

Interpreting the GeneSight® Psychotropic Report



Clinician Report Interpretation Resource

GeneSight® Psychotropic
Pharmacogenomic Test

Patient, Sample
Date of Birth: MM/DD/YYYY
Clinician: Sample Clinician

Order Number: 0000000
Report Date: MM/DD/YYYY
Reference: 000000

Questions about report interpretation?
Contact our Medical Information team:
855.891.9415 | medinfo@genesight.com

Non-Smokers
Smoking is defined as the daily inhalation of burning plant material (cigarettes, marijuana), and excludes vaping and e-cigarettes. This is used to determine medication results.

Use as Directed

desipramine (Norpramin®)	7
desvenlafaxine (Pristiq®)	7
levomilnacipran (Fetzima®)	
nortriptyline (Pamelor®)	
trazodone (Desyre®)	
vortioxetine (Vibryd®)	
duloxetine (Trintellix®)	
mirtazapine (Remeron®)	

Antidepressants

Moderate Gene-drug Interaction

venlafaxine (Effexor®)	1
selegiline (Elavil®)	3
fluoxetine (Prozac®)	1.4
clomipramine (Anafranil®)	1.7
fluvoxamine (Luvox®)	4.7

Significant Gene-drug Interaction

bupropion (Wellbutrin®)	2
amitriptyline (Elavil®)	3
paroxetine (Paxil®)	4.6
escitalopram (Lexapro®)	1.4, 6
sertraline (Zoloft®)	1.4, 6
imipramine (Tofranil®)	1.6, 7
citalopram (Celexa®)	1.4, 6, 8
doxepin (Sinequan®)	1.6, 7, 8

Clinical Considerations

1. Serum level may be too high, lower doses may be required.
2. Serum level may be too low, higher doses may be required.
3. Difficult to predict dose adjustments due to conflicting variations in metabolism.
4. Genotype may impact drug mechanism of action and result in moderately reduced efficacy.
5. Use of this drug may increase risk of side effects.
6. Smoking status changes the results of this medication. See next section labeled Smokers for smoking results.
7. FDA label identifies a potential gene-drug interaction for this medication.

All psychotropic medications require clinical monitoring. Medications should not be changed based solely on the test results. The results of this test are not intended to be used as the sole basis for clinical decisions. The results of this test are not intended to be used as the sole basis for clinical decisions. The results of this test are not intended to be used as the sole basis for clinical decisions.

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The GeneSight Test may help inform medication selection and dosing. The results of the test show which medications may require dose adjustments, may be less likely to work, or may have an increased risk of side effects based on the patient's genetic results. The GeneSight Test provides pharmacogenomic information for antidepressants, anxiolytics & hypnotics, antipsychotics, medications for tardive dyskinesia, mood stabilizers, and stimulants & non-stimulants used to treat ADHD.

GeneSight® Psychotropic

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Antidepressants

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Use as Directed	Moderate Gene-drug Interaction	Significant Gene-drug Interaction
desipramine (Norpramin®) desvenlafaxine (Pristiq®) levomilnacipran (Fetzima®) nortriptyline (Pamelor®) trazodone (Desyrel®) vilazodone (Viibryd®) vortioxetine (Trintellix®)	venlafaxine (Effexor®) 1 selegiline (Emsam®) 3 fluoxetine (Prozac®) 1.4 nortriptyline (Pamelor®) 1.7 clomipramine (Anafranin®) 1.4 fluvoxamine (Luvox®) 4.7	bupropion (Wellbutrin®) 2 amitriptyline (Elavil®) 3 paroxetine (Paxil®) 4.6 escitalopram (Lexapro®) 1.4 sertraline (Zoloft®) 1.6 imipramine (Tofranil®) 1.6,7 citalopram (Celexa®) 1.4,6,8 doxepin (Sinequan®) 1.6,7,8

Clinical Considerations

- Serum level may be too high.
- Serum level may be too low.
- Difficult to predict dose adjust
- Genotype may impact drug m
- Use of this drug may increase
- FDA label identifies a potenti

All psychotropic medications require clinical supplement other clinical information consult. This report is not intended to imply that the drug purposes only other brand names may be available based on the patient's individual needs, the characteristics of the drug prescribed, and the risk and safety information provided in the drug's labeling. Propranolol and oxcarbazepine prescribed for neuropsychiatric disorders might be considered off-label. Please consult their respective FDA drug labels for specific guidelines regarding their use.

Clinical Considerations

- Serum level may be too high, lower doses may be required.
- Serum level may be too low, higher doses may be required.
- Difficult to predict dose adjustments due to conflicting variations in metabolism.
- Genotype may impact drug mechanism of action and result in moderately reduced efficacy.
- Use of this drug may increase risk of side effects.
- Smoking status changes the results of this medication. See next section labeled Smokers for smoking results.
- FDA label identifies a potential gene-drug interaction for this medication.

All psychotropic medications require clinical monitoring. Medications should not be changed based solely on the test results. The results of this test are intended to supplement other clinical information considered by a healthcare provider within the context of a comprehensive medical evaluation. This report is not intended to imply that the drugs listed are approved for the same indications or that they are comparable in safety or efficacy. The brand name is shown for illustrative purposes only; other brand names may be available. The prescribing physician should review the prescribing information for the drug(s) being considered and make treatment decisions based on the patient's individual needs, the characteristics of the drug prescribed, and the risk and safety information provided in the drug's labeling. Propranolol and oxcarbazepine prescribed for neuropsychiatric disorders might be considered off-label. Please consult their respective FDA drug labels for specific guidelines regarding their use.

The GeneSight Psychotropic test interpretations are based on a thorough review of published peer-reviewed literature, internal research, and FDA label information when applicable. The clinical validity and utility of the GeneSight Psychotropic test have been evaluated for patients with major depressive disorder who take at least one psychotropic medication in multiple clinical studies.

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Patient, Sample
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These medications are not associated with any known genetic issues that would be expected to change patient medication outcomes. However, these medications are not guaranteed to work and may not always be the best options, as there are many other factors that influence medication response and susceptibility to side effects, including drug-drug interactions, diet, environmental factors, age, etc.

These medications may require dose adjustments in order to have the desired effect, may be less likely to work, or may cause side effects.

These medications are likely to require dose adjustments in order to have the desired effect, may be less likely to work, or may cause side effects. Genetics are expected to have a greater impact on medications in the significant gene-drug interaction category than those that fall into the moderate gene-drug interaction category.

Many factors may influence medication outcomes in addition to genetics. Smoking status is one environmental factor that can contribute to a gene-drug-environment interaction. This is because smoking can increase the activity of CYP1A2 in patients who have the highly inducible CYP1A2 variant. For these patients, the categorization and information provided for certain medications can be affected by smoking status, where smoking is defined as the daily inhalation of burning plant material (cigarettes, marijuana), and excludes vaping and e-cigarettes.

For patients who carry the highly inducible CYP1A2 variant, the CYP1A2 phenotype is dependent on their smoking status.

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Use as Directed

Moderate Gene-drug Interaction

Significant Gene-drug Interaction

desipramine (Norpramin®)	venlafaxine (Effexor®)	1	bupropion (Wellbutrin®)	2
desvenlafaxine (Pristiq®)	selegiline (Emsam®)	3	amitriptyline (Elavil®)	3
levomilnacipran (Fetzima®)	fluoxetine (Prozac®)	1,4	paroxetine (Paxil®)	4,6
nortriptyline (Pamelor®)	clomipramine (Anafranil®)	1,7	escitalopram (Lexapro®)	1,4,6
trazodone (Desyrel®)	fluvoxamine (Luvox®)	4,7	sertraline (Zoloft®)	1,4,6
vilazodone (Viibryd®)			imipramine (Tofranil®)	1,6,7
vortioxetine (Trintellix®)			citalopram (Celexa®)	1,4,6,8
duloxetine (Cymbalta®)			doxepin (Sinequan®)	1,6,7,8
mirtazapine (Remeron®)				



Non-smokers who have the highly inducible variant would be expected to have normal CYP1A2 enzyme activity, and the sections labeled Non-Smokers should be used.

It is important to be aware of smoking status and refer to the correct sections of the report for individuals with the highly inducible CYP1A2 variant.

Some patients (approximately 9%) do not have the highly inducible variant found in CYP1A2.

For these individuals, their reports will note that the results provided are for both non-smokers and smokers since the individual does not have the highly inducible CYP1A2 variant. Therefore, smoking status does not have to be considered as it does for those with the highly inducible variant.

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Use as Directed

Moderate Gene-drug Interaction

Significant Gene-drug Interaction

desipramine (Norpramin®)	venlafaxine (Effexor®)	1	bupropion (Wellbutrin®)	2
desvenlafaxine (Pristiq®)	clomipramine (Anafranil®)	3	duloxetine (Cymbalta®)	2
levomilnacipran (Fetzima®)	selegiline (Emsam®)	3	mirtazapine (Remeron®)	2
nortriptyline (Pamelor®)	fluoxetine (Prozac®)	1,4	amitriptyline (Elavil®)	3
trazodone (Desyrel®)	imipramine (Tofranil®)	1,6	fluvoxamine (Luvox®)	2,4
vilazodone (Viibryd®)	doxepin (Sinequan®)	1,8	paroxetine (Paxil®)	4,6
vortioxetine (Trintellix®)			escitalopram (Lexapro®)	1,4,6
			sertraline (Zoloft®)	1,4,6
			citalopram (Celexa®)	1,4,6,8



Smokers with the highly inducible variant may have increased CYP1A2 enzyme activity, and the sections labeled Smokers should be used.

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Antidepressants

This section shows the results for non-smokers and smokers due to the absence of the highly inducible CYP1A2 variant.

Use as Directed

Moderate Gene-drug Interaction

Significant Gene-drug Interaction

desvenlafaxine (Pristiq®)	selegiline (Emsam®)	1	fluoxetine (Prozac®)	1,6
levomilnacipran (Fetzima®)	trazodone (Desyrel®)	1	mirtazapine (Remeron®)	1,6
	vilazodone (Viibryd®)	1	venlafaxine (Effexor®)	1,6
	sertraline (Zoloft®)	1,4	citalopram (Celexa®)	1,4,6
	bupropion (Wellbutrin®)	1,6	escitalopram (Lexapro®)	1,4,6
			amitriptyline (Elavil®)	1,6,8
			clomipramine (Anafranil®)	1,6,8
			desipramine (Norpramin®)	1,6,8
			doxepin (Sinequan®)	

Interpreting the Clinical Considerations

All medications start in the “Use as Directed” category on the report. Based on an individual’s genetic variation, medications may be moved to the Moderate or Significant Gene-drug Interaction categories, depending on how significantly the variation is expected to impact outcomes with that medication. Medications in the Moderate or Significant Gene-drug Interaction categories on the report do not necessarily need to be avoided. The clinical considerations, which are denoted by numbers next to the medications, explain the rationale for a medication’s classification and can be used to help inform treatment decisions.

Clinical Considerations

- 1: Serum level may be too high, lower doses may be required.
- 2: Serum level may be too low, higher doses may be required.
- 3: Difficult to predict dose adjustments due to conflicting variations in metabolism.
- 4: Genotype may impact drug mechanism of action and result in moderately reduced efficacy.
- 5: CYP2D6 genotype indicates that this patient may experience increased frequency of side effects but also greater symptom improvement in those who find the treatment tolerable.
- 6: Use of this drug may increase risk of side effects.
- 7: Smoking status changes the results of this medication. See next section labeled Smokers for smoking results.
- 8: FDA label identifies a potential gene-drug interaction for this medication.
- 9: Per FDA label, this medication is contraindicated for this genotype.
- 10: While this medication does not have clinically proven genetic markers that allow it to be categorized, it may be an effective choice based on other clinical factors.

Clinical Considerations 1 and 2 reflect an issue with how a medication is metabolized, which means the individual has variation in one or more pharmacokinetic genes. This may affect how much medication is in an individual’s system and may require a lower or higher dose. For medications that may require a higher dose, it is not advised to start the individual on a high dose of the medication, but rather to start at a standard dose while being aware that dose increases may be needed to get the desired benefit.

Clinical Consideration 3 also provides information about a medication’s metabolism. This clinical consideration indicates that there are conflicting variations in the metabolism of that medication, thus making it difficult to predict a dose adjustment. The most common example is when there are two genes that contribute a relatively similar amount to the overall metabolism of the medication, but one has increased activity and the other has decreased activity. If a medication with clinical consideration 3 is chosen, it may be appropriate to consider starting on a lower dose and increasing the dose based upon how the individual is doing.

Clinical Consideration 4 indicates that an individual may have a moderately reduced response to certain medications. This is due to variation in a pharmacodynamic gene, and thus reflects an issue with the drug’s mechanism of action. Since pharmacodynamic genes do not affect metabolism, it is unlikely that adjusting the dose will improve efficacy.

Clinical Consideration 5 specifically applies to atomoxetine (Strattera®) when an individual is a CYP2D6 poor metabolizer. This is based on evidence which shows that CYP2D6 poor metabolizers treated with atomoxetine experienced an increased frequency of side effects but also significantly greater improvement in ADHD symptoms compared to extensive (normal) metabolizers.¹

Clinical Consideration 6 indicates that an individual may have an increased risk for side effects when taking this medication. This can be due to variation in one or more pharmacokinetic genes that may predict slower than normal metabolism of certain medications, which may result in higher levels of medication in the system. It could also be due to an effect caused by one of the pharmacodynamic genes.

Clinical Consideration 7 specifically applies to individuals who have the highly inducible variant of the CYP1A2 gene, which may result in increased activity for CYP1A2 when the patient is a smoker. This increased activity may change how this medication is metabolized. For individuals who are smokers, it is important to refer to the results in the following section labeled Smokers.

When medications have **Clinical Consideration 8**, it means the FDA label contains information that may be relevant for the patient based on their genetic results. This may include information about dosing or more general information to consider.

Clinical Consideration 9 indicates that per the FDA label, this medication is contraindicated for an individual with this genotype.

Clinical Consideration 10 is associated with medications in the gray “No Proven Genetic Markers” category, meaning that genetic markers have not yet been discovered to reliably predict treatment outcomes with these medications. Due to this lack of evidence, we are not currently able to categorize them or provide actionable recommendations. However, these medications may be effective choices based on other clinical factors. Medications that fall into the “No Proven Genetic Markers” category and receive a clinical consideration of 10 are the same for all individuals.

No Proven Genetic Markers

gabapentin (Neurontin®)	10	lithium (Eskalith®)	10	topiramate (Topamax®)	10
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Interpreting the Patient Genotypes and Phenotypes

The patient genotypes and phenotypes section of the report provides a list of all the genes tested on the GeneSight® Psychotropic report along with the individual’s specific genetic result (genotype) and the characteristic associated with that genetic result (phenotype). **The patient genotypes and phenotypes are broken down into two categories:**

Pharmacodynamic Genes



Pharmacodynamic genes provide information about how a medication works on the body. Variation in these genes may affect likelihood of response or risk of side effects with certain medications.

ADRA2A

ADRA2A encodes the alpha-2A adrenergic receptor, which is a norepinephrine receptor. Individuals with the C/C genotype may have a moderately reduced response to certain stimulants, specifically methylphenidate and dexamethylphenidate. If an individual with this genotype is having a less than optimal response to these medications, increasing the dose may not necessarily improve efficacy.

HLA-A*3101

The human leukocyte antigen (HLA) complex, encoded by the HLA gene family, plays a critical role in immunity. Presence of the T allele (either T/T or A/T genotype) is associated with a higher risk of serious hypersensitivity reactions, including Stevens-Johnson syndrome (SJS), toxic epidermal necrolysis (TEN), maculopapular eruptions, and Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS), when taking certain mood stabilizers, specifically carbamazepine.

HLA-B*1502

Presence of genetic variation in HLA-B*1502 is associated with a higher risk of serious dermatologic reactions, including toxic epidermal necrolysis (TEN) and Stevens-Johnson syndrome (SJS), when taking certain mood stabilizers, including carbamazepine, oxcarbazepine, and lamotrigine.

HTR2A

HTR2A encodes for the Serotonin Receptor Type 2A, which is responsible for serotonin signaling. Studies show an increased risk of adverse effects with paroxetine in individuals who have the G/G genotype.

SLC6A4

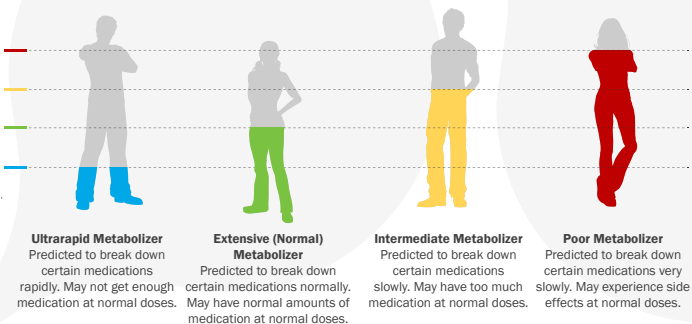
SLC6A4 encodes for the serotonin transporter, which is the main site of action for SSRIs. SLC6A4 has two main versions: the long (L) allele and the short (S) allele. Studies have shown that the short allele results in less serotonin transporters than the long allele. Individuals who have the S allele may be less likely to respond to certain SSRIs based on this genotype. Additionally, due to the lower number of transporters, increasing the dose may not necessarily improve efficacy.

Pharmacokinetic Genes



Pharmacokinetic genes provide information about how the body works on the medication. Variation in these genes may affect the metabolism of a medication. The following PK genes are included on The GeneSight Test: CES1A1, CYP1A2, CYP2B6, CYP2C19, CYP2C9, CYP2D6, CYP3A4, UGT1A4, and UGT2B15.

These genes are categorized based on average enzyme activity into one of four metabolizer phenotypes on the GeneSight Psychotropic report.



The GeneSight Test’s weighted multi-gene approach measures multiple genomic variants for each individual and weighs them together in order to categorize the medications on the report and provide clinical considerations when applicable.

Additional Genotypes

The genotype for COMT, or catechol-o-methyltransferase, is provided for informational purposes only. This gene has not been shown to be a reliable marker of medication outcomes, and therefore, it is not used to categorize medications on the report.

Interpreting the Gene-drug Interaction Chart

The gene-drug interaction chart provides supplementary information about which pharmacokinetic genes are involved in the metabolism of each medication.

The pharmacokinetic genes on the GeneSight® report are listed across the top of the chart.

Medications are listed in a column on the left side.

- Any dot (either shaded, unshaded, or half-shaded) signifies that the enzyme is involved in the metabolism of the associated medication.
- A shaded dot means that variation was found in the patient's genotype that may impact medication metabolism.
- An unshaded dot indicates that the gene is associated with medication metabolism, but the predicted patient phenotype is normal.
- A half-shaded dot means that the phenotype is dependent upon smoking status, with smokers being ultrarapid metabolizers and non-smokers being normal metabolizers. Medication categorization in this chart is according to non-smokers. Refer to sections labeled Smokers to see medication categories for individuals who smoke.

Gene-drug Interactions									
Use as Directed	CES1A1	CYP1A2	CYP2B6	CYP2C19	CYP2C9	CYP2D6	CYP3A4	UGT1A4	UGT2B15
	Normal	Smoking Dependent	Ultrarapid	Poor	Normal	Normal	Normal	Normal	Poor
Antidepressants									
desipramine (Norpramin®)						○			
desvenlafaxine (Pristiq®)				●			○		
duloxetine (Cymbalta®)		●				○			
levomilnacipran (Fetzima®)				●		○	○		
mirtazapine (Remeron®)		●			○	○	○		
nortriptyline (Pamelor®)						○			
trazodone (Desyre®)		●				○	○		
vilazodone (Viibryd®)				●		○	○		
vortioxetine (Trintellix®)			●	●	○	○	○		
Anxiolytics and Hypnotics									
alprazolam (Xanax®)							○		
buspirone (BuSpar®)						○	○		
chlordiazepoxide (Librium®)							○		
clonazepam (Klonopin®)							○		
eszopiclone (Lunesta®)					○		○		
lemborexant (Dayvigo®)							○		
propranolol (Inderal®)		●				○			
suvorexant (Belsomra®)							○		
zolpidem (Ambien®)		●		●			○		
Antipsychotics									
aripiprazole (Abilify®)						○	○		
asenapine (Saphris®)		●				○	○	○	
brexpiprazole (Rexulti®)						○	○		
cariprazine (Vraylar®)						○	○		
chlorpromazine (Thorazine®)		●				○	○		
clozapine (Clozaril®)		●				○	○	○	
haloperidol (Haldol®)		●				○	○	○	
iloperidone (Fanapt®)						○	○		
lumateperone (Caplyta®)							○		

It is not mandatory to refer to the gene-drug interaction chart when using The GeneSight Test to inform patient medication selection. It is simply intended to augment the patient results on the pages where the medications are categorized and the clinical considerations are provided, which serve as the primary resource to help inform treatment decisions.

The format of the gene-drug interaction chart is similar to the earlier pages of the GeneSight report where medications are categorized into the green, yellow, and red categories.

There are situations where an individual may have shaded dots in the green “Use as Directed” category. This means that although the individual has variation in one or more genes, it is unlikely to impact their overall

metabolism of that particular medication. This could mean that while an enzyme is involved in the metabolism of the medication, its role is not clinically significant enough to warrant a change in dosing or there may be compensation among the enzymes known to metabolize the medication, which results in the medication not being moved from the green “Use as Directed” category.

The results of The GeneSight Test are intended to supplement other clinical factors considered by a healthcare provider during a comprehensive medical assessment to help inform treatment decisions.

Frequently Asked Questions

Will green (Use as Directed) category medications work best?

A medication falls in the green category when no variation was found in the genes tested by the GeneSight® Test that is expected to impact how a patient may break down or respond to that medication. However, medications that fall in the green category may not always work for a patient. Pharmacogenomics identifies potential genetically related issues with medications (i.e., those that fall in the yellow or red category); it does not identify medications that will work. Additionally, while genetics provides an important piece of the puzzle, there are many factors that can influence medication response and susceptibility to side effects other than the genes that are tested on the GeneSight panel, including drug-drug interactions, diet, psychological factors, environmental factors, age, etc. Genetics can be a helpful tool, but it is important to take into consideration a patient's entire clinical profile for medication selection.

Are the medications listed first in a category better than the ones listed lower?

Medications are listed in alphanumeric order. Those with the fewest numbers, or clinical considerations, are placed at the top of the list in alphabetical order, and the medications with more clinical considerations are placed further down the list. It is important to consider what the clinical considerations are saying as they may help inform on medication selection and treatment decisions for medications affected by gene-drug interactions.

What does it mean if a medication in the red (Significant Gene-drug Interaction) category is currently working?

While clinical studies have shown that patients on red category medications had less improvement in depression treatment outcomes, this may not be the case for all patients who undergo GeneSight testing. It is important to acknowledge that medication response is multifactorial, and genetics is one factor that may influence how a medication works alongside other non-genetic factors such as drug-drug interactions, diet, psychological factors, environmental factors, age, etc. GeneSight Test results are intended to supplement other clinical and patient specific factors to help inform medication selection and treatment decisions.

Why do some medications have the same clinical considerations, but some are in the yellow (Moderate Gene-drug Interaction) category and some are in the red (Significant Gene-drug Interaction) category?

Medications are categorized by the GeneSight algorithm based on the accumulation of genetic issues (gene-drug interactions) detected. The GeneSight Test takes into account all relevant genes known to affect metabolism or response to each medication and the relative influence of each gene. Thus, a medication in the red category has greater cumulative genetic issues than a medication in the yellow category, and this may be because different genes are involved in predicting changes in metabolism or response to it, or because the same gene plays a larger role in the red category medication than in the yellow category medication.

What is the utility of an all-green (Use as Directed) report?

An all-green report indicates that no significant variation was detected in the patient's genes tested by the GeneSight Test that would be expected to affect their outcomes with medications on the GeneSight report. This occurs <1% of the time for GeneSight Psychotropic. An all-green report may be useful in helping 'rule out' genetic causes of medication intolerance or lack of response, and may lead to alternative treatment approaches, such as psychotherapy.

Do medications the patient is taking affect the GeneSight Test results?

From a genetic perspective, the genetic results shown on the GeneSight report will be accurate regardless of what medications a patient is taking. The GeneSight Test evaluates only gene-drug interactions. In other words, it looks at potential changes in a patient's DNA to predict how this might alter the patient's expected outcomes with certain medications. Since a person's genes will not change based on what medications they are taking, the GeneSight results will not change. However, it is possible for certain medications to interact with medications on the report, and this is called a drug-drug interaction. Drug-drug interaction potential should be taken into consideration alongside GeneSight Test results as they may affect medication outcomes in addition to the genetic information provided on the GeneSight report.



Genetic insights to help navigate your patient's next treatment.

Do you or your patient have comments or questions? Patients and clinicians can all our medical information team to assist with report interpretation.

Medical Information
855.891.9415
medinfo@genesight.com

Visit genesight.com to learn more.

Not all patients who receive The GeneSight Test will have improved outcomes. The GeneSight Test is intended to supplement a clinician's comprehensive medical assessment.

1. Michelson D, et al. 2007. J Am Acad Child Adolesc Psychiatry.

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