



## **Choactase Polyclonal Antibody**

| Catalog No         | BYab-12703   |
|--------------------|--|
| Isotype            | lgG  |
| Reactivity         | Human;Mouse;Rat  |
| Applications       | WB;ELISA   |
| Gene Name          | CHAT   |
| Protein Name       | Choline O-acetyltransferase  |
| Immunogen          | The antiserum was produced against synthesized peptide derived from human Choactase. AA range:334-383  |
| Specificity        | Choactase Polyclonal Antibody detects endogenous levels of Choactase protein.  |
| Formulation        | Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.  |
| Source             | Polyclonal, Rabbit,IgG   |
| Purification       | The antibody was affinity-purified from rabbit antiserum by affinity-chromatography using epitope-specific immunogen.  |
| Dilution           | Western Blot: 1/500 - 1/2000. ELISA: 1/10000. Not yet tested in other applications.  |
| Concentration      | 1 mg/ml  |
| Purity             | ≥90%   |
| Storage Stability  | -20°C/1 year   |
| Synonyms           | CHAT; Choline O-acetyltransferase; CHOACTase; ChAT; Choline acetylase  |
| Observed Band      | 82,70kD  |
| Cell Pathway       | nucleus,cytoplasm,cytosol,presynapse,  |
| Tissue Specificity | Brain,Lymphocyte,Placenta,Spinal cord,   |
| Function           | catalytic activity:Acetyl-CoA + choline = CoA + O-acetylcholine.,disease:Defects<br>in CHAT are the cause of congenital myasthenic syndrome with episodic apnea<br>(CMSEA) [MIM:254210]; formerly known as familial infantile myasthenia gravis 2<br>(FIMG2). CMSEA is an autosomal recessive congenital myasthenic syndrome.<br>Patients have myasthenic symptoms since birth or early infancy, negative tests fo<br>anti-AChR antibodies, and abrupt episodic crises with increased weakness,<br>bulbar paralysis, and apnea precipitated by undue exertion, fever, or<br>excitement.,function:Catalyzes the reversible synthesis of acetylcholine (ACh)<br>from acetyl CoA and choline at cholinergic synapses.,online information:Choline<br>acetyltransferase entry,similarity:Belongs to the carnitine/choline<br>acetyltransferase family., |
|                    |  |

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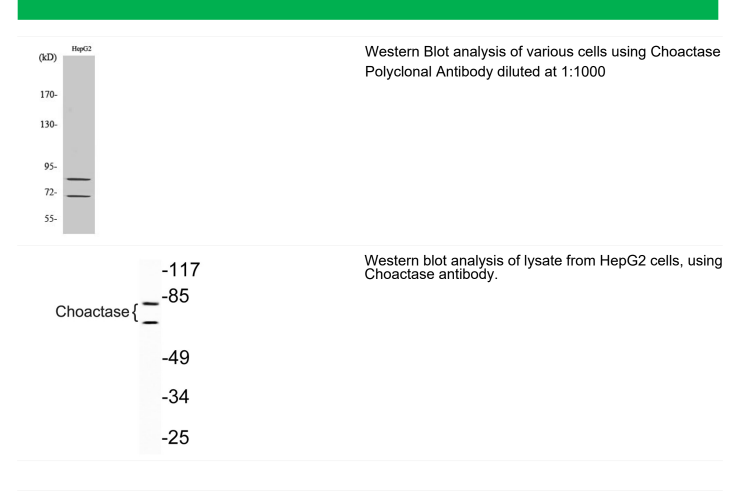
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| Background                | This gene encodes an enzyme which catalyzes the biosynthesis of the<br>neurotransmitter acetylcholine. This gene product is a characteristic feature of<br>cholinergic neurons, and changes in these neurons may explain some of the<br>symptoms of Alzheimer's disease. Polymorphisms in this gene have been<br>associated with Alzheimer's disease and mild cognitive impairment.<br>Mutations in this gene are associated with congenital myasthenic syndrome<br>associated with episodic apnea. Multiple transcript variants encoding different<br>isoforms have been found for this gene, and some of these variants have been<br>shown to encode more than one isoform. [provided by RefSeq, May 2010], |
|---------------------------|---|
| matters needing attention | Avoid repeated freezing and thawing!  |
| Usage suggestions         | This product can be used in immunological reaction related experiments. For more information, please consult technical personnel.   |

## **Products Images**



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