

CD45 MicroBeads

mouse

Order no. 130-052-301

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1. Description

This product is for research use only.

Components 2 mL CD45 MicroBeads, mouse:

MicroBeads conjugated to monoclonal antimouse CD45 antibodies (isotype: rat IgG2b;

clone: 30F11.1).

Capacity For 2×10^9 total cells, up to 200 separations.

Product format CD45 MicroBeads, mouse are supplied in buffer

containing stabilizer and 0.05% sodium azide.

Storage Store protected from light at +2 to +8 °C. Do not

freeze. The expiration date is indicated on the

vial label.

1.1 Principle of the MACS Separation

First, the CD45⁺ cells are magnetically labeled with CD45 MicroBeads, mouse. Then, the cell suspension is loaded onto a MACS Column, which is placed in the magnetic field of a MACS Separator. The magnetically labeled CD45⁺ cells are retained within the column. The unlabeled cells run through; this cell fraction is thus depleted of CD45⁺ cells. After removing the column from the magnetic field, the magnetically retained CD45⁺ cells can be eluted as the positively selected cell fraction. To increase the purity, the positively selected cell fraction containing the CD45⁺ cells must be separated over a second column.

1.2 Background information

CD45 MicroBeads, mouse are used for the positive selection or depletion of leukocytes from lymphoid and non-lymphoid tissues. The CD45 antigen is expressed on all cells of hematopoietic origin except erythrocytes and platelets.

1.3 Applications

- Positive selection or depletion of cells expressing mouse CD45 antigen
- Isolation or depletion of CD45⁺ leukocytes from spleen, lymph nodes, thymus, bone marrow, peripheral blood, body fluids or non-hematopoietic tissue
- Enrichment of human cells from bone marrow of chimeric mice by depletion of mouse leukocytes in combination with Anti-Ter-119 MicroBeads, mouse (# 130-049-901)

1.4 Reagent and instrument requirements

- Buffer: Prepare a solution containing phosphate-buffered saline (PBS), pH 7.2, 0.5% bovine serum albumin (BSA), and 2 mM EDTA by diluting MACS BSA Stock Solution (#130-091-376) 1:20 with autoMACS Rinsing Solution (#130-091-222). Keep buffer cold (+2 to +8 °C). Degas buffer before use, as air bubbles could block the column.
 - ▲ Note: EDTA can be replaced by other supplements such as anticoagulant citrate dextrose formula-A (ACD-A) or citrate phosphate dextrose (CPD). BSA can be replaced by other proteins such as mouse serum albumin, mouse serum, or fetal bovine serum (FBS). Buffers or media containing Ca²⁺ or Mg²⁺ are not recommended for use.
- MACS Columns and MACS Separators: CD45⁺ cells can be enriched by using MS, LS, or XS Columns or depleted with the use of LD or D Columns. Cells which strongly express the CD45 antigen can also be depleted using MS, LS, or XS Columns. Positive selection or depletion can also be performed by using the MultiMACS™ Cell24 Separator Plus or autoMACS Columns on the autoMACS NEO or autoMACS Pro Separators.

Column	Max. number of labeled cells	Max. number of total cells	Separator			
Positive se	lection					
MS	10 ⁷	2×10 ⁸	MiniMACS, OctoMACS, SuperMACS II			
LS	10 ⁸	2×10 ⁹	MidiMACS, QuadroMACS, SuperMACS II			
	10 ⁸	10 ⁹	MultiMACS Cell24 Separator Plus			
XS	10 ⁹	2×10 ¹⁰	SuperMACS II			
Depletion						
LD	10 ⁸	5×10 ⁸	MidiMACS, QuadroMACS, SuperMACS II			
D	10 ⁹		SuperMACS II			
Positive selection or depletion						
autoMACS	2×10 ⁸	4×10 ⁹	autoMACS NEO Separator, autoMACS Pro Separator			

Multi-24	10 ⁸	10 ⁹	MultiMACS Cell24
Column			Separator Plus
Block (per			
column)			

- ▲ Note: Column adapters are required to insert certain columns into the SuperMACS" II Separators. For details refer to the respective MACS Separator data sheet.
- ▲ Note: If separating with LS or LD Columns and the MultiMACS Cell24 Separator Plus use the Single-Column Adapter. Refer to the user manual for details.
- (Optional) Fluorochrome-conjugated antibodies for flow cytometric analysis, e.g., CD45 Antibody, anti-mouse, FITC. For more information about antibodies refer to www.miltenyibiotec.com/antibodies.
- (Optional) Propidium Iodide Solution (#130-093-233) or 7-AAD Staining Solution (#130-111-568) for flow cytometric exclusion of dead cells.
- (Optional) Dead Cell Removal Kit (#130-090-101) for the depletion of dead cells.
- (Optional) Pre-Separation Filters (30 μ m) (# 130-041-407) to remove cell clumps.

2. Protocol

2.1 Sample preparation

When working with anticoagulated peripheral blood or buffy coat, peripheral blood mononuclear cells (PBMCs) should be isolated by using a MACS PBMC Isolation Kit or by density gradient centrifugation, for example, using Ficoll-Paque™.

▲ Note: To remove platelets after density gradient separation, resuspend cell pellet in buffer and centrifuge at 200×g for 10−15 minutes at +20 °C. Carefully aspirate supernatant. Repeat washing step.

For details refer to the data sheet or the protocols section at www.miltenyibiotec.com/protocols.

When working with tissues, prepare a single-cell suspension using standard methods.

▲ Dead cells may bind non-specifically to MACS MicroBeads. To remove dead cells, it is recommended to use density gradient centrifugation or the Dead Cell Removal Kit (# 130-090-101).



2.2 Magnetic labeling

- ▲ Cells can be labeled with MACS MicroBeads using the autolabeling function of the autoMACS NEO or autoMACS Pro Separators. For more information refer to section 2.4.
- ▲ Work fast, keep cells cold, and use pre-cooled solutions. This will prevent capping of antibodies on the cell surface and non-specific cell labeling. *
- \blacktriangle For optimal performance it is important to obtain a single-cell suspension before magnetic labeling. Pass cells through 30 µm nylon mesh (Pre-Separation Filters (30 µm), #130-041-407) to remove cell clumps which may clog the column. Moisten filter with buffer before use.
- ▲ Volumes for magnetic labeling given below are for up to 10^7 total cells. When working with fewer than 10^7 cells, use the same volumes as indicated. When working with higher cell numbers, scale up all reagent volumes and total volumes accordingly (e.g. for 2×10^7 total cells, use twice the volume of all indicated reagent

volumes and total volumes).

- ightharpoonup The recommended incubation temperature is +2 to +8 °C. Higher temperatures and/or longer incubation times may lead to non-specific cell labeling. Working on ice may require increased incubation times.
- Determine cell number.
- Centrifuge cell suspension at 300×g for 10 minutes. Aspirate supernatant completely.
- 3. Resuspend cell pellet in 90 μ L of buffer per 10⁷ total cells.
- 4. Add 10 μL of CD45 MicroBeads, mouse per 10⁷ total cells.
- Mix well and incubate for 15 minutes in the refrigerator (+2 to +8 °C).
- 6. (Optional) Add staining antibodies, e.g., CD45 Antibody, antimouse, FITC according to manufacturer's recommendation.
- 7. Wash cells by adding 1-2 mL of buffer per 10^7 cells and centrifuge at $300\times g$ for 10 minutes. Aspirate supernatant completely.
- 8. Resuspend up to 10^8 cells in 500 µL of buffer.
 - ▲ Note: For higher cell numbers, scale up buffer volume accordingly.
 - \blacktriangle Note: For depletion with LD Columns, resuspend up to $1.25{\times}10^8$ cells in $500~\mu L$ of buffer.
- 9. Proceed to magnetic separation (2.3).



2.3 Magnetic separation

- ▲ Choose an appropriate MACS Column and MACS Separator according to the number of total cells and the number of CD45⁺ cells. For details refer to the table in section 1.4.
- ▲ Always wait until the column reservoir is empty before proceeding to the next step.

Magnetic separation with MS or LS Columns

- Place column in the magnetic field of a suitable MACS Separator. For details refer to the respective MACS Column data sheet.
- Prepare column by rinsing with the appropriate amount of buffer:

MS:
$$500 \mu L$$
 LS: $3 mL$

- Apply cell suspension onto the column. Collect flow-through containing unlabeled cells.
- Wash column with the appropriate amount of buffer. Collect unlabeled cells that pass through and combine with the flowthrough from step 3.

MS:
$$3\times500~\mu L$$
 LS: $3\times3~mL$

- \blacktriangle Note: Perform washing steps by adding buffer aliquots as soon as the column reservoir is empty.
- Remove column from the separator and place it on a suitable collection tube.
- 6. Pipette the appropriate amount of buffer onto the column. Immediately flush out the magnetically labeled cells by firmly pushing the plunger into the column.

 (Optional) To increase the purity of CD45⁺ cells, the eluted fraction can be enriched over a second MS or LS Column. Repeat the magnetic separation procedure as described in steps 1 to 6 by using a new column.

Magnetic separation with XS Columns

For instructions on the column assembly and the separation refer to the XS Column data sheet.

Depletion with LD Columns

- Place LD Column in the magnetic field of a suitable MACS Separator. For details refer to the LD Column data sheet.
- 2. Prepare column by rinsing with 2 mL of buffer.
- 3. Apply cell suspension onto the column.
- 4. Collect unlabeled cells that pass through and wash column with 2×1 mL of buffer. Collect total flow-through; this is the unlabeled cell fraction. Perform washing steps by adding buffer two times. Only add new buffer when the column reservoir is empty.

Depletion with D Columns

For instructions on column assembly and separation refer to the D Column data sheet.

Magnetic separation with the MultiMACS Cell24 Separator Plus

Refer to the MultiMACS Cell Separator Plus user manual for instructions on how to use the MultiMACS Cell24 Separator Plus.

2.4 Magnetic labeling and separation using autoMACS Separators

- ▲ Refer to the user manual and the short instructions for instructions on how to use the autoMACS Separators.
- ▲ Buffers used for operating the autoMACS Separators should have a temperature of $\geq +10$ °C.
- ▲ Place tubes in the following Chill Rack positions: position A = sample, position B = unlabeled (negative) fraction, position C = labeled (positive) fraction.

2.4.1 Magnetic labeling and separation using the autoMACS NEO Separator

- ▲ The autoMACS NEO Separator enables stage loading to extend column capacity for selected reagents, minimizing the need to divide larger samples.
- ▲ For more information on selecting alternative separation programs, stage loading-compatible reagents, autolabeling-compatible reagents, and the minimal and maximal volumes for each reagent and Chill Rack, refer to www.miltenyibiotec.com/automacs-neo-sample-processing.

Magnetic separation after manual labeling

- 1. Label the sample as described in section 2.2 Magnetic labeling.
- 2. Prepare and prime the instrument.
- 3. Place the Chill Rack on the MACS MiniSampler S.

- 4. Select the same Chill Rack in the **Experiment** tab. An experiment is created automatically. Tap to select sample positions.
- 5. Assign a reagent to each sample.
- Manual labeling is set automatically if autolabeling is not supported or no reagent rack is selected. Alternatively, tap Labeling in the reagent placement dialog and select Manual.
- 7. Tap **Sample volume** in the **Sample process** pane and enter the sample volume. Tap the return key.
- The separation program for highest target cell purity is selected by default. Refer to the Sample process pane for all available programs.
- 9. Place the sample(s) and empty tubes to the Chill Rack.
- 10. Tap **Run** to start the separation process.

Fully automated magnetic labeling and separation

- 1. Prepare and prime the instrument.
- Place the Chill Rack and MACS Reagent Rack 8 on the MACS MiniSampler S.
- Select the same Chill Rack and MACS Reagent Rack 8 in the Experiment tab. An experiment is created automatically.
- 4. Tap to select sample position(s).
- 5. To assign a reagent to each sample, tap **Scan reagent** and scan the reagent barcode. Alternatively, tap on a free position of the MACS Reagent Rack 8 for selection out of the reagent list.
- 6. Unscrew the lids from the reagent vials and place the vials onto the designated positions on the MACS Reagent Rack 8.
- 7. Tap Place reagent(s) on reagent rack button in the dialog box.
- 8. Automated labeling is set automatically if autolabeling is supported and a reagent rack is selected. Alternatively, tap **Labeling** in the reagent placement dialog and select **Auto**.
- 9. Tap **Sample volume** in the **Sample process** pane and enter the sample volume. Tap the return key.
- 10. Tap **Run** to start the separation process.

${\bf 2.4.2\,Magnetic\,labeling\,and\,separation\,using\,the\,autoMACS\,Pro\,Separator}$

Magnetic separation after manual labeling

- 1. Label the sample as described in section 2.2 Magnetic labeling.
- 2. Prepare and prime the instrument.
- 3. Apply tube containing the sample.
- 4. For a standard separation choose one of the following programs:

Positive selection: Possel

Collect positive fraction in row C of the tube rack.

Depletion: Depletes

Collect negative fraction in row B of the tube rack.

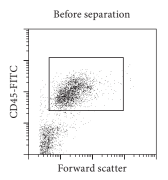
5. Tap **Run** to start the separation process.

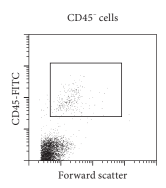
Fully automated magnetic labeling and separation

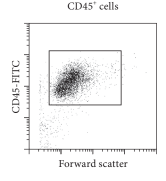
- 1. Switch on the instrument for automatic initialization.
- 2. Go to the **Reagent** menu and select **Read Reagent**. Scan the 2D barcode of each reagent vial with the barcode scanner on the autoMACS Pro Separator. Place the reagent into the appropriate position on the reagent rack.
- 3. Place sample and collection tubes into the Chill Rack.
- Go to the **Separation** menu and select the reagent name for each sample from the **Labeling** submenu. The correct labeling, separation, and wash protocols will be selected automatically.
- 5. Enter sample volume into the **Volume** submenu. Press **Enter**.
- 6. Tap **Run** to start the separation process.

3. Example of a separation using CD45 MicroBeads, mouse

CD45⁺ cells were isolated from mouse spleen using CD45 MicroBeads, mouse, an MS Column, and a MiniMACS[™] Separator. Cells were fluorescently stained with CD45-FITC and analyzed by flow cytometry. Cell debris and dead cells were excluded from the analysis based on scatter signals and propidium iodide fluorescence.







Refer to www.miltenyibiotec.com for all data sheets and protocols. Miltenyi Biotec provides technical support worldwide. Visit www.miltenyibiotec.com for local Miltenyi Biotec Technical Support contact information.

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