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CMC2 Rabbit pAb

CatalogNo: YN0524

Key Features

Host Species

Rabbit

ReactivityHuman,Mouse

ApplicationsWB,ELISA

MW • 74kD (Observed) Isotype • IgG

Recommended Dilution Ratios

WB 1:500-2000 ELISA 1:5000-20000

Storage

Storage*	-15°C to -25°C/1 year(Do not lower than -25°C)
Formulation	Liquid in PBS containing 50% glycerol,0.5% BSA and 0.02% sodium azide.

Basic Information

Clonality Polyclonal

Immunogen Information

Immunogen Synthesized peptide derived from part region of human protein

Specificity CMC2 Polyclonal Antibody detects endogenous levels of protein.

Target Information

Gene name SLC25A13 ARALAR2

Protein Name Calcium-binding mitochondrial carrier protein Aralar2 (Citrin) (Mitochondrial aspartate glutamate carrier 2) (Solute carrier family 25 member 13)

Organism	Gene ID	UniProt ID
Human	<u>10165;</u>	<u>Q9UJS0;</u>
Mouse		Q9QXX4;

Cellular Mitochondrion inner membrane ; Multi-pass membrane protein . **Localization**

Tissue specificity High levels in liver and low levels in kidney, pancreas, placenta, heart and brain.

Function Disease:Defects in SLC25A13 are the cause of citrullinemia type 2 (CTLN2) [MIM:603471]. Citrullinemia belongs to the urea cycle disorders. It is an autosomal recessive disease characterized primarily by elevated serum and urine citrulline levels. Ammonia intoxication is another manifestation. CTLN2 is characterized by neuropsychiatric symptoms including abnormal behaviors, loss of memory, seizures and coma. Death can result from brain edema. Onset is sudden and usually between the ages of 20 and 50 years., Disease: Defects in SLC25A13 are the cause of neonatal intrahepatic cholestasis due to citrin deficiency (NICCD) [MIM:605814]. NICCD is a form of citrullinemia type 2 with neonatal onset. NICCD is characterized by suppression of the bile flow, hepatic fibrosis, low birth weight, growth retardation, hypoproteinemia, variable liver dysfunction. NICCD is generally not severe and symptoms disappear by one year of age with an appropriate diet. Years or even decades later, however, some individuals develop the characteristic features of citrullinemia type 2 with neuropsychiatric symptoms., Function: Calcium-dependent mitochondrial aspartate and glutamate carrier. May have a function in the urea cycle., miscellaneous: Binds calcium., similarity: Belongs to the mitochondrial carrier family., similarity: Contains 3 Solcar repeats., similarity: Contains 4 EF-hand domains., tissue specificity: High levels in liver and low levels in kidney, pancreas, placenta, heart and brain.,

Validation Data

Contact information

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Please scan the QR code to access additional product information: CMC2 Rabbit pAb

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Antibody | ELISA Kits | Protein | Reagents