

Donor 6040

Genetic Testing Summary

Fairfax Cryobank recommends reviewing this genetic testing summary with your healthcare provider to determine suitability.

Last Updated: 04/16/24

Donor Reported Ancestry: German

Jewish Ancestry: No

Genetic Test*	Result	Comments/Donor's Residual Risk**

Chromosome analysis (karyotype)	Normal male karyotype	No evidence of clinically significant chromosome abnormalities
Hemoglobin evaluation	Normal hemoglobin fractionation and MCV/MCH results	Reduced risk to be a carrier for sickle cell anemia, beta thalassemia, alpha thalassemia trait (aa/ and a-/a-) and other hemoglobinopathies
Cystic Fibrosis (CF) carrier screening	Negative by gene sequencing in the CFTR gene	1/440
Spinal Muscular Atrophy (SMA) carrier screening	Negative for deletions of exon 7 in the SMN1 gene	1/894
Expanded Genetic Disease Carrier Screening Panel attached- 283 diseases by gene sequencing	Carrier: Medium Chain Acyl-CoA Dehydrogenase Deficiency (ACADM) Negative for other genes sequenced.	Partner testing is recommended before using this donor.
Special Testing		
Genes: LAMA2, NAGA	Negative by gene sequencing	

*No single test can screen for all genetic disorders. A negative screening result significantly reduces, but cannot eliminate, the risk for these conditions in a pregnancy.

**Donor residual risk is the chance the donor is still a carrier after testing negative.



CARRIER SCREENING REPORT

Patient	Sample	Referring Doctor
Patient Name: Donor 6040 Date of Birth: Reference #: Indication: Carrier Testing Test Type: Expanded Carrier Screen (283) Minus TSE	Specimen Type: Blood Lab #: Date Collected: 5/15/2019 Date Received: 5/16/2019 Final Report: 5/30/2019	Fairfax Cryobank, Inc.

RESULT SUMMARY

THIS PATIENT WAS TESTED FOR 283 DISEASES. Please see Table 1 for list of diseases tested.

POSITIVE for medium chain acyl-CoA dehydrogenase deficiency

A heterozygous (one copy) pathogenic variant, c.464T>C, p.M155T, was detected in the ACADM gene

NEGATIVE for the remaining diseases

Recommendations

Testing the partner for the above positive disorder(s) and genetic counseling are recommended.

Please note that for female carriers of X-linked diseases, follow-up testing of a male partner is not indicated. In addition, CGG repeat analysis of *FMR1* for fragile X syndrome is not performed on males as repeat expansion of premutation alleles is not expected in the male germline.

Individuals of Asian, African, Hispanic and Mediterranean ancestry should also be screened for hemoglobinopathies by CBC and hemoglobin electrophoresis.

Consideration of residual risk by ethnicity after a negative carrier screen is recommended for the other diseases on the panel, especially in the case of a positive family history for a specific disorder.

Interpretation for medium chain acyl-CoA dehydrogenase deficiency

A heterozygous (one copy) pathogenic missense variant, c.464T>C, p.M155T, was detected in the *ACADM* gene (NM_000016.5). When this variant is present in trans with a pathogenic variant, it is considered to be causative for medium chain acyl-CoA dehydrogenase deficiency. Therefore, this individual is expected to be at least a carrier for medium chain acyl-CoA dehydrogenase deficiency. Heterozygous carriers are not expected to exhibit symptoms of this disease.



CARRIER SCREENING REPORT

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What is medium chain acyl-CoA dehydrogenase deficiency?

Medium chain acyl-CoA dehydrogenase (MCAD) deficiency is a pan-ethnic autosomal recessive condition caused by pathogenic variants in the gene *ACADM*. It prevents the body from releasing energy from fats. Symptoms often begin in infancy, although the clinical presentation is highly variable and some affected individuals do not show symptoms until adulthood if at all. MCAD deficiency causes metabolic crises, which present with lethargy and vomiting. Some infants may present with sudden death. Dietary management greatly reduces the risk of metabolic crises and allows affected individuals to live relatively normal lives. Although metabolic crises can be fatal, affected individuals who have a known diagnosis and receive proper care have normal life expectancy. Some *ACADM* variants are known to be associated with milder disease, although it is not possible to exactly predict the severity of disease based on the inherited variants.

This patient was tested for a panel of diseases using a combination of sequencing, targeted genotyping and copy number analysis. Please note that negative results reduce but do not eliminate the possibility that this individual is a carrier for one or more of the disorders tested. Please see Table 1 for a list of genes and diseases tested, and http://go.sema4.com/residualrisk for specific detection rates and residual risk by ethnicity. With individuals of mixed ethnicity, it is recommended to use the highest residual risk estimate. Only variants determined to be pathogenic or likely pathogenic are reported in this carrier screening test.

TEST SPECIFIC RESULTS

Alpha-thalassemia

NEGATIVE for alpha-thalassemia

HBA1 copy number: 2 HBA2 copy number: 2 No pathogenic copy number variants detected HBA1 and HBA2 sequence analysis: No pathogenic or likely pathogenic variants identified Reduced risk of being an alpha-thalassemia carrier (aa/aa)

Genes analyzed: *HBA1* (NM_000558.4) and *HBA2* (NM_000517.4) **Inheritance:** Autosomal Recessive

Recommendations

Individuals of Asian, African, Hispanic and Mediterranean ancestry should also be screened for hemoglobinopathies by CBC and hemoglobin electrophoresis.

Interpretation

No pathogenic or likely pathogenic copy number variants or sequence variants were detected in this patient, suggesting that four copies of the alpha-globin gene are present (aa/aa). Typically, individuals have four



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functional alpha-globin genes: 2 copies of *HBA1* and 2 copies of *HBA2*, whose expression is regulated by a cisacting regulatory element HS-40. Alpha-thalassemia carriers have three (silent carrier) or two (carrier of the alpha-thalassemia trait) functional alpha-globin genes with or without a mild phenotype. Individuals with only one functional alpha-globin gene have HbH disease with microcytic, hypochromic hemolytic anemia and hepatosplenomegaly. Loss of all four alpha-globin genes results in Hb Barts syndrome with the accumulation of Hb Barts in red blood cells and hydrops fetalis, which is fatal in utero or shortly after birth.

This individual was negative for all *HBA* deletions, duplications and variants that were tested. These negative results reduce but do not eliminate the possibility that this individual is a carrier. See *Table of Residual Risks Based on Ethnicity*. With individuals of mixed ethnicity, it is recommended to use the highest residual risk estimate.

Table of Residual Risks Based on Ethnicity

Ethnicity	Carrier Frequency	Detection Rate	Residual Risk
Caucasian	1 in 500	95%	1 in 10,000
African American	1 in 30	95%	1 in 580
Asian	1 in 20	95%	1 in 380
Worldwide	1 in 25	95%	1 in 480

Congenital Adrenal Hyperplasia (21-Hydroxylase Deficiency)

NEGATIVE for congenital adrenal hyperplasia (due to 21-hydroxylase deficiency) *CYP21A2* copy number: 2

No pathogenic copy number variants detected No pathogenic sequence variants detected in *CYP21A2* Reduced risk of being a congenital adrenal hyperplasia carrier

Genes analyzed: *CYP21A2* (NM_000500.6) **Inheritance:** Autosomal Recessive

Recommendations

Consideration of residual risk by ethnicity (see below) after a negative carrier screen is recommended, especially in the case of a positive family history of congenital adrenal hyperplasia.

Interpretation

This individual was negative for all pathogenic *CYP21A2* copy number variants that were tested, and no pathogenic or likely pathogenic variants were identified by sequence analysis. These negative results reduce but do not eliminate the possibility that this individual is a carrier. See *Table of Residual Risks Based on Ethnicity*. With individuals of mixed ethnicity, it is recommended to use the highest residual risk estimate.



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Table of Residual Risk Based On Ethnicity - Classic Congenital Adrenal Hyperplasia Due to 21-Hydroxylase Deficiency

Ethnicity	Carrier Frequency	Detection Rate	Residual Risk
Ashkenazi Jewish	1 in 40	>95%	1 in 780
Caucasian	1 in 67	>95%	1 in 1300
Worldwide	1 in 60	>95%	1 in 1200

Table of Residual Risk Based On Ethnicity - Non-Classic Congenital Adrenal Hyperplasia Due to 21-Hydroxylase Deficiency

Ethnicity	Carrier Frequency	Detection Rate	Residual Risk
Ashkenazi Jewish	1 in 7	>95%	1 in 120
Caucasian	1 in 11	>95%	1 in 200
Worldwide	1 in 16	>95%	1 in 300

Fragile X syndrome

Fragile X CGG triplet repeat expansion testing was not performed at this time, as the patient has either been previously tested or is a male. Sequencing of the *FMR1* gene by next generation sequencing did not identify any clinically significant variants.

Spinal Muscular Atrophy

NEGATIVE for spinal muscular atrophy

SMN1 Copy Number: 2 *SMN2* Copy Number: 2 c.*3+80T>G: Negative

Negative copy number result Decreased risk of being an *SMN1* silent (2+0) carrier (see *SMA Table*)

Genes analyzed: *SMN1* (NM_000344.3) and *SMN2* (NM_017411.3) **Inheritance:** Autosomal Recessive

Recommendations

Consideration of residual risk by ethnicity after a negative carrier screen is recommended, especially in the case of a positive family history for spinal muscular atrophy.



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Interpretation

This patient is negative for loss of *SMN1* copy number. Complete loss of *SMN1* is causative in spinal muscular atrophy (SMA). Two copies of *SMN1* were detected in this individual, which significantly reduces the risk of being an SMA carrier. Parallel testing to assess the presence of an *SMN1* duplication allele was also performed to detect a single nucleotide polymorphism (SNP), c.*3+80T>G, in intron 7 of the *SMN1* gene. This individual was found to be negative for this change and is therefore, at a decreased risk of being a silent (2+0) carrier, see *SMA Table* for residual risk estimates based on ethnicity.

SMA Table: Carrier detection and residual risk estimates before and after testing for c.*3+80T>G

Ethnicity	Carrier Frequency	Detection rate	Residual risk after negative result*	Detection rate with <i>SMN1</i> c.*3+80T>G	Residual risk c.*3+80T>G negative	Residual risk c.*3+80T>G positive
African American	1 in 85	71%	1 in 160	91%	1 in 455	1 in 49
Ashkenazi Jewish	1 in 76	90%	1 in 672	93%	1 in 978	1 in 10
East Asian	1 in 53	94%	1 in 864	95%	1 in 901	1 in 12
Caucasian	1 in 48	95%	1 in 803	95%	1 in 894	1 in 23
Latino	1 in 63	91%	1 in 609	94%	1 in 930	1 in 47
South Asian	1 in 103	87%	1 in 637	87%	1 in 637	1 in 608
Sephardic Jewish	1 in 34	96%	1 in 696	97%	1 in 884	1 in 12

*Residual risk with two copies *SMN1* detected using dosage sensitive methods. The presence of three or more copies of *SMN1* reduces the risk of being an *SMN1* carrier between 5 - 10 fold, depending on ethnicity. *FOR INDIVIDUALS WITH MIXED ETHNICITY, USE HIGHEST RESIDUAL RISK ESTIMATE* ^ Parental follow-up will be requested for confirmation

This case has been reviewed and electronically signed by Yaping Ryan Qian, Ph.D., FACMG, Laboratory Director

Laboratory Medical Consultant: George A. Diaz, M.D., Ph.D.



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Test Methods and Comments

Genomic DNA isolated from this patient was analyzed by one or more of the following methodologies, as applicable:

Fragile X CGG Repeat Analysis (Analytical Detection Rate >99%)

PCR amplification using Asuragen, Inc. AmplideX[®] *FMR1* PCR reagents followed by capillary electrophoresis for allele sizing was performed. Samples positive for *FMR1* CGG repeats in the premutation and full mutation size range were further analyzed by Southern blot analysis to assess the size and methylation status of the *FMR1* CGG repeat.

Genotyping (Analytical Detection Rate >99%)

Multiplex PCR amplification and allele specific primer extension analyses using the MassARRAY[®] System were used to identify variants that are complex in nature or are present in low copy repeats. Rare sequence variants may interfere with assay performance.

Multiplex Ligation-Dependent Probe Amplification (MLPA) (Analytical Detection Rate >99%)

MLPA[®] probe sets and reagents from MRC-Holland were used for copy number analysis of specific targets versus known control samples. False positive or negative results may occur due to rare sequence variants in target regions detected by MLPA probes. Analytical sensitivity and specificity of the MLPA method are both 99%.

For alpha thalassemia, the copy numbers of the *HBA1* and *HBA2* genes were analyzed. Alpha-globin gene deletions, triplications, and the Constant Spring (CS) mutation are assessed. This test is expected to detect approximately 90% of all alpha-thalassemia mutations, varying by ethnicity. Carriers of alpha-thalassemia with three or more *HBA* copies on one chromosome, and one or no copies on the other chromosome, may not be detected. With the exception of triplications, other benign alpha-globin gene polymorphisms will not be reported. Analyses of *HBA1* and *HBA2* are performed in association with long-range PCR of the coding regions followed by short-read sequencing.

For Duchenne muscular dystrophy, the copy numbers of all *DMD* exons were analyzed. Potentially pathogenic single exon deletions and duplications are confirmed by a second method. Analysis of *DMD* is performed in association with sequencing of the coding regions.

For congenital adrenal hyperplasia, the copy number of the *CYP21A2* gene was analyzed. This analysis can detect large deletions due to unequal meiotic crossing-over between *CYP21A2* and the pseudogene *CYP21A1P*. These 30-kb deletions make up approximately 20% of *CYP21A2* pathogenic alleles. This test may also identify certain point mutations in *CYP21A2* caused by gene conversion events between *CYP21A2* and *CYP21A1P*. Some carriers may not be identified by dosage sensitive methods as this testing cannot detect individuals with two copies (duplication) of the *CYP21A2* gene on one chromosome and loss of *CYP21A2* (deletion) on the other chromosome. Analysis of *CYP21A2* is performed in association with long-range PCR of the coding regions followed by short-read sequencing.

For spinal muscular atrophy (SMA), the copy numbers of the *SMN1* and *SMN2* genes were analyzed. The individual dosage of exons 7 and 8 as well as the combined dosage of exons 1, 4, 6 and 8 of *SMN1* and *SMN2* were assessed. Copy number gains and losses can be detected with this assay. Depending on ethnicity, 6 - 29 % of carriers will not be identified by dosage sensitive methods as this testing cannot detect individuals with two copies (duplication) of the *SMN1* gene on one chromosome and loss of *SMN1* (deletion) on the other chromosome (silent 2+0 carrier) or individuals that carry an intragenic mutation in *SMN1*. Please also note that 2% of individuals with SMA have an *SMN1* mutation that occurred *de novo*. Typically in these cases, only one parent is an SMA carrier.

The presence of the c.*3+80T>G (chr5:70,247,901T>G) variant allele in an individual with Ashkenazi Jewish or Asian ancestry is typically indicative of a duplication of *SMN1*. When present in an Ashkenazi Jewish or Asian individual with two copies of *SMN1*, c.*3+80T>G is likely indicative of a silent (2+0) carrier. In individuals with two copies of *SMN1* with African American, Hispanic or Caucasian ancestry, the presence or absence of c.*3+80T>G significantly increases or decreases, respectively, the likelihood of being a silent 2+0 silent carrier.

Pathogenic or likely pathogenic sequence variants in exon 7 may be detected during testing for the c.*3+80T>G variant allele; these will be reported if confirmed to be located in SMN1 using locus-specific Sanger primers

Pathogenic or likely pathogenic sequence variants in exon 7 may be detected during testing for the c.*3+80T>G variant allele; these will be reported if confirmed to be located in SMN1 using locus-specific Sanger primers.

MLPA for Gaucher disease (*GBA*), cystic fibrosis (*CFTR*), and non-syndromic hearing loss (*GJB2/GJB6*) will only be performed if indicated for confirmation of detected CNVs. If *GBA* analysis was performed, the copy numbers of exons 1, 3, 4, and 6 - 10 of the *GBA* gene (of 11 exons total) were analyzed. If *CFTR* analysis was performed, the copy numbers of all 27 *CFTR* exons were analyzed. If *GJB2/GJB6* analysis was

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performed, the copy number of the two *GJB*2 exons were analyzed, as well as the presence or absence of the two upstream deletions of the *GJB*2 regulatory region, del(*GJB*6-D13S1830) and del(*GJB*6-D13S1854).

Next Generation Sequencing (NGS) (Analytical Detection Rate >95%)

NGS was performed on a panel of genes for the purpose of identifying pathogenic or likely pathogenic variants.

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Agilent SureSelectTMQXT technology was used with a custom capture library to target the exonic regions and intron/exon splice junctions of the relevant genes, as well as a number of UTR, intronic or promoter regions that contain previously reported mutations. Samples were pooled and sequenced on the Illumina HiSeq 2500 platform in the Rapid Run mode or the Illumina NovaSeq platform in the Xp workflow, using 100 bp paired-end reads. The sequencing data was analyzed using a custom bioinformatics algorithm designed and validated in house.

The coding exons and splice junctions of the known protein-coding RefSeq genes were assessed for the average depth of coverage (minimum of 20X) and data quality threshold values. Most exons not meeting a minimum of >20X read depth across the exon are further analyzed by Sanger sequencing. Please note that several genomic regions present difficulties in mapping or obtaining read depth >20X. The exons contained within these regions are noted within Table 1 (as "Exceptions") and will not be reflexed to Sanger sequencing if the mapping quality or coverage is poor. Any variants identified during testing in these regions are confirmed by a second method and reported if determined to be pathogenic or likely pathogenic. However, as there is a possibility of false negative results within these regions, detection rates and residual risks for these genes have been calculated with the presumption that variants in these exons will not be detected, unless included in the MassARRAY[®] genotyping platform.

This test will detect variants within the exons and the intron-exon boundaries of the target regions. Variants outside these regions may not be detected, including, but not limited to, UTRs, promoters, and deep intronic areas, or regions that fall into the Exceptions mentioned above. This technology may not detect all small insertion/deletions and is not diagnostic for repeat expansions and structural genomic variation. In addition, a mutation(s) in a gene not included on the panel could be present in this patient.

Variant interpretation and classification was performed based on the American College of Medical Genetics Standards and Guidelines for the Interpretation of Sequence Variants (Richards et al, 2015). All potentially pathogenic variants may be confirmed by either a specific genotyping assay or Sanger sequencing, if indicated. Any benign variants, likely benign variants or variants of uncertain significance identified during this analysis will not be reported.

Copy Number Variant Analysis (Analytical Detection Rate >95%)

Large duplications and deletions were called from the relative read depths on an exon-by-exon basis using a custom exome hidden Markov model (XHMM) algorithm. Deletions or duplications determined to be pathogenic or likely pathogenic were confirmed by either a custom arrayCGH platform, quantitative PCR, or MLPA (depending on CNV size and gene content). While this algorithm is designed to pick up deletions and duplications of 2 or more exons in length, potentially pathogenic single-exon CNVs will be confirmed and reported, if detected.

Exon Array (Confirmation method) (Accuracy >99%)

The customized oligonucleotide microarray (Oxford Gene Technology) is a highly-targeted exon-focused array capable of detecting medically relevant microdeletions and microduplications at a much higher resolution than traditional aCGH methods. Each array matrix has approximately 180,000 60-mer oligonucleotide probes that cover the entire genome. This platform is designed based on human genome NCBI Build 37 (hg19) and the CGH probes are enriched to target the exonic regions of the genes in this panel.

Quantitative PCR (Confirmation method) (Accuracy >99%)

The relative quantification PCR is utilized on a Roche Universal Library Probe (UPL) system, which relates the PCR signal of the target region in one group to another. To test for genomic imbalances, both sample DNA and reference DNA is amplified with primer/probe sets that specific to the target region and a control region with known genomic copy number. Relative genomic copy numbers are calculated based on the standard $\Delta\Delta$ Ct formula.

Long-Range PCR (Analytical Detection Rate >99%)

Long-range PCR was performed to generate locus-specific amplicons for *CYP21A2*, *HBA1* and *HBA2* and *GBA*. The PCR products were then prepared for short-read NGS sequencing and sequenced. Sequenced reads were mapped back to the original genomic locus and run through the bioinformatics pipeline. If indicated, copy number from MLPA was correlated with the sequencing output to analyze the results. For *CYP21A2*, a certain percentage of healthy individuals carry a duplication of the *CYP21A2* gene, which has no clinical consequences. In cases where two copies of a gene are located on the same chromosome in tandem, only the second copy will be amplified and assessed for potentially pathogenic



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variants, due to size limitations of the PCR reaction. However, because these alleles contain at least two copies of the *CYP21A2* gene in tandem, it is expected that this patient has at least one functional gene in the tandem allele and this patient is therefore less likely to be a carrier. When an individual carries both a duplication allele and a pathogenic variant, or multiple pathogenic variants, the current analysis may not be able to determine the phase (cis/trans configuration) of the *CYP21A2* alleles identified. Family studies may be required in certain scenarios where phasing is required to determine the carrier status.

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Residual Risk Calculations

Carrier frequencies and detection rates for each ethnicity were calculated through the combination of internal curations of >28,000 variants and genomic frequency data from >138,000 individuals across seven ethnic groups in the gnomAD database. Additional variants in HGMD and novel deleterious variants were also incorporated into the calculation. Residual risk values are calculated using a Bayesian analysis combining the *a priori* risk of being a pathogenic mutation carrier (carrier frequency) and the detection rate. They are provided only as a guide for assessing approximate risk given a negative result, and values will vary based on the exact ethnic background of an individual. This report does not represent medical advice but should be interpreted by a genetic counselor, medical geneticist or physician skilled in genetic result interpretation and the relevant medical literature.

Sanger Sequencing (Confirmation method) (Accuracy >99%)

Sanger sequencing, as indicated, was performed using BigDye Terminator chemistry with the ABI 3730 DNA analyzer with target specific amplicons. It also may be used to supplement specific guaranteed target regions that fail NGS sequencing due to poor quality or low depth of coverage (<20 reads) or as a confirmatory method for NGS positive results. False negative results may occur if rare variants interfere with amplification or annealing.

Please note these tests were developed and their performance characteristics were determined by Mount Sinai Genomics, Inc. They have not been cleared or approved by the FDA. These analyses generally provide highly accurate information regarding the patient's carrier or affected status. Despite this high level of accuracy, it should be kept in mind that there are many potential sources of diagnostic error, including misidentification of samples, polymorphisms, or other rare genetic variants that interfere with analysis. Families should understand that rare diagnostic errors may occur for these reasons.

SELECTED REFERENCES

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Fragile X syndrome:

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Spinal Muscular Atrophy:

Luo M et al. An Ashkenazi Jewish SMN1 haplotype specific to duplication alleles improves pan-ethnic carrier screening for spinal muscular atrophy. *Genet Med.* 2014 16:149-56.

Ashkenazi Jewish Disorders:

Scott SA et al. Experience with carrier screening and prenatal diagnosis for sixteen Ashkenazi Jewish Genetic Diseases. *Hum. Mutat.* 2010 31:1-11.

Duchenne Muscular Dystrophy:

Flanigan KM et al. Mutational spectrum of DMD mutations in dystrophinopathy patients: application of modern diagnostic techniques to a large cohort. *Hum Mutat.* 2009 30:1657-66.

Variant Classification:

Richards S et al. Standards and guidelines for the interpretation of sequence variants: a joint consensus recommendation of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology. *Genet Med.* 2015 May;17(5):405-24

Additional disease-specific references available upon request.

Table 1. List of genes and diseases tested.

Please see http://go.sema4.com/residualrisk for specific detection rates and residual risk by ethnicity.



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Gene	Disease
ACADM	Medium Chain Acyl-CoA Dehydrogenase Deficiency
ABCB11	Progressive Familial Intrahepatic Cholestasis, Type 2
ABCC8	Familial Hyperinsulinism (ABCC8-Related)
ABCD1	Adrenoleukodystrophy, X-Linked
ACAD9	Mitochondrial Complex I Deficiency (ACAD9-Related)
ACADVL	Very Long Chain Acyl-CoA Dehydrogenase Deficiency
ACAT1	Beta-Ketothiolase Deficiency
ACOX1	Acyl-CoA Oxidase I Deficiency
ACSF3	Combined Malonic and Methylmalonic Aciduria
ADA	Adenosine Deaminase Deficiency
ADAMTS2	Ehlers-Danlos Syndrome, Type VIIC
AGA	Aspartylglycosaminuria
AGL	Glycogen Storage Disease, Type III
AGPS	Rhizomelic Chondrodysplasia Punctata, Type 3
AGXT	Primary Hyperoxaluria, Type 1
AIRE	Polyglandular Autoimmune Syndrome, Type 1
ALDH3A2	Sjogren-Larsson Syndrome
ALDOB	Hereditary Fructose Intolerance
ALG6	Congenital Disorder of Glycosylation, Type Ic
ALMS1	Alstrom Syndrome
ALPL	Hypophosphatasia
AMT	Glycine Encephalopathy (AMT-Related)
AQP2	Nephrogenic Diabetes Insipidus, Type II
ARSA	Metachromatic Leukodystrophy
ARSB	Mucopolysaccharidosis type VI
ASL	Argininosuccinic Aciduria
ASNS	Asparagine Synthetase Deficiency
ASPA	Canavan Disease
ASS1	Citrullinemia, Type 1
ΑΤΜ	Ataxia-Telangiectasia
ATP6V1B1	Renal Tubular Acidosis and Deafness
ΑΤΡΤΑ	Menkes Disease
ATP7B	Wilson Disease
ATRX	Alpha-Thalassemia Mental Retardation Syndrome
BBS1	Bardet-Biedl Syndrome (BBS1-Related)
BBS10	Bardet-Biedl Syndrome (BBS10-Related)
BBS12	Bardet-Biedl Syndrome (BBS12-Related)
BBS2	Bardet-Biedl Syndrome (BBS2-Related)
BCKDHA	Maple Syrup Urine Disease, Type 1a
BCKDHB	Maple Syrup Urine Disease, Type 1b
BCS1L	GRACILE Syndrome and Other BCS1L-Related Disorders
BLM	Bloom Syndrome
BSND	Bartter Syndrome, Type 4A
BTD	Biotinidase Deficiency
CAPN3	Limb-Girdle Muscular Dystrophy, Type 2A
CBS	Homocystinuria (CBS-Related)
CDH23	Usher Syndrome, Type ID
CEP290	Leber Congenital Amaurosis 10 and Other CEP290-Related Ciliopathies
CERKL	Retinitis Pigmentosa 26

Gene	Disease
CFTR	Cystic Fibrosis
СНМ	Choroideremia
CHRNE	Congenital Myasthenic Syndrome (CHRNE-Related)
CIITA	Bare Lymphocyte Syndrome, Type II
CLN3	Neuronal Ceroid-Lipofuscinosis (CLN3-Related)
CLN5	Neuronal Ceroid-Lipofuscinosis (CLN5-Related)
CLN6	Neuronal Ceroid-Lipofuscinosis (CLN6-Related)
CLN8	Neuronal Ceroid-Lipofuscinosis (CLN8-Related)
CLRN1	Usher Syndrome, Type III
CNGB3	Achromatopsia
COL27A1	Steel Syndrome
COL4A3	Alport Syndrome (COL4A3-Related)
COL4A4	Alport Syndrome (COL4A4-Related)
COL4A5	Alport Syndrome (COL4A5-Related)
COL7A1	Dystrophic Epidermolysis Bullosa
CPS1	Carbamoylphosphate Synthetase I Deficiency
CPT1A	Carnitine Palmitoyltransferase IA Deficiency
CPT2	Carnitine Palmitoyltransferase II Deficiency
CRB1	Leber Congenital Amaurosis 8 / Retinitis Pigmentosa 12 /
	Pigmented Paravenous Chorioretinal Atrophy
CTNS	Cystinosis
CTSK	Pycnodysostosis
CYBA	Chronic Granulomatous Disease (CYBA-related)
СҮВВ	Chronic Granulomatous Disease (CYBB-related)
CYP11B2	Corticosterone Methyloxidase Deficiency
CYP17A1	Congenital Adrenal Hyperplasia due to 17-Alpha-Hydroxylase Deficiency
CYP21A2	Classic Congenital Adrenal Hyperplasia due to 21- Hydroxylase Deficiency
CYP19A1	Aromatase Deficiency
CYP27A1	Cerebrotendinous Xanthomatosis
DCLRE1C	Omenn Syndrome / Severe Combined Immunodeficiency, Athabaskan-Type
DHCR7	Smith-Lemli-Opitz Syndrome
DHDDS	Retinitis Pigmentosa 59
DLD	Lipoamide Dehydrogenase Deficiency
DMD	Duchenne Muscular Dystrophy / Becker Muscular Dystrophy
DNAH5	Primary Ciliary Dyskinesia (DNAH5-Related)
DNAI1	Primary Ciliary Dyskinesia (DNAI1-Related)
DNAI2	Primary Ciliary Dyskinesia (DNAI2-related)
DYSF	Limb-Girdle Muscular Dystrophy, Type 2B
EDA	Hypohidrotic Ectodermal Dysplasia 1
EIF2B5	Leukoencephalopathy with Vanishing White Matter
EMD	Emery-Dreifuss Myopathy 1
ESCO2	Roberts Syndrome
ETFA	Glutaric Acidemia, Type IIa
ETFDH	Glutaric Acidemia, Type IIc
ETHE1	Ethylmalonic Encephalopathy
EVC	Ellis-van Creveld Syndrome (EVC-Related)
EYS	Retinitis Pigmentosa 25
F11	Factor XI Deficiency
F9	
FAH	Factor IX Deficiency Tyrosinemia, Type I

Gene Disease

Mail: One Gustave L. Levy Place, Box 1497 Specimens: 1428 Madison Ave, Atran Bldg, Rm 2-25 New York, NY 10029

Disease

Gene



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FAM161A FANCC FANCC FANCG FH FKRP FKTN FMR1 G6PC GAA GALC GALK1	Retinitis Pigmentosa 28 Fanconi Anemia, Group A Fanconi Anemia, Group C Fanconi Anemia, Group G Fumarase Deficiency Limb-Girdle Muscular Dystrophy, Type 2I Walker-Warburg Syndrome and Other FKTN-Related Dystrophies Fragile X Syndrome Glycogen Storage Disease, Type Ia Glycogen Storage Disease, Type II Krabbe Disease Galactokinase Deficiency	
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FKTN FMR1 G6PC GAA GALC	Walker-Warburg Syndrome and Other FKTN-Related Dystrophies Fragile X Syndrome Glycogen Storage Disease, Type Ia Glycogen Storage Disease, Type II Krabbe Disease Galactokinase Deficiency	
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G6PC GAA GALC	Glycogen Storage Disease, Type Ia Glycogen Storage Disease, Type II Krabbe Disease Galactokinase Deficiency	
GAA GALC	Glycogen Storage Disease, Type II Krabbe Disease Galactokinase Deficiency	
GALC	Krabbe Disease Galactokinase Deficiency	
	Galactokinase Deficiency	
GALK1		
GALT	Galactosemia	
GAMT	Cerebral Creatine Deficiency Syndrome 2	
GBA	Gaucher Disease	
GBE1	Glycogen Storage Disease, Type IV / Adult Polyglucosan Body Disease	
GCDH	Glutaric Acidemia, Type I	
GFM1	Combined Oxidative Phosphorylation Deficiency 1	
GJB1	Charcot-Marie-Tooth Disease, X-Linked	
GJB2†	Non-Syndromic Hearing Loss (GJB2-Related)	
GLA	Fabry Disease	
GLB1	Mucopolysaccharidosis Type IVb / GM1 Gangliosidosis	
GLDC	Glycine Encephalopathy (GLDC-Related)	
GLE1	Lethal Congenital Contracture Syndrome 1 / Lethal Arthrogryposis with Anterior Horn Cell Disease	
GNE	Inclusion Body Myopathy 2	
GNPTAB	Mucolipidosis II / IIIA	
GNPTG	Mucolipidosis III Gamma	
GNS	Mucopolysaccharidosis Type IIID	
GP1BA	Bernard-Soulier Syndrome, Type A1	
GP9	Bernard-Soulier Syndrome, Type C	
GPR56	Bilateral Frontoparietal Polymicrogyria	
GRHPR	Primary Hyperoxaluria, Type 2	
HADHA	Long-Chain 3-Hydroxyacyl-CoA Dehydrogenase Deficiency	
HAX1	Congenital Neutropenia (HAX1-Related)	
HBA1/HBA2	Alpha-Thalassemia	
HBB	Beta-Globin-Related Hemoglobinopathies	
HEXA	Tay-Sachs Disease	
HEXB	Sandhoff Disease	
HFE2	Hemochromatosis, Type 2A	
HGSNAT	Mucopolysaccharidosis Type IIIC	
HLCS	Holocarboxylase Synthetase Deficiency	
HMGCL	HMG-CoA Lyase Deficiency	
HOGA1	Primary Hyperoxaluria, Type 3	
HPS1	Hermansky-Pudlak Syndrome, Type 1	
HPS3	Hermansky-Pudlak Syndrome, Type 3	
HSD17B4	D-Bifunctional Protein Deficiency	
HSD3B2	3-Beta-Hydroxysteroid Dehydrogenase Type II Deficiency	
HYAL1	Mucopolysaccharidosis type IX	
HYLS1	Hydrolethalus Syndrome	
IDS	Mucopolysaccharidosis Type II	
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IDUA	Mucopolysaccharidosis Type I
IKBKAP	Familial Dysautonomia
IL2RG	X-Linked Severe Combined Immunodeficiency
IVD	Isovaleric Acidemia
KCNJ11	Familial Hyperinsulinism (KCNJ11-Related)
LAMA3	Junctional Epidermolysis Bullosa (LAMA3-Related)
LAMB3	Junctional Epidermolysis Bullosa (LAMB3-Related)
LAMC2	Junctional Epidermolysis Bullosa (LAMC2-Related)
LCA5	Leber Congenital Amaurosis 5
LDLR	Familial Hypercholesterolemia
LDLRAP1	Familial Autosomal Recessive Hypercholesterolemia
LHX3	Combined Pituitary Hormone Deficiency 3
LIFR	Stuve-Wiedemann Syndrome
LIPA	Wolman Disease / Cholesteryl Ester Storage Disease
LOXHD1	Deafness, Autosomal Recessive 77
LPL	Lipoprotein Lipase Deficiency
LRPPRC	Leigh Syndrome, French-Canadian Type
MAN2B1	Alpha-Mannosidosis
MCCC1	3-Methylcrotonyl-CoA Carboxylase Deficiency (MCCC1-Related)
MCCC2	3-Methylcrotonyl-CoA Carboxylase Deficiency (MCCC2-Related)
MCOLN1	Mucolipidosis IV
MED17	Infantile Cerebral and Cerebellar Atrophy
MEFV	Familial Mediterranean Fever
MESP2	Spondylothoracic Dysostosis
MFSD8	Neuronal Ceroid-Lipofuscinosis (MFSD8-Related)
MKS1	Meckel syndrome 1 / Bardet-Biedl Syndrome 13
MLC1	Megalencephalic Leukoencephalopathy with Subcortical Cysts
MMAA	Methylmalonic Acidemia (MMAA-Related)
ММАВ	Methylmalonic Acidemia (MMAB-Related)
MMACHC	Methylmalonic Aciduria and Homocystinuria, Cobalamin C Type
MMADHC	Methylmalonic Aciduria and Homocystinuria, Cobalamin D Type
MPI	Congenital Disorder of Glycosylation, Type Ib
MPL	Congenital Amegakaryocytic Thrombocytopenia
MPV17	Mitochondrial DNA Depletion Syndrome 6 / Navajo
	Neurohepatopathy
MTHFR	Homocystinuria due to MTHFR Deficiency
MTM1	Myotubular Myopathy 1
MTRR	Homocystinuria, cbIE Type
MTTP	Abetalipoproteinemia
MUT	Methylmalonic Acidemia (MUT-Related)
MYO7A	Usher Syndrome, Type IB
NAGLU	Mucopolysaccharidosis Type IIIB
NAGS	N-Acetylglutamate Synthase Deficiency
NBN	Nijmegen Breakage Syndrome
NDRG1	Charcot-Marie-Tooth Disease, Type 4D
NDUFAF5	Mitochondrial Complex I Deficiency (NDUFAF5-Related)
NDUFS6	Mitochondrial Complex I Deficiency (NDUFS6-Related)
NEB	Nemaline Myopathy 2
NPC1	Niemann-Pick Disease, Type C (NPC1-Related)
NPC2	Niemann-Pick Disease, Type C (NPC2-Related)
NPHS1	Nephrotic Syndrome (NPHS1-Related) / Congenital Finnish Nephrosis



CARRIER SCREENING REPORT

Patient: Donor 6040

DOB:

Lab #:

NPHS2 Nephrotic Syndrome (NPHS2-Related) / Steroid-Resistant Nephrotic Syndrome NR2E3 Enhanced S-Cone Syndrome NR7K1 Congenital Insensitivity to Pain with Anhidrosis OAT Ornithine Aminotransferase Deficiency OPA3 3-Methylglutaconic Aciduria, Type III OTC Ornithine Transcarbomylase Deficiency PAH Phenylalanine Hydroxylase Deficiency PACA Propionic Acidemia (PCCA-Related) PCCB Propionic Acidemia (PCCB-Related) PCDH15 Usher Syndrome, Type IF PDHA1 Pyruvate Dehydrogenase E1-Alpha Deficiency PDHB Pyruvate Dehydrogenase E1-Beta Deficiency PEX1 Zellweger Syndrome Spectrum (PEX1-Related) PEX2 Zellweger Syndrome Spectrum (PEX2-Related) PEX6 Zellweger Syndrome Spectrum (PEX4-Related) PEX7 Rhizonelic Chondrodysplasia Punctata, Type 1 PFKM Glycogen Storage Disease, Type VII PHGDH 3-Phosphoglycorate Dehydrogenase Deficiency PKM2 Congenital Disorder of Glycosylation, Type Ia PMM2 Congenital Disorder of Glycosylation, Type Ia PMM2 Congenital Muscu	Gene	Disease
NTRK1 Congenital Insensitivity to Pain with Anhidrosis OAT Ornithine Aminotransferase Deficiency OPA3 3-Methylglutaconic Aciduria, Type III OTC Ornithine Transcarbomylase Deficiency PAH Phenylalanine Hydroxylase Deficiency PCCA Propionic Acidemia (PCCA-Related) PCCB Propionic Acidemia (PCCB-Related) PCCB Propionic Acidemia (PCCB-Related) PCDH15 Usher Syndrome, Type IF PDHA1 Pytruvate Dehydrogenase E1-Alpha Deficiency PEX1 Zellweger Syndrome Spectrum (PEX1-Related) PEX2 Zellweger Syndrome Spectrum (PEX6-Related) PEX6 Zellweger Syndrome Spectrum (PEX6-Related) PEX7 Rhizomelic Chondrodysplasia Punctata, Type 1 PFKM Glycogen Storage Disease, Type VII PHGDH Polycystic Kidney Disease, Autosomal Recessive PMM2 Congenital Disorder of Glycosylation, Type Ia PMM2 Congenital Muscular Dystrophy-Dystroglycanopathies PPT1 Neuronal Cercid-Lipofuscinosis (PT11-Related) PROP1 Combined SAP Deficiency PYS 6-Pyruvoyl-Tetrahydropterin Synthase Deficiency PTS1 Mitochondrial Myop	NPHS2	
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OPA3 3-Methylglutaconic Aciduria, Type III OTC Ornithine Transcarbomylase Deficiency PAH Phenylalanine Hydroxylase Deficiency PCCA Propionic Acidemia (PCCA-Related) PCCB Propionic Acidemia (PCCB-Related) PCDH15 Usher Syndrome, Type IF PDHA1 Pyruvate Dehydrogenase E1-Alpha Deficiency PDHB Pyruvate Dehydrogenase E1-Beta Deficiency PEX1 Zellweger Syndrome Spectrum (PEX1-Related) PEX2 Zellweger Syndrome Spectrum (PEX2-Related) PEX6 Zellweger Syndrome Spectrum (PEX4-Related) PEX7 Rhizomelic Chondrodysplasia Punctata, Type 1 PFKM Glycogen Storage Disease, Type VII PHGDH 3-Phosphoglycerate Dehydrogenase Deficiency PKMD1 Polycystic Kidney Disease, Autosomal Recessive PMM2 Congenital Disorder of Glycosylation, Type Ia POMGNT1 Muscle-Eye-Brain Disease and Other POMGNT1-Related Congenital Muscular Dystrophy-Dystroglycanopathies PPT1 Neuronal Ceroid-Lipofuscinosis (PPT1-Related) PROP1 Combined SAP Deficiency PVS6 Glycogen Storage Disease, Type V RAB23 Carpenter Syndrome <th>NTRK1</th> <th>Congenital Insensitivity to Pain with Anhidrosis</th>	NTRK1	Congenital Insensitivity to Pain with Anhidrosis
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PDHBPyruvate Dehydrogenase E1-Beta DeficiencyPEX1Zellweger Syndrome Spectrum (PEX1-Related)PEX10Zellweger Syndrome Spectrum (PEX10-Related)PEX2Zellweger Syndrome Spectrum (PEX2-Related)PEX6Zellweger Syndrome Spectrum (PEX6-Related)PEX7Rhizomelic Chondrodysplasia Punctata, Type 1PFKMGlycogen Storage Disease, Type VIIPHGDH3-Phosphoglycerate Dehydrogenase DeficiencyPKHD1Polycystic Kidney Disease, Autosomal RecessivePMM2Congenital Disorder of Glycosylation, Type IaPOMGNT1Muscle-Eye-Brain Disease and Other POMGNT1-Related Congenital Muscular Dystrophy-DystroglycanopathiesPPT1Neuronal Ceroid-Lipofuscinosis (PPT1-Related)PROP1Combined Pituitary Hormone Deficiency 2PRS1Charcot-Marie-Tooth Disease, Type 5 / Arts syndromePSAPCombined SAP DeficiencyPTS6-Pyruvoyl-Tetrahydropterin Synthase DeficiencyPUS1Mitochondrial Myopathy and Sideroblastic Anemia 1PYGMGlycogen Storage Disease, Type VRAB23Carpenter SyndromeRAG2Omenn Syndrome (RAG2-Related)RARS2Pontocerebellar Hypoplasia, Type 6RDH12Leber Congenital Amaurosis 13RMRPCartilage-Hair HypoplasiaRPE65Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20RPGRIP1LJoubert Syndrome 7 / Meckel Syndrome 5 / COACH SyndromeRS1X-Linked Juvenile RetinoschisisRTEL1Dyskeratosis Congenita (RTEL1-Related)SACSAutosomal Recessive Spastic Ataxia of Ch	PCDH15	Usher Syndrome, Type IF
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PEX7 Rhizomelic Chondrodysplasia Punctata, Type 1 PFKM Glycogen Storage Disease, Type VII PHGDH 3-Phosphoglycerate Dehydrogenase Deficiency PKHD1 Polycystic Kidney Disease, Autosomal Recessive PMM2 Congenital Disorder of Glycosylation, Type Ia POMGNT1 Muscle-Eye-Brain Disease and Other POMGNT1-Related Congenital Muscular Dystrophy-Dystroglycanopathies PPT1 Neuronal Ceroid-Lipofuscinosis (PPT1-Related) PROP1 Combined Pituitary Hormone Deficiency 2 PRPS1 Charcot-Marie-Tooth Disease, Type 5 / Arts syndrome PSAP Combined SAP Deficiency PTS 6-Pyruvoyl-Tetrahydropterin Synthase Deficiency PUS1 Mitochondrial Myopathy and Sideroblastic Anemia 1 PYGM Glycogen Storage Disease, Type V RAB23 Carpenter Syndrome RAG2 Omenn Syndrome (RAG2-Related) RARS2 Pontocerebellar Hypoplasia, Type 6 RDH12 Leber Congenital Amaurosis 13 RMRP Cartilage-Hair Hypoplasia RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis	PEX2	Zellweger Syndrome Spectrum (PEX2-Related)
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PKHD1 Polycystic Kidney Disease, Autosomal Recessive PMM2 Congenital Disorder of Glycosylation, Type Ia POMGNT1 Muscle-Eye-Brain Disease and Other POMGNT1-Related Congenital Muscular Dystrophy-Dystroglycanopathies PPT1 Neuronal Ceroid-Lipofuscinosis (PPT1-Related) PROP1 Combined Pituitary Hormone Deficiency 2 PRPS1 Charcot-Marie-Tooth Disease, Type 5 / Arts syndrome PSAP Combined SAP Deficiency PTS 6-Pyruvoyl-Tetrahydropterin Synthase Deficiency PUS1 Mitochondrial Myopathy and Sideroblastic Anemia 1 PYGM Glycogen Storage Disease, Type V RAB23 Carpenter Syndrome RAG2 Omenn Syndrome (RAG2-Related) RARS2 Pontocerebellar Hypoplasia, Type 6 RDH12 Leber Congenital Amaurosis 13 RMRP Cartilage-Hair Hypoplasia RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMH	PFKM	Glycogen Storage Disease, Type VII
PMM2 Congenital Disorder of Glycosylation, Type Ia POMGNT1 Muscle-Eye-Brain Disease and Other POMGNT1-Related Congenital Muscular Dystrophy-Dystroglycanopathies PPT1 Neuronal Ceroid-Lipofuscinosis (PPT1-Related) PROP1 Combined Pituitary Hormone Deficiency 2 PRPS1 Charcot-Marie-Tooth Disease, Type 5 / Arts syndrome PSAP Combined SAP Deficiency PTS 6-Pyruvoyl-Tetrahydropterin Synthase Deficiency PUS1 Mitochondrial Myopathy and Sideroblastic Anemia 1 PYGM Glycogen Storage Disease, Type V RAB23 Carpenter Syndrome RAG2 Omenn Syndrome (RAG2-Related) RARS2 Pontocerebellar Hypoplasia, Type 6 RDH12 Leber Congenital Amaurosis 13 RMRP Cartilage-Hair Hypoplasia RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	PHGDH	3-Phosphoglycerate Dehydrogenase Deficiency
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PRPS1Charcot-Marie-Tooth Disease, Type 5 / Arts syndromePSAPCombined SAP DeficiencyPTS6-Pyruvoyl-Tetrahydropterin Synthase DeficiencyPUS1Mitochondrial Myopathy and Sideroblastic Anemia 1PYGMGlycogen Storage Disease, Type VRAB23Carpenter SyndromeRAG2Omenn Syndrome (RAG2-Related)RARS2Pontocerebellar Hypoplasia, Type 6RDH12Leber Congenital Amaurosis 13RMRPCartilage-Hair HypoplasiaRPE65Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20RPGRIP1LJoubert Syndrome 7 / Meckel Syndrome 5 / COACH SyndromeRS1X-Linked Juvenile RetinoschisisRTEL1Dyskeratosis Congenita (RTEL1-Related)SACSAutosomal Recessive Spastic Ataxia of Charlevoix-SaguenaySAMHD1Aicardi-Goutières Syndrome (SAMHD1-Related)	PPT1	Neuronal Ceroid-Lipofuscinosis (PPT1-Related)
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PYGM Glycogen Storage Disease, Type V RAB23 Carpenter Syndrome RAG2 Omenn Syndrome (RAG2-Related) RAPSN Congenital Myasthenic Syndrome (RAPSN-Related) RARS2 Pontocerebellar Hypoplasia, Type 6 RDH12 Leber Congenital Amaurosis 13 RMRP Cartilage-Hair Hypoplasia RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	PTS	6-Pyruvoyl-Tetrahydropterin Synthase Deficiency
RAB23 Carpenter Syndrome RAG2 Omenn Syndrome (RAG2-Related) RAPSN Congenital Myasthenic Syndrome (RAPSN-Related) RARS2 Pontocerebellar Hypoplasia, Type 6 RDH12 Leber Congenital Amaurosis 13 RMRP Cartilage-Hair Hypoplasia RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	PUS1	Mitochondrial Myopathy and Sideroblastic Anemia 1
RAG2 Omenn Syndrome (RAG2-Related) RAPSN Congenital Myasthenic Syndrome (RAPSN-Related) RARS2 Pontocerebellar Hypoplasia, Type 6 RDH12 Leber Congenital Amaurosis 13 RMRP Cartilage-Hair Hypoplasia RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	PYGM	Glycogen Storage Disease, Type V
RAPSN Congenital Myasthenic Syndrome (RAPSN-Related) RARS2 Pontocerebellar Hypoplasia, Type 6 RDH12 Leber Congenital Amaurosis 13 RMRP Cartilage-Hair Hypoplasia RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	RAB23	Carpenter Syndrome
RARS2 Pontocerebellar Hypoplasia, Type 6 RDH12 Leber Congenital Amaurosis 13 RMRP Cartilage-Hair Hypoplasia RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	RAG2	Omenn Syndrome (RAG2-Related)
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RPE65 Leber Congenital Amaurosis 2 / Retinitis pigmentosa 20 RPGRIP1L Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome RS1 X-Linked Juvenile Retinoschisis RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	RDH12	Leber Congenital Amaurosis 13
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RTEL1 Dyskeratosis Congenita (RTEL1-Related) SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	RPGRIP1L	Joubert Syndrome 7 / Meckel Syndrome 5 / COACH Syndrome
SACS Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	RS1	X-Linked Juvenile Retinoschisis
SAMHD1 Aicardi-Goutières Syndrome (SAMHD1-Related)	RTEL1	Dyskeratosis Congenita (RTEL1-Related)
	SACS	Autosomal Recessive Spastic Ataxia of Charlevoix-Saguenay
SEBSECS Programsive Careballe Carebral Atrantiv	SAMHD1	Aicardi-Goutières Syndrome (SAMHD1-Related)
SEFSESS FIDGRESSIVE CEREBEIIO-CEREDIAL ALTOPHY	SEPSECS	Progressive Cerebello-Cerebral Atrophy

Gene	Disease
SGCA	Limb-Girdle Muscular Dystrophy, Type 2D
SGCB	Limb-Girdle Muscular Dystrophy, Type 2E
SGCG	Limb-Girdle Muscular Dystrophy, Type 2C
SGSH	Mucopolysaccharidosis Type IIIA
SLC12A3	Gitelman Syndrome
SLC12A6	Andermann Syndrome
SLC17A5	Salla Disease
SLC22A5	Primary Carnitine Deficiency
SLC25A13	Citrin Deficiency
SLC25A15	Hyperornithinemia-Hyperammonemia-Homocitrullinuria Syndrome
SLC26A2	Sulfate Transporter-Related Osteochondrodysplasia
SLC26A4	Pendred Syndrome
SLC35A3	Arthrogryposis, Mental Retardation, and Seizures
SLC37A4	Glycogen Storage Disease, Type Ib
SLC39A4	Acrodermatitis Enteropathica
SLC4A11	Corneal Dystrophy and Perceptive Deafness
SLC6A8	Cerebral Creatine Deficiency Syndrome 1
SLC7A7	Lysinuric Protein Intolerance
SMARCAL1	Schimke Immunoosseous Dysplasia
SMN1	Spinal Muscular Atrophy
SMPD1	Niemann-Pick Disease (SMPD1-Related)
STAR	Lipoid Adrenal Hyperplasia
SUMF1	Multiple Sulfatase Deficiency
TCIRG1	Osteopetrosis 1
TECPR2	Hereditary Spastic Paraparesis 49
TFR2	Hemochromatosis, Type 3
TGM1	Lamellar Ichthyosis, Type 1
тн	Segawa Syndrome
TMEM216	Joubert Syndrome 2
TPP1	Neuronal Ceroid-Lipofuscinosis (TPP1-Related)
TRMU	Acute Infantile Liver Failure
TSFM	Combined Oxidative Phosphorylation Deficiency 3
ΤΤΡΑ	Ataxia With Isolated Vitamin E Deficiency
ТҮМР	Myoneurogastrointestinal Encephalopathy
USH1C	Usher Syndrome, Type IC
USH2A	Usher Syndrome, Type IIA
VPS13A	Choreoacanthocytosis
VPS13B	Cohen Syndrome
VPS45	Congenital Neutropenia (VPS45-Related)
VRK1	Pontocerebellar Hypoplasia, Type 1A
VSX2	Microphthalmia / Anophthalmia
WNT10A	Odonto-Onycho-Dermal Dysplasia / Schopf-Schulz-Passarge Syndrome

† Please note that GJB2 testing includes testing for the two upstream deletions, del(GJB6-D13S1830) and del(GJB6-D13S1854) (PMID: 11807148 and 15994881)

PREVENTION GENETICS

NAME 6040, Donor

PATIENT INFORMATION	SPECIMEN INFORMATION	PROVIDER INFORMATION
6040, Donor ID#: DOB: Sex: Male	Type: Whole Blood Collected: March 09, 2021 Received: March 12, 2021 PG ID:	Harvey Stern, MD, PhD Suzanne Seitz, MS, MPA Fairfax Cryobank

MOLECULAR GENETICS REPORT: LAMA2 Gene Sequencing with CNV Detection

SUMMARY OF RESULTS NEGATIVE

RESULTS AND INTERPRETATIONS: In this patient, for the *LAMA2* gene, we found no sequence variants that are likely to be a primary cause of disease.

This patient is apparently negative for copy number variants (CNVs) within the genomic regions of this test.

These results should be interpreted in context of clinical findings, family history and other laboratory data. All genetic tests have limitations. See limitations and other information for this test on the following page(s).

NOTES: Since this test is performed using exome capture probes, a reflex to any of our exome-based tests is available (PGxome, PGxome Custom Panels).

GENE(S) ANALYZED: LAMA2

SUMMARY STATISTICS:

Pipeline	Version	Average NGS Coverage	Fraction Bases Covered with NGS
Infinity_Pipeline	1.8.3	81x	99.1%

Minimum NGS coverage is ≥20x for all exons and +/-10bp of flanking DNA.

nically signed and reported on April 02, 2021 by:
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Molecular Geneticist

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 CAP 7185561
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 AU ID 1407125
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 IS920212 #3950.01



SUPPLEMENTAL INFORMATION V.19.04 SEQUENCING WITH CNV DETECTION

Limitations and Other Test Notes

Interpretation of the test results is limited by the information that is currently available. Better interpretation should be possible in the future as our knowledge about human genetics and the patient's condition improve.

When Next Gen or Sanger sequencing does not reveal any difference from the reference sequence, or when a sequence variant is homozygous, we cannot be certain that we were able to detect both patient alleles. Occasionally, a patient may carry an allele which does not capture or amplify due for example to a large deletion or insertion.

Copy number variants (CNVs) of four exons or more in size are detected with sensitivity approaching 100% through analysis of Next Gen sequence data. However, sensitivity for detection of CNVs smaller than four exons is lower (we estimate ~75%).

Coverage includes all coding exons of the gene(s) analyzed plus 10 bases of flanking noncoding DNA in all available transcripts along with other non-coding regions in which pathogenic variants have been identified at PreventionGenetics or reported elsewhere.

In most cases, we are unable to determine the phase of sequence variants. In particular, when we find two likely causative variants for recessive disorders, we cannot be certain that the variants are on different chromosomes.

Our ability to detect minor sequence variants due to somatic mosaicism is limited. Sequence variants that are present in less than 50% of the patient's nucleated cells may not be detected.

Unless present within coding regions, runs of mononucleotide repeats (eg (A)_n or (T)_n) with n > 8 in the reference sequence) are generally not analyzed because of strand slippage during amplification.

Unless otherwise indicated, DNA sequence data is obtained from a specific cell type (often leukocytes from whole blood). Test reports contain no information about the DNA sequence in other cell types.

We cannot be certain that the reference sequences are correct. Genome build hg19, GRCh37 (Feb2009) is currently used as our reference in nearly all cases.

We have confidence in our ability to track a specimen once it has been received by PreventionGenetics. However, we take no responsibility for any specimen labeling errors that occur before the sample arrives at PreventionGenetics.

Genetic counseling to help to explain test results to the patients and to discuss reproductive options is recommended.

Reported results will typically not contain any additional information regarding pharmacogenetic analysis of genes, nor are these tests designed to help guide dosage requirements. Pharmacogenetic variant analysis is available, for a select list of genes, as an opt-in with PGxome® tests.

Test Methods

We use Next Generation Sequencing (NGS) technologies to cover the coding regions of the targeted genes plus 10 bases of non-coding DNA flanking each exon. As required, genomic DNA is extracted from the specimen. The DNA corresponding to these regions is captured using Agilent Clinical Research Exome hybridization



NAME 6040, Donor

PREVENTION > GENETICS

using 150 by 150 bp paired end reads (Illumina, San Diego, CA, USA).

probes. Captured DNA is sequenced using Illumina's Reversible Dye Terminator (RDT) platform NovaSeq 6000

The following quality control metrics are generally achieved: >98% of target bases are covered at >20x, and mean coverage of target bases >120x. Data analysis is performed using the internally developed software Titanium-Exome. Specified genes for which the enhance option is selected are backfilled with Sanger sequencing to achieve 100% coverage.

For Sanger sequencing, Polymerase Chain Reaction (PCR) is used to amplify the necessary exons plus additional flanking non-coding sequence. After purification of the PCR products, cycle sequencing is carried out using the ABI Big Dye Terminator v.3.1 kit. PCR products are resolved by electrophoresis on an ABI 3730xl capillary sequencer. In most cases, cycle sequencing is performed separately in both the forward and reverse directions; in some cases, sequencing is performed twice in either the forward or reverse directions.

Copy number variants (CNVs) are also detected from NGS data. We utilize a CNV calling algorithm that compares mean read depth and distribution for each target in the test sample against multiple matched controls. Neighboring target read depth and distribution and zygosity of any variants within each target region are used to reinforce CNV calls. All reported CNVs are confirmed using another technology such as aCGH, MLPA, or PCR. On occasion, it will not be technically possible to confirm a smaller CNV called by NGS. In these instances, the CNV will not be included on the report.

All differences from the reference sequences (sequence variants) are assigned to one of five interpretation categories (Pathogenic, Likely Pathogenic, Variant of Uncertain Significance, Likely Benign and Benign) per ACMG Guidelines (Richards et al. 2015). Rare and undocumented synonymous variants are nearly always classified as likely benign if there is no indication that they alter protein sequence or disrupt splicing. Benign variants are not listed in the reports, but are available upon request.

Human Genome Variation Society (HGVS) recommendations are used to describe sequence variants (http://www.hgvs.org).

FDA Notes

These results should be used in the context of available clinical findings, and should not be used as the sole basis for treatment. This test was developed and its performance characteristics determined by PreventionGenetics. US Food and Drug Administration (FDA) does not require this test to go through premarket FDA review. This test is used for clinical purposes. It should not be regarded as investigational or for research. This laboratory is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA) as qualified to perform high complexity clinical laboratory testing.



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Patient Information:
6040, Donor
DOB:
Sex: M
MR#: 6040
Patient#:

Partner Information: Not Tested

Specimen Type: DNA Collected: May 17,2019

Accession:

Test#: F

Accession: N/A

FINAL RESULTS



No carrier mutations identified

Physician: Seitz, Suzanne ATTN: Seitz, Suzanne Fairfax Cryobank 3015 Williams Drive Fairfax, VA 22031

Laboratory: **Fulgent Therapeutics LLC** CAP#: 8042697 CLIA#: 05D2043189 Laboratory Director: Dr. Hanlin (Harry) Gao Report Date: Apr 10,2024

TEST PERFORMED

Single Gene Carrier Screening: NAGA

(1 Gene Panel: NAGA; gene sequencing with deletion and duplication analysis)

INTERPRETATION:

Notes and Recommendations:

- No carrier mutations were identified in the submitted specimen. A negative result does not rule out the possibility of a genetic predisposition nor does it rule out any pathogenic mutations in areas not assessed by this test or in regions that were covered at a level too low to reliably assess. Also, it does not rule out mutations that are of the sort not queried by this test; see Methods and Limitations for more information. A negative result reduces, but does not eliminate, the chance to be a carrier for any condition included in this screen. Please see the supplemental table for details.
- This carrier screening test does not screen for all possible genetic conditions, nor for all possible mutations in every gene tested. This report does not include variants of uncertain significance; only variants classified as pathogenic or likely pathogenic at the time of testing, and considered relevant for reproductive carrier screening, are reported. Please see the gene specific notes for details. Please note that the classification of variants can change over time.
- Patients may wish to discuss any carrier results with blood relatives, as there is an increased chance that they are also carriers. . These results should be interpreted in the context of this individual's clinical findings, biochemical profile, and family history.
- X-linked genes are not routinely analyzed for male carrier screening tests. Gene specific notes and limitations may be present. See below.
- Genetic counseling is recommended. Available genetic counselors and additional resources can be found at the National Society of Genetic Counselors (NSGC; https://www.nsgc.org)

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GENES TESTED:

Custom Beacon Carrier Screening Panel - Gene

This analysis was run using the Custom Beacon Carrier Screening Panel gene list. 1 genes were tested with 100.0% of targets sequenced at >20x coverage. For more gene-specific information and assistance with residual risk calculation, see the SUPPLEMENTAL TABLE.

NAGA

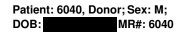
METHODS:

Genomic DNA was isolated from the submitted specimen indicated above (if cellular material was submitted). DNA was barcoded, and enriched for the coding exons of targeted genes using hybrid capture technology. Prepared DNA libraries were then sequenced using a Next Generation Sequencing technology. Following alignment to the human genome reference sequence (assembly GRCh37), variants were detected in regions of at least 10x coverage. For this specimen, 100.00% and 100.00% of coding regions and splicing junctions of genes listed had been sequenced with coverage of at least 10x and 20x, respectively, by NGS or by Sanger sequencing. The remaining regions did not have 10x coverage, and were not evaluated. Variants were interpreted manually using locus specific databases, literature searches, and other molecular biological principles. To minimize false positive results, any variants that do not meet internal guality standards are confirmed by Sanger sequencing. Variants classified as pathogenic, likely pathogenic, or risk allele which are located in the coding regions and nearby intronic regions (+/- 20bp) of the genes listed above are reported. Variants outside these intervals may be reported but are typically not guaranteed. When a single pathogenic or likely pathogenic variant is identified in a clinically relevant gene with autosomal recessive inheritance, the laboratory will attempt to ensure 100% coverage of coding sequences either through NGS or Sanger sequencing technologies ("fill-in"). All genes listed were evaluated for large deletions and/or duplications. However, single exon deletions or duplications will not be detected in this assay, nor will copy number alterations in regions of genes with significant pseudogenes. Putative deletions or duplications are analyzed using Fulgent Germline proprietary pipeline for this specimen. Bioinformatics: The Fulgent Germline v2019.2 pipeline was used to analyze this specimen.

LIMITATIONS:

General Limitations

These test results and variant interpretation are based on the proper identification of the submitted specimen, accuracy of any stated familial relationships, and use of the correct human reference sequences at the queried loci. In very rare instances, errors may result due to mix-up or co-mingling of specimens. Positive results do not imply that there are no other contributors, genetic or otherwise, to future pregnancies, and negative results do not rule out the genetic risk to a pregnancy. Official gene names change over time. Fulgent uses the most up to date gene names based on HUGO Gene Nomenclature Committee (https://www.genenames.org) recommendations. If the gene name on report does not match that of ordered gene, please contact the laboratory and details can be provided. Result interpretation is based on the available clinical and family history information for this individual, collected published information, and Alamut annotation available at the time of reporting. This assay is not designed or validated for the detection of low-level mosaicism or somatic mutations. This assay will not detect certain types of genomic aberrations such as translocations, inversions, or repeat expansions other than specified genes. DNA alterations in regulatory regions or deep intronic regions (greater than 20bp from an exon) may not be detected by this test. Unless otherwise indicated, no additional assays have been performed to evaluate genetic changes in this specimen. There are technical limitations on the ability of DNA sequencing to detect as reliably as single nucleotide variants. Rarely, due to systematic chemical, computational, or human error, DNA variants may be missed.



Accession#: For the second sec





of pseudogene sequences or other highly-homologous sequences, sometimes these may still interfere with the technical ability of the assay to identify pathogenic alterations in both sequencing and deletion/duplication analyses. Deletion/duplication analysis can identify alterations of genomic regions which include one whole gene (buccal swab specimens and whole blood specimens) and are two or more contiguous exons in size (whole blood specimens only); single exon deletions or duplications may occasionally be identified, but are not routinely detected by this test. When novel DNA duplications are identified, it is not possible to discern the genomic location or orientation of the duplicated segment, hence the effect of the duplication cannot be predicted. Where deletions are detected, it is not always possible to determine whether the predicted product will remain in-frame or not. Unless otherwise indicated, deletion/duplication analysis has not been performed in regions that have been sequenced by Sanger.

Gene Specific Notes and Limitations

No gene specific limitations apply to the genes on the tested panel.

SIGNATURE:

Z Gao

Dr. Harry Gao, DABMG, FACMG on 4/10/2024 Laboratory Director, Fulgent

DISCLAIMER:

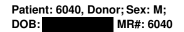
This test was developed and its performance characteristics determined by **Fulgent Therapeutics LLC**. It has not been cleared or approved by the FDA. The laboratory is regulated under CLIA as qualified to perform high-complexity testing. This test is used for clinical purposes. It should not be regarded as investigational or for research. Since genetic variation, as well as systematic and technical factors, can affect the accuracy of testing, the results of testing should always be interpreted in the context of clinical and familial data. For assistance with interpretation of these results, healthcare professionals may contact us directly at (626) 350-0537 or info@fulgentgenetics.com. It is recommended that patients receive appropriate genetic counseling to explain the implications of the test result, including its residual risks, uncertainties and reproductive or medical options.

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To view the supplemental table describing the carrier frequencies, detection rates, and residual risks associated with the genes on this test please visit the following link: <u>Beacon Expanded Carrier Screening Supplemental Table</u>







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Patient	Sample	Referring Doctor
Patient Name: Donor 6040 Date of Birth: Reference #: Indication: Encounter of male for testing for genetic disease carrier status for procreative management Test Type: Chromosome Analysis - Peripheral Blood	Specimen Type: Peripheral Blood Lab #: Date Collected: 5/15/2019 Date Received: 5/16/2019 Final Report: 6/4/2019	Fairfax Cryobank, Inc.

CYTOGENETIC ANALYSIS

Results

Staining:G-bands by trypsin using Giemsa (GTG)Band level:550

Chromosome count: **46** Cells analyzed: **20** Cells captured: 5

Cells karyotyped: 3

Cultures examined: 2

Karyotype: 46,XY

Interpretation

Cytogenetic analysis revealed the presence of a **normal male** karyotype in peripheral blood lymphocytes. This analysis does not show any evidence of a clinically significant numerical or structural chromosome abnormality.

The standard procedures used in this analysis do not routinely detect microdeletions, small rearrangements or low level mosaicism.

This case has been reviewed and electronically signed by Arvind Babu, Ph.D., FACMG, Laboratory Director Laboratory Medical Consultant: Bryn Webb, M.D.





Patient Information	Specimen Information	Client Information
ID 6040, DONOR DOB: AGE: Gender: M Phone: NG Patient ID: M	Specimen: Requisition: Lab Ref #:Image: Collected: Collected: 05/15/2019 Received: 05/17/2019 / 02:51 EDT Reported: 05/18/2019 / 11:43 EDT	Client #: 41578 1000 UNKNOWN SEMA4 1428 MADISON AVE 2ND FL ATRAN BLDG RM 25 NEW YORK, NY 10029
Test Name HEMOGLOBINOPATHY EVAL INTERPRETATION	In Range Out Of Range	Reference RangeLabTBR
Normal P	Pattern	
HEMOGLOBIN A HEMOGLOBIN F HEMOGLOBIN A2 HEMOGLOBIN S HEMOGLOBIN C HEMOGLOBIN VARIANT HEMOGLOBINOPATHY INDICES RBC HEMOGLOBIN HEMATOCRIT MCV MCH RDW	96.4 <1.0 2.6 0.0 0.0 0.0 5.26 16.8 49.7 94.5 31.9 12.5	<pre>>96.0 Percent <2.0 Percent 1.8-3.5 Percent 0.0-0.0 Percent 0.0-0.0 Percent TBR 4.20-5.80 Million/uL 13.2-17.1 g/dL 38.5-50.0 % 80.0-100.0 fL 27.0-33.0 pg 11.0-15.0 %</pre>

PERFORMING SITE:

TBR Quest Diagnostics, One Malcolm Avenue, Teterboro, NJ 07608 Laboratory Director: Lawrence Tsao, M.D., CLIA: 31D0696246

