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Product datasheet for CF804944

Lipin 3 (LPIN3) Mouse Monoclonal Antibody [Clone ID: OTI1D7]

Product data:

Product Type:	Primary Antibodies
Clone Name:	OTI1D7
Applications:	IHC, WB
Recommended Dilution:	WB 1:500, IHC 1:150
Reactivity:	Human
Host:	Mouse
lsotype:	lgG1
Clonality:	Monoclonal
Immunogen:	Full length human recombinant protein of human LPIN3 (NP_075047) produced in HEK293T cell.
Formulation:	Lyophilized powder (original buffer 1X PBS, pH 7.3, 8% trehalose)
Reconstitution Method:	For reconstitution, we recommend adding 100uL distilled water to a final antibody concentration of about 1 mg/mL. To use this carrier-free antibody for conjugation experiment, we strongly recommend performing another round of desalting process. (OriGene recommends Zeba Spin Desalting Columns, 7KMWCO from Thermo Scientific)
Purification:	Purified from mouse ascites fluids or tissue culture supernatant by affinity chromatography (protein A/G)
Conjugation:	Unconjugated
Storage:	Store at -20°C as received.
Stability:	Stable for 12 months from date of receipt.
Predicted Protein Size:	93.4 kDa
Gene Name:	lipin 3
Database Link:	<u>NP_075047</u> <u>Entrez Gene 64900 Human</u> <u>Q9BQK8</u>



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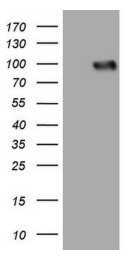
Serigene Lipin 3 (LPIN3) Mouse Monoclonal Antibody [Clone ID: OTI1D7] – CF804944

Background:Humans lipodystrophy is characterized by loss of body fat, fatty liver, hypertriglyceridemia,
and insulin resistance. Mice carrying mutations in the fatty liver dystrophy (fld) gene have
similar phenotypes. Through positional cloning, the mouse gene responsible for fatty liver
dystrophy was isolated and designated Lpin1. The nuclear protein encoded by Lpin1 was
named lipin. Lpin1 mRNA was expressed at high levels in adipose tissue and was induced
during differentiation of preadipocytes. These results indicated that lipin is required for
normal adipose tissue development and provided a candidate gene for human lipodystrophy.
Through database searches, mouse and human EST and genomic sequences with similarities
to Lpin1 were identified. These included two related mouse genes (Lpin2 and Lpin3) and
three human homologs (LPIN1, LPIN2, and LPIN3). Human LPIN1 gene has been mapped to
2p25.; linkages of fat mass and serum leptin levels to this same region have been noted.
Human LPIN2 and LPIN3 mapped to chromosomes 18p11 and 20q11-q12, respectively. The
mouse genes encoding Lpin1, Lpin2, and Lpin3 mapped to chromosome 12, 17, and 2,
respectively. [provided by RefSeq, Jul 2008]

Synonyms:

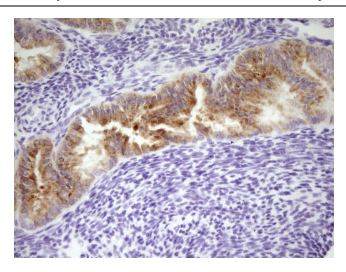
LIPN3L; SMP2

Product images:



HEK293T cells were transfected with the pCMV6-ENTRY control (Cat# [PS100001], Left lane) or pCMV6-ENTRY LPIN3 (Cat# [RC215286], 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-LPIN3(Cat# [TA804944]). Positive lysates [LY411469] (100ug) and [LC411469] (20ug) can be purchased separately from OriGene.

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Immunohistochemical staining of paraffinembedded Human endometrium tissue within the normal limits using anti-LPIN3 mouse monoclonal antibody. Heat-induced epitope retrieval by EDTA solution buffer pH 8.0 at 120°C for 3 min.

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