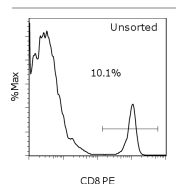
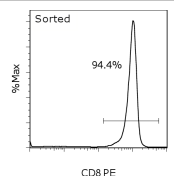


MagniSort™ Mouse CD8 Positive Selection Kit

Catalog Number: 8802-6842

RUO: For Research Use Only. Not for use in diagnostic procedures.





Mouse splenocytes were unsorted (left) or sorted with the MagniSort® Mouse CD8 Positive Selection Kit (right) then stained with Anti-Mouse CD8b FITC (cat. 11-0083). Total viable cells were used for analysis.

Product Information

Contents: MagniSort™ Mouse CD8 Positive

Selection Kit

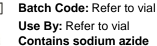
REF Catalog Number: 8802-6842

Handling Conditions: For sorting sterile cells, perform all steps in the hood.



Formulation: aqueous buffer, 0.09% sodium azide, may contain carrier protein/stabilizer **Temperature Limitation:** Store at 2-8°C. Do not







Description

The MagniSort® Mouse CD8 Positive Selection Kit is designed for the magnetic separation of CD8+ cells by positive selection. It has been optimized for the isolation of CD8+ cells from mouse spleens or lymph nodes utilizing a biotinylated Anti-Mouse CD8a antibody and streptavidin-coated magnetic beads. CD8+ cells are bound by antibody and then magnetic beads. When placed in a magnetic field, the undesired cells can be separated from CD8+ cells by decanting.

After positive selection, the purity of selected cells can be verified by staining with Anti-Mouse CD8a, clone 53-6.7.

Components

MagniSort® Anti-Mouse CD8 Biotin (cat. MS13-0081): 200 tests, 20 μ L/test; store at 2-8°C. MagniSort® Positive Selection Beads B (cat. PB-6004): 4 mL; store at 2-8°C.

Applications Reported

The MagniSort® Mouse CD8 Positive Selection Kit has been reported for use in magnetic cell separation.

Applications Tested

The MagniSort® Mouse CD8 Positive Selection Kit has been tested by magnetic cell separation followed by flow cytometric analysis of cells from mouse secondary lymphoid tissues. A test is defined as the amount of antibody or beads to be used to stain $1x10^7$ cells in $100 \, \mu L$.

This MagniSort® kit can sort 2x109 total cells.

Related Products

01-1234 123count™ eBeads Counting Beads 11-0083 eBioscience™ Anti-Mouse CD8b FITC (eBioH35-17.2 (H35-17.2)) MAG-4902 MagniSort™ Magnet

Not for further distribution without written consent.

invitrogen

MagniSort™ Positive Selection Protocol

Introduction

The following protocol is a general guideline for the MagniSort[™] Positive Selection Kits. In positive selection, desired cells are labeled with biotinylated antibodies, followed by streptavidin-coated magnetic beads. When cells are placed in the MagniSort[™] magnet, the desired cells will be held in place by the magnetic field while the undesired cells remain free in solution and can be removed by decanting. For each kit, the biotinylated antibody and the magnetic beads have been pre-titrated and diluted to test size.

General Notes

Caution

The MagniSort™ Magnet, 5 mL, generates a strong magnetic field. Keep away from pacemakers, credit cards, magnetic I.D. cards, watches, computer monitors and hard disks to prevent damage to these devices.

Cell preparation

- 1. The MagniSort™ Positive Selection Kits have been optimized for use with single-cell suspensions of either mouse secondary lymphoid organs or normal human peripheral blood mononuclear cells, unless otherwise noted.
- 2. For mouse cells, removal of debris by passing through a 40 µm nylon filter is recommended for optimal performance of the kits.
- 3. For preparation of normal human peripheral blood mononuclear cells, please refer to Best Protocols: Protocol D: Isolation of PBMC from whole blood located under the Resources Tab online. It is recommended to thoroughly wash the buffy coat cells to remove platelets for optimal performance in the MagniSort™ kits.
- 4. Addition of EDTA to buffers will reduce cell clumping.

Use in sterile cultures

- 1. MagniSortTM Biotin Antibody and Positive Selection Beads contain small amounts of sodium azide as preservative. This does not interfere with cellular functions when used in conjunction with sterile buffers that do not contain sodium azide. Performance in a given assay should be determined empirically.
- 2. For sorting sterile cells, perform all steps in a hood and use sterile polystyrene tubes with caps and sterile buffers.

Protocol:

Materials Provided

- MagniSortTM Positive Selection Antibody, 200 tests, 20 μL/test. Store at 2-8°C.
- MagniSort™ Positive Selection Beads, 4 mL. Store at 2-8°C.

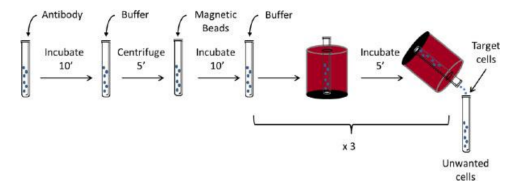
Additional Materials Required

- Recommended buffer for cell separation: PBS or HBSS supplemented with 3% FBS and 10 mM EDTA. Store at 2-8°C. **Note:** We do not recommend the use of tissue culture media, such as RPMI-1640 or DMEM, for use during cell separation.
- MagniSortTM Magnet, 5 mL
- 12 x 75 mm round bottom polystyrene tubes (5 mL, BD Falcon, cat no. 352008, or equivalent)

Experiment Duration

- 40 minutes
- Work flow:





Experimental Procedure

- 1. Prepare a single-cell suspension of lymphocytes at a concentration of 1x10⁷ cells/100 μL (1x10⁸/mL) in desired cell separation buffer. **Note:** Cells must be in a single-cell suspension. Inspect sample and pulse vortex or pipet to remove clumps, if necessary, before proceeding.
- 2. Place desired number of cells, but no more than 2x108 cells, in a 12 x 75 mm, 5 mL tube.
- 3. Add 20 μ L of MagniSortTM Positive Selection Antibody per 100 μ L of cells. Mix well by pulse vortexing 5 times. Incubate at room temperature for 10 minutes.
- 4. Wash cells by bringing the volume up to 4 mL with desired cell separation buffer and then centrifuge at 300 x g for 5 minutes.
- $5. \ \ Discard \ the \ supernatant \ and \ thoroughly \ resuspend \ the \ cells \ to \ their \ original \ volume \ with \ desired \ cell \ separation \ buffer.$
 - **Note:** Cells must be in a single-cell suspension. Inspect sample and pulse vortex or pipet to remove clumps, if necessary, before proceeding.
- 6. Add 20 μ L of MagniSortTM Positive Selection Beads per 100 μ L of cells. Mix well by pulse vortexing 5 times. Incubate at room temperature for 10 minutes.
 - **Note:** The MagniSortTM Positive Selection Beads must be uniformly resuspended before adding to cells to ensure optimal performance. Thoroughly resuspend the beads by pipetting up and down 5 times with a P1000 pipette set to 1 mL or by vortexing.
- 7. Bring the volume up to 2.5 mL with desired cell separation buffer. Mix by pipetting up and down 3 times with a P1000 pipette set to 1 mL. Avoid vortexing.
- 8. Insert the tube into the magnet until the bottom of the tube is touching the bench top through the hole in the bottom of the magnet. Incubate at room temperature for 5 minutes.
- 9. Pick up the magnet and in a continuous motion pour the supernatant into a waste or secondary receptacle; these are the undesired (unbound) cells. Hold the inverted tube for 1 second and then return it to the upright position.
 - **Note:** Do not blot or shake the inverted tube as this may reduce the recovery rate. The unbound cells can be collected and pooled, if needed.
- 10. Remove the tube from the magnet and repeat Steps 7-9 two more times for a total of 3 washes.
- 11. Remove the tube containing target cells from the magnet and add 1 mL of desired cell separation buffer. Wash the sides of the tube by pipetting the buffer down the sides. The positively selected cells are ready to use.

Documentation and support

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 - Certificates of Analysis
 - Safety Data Sheets (SDSs; also known as MSDSs)

Note: For SDSs for reagents and chemicals from other manufacturers, contact the manufacturer.

Limited product warranty

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