

TBLR1 Rabbit mAb

货号: B28662

产品信息

反应	Human,Mouse,Rat
宿主	Rabbit
克隆性	Monoclonal
预测反应	
应用	WB IHC IF/ICC
推荐浓度	WB: 1:500 - 1:2000 IHC: 1:50 - 1:200 IF/ICC: 1:50 - 1:200
理论分子量	55kDa
实测分子量	56kDa
形式	Liquid
保存条件	Store at -20°C. Avoid freeze / thaw cycles. Buffer: PBS with 0.75% BSA,50% glycerol,pH7.3.
偶联物	Unconjugated
阳性对照	Jurkat,HepG2,NCI-H460,THP-1,Mouse brain,Mouse spleen
细胞定位	Nucleus
纯化	Affinity purification

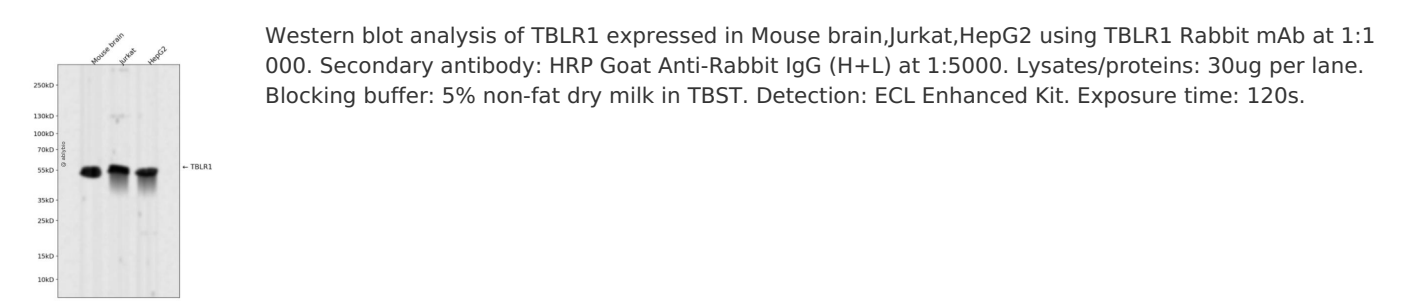
抗原信息

抗原信息	Recombinant fusion protein corresponding to Human TBLR1.
序列	MSISSDEVNFLVYRYLQESGFSHSAFTFGIESHISQSNINGALVPPAALISIIKGLQYVEAEVSINEDGTLFDGRPIESLSLID AVMPDVVQTRQQAYRDKLAQQQAAAAAAAAAAAAASQQGSAKNGENTANGEENGAHTIANNHTDMMMEVDGDVEIPPNKA VVLRGHESEVFICAWNPV

靶点信息

研究背景	This gene is a member of the WD40 repeat-containing gene family and shares sequence similarity with transducin (beta)-like 1X-linked (TBL1X). The protein encoded by this gene is thought to be a component of both nuclear receptor corepressor (N-CoR) and histone deacetylase 3 (HDAC 3) complexes, and is required for transcriptional activation by a variety of transcription factors. Mutations in these gene have been associated with some autism spectrum disorders, and one finding suggests that haploinsufficiency of this gene may be a cause of intellectual disability with dysmorphism. Mutations in this gene as well as recurrent translocations involving this gene have also been observed in some tumors.
基因ID	79718
基因名	TBL1XR1
Swiss	Q9BZK7
别名	TBL1XR1;C21;DC42;IRA1;MRD41;TBLR1

产品验证



实验步骤

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