

Product datasheet for AR50820PU-S

OriGene Technologies, Inc.

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TUBB3 / TUBB4 (1-450, His-tag) Human Protein

Product data:

Product Type: Recombinant Proteins

Description: TUBB3 / TUBB4 (1-450, His-tag) human recombinant protein, 50 µg

Species: Human
Expression Host: E. coli

Expression cDNA Clone

or AA Sequence:

MGSSHHHHHH SSGLVPRGSH MGSMREIVHI QAGQCGNQIG AKFWEVISDE HGIDPSGNYV GDSDLQLERI SVYYNEASSH KYVPRAILVD LEPGTMDSVR SGAFGHLFRP DNFIFGQSGA GNNWAKGHYT EGAELVDSVL DVVRKECENC DCLQGFQLTH SLGGGTGSGM GTLLISKVRE

EYPDRIMNTF SVVPSPKVSD TVVEPYNATL SIHQLVENTD ETYCIDNEAL YDICFRTLKL ATPTYGDLNH

LVSATMSGVT TSLRFPGQLN ADLRKLAVNM VPFPRLHFFM PGFAPLTARG SQQYRALTVP ELTQQMFDAK NMMAACDPRH GRYLTVATVF RGRMSMKEVD EQMLAIQSKN SSYFVEWIPN NVKVAVCDIP PRGLKMSSTF IGNSTAIQEL FKRISEQFTA MFRRKAFLHW YTGEGMDEME

FTEAESNMND LVSEYQQYQD ATAEEEGEMY EDDEEESEAQ GPK

Tag: His-tag

Predicted MW: 52.8 kDa

Concentration: lot specific

Purity: >90% by SDS - PAGE

Buffer: Presentation State: Purified

State: Liquid purified protein

Buffer System: 20 mM Tris-HCl buffer (pH 8.0) containing 0.15M NaCl, 10% glycerol, 1 mM

DTT

Preparation: Liquid purified protein

Protein Description: Recombinant human TUBB3 protein, fused to His-tag at N-terminus, was expressed in E.coli

and purified by using conventional chromatography techniques.

Storage: Store undiluted at 2-8°C for one week or (in aliquots) at -20°C to -80°C for longer. Avoid

repeated freezing and thawing.

Stability: Shelf life: one year from despatch.

RefSeq: NP 001184110

Locus ID: 10381



UniProt ID: Q13509
Cytogenetics: 16q24.3

Synonyms: beta-4; CDCBM; CDCBM1; CFEOM3; CFEOM3A; FEOM3; TUBB4

Summary: This gene encodes a class III member of the beta tubulin protein family. Beta tubulins are one

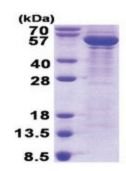
of two core protein families (alpha and beta tubulins) that heterodimerize and assemble to form microtubules. This protein is primarily expressed in neurons and may be involved in neurogenesis and axon guidance and maintenance. Mutations in this gene are the cause of congenital fibrosis of the extraocular muscles type 3. Alternate splicing results in multiple transcript variants. A pseudogene of this gene is found on chromosome 6. [provided by

RefSeq, Oct 2010]

Protein Families: Druggable Genome, ES Cell Differentiation/IPS

Protein Pathways: Gap junction, Pathogenic Escherichia coli infection

Product images:



15% SDS-PAGE (3ug)