KOMBITEST CD3 FITC / **HLA-DR PE**

Cat.No. ED7055

1. Intended purpose

The KOMBITEST CD3 FITC / HIA-DR PE reagent is designed for percentage enumeration of mature human activated T lymphocytes in erythrocyte-lysed whole blood using Flow Cytometry

2. Test principle

This test is based on the specific binding of monoclonal antibodies to the antigenic determinants expressed on the surface of leukocytes. The monoclonal antibodies are labeled with different fluorochromes which are excited via laser beam from a flow cytometer during analysis. Subsequent emissions of light from the fluorochromes of each cell are collected and analyzed by a flow cytometer. The fluorescence intensity differences enable the separation of cell subsets based on the expression of analyzed antigens.

The specific staining of blood cells is performed by the incubation of blood samples with the reagent followed by a lysis of red blood cells. Afterwards, unaffected leukocytes are subjected to analysis by a flow cytometer.

3. Reagents provided

The reagent contains a premixed combination of mouse monoclonal antibody against human CD3 antigen (clone UCHT1) labeled with Fluorescein isothiocyanate (FITC), and mouse monoclonal antibody against human HLA-DR antigen (clone MEM-12) labeled with R-Phycoerythrin (PE). Labeled antibodies are diluted at optimum concentration in stabilizing phosphate buffered saline (PBS) solution containing 15mM sodium azide. The content of a vial (1 ml) is sufficient for 50 tests.

Product specification

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Content	50 tests, 1 ml	
Usage	20 µl per test	
Specificity	CD3	HLA-DR
Clone	UCHT1	MEM-12
Isotype (mouse)	lgG1	lgG1
Fluorochrome	FITC	PE
λ excitation	488 nm	488 nm
Emission maximum	525 nm	575 nm

4. Materials required but not provided

Test tubes for blood staining (e.g. 12 × 75 mm) EXCELLYSE Easy lysing solution

(Cat.No. ED7066)

. PBS buffer Deionized water

5. Equipment required

Automatic pipettes with disposable tips

Vortex mixer

Flow cytometer with excitation laser 488 nm and proper filters

6. Storage and handling

Store the vial at 2-8 °C. Keep away from sunlight. Do not freeze. Do not aliquote. Expiration date is stated on a vial label and on outer packaging.

7. Warnings, precautions and limitations of use

- Intended for In Vitro Diagnostic use in laboratories outside USA and Canada. This CE-IVD kit is in conformity with the European Directive 98/79/FC
- Do not use reagent after expiration date.
- Avoid reagents contamination. Avoid prolonged exposure to light.
- The content of the vial must not freeze.
- The flow cytometer should be calibrated on a routine basis using fluorescent microbeads to ensure stable sensitivity of detectors. Any non-performance of the staining protocol
- may produce false results.
- The reagent contains sodium azide (NaN3) which is highly toxic in pure form. However, the concentration in the reagent (15mM) is not considered as hazardous. When disposing the reagent, flush the sink with a large volume of water.
- Concentrations of labeled antibodies in this reagent were optimized to offer the best specific signal/non-specific signal ratio. Therefore, it is important to adhere to the reagent volume/sample volume ratio in every test. Do not dilute the reagent.

- · Do not use reagent volumes other than specified in this IFU.
- Blood samples are considered as potentially infectious and must be handled with care. Avoid all contact of the sample with the skin, eves and mucosa.
- In case of a hyperleukocytose sample, it is recommended to dilute the blood sample with PBS to obtain leukocyte density approximately 5×10^6 leukocytes/ml. Blood samples from abnormal patients may
- exhibit abnormal values of positive cells.
- Data may be incorrectly interpreted if fluorescent signals were compensated wrongly or if gates were positioned inaccurately.
- Flow cytometer may produce false results if the device has not been aligned and maintained appropriately.

8. Specimen

Use the peripheral human blood in a sterile tube with an anticoagulant (Heparin or EDTA).

Blood must be stored at room temperature

Use the blood sample no later than 48 hours after collection.

9. Procedure

Staining protocol

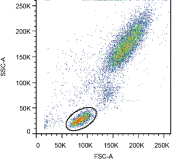
- 1. Add 20 μI of KOMBITEST CD3 FITC / HLA-DR PE reagent to a test tube. Add 100 μ l of blood sample to the tube.
- Vortex the tube
- Incubate the tube for 15 20 minutes at 3. room temperature in the dark. Perform lysis of red cells using EXCELLYSE
- 4. Easy lysing solution (Cat.No. ED7066) or any other commercial lysing solution containing formaldehyde as a fixative using Lyse/Wash protocol. Follow the instructions of the lysing . solution manufacturer. 5.
- Analyze the sample immediately using a flow cytometer or store sample at 2 8 °C in the dark and analyze within 24 hours provided that cells were fixed.

Flow Cytometric Analysis

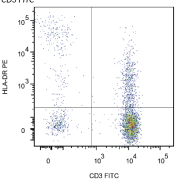
Analyze stained samples using a flow cytometer with excitation laser 488 nm and proper filters. Compensate fluorescent signals prior to or after data acquisition

Visualize compensated data on the side-scatter (SSC) versus forward-scatter (FSC) plot. Set the gate for lymphocyte population (Figure 1). Alternatively, set the optimal lymphocyte gate using KOMBITEST CD45 FITC / CD14 PE (Cat. No. ED7056, refer to the datasheet for lymphocyte gate assessment procedure).

Fig. 1: Delimitation of lymphocyte population



Then make a CD3 FITC versus HLA-DR PE dot-plot of lymphocyte population (Figure 2). Separate populations using appropriate gate and calculate the percentage of activated T lymphocytes situated in upper-right quadrant (CD3+HLA-DR+ subpopulation) on the dot-plot. Fig. 2: Lymphocytes in a dot-plot HLA-DR PE vs. CD3 FITC



10. Analytical performance

Specificity

The antibody UCHT1 recognizes the CD3 antigen of the TCR/CD3 complex on mature human T cells. The UCHT1 antibody reacts with the epsilon chain of the CD3 complex.

HLDA I; WS Code T 3 HLDA III; WS Code T 126

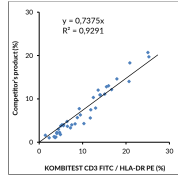
HLDA III; WS Code T 471 HLDA VI; WS Code T 6T-CD3.1

The antibody MEM-12 recognizes common epitope on human HLA-DR which is dependent on the association of alpha and beta chains. DR is the isotype of human MHC Class II molecules expressed on antigen-presenting cells, mainly on B lymphocytes, monocytes, macrophages, activated T lymphocytes, and activated NK lymphocytes.

Accuracy

The accuracy of the method was studied by the comparison of KOMBITEST CD3 FITC / HLA-DR PE with competitor's product in parallel staining of 43 blood samples. The regression analysis is given below.

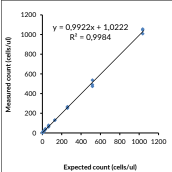
Regression Analysis of CD3+HLA-DR+ Lymphocytes



Linearity

The linearity of the method was determined on 10 serial dilutions of leukocyte-enriched blood sample (buffy coat). Cell samples were stained by KOMBITEST CD3 FITC / HLA-DR PE in triplicates. Measured and expected values were expressed in terms of absolute count (cells/µl) in graphs given below.

Range of CD3+HLA-DR+ Lymphocytes



Repeatability

The repeatability of the assay was measured on stabilized blood sample (Immuno-Troll™ Cell Beckman-Coulter) in ten tubes in parallel. Coefficient of variation (CV) is given in the table below.

Lymphocyte Subset	n	AVG	SD	cv
CD3+HLA-DR+ (%)	10	8.8	0.7	8.4

Reproducibility

The reproducibility of the assay was measured on stabilized blood sample (Immuno-Troll[™] Cells, Beckman-Coulter) under the same conditions for four weeks. Coefficient of variation (CV) is given in the table below.

Lymphocyte Subset	n	AVG	SD	cv
CD3+HLA-DR+ (%)	14	8.1	1.2	15.2

11. Clinical performance

Expected values

Results obtained in different laboratories may vary. Each laboratory should establish a normal range of cell subsets using its own test conditions. Results obtained in our laboratory are given in the table below.

Lymphocyte Subset	n	Mean	95% Range
CD3+HLA-DR+ (%)	43	9.6	2.3 - 25.0

12. References

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Huang Y, Wange RL: T cell receptor signaling: beyond complex complexes. J Biol Chem. 2004 Jul 9;279(28):28827-30.

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Leukocyte Typing III., McMichael A. J. et al (Eds.), Oxford University Press (1987).

Leukocyte Typing VI., Kishimoto T. et al. (Eds.), Garland Publishing Inc. (1997).

13. Manufacturer

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14. Trademarks

Immuno-Troll[™] Cells is registered trademark of Beckman-Coulter.

15. Revision History

 Version 1, ED7055_IFU_v1 Initial Release

• Version 2, ED7055_IFU_v2

The text removed in in the reagent provided: "0.2% (w/v) high-grade protease free Bovine Serum Albumin (BSA) as a stabilizing agent" Version 3 ED7055 IEU v3

The company logo changed. IFU layout changed. "Keep away from sunlight", "Do not aliquote" and "Expiration date is stated on a vial labels and on outer packaging" added in Storage section. Texts "Do not dilute the reagent" and "Do not use reagent volumes other than specified in this

IFU" were added in warning section. Staining protocol specified to the use o Excellyse Easy lysing solution (Cat.No. ED7066). of



KOMBITEST CD3 FITC / HLA-DR PE

50 tests | Cat.No. ED7055



Instructions for Use

Version: ED7055_IFU_v3_EN Date of Issue: 24-02-2020

EN

Symbols

REF	Catalogue number
LOT	Batch code
\square	Use-by date
X	Temperature limits
淤	Keep away from sunlight
IVD	In vitro diagnostic medical device
CE	CE marking of conformity
i	Consult instructions for use
***	Manufacturer

The product is intended for In Vitro Diagnostic Use. In vivo diagnostic or therapeutic applications are strictly forbidden. Products shall not be used for resale or transfer to third parties either as a stand-alone product or

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