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# 1. Description

## This product is for research use only.

Components	2 mL CD19 MicroBeads, human : MicroBeads conjugated to monoclonal anti- human CD19 antibodies (isotype: mouse IgG1).
Capacity	For 10 <sup>9</sup> total cells, up to 100 separations.
Product format	CD19 MicroBeads, human 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 CD19<sup>+</sup> cells are magnetically labeled with CD19MicroBeads, human. Then, the cell suspension is loaded onto a MACS Column, which is placed in the magnetic field of a MACS Separator. The magnetically labeled CD19<sup>+</sup> cells are retained within the column. The unlabeled cells run through; this cell fraction is thus depleted of CD19<sup>+</sup> cells. After removing the column from the magnetic field, the magnetically retained CD19<sup>+</sup> cells can be eluted as the positively selected cell fraction. To increase the purity, the positively selected cell fraction containing the CD19<sup>+</sup> cells must be separated over a second column.

# **CD19 MicroBeads** human

Order no. 130-050-301

## 1.2 Background information

CD19 MicroBeads have been developed for the separation of human B cells based on the expression of the CD19 antigen. CD19 is a 95–120 kDa glycosylated transmembrane protein that is critically involved in signal transduction processes that regulate development, activation, and differentiation of B lymphocytes. CD19 is expressed on lineage B cells from the early lineagecommitted pro-B cell stage to the B cell blast stage including most malignant B cells. Expression is down-regulated during the differentiation into plasma cells. Furthermore, CD19 is expressed on follicular dendritic cells.

### 1.3 Applications

- Positive selection or depletion of cells expressing human CD19 antigen.
- Isolation or depletion of B cells from peripheral blood mononuclear cells (PBMCs), bodily fluids (e.g. bronchial lavage), or single-cell suspensions from tissue (e.g. lymphoid and tumor tissue).

### 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 human serum albumin, human serum, or fetal bovine serum (FBS). Buffers or media containing Ca<sup>2+</sup> or Mg<sup>2+</sup> are not recommended for use.

 MACS Columns and MACS Separators: CD19<sup>+</sup> 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 CD19 antigen can also be depleted using MS, LS, or XS Columns. Positive selection or depletion can also be performed by using the MultiMACS<sup>™</sup> 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 selection				
MS	10 <sup>7</sup>	2×10 <sup>8</sup>	MiniMACS, OctoMACS, SuperMACS II	
LS	10 <sup>8</sup>	2×10 <sup>9</sup>	MidiMACS, QuadroMACS, SuperMACS II	
	10 <sup>8</sup>	10 <sup>9</sup>	MultiMACS Cell24 Separator Plus	
XS	10 <sup>9</sup>	2×10 <sup>10</sup>	SuperMACS II	
Depletion				
LD	10 <sup>8</sup>	5×10 <sup>8</sup>	MidiMACS, QuadroMACS, SuperMACS II	
D	10 <sup>9</sup>		SuperMACS II	
Positive selection or depletion				
autoMACS	2×10 <sup>8</sup>	4×10 <sup>9</sup>	autoMACS NEO Separator autoMACS Pro Separator	
Multi-24 Column Block (per column)	10 <sup>8</sup>	10 <sup>9</sup>	MultiMACS Cell24 Separator Plus	

SuperMACS<sup>™</sup> 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., CD19 Antibody, anti-human, PE. 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 µm) (# 130-041-407) to remove cell clumps.

# 2. Protocol

### 2.1 Sample preparation

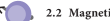
When working with anticoagulated peripheral blood or buffy coat, peripheral blood mononuclear cells (PBMCs) should be isolated by using a MACS PBMC Isolation Kit or by density gradient centrifugation, for example, using Ficoll-Paque™.

▲ 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 or lysed blood, 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 nonspecific cell labeling.

▲ For optimal performance it is important to obtain a single-cell suspension before magnetic labeling. Pass cells through 30  $\mu m$ 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<sup>7</sup> total cells. When working with fewer than 10<sup>7</sup> 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  $2 \times 10^7$  total cells, use twice the volume of all indicated reagent volumes and total volumes).

 $\blacktriangle$  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.

- 1. Determine cell number.
- Centrifuge cell suspension at 300×g for 10 minutes. Aspirate 2. supernatant completely.
- Resuspend cell pellet in 80 µL of buffer per 107 total cells. 3.
- Add 20 µL of CD19 MicroBeads, human per 10<sup>7</sup> total cells. 4.
- Mix well and incubate for 15 minutes in the refrigerator 5. (+2 to +8 °C).
- (Optional) Add staining antibodies, e.g., CD19 Antibody, anti-6. human, PE, according to manufacturer's recommendation.
- 7. Wash cells by adding 1-2 mL of buffer per 107 cells and centrifuge at 300×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. ▲ Note: For depletion with LD Columns, resuspend up to 1.25×10<sup>8</sup> cells in 500 µL 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 CD19<sup>+</sup> 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

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

MS: 500 μL LS: 3 mL

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

MS: 3×500 μL LS: 3×3 mL

▲ Note: Perform washing steps by adding buffer aliquots as soon as the column reservoir is empty.

- 5. 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.

MS: 1 mL LS: 5 mL

 (Optional) To increase the purity of CD19<sup>+</sup> 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

- 1. 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  $\ge$  +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.
- 8 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.
- 2. Place the Chill Rack and MACS Reagent Rack 8 on the MACS MiniSampler S.
- 3. 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.

# 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.

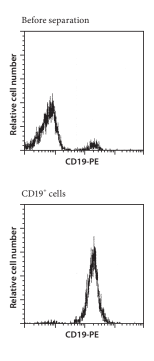
5. Tap **Run** to start the separation process.

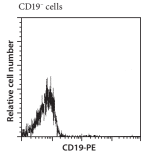
### Fully automated magnetic labeling and separation

- 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.
- 4. 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 CD19 MicroBeads, human

CD19<sup>+</sup> cells were isolated from human PBMCs using CD19 MicroBeads, human, an MS Column, and a MiniMACS<sup>™</sup> Separator. Cells were fluorescently stained with CD19-PE 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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