

## **Product datasheet for TA501038S**

**TUBA8 Mouse Monoclonal Antibody [Clone ID: OTI2E9]** 

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## Product data:

**Product Type:** Primary Antibodies

Clone Name: OTI2E9

**Applications:** FC, IF, IP, WB

Recommended Dilution: WB 1:2000; IF 1:100; IP: 4ug/mL

Reactivity: Human, Mouse, Rat

Host: Mouse Isotype: IgG2a

Clonality: Monoclonal

Immunogen: Full length human recombinant protein of human TUBA8(NP\_061816) produced in HEK293T

cell

**Formulation:** PBS (pH 7.3) containing 1% BSA, 50% glycerol and 0.02% sodium azide.

Concentration: 1 mg/ml

**Purification:** Purified from mouse ascites fluids or tissue culture supernatant by affinity chromatography

(protein A/G)

Conjugation: Unconjugated

**Storage:** Store at -20°C as received.

**Stability:** Stable for 12 months from date of receipt.

**Predicted Protein Size:** 50.1 kDa

**Gene Name:** tubulin alpha 8

Database Link: NP 061816

Entrez Gene 500377 RatEntrez Gene 51807 Human

O9NY65





Background:

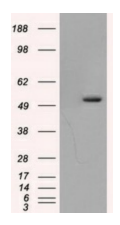
Microtubules are cylindrical tubes of 20-25 nm in diameter. They are composed of protofilaments which are in turn composed of alpha- and beta-tubulin polymers. Each microtubule is polarized, at one end alpha-subunits are exposed (-) and at the other beta-subunits are exposed (+). Microtubules act as a scaffold to determine cell shape, and provide a backbone for cell organelles and vesicles to move on, a process that requires motor proteins. The major microtubule motor proteins are kinesin, which generally moves towards the (+) end of the microtubule, and dynein, which generally moves towards the (-) end. Microtubules also form the spindle fibers for separating chromosomes during mitosis.

Synonyms: TUBAL2

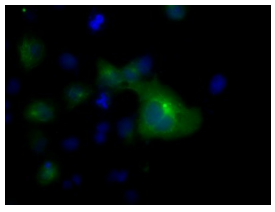
**Protein Families:** Druggable Genome

**Protein Pathways:** Gap junction, Pathogenic Escherichia coli infection

## **Product images:**

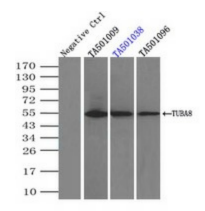


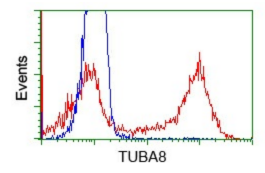
HEK293T cells were transfected with the pCMV6-ENTRY control (Left lane) or pCMV6-ENTRY TUBA8 ([RC211175], Right lane) cDNA for 48 hrs and lysed. Equivalent amounts of cell lysates (5 ug per lane) were separated by SDS-PAGE and immunoblotted with anti-TUBA8. Positive lysates [LY412867] (100ug) and [LC412867] (20ug) can be purchased separately from OriGene.



Anti-TUBA8 mouse monoclonal antibody ([TA501038]) immunofluorescent staining of COS7 cells transiently transfected by pCMV6-ENTRY TUBA8 ([RC211175]).







Immunoprecipitation (IP) of TUBA8 by using TrueMab monoclonal anti-TUBA8 antibodies (Negative control: IP without adding anti-TUBA8 antibody.). For each experiment, 500ul of DDK tagged TUBA8 overexpression lysates (at 1:5 dilution with HEK293T lysate), 2ug of anti-TUBA8 antibody and 20ul (0.1mg) of goat anti-mouse conjugated magnetic beads were mixed and incubated overnight. After extensive wash to remove any non-specific binding, the immunoprecipitated products were analyzed with rabbit anti-DDK polyclonal antibody.

HEK293T cells transfected with either pCMV6-ENTRY TUBA8 ([RC211175]) (Red) or empty vector control plasmid (Blue) were immunostained with anti-TUBA8 mouse monoclonal ([TA501038]), and then analyzed by flow cytometry.