

MaxPab®

SLC46A1 purified MaxPab mouse polyclonal antibody (B01P)

Catalog # H00113235-B01P Size 500 ug

Specification

Product Description	Mouse polyclonal antibody raised against a full-length human SLC46A1 protein.
Immunogen	SLC46A1 (ADZ16056.1, 1 a.a. ~ 431 a.a) full-length human protein.
Sequence	MEGSASPPEKPRARPAAAVLCRGPVEPLVFLANFALVLQGPLTTQYLWHRFSADLGYNGTRQRG GCSNRSADPTMQEVETLTSHWTL YMNVGGLVGLFSSLLGAWSDSVGRRPLLVLASLGLLLQA LVSVFVVQLQLHVG YFVLGRILCALLGDFGGLLAASFASVADVSSSRSTFRMALLEASIGVAGM LASLLGGHWLRAQGYANPFWLALALLIAMTL YAAFCFGETLKEPKSTRLF TFRHHR SIVQLYVAPA PEKSRKHLAL YSLAIFVVITVHFGAQDIL TYELSTPLCWDSKLIGYGSA AQHLPYL TSL LALKLLQYC LADAWVAEIGLAFN LGMVVFATITPLMFTGALFSAVACVNSLAML TASGIFNSLYPATL NFMKG FPFLLGAGLLLIPAVLIGMLEKADPHLEFQQFPQSP
Host	Mouse
Reactivity	Human
Interspecies Antigen Sequence	Mouse (81); Rat (81)
Quality Control Testing	Antibody reactive against mammalian transfected lysate.
Storage Buffer	In 1x PBS, pH 7.4
Storage Instruction	Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing.

Applications

- Western Blot (Transfected lysate)

[Protocol Download](#)

Gene Info — SLC46A1

Entrez GeneID	113235
GeneBank Accession#	JF432839.1
Protein Accession#	ADZ16056.1
Gene Name	SLC46A1
Gene Alias	HCP1, MGC9564, PCFT
Gene Description	solute carrier family 46 (folate transporter), member 1
Omim ID	611672
Gene Ontology	Hyperlink
Gene Summary	This gene encodes a transmembrane proton-coupled folate transporter protein that facilitates the movement of folate and antifolate substrates across cell membranes optimally in acidic pH environments. This protein is also expressed in the brain and choroid plexus where it transports folates into the central nervous system. This protein further functions as a transmembrane heme transporter in duodenal enterocytes and, potentially, in other tissues like liver and kidney. Its localization to the apical membrane or cytoplasm of intestinal cells is modulated by dietary iron levels. Mutations in this gene cause the autosomal recessive hereditary folate malabsorption (HFM) disease. HFM is characterized by folate deficiency due to reduced intestinal folate absorption and subsequent anemia, hypimmunoglobulinemia, and recurrent infections. [provided by RefSeq]
Other Designations	heme carrier protein 1 proton-coupled folate transporter solute carrier family 46, member 1

Disease

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