



Rabbit Anti-TSPAN9/FITC Conjugated antibody

SL9448R-FITC

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| Product Name: | Anti-TSPAN9/FITC |
| Chinese Name: | FITC标记的四分子交联体9/四旋蛋白抗体 |
| Alias: | Tetraspanin 9; NET 5; NET5; PP1057; Tetraspan NET 5; Tetraspan NET-5; Tetraspanin-9; Transmembrane 4 superfamily member tetraspan NET 5; TSN9 HUMAN; Tspan-9; TSPAN9. |
| Organism Species: | Rabbit |
| Clonality: | Polyclonal |
| React Species: | Human,Mouse,Rat,Dog,Pig,Rabbit, |
| Applications: | IF=1:50-200 not yet tested in other applications. optimal dilutions/concentrations should be determined by the end user. |
| Molecular weight: | 27kDa |
| Cellular localization: | The cell membrane |
| Form: | Lyophilized or Liquid |
| Concentration: | 1mg/ml |
| immunogen: | KLH conjugated synthetic peptide derived from human TSPAN9 |
| Lsotype: | IgG |
| Purification: | affinity purified by Protein A |
| Storage Buffer: | 0.01M TBS(pH7.4) with 1% BSA, 0.03% Proclin300 and 50% Glycerol. |
| Storage: | Store at -20 °C for one year. Avoid repeated freeze/thaw cycles. The lyophilized antibody is stable at room temperature for at least one month and for greater than a year when kept at -20°C. When reconstituted in sterile pH 7.4 0.01M PBS or diluent of antibody the antibody is stable for at least two weeks at 2-4 °C. |
| Product Detail: | background: The tetraspanin family is a group of cell surface proteins that regulate cell development, activation, growth and motility. Each member contains four hydrophobic domains and participates in the mediation of signal transduction. NET-5, also known as TSPAN9 (tetraspanin 9), is a 239 amino acid multi-pass membrane protein that belongs to the tetraspanin (TM4SF) family. NET-5 forms a complex with GPVI in the tetraspanin microdomains on the platelet surface, and is encoded by a gene that maps to human |

chromosome 12p13.33. Chromosome 12 encodes over 1,100 genes and comprises approximately 4.5% of the human genome. Chromosome 12 is associated with a variety of diseases and afflictions, including hypochondrogenesis, achondrogenesis, Kniest dysplasia, Noonan syndrome and trisomy 12p, which causes facial developmental defects and seizure disorders.

Subunit:

Found in a complex with GP6.

Subcellular Location:

Membrane; Multi-pass membrane protein.

Tissue Specificity:

Expressed in megakaryocytes and platelets.

Similarity:

Belongs to the tetraspanin (TM4SF) family.

Database links:

[Entrez Gene: 10867](#)Human

[Omid: 613137](#)Human

[SwissProt: O75954](#)Human

[Unigene: 504517](#)Human

Important Note:

This product as supplied is intended for research use only, not for use in human, therapeutic or diagnostic applications.