Catalog Number:PKSH031026



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

LAM Family Member 7;CD2 Subset 1;CD2-Like Receptor-Activating Cytotoxic ells;CRACC;Membrane Protein FOAP-12;Novel Ly9;Protein PA;CD319;SLAMF7;CS1;SLAM7 uman EK293 Cells et 1-Met 226 P_067004.3 8.8 kDa -His easured by its ability to bind biotinylated human SH2D1A-His in a functional LISA.
EK293 Cells et 1-Met 226 P_067004.3 8.8 kDa -His easured by its ability to bind biotinylated human SH2D1A-His in a functional LISA.
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LISA.
93 % as determined by reducing SDS-PAGE.
93 % as determined by reducing SDS-PAGE.
1.0 EU per μ g of the protein as determined by the LAL method.
enerally, lyophilized proteins are stable for up to 12 months when stored at -20 to 0°C. Reconstituted protein solution can be stored at 4-8°C for 2-7 days. Aliquots reconstituted samples are stable at < -20°C for 3 months.
nis product is provided as lyophilized powder which is shipped with ice packs.
yophilized from sterile PBS, pH 7.4 ormally 5 % - 8 % trehalose, mannitol and 0.01% Tween80 are added as otectants before lyophilization. ease refer to the specific buffer information in the printed manual.
ease refer to the printed manual for detailed information.

45.0 35.0 25.0 18.4 14.4

> 93 % as determined by reducing SDS-PAGE.

Background

SLAM family member 7 (SLAMF7), also known as CRACC, CD319, CD2-like receptor-activating cytotoxic cells, and CS1, is a single-pass type I membrane protein and a member of the CD2 family of cell surface receptors. SLAMF7 is expressed in NK cells, activated B-cells, NK-cell line but not in promyelocytic, B-cell lines, or T-cell lines. Although the cytoplasmic domain of CS1 contains immunoreceptor tyrosine-based switch motifs (ITSM), which enables to recruite signaling lymphocyte activation molecule (SLAM)-associated protein (SAP/SH2D1A), it activates NK cells in the absence

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of a functional SAP. CS1 is a self ligand and homophilic interaction of CS1 regulates NK cell cytolytic activity. CRACC positively regulated natural killer cell functions by a mechanism dependent on the adaptor EAT-2 but not the related adaptor SAP. However, in the absence of EAT-2, CRACC potently inhibited natural killer cell function. It was also inhibitory in T cells, which are typically devoid of EAT-2. Thus, CRACC can exert activating or inhibitory influences on cells of the immune system depending on cellular context and the availability of effector proteins.

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