Acute Myelogenous Leukemia without Maturation with a Retinoic Alpha-Receptor Deletion: A Case Report

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Acute myelogenous leukemia without maturation · PML/RAR? · Deletion · t(15;17)(q22;q21) · Acute promyelocytic leukemia · AML-M3 · AML-M1

Abstract
Acute promyelocytic leukemia (APL) is characterized by a t(15;17) which fuses the 17q retinoic acid alpha-receptor sequence to the 15q PML gene sequence. The resulting fusion product plays a role in the development and maintenance of APL, and is very rarely found in other acute myeloid leukemia (AML) subtypes. Rare complex APL genomic rearrangements have retinoic acid alpha-receptor sequence deletions. Here we report a retinoic acid alpha-receptor sequence deletion in a case of AML without differentiation. To our knowledge, this is the first example of a retinoic acid alpha-receptor sequence deletion in this AML subtype.

Introduction

Approximately 95% of acute promyelocytic leukemias (APLs, FAB AML-M3) carry the t(15;17)(q22;q21), which fuses the retinoic acid alpha-receptor (RARα) sequence to the PML gene sequence [1]. The remaining APL gene fusions are composed of rare variant fusions of RARα with STAT5, NUMA, PLZF, and NPM [2]. The resulting fusion gene products play a pivotal role in the pathogenesis of APL and APL sensitivity to all-trans retinoic acid and arsenic trioxide [1, 3]. Occasional APL cases carry complex genomic rearrangements which include RARα sequence deletions. These deletions are apparently very rare compared to
similar deletions in other leukemias associated with disease-specific cytogenetic changes [4–8]. Additionally, t(15;17)(q22;q21) is rare in non-APL acute myeloid leukemias (AMLs), with only rare examples of this translocation identified in AML without maturation and AML with maturation (FAB M1 and M2 subtypes, respectively) [9]. To our knowledge, what we describe here is the first case of a RARα deletion in a case of AML without maturation (FAB AML-M1).

Case Report

A 51-year-old male with a history of hepatitis C and a forty pack-year smoking history presented with neutropenia over the course of several months. A peripheral blood smear confirmed neutropenia with conspicuous atypical lymphocytes and large granular lymphocytes (fig. 1a). Bone marrow biopsy revealed an erythroid predominant hypocellular marrow with a fat-to-cell ratio of 65:35. CD34 immunohistochemical staining showed approximately 25% blast forms in the cellular marrow (fig. 1b). The marrow blasts were negative for myeloperoxidase on immunoperoxidase staining, were small with scant cytoplasm, and had no significant granularity. On aspirate smears, blasts comprised approximately 50% of the marrow cellularity. The blast forms were small with scant cytoplasm and no significant cytoplasmic granularity. Minimal maturation was evident on the myeloperoxidase staining, reiterating the morphologic features of the marrow cells. Blast percentages were determined on a 500-cell differential in bone marrow and/or a 200-cell peripheral blood smear differential. Erythroid maturation was evident with CD71 staining and appeared macronormoblastic and megaloblastoid. Based on these findings, a diagnosis of AML-M1 (AML without maturation) was rendered.

Molecular analyses were performed on the bone marrow aspirate. Cytogenetic analysis of metaphase cells revealed a normal male karyotype, 46,XY. No clonal T-cell gene rearrangements, FLT3 internal duplications or D835 mutations, or NPM1 exon 12 mutations were detected. AML/myelodysplastic syndrome FISH panels revealed no markers of myelodysplasia. However, the AML panel revealed loss of the RARα/17q locus as detected by the RARA(F) probe. No evidence of a t(15;17)(q22;q21) was identified by molecular diagnostics.

Discussion

t(15;17)(q22;q21) characterizes APL and plays an important role in the pathogenesis of this disease and its sensitivity to all-trans retinoic acid and arsenic trioxide [1, 3]. Very rare examples of non-APL with this translocation have been identified [9]. Additionally, a small number of APL with complex chromosomal rearrangements involving deletions at the RARα/17q locus have been described [4–8]. To our knowledge, RARα/17q locus deletions have not previously been detected in AML without differentiation (AML-M1). The significance of the RARα/17q locus deletion in this AML type is currently unknown, although loss of the retinoic acid alpha-receptor is associated with increased cellular growth potential [10]. Thus this deletion may have a role in increasing tumor growth. Further analysis of the function of this deletion in the AML-M1 subtype would be difficult due to the rarity of this molecular event.
References


Fig. 1. Representative H&E and immunohistochemical staining results for the patient's case of AML-M1. a High-power H&E of the conspicuous atypical lymphocytes and large granular lymphocytes in the peripheral smear preparation. b Immunohistochemical staining results for CD34 showing approximately 25% blast forms in the cellular marrow.