



Sample: 16012132564

Clinical Description

a description goes here

Clinical Notes

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

Significant Variants

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:

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Test note

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

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)
Phenotypes searched GTR: • Dejerine-Sottas disease Phenolyzer: • Dejerine sottas disease HPO:	Genes (100) 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

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.