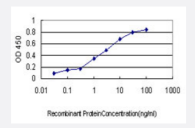


GCSH monoclonal antibody (M02), clone M2

Catalog # H00002653-M02 Size 100 ug

Applications



Sandwich ELISA (Recombinant protein)

Detection limit for recombinant GST tagged GCSH is approximately 0.1ng/ml as a capture antibody.



Western Blot detection against Immunogen (44.77 KDa).

Specification	
Product Description	Mouse monoclonal antibody raised against a full length recombinant GCSH.
Immunogen	GCSH (AAH00790.1, 1 a.a. ~ 173 a.a) full-length recombinant protein with GST tag. MW of the GST tag alone is 26 KDa.
Sequence	MALRVVRSVRALLCTLRAVPLPAAPCPPRPWQLGVGAVRTLRTGPALLSVRKFTEKHEWVTTEN GIGTVGISNFAQEALGDVVYCSLPEVGTKLNKQDEFGALESVKAASELYSPLSGEVTEINEALAEN PGLVNKSCYEDGWLIKMTLSNPSELDELMSEEAYEKYIKSIEE
Host	Mouse
Reactivity	Human



Product Information

Interspecies Antigen Sequence	Mouse (80); Rat (85)
Isotype	lgG1 Kappa
Quality Control Testing	Antibody Reactive Against Recombinant Protein. Western Blot detection against Immunogen (44.77 KDa).
Storage Buffer	In 1x PBS, pH 7.4
Storage Instruction	Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing.

Applications

Western Blot (Recombinant protein)

Protocol Download

Sandwich ELISA (Recombinant protein)

Detection limit for recombinant GST tagged GCSH is approximately 0.1ng/ml as a capture antibody.

Protocol Download

ELISA

Gene Info — GCSH	
Entrez GeneID	<u>2653</u>
GeneBank Accession#	BC000790
Protein Accession#	<u>AAH00790.1</u>
Gene Name	GCSH
Gene Alias	GCE, NKH
Gene Description	glycine cleavage system protein H (aminomethyl carrier)
Omim ID	238330 605899
Gene Ontology	<u>Hyperlink</u>



Product Information

Gene Summary

The enzyme system for cleavage of glycine (glycine cleavage system; EC 2.1.2.10), which is confined to the mitochondria, is composed of 4 protein components: P protein (a pyridoxal phosphate-dependent glycine decarboxylase; MIM 238300), H protein (a lipoic acid-containing protein), T protein (a tetrahydrofolate-requiring enzyme; MIM 238310), and L protein (a lipoamide dehydrogen ase; MIM 238331). Glycine encephalopathy (GCE; MIM 605899), also called nonketotic hyperglyc inemia (NKH), may be due to a defect in any one of these enzymes.[supplied by OMIM

Other Designations

lipoic acid-containing protein|mitochondrial glycine cleavage system H-protein|part of mitochondri al matrix glycine cleavage enzyme complex of 4 proteins: H-, L-, P-, and T-proteins

Disease

- Genetic Predisposition to Disease
- Hyperglycinemia
- Kidney Failure