

# RightMed® Gene Report

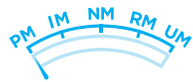
## Patient and report summary

Patient name: **John Doe**  
Patient date of birth: **1970-01-01**  
OneOme report date: **2021-01-06**

Ordering provider: **Sample Doctor**  
Ordering facility: **Healthcare Institution**  
Report type: **Original**

## Phenotype icon legend for CYP genes

CYP phenotype, or metabolizer status, is determined by the total predicted activity of the gene based on the genotype, and is represented by a gauge icon. Total predicted activity which falls between phenotypes will be reported as a range phenotype.



### Poor metabolizer

No to very low activity. Drugs converted to active metabolite(s) may have reduced efficacy. Active drugs converted to inactive metabolite(s) may cause side effects or toxicity.



### Intermediate metabolizer

Decreased activity. Drugs converted to active metabolite(s) may have reduced efficacy. Active drugs converted to inactive metabolite(s) may cause side effects or toxicity.



### Normal metabolizer

Normal level of activity. Drugs metabolized at a normal rate.



### Rapid metabolizer

Increased activity. Drugs converted to active metabolite(s) may cause side effects or toxicity. Active drugs converted to inactive metabolite(s) may lack efficacy.



### Ultrarapid metabolizer

Increased activity. Drugs converted to active metabolite(s) may cause side effects or toxicity. Active drugs converted to inactive metabolite(s) may lack efficacy.

## Phenotype icon legend for other genes

Phenotype icons for other genes represent the extent of impact of the genotype on protein activity, expression, or function, and/or observed clinical impact (e.g., adverse event risk).



### Atypical

Genotype indicates an absence of or major increase in protein activity, expression, or function.



### Atypical

Genotype indicates a moderate loss of or increase in protein activity, expression, or function.



### Typical

Genotype indicates normal or typical protein activity, expression, or function.

## Gene and phenotype summary

Gene	Genotype		Phenotype summary / Metabolic status
CYP1A2	*1A/*1F		<b>Rapid</b> Increased activity. Drugs converted to active metabolite(s) may have increased exposure. Active drugs converted to inactive metabolite(s) may have decreased exposure.
CYP2B6	*1/*5		<b>Intermediate to Normal</b> Decreased activity. Drugs converted to active metabolite(s) may have reduced exposure. Active drugs converted to inactive metabolite(s) may have increased exposure.
CYP2C9	*1/*3		<b>Intermediate</b> Decreased activity. Drugs converted to active metabolite(s) may have reduced exposure. Active drugs converted to inactive metabolite(s) may have increased exposure.
CYP2C19	*17/*17		<b>Ultrarapid</b> Increased activity. Drugs converted to active metabolite(s) may have increased exposure. Active drugs converted to inactive metabolite(s) may have decreased exposure.
CYP2C Cluster	rs12777823 GG		<b>Normal</b> CYP2C rs12777823 homozygous wild-type genotype consistent with normal clearance of a certain medication, independent of the impact of CYP2C9*2 and *3. CYP2C rs12777823, together with CYP4F2, CYP2C9, and VKORC1, may affect treatment management of a certain medication.
CYP2D6	*1/*1		<b>Normal</b> Normal activity. Drugs metabolized at a normal rate.
CYP3A4	*1/*1		<b>Normal</b> Normal activity. Drugs metabolized at a normal rate.
CYP3A5	*3/*3		<b>Poor</b> This CYP3A5 genotype is associated with the phenotype most prevalent in studies used to define standard dosing guidelines.
CYP4F2	*1/*1		<b>Normal activity</b> Genotype consistent with normal activity of the CYP4F2 enzyme, which catalyzes the metabolism of vitamin K, in counterpoint to the activity of VKORC1. CYP4F2, together with CYP2C9, VKORC1, and a variant in CYP2C Cluster, may affect treatment management of a certain medication.

## Gene and phenotype summary (cont.)

COMT	rs4680 GG		<p><b>High activity</b></p> <p>The COMT GG (Val/Val) genotype is predicted to yield higher COMT activity than the AA (Met/Met) or GA (Val/Met) genotypes at rs4680.</p>
DPYD	*1/*1		<p><b>DPD activity score: 2</b></p> <p>Genotype consistent with normal dihydropyrimidine dehydrogenase (DPD) activity with an activity score of 2, or a normal metabolizer phenotype.</p>
DRD2	rs1799978 GG		<p><b>Reduced receptor expression</b></p> <p>Homozygous variant dopamine receptor D2 (DRD2) rs1799978 GG genotype is consistent with reduced receptor expression.</p>
F2	rs1799963 GG		<p><b>Normal risk</b></p> <p>Normal risk of thrombosis associated with Factor II (prothrombin). Other genetic and clinical factors contribute to the risk for thrombosis.</p>
F5	rs6025 GG		<p><b>Normal risk</b></p> <p>Normal risk of thrombosis associated with Factor V. Other genetic and clinical factors contribute to the risk for thrombosis.</p>
GRIK4	rs1954787 CC		<p><b>Normal receptor function</b></p> <p>Glutamate ionotropic receptor kainate type subunit 4 (GRIK4) genotype is consistent with normal receptor function.</p>
HLA-A	Positive for *31:01		<p><b>Increased risk</b></p> <p>Positive for presence of the HLA-A*31:01 allele. Increased risk of hypersensitivity induced by a certain medication, and possibly others of structural similarity. Hypersensitivity and severe cutaneous reactions may occur regardless of the presence of the HLA-A*31:01 allele, in particular the presence of the HLA-B*15:02 allele is associated with severe cutaneous reactions induced by certain medications.</p>
HLA-B	Negative		<p><b>Normal risk</b></p> <p>Negative for presence of the HLA-B*15:02, HLA-B*57:01, and HLA-B*58:01 alleles. Normal risk of hypersensitivity, severe cutaneous reactions, and severe hepatotoxicity induced by certain medications. Hypersensitivity, severe cutaneous reactions, and severe hepatotoxicity may occur regardless of the presence of HLA-B*15:02, HLA-B*57:01, or HLA-B*58:01 alleles. In particular, the presence of the HLA-A*31:01 allele is associated with hypersensitivity reactions induced by a certain medication, and possibly other medications of structural similarity.</p>
HTR2A	rs7997012 AA		<p><b>Intron 2 genotype AA</b></p> <p>Homozygous wild-type HTR2A (5-hydroxytryptamine receptor 2A) genotype is consistent with normal HTR2A receptor function.</p>

## Gene and phenotype summary (cont.)

HTR2C	rs3813929 CC		<p><b>Normal influence</b></p> <p>Homozygous wild-type HTR2C [5-hydroxytryptamine (serotonin) receptor 2C] genotype is associated with a normal influence on weight gain related to certain medications. The HTR2C gene is located on the X chromosome. In patients with only one X, result should read rs3813929 C;-.</p>
IFNL4	rs12979860 CT		<p><b>Variant present</b></p> <p>Genotype consistent with a reduced likelihood of hepatitis C sustained virologic response (SVR) with certain treatment options.</p>
NUDT15	rs116855232 CC		<p><b>Normal risk</b></p> <p>Genotype consistent with normal NUDT15 activity and is not associated with an increased risk of thiopurine-induced toxicities. Reduced metabolizer phenotypes of TPMT are associated with an increased risk of thiopurine-induced toxicities, independent of NUDT15 activity.</p>
OPRM1	rs1799971 GG		<p><b>Asp/Asp isoform</b></p> <p>OPRM1 Asp/Asp (GG) genotype consistent with altered mu-1 opioid receptor function, and decreased sensitivity to the effects of certain substrates has been observed when compared to OPRM1 Asn/Asn (AA) and Asn/Asp (AG) genotypes at rs1799971. Decreased sensitivity has not been consistently observed in this genotype for all substrates that activate the mu-1 receptor.</p>
SLC6A4	L/L (La/La)		<p><b>Typical to increased expression</b></p> <p>Genotype consistent with a typical to increased expression of the SLC6A4 transporter compared to other genotypes. This genotype was shown to exhibit different phenotypes in East Asian populations, as opposite outcomes were observed for this genotype in East Asian populations when compared to Caucasian populations.</p>
SLCO1B1	*1A/*1A		<p><b>Normal function</b></p> <p>SLCO1B1 genotype consistent with normal function of the OATP1B1 transporter.</p>
TPMT	*1/*4		<p><b>Intermediate metabolizer</b></p> <p>TPMT genotype consistent with an intermediate metabolizer phenotype and is associated with an increased risk of thiopurine-induced toxicities. Impaired NUDT15 activity is also associated with an increased risk of thiopurine-induced toxicities, independent of TPMT phenotype.</p>
UGT1A1	*1/*1		<p><b>Normal metabolizer</b></p> <p>Genotype consistent with fully functional UGT1A1 enzyme activity, or a normal metabolizer phenotype.</p>

## Gene and phenotype summary (cont.)

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VKORC1

rs9923231 GG



### Normal activity

Genotype consistent with normal activity of the vitamin K epoxide reductase enzyme, associated with c.-1639GG (rs9923231). VKORC1, together with CYP2C9, CYP4F2, and a variant in CYP2C Cluster, may affect treatment management of a certain medication.

## Test information

Specimen ID: <b>2286156355623</b>	Clinical testing performed by: <b>OneOme</b>	Lab director: <b>Bronwyn R. Hartung, PhD</b>
Specimen type: <b>Buccal swab</b>	<b>807 Broadway St. NE Suite 100</b>	CLIA: <b>24D2109855</b>
Collection date: <b>2021-01-06</b>	<b>Minneapolis, MN 55412, United States</b>	CAP: <b>9432670</b>
Receive date: <b>2021-01-06</b>		NY PFI: <b>9226</b>

## Test results

The following analytical results were interpreted by OneOme to produce the pharmacogenomic interpretations and annotations described in the *Gene and phenotype summary*. Method-specific analytical limitations or inferred haplotypes may limit the ability to produce a definitive phenotype interpretation. See *Methodology and limitations* and/or the *Report and laboratory comments* sections for additional information.

<b>CYP1A2 *1A/*1F</b>			rs3892097	NM_000106.5:c.506-1G>A	GG
rs2069514	NG_008431.2:g.28338G>A	GG	rs72549353	NM_000106.5:c.765_768delAACT	AACTAACT
rs2069526	NM_000761.4:c.-10+103T>G	TT	rs35742686	NM_000106.5:c.775delA	AA
rs12720461	NM_000761.4:c.-10+113C>T	CC	rs5030656	NM_000106.5:c.841_843delAAG	AAGAAG
rs35694136	NM_000761.4:c.-1635delT	TT	rs16947	NM_000106.5:c.886C>T	CC
rs762551	NM_000761.4:c.-9-154C>A	CA	rs5030867	NM_000106.5:c.971A>C	AA
			rs79292917	NM_000106.5:c.975G>A	GG
			rs28371725	NM_000106.5:c.985+39G>A	GG
<b>CYP2B6 *1/*5</b>			<b>CYP3A4 *1/*1</b>		
rs3211371	NM_000767.4:c.1459C>T	CT	rs2740574	NM_017460.5:c.-392G>A	AA
rs3745274	NM_000767.4:c.516G>T	GG	rs35599367	NM_017460.5:c.522-191C>T	CC
rs2279343	NM_000767.4:c.785A>G	AA	<b>CYP3A5 *3/*3</b>		
rs28399499	NM_000767.4:c.983T>C	TT	rs41303343	NM_000777.4:c.1035_1036insT	--
<b>CYP2C9 *1/*3</b>			rs776746	NM_000777.4:c.219-237G>A	GG
rs28371685	NM_000771.3:c.1003C>T	CC	rs10264272	NM_000777.4:c.624G>A	GG
rs1057910	NM_000771.3:c.1075A>C	AC	<b>CYP4F2 *1/*1</b>		
rs56165452	NM_000771.3:c.1076T>C	TT	rs2108622	NM_001082.4:c.1297G>A	GG
rs28371686	NM_000771.3:c.1080C>G	CC	<b>COMT rs4680 GG</b>		
rs1799853	NM_000771.3:c.430C>T	CC	rs4680	NM_000754.3:c.472G>A	GG
rs7900194	NM_000771.3:c.449G>A	GG	<b>DPYD *1/*1</b>		
rs9332131	NM_000771.3:c.817delA	AA	rs55886062	NM_000110.3:c.1679T>G	TT
<b>CYP2C19 *17/*17</b>			rs3918290	NM_000110.3:c.1905+1G>A	GG
rs12248560	NM_000769.2:c.-806C>T	TT	rs67376798	NM_000110.3:c.2846A>T	TT
rs28399504	NM_000769.2:c.1A>G	AA	<b>DRD2 rs1799978 GG</b>		
rs4986893	NM_000769.2:c.636G>A	GG	rs1799978	NM_000795.3:c.-585A>G	GG
rs6413438	NM_000769.2:c.680C>T	CC	<b>F2 rs1799963 GG</b>		
rs4244285	NM_000769.2:c.681G>A	GG	rs1799963	NM_000506.4:c.*97G>A	GG
<b>CYP2C Cluster rs12777823 GG</b>			<b>F5 rs6025 GG</b>		
rs12777823	NC_000010.10:g.96405502G>A	GG	rs6025	NM_000130.4:c.1601G>A	GG
<b>CYP2D6 *1/*1</b>			<b>GRIK4 rs1954787 CC</b>		
rs1080985	NM_000106.5:c.-1584C>G	CC	rs1954787	NM_001282470.2:c.83-10039T>C	CC
rs1065852	NM_000106.5:c.100C>T	CC	<b>HLA-A Positive for *31:01</b>		
rs59421388	NM_000106.5:c.1012G>A	GG	HLA00097	NM_002116 (interrogated at exon 2)	Positive
rs72549346	NM_000106.5:c.1088_1089insGT	--	<b>HLA-B Negative</b>		
rs5030862	NM_000106.5:c.124G>A	GG			
rs267608319	NM_000106.5:c.1319G>A	GG			
rs774671100	NM_000106.5:c.137_138insT	--			
rs765776661	NM_000106.5:c.1411_1412insTGCCCACTG	GTGCCCACTG			
		AC			
rs1135840	NM_000106.5:c.1457G>C	GG			
rs201377835	NM_000106.5:c.181-1G>C	GG			
rs769258	NM_000106.5:c.31G>A	GG			
rs28371706	NM_000106.5:c.320C>T	CC			
rs5030655	NM_000106.5:c.454delT	TT			
rs5030865	NM_000106.5:c.505G>[A,T]	GG			

## Test results (cont.)

HLA00386 NM\_005514 (interrogated at exon 2 and intron 2) Negative  
 HLA00381 NM\_005514 (interrogated at exon 3) Negative  
 rs144012689 NM\_005514.7:c.1012+104A>T AA

### HTR2A rs7997012 AA

rs7997012 NM\_000621.4:c.614-221T>C TT

### HTR2C rs3813929 CC

rs3813929 NM\_000868.3:c.-759C>T CC

### IFNL4 rs12979860 CT

rs12979860 NM\_001276254.2:c.151-152G>A CT

### NUDT15 rs116855232 CC

rs116855232 NM\_018283.3:c.415C>T CC

### OPRM1 rs1799971 GG

rs1799971 NM\_000914.4:c.118A>G GG

### SLC6A4 L/L (La/La)

rs774676466 NM\_001045.5:c.-1917\_-1875del43 LL  
 rs25531 NM\_001045.5:c.-1936A>G AA

### SLCO1B1 \*1A/\*1A

rs4149015 NM\_006446.4:c.-910G>A GG  
 rs2306283 NM\_006446.4:c.388A>G AA  
 rs4149056 NM\_006446.4:c.521T>C TT

### TPMT \*1/\*4

rs1800462 NM\_000367.3:c.238G>C GG  
 rs1800460 NM\_000367.3:c.460G>A GG  
 rs1800584 NM\_000367.3:c.626-1G>A CT  
 rs1142345 NM\_000367.3:c.719A>G AA

### UGT1A1 \*1/\*1

rs4148323 NM\_001072.3:c.862-6536G>A GG  
 rs1976391 NM\_001072.3:c.862-9697A>G AA

### VKORC1 rs9923231 GG

rs9923231 NM\_001311311.1:c.-1639G>A GG  
 rs7200749 NM\_024006.5:c.358C>T GG

## Methodology and limitations

Analytical results were produced using tests developed and validated by OneOme, LLC, a clinical laboratory located at 807 Broadway Street NE Suite 100, Minneapolis, MN 55413. These tests have not been cleared or approved by the U.S. Food and Drug Administration. OneOme is certified under CLIA-88 and accredited by the College of American Pathologists as qualified to perform high-complexity testing. This test is used for clinical purposes and should not be regarded as investigational or for research.

Genomic DNA was analyzed by PCR using Thermo Fisher TaqMan® and/or LGC Biosearch BHQ® probe-based methods to interrogate the variant locations listed in the *Test results* table above. In addition, CYP2D6 copy number status was assessed at sites within the promoter, intron 2, intron 6, and exon 9. The test detects CYP2D6 deletions, duplications/multiplications, and hybrid alleles, but cannot differentiate duplications in the presence of a deletion.

Haplotypes, or combinations of inherited variants on a chromosome, are annotated according to legacy nomenclature for the genes and alleles in the table below. Less frequent haplotypes or novel alleles may be reported when appropriate.

CYP1A2	*1C, *1D, *1E, *1F, *1J, *1K, *1L, *1V, *1W
CYP2B6	*4, *5, *6, *7, *9, *16, *18
CYP2C9	*2, *3, *4, *5, *6, *8, *11
CYP2C19	*2, *3, *4, *4B, *10, *17
CYP2D6	*2, *2A, *3, *4, *4M, *4N, *5, *6, *6C, *7, *8, *9, *10, *11, *12, *13, *14, *15, *17, *18, *19, *20, *29, *31, *34, *35, *36, *39, 41, *42, *59, *63, *64, *68, *69, *70, *91, *109, *114
CYP3A4	*1B, *22
CYP3A5	*3, *6, *7
CYP4F2	*3
DPYD	*2A, *13
SLCO1B1	*5, *15, *17, *21
TPMT	*2, *3A, *3B, *3C, *4
UGT1A1	*6, *28

The test does not detect all known and unknown variations in the gene(s) tested, nor does absence of a detectable variant (designated as \*1 for genes encoding drug metabolizing enzymes) rule out the presence of other, non-detected variants.

As with other common SNP genotyping techniques, these assays cannot differentiate between the maternal and paternal chromosomes. In cases where observed variants are associated with more than one haplotype, OneOme infers and reports the most likely diplotype based on published allele frequency and/or ethnicity data. Inferences with potential clinical impact are reported in the *Report and laboratory comments* section.

The variant detection methods validated by OneOme provide >99.9% accuracy; however, PCR may be subject to general interference by factors such as reaction inhibitors and low quality or quantity of extracted DNA. When present, these interferents typically yield no result rather than an inaccurate one. Very infrequent variants or polymorphisms occurring in primer- or probe-binding regions may also affect testing and could produce an erroneous result or assay failure. Variant locations tested by the assay but not assigned a genotype call are reported as “No Call.” Test results and clinical interpretation may be inaccurate for individuals who have undergone or are receiving non-autologous blood transfusions, tissue, and/or organ transplant therapies. Although extremely rare, results could also be impacted by other factors not addressed above, such as laboratory error.

Due to the complexity of interpreting some genetic test results, such as those that may carry a probabilistic risk of disease, patients and providers should consider the benefits of consulting with a trained genetic counseling professional, physician, or pharmacogenomic specialist. For additional support, contact OneOme through the website or by calling 844-663-6635.

## OneOme liability disclaimer

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