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Mouse Monoclonal Antibody MYL2 conjugated to Sepharose Beads

CatalogNo: ANT8267-S

Size 200ul

Storage Store at 4 °C for frequent use

Description

This Antagene antibody is immobilized via covalent binding of primary amino groups to N-hydroxysuccinimide (NHS)-activated sepharose beads. It is useful for immunoprecipitation assays.

MYL2 (ANT0025R) Rabbit mAb

Formulation: 50% slurry in PBS pH 7.2 with 0.01mg NaN3a3 preservative.

Host Species Reactivity Applications

• Rabbit • Human, Mouse, Rat, • WB, IHC, IF, IP, ELISA

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MW Isotype

18kD (Calculated)
 IgG,Kappa
 18kD (Observed)

#### **Recommended Dilution Ratios**

#### IP

#### **Basic Information**

**Clonality** Monoclonal

Clone Number ANT0025R

### Immunogen Information

**Specificity** Endogenous

Gene name MYL2 **Protein Name** MYL2

Organism	Gene ID	UniProt ID
Human	<u>4633</u> ;	<u>P10916</u> ;
Mouse	<u>17906</u> ;	<u>P51667</u> ;

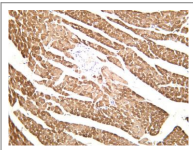
Cellular Localization Cytoplasm

**Tissue specificity** Highly expressed in type I muscle fibers.

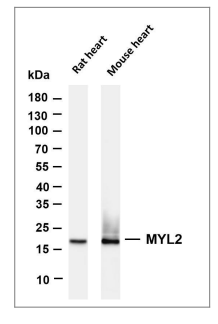
**Function** 

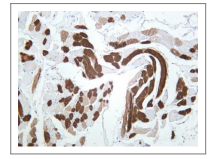
Disease:Defects in MYL2 are the cause of cardiomyopathy familial hypertrophic type 10 (CMH10) [MIM:608758]. Familial hypertrophic cardiomyopathy is a hereditary heart disorder characterized by ventricular hypertrophy, which is usually asymmetric and often involves the interventricular septum. The symptoms include dyspnea, syncope, collapse, palpitations, and chest pain. They can be readily provoked by exercise. The disorder has inter- and intrafamilial variability ranging from benign to malignant forms with high risk of cardiac failure and sudden cardiac death., Disease: Defects in MYL2 are the cause of cardiomyopathy hypertrophic with mid-left ventricular chamber type 2 (MVC2) [MIM:608758]. MVC2 is a very rare variant of familial hypertrophic cardiomyopathy, characterized by mid-left ventricular chamber thickening., miscellaneous: This chain binds calcium., similarity: Contains 3 EF-hand domains., subunit: Myosin is an hexamer of 2 heavy chains and 4 light chains.,

## **Validation Data**

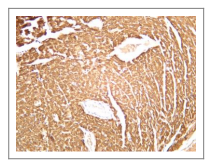


Rat cardiac muscle was stained with anti-MYL2 (ANT0025R) rabbit antibody Various whole cell lysates were separated by 4-20% SDS-PAGE, and the membrane was blotted with anti-MYL2 (ANT0067R) antibody. The HRPconjugated Goat anti-Rabbit IgG(H + L) antibody was used to detect the antibody. Lane 1: Rat heart Lane 2: Mouse heart Predicted band size: 18kDa Observed band size: 18kDa





Human skeletal muscle was stained with anti-MYL2 (ANT0025R) rabbit antibody



Mouse cardiac muscle was stained with anti-MYL2 (ANT0025R) rabbit antibody

# For Research use only, not for diagnostics and clinical use Contact Antagene Inc Tel 1-866-964-2589 Email: info@antageneinc.com