

Product Data Sheet

Anti-Ly-49C/F/H/I Antibody-FITC labled

Catalog # Source Reactivity Applications

CFE8993 Syrian M IF, FC

Hamster

Description Syrian Hamster monoclonal antibody FITC labled to Ly-49C/F/H/I

Immunogen IL-2-activated killer cells (LAK) from C57BL/6 mice

Purification The antibody was purified by affinity chromatography.

Specificity Recognizes mouse Ly-49C/F/H/I

Clonality Monoclonal (clone: 14B11)

Conjugation FITC

Form Syrian Hamster IgG. Liquid in PBS, pH 7.3, and 0.02% sodium azide.

Dilution 10 μl / assay

Gene Symbol Klra3; Klra6; Klra8; Klra9

Alternative Names Ly-49c; Ly49C; Killer cell lectin-like receptor 3; 5E6; Lymphocyte antigen 49c; Ly-49c;

Nk2.1; T-cell surface glycoprotein Ly-49C; Ly-49f; Ly49-f; Ly49F; Killer cell lectin-like

receptor 6; Lymphocyte antigen 49f; Ly-49f; T-cell surface glycoprotein Ly-49F;

Ly-49h; Ly49-h; Ly49H; Killer cell lectin-like receptor 8; Lymphocyte antigen 49h;

Ly-49h; T-cell surface glycoprotein Ly-49H; Killer cell lectin-like receptor subfamily A

member 9; Killer cell lectin-like receptor subfamily A member 9; Natural killer cell

recptor Ly49C

Entrez Gene 16634, 16637, 16639, 16640 (Mouse)

SwissProt Q64329, Q60653, Q60682, Q2TJJ8 (Mouse)

Directions for Use 1. Take 100 μl peripheral blood anticoagulated by EDTA and add to the bottom of 5

ml tube.

Application key: E- ELISA, WB- Western blot, IH- Immunohistochemistry, IF- Immunofluorescence, FC- Flow cytometry, IC-Immunocytochemistry, IP- Immunoprecipitation, ChIP- Chromatin Immunoprecipitation, EMSA- Electrophoretic Mobility Shift Assay, BL- Blocking, SE- Sandwich ELISA, CBE- Cell-based ELISA, RNAi- RNA interference

Species reactivity key: H- Human, M- Mouse, R- Rat, B- Bovine, C- Chicken, D- Dog, G- Goat, Mk- Monkey, P- Pig, Rb-Rabbit, S- Sheep, Z- Zebrafish

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- 2. Add 10 µl labeled antibody to the bottom of flow tube mixing with the whole blood, incubate for 20 minutes at room temperature away from light.
- 3. Add 2 ml RBC lysis buffer, incubate for 10 minutes away from light after mixing, dissolve red blood cells.
- 4. Sample tube is set to 1000 rpm centrifugation for 5 minutes, discard the supernatant.
- 5. Add 2 ml PBS wash buffer to resuspend the cells, then 1000 rpm centrifugation for 5 minutes, discard the supernatant.
- 6. Add 0.5 ml PBS wash buffer to resuspend the cells and detect by flow cytometry (sample should be determined on the day on the machine and can also be added fixation overnight at 4 °C then measured).

Storage/Stability

Shipped and store at 4°C for one year. Do not freeze.

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