

## **Product datasheet**

# PE Anti-Amyloid Precursor Protein antibody [EPR5118-34] ab306412

Recombinant RabMAb

### 1 Image

| Overview            |   |  |
|---------------------|---|--|
| Product name        | PE Anti-Amyloid Precursor Protein antibody [EPR5118-34]   |  |
| Description         | PE Rabbit monoclonal [EPR5118-34] to Amyloid Precursor Protein  |  |
| Host species        | Rabbit  |  |
| Conjugation         | PE. Ex: 488nm, Em: 575nm  |  |
| Tested applications | Suitable for: Target binding affinity, Antibody labelling   |  |
| Immunogen           | Synthetic peptide. This information is proprietary to Abcam and/or its suppliers.   |  |
| General notes       | This <b>conjugated primary antibody</b> is released using a quantitative quality control method that evaluates binding affinity post-conjugation and efficiency of antibody labeling.   |  |
|                     | For suitable applications and species reactivity, please refer to the unconjugated version of this clone. This conjugated antibody is eligible for Abtrial: learn more <b>here</b> .  |  |
|                     | This product is a recombinant monoclonal antibody, which offers several advantages including:<br>- High batch-to-batch consistency and reproducibility<br>- Improved sensitivity and specificity<br>- Long-term security of supply<br>- Animal-free production<br>For more information <u>see here</u> .<br>Our RabMAb <sup>®</sup> technology is a patented hybridoma-based technology for making rabbit<br>monoclonal antibodies. For details on our patents, please refer to <u>RabMAb<sup>®</sup> patents</u> . |  |

| Properties           |   |
|----------------------|---|
| Form                 | Liquid  |
| Storage instructions | Shipped at 4°C. Store at +4°C short term (1-2 weeks). Upon delivery aliquot. Store at +4°C. Store<br>In the Dark. |
| Storage buffer       | pH: 7.40<br>Preservative: 0.02% Sodium azide<br>Constituents: 98% PBS, 1% BSA                                     |
| Purity               | Protein A purified  |

| Clonality    | Monoclonal |
|--------------|------------|
| Clone number | EPR5118-34 |
| lsotype      | lgG        |

#### **Applications**

The Abpromise guarantee Our <u>Abpromise guarantee</u> covers the use of ab306412 in the following tested applications.

The application notes include recommended starting dilutions; optimal dilutions/concentrations should be determined by the end user.

| Application             | Abreviews | Notes                                    |
|-------------------------|-----------|--|
| Target binding affinity |           | Use at an assay dependent concentration. |
| Antibody labelling      |           | Use at an assay dependent concentration. |

#### Target

#### Function

Functions as a cell surface receptor and performs physiological functions on the surface of neurons relevant to neurite growth, neuronal adhesion and axonogenesis. Involved in cell mobility and transcription regulation through protein-protein interactions. Can promote transcription activation through binding to APBB1-KAT5 and inhibits Notch signaling through interaction with Numb. Couples to apoptosis-inducing pathways such as those mediated by G(O) and JIP. Inhibits G(o) alpha ATPase activity (By similarity). Acts as a kinesin I membrane receptor, mediating the axonal transport of beta-secretase and presenilin 1. Involved in copper homeostasis/oxidative stress through copper ion reduction. In vitro, copper-metallated APP induces neuronal death directly or is potentiated through Cu(2+)-mediated low-density lipoprotein oxidation. Can regulate neurite outgrowth through binding to components of the extracellular matrix such as heparin and collagen I and IV. The splice isoforms that contain the BPTI domain possess protease inhibitor activity. Induces a AGER-dependent pathway that involves activation of p38 MAPK, resulting in internalization of amyloid-beta peptide and leading to mitochondrial dysfunction in cultured cortical neurons. Provides Cu(2+) ions for GPC1 which are required for release of nitric oxide (NO) and subsequent degradation of the heparan sulfate chains on GPC1.

Beta-amyloid peptides are lipophilic metal chelators with metal-reducing activity. Bind transient metals such as copper, zinc and iron. In vitro, can reduce Cu(2+) and Fe(3+) to Cu(+) and Fe(2+), respectively. Beta-amyloid 42 is a more effective reductant than beta-amyloid 40. Beta-amyloid peptides bind to lipoproteins and apolipoproteins E and J in the CSF and to HDL particles in plasma, inhibiting metal-catalyzed oxidation of lipoproteins. Beta-APP42 may activate mononuclear phagocytes in the brain and elicit inflammatory responses. Promotes both tau aggregation and TPK II-mediated phosphorylation. Interaction with overexpressed HADH2 leads to oxidative stress and neurotoxicity. Also binds GPC1 in lipid rafts.

Appicans elicit adhesion of neural cells to the extracellular matrix and may regulate neurite outgrowth in the brain.

The gamma-CTF peptides as well as the caspase-cleaved peptides, including C31, are potent enhancers of neuronal apoptosis.

N-APP binds TNFRSF21 triggering caspase activation and degeneration of both neuronal cell bodies (via caspase-3) and axons (via caspase-6).

#### **Tissue specificity**

Expressed in all fetal tissues examined with highest levels in brain, kidney, heart and spleen. Weak expression in liver. In adult brain, highest expression found in the frontal lobe of the cortex

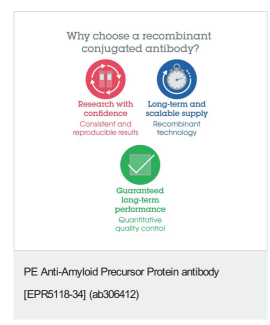
| Involvement in disease              | and in the anterior perisylvian cortex-opercular gyri. Moderate expression in the cerebellar cortex,<br>the posterior perisylvian cortex-opercular gyri and the temporal associated cortex. Weak<br>expression found in the striate, extra-striate and motor cortices. Expressed in cerebrospinal fluid,<br>and plasma. Isoform APP695 is the predominant form in neuronal tissue, isoform APP751 and<br>isoform APP770 are widely expressed in non-neuronal cells. Isoform APP751 is the most<br>abundant form in T-lymphocytes. Appican is expressed in astrocytes.<br>Alzheimer disease 1<br>Cerebral amyloid angiopathy, APP-related   |
|-------------------------------------|--|
| Sequence similarities               | Belongs to the APP family.<br>Contains 1 BPT/Kunitz inhibitor domain.  |
| Domain                              | The basolateral sorting signal (BaSS) is required for sorting of membrane proteins to the basolateral surface of epithelial cells.<br>The NPXY sequence motif found in many tyrosine-phosphorylated proteins is required for the specific binding of the PID domain. However, additional amino acids either N- or C-terminal to the NPXY motif are often required for complete interaction. The PID domain-containing proteins which bind APP require the YENPTY motif for full interaction. These interactions are independent of phosphorylation on the terminal tyrosine residue. The NPXY site is also involved in clathrin-mediated endocytosis.  |
| Post-translational<br>modifications | Proteolytically processed under normal cellular conditions. Cleavage either by alpha-secretase,<br>beta-secretase or theta-secretase leads to generation and extracellular release of soluble APP<br>peptides, S-APP-alpha and S-APP-beta, and the retention of corresponding membrane-<br>anchored C-terminal fragments, C80, C83 and C99. Subsequent processing of C80 and C83 by<br>gamma-secretase yields P3 peptides. This is the major secretory pathway and is non-<br>amyloidogenic. Alternatively, presenilin/nicastrin-mediated gamma-secretase processing of C99<br>releases the amyloid beta proteins, amyloid-beta 40 (Abeta40) and amyloid-beta 42 (Abeta42),<br>major components of amyloid plaques, and the cytotoxic C-terminal fragments, gamma-CTF(50),<br>gamma-CTF(57) and gamma-CTF(59). Many other minor beta-amyloid peptides, beta-amyloid 1-<br>X peptides, are found in cerebral spinal fluid (CSF) including the beta-amyloid X-15 peptides,<br>produced from the cleavage by alpha-secretase and all terminating at GIn-686.<br>Proteolytically cleaved by caspases during neuronal apoptosis. Cleavage at Asp-739 by either<br>caspase-6, -8 or -9 results in the production of the neurotoxic C31 peptide and the increased<br>production of beta-amyloid peptides.<br>N- and O-glycosylated. O-glycosylation on Ser and Thr residues with core 1 or possibly core 8<br>glycans. Partial tyrosine glycosylation (Tyr-681) is found on some minor, short beta-amyloid 38,<br>beta-amyloid 40 nor on beta-amyloid 42. Modification on a tyrosine is unusual and is more<br>prevelant in AD patients. Glycans had Neu5AcHex(Neu5AcHex(Neu5AcHex(Neu5Ac)HexNAc-O-<br>Tyr structures, where O-Ac is O-acetylation of Neu5Ac. Neu5AcNeu5AcHex(Neu5Ac)HexNAc-O-<br>Tyr structures, where O-Ac is O-acetylation of Neu5Ac. Neu5AcNeu5AcHex(Neu5Ac)HexNAc-O-<br>Tyr structures, where O-Ac is O-acetylation of Neu5Ac. Neu5AcNeu5Ac: is most likely Neu5Ac<br>2,8Neu5Ac linked. O-glycosylations in the vicinity of the cleavage sites may influence the<br>proteolytic processing. Appicans are L-APP isoforms with O-linked chondroitin sulfate.<br>Phosphorylation in the |

Cys-158, and the formation of a disulfide bond. In vitro, the APP-Cu(+) complex in the presence of hydrogen peroxide results in an increased production of beta-amyloid-containing peptides. Trophic-factor deprivation triggers the cleavage of surface APP by beta-secretase to release sAPP-beta which is further cleaved to release an N-terminal fragment of APP (N-APP). Beta-amyloid peptides are degraded by IDE.

#### **Cellular localization**

Membrane. Membrane, clathrin-coated pit. Cell surface protein that rapidly becomes internalized via clathrin-coated pits. During maturation, the immature APP (N-glycosylated in the endoplasmic reticulum) moves to the Golgi complex where complete maturation occurs (O-glycosylated and sulfated). After alpha-secretase cleavage, soluble APP is released into the extracellular space and the C-terminal is internalized to endosomes and lysosomes. Some APP accumulates in secretory transport vesicles leaving the late Golgi compartment and returns to the cell surface. Gamma-CTF(59) peptide is located to both the cytoplasm and nuclei of neurons. It can be translocated to the nucleus through association with APBB1 (Fe65). Beta-APP42 associates with FRPL1 at the cell surface and the complex is then rapidly internalized. APP sorts to the basolateral surface in epithelial cells. During neuronal differentiation, the Thr-743 phosphorylated form is located mainly in growth cones, moderately in neurites and sparingly in the cell body. Casein kinase phosphorylation can occur either at the cell surface or within a post-Golgi compartment. Associates with GPC1 in perinuclear compartments. Colocalizes with SORL1 in a vesicular pattern in cytoplasm and perinuclear regions.

#### Images



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