

## **Product datasheet for TP318343**

## OriGene Technologies, Inc.

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## CD59 (NM\_000611) Human Recombinant Protein

**Product data:** 

**Product Type:** Recombinant Proteins

**Description:** Recombinant protein of human CD59 molecule, complement regulatory protein (CD59),

transcript variant 2, 20 µg

Species: Human
Expression Host: HEK293T

**Expression cDNA Clone** 

or AA Sequence:

>RC218343 protein sequence Red=Cloning site Green=Tags(s)

MGIQGGSVLFGLLLVLAVFCHSGHSLQCYNCPNPTADCKTAVNCSSDFDACLITKAGLQVYNKCWKFEHC

NFNDVTTRLRENELTYYCCKKDLCNFNEQLENGGTSLSEKTVLLLVTPFLAAAWSLHP

**TRTRPLEQKLISEEDLAANDILDYKDDDDKV** 

Tag: C-Myc/DDK
Predicted MW: 11.6 kDa

**Concentration:** >0.05 μg/μL as determined by microplate BCA method

**Purity:** > 80% as determined by SDS-PAGE and Coomassie blue staining

**Buffer:** 25 mM Tris-HCl, 100 mM glycine, pH 7.3, 10% glycerol

**Preparation:** Recombinant protein was captured through anti-DDK affinity column followed by

conventional chromatography steps.

**Note:** For testing in cell culture applications, please filter before use. Note that you may experience

some loss of protein during the filtration process.

Storage: Store at -80°C.

**Stability:** Stable for 12 months from the date of receipt of the product under proper storage and

handling conditions. Avoid repeated freeze-thaw cycles.

RefSeq: NP 000602

Locus ID: 966

UniProt ID: P13987

RefSeq Size: 7635





**Cytogenetics:** 11p13

RefSeg ORF: 384

**Synonyms:** 1F5; 16.3A5; EJ16; EJ30; EL32; G344; HRF-20; HRF20; MAC-IP; MACIF; MEM43; MIC11; MIN1;

MIN2; MIN3; MIRL; MSK21; p18-20

**Summary:** This gene encodes a cell surface glycoprotein that regulates complement-mediated cell lysis,

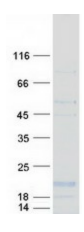
and it is involved in lymphocyte signal transduction. This protein is a potent inhibitor of the complement membrane attack complex, whereby it binds complement C8 and/or C9 during the assembly of this complex, thereby inhibiting the incorporation of multiple copies of C9 into the complex, which is necessary 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, Jul 2008]

**Protein Families:** Druggable Genome

**Protein Pathways:** Complement and coagulation cascades, Hematopoietic cell lineage

## **Product images:**



Coomassie blue staining of purified CD59 protein (Cat# TP318343). The protein was produced from HEK293T cells transfected with CD59 cDNA clone (Cat# [RC218343]) using MegaTran 2.0 (Cat# [TT210002]).