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# Product datasheet for CF805037

## Lipin 3 (LPIN3) Mouse Monoclonal Antibody [Clone ID: OTI1B10]

## **Product data:**

Product Type:	Primary Antibodies
Clone Name:	OTI1B10
Applications:	WB
Recommended Dilution:	WB 1:2000
Reactivity:	Human
Host:	Mouse
lsotype:	lgG2b
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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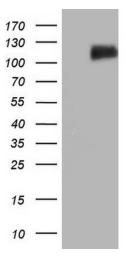
#### STATES CRIGENE Lipin 3 (LPIN3) Mouse Monoclonal Antibody [Clone ID: OTI1B10] – CF805037

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

#### Synonyms:

LIPN3L; SMP2

### **Product images:**



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

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