

P91042Hu02

Integrin Beta 1 (ITGb1)

Organism: Homo sapiens (Human)

Instruction manual

FOR IN VITRO USE AND RESEARCH USE ONLY
NOT FOR USE IN DIAGNOSTIC OR THERAPEUTIC PROCEDURES

4th Edition (Revised in August, 2012)

## [ DESCRIPTION ]

Protein Names: Integrin Beta 1

**Synonyms:** ITGb1, FNRB, MDF2, MSK12

**Species:** Human

Size: 100µg

Source: Escherichia coli-derived

Subcellular Location: Cell membrane; Single-pass type I membrane protein.

Melanosome. Cleavage furrow.

# [PROPERTIES]

Residues: Gln467~Thr635 (Accession # P05556), with two N-terminal Tags, His-tag

and GST-tag.

**Grade & Purity:** >95%, 45 kDa as determined by SDS-PAGE reducing conditions. **Formulation:** Supplied as liquid form in Phosphate buffered saline(PBS), pH 7.4.

**Endotoxin Level:** <1.0 EU per 1µg (determined by the LAL method).

Applications: SDS-PAGE; WB; ELISA; IP.

(May be suitable for use in other assays to be determined by the end user.)

Predicted Molecular Mass: 45.3 kDa Predicted isoelectric point: 5.6

## [PREPARATION]

Reconstitute in sterile PBS, pH7.2-pH7.4.





### [STORAGE AND STABILITY]

Storage: Avoid repeated freeze/thaw cycles.

Store at 2-8°C for one month.

Aliquot and store at -80°C for 12 months.

**Stability Test:** The thermal stability is described by the loss rate of the target protein. The loss rate was determined by accelerated thermal degradation test, that is, incubate the protein at 37°C for 48h, and no obvious degradation and precipitation were observed. (Referring from China Biological Products Standard, which was calculated by the Arrhenius equation.) The loss of this protein is less than 5% within the expiration date under appropriate storage condition.

#### [SEQUENCES]

The target protein is fused with two N-terminal Tags, His-tag and GST-tag, its sequence is listed below.

MRNKKFELGLEFPNLPYYIDGDVKLTQSMAIIRYIADKHNMLGGCPKERAEISMLEGAVLDIRYGVSRIAYSKDFETLKVDFLSKLPEML

KMFEDRLCHKTYLNGDHVTHPDFMLYDALDVVLYMDPMCLDAFPKLVCFKKRIEAIPQIDKYLKSSKYIAWPLQGWQATFGGGDHPP

KSDGSTSGSGHHHHHHSAGLVPRGSTAIGMKETAAAKFERQHMDSPDLGTLEVLFQGPLGSEF-QSEG IPESPKCHEG

NGTFECGACR CNEGRVGRHC ECSTDEVNSE DMDAYCRKEN SSEICSNNGE CVCGQCVCRK RDNTNEIYSG KFCECDNFNC

DRSNGLICGG NGVCKCRVCE CNPNYTGSAC DCSLDTSTCE ASNGQICNGR GICECGVCKC TDPKFQGQTC EMCQT