

# Monoclonal Antibody to MHC Class II (I-Ek) - FITC

Alternate names: MHC class II antigen I-Ek

Catalog No.: CL074F
Quantity: 0.1 mg
Concentration: 0.1 mg/ml
Host / Isotype: Mouse / IgG2a

Clone: 14-4-4S Immunogen: C3H

Donor: C3H.SW

Fusion Partner: SP2/0-Ag 14

Format: State: Liquid purified

**Purification:** Protein G Chromatography

Buffer System: PBS, 0.02% NaN3 and EIA grade BSA as a stabilizing protein to bring total

protein concentration to 4-5 mg/ml.

Label: FITC

**Applications:** Flow Cytometry.

Other applications not tested. Optimal dilutions are dependent on conditions and should

be determined by the user.

Specificity: This monoclonal antibody is specific for cells expressing the la antigen coded for by the Ea

subregion. The reaction pattern of this antibody with a panel of standard and recombinant haplotypes demonstrates that this antibody reacts with antigen Ia.m7, which is expressed by all haplotypes except b,f,q and s. This antibody can be used to quantitate or eliminate cells bearing the Ia.m7 antigen and is well suited for identifying Ia cell populations of

positive mouse strains.

Species: Mouse.

Other species not tested.

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 product is photosensitive and should protected from light.

Shelf life: one year from despatch.

General References: 1. Ozato, K., Mayer, N., and Sachs, D.H. 1980. Hybridoma cell line secreting monoclonal

antibodies to mouse H-2 and Ia Antigens. J. Immunol.: 214:533-540.

**Protocols:** FLOW CYTOMETRY ANALYSIS:

Method:

1. Prepare a cell suspension in media A. For cell preparations, deplete the red blood cell population with Lympholyte®-M cell separation medium.

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Material Safety Datasheets are available at www.acris-antibodies.com or on request.



- 2. Wash 2 times.
- 3. Resuspend the cells to a concentration of 2x10e7 cells/ml in media A. Add 50  $\mu$ l of this suspension to each tube (each tube will then contain 1 x 10e6 cells, representing 1 test).
- 4. To each tube, add  $0.2 0.5 \mu g^*$  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 recommmended that the tubes are protected from light, since most flurochromes are light sensitive.)
- 7. Wash 2 times at 4°C.
- 8. Resuspend the cell pellet in 50 µl ice cold media B.
- 9. Transfer to suitable tubes for flow cytometric analysis containing 15  $\mu$ 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  $\mu$ l of 2M sodium azide in 100 mls).

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

### Results - Tissue Distribution by Flow Cytometry Analysis:

Mouse Strain: CBA/J

<u>Cell Concentration</u>: 1x10e6 cells per tests <u>Antibody Concentration Used</u>: 0.2 μg/10e6 cells

Isotypic Control: FITC Mouse IgG2a

## Cell Source Percentage of cells stained above control:

Spleen: 53.5% see FIGURE 1

Lymph Node: 26.6% Bone Marrow: 38.9%

## Strain Distribution by Flow Cytometry Analysis:

Procedure: As above

Antibody Concentration: 0.2 µg/106 cells

Strains Tested: see FIGURE 2; For a more detailed strain distribution - see reference 1.

#### **Pictures:**

<u>Strain</u>	H-2 Loci Alleles	+/-
	$\underline{K} \underline{A}_{\beta} \underline{A}_{\alpha} \underline{E}_{\beta} \underline{E}_{\alpha} \underline{C4} \underline{C4S} \underline{D}$	
A.TH	s s s s s s d	-
B10.A(3R)	bbb/kkddd	+
AKR	k k k k k k k	+
C3H/He	k k k k k k k	+
A.TL	s k k k k k d	+
C57BL/6	b b b b b b b	-
BALB/c	dddddddd	+

Figure 2





