

SNiPAcad

Block annotations

Block info

genomic range	chr4:88,400,924-88,431,892 <i>e!</i>
block size	30,969 bp
variant count	37 variants

Basic features

Conservation/deleteriousness		Linked genes	
phyloP	$\mu = -0.799$ [-3.224 – 0.94]	gene(s) hit or close-by	SPARCL1 <i>e!</i>
phastCons	$\mu = 0.045$ [0 – 0.942]	eQTL gene(s)	HSD17B11 <i>e!</i> , HSD17B13 <i>e!</i> , NUDT9 <i>e!</i> , PPM1K <i>e!</i> , RP11-529H2.2 <i>e!</i> , SPARCL1 <i>e!</i>
GERP++	$\mu = 0.298$ [-4.01 – 4.25]	potentially regulated gene(s)	RP11-529H2.1 <i>e!</i> , SPARCL1 <i>e!</i>
CADD score	$\mu = 3.588$ [0.322 – 10.4]	disease gene(s)	PPM1K <i>e!</i>

Trait annotations

Disease gene annotation

gene	trait	source DB	source entry/link
PPM1K <i>e!</i>	MAPLE SYRUP URINE DISEASE, MILD VARIANT	OMIM	MIM:615135 
PPM1K <i>e!</i>	Intermediate maple syrup urine disease	OrphaNet	OrphaNet:268162 



Direct effect on transcript

Amino acid sequence alteration

gene	effect type	affected transcript	RefSeq id	protein	exchanged AA's	exchanged codons	SIFT prediction	PolyPhen prediction	variant(s)
SPARCL1 <i>e!</i>	missense variant	ENST00000458304 <i>e!</i>	?	ENSP00000406251 <i>e!</i>	A/D	gCt/gAt	?	benign	1
SPARCL1 <i>e!</i>	missense variant	ENST00000418378 <i>e!</i>	NM_001128310.1	ENSP00000414856 <i>e!</i>	2	2			2
SPARCL1 <i>e!</i>	missense variant	ENST00000434434 <i>e!</i>	?	ENSP00000416971 <i>e!</i>	2	2			2
SPARCL1 <i>e!</i>	missense variant	ENST00000282470 <i>e!</i>	NM_004684.4	ENSP00000282470 <i>e!</i>	2	2			2
SPARCL1 <i>e!</i>	missense variant	ENST00000535835 <i>e!</i>	?	ENSP00000438188 <i>e!</i>	2	2			2
SPARCL1 <i>e!</i>	missense variant	ENST00000509407 <i>e!</i>	?	ENSP00000423483 <i>e!</i>	2	2			2
SPARCL1 <i>e!</i>	missense variant	ENST00000512317 <i>e!</i>	?	ENSP00000423448 <i>e!</i>	2	2			2
SPARCL1 <i>e!</i>	missense variant	ENST00000543631 <i>e!</i>	?	ENSP00000444832 <i>e!</i>	A/D	gCt/gAt	tolerated	benign	1

Direct effect on regulation

cis-eQTL

gene	transcript	probe	tissue	min(statistic) (type)	source	variant(s)
SPARCL1 <i>e!</i>	ENST00000282470 <i>e!</i>	ILMN_2218208 <i>e!</i>	adipocyte	1.24×10 ⁻⁶ (p-value)	MuTHER consortium 	7
SPARCL1 <i>e!</i>	ENST00000503414 <i>e!</i>					
SPARCL1 <i>e!</i>	ENST00000418378 <i>e!</i>					
SPARCL1 <i>e!</i>	ENST00000282470 <i>e!</i>	ILMN_1795251 <i>e!</i>	adipocyte	1.65×10 ⁻⁶ (p-value)	MuTHER consortium 	7
SPARCL1 <i>e!</i>	ENST00000503414 <i>e!</i>					
SPARCL1 <i>e!</i>	ENST00000418378 <i>e!</i>					

NUDT9 <i>e!</i>	ENST00000302174 <i>e!</i>	ILMN_1680239 <i>e!</i>	blood	5.72×10 ⁻⁶ (p-value)	Westra et al. <i>!</i>	6
NUDT9 <i>e!</i>	ENST00000473942 <i>e!</i>					
NUDT9 <i>e!</i>	ENST00000440591 <i>e!</i>					
HSD17B11 <i>e!</i>	ENST00000508413 <i>e!</i>	ILMN_1675117 <i>e!</i>	blood	5.44×10 ⁻⁷ (p-value)	Westra et al. <i>!</i>	4
HSD17B11 <i>e!</i>	ENST00000358290 <i>e!</i>					
HSD17B11 <i>e!</i>	?	ENSG00000198189 <i>e!</i>	tibial nerve	3.20×10 ⁻⁶ (p-value)	GTEx Portal V6 <i>!</i>	25
SPARCL1 <i>e!</i>	?	ENSG00000152583 <i>e!</i>	sun exposed skin	5.36×10 ⁻⁷ (p-value)	GTEx Portal V6 <i>!</i>	26
HSD17B11 <i>e!</i>	?	ENSG00000198189 <i>e!</i>	esophagus mucosa	1.01×10 ⁻⁷ (p-value)	GTEx Portal V6 <i>!</i>	36
SPARCL1 <i>e!</i>	?	ENSG00000152583 <i>e!</i>	subcutaneous adipocytes	5.55×10 ⁻⁶ (p-value)	GTEx Portal V6 <i>!</i>	14
HSD17B13 <i>e!</i>	?	ENSG00000170509 <i>e!</i>	thyroid	2.57×10 ⁻⁶ (p-value)	GTEx Portal V6 <i>!</i>	12
PPM1K <i>e!</i>	ENST00000608933 <i>e!</i>	ILMN_2070043 <i>e!</i>	b-cell	6.14×10 ⁻⁴ (p-value)	Fairfax et al. <i>!</i>	1
PPM1K <i>e!</i>	ENST00000295908 <i>e!</i>					
HSD17B11 <i>e!</i>	?	ENSG00000198189 <i>e!</i>	pancreas	5.48×10 ⁻⁶ (p-value)	GTEx Portal V6 <i>!</i>	3
RP11-529H2.2 <i>e!</i>	?	ENSG00000255723 <i>e!</i>	pancreas	1.05×10 ⁻⁵ (p-value)	GTEx Portal V6 <i>!</i>	1

Putative effect on regulation

FANTOM5 expressed promoter

SNiPA promoter id	variant(s)	associated transcript(s)	gene
FFCP00000660191 <i>e!</i>	1	ENST00000503414 <i>e!</i> , ENST00000541496 <i>e!</i>	SPARCL1 <i>e!</i>

ENCODE promoter-associated distal DHS (Enhancer)

SNiPA enhancer id	variant(s)	associated SNiPA promoter id	associated gene(s)
ENCE00000384981 <i>e!</i>	1	ENCP00000041684	RP11-529H2.1 <i>e!</i>

Regulatory feature cluster

element id	variant(s)	tissue/cell	factors
ENSR00001244943 <i>e!</i> (CTCF binding site)	1	embryonic stem cell (H1ESC)	Rad21, CTCF, DNase1
		blood (K562)	CTCF
		skin (NHDF-AD)	CTCF, DNase1, H3K4me2
		muscle (HSMM)	CTCF
		breast (HMEC)	CTCF
		cervix (HeLa-S3)	CTCF
		endothelium (HUVEC)	H3K36me3, CTCF
		liver (HepG2)	CTCF
		blood (GM12878)	CTCF
		skin (NHEK)	CTCF
ENSR00001244946 <i>e!</i> (promoter flanking region)	4	cervix (HeLa-S3)	DNase1
		muscle (HSMM)	DNase1
ENSR00001686554 <i>e!</i> (CTCF binding site)	2	cervix (HeLa-S3)	DNase1
		muscle (HSMM)	DNase1
ENSR00001244947 <i>e!</i> (CTCF binding site)	1	cervix (HeLa-S3)	CTCF
		endothelium (HUVEC)	H3K36me3, CTCF
		embryonic stem cell (H1ESC)	Rad21, CTCF, DNase1
		liver (HepG2)	CTCF
		blood (K562)	CTCF
		skin (NHEK)	CTCF
ENSR00001432957 <i>e!</i> (promoter flanking region)	8	NHLF	H3K27ac, DNase1
		embryonic stem cell (H1ESC)	CTCF, DNase1
		HSMMtube	DNase1
		Osteobl	H3K4me2, H3K27ac

blood (K562)	CTCF
skin (NHDF-AD)	H3K4me1, DNase1, H3K4me2, H3K27ac
muscle (HSMM)	H3K4me1, H2AZ, H3K27ac, H3K4me2, DNase1
endothelium (HUVEC)	CTCF
nervous (NH-A)	DNase1
skin (NHEK)	CTCF

Variation proximal to gene

gene	variant type	min(distance)	transcript	RefSeq id	protein	variant(s)
SPARCL1 <i>e!</i>	downstream gene variant	2545	ENST00000535835 <i>e!</i>	?	ENSP00000438188 <i>e!</i>	1
SPARCL1 <i>e!</i>	downstream gene variant	2504	ENST00000434434 <i>e!</i>	?	ENSP00000416971 <i>e!</i>	1
SPARCL1 <i>e!</i>	downstream gene variant	49	ENST00000543631 <i>e!</i>	?	ENSP00000444832 <i>e!</i>	2
SPARCL1 <i>e!</i>	downstream gene variant	529	ENST00000458304 <i>e!</i>	?	ENSP00000406251 <i>e!</i>	2
SPARCL1 <i>e!</i>	downstream gene variant	2568	ENST00000512317 <i>e!</i>	?	ENSP00000423448 <i>e!</i>	1
SPARCL1 <i>e!</i>	downstream gene variant	2497	ENST00000541496 <i>e!</i>	?	ENSP00000445678 <i>e!</i>	1
SPARCL1 <i>e!</i>	downstream gene variant	2497	ENST00000509407 <i>e!</i>	?	ENSP00000423483 <i>e!</i>	1

Putative effect on transcript

Intron variant

gene	affected transcript	RefSeq id	protein	variant(s)
SPARCL1 <i>e!</i>	ENST00000543631 <i>e!</i>	?	ENSP00000444832 <i>e!</i>	31
SPARCL1 <i>e!</i>	ENST00000503414 <i>e!</i>	?	ENSP00000422903 <i>e!</i>	35
SPARCL1 <i>e!</i>	ENST00000282470 <i>e!</i>	NM_004684.4	ENSP00000282470 <i>e!</i>	35
SPARCL1 <i>e!</i>	ENST00000512317 <i>e!</i>	?	ENSP00000423448 <i>e!</i>	31
SPARCL1 <i>e!</i>	ENST00000418378 <i>e!</i>	NM_001128310.1	ENSP00000414856 <i>e!</i>	35
SPARCL1 <i>e!</i>	ENST00000458304 <i>e!</i>	?	ENSP00000406251 <i>e!</i>	31
SPARCL1 <i>e!</i>	ENST00000541496 <i>e!</i>	?	ENSP00000445678 <i>e!</i>	31
SPARCL1 <i>e!</i>	ENST00000535835 <i>e!</i>	?	ENSP00000438188 <i>e!</i>	31
SPARCL1 <i>e!</i>	ENST00000434434 <i>e!</i>	?	ENSP00000416971 <i>e!</i>	31
SPARCL1 <i>e!</i>	ENST00000509407 <i>e!</i>	?	ENSP00000423483 <i>e!</i>	31

5'-UTR variant

gene	affected transcript	RefSeq id	protein	variant(s)
SPARCL1 <i>e!</i>	ENST00000503414 <i>e!</i>	?	ENSP00000422903 <i>e!</i>	2
SPARCL1 <i>e!</i>	ENST00000541496 <i>e!</i>	?	ENSP00000445678 <i>e!</i>	2

