

ALS2 rabbit monoclonal antibody

Catalog # H00057679-K Size 100 ug x up to 3

| Specification | |
|-------------------------|---|
| Product Description | Rabbit monoclonal antibody raised against a human ALS2 peptide using ARM Technology. |
| Immunogen | A synthetic peptide of human ALS2 is used for rabbit immunization. Customer or Abnova will decide on the preferred peptide sequence. |
| Host | Rabbit |
| Library Construction | Non-fusion antibody library from rabbit spleen (ARM Technology). |
| Expression | Overexpression vector and transfection into 293H cell line. |
| Reactivity | Human |
| Purification | Protein A |
| Isotype | lgG |
| Quality Control Testing | Antibody reactive against human ALS2 peptide by ELISA and mammalian transfected lysate by Wes tern Blot. |
| Storage Buffer | In 1x PBS, pH 7.4 |
| Storage Instruction | Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing. |
| Deliverable | Up to three rabbit lgG clones of 100 ug each will be delivered to customer. |
| Note | Customer may provide cell or tissue lysate for antibody screening. Rabbit monoclonal antibody generated by ARM technology is amenable to antibody engineering in cluding F(ab)₂, lgG, scFv and different Fc and non-Fc conjugates per customer request. |

Applications

Western Blot (Transfected lysate)

Protocol Download



ELISA

| Gene Info — ALS2 | |
|---------------------|---|
| Entrez GenelD | <u>57679</u> |
| GeneBank Accession# | ALS2 |
| Gene Name | ALS2 |
| Gene Alias | ALS2CR6, ALSJ, FLJ31851, IAHSP, KIAA1563, MGC87187, PLSJ |
| Gene Description | amyotrophic lateral sclerosis 2 (juvenile) |
| Omim ID | <u>205100</u> <u>606352</u> <u>606353</u> <u>607225</u> |
| Gene Ontology | <u>Hyperlink</u> |
| Gene Summary | The protein encoded by this gene contains an ATS1/RCC1-like domain, a RhoGEF domain, and a vacuolar protein sorting 9 (VPS9) domain, all of which are guanine-nucleotide exchange factors that activate members of the Ras superfamily of GTPases. The protein functions as a guanine nucleotide exchange factor for the small GTPase RAB5. The protein localizes with RAB5 on early en dosomal compartments, and functions as a modulator for endosomal dynamics. Mutations in this gene result in several forms of juvenile lateral sclerosis and infantile-onset ascending spastic para lysis. Multiple transcript variants encoding different isoforms have been found for this gene. [provided by RefSeq |
| Other Designations | alsin |

Pathway

• Amyotrophic lateral sclerosis (ALS)

Disease

- Amyotrophic lateral sclerosis
- Genetic Predisposition to Disease
- Multiple Sclerosis
- Tobacco Use Disorder