

THE STATE OF HEMATOLOGICAL PARAMETERS IN APLASTIC ANEMIA

Akhmedova Zukhra Bakhtiyorovna,

Matkarimova Dilfuza Saburovna,

Boboev Kodirzhon Tukhtaboevich

*Republican specialized scientific - practical medical center of
Hematology Tashkent medical academy*

Summary

Purpose of the study. To analyze laboratory changes in peripheral blood and myelogram in adult patients with aplastic anemia.

Methods. The material for clinical and laboratory studies in the work were patients with AA (n=86) who sought diagnostic help and subsequent inpatient examination at the republican specialized Scientific and Practical Medical Center of Hematology (RSNPMCG, Tashkent) from 2019 to 2023. Patients with AA ranged in age from 18 to 79 years, while the median age was 40.8 ± 1.8 years. The diagnosis was made taking into account clinical and laboratory data.

The research methods included laboratory examination (general blood test (UAC) and myelogram) and statistical methods of processing the results using the PC application package "OpenEpi 2009, Version 2.3".

Conclusions. Despite the severity of form A, UAC is characterized by a decrease in hemoglobin concentration, the number of erythrocytes, platelets, and leukocytes due to neutropenia, lymphocytosis, and acceleration of ESR. The myelogram shows a decrease in the cellularity of the red bone marrow, three-stage cytopenia and replacement of the red brain with a fatty brain.

Key words: UAC, myelogram, three-stage cytopenia, fatty brain.

Introduction. AA is a rare type of bone marrow insufficiency syndrome (BCM), which is characterized by severe pancytopenia and bone marrow

hypoplasia of varying severity [1,2,4]. Acquired AA, also called idiopathic AA, accounts for the majority (~70%) of all newly diagnosed cases. [5,7,10].

The severity of AA was assessed in accordance with the parameters of the blood test and the results of the bone marrow examination. Severe AA (SAA) was defined as a BM cell count of < 25% or 25-50% with < 30% residual hematopoietic cells and at least two of the following: (I) absolute neutrophil count < $0.5 \times 10^9/l$, (II) platelets < $20 \times 10^9/L$ and (III) the number of reticulocytes < $20 \times 10^9 / l$. Patients with AA who did not meet the SAA criteria were classified as non-severe AA (NSAA). The result of the treatment was based on previous literature [3,6,8,9].

Purpose of the study. Провести анализ лабораторных изменений в периферической крови и миелограмме у взрослых пациентов с апластической анемии.

Purpose of the study. To analyze laboratory changes in peripheral blood and myelogram in adult patients with aplastic anemia.

Methods. The material for clinical and laboratory studies in the work were patients with AA (n=86) who sought diagnostic help and subsequent inpatient examination at the republican specialized Scientific and Practical Medical Center of Hematology (RSNPMCG, Tashkent) from 2019 to 2023. Patients with AA ranged in age from 18 to 79 years, while the median age was 40.8 ± 1.8 years. The diagnosis was made taking into account clinical and laboratory data.

The research methods included laboratory examination (general blood test (UAC) and myelogram) and statistical methods of processing the results using the PC application package "OpenEpi 2009, Version 2.3".

Results. In patients with AA, a decrease in hemoglobin concentration of cytopenia (decrease in the number of erythrocytes, platelets and leukocytes), absolute neutropenia, relative lymphocytosis and acceleration of ESR were found in clinical blood analysis compared with healthy ones.

Analyzing the average values of the indicators of the general blood test in the main group of patients with AA in comparison with the control, the changes

characteristic of the disease manifested by cytopenia were established. In patients with AA, there was a significant decrease in hemoglobin by 2.5 times (52.8 ± 1.7 g/l versus 134.2 ± 2.4 g/l; $P < 0.01$) and erythrocytes by 2.0 times ($1.7 \pm 0.07 \times 10^{12}/l$ versus $3.4 \pm 0.5 \times 10^{12}/l$; $P < 0.05$).

At the same time, the median platelet count was statistically significantly reduced by 21.1 times ($13.6 \pm 3.0 \times 10^{12}/l$ versus $287.2 \pm 2.1 \times 10^{12}/l$; $P < 0.001$), and leukocytes by 6.2 times ($1.2 \pm 0.05 \times 10^{12}/l$ versus $7.4 \pm 1.2 \times 10^{12}/l$; $P < 0.001$).

The above data show some of the main changes in the UAC characteristic of AA. Meanwhile, the key factor in the UAC for confirming the diagnosis of AA was a decrease in the absolute number of neutrophils among patients by 11.2 times compared with the control ($0.423 \pm 0.03 \times 10^{12}/l$ versus $4,743 \pm 0.25 \times 10^{12}/l$; $P < 0.001$).

A 1.7-fold increase in the number of lymphocytic cells was also specific for this pathology ($61.0 \pm 1.1\%$ vs. $35.5 \pm 1.4\%$; $P < 0.05$), as well as a 7.2-fold increase in the level of ESR (52.0 ± 2.1 mm/h vs. 7.2 ± 1.4 mm/h; $P < 0.001$).

Thus, the laboratory signs of AA in the UAC were normocytic anemia, severe cytopenia (erythro-, thrombo- and leukopenia), neutropenia, relative lymphocytosis and accelerated ESR. Moreover, these signs differed in severity depending on the severity of AA, respectively, having deeper disorders in severe and superheavy forms, which were associated with the severity of clinical manifestations in patients with AA.

The results of morphological and quantitative analysis of the bone marrow showed a picture of small cells in the normoblastic type of hematopoiesis with a content of $0.18 \pm 0.009\%$ blast cells in the main AA group, which showed no increase in their content.

Microscopy revealed fatty voids, as well as single areas containing reticular, erythroid and lymphoid cells. The sum of erythroid elements was reduced to $6.4 \pm 0.2\%$ of their minimum normal number by 2.3 times (14.5% ; $P < 0.01$), and from the maximum by 4.1 times (26.5% ; $P < 0.001$). The number of lymphocytes, which amounted to $24.3 \pm 0.4\%$ of the maximum normal number, was increased by

1.8 times (13.7%; $P < 0.05$), and from the minimum allowable by 5.6 (4.3%; $P < 0.001$).

Meanwhile, the content of plasma cells, eosinophils, basophils and monocytes were within the reference norms (see Table 3.13). While the number of neutrophils and megakaryocytes (MCC) significantly decreased below the permissible normal values by 2.0 (26.1-1.0% vs. 52.7%; $P < 0.05$) and 2.5 times (2.0 ± 0.1 against 5; $P < 0.01$), respectively.

Thus, the myelogram picture was characterized by bone marrow hypocellularity, manifested by three-stage cytopenia, narrowing of granulocytic and erythroid sprouts with normoblastic type of hematopoiesis, the presence of voids and the replacement of the red brain with yellow. All these signs are key characteristics of AA.

Conclusion. In the UAC, all degrees of severity of AA are characterized by a decrease in hemoglobin concentration, the number of erythrocytes, platelets, and leukocytes due to neutropenia, lymphocytosis, and acceleration of ESR. The myelogram shows a decrease in the cellularity of the red bone marrow, three-stage cytopenia and replacement of the red brain with a fatty brain.

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Akhmedova Zukhra Bakhtiyarovna, basic doctoral student RSSPMCH, number: +99894 726 87 88.

Matkarimova Dilfuza Saburova, Professor of the Department of Hematology, Transfusiology and Laboratory Science of TMA, number: +99897 412 91 18.

Boboev Kodirjon Tukhtabaevich, Head of the Department of Molecular Medicine and Cellular Technologies of the RSSPMCH, number: +99890 319 39 57.