

Troponin T-C Polyclonal Antibody

Catalog No: YT5362

Reactivity: Human; Mouse; Rat

Applications: WB;IHC;IF;ELISA

Target: Troponin T-C

Fields: >> Cardiac muscle contraction;>> Adrenergic signaling in

cardiomyocytes;>>Hypertrophic cardiomyopathy;>>Dilated cardiomyopathy

Gene Name: TNNT2

Protein Name: Troponin T cardiac muscle

P45379

P50752

Human Gene Id: 7139

Human Swiss Prot

No:

Mouse Gene Id: 21956

Mouse Swiss Prot

No:

Rat Gene Id: 24837

Rat Swiss Prot No: P50753

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

Internal region of human TNNT2. AA range:131-180

Specificity: Troponin T-C Polyclonal Antibody detects endogenous levels of Troponin T-C

protein.

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

Source: Polyclonal, Rabbit, IgG

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

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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: 35kD

Cell Pathway: Cardiac muscle contraction; Hypertrophic cardiomyopathy (HCM); Dilated

cardiomyopathy;

Background: The protein encoded by this gene is the tropomyosin-binding subunit of the

troponin complex, which is located on the thin filament of striated muscles and regulates muscle contraction in response to alterations in intracellular calcium ion

concentration. Mutations in this gene have been associated with familial

hypertrophic cardiomyopathy as well as with dilated cardiomyopathy. Transcripts for this gene undergo alternative splicing that results in many tissue-specific isoforms, however, the full-length nature of some of these variants has not yet

been determined. [provided by RefSeq, Jul 2008],

Function: alternative products:Additional isoforms seem to exist. Experimental

confirmation may be lacking for some isoforms, disease: Defects in TNNT2 are the cause of cardiomyopathy dilated type 1D (CMD1D) [MIM:601494]. Dilated cardiomyopathy is a disorder characterized by ventricular dilation and impaired systolic function, resulting in congestive heart failure and arrhythmia. Patients are

at risk of premature death.,disease:Defects in TNNT2 are the cause of cardiomyopathy familial hypertrophic type 2 (CMH2) [MIM:115195]. 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

Subcellular Location:

cytosol,troponin complex,striated muscle thin filament,sarcomere,

Expression: Heart. The fetal heart shows a greater expression in the atrium than in the

ventricle, while the adult heart shows a greater expression in the ventricle than in the atrium. Isoform 6 predominates in normal adult heart. Isoforms 1, 7 and 8 are

expressed in fetal heart. Isoform 7 is also expressed in failing adult heart.

Sort: 23596

No4:

Host: Rabbit



Modifications: Unmodified

Products Images

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