abcam

Product datasheet

Recombinant Human SQSTM1 / p62 protein ab95320

4 References 1 Image

Description

Product name Recombinant Human SQSTM1 / p62 protein

Purity > 85 % SDS-PAGE.

ab95320 is purified using conventional chromatography techniques

Expression system Escherichia coli

Accession Q13501

Protein length Protein fragment

Animal free No

Nature Recombinant

Species Human

Sequence MAMSYVKDDI FRIYIKEKKE CRRDHRPPCA

QEAPRNMVHP NVICDGCNGP VVGTRYKCSV
CPDYDLCSVC EGKGLHRGHT KLAFPSPFGH
LSEGFSHSRW LRKVKHGHFG WPGWEMGPPG
NWSPRPPRAG EARPGPTAES ASGPSEDPSV
NFLKNVGESV AAALSPLGIE VDIDVEHGGK
RSRLTPVSPE SSSTEEKSSS QPSSCCSDPS
KPGGNVEGAT QSLAEQMRKI ALESEGRPEE
QMESDNCSGG DDDWTHLSSK EVDPSTGELQ
SLQMPESEGP SSLDPSQEGP TGLKEAALYP
HLPPEADPRL IESLSQMLSM GFSDEGGWLT
RLLQTKNYDI GAALDTIQYS KHPPPLLEHH HHHH

Amino acids 85 to 440

Tags His tag C-Terminus

Specifications

Our Abpromise guarantee covers the use of ab95320 in the following tested applications.

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

Applications SDS-PAGE

Mass Spectrometry

Form Liquid

1

Preparation and Storage

Stability and Storage

Shipped at 4°C. Upon delivery aliquot and store at -20°C or -80°C. Avoid repeated freeze / thaw

cycles.

pH: 8.00

Constituents: 0.0154% DTT, 0.316% Tris HCI, 10% Glycerol (glycerin, glycerine)

General Info

Function Adapter protein which binds ubiquitin and may regulate the activation of NFKB1 by TNF-alpha,

nerve growth factor (NGF) and interleukin-1. May play a role in titin/TTN downstream signaling in muscle cells. May regulate signaling cascades through ubiquitination. Adapter that mediates the interaction between TRAF6 and CYLD (By similarity). May be involved in cell differentiation,

apoptosis, immune response and regulation of K(+) channels.

Tissue specificity

Ubiquitously expressed.

Involvement in disease

Defects in SQSTM1 are a cause of Paget disease of bone (PDB) [MIM:602080]. PDB is a metabolic bone disease affecting the axial skeleton and characterized by focal areas of increased and disorganized bone turn-over due to activated osteoclasts. Manifestations of the disease include bone pain, deformity, pathological fractures, deafness, neurological

complications and increased risk of osteosarcoma. PDB is a chronic disease affecting 2 to 3% of

the population above the age of 40 years.

Sequence similarities

Contains 1 OPR domain.
Contains 1 UBA domain.

Contains 1 ZZ-type zinc finger.

Domain

The UBA domain binds specifically 'Lys-63'-linked polyubiquitin chains of polyubiquitinated

substrates. Mediates the interaction with TRIM55.

The OPR domain mediates homooligomerization and interactions with PRKCZ, PRKCI, MAP2K5

and NBR1.

The ZZ-type zinc finger mediates the interaction with RIPK1.

Post-translational modifications

Phosphorylated. May be phosphorylated by PRKCZ (By similarity). Phosphorylated in vitro by

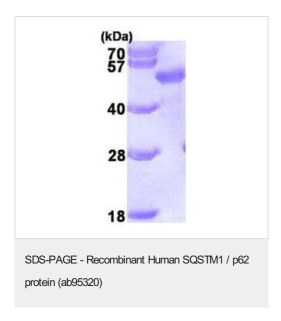
TTN.

Cellular localization

Cytoplasm. Late endosome. Nucleus. Sarcomere (By similarity). In cardiac muscles localizes to the sarcomeric band (By similarity). Localizes to late endosomes. May also localize to the nucleus. Accumulates in neurofibrillary tangles and in Lewy bodies of neurons from individuals with Alzheimer and Parkinson disease respectively. Enriched in Rosenthal fibers of pilocytic astrocytoma. In liver cells, accumulates in Mallory bodies associated with alcoholic hepatitis, Wilson disease, indian childhood cirrhosis and in hyaline bodies associated with hepatocellular

carcinoma.

Images



ab95320 at 3 µg on an SDS-PAGE gel (15%).

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