

Name: John Doe

Medical Request No. 1234567

D.O.B: dd/mm/yyyy Referring physician: Dr. Doctor name

 Sex:
 Male
 Referring facility:
 Medical Center, Country

 Subject ID:
 8xxxx
 Email physician:
 doctorname@email.com

Order ID: 2xxxxx Report type: myLifeGenomeTM Device/ Material ID: ARCxxxxxx Date of report: 11/05/2024

Specimen type: Buccal swab
Specimen arrival date: dd/mm/2024

Requested Test: myLifeGenome[™] -Solo

Indication for test: Elevated circulating alkaline phosphatase concentration, Elevated

gamma-glutamyltransferase level, Increased circulating IgE level,

Pancytopenia, Severe generalized osteoporosis

Consanguineous parents: Yes

Consent for evaluation: Primary findings: Yes

Incidental findings: Yes Carrier findings: Yes

Pharmacogenomics findings: Yes

SUMMARY OF GENETIC FINDINGS

PRIMARY INCIDENTAL CARRIER STATUS PHARMACOGENOMICS

Positive Positive Positive Positive

Primary Findings

(i) Primary findings describe genetic variants that are relevant to the indication for which sequencing was ordered.

Results: A homozygous pathogenic variant was identified in the SGSH gene. This result is consistent with the genetic diagnosis of autosomal recessive mucopolysaccharidosis type 3A.

Gene	Variant	Zygosity	Variant class*	Disease name	Disease MOI*
SGSH	c.697C>T; p.Arg233*	Hom	Р	Mucopolysaccharidosis type 3A	AR

Incidental Findings

Incidental findings describe actionable variants in gene(s) that are unrelated to the indication for which sequencing was ordered. These findings are reported based on ACMG guidelines and ClinGen recommendations.

Results: A heterozygous likely pathogenic variant was identified in the TTN gene, A-band. This result is consistent with increased risk to develop autosomal dominant dilated cardiomyopathy type 1G.

Gene	Variant	Zygosity	Variant class*	Disease name	Disease MOI*
TTN	c.107641G>T; p.(Glu35881*)	Het	LP	Dilated cardiomyopathy type 1G (CMD1G)	AD

AD: autosomal dominant; AR: autosomal recessive; XL- X-linked, DR: digenic recessive; Het: Heterozygous; Hom: Homozygous; Hem: Hemizygous; LP: Likely Pathogenic; P: Pathogenic; RF: risk factor VUS: Variant of Uncertain Significance.

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Carrier Status

<u>(i)</u>

Carrier Status includes pathogenic or likely pathogenic variants which have a direct impact on reproductive risk (heterozygous variants in a gene associated with a recessive or X-linked disorder).

Results: This proband is found to be a carrier for 3 genetic conditions.

Gene	Variant	Zygosity	Variant class*	Disease name	Disease MOI*
CEP250	c.6913C>T; p.Arg2305*	Het	LP	Cone-rod dystrophy and hearing loss type 2	AR
DHTKD1	c.1671+1G>A; p.?	Het	LP	Alpha-aminoadipic and alpha- ketoadipic aciduria	AR
TPRN	c.25delT; p.Ser9fs*22	Het	LP	Deafness type 79	AR

Pharmacogenomic Associations



Pharmacogenomic (PGx) Associations are representations of the relationship between specific genes and drugs based on their drug metabolizing status defined according to publicly available data sets provided by Clinical Pharmacogenetics Implementation Consortium (CPIC) (mainly levels A/B or 1/2) using Pharmacogenomics Clinical Annotation Tool (PharmCAT).

Results: Genetic variants associated with drug use and dosing were identified.

See the Pharmacogenomic Associations section under Detailed Insights to understand the identified gene-drug pairs that could lead to treatment modifications based on the individual's genetic variants. See Technical Information section for the list of all genes and drugs supported by PharmCAT.

RECOMMENDATIONS

- Genetic counselling is recommended to further explain the implications of this test result and assess family health history, which may point to health information that merits additional consideration.
- The medical genetics field is continuously evolving, so updates related to your genetic results, medical recommendations, and potential treatments may be available over time.
- Mucopolysaccharidosis type 3A is inherited in an autosomal recessive manner. The parents of an affected child are obligate heterozygotes (i.e. carriers). At conception, each sib of an affected individual has a 25% chance of being affected, a 50% chance of being an asymptomatic carrier, and a 25% chance of being unaffected and not a carrier. Once pathogenic variants in the family are known, carrier testing for at-risk relatives, prenatal testing for a pregnancy at increased risk, and preimplantation genetic testing is possible. We will proceed with Sanger sequencing in the family members (affected and unaffected siblings) to establish the genetic diagnosis and further identify at-risk carriers.
- Truncating TTN variants localized in the A-band of titin protein have been associated with dilated cardiomyopathy. The transmission pattern of CMD1G in the families reported by Gerull et al. (2002) was consistent with autosomal dominant inheritance with incomplete penetrance. Cardiovascular screening of asymptomatic first-degree family members of an individual with genetic increased susceptibility risk to develop DCM can allow early detection of DCM, prompt initiation of treatment, and improvement in long-term outcome (Morales & Hershberger 2015). Clarification of the genetic status of first-degree family members of an individual with DCM can inform who is at risk and the recommended frequency of subsequent cardiovascular screening (Hershberger et al 2018).

Signatures



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DETAILED INSIGHTS

Primary Findings

A homozygous pathogenic variant was identified in the SGSH gene. This result is consistent with the genetic diagnosis of autosomal recessive mucopolysaccharidosis type 3A.



SGSH (N-Sulfoglucosamine Sulfohydrolase) is a protein coding gene. This gene encodes the enzyme sulfamidase; one of several enzymes involved in the lysosomal degradation of heparan sulfate. Mutations in this gene are associated with the lysosomal storage disease mucopolysaccaridosis 3A, also known as Sanfilippo syndrome A, which results from impaired degradation of heparan sulfate. Transcripts of varying sizes have been reported but their biological validity has not been determined. An important paralog of this gene is ARSA.

Gene/OMIM	SGSH/605270
Genomic coordinate (GRCh38)	chr17:80213852G>A
ID Transcript	NM_000199.5
HGVS nomenclature	c.697C>T
Protein change	p.Arg233*
Location	exon 6/8
Zygosity	Hom
Function	Nonsense
Impact	High
ClinVar	Pathogenic, Likely Pathogenic
Local Database	N/A
gnomAD	0.0000263
REVEL	N/A
CADD (PHRED)	42.0
Splice-Al	0.04
Clinical significance	Pathogenic
ACMG Criteria	PVS1, PM2, PM3
	Genomic coordinate (GRCh38) ID Transcript HGVS nomenclature Protein change Location Zygosity Function Impact ClinVar Local Database gnomAD REVEL CADD (PHRED) Splice-Al Clinical significance

HGVS= Human Genome Variation Society; gnomAD= Genome Aggregation Database; ACMG= American College of Medical Genetics and Genomics; REVEL score (combination from 13 individual tools; ranges from 0 to 1)= higher scores reflect greater likelihood that variant is disease- causing; CADD (PHRED)= Combined Annotation Dependent Depletion scoring, ranging from 1 to 99, Splice-AI= deep neural network that accurately predicts splice junctions from an pre-mRNA transcript (using 0.8 as high-precision cut-off).

*: The ACMG criteria are described under Methods /Variant interpretation section.

Disease description: Mucopolysaccharidosis type 3 (MPS 3) is a multisystem lysosomal storage disease characterized by progressive central nervous system degeneration manifest as severe intellectual disability (ID), developmental regression, and other neurologic manifestations including autism spectrum disorder (ASD), behavioral problems, and sleep disturbances. Disease onset is typically before age ten years. Disease course may be rapidly or slowly progressive; some individuals with an extremely attenuated disease course present in mid-to-late adulthood with early-onset dementia with or without a history of ID. Systemic manifestations can include musculoskeletal problems (joint stiffness, contractures, scoliosis, and hip dysplasia), hearing loss, respiratory tract and sinopulmonary infections, and cardiac disease (valvular thickening, defects in the cardiac conduction system). Neurologic decline is seen in all affected individuals; however, clinical severity can vary even among members of the same family. The subtypes of MPS 3 (MPS 3A, MPS 3B, MPS 3C, MPS 3D) are distinguished by their associated enzymatic deficiencies rather than phenotypic differences. However, MPS 3A typically have the most severe and rapidly progressing disease course (PMID: 31536183).

Treatment of manifestations is based on supportive therapies for neurodevelopmental delays, hearing loss, and visual impairment; medications (rather than behavioral therapy) for psychiatric/behavioral issues; physical therapy and/or orthopedic management of musculoskeletal manifestations; and management as prescribed by consulting specialists for seizures, cardiac involvement, sleep disorders, feeding difficulties. Surveillance is through routine monitoring of developmental capabilities and educational needs, destructive or disruptive behaviors; musculoskeletal involvement; hearing; cardiac involvement.

Individuals with mucopolysaccharidosis type 3 may participate in the clinical trials: https://clinicaltrials.gov/search?cond=MUCOPOLYSACCHARIDOSIS,%20TYPE%20IIIA

Variant description: This variant creates a premature stop codon at position 233. It is expected to result in a truncated or disrupted protein. The variant is present in gnomAD (allele frequency: 0.0000263) and is absent from the local database. This variant is listed in ClinVar as pathogenic/likely Pathogenic (Accession ID: 370732). This variant is classified as pathogenic based on ACMG/ClinGen recommendations.



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Incidental Findings

A heterozygous likely pathogenic variant was identified in the TTN gene. This result is consistent with increased risk to develop autosomal dominant dilated cardiomyopathy type 1G.

(i)

The TTN gene encodes a large abundant protein of striated muscle. The product of this gene is divided into two regions, a N-terminal I-band and a C-terminal A-band. The I-band, which is the elastic part of the molecule, contains two regions of tandem immunoglobulin domains on either side of a PEVK region that is rich in proline, glutamate, valine and lysine. The A-band, which is thought to act as a protein-ruler, contains a mixture of immunoglobulin and fibronectin repeats, and possesses kinase activity. An N-terminal Z-disc region and a Cterminal M-line region bind to the Z-line and M-line of the sarcomere, respectively, so that a single titin molecule spans half the length of a sarcomere. Titin also contains binding sites for muscle associated proteins so it serves as an adhesion template for the assembly of contractile machinery in muscle cells. It has also been identified as a structural protein for chromosomes. Alternative splicing of this gene results in multiple transcript variants. Considerable variability exists in the I-band, the M-line and the Z-disc regions of titin. Variability in the I-band region contributes to the differences in elasticity of different titin isoforms and, therefore, to the differences in elasticity of different muscle types.

	Gene/OMIM	TTN/602851
	Genomic coordinate (GRCh38)	Chr2:178527485 C>T
	ID Transcript	NM_001267550.2
	HGVS nomenclature	c.107641G>T
	Protein change	p.(Glu35881*)
	Location	exon 179 / 360
	Zygosity	Het
	Function	stop_gained
	Impact	HIGH
	ClinVar	-
Allele	Local Database	-
Frequency	gnomAD	-
In silico	REVEL	-
Predictors	CADD (PHRED)	-
	Splice-Al	-
	Clinical significance	Likely pathogenic
	ACMG Criteria*	PVS1. PM2_SUP

HGVS= Human Genome Variation Society; gnomAD= Genome Aggregation Database; ACMG= American College of Medical Genetics and Genomics; REVEL score (combination from 13 individual tools; ranges from 0 to 1)= higher scores reflect greater likelihood that variant is disease-causing; CADD (PHRED)= Combined Annotation Dependent Depletion scoring, ranging from 1 to 99, Splice-AI= deep neural network that accurately predicts splice junctions from an pre-mRNA transcript (using 0.8 as high-precision cut-off). *: The ACMG criteria are described under Methods / Variant interpretation section.

Variant description: This changes the amino acid from a Glu to a

stop codon within coding exon 179. This exon is in the A-band

region of the N2-B isoform of the titin protein and is constitutively

expressed in TTN transcripts (percent spliced in or PSI 100%). This

alteration is expected to result in loss of function by premature

protein truncation or nonsense-mediated mRNA decay. This

variant is rare based on population cohorts in the Genome

Disease Disease description: Dilated cardiomyopathy type 1D. Dilated cardiomyopathy (DCM) is a heart muscle disease characterized by left ventricular dilation and systolic dysfunction, and typically it presents with heart failure with symptoms of congestion (edema, orthopnea, paroxysmal nocturnal dyspnea) and/or reduced cardiac output (fatigue, dyspnea on exertion)], arrhythmias and/or conduction system disease, and thromboembolic disease including stroke. Patients with DCM are at risk of premature death (ORPHA:217604. DCM may be asymptomatic with only mild ventricular dilation and DCM may be asymptomatic with only mild ventricular dilation and dysfunction for years. Patients with severe heart failure, severe reduction of the functional capacity and depressed left ventricular ejection fraction have a low survival rate and may require heart transplant.

The management of DCM aims at reducing symptoms of heart failure and improving cardiac function. Clinical management of a patient with symptomatic DCM starts with standard heart failure medications. Specific recommendations regarding DCM for individuals involved in sports can be found in the relevant guidelines (PMID: 32860412, PMID: 32845299).

Individuals with DCM my participate in the clinical trials: https://clinicaltrials.gov/ct2/show/NCT04572893 https://www.clinicaltrialsregister.eu/ctr-search/search/query=Dilated+cardiomyopathy

Aggregation Database (gnomAD). While truncating variants in TTN are present in 1-3% of the general population, truncating variants in the A-band are the most common cause of dilated cardiomyopathy (DCM) (Herman DS et al. N. Engl. J. Med., 2012 Feb;366:619-28; Roberts AM et al. Sci Transl Med, 2015 Jan;7:270ra6). This variant is classified as likely pathogenic based on ACMG and ClinGen recommendations.



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Carrier Status Findings

(i)

Carrier status determines the proband's risk for passing inherited genetic condition(s) to the children. Carriers are typically healthy/ asymptomatic. When an individual is found to be a carrier of a genetic condition, his or her relatives are at risk of carrying the same mutation. The patient should be encouraged to inform his or her relatives of the risk and the availability of carrier screening. If an individual is found to be a carrier of a specific condition, the patient's reproductive partner should be offered testing to receive informed genetic counseling about potential reproductive outcomes. If both partners are found to be carriers of a genetic condition, genetic counseling should be offered.

This proband is found to be a carrier for 3 genetic conditions.

Gene	Variant details	Zygosity	Annotations	Related disease (OMIM)- MOI Clinical assessment	Clinical significance (ACMG criteria*)
CEP250	chr20:35508949C>T NM_007186.6 c.6913C>T p.Arg2305* Exon/Intron rank:33/35 Nonsense Impact:High	Het	-ClinVar:N/A -Mastermind ID:N/A -gnomAD Total: N/A -Internal DB:N/A -Total MAF:N/A -REVEL:N/A -CADD (PHRED):48.0 -Splice-AI:0.21	Cone-rod dystrophy and hearing loss type 2 AR Carrier	Likely pathogenic PVS1, PM2
DHTKD1	chr10:12097997G>A NM_018706.7 c.1671+1G>A p.? Exon/Intron rank:8/17 Intron, Splice site donor Impact:High	Het	-ClinVar:N/A -Mastermind ID:N/A -gnomAD Total: 0.0000167 -Internal DB:N/A -Total MAF:N/A -REVEL:N/A -CADD (PHRED):33.0 -Splice-AI:1.0	Alpha-aminoadipic and alpha- ketoadipic aciduria AR Carrier	Likely pathogenic PVS1, PM2
TPRN	chr9:137200686GA>G NM_001128228.3 c.25delT p.Ser9fs*22 Exon/Intron rank:1/4 Frameshift Impact:High	Het	-ClinVar:N/A -Mastermind ID:N/A -gnomAD Total: N/A -Internal DB:N/A -Total MAF:N/A -REVEL:N/A -CADD (PHRED):N/A -Splice-AI:0.0	Deafness type 79 AR Carrier	Likely pathogenic PVS1, PM2

^{*:} The ACMG criteria are described under Methods /Variant interpretation section.



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Pharmacogenomic Associations



This test identified the following variants associated with drug use and dosing generated based on PharmCAT v2.8 (according to CPIC Guidelines (https://cpicpgx.org/guidelines). Pharmacogenetic tests, along with other information about patients and their disease or condition, can play an important role in drug therapy. When a health care provider is considering prescribing a drug, knowledge of a patient's genotype may be used to aid in determining a therapeutic strategy, determining an appropriate dosage, or assessing the likelihood of benefit or toxicity.

Genetic variants associated with drug use and dosing were identified.

Drugs	PGx Phenotype	Genes / Genotype	CPIC recommendations
Acenocoumarol	Not assigned	VKORC1: rs9923231 reference (C)/ rs9923231 variant (T)	No CPIC recommendations available for this drug-gene interaction.
Allopurinol	Normal metabolizer	ABCG2: rs2231142 reference (G)/ rs2231142 reference (G)	No CPIC recommendations available for this drug-gene interaction.
Amitriptyline	Normal metabolizer	CYP2C19:*1/*1	Initiate therapy with recommended starting dose.
Atazanavir	Intermediate metabolizer	UGT1A1:*1/*80+*28	There is no need to avoid prescribing of atazanavir based on UGT1A1 genetic test result. Inform the patient that some patients stop atazanavir because of jaundice (yellow eyes and skin), but that this patient's genotype makes this unlikely (less than about a 1 in 20 chance of stopping atazanavir because of jaundice).
Atorvastatin	Normal metabolizer	SLCO1B1:*1/*14	Prescribe desired starting dose and adjust doses based on disease-specific guidelines.
Azathioprine	Normal metabolizer	NUDT15:*1/*1 TPMT:*1/*1	Start with normal starting dose (e.g., 2-3 mg/kg/day) and adjust doses of azathioprine based on disease-specific guidelines. Allow 2 weeks to reach steady-state after each dose adjustment (PMID 20354201, 11302950, 15606506).
Capecitabine	Not assigned	DPYD:c.1601G>A (*4) DPYD:c.1627A>G (*5)	Based on genotype, there is no indication to change dose or therapy. Use label-recommended dosage and administration.
Celecoxib	Normal metabolizer	CYP2C9:*1/*1	Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals.
Citalopram	Normal metabolizer	CYP2C19:*1/*1	Initiate therapy with recommended starting dose
Clomipramine	Normal metabolizer	CYP2C19:*1/*1	Initiate therapy with recommended starting dose.
Clopidogrel	Normal metabolizer	CYP2C19:*1/*1	If considering clopidogrel, use at standard dose (75 mg/day)
Dapsone	Normal metabolizer	G6PD:B (reference)/ B (reference)	No reason to avoid based on G6PD status
Desflurane	Uncertain susceptibility	CACNA1S:Reference/ Reference RYR1:Reference/ Reference	Clinical findings, family history, further genetic testing and other laboratory data should guide use of halogenated volatile anesthetics or depolarizing muscle relaxants.
Dexlansoprazole	Normal metabolizer	CYP2C19:*1/*1	Initiate standard starting daily dose. Consider increasing dose by 50-100% for the treatment of H. pylori infection and erosive esophagitis. Daily dose may be given in divided doses. Monitor for efficacy.
Doxepin	Normal metabolizer	CYP2C19:*1/*1	Initiate therapy with recommended starting dose.
Efavirenz	Normal metabolizer	CYP2B6:*1/*2	Initiate efavirenz with standard dosing (600 mg/day)
Enflurane	Uncertain susceptibility	CACNA1S:Reference/ Reference RYR1:Reference/ Reference	Clinical findings, family history, further genetic testing and other laboratory data should guide use of halogenated volatile anesthetics or depolarizing muscle relaxants.
Escitalopram	Normal metabolizer	CYP2C19:*1/*1	Initiate therapy with recommended starting dose



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rouracil	Not assigned	DPYD:c.1601G>A (*4) DPYD:c.1627A>G (*5)	No CPIC recommendations available for this drug-gene interaction.
	Not assigned	DPYD:c.1601G>A (*4) DPYD:c.1627A>G (*5)	Based on genotype, there is no indication to change dose or therapy. Use label-recommended dosage and administration.
	Normal metabolizer	CYP2C9:*1/*1	Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals.
ao ca cirr	Normal metabolizer	CYP2C9:*1/*1 SLCO1B1:*1/*14	Prescribe desired starting dose and adjust doses of fluvastatin based on disease-specific guidelines.
,	Normal metabolizer	CYP2C9:*1/*1	No adjustments needed from typical dosing strategies. Subsequent doses should be adjusted according to therapeutic drug monitoring, response, and side effects. An HLA-B*15:02 negative test does not eliminate the risk of phenytoin-induced SJS/TEN and patients should be carefully monitored according to a usual standard.
	Uncertain susceptibility	CACNA1S:Reference/ Reference RYR1:Reference/ Reference	Clinical findings, family history, further genetic testing and other laboratory data should guide use of halogenated volatile anesthetics or depolarizing muscle relaxants.
	Normal metabolizer	CYP2C9:*1/*1	Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals.
	Normal metabolizer	CYP2C19:*1/*1	Initiate therapy with recommended starting dose.
	Intermediate metabolizer	UGT1A1:*1/*80+*28	No CPIC recommendations available for this drug-gene interaction.
	Uncertain susceptibility	CACNA1S:Reference/ Reference RYR1:Reference/ Reference	Clinical findings, family history, further genetic testing and other laboratory data should guide use of halogenated volatile anesthetics or depolarizing muscle relaxants.
	Ivacaftor non- responsive in cf patients	CFTR:Reference/ Reference	Ivacaftor is not recommended
	Normal metabolizer	CYP2C19:*1/*1	Initiate standard starting daily dose. Consider increasing dose by 50-100% for the treatment of H. pylori infection and erosive esophagitis. Daily dose may be given in divided doses. Monitor for efficacy.
	Normal metabolizer	CYP2C9:*1/*1	Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals.
	Normal metabolizer	SLCO1B1:*1/*14	Prescribe desired starting dose and adjust doses based on disease-specific guidelines.
	Normal metabolizer	CYP2C9:*1/*1	Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals.
1 1	Normal metabolizer	NUDT15:*1/*1 TPMT:*1/*1	Start with normal starting dose (e.g., 75 mg/m2/day or 1.5 mg/kg/day) and adjust doses of mercaptopurine (and of any other myelosuppressive therapy) without any special emphasis on mercaptopurine compared to other agents. Allow at least 2 weeks to reach steady-state after each dose adjustment (PMID 20354201, 16401827, 11302950).
	Uncertain susceptibility	CACNA1S:Reference/ Reference RYR1:Reference/ Reference	Clinical findings, family history, further genetic testing and other laboratory data should guide use of halogenated volatile anesthetics or depolarizing muscle relaxants.
,			
hylene blue	Normal metabolizer	G6PD:B (reference)/ B (reference)	No reason to avoid based on G6PD status
captopurine	Normal metabolizer Uncertain	NUDT15:*1/*1 TPMT:*1/*1 CACNA1S:Reference/ Reference	accordance with the prescribing information, use effective dosage for shortest duration consistent individual patient treatment goals. Start with normal starting dose (e.g., 75 mg/m2/mg/kg/day) and adjust doses of mercaptopurine other myelosuppressive therapy) without any speemphasis on mercaptopurine compared to other Allow at least 2 weeks to reach steady-state after adjustment (PMID 20354201, 16401827, 113029). Clinical findings, family history, further genetic to other laboratory data should guide use of haloge



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Omeprazole	Normal metabolizer	CYP2C19:*1/*1	Initiate standard starting daily dose. Consider increasing dose by 50-100% for the treatment of H. pylori infection and erosive esophagitis. Daily dose may be given in divided doses. Monitor for efficacy.
Pantoprazole	Normal metabolizer	CYP2C19:*1/*1	Initiate standard starting daily dose. Consider increasing dose by 50-100% for the treatment of H. pylori infection and erosive esophagitis. Daily dose may be given in divided doses. Monitor for efficacy.
Peginterferon alfa-2a	Not assigned	IFNL3:rs12979860 reference (C)/rs12979860 variant (T)	CPIC provides no genotype-based recommendations for the following genotype, after evaluating the evidence.
Peginterferon alfa-2b	Not assigned	IFNL3:rs12979860 reference (C)/rs12979860 variant (T)	CPIC provides no genotype-based recommendations for the following genotype, after evaluating the evidence.
Pegloticase	Normal metabolizer	G6PD:B (reference)/ B (reference)	No reason to avoid based on G6PD status
Phenprocoumon	Not assigned	VKORC1: rs9923231 reference (C)/ rs9923231 variant (T)	No CPIC recommendations available for this drug-gene interaction.
Phenytoin	Normal metabolizer	CYP2C9:*1/*1	No adjustments needed from typical dosing strategies. Subsequent doses should be adjusted according to therapeutic drug monitoring, response, and side effects. An HLA-B*15:02 negative test does not eliminate the risk of phenytoin-induced SJS/TEN and patients should be carefully monitored according to a usual standard.
Piroxicam	Normal metabolizer	CYP2C9:*1/*1	Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest effective dosage for shortest duration consistent with individual patient treatment goals.
Pitavastatin	Normal metabolizer	SLCO1B1:*1/*14	Prescribe desired starting dose and adjust doses based on disease-specific guidelines.
Pravastatin	Normal metabolizer	SLCO1B1:*1/*14	Prescribe desired starting dose and adjust doses based on disease-specific guidelines.
Primaquine	Normal metabolizer	G6PD:B (reference)/ B (reference)	No reason to avoid based on G6PD status
Quetiapine	Normal metabolizer	CYP3A4:*1/*1	No CPIC recommendations available for this drug-gene interaction.
Rasburicase	Normal metabolizer	G6PD:B (reference)/ B (reference)	No reason to avoid based on G6PD status
Ribavirin	Not assigned	IFNL3:rs12979860 reference (C)/rs12979860 variant (T)	CPIC provides no genotype-based recommendations for the following genotype, after evaluating the evidence.
Rosuvastatin	Normal metabolizer	ABCG2: rs2231142 reference (G)/ rs2231142 reference (G) SLC01B1:*1/*14	Prescribe desired starting dose and adjust doses of rosuvastatin based on disease-specific and specific population guidelines.
Sertraline	Normal metabolizer	CYP2B6:*1/*2 CYP2C19:*1/*1	Initiate therapy with recommended starting dose.
Sevoflurane	Uncertain susceptibility	CACNA1S:Reference/ Reference RYR1:Reference/ Reference	Clinical findings, family history, further genetic testing and other laboratory data should guide use of halogenated volatile anesthetics or depolarizing muscle relaxants.
Simvastatin	Normal metabolizer	SLCO1B1:*1/*14	Prescribe desired starting dose and adjust doses based on disease-specific guidelines.
Siponimod	Normal metabolizer	CYP2C9:*1/*1	No CPIC recommendations available for this drug-gene interaction.
Succinylcholine	Uncertain susceptibility	CACNA1S:Reference/ Reference RYR1:Reference/ Reference	Clinical findings, family history, further genetic testing and other laboratory data should guide use of halogenated volatile anesthetics or depolarizing muscle relaxants.
Tacrolimus	Poor metabolizer	CYP3A5:*3/*3	Initiate therapy with standard recommended dose. Use therapeutic drug monitoring to guide dose adjustments.
Tafenoquine	Normal metabolizer	G6PD:B (reference)/ B (reference)	No reason to avoid based on G6PD status
Tegafur	Not assigned	DPYD:c.1601G>A (*4) DPYD:c.1627A>G (*5)	No CPIC recommendations available for this drug-gene interaction.
Tenoxicam	Normal metabolizer	CYP2C9:*1/*1	Initiate therapy with recommended starting dose. In accordance with the prescribing information, use the lowest

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			effective dosage for shortest duration consistent with individual patient treatment goals.
Thioguanine	Normal metabolizer	NUDT15:*1/*1 TPMT:*1/*1	Start with normal starting dose (e.g., 40-60 mg/m2/day) and adjust doses of thioguanine and of other myelosuppressive therapy without any special emphasis on thioguanine. Allow 2 weeks to reach steady-state after each dose adjustment (PMID 20354201, 11037857).
Toluidine blue	Normal metabolizer	G6PD:B (reference)/ B (reference)	No reason to avoid based on G6PD status
Trimipramine	Normal metabolizer	CYP2C19:*1/*1	Initiate therapy with recommended starting dose.
Voriconazole	Normal metabolizer	CYP2C19:*1/*1	Initiate therapy with recommended standard of care dosing
Warfarin	Normal metabolizer	CYP2C9:*1/*1 CYP4F2:*1/*1 VKORC1: rs9923231 reference (C)/ rs9923231 variant (T) rs12777823:G/G	The updated guideline for pharmacogenetics-guided warfarin dosing is published by the <i>Clinical Pharmacogenetics Implementation Consortium</i> . The recommendations for dosing are for adult and pediatric patients that are specific to continental ancestry, and are based on genotypes from <i>CYP2C9</i> , <i>VKORC1</i> , <i>CYP4F2</i> , and rs12777823. For more information please visit: https://www.pharmgkb.org/guidelineAnnotation/PA16610494949





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TECHNICAL INFORMATION

Methods

Whole genome sequencing and data analysis. DNA was extracted from a biological sample and TruSeq Nano DNA High Throughput Library Prep Kit (Illumina®) was used to prepare libraries, which were sequenced using the 150nt pair-end protocol on an Illumina platform to yield an average coverage depth of 30x for the nuclear genome and at least 1000x for the mitochondrial genome. Bacterial contamination of a sample may impact the depth of coverage. Raw read alignment to reference genome GRCh38 and variant calling, including single nucleotide substitutions (SNVs), small insertions/deletions (Indels) and structural variants (SVs) with default parameters were performed using DRAGEN (version 4.2.4, Illumina). SNV and Indel variant annotation was performed by Geneyx (https://geneyx.com). Structural variants were annotated with ANNOTSV3.1 and in-house structural variant database to obtain allele frequencies. For the mitochondrial genome, variants with frequencies/heteroplasmy level ≥5% are detected. Genetic variants are described following the Human Genome Variation Society (HGVS) recommendations (www.hgvs.org). In case of complex DNA changes both International System for Human Cytogenomic Nomenclature (ISCN; https://iscn.karger.com/) and complex HGVS/ISCN recommendations (https://hgvs-nomenclature.org/stable/recommendations/ DNA/ complex/) are indicated.

Incidental genes: The gene panel is based on the ACMG (American College of Medical Genetics and Genomics) SF v.3.2 recommendations (https://www.gimjournal.org/article/S1098-3600(23)00879-1/fulltext).

ACTA2, ACTC1, ACVRL1, APC, APOB, ATP7B, BAG3, BMPR1A, BRCA1, BRCA2, BTD, CACNA1S, CALM1, CALM2, CALM3, CASQ2, COL3A1, DES, DSC2, DSG2, DSP, ENG, FBN1, FLNC, GAA, GLA, HFE, HNF1A, KCNH2, KCNQ1, LDLR, LMNA, MAX, MEN1, MLH1, MSH2, MSH6, MUTYH, MYBPC3, MYH11, MYH7, MYL2, MYL3, NF2, OTC, PALB2, PCSK9, PKP2, PMS2, PRKAG2, PTEN, RB1, RBM20, RET, RPE65, RYR1, RYR2, SCN5A, SDHAF2, SDHB, SDHC, SDHD, SMAD3, SMAD4, STK11, TGFBR1, TGFBR2, TMEM127, TMEM43, TNNC1, TNNI3, TNNT2, TP53, TPM1, TRDN, TSC1, TSC2, TTN, TTR, VHL, WT1.

Variant interpretation: All candidate variants were evaluated with respect to their pathogenicity and causality significance, which are categorized following ACMG guidelines (PMID: 25741868) and ClinGen recommendations (https://www.clinicalgenome.org). Only those variants with evidence for causing or contributing to disease are reported as primary findings. All variants are visually inspected in IGV prior to reporting. The variants are classified following the 5-tier classes: pathogenic, likely pathogenic, variants of uncertain significance (VUS), likely benign and benign variants are not reported.

VUSs are classified as "strong variants of unclear significance" when there is limited supporting evidence of pathogenicity (e.g., rare or absent from general population databases BUT in silico tools predict a deleterious effect on the protein consistent with the mechanism of disease; AND the gene has already been confirmed to be associated with the patient's observed phenotype). Incidental findings that do not correlate with the provided phenotype(s) are reported according to ACMG recommendations for reporting of incidental findings in using clinical exome and genome sequencing (PMID: 37347242), if consented. Only variants classified as pathogenic, likely pathogenic or uncertain (variants of unknown significance or VUS) according to the ACMG guidelines and associated with the patient's phenotype are listed among the primary findings. Variants of uncertain significance are categorized as strong candidates when they are extremely rare or absent in external and internal databases, are predicted to be deleterious, and the respective gene matches patient's phenotype (i.e. insufficient evidence available). Variants like risk factors (or risk alleles) and genetic modifiers, impacting the disease severity are reported ONLY when extensive scientific and clinical evidence is established.

ACMG criteria for classifying SNV/ Indels pathogenic variants: PVS1- Null variant (nonsense, frameshift, canonical +/-1 or 2 splice sites, initiation codon, single or multi-exon deletion) in a gene where loss of function (LOF) is a known mechanism of disease; PS1- Same amino acid change as a previously established pathogenic variant regardless of nucleotide change; PS2- De novo (both maternity and paternity confirmed) in a patient with the disease and no family history; PS3- Well-established in vitro or in vivo functional studies supportive of a damaging effect on the gene or gene product; PS4- The prevalence of the variant in affected individuals is significantly increased compared to the prevalence in controls; PM1- Located in a mutational hot spot and/or critical and well-established functional domain (e.g. active site of an enzyme) without benign variation; PM2- Absent from controls (or at extremely low frequency if recessive) in Exome Sequencing Project, 1000 Genomes, ExAC or gnomAD databases; PM3- For recessive disorders, detected in trans with a pathogenic variant; PM4-Protein length changes due to in-frame deletions/insertions in a non-repeat region or stop-loss variants; PM5- Novel missense change at an amino acid residue where a different missense change determined to be pathogenic has been seen before at the acid/protein level; PM6- Assumed de novo, but without confirmation of paternity and maternity; PP1- Co-segregation with disease in multiple affected family members in a gene definitively known to cause the disease; PP2- Missense variant in a gene that has a low rate of benign missense variation and where missense variants are a common mechanism of disease; PP3- Multiple lines of computational evidence support a deleterious effect on the gene or gene product (conservation, evolutionary, splicing impact, etc); PP4- Patient's phenotype or family history is highly specific for a disease with a single genetic etiology.



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ACMG criteria for classifying benign variants: BA1- Allele frequency is above 5% in Exome Sequencing Project, 1000 Genomes, ExAC or gnomAD databases; BS1- Allele frequency is greater than expected for the disorder; BS2- Observed in a healthy adult individual for a recessive (homozygous), dominant (heterozygous), or X-linked (hemizygous) disorder with full penetrance expected at an early age; BS3- Well-established in vitro or in vivo functional studies shows no damaging effect on protein function or splicing; BS4- Lack of segregation in affected members of a family; BP1- Missense variant in a gene for which primarily truncating variants are known to cause disease; BP2- Observed in trans with a pathogenic variant for a fully penetrant dominant gene/disorder; or observed in cis with a pathogenic variant in any inheritance pattern; BP3- In-frame deletions/insertions in a repetitive region without a known function; BP4- Multiple lines of computational evidence suggest no impact on gene or gene product (conservation, evolutionary, splicing impact, etc); BP5- Variant found in a case with an alternate molecular basis for disease; BP7- A synonymous (silent) variant for which splicing prediction algorithms predict no impact to the splice consensus sequence nor the creation of a new splice site AND the nucleotide is not highly conserved.

ACMG criteria for classifying CNV/ SV pathogenic variants: LOSS 1A. Contains protein-coding or other known functionally important elements; 1B. Does NOT contain protein-coding or any known functionally important elements; 2A. Complete overlap of an established HI gene/genomic region; 2B. Partial overlap of an established HI genomic region; 2C. Partial overlap with the 5' end of an established HI gene (3' end of the gene not involved); 2D. Partial overlap with the 3' end of an established HI gene (5' end of the gene not involved); 2E. Both breakpoints are within the same gene (intragenic CNV; gene-level sequence variant); 2F. Completely contained within an established benign CNV region; 2G. Overlaps an established benign CNV but includes additional genomic material; 2H. Two or more HI predictors suggest that AT LEAST ONE gene in the interval is haploinsufficient (HI); 3A. Number of protein-coding RefSeq genes wholly or partially included in the copy number loss (0-24 genes); 3B. Number of protein-coding RefSeq genes wholly or partially included in the copy number loss (25-34 genes); 3C. Number of protein-coding RefSeq genes wholly or partially included in the copy number loss (35+ genes); 4A-4C. Individual case evidence – de novo occurrences; 4D. Individual case evidence - inconsistent phenotype; 4E. Individual case evidence - unknown inheritance; 4F-4H. Individual case evidence – segregation among similarly affected family members; 4I-4K. Individual case evidence - Non-Segregations; 4L-4O. Case-control and population evidence; 5A. Observed copy number loss is DE NOVO; 5B-5D. Observed copy number loss is INHERITED; 5E. Observed copy number loss – NON-SEGREGATIONS; 5F-5H Other. GAIN: 1A. Contains protein-coding or other known functionally important elements; 1B. Does NOT contain protein-coding or any known functionally important elements; 2A. Complete overlap; the TS gene or minimal critical region is fully contained within the observed copy number gain; 2B. Partial overlap of an established TS region; 2C. Identical in gene content to the established benign copy number gain; 2D. Smaller than established benign copy number gain, breakpoint(s) does not interrupt protein-coding genes; 2E. Smaller than established benign copy number gain, breakpoint(s) potentially interrupts protein-coding gene; 2F. Larger than known benign copy number gain, does not include additional protein-coding genes; 2G. Overlaps a benign copy number gain but includes additional genomic material; 2H. HI gene fully contained within observed copy number gain; 2I. Both breakpoints are within the same gene (gene-level sequence variant, possibly resulting in loss of function (LOF)); 2J. One breakpoint is within an established HI gene, patient's phenotype is either inconsistent with what is expected for LOF of that gene OR unknown; 2K. One breakpoint is within an established HI gene, patient's phenotype is highly specific and consistent with what is expected for LOF of that gene; 2L. One or both breakpoints are within gene(s) of no established clinical significance; 3A. Number of protein-coding RefSeq genes wholly or partially included in the copy number gain (0-34 genes); 3B. Number of protein-coding RefSeq genes wholly or partially included in the copy number gain (35-49 genes); 3C. Number of protein-coding RefSeq genes wholly or partially included in the copy number gain (50+ genes); 4A-4C. Individual case evidence – de novo occurrences; 4D. Individual case evidence – inconsistent phenotype; 4E. Individual case evidence - unknown inheritance; 4F-4H. Individual case evidence - segregation among similarly affected family members; 4I-4K. Individual case evidence - Non-Segregations; 4L-4O. Case-control and population evidence; 5A. Observed copy number loss is DE NOVO; 5B-5D. Observed copy number loss is INHERITED; 5E. Observed copy number loss – NON-SEGREGATIONS; 5F-5H Other.

Point- based scoring framework: Pathogenic for 0.99 or more points; Liley pathogenic for 0.90 to 0.98 points; Variant of Uncertain Significance for scores between -0.89 to 0.89 points; Likely benign for scores between -0.90 to - 0.98; Benign for -0.99 or fewer points.

Consanguinity score (CS): this score is obtained from the DRAGEN's region of homozygosity calculation that considers homozygous single nucleotide variants on the autosomes. The higher the score, the closer the biological relationship of subjects' parents. A score above two (2) suggests consanguinity.

Pharmacogenomic variants

The design of the pharmacogenomics panel is based on PharmCAT v2.8.3 (https://pharmcat.org/) and CPIC v1.30.0 (https://cpicpgx.org/guidelines/). The list of genes and drugs covered by PharmCAT is provided under: https://pharmcat.org/Genes-Drugs

The covered drugs based on Anatomical Therapeutic Chemical (ATC) classification by PharmCAT v2.8.3 are:

i. Anti-cancer and immune response: Azathioprine, Capecitabine, Fluorouracil, Irinotecan, Mercaptopurine, Peginterferon alfa-2a, Peginterferon alfa-2b, Rasburicase, Siponimod, Tacrolimus, Tegafur, Thioguanine



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- ii. Blood and cardiovascular system: Acenocoumarol, Atorvastatin, Clopidogrel, Fluvastatin, Lovastatin, Methylene blue, Phenprocoumon, Pitavastatin, Pravastatin, Rosuvastatin, Simvastatin, Warfarin
- iii. Digestive system: Dexlansoprazole, Lansoprazole, Omeprazole, Pantoprazole
- iv. Infection control: Atazanavir, Dapsone, Efavirenz, Flucytosine, Nitrofurantoin, Primaquine, Ribavirin, Tafenoquine, Voriconazole
- v. Musculo-skeletal system: Allopurinol, Celecoxib, Flurbiprofen, Ibuprofen, Lornoxicam, Meloxicam, Pegloticase, Piroxicam, Succinylcholine, Tenoxicam
- vi. Nervous system: Amitriptyline, Citalopram, Clomipramine, Desflurane, Doxepin, Enflurane, Escitalopram, Fosphenytoin, Halothane, Imipramine, Isoflurane, Methoxyflurane, Phenytoin, Quetiapine, Sertraline, Sevoflurane, Trimipramine
- vii. Respiratory system: Ivacaftor
- viii. Other: Toluidine blue

The covered genes by PharmCAT v2.8.3 are: ABCG2, CACNA1S, CFTR, CYP2B6, CYP2C19, CYP2C9, CYP3A4, CYP3A5, CYP4F2, DPYD, G6PD, IFNL3, NUDT15, RYR1, SLCO1B1, TPMT, UGT1A1, VKORC1

PharmCAT is only able to generate recommendations based on the information provided to the software. The gene and variant information for all reported sections are interpreted directly from Arcensus-supplied data. Reported genotype calls are displayed with respect to the reference genome. Variants indicated as homozygous or heterozygous differ from the GRCh38/hg38 reference sequence (wild type).

For all genes, variation reported in the VCF file but NOT included in the gene definition table will not be considered during allele assignment. There is a possibility that any such variation results in a reduced or no activity allele which could lead to inaccurate phenotype and CPIC recommendation.

Structural variation star alleles that cannot be detected using VCF file data: CYP2B7-CYP2B6 hybrids: CYP2B6*29, CYP2B6*30; Partial and whole gene deletions: CYP2C19*36, CYP2C19*37, CYP4F2*16, SLCO1B1*48, SLCO1B1*49.

This test does not report polymorphisms other than those specifically listed, and mutations in other genes associated with drug metabolism will not be reported. All content is sourced from the CPIC database.

Limitations

myLifeGenome is NOT indicated for somatic variants in tumor samples, Alzheimer's risk assessment, prenatal samples, partial UPD (uniparental disomy), epigenetic modifications like methlyation (known to cause Prader-Willi, Russell Silver or Beckwith-Wiedemann syndromes), gene conversions (GBA/GBAP1, CYP21A2, NCF1 or VWF), D4Z4 repeat expansion (known to cause facioscapulohumeral muscular dystrophy), or low level of mosaicims (VAF<10%).

Pathogenic repeat expansions within the following genes can be determined accurately. Repeat expansion in genes outside the list (for eg. but not limited to ATN1, ATXN1, ATXN10, ATXN2, ATXN3, ATXN7, C9orf72, CACNA1A, CNBP, CSTB, FMR1, FXN, HTT, PABPN1, PHOX2B, PPP2R2B, PRNP, TBP) may not be reliably detected. Variants within high complexity genes like CFH, HLA, MUC5B, NOTCH2, NCL, PKD1 or RPGR genes might be detected, but recommendations to be confirmed by an orthogonal method provided.

The genetic result's interpretation is strongly dependent on the provided clinical information and family history. Misinterpretation may occur if this data is provided incorrectly or incompletely. Variant frequencies are subjected to changes due to growing variant databases and may result in reclassification of previously reported variants. The variants detected with this assay cover the whole genome, within (intragenic) and beyond (intergenic) genes. The detectable variant types include nucleotide substitution, small insertions/deletions, copy number variants (CNVs), inversions, translocations, and complex rearrangements. Variants may not be detected in low complexity genomic regions due to high sequence homology, pseudogenes, or highly repetitive sequences. This methodology detects events of mosaicism of single nucleotide variants (SVN) with an minor allele fraction of minimum of at least 10%.

It is possible that a particular genetic variant may not be recognized as the underlying cause of the genetic disorder due to incomplete scientific knowledge about the biological function of the gene and/or the impact of the variant on the expression and/or function of the gene.

Test Performance

Total number of reads: 618,227,002 Percentage of mapped reads to hg38: 97.52% Median coverage: 30.74x

Annotation Datasets:

1kGenome:2019-02 ACMG:2024-12 AcmgSv:2024-12 AlphaMissense:v2-t113 CADD:1.6 CamouflagedGenes:v1 ClinGen:2024-12 ClinVar:2024-12 CoLoRSdb:v1.0.0 Cytogenetic:2022-10 DbscSNV:1.1 DGVGold:v107_2020-03-02 DarkGenes:v1 DbNsfp:4.4a DbSnp:1405 DGVSV:v107_2020-02-25 ESP6500:2 GeneEnhancerSv:v6.1 GeneYX Version:v6.1 GeneyxRepeats:v1.1 Gerp:2010 gnomAD-exomes:2.1.1 gnomAD-genomes:3.1.2 gnomAD-mit:2.1.1 GnomADSV:v2.1 GnomADv4-exomes:4.1.0 GnomADv4-genomes:4.1.0 LitVar2:2024-12 MANE:v1.4 MasterMind:2024-07 MitoMap:2021-10 OMIM:2024-12 Phylop:2015-05 Revel:2016-03 Rmsk:2022-10 SnpEff:v5.2-2024-12 SpliceAI:1.3

Arcensus GmbH Friedrich-Barnewitz-Str.9, 18119 Rostock www.arcensus-diagnostics.com



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