

Produktinformation



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Zellkultur & Verbrauchsmaterial



Diagnostik & molekulare Diagnostik



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for the Science of Tomorrow™

Lympholyte®-Rabbit

CL5050 CL5055

DESCRIPTION:

Lympholyte®-Rabbit is a density separation medium specifically designed for the isolation of viable lymphocytes from rabbit lymphoid cell suspensions.

APPLICATIONS:

Lympholyte®-Rabbit can be utilized with a simple protocol for the elimination of erythrocytes, dead cells and debris from rabbit spleen, lymph node and thymus suspensions. The resulting cell population demonstrates a high and non-selective recovery of viable lymphocytes that are suitable for use as target cells in cytotoxicity and FACS assays, as well as in <u>in vivo</u> and <u>in vitro</u> functional studies. Other successful applications include:

- i) the removal of dead cells in sequential cytotoxicity studies eg. B-cell depletion.
- ii) the removal of erythrocytes, dead cells and debris from other rabbit tissue suspensions including bone marrow, liver and lung.

PRESENTATION:

0.22 µm filtered liquid. CL5050, 5 x 30 ml CL5055, 1 x 500 ml

STORAGE/STABILITY:

Store at room temperature unopened. Store at +4°C once opened., Always store protected from

Note: Phase separation may occur with long-term storage. SHAKE WELL BEFORE USE. ALLOW TO STAND UNTIL NO AIR BUBBLES REMAIN. **USE AT ROOM** TEMPERATURE.

Visit our website for your local distributor.



In CANADA: Toll Free: 1-800-268-5058

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1210 Turrentine Street, Burlington, NC 27215 ph: (336) 513-5135, fax: (336) 513-5138 e-mail: service@cedarlanelabs.com

SPECIFICATIONS:

<u>Composition</u>: NycogradeTM Polysucrose 400 and Sodium Diatrizoate

<u>Density</u>: $1.0965 \pm 0.001 \text{ g/cm}^3$ @ 22°C.

<u>pH</u>: 6.9 ± 0.3

<u>Viability/ Purity</u>: Recovery of viable lymphocytes $\geq 70\%$.

Results obtained on a rabbit spleen suspension:

Fraction	Viable Lymphocytes	Erythrocyte Contamination
upper	<1%	<1%
interphase	>70%	< 15%
lower	<10%	< 5%
pellet	<20%	> 80%

METHOD OF USE:

Use Lympholyte[®]-Rabbit and preferably a serum-free medium (Phosphate Buffered Saline, Modified McCoy's Medium, etc) at room temperature (approximately 22°C).

1. Prepare a lymphocyte suspension using your preferred method and medium. Spleen has a high membrane content and a clean suspension is required for proper separation.

Suggested method: a) cut up spleen into small pieces

b) homogenize

c) pass suspension through a fine screen mesh

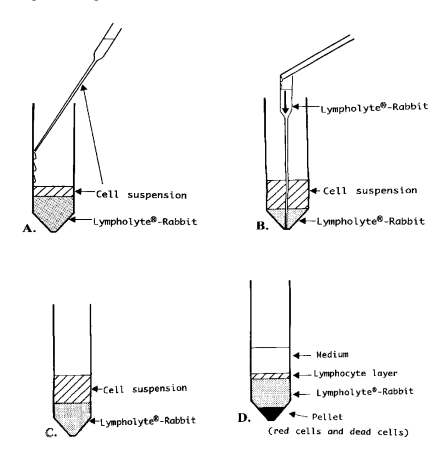
Other tissues: homogenize thoroughly to obtain a clean suspension.

- 2. Adjust the cell concentration to a **maximum** of 5×10^6 nucleated cells per ml. **Note**: If cell suspension contains a large amount of debris or erythrocytes, a cleaner separation will be obtained if the cell concentration is set at 2.5×10^6 cells/ml.
- 3. Layer the cell suspension over Lympholyte[®]-Rabbit according to Method A or Method B (see figures). Use a 10-15 ml centrifuge tube.

Method A: Add 5 ml of Lympholyte[®]-Rabbit to the centrifuge tube. Using a pipette, carefully layer 5 ml of the cell suspension over the Lympholyte[®]-Rabbit with as little mixing as possible at the interface (Figure A). Since Lympholyte[®]-Rabbit is of greater density than the cell suspension, a distinct interface will be formed (Figure C).

Method B: Add 5 ml of the cell suspension to the centrifuge tube. Place a large (23 cm) Pasteur pipette to the bottom of the tube (Figure B). Slowly add Lympholyte[®]-Rabbit to the Pasteur pipette allowing gravity to layer it under the cell suspension. Continue until 5 ml of Lympholyte[®]-Rabbit has been layered under the cell suspension. Since Lympholyte[®]-Rabbit is of greater density than the cell suspension, the cell suspension will form a layer above the Lympholyte[®]-Rabbit with a distinct interface (Figure C).

- 4. Centrifuge for 30 minutes at 1500g at **room temperature**.
- 5. After centrifugation, there will be a well-defined lymphocyte layer at the interface (Figure D). Using a Pasteur pipette, carefully remove the cells from the interface and transfer to a new centrifuge tube.
- 6. Dilute the transferred cells with medium and centrifuge at 800g for 10 minutes to pellet the lymphocytes; discard the supernatant.
- 7. Wash the lymphocytes 2-3 times in medium (can use media containing serum at this point) before further processing.



FOR RESEARCH USE ONLY

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JK 11/11/20