

Product datasheet for TA503423BM

OriGene Technologies, Inc.

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Lipoamide Dehydrogenase (DLD) Mouse Monoclonal Antibody (HRP conjugated) [Clone ID: OTI4D5]

Product data:

Product Type: Primary Antibodies

Clone Name: OTI4D5

Applications: FC, IF, IHC, WB

Recommended Dilution: WB 1:2000, IHC 1:150, IF 1:100, FLOW 1:100

Reactivity: Human, Mouse, Rat

Host: Mouse Isotype: IgG1

Clonality: Monoclonal

Immunogen: Full length human recombinant protein of human DLD(NP_000099) produced in HEK293T

cell.

Formulation: PBS (pH 7.3) containing 1% BSA, 50% glycerol.

Concentration: 0.5 mg/ml

Purification: Purified from mouse ascites fluids or tissue culture supernatant by affinity chromatography

(protein A/G)

Conjugation: HRP

Storage: Store at -20°C as received.

Stability: Stable for 12 months from date of receipt.

Predicted Protein Size: 50.1 kDa

Gene Name: dihydrolipoamide dehydrogenase

Database Link: NP 000099

Entrez Gene 13382 MouseEntrez Gene 298942 RatEntrez Gene 1738 Human

P09622



Lipoamide Dehydrogenase (DLD) Mouse Monoclonal Antibody (HRP conjugated) [Clone ID: OTI4D5] – TA503423BM

Background: This gene encodes the L protein of the mitochondrial glycine cleavage system. The L protein,

also named dihydrolipoamide dehydrogenase, is also a component of the pyruvate dehydrogenase complex, the alpha-ketoglutarate dehydrogenase complex, and the branched-chain alpha-keto acide dehydrogenase complex. Mutations in this gene have been

identified in patients with E3-deficient maple syrup urine disease and lipoamide

dehydrogenase deficiency. [provided by RefSeq]

Synonyms: DLDD; DLDH; E3; GCSL; LAD; PHE3

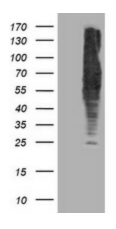
Protein Families: Druggable Genome

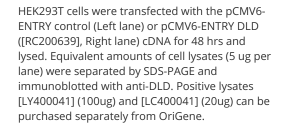
Protein Pathways: Citrate cycle (TCA cycle), Glycine, serine and threonine metabolism, Glycolysis /

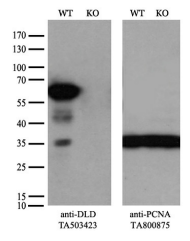
Gluconeogenesis, Metabolic pathways, Pyruvate metabolism, Valine, leucine and isoleucine

degradation

Product images:

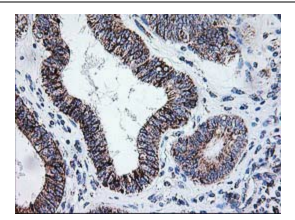




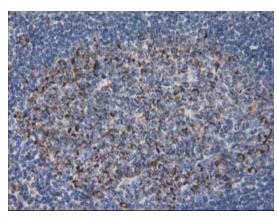


Equivalent amounts of cell lysates (10 ug per lane) of wild-type HeLa cells (WT, Cat# LC810HELA) and DLD-Knockout HeLa cells (KO, Cat# [LC832718]) were separated by SDS-PAGE and immunoblotted with anti-DLD monoclonal antibody [TA503423] (1:500). Then the blotted membrane was stripped and reprobed with anti-PCNA antibody as a loading control.

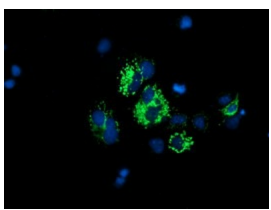
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Immunohistochemical staining of paraffinembedded Adenocarcinoma of Human endometrium tissue using anti-DLD mouse monoclonal antibody. (Heat-induced epitope retrieval by 10mM citric buffer, pH6.0, 100°C for 10min, [TA503423])

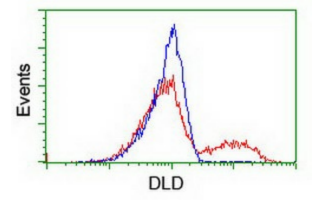


Immunohistochemical staining of paraffinembedded Human lymph node tissue within the normal limits using anti-DLD mouse monoclonal antibody. (Heat-induced epitope retrieval by 10mM citric buffer, pH6.0, 100°C for 10min, [TA503423])



Anti-DLD mouse monoclonal antibody ([TA503423]) immunofluorescent staining of COS7 cells transiently transfected by pCMV6-ENTRY DLD ([RC200639]).





HEK293T cells transfected with either [RC200639] overexpress plasmid (Red) or empty vector control plasmid (Blue) were immunostained by anti-DLD antibody ([TA503423]), and then analyzed by flow cytometry.