

Produktinformation



Forschungsprodukte & Biochemikalien
Zellkultur & Verbrauchsmaterial
Diagnostik & molekulare Diagnostik
Laborgeräte & Service

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SZABO-SCANDIC HandelsgmbH

Quellenstraße 110, A-1100 Wien T. +43(0)1 489 3961-0 F. +43(0)1 489 3961-7 <u>mail@szabo-scandic.com</u> www.szabo-scandic.com



Address: Avda. Universidad de Coimba, s/n Cancer Research Center (C.I.C.) Campus Miguel de Unamuno 37007 Salamanca (Spain) Tel. / Fax: (+34) 923 294 827

E-mail: info@immunostep.com

www.immunostep.com

Product: FITC Anti-Human CD10 Cat. Ref: 10F-100T Reagent provided: 100 test (20µl / test) Description: Monoclonal Mouse Anti-Human CD10 FITC, is recommended for use in flow cytometry for identification of human common acute lymphoblastic leukaemia antigen (CALLA). The conjugate is provided in aqueous buffered solution containing protein stabilizer, and ≤0.09% sodium Azide Clone: TP1/31 Isotype: IgG1 Fluorochrome: Fluorescein isothiocyanate (Molecular Probes)

INTENDED PURPOSE.

CDIO FITC is a monoclonal antibody conjugated that may be used to enumerate early B lymphocytes, stem-cell of the lymphocyte lineage and immature thymocytes.

TECHNICAL SUMMARY.

Reactivity: The monoclonal antibody is directed against the CDIO- antigen (CALL-antigen), which is expressed on human lymphoblasts. The antibody reacts with early B lymphocytes (stem-cell, pre-B) and with the stem-cell of the lymphocyte lineage and immature thymocytes. Lymphoblasts of a patient with an Acute Lymphocytic Leukaemia of the C-ALL type were found to be positive. Normal B- and T lymphocytes, monocytes and platelets were found to be negative.

Specificity: CD10 clon HI10a recognizes neutral endopeptidase, a 95 kD type II transmembrane glycoprotein, which is refered as the Common Acute Lymphoblastic Leukemia Antigen (CALLA) expressed on common acute lymphoblastic leukaemia cells which is seen on early B-lymphocytes and with stem cells of the lymphocyte lineage and immature thymocytes. The antigen is a neutral endopeptidase.

CLINICAL RELEVANCE

Monoclonal Mouse Anti-Human, can be used to define malignant cells designated as common acute lymphoblastic leukaemia B cells, T-cell leukemias, lymphoma, melanoma, and glioma cell lines.

PRINCIPLES OF THE TEST.

Immunostep CDIO FITC monoclonal antibodies bind to the surface of cells that express the CDIO antigen. To identify these cells, peripheral blood leucocytes are incubated with the antibodies and red blood cells are lysed before washing to remove unbound antibodies. An appropriate fixative solution is added to lysed, washed cells before the stained and fixed cells are analysed by flow cytometry with an argon ion laser at 488nm.

REAGENTS.

Cluster Designation:	CDIO
WHO Classification:	Leukocyte Workshop V.
Clone:	HIIOa
lsotype:	lgG1
Species:	Mouse
Composition:	IgG1 heavy chain
	Kappa light chain
Source:	Hybridome Cells
Method of Purification:	Affinity chromatography
Fluorochrome:	Fluorescein isothiocyanate
	Excitation wavelength 488 nm
	Emission wavelength 520 nm
Molar composition:	FITC/protein 6,0 – 8,0

Revision Nº2

Emission Date: 20/03/2013

HT-FITC-0010-1



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Reagents contents:

2 ml vial containing monoclonal antibody for 100 tests. The conjugate is provided in aqueous buffered solution containing protein stabilizer, and $\leq 0.09\%$ sodium Azide Ready to use.

Reagent preparation:

1. STATEMENTS, SETTINGS AND WARNINGS.

- Reagents contain sodium azide. Sodium azide under acid conditions yields hydrazoic acid, an
 extremely toxic compound. Azide compounds should be diluted with running water before
 being discarded. These conditions are recommended to avoid deposits in plumbing where
 explosive conditions may develop.
- Light exposure should be avoided. Use dim light during handling, incubation with cells and prior to analysis.
- Do not pipet by mouth.
- Samples should be handled as if capable of transmitting infection. Appropriate disposal methods should be used.
- The sample preparation procedure employs a fixative (formaldehyde). Contact is to be avoided with skin or mucous membranes.
- Do not use antibodies beyond the stated expiration dates of the products.
- Deviations from the recommended procedure enclosed within this product insert may invalidate the results of testing.
- FOR IN VITRO DIAGNOSTIC USE
- For professional use only.

2. APPROPIATE STORAGE CONDITIONS.

- Fluorescein isothiocyanate (Molecular Probes)
- Keep in dark place at 2-8°C. DO NOT FREEZE.

*Note: it's been described stored conjugated monoclonal antibodies on FITC at -20°C. This can affect to the conjugated intense.

3. EVIDENCE OF DETERIORATION.

- Reagents should not be used if any evidence of deterioration or substantial loss of reactivity is observed. For more information, please contact with our technical service: tech@immunostep.com
- The normal appearance of the FITC conjugated monoclonal antibody is a clear, yellow-orange liquid.

4. SPECIMEN COLLECTION.

Collect venous blood samples into blood collection tubes using an appropriate anticoagulant (EDTA or heparin). For optimal results the sample should be processed within 6 hours of venipuncture. EDTA, ACD or heparin may be used if the blood sample is processed for analysis within 30 hours of venipuncture. ACD or heparin, but not EDTA, may be used if the sample is not processed within 30 hours of venipuncture. Samples that cannot be processed within 48 hours should be discarded.

If venous blood samples are collected into ACD for flow cytometric analysis, a separate venous blood sample should be collected into EDTA if a CBC is required.

Unstained anticoagulated blood should be retained at 20-25° C prior to sample processing. Blood samples that are hemolyzed, clotted or appear to be lipemic, discoloured or to contain interfering substances should be discarded.

Refer to "*Standard Procedures for the Collection of Diagnostic Blood Specimens*" published by the National Committee for Clinical Laboratory Standards (NCCLS) for additional information on the collection of blood specimens.

5. SAMPLE PREPARATION.

1. From a collect blood into an appropriate anticoagulan mixed with EDTA (until the process moment, keep in cold). Determine cell viability using Trypan Blue or propidium iodide. If the cell viability is not at least 85%, the blood sample should be discarded.



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- 2. Pipette 100µl of well mixed blood into 12 x 75 mm polypropylene centrifuge tubes marked unknown and control.
- 3. Add 20µl of Immunostep CDIO FITC-conjugated monoclonal antibody and 180µl of phosphate buffered saline (PBS) to tubes marked unknown. In other control tube add 10µl of corresponding Immunostep IgG1 PE-conjugated isotypic control reagent. Mix gently.
- 4. Incubate all tubes for 15 minutes at room temperature ($22 \pm 3^{\circ}$ C) in the dark.
- 5. Add lysing solution to all tubes according to the manufacturer's directions.
- 6. Centrifuge all tubes at 400 x g for 3 minutes at room temperature.
- 7. Add fixing solution to all tubes according to the manufacturer protocol. Retain cells in fixing solution for not less than 30 minutes at room temperature (22 ±3°C) in the dark.
- 8. Wash the cells in all tubes twice with 4mL of PBS. Centrifuge at 400 x g for 3 minutes after each wash procedure.
- 9. Resuspend the cells from the final wash in 1 ml of PBS and store tubes at 2-8°C in the dark until flow cytometric analysis is performed. It is recommended that analysis be performed within 24-48 hours of staining and fixation.
- 10. Analyze on a flow cytometer according to the manufacturer instructions. For alternate methods of whole blood lysis, refer to the manufacturer recommended procedure.

6. MATERIALS REQUIRED BUT NOT SUPPLIED.

ISOTYPE CONTROL REAGENTS:

Leucocyte gating reagent:

Serofuge or equivalent centrifuge

12 x 75 mm polypropylene centrifuge tubes

Micropipette capable of dispensing 5 μ l, 20 μ l, 100 μ l, and 500 μ l volumes

Blood collection tubes with anticoagulant

Phosphate buffered saline (PBS)

Trypan Blue or propidium iodide, 0.25% (w/v) in PBS for the determination of cell viability

Lysing Solution

Fixing Solution

Flow cytometer:

FITC (Ex-Max: 494 nm/Em-Max: 520 nm). Recommended 488 nm ion argon laser, 502 LP filter and 530/30 detector-equipped flow cytometer.

MOUSE IGG1: FITC

Mouse anti-human CD45: PE/CD14: PerCP

7. INTERPRETATION OF RESULTS.

a. <u>FLOW CYTOMETRY</u>

Analyze antibody-stained cells on an appropriate flow cytometer analyzer according to the manufacturer instructions. The right angle light scatter or other scatter (SSC) versus forward angle light scatter (FSC) is collected to reveal the lymphocyte cell cluster. A gate is drawn for the lymphocyte cluster (lymphocyte bitmap). The fluorescence attributable to the FITC- conjugated monoclonal antibody is collected, and the percentage of antibody-stained T lymphocytes is determined. An appropriate FITC- conjugated isotypic control of the same heavy chain immunoglobulin class and antibody concentration must be used to estimate and correct for non-specific binding to lymphocytes. An analysis region is set to exclude background fluorescence and



to include positively stained cells. The following histograms are representative of cells stained and gated on the lymphocyte region from a normal donor.



The histogram is a representations of a Nalm-6 cell line. Cells were stained with CD10 FITC (yellow histogram) and FITC IgG1 Mouse Isotype Control (red histogram). Cells were analyzed on a FACSAria (Becton Dickinson, San Jose, CA) flow cytometer, using BD FACSDiva software.

8. QUALITY CONTROL PROCEDURES.

Non-specific fluorescence identified by the FITC conjugated isotypic control is usually less than 2% in normal individuals. Non-specific fluorescence identified by the PE and APC conjugated isotypic controls are usually less than 4% in normal individuals. If the background level exceeds these values, test results may be in error. Increased non-specific fluorescence may be seen in some disease states.

A blood sample from each normal and abnormal donor should be stained with the CD45 Panlymphocyte and CD14 Pan-monocyte monoclonal antibodies. When used in combination, these reagents assist in identifying the lymphocyte analysis region, and distinguish lymphocytes from monocytes, granulocytes and unlysed or nucleated red cells and cellular debris.

A blood sample from a healthy normal donor should be analyzed as a positive control on a daily basis or as frequently as needed to ensure proper laboratory working conditions. Each laboratory should establish their own normal ranges, since values obtained from normal samples may vary from laboratory to laboratory.

An appropriate isotype control should be used as a negative control with each patient sample to identify non-specific Fc binding to lymphocytes. An analysis region should be set to exclude the non-specific fluorescence identified by the isotypic control, and to include the brighter fluorescence of the lymphocyte and granulocyte population that is identified by the specific antibody.

Refer to the appropriate flow cytometer instrument manuals and other available references for recommended instrument calibration procedures.

9. LIMITATIONS OF THE PROCEDURE.

- 1. Incubation of antibody with cells for other than the recommended time and temperature may result in capping or loss of antigenic determinants from the cell surface.
- 2. The values obtained from normal individuals may vary from laboratory to laboratory; therefore, it is recommended that each laboratory establish its own normal range.
- 3. Abnormal cells or cell lines may have a higher antigen density than normal cells. This could, in some cases, require the use of a larger quantity of monoclonal antibody than is indicated in the procedures for Sample Preparation.
- 4. Blood samples from abnormal donors may not always show abnormal values for the percentage of lymphocytes stained with a given monoclonal antibody. Results obtained by flow cytometric analysis should be considered in combination with results from other diagnostic procedures.
- 5. When using the whole blood method, red blood cells found in some abnormal donors, as well as nucleated red cells found in normal and abnormal donors may be resistant to lysis by lysing solutions. Longer red cell lysis periods may be needed to avoid the inclusion of unlysed red cells in the lymphocyte gated region.
- 6. Blood samples should not be refrigerated or retained at ambient temperature for an extensive period (longer than 24-30 hours) prior to incubating with monoclonal antibodies.
- 7. Accurate results with flow cytometric procedures depend on correct alignment and calibration of the laser, as well as proper gate settings.
- 8. Due to an unacceptable variance among the different laboratory methods for determining absolute lymphocyte counts, an assessment of the accuracy of the method used is necessary.



9. Al results need to be interpreted in the context of clinical features, complete immunophenotype and cell morphology, taking due account of samples containing a mixture of normal and neoplastic cells.

10. REFERENCE VALUES.

The cellular elements of human Bone Marrow include lymphocytes, monocytes, granulocytes, red blood cells and platelets.

Cell type	Percentage
Progranulocytes	56,7
Neutrophils	53,6
Myeloblasts	0,9
Promyeloblasts	3,3
Promyelocytes	12,7
Metamyelocytes	15,9
Eosinophils	3,1
Basophils	<0,1
Proerythrocyte	25,6
Proerythrblasts	0,6
Basophil Erythroblast	1,4
Polycromatic Erythroblast	21,6
Ortocromatic Erythroblast	2
Megakaryocytes	<0,2
Lymphocytes	16,2
Plasma cells	2,3
Reticular cells	0,4

Nucleated cells Percentage in the Bone Marrow

Normal human peripheral blood lymphocytes 20-47% (n=150% confidence interval)

Nucleated cells Percentage in Peripheral Blood of a Normal Patient

Cell type	Percentage	Number of event.
Red Blood Count		3,8 - 5,6 X10 ⁶ /µL
Platelets		150 - 450 X10 ³ /µL
White Blood Count		4.3 - 10.0 X10 ³ /µL
(WBC)		
Neutrophils	57 – 67 %	1,5 - 7.0 X10³/µL
Lymphocytes*	25 – 33 %	1.0 - 4.8 X10 ³ /μL
T cell	56 – 82 % of lymphocytes	
T cell CD4+	60 % of T cells	
T cell CD8+	40 % of T cells	
Cell NK+	6 – 33 of lymphocytes	
B cell	7.7 – 22 of lymphocytes	
Monocytes	3 – 7 %	0.28 - 0.8 X10 ³ /µL
Eosinophils	1 – 3 %	0.05 – 0,25 X10 ³ /µL
Basophils	0 – 0,075 %	0,015 – 0,05
		X10 ³ /µL
Reticulocyte	0,5 – 1,5 % of total Red	
	Blood Cell	

Expected values for pediatrics and adolescents have not been established.

The values obtained from normal individuals may vary from laboratory to laboratory; therefore, it is recommended that each laboratory establish its own normal range.



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Cancer Research Center (C.I.C.)
Campus Miguel de Unamuno
37007 Salamanca (Spain)Tel. / Fax:(+34) 923 294 827E-mail:info@immunostep.com

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11. PERFORMANCE CHARACTERISTICS.

a. <u>SPECIFICITY</u>

Blood samples were obtained from healthy normal donors of Caucasian were stained with Immunostep CDIO FITC monoclonal antibody. Cells contained in the lymphocyte, monocyte and granulocyte regions were selected for analysis. Blood samples were processed by a leukocyte method, with a direct immunofluorescence staining for flow cytometric analysis.

To evaluate the reagent's Specificity (cross-reactivity with other cell populations), 10 blood samples from healthy donors were studied, stained with an adequate isotype control and the MAb to study. The percentage of lymphocytes, monocytes and granulocytes stained with the mentioned MAb was evaluated. The results obtained are shown in the following table:

Case Summaries

	% Lymphocytes	% Monocytes	% Granulocytes
1	2,01	40,52	87,09
2	2,51	48,25	87,17
3	1,36	55,50	98,52
4	3,38	46,99	98,93
5	,69	39,17	97,63
6	2,47	47,09	85,08
7	1,40	62,35	92,14
8	2,12	51,60	95,37
9	1,15	44,98	96,73
10	,94	46,45	94,53
Total N	10	10	10

Statistics

-	-	% Lymphocytes	% Monocytes	% Granulocytes			
N	Valid	10	10	10			
	Missing	0	0	0			
Mear	ı	1,8030	48,2900	93,3190			
Medi	an	1,7050	47,0400	94,9500			
Mode	2	,69ª	39,17ª	85,08ª			
Std. D	Deviation	,83955	6,84466	5,16729			
Varia	ince	,705	46,849	26,701			
Rang	e	2,69	23,18	13,85			

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Minimum	,69	39,17	85,08
Maximum	3,38	62,35	98,93

b. <u>SENSIBILITY</u>

Sensitivity of the Immunostep CDIO monoclonal antibodies was determined by staining a blood sample from donor. Dilutions of a peripheral blood sample were made to check the concentration scale of stained cells obtained. The results show an excellent correlation level between the results obtained and expected based on the dilution used.

To determine the consistency of the conjugated monoclonal antibody as opposed to small variations (but deliberate). It provides an indication of its reliability during its normal use

Case Summaries									
	Sample	Dilution	% Expected	% Obtained					
1	400A + 0B	100,0	68,23	68,23					
2	350A + 50B	87,5	59,70	60,56					
3	300A + 100B	75,0	51,11	51,31					
4	250A + 150B	62,5	42,64	48,21					
5	200A + 200B	50,0	34,11	36,05					
6	150A + 250B	37,5	25,58	23,11					
7	100A + 300B	25,0	17,05	19,71					
8	50A + 350B	12,5	8,52	8,69					
9	0A + 400B	,0	,00	,00					
Total N	9	9	9	9					

Model Summarv

-	Model Samilary							
				Adjusted R	Std. Error of the			
	Model	R	R Square	Square	Estimate			
	1	,996ª	,991	,990	2,34014			

a Predictors: (Constant), Obtained



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c. <u>REPRODUCIBILITY</u>

Reproducibility for the Immunostep CDIO: FITC-conjugated monoclonal antibodies was determined by performing 10 replicated determinations of each antibody in each of three CDIO+ ranges, high, medium and low. Thus, a total of 30 determinations were performed for each form of CDIO. In this manner, reproducibility was demonstrated throughout the entire measuring range.

The 10 determinations for each range were performed by the staining, processing and analysis of 10 separate samples. Lymphocytes were selected for the analysis of percent cells stained in each of the three ranges.

To perform this study, anticoagulated blood was obtained from a normal donor expressing a high percentage of CDIO+ cells. Mid-range and low range samples were obtained by mixing known CDIO- cells in appropriate ratios, while maintaining the same total cell concentration for the three ranges.

The study was performed in each of three independent laboratories, in the manner that each laboratory obtained, stained and analyzed separate blood samples.

	High	Medium	Low
1	73,20	58,17	46,21
2	75,36	58,63	46,98
3	72,17	57,24	47,48
4	72,50	58,84	46,66
5	74,20	57,84	47,30
6	72,67	58,22	46,80
7	74,39	57,07	47,61
8	74,02	57,90	46,09
9	73,81	56,33	46,39

Case Summaries(a)



10		72,66	56,10	46,13
Total	10	10	10	

Descriptive Statistics

	N	Minimum	Maximum	Mean	Std. Deviation	variation coefficient
High	10	72,17	72,17	73,4980	1,01989	1,040
Medium	10	56,10	56,10	57,6340	,92600	,857
Low	10	46,09	46,09	46,7650	,56543	,320
Valid N (listwise)	10					

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