abcam

Product datasheet

Recombinant Human Hsp27 protein ab91578

画像数 2

製品の詳細	
製品名	Recombinant Human Hsp27 protein
精製度	> 95 % SDS-PAGE.
	Purity > 95% pure as determined by SDS-PAGE and Western Blot analysis.
発現系	Escherichia coli
タンパク質長	Full length protein
Animal free	No
由来	Recombinant
生物種	Human

特性

Our Abpromise guarantee covers the use of ab91578 in the following tested applications.

The application notes include recommended starting dilutions; optimal dilutions/concentrations should be determined by the end user.

アプリケーション	Western blot
	SDS-PAGE
製品の状態	Liquid
前処理および保存	

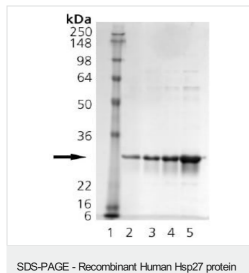
保存方法および安定性	Shipped on dry ice. Upon delivery aliquot and store at -80°C. Avoid freeze / thaw cycles.
	Constituents: 0.0154% (R*,R*)-1,4-Dimercaptobutan-2,3-diol, 0.242% Tris, 0.0292% EDTA,
	0.058% Sodium chloride

関連情報

機能	Involved in stress resistance and actin organization.
組織特異性	Detected in all tissues tested: skeletal muscle, heart, aorta, large intestine, small intestine,
	stomach, esophagus, bladder, adrenal gland, thyroid, pancreas, testis, adipose tissue, kidney,
	liver, spleen, cerebral cortex, blood serum and cerebrospinal fluid. Highest levels are found in the
	heart and in tissues composed of striated and smooth muscle.

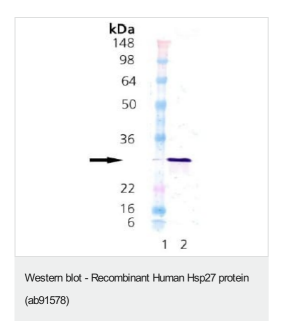
関連疾患	Defects in HSPB1 are the cause of Charcot-Marie-Tooth disease type 2F (CMT2F) [MIM:606595]. CMT2F is a form of Charcot-Marie-Tooth disease, the most common inherited disorder of the peripheral nervous system. Charcot-Marie-Tooth disease is classified in two main groups on the basis of electrophysiologic properties and histopathology: primary peripheral demyelinating neuropathy or CMT1, and primary peripheral axonal neuropathy or CMT2. Neuropathies of the CMT2 group are characterized by signs of axonal regeneration in the absence of obvious myelin alterations, normal or slightly reduced nerve conduction velocities, and progressive distal muscle weakness and atrophy. Nerve conduction velocities are normal or slightly reduced. CMT2F onset is between 15 and 25 years with muscle weakness and atrophy usually beginning in feet and legs (peroneal distribution). Upper limb involvement occurs later. CMT2F inheritance is autosomal dominant. Defects in HSPB1 are a cause of distal hereditary motor neuronopathy type 2B (HMN2B) [MIM:608634]. Distal hereditary motor neuronopathies constitute a heterogeneous group of neuromuscular disorders caused by selective impairment of motor neurons in the anterior horn of the spinal cord, without sensory deficit in the posterior horn. The overall clinical picture consists of a classical distal muscular atrophy syndrome in the legs without clinical sensory loss. The disease starts with weakness and wasting of distal muscles of the anterior tibial and peroneal compartments of the legs. Later on, weakness and atrophy may expand to the proximal muscles of the lower limbs and/or to the distal upper limbs.
配列類似性	Belongs to the small heat shock protein (HSP20) family.
翻訳後修飾	Phosphorylated in MCF-7 cells on exposure to protein kinase C activators and heat shock.
細胞内局在	Cytoplasm. Nucleus. Cytoplasm > cytoskeleton > spindle. Cytoplasmic in interphase cells. Colocalizes with mitotic spindles in mitotic cells. Translocates to the nucleus during heat shock and resides in sub-nuclear structures known as SC35 speckles or nuclear splicing speckles.

画像



SDS-PAGE Analysis stained with Imperial stain Lane 1: MW Marker Lane 2: 0.5 µg Lane 3: 1 µg, Lane 4: 2 µg Lane 5 5 µg

(ab91578) SDS-PAGE - Recombinant Human Hsp27 protein



Western Blot Analysis Lane 1 MW Marker Lane 2 ab91578 probed with anti-Hsp27 monoclonal antibody

Please note: All products are "FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES"

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