White Spleen in Hairy Cell Leukemia – Case Report and Review of the Literature

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Abstract: Hairy cell leukemia is a malignant chronic lymphoproliferative disease that covers for about 2% of all leukemia and occurs four times more often in males. It is most often manifested by splenomegaly. Splenomegaly often becomes symptomatic and manifests as pain in the epigastrium and feeling of flatulence and discomfort in the abdomen. Cladribine and Pentostatin are purine analogues and they represent the first line in the treatment of this disease. Splenectomy is applied in patients with massive symptomatic splenomegaly, severe pancytopenia, as well as a temporary measure in pregnant women when symptomaticity is expressed. In our work, we present a patient who experienced a relapse of the disease after two cycles of chemotherapy, as confirmed by biopsies of the bone marrow. From this case can be concluded that even in the era of the use of purine analogues, splenectomy can be of great benefit in the treatment, and determinant of its further outcome for those affected by Hairy cell leukemia.

Keywords: spleen, splenomegaly, splenectomy, hairy cell leukemia

1. INTRODUCTION

The first researches by the haematologist and the first indications regarding Hairy cell leukemia date back to the 1923. Since then, there have been numerous studies related to this disease and described in the literature (1,2). In 1958, Bouroncle and associates described the leukemic reticuloendotheliosis as a separate entity with a typical (Hairy cell) cell appearance (3).

Hairy cell leukemia (HCL) is a malignant chronic lymphoproliferative disease that covers 2% of all leukemia and occurs four times more often in males than in females. The average age of patients diagnosed with this disease is about 55 years (4.5). The disease is manifested by pancytopenia, absence of peripheral lymphadenopathy, enlarged spleen and often absolute monocytopenia. Peripheral blood contains mature B cells with voluminous cytoplasmic projections that resemble hairs. Histologically, HCL cells biopsied from the bone marrow have a cytoplasm that surrounds the nucleus and that way resembles an egg (6).

The most common manifestation of this disease is splenomegaly and occurs in about 96% of patients. Often, splenomegaly becomes symptomatic and manifests as pain in the epigastrium and feeling of flatulence and discomfort in the stomach (7,8).

Until 1984, splenectomy was the method of choice in the treatment of this disease (9). The start of administration of purine analogues in the treatment of HCL was a revolutionary treatment, with a total response of 85% of patients and survival up to 15 years without a progression (10.11).

Cladribine and Pentostatin, 2 analogues of purine, are the first line in the treatment of HCL. They are equally effective in encouraging and maintaining a remission. Thanks to its favorable toxicity profile, most hematologists consider that Cladribin is the first line of chemotherapy. Interferon alfa is an option for treatment of this disease during pregnancy, when the treatment is justified. It can also be used as initial treatment in patients with severe pancytopenia or an active infection, in order to improve blood test results and to allow further therapy of purine analogues (12,13,14).

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due to cell sequestration, as well as a temporary measure in pregnant women when symptomatology is expressed (14).

In our work, we present a patient who experienced a relapse of the disease after two cycles of chemotherapy, as confirmed by biopsies of the bone marrow. Because of massive symptomatic splenomegaly, leukocytopenia and, thrombocytopenia, the consilium, consisted of surgeons and hematologist, decided to perform a splenectomy.

2. CASE REPORT

61 years old patient was admitted to the Clinic for Digestive Surgery - First Surgical Clinic, Clinical Center of Serbia, due to pains in the upper parts of the abdomen, feeling of flatulence, weight and discomfort in the abdomen.

At the admission, we discover that his enlarged spleen was diagnosed 16 years ago, when he was also supposed to have a bone marrow biopsy to which he did not go, and hasn't report to a doctor since then. His first admission to our institution was at the end of 2014 as an emergency case due to abdominal pain and the presence of splenomegaly. After initial diagnostics, he was transferred to the Clinic for Hematology where he was diagnosed with Hairy Cell Leukemia and ordinated with chemotherapy (Cladribine). After 6 months, a control biopsy of the bone marrow showed a significant reduction in bone marrow infiltration from 40% to 10%. In July 2015, he got another cycle of the (Cladribine) chemotherapy. The bone marrow biopsy in January 2016 detected a complete haematological remission of the disease in a bone marrow, but there was still a splenomegaly with expressed symptoms. The value of the platelets at the admission to our Clinic were 38x10^9/L. Computerized tomography (CT) confirmed the previous finding of splenomegaly. The length of the spleen in the cranial - caudal direction (longitudinal diameter) was 29.09 cm (Figure 1), while its transversal diameter was 21.81 cm (Figure 2). Due to the existence of a massive splenomegaly (> 20 cm in the cranial - caudal direction) we decided for the classic splenectomy. After surgery, the weight of the spleen was 1950 gr. The macroscopic preparation of the spleen was largely covered with a light-colored hyaline capsule with a thickness of 5-15 mm (Figure 3).

![Preoperative imaging (CT), Craniocaudal diameter of the spleen](image1)

**Figure1.** Preoperative imaging (CT), Craniocaudal diameter of the spleen

![Preoperative imaging (CT), transversal diameter of the spleen](image2)

**Figure2.** Preoperative imaging (CT), transversal diameter of the spleen
Histopathological examination found that splenic architecture was changed to the account of highly expanded red pulp with very rare white pulp residues, and that corresponds to infiltrative leukemic infiltration. This examination also the presence of Residual splenic disease in hairy cell leukaemia after treatment with chemotherapy (Cladribin). The infiltrates are confined to the red pulp (Figure 4). And Hyaline perisplenitis known as “icing sugar spleen” with capsule thickening ranging 5 to 15mm in this patient. Hairy cell infiltrate and blood lakes are seen in splenic parenchyma (Figure 5).

Additional immunohistochemical examination confirmed strong, diffuse immunoreactivity for CD20 (Figure 6). And Immunoreactivity for DAB 44 in splenic hairy cell infiltrate (Figure 7).
3. DISCUSSION

Introduction of purine analogues in the treatment of HCL significantly contributed to improving the patients’ survival. In some HCL patients, a long perennial remission can be achieved only with the usage of the purine analogues and without additional therapy (10).

Some authors believe that the length of the disease-free interval is related to achieving a complete response to therapy (15,16). Although Hair Cell Leukemia is an incurable disease, with current therapy, patients can achieve permanent remission without further continuous application of therapy, except in cases of symptomatic relapse (17).

Despite the complete therapeutic response after two cycles of chemotherapy, as confirmed by biopsies of the bone marrow, a permanent splenomegaly remained with pronounced symptoms. By agreement with a surgeon and hematologist, the consilium decided to perform splenectomy. Histopathological examination and immunohistochemical verification confirmed the residual HCL in the spleen. It is not clear if splenomegaly would have spontaneously regressed anyway, but after a histopathological examination, it turned out that performing of splenectomy as a strategy proved to be the real therapeutic option.

Patients with complete therapeutic response should have hemoglobin > 110 g/L, no additional transfusion, platelets > 100 x 10^9/L and neutrophils > 1500 / μL (18).

After surgery, and just before hospital release, the patient we operated on had hemoglobin 108 g/L, platelets 318 x 10^9/ L and neutrophils 2100 / μL.

In patients with HCL, the most common cause is infection. These patients have neutrophilia and monocytopenia, so bacterial, viral and opportunistic infections are expected. Purine analogues provide long-term suppression of immune system cells and induce prolonged neutropenia (19).

It is very important to inform and educate patients about infections prevention. The results of the use of specific preventive options has not been confirmed in some controlled studies, but the practice differs from patient to patient, thus the strategies of prevention and treatment vary (18,20).

After splenectomy, antibiotic prophylaxis, as well as postoperative vaccination, were performed in accordance with the current directives and guidelines for the treatment and prevention of postsplenectomy infections.

After surgery, the patient is in remission for 2 years. His period of monitoring and remission continues.

4. CONCLUSION

Although splenectomy does not represent the first measure of treatment of HCL, from this case can be concluded that even in the era of the use of purine analogues, splenectomy can be of great benefit in the treatment and a determinant of it's success. From the literature can be concluded that HCL has many unusual manifestations of the disease and it is necessary to continuously monitor and improve the guidelines and strategies for the treatment, and guidelines should also be of great benefit in improving the treatment and should not be used for refusal of the necessary diagnostic and therapeutic treatment options.

REFERENCES


