

Block annotations

Block info

genomic range	chr5:149,519,260-149,537,194 <i>e!</i>
block size	17,935 bp
variant count	3 variants

Basic features

Conservation/deleteriousness		Linked genes	
phyloP	$\mu = -0.217$ [-0.624 – 0.211]	gene(s) hit or close-by	PDGFRB <i>e!</i>
phastCons	$\mu = 0.001$ [0 – 0.002]	eQTL gene(s)	CDX1 <i>e!</i> , CSF1R <i>e!</i> , PDGFRB <i>e!</i>
GERP++	$\mu = 0.540$ [-0.121 – 1.03]	potentially regulated gene(s)	-
CADD score	$\mu = 6.404$ [4.963 – 8.056]	disease gene(s)	PDGFRB <i>e!</i> , CSF1R <i>e!</i>

Trait annotations

Disease gene annotation

gene	trait	source DB	source entry/link
PDGFRB <i>e!</i>	Familial Infantile Myofibromatosis	DECIPHER	MIM:228550 <i>OMIM</i>
PDGFRB <i>e!</i>	JUVENILE MYELOMONOCYTIC LEUKEMIA	OMIM	MIM:607785 <i>OMIM</i>
PDGFRB <i>e!</i>	BASAL GANGLIA CALCIFICATION, IDIOPATHIC, 4	OMIM	MIM:615007 <i>OMIM</i>
PDGFRB <i>e!</i>	MYOFIBROMATOSIS, INFANTILE, 1	OMIM	MIM:228550 <i>OMIM</i>
PDGFRB <i>e!</i>	LEUKEMIA, ACUTE MYELOID	OMIM	MIM:601626 <i>OMIM</i>
PDGFRB <i>e!</i>	MYELOPROLIFERATIVE DISORDER, CHRONIC, WITH EOSINOPHILIA	OMIM	MIM:131440 <i>OMIM</i>
CSF1R <i>e!</i>	LEUKOENCEPHALOPATHY, DIFFUSE HEREDITARY, WITH SPHEROIDS	OMIM	MIM:221820 <i>OMIM</i>
PDGFRB <i>e!</i>	Bilateral striopallidodentate calcinosis	OrphaNet	OrphaNet:1980 <i>orphanet</i>
PDGFRB <i>e!</i>	Unclassified chronic myeloproliferative disease	OrphaNet	OrphaNet:86830 <i>orphanet</i>
PDGFRB <i>e!</i>	Chronic myelomonocytic leukemia	OrphaNet	OrphaNet:98823 <i>orphanet</i>
PDGFRB <i>e!</i>	Idiopathic hypereosinophilic syndrome	OrphaNet	OrphaNet:3260 <i>orphanet</i>
PDGFRB <i>e!</i>	Infantile myofibromatosis 1	OrphaNet	OrphaNet:2591 <i>orphanet</i>
PDGFRB <i>e!</i>	Myeloid neoplasm associated with PDGFRB rearrangement	OrphaNet	OrphaNet:168950 <i>orphanet</i>
CSF1R <i>e!</i>	Adult-onset leukoencephalopathy with axonal spheroids and pigmented glia	OrphaNet	OrphaNet:313808 <i>orphanet</i>

Direct effect on regulation

cis-eQTL

gene	transcript	probe	tissue	min(statistic) (type)	source	variant(s)
PDGFRB <i>e!</i>	ENST00000520579 <i>e!</i>	ILMN_1815057 <i>e!</i>	blood	8.08×10^{-39} (p-value)	MuTHER consortium <i>IM</i>	1
PDGFRB <i>e!</i>	ENST00000261799 <i>e!</i>					
CSF1R <i>e!</i>	ENST00000509861 <i>e!</i>	ILMN_1686623 <i>e!</i>	blood	5.51×10^{-7} (p-value)	Westra et al. <i>IM</i>	1
CSF1R <i>e!</i>	ENST00000504875 <i>e!</i>					
CSF1R <i>e!</i>	ENST00000286301 <i>e!</i>					
CDX1 <i>e!</i>	?	ENSG00000113722 <i>e!</i>	tibial nerve	5.45×10^{-13} (p-value)	GTEx Portal V6 <i>IM</i>	2
PDGFRB <i>e!</i>	?	ENSG00000113721 <i>e!</i>	EBV lymphocytes	6.13×10^{-8} (p-value)	GTEx Portal V6 <i>IM</i>	2
PDGFRB <i>e!</i>	ENST00000520579 <i>e!</i>	ILMN_1815057 <i>e!</i>	b-cell	5.54×10^{-6} (p-value)	Fairfax et al. <i>IM</i>	1
PDGFRB <i>e!</i>	ENST00000261799 <i>e!</i>					

Putative effect on regulation

Regulatory feature cluster

element id	variant(s)	tissue/cell	factors		
ENSR00001293753 <i>e!</i> (promoter flanking region)	1	embryonic stem cell (H1ESC)	DNase1, H3K4me1, USF1		
		HSMMtube	H3K4me1, H3K9ac, H3K27ac, H3K36me3, DNase1		
		blood (K562)	H3K27me3, DNase1, Egr1, PU1, Cjun		
		skin (NHDF-AD)	H3K27ac, H3K4me2, H3K9ac, DNase1, H3K4me3, H3K4me1, H3K36me3		
		muscle (HSMM)	H3K27ac, H3K9ac, H3K36me3, DNase1		
		liver (HepG2)	FOSL2, Jund, H3K4me3, PolII, TAF1, ELF1, USF1, H3K4me1, p300, H3K4me2, FOXA1, H3K9ac, H3K27ac, CTCF, DNase1		
		lung (IMR90)	H3K56ac, H3K79me2, H3K18ac, H3K27ac, H3K4me2, H4K5ac, H3K36me3, H3K4ac, H2BK120ac, H2BK20ac, H2AK5ac, H3K9ac, H3K4me1, H2BK12ac, DNase1, H4K8ac, H4K91ac		
		nervous (NH-A)	DNase1, H3K9ac, H3K4me1, H3K4me2, H3K36me3, H3K27ac		
		skin (NHEK)	H3K27me3, DNase1		
		NHLF	H3K27ac, H3K36me3, H3K9ac, H3K4me1, DNase1		
		Osteobl	H3K36me3, H3K4me1, H3K4me2, H3K27ac		
		blood (DND-41)	H3K27me3		
		breast (HMEC)	H3K27ac, H3K4me2		
		cervix (HeLa-S3)	DNase1, Cjun, Jund, H3K4me2, H3K27ac, H3K4me1, Max		
		monocytes (Monocytes-CD14+)	H3K27ac, DNase1		
		endothelium (HUVEC)	H3K27me3		
		A549	H3K4me3, H3K4me2, DNase1, H3K27ac		
		ENSR00001698907 <i>e!</i> (promoter flanking region)	1	embryonic stem cell (H1ESC)	DNase1, H3K27me3, Rad21, CTCF
				HSMMtube	DNase1, CTCF
				blood (K562)	H3K27me3, PolII, Egr1, Rad21, H2AZ, H3K4me2, CTCF, DNase1
skin (NHDF-AD)	H3K27ac, H3K4me2, H3K9ac, DNase1, CTCF, H3K4me1				
muscle (HSMM)	H2AZ, CTCF				
liver (HepG2)	Rad21, H2AZ, H3K4me2, CTCF, H3K27me3, DNase1, H3K4me3				
blood (GM12878)	DNase1, EBF1, H2AZ, Rad21, CTCF				
lung (IMR90)	DNase1, CTCF				
nervous (NH-A)	DNase1, CTCF				
skin (NHEK)	H3K27me3, CTCF				
NHLF	DNase1, H3K4me1, CTCF, H3K27ac				
Osteobl	CTCF, H3K27ac				
blood (DND-41)	CTCF, H3K27ac, H3K27me3, H3K4me1				
breast (HMEC)	CTCF				
cervix (HeLa-S3)	H3K9ac, H3K4me2, H3K4me1, CTCF, DNase1				
monocytes (Monocytes-CD14+)	H3K27me3, CTCF				
endothelium (HUVEC)	H3K36me3, DNase1, H3K27me3, CTCF				
A549	H3K4me3, H3K4me2, CTCF, DNase1				

Variation proximal to gene

gene	variant type	min(distance)	transcript	RefSeq id	protein	variant(s)
PDGFRB <i>e!</i>	upstream gene variant	2260	ENST00000517957 <i>e!</i>	?	ENSP00000430715 <i>e!</i>	1
PDGFRB <i>e!</i>	upstream gene variant	1771	ENST00000517660 <i>e!</i>	?	?	1
PDGFRB <i>e!</i>	downstream gene variant, upstream gene variant	867	ENST00000523456 <i>e!</i>	?	?	2
PDGFRB <i>e!</i>	upstream gene variant	1772	ENST00000520579 <i>e!</i>	?	ENSP00000430026 <i>e!</i>	1
PDGFRB <i>e!</i>	upstream gene variant	2004	ENST00000517488 <i>e!</i>	?	ENSP00000429218 <i>e!</i>	1
PDGFRB <i>e!</i>	upstream gene variant	1771	ENST00000261799 <i>e!</i>	NM_002609.3	ENSP00000261799 <i>e!</i>	1

Putative effect on transcript

Intron variant

gene	affected transcript	RefSeq id	protein	variant(s)
PDGFRB <i>e!</i>	ENST00000517660 <i>e!</i>	?	?	2
PDGFRB <i>e!</i>	ENST00000517488 <i>e!</i>	?	ENSP00000429218 <i>e!</i>	2

PDGFRB <i>e!</i>	ENST00000520579 <i>e!</i>	?	ENSP00000430026 <i>e!</i>	2
PDGFRB <i>e!</i>	ENST00000261799 <i>e!</i>	NM_002609.3	ENSP00000261799 <i>e!</i>	2
