Catalog Number: PKSH030304



Note: Centrifuge before opening to ensure complete recovery of vial contents.

Description	
Synonyms	B1;Bp35;CD20;CVID5;LEU-16;MS4A1;MS4A2;S7
Species	Human
Expression Host	E.coli
Sequence	Ile 141-Ser 188
Accession	NP_068769.2
Calculated Molecular Weight	23.9 kDa
Tag	N-Trx
Properties	
Purity	> 80 % as determined by reducing SDS-PAGE.
Endotoxin	Please contact us for more information.
Storage	Generally, lyophilized proteins are stable for up to 12 months when stored at -20 to -80°C. Reconstituted protein solution can be stored at 4-8°C for 2-7 days. Aliquots of reconstituted samples are stable at < -20°C for 3 months.
Shipping	This product is provided as lyophilized powder which is shipped with ice packs.
Formulation	Lyophilized from sterile 50 mM Tris, 150 mM NaCl, 1 mM EDTA, pH 8.0 Normally 5 % - 8 % trehalose, mannitol and 0.01% Tween80 are added as protectants before lyophilization. Please refer to the specific buffer information in the printed manual.
Reconstitution	Please refer to the printed manual for detailed information.
Data	

KDa MK 116 66.2 45.0 35.0

> 25.0 18.4 14.4

> 80 % as determined by reducing SDS-PAGE.

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Background

CD20 (membrane-spanning 4-domains; subfamily A; member 1); also known as MS4A1; is a member of the membranespanning 4A gene family. Members of this nascent protein family are characterized by common structural features and similar intron/exon splice boundaries and display unique expression patterns among hematopoietic cells and nonlymphoid tissues. CD20 / MS4A1 is expressed on all stages of B cell development except the first and last. CD20 / MS4A1 is present from pre-pre B cells through memory cells; but not on either pro-B cells or plasma cells. It is a B-lymphocyte surface molecule which plays a role in the development and differentiation of B-cells into plasma cells. CD20 / MS4A1may be involved in the regulation of B-cell activation and proliferation. Defects in CD20 / MS4A1 are the cause of immunodeficiency common variable type 5(CVID5). CVID5 is a primary immunodeficiency characterized by antibody deficiency; hypogammaglobulinemia; recurrent bacterial infections and an inability to mount an antibody response to antigen. The defect results from a failure of B-cell differentiation and impaired secretion of immunoglobulins; the

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numbers of circulating B-cells is usually in the normal range; but can be low.

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