PSAP Antibody

Catalog No: #32449

Description

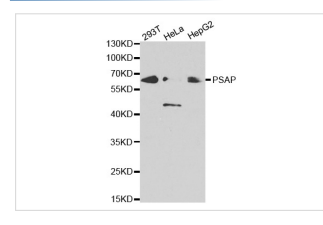


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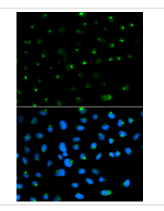
Product Name **PSAP** Antibody Rabbit Host Species Clonality Polyclonal Purification Antibodies were purified by affinity purification using immunogen. WB IHC IF Applications Species Reactivity Hu Ms Rt Specificity The antibody detects endogenous level of total PSAP protein. **Recombinant Protein** Immunogen Type Immunogen Description Recombinant protein of human PSAP. Target Name PSAP Other Names FLJ00245; GLBA; MGC110993; SAP1; Accession No. Swiss-Prot:P07602NCBI Gene ID:5660 SDS-PAGE MW 58KD Concentration 1.0mg/ml Formulation Supplied at 1.0mg/mL in phosphate buffered saline (without Mg2+ and Ca2+), pH 7.4, 150mM NaCl, 0.02% sodium azide and 50% glycerol. Storage Store at -20°C

Application Details Western blotting: 1:500 - 1:2000 Immunohistochemistry: 1:50 - 1:100 Immunofluorescence: 1:50 - 1:200

Images



Western blot analysis of extracts of various cell lines, using PSAP antibody.



Immunofluorescence analysis of MCF7 cell using PSAP antibody. Blue: DAPI for nuclear staining.

Background

The PSAP gene encodes prosaposin, a precursor of four small nonenzymatic glycoproteins termed 'sphingolipid activator proteins' (SAPs) that assist in the lysosomal hydrolysis of sphingolipids. After proteolytic processing of the presaposin protein, these 4 released polypeptides are functional activators. Saposin A is encoded by residues 60 to 143 of PSAP, saposin B by 195 to 275, saposin C by 311 to 390, and saposin D by 405 to 487. They are four 12-14 kDa heatstable glycoproteins. Saposins A-D localize primarily to the lysosomal compartment where they facilitate the catabolism of glycosphingolipids with short oligosaccharide groups. Saposins A-D are required for the hydrolysis of certain sphingolipids by specific lysosomal hydrolases. (PMID: 2001789) Defects in PSAP are the cause of Gaucher disease, Tay-Sachs disease, and metachromatic leukodystrophy (PubMed: 2060627, PMID: 15773042). This PSAP antibody (10801-1-AP) is expected to recognize both saposin A and B.

Note: This product is for in vitro research use only and is not intended for use in humans or animals.