Insertion of the 3' ABL region into the long arm of chromosome 1 in a Philadelphia chromosome-negative chronic myeloid leukemia case

WALID AL-ACHKAR¹, THOMAS LIEHR² and ABDULSAMAD Wafa¹

¹Molecular Biology and Biotechnology Department, Human Genetics Division, Atomic Energy Commission of Syria, Damascus, Syria; ²Jena University Hospital, Institute of Human Genetics and Anthropology, Jena, Germany

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Abstract. Chronic myeloid leukemia (CML) is a pluripotent hematopoietic stem cell disorder almost always characterized by the presence of the Philadelphia chromosome (Ph), usually due to t(9;22)(q34;q11). The presence of Ph results in the formation of the BCR/ABL fusion gene, which is a constitutively activated tyrosine kinase. Approximately 1% of CML patients appear to have a Ph-negative karyotype but carry a cryptic BCR/ABL fusion that can be located by fluorescence in situ hybridization (FISH) at chromosome 22q11, 9q34 or a third chromosome. This study investigated a rare Ph-negative CML case with insertion of the 3' ABL region into the long arm of derivative chromosome 1 but lacking the 5' BCR region on der(22).

Introduction

Chronic myeloid leukemia (CML) is a pluripotent hematopoietic stem cell disorder defined by expression of the BCR/ABL fusion gene, a constitutively activated tyrosine kinase, harbored by the Philadelphia chromosome (Ph), which is the result of a t(9;22)(q34;q11) or a related variant translocation (1). In approximately 1% of CML patients, bone marrow cells appear to be Ph-negative by G-banding, although the BCR/ABL fusion gene can be identified by molecular means and can be located by fluorescence in situ hybridization (FISH) on chromosome 22q11, 9q34 or even a third chromosome (2-5). Cases of Ph-negative BCR/ABL-positive CML with the chimeric gene present on derivative chromosome der(22), as in the majority of CML cases, or alternatively on der(9) appear to have the same clinical and molecular characteristics as Ph-positive patients. However, a worse prognosis associated with the location of BCR/ABL on der(9) was previously noted (6-9).

The biology and clinical significance of genetic rearrangements in Ph-negative BCR/ABL-positive disease was evaluated following initial descriptions (2-5). Two mechanisms involved in the formation of the chimeric gene in masked Ph-positive cells have been postulated: insertion of ABL into the BCR region (or vice versa), or by a multiple-step model where a classical t(9;22) is followed by translocation of the two products and/or another autosome, thereby restoring the normal chromosome morphology. In both instances, more than the 2 breaks associated with classical t(9;22) are implicated (10).

This study investigated a rare CML case, Ph-negative, with an insertion of the 3' ABL region into the long arm of derivative chromosome 1 and lacking the 5' BCR region on der(22).

Materials and methods

Case report. In July 2005, a female patient, 31 years of age, presented for the first time with a whole blood cell count (WBC) of 35.07x10⁹/l (72.3% neutrophils, 13% lymphocytes, 1.16% eosinophiles, 5.7% monocytes, 4.1% basophiles and 3.6% immature cells). The platelet count was 381x10⁹/l, and the hemoglobin level was 9.8 g/dl. A physical examination showed no splenomegaly, although loss of weight was noted.

Results of a chromosome analysis using banding cytogenetics showed a karyotype in concordance with the clinical diagnosis of CML in the chronic phase (CP). The patient was treated with hydroxyurea (1500 mg daily dose) for a duration of two years and eleven months. In June 2008, the patient presented for the second time with a WBC count of 35.07x10⁹/l (72.3% neutrophils, 13% lymphocytes, 1.16% eosinophiles, 5.7% monocytes, 4.1% basophiles and 3.6% immature cells). The platelet count was 381x10⁹/l, and the hemoglobin level was 9.8 g/dl. A physical examination showed no splenomegaly, although loss of weight was noted. Results of a chromosome analysis using banding cytogenetics showed a karyotype in concordance with the clinical diagnosis of CML in the chronic phase (CP). The patient was treated with hydroxyurea (1500 mg daily dose) for a duration of two years and eleven months. In June 2008, the patient presented for the second time with a WBC count of 35.07x10⁹/l (72.3% neutrophils, 13% lymphocytes, 1.16% eosinophiles, 5.7% monocytes, 4.1% basophiles and 3.6% immature cells). The platelete count was 381x10⁹/l, and the hemoglobin level was 9.8 g/dl. A physical examination showed no splenomegaly, although loss of weight was noted.
**Band configurations.** Banding cytogenetics using the GTG-method was conducted according to standard procedures (11). A total of 20 metaphases derived from the unstimulated

Figure 1. GTG-banding revealed a karyotype 46,XX.

Figure 2. Karyotype and chromosomal aberrations were unconfirmed using molecular cytogenetic approaches. (A) The BCR/ABL fusion gene was absent on der(22), while the ABL/BCR fusion gene was observed on der(9). Insertion of the 3’ ABL region into the long arm on der(1) and lack of the 5’ BCR region on der(22) were noted. (B) The BCR/ABL fusion gene was absent on der(22), while the ABL/BCR fusion gene was observed on der(9). Insertion of the 3’ ABL region into the long arm on der(1) confirmed by using wcp1 and lack of the 5’ BCR region on der(22) were noted. (C) The application of FISH using wcp for chromosomes 1 and 9 did not reveal a translocation between these chromosomes. (D) The application of FISH using wcp for chromosomes 1 and 22 did not reveal a translocation between these chromosomes. #, chromosome; der, derivative chromosome; Ph, Philadelphia-chromosome.

Figure 3. Schematic representation of chromosomal changes that occurred in the patient. The BCR/ABL probe showed a fusion signal on der(9) and insertion of 3’ ABL on der(1). The 5’ region of BCR was deleted on der(22) while the 3’ region of BCR remained on chromosome 22. #, chromosome; der, derivative chromosome.
bone marrow of the patient were analyzed individually. Karyotypes were described according to the International System for Human Cytogenetic Nomenclature (12).

Fluorescence in situ hybridization. FISH using BCR/ABL dual-color dual fusion translocation probe (Abbott Molecular/Vysis, USA) and whole chromosome painting (wcp) probe for chromosomes 1, 9 and 22 (MetaSystems, Germany) were applied as previously described (11). A total of 50 metaphase spreads were analyzed, each using a fluorescence microscope (Axio Imager.Z1 mot; Zeiss) equipped with appropriate filter sets to discriminate between a maximum of five fluorochromes and the counterstain DAPI (4',6-diamino-2-phenylindole). Image capturing and processing were carried out using an image analysis system (MetaSystems, Altussheim, Germany).

Results

A normal karyotype 46,XX was determined in the GTG-banding (Fig. 1) and was further studied by molecular cytogenetics (Fig. 2). Dual-color-FISH using probes specific for BCR and ABL revealed that the BCR/ABL-translocation fusion gene was absent on der(22). However, the presence of the ABL/BCR-translocation fusion gene on der(9), insertion of the 3’ ABL region on der(1) and the 5’ BCR region on der(22) were noted (Fig. 2A and B). Moreover, FISH using wcp1, wcp9 and wcp22 probes did not show a translocation between chromosome 1 and 9 or chromosome 1 and 22 (Fig. 2C and D). Thus, the result obtained was: 46,XX,ins(1;9)(q21.3;q34;q34),ins(9;22)(q34q11q11),del(22)(q11q11).

Discussion

The present study identified one additional chromosomal alteration, insertion of the 3’ ABL region on der(1), in a Ph-negative BCR/ABL-positive CML-CP case. To the best of our knowledge, the insertion of the ABL region into der(1) has never been described in CML (13).

According to the literature, two alternative mechanisms were postulated to elucidate the formation of a fusion gene in Ph-negative BCR/ABL-positive CML patients (10,14). The first mechanism involves a one-step model where BCR/ABL results from a simple insertion of either 3’ ABL into BCR or 5’ BCR into ABL after three genomic breaks. The second mechanism is a multiple-step model involving an initial classical t(9;22)(q34;q11) followed by a second translocation of the two products and/or a third chromosome, requiring a minimum of 4 genomic breaks (14).

In the present case, FISH results led to the conclusion that insertion of 5’ BCR sequences within the ABL gene transpires while 3’ BCR sequences remained on chromosome 22 (Fig. 3). Previous studies on other Ph-negative patients with the BCR-ABL fusion gene at 9q34 also proposed the insertion of chromosome 22 sequences into chromosome 9 as the more likely mechanism, as opposed to one involving two consecutive translocations. This event appears more likely as it requires only two breaks at chromosome 22 and one at chromosome 9 instead of a total of four breaks involved in the double-consecutive translocation (15).

A second event, insertion of 3’ ABL sequences on der(1), possibly shifted ABL from der(22) to the long arm of chromosome 1. The 5’ region of the BCR gene was deleted on der(22). The patient presented with CML-CP. Thus, the second event may be the evolution of a Ph-negative karyotype.

In conclusion, regarding the karyotype observed in the present case with the insertion mechanism, a minimum of five breaks are required; one on chromosome 1, two on chromosome 9 and two on chromosome 22, to explain the insertion and deletion.

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