

Product no **AS10 681**

## Tubulin beta chain

### Product information

**Immunogen** | KLH-conjugated peptide derived from available tubulin beta chain sequences (located at a surface loop) including all *Arabidopsis thaliana* tubulin beta-2/beta-3 chain [P29512](#), beta-4 chain [P24636\(At5g44340\)](#), beta-5 chain [P29513\(At1g20010\)](#), beta-6 chain [P29514\(At5g12250\)](#), beta-7 chain [P29515\(At2g29550\)](#), beta-8 chain [P29516\(At5g23860\)](#), beta-9 chain [P29517\(At4g20890\)](#).

**Host** | Rabbit

**Clonality** | Polyclonal

**Purity** | Affinity purified serum in PBS, pH 7.4

**Format** | Lyophilized in PBS pH 7.4

**Quantity** | 100 µg

**Reconstitution** | For reconstitution add 100 µl of sterile water

**Storage** | Store lyophilized/reconstituted at -20°C; once reconstituted make aliquots to avoid repeated freeze-thaw cycles. Please, remember to spin tubes briefly prior to opening them to avoid any losses that might occur from lyophilized material adhering to the cap or sides of the tubes.

**Additional information** | Please, note that tubulin beta is only detected in actively dividing meristematic cells (see immunolocalization image below). Therefore to allow detection on a western blot, analyzed material must contain enough meristematic cells with dividing activity.

Signal in a western blot application in *Chlamydomonas reinhardtii* is obtained with a load of 100 µg/well and a dilution of 1: 500.

### Application information

**Recommended dilution** | 1 : 1000 (IF), 1 : 500 (WB)

**Expected | apparent MW** | 49 | 49 kDa (*Arabidopsis thaliana*)

**Confirmed reactivity** | *Arabidopsis thaliana* (including suspension cells), *Chlamydomonas reinhardtii*, *Medicago sativa*

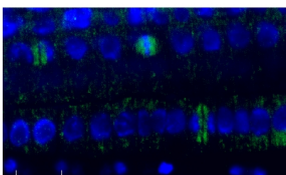
**Predicted reactivity** | *Brassica napus*, *Glycine max*, *Hordeum vulgare*, *Micromonosas pusilla*, *Oryza sativa*, *Ostreococcus lucimarinus*, *Picea sitchHensis*, *Pisum sativum*, *Physcomitrella patens*, *Populus trichocarpa*, *Saccharomyces cerevisiae*, *Solanum tuberosum*, *Sorghum bicolor*, *Ricinus communis*, *Zea mays*, *Vigna radiata*, *Vitis vinifera*  
Species of your interest not listed? [Contact us](#)

**Not reactive in** | No confirmed exceptions from predicted reactivity are currently known.

**Additional information** | Metal induced stress affected the expression of tubulin, and that therefore, this protein cannot be used as a loading control under that type of conditions. More information can be found [here](#). For high resolution images, please visit the specific product page at [www.agrisera.com](http://www.agrisera.com)

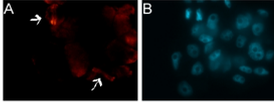
**Selected references** | [Durian et al. \(2019\)](#). PROTEIN PHOSPHATASE 2A-B' gamma controls Botrytis cinerea resistance and developmental leaf senescence. *Plant Physiol.* 2019 Oct 28. pii: pp.00893.2019. doi: 10.1104/pp.19.00893.  
[Wang et al. \(2017\)](#). The inhibition of protein translation mediated by AtGCN1 is essential for cold tolerance in *Arabidopsis thaliana*. *Plant Cell Environ.* 2017 Jan;40(1):56-68. doi: 10.1111/pce.12826.  
[Heinricke et al. \(2016\)](#). Tetratricopeptide repeat protein protects photosystem I from oxidative disruption during assembly. *Proc Natl Acad Sci U S A.* 2016 Mar 8;113(10):2774-9. doi: 10.1073/pnas.1524040113

### Application example



Tubulin beta localization in *Arabidopsis thaliana* epidermis cells. Tubulin beta localized to division plates visualized in green, nucleus with DAPI in blue. Plant material has been fixed in para-formaldehyde for 30 minutes. Tissue cleaning has been performed before immunolocalization. Primary antibodies: Agrisera anti-tubulin beta primary antibody in dilution 1: 1000 and Agrisera [goat anti-rabbit IgG DyLight® 488](#), secondary antibody in

dilution 1: 2000. Scale bar – 10  $\mu$ m.



**(A)** The 3 days old suspension cells of *Arabidopsis thaliana* have been fixed for 25 minutes with 2% FA, cell wall was partially digested with 0.2% Dricelase in MES buffer (pH 5.1) for 30 min at 37 °C and the cells were dried on coated slide to omit PM extraction. Following by incubation for 45 minutes with Agrisera anti-tubulin beta chain primary antibody in a dilution of 1: 1000, washing once and 45 min incubation with a secondary goat anti-rabbit Alexa 555 antibody, in a dilution of 1:1000 dilution (Invitrogen). Cells were counterstained with DAPI and visualized under epi/fluorescence microscope **(B)**.

Courtesy Dr. Taras Pasternak, Freiburg University, Germany