

Beta 氨基己糖苷酶 beta 亚基蛋白 B 链 抗体

产品货号: mIR15468

英文名称: HEXB chain B

中文名称: Beta 氨基己糖苷酶 beta 亚基蛋白 B 链抗体

别 名: Beta hexosaminidase beta chain; Beta hexosaminidase subunit beta; Beta N acetylhexosaminidase; Beta-hexosaminidase subunit beta chain B; Beta-N-acetylhexosaminidase subunit beta; Cervical cancer proto oncogene 7 protein; Cervical cancer proto-oncogene 7 protein; ENC 1AS; HCC 7; HCC-7; HCC-7; HCC7; HEX B; Hexb; HEXB_HUMAN; Hexosaminidase B (beta polypeptide); Hexosaminidase B; Hexosaminidase subunit B; HexosaminidaseB; N acetyl beta glucosaminidase; N-acetyl-beta-glucosaminidase subunit beta; HEXB chain B.

研究领域: 细胞生物 免疫学 信号转导 新陈代谢

抗体来源: Rabbit

克隆类型: Polyclonal

交叉反应: Human, Mouse, Rat, Dog, Cow, Sheep,

产品应用: WB=1:500-2000 ELISA=1:500-1000 IHC-P=1:400-800 IHC-F=1:400-800 ICC=1:100-500 IF=1:100-500

(石蜡切片需做抗原修复)

not yet tested in other applications.

optimal dilutions/concentrations should be determined by the end user.

分子量: 22/50kDa

细胞定位: 细胞浆

性 状: Lyophilized or Liquid



浓 度: 1mg/ml

免疫原: KLH conjugated synthetic peptide derived from human HEXB:201-300/556

亚 型: IgG

纯化方法: affinity purified by Protein A

储存液: 0.01M TBS(pH7.4) with 1% BSA, 0.03% Proclin300 and 50% Glycerol.

保存条件: Store at -20 °C for one year. Avoid repeated freeze/thaw cycles. The lyophilized antibody is stable at room temperature for at least one month and for greater than a year when kept at -20 °C. When reconstituted in sterile pH 7.4 0.01M PBS or diluent of antibody the antibody is stable for at least two weeks at 2-4 °C.

PubMed: PubMed

产品介绍: Hexosaminidase B is the beta subunit of the lysosomal enzyme beta-hexosaminidase that, together with the cofactor GM2 activator protein, catalyzes the degradation of the ganglioside GM2, and other molecules containing terminal N-acetyl hexosamines. Beta-hexosaminidase is composed of two subunits, alpha and beta, which are encoded by separate genes. Both beta-hexosaminidase alpha and beta subunits are members of family 20 of glycosyl hydrolases. Mutations in the alpha or beta subunit genes lead to an accumulation of GM2 ganglioside in neurons and neurodegenerative disorders termed the GM2 gangliosidoses. Beta subunit gene mutations lead to Sandhoff disease (GM2-gangliosidosis type II). [provided by RefSeq, Jul 2008].

Function:

Responsible for the degradation of GM2 gangliosides, and a variety of other molecules containing terminal Nacetyl hexosamines, in the brain and other tissues.

Subunit:

There are 3 forms of beta-hexosaminidase: hexosaminidase A is a trimer composed of one subunit alpha, one subunit beta chain A and one subunit beta chain B; hexosaminidase B is a tetramer of two subunit beta chains A



and two subunit beta chains B; hexosaminidase S is a homodimer of two alpha subunits. The two beta chains are derived from the cleavage of the beta subunit.

Subcellular Location:
Lysosome.
DISEASE:
GM2-gangliosidosis 2 (GM2G2) [MIM:268800]: An autosomal recessive lysosomal storage disease marked by the
accumulation of GM2 gangliosides in the neuronal cells. Clinically indistinguishable from GM2-gangliosidosis type
1, presenting startle reactions, early blindness, progressive motor and mental deterioration, macrocephaly and
cherry-red spots on the macula. Note=The disease is caused by mutations affecting the gene represented in this
entry.
Similarity:
Belongs to the glycosyl hydrolase 20 family.
SWISS:
P07686
Gene ID:
3074
Important Note:

This product as supplied is intended for research use only, not for use in human, therapeutic or diagnostic applications.



产品图片

