Onset of chronic myeloid leukemia with complex karyotype in a pregnant patient: case report and revision of literature

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Abstract: Approximately 10%–12% of patients in chronic-phase chronic myeloid leukemia (CP-CML) have additional chromosomal aberrations at diagnosis; moreover, CML occurs in up to 10% of pregnancy-associated leukemias, with an annual incidence of 1 per 100,000 pregnancies. In this report we describe the case of a 36-year-old female with CP-CML diagnosed in the 18th week of pregnancy and with a new complex variant translocation t(4;9;22;21)(q24;q34;q11;q22) and an additional chromosomal aberration t(1;20)(p36;p11). In consideration of her pregnancy, the patient strictly monitored her blood cell count without any specific treatment. At 32 weeks of pregnancy, the patient delivered via cesarean section a healthy baby girl. After 10 days from childbirth, dasatinib was started at a standard dosage of 100 mg/day and 3 months later complete cytogenetic response and major molecular response were obtained, with the achievement of an optimal response according to European Leukemia Net recommendations and showing efficacy of this tyrosine kinase inhibitor (TKI) in the presence of a complex karyotype.

Keywords: chronic myeloid leukemia, complex karyotype, pregnancy, additional chromosomal aberration, variant translocation, tyrosine kinase inhibitor

Introduction

Chronic myeloid leukemia (CML) is a myeloproliferative neoplasm characterized by the presence of the Philadelphia chromosome (Ph) resulting from the reciprocal translocation t(9;22)(q34;q11). 1 Approximately 10%–12% of patients in chronic-phase CML (CP-CML) present with additional chromosomal aberrations (ACAs) at diagnosis.2 Recently different researchers have studied this subgroup, including European Leukemia Net (ELN) 3 and have classified CML ACAs into “major” and “minor” route changes.4 The major route ACAs are the most common chromosomal abnormalities (>10% of cases with ACAs) and include trisomy 8, an extra Ph (1der(22)(t(9;22)q34;q11)), isochromosome 17(i(17)(q10)), trisomy 19 and ider(22)(q10)t(9;22)(q34;q11). Other less common ACAs belong to minor route ACAs. Moreover, microdeletions can be present too and the occurrence of genomic microdeletions proximally to ABL1 or distally to BCR have been reported in CML cases with variant translocations with a greater frequency (30%–40%) than in cases with classic t(9;22) (10%–18%).5 When the chromosome changes are submicroscopic, the translocation can be masked and revealed only by fluorescence in situ hybridization (FISH) or by molecular analysis.6

Chromosome 9 deletions and variant translocations have no value for prognosis whereas ACA/Ph1 have been reported to have an adverse prognostic value, particularly in the case of the major route abnormalities, including, as described previously, trisomy 8, an extra Ph (1der(22)(t(9;22)(q34;q11)), isochromosome 17 (i(17)(q10)),...
trisomy 19, and ider(22)(q10)t(9;22)(q34;q11). Major route ACA/Ph1 at diagnosis do not mandate different initial treatments but represent a warning: this implies that the characteristics of the disease and the response to treatment require more frequent monitoring to permit timely changes in therapy in case of treatment failure. Major route ACA/Ph1 developing during treatment were confirmed to be a signal of acceleration. CML occurs in up to 10% of pregnancy-associated leukemias, with an annual incidence of 1 per 100,000 pregnancies. In this report we describe the case of a 36-year-old female diagnosed in her 18th week of pregnancy with CP-CML and presenting with a new complex variant translocation t(4;9;22;21)(q24;q34;q11;q22) and an ACA t(1;20)(p36;p11). The rearrangement was analyzed by cytogenetic and FISH tests. To the best of our knowledge, this translocation has not been described in CML previously and specifically in a pregnant patient.

Case report
Patient presentation
A 36-year-old female, 18 weeks pregnant, presented in October 2015 with abnormal blood cell counts (white blood count 29.87×10^9/L, hemoglobin 11.5 g/dL and platelets 592×10^9/L); evaluation of the peripheral blood smear revealed basophils (2%), myelocytes (7%), metamyelocytes (13%), and blasts (1%). Bone marrow aspiration was performed and its examination revealed a slightly hypercellular marrow with granulocytic hyperplasia. Cytogenetic analysis revealed the presence of a complex karyotype in all examined metaphases: 46 XX, t(1;20)(p36;p11), t(4;9;22;21)(q24;q34;q11;q22). Reverse transcriptase-polymerase chain reaction (RT-PCR) done on peripheral blood and bone marrow showed a b3a2 BCR/ABL fusion gene. The patient was diagnosed with Ph+ CML (low risk Sokal score). In consideration of her pregnancy and leukocyte and platelet values she was advised to strictly monitor her blood cell count, with no other treatment apart from a low dose of aspirin. At 32 weeks (January 2016), the patient delivered via cesarean section a healthy baby girl (weight 2,120 g, height 43 cm; APGAR 9). At the time of childbirth, the hematological analysis revealed a white blood cell count of 73.51×10^9/L, a hemoglobin level of 10 g/dL, a platelet count of 494×10^9/L. After 10 days from childbirth, the patient started dasatinib at a standard dosage of 100 mg/day. After 3 months on dasatinib the patient obtained complete cytogenetic response and major molecular response (MMR), achieving an optimal response according to the ELN recommendations.

Cytogenetic analysis
Cytogenetic analysis was performed on bone marrow culture using a standard technique. In total, 20 GTG banded bone marrow metaphase cells were analyzed. The karyotypes were named according to the International System for Human Cytogenetic Nomenclature. The cytogenetic analysis performed on 20 metaphase cells identified the presence of a complex, four-way (4;9;22;21)(q24;q34;q11;q22) Ph chromosome translocation (Figure 1) and an additional chromosomal aberration t(1;20)(p36;p11). In the present case, the Philadelphia translocation is likely to be the first event, followed by the further rearrangement involving chromosomes 19 and 22. Reverse transcriptase-polymerase chain reaction (RT-PCR) done on peripheral blood and bone marrow showed a b3a2 BCR/ABL fusion gene. The patient was diagnosed with Ph+ CML (low risk Sokal score). In consideration of her pregnancy and leukocyte and platelet values she was advised to strictly monitor her blood cell count, with no other treatment apart from a low dose of aspirin. At 32 weeks (January 2016), the patient delivered via cesarean section a healthy baby girl (weight 2,120 g, height 43 cm; APGAR 9). At the time of childbirth, the hematological analysis revealed a white blood cell count of 73.51×10^9/L, a hemoglobin level of 10 g/dL, a platelet count of 494×10^9/L. After 10 days from childbirth, the patient started dasatinib at a standard dosage of 100 mg/day. After 3 months on dasatinib the patient obtained complete cytogenetic response and major molecular response (MMR), achieving an optimal response according to the ELN recommendations.

Figure 1 Cytogenetic analysis.
Notes: Chromosome analysis by G banding showing a complex, four-way (4;9;22;21)(q24;q34;q11;q22) Philadelphia chromosome translocation and an additional chromosomal aberration t(1;20)(p36;p11). Of the two chromosomes 21, is visible only the normal one; the other is split among 4,9 and 22 chromosomes. Arrows indicate rearranged chromosomes.
4q and 21q, in a step by step sequence. The additional t(1;20) (p36;p11) can be considered an independent event.

**FISH analysis**

FISH analysis using a BCR/ABL dual color dual fusion probe (Figure 2), showed the presence of a variant BCR/ABL translocation in approximately 80% of interphase nuclei with a single fusion signal; two BCR and two ABL signals showing the splitting of one of the two expected fusion signals, thus suggested the involvement of additional chromosomes and, therefore, we extended the analysis to metaphase spreads. Painting probes for chromosomes 4 and 22 confirmed the presence of a translocation involving chromosomes 4 and 22, generating two derivative chromosomes, one normal chromosome 4 and one normal 22 (Figure 3).

**Molecular testing**

At diagnosis, nested RT-PCR was done on peripheral blood and bone marrow showing a b3a2 BCR/ABL fusion gene; at 3 months real-time quantitative RT-PCR was perfomed using TaqMan system and a BCR-ABL1 transcripts’ level <10% (0.0829%) according to the International Scale was found (MMR).

**Cytological analysis**

Examination of aspirated bone marrow revealed a slightly hypercellular marrow with granulocytic hyperplasia. Histological examination of the placenta revealed the presence of chorionic villus with characters of development corresponding to gestational age (32 weeks), with intervillous space characterized by the presence of increased lympho-monocytic and granulocytic elements. Membranes and umbilical cord were without significant histological alterations.

**Discussion**

In this report we describe a unique case of a 36-year-old female with CP-CML diagnosed at 18 weeks of pregnancy, characterized by a new complex variant translocation t(4;9;22;21)(q24;q34;q11;q22) and t(1;20)(p36;p11) as ACA. The rearrangement was analyzed by cytogenetic and FISH tests. To the best of our knowledge, this translocation has not been observed in CML previously and particularly in a pregnant patient. The management of CML diagnosed during pregnancy is very challenging due to the physiological changes, including those in hematological parameters which accompany the pregnancy, that may mask the symptoms. Previously, it seemed that CML diagnosed during pregnancy was associated with low birth weight and preterm birth, but this is no longer described in more recent reports. Reassuringly, the course of the disease does not appear to be affected by pregnancy. The prothrombotic potential of a normal pregnancy is well recognized as a result of a physiological increase in hemostatic factors and prothrombotic proteins in addition to the physical obstruction of venous blood flow. As a result, thrombosis continues to be the most common
cause of maternal morbidity and this may be compounded in the myeloproliferative diseases where there is an associated elevation in the platelet count. Therapeutic approaches for CML diagnosed in pregnancy include supportive care in the form of leukapheresis and platelet pheresis and interferon-a (IFN-a), while the use of tyrosine kinase inhibitors (TKIs) is controversial. Leukapheresis and platelet pheresis allow avoidance of potentially teratogenic drugs but are not easily available and tolerable. The frequency of these procedures depends on white cell and platelet count. IFN-a, which has a high molecular weight, does not cross the placenta and does not inhibit DNA synthesis; for these reasons it is considered safe in pregnancy. Among TKIs, imatinib is the most studied. It does not cross the placenta but various congenital abnormalities have been described after exposure to imatinib in the first trimester, when placenta formation is not completed, probably due to platelet derived growth factor receptor alpha inhibition. Thus, imatinib and generally all TKIs should be avoided in the first trimester and during organogenesis. Dasatinib crosses the placenta and leads to considerable levels in fetal plasma. It should be avoided in all pregnant patients since it is responsible for fetal hydrops and severe fetal bicytopenia even if normal pregnancies during exposure have been reported. Nilotinib does not cross the placenta in a significant concentration and does not seem to be teratogenic, but data are limited. A recent review summarized reported cases and provided recommendations of management of pregnancy in CML; so far no case of pregnancy in a CML patient associated with bosutinib and ponatinib therapy has been described. However, as in our case, treatment is not always mandatory; it is necessary if white cell count exceeds 100×10⁹/L and platelet count exceeds 500×10⁹/L, as reported by Milojkovic and Apperley. Low-molecular-weight heparin, as well as aspirin, can be used. Regarding the complex karyotype of our patient, Philadelphia translocation is likely to be the first event, followed by the further rearrangement involving chromosomes 4q and 21q, in a step by step sequence. The additional t(1;20)(p36;p11) can be considered an independent event. t(1;20) has been reported associated with a high rate of recurrent first trimester abortions in a large family; in our case, the mother did not spontaneously abort her child during the pregnancy despite the presence of this aberration. suggested two possible mechanisms which may be involved in the formation of variant or complex translocations. The first is a single event in which rearrangement due the simultaneous breakage of several chromosomes is by mismatched joining. The second is a multi-step mechanism in which a classical Ph translocation is followed by further translocation events involving chromosomes 9 and 22, plus a second, third, and subsequent event leading to a multiple-way translocation. Concerning treatment, after strict observation, low dose aspirin and a safe delivery, our patient was treated with dasatinib, a second-generation TKI, at a dosage of 100 mg per day, considering early age and the complex, never described before, translocation of uncertain prognostic significance. Dasatinib is a multi-targeted kinase inhibitor of BCR/ABL, SRK, c-KIT, ephrin receptors, and PDGFRB and from 2012 it can be used also in first line treatment of CML. Treatment was well tolerated with no significant side effects. After 3 months the patient showed a complete cytogenetic response and MMR achieving an optimal response according to ELN recommendations, proving dasatinib’s efficacy also in the case of complex karyotype.

Conclusion
We report a novel case of CP-CML with a new complex variant translocation t(4;9;22;21)(q24;q34;q11;q22) and an ACA t(1;20)(p36;p11). Notably, the patient concerned showed a good tolerability and a good response to dasatinib also, in the presence of a complex karyotype.

Acknowledgment
Written informed consent was obtained from the patient for publication of this case report and any accompanying images.

Disclosure
The authors report no conflicts of interest in this work.

References