



Patient:	Nora Syke	Sample ID:	P000034875
Patient DOB:	7/12/1977	Accession ID:	255836
Ordering Clinician:	August Edwards	Sample Collection Date:	7/20/2026
Sample Type:	Buccal	Sample Received Date:	7/21/2026
Assay Ordered:	PGx (v3.2)	Report Date:	7/21/2026 2:20 PM

Understanding this report:

1. Current medications that have a higher risk drug-drug or drug-gene interaction
2. Pharmacogenetic (PGx) risk categorization of common medications across different conditions
3. Gene result overview

Actionable interactions

Usually have specific recommendations based on PGx (dose adjustment, alternative drug considerations or increased monitoring)



Higher Risk: large changes in drug blood levels



Contraindication



Responsible for major drug-drug interactions

No to moderate interactions

These drugs have lower PGx risk and are less likely to require intervention due to pharmacogenetics



No detected interactions



Lower Risk: small changes in drug blood levels



Moderate Risk: moderate changes in drug blood levels

Legend:

Additional information about the nature of the gene-drug interaction



Increased Efficacy



Decreased Efficacy



Weight Gain



Actionable PGx Guidelines



Therapeutic Duplication

Contact us: **877-895-8658** • vantadiagnostics.com • pgxsupport@vantadx.com

Disclaimer: This summary page sorts medications based solely on pharmacogenetic parameters and drug category. Pharmacogenetics is only one factor to consider when prescribing medications and you, as the prescriber, are ultimately responsible for evaluating other patient characteristics. This includes, but is not limited to, proper diagnosis, comorbidities, drug-drug interactions, medical history, contraindications, and other factors. Never rely solely on PGx to make any medication decisions. This test was developed, and its performance characteristics determined by Vanta Diagnostics. It has not been cleared or approved by the US Food and Drug Administration.

Electronically signed by: Juel Meya, Quality Assurance

1. Current risk medications

The following medications were listed as current on the patient requisition form and have a higher risk drug-drug or drug-gene interaction. We have identified up to 5 alternative medications, ranked by their interaction risk.

RISK LEVEL	MEDICATION ASSESSMENT	POSSIBLE ALTERNATIVES (Based on USP classification)	
	ZOLOFT Potential increase in ZOLOFT serum levels: May require lower doses Due to: 2C19 Poor metabolizer		fluvoxamine
			Citalopram
			Escitalopram
			FLUoxetine
			PARoxetine

Additional medications the patient is taking that have no to moderate interactions:

 **Codeine**

This medication list was provided on 7/20/2026.

2. PGx risk categorization

Actionable gene-drug interactions

Antidepressants (1st line)

 Citalopram (Celexa®) 	Escitalopram (Lexapro®) 
Sertraline (Zoloft®) 	

Antidepressants (2nd line or augmentation)

 Amitriptyline (Elavil®) 	Clomipramine (Anafranil®) 
Doxepin (Sinequan®) 	Imipramine (Tofranil®) 

Anti-anxiety medications (1st line)

 Citalopram (Celexa®) 	Escitalopram (Lexapro®) 
Sertraline (Zoloft®) 	

No to moderate gene-drug interactions

Antidepressants (1st line)

 Bupropion (Wellbutrin®)	Duloxetine (Cymbalta®)
Fluvoxamine (Luvox®)	Mirtazapine (Remeron®)
Paroxetine (Paxil®)	Vortioxetine (Trintellix®)
 Fluoxetine (Prozac®)	
 Venlafaxine (Effexor®)	

Antidepressants (2nd line or augmentation)

 L-methylfolate (Deplin®) 	Cariprazine (Vraylar®)
Desipramine (Norpramin®)	Desvenlafaxine (Pristiq®)
Dextromethorphan/ Bupropion (Auvelity®)	Levomilnacipran (Fetzima®)
Nortriptyline (Pamelor®)	Selegeline Transdermal (Emsam®)
Trazodone (Desyrel®)	Vilazodone (Viibryd®)
Aripiprazole (Abilify®) 	Brexipiprazole (Rexulti®) 
Quetiapine (Seroquel®) 	

Anti-anxiety medications (1st line)

 Buspirone (Buspar®)	Duloxetine (Cymbalta®)
Fluvoxamine (Luvox®)	Hydroxyzine (Vistaril®)
Paroxetine (Paxil®)	
 Fluoxetine (Prozac®)	
 Venlafaxine (Effexor®)	

Anti-anxiety medications (2nd line)

 **Diazepam** (Valium®) **Amitriptyline** (Elavil®) 

Clomipramine (Anafranil®)  **Doxepin** (Sinequan®) 

ADHD

No major interactions.

Statin

 **Fluvastatin** (Lescol®)  **Rosuvastatin** (Crestor®) 

B-blocker

No major interactions.

Non-opioid Analgesics

 **Celecoxib** (Celebrex®)  **Flurbiprofen** (Ansaid®) 

Ibuprofen (Motrin®)  **Meloxicam** (Mobic®) 

Piroxicam (Feldene®) 

Anti-anxiety medications (2nd line)

 **Alprazolam** (Xanax®) **Clonazepam** (Klonopin®)

Lorazepam (Ativan®) **Nortriptyline** (Pamelor®)

Pregabalin (Lyrica®) **Quetiapine** (Seroquel®) 

ADHD

 **Amphetamine-Dextroamphetamine** (Adderall®) **Atomoxetine** (Strattera®)

Clonidine ER (Kapvay®) **Dextroamphetamine** (Dexedrine®)

Guanfacine ER (Intuniv®) **Lisdexamfetamine** (Vyvanse®)

Viloxazine (Qelbree®) **Dexmethylphenidate** (Focalin®) 

Methylphenidate (Ritalin®) 

Statin

 **Lovastatin** (Mevacor®) **Pitavastatin** (Livalo®)

Pravastatin (Pravachol®) **Simvastatin** (Zocor®)

 **Atorvastatin** (Lipitor®)

B-blocker

 **Atenolol** (Tenormin®) **Bisoprolol** (Ziac®)

Carvedilol (Coreg®) **Metoprolol** (Toprol®)

Nebivolol (Bystolic®)

Non-opioid Analgesics

 **Acetaminophen** (Tylenol®) **Aspirin** (Durlaza®)

Ketorolac (Toradol®) **Nabumetone** (Relafen®)

 **Diclofenac** (Voltaren®) **Indomethacin** (Indocin®)

Naproxen (Aleve®)

Opioids

No major interactions.

Opioids

 Codeine (Unbranded)	Fentanyl (Duragesic®)
Hydrocodone (Hysingla ER®)	Hydromorphone (Dilaudid®)
Morphine (MS Contin®)	Oxycodone (Oxycontin®)
Oxymorphone (Unbranded)	Tapentadol (Nucynta®)
Tramadol (Ultram®)	

Proton Pump Inhibitors

 Dexlansoprazole (Dexilant®) 	Lansoprazole (Prevacid®) 
Omeprazole (Prilosec®) 	Pantoprazole (Protonix®) 

Proton Pump Inhibitors

 Esomeprazole (Nexium®)	Rabeprazole (Aciphex®)
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Antiplatelet

 Clopidogrel (Plavix®) 
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Antiplatelet

 Prasugrel (Effient®)	Ticagrelor (Brilinta®)
Ticlopidine (Unbranded)	

Anticoagulants

 Warfarin (Coumadin®) 

Anticoagulants

 Dabigatran (Pradaxa®)	Edoxaban (Savaysa®)
Rivaroxaban (Xarelto®)	
 Apixaban (Eliquis®)	

Genitourinary

No major interactions.

Genitourinary

 Darifenacin (Enablex®)	Fesoterodine (Toviaz®)
Flavoxate (Urispas®)	Mirabegron (Myrbetriq®)
Oxybutynin (Ditropan®)	Solifenacin (Vesicare®)
Tolterodine (Detrol®)	Trospium (Sanctura®)

Nausea / Vomiting

 **Dronabinol** (Marinol®) 

Nausea / Vomiting

- 
Aprepitant (Emend®)
- Dolasetron** (Anzemet®)
- Granisetron** (Sancuso®)
- Metoclopramide** (Reglan®)
- Ondansetron** (Zofran®)
- Palonosetron** (Aloxi®)
- Prochlorperazine** (Compazine®)
- Promethazine** (Promethegan®)
- Rolapitant** (Varubi®)

3. Gene results overview

Pharmacokinetic Genes (Drug Metabolism / Drug Absorption)	Gene	Genotype	Phenotype	Impact
	ABCB1	A/A	NF	Normal exposure is expected
	ABCB1 C3435T	G/G	NF	Normal exposure is expected
	ABCG2	T/T	PF	Increased exposure to certain medications
	CYP1A2	*1B/H7	NM	Normal metabolism is expected
	CYP2B6	*1/*1	NM	Normal metabolism is expected
	CYP2C19	*4A/*4A	PM	Risk of increased (↑) drug levels
	CYP2C9	*1/*3	IM	Risk of increased (↑) drug levels
	CYP2D6	*1/*1	NM	Normal metabolism is expected
	CYP3A4/5	*1/*1, *3/*3	NA	Normal metabolism is expected
	SLCO1B1	*1/*1	NF	Normal exposure is expected
	UGT1A4	*1a/*3b	UM	Risk of decreased (↓) drug levels
	UGT2B15	*1/*2	NM	Normal metabolism is expected

Pharmacodynamic Genes (Drug Targets / Mechanisms)	Antidepressant Response		
	Gene	Result	Impact
	SLC6A4	L(A)/S	Higher odds of gastrointestinal side effects with SSRIs in individuals of European descent
	BDNF	Val/Met	More pronounced effect to exercise; Possible higher odds of response to SNRIs
	HTR2A	G/A	No known significant clinical impact
	MTHFR	C677T: C/T A1298C: A/C	Reduced MTHFR activity and methylfolate production
	Attention-deficit/hyperactivity disorder Response		
	Gene	Result	Impact
	ADRA2A	C/C	Lower odds of response to methylphenidate for inattentive symptoms of ADHD
	COMT	Val/Met	No known significant clinical impact
	Antipsychotic Response and Tolerability		
	Gene	Result	Impact
	DRD2	C/C	No known significant clinical impact
	HTR2C	C/C	No known significant clinical impact
	MC4R	A/A	Higher risk of weight gain with certain 2nd generation antipsychotics
	Other		
	Gene	Result	Impact
	ANK3	C/C	No known significant clinical impact
	CACNA1C	G/G	No known significant clinical impact
	GRIK1	A/A	No known significant clinical impact
	HLA-A *31:01	Positive	Higher risk of skin reactions with carbamazepine
	HLA-B *15:02	Negative	No known significant clinical impact
	OPRM1	A/A	No known significant clinical impact

Test Methodology

This test was developed and performance characteristics were validated in the Vanta Diagnostics clinical laboratory. This test has not been cleared or approved by the U.S. Food and Drug Administration (FDA). This test is used for clinical purposes and should not be regarded as investigational or for research use. Vanta Diagnostics' laboratory is certified under the Clinical Laboratory Improvement Amendments (CLIA), as qualified to perform high complexity clinical laboratory testing. Vanta Diagnostics performed the testing using standard and custom TaqMan reagents for all variants. The test results are intended to be used as prognostic and not diagnostic and are not intended as the sole means for patient management decisions.

Test Methodology Limitations: Factors influencing the amount and quality of DNA extracted include but are not limited to the amount of buccal cells extracted, patient oral hygiene, collection technique, and the presence of dietary or microbial sources of nucleic acids and nucleases. DNA quality and quantity are subject to matrix dependent influences. PCR inhibitors, extraneous DNA and nucleic acid degrading enzymes are all factors which may affect the evaluation of assay results. Some single nucleotide polymorphism (SNP) assays are problematic due to multiple base repeats and other sequence aberrations, which may hinder proper amplification and analysis. DNA purity can influence the assay. SLC6A4 contains many polymorphisms, and the assay was developed and validated according to the current available scientific information. For pharmacogenetic tests, undetected genetic and/or non-genetic factors such as drug-drug interactions may impact the phenotype. In liver transplant recipients, certain genotypes of the donor liver may not be the same as those of the recipient. In these cases, it may be necessary to account for both the donor and recipient genotypes when evaluating drug metabolism genes. However, studies to date have been inconclusive as to the relative influence of the donor and recipient genotypes. This pharmacogenetic report is based on a current understanding of the clinical relevance of the variant identified, penetrance, phenotype predictions, and recurrence risks.

Variants tested include ABCB1 C3435T rs1045642; ABCB1 rs2032583; ABCG2 rs2231142, ADRA2A rs1800544; ANK3 rs10994336; BDNF rs6265; CACNA1C rs1006737; COMT rs4680; CYP1A2 *1B, *1C, *1D, *1E, *1F, *1K and *11; CYP2B6 *4, *5, *6 and *9; CYP2C19 *2, *3, *4, *5, *6, *7, *8, *9, *10, *17, and *35; CYP2C9 *2, *3, *4, *5, *6, *8, *11, *13, and *27; CYP2D6 *2, *3, *4, gene deletion (*5), gene duplication, *6, *7, *8, *9, *10, *11, *12, *14, *15, *17, *29 and *41; CYP3A4 *22; CYP3A5 *3, *6, *7; DRD2 rs1799732; GRIK1 rs2832407; HLA-B*15:02 presence and HLA-A*31:01 presence detected by qPCR; HTR2A rs7997012; HTR2C rs3813929; MC4R rs489693; MTHFR rs1801131 and rs1801133; OPRM1 rs1799971; SLC6A4 rs25531 and rs63749047; SLCO1B1*5, UGT2B15 rs1902023; and UGT1A4 rs2011425. Other known variants that are not listed are not detected and will not be included in the test report.

Important Notice

This report is based on pharmacogenomic (PGx) analysis performed by Vanta Diagnostics, a CLIA certified laboratory. The laboratory is responsible for the accuracy, validity, and interpretation of the test results. This report has been generated, formatted, or transmitted using a third party reporting and informatics platform ("Reporting Vendor"). The Reporting Vendor does not perform laboratory testing and does not independently verify, modify, or interpret raw laboratory data. Validation and interpretation of laboratory data are the sole responsibility of the testing laboratory director. The director has reviewed and approved this report.

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