



Pharmacogenetic Testing

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HMO; PPO; QUEST Integration; Medicare; FEP	Refer to the GTM Utilization Review Matrix

I. Policy Description

Pharmacogenetics aims to study the influence of genetic variation on drug response and drug toxicity, which allows physicians to select a more targeted therapeutic strategy to suit each patient's genetic profile (Aka et al., 2017). Genetic variations in human proteins, such as, cytochrome P450 enzymes, Thiopurine methyltransferase (*TPMT*), dihydropyrimidine dehydrogenase (*DPD*), and cell surface proteins, highlights the clinical importance of pharmacogenetic testing.

Cytochrome (CYP) P450 enzymes are a class of enzymes essential in the synthesis and breakdown metabolism of various molecules and chemicals. Found primarily in the liver, these enzymes are also essential for the metabolism of many medications. CYP P450 enzymes, approximately 58 CYP human genes, are essential to produce many biochemical building blocks, such as cholesterol, fatty acids, and bile acids. Additional cytochrome P450 are involved in the metabolism of drugs, carcinogens, and internal substances, such as toxins formed within cells. Mutations in CYP P450 genes can result in the inability to properly metabolize medications and other substances, leading to increased levels of toxic substances in the body (Bains, 2013; Tantisira & Weiss, 2023).

Thiopurine methyltransferase (*TPMT*) is an enzyme that methylates azathioprine, mercaptopurine and thioguanine into active thioguanine nucleotide metabolites. Azathioprine and mercaptopurine are used for treatment of nonmalignant immunologic disorders; mercaptopurine is used for treatment of lymphoid malignancies; and thioguanine is used for treatment of myeloid leukemias (Relling et al., 2013).

Dihydropyrimidine dehydrogenase (DPD), encoded by the gene *DPYD*, is a rate-limiting enzyme responsible for fluoropyrimidine catabolism. The fluoropyrimidines (5-fluorouracil and capecitabine) are drugs used in the treatment of solid tumors, such as colorectal, breast, and aerodigestive tract tumors (Amstutz et al., 2018).

A variety of cell surface proteins, such as antigen-presenting molecules and other proteins, are encoded by the human leukocyte antigen genes (*HLAs*). HLAs are also known as major histocompatibility complex (MHC) (Viatte, 2023).





II. Indications and/or Limitations of Coverage

Application of coverage criteria is dependent upon an individual's benefit coverage at the time of the request. Specifications pertaining to Medicare and Medicaid can be found in the "Applicable State and Federal Regulations" section of this policy document.

- To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with any of the medications listed below, testing for the CYP2D6 genotype once per lifetime (see Note 1) MEETS COVERAGE CRITERIA:
 - a) Amphetamine
 - b) Aripiprazole
 - c) Aripiprazole Lauroxil
 - d) Atomoxetine
 - e) Brexpiprazole
 - f) Carvedilol
 - g) Cevimeline
 - h) Clozapine
 - i) Codeine
 - j) Desipramine
 - k) Deutetrabenazine
 - I) Eliglustat
 - m) Fluvoxamine
 - n) Gefitinib
 - o) Iloperidone
 - p) Lofexidine
 - q) Meclizine
 - r) Metoclopramide
 - s) Nortriptyline
 - t) Oliceridine
 - u) Ondansetron
 - v) Paroxetine
 - w) Perphenazine
 - x) Pimozide
 - y) Pitolisant
 - z) Propafenone
 - aa) Tamoxifen
 - bb) Tetrabenazine
 - cc) Thioridazine
 - dd) Tolterodine
 - ee) Tramadol





- ff) Tropisetron
- gg)Valbenazine
- hh) Venlafaxine
- ii) Vortioxetine
- 2) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with any of the medications listed below, testing for the *CYP2D6* and *CYP2C19* genotype once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA:**
 - a) Amitriptyline
 - b) Clomipramine
 - c) Doxepin
 - d) Imipramine
 - e) Trimipramine
- 3) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy, testing for the CYP2C19 genotype once per lifetime (see Note 1) MEETS COVERAGE CRITERIA:
 - a) Abrocitinib
 - b) Brivaracetam
 - c) Citalopram
 - d) Clobazam
 - e) Clopidogrel
 - f) Dexlansoprazole (see Note 2)
 - g) Escitalopram
 - h) Flibanserin
 - i) Lansoprazole (see Note 2)
 - j) Mavacamten
 - k) Omeprazole (see Note 2)
 - I) Pantoprazole (in pediatric individuals) (see Note 2)
 - m) Sertraline
 - n) Voriconazole (see Note 2)
- 4) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with any of the medications listed below, testing for the *CYP2C9* genotype once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA**:
 - a) Celecoxib
 - b) Dronabinol
 - c) Erdafitinib
 - d) Flurbiprofen
 - e) Lornoxicam
 - f) Meloxicam
 - g) Nateglinide
 - h) Piroxicam





- i) Siponimod
- j) Tenoxicam
- 5) For individuals being considered for warfarin therapy, testing for the *CYP2C9*, *CYP4F2*, *VKORC1*, and rs12777823 genotype once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA**.
- 6) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with the below medications, testing for the *TPMT* and *NUDT15* genotype once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA**:
 - a) Azathioprine
 - b) Mercaptopurine
 - c) Thioguanine
- 7) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with the below medications, testing for the *DPYD* genotype once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA**:
 - a) Capecitabine
 - b) Flucytosine
 - c) Fluorouracil
 - d) Tegafur
- 8) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with the below medications, testing for the following human leukocyte antigens (HLAs) genotypes once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA**:
 - a) HLA-B*57:01 before treatment with Abacavir
 - b) HLA-B*58:01 before treatment with Allopurinol
 - c) HLA-B*15:02 for treatment with Oxcarbazepine
 - d) HLA-B*15:02 and HLA-A*31:01 for treatment with Carbamazepine
- 9) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with phenytoin/fosphenytoin, testing for the *CYP2C9* and *HLA-B*15:02* genotype once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA**.
- 10) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with the medications listed below, testing for the G6PD genotype once per lifetime (see Note 1) MEETS COVERAGE CRITERIA:
 - a) Pegloticase
 - b) Primaguine
 - c) Rasburicase
 - d) Tafenoquine
- 11) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with the below medications, testing for the following genotypes once per lifetime (see Note 1) MEETS COVERAGE CRITERIA:
 - a) BCHE for treatment with mivacurium or succinylcholine.
 - b) CFTR for treatment with ivacaftor, elexacaftor and tezacaftor, ivacaftor and lumacaftor, or ivacaftor and tezacaftor.
 - c) CYP2B6 for treatment with efavirenz.





- d) CYP3A5 for treatment with tacrolimus.
- IFNL3 treatment with peginterferon alfa-2a, peginterferon alfa-2b or ribavirin.
- f) NAT2 for treatment with amifampridine or amifampridine phosphate.
- g) UGT1A1 for treatment with atazanavir, belinostat, irinotecan, nilotinib, pazopanib, or sacituzumab govitecan-hziy.
- 12) To aid in therapy selection and/or dosing for individuals being considered for therapy or who are in their course of therapy with belzutifan, testing for the *CYP2C19* and *UGT2B17* genotype once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA**.
- 13) For individuals being considered for the use of halogenated volatile anesthetics or depolarizing muscle relaxants, testing for the *RYR1* and *CACNA1S* genotype once per lifetime (see Note 1) **MEETS COVERAGE CRITERIA**.
- 14) When formulary coverage allows a pharmacotherapy that is dependent on a known genetic status (e.g., APOE testing prior to lecanemab-irmb treatment), gene specific testing **MEETS COVERAGE CRITERIA.**
- 15) To identify patients at risk of statin-induced myopathy, genetic testing for the presence of variants in the *SLCO1B1* gene **DOES NOT MEET COVERAGE CRITERIA**.

The following does not meet coverage criteria due to a lack of available published scientific literature confirming that the test(s) is/are required and beneficial for the diagnosis and treatment of an individual's illness.

- 16) The following pharmacogenetic testing **DOES NOT MEET COVERAGE CRITERIA**:
 - a) Genotyping more than once per lifetime (see Note 1) for any medication therapy.
 - b) Genotyping of the general population.
 - c) Pharmacogenetic testing (e.g., single nucleotide polymorphism [SNP] testing or SNP panel testing; single gene or multi-gene panel testing [see Note 3]) for all other situations not addressed above.

NOTES:

Note 1: Any gene may only be tested **once** per lifetime, regardless of the indication (an exception would be for *HLA* where a specific variant is tested for the medication). For example, if *CYP2C19* was tested for therapy with citalopram, additional testing for *CYP2C19* for treatment with clopidogrel is not needed and **DOES NOT MEET COVERAGE CRITERIA**. Testing in a patient post-liver transplant is not indicated.

Note 2: Pharmacogenetic testing for proton pump inhibitor therapies (PPIs) **ONLY MEETS COVERAGE CRITERIA** if the patient has an active *H. pylori* infection.

Note 3: For 2 or more gene tests being run on the same platform, please refer to AHS-R2162-Reimbursement Policy.





III. Table of Terminology

Term	Definition
AACAP	American Academy of Child and Adolescent Psychiatry
AACC	American Association for Clinical Chemistry
AACF	American College of Cardiology Foundation
AAFP	American Family Physician
AAN	American Academy of Neurology
ACMG	American College of Medical Genetics and Genomics
AHA	American Heart Association
AMP	Association For Molecular Pathology
APOE	Apolipoprotein E
ARIA	Amyloid related imaging abnormalities
AS	Activity score
ASCPT	American Society for Clinical Pharmacology and Therapeutics
ASHP	American Society of Health System Pharmacists
ВСНЕ	Butyrylcholinesterase
BDI	Beck's Depression Inventory
CACNA1S	Calcium voltage-gated channel subunit alpha1 S
CFTR	Cystic fibrosis transmembrane conductance regulator
COMT	Catechol-O-methyltransferase
CPIC	Clinical Pharmacogenetics Implementation Consortium
CYP	Cytochrome
CYP1A2	Cytochrome P450 family 1 subfamily A member 2
CYP2B6	Cytochrome P450 family 2 subfamily B member 6
CYP2C9	Cytochrome P450 family 2 subfamily C member 9
CYP2C19	Cytochrome P450 family 2 subfamily C member 19
CYP2D6	Cytochrome P450 family 2 subfamily D member 6
CYP3A4	Cytochrome P450 family 3 subfamily A member 4
CYP3A5	Cytochrome P450 family 3 subfamily A member 5
CYP4F2	Cytochrome P450 family 4 subfamily F member 2
DAT1	Former gene name of solute carrier family 6 member 3
DBH	Dopamine beta-hydroxylase
DPD	Dihydropyrimidine dehydrogenase
DPWG	Dutch Pharmacogenetics Working Group
DPYD	Dihydropyrimidine dehydrogenase gene
DRD1/2/4	Dopamine receptor gene 1/2/4
EMA	European Medicines Agency



FDA	Food and Drug Administration
G6PD	Glucose-6-phosphate dehydrogenase gene
HLA	Human leukocyte antigen
HLA-A	Major histocompatibility complex, class I, A
HLA-B	Major histocompatibility complex, class I, B
IFNL3	Interferon lambda 3
IM	Intermediate metabolizer
ISPG	The International Society of Psychiatric Genetics
MDD	Major depressive disorder
MHC	Major histocompatibility complex
MP	Mercaptopurine
MTHFR	Methylenetetrahydrofolate reductase
NAT2	N-acetyltransferase 2
NM	Normal metabolizer
NUDT15	Nudix hydrolase 15
OPRK1	Opioid receptor kappa 1
OPRM1	Opioid receptor mu 1
PM	Poor metabolizer
PPIs	Proton pump inhibitor therapies
RM	Rapid metabolizer
rs12777823	SNP that can affect warfarin sensitivity
RYR1	Ryanodine receptor 1 gene
SJS	Stevens Johnson Syndrome
SLC6A3	Solute carrier family 6 member 3
SLC6A4	Solute carrier family 6 member 4
SLCO1B1	Solute carrier organic anion transporter family member 1B1
SNP	Single nucleotide polymorphism
TAU	Treatment as usual
TCAs	Tricyclic antidepressants
TEN	Toxic epidermal necrolysis
TG	Thioguanine
TPMT	Thiopurine methyltransferase
TYMS	Thymidylate synthetase
UGT1A1	UDP glucuronosyltransferase family 1 member A1
UGT2B15	Uridine diphosphate glycosyltransferase 2 family, member 15
UGT2B17	UDP glucuronosyltransferase family 2 member B17
URM	Ultra-rapid metabolizer
VKORC1	Vitamin K epoxide reductase complex subunit 1





IV. Scientific Background

Genetic variations play a potentially large role in an individual's response to medications. However, drug metabolism and responses are affected by many other factors, including age, sex, interactions with other drugs, and disease states (Tantisira & Weiss, 2023). Nonetheless, inherited differences in the metabolism and disposition of drugs and genetic polymorphisms in the targets of drug therapy can have a significant influence on the efficacy and toxicity of medications potentially even more so than clinical variables such as age and organ function (Kapur et al., 2014; Ting & Schug, 2016). Genetic variation can influence pharmacodynamic factors through variations affecting drug target receptors and downstream signal transduction, or pharmacokinetic factors, affecting drug metabolism and/or elimination (Tantisira & Weiss, 2023).

The Cytochrome P450 (CYP 450) system is a group of enzymes responsible for the metabolism of many endogenous and exogenous substances, including many pharmaceutical agents. This system may serve to "activate" an inactive form of a drug, as well as inactivate and/or clear a drug from circulation. The CYP 450 enzymes are responsible for the clearance of over half of all drugs, and their activity can be affected by diet, age, and other medications. The genes encoding for the CYP 450 enzymes are highly variable with multiple alleles that confer various levels of metabolic activity for specific substrates. In some cases, alleles can be highly correlated with ethnic background. Generally, there are three categories of metabolizer; ultra-rapid metabolizers, normal metabolizers, and poor metabolizers (Tantisira & Weiss, 2023).

Due to the variations in enzyme activity conferred by allelic differences, some CYP 450 alleles are associated with an increased risk for certain conditions or adverse outcomes with certain drugs. Knowledge of the allele type may assist in the selection of a drug, or in drug dosing. Three CYP 450 enzymes are most often considered regarding clinical use for drug selection and/or dosing. Phenotypes, such as CYP2D6, CYP2C9 and CYP2C19, have been associated with the metabolism of several therapeutic drugs, and various alleles of the *CYP450* gene confer differences in metabolic function. For these CYP 450 enzymes, it is thought that "poor metabolizers" could have less efficient elimination of a drug, and therefore may be at risk for side effects due to drug accumulation. For drugs that require activation by a specific CYP 450 enzyme, lower activity may yield less of the biologically active drug, which could result in lower drug efficacy. Individuals considered as "ultra-rapid metabolizers" may clear the drug more quickly than normal, and therefore may require higher doses to yield the desired therapeutic effect. Likewise, for drugs that require activation, these individuals may produce higher levels of the active drug, potentially causing unwanted side effects. Due to these differences in enzyme activity, some alleles are associated with a higher risk of adverse outcomes depending on the drug prescribed (Tantisira & Weiss, 2023).

ApoE

Apolipoprotein E (APOE) is the gene most strongly associated as a genetic risk factor for late-onset Alzheimer disease. APOE can have three alleles: $\varepsilon 1$, $\varepsilon 2$, and $\varepsilon 4$ (Sherva & Kowall, 2022). APOE $\varepsilon 4$ is a susceptibility gene, meaning it is associated with increased risk but does not cause Alzheimer disease,





and not all patients with Alzheimer disease will carry APOE $\varepsilon 4$. In one study of 1303 patients, 55% of those homozygous for $\varepsilon 4$ developed Alzheimer disease, while 27% of those heterozygous and 9% with no $\varepsilon 4$ allele also developed Alzheimer disease (Myers et al., 1996). APOE, as well as CYP2D6, carrier status may have an effect on a patient's response to drugs, "with CYP2D6-PMs [poor metabolizers], CYP2D6-Ums [ultrarapid metabalizers], and APOE-4/4 carriers acting as the worst responders" (Cacabelos et al., 2012).

In 2023, the FDA approved Lecanemab (brand name Leqembi), an amyloid beta-directed antibody, for the treatment of Alzheimer disease in adult patients (FDA, 2023). One potential side effect of Leqembi is amyloid related imaging abnormalities (ARIA), which may be more likely to occur in people who are homozygous $APOE \, \varepsilon 4$ carries (Leqembi, 2024). The FDA includes that "the prescribing information states that testing for ApoE $\varepsilon 4$ status should be performed before starting treatment with Leqembi to inform the risk of developing ARIA" (FDA, 2023).

CYP2C9

Warfarin (brand name Coumadin) is widely used as an anticoagulant in the treatment and prevention of thrombotic disorders. *CYP2C9* participates in warfarin metabolism, and several *CYP2C9* alleles have reduced activity, resulting in a higher circulating drug concentration. *CYP2C9*2* and *CYP2C9*3* are the most common variants with reduced activity. Variations in a second gene, *VKORC1*, also can impact warfarin's effectiveness. This gene codes for the enzyme that is the target for warfarin. Genotypes resulting in reduced metabolism may need a higher dose to achieve the desired efficacy (Tantisira & Weiss, 2023).

CYP2C19

Clopidogrel (brand name Plavix) is used to inhibit platelet aggregation and is given as a pro-drug that is metabolized to its active form by *CYP2C19*. Alleles *CYP2C19*2* and *CYP2C19*3* are associated with reduced metabolism of clopidogrel. Individuals with the "poor metabolizer" alleles may not benefit from clopidogrel treatment at standard doses (Tantry et al., 2021). Tuteja et al. (2020) studied *CYP2C19* genotyping to guide antiplatelet therapy. A total of 504 participants contributed to this study, with only 249 participants genotyped. The authors noted that genotyping results "significantly influenced antiplatelet drug prescribing; however, almost half of *CYP2C19* LOF [loss-of-function] carriers continued to receive clopidogrel. Interventional cardiologists consider both clinical and genetic factors when selecting antiplatelet therapy following PCI [percutaneous coronary intervention]" (Tuteja et al., 2020).

CYP2D6

Tetrabenazine (brand name Xenazine) is used in the treatment of chorea associated with Huntington disease. This drug is metabolized for clearance primarily by *CYP2D6*. Poor metabolizers are considered to be those individuals with impaired *CYP2D6* function, and dosing is often influenced by how well a patient metabolizes the drug. For example, a poor metabolizer will often have a maximum dose of 50 mg daily whereas an extensive metabolizer has a maximum dose of 100 mg daily (Suchowersky, 2023).





Tamoxifen, a drug commonly used for the treatment and prevention of recurrence of estrogen receptor positive breast cancer, is metabolized by *CYP2D6*. Polymorphisms of *CYP2D6* have been noted to affect the efficacy of tamoxifen by affecting the amount of active metabolite produced. Endoxifen, which is the primary active metabolite of tamoxifen, has a 100-fold affinity for the estrogen receptor compared to tamoxifen, but poor metabolizers have been demonstrated to show lower than expected levels of plasma endoxifen (Ahern et al., 2017).

Codeine, which is commonly used to treat mild to moderate pain, is metabolized to morphine, a much more powerful opioid, by *CYP2D6*. Individuals with varying CYP2D6 activity may see negative side effects or a shorter duration of pain relief. The effect is significant enough to have caused fatalities in unusual metabolizers; for instance, an ultra-rapid metabolizing toddler was reported to have passed away after being given codeine for a routine dental operation (Kelly et al., 2012; Tantisira & Weiss, 2023).

TPMT

Thiopurine methyltransferase (TPMT) is an enzyme that methylates thiopurines into active thioguanine nucleotides. The *TPMT* gene is inherited as a monogenic co-dominant trait with ethnic differences in the frequencies of low-activity variant alleles. Individuals who inherit two inactive *TPMT* alleles will develop severe myelosuppression. Individuals that inherit only one inactive *TPMT* allele will develop moderate to severe myelosuppression, and those individuals who inherit both active *TPMT* alleles will have a lower risk of myelosuppression. Therefore, genotyping for *TPMT* is critical before starting therapy with thiopurine drugs (Relling et al., 2013).

DPYD

The dihydropyrimidine dehydrogenase (*DPYD*) gene encodes for the rate-limiting enzyme dihydropyrimidine dehydrogenase, which is involved in catabolism of fluoropyrimidine drugs used in the treatment of solid tumors. Decreased DPD activity increases the risk for severe or even fatal drug toxicity when patients are being treated with fluoropyrimidine drugs. Numerous genetic variants in the *DPYD* gene have been identified that alter the protein sequence or mRNA splicing; however, some of these variants have no effect on DPD enzyme activity. The most studied causal variant of *DPYD* haplotype (HapB3) spans intron 5 to exon 11 and affects protein function. The most common variant in Europeans is HapB3 with a c.1129–5923C>G *DPYD* variant which demonstrates decreased function with carrier frequency of 4.7%, followed by c.190511G>A (carrier frequency: 1.6%) and c.2846A>T (carrier frequency: 0.7%). Approximately 7% of Europeans carry at least one decreased function *DPYD* variant. In people with African ancestry, the most common variant is c.557A>G (rs115232898, p.Y186C) and is relatively common (3–5% carrier frequency). Other *DPYD* decreased function variants are rare. Therefore, most available genetic tests focus on identifying the most common variants with well-established risk: (c.190511G>A, c.1679T>G, c.2846A>T, c.1129–5923C>G) (Amstutz et al., 2018).

TYMS

TYMS (thymidylate synthetase) encodes an enzyme necessary for thymidine production. As with *DPYD*, *TYMS* is thought to be involved with the toxicity of fluoropyrimidines. Fluorouracil (FU)'s primary metabolite inhibits thymidylate synthetase by forming a stable complex with thymidylate synthetase





and folate, thereby blocking activity of the enzyme. Polymorphisms in the *TYMS* gene further affect the interaction between *TYMS* and FU, potentially increasing the toxicity of FU. Genotyping of *TYMS* prior to treatment with FU or capecitabine has been suggested for clinical practice, but data has been varied (Krishnamurthi & Kamath, 2024).

Castro-Rojas et al. (2017) evaluated *TYMS* genotypes as predictors of both clinical response and toxicity to fluoropyrimidine-based treatment for colorectal cancer. A total of 105 patients were genotyped. The authors noted that while the 2R/2R genotype was associated with clinical response (odds ratio = 3.45), the genotype was also associated with severe toxicity (odds ratio = 5.21). The genotype was thought to be associated with low *TYMS* expression. The authors further identified the rs2853542 and rs151264360 alleles to be independent predictors of response failure to chemotherapy (Castro-Rojas et al., 2017).

HLAs

Human Leukocyte antigens (HLAs) are divided into three regions, such as class I, class II and class III. Each class has many gene loci, expressed genes and pseudogenes. The class I encodes HLA-A, HLA-B, HLA-C and other antigens. The class II encodes HLA-DP, DQ and DR. The class III region is located between class I and class II and does not encode any HLAs, but other immune response proteins (Viatte, 2023).

An article published by van der Wouden et al. (2019) reports on the development of the new PGx-Passport panel (pre-emptive pharmacogenetics-passport panel), which is able to test "58 germline variant alleles, located within 14 genes (*CYP2B6, CYP2C9, CYP2C19, CYP2D6, CYP3A5, DPYD, F5, HLA-A, HLA-B, NUDT15, SLCO1B1, TPMT, UGT1A1*, and *VKORC1*)"; this standardized panel is based on the Dutch Pharmacogenetics Working Group (DPWG) guidelines and will help physicians to optimize drug prescription in 49 common drugs. It is recommended by the authors that commercial and hospital laboratories utilize this panel for personalized medicinal purposes. Drug optimization in the 49 commonly prescribed drugs includes ten antidepressants, five immunosuppressants, five anti-cancer drugs, four anti-infectives, four anticoagulants, four antiepileptics, four antipsychotics, three proton pump inhibitors, two anti-arrhythmics, two analgesics, two antilipidemics, one antihypertensive, one psychostimulant, one anticontraceptive and one Gaucher disease drug (van der Wouden et al., 2019).

Proprietary Testing

Due to the increase in pharmacogenetic genotyping, proprietary gene panels have become commercially available. Panels encompassing the most common genes that influence drug metabolism have increased in usage. For example, Myriad's new proprietary panel "GeneSight" proposes it can "predict poorer antidepressant outcomes and to help guide healthcare providers to more genetically optimal medications," thereby leading to better patient outcomes. The test assesses every known metabolic pathway (CYP450 or otherwise) for a given drug and their metabolites, as well as the pharmacodynamic activity of the compound and its metabolites, any FDA information on that drug, and other validated research on the relevant alleles; this information is then integrated with the genetic test results. This allows the test to categorize the 64 medications into three categories: "green (use as directed), yellow (some moderate gene-drug interaction) and red (significant gene-drug interaction)." Myriad states that this allows every metabolic pathway of a drug to be evaluated instead of the "one gene, one drug" view.







Other GeneSight variations, such as GeneSight Psychotropic used for psychotropic medications, exist as well (Myriad, 2016, 2019, 2022). Still, other companies such as Mayo Clinic and Sema4 have developed their own pharmacogenetic panels, each with individually chosen analytes (Mayo, 2023; Sema4, 2022).

Benitez et al. (2018) assessed the cost-effectiveness of pharmacogenomics in treating psychiatric disorders. The authors compared 205 members that received guidance from GeneSight's Psychotropic to 478 members that received "treatment-as-usual" (TAU). Reimbursement costs were calculated over the 12 months pre- and post-index event periods. The authors found a total post-index cost savings of \$5505, which was equivalent to a savings of \$0.07 per-member-per-month (PMPM). The authors also evaluated the savings at different adoption rates of the GeneSight test. At 5% adoption, commercial payer savings was calculated at \$0.02 PMPM and at 40% adoption, savings was \$0.15 PMPM (Benitez et al., 2018).

The AmpliChip® (Roche Molecular Systems, Inc.) is the FDA-cleared test for CYP450 genotyping. This test genotypes *CYP2D6* and *CYP2C19*. From the FDA website: "The AmpliChip CYP4502C19 Test is designed to identify specific nucleic acid sequences and query for the presence of certain known sequence polymorphisms through analysis of the pattern of hybridization to a series of probes that are specifically complementary either to wild-type or mutant sequences" (FDA, 2005). The analytical accuracy was evaluated at 99.6%, or 806 of 809 samples identified correctly. This test assesses a total of 30 alleles, three for CYP219 and 27 for *CYP2D6* (FDA, 2005).

The OneOme RightMed Pharmacogenomic Test analyzes more than 100 variants in 27 genes to study how a patient may respond to certain medications. The test covers *CYP1A2*, *CYP2B6*, *CYP2C Cluster*, *CYP2C9*, *CYP2C19*, *CYP2D6*, *CYP3A4*, *CYP3A5*, *CYP4F2*, *COMT*, *DPYD*, *DRD2*, *F2*, *F5*, *GRIK4*, *HLA-A*, *HLA-B*, *HTR2A*, *HTR2C*, *IL28B* (*IFNL4*), *MTHFR**, *NUDT15*, *OPRM1*, *SLC6A4*, *SLC01B1*, *TPMT*, *UGT1A1*, and *VKORC1* (OneOme, 2021). Analytical validity of the test was assessed by comparing RightMed test results with bi-directional Sanger sequencing results, which resulted in 100% concordance. The RightMed test detects *CYP2D6* deletions, duplications, and hybrid alleles, but cannot differentiate duplications in the presence of a deletion (GTR, 2017).

Clinical Utility and Validity

A study evaluating GeneSight Psychotropic's clinical utility was performed by Greden et al. (2019); a total of 1167 patients with major depressive disorder were split into two randomized groups: treatment as usual (TAU) and pharmacogenetic-guided. Medications were classified as "congruent" (use as directed' or 'use with caution' test categories) or "incongruent" ('use with increased caution and with more frequent monitoring' test category) with test results. After eight weeks, the authors found a statistically significant improvement in response and remission; 26% for the pharmacogenetic arm compared 19.9% for TAU and 15.3% for remission compared to 10.1% for TAU (Greden et al., 2019). The authors concluded that pharmacogenetic testing did not improve results, but significantly improved response and remission rates for "difficult-to-treat depression patients over standard of care" (Greden et al., 2019).

Kekic et al. (2020) studied genetic variants that commonly affect supportive care medications, which include, antidepressants, antiemetics, and analgesics, used in oncology practice. A total of 196 cancer patients were genotyped using a multi-gene panel, OneOme RightMed. The panel assessed 27 genes,





including CYP2C9, CYP2C19, CYP2D6, CYP3A4, COMT, OPRM1, GRIK4, HTR2A, SLC6A4, associated with pain medications, antidepressants, and antiemetics. Of the 196 patients, 19.9% had prostate cancer, 17.9% had colorectal cancer, 14.8% had melanoma, and 47.4% had other cancer types. All 196 patients had at least one actionable polymorphism related to these supportive care medications, specifically, in CYP2C19 and CYP2D6. Specifically, 67.3% of the patients had other than normal CYP2D6 metabolizer phenotype and 57.1% had other than normal CYP2C19 metabolizer phenotype. Based on the results, 37 patients were recommended an alternative analgesic, nine were recommended an alternative anti-depressant (Kekic et al., 2020).

Plumpton et al. (2019) evaluated the cost-effectiveness of panel tests with various pharmacogenes. The constructed multigene panel included *HLA-A*31:01*, *HLA-B*15:02*, *HLA-B*57:01*, *HLA-B*58:01*, *HLA-B* (158T), and *HLA-DQB1* (126Q), which are involved with various treatments (abacavir, carbamazepine, et al). The constructed multigene panel was found to provide a cost savings of \$491 if all findings for all alleles were acted on, regardless of an allele's individual cost-effectiveness. Testing for patients eligible for abacavir (*HLA-B*57:01*) and clozapine (*HLA-B* (158T) and *HLA-DQB1* (126Q)) was found to be cost-effective. However, testing for patients eligible for allopurinol (*HLA-B*58:01*) was not found to be cost-effective. Furthermore, testing for *HLA-A*31:01* for carbamazepine was found to be cost-effective, but not testing for *HLA-B*15:02* (Plumpton et al., 2019).

Braten et al. (2020) researched the impact of *CYP2C19* genotyping on the antidepressant drug sertraline, which is metabolized by the polymorphic *CYP2C19* enzyme. A total of 1202 patients participated and submitted 2190 sertraline serum samples. All patients were categorized based on *CYP2C19* genotype-predicted phenotype subgroups; these groups include normal (NM), ultra-rapid (UM), intermediate (IM), and poor metabolizer (PM). Serum samples showed that *CYP2C19* IM and PM patients had significantly higher sertraline concentrations compared to NMs; "Based on the relative differences in serum concentrations compared to NMs, dose reductions of 60% and 25% should be considered in PMs and IMs, respectively, to reduce the risk of sertraline overexposure in these patients" (Braten et al., 2020).

Roscizewski et al. (2021) conducted a retrospective observational study to determine what effect pharmacogenomic testing had on "treatment decisions in patients with depressive symptoms in an interprofessional primary care setting." From April 2019 to March 2021, they identified 78 patients who underwent pharmacogenomic testing for psychotropic medications. They found that 53.8% of patients "experienced a change to their antidepressant regiment after [pharmacogenomic] testing," with the most cited change being addition of another antidepressant, followed by switching the antidepressant, then increased dose. This demonstrated how pharmacogenomic testing could be useful in informing clinical decision making at the beginning of treatment or "in those who experience an inadequate response to their prescribed regimen" and ensuring optimal patient recovery.

Stevenson et al. (2021) aimed to assess the potential impact of multigene pharmacogenomic testing among those hospitalized with COVID-19 in the United States. Through a cross-sectional analysis with electronic health records, researchers "characterized medication orders, focusing on medications with actionable guidance related to 14 commonly assayed genes (CYP2C19, CYP2C9, CYP2D6, CYP3A5, DPYD, G6PD, HLA-A, HLA-B, IFNL3, NUDT15, SLCO1B1, TPMT, UGT1A1, and VKORC1)." From their cohort, they found that 64 unique medications with pharmacogenomic guidance were ordered at least once, and about 89.7% of patients "had at least one order for a medication with PGx guidance and... (23.1%) had





orders for 4 or more actionable medications." Through a simulation analysis, they estimated that "17 treatment modifications per 100 patients would be enabled if [pharmacogenomic] results were available," and that the genes *CYP2D6* and *CYP2C19* were responsible for most of the treatment modifications. Medications most affected included ondansetron, oxycodone, and clopidogrel. With additional investigations that support these findings, pharmacogenomic testing would better inform the curation of individualized treatment plans for patients suffering from severe COVID-19.

Galli et al. (2021) studied the use of guided selection of antiplatelet therapy for patients undergoing percutaneous coronary intervention. The authors conducted a meta-analysis that included 3656 relevant articles with 20743 patients. Overall, "guided selection of antiplatelet therapy was associated with a reduction in major adverse cardiovascular events and reduced bleeding, although not statistically significant." Additionally, cardiovascular death, myocardial infraction, stent thrombosis, and minor bleeding were all reduced with guided therapy compared to standard therapy, but the risks of all-cause death and major bleeding did not differ. The authors concluded that "guided selection of antiplatelet therapy improved both composite and individual efficacy outcomes with a favourable safety profile, driven by a reduction in minor bleeding, supporting the use of platelet function or genetic testing to optimise the choice of agent in patients undergoing PCI" (Galli et al., 2021).

Oslin et al. (2022) conducted a randomized clinical trial that compared treatment guided by pharmacogenomic testing vs. usual care "to determine whether pharmacogenomic testing affects antidepressant medication selection and whether such testing leads to better clinical outcomes". Participants of this clinical trial included 676 clinicians and 1944 patients. Criteria for patient enrollment were those with major depressive disorder who were initiating or switching treatment with a single antidepressant and exclusion included those who have active substance use disorder, mania, psychosis, or concurrent treatment with a specified list of medications. Results of this study determined "remission rates over 24 weeks were higher among patients whose care was guided by pharmacogenomic testing than those in usual care (OR, 1.28 [95% CI, 1.05 to 1.57]; P = .02; risk difference, 2.8% [95% CI, 0.6% to 5.1%]) but were not significantly higher at week 24 when 130 patients in the pharmacogenomic-guided group and 126 patients in the usual care group were in remission (estimated risk difference, 1.5% [95%CI, -2.4% to 5.3%]; P = .45)". In conclusion, in provision of pharmacogenomic testing for drug-gene interaction amongst patients with major depressive disorder, pharmacogenomic testing "reduced prescription of medications with predicted drug-gene interactions compared to usual care. Provision test results had small nonpersistent effects on symptom remission" (Oslin et al., 2022).

Ghanbarian et al. (2023) studied the cost-effectiveness of pharmacogenetic testing used to guide prescription of antidepressants. The authors looked at data from patients with major depressive disorder in British Columbia, Canada. The data included unique patient characteristics, including metabolizer phenotypes, incremental costs, life-years, and quality-adjusted life-years. "Pharmacogenomic-guided care was associated with 37% fewer patients with refractory depression over 20 years." The costs of pharmacogenetic testing were estimated to be offset within about two years of use, with an overall saving of 956 million Canadian dollars (4926 Canadian dollars per patient) (Ghanbarian et al., 2023).





The 2023 PREPARE (preemptive pharmacogenomic testing for preventing adverse drug reactions) trial investigated the effects of pre-emptive genotyping using a pharmacogenetic panel on adverse drug reactions. Swen et al. (2023) conducted an "open-label, multicentre, controlled, cluster-randomised, crossover implementation study of a 12-gene pharmacogenetic panel in 18 hospitals, nine community health centres, and 28 community pharmacies in seven European countries." A total of 6944 patients receiving their first prescription for a clinically recommended drug were included in the study. The participants were divided into a study group, which received genotyping and recommended treatment adjustments, and a control group, which received standard care. The primary outcome measured was the occurrence of clinically relevant adverse drug reactions within 12-weeks. A clinically relevant adverse drug reactions occurred in 21.5% of patients in the study group (N=2923), and 28.6% of patients in the control group (N=3270). The authors concluded that "genotype-guided treatment using a 12-gene pharmacogenetic panel significantly reduced the incidence of clinically relevant adverse drug reactions and was feasible across diverse European health-care system organisations and settings" (Swen et al., 2023)

V. Guidelines and Recommendations

Clinical Pharmacogenetics Implementation Consortium (CPIC)

CPIC guidelines provide guidance to physicians on how to use genetic testing to help them to optimize drug therapy. The guidelines and projects were endorsed by several professional societies including The Association for Molecular Pathology (AMP), The American Society for Clinical Pharmacology and Therapeutics (ASCPT) and The American Society of Health-System Pharmacists (ASHP) (CPIC, 2023b).

In their list of guidelines, CPIC provides specific therapeutic recommendations for drugs metabolized by Cytochrome P450 enzymes and other important metabolic enzymes.

CYP2C9 Genotypes

Drug	CYP2C9/	Summary of CPIC Therapeutic	Level of	Reference
	Phenotype	Recommendations	Recommendatio	
			ns	
Phenytoin/	HLA-B*15:02	If patient is phenytoin-naïve, do not	Strong	(Caudle et
fosphenytoi	Positive- Normal	use phenytoin/fosphenytoin. Avoid		al., 2014;
n based on	Metabolizer (NM),	carbamazepine and oxcarbazepine.		Karnes et
HLA-	Intermediate	If the patient has previously used		al., 2021)
B*15:02	Metabolizer (EM),	phenytoin continuously for longer		
	and Poor	than three months without		
	Metabolism (M)	incidence of cutaneous adverse		
		reactions, cautiously consider use		
		of phenytoin in the future.		





	HLA-B*15:02 Negative-Normal Metabolizer (NM)	No adjustments needed from typical dosing strategies. Subsequent doses should be adjusted according to therapeutic drug monitoring, response, and side effects. An HLA-B*15:02 negative test does not eliminate the risk of phenytoin-induced SJS/TEN, and patients should be carefully monitored according to standard practice.	Strong	
	HLA-B*15:02 Negative- Intermediate Metabolizer (IM)	No adjustments needed from typical dosing strategies. Subsequent doses should be adjusted according to therapeutic drug monitoring, response and side effects. An HLA-B*15:02 negative test does not eliminate the risk of phenytoin-induced SJS/TEN, and patients should be carefully monitored according to standard practice. For first dose, use typical initial or loading dose. For subsequent doses, use approximately 25% less than typical maintenance dose. Subsequent doses should be adjusted according to therapeutic drug monitoring, response and side effects.	Moderate	
	HLA-B*15:02 Negative-Poor Metabolizer (PM)	For first dose, use typical initial or loading dose. For subsequent doses use approximately 50% less than typical maintenance dose. Subsequent doses should be adjusted according to therapeutic drug monitoring, response, and side effects. An HLA-B*15:02 negative test does not eliminate the risk of phenytoin-induced SJS/TEN, and patients should be carefully monitored according to standard practice.	Strong	
Warfarin	Various phenotypes	Genotype-guided warfarin dosing is very complex and involves a	Multiple	(Johnson et al., 2017)





		combination of <i>CYP2C9</i> , <i>VKORC1</i> , <i>CYP4F2</i> and rs12777823 as well as an algorithm including ancestry information.		
Celecoxib, flurbiprofen, ibuprofen, lornoxicam	NM	"In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals"	Strong	(Theken et al., 2020)
	IM (Activity Score [AS] = 1.5)	"Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals."	Moderate	
	IM (AS = 1)	"Initiate therapy with lowest recommended starting dose. Titrate dose upward to clinical effect or maximum recommended dose with caution. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals. Carefully monitor adverse events, such as blood pressure and kidney function during course of therapy."	Moderate	
	PM	"Initiate therapy with 25–50% of the lowest recommended starting dose. Titrate dose upward to clinical effect or 25–50% of the maximum recommended dose with caution. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals. Upward dose titration should not occur until after steady-state is reached (at	Moderate	





		least 8 days for celecoxib and 5 days for ibuprofen, flurbiprofen, and lornoxicam after first dose in PMs). Carefully monitor adverse events such as blood pressure and kidney function during course of therapy. Alternatively, consider an alternate therapy not metabolized by CYP2C9 or not significantly impacted by CYP2C9 genetic variants in vivo"		
Meloxicam	NM	"Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals"	Strong	(Theken et al., 2020)
	IM, AS 1.5	See NM	Moderate	
	IM, AS 1	"Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals. Upward dose titration should not occur until after steady-state is reached (at least 7 days). Carefully monitor adverse events, such as blood pressure and kidney function during course of therapy. Alternatively, consider alternative therapy not metabolized by CYP2C9 or not significantly impacted by CYP2C9 genetic variants in vivo or choose an NSAID metabolized by CYP2C9 but with a shorter half-life"	Moderate	
	PM	"Choose an alternative therapy not metabolized by CYP2C9 or not	Moderate	





	significantly impacted by CYP2C9		
	•		
	with a shorter half-life"		
NM	"Initiate therapy with	Strong	(Theken et
	recommended starting dose. In		al., 2020)
	accordance with the prescribing		
	information, use the lowest		
	effective dosage for shortest		
	duration consistent with individual		
	patient treatment goals."		
IM AS 1.5	"Initiate therapy with	Moderate	
	recommended starting dose. In		
	accordance with the prescribing		
	information, use the lowest		
	effective dosage for shortest		
	duration consistent with individual		
	patient treatment goals."		
IM AS 1	"Choose an alternative therapy not	Moderate	
	metabolized by CYP2C9 or not	(Optional for	
	significantly impacted by CYP2C9	Tenoxicam)	
	genetic variants in vivo or choose an	·	
	NSAID metabolized by CYP2C9 but		
	with a shorter half-life"		
PM	"Choose an alternative therapy not	Moderate	
		(Optional for	
	significantly impacted by CYP2C9	' '	
	genetic variants in vivo or choose an	,	
	•		
	with a shorter half-life"		
	IM AS 1.5	genetic variants in vivo or choose an NSAID metabolized by CYP2C9 but with a shorter half-life" NM "Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals." IM AS 1.5 "Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals." IM AS 1 "Choose an alternative therapy not metabolized by CYP2C9 or not significantly impacted by CYP2C9 genetic variants in vivo or choose an NSAID metabolized by CYP2C9 but with a shorter half-life" PM "Choose an alternative therapy not metabolized by CYP2C9 or not significantly impacted by CYP2C9 genetic variants in vivo or choose an NSAID metabolized by CYP2C9 genetic variants in vivo or choose an NSAID metabolized by CYP2C9 genetic variants in vivo or choose an NSAID metabolized by CYP2C9 but	genetic variants in vivo or choose an NSAID metabolized by CYP2C9 but with a shorter half-life" NM "Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals." IM AS 1.5 "Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals." IM AS 1 "Choose an alternative therapy not metabolized by CYP2C9 or not significantly impacted by CYP2C9 but with a shorter half-life" PM "Choose an alternative therapy not metabolized by CYP2C9 or not significantly impacted by CYP2C9 genetic variants in vivo or choose an NSAID metabolized by CYP2C9 but with a shorter half-life" PM "Choose an alternative therapy not metabolized by CYP2C9 or not significantly impacted by CYP2C9 genetic variants in vivo or choose an NSAID metabolized by CYP2C9 but with a shorter half-life" PM "Choose an alternative therapy not metabolized by CYP2C9 or not significantly impacted by CYP2C9 genetic variants in vivo or choose an NSAID metabolized by CYP2C9 but with a shorter half-life" PM "Choose an alternative therapy not metabolized by CYP2C9 but with a shorter half-life" PM "Choose an alternative therapy not metabolized by CYP2C9 but with a shorter half-life" PM "Choose an alternative therapy not metabolized by CYP2C9 but with a shorter half-life" PM "Choose an alternative therapy not metabolized by CYP2C9 but with a shorter half-life"

CYP2D6 Genotype

Drug	CYP2D6 Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio ns	Reference
Amitriptylin	Ultra-rapid	Avoid tricyclic use due to potential	Strong	(Hicks et
e and	Metabolizer (URM)	lack of efficacy. Consider alternative	(recommendatio	al., 2016)
Nortripyline		drug not metabolized by CYP2D6. If	n for other TCAs	





Drug	CYP2D6	Summary of CPIC Therapeutic	Level of	Reference
	Phenotype	Recommendations	Recommendatio	
			ns	
Other TCAs (tricyclic antidepress ants):		a TCA is warranted, consider titrating to a higher target dose (compared to normal metabolizers). Utilize therapeutic drug monitoring to guide dose adjustments.	is Optional)	
clomipramin	Normal	Initiate therapy with recommended	Strong	
e, desipramine , doxepin,	Metabolizer (NM)	starting dose.	(recommendatio n for other TCAs is Strong)	
imipramine, and trimipramin e	IM	Consider a 25% reduction of recommended starting dose. Utilize therapeutic drug monitoring to guide dose adjustments.	Moderate (recommendatio n for other TCAs is Optional)	
	PM	Avoid tricyclic use due to potential for side effects. Consider alternative drug not metabolized by <i>CYP2D6</i> . If a TCA is warranted, consider a 50% reduction of recommended starting dose. Utilize therapeutic drug monitoring to guide dose adjustments.	Strong (recommendatio n for other TCAs is Optional)	
Codeine	URM	Avoid codeine use due to potential for toxicity.	Strong	(Crews et al., 2021)
	EM	Use label-recommended age or weight-specific dosing.	Strong	
	IM	Use label-recommended age or weight-specific dosing. If no response, consider alternative analgesics such as morphine or a nonopioid.	Moderate	
	PM	Avoid codeine use because of possibility of diminished analgesia	Strong	
Paroxetine	URM	Select alternative drug not predominantly metabolized by CYP2D6	Moderate	(Bousman et al., 2023)





Drug	CYP2D6 Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio ns	Reference
	М	Initiate therapy with recommended starting dose.	Strong	
	IM	Consider a lower starting dose and slower titration schedule as compared with normal metabolizers	Optional	
	PM	Consider a 50% reduction in recommended starting dose, slower titration schedule, and a 50% lower maintenance dose as compared with normal metabolizers.	Moderate	
Fluvoxamin e	URM	No recommendation due to lack of evidence.	No recommendation	(Bousman et al., 2023)
	EM	Initiate therapy with recommended starting dose.	Strong	
	IM	Initiate therapy with recommended starting dose.	Moderate	
	PM	Consider a 25–50% lower starting dose and slower titration schedule as compared with normal metabolizers or consider a clinically appropriate alternative antidepressant not predominantly metabolized by CYP2D6	Optional	
Ondansetro n and Tropisetron	URM	Select alternative drug not predominantly metabolized by <i>CYP2D6</i> (i.e., granisetron).	Moderate	(Bell et al., 2016)
	NM	Initiate therapy with recommended starting dose.	Strong	
	IM	Insufficient evidence demonstrating clinical impact based on <i>CYP2D6</i> genotype. Initiate therapy with recommended starting dose.	No recommendation	
	PM	Insufficient evidence demonstrating clinical impact based on CYP2D6	No recommendation	





Drug	CYP2D6	Summary of CPIC Therapeutic	Level of	Reference
	Phenotype	Recommendations	Recommendatio ns	
		genotype. Initiate therapy with recommended starting dose.		
Tamoxifen	URM	Avoid moderate and strong <i>CYP2D6</i> inhibitors. Initiate therapy with recommended standard of care dosing (tamoxifen 20 mg/day).	Strong	(Goetz et al., 2018)
	NM	Avoid moderate and strong <i>CYP2D6</i> inhibitors. Initiate therapy with recommended standard of care dosing (tamoxifen 20 mg/day).	Strong	
	NM/IM	Consider hormonal therapy such as an aromatase inhibitor for postmenopausal [individuals] or aromatase inhibitor along with ovarian function suppression in premenopausal [individuals], given that these approaches are superior to tamoxifen regardless of <i>CYP2D6</i> genotype. If aromatase inhibitor use is contraindicated, consideration should be given to use a higher but FDA approved tamoxifen dose (40 mg/day).45 Avoid <i>CYP2D6</i> strong to weak inhibitors.	Optional (Controversy remains)	
	IM	Consider hormonal therapy such as an aromatase inhibitor for postmenopausal [individuals] or aromatase inhibitor along with ovarian function suppression in premenopausal [individuals], given that these approaches are superior to tamoxifen regardless of <i>CYP2D6</i> genotype. If aromatase inhibitor use is contraindicated, consideration should be given to use a higher but FDA approved tamoxifen dose (40	Moderate	





Drug	CYP2D6	Summary of CPIC Therapeutic	Level of	Reference
	Phenotype	Recommendations	Recommendatio	
			ns	
		mg/day). Avoid CYP2D6 strong to		
		weak inhibitors.		
	PM	Recommend alternative hormonal	Strong	
		therapy such as an aromatase		
		inhibitor for postmenopausal		
		[individuals] or aromatase inhibitor		
		along with ovarian function		
		suppression in premenopausal		
		[individuals] given that these approaches are superior to		
		tamoxifen regardless of <i>CYP2D6</i>		
		genotype and based on knowledge		
		that <i>CYP2D6</i> poor metabolizers		
		switched from tamoxifen to		
		anastrozole do not have an		
		increased risk of recurrence. Note,		
		higher dose tamoxifen (40 mg/day)		
		increases but does not normalize		
		endoxifen concentrations and can		
		be considered if there are		
		contraindications to aromatase		
		inhibitor therapy.		
Atomoxetin	URM	Initiate with a dose of 0.5	Moderate	(Brown et
e (for		mg/kg/day and increase to 1.2		al., 2019)
children)		mg/kg/day after 3 days. If no clinical		
		response and in the absence of adverse events after 2 weeks,		
		consider obtaining a peak plasma		
		concentration (1–2 hours after dose		
		administered). If < 200 ng/mL,		
		consider a proportional increase in		
		dose to approach 400 ng/mL		
	NM	Initiate with a dose of 0.5 mg/kg and	Moderate	
		increase to 1.2 mg/kg/day after 3		
		days. If no clinical response and in		
		the absence of adverse events after		
		2 weeks, consider obtaining a peak		





Drug	CYP2D6 Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio	Reference
		plasma concentration (1–2 hours after dose administered). If < 200 ng/mL, consider a proportional increase in dose to approach 400 ng/mL.	ns	
	IM	Initiate with a dose of 0.5 mg/kg/day and if no clinical response and in the absence of adverse events after 2 weeks, consider obtaining a plasma concentration 2–4 hours after dosing. If response is inadequate and concentration is < 200 ng/mL, consider a proportional dose increase to achieve a concentration to approach 400 ng/mL.b,c If unacceptable side effects are present at any time, consider a reduction in dose	Moderate	
	PM	Initiate with a dose of 0.5 mg/kg/day and if no clinical response and in the absence of adverse events after 2 weeks, consider obtaining a plasma concentration 4 hours after dosing. If response is inadequate and concentration is < 200 ng/mL, consider a proportional dose increase to achieve a concentration to approach 400 ng/mL.b,c If unacceptable side effects are present at any time, consider a reduction in dose	Strong	
Atomoxetin e (for adults)	URM	Initiate with a dose of 40 mg/day and increase to 80 mg/ day after 3 days. If no clinical response and in the absence of adverse events after	Moderate	





Drug	CYP2D6	Summary of CPIC Therapeutic	Level of	Reference
	Phenotype	Recommendations	Recommendatio ns	
		2 weeks, consider increasing dose to	115	
		100 mg/day. If no clinical response		
		observed after 2 weeks, consider		
		obtaining a peak plasma		
		concentration (1–2 hours after dose		
		administered). If < 200 ng/mL,		
		consider a proportional increase in		
		dose to approach 400 ng/mL.b,c		
		Dosages > 100 mg/day may be		
		needed to achieve target		
		concentrations.		
	NM	Initiate with a dose of 40 mg/day	Moderate	1
		and increase to 80 mg/ day after 3		
		days. If no clinical response and in		
		the absence of adverse events after		
		2 weeks, consider increasing dose to		
		100 mg/day. If no clinical response		
		observed after 2 weeks, consider		
		obtaining a peak plasma		
		concentration (1–2 hours after dose		
		administered). If < 200 ng/mL,		
		consider a proportional increase in		
		dose to approach 400 ng/mL.b,c		
		Dosages > 100 mg/day may be		
		needed to achieve target		
		concentrations.		
	IM	Initiate with a dose of 40 mg/day	Moderate	
		and if no clinical response and in the		
		absence of adverse events after 2		
		weeks increase dose to 80 mg/day.		
		If response is inadequate after 2		
		weeks consider obtaining a plasma		
		concentration 2–4 hours after		
		dosing. If concentration is < 200		
		ng/mL, consider a proportional dose		
		increase to achieve a concentration		
		to approach 400 ng/mL.b,c If		





Drug	CYP2D6 Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio	Reference
			ns	
		unacceptable side effects are		
		present at any time, consider a reduction in dose.		
	PM	Initiate with a dose of 40 mg/day and if no clinical response and in the	Moderate	
		absence of adverse events after 2 weeks increase dose to 80 mg/day.		
		If response is inadequate after 2		
		weeks, consider obtaining a plasma concentration 2–4 hours after		
		dosing. If concentration is < 200 ng/mL, consider a proportional dose		
		increase to achieve a concentration to approach 400 ng/mL.b,c If		
		unacceptable side effects are		
		present at any time, consider a reduction in dose		

CYP2B6 Genotypes

Drug	CYP2B6	Summary of CPIC Therapeutic	Level of	Reference
	Phenotype	Recommendations	Recommendations	
Efavirenz (for children >40	URM	Initiate efavirenz with standard dosing (600 mg/day)	Strong	(Desta et al., 2019)
kg and adults)	Rapid Metabolizer (RM)	Initiate efavirenz with standard dosing (600 mg/day)	Strong	
	NM	Initiate efavirenz with standard dosing (600 mg/day)	Strong	
	IM	Consider initiating efavirenz with decreased dose of 400 mg/day	Moderate	
	PM	Consider initiating efavirenz with decreased dose of 400 or 200 mg/day.	Moderate	





CYP2C19 Genotype

Drug	Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio ns for amitriptyline	Reference
Amitriptyline and Nortripyline Other TCAs: clomipramin e, doxepin, imipramine, and	URM, RM	Avoid tertiary amine use due to potential for sub-optimal response. Consider alternative drug not metabolized by <i>CYP2C19</i> . TCAs without major <i>CYP2C19</i> metabolism include the secondary amines nortriptyline and desipramine. If a tertiary amine is warranted, utilize therapeutic drug monitoring to guide dose adjustments.	Optional (Recommendatio n for other TCAs is Optional)	(Hicks et al., 2016)
trimipramine	NM	Initiate therapy with recommended starting dose.	Strong (Recommendatio n for other TCAs is Strong)	
	IM	Initiate therapy with recommended starting dose.	Strong (Recommendatio n for other TCAs is Optional)	
	PM	Avoid tertiary amine use due to potential for sub-optimal response. Consider alternative drug not metabolized by <i>CYP2C19</i> . TCAs without major <i>CYP2C19</i> metabolism include the secondary amines nortriptyline and desipramine. For tertiary amines, consider a 50% reduction of the recommended starting dose. Utilize therapeutic drug monitoring to guide dose adjustments.	Moderate (Recommendatio n for other TCAs is Optional)	
	URM	Consider a clinically appropriate alternative antidepressant not predominantly metabolized by	Strong	





Drug	Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendations for amitriptyline	Reference
Citalopram and Escitalopram		CYP2C19. If citalopram or escitalopram are clinically appropriate, and adequate efficacy is not achieved at standard maintenance dosing, consider titrating to a higher maintenance dose		(Bousman et al., 2023)
	EM	Initiate therapy with recommended starting dose.	Strong	
	IM	Initiate therapy with recommended starting dose. Consider a slower titration schedule and lower maintenance dose than normal metabolizers	Moderate	
	PM	Consider a clinically appropriate antidepressant not predominantly metabolized by CYP2C19. If citalopram or escitalopram are clinically appropriate, consider a lower starting dose, slower titration schedule, and 50% reduction of the standard maintenance dose as compared with normal metabolizers	Strong	
Sertraline	URM	Initiate therapy with recommended starting dose.	Strong	(Bousman et al.,
	EM	Initiate therapy with recommended starting dose.	Strong	2023)
	IM	Initiate therapy with recommended starting dose. Consider a slower titration schedule and lower maintenance dose than CYP2C19 normal metabolizers	Moderate	





Drug	Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio ns for amitriptyline	Reference
	PM	Consider a lower starting dose, slower titration schedule, and 50% reduction of standard maintenance dose as compared with CYP2C19 normal metabolizers or select a clinically appropriate alternative antidepressant not predominantly metabolized by CYP2C19	Moderate	
Clopidogrel	URM, RM, NM	If considering clopidogrel, use at standard dose (75 mg/day)	Strong	(Lee et al., 2022)
	IM, Likely IM	Avoid standard dose clopidogrel (75mg) if possible. Use prasugrel or ticagrelor at standard dose if no contraindication.	Strong	
	PM, Likely PM	Avoid clopidogrel if possible. Use prasugrel or ticagrelor at standard dose if no contraindication.	Strong	
Voriconazole	URM	Choose an alternative agent that is not dependent on <i>CYP2C19</i> metabolism as primary therapy in lieu of voriconazole. Such agents include isavuconazole, liposomal amphotericin B, and posaconazole.	Moderate	(Moriyam a et al., 2017)
	RM	Choose an alternative agent that is not dependent on <i>CYP2C19</i> metabolism as primary therapy in lieu of voriconazole. Such agents include isavuconazole, liposomal amphotericin B, and posaconazole.	Moderate	
	NM	Initiate therapy with recommended starting dose.	Strong	
	IM	Initiate therapy with recommended starting dose.	Moderate	





Drug	Phenotype	Summary of CPIC Therapeutic	Level of	Reference
		Recommendations	Recommendations for	
			ns for amitriptyline	
	PM	Choose an alternative agent that is not dependent on <i>CYP2C19</i> metabolism as primary therapy in lieu of voriconazole. Such agents include isavuconazole, liposomal amphotericin B, and posaconazole. In the event that voriconazole is	Moderate	
		considered to be the most appropriate agent, based on clinical advice, for a patient with poor metabolizer genotype, voriconazole should be administered at a preferably lower than standard dosage with careful therapeutic drug monitoring.		
Proton Pump Inhibitors (omeprazole,	URM	"Increase starting daily dose by 100%. Daily dose may be given in divided doses. Monitor for efficacy"	Optional	(Lima et al., 2020)
lansoprazole, and pantoprazole) All recommenda	RM	"Initiate standard starting daily dose. Consider increasing dose by 50– 100% for the treatment of Helicobacter pylori infection and erosive esophagitis. Daily dose may be given in divided doses. Monitor for efficacy"	Moderate	
tions here are "Optional" for dexlansopraz ole	Normal	"Initiate standard starting daily dose. Consider increasing dose by 50– 100% for the treatment of H. pylori infection and erosive esophagitis. Daily dose may be given in divided doses. Monitor for efficacy"	Moderate	
	Likely IM/IM	"Initiate standard starting daily dose. For chronic therapy (> 12 weeks) and efficacy achieved,	Optional	





Drug	Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio ns for amitriptyline	Reference
		consider 50% reduction in daily dose and monitor for continued efficacy"		
	Likely PM/PM	"Initiate standard starting daily dose. For chronic therapy (> 12 weeks) and efficacy achieved, consider 50% reduction in daily dose and monitor for continued efficacy"	Moderate	

CYP2D6 and CYP2C19 Genotypes (Caudle et al., 2020; Hicks et al., 2016) for Amitriptyline, Clomipramine, Doxepin, Imipramine, and Trimipramine

Phenotype	CYP2D6	CYP2D6	CYP2D6	CYP2D6
CYP2C19	UM	NM	IM	PM
URM	Avoid amitriptyline use Recommendatio n: Optional	Consider alternative drug not metabolized by <i>CYP2C19</i> . Recommendation: Optional	Consider alternative drug not metabolized by <i>CYP2C19</i> . Recommendation: Optional	Avoid amitriptyline use Recommendation: Optional
NM	Avoid amitriptyline use. If amitriptyline is warranted, consider titrating to a higher target dose (compared to normal metabolizers) Recommendatio n: Strong	Initiate therapy with recommended starting dose. Recommendation: Strong	Consider a 25% reduction of recommended starting dose. Recommendation: Moderate	Avoid amitriptyline use. If Amitriptyline is warranted, consider a 50% reduction of recommended starting dose. Recommendation: Strong
IM	Avoid amitriptyline use	Initiate therapy with recommended starting dose.	Consider a 25% reduction of recommended	Avoid amitriptyline use. If Amitriptyline is warranted, consider a
		starting dose.	recommended	warranteu, consider a





	Recommendatio n: Optional	Recommendation: Strong	starting dose. Recommendation:	50% reduction of recommended
			Optional This recommendation may also be considered for diplotypes with an activity score of 1.	starting dose. Recommendation: Optional
PM	Avoid amitriptyline use Recommendatio n: Optional	Avoid amitriptyline use. If Amitriptyline is warranted, consider a 50% reduction of recommended starting dose. Recommendation: Moderate	Avoid amitriptyline use Recommendation: Optional	Avoid amitriptyline use Recommendation: Optional

TPMT Genotype

Drug	TPMT	Summary of CPIC Therapeutic	Level of	Reference
	Phenotyp	Recommendations	Recommendatio	
	е		ns	
Mercaptopurin	NM	Start with normal starting dose (e.g., 75	Strong	(Relling et
e (MP)		mg/m ² /d or 1.5 mg/kg/d) and adjust		al., 2018)
		doses of MP (and of any other		
		myelosuppressive therapy) without any		
		special emphasis on MP compared to		
		other agents. Allow 2 weeks to reach		
		steady state after each dose		
		adjustment. Consider evaluating TPMT		
		erythrocyte activity to assess TPMT		
		phenotype. If thiopurines are required		
		and either TPMT or NUDT15 status is		
		unknown, monitor closely for toxicity.		
	IM	Start with reduced doses (start at 30-	Strong	
		70% of full dose: e.g., at 50 mg/m²/d or		
		0.75 mg/kg/d) and adjust doses of MP		
		based on degree of myelosuppression		





			,
	and disease-specific guidelines. Allow 2–4 weeks to reach steady state after each dose adjustment. In those who require a dosage reduction based on myelosuppression, the median dose may be ~40% lower (44 mg/m²) than that tolerated in wild-type patients (75 mg/m²). In setting of myelosuppression, and depending on other therapy, emphasis should be on reducing MP over other agents. Consider evaluating TPMT erythrocyte activity to assess TPMT phenotype. If thiopurines are required and either TPMT or NUDT15 status is unknown, monitor closely for toxicity.		
PM	For malignancy, start with drastically reduced doses (reduce daily dose by 10-fold and reduce frequency to thrice weekly instead of daily, e.g., 10 mg/m²/d given just 3 days/week) and adjust doses of MP based on degree of myelosuppression and disease-specific guidelines. Allow 4–6 weeks to reach steady state after each dose adjustment. In setting of myelosuppression, emphasis should be on reducing MP over other agents. For nonmalignant conditions, consider alternative nonthiopurine immunosuppressant therapy. Consider evaluating TPMT erythrocyte activity to assess TPMT phenotype. If thiopurines are required and either TPMT or NUDT15 status is unknown, monitor	Strong	
NM	Start with normal starting dose (e.g., 2–3 mg/kg/d) and adjust doses of azathioprine based on disease-specific	Strong	(Relling et al., 2018)
		2–4 weeks to reach steady state after each dose adjustment. In those who require a dosage reduction based on myelosuppression, the median dose may be ~40% lower (44 mg/m²) than that tolerated in wild-type patients (75 mg/m²). In setting of myelosuppression, and depending on other therapy, emphasis should be on reducing MP over other agents. Consider evaluating TPMT erythrocyte activity to assess TPMT phenotype. If thiopurines are required and either TPMT or NUDT15 status is unknown, monitor closely for toxicity. PM For malignancy, start with drastically reduced doses (reduce daily dose by 10-fold and reduce frequency to thrice weekly instead of daily, e.g., 10 mg/m²/d given just 3 days/week) and adjust doses of MP based on degree of myelosuppression and disease-specific guidelines. Allow 4–6 weeks to reach steady state after each dose adjustment. In setting of myelosuppression, emphasis should be on reducing MP over other agents. For nonmalignant conditions, consider alternative nonthiopurine immunosuppressant therapy. Consider evaluating TPMT erythrocyte activity to assess TPMT phenotype. If thiopurines are required and either TPMT or NUDT15 status is unknown, monitor closely for toxicity. NM Start with normal starting dose (e.g., 2–3 mg/kg/d) and adjust doses of	2–4 weeks to reach steady state after each dose adjustment. In those who require a dosage reduction based on myelosuppression, the median dose may be ~40% lower (44 mg/m²) than that tolerated in wild-type patients (75 mg/m²). In setting of myelosuppression, and depending on other therapy, emphasis should be on reducing MP over other agents. Consider evaluating TPMT erythrocyte activity to assess TPMT phenotype. If thiopurines are required and either TPMT or NUDT15 status is unknown, monitor closely for toxicity. PM For malignancy, start with drastically reduced doses (reduce daily dose by 10-fold and reduce frequency to thrice weekly instead of daily, e.g., 10 mg/m²/d given just 3 days/week) and adjust doses of MP based on degree of myelosuppression and disease-specific guidelines. Allow 4–6 weeks to reach steady state after each dose adjustment. In setting of myelosuppression, emphasis should be on reducing MP over other agents. For nonmalignant conditions, consider alternative nonthiopurine immunosuppressant therapy. Consider evaluating TPMT erythrocyte activity to assess TPMT phenotype. If thiopurines are required and either TPMT or NUDT15 status is unknown, monitor closely for toxicity. NM Start with normal starting dose (e.g., 2– 3 mg/kg/d) and adjust doses of azathioprine based on disease-specific





		steady state after each dose adjustment. Consider evaluating erythrocyte TPMT activity to assess TPMT phenotype. If thiopurines are required and TPMT status is unknown, monitor closely for toxicity.		
	IM	Start with reduced starting doses (30%-80% of normal dose) if normal starting dose is 2-3 mg/kg/day, (e.g. 0.6 – 2.4 mg/kg/day), and adjust doses of azathioprine based on degree of myelosuppression and disease-specific guidelines. Allow 2-4 weeks to reach steady-state after each dose adjustment. Consider evaluating erythrocyte TPMT activity to assess TPMT phenotype. If thiopurines are required and TPMT status is unknown, monitor closely for toxicity.	Strong	
	PM	For non-malignant conditions, consider alternative-nonthiopurine immunosuppressant therapy or malignancy, start with drastically reduced doses (reduce daily dose by 10-fold and dose thrice weekly instead of daily) and adjust doses of azathioprine based on degree of myelosuppression and disease specific guidelines. Allow 4-6 weeks to reach steady state after each dose adjustment. Consider evaluating erythrocyte TPMT activity to assess TPMT phenotype. If thiopurines are required and TPMT status is unknown, monitor closely for toxicity.	Strong	
Thioguanine	NM	Start with normal starting dose (e.g. 40-60 mg/m2 /day). Adjust doses of thioguanine (TG) and of other myelosuppressive therapy without any special emphasis on TG. Allow 2 weeks to reach steady state after each dose adjustment. Consider evaluating erythrocyte TPMT activity to assess TPMT phenotype. If thiopurines are	Strong	(Relling et al., 2013; Relling et al., 2018)





	required and TPMT status is unknown,	
	monitor closely for toxicity.	
IM	Start with reduced doses (50% to 80% of	Moderate
	normal dose) if normal starting dose is	
	≥40-60 mg/m2 /day (e.g. 20-48 mg/m2	
	/day) and adjust doses of TG based on	
	degree of myelosuppression and	
	disease-specific guidelines. Allow 2–4	
	weeks to reach steady state after each	
	dose adjustment. In setting of	
	myelosuppression, and depending on	
	other therapy, emphasis should be on	
	reducing TG over other agents. Consider	
	evaluating erythrocyte TPMT activity to	
	assess TPMT phenotype. If thiopurines	
	are required and TPMT status is	
	unknown, monitor closely for toxicity.	
55.4		C 1
PM	Start with drastically reduced	Strong
	doses (reduce daily dose by 10-fold and	
	dose thrice weekly instead of daily) and	
	adjust doses of TG based on degree of	
	myelosuppression and disease-specific	
	guidelines. Allow 4–6 weeks to reach	
	steady state after each dose	
	adjustment. In setting of	
	myelosuppression, emphasis should be	
	on reducing TG over other agents. For	
	nonmalignant conditions, consider	
	alternative nonthiopurine	
	immunosuppressant therapy. Consider	
	evaluating erythrocyte TPMT activity to	
	assess TPMT phenotype. If thiopurines	
	- , , ,	





NUDT15 Genotype

Drug	NUDT15	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio	Reference
	Phenotyp e	Recommendations	ns	
Mercaptopurin e	NM	Start with normal starting dose (e.g., 75 mg/m²/d or 1.5 mg/kg/d) and adjust doses of MP (and of any other myelosuppressive therapy) without any special emphasis on MP compared to other agents. Allow 2 weeks to reach steady state after each dose adjustment.	Strong	(Relling et al., 2018)
	IM	Start with reduced doses (start at 30–80% of normal dose: if normal starting dose is ≥75 mg/m2 /day or ≥ 1.5 mg/kg/day (e.g. start at 25-60 mg/m2 /day or 0.45-1.2 mg/kg/day) and adjust doses of MP based on degree of myelosuppression and disease-specific guidelines. Allow 2–4 weeks to reach steady state after each dose adjustment. If myelosuppression occurs, and depending on other therapy, emphasis should be on reducing mercaptopurine over other agents. If normal starting dose is already < 1.5mg/kg/day, dose reduction may not be recommended.	Strong	
	PM	For malignancy, initiate dose at 10 mg/m2 /day and adjust dose based on myelosuppression and disease specific guidelines. Allow 4-6 weeks to reach steady state after each dose adjustment. If myelosuppression occurs, emphasis should be on reducing mercaptopurine over other agents. For nonmalignant conditions, consider alternative nonthiopurine immunosuppressant therapy.	Strong	





Drug	NUDT15	Summary of CPIC Therapeutic	Level of	Reference
	Phenotyp	Recommendations	Recommendatio	
	е		ns	
Azathioprine	NM	Start with normal starting dose (e.g., 2–3 mg/kg/day) and adjust doses of azathioprine based on disease-specific guidelines. Allow 2 weeks to reach steady state after each dose adjustment.	Strong	(Relling et al., 2018)
	IM	Start with reduced doses (start at 30–80% of normal dose: if normal starting dose is 2-3 mg/kg/day, (e.g. 0.6 – 2.4 mg/kg/day) and adjust doses of MP based on degree of myelosuppression and disease-specific guidelines. Allow 2–4 weeks to reach steady state after each dose adjustment.	Strong	
	PM	For nonmalignant conditions, consider alternative nonthiopurine immunosuppressant therapy. For malignant conditions, start with drastically reduced normal daily doses (reduce daily dose by 10-fold) and adjust doses of azathioprine based on degree of myelosuppression and disease specific guidelines. Allow 4-6 weeks to reach steady-state after each dose adjustment.	Strong	
Thioguanine	NM	Start with normal starting dose (40- 60 mg/day). Adjust doses of thioguanine and of other myelosuppressive therapy without any special emphasis on thioguanine. Allow 2 weeks to reach steady-state after each dose adjustment	Strong	(Relling et al., 2018)
	IM	Start with reduced doses (50% to 80% of normal dose) if normal starting dose is ≥40-60 mg/m2 /day (e.g. 20-48 mg/m2 /day) and adjust doses of thioguanine based on degree of myelosuppression	Moderate	





Drug	NUDT15 Phenotyp e	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio ns	Reference
		and disease specific guidelines. Allow 2-4 weeks to reach steady-state after each dose adjustment. If myelosuppression occurs, and depending on other therapy, emphasis should be on reducing thioguanine over other agents.		
	PM	Reduce doses to 25% of normal dose and adjust doses of thioguanine based on degree of myelosuppression and disease specific guidelines. Allow 4-6 weeks to reach steady-state after each dose adjustment. In setting of myelosuppression, emphasis should be on reducing thioguanine over other agents. For non-malignant conditions, consider alternative nonthiopurine immunosuppressant therapy.	Strong	

DPYD Genotypes

Drug	Phenotype	Summary of CPIC Therapeutic	Level of	Reference
		Recommendations	Recommendations	
5-	NM	Based on genotype, there is no	Strong	(Amstutz et
Fluorouracil		indication to change dose or therapy.		al., 2018)
Capecitabine		Use label recommended dosage and		
		administration.		
	IM	Reduce starting dose based on activity	Activity score 1:	
		score followed by titration of dose	Strong Activity score	
		based on toxicity or therapeutic drug	1.5: Moderate	
		monitoring (if available). Activity score		
		1: Reduce dose by 50% Activity score		
		1.5: Reduce dose by 25% to 50%		
	PM	Activity score 0.5: Avoid use of 5-	Strong	
		fluorouracil or 5-fluorouracil prodrug-		
		based regimens. In the event, based on		
		clinical advice, alternative agents are		





not considered a suitable therapeutic	
option, 5-fluorouracil should be	
administered at a strongly reduced	
dosed with early therapeutic drug	
monitoring. Activity score 0: Avoid use	
of 5-fluorouracil or 5-fluorouracil	
prodrug-based regimens.	

HLA-B Genotypes

Drug	Phenotype	Summary of CPIC Therapeutic	Level of	Reference
		Recommendations	Recommendatio	
			ns	
Abacavir	Noncarrier of	Low or reduced risk of abacavir	Strong	(Martin et al.,
	HLA-B*57:01	hypersensitivity		2014)
	Carrier of	Abacavir is not recommended	Strong	
	HLA-B*57:01			
Allopurinol	Noncarrier of	Use allopurinol per standard	Strong	(Hershfield et
	HLA-B*5801	dosing guidelines		al., 2013; Saito
	(*X/*X)			et al., 2016)
	Carrier of	Allopurinol is contraindicated	Strong	
	HLA-B*5801			
	(HLA-			
	<i>B*5801/</i> *X,b			
	HLA-			
	B*5801/HLA-			
	B*5801)			
Oxcarbazepin	HLA-B*15:02	Use oxcarbazepine per standard	Strong	(Phillips et al.,
е	negative	dosing guidelines		2018)
	HLA-B*15:02	If patient is oxcarbazepine naıve,	Strong	
	positive	do not use oxcarbazepine.		
Carbamazepin	HLA-B*15:02	Use carbamazepine per standard	Strong	(Phillips et al.,
е	negative and	dosing guidelines.		2018)
	HLA-A*31:01			
	negative			
	HLA-B*15:02	If patient is carbamazepine-naive	Strong	
	negative and	and alternative agents are		





Drug	Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommendatio ns	Reference
	HLA-A*31:01 positive	available, do not use carbamazepine.		
	HLA-B*15:02 positive and any HLA- A*31:01 genotype (or HLA-A*31:01 genotype unknown)	If patient is carbamazepine-naive, do not use carbamazepine.	Strong	

Additional Genotypes

Drug/Genotype	Phenotype	Summary of CPIC Therapeutic	Level of	Reference
		Recommendations	Recommend	
			ations	
UGT1A1 for Atazanavir	EM	There is no need to avoid prescribing of atazanavir based on <i>UGT1A1</i> genetic test result. Inform the patient that some patients stop atazanavir because of jaundice (yellow eyes and skin), but that this patient's genotype makes this unlikely (less than about a 1 in 20 chance of stopping atazanavir because of jaundice).	Strong	(Gammal et al., 2016)
	IM	There is no need to avoid prescribing of atazanavir based on <i>UGT1A1</i> genetic test result. Inform the patient that some patients stop atazanavir because of jaundice (yellow eyes and skin), but that this patient's genotype makes this unlikely (less than about a 1 in 20 chance of stopping atazanavir because of jaundice).	Strong	





Drug/Genotype	Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommend ations	Reference
	PM	Consider an alternative agent particularly where jaundice would be of concern to the patient. If atazanavir is to be prescribed, there is a high likelihood of developing jaundice that will result in atazanavir discontinuation (at least 20% and as high as 60%).	Strong	
UGT1A1 for Irinotecan	N/A	N/A	A, 1A level of evidence	(CPIC, 2023a)
CFTR for Ivacaftor	Homozygous or heterozygous G551D-CFTR— e.g. G551D/ F508del, G551D/G551D, rs75527207 genotype AA or AG	Use ivacaftor according to the product label (e.g., 150 mg every 12h for patients aged 6 years and older without other diseases; modify dose in patients with hepatic impairment)	Strong	(Clancy et al., 2014)
	Noncarrier of G551D-CFTR— e.g. F508del/R553X , rs75527207 genotype GG	Ivacaftor is not recommended	Moderate	
	Homozygous for F508del- CFTR (F508del/F508 del), rs113993960, or rs199826652 genotype del/ del	Ivacaftor is not recommended	Moderate	
G6PD for high- risk drugs	Normal	No reason to avoid high-risk drugs based on G6PD status	Strong	(Gammal et al., 2023)





Drug/Genotype Phenotype Summary of CPIC Therapeutic Level Reference Recommendations Recommend ations (rasburicase and Avoid use of high-risk drugs Deficient Strong pegloticase) deficient with **CNSHA** Variable To ascertain G6PD status, enzyme Moderate activity must be measured. Drug use should be guided per the recommendations based on the activity-based phenotype SLCO1B1 for SLCO1B1 Prescribe an alternative statin Strong (Cooper-DeHoff Simvastatin decreased depending on the desired potency. al., 2022) function or simvastatin therapy SLCO1B1 warranted, limit dose to <20 possible mg/day decreased function SLCO1B1 poor Prescribe an alternative statin Strong function depending on the desired potency CYP3A5 for EM Increase starting dose 1.5–2 times Strong (Birdwell al., et recommended starting dose. Total treatment with 2015) **Tacrolimus** starting dose should not exceed 0.3 mg/kg/day. Use therapeutic drug monitoring to guide dose adjustments. IM Increase starting dose 1.5–2 times Strong recommended starting dose. Total starting dose should not exceed 0.3 mg/kg/day. Use therapeutic drug monitoring to guide dose adjustments. PM Initiate therapy with standard Strong recommended dose. Use therapeutic drug monitoring to guide dose adjustments. Favorable Approximately 90% chance for SVR IFNL3 treatment Strong (Muir et al., 2014) with response after 24-48 weeks of treatment. Approximately 80-90% of patients Peginterferon genotype





Drug/Genotype	Phenotype	Summary of CPIC Therapeutic Recommendations	Level of Recommend ations	Reference
alfa-2a, Peginterferon alfa-2b or Ribavirin	Unfavorable	are eligible for shortened therapy (24–28 weeks vs. 48 weeks). Weighs in favor of using PEG-IFN- α - and RBV- containing regimens. Approximately 60% chance of SVR	Strong	
	response genotype	after 24–48 weeks of treatment. Approximately 50% of patients are eligible for shortened therapy regimens (24–28 weeks). Consider implications before initiating PEG-IFN-α- and RBV-containing regimens.	Strong	
RYR1 and CACNA1S genotypes for Potent Volatile Anesthetic Agents and Succinylcholine	Malignant Hyperthermia Susceptible	Halogenated volatile anesthetics or depolarizing Halogenated volatile anesthetics or depolarizing muscle relaxants succinylcholine are relatively contraindicated in persons with MHS. They should not be used, except in extraordinary circumstances in which the benefits outweigh the risks. In general, alternative anesthetics are widely available and effective in patients with MHS	Strong	(Gonsalves et al., 2019)
	Uncertain susceptibility	Clinical findings, family history, further genetic testing and other laboratory data should guide use of halogenated volatile anesthetics or depolarizing muscle relaxants.	Strong	

CPIC notes that evidence for *TYMS* testing is unclear or weak and have assigned *TYMS* a "D" level recommendation. CPIC does not recommend any change in prescription based on *TYMS* genotype (CPIC, 2023a).





American College of Medical Genetics and Genomics (ACMG)

ACMG notes that *CYP2C9* and *VKORC1* testing may be useful for assessing unusual responses to warfarin, but cannot recommend for or against routine genotyping (ACMG, 2007).

American College of Cardiology Foundation (AACF) and the American Heart Association (AHA) Joint Guidelines

A report by the ACCF and the AHA on genetic testing for selection and dosing of clopidogrel provided the following recommendations for practice:

- "Clinicians must be aware that genetic variability in CYP enzymes alter clopidogrel metabolism, which in turn can affect its inhibition of platelet function. Diminished responsiveness to clopidogrel has been associated with adverse patient outcomes in registry experiences and clinical trials."
- "The specific impact of the individual genetic polymorphisms on clinical outcome remains to be determined (e.g., the importance of *CYP2C19*2* versus *3 or *4 for a specific patient), and the frequency of genetic variability differs among ethnic groups."
- "Information regarding the predictive value of pharmacogenomic testing is very limited at this time; resolution of this issue is the focus of multiple ongoing studies."
- "The evidence base is insufficient to recommend either routine genetic or platelet function testing at the present time. There is no information that routine testing improves outcome in large subgroups of patients. In addition, the clinical course of the majority of patients treated with clopidogrel without either genetic testing or functional testing is excellent. Clinical judgment is required to assess clinical risk and variability in patients considered to be at increased risk. Genetic testing to determine if a patient is predisposed to poor clopidogrel metabolism ("poor metabolizers") may be considered before starting clopidogrel therapy in patients believed to be at moderate or high risk for poor outcomes. This might include, among others, patients undergoing elective high-risk PCI procedures (e.g., treatment of extensive and/or very complex disease). If such testing identifies a potential poor metabolizer, other therapies, particularly prasugrel for coronary patients, should be considered. (Holmes et al., 2010).

American Academy of Neurology (AAN)

The AAN published a position paper on the use of opioids for chronic non-cancer pain. Regarding pharmacogenetic testing, the guidelines state "genotyping to determine whether response to opioid therapy can/should be more individualized will require critical original research to determine effectiveness and appropriateness of use" (Franklin, 2014).

American Association for Clinical Chemistry (AACC) Academy Laboratory Medicine Practice Guidelines

AACC Academy issued laboratory medicine practice guidelines on using clinical laboratory tests to monitor drug therapy in pain management. Their guidelines have a total of 26 recommendations and seven expert opinions. Regarding pharmacogenetic testing for pain management, they stated in the recommendation #20 (Level A, II) that: "While the current evidence in the literature doesn't support routine genetic testing for all pain management patients, it should be considered to predict or explain





variant pharmacokinetics, and/ or pharmacodynamics of specific drugs as evidenced by repeated treatment failures, and/or adverse drug reactions/toxicity" (AACC, 2017).

American Family Physician (AAFP)

The AAFP has published guidelines on pharmacogenetics: using genetic information to guide drug therapy. CPIC guidelines are cited for many medication/allele combinations in this article. The recommendations by the AAFP are listed in the table below taken from Chang et al. (2015):

Allele	Medications	Test Results and Clinical Implications	Comments
CYP2D6	Codeine, hydrocodone, oxycodone, tramadol	Ultrarapid metabolizer: Avoid codeine because of potential for toxicity Poor metabolizer: Avoid codeine and possibly tramadol because of possible lack of effectiveness	CPIC guidance limits genotype-guided dosing recommendations to codeine. Alternative analgesics not affected by CYP2D6 variability include morphine, oxymorphone, and nonopioid analgesics. Oxycodone may also have reduced effectiveness in poor CYP2D6 metabolizers.
CYP2C19	Clopidogrel (Plavix)	Intermediate metabolizer: Use alternative antiplatelet therapy if no contraindications Poor metabolizer: Use alternative antiplatelet therapy if no contraindications	Clopidogrel prescribing information states that CYP2C19 tests can be used as an aid to determine therapeutic strategy in patients with acute coronary syndromes who are undergoing percutaneous coronary intervention. CPIC guidance limits genotype-guided dosing recommendations to patients undergoing percutaneous coronary intervention for acute coronary syndromes (excluding medical management of acute coronary syndromes, stroke, and peripheral artery disease). ACCF/AHA guidelines state that genotyping may be considered in patients with unstable angina/non-ST segment elevation myocardial infarction (or after percutaneous coronary intervention for acute coronary syndromes) if test results could alter management. Alternative antiplatelet therapy not affected by CYP2C19 variability includes prasugrel (Effient) and ticagrelor (Brilinta).
CYP2C19	Amitriptyline	Poor metabolizer: Consider 50% reduction	CPIC guidance is available for CYP2D6- and CYP2C19-genotype guided tricyclic antidepressant therapy.





Allele	Medications	Test Results and Clinical Implications	Comments
		in recommended starting dose	Although limited data exist for other tricyclic antidepressants, most supporting evidence of clinically relevant gene-drug effects is for amitriptyline and nortriptyline (Pamelor).
CYP2C19	Citalopram (Celexa), escitalopram (Lexapro)	Ultrarapid metabolizer: Consider alternative Poor metabolizer: Consider 50% starting dose reduction and titrate to response, or use alternative	cPIC guidance is available for CYP2C19- genotype guided citalopram and escitalopram therapy. FDA label for citalopram states that 20 mg per day is the maximum recommended dosage for patients older than 60 years, patients with hepatic impairment, and CYP2C19 poor metabolizers or patients taking cimetidine (Tagamet) or another CYP2C19 inhibitor.
CYP2C19	Sertraline (Zoloft)	Ultrarapid metabolizer: If patient does not respond to recommended dose, consider alternative Poor metabolizer: Consider 50% dose reduction or alternative	CPIC guidance is available for <i>CYP2C19</i> -genotype guided sertraline therapy.
CYP2D6	Amitriptyline, nortriptyline	Ultrarapid metabolizer: Avoid because of possible lack of effectiveness Poor metabolizer: Avoid because of possible adverse effects; if use is warranted, consider 50% reduction in recommended starting dose	CPIC guidance is available for CYP2D6- and CYP2C19-genotype guided tricyclic antidepressant therapy. Although limited data exist for other tricyclic antidepressants, most supporting evidence of clinically relevant gene-drug effects is for amitriptyline and nortriptyline.
CYP2D6	Aripiprazole (Abilify)	Poor metabolizer: Decrease dose	Quality of supporting evidence is classified as low by PharmGKB FDA label for aripiprazole states that in poor metabolizers, the usual dose should initially be reduced to 50% and then adjusted to achieve a favorable clinical response; in poor metabolizers receiving a strong CYP3A4 inhibitor, the usual dose should be reduced to 25%.





Allele	Medications	Test Results and Clinical Implications	Comments
CYP2D6	Atomoxetine (Strattera)	Poor metabolizer: Adjust dose	Quality of supporting evidence is classified as moderate (Level 2a) by PharmGKB.
			FDA label for atomoxetine states that in poor metabolizers, the initial dosage should be 0.5 mg per kg per day and then increased to the the usual target dosage of 1.2 mg per kg per day only if symptoms do not improve after 4 weeks and the initial dose is well tolerated.
CYP2D6	Paroxetine (Paxil)	Ultrarapid metabolizer: Select alternative because of possible lack of effectiveness. Poor metabolizer: Select alternative or if use is warranted, consider 50% starting dose reduction	CPIC guidance is available for CYP2D6-genotype guided paroxetine therapy.

Dutch Pharmacogenetics Working Group (DPWG)

The DPWG has published guidelines for the gene-drug interaction of *DPYD* and fluoropyrimidines. Conclusions state that "four variants have sufficient evidence to be implemented into clinical care: DPYD*2A (c.1905+1G>A, IVS14+1G>A), DPYD*13 (c.1679T>G), c.2846A>T and c.1236G>A (in linkage disequilibrium with c.1129–5923C>G). The current guideline only reports recommendations for these four variants; no recommendations are provided for other variants in *DPYD* or other genes" (Lunenburg et al., 2020).

Food and Drug Administration

The FDA published several tables of pharmacogenetic associations with "sufficient scientific evidence to suggest that subgroups of patients with certain genetic variants, or genetic variant-inferred phenotypes (i.e., affected subgroup in the table below), are likely to have altered drug metabolism, and in certain cases, differential therapeutic effects, including differences in risks of adverse events".

The table below lists associations "for which the data support therapeutic management recommendations" (FDA, 2022).

Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Abacavir	HLA-B	*57:01 allele positive	Results in higher adverse reaction risk (hypersensitivity reactions). Do not use abacavir in patients positive for HLA-B*57:01.





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Abrocitinib	CYP2C19	poor metabolizers	Results in higher systemic concentrations and may result in higher adverse reaction risk. Dosage adjustment is recommended. Refer to FDA labeling for specific dosing recommendations.
Amifampridine	NAT2	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk. Use lowest recommended starting dosage and monitor for adverse reactions. Refer to FDA labeling for specific dosing recommendations.
Amifampridine Phosphate	NAT2	poor metabolizers	Results in higher systemic concentrations. Use lowest recommended starting dosage (15 mg/day) and monitor for adverse reactions.
Amphetamine	CYP2D6	poor metabolizers	May affect systemic concentrations and adverse reaction risk. Consider lower starting dosage or use alternative agent.
Aripiprazole	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk. Dosage adjustment is recommended. Refer to FDA labeling for specific dosing recommendations.
Aripiprazole Lauroxil	CYP2D6	poor metabolizers	Results in higher systemic concentrations. Dosage adjustment is recommended. Refer to FDA labeling for specific dosing recommendations.
Atomoxetine	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk. Adjust titration interval and increase dosage if tolerated. Refer to FDA labeling for specific dosing recommendations.
Azathioprine	TPMT and/or NUDT15	intermediate or poor metabolizers	Alters systemic active metabolite concentration and dosage requirements. Results in higher adverse reaction risk (myelosuppression). Consider alternative therapy in poor metabolizers. Dosage reduction is recommended in intermediate metabolizers for NUDT15 or TPMT. Intermediate metabolizers for both genes may require more substantial dosage reductions. Refer to FDA labeling for specific dosing recommendations.





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Belinostat	UGT1A1	*28/*28 (poor metabolizers)	May result in higher systemic concentrations and higher adverse reaction risk. Reduce starting dose to 750 mg/m2 in poor metabolizers.
Belzutifan	CYP2C19 and/or UGT2B17	Poor metabolizers	Results in higher systemic concentrations and may result in higher adverse reaction risk (anemia, hypoxia). Monitor patients who are poor metabolizers for both genes for adverse reactions.
Brexpiprazole	CYP2D6	poor metabolizers	Results in higher systemic concentrations. Dosage adjustment is recommended. Refer to FDA labeling for specific dosing recommendations.
Brivaracetam	CYP2C19	intermediate or poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk. Consider dosage reductions in poor metabolizers.
Capecitabine	DPYD	intermediate or poor metabolizers	Results in higher adverse reaction risk (severe, life-threatening, or fatal toxicities). No dosage has proven safe in poor metabolizers, and insufficient data are available to recommend a dosage in intermediate metabolizers. Withhold or discontinue in the presence of early-onset or unusually severe toxicity.
Carbamazepine	HLA-B	*15:02 allele positive	Results in higher adverse reaction risk (severe skin reactions). Avoid use unless potential benefits outweigh risks and consider risks of alternative therapies. Patients positive for HLA-B*15:02 may be at increased risk of severe skin reactions with other drugs that are associated with a risk of Stevens Johnson Syndrome/Toxic Epidermal necrolysis (SJS/TEN). Genotyping is not a substitute for clinical vigilance.
Celecoxib	CYP2C9	poor metabolizers	Results in higher systemic concentrations. Reduce starting dose to half of the lowest recommended dose in poor metabolizers. Consider alternative therapy in patients with juvenile rheumatoid arthritis.
Citalopram	CYP2C19	poor metabolizers	Results in higher systemic concentrations and adverse reaction risk (QT prolongation). The maximum recommended dose is 20 mg.





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Clobazam	CYP2C19	intermediate or poor metabolizers	Results in higher systemic active metabolite concentrations. Poor metabolism results in higher adverse reaction risk. Dosage adjustment is recommended. Refer to FDA labeling for specific dosing recommendations.
Clopidogrel	CYP2C19	intermediate or poor metabolizers	Results in lower systemic active metabolite concentrations, lower antiplatelet response, and may result in higher cardiovascular risk. Consider use of another platelet P2Y12 inhibitor.
Clozapine	CYP2D6	poor metabolizers	Results in higher systemic concentrations. Dosage reductions may be necessary.
Codeine	CYP2D6	ultrarapid metabolizers	Results in higher systemic active metabolite concentrations and higher adverse reaction risk (life-threatening respiratory depression and death). Codeine is contraindicated in children under 12 years of age.
Deutetrabenazine	CYP2D6	poor metabolizers	Results in higher systemic concentrations and adverse reaction risk (QT prolongation). The maximum recommended dosage should not exceed 36 mg (maximum single dose of 18 mg).
Dronabinol	CYP2C9	intermediate or poor metabolizers	May result in higher systemic concentrations and higher adverse reaction risk. Monitor for adverse reactions.
Eliglustat	CYP2D6	ultrarapid, normal, intermediate, or poor metabolizers	Alters systemic concentrations, effectiveness, and adverse reaction risk (QT prolongation). Indicated for normal, intermediate, and poor metabolizer patients. Ultrarapid metabolizers may not achieve adequate concentrations to achieve a therapeutic effect. The recommended dosages are based on CYP2D6 metabolizer status. Coadministration with strong CYP3A inhibitors is contraindicated in intermediate and poor CYP2D6 metabolizers. Refer to FDA labeling for specific dosing recommendations.
Erdafitinib	CYP2C9	*3/*3 (poor metabolizers)	May result in higher systemic concentrations and higher adverse reaction risk. Monitor for adverse reactions.





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Flibanserin	CYP2C19	poor metabolizers	May result in higher systemic concentrations and higher adverse reaction risk. Monitor patients for adverse reactions.
Flurbiprofen	CYP2C9	poor metabolizers	Results in higher systemic concentrations. Use a reduced dosage.
Fluorouracil	DPYD	intermediate or poor metabolizer	Results in higher adverse reaction risk (severe, life-threatening, or fatal toxicities). No dosage has proven safe in poor metabolizers and insufficient data are available to recommend a dosage in intermediate metabolizers. Withhold or discontinue in the presence of early-onset or unusually severe toxicity.
Fosphenytoin	CYP2C9	Intermediate or poor metabolizers	May result in higher systemic concentrations and higher adverse reaction risk (central nervous system toxicity). Consider starting at the lower end of the dosage range and monitor serum concentrations. Refer to FDA labeling for specific dosing recommendations. Carriers of CYP2C9*3 alleles may be at increased risk of severe cutaneous adverse reactions. Consider avoiding fosphenytoin as an alternative to carbamazepine in patients who are CYP2C9*3 carriers. Genotyping is not a substitute for clinical vigilance and patient management.
Fosphenytoin	HLA-B	*15:02 allele positive	May result in higher adverse reaction risk (severe cutaneous reactions). Patients positive for HLA-B*15:02 may be at increased risk of Stevens Johnson Syndrome/Toxic Epidermal necrolysis (SJS/TEN). Consider avoiding fosphenytoin as an alternative to carbamazepine in patients who are positive for HLA-B*15:02. Genotyping is not a substitute for clinical vigilance and patient management.
Gefitinib	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk. Monitor for adverse reactions.
lloperidone	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (QT prolongation). Reduce dosage by 50%.





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Irinotecan	UGT1A1	*28/*28 (poor metabolizers)	Results in higher systemic active metabolite concentrations and higher adverse reaction risk (severe neutropenia). Consider reducing the starting dosage by one level and modify the dosage based on individual patient tolerance.
Lofexidine	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk. Monitor for orthostatic hypotension and bradycardia.
Meclizine	CYP2D6	ultrarapid, intermediate, or poor metabolizers	May affect systemic concentrations. Monitor for adverse reactions and clinical effect.
Meloxicam	CYP2C9	Poor metabolizers or *3 carriers	Results in higher systemic concentrations. Consider dose reductions in poor metabolizers. Monitor patients for adverse reactions.
Metoclopramide	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk. The recommended dosage is lower. Refer to FDA labeling for specific dosing recommendations.
Mercaptopurine	TPMT and/or NUDT15	intermediate or poor metabolizers	Alters systemic active metabolite concentration and dosage requirements. Results in higher adverse reaction risk (myelosuppression). Initial dosages should be reduced in poor metabolizers; poor metabolizers generally tolerate 10% or less of the recommended dosage. Intermediate metabolizers may require dosage reductions based on tolerability. Intermediate metabolizers for both genes may require more substantial dosage reductions. Refer to FDA labeling for specific dosing recommendations.
Mivacurium	ВСНЕ	intermediate or poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (prolonged neuromuscular blockade). Avoid use in poor metabolizers.
Nateglinide	CYP2C9	Poor metabolizers	Results in higher systemic concentrations and may result in higher adverse reaction risk (hypoglycemia). Dosage reduction is recommended. Increase monitoring frequency





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
			for adverse reactions. Refer to FDA labeling for specific dosing recommendations.
Oliceridine	CYP2D6	Poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (respiratory depression and sedation). May require less frequent dosing.
Pantoprazole	CYP2C19	poor metabolizers	Results in higher systemic concentrations. Consider dosage reduction in children who are poor metabolizers. No dosage adjustment is needed for adult patients who are poor metabolizers.
Phenytoin	CYP2C9	Intermediate or poor metabolizers	May result in higher systemic concentrations and higher adverse reaction risk (central nervous system toxicity). Refer to FDA labeling for specific dosing recommendations. Carriers of CYP2C9*3 alleles may be at increased risk of severe cutaneous adverse reactions. Consider avoiding phenytoin as an alternative to carbamazepine in patients who are CYP2C9*3 carriers. Genotyping is not a substitute for clinical vigilance and patient management.
Phenytoin	HLA-B	*15:02 allele positive	May result in higher adverse reaction risk (severe cutaneous reactions). Patients positive for HLA-B*15:02 may be at increased risk of Stevens Johnson Syndrome/Toxic Epidermal necrolysis (SJS/TEN). Consider avoiding phenytoin as an alternative to carbamazepine in patients who are positive for HLA-B*15:02. Genotyping is not a substitute for clinical vigilance and patient management.
Pimozide	CYP2D6	poor metabolizers	Results in higher systemic concentrations. Dosages should not exceed 0.05 mg/kg in children or 4 mg/day in adults who are poor metabolizers and dosages should not be increased earlier than 14 days.
Piroxicam	CYP2C9	intermediate or poor metabolizers	Results in higher systemic concentrations. Consider reducing dosage in poor metabolizers.
Pitolisant	CYP2D6	Poor metabolizers	Results in higher systemic concentrations. Use lowest recommended starting dosage. Refer





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
			to FDA labeling for specific dosing recommendations.
Propafenone	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (arrhythmia). Avoid use in poor metabolizers taking a CYP3A4 inhibitor.
Sacituzumab Govitecan-hziy	UGT1A1	*28/*28 (poor metabolizers)	May result in higher systemic concentrations and adverse reaction risk (neutropenia). Monitor for adverse reactions and tolerance to treatment.
Siponimod	CYP2C9	intermediate or poor metabolizers	Results in higher systemic concentrations. Adjust dosage based on genotype. Do not use in patients with CYP2C9 *3/*3 genotype. Refer to FDA labeling for specific dosing recommendations.
Succinylcholine	ВСНЕ	intermediate or poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (prolonged neuromuscular blockade). Avoid use in poor metabolizers. May administer test dose to assess sensitivity and administer cautiously via slow infusion.
Tacrolimus	CYP3A5	intermediate or normal metabolizers	Results in lower systemic concentrations and lower probability of achieving target concentrations. Measure drug concentrations and adjust dosage based on trough whole blood tacrolimus concentrations.
Tetrabenazine	CYP2D6	poor metabolizers	Results in higher systemic concentrations. The maximum recommended single dose is 25 mg and should not exceed 50 mg/day.
Thioguanine	TPMT and/or NUDT15	intermediate or poor metabolizers	Alters systemic active metabolite concentration and dosage requirements. Results in higher adverse reaction risk (myelosuppression). Initial dosages should be reduced in poor metabolizers; poor metabolizers generally tolerate 10% or less of the recommended dosage. Intermediate metabolizers may require dosage reductions based on tolerability. Intermediate metabolizers for both genes may require more substantial dosage reductions. Refer to FDA labeling for specific dosing recommendations.





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Thioridazine	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (QT prolongation). Predicted effect based on experience with CYP2D6 inhibitors. Contraindicated in poor metabolizers.
Tramadol	CYP2D6	Ultrarapid metabolizers	Results in higher systemic and breast milk active metabolite concentrations, which may result in respiratory depression and death. Contraindicated in children under 12 and in adolescents following tonsillectomy/adenoidectomy. Breastfeeding is not recommended during treatment.
Valbenazine	CYP2D6	poor metabolizers	Results in higher systemic active metabolite concentrations and higher adverse reaction risk (QT prolongation). Dosage reductions may be necessary.
Venlafaxine	CYP2D6	poor metabolizers	Alters systemic parent drug and metabolite concentrations. Consider dosage reductions.
Vortioxetine	CYP2D6	poor metabolizers	Results in higher systemic concentrations. The maximum recommended dose is 10 mg.
Warfarin	CYP2C9	intermediate or poor metabolizers	Alters systemic concentrations and dosage requirements. Select initial dosage, taking into account clinical and genetic factors. Monitor and adjust dosages based on INR.
Warfarin	CYP4F2	V433M variant carriers	May affect dosage requirements. Monitor and adjust doses based on INR.
Warfarin	VKORC1	-1639G>A variant carriers	Alters dosage requirements. Select initial dosage, taking into account clinical and genetic factors. Monitor and adjust dosages based on INR.

The table below lists associations "for which the data indicate a potential impact on safety or response" (FDA, 2022).

Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Allopurinol	HLA-B	*58:01 allele positive	Results in higher adverse reaction risk (severe skin reactions).
Carbamazepine	HLA-A	*31:01 allele positive	Results in higher adverse reaction risk (severe skin reactions). Consider risk and benefit of carbamazepine use in patients





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
			positive for HLA-A*31:01. Genotyping is not a substitute for clinical vigilance.
Carvedilol	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (dizziness).
Cevimeline	CYP2D6	poor metabolizers	May result in higher adverse reaction risk. Use with caution.
Codeine	CYP2D6	poor metabolizers	Results in lower systemic active metabolite concentrations and may result in reduced efficacy.
Efavirenz	CYP2B6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (QT prolongation).
Isoniazid	Nonspecific (NAT)	poor metabolizers	May result in higher systemic concentrations and adverse reaction risk.
Lapatinib	HLA-DRB1	*07:01 allele positive	Results in higher adverse reaction risk (hepatotoxicity). Monitor liver function tests regardless of genotype.
Lapatinib	HLA-DQA1	*02:01 allele positive	Results in higher adverse reaction risk (hepatotoxicity). Monitor liver function tests regardless of genotype.
Mavacamten	CYP2C19	Intermediate or poor metabolizers	Results in higher systemic concentrations and may have higher adverse reaction risk (heart failure). Dosage is based on individual response. The dose titration and monitoring schedule accounts for differences due to CYP2C19 genetic variation, so adjustments based on CYP2C19 genotype are not necessary. Refer to FDA labeling for specific dosing recommendations and monitoring.
Nilotinib	UGT1A1	*28/*28 (poor metabolizers)	Results in higher adverse reaction risk (hyperbilirubinemia).
Oxcarbazepine	HLA-B	*15:02 allele positive	Results in higher adverse reaction risk (severe skin reactions). Patients positive for HLA-B*15:02 may be at increased risk of severe skin reactions with other drugs that are associated with a risk of Stevens Johnson Syndrome/Toxic Epidermal necrolysis (SJS/TEN). Genotyping is not a substitute for clinical vigilance.





Drug	Gene	Affected Subgroups	Description of Gene-Drug Interaction
Pazopanib	HLA-B	*57:01 allele positive	May result in higher adverse reaction risk (liver enzyme elevations). Monitor liver function tests regardless of genotype.
Pazopanib	UGT1A1	*28/*28 (poor metabolizers)	Results in higher adverse reaction risk (hyperbilirubinemia).
Perphenazine	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk.
Procainamide	Nonspecific (NAT)	poor metabolizers	Alters systemic parent drug and metabolite concentrations. May result in higher adverse reaction risk.
Simvastatin	SLCO1B1	521 TC or 521 CC (intermediate or poor function transporters)	Results in higher systemic concentrations and higher adverse reaction risk (myopathy). The risk of adverse reaction (myopathy) is higher for patients on 80 mg than for those on lower doses.
Sulfamethoxazole and Trimethoprim	Nonspecific (NAT)	poor metabolizers	May result in higher adverse reaction risk.
Sulfasalazine	Nonspecific (NAT)	poor metabolizers	Results in higher systemic metabolite concentrations and higher adverse reaction risk.
Tolterodine	CYP2D6	poor metabolizers	Results in higher systemic concentrations and higher adverse reaction risk (QT prolongation).
Tramadol	CYP2D6	poor metabolizers	Results in lower systemic active metabolite concentrations and may result in reduced efficacy.
Voriconazole	CYP2C19	Intermediate or poor metabolizers	Results in higher systemic concentrations and may result in higher adverse reaction risk.

The International Society of Psychiatric Genetics

The International Society of Psychiatric Genetics (ISPG) released recommendations on the use of pharmacogenetic testing to guide psychiatric treatment. ISPG recommends that pharmacogenetic testing should be used as a decision-support tool. *HLA-A* and *HLA-B* testing is recommended before the use of carbamazepine and oxcarbazepine. *CYP2C19* and *CYP2D6* testing would be beneficial for those who experienced an inadequate response or adverse reaction to a previous antidepressant or antipsychotic medication (ISPG, 2019).





The American Academy of Child and Adolescent Psychiatry

AACAP does not recommend the use of pharmacogenetic testing to select psychotropic medications for children and adolescents (AACAP, 2020)

Association for Molecular Pathology PGx Working Group (AMP)

AMP released clinical practice guidelines to define a minimum set of CYP2C19 allele variants that should be included in the pharmacogenomic genotyping assay. Tier 1 represents alleles that have been shown to affect drug response and should be included, while Tier 2 represents alleles which meet at least one but not all the criteria for inclusion in Tier 1 and are considered optional for inclusion in expanded clinical genotyping panels. Those in Tier 1 include alleles *2, *3, and *17. The following CYP2C19 alleles were recommended as Tier 2: *4A, *4B, *5, *6, *7, *8, *9, *10, and *35 (Pratt et al., 2018). Regarding CYP2C9 variant alleles, Tier 1 alleles include CYP2C9 *2, *3, *5, *6, *8, and *11. The following CYP2C9 alleles are recommended for inclusion in Tier 2: CYP2C9*12, *13, and *15 (Pratt et al., 2019). For testing genes and alleles specific to warfarin, AMP recommends including VKORC1 c.-1639G>A in Tier 1 and VKORC1 c.196G>A and c.106G>A in Tier 2 (Pratt et al., 2020). In a joint recommendation endorsed by the AMP, College of American Pathologists, Dutch Pharmacogenetics Working Group of the Royal Dutch Pharmacists Association, and the European Society for Pharmacogenomics and Personalized Therapy, CYP2D6 variant alleles were elucidated. Tier 1 alleles include CYP2D6 *2 to *6, *9, *10, *17, *29, and *41. Tier 2 *CYP2D6* alleles include *CYP2D6* *7, *8, *12, *14, *15, *21, *31, *40, *42, *49, *56, and *59, and hybrid genes containing portions of CYP2D6 and CYP2D7 (Pratt et al., 2021). These recommendations should help to standardize testing and genotyping concordance among laboratories.

European Medicines Agency

EMA released recommendations on *DPD* testing before treatment with fluorouracil, capecitabine, tegafur, and flucytosine. EMA recommends testing for the lack of *DPD* before starting cancer treatment with fluorouracil, capecitabine, or tegafur. Patients who completely lack *DPD* should not be given these medications. For patients with partial deficiency, the physician may consider beginning treatment at a lower dose and terminating treatment if severe side effects occur. These recommendations do not apply to fluorouracil medications used for skin conditions or flucytosine used for fungal infection (EMA, 2020).

VI. Applicable State and Federal Regulations

DISCLAIMER: If there is a conflict between this Policy and any relevant, applicable government policy for a particular member [e.g., Local Coverage Determinations (LCDs) or National Coverage Determinations (NCDs) for Medicare and/or state coverage for Medicaid], then the government policy will be used to make the determination. For the most up-to-date Medicare policies and coverage, please visit the Medicare search website. For the most up-to-date Medicaid policies and coverage, visit the applicable state Medicaid website.

Food and Drug Administration (FDA)

Diagnostic genotyping tests for certain drug metabolizing enzymes are FDA-approved. Many labs have developed specific tests that they must validate and perform in house. These laboratory-developed tests







(LDTs) are regulated by the Centers for Medicare and Medicaid (CMS) as high-complexity tests under the Clinical Laboratory Improvement Amendments of 1988 (CLIA'88). As an LDT, the U. S. Food and Drug Administration has not approved or cleared this test; however, FDA clearance or approval is not currently required for clinical use.

Currently, there are over 14 other FDA-approved tests for the drug metabolizing enzymes that are nucleic acid-based tests including xTAG *CYP2D6* Kit v3 and XTAG *CYP2C19* KIT V3 (Luminex Molecular Diagnostics, Inc), Spartan RX *CYP2C19* Test System (Spartan Bioscience, Inc), Verigene *CYP2C19* Nucleic Acid Test (Nanosphere, Inc), INFINITI *CYP2C19* Assay (AutoGenomics, Inc), Invader *UGT1A1* (Third Wave Technologies Inc.), eSensor Warfarin Sensitivity Saliva Test (GenMark Diagnostics), eQ-PCR LC Warfarin Genotyping kit (TrimGen Corporation), eSensor Warfarin Sensitivity Test and XT-8 Instrument (Osmetech Molecular Diagnostics), Gentris Rapid Genotyping Assay-*CYP2C9*&VKORCI (ParagonDx, LLC), INFINITI 2C9 & *VKORC1* Multiplex Assay for Warfarin (AutoGenomics, Inc), Verigene Warfarin Metabolism Nucleic Acid Test and Verigene System (Nanosphere, Inc), TruDiagnosis System (Akonni Systems, Inc), Roche AmpliChip CYP450 microassay (Roche Molecular Systems, Inc) (FDA, 2021a).

FDA Notes

The Office of Clinical Pharmacology within FDA includes The Genomics and Targeted Therapy Group responsible for applying pharmacogenomics and other biomarkers in drug development and clinical practice. The FDA scientists review current pharmacogenomic information and ensure that pharmacogenomic strategies are utilized appropriately in all phases of drug development (FDA, 2022).

The current list of pharmacogenomic biomarkers in drug labeling by FDA contain numerous medications that have genotypes related to metabolism dosage recommendations or warnings. These medications are involved in different therapeutic areas and the list includes the following genes and medications:

CYP1A2: Rucaparib

CYP2B6: Efavirenz, Prasugrel, Ospemifene

CYP2C19 contains 22 different medications: Clopidogrel, Prasugrel, Ticagrelor, Lansoprazole, Omeprazole, Esomeprazole, Rabeprazole, Pantoprazole, Dexlansoprazole, Flibanserin, Drospirenone and Ethinyl Estradiol, Voriconazole, Lacosamide, Brivaracetam, Clobazam, Phenytoin, Diazepam, Citalopram, Escitalopram, Doxepin, Formoterol, Carisoprodol

CYP2C9 contains 15 different medications: Prasugrel, Dronabinol, Flibanserin, Warfarin, Phenytoin, Celecoxib, Piroxicam, Flurbiprofen, Lesinurad, Avatrombopag, Erdafitinib, Ospemifene, Siponimod, Meloxicam, Rimegepant

CYP2D6 contains 70 different medications: Tramadol, Metoprolol, Nebivolol, Propafenone, Propranolol, Ondansetron, Palonosetron, Flibanserin, Eliglustat, Deutetrabenazine, Dextromethorphan and Quinidine, Galantamine, Tetrabenazine, Valbenazine, Rucaparib, Aripiprazole, Aripiprazole Lauroxil, Atomoxetine, Brexpiprazole, Cariprazine, Citalopram, Clozapine, Desvenlafaxine, Doxepin, Escitalopram, Fluoxetine, Fluvoxamine, Iloperidone, Modafinil, Paroxetine, Perphenazine, Risperidone, Venlafaxine, Vortioxetine, Arformoterol, Formoterol, Umeclidinium, Darifenacin, Mirabegron, Tolterodine, Amphetamine, Donepezil, Fesoterodine, Gefitinib, Metoclopramide, Paliperidone, Tamoxifen,





Carvedilol, Amitriptyline, Amoxapine, Clomipramine, Codeine, Desipramine, Duloxetine, Imipramine, Meclizine, Metoclopramide, Nefazodone, Nortriptyline, Pimozide, Protriptyline, Quinine Sulfate, Tamsulosin, Thioridazine, Trimipramine, Pitolisant, Upadacitinib, Bupropion.

CYP3A5: Prasugrel

TPMT: Thioguanine, Azathioprine, Mercaptopurine, Cisplatin

NUDT15: Thioguanine, Azathioprine, Mercaptopurine

UGT1A1: Arformoterol, Belinostat, Binimetinib, Dolutegravir, Indacaterol, Irinotecan, Nilotinib, Pazopanib, Raltegravir, Sacituzumab Govitecan-hziy (FDA, 2021b).

FDA Recommendations

The FDA package insert for Plavix (clopidogrel) carries the following "Black Box" warning: "The effectiveness of Plavix results from its antiplatelet activity which is dependent on its conversion to an active metabolite by the cytochrome P450 (CYP) system, principally *CYP2C19*. Plavix at recommended doses forms less of the active metabolite and so has a reduced effect on platelet activity in patients who are homozygous for nonfunctional alleles of the *CYP2C19* gene, (termed "*CYP2C19* poor metabolizers"). Tests are available to identify patients who are *CYP2C19* poor metabolizers. Consider another platelet P2Y12 inhibitor in patients identified as *CYP2C19* poor metabolizers." (FDA, 2016)

The FDA package insert for Xenazine (tetrabenazine) indicates, "Patients who require doses of Xenazine greater than 50 mg per day should be first tested and genotyped to determine if they are poor metabolizers (PMs) or extensive metabolizers (EMs) by their ability to express the drug metabolizing enzyme, *CYP2D6*. The dose of XENAZINE should then be individualized accordingly to their status as PMs or EMs. (FDA, 2008)

The Coumadin (warfarin) highlights of prescription information notes that "The appropriate initial dosing of COUMADIN varies widely for different patients. Not all factors responsible for warfarin dose variability are known, and the initial dose is influenced by: Genetic factors (CYP2C9 and VKORC1 genotypes)." Although dosage suggestions based on CYP2C9 and VKORC1 genotypes are provided in the package insert, the requirement for genetic testing is not included (FDA)

The eligibility and dosing of Eliglustat is dependent on cytochrome P450 *CYP2D6* genotype as eliglustat is extensively metabolized by *CYP2D6*. The FDA contraindicates this medication in the following patients due "to the risk of cardiac arrhythmias from prolongation of the PR, QTc, and/or QRS cardiac Intervals":

EMs of CYP2D6

- Taking a strong or moderate CYP2D6 inhibitor concomitantly with a strong or moderate CYP3A inhibitor
- Moderate or severe hepatic impairment
- Mild hepatic impairment and taking a strong or moderate CYP2D6 inhibitor





IMs

- Taking a strong or moderate CYP2D6 inhibitor concomitantly with a strong or moderate CYP3A inhibitor
- Taking a strong CYP3A inhibitor
- Any degree of hepatic impairment

PMs

- Taking a strong CYP3A inhibitor
- Any degree of hepatic impairment (FDA, 2014)

The FDA also includes a warning for irinotecan's interaction with *UGT1A1*, stating "When administered in combination with other agents, or as a single-agent, a reduction in the starting dose by at least one level of CAMPTOSAR [irinotecan] should be considered for patients known to be homozygous for the *UGT1A1*28* allele" (FDA, 2021b).

Many labs have developed specific tests that they must validate and perform in house. These laboratory-developed tests (LDTs) are regulated by the Centers for Medicare and Medicaid (CMS) as high-complexity tests under the Clinical Laboratory Improvement Amendments of 1988 (CLIA '88). LDTs are not approved or cleared by the U. S. Food and Drug Administration; however, FDA clearance or approval is not currently required for clinical use.

VII. Important Reminder

The purpose of this Medical Policy is to provide a guide to coverage. This Medical Policy is not intended to dictate to providers how to practice medicine. Nothing in this Medical Policy is intended to discourage or prohibit providing other medical advice or treatment deemed appropriate by the treating physician.

Benefit determinations are subject to applicable member contract language. To the extent there are any conflicts between these guidelines and the contract language, the contract language will control.

This Medical Policy has been developed through consideration of the medical necessity criteria under Hawaii's Patients' Bill of Rights and Responsibilities Act (Hawaii Revised Statutes §432E-1.4) or for QUEST members, under Hawaii Administrative Rules (HAR 1700.1-42), generally accepted standards of medical practice and review of medical literature and government approval status.

HMSA has determined that services not covered under this Medical Policy will not be medically necessary under Hawaii law in most cases. If a treating physician disagrees with HMSA's determination as to medical necessity in a given case, the physician may request that HMSA reconsider the application of the medical necessity criteria to the case at issue in light of any supporting documentation.

Genetic testing is covered for level 1 or 2A recommendations of the National Comprehensive Cancer Network (NCCN and in accordance with Hawaii's Patients' Bill of Rights and Responsibilities Act (Hawaii Revised Statutes §432E-1.4) or for QUEST members, the Hawaii Administrative Rules (HAR 1700.1-42).





VIII. Evidence-based Scientific References

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IX. Policy History

Action Date	Action
June 01, 2023	Policy created
December 03, 2024	Policy approved by Medical Directors
December 20, 2024	Policy approved at UMC
February 01, 2025	Policy effective date following notification period