abcam

Product datasheet

Anti-Acid sphingomyelinase antibody [mAbcam74281] ab74281

★★★★★ 1 Abreviews 5 References 画像数 2

製品名	Anti-Acid sphingomyelinase antibody [mAbcam74281]	
製品の詳細	Mouse monoclonal [mAbcam74281] to Acid sphingomyelinase	
由来種	Mouse	
アプリケーション	適用あり: WB, Flow Cyt (Intra)	
種交差性	交差種: Human	
免疫原	Synthetic peptide corresponding to Human Acid sphingomyelinase aa 1-100 conjugated to keyhole limpet haemocyanin.	
ポジティブ・コントロール	This antibody gave a positive signal in the following whole cell lysates: HepG2; A431; MCF7; HeLa; THP1.	
特記事項	We can conjugate this antibody to FITC for you (please see <u>ab150251</u> for details).	
	This antibody clone is manufactured by Abcam. If you require a custom buffer formulation or conjugation for your experiments, please contact orders@abcam.com .	
	The Life Science industry has been in the grips of a reproducibility crisis for a number of years. Abcam is leading the way in addressing this with our range of recombinant monoclonal antibodies and knockout edited cell lines for gold-standard validation. Please check that this product meets your needs before purchasing.	
	If you have any questions, special requirements or concerns, please send us an inquiry and/or contact our Support team ahead of purchase. Recommended alternatives for this product can be found below, along with publications, customer reviews and Q&As	
製品の特性		

製品の状態	Liquid
保存方法	Shipped at 4°C. Store at +4°C short term (1-2 weeks). Upon delivery aliquot. Store at -20°C or - 80°C. Avoid freeze / thaw cycle.
バッファー	pH: 7.40 Preservative: 0.02% Sodium azide Constituent: PBS
精製度	Protein G purified
ポリモノ	モノクローナル

クローン名	mAbcam74281
₹ID-7	Sp2/0-Ag14
アイソタイプ	lgG2a
軽鎖の種類	kappa

アプリケーション

The Abpromise guarantee Abpromise保証は、次のテスト済みアプリケーションにおけるab74281の使用に適用されます

アプリケーションノートには、推奨の開始希釈率がありますが、適切な希釈率につきましてはご検討ください。

アプリケーション	Abreviews	特記事項
WB	★ ★ ★ ★ ★ <u>(1)</u>	Use a concentration of 5 μ g/ml. Detects a band of approximately 65 kDa (predicted molecular weight: 70 kDa).
Flow Cyt (Intra)		Use $1\mu g$ for 10^6 cells. <u>ab170191</u> - Mouse monoclonal lgG2a, is suitable for use as an isotype control with this antibody.

ターゲット情報

機能

関連疾患

Converts sphingomyelin to ceramide. Also has phospholipase C activities toward 1,2diacylglycerolphosphocholine and 1,2-diacylglycerolphosphoglycerol. Isoform 2 and isoform 3 have lost catalytic activity.

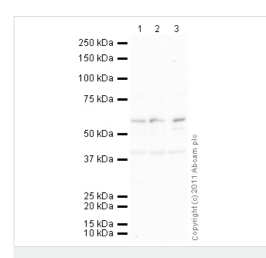
Defects in SMPD1 are the cause of Niemann-Pick disease type A (NPDA) [MIM:257200]; also known as Niemann-Pick disease classical infantile form. It is an early-onset lysosomal storage disorder caused by failure to hydrolyze sphingomyelin to ceramide. It results in the accumulation of sphingomyelin and other metabolically related lipids in reticuloendothelial and other cell types throughout the body, leading to cell death. Niemann-Pick disease type A is a primarily neurodegenerative disorder characterized by onset within the first year of life, mental retardation, digestive disorders, failure to thrive, major hepatosplenomegaly, and severe neurologic symptoms. The severe neurological disorders and pulmonary infections lead to an early death, often around the age of four. Clinical features are variable. A phenotypic continuum exists between type A (basic neurovisceral) and type B (purely visceral) forms of Niemann-Pick disease, and the intermediate types encompass a cluster of variants combining clinical features of both types A and B.

Defects in SMPD1 are the cause of Niemann-Pick disease type B (NPDB) [MIM:607616]; also known as Niemann-Pick disease visceral form. It is a late-onset lysosomal storage disorder caused by failure to hydrolyze sphingomyelin to ceramide. It results in the accumulation of sphingomyelin and other metabolically related lipids in reticuloendothelial and other cell types throughout the body, leading to cell death. Clinical signs involve only visceral organs. The most constant sign is hepatosplenomegaly which can be associated with pulmonary symptoms. Patients remain free of neurologic manifestations. However, a phenotypic continuum exists between type A (basic neurovisceral) and type B (purely visceral) forms of Niemann-Pick disease, and the intermediate types encompass a cluster of variants combining clinical features of both types A and B. In Niemann-Pick disease type B, onset of the first symptoms occurs in early childhood and patients can survive into adulthood.

Belongs to the acid sphingomyelinase family.

細胞内局在

画像



Western blot - Anti-Acid sphingomyelinase antibody [mAbcam74281] (ab74281) **All lanes :** Anti-Acid sphingomyelinase antibody [mAbcam74281] (ab74281) at 5 µg/ml

Lane 1 : HepG2 (Human hepatocellular liver carcinoma cell line) Whole Cell Lysate

Lane 2 : A431 (Human epithelial carcinoma cell line) Whole Cell Lysate

Lane 3 : MCF7 (Human breast adenocarcinoma cell line) Whole Cell Lysate

Lysates/proteins at 10 µg per lane.

Secondary

All lanes : Goat Anti-Mouse IgG H&L (HRP) preadsorbed (<u>ab97040</u>) at 1/5000 dilution

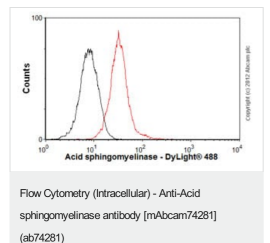
Developed using the ECL technique.

Performed under reducing conditions.

Predicted band size: 70 kDa Observed band size: 65 kDa

Exposure time: 1 minute

The predicted molecular weight of Acid Sphingomyelinase protein is 70 kDa. However, the protein sequence contains a 46-residue signal sequence at the amino-terminal, which could explain the band observed.



Overlay histogram showing HeLa cells stained with ab74281 (red line). The cells were fixed with 80% methanol (5 min) and then permeabilized with 0.1% PBS-Tween for 20 min. The cells were then incubated in 1x PBS / 10% normal goat serum / 0.3M glycine to block non-specific protein-protein interactions followed by the antibody (ab74281, 1µg/1x10⁶ cells) for 30 min at 22°C. The secondary antibody used was DyLight® 488 goat anti-mouse IgG (H+L) (**ab96879**) at 1/500 dilution for 30 min at 22°C. Isotype control antibody (black line) was mouse IgG2a [ICIGG2A] (**ab91361**, 1µg/1x10⁶ cells) used under the same conditions. Acquisition of >5,000 events was performed.

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