Species Reactivity: Mouse

Specificity: Detects mouse MGL1/CD301a in direct ELISAs and Western blots. In direct ELISAs and Western blots, less than 1% cross-reactivity with recombinant mouse MGL2 is observed.

Source: Polyclonal Goat IgG

Purification: Antigen Affinity-purified

Immunogen: Mouse myeloma cell line NS0-derived recombinant mouse MGL1/CD301a Gln57-Ser304

Accession #: AAH14811

Conjugate: Allophycocyanin

Excitation Wavelength: 620-650 nm

Emission Wavelength: 660-670 nm

Formulation: Supplied in a saline solution containing BSA and Sodium Azide. See Certificate of Analysis for details.

*Contains <0.1% Sodium Azide, which is not hazardous at this concentration according to GHS classifications. Refer to the Safety Data Sheet (SDS) for additional information and handling instructions.

APPLICATIONS

Please Note: Optimal dilutions should be determined by each laboratory for each application. General Protocols are available in the Technical Information section on our website.

Recomended Concentration | Sample
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10 µL/10⁶ cells | See Below

DATA

Detection of MGL1/CD301a in RAW 264.7 Mouse Cell Line by Flow Cytometry.

RAW 264.7 mouse macrophage cell line was stained with Goat Anti-Mouse MGL1/CD301a APC-conjugated Antibody (Catalog # FAB4938A, filled histogram) or isotype control antibody (Catalog # IC108A, open histogram). View our protocol for Staining Membrane-associated Proteins.

PREPARATION AND STORAGE

Shipping: The product is shipped with polar packs. Upon receipt, store it immediately at the temperature recommended below.

Stability & Storage: Protect from light. Do not freeze.

- 12 months from date of receipt, 2 to 8 °C as supplied.
Mouse MGL1 (macrophage galactose N-acetyl-galactosamine (GalNAc) specific Lectin 1, CD301a), also called ASGP-BP (asialoglycoprotein binding protein), is a 38 kDa type II transmembrane glycoprotein of the C-type lectin family (1). Two MGL proteins are encoded by separate genes in the mouse, but share 91% amino acid (aa) identity in the extracellular domain (ECD) (2). Only one MGL occurs in human and rat, and this is more structurally similar to mouse MGL1 than MGL2. However, mouse MGL1 binds Lewis X, in contrast to human MGL and mouse MGL2 which both bind specifically to terminal GalNAc residues (2). Lewis X is a trisaccharide commonly found on leukocytes and some tumor cells. Both mouse MGL proteins are expressed on immature dendritic cells. Mouse MGL1 and MGL2 are markers for connective tissue macrophages of a type termed alternately activated macrophages. These macrophages are induced by IL-4 that is produced during Th2-mediated inflammatory responses to parasitic infections or allergic airway inflammation (3, 4). Quantitative RT-PCR after helminth infection shows a peak of MGL1 expression at 7 days, while MGL2 shows increasing expression for at least 29 days (3). This, and data from MGL1 knockout mice (5), indicates that MGL1 is critical during the formation of granulation tissue, with MGL2 remaining involved during chronic infection. Mouse MGL1 is synthesized with an N-terminal 35 aa cytoplasmic region, a 21 aa transmembrane segment and a 248 aa ECD. The ECD contains one 129 aa carbohydrate recognition domain (CRD) that shows 78% and 63% aa identity with rat and human MGL, respectively.

References: