



Sample: 16012132564

Inputs

The patient is diagnosed Dejerine sottas disease. The suspected phenotypes for this patient are hammertoes; distal muscle weakness; pes cavus.



Gene	PRX
Mode of inheritance	recessive
Consequence	stop gained
HGVSp	ENSP00000326018.6:p.Arg953Ter
HGVSc	ENST00000324001.7:c.2857C>T
rsld	rs104894714
gnomAD allele frequency	3.183e-05
ExAC allele frequency	
REVEL	
Impact	high
Location	chr19: 40901402-40901403
Variant type	snp
Variant quality	1933.95

■ Notes:

Test note

— Demo User

Gene	SCN8A
Mode of inheritance	de novo
Consequence	missense variant
HGVSp	ENSP00000346534.4:p.Arg1872Trp
HGVSc	ENST00000354534.6:c.5614C>T
rsId	rs796053228

gnomAD allele frequency	0
ExAC allele frequency	
REVEL	0.901
Impact	moderate
Location	chr12: 52200884-52200885
Variant type	snp
Variant quality	429.716

Gene	DLL4
Mode of inheritance	de novo
Consequence	missense variant
HGVSp	ENSP00000249749.5:p.Cys653Trp
HGVSc	ENST00000249749.5:c.1959T>G
rsld	rs533126562
gnomAD allele frequency	0
ExAC allele frequency	
REVEL	0.785
Impact	moderate
Location	chr15: 41229631-41229632
Variant type	snp
Variant quality	383.909

Gene	ABCA3
Mode of inheritance	compound het

Consequence	missense variant
HGVSp	ENSP00000301732.5:p.Pro766Ser
HGVSc	ENST00000301732.5:c.2296C>T
rsId	rs45592239
gnomAD allele frequency	0.001708
ExAC allele frequency	
REVEL	0.113
Impact	moderate
Location	chr16: 2345709-2345710
Variant type	snp
Variant quality	1166.76

Gene	ABCA3
Mode of inheritance	compound het
Consequence	missense variant
HGVSp	ENSP00000301732.5:p.Tyr247Ser
HGVSc	ENST00000301732.5:c.740A>C
rsld	rs775442517
gnomAD allele frequency	0.001114
ExAC allele frequency	
REVEL	0.508
Impact	moderate
Location	chr16: 2369715-2369716
Variant type	snp
Variant quality	1.09117e-06

Gene	DVL3
Mode of inheritance	de novo
Consequence	missense variant
HGVSp	ENSP00000316054.3:p.Arg221Gly
HGVSc	ENST00000313143.3:c.661C>G
rsld	rs76594728
gnomAD allele frequency	0
ExAC allele frequency	
REVEL	0.296
Impact	moderate
Location	chr3: 183882962-183882963
Variant type	snp
Variant quality	487.882

Gene	NOTCH4
Mode of inheritance	compound het
Consequence	missense variant
HGVSp	ENSP00000364163.3:p.Gly1121Arg
HGVSc	ENST00000375023.3:c.3361G>A
rsld	rs72846312
gnomAD allele frequency	7.417e-05
ExAC allele frequency	
REVEL	0.204
Impact	moderate

Location	chr6: 32170247-32170248
Variant type	snp
Variant quality	1096.51

Gene	NOTCH4
Mode of inheritance	compound het
Consequence	missense variant
HGVSp	ENSP00000364163.3:p.Asp272Gly
HGVSc	ENST00000375023.3:c.813_815delinsGGG
rsId	rs71556915
gnomAD allele frequency	0
ExAC allele frequency	
REVEL	
Impact	moderate
Location	chr6: 32188640-32188643
Variant type	complex
Variant quality	1354.34

? Unknown Significance

Gene	LMNA
Mode of inheritance	de novo
Consequence	missense variant
HGVSp	ENSP00000357283.4:p.His565Pro
HGVSc	ENST00000368300.4:c.1694A>C
rsld	Not found
gnomAD allele frequency	0
ExAC allele frequency	
REVEL	0.560
Impact	moderate
Location	chr1: 156107530-156107531
Variant type	snp
Variant quality	1.36779e-10

Genes

Phenotypes searched	Genes (100)
GTR: Dejerine-Sottas disease	MPZ, EGR2, PMP22, PRX, GDAP1, FGD4, DNM2, HSPB1, FIG4, DYNC1H1, NEFL, MFN2, LMNA, MTMR2, NDRG1, SH3TC2, SBF2, RAB7A, LITAF, HSPB8, TRPV4, MED25, LRSAM1, BSCL2, SBF1, SPG11, HK1, PEX7, PHYH, SPAST, GAN, GJB1, AARS1, GARS1, GNB4, YARS1, AIFM1, COX6A1, PRPS1, PDK3, DHTKD1, PLEKHG5, KARS1, KIF1B, TRIM2, INF2, C12orf65, ATP1A1, HADHB, IGHMBP2, TFG, GLA, HINT1, DNAJB2, MARS1, HARS1, KIF5A, TTR, CTDP1, FBLN5, SPTLC1, MORC2, DNMT1, HOXD10, SCN9A, SPTLC2, SLC12A6, ATL1, REEP1, TSEN54, ALMS1, RARS2, FA2H, L1CAM, SEPT9, SMCHD1, PLP1, ALS2, DUX4, ETS1, SPG7, FGF3, ISCU, MARS2, GARS, TSEN2, TSEN34, VRK1, GRN, SPG20, KIAA0196, AARS, EXOSC3, OPA3, GATA2, SEPSECS, FRG1, PRKCA, YBX1, AKT1
Phenolyzer: dejerine sottas disease	
HPO:	

Disclaimers

Genetic testing information has caveats and should not be considered a definitive diagnosis.

References/Methodology

DNA sequencing was performed in accordance with established Utah Genome Project (UGP) methodologies including sample preparation, sequencing and data analysis.