

OriGene Technologies, Inc.

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Product datasheet for TA803208AM

ABCD1 Mouse Monoclonal Antibody (Biotin conjugated) [Clone ID: OTI4C2]

Product data:

Product Type:	Primary Antibodies
Clone Name:	OTI4C2
Applications:	IHC, WB
Recommended Dilution:	WB 1:2000, IHC 1:150
Reactivity:	Human, Mouse, Rat
Host:	Mouse
lsotype:	lgG2a
Clonality:	Monoclonal
Immunogen:	Human recombinant protein fragment corresponding to amino acids 508-745 of human ABCD1 (NP_000024) produced in E.coli.
Formulation:	PBS (pH 7.3) containing 1% BSA, 50% glycerol and 0.02% sodium azide.
Concentration:	0.5 mg/ml
Purification:	Purified from mouse ascites fluids or tissue culture supernatant by affinity chromatography (protein A/G)
Conjugation:	Biotin
Storage:	Store at -20°C as received.
Stability:	Stable for 12 months from date of receipt.
Predicted Protein Size:	82.8 kDa
Gene Name:	ATP binding cassette subfamily D member 1
Database Link:	<u>NP_000024</u> <u>Entrez Gene 215 Human</u> <u>P33897</u>



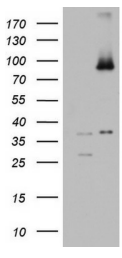
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Sourigene ABCD1 Mouse Monoclonal Antibody (Biotin conjugated) [Clone ID: OTI4C2] – TA803208AM

Background:The protein encoded by this gene is a member of the superfamily of ATP-binding cassette
(ABC) transporters. ABC proteins transport various molecules across extra- and intra-cellular
membranes. ABC genes are divided into seven distinct subfamilies (ABC1, MDR/TAP, MRP,
ALD, OABP, GCN20, White). This protein is a member of the ALD subfamily, which is involved
in peroxisomal import of fatty acids and/or fatty acyl-CoAs in the organelle. All known
peroxisomal ABC transporters are half transporters which require a partner half transporter
molecule to form a functional homodimeric or heterodimeric transporter. This peroxisomal
membrane protein is likely involved in the peroxisomal transport or catabolism of very long
chain fatty acids. Defects in this gene have been identified as the underlying cause of
adrenoleukodystrophy, an X-chromosome recessively inherited demyelinating disorder of the
nervous system. [provided by RefSeq, Jul 2008]

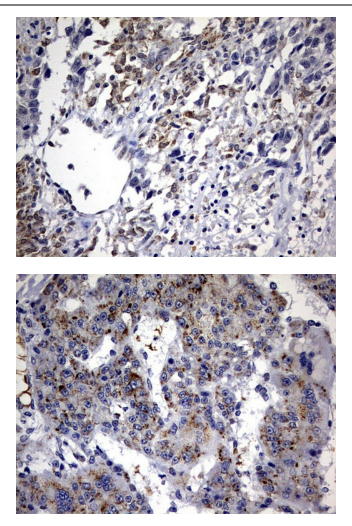
Synonyms:	ABC42; ALD; ALDP; AMN
Protein Families:	Druggable Genome
Protein Pathways:	ABC transporters

Product images:



HEK293T cells were transfected with the pCMV6-ENTRY control (Cat# [PS100001], Left lane) or pCMV6-ENTRY ABCD1 (Cat# [RC206885], Right lane) cDNA for 48 hrs and lysed. Equivalent amounts of cell lysates (5 ug per lane) were separated by SDS-PAGE and immunoblotted with anti-ABCD1(Cat# [TA803208]). Positive lysates [LY424967] (100ug) and [LC424967] (20ug) can be purchased separately from OriGene.

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Immunohistochemical staining of paraffinembedded Adenocarcinoma of Human ovary tissue using anti-ABCD1 mouse monoclonal antibody. (Heat-induced epitope retrieval by 10mM citric buffer, pH6.0, 120°C for 3min, [TA803208])

Immunohistochemical staining of paraffinembedded Carcinoma of Human liver tissue using anti-ABCD1 mouse monoclonal antibody. (Heat-induced epitope retrieval by 10mM citric buffer, pH6.0, 120°C for 3min, [TA803208])

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