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Myobrevin

Cat.No. 176 003; Polyclonal rabbit antibody, 50 µg specific antibody (lyophilized)

Data Sheet

| Reconstitution/ Storage | 50 μg specific antibody, lyophilized. Affinity purified with the immunogen. Albumin was added for stabilization. For reconstitution add 50 μl H ₂ O to get a 1mg/ml solution in PBS. Then aliquot and store at -20°C to -80°C until use. Antibodies should be stored at +4°C when still lyophilized. Do not freeze! For detailed information, see back of the data sheet. |
|----------------------------|---|
| Applications | WB: 1: 1000 up to 1: 5000 (AP staining) IP: not tested yet ICC: 1: 1000 IHC: yes IHC_P: 1: 200 |
| Immunogen | Recombinant protein corresponding to AA 1 to 70 from mouse Myobrevin (UniProt Id: Q9Z2P8) |
| Reactivity | Reacts with: mouse (Q9Z2P8). No signal: rat. Other species not tested yet. |
| Specificity | K.O. PubMed: <u>29330887</u> |
| Matching control | 176-0P |

TO BE USED IN VITRO / FOR RESEARCH ONLY NOT TOXIC, NOT HAZARDOUS, NOT INFECTIOUS, NOT CONTAGIOUS

Background

Myobrevin, also known as **VAMP 5** belongs to the family of **v**esicle-**a**ssociated **m**embrane **p**roteins and has a theoretical molecular weight of 11.4 kDa. Like other VAMP isoforms it is composed of an N-terminal cytoplasmic region and a C-terminal transmembrane domain.

Vamp 5 is preferentially expressed in skeletal muscle and heart tissue and is upregulated during the differentiation of C2C12 cells into myotubes.

Selected References for 176 003

The localization of VAMP5 in skeletal and cardiac muscle.
Takahashi M, Tajika Y, Khairani AF, Ueno H, Murakami T, Yorifuji H
Histochemistry and cell biology (2013) 1394: 573-82. . WB, IHC

Vesicular transport system in myotubes: ultrastructural study and signposting with vesicle-associated membrane proteins. Taiika Y. Takahashi M. Khairani AF. Ueno H. Murakami T. Yorifuii H

Histochemistry and cell biology (2014) 1414: 441-54. . WB, ICC; tested species: mouse

Loss of VAMP5 in mice results in duplication of the ureter and insufficient expansion of the lung.

Ikezawa M. Tajika Y. Ueno H. Murakami T. Inoue N. Yorifuji H

Developmental dynamics: an official publication of the American Association of Anatomists (2018):.. WB, IHC; KO verified; tested species: mouse

Lysosomal exocytosis releases pathogenic a-synuclein species from neurons in synucleinopathy models.

Xie YX, Naseri NN, Fels J, Kharel P, Na Y, Lane D, Burré J, Sharma M

Nature communications (2022) 131: 4918. . WB; tested species: mouse

SNAP23 deficiency causes severe brain dysplasia through the loss of radial glial cell polarity.

Kunii M, Noguchi Y, Yoshimura SI, Kanda S, Iwano T, Avriyanti E, Atik N, Sato T, Sato K, Ogawa M, Harada A, et al.

The Journal of cell biology (2021) 2201: .. ICC; KD verified; tested species: mouse

Characterisation of GLUT4 trafficking in HeLa cells: comparable kinetics and orthologous trafficking mechanisms to 3T3-L1 adipocytes.

Morris S, Geoghegan ND, Sadler JBA, Koester AM, Black HL, Laub M, Miller L, Heffernan L, Simpson JC, Mastick CC, Cooper J, et al.

PeerJ (2020) 8: e8751.. WB; tested species: mouse

Cardiac SNARE Expression in Health and Disease.

Bowman PRT, Smith GL, Gould GW

Frontiers in endocrinology (2019) 10: 881.. WB; tested species: mouse

Characterization of VAMP isoforms in 3T3-L1 adipocytes: implications for GLUT4 trafficking.

Sadler JB, Bryant NJ, Gould GW

Molecular biology of the cell (2015) 263: 530-6. . WB

Effects of contraction on localization of GLUT4 and v-SNARE isoforms in rat skeletal muscle.

Rose AJ, Jeppesen J, Kiens B, Richter EA

American journal of physiology. Regulatory, integrative and comparative physiology (2009) 2975: R1228-37. . WB

Selected General References

VAMP5 and VAMP8 are most likely not involved in primary open-angle glaucoma. Brinkman JF, Ottenheim CP, de Jong LA, Zegers RH, de Smet MD, de Jong PT, Bergen AA Molecular vision (2005) 11: 582-6. .

The cytoplasmic domain of Vamp4 and Vamp5 is responsible for their correct subcellular targeting: the N-terminal extenSion of VAMP4 contains a dominant autonomous targeting signal for the trans-Golgi network.

Zeng Q, Tran TT, Tan HX, Hong W

The Journal of biological chemistry (2003) 27825: 23046-54. .

Access the online factsheet including applicable protocols at https://sysy.com/product/176003 or scan the QR-code.



FAQ - How should I store my antibody?

Shipping Conditions

 All our antibodies and control proteins / peptides are shipped lyophilized (vacuum freezedried) and are stable in this form without loss of quality at ambient temperatures for several weeks.

Storage of Sealed Vials after Delivery

- Unlabeled and biotin-labeled antibodies and control proteins should be stored at 4°C before reconstitution. They must not be stored in the freezer when still lyophilized!
 Temperatures below zero may cause loss of performance.
- Fluorescence-labeled antibodies should be reconstituted immediately upon receipt. Long term storage (several months) may lead to aggregation.
- **Control peptides** should be kept at -20°C before reconstitution.

Long Term Storage after Reconstitution (General Considerations)

- The storage freezer must not be of the frost-free variety ("no-frost freezer"). This cycle
 between freezing and thawing (to reduce frost-build-up), which is exactly what should be
 avoided. For the same reason, antibody vials should be placed in an area of the freezer that
 has minimal temperature fluctuations, for instance towards the back rather than on a door
 shelf.
- Aliquot the antibody and store frozen (-20°C to -80°C). Avoid very small aliquots (below 20 µl)
 and use the smallest storage vial or tube possible. The smaller the aliquot, the more the stock
 concentration is affected by evaporation and adsorption of the antibody to the surface of the
 storage vial or tube. Adsorption of the antibody to the surface leads to a substantial loss of
 activity.
- The addition of glycerol to a final concentration of 50% lowers the freezing point of your stock and keeps your antibody at -20°C in liquid state. This efficiently avoids freeze and thaw cycles.

Product Specific Hints for Storage

Control proteins / peptides

• Store at -20°C to -80°C.

Monoclonal Antibodies

- Ascites and hybridoma supernatant should be stored at -20°C up to -80°C. Prolonged storage at 4°C is not recommended! Unlike serum, ascites may contain proteases that will degrade the antibodies.
- **Purified IgG** should be stored at -20°C up to -80°C. Adding a carrier protein like BSA will increase long term stability. Many of our antibodies already contain carrier proteins. Please refer to the data-sheet for detailed information.

Polyclonal Antibodies

- Crude antisera: With anti-microbials added, they may be stored at 4°C. However, frozen storage (-20°C up to -80°C) is preferable.
- Affinity purified antibodies: Less robust than antisera. Storage at -20°C up to -80°C is
 recommended. Adding a carrier protein like BSA will increase long term stability. Most of our
 antibodies already contain carrier proteins. Please refer to the data-sheet for detailed
 information.

Fluorescence-labeled Antibodies

• Store as a liquid with 1:1 (v/v) glycerol at -20°C. Protect these antibodies from light exposure.

Avoid repeated freeze-thaw cycles for all antibodies!

FAQ - How should I reconstitute my antibody?

Reconstitution

- All our purified antibodies are lyophilized from PBS. To reconstitute the antibody in PBS, add
 the amount of deionized water given in the respective datasheet. If higher volumes are
 preferred, add water as mentioned above and then the desired amount of PBS and a
 stabilizing carrier protein (e.g. BSA) to a final concentration of 2%. Some of our antibodies
 already contain albumin. Take this into account when adding more carrier protein.
 For complete reconstitution, carefully remove the lid. After adding water, briefly vortex the
 solution. You can spin down the liquid by placing the vial into a 50 ml centrifugation tube filled
 with paper.
- If desired, add small amounts of azide or thimerosal to prevent microbial growth. This is especially recommended if you want to keep an aliquot a 4°C.
- After reconstitution of fluorescence-labeled antibodies, add 1:1 (v/v) glycerol to a final
 concentration of 50%. This lowers the freezing point of your stock and keeps your antibody in
 liquid state at -20°C.
- Glycerol may also be added to unlabeled primary antibodies. It is a suitable way to avoid freezethaw cycles.
- Please refer to our tips and hints for subsequent storage of reconstituted antibodies and control peptides and proteins.