

## Neurofilament (PT0124R) rabbit mAb

YM7175 Catalog No:

Reactivity: Human;

**Applications:** IHC;WB; ELISA

NF-L **Target:** 

Fields: >>Amyotrophic lateral sclerosis;>>Pathways of neurodegeneration - multiple

diseases

P07196

Gene Name: **NEFL** 

**Protein Name:** Neurofilament

**Human Gene Id:** 4747

**Human Swiss Prot** 

No:

Synthesized peptide derived from human Neurofilament AA range:400-543 Immunogen:

This antibody detects endogenous levels of NF-L **Specificity:** 

PBS, 50% glycerol, 0.05% Proclin 300, 0.05% BSA Formulation:

Monoclonal, Rabbit IgG1, Kappa Source:

IHC 1:100-500, WB 1:500-1000, ELISA 1:5000-20000 **Dilution:** 

**Purification:** Recombinant Expression and Affinity purified

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

**Molecularweight:** 62kD

Neurofilaments are type IV intermediate filament heteropolymers composed of **Background:** 

> light, medium, and heavy chains. Neurofilaments comprise the axoskeleton and they functionally maintain the neuronal caliber. They may also play a role in



intracellular transport to axons and dendrites. This gene encodes the light chain neurofilament protein. Mutations in this gene cause Charcot-Marie-Tooth disease types 1F (CMT1F) and 2E (CMT2E), disorders of the peripheral nervous system that are characterized by distinct neuropathies. A pseudogene has been identified on chromosome Y. [provided by RefSeq, Oct 2008],

## **Function:**

caution:The sequence shown here is derived from an Ensembl automatic analysis pipeline and should be considered as preliminary data., disease:Defects in NEFL are the cause of Charcot-Marie-Tooth disease type 1F (CMT1F) [MIM:607734]. CMT1F is a form of Charcot-Marie-Tooth disease, the most common inherited disorder of the peripheral nervous system. Charcot-Marie-Tooth disease is classified in two main groups on the basis of electrophysiologic properties and histopathology: primary peripheral demyelinating neuropathy or CMT1, and primary peripheral axonal neuropathy or CMT2. Neuropathies of the CMT1 group are characterized by severely reduced nerve conduction velocities (less than 38 m/sec), segmental demyelination and remyelination with onion bulb formations on nerve biopsy, slowly progressive distal muscle atrophy and weakness, absent deep tendon reflexes, and hollow feet. CMT1F is charac

Subcellular Location:

Cytoplasmic

**Expression:** Cytoplasmic

## **Products Images**

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