

Test Information Sheet

GeneDx

Custom Cardiology Panel

Panel Gene List

ABCC6, ABCC9, ACADVL, ACTA1, ACTA2, ACTC1, ACTN2, ACVR1, ACVRL1, ADAMTS10, ADAMTS17, ADAMTS2, ADAMTSL4, AEBP1, AGL, AKAP9, ALDH18A1, ALMS1, ALPK3, ANK2, ANKRD1, APOB, ASPH, ATP6V0A2, ATP6V0D2, ATP6V1E1, ATP7A, B3GALT6, B3GAT3, B4GALT7, BAG3, BGN, BMPR1B, BMPR2, BRAF, CACNA1C, CACNA2D1, CACNB2, CALM1, CALM2, CALM3, CALR, CALR3, CASQ2, CAV1, CAV3, CBLN2, CBS, CHRM2, CHST14, COL11A1, COL11A2, COL12A1, COL1A1, COL1A2, COL2A1, COL3A1, COL4A1, COL5A1, COL5A2, COL9A1, COL9A2, COL9A3, COX15, CPT1A, CRYAB, CSRP3, CTF1, CTNNA3, DES, DMD, DNAJC19, DOLK, DSC2, DSE, DSG2, DSP, DTNA, EFEMP2, EIF2AK4, ELAC2, ELN, EMD, ENG, EYA4, FBLN5, FBN1, FBN2, FGF12, FHL1, FHL2, FHOD3, FKBP14, FKRP, FKTN, FLNA, FLNC, FOXC2, FOXE3, FOXF1, FOXRED1, GAA, GATA4, GATA5, GATA6, GATAD1, GDF2, GJA5, GLA, GLB1, GNB5, GPD1L, HCN4, HFE, HRAS, ILK, JAG1, JPH2, JUP, KCNA5, KCNB2, KCND3, KCNE1, KCNE1L (KCNE5), KCNE2, KCNE3, KCNH2 (HERG), KCNJ16, KCNJ2, KCNJ5, KCNJ8, KCNK3, KCNQ1, KCNT1, KLF10, KRAS, LAMA4, LAMP2, LDB3, LDLR, LDLRAP1, LMNA, LOX, LRRC10, LTBP2, LTBP4, LZTR1, LZTS1, MAP2K1, MAP2K2, MAP3K8, MAT2A, MED12, MFAP5, MIB1, MRPL3, MRPS22, MURC, MTO1, MYBPC3, MYH11, MYH6, MYH7, MYL2, MYL3, MYL4, MYLK, MYLK2, MYO6, MYOM1, MYOZ2, MYPN, NEBL, NEXN, NKX2-5, NKX2-6, NOS1AP, NOTCH1, NPPA, NRAS, PCSK9, PDLIM3, PI4KA, PKP2, PLEC, PLEKHM2, PLN, PLOD1, PLOD3, PPA2, PRDM16, PRDM5, PRKAG2, PRKG1, PTPN11, PYCR1, RAF1, RANGRF, RASA1, RASA2, RBM20, RIN2, RIT1, RRAS, RYR2, SCARF2, SCN10A, SCN1B, SCN2B, SCN3B, SCN4A, SCN4B, SCN5A, SCNN1A, SCO2, SGCD, SHOC2, SKI, SLC25A20, SLC25A3, SLC25A4, SLC2A10, SLC2A5, SLC39A13, SLMAP, SMAD1, SMAD2, SMAD3, SMAD4, SMAD6, SMAD9, SMS, SNTA1, SOS1, SOS2, SPRY1, SYNE1, SYNE2, TAB2, TANGO2, TAZ, TBX1, TBX20, TBX5, TCAP, TECRL, TGFB2, TGFB3, TGFB1, TGFB2, TMEM43, TMEM70, TMPO, TNNC1, TNNI3, TNNI3K, TNNT2, TNXB, TOR1AIP1, TPM1, TRDN, TRIM63, TRPM4, TSFM, TTN, TTR, TXNRD2, UPF3B, VCL, XK, ZNF469

Clinical Features

This panel can assess for cardiac arrhythmias, cardiomyopathies, sudden unexplained death syndrome, Danon disease, syndromic cardiac disorders, Fabry disease, mitochondrial myopathy, or muscular dystrophy, heritable disorders of connective tissue, PAH, HHT, and familial hypercholesterolemia.

Inheritance Pattern/Genetics

Autosomal Dominant, Autosomal Recessive, X-linked, or Mitochondrial

Test Methods

Using genomic DNA from the submitted specimen, the complete coding regions and splice site junctions of the genes on this panel are enriched using a proprietary targeted capture system developed by GeneDx for next-generation sequencing with CNV calling (NGS-CNV) (excluding exon 6 of the PKP2 gene and the following genomic regions of the TTN gene: chr2:179527692- 179527782, 179523898-179523982, 179523731-179523815). The enriched targets are simultaneously sequenced with paired-end reads on an Illumina platform. Bi-directional sequence reads are assembled and aligned to reference sequences based on NCBI RefSeq transcripts and human genome build GRCh37/UCSC hg19. After gene specific filtering, data are analyzed to identify sequence variants and most deletions and duplications involving coding exons. Alternative sequencing or copy number detection methods are used to analyze regions with inadequate sequence or copy number data. Reportable variants include pathogenic variants, likely pathogenic variants and variants of uncertain significance. Likely benign and benign variants, if present, are not routinely reported but are available upon request.

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Test Sensitivity

The technical sensitivity of sequencing is estimated to be >99% at detecting single nucleotide events. It will not reliably detect deletions greater than 20 base pairs, insertions or rearrangements greater than 10 base pairs, or low-level mosaicism. The copy number assessment methods used with this test cannot reliably detect copy number variants of less than 500 base pairs or mosaicism and cannot identify balanced chromosome aberrations. Assessment of exon-level copy number events is dependent on the inherent sequence properties of the targeted regions, including shared homology and exon size. For B3GALT6, CTF1, FKRP, FOXE3, HRAS, SCO2, TBX1 genes, sequencing but not deletion/duplication analysis, is performed. Gene specific exclusions for exon- level deletion/duplication testing for this panel are: APOA1, CALM1, GATA5, KCNT1, LCAT, LMF1, SCN1B, TAZ and TBX20 genes only whole gene deletions or duplications may be detected. Recombination of TNXB with its pseudogene (gene conversion or TNXB/XA fusion), is not evaluated.

Gene	Protein	Inheritance	Disease Association(s)
ABCC6	ATP-BINDING CASSETTE, SUBFAMILY C, MEMBER 6	AR	Pseudoxanthoma elasticum
ABCC9	ATP-BINDING CASSETTE, SUBFAMILY C, MEMBER 9	AD	DCM, BrS, Cantu syndrome and related disorders
ACADVL	ACYL-CoA DEHYDROGENASE, VERY LONG-CHAIN	AR	neonatal HCM/VLCAD deficiency
ACTA1	ACTIN, ALPHA, SKELETAL MUSCLE 1	AD/AR	Cardiomyopathy, myopathy
ACTA2	ACTIN, ALPHA-2, SMOOTH MUSCLE, AORTA	AD	fTAAD
ACTC1	ACTIN, ALPHA, CARDIAC MUSCLE	AD	CHD, DCM, HCM, LVNC
ACTN2	ACTININ, ALPHA-2	AD	DCM, HCM
ACVR1	ACTIVIN A RECEPTOR, TYPE II-LIKE KINASE 2	AD	Fibrodysplasia ossificans progressiva (FOP)
ACVRL1	ACTIVIN A RECEPTOR TYPE II-LIKE 1	AD	HHT, PAH
ADAMTS10	A DISINTEGRIN-LIKE AND METALLOPROTEINASE WITH THROMBOSPONDIN TYPE 1 MOTIF, 10	AR	Weill-Marchesani syndrome 1
ADAMTS2	ADAM METALLOPEPTIDASE WITH THROMBOSPONDIN TYPE 1 MOTIF 2	AR	dEDS
ADAMTS4	ADAMTS-LIKE 4	AR	Ectopia lentis
AEBP1	AE-BINDING PROTEIN 1	AR	EDS, unclassified
AGL	AMYLO-1,6-GLUCOSIDASE, 4-ALPHA-GLUCANOTRANSFERASE	AR	GSD, type IIIa GSD, Type IIIb
AKAP9	A-KINASE ANCHOR PROTEIN 9	AD	LQTS

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Gene	Protein	Inheritance	Disease Association(s)
<i>ALDH18A1</i>	ALDEHYDE DEHYDROGENASE 18 FAMILY MEMBER A1	AD	Cutis laxa
<i>ALMS1</i>	CENTROSOME AND BASAL BODY ASSOCIATED PROTEIN	AR	Alstrom syndrome, infantile DCM
<i>ALPK3</i>	ALPHA-KINASE 3	AR	Pediatric Cardiomyopathy
<i>ANK2</i>	ANKYRIN 2	AD	Arrhythmia, LQTS
<i>ANKRD1</i>	ANKYRIN REPEAT DOMAIN-CONTAINING PROTEIN 1	AD	HCM, DCM
<i>APOB</i>	APOLIPOPROTEIN	AD	HeFH/HoFH
<i>ASPH</i>	ASPARTATE BETA-HYDROXYLASE	AR	Ectopia lentis, spontaneous filtering blebs, and craniofacial dysmorphism
<i>ATP6V0A2</i>	ATPASE H ⁺ TRANSPORTING V0 SUBUNIT A2	AR	Cutis laxa
<i>ATP6V0D2</i>	ATPase, H ⁺ TRANSPORTING, LYSOSOMAL, 38-KD, V0 SUBUNIT D, ISOFORM 2	AR	Cutis laxa
<i>ATP6V1E1</i>	ATPASE H ⁺ TRANSPORTING V1 SUBUNIT E	AR	Cutis laxa
<i>ATP7A</i>	ATPASE COPPER TRANSPORTING ALPHA	XL	Menkes, OHS
<i>B3GALT6</i>	BETA-1,3-GALACTOSYLTRANSFERASE 6	AR	spEDS
<i>B3GAT3</i>	BETA-1,3-GLUCURONYLTRANSFERASE 3	AR	Joint dislocations, short stature, dysmorphisms, CHD
<i>B4GALT7</i>	BETA-1,4-GALACTOSYLTRANSFERASE 7	AR	spEDS
<i>BAG3</i>	BCL2-ASSOCIATED ATHANOGENE 3	AD	DCM, myofibrillar myopathy
<i>BGN</i>	BIGLYCAN	XL	Meester-Loeys syndrome Spondyloepimetaphyseal dysplasia
<i>BMPR1B</i>	BONE MORPHOGENETIC PROTEIN RECEPTOR, TYPE IB	AD, AR	PAH
<i>BMPR2</i>	BONE MORPHOGENETIC PROTEIN RECEPTOR, TYPE II	AD	PAH
<i>BRAF</i>	V-RAF MURINE SARCOMA VIRAL ONCOGENE HOMOLOG B1	AD	Noonan/CFC/Costello
<i>CACNA1C</i>	CALCIUM CHANNEL, VOLTAGE-DEPENDENT, L TYPE, ALPHA-1C SUBUNIT	AD	BrS, Timothy syndrome, LQTS

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CACNA2D1	CALCIUM CHANNEL, VOLTSGE-DEPENDENT ALPHA-2/DELTA SUBUNIT 1	AD	BrS
CACNB2	CALCIUM CHANNEL, VOLTAGE-DEPENDENT, BETA-2 SUBUNIT	AD	BrS
CALM1	CALMODULIN 1	AD	LQTS,CPVT
CALM2	CALMODULIN 2	AD	LQTS, CPVT
CALM3	CALMODULIN 3	AD	LQTS, CPVT
CALR	CALRETICULIN	AD	Arrhythmia
CALR3	CALRETICULIN 3	AD	HCM
CASQ2	CALSEQUESTRIN 2	AR	CPVT
CAV1	CAVEOLIN 1	AD	PAH, lipodystrophy
CAV3	CAVEOLIN 3	AD, AR	HCM, LQTS, LGMD, Tateyama-type distal myopathy, SIDS, rippling muscle disease
CBS	CYSTATHIONINE BETA-SYNTHASE	AR	Homocystinuria
CHRM2	M2-MUSCARINIC ACETYLCHOLINE RECEPTOR	AD	DCM
CHST14	CARBOHYDRATE (DERMATAN 4) SULFOTRANSFERASE 14	AR	mcEDS
COL11A1	COLLAGEN TYPE XI ALPHA 1	AD	Fibrochondrogenesis Stickler syndrome
COL11A2	COLLAGEN TYPE XI ALPHA 2	AD	Fibrochondrogenesis Stickler syndrome, non-ocular
COL12A1	COLLAGEN TYPE XIIALPHA 1	AD	mEDS
COL1A1	COLLAGEN TYPE I ALPHA 1	AD	aEDS cEDS Osteogenesis Imperfecta
COL1A2	COLLAGEN TYPE I ALPHA 2	AD, AR	aEDS Osteogenesis Imperfecta cvEDS
COL2A1	COLLAGEN TYPE II ALPHA 1	AD, AR	OSMED; Stickler syndrome
COL3A1	COLLAGEN TYPE III ALPHA 1	AD	vEDS
COL4A1	COLLAGEN TYPE IV ALPHA 1	AD	fTAAD
COL5A1	COLLAGEN TYPE V ALPHA 1	AD	cEDS
COL5A2	COLLAGEN TYPE V ALPHA 2	AD	cEDS
COL9A1	COLLAGEN TYPE IX ALPHA 1	AD, AR	Stickler syndrome
COL9A2	COLLAGEN TYPE IX ALPHA 2	AD, AR	Stickler syndrome
COL9A3	COLLAGEN TYPE IX ALPHA-3	AD, AR	multiple epiphyseal dysplasia (MED)/Stickler syndrome
COX15	CYTOCHROME c OXIDASE ASSEMBLY FACTOR COX15	AR	HCM/COX deficiency
CPT1A	CARNITINE PALMITOYLTRANSFERASE I, LIVER	AR	carnitine palmitoyltransferase 1A (CPT1A) deficiency
CRYAB	CRYSTALLIN, ALPHA-B	AD, AR	DCM, myofibrillar myopathy

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Gene	Protein	Inheritance	Disease Association(s)
<i>CSRP3</i>	CYSTEINE- AND GLYCINE-RICH PROTEIN 3	AD	HCM, DCM
<i>CTF1</i>	CARDIOTROPHIN 1 I	AD	DCM
<i>CTNNA3</i>	CATENIN, ALPHA-3	AD	ARVC
<i>DES</i>	DESMIN	AD	DCM, ARVC, myopathy, AV block, LGMD
<i>DMD</i>	DYSTROPHIN	XL	DMD, BMD, DCM
<i>DNAJC19</i>	DNAJ/HSP40 HOMOLOG, SUBFAMILY C, MEMBER 19	AR	DCM with ataxia
<i>DOLK</i>	DOLICHOL KINASE	AR	DCM, congenital disorder of glycosylation type Im
<i>DSC2</i>	DESMOCOLLIN	AD, AR	ARVC, ARVC+skin and hair findings , DCM
<i>DSE</i>	DERMATAN SULFATE EPIMERASE	AR	mcEDS
<i>DSG2</i>	DESMOGLEIN	AD	ARVC, DCM
<i>DSP</i>	DESMOPLAKIN	AD, AR	ARVC, DCM, Carvajal syndrome
<i>DTNA</i>	DYSTROBREVIN, ALPHA	AD	LVNC, CHD
<i>EFEMP2</i>	EGF CONTAINING FIBULIN-LIKE EXTRACELLULAR MATRIX PROTEIN 2	AR	Cutis laxa
<i>EIF2AK4</i>	EUKARYOTIC TRANSLATION INITIATION FACTOR 2-ALPHA KINASE 4	AR	PVOD2, PCH, PAH
<i>ELAC2</i>	ELAC, E. COLI, HOMOLOG OF, 2	AR	infantile HCM
<i>ELN</i>	ELASTIN	AD	Cutis laxa
<i>EMD</i>	EMERIN	XL	EMD
<i>ENG</i>	ENDOGLIN	AD	HHT +/- PAH
<i>EYA4</i>	EYES ABSENT 4	AD	DCM
<i>FBLN5</i>	FIBULIN 5	AD, AR	Cutis laxa
<i>FBN1</i>	FIBRILLIN 1	AD	Marfan syndrome
<i>FBN2</i>	FIBRILLIN 2	AD	Congenital contractual arachnodactyly
<i>FGF12</i>	FIBROBLAST GROWTH FACTOR 12	AD	BrS, VT
<i>FHL1</i>	FOUR-AND-A-HALF LIM DOMAINS 1	XL	HCM, LVH, EMD, skeletal muscle, muscle hypertrophy, Myofibrillar myopathy
<i>FHL2</i>	FOUR-AND-A-HALF LIM DOMAINS 2	AD	HCM
<i>FHOD3</i>	FORMIN HOMOLOGY-2 DOMAIN-CONTAINING PROTEIN 3	AD	DCM
<i>FKBP14</i>	FK506 BINDING PROTEIN 14	AR	kEDS, myopathy, and hearing loss
<i>FKRP</i>	FUKUTIN RELATED PROTEIN	AR	DCM, muscular dystrophy
<i>FKTN</i>	FUKUTIN	AR	DCM, LGMD, Fukuyama Congenital Muscular Dystrophy

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Gene	Protein	Inheritance	Disease Association(s)
<i>FLNA</i>	FILAMIN A	XL	EDS with periventricular heterotopia
<i>FLNC</i>	FILAMIN C	AD	RCM, HCM, ARVC
<i>FOXE3</i>	FORKHEAD BOX E3	AD	ftAAD
<i>FOXF1</i>	FORKHEAD, DROSOPHILA, HOMOLOG-LIKE 5	AD	PAH
<i>FOXRED1</i>	FAD-DEPENDENT OXIDOREDUCTASE DOMAIN-CONTAINING PROTEIN 1	AR	Cardiomyopathy, myopathy
<i>GAA</i>	GLUCOSIDASE, ALPHA, ACID	AR	Cardiomyopathy, GSD II
<i>GATA4</i>	GATA-BINDING PROTEIN 4	AD	AF, CHD, cardiomyopathy, SUDS
<i>GATA5</i>	GATA-BINDING PROTEIN 5	AD	AF, CHD, cardiomyopathy
<i>GATA6</i>	GATA-BINDING PROTEIN 6	AD	AF, CHD, cardiomyopathy
<i>GATAD1</i>	GATA ZINC FINGER DOMAIN-CONTAINING PROTEIN 1	AR	DCM
<i>GDF2</i>	GROWTH/DIFFERENTIATION FACTOR 2	AD	HHT +/- PAH
<i>GJA5</i>	GAP JUNCTION PROTEIN, ALPHA-5	AD	AF, HB, SADS, SIDS, CHD
<i>GLA</i>	GALACTOSIDASE, ALPHA	XL	Fabry disease
<i>GLB1</i>	GALACTOSIDASE, BETA-1	AR	HCM, DCM
<i>GNB5</i>	GUANINE NUCLEOTIDE-BINDING PROTEIN, BETA-5	AR	Intellectual developmental disorder with cardiac, arrhythmia
<i>GDPIL</i>	GLYCEROL-3-PHOSPHATE DEHYDROGENASE 1-LIKE	AD	BrS
<i>HCN4</i>	HYPERPOLARIZATION-ACTIVATED CYCLIC NUCLEOTIDE-GATED POTASSIUM CHANNEL 4	AD	BrS, SSS
<i>HFE</i>	HUMAN HEMOCHROMATOSIS PROTEIN	AR	Hereditary Hemochromatosis
<i>HRAS</i>	V-HA-RAS HARVEY RAT SARCOMA VIRAL ONCOGENE HOMOLOG	AD	Costello syndrome
<i>ILK</i>	INTEGRIN-LINKED KINASE	AD	DCM
<i>JAG1</i>	JAGGED 1	AD	Allagile syndrome
<i>JPH2</i>	JUNCTOPHILIN 2	AD	HCM
<i>JUP</i>	JUNCTION PLAKOGLOBIN	AD, AR	ARVC, Naxos Disease
<i>KCNA5</i>	POTASSIUM CHANNEL, VOLTAGE-GATED, SHAKER-RELATED SUBFAMILY, MEMBER 5	AD	Arrhythmia, AF, PAH

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Gene	Protein	Inheritance	Disease Association(s)
KCNB2	POTASSIUM CHANNEL, VOLTAGE-GATED, SHAB-RELATED SUBFAMILY, MEMBER 2	AD	BrS
KCND3	POTASSIUM CHANNEL, VOLTAGE-GATED, SHAL-RELATED SUBFAMILY, MEMBER 3	AD	BrS, SIDS, Spinocerebellar ataxia
KCNE1	POTASSIUM CHANNEL, VOLTAGE-GATED, ISK-RELATED SUBFAMILY, MEMBER 1	AD, AR	LQTS, JLNS
KCNE2	POTASSIUM CHANNEL, VOLTAGE-GATED, ISK-RELATED SUBFAMILY, MEMBER 2	AD	LQTS
KCNE3	POTASSIUM CHANNEL, VOLTAGE-GATED, ISK-RELATED SUBFAMILY, MEMBER 3	AD	BrS
KCNE1L (KCNE5)	POTASSIUM CHANNEL, VOLTAGE-GATED, ISK-RELATED FAMILY, MEMBER 1-LIKE	XL	BrS, AF, VF
KCNE2	POTASSIUM CHANNEL, VOLTAGE-GATED, ISK-RELATED SUBFAMILY, MEMBER 2	AD	LQTS
KCNE3	POTASSIUM CHANNEL, VOLTAGE-GATED, ISK-RELATED SUBFAMILY, MEMBER 3	AD	BrS
KCNH2 (HERG)	POTASSIUM CHANNEL, VOLTAGE-GATED, SUBFAMILY H, MEMBER 2	AD	LQTS, SQTS
KCNJ16	POTASSIUM CHANNEL, INWARDLY RECTIFYING, SUBFAMILY J, MEMBER 16	AD	BrS
KCNJ2	POTASSIUM CHANNEL, INWARDLY RECTIFYING, SUBFAMILY J, MEMBER 2	AD	Andersen-Tawil syndrome, SQTS
KCNJ5	POTASSIUM CHANNEL, INWARDLY RECTIFYING, SUBFAMILY J, MEMBER 5	AD	LQTS
KCNJ8	POTASSIUM CHANNEL, INWARDLY RECTIFYING, SUBFAMILY J, MEMBER 8	AD	ERS, SIDS
KCNK3	POTASSIUM CHANNEL, SUBFAMILY K, MEMBER 3	AD	PAH, AF
KCNQ1	POTASSIUM CHANNEL, VOLTAGE-GATED, KQT-LIKE SUBFAMILY, MEMBER	AD, AR	JLNS, LQTS, SQTS

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Gene	Protein	Inheritance	Disease Association(s)
	1		
<i>KCNT1</i>	POTASSIUM CHANNEL, SUBFAMILY T, MEMBER 1	AD	BrS
<i>KLF10</i>	KRUPPEL-LIKE FACTOR 10	AD	HCM
<i>KRAS</i>	V-KI-RAS2 KIRSTEN RAT SARCOMA VIRAL ONCOGENE HOMOLOG	AD	Noonan/CFC/Costello
<i>LAMA4</i>	LAMININ, ALPHA-4	AD	DCM
<i>LAMP2</i>	LYSOSOME-ASSOCIATED MEMBRANE PROTEIN 2	XL	Danon disease
<i>LDB3</i>	LIM DOMAIN-BINDING 3	AD	DCM, LVNC, myopathy
<i>LDLR</i>	LOW-DENSITY LIPOPROTEIN RECEPTOR	AD	HeFH/HoFH
<i>LDLRAP1</i>	LOW-DENSITY LIPOPROTEIN RECEPTOR ADAPTOR PROTEIN 1	AR	ARFH
<i>LMNA</i>	LAMIN A/C	AD, AR	DCM, congenital muscular dystrophy, EMD, ARVC
<i>LOX</i>	LYSYL OXIDASE	AD	FTAAD
<i>LRRC10</i>	LEUCINE-RICH REPEAT-CONTAINING PROTEIN 10	AD, AR	DCM, HCM, congenital muscular dystrophy, EMD
<i>LTBP2</i>	LATENT TRANSFORMING GROWTH FACTOR-BETA-BINDING PROTEIN 2	AD, AR	Ectopia lentis, Weill-Marchesani syndrome, Marfan syndrome
<i>LTBP4</i>	LATENT TRANSFORMING GROWTH FACTOR BETA BINDING PROTEIN 4	AR	Cutis laxa, autosomal recessive
<i>LZTR1</i>	LEUCINE ZIPPER-LIKE TRANSCRIPTIONAL REGULATOR 1	AD, AR	Noonan syndrome
<i>LZTS1</i>	LEUCINE ZIPPER, PUTATIVE TUMOR SUPPRESSOR 1	AD	EDS, hypermobile
<i>MAP2K1</i>	MITOGEN-ACTIVATED PROTEIN KINASE KINASE 1	AD	Noonan/CFC/Costello
<i>MAP2K2</i>	MITOGEN-ACTIVATED PROTEIN KINASE KINASE 2	AD	Noonan/CFC/Costello
<i>MAP3K8</i>	MITOGEN-ACTIVATED PROTEIN KINASE KINASE KINASE 8	AD	Noonan syndrome
<i>MAT2A</i>	METHIONINE ADENOSYLTRANSFERASE II, ALPHA	AD	FTAAD
<i>MED12</i>	MEDIATOR COMPLEX SUBUNIT 12	AD	Lujan syndrome, fTAAD
<i>MFAP5</i>	MICROFIBRILLAR-ASSOCIATED PROTEIN 5	AD	FTAAD

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Gene	Protein	Inheritance	Disease Association(s)
<i>MIB1</i>	MINDBOMB E3 UBIQUITIN PROTEIN LIGASE 1	AD	LVNC
<i>MRPL3</i>	MITOCHONDRIAL RIBOSOMAL PROTEIN L3	AR	HCM
<i>MRPS22</i>	MITOCHONDRIAL RIBOSOMAL PROTEIN S22	AR	Cardiomyopathy
<i>MTO1</i>	MITOCHONDRIAL TRANSLATION OPTIMIZATION 1, <i>S. CEREVIAE</i> , HOMOLOG OF	AR	HCM
<i>MURC</i>	MUSCLE-RELATED COILED-COIL PROTEIN	AD	DCM
<i>MYBPC3</i>	MYOSIN-BINDING PROTEIN C, CARDIAC	AD	HCM, DCM
<i>MYH11</i>	MYOSIN, HEAVY CHAIN 11, SMOOTH MUSCLE	AD	fTAAD
<i>MYH6</i>	MYOSIN, HEAVY CHAIN 6, CARDIAC MUSCLE, ALPHA	AD	CHD, DCM, HCM, SSS
<i>MYH7</i>	MYOSIN, HEAVY CHAIN 7, CARDIAC MUSCLE, BETA	AD	DCM, HCM, myopathy
<i>MYL2</i>	MYOSIN, LIGHT CHAIN 2, REGULATORY, CARDIAC, SLOW	AD	HCM
<i>MYL3</i>	MYOSIN, LIGHT CHAIN 3, ALKALI, VENTRICULAR, SKELETAL, SLOW	AD, AR	HCM
<i>MYL4</i>	MYOSIN, LIGHT CHAIN 4, ALKALI, ATRIAL, EMBRYONIC	AD, AR	AF
<i>MYLK</i>	MYOSIN LIGHT CHAIN KINASE	AD	fTAAD
<i>MYLK2</i>	MYOSIN LIGHT CHAIN KINASE 2	AD	HCM
<i>MYO6</i>	MYOSIN VI	AR	HCM and hearing loss
<i>MYOM1</i>	MYOMESIN 1	AD	HCM, DCM
<i>MYOZ2</i>	MYOZENIN 2	AD	HCM
<i>MYPN</i>	MYOPALLADIN	AD	DCM, RCM, HCM
<i>NEBL</i>	NEBULETTE	AD	DCM, endocardial fibroelastosis
<i>NEXN</i>	NEXILIN	AD	DCM, HCM
<i>NKX2-5</i>	NK2 HOMEOBOX 5	AD	CHD, CCD
<i>NKX2-6</i>	NK2, DROSOPHILA, HOMOLOG OF, 6	AD, AR	CHD, AF, HB
<i>NOS1AP</i>	NITRIC OXIDE SYNTHASE 1 (NEURONAL) ADAPTOR PROTEIN	AD	LQTS

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Gene	Protein	Inheritance	Disease Association(s)
<i>NOTCH1</i>	NOTCH, DROSOPHILA, HOMOLOG OF, 1	AD	fTAAD
<i>NPPA</i>	NATRIURETIC PEPTIDE PRECURSOR A	AD, AR	AF, atrial DCM
<i>NRAS</i>	NEUROBLASTOMA RAS VIRAL ONCOGENE HOMOLOG	AD	Noonan/CFC/Costello
<i>PCSK9</i>	PROPROTEIN CONVERTASE SUBTILISIN/KEXIN TYPE 9	AD	HeFH/HoFH
<i>PDLIM3</i>	PDZ AND LIM DOMAIN PROTEIN 3	AD	HCM, DCM
<i>PI4KA</i>	PHOSPHATIDYLINOSITOL 4-KINASE, CATALYTIC, ALPHA	AR	LQTS
<i>PKP2</i>	PLAKOPHILIN 2	AD	ARVC, DCM, BrS
<i>PLEC</i>	PLECTIN	AR	Cardiomyopathy, muscular dystrophy
<i>PLEKHM2</i>	PLECKSTRIN HOMOLOGY DOMAIN-CONTAINING PROTEIN, FAMILY M, MEMBER 2	AR	DCM, LVNC
<i>PLN</i>	PHOSPHOLAMBAN	AD	DCM, HCM, ARVC
<i>PLOD1</i>	PROCOLLAGEN-LYSINE, 2-OXOGLUTARATE 5-DIOXYGENASE	AR	kEDS, fTAAD
<i>PLOD3</i>	PROCOLLAGEN-LYSINE, 2-OXOGLUTARATE 5-DIOXYGENASE 3	AR	Connective Tissue disorder
<i>PPA2</i>	PYROPHOSPHATASE, INORGANIC, 2	AR	Sudden cardiac arrest, infancy Infantile cardiomyopathy
<i>PRDM16</i>	PR DOMAIN CONTAINING 16	AD	DCM, LVNC
<i>PRDM5</i>	PR DOMAIN 5	AR	BCS
<i>PRKAG2</i>	PROTEIN KINASE, AMP-ACTIVATED, NONCATALYTIC, GAMMA2	AD	HCM, Wolff-Parkinson-White syndrome
<i>PRKG1</i>	PROTEIN KINASE, cGMP-DEPENDENT, REGULATORY, TYPE I	AD	fTAAD
<i>PTPN11</i>	PROTEIN-TYROSINE PHOSPHATASE, NONRECEPTOR-TYPE 11	AD	Noonan/CFC/Costello
<i>PYCR1</i>	PYRROLINE-5-CARBOXYLATE REDUCTASE 1	AR	Cutis laxa, autosomal recessive
<i>RAF1</i>	V-RAF-1 MURINE LEUKEMIA VIRAL ONCOGENE HOMOLOG 1	AD	Noonan/CFC/Costello
<i>RANGRF</i>	RAN GUANINE NUCLEOTIDE RELEASE FACTOR	AD	BrS

Test Information Sheet

Gene	Protein	Inheritance	Disease Association(s)
RASA1	RAS P21 PROTEIN ACTIVATOR 1	AD	Capillary malformation-arteriovenous malformation, Parkes Weber syndrome, Basal cell carcinoma
RASA2	RAS p21 PROTEIN ACTIVATOR 2	AD	Noonan syndrome
RBM20	RNA-BINDING MOTIF PROTEIN 20	AD	DCM
RIN2	RAS AND RAB INTERACTOR 2	AR	MACS
RIT1	RAS-LIKE WITHOUT CAAX 1	AD	Noonan syndrome
RRAS	RELATED RAS VIRAL ONCOGENE HOMOLOG	AD	Noonan syndrome-like
RYR2	RYANODINE RECEPTOR 2	AD	ARVC, CPVT, LQTS
SCARF2	SCAVENGER RECEPTOR CLASS F, MEMBER 2	AR	Van den Ende-Gupta syndrome
SCN10A	SODIUM CHANNEL, VOLTAGE-GATED, TYPE X, ALPHA SUBUNIT	AD	BrS
SCN1B	SODIUM CHANNEL, VOLTAGE-GATED, TYPE I, BETA SUBUNIT	AD, AR	BrS, Cardiac conduction disease
SCN2B	SODIUM CHANNEL, VOLTAGE-GATED, TYPE II, BETA SUBUNIT	AD	BrS, AF
SCN3B	SODIUM CHANNEL, VOLTAGE-GATED, TYPE III, BETA SUBUNIT	AD	BrS, AF, VF, SIDS
SCN4A	SODIUM CHANNEL, VOLTAGE-GATED, TYPE IV, ALPHA SUBUNIT	AD, AR	BrS w/ muscle stiffness
SCN4B	SODIUM CHANNEL, VOLTAGE-GATED, TYPE IV, BETA SUBUNIT	AD	LQTS
SCN5A	SODIUM CHANNEL, VOLTAGE-GATED, TYPE V, ALPHA SUBUNIT	AD, AR	BrS, DCM, Heart block, LQTS, SSS, SIDS
SCNN1A	SODIUM CHANNEL, NONVOLTAGE-GATED 1, ALPHA SUBUNIT	AD	BrS
SCO2	SCO2 CYTOCHROME c OXIDASE ASSEMBLY PROTEIN	AD, AR	HCM
SGCD	SARCOGLYCAN, DELTA	AD, AR	DCM, LGMD
SHOC2	SOC-2 HOMOLOG	AD	Noonan-like syndrome with loose Anagen Hair 1
SKI	V-SKI AVIAN SARCOMA VIRAL ONCOGENE HOMOLOG	AD	Shprintzen-Goldberg syndrome
SLC25A20	SOLUTE CARRIER FAMILY 2 (FACILITATED GLUCOSE TRANSPORTER), MEMBER 10	AR	Arterial tortuosity syndrome

Test Information Sheet

Gene	Protein	Inheritance	Disease Association(s)
SLC25A3	SOLUTE CARRIER FAMILY 25 (MITOCHONDRIAL CARRIER), MEMBER 3	AR	Cardiomyopathy, myopathy
SLC25A4	SOLUTE CARRIER FAMILY 25 (MITOCHONDRIAL CARRIER, ADENINE NUCLEOTIDE TRANSLOCATOR), MEMBER 4	AD, AR	HCM, myopathy
SLC2A10	SOLUTE CARRIER FAMILY 2 (FACILITATED GLUCOSE TRANSPORTER), MEMBER 10	AR	Arterial tortuosity syndrome
SLC2A5	SOLUTE CARRIER FAMILY 2 (FACILITATED GLUCOSE/FRUCTOSE TRANSPORTER), MEMBER 5	AD	LQTS
SLC39A13	SOLUTE CARRIER FAMILY 39 MEMBER 13	AR	spEDS
SLMAP	SARCOLEMMAL-ASSOCIATED PROTEIN	AD	BrS
SMAD1	MOTHERS AGAINST DECAPENTAPLEGIC, DROSOPHILA, HOMOLOG OF, 1	AD	PAH
SMAD2	MOTHERS AGAINST DECAPENTAPLEGIC, DROSOPHILIA, HOMOLOG OF, 2	AD	FTAAD, LDS, CHD
SMAD3	MOTHERS AGAINST DECAPENTAPLEGIC, DROSOPHILA, HOMOLOG OF, 3	AD	LDS
SMAD4	MOTHERS AGAINST DECAPENTAPLEGIC, DROSOPHILA, HOMOLOG OF, 4	AD	Juvenile polyposis/HHT; Myhre syndrome
SMAD6	MOTHERS AGAINST DECAPENTAPLEGIC, DROSOPHILA, HOMOLOG OF, 6	AD	PAH
SMAD9	MOTHERS AGAINST DECAPENTAPLEGIC, DROSOPHILA, HOMOLOG OF, 9	AD	PAH
SMS	SPERMINE SYNTHASE	XL	Connective Tissue disorder
SNTA1	ALPHA SYNTROPHIN	AD	LQTS
SOS1	SON OF SEVENLESS, DROSOPHILA, HOMOLOG 1	AD	Noonan/CFC/Costello
SOS2	SON OF SEVENLESS, DROSOPHILA, HOMOLOG 2	AD	Noonan syndrome
SPRY1	SPROUTY, DROSOPHILA, HOMOLOG OF, 1	unknown	Noonan syndrome
SYNE1	SPECTRIN REPEAT-CONTAINING NUCLEAR ENVELOPE PROTEIN 1	AD	EMD
SYNE2	SPECTRIN REPEAT-CONTAINING NUCLEAR ENVELOPE PROTEIN 2	AD	EMD

Test Information Sheet

Gene	Protein	Inheritance	Disease Association(s)
<i>TAB2</i>	TAK1-BINDING PROTEIN 2	AD	CHD
<i>TANGO2</i>	TRANSPORT AND GOLGI ORGANIZATION 2, DROSOPHILA, HOMOLOG OF	AR	Noonan syndrome
<i>TAZ</i>	TAFAZZIN	XL	DCM, LVNC, Barth syndrome
<i>TBX1</i>	T-BOX 1	AD	Velocardiofacial syndrome, CHD
<i>TBX5</i>	T-BOX 5	AD	Holt-Oram syndrome
<i>TBX20</i>	T-BOX 20	AD	CHD, DCM, LVNC
<i>TCAP</i>	TITIN-CAP (TELETHONIN)	AD, AR	HCM, DCM, LGMD
<i>TECRL</i>	TRANS-2,3-ENOYL-CoA REDUCTASE-LIKE PROTEIN	AR	CPVT3
<i>TGFB2</i>	TRANSFORMING GROWTH FACTOR, BETA-2	AD	LDS
<i>TGFB3</i>	TRANSFORMING GROWTH FACTOR BETA 3	AD	ARVC, Loeys-Dietz syndrome
<i>TGFBR1</i>	TRANSFORMING GROWTH FACTOR-BETA RECEPTOR, TYPE I	AD	LDS
<i>TGFBR2</i>	TRANSFORMING GROWTH FACTOR-BETA RECEPTOR, TYPE II	AD	LDS
<i>TMEM43</i>	TRANSMEMBRANE PROTEIN 43	AD	ARVC, EMD
<i>TMPO</i>	THYMOPOIETIN	AD	DCM
<i>TNNC1</i>	TROPONIN C, SLOW	AD	DCM, HCM
<i>TNNI3</i>	TROPONIN I, CARDIAC	AD, AR	DCM, HCM, RCM
<i>TNNI3K</i>	TNNI3-INTERACTING KINASE	AD	DCM
<i>TNNT2</i>	TROPONIN T2, CARDIAC	AD	DCM, HCM, RCM, LVNC
<i>TNXB</i>	TENASCIN XB	AR	Ehlers-Danlos syndrome, classic-like, 1
<i>TOR1AIP1</i>	TORSIN-1A-INTERACTING PROTEIN 1	AR	LGMD, Contractures, DCM
<i>TPM1</i>	TROPOMYOSIN 1	AD	DCM, HCM
<i>TRDN</i>	TRIADIN	AR	CPVT, LQTS
<i>TRIM63</i>	TRIPARTITE MOTIF-CONTAINING PROTEIN 63	AD	HCM
<i>TRPM4</i>	TRANSIENT RECEPTOR POTENTIAL CATION CHANNEL, SUBFAMILY M, MEMBER 4	AD	HB, BrS
<i>TSFM</i>	Ts TRANSLATION ELONGATION FACTOR, MITOCHONDRIAL	AR	HCM
<i>TTN</i>	TITIN	AD	DCM, ARVC, TTN-related myopathies and muscular dystrophies
<i>TTR</i>	TRANSTHYRETIN	AD	TTR-related amyloidosis
<i>TXNRD2</i>	THIOREDOXIN REDUCTASE 2	AD, AR	DCM
<i>UPF3B</i>	UPF3, YEAST, HOMOLOG OF, B	XL	Lujan syndrome
<i>VCL</i>	VINCULIN	AD	HCM, DCM, LVNC
<i>XK</i>	KELL BLOOD GROUP PROTEIN, MCLEOD SYNDROME-ASSOCIATED	XL	Cardiomyopathy, muscular dystrophy, AF
<i>ZNF469</i>	ZINC FINGER PROTEIN 469	AR	BCS

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GeneDx

Abbreviations: AD- Autosomal dominant; aEDS- arthrochalasia Ehlers-Danlos syndrome; AF- Atrial Fibrillation; AR- Autosomal recessive; ARVC – Arrhythmogenic Right Ventricular Cardiomyopathy; ARFH – Autosomal recessive familial hypercholesterolemia; BCS – Brittle Cornea Syndrome; BMD – Becker Muscular Dystrophy; BrS – Brugada Syndrome; cEDS-classical Ehlers-Danlos syndrome; CHD – Congenital Heart Defects; CPVT – Catecholaminergic Polymorphic Ventricular Tachycardia; cvEDS- Cardiac-valvular Ehlers-Danlos syndrome; DCM – Dilated Cardiomyopathy; dEDS- dermatosparaxis Ehlers-Danlos syndrome; DMD- Duchenne Muscular Dystrophy; EMD – Emery Dreifuss Muscular Dystrophy; ERS-Early repolarization syndrome; FTAAD – familial thoracic aortic aneurysm and dissection; GSD- Glycogen storage disease, HB- Heart Block; HCM – Hypertrophic Cardiomyopathy; HeFH – Heterozygous familial Hypercholesterolemia (FH); HoFH – Homozygous FH; JLNS – Jervell and Lange-Nielsen Syndrome; JP/HHT – juvenile polyposis/hereditary hemorrhagic telangiectasia; kEDS- kyphoscoliotic Ehlers-Danlos syndrome; LDS – Loeys-Dietz syndrome; LGMD – Limb Girdle Muscular Dystrophy; LQTS – Long QT Syndrome; LVNC – Left Ventricular Non-Compaction; MACS - Macrocephaly, alopecia, cutis laxa, and scoliosis; mcEDS- musculocontractural Ehlers-Danlos syndrome; OHS – Occipital horn syndrome; RCM – Restrictive Cardiomyopathy; SIDS – Sudden Infant Death Syndrome; spEDS- Spondylocheirodysplasia type Ehlers-Danlos syndrome; SSS – Sick Sinus Syndrome; vEDS- vascular Ehlers-Danlos syndrome; VF- Ventricular fibrillation; XL- X-linked

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