STEMSEP® PROCEDURE:

This protocol is recommended for use with commercially available magnetic positive selection columns*.

- Prepare nucleated cell suspension at a concentration of 10⁸ cells/mL in recommended medium (see Notes and Tips).
- 2. Add StemSep® selection cocktail at 100 μ L/mL of cells (e.g. for 1 mL of cells add 100 μ L of cocktail). Mix well and incubate for 10 minutes at 4 8°C.
- 3. Add CD14-PE labeled antibody if desired (see Notes and Tips).
- Add magnetic colloid at 60 μL/mL cells (e.g. for 1 mL of cells add 60 μL of colloid). Mix well and incubate for 10 minutes at 4 - 8°C.
- 5. Wash cells by adding 10 20X the original cell suspension volume of recommended medium. Centrifuge at 300 x g for 8 minutes, remove supernatant and resuspend pellet to original cell volume or the maximum volume recommended for the separation column of choice.
- 6. Cells are ready for separation. Follow directions recommended by positive selection column manufacturer.

*Not for use with StemSep® negative selection columns.

NOTES AND TIPS:

PREPARING A MONONUCLEAR CELL SUSPENSION. Prepare a mononuclear cell suspension from whole peripheral blood by Ficoll-PaqueTM PLUS density separation (Catalog #07957). For previously frozen mononuclear cells, we recommend incubating the cells with 100 μg/mL DNase I (Catalog #07900) for at least 15 minutes at room temperature prior to labeling and separation. Filter clumpy suspensions through a 30 μm mesh nylon strainer to avoid blocking the separation column.

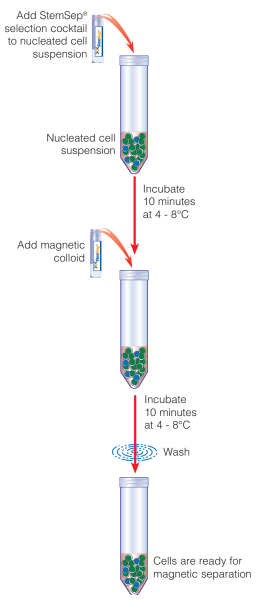
RECOMMENDED MEDIUM. The recommended medium is PBS containing 2 mM EDTA and 0.5% BSA. Medium should be Ca⁺⁺ and Mg⁺⁺ free. Using degassed medium reduces the chance of developing air bubbles in the column. Air bubbles cause channeling in the column, reducing the capacity of the column and potentially compromising purity.

ASSESSING PURITY. The CD14 Positive Selection Cocktail uses the antihuman CD14 antibody clone MEM-15. We recommend one of the following antibody clones to assess purity by flow cytometry: UCHM1 or MO-P9 (Catalog #10506) (both non-blocking). One of the following methods can also be used to assess purity:

- Add PE-labeled antibodies at the same time as the cocktail: Add the fluorochrome-conjugated anti-human CD14 antibody (Catalog #10506) at a concentration of 0.4 μg/mL immediately after adding the cocktail to provide a strong detection signal without affecting separation performance. This method labels the positive cells in the entire sample.
- 2. Use alternative markers after separation: Detect CD36+ cells.
- 3. Use a fluorochrome-conjugated secondary antibody, such as a FITC-labeled sheep anti-mouse IgG.

Ficoll™ and Ficoll-Paque™ PLUS are trademarks of GE Healthcare Ltd.







Components:

• StemSep® Human CD14 Positive Selection Cocktail

StemSep® Magnetic Colloid

1.0 mL 1.5 mL



PRODUCT INFORMATION SHEET.

REQUIRED EQUIPMENT:

StemSep® Magnet (Catalog #11030, 11050, 11060, or 11070) or a magnet with the strength of at least 0.6 Tesla, and commercially available positive selection columns.

PRODUCT DESCRIPTION AND APPLICATIONS:

StemSep® Human CD14 Positive Selection Cocktail and StemSep® Magnetic Colloid label CD14+ cells for magnetic separation. These positive selection reagents are designed to positively select CD14+ cells (cells expressing the CD14 antigen) from fresh or previously frozen peripheral blood mononuclear cells.

STEMSEP® LABELING OF HUMAN CELLS:

Target cells are specifically labeled with dextran-coated magnetic colloid using bispecific Tetrameric Antibody Complexes (TAC). These complexes recognize both dextran and the target cell surface antigen (Figure 1). The use of bispecific TAC avoids expensive and inefficient covalent coupling of antibodies to magnetic particles. The small size of the colloidal magnetic dextran iron particles allows for efficient binding to the TAC-labeled cells. Magnetically labeled cells are then separated from unlabeled cells by passing them through a magnetic separation column placed in a magnet.

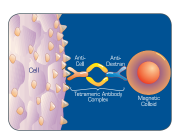
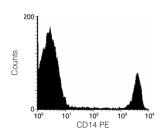
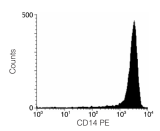


FIGURE 1.Schematic Drawing of StemSep® TAC Magnetic Labeling of Human Cells.

TYPICAL STEMSEP® CD14 SELECTION PROFILE:

Start: 16.7% CD14⁺ Cells Selected: 98.6% CD14⁺ Cells





The expected CD14⁺ content of the enriched fraction is $97.5 \pm 0.5\%$ with expected recoveries of $58 \pm 9\%$.

COMPONENT DESCRIPTIONS:

STEMSEP® HUMAN CD14 POSITIVE SELECTION COCKTAIL CODE #14758C This cocktail contains a combination of monoclonal antibodies purified from hybridoma culture supernatant by affinity chromatography using Protein A or Protein G Sepharose. These antibodies are bound in bispecific TAC which are directed against CD14 and dextran. The mouse monoclonal antibody subclass is $\lg G_1$. It should be noted that this product is a biological reagent, and as such cannot be completely

STEMSEP® MAGNETIC COLLOID

CODE #10051

A colloidal suspension of magnetic dextran iron particles in USP saline, pH 7.0 - 7.5.

characterized or quantified. Some variability is unavoidable.

STABILITY AND STORAGE:

STEMSEP® HUMAN CD14 POSITIVE SELECTION COCKTAIL

Stable at 2 - 8°C for 2 years. Do not freeze. This product has been sterility tested.

STEMSEP® MAGNETIC COLLOID

This product is shipped at room temperature. Once opened, stable at 2 - 8°C for 6 weeks. Stable at -20°C for 1 year. Repeated freezing and thawing is possible but not recommended. Vortex before re-freezing. This product has been sterility tested.

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- 3. Molday RS, MacKenzie D. Immunospecific ferromagnetic iron dextran reagents for the labelling and magnetic separation of cells. J Immunol Methods 52: 353, 1982



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