

MAO-A Polyclonal Antibody

Catalog No: YT2635

Reactivity: Human; Mouse; Rat

Applications: WB;IHC;IF;ELISA

Target: MAO-A

Fields: >>Glycine, serine and threonine metabolism;>>Arginine and proline

metabolism;>>Histidine metabolism;>>Tyrosine metabolism;>>Phenylalanine metabolism;>>Tryptophan metabolism;>>Drug metabolism - cytochrome P450;>>Metabolic pathways;>>Serotonergic synapse;>>Dopaminergic synapse;>>Parkinson disease;>>Cocaine addiction;>>Amphetamine

addiction;>>Alcoholism

Gene Name: MAOA

Protein Name: Amine oxidase [flavin-containing] A

P21397

Q64133

Human Gene Id: 4128

Human Swiss Prot

No:

Mouse Gene ld: 17161

Mouse Swiss Prot

No:

Rat Swiss Prot No: P21396

Immunogen: The antiserum was produced against synthesized peptide derived from human

MAO-A. AA range:298-347

Specificity: MAO-A Polyclonal Antibody detects endogenous levels of MAO-A protein.

Formulation : Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.

Source: Polyclonal, Rabbit, IgG

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Dilution : WB 1:500 - 1:2000. IHC 1:100 - 1:300. ELISA: 1:20000.. IF 1:50-200

Purification: The antibody was affinity-purified from rabbit antiserum by affinity-

chromatography using epitope-specific immunogen.

Concentration: 1 mg/ml

Storage Stability: -15°C to -25°C/1 year(Do not lower than -25°C)

Observed Band: 61kD

Cell Pathway: Glycine; serine and threonine metabolism; Arginine and proline

metabolism;Histidine metabolism;Tyrosine metabolism;Phenylalanine

metabolism; Tryptophan metabolism; Drug metabolism;

Background: This gene is one of two neighboring gene family members that encode

mitochondrial enzymes which catalyze the oxidative deamination of amines, such as dopamine, norepinephrine, and serotonin. Mutation of this gene results in Brunner syndrome. This gene has also been associated with a variety of other psychiatric disorders, including antisocial behavior. Alternatively spliced transcript variants encoding multiple isoforms have been observed. [provided by RefSeq,

Jul 2012],

Function : catalytic activity:RCH(2)NHR' + H(2)O + O(2) = RCHO + R'NH(2) +

H(2)O(2).,cofactor:FAD.,disease:Defects in MAOA are the cause of Brunner syndrome (BRUNS) [MIM:300615]. Brunner syndrome is a form of X-linked non-dysmorphic mild mental retardation. Male patients are affected by a syndrome of borderline mental retardation and exhibit abnormal behavior, including disturbed

regulation of impulsive aggression. Obligate female carriers have normal intelligence and behavior., function: Catalyzes the oxidative deamination of

biogenic and xenobiotic amines and has important functions in the metabolism of neuroactive and vasoactive amines in the central nervous system and peripheral

tissues. MAOA preferentially oxidizes biogenic amines such as 5-hydroxytryptamine (5-HT), norepinephrine and epinephrine.,mass spectrometry: PubMed:11812236,online information:Monoamine oxidase

entry, similarity: Belongs to

Subcellular Location :

Mitochondrion outer membrane; Single-pass type IV membrane protein;

Cytoplasmic side.

Expression: Heart, liver, duodenum, blood vessels and kidney.

Tag: hot

Sort : 9361

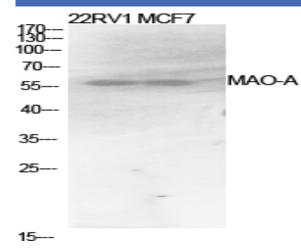


No4: 1

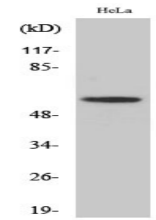
Host: Rabbit

Modifications: Unmodified

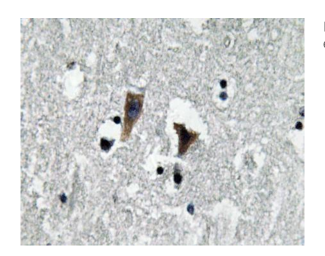




Western Blot analysis of various cells using MAO-A Polyclonal Antibody diluted at 1:1000



Western Blot analysis of HeLa cells using MAO-A Polyclonal Antibody diluted at 1:1000



Immunohistochemistry analysis of MAO-A antibody in paraffinembedded human brain tissue.

