

SNiPACard

Block annotations

Block info





genomic range	chr4:74,331,266-74,483,608 <i>e!</i>
block size	152,343 bp
variant count	52 variants

Basic features

Conservation/deleteriousness		Linked genes	
phyloP	$\mu = -0.421$ [-3.218 – 1.469]	gene(s) hit or close-by	AFM <i>e!</i> , RASSF6 <i>e!</i> , RP11-622A1.1 <i>e!</i> , RP11-622A1.2 <i>e!</i>
phastCons	$\mu = 0.062$ [0 – 0.979]	eQTL gene(s)	RASSF6 <i>e!</i>
GERP++	$\mu = -0.510$ [-6.47 – 3.2]	potentially regulated gene(s)	AFM <i>e!</i> , AFP <i>e!</i>
CADD score	$\mu = 3.743$ [0.044 – 14.92]	disease gene(s)	AFP <i>e!</i>

Trait annotations

Disease gene annotation

gene	trait	source DB	source entry/link
AFP <i>e!</i>	ALPHA-FETOPROTEIN, HEREDITARY PERSISTENCE OF	OMIM	MIM:615970 
AFP <i>e!</i>	ALPHA-FETOPROTEIN DEFICIENCY	OMIM	MIM:615969 
AFP <i>e!</i>	Congenital deficiency in alpha-fetoprotein	OrphaNet	OrphaNet:168612 
AFP <i>e!</i>	Hereditary persistence of alpha-fetoprotein	OrphaNet	OrphaNet:168615 


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)
AFP <i>e!</i>	missense variant	ENST00000226355 <i>e!</i>	NM_001133.2	ENSP00000226355	H/R	cAt/cGt	?	?	1

Direct effect on regulation

cis-eQTL

gene	transcript	probe	tissue	min(statistic) (type)	source	variant(s)
RASSF6 <i>e!</i>	ENST00000395777 <i>e!</i>	ILMN_1657381 <i>e!</i>	b-cell	5.47×10 ⁻¹⁸ (p-value)	Fairfax et al. 	1
RASSF6 <i>e!</i>	ENST00000307439 <i>e!</i>					
RASSF6 <i>e!</i>	ENST00000512591 <i>e!</i>					

Putative effect on regulation

ENCODE promoter-associated DHS

SNiPA promoter id	variant(s)	associated gene(s)
ENCP00000041356 <i>e!</i>		

ENCODE promoter-associated distal DHS (Enhancer)

SNiPA enhancer id	variant(s)	associated SNiPA promoter id	associated gene(s)
ENCE00000382154 <i>e!</i>	1	ENCP00000041353	AFP <i>e!</i>

Regulatory feature cluster

element id	variant(s)	tissue/cell	factors
ENSR00001242774 <i>e!</i> (enhancer)	1	skin (NHEK)	H3K36me3
ENSR00001432007 <i>e!</i> (promoter flanking region)	6	embryonic stem cell (H1ESC) blood (DND-41) skin (NHDF-AD) muscle (HSMM) breast (HMEC) cervix (HeLa-S3) endothelium (HUVEC) liver (HepG2) lung (IMR90) blood (GM12878) nervous (NH-A) skin (NHEK)	DNase1 H3K4me1 DNase1 DNase1 H3K27ac, H3K4me1 DNase1 H3K36me3 DNase1 DNase1 Cjun, H3K4me3, PU1, PolII, BATF, DNase1, IRF4, BCL11A, H3K27ac, H3K4me2, H3K9ac DNase1 H3K4me1, H3K27ac, H3K4me2
ENSR00001685745 <i>e!</i> (CTCF binding site)	4	embryonic stem cell (H1ESC) blood (DND-41) skin (NHDF-AD) breast (HMEC) muscle (HSMM) liver (HepG2) lung (IMR90) blood (GM12878) skin (NHEK)	DNase1 H3K4me1 DNase1 H3K27ac, H3K4me1 DNase1 DNase1 DNase1 PU1, PolII, BATF, DNase1, IRF4, BCL11A, H3K27ac, H3K4me2, Cjun H3K4me1, H3K27ac, H3K4me2
ENSR00001685748 <i>e!</i> (enhancer)	1	A549 breast (HMEC)	H3K36me3 H3K36me3

Variation proximal to gene

gene	variant type	min(distance)	transcript	RefSeq id	protein	variant(s)
AFM <i>e!</i>	upstream gene variant, downstream gene variant	219	ENST00000505794 <i>e!</i>	? ?	?	5
AFM <i>e!</i>	upstream gene variant, downstream gene variant	184	ENST00000226355 <i>e!</i>	NM_001133.2	ENSP00000226355 <i>e!</i>	10
RP11-622A1.1 <i>e!</i>	upstream gene variant, downstream gene variant	3438	ENST00000511117 <i>e!</i>	? ?	?	3
RP11-622A1.2 <i>e!</i>	upstream gene variant, downstream gene variant	1361	ENST00000508147 <i>e!</i>	? ?	?	6
RP11-622A1.2 <i>e!</i>	upstream gene variant, downstream gene variant	1326	ENST00000504368 <i>e!</i>	? ?	?	6
RP11-622A1.2 <i>e!</i>	upstream gene variant, downstream gene variant	1562	ENST00000502291 <i>e!</i>	? ?	?	5

Putative effect on transcript

Intron variant

gene	affected transcript	RefSeq id	protein	variant(s)
AFM <i>e!</i>	ENST00000505794 <i>e!</i>	?	?	3
AFM <i>e!</i>	ENST00000226355 <i>e!</i>	NM_001133.2	ENSP00000226355 <i>e!</i>	10
RASSF6 <i>e!</i>	ENST00000307439 <i>e!</i>	NM_177532.4	ENSP00000303877 <i>e!</i>	15
RASSF6 <i>e!</i>	ENST00000342081 <i>e!</i>	NM_201431.2	ENSP00000340578 <i>e!</i>	15
RASSF6 <i>e!</i>	ENST00000335049 <i>e!</i>	NM_001270392.1	ENSP00000335582 <i>e!</i>	15
RASSF6 <i>e!</i>	ENST00000512591 <i>e!</i>	?	?	1
RASSF6 <i>e!</i>	ENST00000395777 <i>e!</i>	NM_001270391.1	ENSP00000379123 <i>e!</i>	15
RP11-622A1.2 <i>e!</i>	ENST00000508147 <i>e!</i>	?	?	8
RP11-622A1.2 <i>e!</i>	ENST00000502291 <i>e!</i>	?	?	8
RP11-622A1.2 <i>e!</i>	ENST00000504368 <i>e!</i>	?	?	9

Non-coding exon variant

gene	affected transcript	RefSeq id	variant(s)
AFM <i>e!</i>	ENST00000505794 <i>e!</i>	?	2
RP11-622A1.2 <i>e!</i>	ENST00000502291 <i>e!</i>	?	1

