

SNiPACard

Block annotations

Block info

genomic range	chr9:136,115,176-136,184,526 <i>e!</i>
block size	69,351 bp
variant count	25 variants

Basic features

Conservation/deleteriousness		Linked genes	
phyloP	$\mu = -0.373$ [-2.969 – 1.925]	gene(s) hit or close-by	ABO <i>e!</i> , LCN1P2 <i>e!</i> , Y_RNA <i>e!</i>
phastCons	$\mu = 0.069$ [0 – 0.636]	eQTL gene(s)	ABO <i>e!</i> , MED22 <i>e!</i> , OBP2B <i>e!</i> , SLC2A6 <i>e!</i>
GERP++	$\mu = -0.804$ [-8.76 – 2.51]	potentially regulated gene(s)	ABO <i>e!</i> , ADAMTS13 <i>e!</i> , ADAMTS13 <i>e!</i> , ADAMTS13 <i>e!</i> , ADAMTS13 <i>e!</i> , AK8 <i>e!</i> , C9orf9 <i>e!</i> , CELP <i>e!</i> , DBH <i>e!</i> , GF1B <i>e!</i> , GTF3C5 <i>e!</i> , OBP2B <i>e!</i> , OBP2B <i>e!</i> , RALGDS <i>e!</i> , REXO4 <i>e!</i> , REXO4 <i>e!</i> , SARDH <i>e!</i> , SURF1 <i>e!</i> , SURF1 <i>e!</i> , SURF2 <i>e!</i> , SURF2 <i>e!</i> , SURF4 <i>e!</i> , SURF4 <i>e!</i>
CADD score	$\mu = 3.314$ [0.003 – 16.8]	disease gene(s)	SURF1 <i>e!</i> , DBH <i>e!</i> , ADAMTS13 <i>e!</i> , ADAMTS13 <i>e!</i> , SARDH <i>e!</i> , GF1B <i>e!</i>

Trait annotations

Variant association

trait	min(p-value)	source DB	source entry/link	variant(s)
Intraocular pressure	<3.00×10 ⁻¹¹	GWAS Catalog	25173106	1
High serum lipase activity	<1.00×10 ⁻²²	GWAS Catalog	25028398	1
Elevated serum carcinoembryonic antigen levels	<2.00×10 ⁻²⁴	GWAS Catalog	24941225	1
Urinary metabolites (H-NMR features)	<4.00×10 ⁻¹²	GWAS Catalog	24586186	1
End-stage coagulation	<2.00×10 ⁻¹⁷	GWAS Catalog	23381943	1
Tumor biomarkers	<7.00×10 ⁻¹⁰⁵	GWAS Catalog	23300138	1
Coagulation factor levels	<2.00×10 ⁻¹³⁸	GWAS Catalog	23267103	1
Mean corpuscular hemoglobin concentration	<4.00×10 ⁻⁸	GWAS Catalog	20139978	1
Parkinson disease	8.66×10 ⁻⁶	dbGaP	pha002868 dbGaP	1

Variant annotation

trait	type	source DB	source entry/link	Variant(s)
Abo blood group system, b(a) phenotype	association	ClinVar	RCV000019312.3 ClinVar	1
ABO BLOOD GROUP SYSTEM, CIS-AB PHENOTYPE	OMIM curated	OMIM	MIM:110300 OMIM [®]	1

Disease gene annotation

gene	trait	source DB	source entry/link
SURF1 <i>e!</i>	Complex IV deficiency	DECIPHER	MIM:220110 OMIM [®]
SURF1 <i>e!</i>	Leigh Syndrome (nuclear DNA mutation)	DECIPHER	MIM:256000 OMIM [®]
<i>e!</i>	Complex IV deficiency	DECIPHER	MIM:220110 OMIM [®]
<i>e!</i>	Leigh Syndrome (nuclear DNA mutation)	DECIPHER	MIM:256000 OMIM [®]
DBH <i>e!</i>	DOPAMINE BETA-HYDROXYLASE DEFICIENCY, CONGENITAL	OMIM	MIM:223360 OMIM [®]
ADAMTS13 <i>e!</i>	GELEOPHYSIC DYSPLASIA 1	OMIM	MIM:231050 OMIM [®]
SURF1 <i>e!</i>	LEIGH SYNDROME	OMIM	MIM:256000 OMIM [®]
ADAMTS13 <i>e!</i>	THROMBOTIC THROMBOCYTOPENIC PURPURA, CONGENITAL	OMIM	MIM:274150 OMIM [®]
SARDH <i>e!</i>	SARCOSINEMIA	OMIM	MIM:268900 OMIM [®]
GF1B <i>e!</i>	BLEEDING DISORDER, PLATELET-TYPE, 17	OMIM	MIM:187900 OMIM [®]
DBH <i>e!</i>	DOPAMINE BETA-HYDROXYLASE DEFICIENCY	OrphaNet	OrphaNet:230 orphanet

ADAMTSL2	e!	Geleophysic dysplasia	OrphaNet	OrphaNet:2623	orphanet
SURF1	e!	Fatal infantile cytochrome C oxidase deficiency	OrphaNet	OrphaNet:1561	orphanet
SURF1	e!	Leigh syndrome with cardiomyopathy	OrphaNet	OrphaNet:70474	orphanet
SURF1	e!	Leigh syndrome with leukodystrophy	OrphaNet	OrphaNet:255241	orphanet
SURF1	e!	SURF1-related Charcot-Marie-Tooth disease type 4	OrphaNet	OrphaNet:391351	orphanet
ADAMTSL3	e!	Congenital thrombotic thrombocytopenic purpura due to ADAMTSL3 deficiency	OrphaNet	OrphaNet:93583	orphanet
SARDH	e!	SARCOSINEMIA	OrphaNet	OrphaNet:3129	orphanet
GF1B	e!	GRAY PLATELET SYNDROME	OrphaNet	OrphaNet:721	orphanet

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)
ABO	missense variant	ENST00000538324	?	ENSP00000483018	3	3			3
ABO	missense variant	ENST00000611156	NM_020469.2	ENSP00000483265	3	3			3

Direct effect on regulation

cis-eQTL

gene	transcript	probe	tissue	min(statistic) (type)	source	variant(s)			
ABO	e!	?	ENSG00000175164	e!	stomach	1.87×10 ⁻¹¹ (p-value)	GTEx Portal V6	23	
ABO	e!	?	ENSG00000175164	e!	lung	1.34×10 ⁻²² (p-value)	GTEx Portal V6	24	
ABO	e!	?	ENSG00000175164	e!	tibial nerve	2.83×10 ⁻¹⁸ (p-value)	GTEx Portal V6	24	
ABO	e!	?	ENSG00000175164	e!	thyroid	2.25×10 ⁻³⁴ (p-value)	GTEx Portal V6	24	
ABO	e!	?	ENSG00000175164	e!	esophagus mucosa	4.41×10 ⁻²⁶ (p-value)	GTEx Portal V6	24	
ABO	e!	?	ENSG00000175164	e!	skeletal muscle	2.99×10 ⁻¹¹ (p-value)	GTEx Portal V6	24	
ABO	e!	?	ENSG00000175164	e!	transverse colon	2.25×10 ⁻⁸ (p-value)	GTEx Portal V6	22	
ABO	e!	?	ENSG00000175164	e!	unexposed skin	1.58×10 ⁻²⁶ (p-value)	GTEx Portal V6	24	
ABO	e!	?	ENSG00000175164	e!	sun exposed skin	4.86×10 ⁻²⁷ (p-value)	GTEx Portal V6	24	
OBP2B	e!	?	ENSG00000171102	e!	sun exposed skin	5.09×10 ⁻⁸ (p-value)	GTEx Portal V6	24	
ABO	e!	?	ENSG00000175164	e!	subcutaneous adipocytes	7.24×10 ⁻¹⁰ (p-value)	GTEx Portal V6	21	
ABO	e!	?	ENSG00000175164	e!	muscularis mucosae	1.92×10 ⁻⁷ (p-value)	GTEx Portal V6	19	
ABO	e!	?	ENSG00000175164	e!	visceral adipocytes	2.37×10 ⁻⁸ (p-value)	GTEx Portal V6	17	
ABO	e!	?	ENSG00000175164	e!	pancreas	7.12×10 ⁻⁷ (p-value)	GTEx Portal V6	15	
ABO	e!	?	ENSG00000175164	e!	testis	4.43×10 ⁻⁹ (p-value)	GTEx Portal V6	20	
ABO	e!	?	ENSG00000175164	e!	left ventricle	3.30×10 ⁻⁸ (p-value)	GTEx Portal V6	18	
MED22	e!	ENST00000614493	e!	ILMN_1697218	e!	monocyte	1.69×10 ⁻⁴ (p-value)	Fairfax et al.	2
MED22	e!	ENST00000610672	e!						
MED22	e!	ENST00000343730	e!						
MED22	e!	ENST00000610888	e!						

SLC2A6 <i>e!</i>	ENST00000371899 <i>e!</i>	ILMN_1778321 <i>e!</i>	blood	5.84×10 ⁻⁷ (p-value)	Westra et al. <i>l</i> <i>m</i>	6
SLC2A6 <i>e!</i>	ENST00000485978 <i>e!</i>					
SLC2A6 <i>e!</i>	ENST00000371897 <i>e!</i>					
ABO <i>e!</i>	?	ENSG00000175164 <i>e!</i>	adrenal gland	2.08×10 ⁻⁶ (p-value)	GTEx Portal V6 <i>l</i> <i>m</i>	2

Putative effect on regulation

FANTOM5 expressed promoter

SNiPA promoter id	variant(s)	associated transcript(s)	gene
FFCP00000859024 <i>e!</i>	1	ENST00000538324 <i>e!</i> , ENST00000453660 <i>e!</i>	ABO <i>e!</i>

ENCODE promoter-associated distal DHS (Enhancer)

SNiPA enhancer id	variant(s)	associated SNiPA promoter id	associated gene(s)
ENCE00000558431 <i>e!</i>	1	ENCP00000060477	GFI1B <i>e!</i>
		ENCP00000060525	ADAMTSL2 <i>e!</i>
		ENCP00000060529	DBH <i>e!</i>
		ENCP00000060490	RALGDS <i>e!</i>
		ENCP00000060486	CELP <i>e!</i>
		ENCP00000060489	RALGDS <i>e!</i>
		ENCP00000060518	ADAMTSL13 <i>e!</i> ADAMTSL13 <i>e!</i>
		ENCP00000060527	ADAMTSL2 <i>e!</i>
		ENCP00000060535	SARDH <i>e!</i>
		ENCP00000060507	SURF2 <i>e!</i> SURF2 <i>e!</i> SURF1 <i>e!</i> SURF1 <i>e!</i>
		ENCP00000060492	RALGDS <i>e!</i>
ENCE00000558621 <i>e!</i>	1	ENCP00000060495	OBP2B <i>e!</i> OBP2B <i>e!</i>
ENCE00000558373 <i>e!</i>	3	ENCP00000060477	GFI1B <i>e!</i>
		ENCP00000060471	AK8 <i>e!</i> C9orf9 <i>e!</i>
		ENCP00000060525	ADAMTSL2 <i>e!</i>
		ENCP00000060529	DBH <i>e!</i>
		ENCP00000060516	REXO4 <i>e!</i> REXO4 <i>e!</i>
		ENCP00000060509	SURF4 <i>e!</i> SURF4 <i>e!</i>
		ENCP00000060490	RALGDS <i>e!</i>
		ENCP00000060486	CELP <i>e!</i>
		ENCP00000060489	RALGDS <i>e!</i>
		ENCP00000060518	ADAMTSL13 <i>e!</i> ADAMTSL13 <i>e!</i>
		ENCP00000060527	ADAMTSL2 <i>e!</i>
		ENCP00000060535	SARDH <i>e!</i>
		ENCP00000060507	SURF2 <i>e!</i> SURF2 <i>e!</i> SURF1 <i>e!</i> SURF1 <i>e!</i>
		ENCP00000060492	RALGDS <i>e!</i>
ENCE00000558506 <i>e!</i>	1	ENCP00000060483	GTF3C5 <i>e!</i>

Regulatory feature cluster

element id	variant(s)	tissue/cell	factors
ENSR00001734089 <i>e!</i> (enhancer)	2	liver (HepG2)	H3K4me1, H3K4me2
		HSMMtube	H3K27me3
		lung (IMR90)	H3K27me3
		A549	H3K4me3, H3K4me2
		blood (K562)	H3K4me2, H2AZ
		skin (NHDF-AD)	H3K4me2
ENSR00000452813 <i>e!</i> (promoter flanking region)	2	cervix (HeLa-S3)	H3K9ac, H3K4me2, H3K27ac, H3K4me3, PolII, DNase1
		endothelium (HUVEC)	H3K27me3
		HSMMtube	H3K27me3
		lung (IMR90)	H3K27me3
		A549	H3K27me3
ENSR00001475637 <i>e!</i> (TF binding site)	1	embryonic stem cell (H1ESC)	H3K27me3
		endothelium (HUVEC)	H3K27me3
		lung (IMR90)	H3K27me3
		blood (K562)	H3K36me3, H3K4me2, H3K9ac, H3K27ac, DNase1, Brg1, Ini1, PolII
		A549	H3K27me3
ENSR00001475641 <i>e!</i> (open chromatin region)	1	embryonic stem cell (H1ESC)	H3K27me3
		lung (IMR90)	H3K27me3
		blood (K562)	H3K27me3

Variation proximal to gene

gene	variant type	min(distance)	transcript	RefSeq id	protein	variant(s)
ABO <i>e!</i>	downstream gene variant	2696	ENST00000453660 <i>e!</i>	?	?	1
ABO <i>e!</i>	downstream gene variant	248	ENST00000538324 <i>e!</i>	?	ENSP00000483018 <i>e!</i>	7
ABO <i>e!</i>	downstream gene variant	848	ENST00000611156 <i>e!</i>	NM_020469.2	ENSP00000483265 <i>e!</i>	6
LCN1P2 <i>e!</i>	upstream gene variant	103	ENST00000372030 <i>e!</i>	?	?	1
Y_RNA <i>e!</i>	downstream gene variant	4516	ENST00000364973 <i>e!</i>	?	?	1

Putative effect on transcript

Synonymous coding variant

gene	affected transcript	RefSeq id	protein	AA's	exchanged codons	variant(s)
ABO <i>e!</i>	ENST00000611156 <i>e!</i>	NM_020469.2	ENSP00000483265 <i>e!</i>	2	2	2
ABO <i>e!</i>	ENST00000538324 <i>e!</i>	?	ENSP00000483018 <i>e!</i>	2	2	2

Intron variant

gene	affected transcript	RefSeq id	protein	variant(s)
ABO <i>e!</i>	ENST00000453660 <i>e!</i>	?	?	7
ABO <i>e!</i>	ENST00000611156 <i>e!</i>	NM_020469.2	ENSP00000483265 <i>e!</i>	7
ABO <i>e!</i>	ENST00000538324 <i>e!</i>	?	ENSP00000483018 <i>e!</i>	7

3'-UTR variant

gene	affected transcript	RefSeq id	protein	variant(s)
ABO <i>e!</i>	ENST00000611156 <i>e!</i>	NM_020469.2	ENSP00000483265 <i>e!</i>	1

Non-coding exon variant

gene	affected transcript	RefSeq id	variant(s)
ABO <i>e!</i>	ENST00000453660 <i>e!</i>	?	12

