

SNiPACard

Block annotations

Block info

genomic range	chr1:196,696,875-196,696,875 <i>e!</i>
block size	1 bp
variant count	1 variants

Basic features

Conservation/deleteriousness		Linked genes	
phyloP	-1.535	gene(s) hit or close-by	CFH <i>e!</i>
phastCons	0	eQTL gene(s)	-
GERP++	-4.04	potentially regulated gene(s)	-
CADD score	0.742	disease gene(s)	CFH <i>e!</i>

Trait annotations

Variant association

trait	min(p-value)	source DB	source entry/link	variant(s)
Macular degeneration	2.33×10 ⁻⁹	dbGaP	pha002890 dbGaP	1
Macular degeneration	1.06×10 ⁻¹⁰	dbGaP	pha002869 dbGaP	1

Disease gene annotation

gene	trait	source DB	source entry/link
CFH <i>e!</i>	Immunodeficiency with factor H anomaly	OrphaNet	OrphaNet:200421 orphanet
CFH <i>e!</i>	Atypical hemolytic uremic syndrome with H factor anomaly	OrphaNet	OrphaNet:93579 orphanet
CFH <i>e!</i>	Dense deposit disease	OrphaNet	OrphaNet:93571 orphanet
CFH <i>e!</i>	Familial drusen	OrphaNet	OrphaNet:75376 orphanet
CFH <i>e!</i>	Immunoglobulin-mediated membranoproliferative glomerulonephritis	OrphaNet	OrphaNet:329903 orphanet

Putative effect on transcript

Intron variant

gene	affected transcript	RefSeq id	protein	variant(s)
CFH <i>e!</i>	ENST00000466229 <i>e!</i>	?	?	1
CFH <i>e!</i>	ENST00000367429 <i>e!</i>	NM_000186.3	ENSP00000356399 <i>e!</i>	1

