

# KOMBITEST TBNK 6-color (RUO) 50 tests | Cat. No. ED7766

RU0

Not for use in diagnostic or therapeutic procedures.

# Technical Data Sheet (EN)

Version: ED7766\_TDS\_v1\_EN Date of Issue: 20-03-2023

#### Symbols used in the product labeling

RUO	Research Use Only
***	Manufacturer
[]i	Consult instructions for use
Σ	Contains sufficient for <n> tests</n>
REF	Catalogue number
LOT	Batch code
Ω	Use by date
1	Temperature limit
*	Keep away from sunlight

# 1. Intended Purpose

KOMBITEST TBNK 6-color is intended for detection and enumeration of lymphocyte populations and subsets in human whole blood by flow cytometry.

#### What is detected and/or measured

The product KOMBITEST TBNK 6-color detects and measures relative percentages and absolute counts of human T cells (CD3+), B cells (CD3-CD19+), NK cells (CD3-CD16+56+), helper/inducer (CD3+CD4+) and suppressor/cytotoxic (CD3+CD8+) T cell subsets.

## Context of a physiological or pathological state

Frequencies of lymphocyte populations measured by the product can be affected by various pathological conditions and evaluation of their percentages and counts can be used in the assessment of:

- human immunodeficiency virus (HIV) infection progression (1, 4, 7, 9)
- hereditary immunodeficiencies (2, 3, 4, 11, 12, 15, 17)
- autoimmune diseases (5, 6)
- defects in innate immune defense (13, 14)

#### Type of assay

Not automated

Quantitative

# Type of specimen required

Human anticoagulated peripheral whole blood specimen

# **Testing population**

Not intended for a specific population.

# 2. Intended user

The product is intended for professional laboratory use only.

## Requirements on qualification

Intended user shall have a state-of-the-art expertise in flow cytometry analysis of human cells, standard laboratory techniques, including pipetting skills, safe and proper handling of specimens derived from the human body.

# 3. Test principle

The test principle is based on the detection of monoclonal antibody binding to a specific molecule (antigen) expressed by certain human blood cells. Monoclonal antibodies used in the test are labeled with different fluorochromes which are excited by a laser beam from a flow cytometer during acquisition of an antibody-stained blood specimen. Subsequent fluorescence (light emission) from each

fluorochrome present on an acquired blood cell is collected and analyzed by the instrument. Fluorescence intensity is directly proportional to the antigen expression density in a cell allowing for separation of different cell subsets.

# 4. Reagent(s) provided

#### Contents

The product KOMBITEST TBNK 6-color is sufficient for 50 tests and is provided with the following reagent:

1 vial (1 ml) containing a premixed combination of fluorochrome-labeled monoclonal antibodies CD3 FITC / CD16 PE + CD56 PE / CD45 PerCP-Cy $^{\text{TM}}$ 5.5 / CD4 PE-Cy $^{\text{TM}}$ 7 / CD19 APC / CD8 APC-Cy $^{\text{TM}}$ 7, diluted at optimum concentrations in a stabilizing phosphate buffered saline (PBS) solution containing 15mM sodium azide.

#### Composition

 Table 1
 Description of active components

Antigen	Flurochrome	Clone	Isotype	Concentration (μg/ml)
CD3	FITC	TB3	lgG2b	2
CD4	PE-Cy™7	MEM-241	lgG1	1.5
CD8	APC-Cy™7	LT8	lgG1	1.8
CD16	PE	3G8	lgG1	1.5
CD56	PE	LT56	lgG2a	1.5
CD19	APC	LT19	lgG1	2
CD45	PerCP-Cy™5.5	MEM-28	lgG1	3

# 5. Materials required but not provided

12 x 75 mm round bottom test tubes

Erythrocyte lysing solution (EXCELLYSE Easy, EXBIO Praha, a.s.)

Deionized water (Reagent-grade)

Process control cells (Streck CD-Chex Plus®, Cat. No. 213323 or equivalent lysable cell control)

# 6. Equipment required

Automatic pipette with disposable tips (20 - 100  $\mu\text{l})$  for pipetting specimen and reagents

Liquid dispenser or pipette with disposable tips (0.5 – 2 ml) for dispensing erythrocyte lysing solution

Vortex mixer

Hematology analyzer (for absolute cell counts) capable of white blood cell (WBC) and lymphocyte count per  $\mu$ I of specimen

Flow cytometer with two laser excitation sources (488 nm and ~635 nm), detectors for scattered light, optical filters and emission detectors appropriate to collect signals from fluorochromes provided in Table 2.

Flurochrome	Excitation [nm]	Emission [nm]
FITC	488	525
PE	488	576
PerCP-Cy™5.5	488	695
PE-Cy™7	488	780
APC	630 - 640	660
APC-Cy™7	630 - 640	780

 Table 2
 Spectral characteristic of fluorochromes used in the product

**NOTICE:** The product was tested on flow cytometers BD FACSCanto™ II (BD Biosciences), BD FACSLyric™ (BD Biosciences), Navios EX (Beckman Coulter), DxFLEX (Beckman Coulter) and Sysmex™ XF-1600 (Sysmex Corporation).

# 7. Storage and handling

Store at 2-8 °C.

Avoid prolonged exposure to light.

Do not freeze.

See Section 10 Procedure (Reagent Preparation) for information about In-Use stability and shelf-life following the first opening, together with the storage conditions and stability of working solutions (where applicable).

# 8. Warnings, precautions and limitations of use

#### **GHS Hazard Classification**

Consult Safety Data Sheet (SDS) available on the product page at www.exbio.cz for the full information on the risks posed by chemical substances and mixtures contained in the Product and how they should be handled and disposed.

## **Biological Hazard**

Human biological samples and blood specimens and any materials coming into contact with them are always considered as infectious materials.

Use personal protective and safety equipment to avoid contact with skin, eyes and mucous membranes.

Follow all applicable laws, regulations and procedures for handling and disposing of infectious materials.

#### **Evidence of deterioration**

Normal appearance of the reagent provided is a clear liquid. Do not use the reagent if you observe any change in appearance, for example turbidity or signs of precipitation.

#### Limitation of use

Do not use after the expiry date stated on the product labels.

# 9. Specimen

Use venous peripheral blood collected into specimen receptacle classified as a medical device, with the anticoagulant EDTA.

**NOTICE:** Determine WBC absolute cell count and lymphocyte count in the collected blood specimen by a hematology analyzer. The product KOMBITEST TBNK 6-color alone does not provide enumeration of absolute cell counts.

Blood specimen with WBC count exceeding  $40x10^3$  cells/ $\mu$ l will require dilution with PBS before sample processing.

Process the blood specimen no later than 24 hours after collection.

#### 10. Procedure

## Preparation of reagent(s) provided

No reagent preparation is necessary.

Bring the reagent to the room temperature prior to use. Keep the product primary container dry.

Use the reagent directly from its original primary container. Time, for which the reagent is in use (exposed to light and elevated temperature), shall not exceed 4 hours per day.

Following the first opening, the reagent retains its performance characteristics until the expiry date when stored under the stated conditions in its original primary container.

**CAUTION:** Do not dilute the reagent.

## Preparation of materials required but not provided

Dilute concentrated erythrocyte lysing solution with deionized water according to the manufacturer's instructions. Diluted (1X) erythrocyte lysing solution is stable for 1 month when stored in a liquid dispenser or closed container at room temperature.

#### **Quality control**

Use Streck CD-Chex Plus® or equivalent control cells as positive procedural control to ensure proper performance of the product as intended. Streck CD-Chex Plus® provides established values for percent positive and absolute counts of T cells, B cells, granulocytes, monocytes and NK cells, including two clinically relevant levels of CD4+ cells.

Stain the control cells using KOMBITEST TBNK 6-color reagent according to sample processing as specified in the TDS. Verify that the obtained results (% Positive Cells) are within the Expected range reported for the used lot of control cells.

# Specimen staining

- 1. For each specimen, label a  $12 \times 75$  mm round bottom test tube with the appropriate sample identification.
- 2. Pipette 20  $\mu$ I of KOMBITEST TBNK 6-color reagent into the bottom of the 12 x 75 mm test tube.
- 3. Pipette 50  $\mu l$  of well-mixed blood specimen to the bottom of the test tube.

**CAUTION:** Avoid pipetting blood on the side of the test tube. If blood smear or droplet remains on the side of the tube, it may not be stained with the reagent or erythrocytes may not be lysed and the test result may not be valid.

- 4. Vortex and incubate the test tube for 20 minutes at room temperature in the dark.
- 5. Add 500  $\mu$ l of diluted (1X) lysing solution to the test tube.
- 6. Vortex and incubate the test tube for 10 minutes at room temperature in the dark.

Acquire the stained sample immediately on the flow cytometer. If the stained sample will not be acquired immediatelly, store at 2-8 °C in the dark and analyze within 24 hours.

**CAUTION:** Vortex the stained sample immediately before acquisition on the flow cytometer to avoid aggregates.

#### Flow cytometry analysis

The flow cytometer selected for use with the product KOMBITEST TBNK 6-color shall be calibrated on a routine basis using fluorescent microbeads to ensure stable sensitivity of detectors according to the cytometer manufacturers instructions.

If not maintained properly the flow cytometer may produce false results.

Refer to the manufacturer's cytometer specifications for lasers and fluorescence detectors according to the excitation and emission characteristics of the fluorochromes in Section 6 Equipment required.

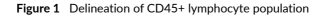
Set voltages on the fluorescence detectors of interest prior to stained specimen analysis. Voltage on a PMT detector should be set high enough, so that minimum of negatively stained events interfere with 0th channel on the fluorescence axis. Also, PMT detector voltage should not exceed values at which positive events are pressed to the right axis.

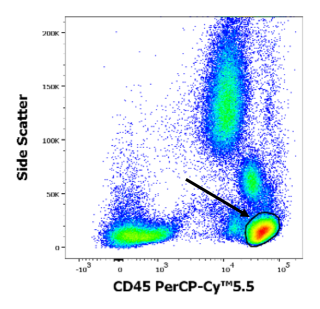
Compensate fluorescence signals between detectors prior to or after data acquisition. Data may be incorrectly interpreted if fluorescence signals are compensated improperly or if gates are positioned inaccurately.

For measured data analysis, it is possible to use cytometer software developed by the manufacturer, or software dedicated for offline cytometry data analysis (for example FlowJo™, VenturiOne®, Infinicyt™).

## Data analysis of the KOMBITEST TBNK 6-color stained specimen

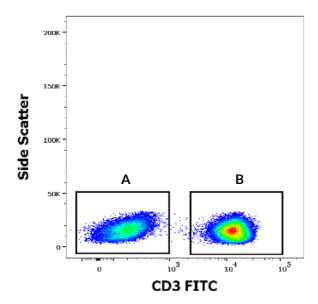
Visualize compensated data in a side-scatter (SSC) versus CD45 PerCP-Cy<sup>™</sup>5.5 plot. Set the gate for CD45+ lymphocyte population as shown in Figure 1.





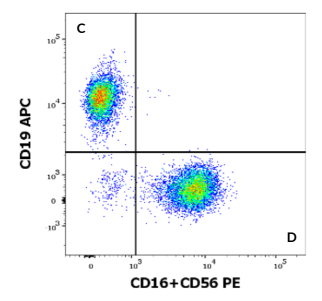
Plot the gated CD45+ lymphocytes in a side-scatter (SSC) versus CD3 FITC plot as shown in Figure 2. Separate CD3+ and CD3- lymphocytes using appropriate gates. Calculate the percentage of T cells (CD3+; region B on the Figure 2) from all lymphocytes.



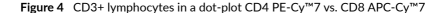


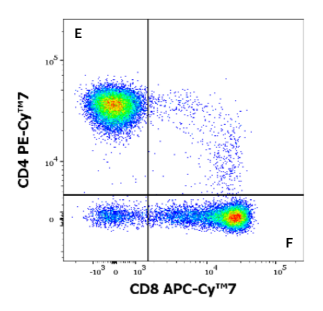
Plot the gated CD3- lymphocytes (region A on the Figure 2) as CD19 APC versus CD16+CD56 PE as shown in Figure 3. Set appropriate gates and calculate the percentage of B cells (CD16-CD56-CD19+; region C on the Figure 3) and natural killer (NK) cells (CD16+CD56+CD19-; region D on the Figure 3) from all lymphocytes.

Figure 3 CD3- lymphocytes in a dot-plot CD19 APC vs. CD16+CD56 PE



Plot the gated T cells (CD3+; region B on the Figure 2) as CD4 PE-Cy<sup>™</sup>7 versus CD8 APC-Cy<sup>™</sup>7 as shown in Figure 4. Set appropriate gates and calculate the percentage of helper/inducer T cells (CD4+CD8-; region E on the Figure 4) and suppressor/cytotoxic T cells (CD4-CD8+; region F on the Figure 4) from all lymphocytes.





## **Calculation and interpretation of analytical results**

To have absolute counts, use the absolute lymphocyte count as determined by a hematology analyzer. Refer to hematology analyzer manufacturer's instructions. Use the equations below for absolute count enumeration of required lymphocyte subset.

Ax 
$$\frac{B(\%)}{100(\%)}$$
 = Absolute count of required lymphocyte subset

A = absolute lymphocyte count (data from hematology analyzer; cells /  $\mu$ l)

B = relative percentages of required lymphocyte subset from all lymphocytes (data from flow cytometer; %)

# 11. Analytical performance

**NOTICE:** All analytical performance data were measured using erythrocyte lysing solution EXCELLYSE Easy (EXBIO Praha, a.s., Cat. No. ED7066).

#### Specificity

The antibody TB3 recognizes human CD3 antigen of the TCR/CD3 complex. Specificity of the antibody has been confirmed by HCDM Council (HLDA X and HLDA XI workshop).

The antibody MEM-241 recognizes human CD4 antigen (T cell surface glycoprotein CD4). Specificity of the antibody has been confirmed by HCDM Council (HLDA VIII workshop).

The antibody LT8 recognizes human CD8 antigen (disulfide-linked dimer expressed as two CD8 alpha chain homodimers or CD8 alpha/beta chain heterodimers). Specificity of the antibody has been confirmed by HLDA workshops (HLDA V workshop<sup>(16)</sup> and HLDA VII workshop<sup>(8)</sup>).

The antibody 3G8 recognizes human CD16 antigen (low affinity immunoglobulin type III Fc-gamma receptor). Specificity of the antibody has been confirmed by HLDA workshop (HLDA V workshop (16)).

The antibody LT56 recognizes the leukocyte isoform of human CD56 antigen (Neural cell adhesion molecule 1). Specificity of the antibody has been confirmed by HCDM Council (HLDA X workshop).

The antibody LT19 recognizes human CD19 antigen (B cell transmembrane glycoprotein CD19). Specificity of the antibody has been confirmed by HCDM Council (HLDA X workshop).

The antibody MEM-28 recognizes all leukocyte isoforms of human CD45 (Protein tyrosine phosphatase receptor type C). Specificity of the antibody has been confirmed by HLDA workshop (HLDA III workshop (10)).

## Accuracy

Accuracy of the method was determined as a comparison of the product KOMBITEST TBNK 6-color with similar products available on the market or with other well-documented methods by parallel staining of 30 healthy donors and 134 patients suspected of having immune system pathological condition. Linear regression analysis parameters are provided in Table 3 and 4.

**Table 3** Linear regression analysis for lymphocyte subsets in healthy donors (comparison of the product KOMBITEST TBNK 6-color with product BD Multitest™ 6-Color TBNK Reagent (Cat. No. 644611))

Lymphocyte Subset	Unit	n	Slope	Intercept	R <sup>2</sup>	Range
CD3+	%	30	1.00	-0.006	0.99	49.03 - 84.87
CDS	cells/µl	30	0.99	5.464	1.00	608 - 2137
CD3+CD8+	%	30	0.98	0.003	1.00	10.43 - 40.17
CD31CD01	cells/μl	30	0.97	8.959	1.00	145 - 1016
CD3+CD4+	%	30	0.99	0.004	0.99	29.70 - 56.37
CD31CD41	cells/µl	30	0.99	6.588	1.00	321 - 1407
CD3-CD16+CD56+	%	30	1.01	-0.002	1.00	5.15 - 38.93
CD3-CD10+CD30+	cells/µl	30	1.02	-6.186	1.00	97 - 1036
CD3-CD19+	%	30	1.00	0.003	0.99	5.42 - 25.00
CDG CD171	cells/μl	30	1.02	1.611	0.99	74 - 352

n = number of blood samples

Table 4 Linear regression analysis for lymphocyte subsets in patients suspected of having immune system pathological conditions (comparison of the product KOMBITEST TBNK 6- color with AQUIOS CL Flow Cytometry System - Beckman Coulter, Inc. and accredited clinical laboratory in-house method - a cocktail of single color conjugated antibodies from different manufacturers and analysis on the BD FACSCanto™ II)

Lymphocyte Subset	Unit	n	Slope	Intercept	R <sup>2</sup>	Range
CD3+	%	134	1.032	-2.655	0.98	23.9 - 94.5
CDOT	cells/μl	134	1.023	-0.047	0.97	140 - 5178
CD3+CD8+	%	134	1.020	-0.803	0.98	9.1 - 80.7
CDSTCDOT	cells/μl	134	1.055	-0.041	0.96	60 - 3546
CD3+CD4+	%	134	1.014	-0.651	0.98	1.4 - 67.5
CD3+CD4+	cells/μl	134	0.994	-0.005	0.98	8 - 2826
CD3-CD16+CD56+	%	134	1.064	-0.400	0.98	1.6 - 68.2
CD3-CD10+CD30+	cells/μl	134	1.080	-0.014	0.99	10 - 2612
CD3-CD19+	%	134	1.027	-0.376	0.99	0.0 - 69.7
CDU CD171	cells/µl	134	1.043	-0.010	1.00	0 - 4586

## Linearity

The linearity of the method was verified on 10 serial dilutions of a leukocyte-enriched blood sample (buffy coat). Cell samples were stained with KOMBITEST TBNK 6-color in hexaplicates. Samples were analyzed using BD FACSCanto™ II flow cytometer and Beckman Coulter DxFLEX flow cytometer. Measured data for the indicated lymphocyte subsets were observed to be linear across the lymphocyte range 333 - 9492 cells/µl using BD FACSCanto™ II and 309 - 8693 cells/µl using Beckman Coulter DxFLEX. Cell subsets were in the ranges found in Tables 5 and 6.

**Table 5** Linear ranges of lymphocyte subsets analysed by BD FACSCanto™ II

BD FACSCanto™ II				
Lymphocyte Subset	Range (cells/μl)			
CD3+	249 - 6594			
CD3+CD8+	96 - 2560			
CD3+CD4+	136 - 3628			
CD3-CD16+CD56+	55 - 1525			
CD3-CD19+	44 - 1342			

 Table 6
 Linear ranges of lymphocyte subsets analysed by Beckman Coulter DxFLEX

Beckman Coulter DxFLEX				
Lymphocyte Subset	Range (cells/μl)			
CD3+	243 - 6565			
CD3+CD8+	102 - 2652			
CD3+CD4+	128 - 3517			
CD3-CD16+CD56+	64 - 1588			
CD3-CD19+	41 - 1280			

# Repeatability

The repeatability of the assay was measured on ten blood samples in hexaplicates. Samples were analyzed using BD FACSCanto™ II flow cytometer and Beckman Coulter DxFLEX flow cytometer. Coefficients of variation (CV) are provided in the tables below (Table 7 and 8).

 Table 7
 Repeatability of the product on BD FACSCanto™ II

BD FACSCanto™ II							
Lymphocyte Subset	Lymphocyte Subset Unit n Average SD %CV						
CD3+	%	10	70.34	0.56	0.91		
CD31	cells/µl	10	1396	10.22	0.91		
CD3+CD8+	%	10	23.11	0.27	1.25		
СВЭТСВОТ	cells/µl	10	453	5.23	1.25		
CD3+CD4+	%	10	41.06	0.53	1.36		
CDSTCD41	cells/µl	10	808	9.71	1.36		
CD3-CD16+CD56+	%	10	16.35	0.40	2.43		
CD3-CD10+CD30+	cells/μl	10	289	7.20	2.43		
CD3-CD19+	%	10	11.63	0.25	2.31		
CD3 CD171	cells/μl	10	227	4.78	2.31		

 Table 8
 Repeatability of the product on Beckman Coulter DxFLEX

Beckman Coulter DxFLEX						
Lymphocyte Subset Unit n Average SD %CV						
CD3+	%	10	70.80	0.61	0.95	
CD31	cells/µl	10	1406	11.19	0.95	
CD3+CD8+	%	10	23.80	0.32	1.42	
CDSTCDST	cells/µl	10	468	6.12	1.42	
CD3+CD4+	%	10	40.81	0.56	1.47	
CD3+CD4+	cells/µl	10	803	10.52	1.47	
CD3-CD16+CD56+	%	10	15.89	0.40	2.72	
CD3-CD10+CD30+	cells/μl	10	282	7.23	2.72	
CD3-CD19+	%	10	11.68	0.32	2.83	
CD3-CD17+	cells/μl	10	227	5.94	2.83	

## Reproducibility

The reproducibility of the assay was measured on 2 stabilized blood samples (CD-Chex Plus® and CD-Chex Plus® CD4 Low) under the same conditions for 15 days using 3 lots of the product (5 days each). Samples were analyzed using BD FACSCanto™ II flow cytometer and Beckman Coulter DxFLEX flow cytometer. Coefficients of variation (CV) are given in the tables below (Table 9 and 10).

**Table 9** Reproducibility of the product on BD FACSCanto™ II

Lymphocyte Subset	Material	Unit	Average	SD	%CV
	CD-Chex Plus®	%	76.84	0.18	0.23
CD3+	CD-Cliex Flus	cells/μl	1896	4.39	0.23
CD3+	CD-Chex Plus®	%	60.61	0.32	0.53
	CD4 Low	cells/μl	879	4.65	0.53
	CD-Chex Plus®	%	23.45	0.23	0.97
CD3+CD8+	CD-Cliex Flus®	cells/μl	578	5.62	0.97
CDSTCDST	CD-Chex Plus®	%	42.17	0.31	0.73
	CD4 Low	cells/μl	612	4.55	0.73
	CD-Chex Plus®	%	48.78	0.45	0.93
CD3+CD4+		cells/μl	1203	11.15	0.93
CDOTCD41	CD-Chex Plus® CD4 Low	%	12.53	0.26	2.11
		cells/μl	182	3.84	2.11
	CD-Chex Plus®	%	10.76	0.22	2.03
CD3-CD16+CD56+	CD-Cliex Flus®	cells/μl	265	5.39	2.03
CD3-CD10+CD30+	CD-Chex Plus®	%	19.51	0.38	1.94
	CD4 Low	cells/μl	283	5.49	1.94
	CD-Chex Plus®	%	11.30	0.16	1.45
CD3-CD19+	CD-Cliex Plus®	cells/μl	279	4.03	1.45
CD3-CD17+	CD-Chex Plus®	%	18.05	0.32	1.75
	CD4 Low	cells/μl	262	4.58	1.75

 Table 10
 Reproducibility of the product on Beckman Coulter DxFLEX

Lymphocyte Subset	Material	Unit	Average	SD	%CV
	CD-Chex Plus®	%	77.17	0.21	0.27
CD3+	CD-Cliex Flus®	cells/μl	1904	5.23	0.27
CD31	CD-Chex Plus®	%	60.85	0.43	0.71
	CD4 Low	cells/μl	883	6.24	0.71
	CD-Chex Plus®	%	23.87	0.20	0.85
CD3+CD8+	CD-Cliex Flus®	cells/μl	589	4.99	0.85
CD31CD01	CD-Chex Plus®	%	42.81	0.32	0.75
	CD4 Low	cells/μl	621	4.65	0.75
	CD-Chex Plus®	%	46.47	1.41	3.03
CD3+CD4+	CD-Cliex Flus®	cells/μl	1146	34.77	3.03
CD31CD41	CD-Chex Plus® CD4 Low	%	12.16	0.53	4.37
		cells/μl	176	7.71	4.37
	CD-Chex Plus®	%	10.59	0.20	1.88
CD3-CD16+	CD-Cliex Flus®	cells/μl	261	4.92	1.88
CD56+	CD-Chex Plus®	%	19.38	0.32	1.63
	CD4 Low	cells/μl	281	4.59	1.63
	CD-Chex Plus®	%	11.07	0.17	1.54
CD0 CD40:	CD-Cliex Plus®	cells/μl	273	4.19	1.54
CD3-CD19+	CD-Chex Plus®	%	17.85	0.35	1.95
	CD4 Low	cells/μl	259	5.05	1.95

# 12. Expected values

#### Reference Interval

Reference intervals for the product KOMBITEST TBNK 6-color were determined in a cohort of subjects using erythrocyte lysing solution EXCELLYSE Easy (EXBIO Praha, a.s., Cat. No. ED7066) and the BD FACSCanto $^{\text{TM}}$  II flow cytometer. Subjects were healthy normal adults (blood donors).

LymphocyteSubset	Unit	n	Mean	95% Range
CD3+	%	30	69.49	50.81 - 88.17
CDS1	cells/μl	30	1308	573 - 2043
CD3+CD8+	%	30	22.68	9.29 - 36.07
CD31CD01	cells/μl	30	431	53 - 810
CD3+CD4+	%	30	42.70	27.72 - 57.67
CD3+CD4+	cells/µl	30	802	352 - 1252
CD3-CD16+CD56+	%	30	18.70	0.60 - 35.53
CD3-CD10+CD30+	cells/μl	30	351	0 - 822
CD3-CD19+	%	30	11.62	2.65 - 20.59
CD3 CD171	cells/µl	30	211	72 - 351

**Table 11** Representative reference intervals for the KOMBITEST TBNK 6-color

**CAUTION:** Indicated values using the product are intended to be representative only. Each laboratory must establish its own reference intervals from the local population of normal donors.

# 13. Interfering substances and limitations

The product KOMBITEST TBNK 6-color has not been validated for use in specimens collected with heparin or acid citrate dextrose (ACD) anticoagulants in determining relative and absolute counts.

The product KOMBITEST TBNK 6-color is not intended for screening and/or phenotyping of leukemia and lymphoma samples.

Absolute counts are not comparable between laboratories using different equipment from various manufacturers.

# 14. References

 Bensussan, A et al. Significant enlargement of a specific subset of CD3+CD8+ peripheral blood leukocytes mediating cytotoxic T-lymphocyte activity during human immunodeficiency virus infection. Proc Natl Acad Sci U S A. 1993 15;90(20):9427-30. doi: 10.1073/pnas.90.20.9427.

- 2) Boldt, A et al. Eight-color immunophenotyping of T-, B-, and NK-cell subpopulations for characterization of chronic immunodeficiencies. Cytometry B Clin Cytom 2014 May;86(3):191-206. doi:10.1002/cyto.b.21162.
- 3) de Saint Basile, G et al. Severe combined immunodeficiency caused by deficiency in either the delta or the epsilon subunit of CD3. J Clin Invest. 2004 Nov;114(10):1512-7. doi: 10.1172/JCl22588.
- 4) Giorgi, J V. Characterization of T lymphocyte subset alterations by flow cytometry in HIV disease. Ann N Y Acad Sci. 1993 Mar 20;677:417-9. doi: 10.1111/j.1749-6632.1993.tb38803.x.
- 5) Iwatani, Y et al. Decreases in alpha beta T cell receptor negative T cells and CD8 cells, and an increase in CD4+ CD8+ cells in active Hashimoto's disease and subacute thyroiditis. Clin Exp Immunol. 1992 Mar;87(3):444-9. doi: 10.1111/j.1365-2249.1992.tb03017.x.
- 6) Kucuksezer, U C et al. The Role of Natural Killer Cells in Autoimmune Diseases. Front Immunol. 2021 Feb 25;12:622306. doi: 10.3389/fimmu.2021.622306.
- 7) Li, Y et al. AIDS prevention and control in the Yunnan region by T cell subset assessment. PLoS One. 2019 Apr 18;14(4):e0214800. doi: 10.1371/journal.pone.0214800.
- 8) Mason, D et al, eds.: Leucocyte Typing VII: White Cell Differentiation Antigens: Proceedings of the Seventh International Workshop and Conference Held in Harrogate, United Kindom: Oxford University Press; 2002.
- 9) McCarty, B et al. Low Peripheral T Follicular Helper Cells in Perinatally HIV-Infected Children Correlate With Advancing HIV Disease. Front Immunol. 2018 Aug 24;9:1901. doi: 10.3389/fimmu.2018.01901.
- 10) McMichael AJ, ed. Leucocyte Typing III: 54 White Cell Differentiation Antigens. New York, NY: Oxford University Press; 1987.
- 11) Monafo, W J et al. A hereditary immunodeficiency characterized by CD8+ T lymphocyte deficiency and impaired lymphocyte activation. Clin Exp Immunol. 1992 Dec;90(3):390-3. doi: 10.1111/j.1365-2249.1992.tb05856.x.
- 12) North, M E et al. Primary defect in CD8+ lymphocytes in the antibody deficiency disease (common variable immunodeficiency): abnormalities in intracellular production of interferon-gamma (IFN-gamma) in CD28+ ('cytotoxic') and CD28- ('suppressor') CD8+ subsets. Clin Exp Immunol. 1998 Jan;111(1):70-5. doi: 10.1046/j.1365-2249.1998.00479.x.
- 13) Orange, J S. Natural killer cell deficiency. J Allergy Clin Immunol. 2013 Sep;132(3):515-525. doi: 10.1016/j.jaci.2013.07.020.
- 14) Orange, J S. How I Manage Natural Killer Cell Deficiency. J Clin Immunol. 2020 Jan;40(1):13-23. doi: 10.1007/s10875-019-00711-7.

- 15) Picat, M Q et al. T-cell activation discriminates subclasses of symptomatic primary humoral immunodeficiency diseases in adults. BMC Immunol. 2014 Mar 12;15:13. doi: 10.1186/1471-2172-15-13.
- 16) Schlossman SF, Boumsell L, Gilks W, et al, eds.: Leucocyte Typing V: White Cell Differentiation Antigens. New York, NY: Oxford University Press; 1995.
- 17) van Dongen, J J M et al. EuroFlow-Based Flowcytometric Diagnostic Screening and Classification of Primary Immunodeficiencies of the Lymphoid System. Front Immunol. 2019 Jun 13;10:1271. doi: 10.3389/fimmu.2019.01271.

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# **16. Revision History**

Version 1, ED7766\_TDS\_v1 Initial release

## 17. Manufacturer

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**NOTICE**: Any serious incident that has occured in relation to the product shall be reported to the manufacturer and the local competent authority.