**CASE REPORT**

A diagnostically difficult case of chronic myeloid neoplasm with eosinophilia and abnormalities of PDGFRA effectively treated with imatinib in accelerated phase

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**KEY WORDS**
chronic myeloid neoplasm with eosinophilia and abnormalities of PDGFRA, imatinib

**ABSTRACT**

Chronic myeloid neoplasm with eosinophilia and abnormalities of platelet-derived growth factor receptor α (PDGFRA), referred to until 2008 as chronic eosinophilic leukemia, is distinguished from hypereosinophilic syndrome (HES), if accompanied by genetic abnormalities that enable to determine eosinophil clonality. Typically, HES has a benign course and glucocorticosteroids suffice to achieve remission. In chronic myeloid neoplasm with eosinophilia and abnormalities of PDGFRA the FIP1L1-PDGFRα fusion gene can be detected. Its product is a protein showing tyrosine kinase activity leading to malignant proliferation of eosinophil precursors. Differential diagnosis of HES is often difficult because hypereosinophilia may also be reactive and may occur in many nonhematological as well as hematological disorders. Thus, reverse-transcription polymerase chain reaction (RT-PCR) is indicated in all patients with HES in order to detect the FIP1L1-PDGFRα transcript. Traditional treatment of chronic myeloid neoplasm with cytostatic drugs results in a short-term and transient remission or stabilization of the disease.

We present the case of a 52-year-old patient with chronic myeloid neoplasm with eosinophilia and abnormalities of PDGFRA, in whom acceleration occurred after a year of cytostatic therapy with hydroxyurea and was successfully treated with imatinib. It was impossible to unequivocally determine the type of bone marrow disease based on histologic criteria, and a wide spectrum of molecular tests differentiating the type of myeloid proliferation were necessary to establish the diagnosis. RT-PCR did not reveal BCR-ABL or JAK2 V617F mutation. Further molecular testing showed rearrangement involving the FIP1L1 gene, thus enabling implementation of targeted therapy.

**INTRODUCTION**

Chronic myeloid neoplasm with eosinophilia and abnormalities of platelet-derived growth factor receptor α (PDGFRA) had been referred to as chronic eosinophilic leukemia (CEL), until the World Health Organization published a new classification of tumors of hematopoietic and lymphoid tissues in 2008. It occurs very rarely and is distinguished from hypereosinophilic syndrome (HES), if accompanied by genetic disorders that indicate eosinophil clonality. The term “HES” was introduced in 1968 by Hardy and Anderson.¹,² It occurs more often in men than in women (9:1), mainly in subjects aged 20 to 50 years.¹,³

The disease is characterized by deletion on chromosome 4: del(4)(q12;q12), which accounts for the formation of a fusion gene consisting of the PDGFRα and FIP1L1 (FIP1-like-1) genes.¹,⁴ Del(4) is not detectable using conventional cytogenetic techniques. Reverse-transcription polymerase chain reaction (RT-PCR) demonstrated the prevalence of the FIP1L1-PDGFRα fusion gene...
in about 10% to 15% of patients with chronic myeloid neoplasm with eosinophilia and abnormalities of PDGFRA. The protein which is produced as a result of this chromosomal aberration has an increased tyrosine kinase activity, which causes neoplastic transformation of hematopoietic cells. Eosinophil precursors release proinflammatory cytokines, mediators derived from arachidonic acid, and enzymes toxic to surrounding tissues. Eosinophilic infiltration occurs in various organs leading to their damage. This causes a whole spectrum of accompanying symptoms, both general and related to pathologies of the cardiovascular, respiratory, nervous, and gastrointestinal systems and to skin lesions.

Eosinophilia may be reactive and may occur in numerous nonhematological diseases (allergies, parasitic diseases, systemic connective tissue diseases, and others). It may also accompany a number of hematological diseases, including T-cell lymphomas, Hodgkin’s lymphoma, mastocytosis, myelodysplastic syndromes, and other myeloproliferative disorders. For this reason, differential diagnosis may be challenging and painstaking.

**CASE REPORT** In December 2006, a 52-year-old patient was diagnosed with splenomegaly, leukocytosis (27,000/mm³) with an increase in younger forms including promyelocytes and eosinophils (33%), and lactate dehydrogenase (LDH) of 215 U/l. Abdominal ultrasound showed enlarged spleen up to 20 cm, and a 15 × 12 mm lymph node in the hepatic hilus. There were no abnormalities in other organs (chest X-ray was normal, there were no skin lesions). Histologic examination of a bone marrow biopsy showed features of a bone marrow neoplasm, suggesting either myelodysplastic syndrome or myeloproliferative disease, with emphasis on chronic myelomonocytic leukemia or an atypical chronic myeloid leukemia (CML), but with an increased percentage of eosinophilic granulocyte forms (there was an increased percentage of eosinophils, but neutrophils were predominant) and mast cells (figure). It was impossible to unequivocally diagnose the type of disease based on histologic criteria, and it was necessary to perform a wide spectrum of molecular tests. Cytogenetic analysis of the bone marrow by G-banding method revealed a normal karyotype. RT-PCR did not detect BCR-ABL p210 t(9;22)(q34;q11). The examination of fine-needle biopsy of the spleen yielded inconclusive results. Granulocyte alkaline phosphatase activity was 6 points.

Given an ambiguous clinical presentation, an unclassified myeloproliferative syndrome was diagnosed in April 2007. Cytoreductive therapy was initiated (1 g hydroxyurea [HU]) due to the size of the spleen and hyperleukocytosis with younger stages including myeloblasts (4%). Complementary genetic tests for the JAK2 V617F mutation and FIP1L1 gene rearrangement were performed. Cytogenetics was repeated.

Further molecular testing using RT-PCR enabled to identify the FIP1L1-PDGFRα fusion gene associated with del(4)(q12;q12). JAK2 V617F mutation was not detected. Cytogenetic analysis of peripheral blood using the G-banding method revealed 46XY, del(7)(q22) karyotype. Repeat histologic evaluation of the bone marrow and analysis of genetic tests resulted in the diagnosis of chronic myeloid neoplasm with eosinophilia and abnormalities of PDGFRA in March 2008. It is important to note that this diagnosis was based, above all, on molecular analysis, which revealed the presence of aberration that was typical and highly characteristic of CEL, with coexisting peripheral hyperleukocytosis (it rose to 54,000/μl) and hypereosinophilia (it increased from 33% to 41%). The histologic examination of the bone marrow (despite an increase in the percentage of eosinophilic forms) did not provide conclusive evidence to establish the diagnosis. Detection of FIP1L1-PDGFRα rearrangement results in effective treatment with tyrosine kinase inhibitors (imatinib, dasatinib, nilotinib) because the product of this gene is 100 times more sensitive to imatinib than BCR-ABL.

In April 2008, despite HU therapy, the patient’s state deteriorated. The following symptoms were observed: weight loss, body temperature 38°C, fatigue, dyspnea at rest, dry cough, and peripheral edema. Laboratory tests revealed increased leukocytosis to 168,000/mm³, hemoglobin 9.4 g/dl, a decrease in platelet count to 61,000/mm³, an increase in the percentage of peripheral blastic cells to 8%, and a decrease in the percentage of eosinophils to 4%. Cytological assessment of the marrow showed signs of disease acceleration, abnormal plasma coagulation tests (prothrombin index – 72%, international normalized ratio – 1.31, activated partial thromboplastin time – 36.6 seconds, fibrinogen concentration – 4.63 g/l), hypalbuminemia, an elevation of LDH to 482 U/l. Radiogram of the chest demonstrated a small amount of fluid in the pleural cavities, as well as
as hepatosplenomegaly. During escalation of cytoreductive therapy (3 g HU), further progression of the disease was noted, with deterioration of the patient’s condition, and a rise in leukocytosis to 255,000/mm³. 6-merkaptopurine was administered at a dose of 100 mg/day.

We decided that the patient should receive 100 mg of imatinib daily. This led to improvement of his state, resolution of all symptoms, a decrease in leukocytosis to 1500/mm³, and reduced hepatosplenomegaly. Subsequently, the dose of imatinib was reduced to 100 mg twice a week, which improved the patient’s general condition, normalized blood cell counts, and reduced spleen size (to 136 mm).

**DISCUSSION**

Imatinib, due to inhibition of tyrosine kinase, proved useful not only in the treatment of chronic myeloid leukemia, but also in the treatment of chronic myeloid neoplasm associated with eosinophilia and abnormalities of PDGFR, chronic myeloproliferative syndromes with ETV6-PDGFRA gene rearrangement, Pb+ acute lymphoblastic leukemia, and gastrointestinal stromal tumors. However, in some patients with HES, FIP1L1-PDGFRA and ETV6-PDGFRA gene rearrangement was not detected. Despite this, imatinib therapy was successful, which might indicate the presence of undiagnosed chromosomal aberrations that increase tyrosine kinase activity. Detection of FIP1L1-PDGFRA mutation allows to identify a group of patients with a good response to treatment with low doses of imatinib.

The accessibility of modern therapy poses a greater diagnostic challenge – it is necessary to precisely determine the type of neoplastic disorders, which sometimes requires molecular diagnostic testing, as was the case in our patient. It should be emphasized that these tests are particularly important and largely conclusive in the diagnosis of eosinophil clonality, because these disorders quite often do not display any cytological or histological bone marrow abnormalities that are characteristic enough to distinguish them from reactive eosinophilia. They may also be accompanied by abnormalities that suggest a quite different disease, e.g., an increase in mast cells, including those showing abnormal fusiform morphologic features (which in our patient was detectable in histopathologic examination of the bone marrow). This pathology may be associated with increased serum mast cell tryptase levels, and it might be entirely reactive by nature in chronic myeloid neoplasm with eosinophilia and abnormalities of PDGFR.

Progress in genetic testing, increased accessibility of genetic tests, and the introduction of new molecularly targeted agents allow us to prolong the lives of patients while ensuring a significantly lower risk of therapeutic failure and less side effects.

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Trudny diagnostycznie przypadek chorego na przewlekły nowotwór mieloproliferacyjny z eozynofilią i rearanżacją PDGFRA, skutecznie leczonego imatynibem w fazie akceleracji

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SŁOWA KLUCZOWE
imatynib, przewlekły nowotwór mieloproliferacyjny z eozynofilią i rearanżacją PDGFRA

STRESZCZENIE
Przewlekły nowotwór mieloproliferacyjny z eozynofilią i rearanżacją PDGFRA (platelet-derived growth factor receptor α), określany do 2008 roku jako przewlekła białaczka eozynofilowa (chronic eosinophilic leukemia – CEL), jest wyodrębniany z zespołu hipereozynofilowego (hypereosinophilic syndrome – HES), jeśli towarzyszą mu zaburzenia genetyczne pozwalające stwierdzić klonalność rozrostu. HES ma zwykle łagodny przebieg, a glikokortykoidy wystarczają do uzyskania poprawy.

W przewlekłym nowotworze mieloproliferacyjnym z eozynofilią i rearanżacją PDGFRA wykrywa się gen fuzyjny FIP1L1-PDGFRA, którego produktem jest białko o aktywności kinazy tyrozynowej powodujące nowotworową proliferację prekursorów eozynofilii. Diagnostyka różnicowa w HES jest często bardzo trudna, ponieważ hipereozynofilia może mieć także charakter odczynowy i występować w wielu niehematologicznych i hematologicznych jednostkach chorobowych. Wskazane jest więc badanie RT-PCR (reverse-transcription polymerase chain reaction) u wszystkich chorych z HES w celu wykrycia transkryptu FIP1L1-PDGFRA. Tradycyjne leczenie choroby lekami cytostatycznymi przynosi krótkotrwałą i przejściową poprawę lub stabilizację choroby.

Poniżej opisujemy przypadek 52-letniego chorego z przewlekłym nowotworem mieloproliferacyjnym z eozynofilią i rearanżacją PDGFRA, u którego po roku leczenia cytostatycznym hydroksymocznikiem wystąpiła akceleracja rozrostu skutecznie leczona imatynibem. Ponieważ w oparciu o kryteria histologiczne niemożliwe było jednoznaczne określenie typu choroby szpiku kostnego, do postawienia diagnozy konieczna okazała się szeroka gama badań molekularnych różnicujących charakter proliferacji szpikowej. W badaniu metodą RT-PCR nie wykryto mutacji BCR-ABL, nie wykryto też mutacji V617F genu JAK2. Dalsza diagnostyka zmian molekularnych pozwoliła wykryć rearanżację z udziałem genu FIP1L1, co pozwoliło wdrożyć leczenie celowane.
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