

AGT Monoclonal Antibody

Catalog No: YM0016

Reactivity: Human

Applications: WB;ELISA

Target: AGT

Fields: >>Phospholipase D signaling pathway;>>Neuroactive ligand-receptor

interaction;>>Adrenergic signaling in cardiomyocytes;>>Vascular smooth muscle contraction;>>Renin-angiotensin system;>>Renin secretion;>>Aldosterone

synthesis and secretion;>>Cortisol synthesis and secretion;>>Insulin

resistance;>>AGE-RAGE signaling pathway in diabetic complications;>>Cushing

syndrome;>>Pathways in cancer;>>Hypertrophic cardiomyopathy;>>Dilated

cardiomyopathy;>>Diabetic cardiomyopathy

Gene Name: AGT

Protein Name: Angiotensinogen

P01019

P11859

Human Gene Id: 183

Human Swiss Prot

No:

Mouse Swiss Prot

No:

Immunogen: Purified recombinant fragment of human AGT expressed in E. Coli.

Specificity: AGT Monoclonal Antibody detects endogenous levels of AGT protein.

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

Source: Monoclonal, Mouse

Dilution: WB 1:500 - 1:2000. ELISA: 1:10000. Not yet tested in other applications.

Purification : Affinity purification

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

Molecularweight: 52kD

Cell Pathway: Renin-angiotensin system;

P References: 1. Am J Physiol Heart Circ Physiol. 2007 Sep;293(3):H1900-7.

2. Regul Pept. 2006 Jan 15;133(1-3):155-9.

Background : The protein encoded by this gene, pre-angiotensinogen or angiotensinogen

precursor, is expressed in the liver and is cleaved by the enzyme renin in response to lowered blood pressure. The resulting product, angiotensin I, is then cleaved by angiotensin converting enzyme (ACE) to generate the physiologically active enzyme angiotensin II. The protein is involved in maintaining blood pressure and in the pathogenesis of essential hypertension and preeclampsia. Mutations in this gene are associated with susceptibility to essential hypertension, and can cause renal tubular dysgenesis, a severe disorder of renal tubular development. Defects in this gene have also been associated with non-familial structural atrial fibrillation, and inflammatory bowel disease. [provided by RefSeq,

Jul 2008],

Function: caution:It is uncertain whether Met-1 or Met-10 is the initiator.,disease:Defects in

AGT are a cause of renal tubular dysgenesis (RTD) [MIM:267430]. RTD is an autosomal recessive severe disorder of renal tubular development characterized by persistent fetal anuria and perinatal death, probably due to pulmonary

hypoplasia from early-onset oligohydramnios (the Potter

phenotype)., disease: Defects in AGT are associated with susceptibility to essential hypertension [MIM:145500]. Hypertension also occurs in 5-7% of all pregnancies where it is a leading cause of maternal, fetal and neonatal morbidity and mortality. Among pregnancy-induced hypertension cases, severe preeclampsia [MIM:189800] is characterized by the development of hypertension and proteinuria after the 20th week of pregnancy and is the most distinctive, life-

threatening form., function: Angiotensin-3 stimulates aldosterone release.

Subcellular Location:

Secreted.

Expression : Expressed by the liver and secreted in plasma.

Sort: 1794

No4: 1

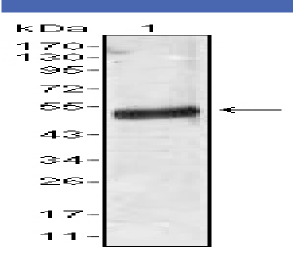
Host: Mouse

Modifications: Unmodified

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Products Images



Western Blot analysis using AGT Monoclonal Antibody against human plasma (1).