Presentation of chronic myeloid leukemia as promyelocytic blastic transformation: A case report and literature review

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Abstract
Chronic myelocytic leukemia (CML) is a clonal disorder involving the pluripotent stem cell and is consistently associated with the BCR-ABL1 fusion gene located on the Philadelphia chromosome. The disease typically evolves in 3 distinct clinical stages: chronic and accelerated phases and blast crisis. The promyelocytic blast crisis is a rare form of transformation during the evolution of CML. Here we report a case of promyelocytic blastic crisis with t(15;17) in addition to t(9;22).

Abbreviations: BMA: Bone Marrow Aspiration; BMB: Bone Marrow Biopsy

Introduction
Chronic myelocytic leukemia (CML) is a clonal disorder of pluripotent stem cell. It is characterized by the dysregulated production and uncontrolled proliferation of mature and maturing granulocytes with fairly normal differentiation. Philadelphia chromosome occurs in 95% of CML cases. CML is associated with the fusion of two genes: BCR (on chromosome 22) and ABL1 (on chromosome 9) resulting in the BCR-ABL1 fusion gene [1]. Acute promyelocytic leukemia (APL) is a biologically and clinically different type of acute myeloid leukemia (AML). It occurs result of reciprocal translocation of PML-RARA fusion gene on chromosome 15 and 17 and positive 90% of AML patients. In CML progression toblastic transformation develops block of differentiation and apoptosis CML stem cells or progenitor cells. Genetic instability and epigenetics modifications of BCR-ABL is probably responsible for this pathogenesis and acute leukemic transformation [2]. The promyelocytic blast crisis is a rare form of transformation during the evolution of CML. The morphology and immunophenotype of the blasts were similar to those seen in acute promyelocytic leukemia APL (3). Here we report a case of promyelocytic blastic crisis with t(15;17) in addition to t(9;22).

Case report
A 19-year old man applied the hematology department with complaints of abdominal pain, fatigue, nausea, vomiting and epistaxis for one month. He had found out through the medical history that he was diagnosed with CML 6 months ago and Imatinib mesylate equivalent to 400 mg/day was appointed (Figure 1a). An irregular intake of Imatinib mesylate was also found out. A physical examination showed that the spleen had crossed costal margin 12 cm and no hepatomegaly. Complete blood count was also found out that the hemoglobin level was 7.4 gr/dl (13.5-17.5 gr/dl), platelet count was 16,000/mcl (150,000-450,000/ mcl) and WBC count was 3.3 × 10^9/L (3.5-10.5 × 10^9/L). The patient had no evidence of coagulopathy with clinical and laboratory findings of disseminated intravascular coagulation (DIC). Promyelocytic blast with Auer rod were observed in the proportion of 70% to hypercellularity after the bone marrow aspiration and biopsy examination (Figure 1b-d). Flow cytometric analysis revealed that more than 20% of blast cells were immunopositive for CD45, CD13, CD33, CD117, MPO and immune-negative for CD34, HLA-DR expression. Translocation of t(9;22)(BCR-ABL) and t(15;17) were positive.

A diagnosis of chronic myelocytic leukemia with promyelocytic blastic transformation was made. Remission induction chemotherapy was initiated with ATRA 45 mg/m2, cytosine arabinoside (200 mg/m^2/day), idarubicin (12 mg/m^2/day) and dasatinib 140 mg/day. After induction chemotherapy a bone marrow biopsy and aspiration being repeated after a month. Normocellular bone marrow and aspiration was seen (Figure 2). T(15;17) translocation was resulted positively in the proportion of 0.0004. T(9;22)(BCR-ABL) area was seen positive in the proportion of 0.004 and t(15;17) was negative in the analysis of blood sample taken through RT-PCR method from a bone marrow after chemotherapy. The second cure consolidation chemotherapy was commenced for the patient with dasatinib 140 mg in a day. There were negativity of t(15;17) and, t(9;22) . Allogeneic bone marrow transplantation was planned in the case and follow-up as well as treatment is being continued.

Discussion
The promyelocytic blastic crisis is an uncommon type of CML blastic transformation. We report a case of promyelocytic blastic crisis with t(15;17) in addition to t(9;22). The morphology and immunophenotype

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CML develops in ranges to 8 months to 9 years normally [4]. Blastic transformation was diagnosed with one year in our patient. We found no chromosomal aberrations and translocations. Genomic instability and less vulnerability of CML stem cell to imatinib may cause to CML progression in a little time. Although imatinib mesylate was used by the patient whom BCR-ABL suppressed or undergo apoptosis while leukemic cells developed to clonal evaluation [5].

Promyelocytic blastic crisis of CML is highly rare, accounting for about 30 cases searching English literature before 2007 [5]. The first case with promyelocytic blastic crisis during Imatinib mesylate therapy was published by Gozetti et al. [6] who is 50 years old male patient.

In the early stages of CML chronic phase a clonal evolution developed by the adding of PML/RARA translocation may result with this clinical illustration. Some novel translocations and chromosomal aberrations, like t(5;13)(q12;p13) and t(15;20)(q22;p13), i(7q), +8, del(7q) may occur during promyelocytic crisis of CML [7]. But, De Botton et al. [8] showed no significant difference in complete remission, event-free survival, relaps or overall survival at 2 years between the prognosis of APL with additional chromosomal changes and t(15;17) in 292 patients who are treated with ATRA and chemotherapy. The most commonly seen chromosomal abnormalities were trisomy 8, deletions and translocations chromosomes 9, 7, 21 and 17 [8]. Rosenthal et al. [9] in a series of 89 cases of blast crisis of CML found two cases classified as APL. The two of CML revealed t(15;17) in addition to t(9;22). Both patients developed disseminated intravascular coagulation at presentation or during treatment of blastic crisis. Oki et al. [10] reported a 66-year-old woman with promyelocytic crisis of CML diagnosed by demonstration of both BCR/ABL and PML/RAR alpha chimeric genes in leukemic cells by karyotype and fluorescence in situ hybridization analysis. Authors claimed that clonal evolution with addition of the PML/RAR alpha translocation may have arisen in the early chronic phase of CML, with expansion of this clone during imatinib treatment [10]. Yamamoto and his friends reported a CML case transformed into APL with T315I mutation in only one CML patient who used dasatinib. In this case combination therapy with imatinib and arsenic trioxide achieves favorable outcomes for promyelocytic blastic crisis finally with allogeneic stem cell transplantation [11].

Additional chromosomal abnormalities had higher rates of coagulopathy, thrombocytopenia and higher relapse rate compared to t(15;17) alone [12]. But, our patients had no findings of coagulopathy which may have relation with no chromosomal abnormalities and achieved complete cytogenetic remission after second chemotherapy cycle with dasatinib. This is the first case who was achieved complete remission with using dasatinib during promyelocytic blastic crisis. It is possible to assume that dasatinib contributed to achieve complete remission in this case with negativity of t(9;22) after treatment. Transformation of CML to APL is a rare clinical setting appropriate treatment approach has yet to be determined.
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