Case Series- Heterogeneity of Primary Myelofibrosis- A Challenge to the Clinician

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Introduction

Primary myelofibrosis (PMF) is a myeloproliferative neoplasm characterized by stem cell derived clonal myeloproliferation and extramedullary haematopoiesis (EMH). Myelofibrosis can present as a de novo disorder or evolve secondary to previous polycythemia vera or essential thrombocythemia (post PVMF or post ETMF) [1]. Primary myelofibrosis (PMF) is the least frequent among the chronic myeloproliferative diseases. One study reported an estimated incidence of 1.5 per 100,000 per year 2 PMF occurs mainly in the middle aged and elderly population. The median age at presentation is 67 years [2]. The primary pathogenic mechanism is a clonal stem-cell disorder that leads to ineffective erythropoiesis, dysplastic-megakaryocytic hyperplasia, and an increase in neutrophils with predominance in immature forms [3,4].

The disease is clinically characterized by progressive anemia, constitutional symptoms and splenomegaly. The laboratory findings show a leuco-erythroblastic blood film, tear drop poikilocytosis, reticulin fibrosis in bone marrow and elevated levels of various inflammatory and pro-angiogenic cytokines [1].

The majority (approximately 45-68%) of the patients harbor JAK2V617F mutation [5]. MPLW515L mutations were described in JAK2V617F mutation-negative PMF. Recent discovery of the CALR reticulin mutation has improved the genetic detection of MF to about 80% [6]. PMF has poor prognosis [7-11]. Causes of death are infections, hemorrhage, thrombosis or progression to leukaemia.

Herein we present the clinico-hematological profile of three cases fitting the diagnosis of PMF and one case with post PVMF. These four cases show the diversity of the disease with regard to laboratory, clinical and prognostic features.

Case 1

This 58 year old male patient presented with fever, loss of appetite and weight loss of two weeks duration. On examination, he had mild pallor and moderate splenomegaly. There was no lymphadenopathy, hepatomegaly or bleeding manifestations. Full blood count revealed haemoglobin 10.2 g/dl, total leucocyte count 29.7 × 10³/µl and platelet count 229 × 10³/µl. Peripheral blood smear showed normochromic normocytic red cells with no tear drop cells or nucleated red cells. White cells showed leukocytosis with complete spectrum of neutrophil elements with a streaming pattern in the intertrabecular area. Reticulin was grade 4. BCR - ABL was negative. The patient was treated with hydroxyurea/hydroxyurea 500 mg daily. At two months of follow up, blood counts were normal, but her spleen was palpable at 18 cm below the costal margin.

Case 2

This 67 year old female presented with loss of appetite and weight loss of six months duration. At presentation she was mildly pallid and had hepatosplenomegaly, no lymphadenopathy or bleeding manifestations. Full blood count showed haemoglobin 10.3 g/dl, total leucocyte count 27.6 × 10³/µl and platelet count 286,000. Peripheral blood smear revealed normochromic normocytic red cells, tear drop cells and nucleated red cells. White cells showed leukocytosis with complete spectrum of granulocytic lineage including myeloblasts (1%). Platelets were normal. Bone marrow trephine biopsy showed marked fibroblastic proliferation (Figure 2), lack of other hematopoietic elements with a streaming pattern in the intertrabecular area. Reticulin was grade 4. BCR - ABL was negative. This patient was treated with hydroxyurea/hydroxyurea 500 mg daily. At two months of follow up, blood counts were normal, but her spleen was palpable at 18 cm below the costal margin.

Case 3

This 69 year old male presented with dyspnea and loss of appetite of 2/52 duration. On examination he had pallor but no organomegaly or lymphadenopathy or bleeding manifestations. Full blood count revealed hemoglobin, 4.9 g/dl, total leucocyte count 4.64 × 10³/µl and a platelet count 58,000. Peripheral blood film revealed normochromic, normocytic red cells, macrocytes, no tear drop cells or nucleated RBCs. There were no abnormal white blood cells. Platelets were low. The
bone marrow trephine biopsy showed dense fibroblastic proliferation (Figure 3) with megakaryocytic clustering and dysplasia resembling idiopathic myelofibrosis. Reticulin was grade 3. This patient was treated with Erythropoietin 10,000 IU weekly for 4 months with no significant response. He was then treated with thalidomide and oral prednisolone, for which he developed disturbing side effects of severe constipation and peripheral neuropathy with no improvement in the blood counts, which warranted us to stop the drug. Finally he was started on Danazol 300 mg daily which we increased to 500 mg later. At 3 months of follow up his hemoglobin was 6.0 g/dl. Results are yet to be shown and still he is on regular blood transfusions once in three weeks.

Case 4

This 55 year old diagnosed patient with JAK 2 positive primary proliferative polycythemia was well controlled on hydroxyurea 500mg daily and aspirin for 6 years. She eventually showed a trilineage increment of blood counts which needed to be controlled with a higher dose of hydroxycarbamide/hydroxyurea. Her blood counts were well controlled with dose adjustments with hydroxyurea for the following 6 years and then, she developed a leukoerythroblastic blood picture with tear drop poikilocytosis. Her counts were WBC -64,300/μL, Hb- 11.0 g/dL, Platelets - 1485,000/mm³. White blood cells showed marked leukocytosis with complete spectrum of neutrophil lineage including myeloblasts. The bone marrow Trephine biopsy was markedly hypercellular with fibroblastic proliferation, megakaryocytic proliferation with dysplastic megakaryocytes, markedly suppressed other haemopoietic elements, vascular sinusoidal proliferation and reticulin stain was grade 3. The patient continued with dose adjustment of hydroxycarbamide. At one year of follow up she maintained hemoglobin between 9-10 g/dL with platelet counts ranging from 400-800x10⁹ and the white cell count was controlled around 15,000 x 10⁹/L. She now has palpable splenomegaly. Abdominal ultrasound shows a splenomegaly 14cm below costal margin. She fulfilled all the diagnostic criteria of MF secondary to PV.

Discussion

Myeloproliferative neoplasms are a collection of diseases that have overlapping clinico-pathological features and this is a challenge to the hematologist distinguishing one from the other. This is particularly true in ET and early phase/pre-fibrotic form of MF.

The diagnosis of primary myelofibrosis is based on the 2008 World Health Organization (WHO) criteria, which include histopathological, morphological, clinical, and molecular-cytogenetic variables. To confirm a diagnosis of PMF, patients must meet all three major criteria plus two minor criteria. The major criteria are largely histopathology-based. For example, the presence of increased megakaryopoiesis with a preponderance of atypical megakaryocytes is an important feature of PMF. Such findings are usually associated with increased bone marrow cellularity. The later, although less frequently, can also occur in cases of ET. The second major criterion is excluding the other MPD (CML,MDS or PV) which have a similar presentation. The third criterion is demonstration of JAK2V617F or the absence of reactive marrow fibrosis [3]. Minor criteria include the presence of a leukoerythroblastic blood picture, a high Lactate Dehydrogenase (LDH) level, splenomegaly, and/or anaemia.

First two patients have massive splenomegaly, leukoerythroblastic blood picture with tear drop poikilocytosis but no cytopenias. In addition to bone marrow fibrosis, megakaryocytic dysplasia was present with clustering of megakaryocytes. BCR-ABL was negative in both cases and JAK2V617F mutation was positive in the first case. They were diagnosed as fibrotic stage of primary myelofibrosis and therefore they have been treated with hydroxycarbamide 500 mg daily. The extra medullary hematopoiesis was well controlled with improvement of their splenomegaly.

In case 3, compared to previous cases, patient had no organomegaly, leukoerythroblastic blood film or tear drop poikilocytosis but had pancytopenia. Bone marrow histology showed features resembling idiopathic bone marrow fibrosis such as clustered megakaryocytes and dense fibrosis. Currently patient is on regular blood transfusions.

In case 4, patient had a previous history of PV with grade 3 bone marrow reticulin fibrosis and increasing splenomegaly. Patient was treated with hydroxycarbamide/hydroxyurea. The International Prognostic Scoring System (IPSS) uses the following 5 risk factors for estimating survival at the time of diagnosis: age > 65 years, hemoglobin level<10 g/dL, leukocyte count >25 x 10⁹/L, circulating blasts ≥ 1%, and the presence of constitutional symptoms. The presence of 0, 1, 2, and ≥ 3 adverse factors define low, intermediate 1, intermediate 2, and high-risk disease with median survivals of 11.3, 7.9, 4, and 2.3 years, respectively. Later IPSS was modified to Dynamic IPSS (DIPSS). Most recently, DIPSS9.10 was upgraded to DIPSS-plus11 by the incorporation of 3 additional IPSS/DIPSS independent risk factors including red cell transfusion need, platelet count <100 x 10⁹/L, and unfavorable karyotype; the latter includes sole complex karyotype or 2 abnormalities that include +8, −7/7q−, I (17q), inv (3), −5/5q−.
12p−, or 11q23 rearrangement. The eight (8) DIPSS-plus risk factors are used to define low (no risk factors), intermediate 1 (1 risk factor), intermediate 2 (2 or 3 risk factors), and high (≥ 4 risk factors) risk groups with respective median survivals of 15.4, 6.5, 2.9, and 1.3 years.

All the cases above come under the high risk category. These cases illustrate the significant heterogeneity that is seen in patients with MF which could cause a challenge in the diagnosis and management of myelofibrosis.

References