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Product datasheet for CL121FX

MHC Class II RT1D Mouse Monoclonal Antibody [Clone ID: OX-17]

Product data:

Product Type:	Primary Antibodies
Clone Name:	OX-17
Applications:	FC
Recommended Dilution:	Flow Cytometry.
Reactivity:	Rat
Host:	Mouse
lsotype:	lgG1
Clonality:	Monoclonal
Immunogen:	Rat spleen membrane glycoproteins depleted of Ia-A antigens. Immunocyte Donor: BALB/c spleen Fusion Partner: X63 Ag8.653
Specificity:	This monoclonal antibody recognizes a monomorphic determinant on the α chain of the rat la antigen and appears to be the rat homologue of mouse la-E. It recognizes the rat la product present on B, but not T cells from lymph node or thoracic duct lymph. It does not bind to thymocytes or erythrocytes. The antibody does not cross-react with rat la-A or mouse la-E antigen, but rabbit antibody raised against the antibody affinity column-purified MRC OX- 17 antigen cross-reacted on tissues of mice expressing la-E mouse antigen but not on those mouse strains that were la-E antigen negative.
Formulation:	PBS, 0.02% NaN3 and EIA grade BSA as a stabilizing protein to bring total protein concentration to 4-5 mg/ml Label: FITC State: Liquid purified Ig
Concentration:	lot specific
Purification:	Protein G Chromatography
Conjugation:	FITC
Storage:	Store the antibody undiluted at 2-8°C for one month or (in aliquots) at -20°C for longer. Avoid repeated freezing and thawing. This antibody is photosensitive and should protected from light.
Stability:	Shelf life: one year from despatch.



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Synonyms:

HLA Class II

Note:

TEX Club.

Protocol: FLOW CYTOMETRY ANALYSIS:

Method:

1. Prepare a cell suspension in media A. For cell preparations, deplete the red blood cell population with Lympholyte®-Rat Cell Separation Medium.

2. Wash 2 times.

3. Resuspend the cells to a concentration of 2x10e7 cells/ml in media A. Add 50 µl of this suspension to each tube (each tube will then contain 1 x 10e6 cells, representing 1 test). 4. To each tube, add 0.05-0.1 µg* of this Ab per 10e6 cells.

5. Vortex the tubes to ensure thorough mixing of antibody and cells.

6. Incubate the tubes for 30 minutes at 4°C. (It is recommended that the tubes are protected from light, since most fluorochromes are light sensitive.)

7. Wash 2 times at 4°C.

8. Resuspend the cell pellet in 50 μ l ice cold media B.

9. Transfer to suitable tubes for flow cytometric analysis containing 15 μ l of propidium iodide at 0.5 mg/ml in PBS. This stains dead cells by intercalating in DNA.

Media:

A. Phosphate buffered saline (pH 7.2) + 5% normal serum of host species + sodium azide (100 μ l of 2M sodium azide in 100 mls).

B. Phosphate buffered saline (pH 7.2) + 0.5% Bovine serum albumin + sodium azide (100 μ l of 2M sodium azide in 100 mls).

Results - Tissue Distribution:

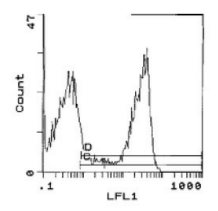
<u>Rat Strain</u>: Fischer <u>Cell Concentration</u>: 1x10e6 cells per test <u>Antibody Concentration Used</u>: 0.1 µg/10e6 cells <u>Isotypic Control</u>: FITC Mouse IgG1

Cell Source Percentage of cells stained above control:

Thymus 10.2% Spleen 49.0% Lymph Node 27.9%

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Product images:



Cell Source: Spleen Percentage of cells stained above control: 49.0%

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