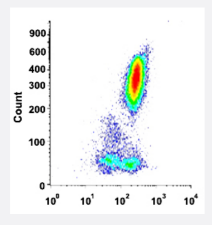


# CD59 monoclonal antibody, clone VJ1/12,2 (PerCP)

Catalog # MAB13957 Size 100 Reactions

## **Applications**



#### Flow Cytometry

Flow cytometric analysis of human normal whole blood with CD59 monoclonal antibody, clone VJ1/12,2 (PerCP) (Cat # MAB13957).

Specification	
Product Description	Mouse monoclonal antibody raised against human CD59.
Immunogen	TNF activated HUVEC cells.
Host	Mouse
Theoretical MW (kDa)	18-20
Reactivity	Human
Form	Liquid
Conjugation	PerCP
Purification	Protein A/G purification
Purity	>90%
Isotype	lgG2a



### **Product Information**

Recommend Usage	Flow Cytometry (20 uL/10 <sup>6</sup> cells) The optimal working dilution should be determined by the end user.
Storage Buffer	In PBS, pH 7.4 (protein stabilizer, 0.09% sodium azide).
Storage Instruction	Store in the dark at 4°C. Avoid prolonged exposure to light.
Note	This product contains sodium azide: a POISONOUS AND HAZARDOUS SUBSTANCE which shoul d be handled by trained staff only.

## **Applications**

Flow Cytometry

Flow cytometric analysis of human normal whole blood with CD59 monoclonal antibody, clone VJ1/12,2 (PerCP) (Cat # MAB13957).

Gene Info — CD59	
Entrez GenelD	<u>966</u>
Protein Accession#	<u>P13987</u>
Gene Name	CD59
Gene Alias	16.3A5, 1F5, EJ16, EJ30, EL32, FLJ38134, FLJ92039, G344, HRF-20, HRF20, MAC-IP, MACI F, MEM43, MGC2354, MIC11, MIN1, MIN2, MIN3, MIRL, MSK21, p18-20
Gene Description	CD59 molecule, complement regulatory protein
Omim ID	107271
Gene Ontology	<u>Hyperlink</u>
Gene Summary	This gene encodes a cell surface glycoprotein that regulates complement-mediated cell lysis, and it is involved in lymphocyte signal transduction. This protein is a potent inhibitor of the complement membrane attack complex, whereby it binds complement C8 and/or C9 during the assembly of this complex, thereby inhibiting the incorporation of multiple copies of C9 into the complex, which is necessary for osmolytic pore formation. This protein also plays a role in signal transduction pathways in the activation of T cells. Mutations in this gene cause CD59 deficiency, a disease resulting in hemolytic anemia and thrombosis, and which causes cerebral infarction. Multiple alternatively spliced transcript variants, which encode the same protein, have been identified for this gene. [provided by RefSeq
Other Designations	20 kDa homologous restriction factor CD59 antigen CD59 antigen p18-20 (antigen identified by



## Pathway

- Complement and coagulation cascades
- Hematopoietic cell lineage

### Disease

- Genetic Predisposition to Disease
- Lymphoma
- Macular Degeneration