenetic is metabolized to r variants in these el meperidine metab ritonavir, meperidi these findings, the increased concent This combination s Normal Respons Pharmacogenetic CYP2D6, CYP2E1, a | 50 enzymes do not play a signif effects of genetic variability in n etically guided drug selection or dominantly renally eliminated, a iteract with memantine. Because the same renal cationic system, ne, and nicotine, could potential se to Meperidine c guidance: no genetically guid normeperidine by multiple CYPs nzymes have not been studied. wolism is increased resulting in h ine's exposure is significantly re- erisk of narcotic-related adverse rations of normeperidine sugge should be avoided is possible. se to Metaxalone c guidance: Metaxalone is exter | icant role in the metabolis netabolizing enzymes or con- dosing recommendation and drugs that are substrate ememantine is eliminated including hydrochlorothia ly result in altered plasma ed drug selection or dosir s, including CYP2B6, CYP3 Polypharmacy guidance igher levels of its neuroto duced while normeperidir effects from this combin st a potential for toxicity of nsively metabolized by mu- nisms of these enzymes an | sm of memani organic cation is are available tes and/or inh d in part by tul azide, triamter a levels of both mg recommen A4, and CYP2 in patients t oxic metabolite the concentrati ation appears with increased | tine. There are no studies ic transporters on memantine e. Polypharmacy Guidance: hibitors of the CYP450 system are bular secretion, coadministration rene, metformin, cimetidine, h agents. INFORMATIV dations are available. Meperidine C19. The effects of genetic taking strong CYP inducers , e normeperidine. In presence of ions are increased. Based on to be minimal. However, d dosages or long-term therapy. INFORMATIV |
| ✓ ✓ ✓ | Demerol Metaxalone | metabolite). CYP45 documenting the e response. No gene Memantine is prec not expected to in of drugs that use t ranitidine, quinidir Normal Respons Pharmacogenetic is metabolized to r variants in these en meperidine metab ritonavir, meperidi these findings, the increased concent This combination s Normal Respons Pharmacogenetic CYP2D6, CYP2E1, a extent. no genetica | 50 enzymes do not play a signif effects of genetic variability in n etically guided drug selection or dominantly renally eliminated, a teract with memantine. Because the same renal cationic system, ne, and nicotine, could potential se to Meperidine c guidance: no genetically guid normeperidine by multiple CYPs nzymes have not been studied. Poolism is increased resulting in h ine's exposure is significantly re- erisk of narcotic-related adverse rations of normeperidine sugge should be avoided is possible. se to Metaxalone c guidance: Metaxalone is exter and CYP3A4. Genetic polymorpl | icant role in the metabolis netabolizing enzymes or of dosing recommendation and drugs that are substrate memantine is eliminated including hydrochlorothia ly result in altered plasma ed drug selection or dosin s, including CYP2B6, CYP3 Polypharmacy guidance igher levels of its neuroto duced while normeperidir effects from this combin st a potential for toxicity of history of these enzymes an using recommendations an | sm of memani organic cation is are available tes and/or inh d in part by tul azide, triamter a levels of both mg recommen A4, and CYP2 in patients t oxic metabolite the concentrati ation appears with increased | tine. There are no studies ic transporters on memantine e. Polypharmacy Guidance: hibitors of the CYP450 system are bular secretion, coadministration rene, metformin, cimetidine, h agents. INFORMATIV dations are available. Meperidine C19. The effects of genetic taking strong CYP inducers , e normeperidine. In presence of ions are increased. Based on to be minimal. However, d dosages or long-term therapy. INFORMATIV zymes, including CYP1A2, |
| ✓ ✓ ✓ | Demerol Metaxalone Skelaxin | metabolite). CYP45 documenting the e response. No gene Memantine is prec not expected to in of drugs that use t ranitidine, quinidir Normal Respons Pharmacogenetic is metabolized to r variants in these er meperidine metab ritonavir, meperidi these findings, the increased concent This combination s Normal Respons Pharmacogenetic CYP2D6, CYP2E1, a extent. no genetica | 50 enzymes do not play a signif effects of genetic variability in n etically guided drug selection or dominantly renally eliminated, a iteract with memantine. Because the same renal cationic system, he, and nicotine, could potential se to Meperidine c guidance: no genetically guid normeperidine by multiple CYPs nzymes have not been studied. bolism is increased resulting in h ine's exposure is significantly re- erisk of narcotic-related adverse rations of normeperidine sugge should be avoided is possible. se to Metaxalone c guidance: Metaxalone is exter and CYP3A4. Genetic polymorpl ally guided drug selection or do | icant role in the metabolis netabolizing enzymes or control of the substration of drugs that are substration including hydrochlorothia ly result in altered plasma ed drug selection or dosir s, including CYP2B6, CYP3 Polypharmacy guidance igher levels of its neuroto duced while normeperidir e effects from this combin st a potential for toxicity of insime of these enzymes and issing recommendations an Normal Metabolizer) | sm of memani organic cation is are available tes and/or inh d in part by tul azide, triamter a levels of both ng recommen A4, and CYP2 in patients to acconcentrati ation appears with increased ultiple CYP enz re unlikely to a re available. | tine. There are no studies ic transporters on memantine e. Polypharmacy Guidance: hibitors of the CYP450 system are bular secretion, coadministration rene, metformin, cimetidine, h agents. INFORMATIV dations are available. Meperidine C19. The effects of genetic taking strong CYP inducers , e normeperidine. In presence of ions are increased. Based on to be minimal. However, d dosages or long-term therapy. INFORMATIV zymes, including CYP1A2, affect its exposure to a significan |
| ✓ ✓ ✓ | Demerol Demerol Metaxalone Skelaxin | metabolite). CYP45 documenting the e response. No gene Memantine is prec not expected to in of drugs that use t ranitidine, quinidir Normal Respons Pharmacogenetic is metabolized to r variants in these er meperidine metab ritonavir, meperidi these findings, the increased concent This combination s Normal Respons Pharmacogenetic CYP2D6, CYP2E1, a extent. no genetica Normal Sensitiv Methadone can be precautions. | 50 enzymes do not play a signif effects of genetic variability in n etically guided drug selection or dominantly renally eliminated, a tteract with memantine. Because the same renal cationic system, ne, and nicotine, could potential se to Meperidine c guidance: no genetically guid normeperidine by multiple CYPs nzymes have not been studied. Polism is increased resulting in h ine's exposure is significantly re- ersis of narcotic-related adverse rations of normeperidine sugge should be avoided is possible. se to Metaxalone c guidance: Metaxalone is exter and CYP3A4. Genetic polymorpl ally guided drug selection or do rity to Methadone (CYP2B6: | icant role in the metabolis netabolizing enzymes or control of the substration of drugs that are substration including hydrochlorothia ly result in altered plasma ed drug selection or dosir s, including CYP2B6, CYP3 Polypharmacy guidance igher levels of its neuroto duced while normeperidir e effects from this combin st a potential for toxicity of insime of these enzymes and issing recommendations an Normal Metabolizer) | sm of memani organic cation is are available tes and/or inh d in part by tul azide, triamter a levels of both ng recommen A4, and CYP2 in patients to acconcentrati ation appears with increased ultiple CYP enz re unlikely to a re available. | tine. There are no studies ic transporters on memantine e. Polypharmacy Guidance: hibitors of the CYP450 system are bular secretion, coadministration rene, metformin, cimetidine, h agents. INFORMATIV dations are available. Meperidine C19. The effects of genetic taking strong CYP inducers , e normeperidine. In presence of ions are increased. Based on to be minimal. However, d dosages or long-term therapy. INFORMATIV zymes, including CYP1A2, affect its exposure to a significan |

| FOR ACADEMIC P Methot: Methot: Trexall Micafur Micafur Micafur Mirabeg Myrbetriq Mirtaza Mirtaza Mirtaza Mirtaza Morphin Mabuma Nabuma Naproxa Aleve | - • | hester | PATIENT INFORMATION NAME: Patient 33169 | SPECIMEN DETAILS | ORDERED BY |
|---|---------|---|---|--|---|
| Methot Trexall Micafur Mycamine Milnaci Savella Mirabeg Myrbetriq Mirtaza Remeron Morphin MS Contin Morphin MS Contin Morphin MS Contin Nabume Relafen | Jnive | | ACC #: 33169 DOB: 1/1/1900 SEX: | COLLECTION DATE: 1/1/1 RECEIVED DATE: 1/1/1 REPORT DATE: 2/8/2 | 1900 |
| Trexall Micafur Mycamine Milnacig Savella Mirabeg Myrbetriq Mirtaza Remeron Morphin MS Contin Morphin MS Contin Morphin MS Contin Nabuma Relafen Naproxe | | | athatravata tavisity (NATH | | v) INFORMATIVE |
| Mycamine Milnacif Savella Mirabeg Myrbetriq Mirtaza Remeron Morphin MS Contin Morphin MS Contin Nabume Relafen Nabume | Juexale | The patient does no | | and unless other risk factors a | are present, the patient is not expected to ded dosage and administration. |
| Milnacig Savella Mirabeg Myrbetriq Mirtaza Remeron Morphin MS Contin Morphin MS Contin Nabuma Relafen Naproxe | ungin | Normal Response | e to Micafungin | | ACTIONABLE |
| Savella Savella Mirabeg Myrbetriq Mirtaza Remeron Morphin MS Contin Morphin MS Contin Morphin MS Contin Morphin MS Contin Nabume Relafen Naproxe | ine | P450 enzymes. Ever | h though micafungin is a substr vay for micafungin metabolism | ate for and a weak inhibitor of | ol-O-methyltransferase and cytochrome f CYP3A in vitro, hydroxylation by CYP3A l drug selection or dosing |
| Myrbetriq Myrbetriq Mirtaza Remeron Morphi MS Contin MS Contin NS Contin | cipran | in urine. No genetic | guidance: milnacipran is minim ally guided drug selection or d | osing recommendations are av | INFORMATIVE ymes and primarily excreted unchanged vailable. Polypharmacy guidance: ly to affect the exposure of milnacipran. |
| Remeron Morphin MS Contin MS Contin | | | y to Mirabegron (CYP2D6: prescribed at standard label-re | | - |
| MS Contin | • | Mirtazapine can be | y to Mirtazapine (CYP2D6: prescribed at standard label-re a favorable response is achieve | commended dosage and adm | |
| MS Contin | | The patient does no experience good an | | itation. Acute postoperative al oses. The dosing regimen nee | INFORMATIVE nd cancer pain: the patient is expected to eds to be individualized for each patient, |
| Relafen | | The patient carries of require average to l | | , which translates to a reduced uate pain control. The dosing | INFORMATIVE d COMT function. The patient may regimen needs to be individualized for nce. |
| | netone | that is further metal (i.e CYP2C9 poor me an altered drug resp Guidance: CYP1A2 the therapeutic effe | guidance: Nabumetone is a pro- polized by CYP2C9 to an inactive etabolizers) may have higher le ponse. No genetically guided de inhibitors may inhibit the active | e metabolite. Theoretically, in vels of the active metabolite, k ug selection or dosing recom ation of nabumetone to its act and, CYP1A2 inducers (i.e smo | INFORMATIVE CYP1A2 to an active metabolite (6-MNA) dividuals with reduced CYP2C9 activity but it is unknown whether this results in mendations are available. Polypharmacy ive metabolite resulting in a reduction in bking) may result in higher levels of |
| , neve | oxen | elimination pathway desmethylnaproxen | y for this drug (60% of total clear but this pathway is not the pri peen found to affect the respon | arance). CYP2C9 and CYP1A2 a mary pathway for the eliminat | INFORMATIVE Ilucuronidation, which is the primary are responsible for the formation of O- ion for naproxen. Genetic polymorphism y guided drug selection or dosing |
| Powered By Translational software | าอไ | | Genetic Test Results For Pati | | Page 29 of 64 |

| $\overline{\mathbf{N}}$ | A Mancl | lester | PATIENT INFORMATION | SPECIMEN DETAILS | ORDERED | BY |
|-------------------------|---------------------------------|---|--|--|---|---|
| V | Univer | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | | 1/1/1900 1/1/1900 2/8/2018 | |
| | FOR ACADEMIC PURPOSES ONLY - NC | | | | | |
| | Nateglinide | | ty to Nateglinide (SLCO1B1: | | | INFORMATIV |
| | Starlix | | two copies of SLCO1B1 rs41490 prescribed at label-recommenc | | | orter function. |
| | Nateglinide | Normal Sensitivi | ty to Nateglinide (CYP2C9: | Normal Metabolizer) | | INFORMATIV |
| | Starlix | The patient's genot dosage and admini | ype predicts a normal exposure stration. | to nateglinide, and this c | drug can be prescribed at | label-recommended |
| | Nefazodone | Normal Sensitivi | ty to Nefazodone (CYP2D6: | Intermediate Metabo | lizer) | INFORMATIV |
| - | Serzone | chlorophenylpipera | abolized by CYP3A4 to its active zine metabolite which may con prescribed standard label reco | tribute to adverse events, | is further metabolized by | |
| | Netupitant- | Normal Respons | e to Netupitant-Palonosetr | on (CYP2D6: Intermed | liate Metabolizer) | INFORMATIV |
| | Palonosetron | | | | | |
| | Akynzeo | derivatives). Metabo guided drug selecti label-recommended | tant is extensively metabolized to olism is mediated primarily by C on or dosing recommendations d dosage and administration. hosetron can be prescribed at s | YP3A4 and to a lesser ext are available for this dru | tent by CYP2C9 and CYP2 g. Netupitant can be pres | D6. No genetically cribed at standard |
| | Olmesartan | Normal Sensitivi | ty to Olmesartan Medoxom | il | | ACTIONABL |
| | Benicar | gastrointestinal trac | guidance: Olmesartan medoxo ct during absorption. There is vi enes is not expected to affect t s are available. | rtually no further metabo | lism of olmesartan. Genet | ic variability of the |
| V | Ondansetron | Normal Respons | e to Ondansetron (CYP2D6 | Intermediate Metabo | blizer) | INFORMATIV |
| | Zofran, Zuplenz | Ondansetron can b | e prescribed at standard label-r | ecommended dosage and | d administration. | |
| | Oxcarbazepine | Normal Respons | e to Oxcarbazepine | | | INFORMATIV |
| - | Trileptal, Oxtellar XR | be used to identify syndrome, Stevens- by a reductase to it eliminated by direc or dosing recomme | guidance: Genotype results ob patients at risk for severe cutan -Johnson syndrome (SJS) and to s active monohydroxylated acti t renal excretion, glucuronidatio endations are available. Polyph e active metabolite (MHD) are d | eous adverse reactions su xic epidermal necrolysis (ve metabolite: 10-hydroxy on, and hydroxylation (mir armacy guidance: In the | uch as anticonvulsant hyp (TEN). Oxcarbazepine (pro ycarbazepine (MHD). This nimal). No genetically guid | ersensitivity drug) in converted active metabolite is ded drug selection |
| | Oxybutynin | Normal Respons | e to Oxybutynin | | | INFORMATIV |
| _ | Ditropan | Polypharmacy gui | guidance: no genetically guide dance: Oxybutynin is extensive bitor (itraconazole) increases ox | ly metabolized in humans | s by CYP3A4, and coadmir | nistration of a |

| | Mancl | iestel. | NAME: Patient 33169 | SPECIMEN TYPE: | | |
|---|---------------------------------|--|--|---|---|---|
| X | Univer | SILY | ACC #: 33169 DOB: 1/1/1900 | COLLECTION DATE: RECEIVED DATE: | 1/1/1900 | |
| | FOR ACADEMIC PURPOSES ONLY - NO | OT FOR CLINICAL USE | SEX: | REPORT DATE: | 2/8/2018 | |
| / | Oxymorphone | Normal Respons | se to Oxymorphone | | | INFORMATI |
| - | Opana, Numorphan | CYPs, and genetic | ded drug selection or dosing r variations in these metabolizin be prescribed at standard lab | g enzymes are not expecte | ed to affect its | efficacy or toxicity profiles. |
| / | Paliperidone | Normal Sensitiv | ity to Paliperidone (CYP2D | 6: Intermediate Metabo | olizer) | ACTIONABI |
| | Invega | Paliperidone can b | e prescribed at standard label | -recommended dosage and | d administratio | on. |
| / | Palonosetron | Normal Respons | se to Palonosetron (CYP2D | 96: Intermediate Metabo | olizer) | INFORMATIV |
| | Aloxi | Palonosetron can b | be prescribed at standard labe | l-recommended dosage an | ıd administrati | on. |
| | Paroxetine | Normal Sensitiv | ity to Paroxetine (CYP2D6 | Intermediate Metaboli | izer) | ACTIONABI |
| | Paxil, Brisdelle | | prescribed at standard label-re til a favorable response is achie | - | administration | . Careful titration is |
| | Perampanel Fycompa | Pharmacogenetic and CYP3A5. No g Enzyme-inducing should be increase Coadministration v | • | on or dosing recommendati asma concentrations by 50 therapy regimen containin thers than antiepileptic dru | ions are availa)-60%, and the g enzyme-ind ugs (e.g., rifam | ucing antiepileptic drugs. pin) should be avoided. |
| | Phenobarbital | Normal Sensitiv | ity to Phenobarbital (CYP2 | C19: Rapid Metabolizer | ·) | INFORMATI |
| | Luminal | | involved in the metabolism of sage and administration. | phenobarbital, and this dru | ıg can be pres | cribed at standard label- |
| / | Phenytoin | Normal Sensitiv | ity to Phenytoin (CYP2C9: | Normal Metabolizer) | | ACTIONABI |
| | Dilantin | | | | | Phenytoin can be prescribed at concentrations 7-10 days after |
| | Pimavanserin | Normal Respons | se to Pimavanserin | | | INFORMATIV |
| | Nuplazid | by CYP2J2, CYP2D6 major active metab Polypharmacy gu | 6, and other CYP and FMO enz bolite (AC-279). There are no a idance: Pimavanserin prolong | ymes. CYP3A4 is the major vailable genetically-guided s the QT interval and its us | enzyme respo drug selection e should be av interval includi | n or dosing recommendations. voided in patients with known ing Class 1A antiarrhythmics |



| V | Unive | hester rsity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
|---|--|--|---|---|--|---|
| | FOR ACADEMIC PURPOSES ONLY - | NOT FOR CLINICAL USE | | | _, _, _, | |
| \checkmark | Pimozide | Normal Sensitivit | ty to Pimozide (CYP2D6: In | itermediate Metaboliz | zer) | ACTIONABL |
| | Orap | | escribed at standard label-recc <g (children).="" b<="" day="" doses="" may="" td=""><td></td><td></td><td></td></g> | | | |
| \ | Piroxicam | Normal Sensitivit | ty to Piroxicam (CYP2C9: N | lormal Metabolizer) | | INFORMATIV |
| | Feldene | Piroxicam can be pr | rescribed at standard label-reco | ommended dosage and a | administration. | |
| \ | Pitavastatin | • • | ny Risk (SLCO1B1: Normal F | | | INFORMATIV |
| | Livalo | are present, pitavas specific guidelines. | The myopathy risk increases w | dard FDA-recommended ith use of the 4 mg daily o | starting doses dose. (Other m | and adjusted based on disease |
| | Posaconazole | Normal Response | e to Posaconazole | | | ACTIONABL |
| | Noxafil | direct glucuronidati | or approximately 17% of the a ion, minor oxidation and dealk | ylation. CYP3A4 (and pose | sibly CYP1A1 a | nd CYP3A5), UGT1A4, and P- |
| | | direct glucuronidati glycoprotein are en drug selection or do inducers may affect avoided unless the | ion, minor oxidation and dealk zymes and transporters that pl osing recommendations are av posaconazole plasma concent benefit to the patient outweigl | ylation. CYP3A4 (and poss lay a role in the eliminatio railable. Polypharmacy g i trations. Concomitant use | sibly CYP1A1 a on of this antifu j uidance: UGT | nd CYP3A5), UGT1A4, and P- ungal. No genetically guided and P-glycoprotein inhibitors o ole and these agents should be |
| ✓ | Prasugrel | direct glucuronidati glycoprotein are en drug selection or de inducers may affect avoided unless the Normal Response | ion, minor oxidation and dealk zymes and transporters that pl osing recommendations are av posaconazole plasma concent benefit to the patient outweigh e to Prasugrel | ylation. CYP3A4 (and poss lay a role in the eliminatio ailable. Polypharmacy g trations. Concomitant use hs the risk. | sibly CYP1A1 a on of this antifu uidance: UGT e of posaconazo | And CYP3A5), UGT1A4, and P- ungal. No genetically guided and P-glycoprotein inhibitors o ole and these agents should be ACTIONABL |
| Image: A start of the start of | | direct glucuronidati glycoprotein are en drug selection or do inducers may affect avoided unless the Normal Response Pharmacogenetic converted to the ac Prasugrel active me efficacy or safety pr drug selection or do | ion, minor oxidation and dealk izymes and transporters that pl osing recommendations are av posaconazole plasma concent benefit to the patient outweigh e to Prasugrel guidance : Prasugrel is a prodr tive metabolite primarily by CY tabolite exposure and platelet ofile are also unaffected by CY | ylation. CYP3A4 (and pose lay a role in the eliminatio railable. Polypharmacy g i trations. Concomitant use hs the risk. ug that is hydrolyzed in th (P3A4 and CYP2B6, and to reactivity are not affected (P2B6, CYP3A5, and CYP2C railable. Polypharmacy g i | sibly CYP1A1 a on of this antifu uidance: UGT e of posaconazo he intestine to o a lesser exten d by CYP2C19 g C9 genetic varia | nd CYP3A5), UGT1A4, and P- ingal. No genetically guided and P-glycoprotein inhibitors o ole and these agents should be ACTIONABL a thiolactone, which is then nt by CYP2C9 and CYP2C19. genetic variants. Prasugrel |
| ✓ ✓ | Prasugrel | direct glucuronidati glycoprotein are en drug selection or do inducers may affect avoided unless the Normal Response Pharmacogenetic converted to the ac Prasugrel active me efficacy or safety pr drug selection or do drugs that are induc | ion, minor oxidation and dealk zymes and transporters that pl osing recommendations are av posaconazole plasma concent benefit to the patient outweigh e to Prasugrel guidance : Prasugrel is a prodr twe metabolite primarily by CY stabolite exposure and platelet rofile are also unaffected by CY osing recommendations are av | ylation. CYP3A4 (and pose lay a role in the eliminatio iailable. Polypharmacy g i trations. Concomitant use hs the risk. ug that is hydrolyzed in th (P3A4 and CYP2B6, and to reactivity are not affected (P2B6, CYP3A5, and CYP2(iailable. Polypharmacy g i le P450 enzymes. | sibly CYP1A1 a on of this antifu uidance: UGT e of posaconazo he intestine to o a lesser exten d by CYP2C19 g C9 genetic varia | ACTIONABL a thiolactone, which is then thy CYP2C9 and CYP2C19. genetic variants. No genetically-guided |
| ✓ ✓ | Prasugrel Effient | direct glucuronidati glycoprotein are en drug selection or do inducers may affect avoided unless the Normal Response Pharmacogenetic converted to the ac Prasugrel active me efficacy or safety pr drug selection or do drugs that are induce Normal Myopath Pravastatin plasma present, pravastatin specific guidelines. | ion, minor oxidation and dealk zymes and transporters that pl osing recommendations are av posaconazole plasma concent benefit to the patient outweigh e to Prasugrel guidance : Prasugrel is a prodr tive metabolite primarily by CY etabolite exposure and platelet rofile are also unaffected by CY osing recommendations are av cers or inhibitors of cytochrom by Risk (SLCO1B1: Normal F | ylation. CYP3A4 (and poss lay a role in the eliminatio ailable. Polypharmacy g trations. Concomitant use hs the risk. ug that is hydrolyzed in th (P3A4 and CYP2B6, and to reactivity are not affected (P2B6, CYP3A5, and CYP2C railable. Polypharmacy g te P450 enzymes. Function) ed to increase, and unless I FDA-recommended start g factors include advanced | sibly CYP1A1 a on of this antifu juidance: UGT of posaconazo he intestine to o a lesser exten d by CYP2C19 of C9 genetic vari- juidance: Prasu | INDERCEMPTION INDERCEMPTION INDERCEMPTION INDERCEMPTION INFORMATIV or circumstantial risk factors are adjusted based on disease- |
| ✓ ✓ ✓ | Prasugrel Effient Pravastatin | direct glucuronidati glycoprotein are en drug selection or do inducers may affect avoided unless the Normal Response Pharmacogenetic converted to the ac Prasugrel active me efficacy or safety pr drug selection or do drugs that are induce Normal Myopath Pravastatin plasma present, pravastatin specific guidelines. | ion, minor oxidation and dealk izymes and transporters that pl osing recommendations are av posaconazole plasma concent benefit to the patient outweigh e to Prasugrel guidance: Prasugrel is a prodr tive metabolite primarily by CY atabolite exposure and platelet ofile are also unaffected by CY osing recommendations are av cers or inhibitors of cytochrom ty Risk (SLCO1B1: Normal F concentrations are not expected of the prescribed at standard (Other myopathy predisposing igh statin dose, comedications | ylation. CYP3A4 (and poss lay a role in the eliminatio ailable. Polypharmacy g trations. Concomitant use hs the risk. ug that is hydrolyzed in th (P3A4 and CYP2B6, and to reactivity are not affected (P2B6, CYP3A5, and CYP2C railable. Polypharmacy g the P450 enzymes. Function) ed to increase, and unless I FDA-recommended start of factors include advanced | sibly CYP1A1 a on of this antifu juidance: UGT of posaconazo he intestine to o a lesser exten d by CYP2C19 of C9 genetic vari- juidance: Prasu | INDERCEMPTION INDERCEMPTION INDERCEMPTION INDERCEMPTION INFORMATIV or circumstantial risk factors are adjusted based on disease- |
| ✓ ✓ ✓ | Prasugrel Effient Pravastatin Pravachol | direct glucuronidati glycoprotein are en drug selection or do inducers may affect avoided unless the Normal Response Pharmacogenetic g converted to the ac Prasugrel active me efficacy or safety pr drug selection or do drugs that are induce Normal Myopath Pravastatin plasma present, pravastatin specific guidelines. renal impairment, h Normal Response Pharmacogenetic g Polypharmacy guid Genetic variations in | ion, minor oxidation and dealk izymes and transporters that pl osing recommendations are av posaconazole plasma concent benefit to the patient outweigh e to Prasugrel guidance : Prasugrel is a prodr tive metabolite primarily by CY etabolite exposure and platelet rofile are also unaffected by CY osing recommendations are av cers or inhibitors of cytochrom hy Risk (SLCO1B1: Normal F concentrations are not expected a can be prescribed at standard (Other myopathy predisposing igh statin dose, comedications e to Pregabalin guidance: No genetically guid dance: Pregabalin is eliminated | ylation. CYP3A4 (and poss lay a role in the eliminatio ailable. Polypharmacy g trations. Concomitant use hs the risk. ug that is hydrolyzed in th (P3A4 and CYP2B6, and to reactivity are not affected (P2B6, CYP3A5, and CYP2C railable. Polypharmacy g the P450 enzymes. Function) ed to increase, and unless I FDA-recommended start g factors include advanced ; and female gender.) ed drug selection or dosin d primarily through renal are not expected to affec | sibly CYP1A1 a on of this antifu juidance: UGT e of posaconazo he intestine to o a lesser extend d by CYP2C19 c C9 genetic varia juidance: Prasu s other genetic ting doses and d age (≥65), un | INFORMATIV Or circumstantial risk factors are adjusted based on disease- icontrolled hypothyroidism, INFORMATIV dations are available. |
| ✓ ✓ ✓ | Prasugrel Effient Pravastatin Pravachol Pregabalin | direct glucuronidati glycoprotein are en drug selection or do inducers may affect avoided unless the Normal Response Pharmacogenetic g converted to the ac Prasugrel active me efficacy or safety pr drug selection or do drugs that are induce Normal Myopath Pravastatin plasma present, pravastatin specific guidelines. renal impairment, h Normal Response Pharmacogenetic g Polypharmacy gui Genetic variations in be prescribed at sta | ion, minor oxidation and dealk izymes and transporters that pl osing recommendations are av posaconazole plasma concent benefit to the patient outweigh e to Prasugrel guidance : Prasugrel is a prodr tive metabolite primarily by CV etabolite exposure and platelet rofile are also unaffected by CV osing recommendations are av cers or inhibitors of cytochrom hy Risk (SLCO1B1: Normal F concentrations are not expected a can be prescribed at standard (Other myopathy predisposing igh statin dose, comedications e to Pregabalin guidance: No genetically guid dance: Pregabalin is eliminated in these metabolizing enzymes | ylation. CYP3A4 (and poss lay a role in the eliminatio iailable. Polypharmacy g trations. Concomitant use hs the risk. ug that is hydrolyzed in th (P3A4 and CYP2B6, and to reactivity are not affected (P2B6, CYP3A5, and CYP2C iailable. Polypharmacy g ie P450 enzymes. Function) ed to increase, and unless I FDA-recommended start factors include advanced i, and female gender.) ed drug selection or dosii d primarily through renal are not expected to affect isage and administration. | sibly CYP1A1 a on of this antifu juidance: UGT e of posaconazo he intestine to o a lesser extend d by CYP2C19 c C9 genetic varia juidance: Prasu s other genetic ting doses and d age (≥65), un | INFORMATIV or circumstantial risk factors are adjusted based on disease- icontrolled hypothyroidism, INFORMATIV dations are available. is not metabolized by CYPs. |

| | / Manc | hester | PATIENT INFORMATION | SPECIMEN DETAILS | | RDERED BY |
|---|--------------------------------|--|---|---|---|---|
| | Univer | rsity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: 1/1/1900 | | 1/1/1900 1/1/1900 2/8/2018 | |
| | FOR ACADEMIC PURPOSES ONLY - N | IOT FOR CLINICAL USE | | | | |
| | Proguanil Malarone | Proguanil is metabolic increased metabolic | e to Proguanil (CYP2C19: R olized to an active metabolite c sm of proguanil to cycloguanil, guanil can be prescribed at sta patient's response. | ycloguanil by CYP2C19. Al there is insufficient data to | o whether such cl | nange has a significant |
| / | Propranolol | Normal Sensitivi | ty to Propranolol (CYP2D6: | Intermediate Metabol | lizer) | ACTIONABL |
| | Inderal | | prescribed at standard label-re avorable response is achieved. | | administration wi | th careful titration and |
| | Quetiapine Seroquel | CYP2D6 are also re- compared to CYP3A effect) is further me CYP3A4, CYP2D6 ar metabolite N-desal genetically guided the clinical respons- reduced to one six itraconazole, indina by 6 fold. Quetiapir treatment (e.g. > 7- | guidance: Quetiapine is predo sponsible for quetiapine is predo sponsible for quetiapine metab A4. N-desalkylquetiapine, a pha etabolized by CYP2D6 and CYP3 nd CYP3A5 enzymes may be re- kylquetiapine. However, the cli drug selection or dosing recom e and tolerability of the individ th of original dose when co-n avir, ritonavir, nefazodone). Whe he dose should be increased up -14 days) of a potent CYP3A4 ir inducer is discontinued, the dos | polism but their role in the armacologically active meta BA4. Preliminary studies has sponsible in variable expos- nical significance of these unendations are available. ual patient. Polypharmacy nedicated with a potent CY en the CYP3A4 inhibitor is to 5 fold of the original d nducer (e.g., phenytoin, car | overall metabolis abolite (responsit ave shown that ge sures to quetiapir changes is not es Quetiapine dose y guidance: Quet (P3A4 inhibitor (e discontinued, the lose when used in rbamazepine, rifa | m of this drug is minor ole of the antidepressant enetic polymorphisms of the and to its active tablished yet and no should be titrated based or claphie dose should be .g., ketoconazole, dose should be increased combination with a chroni- mpin, St. John's wort etc.). |
| / | Rabeprazole Aciphex | • | e to Rabeprazole (CYP2C19 e prescribed at standard dosage | • | | INFORMATIV |
| / | Raltegravir | Normal Respons | e to Raltegravir | | | ACTIONABL |
| _ | Isentress, Dutrebis | metabolizers or pat are not clinically sig UGT1A1. Polyphar | guidance: Raltegravir is elimin tients taking inhibitors of UGT1 gnificant. No dosing adjustment macy guidance: Coadministrat sult in reduced plasma concent | A1 activity have increased ts are required for raltegra tion of raltegravir with drug | plasma levels of wir in patients wh | raltegravir, these changes o carry genetic variants of |
| | Ranolazine | Normal Sensitivi | ty to Ranolazine (CYP2D6: | Intermediate Metaboli | zer) | ACTIONABL |
| - | Ranexa | label-recommended the dose should be | oolized mainly by CYP3A4, and d dosage and administration. T titrated to 500 mg twice daily, imum dose of 1000 mg twice c | he recommended initial do and according to the patie | ose is 375 mg twi | ce daily. After 2–4 weeks, |
| | | | es treatment-related adverse e ng or 375 mg twice daily may b nued. | | | |
| | | | | | | |

| | Manch Univer | ester | PATIENT INFORMATION NAME: Patient 33169 ACC #: 33169 | SPECIMEN DETAILS SPECIMEN TYPE: COLLECTION DATE: | | ORDERED BY |
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| | | SIUY | DOB: 1/1/1900 | RECEIVED DATE: | 1/1/1900 | |
| | FOR ACADEMIC PURPOSES ONLY - NOT | FOR CLINICAL USE | SEX: | REPORT DATE: | 2/8/2018 | |
| | Repaglinide | Normal Sensitivit | y to Repaglinide (SLCO1B1 | Normal Function) | | INFORMATIVE |
| _ | Prandin, Prandimet | | wo copies of SLCO1B1 rs41490 prescribed at label-recommend | | | |
| | Rivaroxaban | Normal Response | e to Rivaroxaban | | | INFORMATIV |
| | Xarelto | (ABCB1) and BCRP (safety profiles of riv strong CYP3A4 inhil concomitant use of phenytoin, rifampin as combined P-gp a increased exposure | ABCG2) transporters. Genetic p aroxaban. Polypharmacy guid pitors (e.g., ketoconazole, itracc rivaroxaban with drugs that are | oolymorphisms of these of lance: Avoid concomitar onazole, lopinavir/ritonav e combined P-gp and str with renal impairment co s (e.g., diltiazem, verapau | genes are not nt use of rivarc vir, ritonavir, in rong CYP3A4 i padministered mil, dronedarc | rivaroxaban with drugs classified one, and erythromycin) have |
| | Rolapitant | Normal Response | e to Rolapitant | | | ACTIONABLE |
| | Varubi | selection or dosing decrease rolapitant moderate CYP2D6 i while others should medication. Rolapit glycoprotein (P-gp) | recommendations are available exposure resulting in a loss of nhibitor and some CYP2D6 sub be closely monitored and their | e. Polypharmacy Guida efficacy. These drugs sho strates (e.g. thioridazine, doing adjusted when co rug efflux transporters: b | nce: Strong C puld be avoide , pimozide) are padministered preast-cancer-r | esistance protein (BCRP) and P- |
| | Rosuvastatin | Normal Myopath | y Risk (SLCO1B1 521T>C T/ | T) | | INFORMATIVE |
| - | Crestor | are present, rosuvas -specific guidelines. | tatin can be prescribed at stan The myopathy risk increases w | dard FDA-recommendec ith use of the 40 mg dos | d starting dose se. (Other myo | tic or circumstantial risk factors as and adjusted based on disease pathy predisposing factors dose, comedications, and female |
| | Rufinamide | Normal Response | e to Rufinamide | | | INFORMATIV |
| - | Banzel | Polypharmacy guid not involved in its m efficacy or toxicity p rufinamide plasma l Patients stabilized c | guidance: No genetically guide dance: Rufinamide is extensive netabolism. Therefore, genetic v rofiles. Coadministration of en evels, while coadministration o n rufinamide should begin valg n valproate should begin rufina | ly metabolized by carbox variations in these metab zyme-inducing antiepiler f valproate increases the proate therapy at a low d | xylesterases. C polizing enzym ptic drugs proc e drug levels ar | ytochrome P450 enzymes are les are not expected to affect its duce modest decreases in nd requires dose adjustment. |
| | Sildenafil | Normal Response | e to Sildenafil | | | INFORMATIV |
| - | Viagra | Pharmacogenetic g CYP3A5*3/*3 genot unknown. Polyphar patients taking str | guidance: Preliminary findings ype compared to those with CN macy guidance: Sildenafil is n ong CYP3A inhibitors, sildena | /P3A5*1/*1 genotype. Th netabolized by CYP3A4 (i a fil exposure is significa | ne clinical sign major route) a antly increase | |

| | 🗸 Mana | ehester | PATIENT INFORMATION | SPECIMEN DETAILS | ORDERED BY |
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| X | | ersity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 |
| | | | | | |
| | Silodosin Rapaflo | metabolites. no ge silodosin is contra | guidance: silodosin is extensive netically guided drug selection c ndicated with potent CYP3A4 in | r dosing recommendation hibitors, as the risk for se | INFORMATIN A4 into pharmacologically inactive ons are available. Polypharmacy guidance: erious adverse events is increased at higher erate inhibitors, as drug levels may increase. |
| ✓ | Simvastatin Zocor | Simvastatin plasma are present, simvas specific guidelines. tolerated this dos | tatin can be prescribed at stand The FDA recommends agains e for 12 months without evide | d to be elevated, and un ard FDA-recommended s : the use of the 80 mg o nce of myopathy. Othe | ACTIONAB less other genetic or circumstantial risk facto starting doses and adjusted based on disease Jaily dose unless the patient had already r myopathy predisposing factors include tatin dose, comedications, and female gende |
| √ | Simvastatin Zocor | The genotype resu | enzyme activity). The patient is | not carry the CYP3A4*2 | INFORMATIN 2 allele (this allele is associated with a ptimal lipid control goal with standard |
| ✓ | Solifenacin Vesicare | Polypharmacy gui concentrations sigr coadministered w at higher concent | guidance: no genetically guide idance: Coadministration of a C nificantly. Therefore, it is recon ith strong CYP3A4 inhibitors, | (P3A4 strong inhibitor in mended not to exceed as the risk for QTc prolo moderate CYP3A4 inhibit | INFORMATIN g recommendations are available. creases solifenacin serum a 5 mg daily dose of solifenacin when ongation induced by this drug is increased tors were not examined, use caution when |
| | | | | | |
| \checkmark | Sufentanil Sufenta | Polypharmacy gui | guidance: No genetically guide | | INFORMATIN ng recommendations are available. nd so should be used with caution when |
| ✓ ✓ | | Pharmacogenetic Polypharmacy gui prescribed with CY Normal Respons Pharmacogenetic including UGT1A3, | guidance: No genetically guide idance: Sufentanil is primarily m P3A4 inhibitors or inducers. e to Sulindac guidance: Sulindac is primarily | etabolized by CYP3A4 ar eliminated by glucuronic of CYP2C9 in sulindac me | ng recommendations are available. |
| √ √ √ | Sufenta Sulindac | Pharmacogenetic Polypharmacy gui prescribed with CY Normal Respons Pharmacogenetic including UGT1A3, guided drug select Normal Respons Pharmacogenetic Polypharmacy gui taking concomitan vardenafil is 10 mg strong inhibitors of studied, other CYP when coadminister | guidance: No genetically guide idance: Sufentanil is primarily m P3A4 inhibitors or inducers. e to Sulindac guidance: Sulindac is primarily UGT1A9 and UGT2B7. The role of ion or dosing recommendations e to Tadalafil guidance: no genetically guide idance: Tadalafil is extensively n t potent inhibitors of CYP3A4, su , not to exceed once every 72 ho CYP3A4, the maximum recomm BA4 moderate inhibitors would l | etabolized by CYP3A4 ar eliminated by glucuronic of CYP2C9 in sulindac me are available. d drug selection or dosin netabolized by CYP3A4. T ch as ketoconazole or rit purs. Tadalafil for Once nended dose is 2.5 mg. A kely increase tadalafil ex 4 inducers. This can be a | INFORMATIN Informations are available. INFORMATIN Information which is catalyzed by several isoforms etabolism is of minor relevance. No genetical INFORMATIN Informations are available. Infadalafil for Use as Needed — For patients conavir, the maximum recommended dose of Daily Use — For patients taking concomitar Ithough specific interactions have not been posure. The exposure of tadalafil is reduced inticipated to decrease the efficacy of tadalafi |

| V | Mancl Univer | sity | | Patient 33169 33169 1/1/1900 | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
|----------|---------------------------------|---|--|--|--|--|--|
| I | FOR ACADEMIC PURPOSES ONLY - NO | OT FOR CLINICAL USE | 52% | | | 2,0,2010 | |
| √ | Tapentadol Nucynta | and genetic variation | led drug s ons in the | selection or dosing re se metabolizing enzy | commendations are availant nes are not expected to a commended dosage and | ffect its effica | |
| √ | Telmisartan Micardis | glucuronide. Telmis | guidance sartan is n | e: Telmisartan is meta ot metabolized by th | e cytochrome P450 isoenz | zymes. Geneti | ACTIONABL nacologically inactive acyl c variability of the cytochrome based dosing adjustments are |
| √ | Terazosin Hytrin | _ | guidance | : no genetically guid | ed drug selection or dosir in metabolizing terazosin | | |
| ✓ | Thiothixene Navane | CYP3A4). No genet likely that strong e | guidance ically guid nzyme inc ed effectiv | e: Thiothixene is meta ded drug selection or lucers may lead to su veness. Consider incre | dosing recommendations ostantial decreases in thic | are available othixene plasn | INFORMATIV 150 enzymes (CYP1A2 and . Polypharmacy guidance: It is na concentrations with the ncomitantly used with strong |
| ✓ | Tiagabine Gabitril | Polypharmacy gui caution when preso | guidance dance: Ti ribed with e drug sho | e: no genetically guid iagabine is extensively h CYP3A4 inhibitors. I ould be considered ca | nducers of CYP3A4 increa | and therefor se tiagabine | INFORMATIV dations are available. e this drug should be used with clearance by 2-fold, and the regimen containing enzyme- |
| ./ | Ticagrelor | Normal Respons | e to Tica | arelor | | | INFORMATIV |
| Y | Brilinta | Pharmacogenetic metabolites, and th P-glycoprotein, end depend on CYP2C1 variants within the profiles. No genetic presence of strong adverse reactions s can significantly de Ticagrelor is a weal | guidance is drug do coded by 9 or CYP3 ABCB1, SI cally-guid CYP3A4 i uch as dy crease tic c inhibitor | Ticagrelor is extens oes not require bioac the ABCB1 gene. Stuc 3A5 metabolizer statu LCO1B1, CYP3A4 and ed drug selection or o nhibitors, significantly spnea or bleeding. Th agrelor exposure (res | ivation to achieve its anti- lies have shown that the e- ses. Moreover, preliminar UGT2B7 genes do not aff losing recommendations r increased exposure to ti- ese drugs should be avoi- ulting in a loss of efficacy) | platelet effect efficacy and sa y studies india ect ticagrelor are available. cagrelor is exp ded with ticag and these do trates of thes | A5 to both active and inactive The drug is also a substrate of afety profile of ticagrelor do not cate that relevant genetic exposure, efficacy or safety Polypharmacy guidance: In bected which may lead to grelor. Strong CYP3A4 inducers ugs should also be avoided. e proteins should be closely |
| ✓ | Tofacitinib Xeljanz | Tofacitinib is metal gene do not signifi | olized pr cantly infl | imarily by CYP3A4 wi | osure. Tofacitinib can be p | | INFORMATIV enetic variations in the CYP2C19 cording to standard label- |
| | Tolbutamide | Normal Sensitivi | tv to Tol | butamide (CYP2CG | : Normal Metabolizer |) | ACTIONABL |
| V | Orinase | | • | | lard label-recommended | | |

| | Manch | octor | PATIENT INFORMATION | SPECIMEN DETAILS | ; | ORDERED BY | |
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| | FOR ACADEMIC PURPOSES ONLY - NOT | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: Image: Comparison of the second | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | | |
| | Tolterodine | | u to Taltaradina (CVD2D6: 1 | ntarmadiata Mataba | lizor) | INFORMATIV | |
| | Detrol | | y to Tolterodine (CYP2D6: 1 | | | | |
| / | Topiramate | Normal Response | e to Topiramate | | | INFORMATIV | |
| | Topamax | Polypharmacy guid is present as metabor elimination when the inducing antiepilept titrated slowly, and o | ic drugs, and may result in redu | topiramate dose appear te metabolism by cytocl py. However, this pathwa iced topiramate plasma dered in presence of ind | rs unchanged i nrome P450 er ay is enhanced concentrations ucers. Concorr | n urine, and an additional 50% nzymes is minor for its by concomitant use of enzyme s. Thus, this drug should be nitant administration of valproic | |
| | Torsemide | Normal Response | e to Torsemide (CYP2C9: No | ormal Metabolizer) | | INFORMATIV | |
| | Demadex | The patient's genoty dosage and adminis | pe predicts a normal exposure tration. | to torsemide and this d | rug can be pre | scribed at label-recommended | |
| / | Trazodone | Normal Response | e to Trazodone | | | INFORMATIV | |
| | Oleptro | Pharmacogenetic guidance: Trazodone is metabolized to its active metabolite m-chlorophenylpiperazine be This metabolite which may contribute to adverse events, is further metabolized by CYP2D6. The impact of guidance polymorphisms of this enzyme on the clinical response to trazodone is not well documented. No genetically selection or dosing recommendations are available. Polypharmacy guidance : It is likely that CYP3A4 inhibit to substantial increases in trazodone plasma concentrations with the potential for adverse effects. If trazodo with a potent CYP3A4 inhibitor, the risk of cardiac arrhythmia may be increased. Therefore coadministration with drugs that are inhibit CYP3A4 should be approached with caution. | | | | 6. The impact of genetic ted. No genetically guided drug hat CYP3A4 inhibitors may lead e effects. If trazodone is used | |
| / | Trifluoperazine | Normal Response | to Trifluoperazine | | | INFORMATIV | |
| | Stelazine | Pharmacogenetic g direct glucuronidation available. Polypharm | uidance: Thrifluoperazine extern on catalyzed by UGT1A4. No ge nacy guidance: It is likely that na concentrations with the pote | netically guided drug se strong enzyme inducers | lection or dosi may lead to s | ing recommendations are | |
| | Trospium | Normal Response | to Trospium | | | INFORMATIV | |
| - | Sanctura | Polypharmacy guid | uidance: no genetically guided lance: CYP enzymes do not con e expected with CYP inhibitors of | ntribute significantly to t | | | |
| | Valbenazine | Normal Sensitivit | y to Valbenazine (CYP2D6: | Intermediate Metabo | olizer) | ACTIONABL | |
| - | Ingrezza | | prescribed at standard label-ren acreased after a week of therap | - | | n. The initial dose is 40 mg onco once daily. | |
| | | coadministered. In p | with comedications: reduce the daily recommended dose to 40 mg if a strong CYP3A4 inhibitor is presence of a CYP2D6 inhibitor, the daily recommended dose may be reduced based on tolerability. with CYP3A4 inducers should be avoided. | | | | |

| | 7) Manak | octor | PATIENT INFORMATION | SPECIMEN DETAILS | ; | ORDERED BY |
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| | FOR ACADEMIC PURPOSES ONLY - NOT | U | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | SPECIMEN TYPE: COLLECTION DATE: RECEIVED DATE: REPORT DATE: | 1/1/1900 1/1/1900 2/8/2018 | |
| | Valarais Asid | Normal Pospons | e to Valproic acid | | | INFORMATIVE |
| V | Valproic Acid Depakote, Depakene | Pharmacogenetic be used to identify contraindicated in | guidance: Genotype results obta patients carrying mutations in m patients known to have mitochon G; e.g., Alpers-Huttenlocher Synd | itochondrial DNA polyr drial disorders caused | nerase γ (POL by mutations | performed in this patient cannot G). Valproic acid is in mitochondrial DNA |
| | | contributions of UC pathway, which inc documenting the ir genetically guided drugs increase valp | ensively metabolized in the liver, 5T1A6, UGT1A9, and UGT2B7. This ludes multiple enzymes such as C npact of genetic polymorphisms drug selection or dosing recomm roic acid clearance 2-fold, and his en added to a therapy regimen co | s drug is also metaboliz (YP2A6, CYP2C9, and C of these metabolizing o endations are available gher doses of this drug | zed by a minc YP2C19. There enzymes on v. e. Polypharm are required | or CYP-dependent oxidation are insufficient studies alproic acid response, and no acy guidance: enzyme-inducing to maintain therapeutic |
| | Valsartan | Normal Sensitivi | ty to Valsartan | | | ACTIONABL |
| | Diovan, Entresto | Pharmacogenetic formation of a mine contribution of CYF | guidance: Valsartan is excreted I or metabolite, valeryl 4-hydroxy v 22C9 in the overall disposition of response to valsartan. No genoty | alsartan, which accoun valsartan, genetic varia | ts for about 9 bility of the C | % of a dose. Given the limited YP2C9 gene is not expected to |
| | Vardenafil | Normal Respons | e to Vardenafil | | | ACTIONABL |
| | Levitra | CYP3A5*3/*3 geno Polypharmacy gui inhibitors such as k patients receiving r should not be exc For itraconazole: 4 24-hour period. For | guidance: Preliminary findings in type compared to those with CYP dance: The dosage of vardenafil etoconazole, itraconazole, ritonav noderate CYP3A4 inhibitors such eeded in a 72-hour period. For 400 mg daily. For clarithromycir or ketoconazole: 200 mg daily. nould not be exceeded in a 24-1 | 3A5*1/*1 genotype. Th may require adjustmer <i>v</i> ir, indinavir, saquinavir as erythromycin. For r indinavir, saquinavir, n: a single dose of 2.5 For itraconazole: 200 | e clinical imp nt in patients i , atazanavir, c itonavir, a sii atazanavir, c mg vardena mg daily. Fo | act of this change is unknown. receiving strong CYP3A4 or clarithromycin, as well as in ngle dose of 2.5 mg vardenafil or ketoconazole: 400 mg daily. fil should not be exceeded in a r erythromycin: a single dose o |
| | Vigabatrin | Normal Respons | e to Vigabatrin | | | INFORMATIV |
| - | Sabril | Polypharmacy gui Therefore, genetic | guidance: no genetically guided dance: Vigabatrin is eliminated p variations in these metabolizing e prescribed at standard label-recor | primarily through renal enzymes are not expect | excretion and ed to affect it | is not metabolized by CYPs. s efficacy or toxicity profiles. |
| | Vilazodone | Normal Respons | e to Vilazodone | | | INFORMATIV |
| 1 | Viibryd | a minor role in the available. Polypha plasma concentrati with a strong inhibi erythromycin), the readjusted to the o to 2-fold when con | guidance: Vilazodone is predom biotransformation of this drug. N rmacy guidance: It is likely that C ons with the potential for adverse itor of CYP3A4 (e.g., ketoconazole dose should be reduced to 20 mg riginal level when the CYP3A4 inh comitantly used with strong CYP3 If CYP3A4 inducers are disconting | o genetically guided di CYP3A4 inhibitors may e effects. Vilazodone sh e). During coadministra g for patients with intol hibitor is discontinued. BA4 inducers (e.g., carb | rug selection lead to substa ould be reduc tion with moc erable advers Consider incre amazepine). T | or dosing recommendations are initial increases in vilazodone ced to 20 mg if co-administered lerate inhibitors of CYP3A4 (e.g., e events. The dose can be easing the dose of vilazodone up 'he maximum daily dose should |



| | Manch | noctor | PATIENT INFORMATION | SPECIMEN DETAILS | ORDERE | D BY |
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| | FOR ACADEMIC PURPOSES ONLY - NOT | sity | NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX: | | 1/1/1900 1/1/1900 2/8/2018 | |
| | Vorapaxar | Normal Respons | se to Voranaxar | | | ACTIONABLE |
| V | Zontivity | Pharmacogenetic polymorphisms of contraindicated in because of the incc CYP3A4 inhibitors increases in vorapa | guidance: vorapaxar is metabol 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 bleed amazepine, phenytoin, rifampin, | affect the efficacy or safe ransient ischemic attack (1 acy guidance: Avoid con lopinavir/ritonavir, ritona ding risk. Avoid concomita | ty profiles of this drug. IIA), or intracranial hem comitant use of vorapa vir, indinavir, and coniv | n CYP2J2. Genetic Vorapaxar is norrhage, (ICH) axar with strong raptan). Significant |
| | Vortioxetine | Normal Sensitiv | ity to Vortioxetine (CYP2D6: | Intermediate Metabol | lizer) | ACTIONABLE |
| - | Trintellix | | e prescribed at standard label-re , which can then be increased to | - | administration. The rec | commended starting |
| | Warfarin | Less than norma | al Sensitivity to Warfarin (CY | P2C9 *1/*1 VKORC1 -16 | 539G>A G/G) | ACTIONABLE |
| _ | Coumadin | FDA-approved lab | a dose increase may be required el: 5-7 mg/day. OR consider usi e to reach steady state is 4-5 day | ng a personalized dose ca | | |
| | Ziprasidone | Normal Respons | se to Ziprasidone | | | INFORMATIVE |
| | Geodon | contributing to the ziprasidone metab reduction involving recommendations adjustments shoul achieved within 1 t improvement for s available, the press compared to sever inhibitors are expe patient's response | guidance: Ziprasidone is primal e oxidative metabolism of ziprasi olic clearance is mediated by cyt g glutathione as well as aldehyde are available. Individualization o d generally occur at intervals of r to 3 days. In order to ensure use everal weeks before upward dos criber should consider the finding ral other antipsychotic drugs. Po I cted to result in modest increase and a dose reduction may be co a chronic treatment of a potent C | done with minor involvem ochrome P450 catalyzed of e oxidase. No genetically of f ziprasidone dose with ca no less than 2 days, as stea of the lowest effective do age adjustment. When de g of ziprasidone's greate lypharmacy guidance: A es in ziprasidone plasma co insidered. Ziprasidone dos | nent from CYP1A2. Less poxidation and approxim guided drug selection o rreful weekly titration is ady-state plasma conce se, patients should ordi ciding among the alter er capacity to prolong Ithough coadministratio oncentrations, a closer se may need to be incre | than one-third of nately two-thirds via or dosing required. Dosage entrations are inarily be observed for native treatments the QT/QTc interval on of strong CYP3A4 monitoring of the eased when used in |
| \checkmark | Zonisamide | Normal Sensitiv | ity to Zonisamide (CYP2C19: | Rapid Metabolizer) | | INFORMATIVE |
| - | Zonegran | CYP2C19 is partly i | nvolved in the metabolism of zo | nisamide, and this drug ca | an be prescribed at star | ndard label- |





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Test Details

| Gene | Genotype | Phenotype | Alleles Tested |
|------------------------------|---------------------------|--|--|
| CYP2C9 | *1/*1 | Normal Metabolizer | *2, *3, *4, *5, *6, *11 |
| CYP2C19 | *1/*17 | Rapid Metabolizer | *2, *3, *4, *4B, *5, *6, *7, *8, *9, *17 |
| CYP2D6 | *4/*17 | Intermediate Metabolizer | *2, *3, *4, *4M, *6, *7, *8, *9, *10, *12, *14A, *14B, *17, *29, *35, *41 |
| CYP3A5 | *1/*1 | Normal Metabolizer | *1D, *2, *3, *3B, *3C, *6, *7, *8, *9 |
| CYP3A4 | *1/*1B | Normal Metabolizer | *1B, *2, *3, *12, *17, *22 |
| VKORC1 | -1639G>A G/G | Low Warfarin Sensitivity | -1639G>A |
| Apolipoprotein E | Indeterminate | Unknown Phenotype | ε2, ε4, (ε3 is reference) |
| CYP2B6 | *1/*1 | Normal Metabolizer | *6, *9 |
| SLCO1B1 | 521T>C T/T | Normal Function | 521T>C, 388A>G |
| COMT | Val158Met A/G | Intermediate COMT Activity | Val158Met |
| OPRM1 | A118G A/A | Normal OPRM1 Function | A118G |
| CYP1A2 | *1L/*1L | Unknown Phenotype | *1C, *1D, *1F, *1K, *1L, *1V, *1W |
| MTHFR | 1298A>C AC 677C>T CC | No Increased Risk of Hyperhomocysteinemia | 1298A>C, 677C>T |
| MTHFR | 677C>T CC | Normal MTHFR Activity | 1298A>C, 677C>T |
| Factor II Factor V Leiden | 20210G>A GG 1691G>A GG | No Increased Risk of Thrombosis | 20210G>A, 1691G>A |

Additional Test Results (added to this original report)

| HLA-B*15:02 | negative/negative | Negative |
|-------------|-------------------|----------|
| HLA-B*57:01 | negative/negative | Negative |
| HLA-B*58:01 | negative/negative | Negative |

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.





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APOE Monograph

Clinical Utility

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 ε3/ε3 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 $\epsilon 2/\epsilon 2$ 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 $\epsilon 2$ homozygous subjects are normolipidemic or even hypolipidemic because of low LDL cholesterol levels. Other non- $\epsilon 2/\epsilon 2$ APOE genotypes ($\epsilon 3/\epsilon 3$, $\epsilon 2/\epsilon 3$ $\epsilon 2/\epsilon 4$ $\epsilon 3/\epsilon 4$ $\epsilon 4/\epsilon 4$) 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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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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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), 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

Clinical Utility

The cytochrome P450 2B6 (CYP2B6) is involved in the metabolism of 4% of clinically important medications. This enzyme is highly polymorphic: to date, 37 different variants have been identified. The CYP2B6 assay identifies some common variants that are associated with variability in enzyme activity.

Assay Interpretation

CYP2B6 enzyme activity defines a normal or an abnormal (intermediate or poor) metabolizer status for a given individual. Several variant alleles have been identified and result in different CYP2B6 isoforms that functionally are fully active, partially active, inactive, or have increased activity. The CYP2B6*1 allele is considered wild-type and encodes a functionally active enzyme (normal). The alleles *6, *7, *9, *11, *16, *18, and *36 encode a decreased activity enzyme. The *22 alleles represents a gain-of-function allele. The functional impact of variants that define the CYP2B6 *4 and *5 alleles is drug dependent.

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

The genotype-phenotype relationship is not well established, and there is a lack of consistency regarding the clinical impact of certain allelic variants. The following provisional genotype-to-phenotype assignment can be used: individuals with two fully active alleles are considered normal (extensive) metabolizers. Individuals with one fully active allele with either a partially active or an inactive allele are considered intermediate metabolizers. Individuals with two partially active alleles or two inactive alleles are considered poor metabolizers. Individuals carrying two increased function alleles or one active allele with a gain-of-function allele are classified as normal/rapid metabolizers.

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 CYP2B*6 polymorphism on the pharmacokinetics and the clinical response have been studied in patients taking methadone, bupropion, and efavirenz. Limited evidence exists regarding the clinical impact of other polymorphisms.

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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SPECIMEN DETAILS

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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 encodes a functionally active enzyme (normal function allele). The alleles *2, *3 *4, *5, *6, and *8 encode an inactive enzyme and are referred to as no function alleles while the *9 and *10 alleles are classified as reduced function alleles. The CYP2C19*17 is an increased function allele.

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.

Clinical Implications

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.

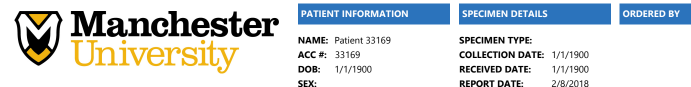
Inhibitors

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: 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, 30 variants 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

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 allele is considered wild-type and encodes a functionally active enzyme (normal). The alleles *2, *3, *4, *5, and *11 encode a partially active enzyme. The allele *6 is a null allele encoding for an inactive enzyme.

The genotype-phenotype relationship is established based on the allele's activity. Individuals with two fully active alleles are considered normal (extensive) metabolizers. Individuals with one fully active allele with either a partially active or an inactive allele are considered intermediate metabolizers. Individuals with two partially active alleles or with two inactive alleles are considered poor metabolizers.

The reference range for CYP2C9 metabolic status is CYP2C9 *1/*1, which is consistent with 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).

Other important drugs metabolized by CYP2C9 include antidiabetics such as tolbutamide, glibenclamide (Micronase), glipizide (Glucotrol), and nateglinide (Starlix).

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: 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 25% 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

CYP2D6 enzyme activity defines a normal or abnormal (intermediate, poor, or ultra-rapid) metabolizer status for a given individual. Commonly tested CYP2D6 variant alleles are classified into functional groups: Full or normal function (e.g., CYP2D6 *1, *2 and *35), reduced function (e.g., CYP2D6*9, *10, *14B, *17, *29, and *41) and no function (e.g., CYP2D6 *3, *4, *5, *6, *7, *8, *11, *12, *14A, *15, *36 and *56). Increased CYP2D6 activity is found in individuals carrying multiple copies of functional alleles. CYP2D6 is subject to gene duplications, and these are denoted "XN", where N represents the number of CYP2D6 gene copies when available.

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 as follows: AS of 0 predicts a poor metabolizer, AS ranging between 1 and 2 predicts a normal (extensive) metabolizer, AS=0.5 predicts an intermediate metabolizer, and AS greater than 2 predicts an ultra-rapid metabolizer. Fully functional alleles are assigned an activity value of 1, reduced function alleles have an activity value of 0.5, while non-functional alleles are assigned an activity value of 0.

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

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), duloxetine (Cymbalta), 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), carvedilol (Coreg), 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), 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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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

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.





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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: carbamazepine (Tegretol), enzalutamide (Xtandi), fosphenytoin (Cerebyx), 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), 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).

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CYP3A5 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

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.





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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: carbamazepine (Tegretol), enzalutamide (Xtandi), fosphenytoin (Cerebyx), 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), 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).

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Factor II Monograph

Clinical Utility

Clotting Factor II, or prothrombin, 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 prothrombin 20210G>A mutation in the Factor II 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 mutations exist for other coagulation factors such 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 Factor II 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 Factor II 20210G>A mutation is associated with a hypercoagulable state. In the United States, the prevalence of this mutation 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 Factor II 20210G>A mutation is Factor II 20210GG.

Clinical Implications

The Factor II 20210G>A mutation 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 Factor II 20210G>A homozygotes is not well defined, but is presumed to be higher than in 20210G>A heterozygotes. Factor II 20210G>A homozygotes for Factor V Leiden and Factor II 20210G>A have an estimated 20-fold increased risk when compared to individuals without either mutation, 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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Factor V Leiden Monograph

Clinical Utility

The Factor V 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 mutation 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 mutations exist in other coagulation factors such as Factor II, or in the presence of nongenetic 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 Factor V Leiden mutation.

Assay Interpretation

Factor V Leiden is the most common known inherited risk factor for thrombosis. Factor V Leiden 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 Factor V Leiden mutation varies by ethnicity, with about 5% of Caucasians, 2% of Hispanics, and 1% of African-Americans having one mutation. Only 1 in 5000 individuals have two Factor V Leiden mutations.

The reference range for Factor V Leiden mutation is Factor V 1691 GG.

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 Factor V Leiden mutation 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 Factor V Leiden 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 Factor V Leiden and the Factor II (compound heterozygote) has an even greater risk of VTE (20-fold) than an individual with a mutation 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 mutations in the MTHFR gene: 677C>T and 1298A> 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 mutations in these conditions is not well established.

Assay Interpretation

The approximate minor allele frequencies for most populations are 30-50% for the MTHFR 1298 A variant and 18-30% for the MTHFR 677 T variant. Heterozygotes and homozygotes for the MTHFR 677C>T mutation have 60% and 30% of normal MTHFR activity, respectively. Heterozygotes and homozygotes for the MTHFR 1298A>C mutation have 80% and 60% of normal MTHFR activity, respectively.

The reference ranges for both mutations of MTHFR are 677CC and 1298AA. 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 677C>T mutation (individual with MTHFR 677 TT 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 677 CT and MTHFR 1298AC genotypes) is not associated with an increase in plasma homocysteine level. Measurement of total plasma homocysteine is informative in this case.
- Homozygosity or heterozygosity for the MTHFR 1298A>C mutation alone (individual with MTHFR 1298AC or MTHFR 1298CC genotypes) does not increase homocysteine levels. Similarly, heterozygosity for the MTHFR 677C>T mutation alone (individuals with MTHFR 677 CT genotype) does not increase homocysteine levels.
- Hyperhomocysteinemia related to MTHFR genetic mutations 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 mutations 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 mutations. Methotrexate intolerance is observed in individuals that are heterozygous or homozygous for the MTHFR 677C>T mutation.





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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 guanine (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 post-surgical 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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 1/1/1900

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 SPECIMEN TYPE:

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 2/8/2018

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

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).

References

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PATIENT INFORMATION

NAME: Patient 33169 ACC #: 33169 DOB: 1/1/1900 SEX:

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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.





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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 University | | REPORT DETAILS | | | | |
|--------------------------|-------------|---|--------------|---|--|--|
| | | Patient: Patient 33169 | VKORC1 | -1639G>A G/G | Low Warfarin Sensitivity | |
| | | DOB: 1/1/1900 ACC #: 33169 | MTHFR | 1298A>C AC 677C>T CC | No Increased Risk of Hyperhomocysteinemia | |
| | Pharmacoger | netic Test Summary | MTHFR | 677C>T CC | Normal MTHFR Activity | |
| CYP2C19 | *1/*17 | Rapid Metabolizer | Factor II | 20210G>A GG | | |
| CYP2C9 | *1/*1 | Normal Metabolizer | Factor V | | No Increased Risk of Thrombosis | |
| CYP2D6 | *4/*17 | Intermediate Metabolizer | Leiden | 1691G>A GG | | |
| CYP3A4 | *1/*1B | Normal Metabolizer | For a comple | ete report contact M | anchester University Master of Science | |
| CYP3A5 *1/*1 | | Normal Metabolizer | | in Pharmacogenomics Program www.manchester.edu/pgx | | |