Magnetic Cell Selection and Separation of Human CD44+ Cells

OVERVIEW

The MAG-iso™ Human CD44+ Cancer Cells Isolation Kit (Cat. # K10134, ProMab Biotechnologies) is designed to isolate CD44+ human cancer stem cells using positive selection. The resulting cell preparation is highly enriched for CD44+ cells. Purity of recovered CD44+ cells can be up to 90%-99% and will vary depending on the preparation.

MATERIALS REQUIRED

1. Magnetic Separator like MagCellect Magnet (R&D System, Catalog # MAG997)
2. 12 x 75 mm (5 mL) tubes (Falcon, Catalog # 352008 or equivalent)
3. 50mL polypropylene centrifuge tubes (Santa Cruz Biotechnology, Inc, Catalog # sc-200251, or equivalent)
4. Sterile serological and Pasteur pipettes or transfer pipettes
5. Bench top centrifuge
6. 2-8° C refrigerator
7. Deionized or distilled water

Cell Selection Principle

1. Positive selection of CD44+ cells is achieved by incubation with biotinylated anti-Human CD44 monoclonal antibody.
2. CD44 monoclonal antibody bound cells are then magnetically tagged with MAG-iso™-Streptavidin 200nm.
3. Magnetically tagged cells are then isolated using magnetic separation. CD44+ tagged cells will migrate toward the magnet (these are the desired cells); unwanted/untagged cells remain in suspension
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Cell Selection Capacity

<table>
<thead>
<tr>
<th>Separator</th>
<th>No. of labeled cells</th>
<th>No. of total cells</th>
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<tbody>
<tr>
<td>PM Magnet</td>
<td>*2x10^6</td>
<td>*5x10^6</td>
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* The Max No. of cells will vary ±20% depending on the preparation.

Components of Kit (up to 20 tests, 10^6 cells).
1. Biotinylated anti-Human CD44 Antibody (Part C10108) – 0.5mL (for up to 20 tests with 5x10^6 cells per test).
2. MAG-isoTM-Streptavidin (Part B10002) - 1mL proprietary formulation (sufficient for 20 selections).
3. 10X MAG-isoTM Buffer (Part S10001) - 50 mL proprietary formulation.
4. DRNase (proprietary formulation of DNase I and RNase) - 700µL (Part M10001).

Storage

Reagents, except DRNase, are stable for 12 months from the date of receipt when stored in the dark at 2 - 8°C. DO NOT FREEZE.
DRNase can be stored in -20°C for long-term storage.

Reagent Preparation

1x MAG-isoTM Buffer: Prepare 20 mL of 1x MAG-isoTM Buffer for each sample by mixing 2 mL of 10X MAG-isoTM Buffer with 18 mL of sterile deionized or distilled water. The 1x MAG-isoTM Buffer is stable for 6 months at 4°C and should be kept on ice or at 4°C, but can be kept at room temperature in Magnetic separation step.

Cell Selection Procedure (5x10^6 cells/mL*)

Procedure has been used to select cells from a starting population as low as 1x10^6 cells/mL.

I. Cell Preparation: Cells and reagents should be kept cold using an ice bath or a refrigerator unless otherwise specified. Incubations must be carried out at 2-8°C in a refrigerator and not in an ice bath to avoid excessively low temperatures that can slow the kinetics of the optimized reactions.

Prepare the reaction buffer ahead of time and keep it refrigerated or on ice --- see Reagent Preparation.
A. Preparing a single cell suspension from frozen cells

1. To a 50 mL conical tube, add 30µL formulated DRNase.
2. Transfer the cell suspension to the 50ml conical tube.
3. Slowly add 10mL pre-warmed (37°C) DMEM medium (with 10% FBS) drop wise to the cells.
4. Centrifuge cell suspension at 200 x g at 4°C for 15 minutes.
5. Carefully remove all but approximately 100µL of the supernatant using a pipette.
6. Gently resuspend the cell pellet in 10mL of fresh medium (pre-chilled to 4°C) to the tube.
7. Centrifuge the suspension at 200 x g at 4°C for 15 minutes.
8. Remove media and resuspend final pellet up to 5 x106 in 500 µL of Selection Buffer.
9. Pass the suspended cells through a 30-50µm nylon cell strainer.

Cells must be resuspended in cold reaction buffer prior to the antibody selection procedure. Buffer has to be kept on ice at all times.

NOTE: For downstream applications that are sensitive to DRNase (eg. hematopoietic colony assays), wash cells once in the appropriate assay buffer (without DRNase) before continuing.

B. Preparing a single cell suspension from fresh adherent cells

NOTE: Cells reaching 80-100% confluence are ready for harvest and can be used for selection.

1. Remove media and rinse adherent cells with PBS.
2. Treat cells with 3mL 1x Trypsin EDTA (0.05%Trypsin/0.53mM EDTA in HBSS) for 2-3 min.
3. Break cell clumps by serological pipetting 8-10 times.
4. Add 12mL culture medium; break the cell clumps to single cell suspension by serological pipetting 8-10 times.
5. Centrifuge in 50ml conical tube for 5min at 300 x g.
6. Remove media and resuspend final pellet up to 5 x106 to a concentration of 1x10⁷cells/mL in 500 µL of in MAG-iso™ Buffer Selection Buffer.
7. Pass the suspended cells through a 30-50µm nylon cell strainer.

C. Preparing a single cell suspension from fresh suspension culture

1. Harvest suspension culture into 50ml conical tube and centrifuge for 5 min at 300 x g.
2. Remove media and rinse pellet with PBS.
3. Centrifuge for 5 min at 300 x g; remove the PBS.
4. Treat the cells with 1mL 1x Trypsin EDTA (0.05%Trypsin/0.53mM EDTA in HBSS) for 2-3 min.
5. Break cell clumps by pipetting 8-10 times with P1000 pipette, or until a single cell suspension is achieved.
6. Add 10mL culture medium; centrifuge for 15 min at 200 x g at 4°C.
7. Remove media and resuspend the pellet in MAG- iso™ Buffer to a concentration of 8x10⁶ cells/mL.
8. Pass the suspended cells through a 30-50µm nylon cell strainer.
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**NOTE:** Regardless of which preparation is undertaken, it is imperative to ensure a single-cell suspension by passing cells through a 30-50μm nylon cell strainer.

**NOTE:** Cells must be suspended in cold reaction buffer to a concentration of 1x10^7 cells/mL prior to beginning the antibody selection procedure.

(Optional): Fc domain blocking
In some applications, this step is required to minimize non-specific binding of the antibodies via their Fc domain to the Fc receptors (FcR) present on various cell types.

1. Transfer 0.5mL suspended cells (5x10^6 adherent or, 4x10^6 suspension cells) into an Eppendorf tube.
2. Add 1-10 μg of IgG or Fc blocking specific antibody (in a volume not exceeding 100 μL) per 1x10^7 cells and incubate 5-10 minutes at 2 - 8° C.

II. Positive selection of CD44+ cells

1. Transfer desired amount CD44+ cells to a micro-centrifuge tube.
2. Add 2 μL (~ 2 ug) of biotinylated anti-human CD44+ antibody per 1x10^7 cells to the tube.
3. Gently mix the cell-antibody suspension, avoiding formation of bubbles, and incubate at 2-8°C on a rotator for 15 minutes.
4. After incubation, dilute the cell suspension with 1mL of cold 1x MAG-isoTM Selection Buffer. Aliquot into Eppendorf tubes and centrifuge at 4°C for 5 minutes at 300g. Repeat washing step once more.
5. Completely remove the supernatant and gently resuspend the cell pellet with 0.5 mL of cold 1x Selection Buffer.
6. Add 50 µL MAG-iso™-Streptavidin (Part B10002) to the cell suspension. Mix gently and incubate at 2 - 8° C on a rotator in a refrigerator for 15 minutes.
7. At the end of the incubation period, dilute the cell content to 1.5mL with cold 1x MAG-iso™ Buffer, centrifuge in a refrigerated Eppendorf centrifuge at 300 x g for 5 minutes or in a single-speed clinical bench top centrifuge for 1 minute.
8. Completely remove the supernatant and resuspend the cell pellet by gently pipetting 1.5mL of cold 1x MAG-iso™ Buffer, and centrifuge in a refrigerated Eppendorf centrifuge at 300 x g for 5 minutes or in a single-speed clinical bench top centrifuge for 1 minute.
9. Completely remove the supernatant and resuspend the cell pellet by gently pipetting 3mL of cold 1x MAG-iso™ Buffer and transfer the cell into a 5mL Falcon tube.
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III. Magnetic Separation

1. Place the 5 mL Falcon tube in the magnetic fields of R&D Magnet or a suitable Separator.
2. Incubate for 6 minutes at room temperature (18-25 °C).
3. Magnetically tagged cells will migrate toward the magnet (these are desired CD44+ cells), leaving the unwanted/non-specific cells in suspension.
4. While the tube remains in the magnet, carefully remove all the reaction supernatant and save to a 15ml tube as Flow Through.
5. Remove the tube containing the desired magnetically bound CD44+ cells from the magnet, and gently resuspend pellet in 3 mL of cold 1x Selection buffer.
6. Repeat steps 1. through 4. 2-3x with the positively selected (bound) cell fraction following cell resuspension; save all the reaction supernatant to the 15ml tube.
7. For the final wash, resuspend cell pellet in 1mL of 1x Selection Buffer.
8. Centrifuge the fractions at 300 x g for 5 minutes.
9. The resulting bound cells contain the desired and enriched CD44+ cells, save the pellet for FACS analysis). The Flow Through and the final Wash contain the undesired CD44- cells and should be discarded following lab protocol.
10. Cells are now ready for further experimentation or FACS analysis.

IV. Cell Harvesting

1. Depending upon the intended use of the cells, resuspend the beads fraction and/or the pellet from the supernatant fraction with your application buffer.
2. For FACS analysis, resuspend the beads fraction and the pellet from the supernatant fraction with 1.5ml cell staining buffer (Part S10002; not included); centrifuge at 300 x g for 3 min; remove the supernatant; (continue to Cell Staining Procedure).
3. For cryopreservation, re-suspend the beads and the pellet in 1x DMEM or RPMI medium (continue to Cell Cryopreservation Procedure).