

# Estimating Normal Values of Rare T-Lymphocyte Populations in Peripheral Blood of Healthy Cuban Adults

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## ABSTRACT

**INTRODUCTION** Flow cytometry allows immunophenotypic characterization of important lymphocyte subpopulations for diagnosis of diseases such as cancer, autoimmune diseases, immunodeficiencies and some infections. Normal values of rare lymphoid cells in blood, quantified by cytometry, vary among different populations; so it is indispensable to obtain normal national values that can be used in clinical practice.

**OBJECTIVE** Characterize distribution of rare T-lymphocyte populations in peripheral blood, specifically double-positive T, natural killer T and activated T lymphocytes, as well as their relationship to sex and age.

**METHODS** A cross-sectional study was carried out in 129 adults (68 women, 61 men) aged >18 years, without chronic diseases or unhealthy habits, who signed informed consent. Peripheral blood was collected for immunophenotyping of lymphocyte subpopulations with monoclonal antibodies specific for CD4<sup>+</sup>CD8<sup>+</sup> double-positive T cells, CD3<sup>+</sup>CD56<sup>+</sup> natural killer T cells, and CD3<sup>+</sup>CD25<sup>+</sup>HLA-DR<sup>+</sup> activated T cells. An eight-color flow cytometer (Beckman Coulter Gallios) was used. The analytic strategy was modified, associating variables of interest in a single graphic, using conventional monoclonal labeling antibodies. Medians and minimum and maximum percentiles (2.5 and 97.5, respectively) were used as descriptive statistics, stratified by

sex, for cell counts and percentages. A linear regression model was applied to assess age effects and a two-tailed Mann-Whitney *U* test for independent samples was used to assess sex differences. The significance threshold was set as  $p \leq 0.05$ .

**RESULTS** Median percentages of total lymphocytes: natural killer T cells 6.3% (1.4%–23%) in men and 4.7% (0.8%–11.3%) in women ( $p = 0.003$ ); activated T cells 1.0% (0.2%–2.2%) in men and 1.2% (0.4%–3.1%) in women, without statistical significance; and double positives 0.8% (0.1%–4.2%) in men and 0.9% (0.3–5.1) in women, also without statistical significance. Median cell counts (cells/mL) were: natural killer T cells, 126 (27–580) in men and 105 (20–279) in women ( $p = 0.023$ ); activated T cells: 20 (4–46) in men and 25 (7–75) in women, ( $p = 0.013$ ) and double-positive T cells: 17 (2–85) in men and 21 (7–154) in women, without statistical significance. Sex influenced natural killer T cells, but age did not.

**CONCLUSIONS** Age does not affect counts and percentages of rare T lymphocyte subpopulations in the blood of healthy Cuban adults. Sex differences found for some phenotypes suggest the need for different reference values for women and men.

**KEYWORDS** Normal values, T-lymphocyte subpopulations, flow cytometry, Cuba

## INTRODUCTION

Flow cytometry (FC) is a useful technology for analytical and quantitative characterization of cells, because it offers rapid, simultaneous and complete information on distinctive cell aspects. Thus, it has become an essential tool in diagnosis, monitoring and management of numerous diseases, such as immunodeficiencies, infections, autoimmune diseases, cancer and others with immunopathogenic components. Because of its high sensitivity, it is the gold standard for assessing leukemias and natural killer (NK)/T-cell lymphoma, particularly in special biological samples such as cerebrospinal fluid. Its use has extended to other areas of medicine such as post-transplant care.[1–8]

FC has enabled discovery of different lymphocyte populations and shown their great heterogeneity. Previously, the classic lymphocyte populations reported were NK B and T lymphocytes. The latter have two types, TCD4+ and TCD4+8, which once were considered unique and mutually exclusive because of limitations in early cytometry technology that did not allow more than two or three markers on the same cell.[2] We now know that several cell

subpopulations form each of these markers (CD4 and CD8). This shows the diversity of lymphocyte subpopulations, each with its own role in immunity; even if rare, they have important functions, so it is essential to consider them for correctly phenotyping peripheral lymphocytes.[6–15]

The lymphoid cells of innate immunity are an example of recently discovered lymphocyte diversity.[9,10,16–24] These cells are rare in the blood (together <10% of circulating T lymphocytes) but are not necessarily in the minority in other compartments, in lymphoid tissues or elsewhere. They have important functions in maintenance of homeostasis and in immunopathogenesis of some diseases. They are quantified in counts and percentages of total lymphocytes by FC.

T lymphocytes are more heterogeneous than B lymphocytes and some subpopulations can be characterized with the same monoclonal antibodies used for conventional T, B and NK cell protocols, using a different reading window, which implies changing the analytic strategy and associating the variables of interest in a single graphic in the flow cytometer.

Thus far, normal values of these lymphocyte populations in healthy Cuban adults have not been established. Immunophenotyping of rare lymphocytes in peripheral blood extends analysis to more circulating variants, avoiding underreporting of cells with demonstrable clinical importance and not requiring other technologies or additional reagents for their characterization.

**IMPORTANCE** This study is an initial step towards the use of flow cytometry and conventional reagents to quantify rare but important lymphocytes in diagnosis and prognosis of adult Cuban patients with cancer, autoimmune diseases, HIV/AIDS and chronic infections.

The study objective was to characterize the distribution of rare T-lymphocyte populations in peripheral blood and their relation to sex and age in healthy Cuban adults, with the purpose of eventually using them as reference values in diagnosis and prognosis of multiple diseases.

## METHODS

**Design, subjects and sample collection** A cross-sectional study was carried out from January through April 2017 at the Hermanos Ameijeiras Clinical–Surgical Teaching Hospital. We included 129 apparently healthy adults, companions of patients who were seen at the immunology service of the Hermanos Ameijeiras Clinical–Surgical Teaching Hospital, and who provided written informed consent to participate. The sample consisted of 68 women and 61 men, aged 18–80 years (average 40 years). Exclusion criteria were habits, diseases and treatments that could structurally and functionally modify the immune system: toxic habits (smoking, alcohol consumption >40 g daily or its equivalent per week,[25] >4 cups of coffee a day); history of infections; use of antibiotics, immunosuppressants, immunostimulants, anti-inflammatories or anticoagulants in the previous 6 months; diabetes, immunodeficiencies, autoimmune or neoplastic diseases; (for women) pregnancy.

Blood was obtained by peripheral venipuncture; 4 mL were deposited in Vacutainer tubes using ethylene-deamine tetraacetic acid as anticoagulant. Samples were processed within six hours after extraction and the process followed good laboratory practice standards.

**Flow cytometry** A Beckman Coulter Gallios 8-color cytometer (Beckman Coulter, France) was used; 100  $\mu$ L of blood were dispensed for staining with fluorochrome-conjugated monoclonal antibodies from the same manufacturer (Beckman Coulter, France): anti-CD45 AA750 (Clone J33), anti-CD19 (Clone J3-119), anti-CD3 FITC (Clone UCYT1), anti-CD4 PC5.5 (Clone 13B8.2), anti-CD8 AA700 (Clone B9.11), anti-CD56 PE (Clone N901) (NKH-1), anti-HLA-DR PE (Clone Immu-357), anti-CD25 PC5 (Clone B1.49.9). Two panels were designed:

1. phenotype of double-positive mature subpopulations and NKT, and
2. phenotype of activated T cells.

To characterize the phenotype of T lymphocyte populations, immunophenotyping was done as follows:

1. mature double-positive alpha beta T cells (CD45<sup>+</sup>, CD3<sup>+</sup>, CD4<sup>+</sup>CD8<sup>+</sup>) and their subtypes—
  - a. CD45<sup>+</sup>CD3<sup>+</sup>CD4<sup>high</sup>CD8<sup>low</sup>
  - b. CD45<sup>+</sup>CD3<sup>+</sup>CD4<sup>low</sup>CD8<sup>high</sup>
2. NKT cells (CD45<sup>+</sup>CD3<sup>+</sup>CD56<sup>+</sup>) and subtypes—
  - a. CD45<sup>+</sup>CD3<sup>+</sup>CD56<sup>+</sup>CD4<sup>+</sup>CD8<sup>-</sup>
  - b. CD45<sup>+</sup>CD3<sup>+</sup>CD56<sup>+</sup>CD4<sup>-</sup>CD8<sup>+</sup>
  - c. CD45<sup>+</sup>CD3<sup>+</sup>CD56<sup>+</sup>CD4<sup>-</sup>CD8<sup>-</sup>
3. activated T cells—
  - a. CD45<sup>+</sup>CD3<sup>+</sup>HLA-DR<sup>+</sup>
  - b. CD45<sup>+</sup>CD3<sup>+</sup>CD25<sup>+</sup>
  - c. CD45<sup>+</sup>CD3<sup>+</sup>HLA-DR<sup>+</sup>CD25<sup>+</sup>

Sample preparation was carried out according to manufacturer's specifications for cell surface immunophenotyping, using a protocol of unwashed red blood cells with Versalyse buffer (Beckman

Coulter, France). Cytometer quality control was performed daily with Flow-Check fluorospheres (Beckman Coulter, France) to align lasers and check the water system. Fluorescence intensity was controlled with Flow-Set fluorospheres from the same company.

Acquisition data were processed using Kaluza Analysis Software V1.5A, with a minimum of 50,000 acquired events, which refers to the number of formed elements contained in a blood sample that pass through the cytometer's laser light beam and are counted and analyzed. In each case >4000 events were obtained in the lymphocyte region characterized by high expression of CD45 and low complexity as indicated by side scatter. Rare subpopulations were identified with modification of the analysis strategy to associate the variables of interest in a single graphic, permitting analysis of six monoclonal antibodies specific to six cell surface antigens at once, using a single 50- $\mu$ L blood sample. This is illustrated in Figure 1, where each dot plot represents two parameters analyzed by the cytometer.

Bidimensional analyses were concatenated hierarchically to enable multiparametric analysis and characterization of several cell population phenotypes in a single tube. Thus, the first dot plot in Figure 1a reflects a window through which single cells passed through the laser beam for cell-by-cell analysis. Next a second dot plot was generated to relate cell size to complexity, distinguishing cells from artifacts or detritus. Once cell events were selected, a third dot plot was created that identified groups of leukocytes based on expression of a pan-leukocyte antigen, CD45. The lymphocyte population for analysis is circled at the bottom of the third dot plot. Then antigens representative of each strain were combined successively, from general to specific, each population displayed in a separate quadrant (Figure 1).

Absolute counts were obtained by double platform; results obtained were combined in an automatic hematological counter and by cytometry. The following formula was applied:

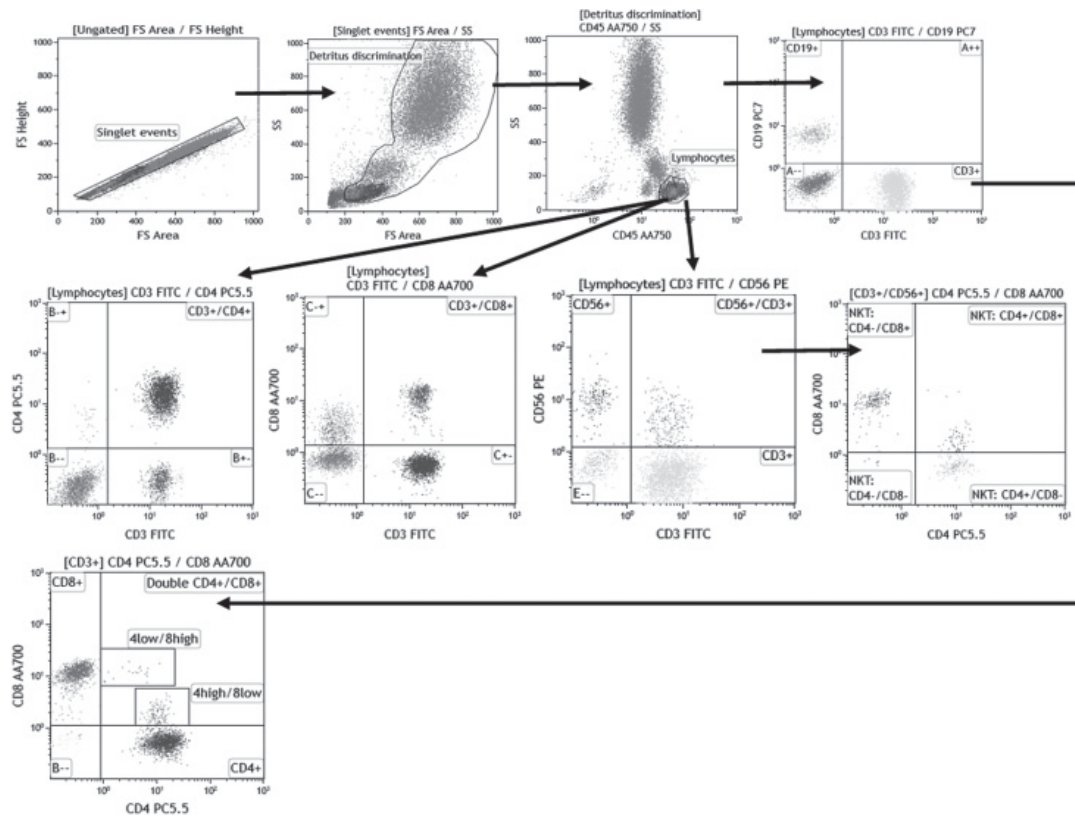
$$\text{Absolute count (cells}/\mu\text{L}) = \text{lymphocyte count (number of cells}/\mu\text{L in blood count}) \times \% \text{ of the cellular subpopulation of interest} \div 100.$$

**Analysis** Descriptive statistics were calculated: absolute and relative frequencies, means, medians and SD. The Kolmogorov–Smirnov test was used to assess normality of distribution of variable values. Because of variable asymmetry, percentiles 2.5 and 97.5 were specified as lower and upper limits, respectively, for reference intervals. A linear regression model was used to assess the effect of age and the two-tailed Mann–Whitney *U* test for independent samples to measure the effect of sex. The significance threshold was set at *p* <0.05.

**Ethics** Study participants provided written informed consent per the Helsinki Declaration.[26] The consent document described the importance of participation and explained the study's characteristics and possible risks and benefits. All data were kept confidential and participant identity was delinked. The selection of diagnostic tools followed the ethical principles of maximum benefit and nonmaleficence.

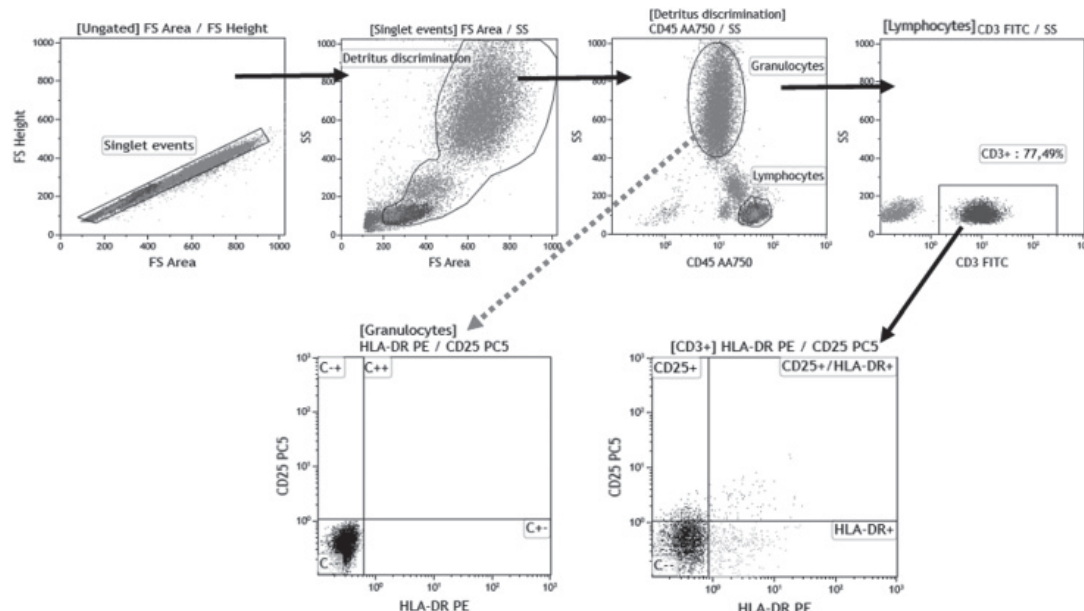
**Figure 1: Reading window strategies**

**a. Design for panel 1 of T and NKT cell phenotypes<sup>a</sup>**



<sup>a</sup>To identify double-positive populations, a CD4 vs. CD8 graphic from the CD3<sup>+</sup> region was added.

**b. Design for activated T lymphocyte panel<sup>b</sup>**



<sup>b</sup>The scatter plot marked with a dotted arrow (which comes from the granulocyte region) was used as a control to establish cutoff points for HLA-DR and CD25 tertiary antigens, since these cells do not express these antigens, which were cloned to obtain values for activated subpopulations in the CD3<sup>+</sup> region.

NKT: natural killer T

(available in color online at [www.mediccreview.org/estimating-normal-values-of-rare-t-lymphocyte-populations-in-peripheral-blood-of-healthy-cuban-adults](http://www.mediccreview.org/estimating-normal-values-of-rare-t-lymphocyte-populations-in-peripheral-blood-of-healthy-cuban-adults))

## RESULTS

Age did not affect counts and percentages of the cell subpopulations studied;  $r^2$  values were distant from unity and differences by age did not meet the specified significance level of  $p < 0.05$ . For some subpopulations, significant differences were found between men and women.

No significant sex differences were found in counts and percentages (of total lymphocytes) of double-positive alpha beta T lymphocytes subpopulations, except for the  $CD3^+CD4^lowCD8^high$  phenotype. Median percentages were less than one percent, but the median was one percentage point higher for women than for men. The upper range limit did not exceed 5.1% (Table 1).

Counts and percentages of NKT cells and their subtypes are summarized in Table 2. The  $CD3^+CD56^+$  phenotype was influenced by sex, with both counts and percentages significantly higher in men ( $p = 0.003$  and  $p = 0.023$ , respectively). Percentages of  $CD3^+CD56^+CD8^+$  were also significantly higher in men. For all other subtypes, values were higher in men, but without statistical significance. Wider ranges in counts and percentages were found in this lymphocyte subpopulation than in other subpopulations studied, especially among men.

Counts and percentages of activated lymphocyte subpopulation phenotypes are summarized in Table 3. Modest sex differences were found in subtype medians, with statistical significance for  $CD3^+HLA-DR^+CD25^+$  ( $p = 0.013$ ) counts, but not percentage of total lymphocytes (Table 3).

## DISCUSSION

Normal blood percentages of rare lymphocytes are influenced by many factors, such as age, sex, viral infections, stress, medications, chronic diseases, lifestyles and even study methodology. Therefore, it is recommended that reference values be established through regional studies and even by countries and that they be updated periodically.[1,2,4,6] To obtain these values and their ranges, immunophenotyping of these subpopulations and their subtypes should be performed using flow cytometry in persons with immune systems unaffected by diseases, treatments or lifestyles. Once determined, they can be used in clinical practice to diagnose and predict the clinical course of immune system diseases and other diseases involving the immune system, such as cancer and infections.[2,4,6]

Values found for the double-positive (DP) alpha beta T population were similar to those published by authors in other countries and showed no differences between men and women.[27–32] However, some studies report an increase in persons aged >60

**Table 1: Reference ranges for peripheral double-positive T lymphocytes in healthy Cuban adults (n = 129)**

Phenotype	Men		Women		p Value <sup>a</sup>
	Median	Percentiles (2.5–97.5)	Median	Percentiles (2.5–97.5)	
$CD3^+CD4^+CD8^+$ (%)	0.8	0.1–4.2	0.9	0.3–5.1	0.173
$CD3^+CD4^+CD8^+$ (cells per $\mu$ L)	17	2–85	21	7–154	0.080
$CD3^+CD4^{high}CD8^{low}$ (%)	0.3	0–3.8	0.4	0–4.7	0.312
$CD3^+CD4^{high}CD8^{low}$ (cells per $\mu$ L)	7	0–77	10	0–142	0.218
$CD3^+CD4^{low}CD8^{high}$ (%)	0.2	0–1.9	0.3	0–2.5	0.044 <sup>b</sup>
$CD3^+CD4^{low}CD8^{high}$ (cells per $\mu$ L)	5	0–33	8	0–57	0.028 <sup>b</sup>

<sup>a</sup>Mann–Whitney *U* test   <sup>b</sup>significant

**Table 2: Reference ranges for NKT lymphocytes and subtypes in healthy Cuban adults (n = 129)**

Phenotype	Men		Women		p Value <sup>a</sup>
	Median	Percentiles (2.5–97.5)	Median	Percentiles (2.5–97.5)	
$CD3^+CD56^+$ (%)	6.3	1.4–23	4.7	0.8–11.3	0.003 <sup>b</sup>
$CD3^+CD56^+$ (cells per $\mu$ L)	126	27–580	105	20–279	0.023 <sup>b</sup>
$CD3^+CD56^+CD4^+$ (%)	0.2	0–3.4	0.2	0–4	0.969
$CD3^+CD56^+CD4^+$ (cells per $\mu$ L)	5	0–86	7	1–100	0.436
$CD3^+CD56^+CD8^+$ (%)	3.7	0.6–18	3.4	0.5–8.5	0.037 <sup>b</sup>
$CD3^+CD56^+CD8^+$ (cells per $\mu$ L)	81	9–395	66	11–212	0.082
$CD3^+CD56^+CD4CD8$ (%)	1.1	0.2–6.1	0.9	0.04–5.6	0.383
$CD3^+CD56^+CD4CD8$ (cells per $\mu$ L)	22	3–125	20	1–141	0.579

<sup>a</sup>Mann–Whitney *U* test   <sup>b</sup>significant   NKT: natural killer T

**Table 3: Reference ranges for activated lymphocytes and subtypes in healthy Cuban adults (n = 95)**

Phenotype	Men		Women		p Value <sup>a</sup>
	Median	Percentiles (2.5–97.5)	Median	Percentiles (2.5–97.5)	
$CD3^+HLA-DR^+$ (%)	12.2	2.7–26.4	10.4	3.1–28.2	0.717
$CD3^+HLA-DR^+$ (cells per $\mu$ L)	219	54–588	253	59–695	0.243
$CD3^+CD25^+$ (%)	3.7	1.4–12.0	3.6	1.0–10.3	0.612
$CD3^+CD25^+$ (cells per $\mu$ L)	82	19–208	80	17–235	0.970
$CD3^+HLA-DR^+CD25^+$ (%)	1.0	0.2–2.2	1.2	0.4–3.1	0.550
$CD3^+HLA-DR^+CD25^+$ (cells per $\mu$ L)	20	4–46	25	7–75	0.013 <sup>b</sup>

<sup>a</sup>Mann–Whitney *U* test   <sup>b</sup>significant

years.[32] Differences from similar studies are explained by the variety of factors influencing the composition of these populations, which cause great variability in the ranges, underscoring the need to establish country-specific reference values.[28,32]

The population of DP T lymphocytes in the peripheral blood of healthy individuals was described by Nascimbeni in 2004.[27] DP lymphocytes have been associated with antitetanus vaccination, good response to influenza vaccination in older adults, and lymphoproliferation (especially when the monoclonal phenotype is expanded after viral infections such as influenza A or *Vaccinia*).[28,32] Absence of lymphocytosis is more common in persistent viral infections, either latent or chronic, caused by cytomegalovirus, Epstein–Barr, herpes simplex, varicella zoster, HIV, or hepatitis B or C.[27,28] An increase in DP lymphocytes has been linked to autoimmune diseases (lupus, multiple sclerosis and rheumatoid

arthritis), cancer (melanoma and Hodgkin lymphoma) and beta thalassemia (especially after splenectomy). Thus quantification of DP cells is important for prognostic purposes. However, a decrease in or absence of these cells has not been associated with any disease and is considered normal.[6,11,29,30]

DP cell functions are varied due to their double phenotype, with production of cytokine patterns whose balance is related to the most strongly expressed coreceptor, as well as cooperative or cytotoxic functions. In addition, they maintain the ability to recognize antigens presented by both class I and II antigen-presenting molecules.[6,11,28–31] This functional capacity, which demonstrates the lymphoid cell plasticity, allows the immune system to adapt to antigenic challenges, especially long-lasting ones, as in chronic infections and cancer. In the case of HIV infection, their increase has been related to good viral control, even in the acute phase.[32]

DP cells have shown a mainly effector memory phenotype combined with markers of replicative senescence, suggesting that they have undergone continued stimulation over time.[28] This may explain the great variability observed, since they will be more or less abundant depending on the number of exposures to different antigens.

If this DP subpopulation in blood is not considered when performing cell counts, there will be incorrect duplication of their markers, biasing calculation of the CD4/CD8 index. This is a serious problem in followup of HIV patients, since this index requires precision to be useful for classifying patients for antiretroviral therapy, assessing antiviral resistance, monitoring treatment adherence and detecting possible evolution towards AIDS.

NKT populations are characterized mainly by a CD3<sup>+</sup>, CD56<sup>+</sup> phenotype and in humans by absence or differential expression of CD4 or CD8. In 1995, the first articles were published about lymphocyte populations that not only had T cell receptor and NK markers, but also showed a unique type of receptor chain rearrangement and a frequency well above the expected for a specific rearrangement. These cells are called invariant NKT cells.[18–19]

The NKT ranges we found are similar to those reported by Rojas-Pandales in Colombia, who did not observe sex or age effects.[33] The great interindividual variability and wide ranges can be related to the biological characteristics of these lymphocytes, which accumulate in blood as they encounter their cognate antigens. Each person's NKT levels depend on the number of previous exposures to different agents capable of stimulating this cell population. However, the range for the normal (healthy) population in a given country has lower and upper limits, which allows detection of abnormal findings. [18,23,34,35]

Increased NKTs have been observed in chronic infections, allergies, cancer and autoimmune diseases. However, values can also be normal or decreased in allergies and cancer. Decreased values indicate poor cancer prognosis. NKTs comprise different subpopulations with functional diversity. NKT CD8<sup>+</sup> and NKT CD4<sup>-</sup>CD8<sup>-</sup> phenotypes are a source of interferon alpha and tumor necrosis factor gamma, which

possess obvious antitumor activity and are cytotoxic due to perforin secretion and binding of Fas/Fas-L (molecules that mediate apoptosis). NKT cells recognize nonprotein antigens coupled to CD1 molecules on cell surfaces, detecting malignant cells by recognizing antigens missed by their conventional T cell counterparts. This shows that NKTs complement cancer defense and immunosurveillance differently than classical T and NK cells.[24,36–39]

Among lymphocyte activation markers, HLA-DR and CD25 showed great variability among individuals, with an especially wide range in HLA-DR. This may be related to the presence of effector memory populations of T lymphocytes in blood, which also express these molecules, and to the fact that the size of this subpopulation is determined, among other factors, by the number of previous personal exposures to different immunogens.[40]

These molecules are classified as T lymphocyte late activation antigens and are synthesized after the cellular activation process has begun;[40] in flow cytometry interpretation, they are considered tertiary. Since expression cannot be predicted until cellular activation has occurred, they could influence range variability. Some authors report that age and sex do not greatly influence these expression levels, but do affect the process triggering such activation.[41] Considering that regulatory T cells characteristically express the CD25 marker, measuring lymphocyte activation by other markers, such as HLA-DR is more reliable.[40–43] One trait that can inform flow cytometry analysis is that regulatory T cells express CD25 more intensely, so that it does not behave like a typical tertiary marker.[44]

Expression of markers that reflect cell function are important to assess the activation state of the T cell compartment in patients with autoimmune diseases, because such markers discriminate between the active and compensated states of the disease.[45] There is evidence that the greater the number of activated T cells, the greater the severity of decompensation, through autoimmune pathogenesis. This has been observed in diseases such as multiple sclerosis, Wegener granulomatosis, Kawasaki disease, systemic lupus erythematosus and autoimmune aplastic anemias.[46,47] HLA-DR is used as a marker to predict therapeutic response in autoimmune diseases such as Kawasaki disease, in which patients with a higher percentage of HLA-DR<sup>+</sup> T cells fail to respond to therapy with intravenous immunoglobulins.[45–47]

In cancer, activation is analyzed by cell population. Activation of suppressor T cells indicates worsened clinical evolution and poorer prognosis. Evidence indicates correspondence between blood and tumor microenvironments, so that, if activated suppressor cells in the blood increase, so do tumor-infiltrating lymphocytes. But if the activated cells are cytotoxic, this increase can be beneficial, because they are known to help eliminate tumor cells.[43,44]

In the course of infections such as HIV, chronic immune activation is part of immunopathogenesis and is one of the most damaging phenomena in disease progression, since it increases viral load and decreases CD4<sup>+</sup> lymphocytes. Monitoring expression of CD25 and HLA-DR, as markers of the

degree of T cell activation in these patients, has been found to be very useful.[47]

It is important to consider activation markers to assess immune system function in many diseases, in some cases as an immunopathogenic factor and in others as pathological or physiological consequence of a previous illness or condition. For example, increased HLA-DR expression has even appeared in patients with chronic spinal cord injuries, possibly contributing to persistence of chronic inflammation or decreased resistance to infection.[44]

The levels of T-lymphocyte populations we observed are similar to those found in healthy controls in published research. [43-47]

## CONCLUSIONS

Age does not affect counts and percentages of rare T lymphocyte subpopulations in the blood of healthy Cuban adults. Sex differences found for some phenotypes suggest the need for different reference values for women and men.

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