

GENETIC TESTING: PHARMACOGENETICS

OVERVIEW

Pharmacogenetic tests are germline genetic tests that are developed to aid in assessing an individual's response to a drug treatment or to predict the risk of toxicity from a specific drug treatment. Testing may be performed prior to initiation of treatment to identify if an individual has genetic variants that could either affect response to a particular drug and/or increase the risk of adverse drug reactions. Testing may also be performed during treatment to assess an individual who has had an adverse drug reaction or to assess response to treatment. Test methodology includes genotyping and single nucleotide variant testing.

POLICY REFERENCE TABLE

The tests and associated laboratories and CPT codes contained within this document serve only as examples to help users navigate claims and corresponding coverage criteria; as such, they are not comprehensive and are not a guarantee of coverage or non-coverage. Please see the [Concert Genetics Platform](#) for a comprehensive list of registered tests.

| Coverage Criteria Sections | Example Tests (Labs) | Common CPT Codes | Common ICD Codes | Ref |
|---|---|------------------|---|---------------------|
| Pharmacogenetic Panel Tests | GeneSight Psychotropic (Myriad Genetics) | 0345U | B20, C00.0-C96.9, D00.0-D49.9, E75.22, F01-F99, G10, G71.14, G89.0-G89.4, I20.0, I21.01-I22.9, I24.1, I25.110, I26.01-I26.99, I48.0, I60.00-I66.99, I73, I82.210-I82.91, K50.00-K50.019 | 1, 2, 3, 4, 5, 6 |
| | Professional PGX (formerly Genecept Assay) (Genomind) | 81418 | | |
| | PGxOne (Admera Health) | | | |
| | Genomind Professional PGX Express CORE | 0175U | | |

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|---|--|----------------------------|---|---|
| | Cytochrome P450 Genotyping Panel (ARUP Laboratories) | 81418 | K51.00-K51.319, R52, R79.9, T46.6X1A-T46.6X6S, Z13.71-Z13.79, Z80.3, Z81.8, Z82.49, Z85.3, Z86.000, Z86.59, Z86.71-Z86.79 | |
| | OneOme RightMed Pharmacogenomic Test (OneOme) | 0347U, 0348U, 0349U, 0350U | | |
| | Focused Pharmacogenomics Panel (Mayo Clinic Laboratories) | 0029U | | |
| | Psych HealthPGx Panel, (RPRD Diagnostics) | 0173U | | |
| | CNT Genotyping Panel (RPRD Diagnostics) | 0286U | | |
| | PersonalisedRX (Lab Genomics LLC) | 0380U | | |
| | Serotonin Receptor Genotype (HTR2A and HTR2C), (Mayo Medical Laboratories) | 0033U | | |
| | EffectiveRX Comprehensive Panel (GENETWORx) | 0438U | | |
| | RightMed Gene Test Exclude F2 and F5 (OneOme LLC) | 0434U | | |
| | Genomind Pharmacogenetics Report (Genomind, Inc) | 0423U | | |
| | Tempus nP (Tempus) | 0419U | | |
| | IDgenetix (Castle Biosciences) | 0411U | | |
| | Medication Management Neuropsychiatric Panel (RCA Laboratory) | 0392U | | |
| <u>Pharmacogenetic Single Gene Tests</u> | | | | |
| BCHE Variant Analysis | BCHE Single Gene Test (Blueprint Genetics) | 81479 | Z01.81, Z01.810, Z01.811, Z01.818, | 8 |

| | | | | |
|--|---|-------|--|------|
| | | | Z01.89 | |
| CYP2C9 Variant Analysis | Cytochrome P450 2C9 Genotype (Quest Diagnostics) | 81227 | E78.00, E78.1, G35, I21.0-I22.9, I26.01-I26.99, I48.0, I60.00-I66.99, I82.210-I82.91, Z86.71-Z86.79 | 8 |
| CYP2C19 Variant Analysis | CYP2C19 Single Gene Test (Blueprint Genetics) | 81225 | C64, F32, I21.0-I22.9, I24.9, I26.01-I26.99, I48.0, I60.00-I66.99, I82.210-I82.91, K21.9, L20, Q85.83, R56.9, R68.82, Z86.71-Z86.79 | 8 |
| CYP2D6 Variant Analysis | CYP2D6 (ARUP Laboratories) | 81226 | C50.011-C50.929, C79.81, D05.00-D05.92, D07.30-D07.39, E11.9, E75.22, F11, F20.9, F31, F33, F84.0, F90, F95.2, G10, G24, G47.419, I10, I20.0, I21.01-I22.9, I24.1, I25.110, I48, I63.50-I63.549, I66.01-I66.9, I73, K21.9, R42, R52, T75.3, Z13.71-Z13.79, Z80.3, Z85.3, Z86.000 | 7, 8 |
| | CYP2D6 Common Variants and Copy Number (Mayo Clinic Laboratories) | 0070U | | |
| | CYP2D6 Full Gene Sequencing (Mayo Clinic Laboratories) | 0071U | | |
| | CYP2D6-2D7 Hybrid Gene Targeted Sequence Analysis (Mayo Clinic Laboratories) | 0072U | | |
| | CYP2D7-2D6 Hybrid Gene Targeted Sequence Analysis (Mayo Clinic Laboratories) | 0073U | | |
| | CYP2D6 CYP2D6 Nonduplicated Gene Analysis (Mayo Clinic Laboratories) | 0074U | | |
| | CYP2D6 5' gene duplication/multiplication targeted sequence analysis (Mayo Clinic Laboratories) | 0075U | | |
| | CYP2D6 3' gene | 0076U | | |

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|--|--|--------------|---|---|
| | duplication/multiplication targeted sequence analysis (Mayo Clinic Laboratories) | | | |
| CYP3A5 Variant Analysis | CYP3A5 single gene test (Blueprint Genetics) | 81231 | T86, Z79.6, Z94 | 8 |
| CYP4F2 Variant Analysis | CYP4F2 Single Gene Test (Blueprint Genetics) | 81479 | I21.0-I22.9, I26.01-I26.99, I48.0, I60.00-I66.99, I82.210-I82.91, Z86.71-Z86.79 | 8 |
| DPYD Variant Analysis | DPD 5-Fluorouracil Toxicity (Labcorp) | 81232 | C00.0-C96.9, D00.0-D49.9 | 8 |
| HLA-B*15:02 Variant Analysis | HLA-B*15:02, Carbamazepine Sensitivity (Labcorp) | 81381 | G40 | 8 |
| HLA-B*57:01 Variant Analysis | HLA B*57:01 Abacavir Hypersensitivity (Labcorp) | 81381 | B20, Z21 | 8 |
| NAT2 Variant Analysis | NAT2 single gene test (Blueprint Genetics) | 81479 | G73, M35.9 | 8 |
| TPMT and NUDT15 Variant Analysis | Thiopurine S-Methyltransferase (<i>TPMT</i>) Genotype (Quest Diagnostics) | 81335 | C91.0, K50.00-K50.90, K51.00-K51.319, M35.9, M05-M06.9, C85.90 | 8 |
| | <i>TPMT</i> and <i>NUDT15</i> (ARUP Laboratories) | 81335, 81306 | | |
| | Thiopurine Methyltransferase (<i>TPMT</i>) and Nudix Hydrolase (<i>NUDT15</i>) Genotyping (Mayo Clinic Laboratories) | 0034U | | |
| | NT (<i>NUDT15</i> and <i>TPMT</i>) genotyping panel (RPRD Diagnostics) | 0169U | | |
| | UGT1A1 Irinotecan Toxicity (Labcorp) | 81350 | | |
| UGT1A1 Variant Analysis | | | | |

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|--|---|--------------|---|------|
| UGT2B17 Variant Analysis | UGT2B17 Single Gene (Fulgent Genetics) | 81479 | C25, C64, C71, C72, Q85.83 | 8 |
| VKORC1 Variant Analysis | VKORC1 Single Gene Test (Blueprint Genetics) | 81355 | I21.0-I22.9, I26.01-I26.99, I48.0, I60.00-I66.99, I82.210-I82.91, Z86.71-Z86.79 | 8 |
| Warfarin Sensitivity Analysis Panels | Warfarin Response Genotype (Mayo Medical Laboratories) | 0030U | I21, I26, I48 | 8, 9 |
| | Accutype Warfarin (Quest) | 81227, 81355 | | |
| Other Single Gene Variant Analysis | Catechol-O-Methyltransferase (COMT) Genotype (Mayo Clinic Laboratories) | 0032U | F01-F69, F80-F99, G20, Z81.8, Z86.59 | 8 |
| | COMT single gene test (Blueprint Genetics) | 81479 | | |
| | Cytochrome P450 1A2 Genotype (Mayo Clinic Laboratories) | 0031U | F01-F69, F80-F99, Z81.8, Z86.59 | |
| | CYP1A2 single gene test (Blueprint Genetics) | 81479 | | |
| | Cardio IQ KIF6 Genotype (Quest Diagnostics) | 81479 | E78.0-E78.9, R79.9, Z82.49 | |
| | Opioid Receptor, mu OPRM1 Genotype, 1 Variant (ARUP Laboratories) | 81479 | G89.0-G89.4 | |
| | SLCO1B1, 1 Variant (ARUP Laboratories) | 81328 | E78.00-E78.5, G71.14, R79.9, T46.6X1A-T46.6X6S, Z82.49 | |
| | TYMS Single Gene (Sequencing & Deletion/Duplication) (Fulgent Genetics) | 81479 | C00.0-C96.9, D00.0-D49.9 | |

OTHER RELATED POLICIES

This policy document provides coverage for tests that determine the dosage of or the selection of a specific drug based on pharmacogenetic testing. For other related testing, please refer to:

- **Oncology: Molecular Analysis of Solid Tumors and Hematologic Malignancies** for coverage criteria related to DNA testing of a solid tumor or a blood cancer.
- **Genetic Testing: Hematologic Conditions (non-cancerous)** for coverage criteria related to diagnostic testing for non-cancerous genetic blood disorders.
- **Genetic Testing: Multisystem Inherited Disorders, Intellectual Disability, and Developmental Delay** for coverage criteria related to diagnostic testing for cystic fibrosis, and related therapies.
- **Genetic Testing: Metabolic, Endocrine, and Mitochondrial Disorders** for coverage criteria related to *MTHFR* testing.
- **Genetic Testing: General Approach to Genetic and Molecular Testing** for coverage criteria related to pharmacogenetic testing that are not specifically discussed in this or other specific policies.

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COVERAGE CRITERIA

PHARMACOGENETIC PANEL TESTS

- I. The use of pharmacogenetic testing panels (81418, 0029U, 0033U, 0173U, 0175U, 0286U, 0345U, 0347U, 0348U, 0349U, 0350U, 0380U, 0392U, 0411U, 0419U, 0423U, 0434U, 0438U) is considered **investigational*** for all indications.

*See *HLA-B*15:02* and *HLA-A*31:01* Variant Analysis and *TPMT* and *NUDT15* Variant Analysis below for coverage criteria. These tests involve analysis of more than one gene, but are not considered experimental/investigational as a panel ("panel" defined as a genetic test analyzing more than one gene)

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PHARMACOGENETIC SINGLE GENE TESTS

BCHE Variant Analysis

- I. *BCHE* variant analysis (81479) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with either of the following:
 1. Mivacurium¹ (e.g., Mivacron), **OR**
 2. Succinylcholine¹ (e.g., Anectine, Suxamethonium).
- II. *BCHE* variant analysis (81479) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly used as a muscle relaxant during surgery or intubation.

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CYP2C9 Variant Analysis

- I. *CYP2C9* variant analysis (81227) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with any of the following:
 1. Siponimod¹ (e.g., Mayzent), **OR**
 2. Celecoxib² (e.g., Celebrex, Elyxyb), **OR**
 3. Dronabinol³ (e.g., Marinol, Syndros), **OR**
 4. Erdafitinib⁴ (e.g., Balversa), **OR**
 5. Flurbiprofen⁵ (e.g., Ansaid), **OR**
 6. Fosphenytoin⁶ (e.g., Cerebyx, Sesquient), **OR**
 7. Meloxicam⁷ (e.g., Anjeso, Mobic, Vivlodex, Qmiiz ODT), **OR**

8. Nateglinide⁸ (e.g., Starlix), **OR**
9. Phenytoin⁹ (e.g., Dilantin, Phenytek), **OR**
10. Piroxicam¹⁰ (e.g., Feldene), **OR**
11. Warfarin¹¹ (e.g., Coumadin, Jantoven).

II. *CYP2C9* variant analysis (81227) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed for individuals diagnosed with multiple sclerosis

² Commonly prescribed for treating pain or inflammation

³ Commonly prescribed for treating loss of appetite and severe nausea and vomiting

⁴ Commonly prescribed for treatment of bladder cancer

⁵ Commonly prescribed for treatment of pain or inflammation

⁶ Commonly prescribed for preventing or controlling seizures

⁷ Commonly prescribed for treating pain, inflammation, or severe pain

⁸ Commonly prescribed for blood sugar control in individuals with type II diabetes

⁹ Commonly prescribed for treatment of seizures

¹⁰ Commonly prescribed to treat pain or inflammation

¹¹ Commonly prescribed to reduce the formation of blood clots

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CYP2C19 Variant Analysis

I. *CYP2C19* variant analysis (81225) to determine drug metabolizer status is considered **medically necessary** when:

A. The member is being considered for or is currently undergoing treatment with any of the following:

1. Clopidogrel¹ (e.g., Plavix), **AND**

a) The member meets all of the following:

- (1) Will be undergoing percutaneous coronary intervention (PCI), **AND**
- (2) Has acute coronary syndromes (ACS), **AND**
- (3) Is at high risk for poor outcomes (e.g., urgent PCI for an ACS event, elective PCI for unprotected left main disease or last patent coronary artery), **OR**

2. Abrocitinib² (e.g., Cibinqo), **OR**
3. Belzutifan³ (e.g., Welireg), **OR**
4. Brivaracetam⁴ (e.g., Briviact, Brivajoy), **OR**
5. Citalopram⁵ (e.g., Celexa), **OR**
6. Cobazam⁶ (e.g., Onfi), **OR**
7. Flibanserin⁷ (e.g., Addyi), **OR**
8. Pantoprazole⁸ (e.g., Protonix).

II. *CYP2C19* variant analysis (81225) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed after a angina or cardiac arrest to lower risk of stroke and blood clots

² Commonly prescribed for eczema

³ Commonly prescribed to treat tumors in individuals with Von Hippel-Lindau syndrome

⁴ Commonly prescribed to treat seizures

⁵ Commonly prescribed for treatment of depression and major depressive disorder

⁶ Commonly prescribed for treatment of seizures caused by Lennox-Gastaut syndrome

⁷ Commonly prescribed for low libido in pre-menopausal women

⁸ Commonly prescribed for treatment of erosive esophagitis caused by GERD, and Zollinger-Ellison syndrome

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CYP2D6 Variant Analysis

- I. CYP2D6 variant analysis (81226, 0070U, 0071U, 0072U, 0073U, 0074U, 0075U, 0076U) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with any of the following:
 1. Eliglustat¹ (e.g., Cerdelga), **OR**
 2. Tetrabenazine² (e.g., Xenazine), **OR**
 3. Amphetamine³ (e.g., Adzenys, Dyanavel, Evekeo), **OR**
 4. Aripiprazole⁴ (e.g., Abilify, Abilify Maintena), **OR**
 5. Aripiprazole lauroxil⁵ (e.g., Aristada), **OR**
 6. T Atomoxetine⁶ (e.g., Strattera), **OR**
 7. Brexpiprazole⁷ (e.g., Rexulti), **OR**
 8. Clozapine⁸ (e.g., Versacloz, FazaClo, Clozaril), **OR**
 9. Deutetrabenazine⁹ (e.g., Austedo), **OR**
 10. Gefitinib¹⁰ (e.g., Iressa), **OR**
 11. Iloperidone¹¹ (e.g., Fanapt), **OR**
 12. Lofexidine¹² (e.g., Lucemyra), **OR**
 13. Meclizine¹³ (e.g., Antivert, Bonine, Dramamine, Verticalm, Zentrip), **OR**
 14. Metoclopramide¹⁴ (e.g., Reglan), **OR**
 15. Oliceridine¹⁵ (e.g., Olinvyk), **OR**
 16. Pimozide¹⁶ (e.g., Orap), **OR**
 17. Pitolisant¹⁷ (e.g., Wakix), **OR**

- 18. Propafenone¹⁸ (e.g., Rythmol), **OR**
- 19. Thioridazine¹⁹ (e.g., Mellaril), **OR**
- 20. Tramadol²⁰ (e.g., ConZip, Ultram), **OR**
- 21. Valbenazine²¹ (e.g., Ingrezza), **OR**
- 22. Venlafaxine²² (e.g., Effexor), **OR**
- 23. Vortioxetine²³ (e.g., Trintellix, Brintellix), **OR**
- 24. Codeine²⁴

- II. *CYP2D6* variant analysis (81226, 0070U, 0071U, 0072U, 0073U, 0074U, 0075U, 0076U) to determine drug metabolizer status is considered **investigational** for all other indications, including:
 - A. For the purpose of managing treatment with tamoxifen for women at high risk for or with breast cancer.

¹ Commonly prescribed for treatment of Gaucher disease

² Commonly prescribed for treatment of involuntary movements (chorea) caused by Huntington disease

³ Commonly prescribed for treatment of hyperactivity, impulse control, and attention deficit hyperactivity disorder (ADHD)

⁴ Commonly prescribed for schizophrenia, bipolar I disorder, and major depressive disorder

⁵ Commonly prescribed for schizophrenia

⁶ Commonly prescribed for treatment of attention deficit hyperactivity disorder (ADHD)

⁷ Commonly prescribed for treatment of schizophrenia and major depressive disorder

⁸ Commonly prescribed for treatment of schizophrenia

⁹ Commonly prescribed for treatment of involuntary muscle movements (chorea) caused by Huntington disease, and tardive dyskinesia

¹⁰ Commonly prescribed for treatment of non-small cell lung cancer

¹¹ Commonly prescribed for treatment of schizophrenia

¹² Commonly prescribed for treatment of opioid withdrawal symptoms

- 13 Commonly prescribed for treatment of motion sickness and vertigo
- 14 Commonly prescribed for treatment of heartburn caused by GERD, gastroparesis, nausea and vomiting, and to aid in certain medical procedures involving the stomach or intestines
- 15 Commonly prescribed for treatment of severe pain
- 16 Commonly prescribed for treatment of Tourette's syndrome
- 17 Commonly prescribed for treatment of excessive daytime sleepiness or sudden loss of muscle strength (cataplexy) related to narcolepsy
- 18 Commonly prescribed for treatment of heart rhythm disorders
- 19 Commonly prescribed for treatment of schizophrenia
- 20 Commonly prescribed for treatment of moderate to severe pain
- 21 Commonly prescribed for treatment of tardive dyskinesia
- 22 Commonly prescribed for treatment of major depressive disorder, anxiety, and panic disorder
- 23 Commonly prescribed for treatment of major depressive disorder
- 24 Commonly prescribed for treatment of mild to moderately severe pain, and to help reduce coughing

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CYP3A5 Variant Analysis

- I. CYP3A5 variant analysis (81231) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with tacrolimus¹ (e.g., Protopic, Envarsus, Astagraf, Prograf).
- II. CYP3A5 variant analysis (81231) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed to individuals who have undergone a heart, kidney, liver, or lung transplant

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CYP4F2 Variant Analysis

- I. *CYP4F2* variant analysis (81479) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with warfarin¹ (e.g., Coumadin, Jantoven).
- II. *CYP4F2* variant analysis (81479) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed to reduce the formation of blood clots

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DPYD Variant Analysis

- I. *DPYD* variant analysis (81232) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with either of the following:
 1. Fluorouracil¹ (e.g., Adrucil), **OR**
 2. Capecitabine¹ (e.g., Xeloda).
- II. *DPYD* variant analysis (81232) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed for individuals diagnosed with colorectal, breast, and aerodigestive tract tumors

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HLA-B*15:02 Variant Analysis

- I. *HLA-B*15:02* variant analysis (81381) to determine drug metabolizer status is considered **medically necessary** when:

- A. The member is being considered for or is currently undergoing treatment with any of the following:
1. Carbamazepine containing therapy¹ (e.g., Tegretol, Carbatrol, Eptol, Equetro), **OR**
 2. Phenytoin² (e.g., Dilantin, Phenytek), **OR**
 3. Fosphenytoin² (e.g., Cerebyx, Sesquient).
- II. *HLA-B*15:02* variant analysis (81381) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed for individuals with epilepsy, trigeminal neuralgia, or bipolar disorder

² Commonly prescribed for treatment of seizures

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***HLA-B*57:01* Variant Analysis**

- I. *HLA-B*57:01* variant analysis (81381) to determine drug metabolizer status is considered **medically necessary** when:
- A. The member is being considered for or is currently undergoing treatment with abacavir¹ (e.g., Ziagen).
- II. *HLA-B*57:01* variant analysis (81381) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed for individuals with HIV

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***NAT2* Variant Analysis**

- I. *NAT2* variant analysis (81479) to determine drug metabolizer status is considered **medically necessary** when:
- A. The member is being considered for or is currently undergoing treatment with amifampridine/amifampridine phosphate¹ (e.g., Firdapse, Ruzurgi).

- II. *NAT2* variant analysis (81479) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed for treatment of Lambert-Eaton myasthenic syndrome

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***TPMT* and *NUDT15* Variant Analysis**

- I. *TPMT* and *NUDT15* variant analysis (81306, 81335, 0034U, 0169U) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with any of the following:
 - 1. Azathioprine¹ (e.g., Imuran and Azasan), **OR**
 - 2. Mercaptopurine² (e.g., Purinethol and Purixan), **OR**
 - 3. Thioguanine³ (e.g., Tabloid), **OR**
 - B. The member is on thiopurine therapy and has had abnormal complete blood count results that do not respond to dose reduction.
- II. *TPMT* and *NUDT15* variant analysis (81306, 81335, 0034U, 0169U) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed for treatment of avoiding rejection of a transplanted organ, and rheumatoid arthritis

² Commonly prescribed for treatment of acute lymphoblastic or lymphocytic leukemia

³ Commonly prescribed for treatment of acute nonlymphocytic leukemia

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***UGT1A1* Variant Analysis**

- I. *UGT1A1* variant analysis (81350) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with any of the following:

1. Irinotecan¹ (e.g., Onivyde, Camptosar), **OR**
 2. Belinostat² (e.g., Beleodaq), **OR**
 3. Sacituzumab govitecan-hziy³ (e.g., Trodelvy).
- II. *UGT1A1* variant analysis (81350) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed for treatment of colon and rectal cancers

² Commonly prescribed for treatment of peripheral T-cell lymphoma

³ Commonly prescribed for treatment of breast and urothelial cancers

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***UGT2B17* Variant Analysis**

- I. *UGT2B17* variant analysis (81479) to determine drug metabolizer status is **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with belzutifan¹ (e.g., Welireg).
- II. *UGT2B17* variant analysis (81479) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed to treat tumors in individuals with Von Hippel-Lindau syndrome

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***VKORC1* Variant Analysis**

- I. *VKORC1* variant analysis (81355) to determine drug metabolizer status is considered **medically necessary** when:
 - A. The member is being considered for or is currently undergoing treatment with warfarin¹ (e.g., Coumadin, Jantoven).

- II. *VKORC1* variant analysis (81355) to determine drug metabolizer status is considered **investigational** for all other indications.

¹ Commonly prescribed to reduce the formation of blood clots

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Warfarin Sensitivity Analysis Panels

- I. Multigene panel analysis to determine drug metabolizer status for warfarin¹ sensitivity (81227, 81355, 0030U) is considered **medically necessary** when:
 - A. The member is undergoing prophylaxis and treatment of venous thrombosis or pulmonary embolism, **OR**
 - B. The member is undergoing prophylaxis and treatment of thromboembolic complications associated with atrial fibrillation and/or cardiac valve replacement, **OR**
 - C. The member has a history of previous myocardial infarction, **AND**
 - D. The member is being considered for or is undergoing treatment with warfarin, **AND**
 - 1. The member has not reached a therapeutic dose.
- II. Multigene panel analysis to confirm drug metabolizer status for warfarin¹ sensitivity (81227, 81355, 0030U) is considered **investigational** for all other indications.

¹ Commonly prescribed to reduce the formation of blood clots

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Other Single Gene Variant Analysis

- I. Variant analysis of all other genes for drug metabolizer status is considered **investigational**, including but not limited to:
 - A. *COMT* (0032U, 81479)

- B. *CYP1A2* (0031U, 81479)
- C. *KIF6* (81479)
- D. *OPRM1* (81479)
- E. *SLCO1B1* (81328)
- F. *TYMS* (81479)

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BACKGROUND AND RATIONALE

Pharmacogenetic Panel Testing

There are no professional society guidelines that address the clinical utility of large pharmacogenetic testing panels for the general population or for a specific population. The US Food and Drug Administration (FDA) also does not address the usage of pharmacogenetic panels.

There are several recent studies that investigated the usefulness of pharmacogenetic panels [for example, Greden et al (2019), Perlis et al (2020), Shan et al (2019), Tiwari et al (2022), Oslin (2022)]. However, these studies had different designs and often conflicting results regarding clinical utility, making it difficult to determine whether there is clinical utility for these types of tests.

A rapid review and meta-analysis by Bunka et al (2023) of 10 randomized controlled trials to evaluate pharmacogenomic-guided care for major depression showed that, while there is likely beneficial effects to adults with moderate to severe major depressive disorder utilizing pharmacogenomic panels, there is “very low certainty in the magnitude of effect.” (p. 1) This analysis also noted the “high risk of bias and inconsistency between trials.” (p. 1)

There are several single gene pharmacogenetic tests in which the FDA describes the clinical utility of the test results for a given gene/drug/testing indication. These are outlined below.

BCHE Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations, which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, lists the following recommendations for *BCHE*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|------------------------|------|-----------------------------------|--|
| Mivacurium | BCHE | intermediate or poor metabolizers | Results in higher systemic concentrations and higher adverse reaction risk (prolonged neuromuscular blockade). Avoid use in poor metabolizers. |
| Succinylcholine | BCHE | intermediate or poor metabolizers | Results in higher systemic concentrations and higher adverse reaction risk (prolonged neuromuscular blockade). Avoid use in poor metabolizers. May administer a test dose to assess sensitivity and administer cautiously via slow infusion. |

CYP2C9 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for *CYP2C9*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|-------------------|--------|----------------------------------|--|
| Celecoxib | CYP2C9 | poor metabolizers or *3 carriers | Results in higher systemic concentrations. Reduce starting dose to half of the lowest recommended dose in poor metabolizers. Consider alternative therapy in poor metabolizers with juvenile rheumatoid arthritis. |
| Dronabinol | CYP2C9 | intermediate or poor | May result in higher systemic |

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| | | metabolizers | concentrations and higher adverse reaction risk. Monitor for adverse reactions. |
| Erdafitinib | CYP2C9 | *3/*3 (poor metabolizers) | May result in higher systemic concentrations and higher adverse reaction risk. Monitor for adverse reactions. |
| Flurbiprofen | CYP2C9 | poor metabolizers or *3 carriers | Results in higher systemic concentrations. Use a reduced dosage in poor metabolizers. |
| 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. |
| Meloxicam | CYP2C9 | poor metabolizers or *3 carriers | Results in higher systemic concentrations. Consider dose reductions in poor metabolizers. Monitor patients for adverse reactions. |
| 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 for adverse reactions. Refer to FDA labeling for specific dosing |

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| | | | recommendations. |
| 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. |
| Piroxicam | CYP2C9 | intermediate or poor metabolizers | Results in higher systemic concentrations. Consider reducing dosage in poor metabolizers. |
| 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. |
| 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. |

CYP2C19 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for CYP2C19:

| 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. |
| 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. |
| Brivaracetam | CYP2C19 | intermediate or poor metabolizers | Results in higher systemic concentrations and higher adverse reaction risk. Consider dosage reductions in poor metabolizers. |
| Citalopram | CYP2C19 | poor metabolizers | Results in higher systemic concentrations and adverse reaction risk (QT prolongation). The maximum recommended dose is 20 mg. |
| 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. |
| Flibanserin | CYP2C19 | poor metabolizers | May result in higher systemic concentrations and higher adverse reaction risk. Monitor patients for adverse reactions. |
| Pantoprazole | CYP2C19 | intermediate or poor metabolizers | Results in higher systemic concentrations. Consider dosage reduction in children who are poor |

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| | | | metabolizers. No dosage adjustment is needed for adult patients who are intermediate or poor metabolizers. |
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CYP2D6 Variant Analysis

National Comprehensive Cancer Network (NCCN)

NCCN Breast Cancer guidelines (1.2024) recommend against CYP2D6 genotype testing for women being considered for tamoxifen treatment. (p. DCIS-2 and p. BINV-K)

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for CYP2D6:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|----------------------------------|--------|--------------------|--|
| Amphetamine | CYP2D6 | poor metabolizers | May affect systemic concentrations and adverse reaction risk. Consider a lower starting dosage or use an 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. |

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| 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. |
| Brexiprazole | CYP2D6 | poor metabolizers | Results in higher systemic concentrations. Dosage adjustment is recommended. Refer to FDA labeling for specific dosing recommendations. |
| 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). |

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| 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. |
| Gefitinib | CYP2D6 | poor metabolizers | Results in higher systemic concentrations and higher adverse reaction risk. Monitor for adverse reactions. |
| Iloperidone | CYP2D6 | poor metabolizers | Results in higher systemic concentrations and higher adverse reaction risk (QT prolongation). Reduce dosage by 50%. |
| 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. |

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| 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. |
| Oliceridine | CYP2D6 | poor metabolizers | Results in higher systemic concentrations and higher adverse reaction risk (respiratory depression and sedation). May require less frequent dosing. |
| 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. |
| Pitolisant | CYP2D6 | poor metabolizers | Results in higher systemic concentrations. Use the lowest recommended starting dosage. Refer 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. |
| Tetrabenazine | CYP2D6 | poor metabolizers | Results in higher systemic concentrations. The maximum recommended single dose is 25 mg and should not exceed 50 mg/day. |

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| 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. |

CYP3A5 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations, which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, lists the following recommendations for *CYP3A5*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|------------|--------|-------------------------------------|--|
| Tacrolimus | CYP3A5 | intermediate or normal metabolizers | Results in lower systemic concentrations, lower probability of achieving target concentrations and may result in higher rejection risk. Measure drug concentrations and adjust dosage based on trough whole blood tacrolimus concentrations. |

CYP4F2 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for *CYP4F2*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|----------|--------|------------------------|--|
| Warfarin | CYP4F2 | V433M variant carriers | May affect dosage requirements. Monitor and adjust doses based on INR. |

DPYD Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for *DPYD*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|--------------|------|-----------------------------------|--|
| 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 |

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| | | | available to recommend a dosage in intermediate metabolizers. Withhold or discontinue in the presence of early-onset or unusually severe toxicity. |
| 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. |

HLA-B*15:02 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for *HLA-B*15:02*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|----------------------|-------------|---------------------------|--|
| 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. |
| 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 |

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| | | | HLA-B*15:02. 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. |

HLA-B*57:01 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for *HLA-B*57:01*:

| 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. |

NAT2 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations, which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, lists the following recommendations for *NAT2*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|-------------|-------------|---------------------------|---|
|-------------|-------------|---------------------------|---|

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|--------------------------------|------|-------------------|--|
| 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. |

***TPMT* and *NUDT15* Variant Analysis**

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for *TPMT* and *NUDT15*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|---------------------|--------------------|-----------------------------------|--|
| 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 |

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|-----------------------|--------------------|-----------------------------------|---|
| | | | 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. |
| 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 |

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| | | | FDA labeling for specific dosing recommendations. |
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UGT1A1 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for *UGT1A1*:

| 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/m ² in poor metabolizers. |
| Irinotecan | UGT1A1 | *1/*6, *1/*28 (intermediate metabolizers) or *6/*6, *6/*28, *28/*28 (poor metabolizers) | Results in higher systemic active metabolite concentrations and higher adverse reaction risk (severe or life-threatening neutropenia, severe diarrhea). Closely monitor for neutropenia during and after treatment. Consider reducing the starting dosage by at least one level in poor metabolizers and modify the dosage based on individual patient tolerance. Refer to FDA labeling for specific dosing recommendations. |
| 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. |

UGT2B17 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations, which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, lists the following recommendations for *UGT2B17*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|-------------------|------------------------------|--------------------|--|
| 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. |

VKORC1 Variant Analysis

Food and Drug Administration (FDA)

The FDA published a Table of Pharmacogenetic Associations which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, list the following recommendations for *VKORC1*:

| Drug | Gene | Affected Subgroups | Description of Gene-Drug Interaction |
|-----------------|--------|---------------------------|---|
| 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. |

Warfarin Sensitivity Analysis Panels

Food and Drug Administration (FDA)

Per the FDA label, the indications and usage for Warfarin include the following:

- Prophylaxis and treatment of venous thrombosis and its extension, pulmonary embolism
- Prophylaxis and treatment of thromboembolic complications associated with atrial fibrillation and/or cardiac valve replacement

- Reduction in the risk of death, recurrent myocardial infarction, and thromboembolic events such as stroke or systemic embolization after myocardial infarction

The FDA published a Table of Pharmacogenetic Associations, which details possible gene-drug interactions. Section 1, entitled Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations, lists the following recommendations for *CYP2C9*, *CYP4F2* and *VKORC1*:

| | | | |
|-----------------|--------|-----------------------------------|---|
| 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. |
| | CYP4F2 | V433M variant carriers | May affect dosage requirements. Monitor and adjust doses based on INR. |
| | 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. |

Other Single Gene Variant Analysis

The Food and Drug Administration (FDA) does not list *COMT*, *CYP1A2*, *KIF6*, *OPRM1*, *SLCO1B1*, or *TYMS* in Section 1 of the Table of Pharmacogenetic Associations (“Pharmacogenetic Associations for which the Data Support Therapeutic Management Recommendations”).

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