

Instructions For Use

A00024-C-IFU-RUO

Rev. Date: June 25, 2013

Revision: 1

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P.O. Box 3286 - Logan, Utah 84323, U.S.A. - Tel. (800) 729-8350 - Tel. (435) 755-9848 - Fax (435) 755-0015 - www.scytek.com

CD45RO, T-Cell; Clone UCHL1 (Concentrate)

Volume

Availability/Contents: <u>Item #</u>

A00024-C 1

Description:

Species: Mouse

Immunogen: BALB/C mice were immunized with IL-2 dependent T cell line, CA1.

Clone: UCHL1 lsotype: lgG2a, Kappa

Format: This antibody is provided in a phosphate buffered saline containing 1% BSA.

Specificity: This antibody reacts with a 180 kD glycoprotein of CD45 family, occurring on most thymocytes

and activated T cells but only on the portion of the resting T cells. It reacts with most

thymocytes, a subpopulation of resting cells within both the CD4 and CD8 subsets and mature activated T cells. This antibody shows no reactivity with normal B or natural killer cells, but reacts with granulocytes and monocytes. Though this antibody is useful to identify T-cell

lymphomas and leukemia, rare staining with B cell lymphomas reported.

Background: The CD45RA and CD45RO isoforms of the leukocyte common antigen identify functionally

distinct naive and memory T cell subsets. In vitro, CD45RA+ / CD45RO - peripheral blood lymphocytes (PBL) can be converted to CD45RA- / CD45RO + phenotype upon activation in the presence of IL-2. Both CD3+ and CD3-/CD56+ lymphocyte subsets can be converted to CD45RO + lymphocytes. Expression of CD45RO was observed only in response to IL-2 and

was not observed during long-term culture in IL-4, IL-6, or IL-7.

Species Reactivity: Human, Rhesus Monkey. Does not react with Rat. Others not tested.

Positive Control: Tonsil or Lymph Node. Cellular Localization: Cell Membrane.

Titer/Working Dilution: Immunohistochemistry: 1:150 – 1:250

Microbiological State: This product is not sterile.

Uses/Limitations: Not to be taken internally.

For Research Use Only.

This product is intended for qualitative immunohistochemistry with normal and neoplastic formalin-fixed, paraffin-embedded tissue sections, to be viewed by light microscopy.

Do not use if reagent becomes cloudy. Do not use past expiration date. Use caution when handling reagents.

Non-Sterile.

Ordering Information and Current Pricing at www.scytek.com

Storage: 2° C 8° C

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Procedure:

- 1. **Tissue Section Pretreatment (Highly Recommended):** Staining of formalin fixed, paraffin embedded tissue sections is significantly enhanced by pretreatment with Citrate Plus (ScyTek catalog# CPL500) or equivalent.
- Primary Antibody Incubation Time: We suggest an incubation period of 30 minutes at room temperature.
 However, depending upon the fixation conditions and the staining system employed, optimal incubation should be determined by the user.
- 3. **Visualization:** For maximum staining intensity we recommend the "UltraTek HRP Anti-Polyvalent Lab Pack" (ScyTek catalog# UHP125, see IFU for instructions) combined with the "DAB Chromogen/Substrate Bulk Pack (High Contrast)" (ScyTek catalog# ACV500, see IFU for instructions).

Precautions:

Contains Sodium Azide as a preservative (0.09% w/v).

Do not pipette by mouth.

Avoid contact of reagents and specimens with skin and mucous membranes.

Avoid microbial contamination of reagents or increased nonspecific staining may occur.

This product contains no hazardous material at a reportable concentration according to U.S. 29 CFR 1910.1200,

OSHA Hazardous Communication Standard and EC Directive 91/155/EC.

References:

- Greene GL; Nolan C; Engler JP; Jensen EV. Proceedings of the National Academy of Sciences of the United States of America, 1980, 77(9):5115-9.
- 2. Greene GL; Gilna P; Waterfield M; Baker A; Hort Y; Shine J. Science, 1986, 231(4742):1150-4.
- 3. Green S; Walter P; Greene G; Krust A; Goffin C; Jensen E; Scrace G; Waterfield M; Chambon P. Journal of Steroid Biochemistry, 1986, 24(1):77-83
- 4. Green S; Chambon P. Nature, 1986, 324(6098):615-7.
- 5. Green S; Gronemeyer H; et al. Growth Factors and Oncogenes in Breast Cancer, 1987. Chichester, England, Ellis Horwood Ltd. 7-28.
- 6. Evans RM. Science, 1988, 240:889-95.
- 7. Danielson M; Northrop JP; et al. EMBO Journal, 1986, 5:2513-22.
- 8. Kumar V; Green S; Stack G; Berry M; Jin JR; Chambon P. Cell, 1987, 51(6):941-51.
- 9. Greene GL; Sobel BN; et al. Molecular Endocrinology, 1988, 2:714-26.
- 10. Jensen EV; Jacobson HI. Recent Progress in Hormone Research, 1962, 18:387-414.
- Walter P; Green S; Greene G; Krust A; Bornert JM; Jeltsch JM; Staub A; Jensen E; Scrace G; Waterfield M; et al. Proc Nat Academy of Sciences of the United States of America, 1985, 82(23):7889-93.

Warranty:

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