

SNiPacard

Block annotations

Block info



genomic range	chr1:168,664,450-168,675,971 <i>e!</i>
block size	11,522 bp
variant count	10 variants

Basic features

Conservation/deleteriousness		Linked genes	
phyloP	$\mu = -0.033$ [-0.449 – 0.632]	gene(s) hit or close-by	DPT <i>e!</i>
phastCons	$\mu = 0.002$ [0 – 0.007]	eQTL gene(s)	DPT <i>e!</i>
GERP++	$\mu = -0.783$ [-2.66 – 1.57]	potentially regulated gene(s)	RP5-1018K9.1 <i>e!</i> , TBX19 <i>e!</i>
CADD score	$\mu = 2.879$ [1.17 – 5.142]	disease gene(s)	TBX19 <i>e!</i>

Trait annotations

Disease gene annotation

gene	trait	source DB	source entry/link
TBX19 <i>e!</i>	ACTH DEFICIENCY, ISOLATED	OMIM	MIM:201400 
TBX19 <i>e!</i>	Congenital isolated ACTH deficiency	OrphaNet	OrphaNet:199296 

Direct effect on regulation

cis-eQTL

gene	transcript	probe	tissue	min(statistic) (type)	source	variant(s)
DPT <i>e!</i>	ENST00000367817 <i>e!</i>	ILMN_1708107 <i>e!</i>	skin	1.29×10 ⁻⁶ (p-value)	MuTHER consortium 	2

Putative effect on regulation

ENCODE promoter-associated distal DHS (Enhancer)

SNiPA enhancer id	variant(s)	associated SNiPA promoter id	associated gene(s)
ENCE00000043270 <i>e!</i>	1	ENCP00000005412	TBX19 <i>e!</i>
		ENCP00000005426	RP5-1018K9.1 <i>e!</i>

Variation proximal to gene

gene	variant type	min(distance)	transcript	RefSeq id	protein	variant(s)
DPT <i>e!</i>	downstream gene variant	247	ENST00000367817 <i>e!</i>	NM_001937.4	ENSP00000356791 <i>e!</i>	1

Putative effect on transcript

Intron variant

gene	affected transcript	RefSeq id	protein	variant(s)
DPT <i>e!</i>	ENST00000367817 <i>e!</i>	NM_001937.4	ENSP00000356791 <i>e!</i>	9

