Chronic eosinophilic leukemia with FIP1L1-PDGFRα transcripts after occupational and therapeutic exposure to radiation

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Abstract

We present for the first time a 40-year-old male patient with a 20 year history of occupational exposure to radiation as a nuclear power plant worker, who developed FIP1L1-PDGFRα-positive chronic eosinophilic leukemia 27 months after radiotherapy for testicular seminoma. After an one-year history of dry cough, itching and night sweats, the patient presented with an elevated leukocyte count with absolute eosinophilia of 14.2x10⁹/L, bone marrow and lymph node involvement. Treatment with imatinib was initiated, resulting in complete hematological remission at the sixth month and complete molecular response by nested primers reverse transcription polymerase chain reaction – at the end of the first year. This case contributes to the clinical heterogeneity of a rare entity such as FIP1L1-PDGFRα-positive myeloproliferative neoplasms, and for the possible role of occupational and therapeutic radiation, raising the question if one or both of them might be the causative factor.

Introduction

The fusion gene FIP1L1-PDGFRα, containing regions of FIP1-like 1 (FIP1L1) gene and the Platelet-Derived Growth Factor Receptor Alpha (PDGFRα) gene as a result of the interstitial cryptic deletion on chromosome 4q12 [del(4)(q12g12)], is considered a recurrent molecular abnormality in patients with eosinophilia-associated myeloproliferative neoplasms (MPNs). The hyemic product of FIP1L1-PDGFRα acts as a constitutively active tyrosine kinase, however the precise underlying molecular mechanisms of FIP1L1-PDGFRα-mediated malignant transformation are still incompletely understood.1 According to the revised World Health Organization (WHO) Classification of Tumors of Hematopoietic and Lymphoid Tissues (2008), neoplasms bearing the FIP1L1-PDGFRα rearrangements are defined as a rare, separate disease entity. Clinically, the disease is manifested most often as chronic eosinophilic leukemia (CEL), but also as acute myeloid leukemia (AML) or T lymphoblastic lymphoma, or both simultaneously.2 As a rule, FIP1L1-PDGFRα-positive MPNs are presented as de novo malignancy, and only two cases following cytotoxic chemotherapy have been reported as therapy-related so far.2,3 To our knowledge, FIP1L1-PDGFRα-positive disease after radiation exposure has not been published yet.

In this study we present for the first time a patient with a 20 year history of occupational exposure to radiation, who developed FIP1L1-PDGFRα-positive CEL 27 months after radiotherapy for testicular seminoma.

Case Report

A 40-year-old male nuclear power plant worker was referred to The National Hospital for Active Treatment of Hematological Diseases in Sofia (September 2009) for diagnostic evaluation of leucocytosis and eosinophilia, detected ten months before his admission to our hospital, and resistant to corticosteroid therapy. The patient had a one-year history of dry cough, predominantly at night, itching and night sweats. Previous medical history included testicular seminoma, diagnosed 27 months earlier (June 2007) and treated with surgery followed by irradiation to the para-aortic and high iliac lymph nodes with a total dose of 36 Gy. No hematological abnormalities were observed during the whole period of regular clinical and laboratory follow up thereafter until December 2008, when the routine blood tests revealed eosinophilia (23% - 1.3x10⁹/L) with a normal white blood cell (WBC) count (5.6x10⁹/L), which progressively increased. On physical examination the patient had no hepato- or splenomegaly, but a single enlarged cervical lymph node. Radiological chest studies showed signs of minor interstitial pulmonary fibrosis in the hilar regions and no lung infiltration.

At the time of admission, laboratory tests revealed elevated WBC count of 18.1x10⁹/L, hemoglobin level of 128 g/L and platelet count of 290x10⁹/L. Peripheral blood differential showed 78% eosinophils (14.2x10⁹/L), 14% mature neutrophils (2.5x10⁹/L), 2% basophils (0.36x10⁹/L), and 6% lymphocytes (1.1x10⁹/L). Laboratory chemistry tests were within normal ranges. On aspirate smears bone marrow was markedly hypercellular due to a proliferation of abnormal eosinophils accounting for 51% of all bone marrow cells, including 31.5% eosinophilic myelocytes and 19.5% mature eosinophils. Most of the cells showed sparse granulation with clear areas of cytoplasm, some had pathological violet granules on Giemsa stains (Figure 1A). Eosinophils were positive for myeloperoxidase and were negative for non-specific esterase and toluidine blue stains. Neutrophils were 7.5% of all cells, erythroblasts - 37.5%, small non-granular blasts were 1% and megakaryocytes were reduced. Immunophenotyping on bone marrow aspirate was performed by flow cytometry using three color combinations of commercially available fluorescent dye labeled monoclonal antibodies (Becton Dickinson) and erythrocyte lyses and wash technique according to manufacturer’s instructions. Samples were acquired and analyzed on a BD FACSCanto II flowcytometer and FACSDiva software (Becton Dickinson). Expansion of eosinophils with high FSC and SSC and high expression of CD45 was found, that were positive for CD13, CD33, and CD15, and negative for CD34, CD117, CD64, CD14, and CD10, as well as for major lymphoid lineage associated markers. No aberrant antigen expression was detected. Blast cells were identified, accounting for...
2.6% of bone marrow cells showing myeloid phenotype - positive for CD34, CD33, CD13, CD117, CD38. Lymphoid populations were within the normal ranges. An excision biopsy of an enlarged cervical lymph node was performed in order to exclude a concomitant lymphoid neoplasm. The microscopic examination revealed nodal architecture almost completely effaced. Though there was a proliferation of preserved follicles with activated germinal centers, interfollicular areas showed massive infiltration by eosinophils at various stages of maturation as well as CD68-positive histiocytes and densely proliferating small vessels confirmed by immunohistochemistry for CD34 (Figure 1/B). Only few cells showed positive immunostaining for Mast cell tryptase. Unfortunately, at the time of admission the cytogenetic study was unsuccessful.

The subsequent analyses during the course of treatment revealed normal karyotype. Molecular analysis on RNA, extracted from bone marrow and peripheral blood cells, using Reverse Transcription Polymerase Chain Reaction (RT-PCR) revealed FIP1L1-PDGFRA rearrangement (Figure 1/C). Screening for BCR-ABL rearrangements and JAK2 gene V617F mutation was negative. Taking into account all above mentioned data, a diagnosis of chronic eosinophilic leukemia with FIP1L1-PDGFRA rearrangement was made and Imatinib treatment initiated with a daily dose of 100 mg. Normal WBC (4.3 × 10⁹/L) and eosinophil counts (1%) were achieved at the second month of therapy, however the bone marrow aspirations were dry until 6 months when a significantly hypocellular bone marrow with a complete remission was found. No FIP1L1-PDGFRA mRNA was detected by a single round RT-PCR and at the end of the first year a complete molecular response by nested primers RT-PCR was registered.

Discussion

Secondary malignancies are the most serious complications of otherwise successful treatment of some solid tumors. Myelodysplastic syndromes (MDS) and AML are the most common therapy-related hematological malignancies and a large amount of data is available regarding their epidemiology, molecular pathogenesis, clinical behavior and response to therapy. Secondary MPNs are significantly rarer. Almost all of these cases, reportedly more than 150 patients, present as secondary BCR-ABL-positive chronic myelogenous leukemia (CML). In contrast, secondary BCR-ABL-negative MPNs have only been occasionally observed. In a survey among 29,356 testicular cancer survivors, 621 secondary cancers were found, and the only MPNs were four cases of CML. Similarly, in another study, a review of 463,618 cases of cancer patients treated with chemotherapy and radiotherapy, revealed 233 patients with hematological malignancies including AML/MDS (n=741), CML (n=178), chronic lymphocytic leukemia (CLL) (n=253) and acute lymphoblastic leukemia (n=61). Notably, no cases of BCR-ABL-negative MPNs were identified.

Herein, we report a patient who developed a secondary FIP1L1-PDGFRα-positive CEL, 27 months after radiotherapy for testicular seminoma and 20 years of occupational exposure at a power plant station. This case is of clinical interest in several aspects. Firstly, although more than a hundred cases of FIP1L1-PDGFRα-positive MPNs have been reported so far, only two of them were secondary to chemotherapy for a primary cancer. In the first case, FIP1L1-PDGFRα fusion gene was detected 11 years after 3 courses of combination chemotherapy because of non-Hodgkin’s lymphoma and 3 years after therapy with cyclophosphamide for hypereosinophilic syndrome. In the second case, the molecular abnormality was found 1 year after the application of several courses of multiagent chemotherapy for Langerhans cell histiocytosis. No radiotherapy-related FIP1L1-PDGFRα-positive MPN has been reported so far. Secondly, our patient has two radiation exposure events - occupational and therapeutic, raising the question if one or both of them might be the causative factor. Though it is well known that abdominal and pelvic irradiation may produce eosinophilia, two facts are against the hypothesis that radiotherapy has a primary role in the secondary malignancy pathogenesis. In the reported case, the diagnosis of CEL was made only 27 months after radiotherapy and this period was significantly shorter than the 5-9 years or longer, reported in the literature, though secondary myeloid malignancies have been reported after similar or even shorter latency period. Besides, our patient received a high dose of 366 Gy, though for a larger volume including parts of the abdomen and pelvis and the radiation-related leukemia risk has proven to be considerably lower at high doses due to cell killing.

On the other hand, the patient’s long-term occupational radiation exposure may not be the only culprit as many studies have failed to prove greater risk for hematological neoplasms among nuclear power plant workers compared with national rates. Similarly, no strong evidence for increased risk of leukemia, or other malignant disease was detected 11 years after 3 courses of combination chemotherapy because of non-Hodgkin’s lymphoma and 3 years after therapy with cyclophosphamide for hypereosinophilic syndrome. In the second case, the molecular abnormality was found 1 year after the application of several courses of multiagent chemotherapy for Langerhans cell histiocytosis. No radiotherapy-related FIP1L1-PDGFRα-positive MPN has been reported so far. Secondly, our patient has two radiation exposure events - occupational and therapeutic, raising the question if one or both of them might be the causative factor. Though it is well known that abdominal and pelvic irradiation may produce eosinophilia, two facts are against the hypothesis that radiotherapy has a primary role in the secondary malignancy pathogenesis. In the reported case, the diagnosis of CEL was made only 27 months after radiotherapy and this period was significantly shorter than the 5-9 years or longer, reported in the literature, though secondary myeloid malignancies have been reported after similar or even shorter latency period. Besides, our patient received a high dose of 366 Gy, though for a larger volume including parts of the abdomen and pelvis and the radiation-related leukemia risk has proven to be considerably lower at high doses due to cell killing. On the other hand, the patient’s long-term occupational radiation exposure may not be the only culprit as many studies have failed to prove greater risk for hematological neoplasms among nuclear power plant workers compared with national rates. Similarly, no strong evidence for increased risk of leukemia, or other malignant disease was
found among adult populations with low dose-rate fractioned exposures after the Chernobyl accident. However, the understanding of radiation biology has undergone a fundamental shift in paradigms away from deterministic hit-effect relationships and towards complex ongoing cellular responses such as increased genomic instability, and decreased adaptive responses observed at very low doses, and particularly relevant when exposure is spread over a period of time. These are circumstances that are important to understanding cancer risk associated with occupational radiation exposures due to increased sensitivity to physical, radiation and chemical agents and modified biological effects. Therefore, in accordance with the hypothesis of two/multiple-hits carcinogenesis, it seems more probable that in our patient, the cumulative dose or sequential exposure to both occupational and therapeutic radiation have contributed to the development of a FIP1L1-PDGFRα-positive neoplasm. However, a question still remains why no other similar cases have been reported so far in thousands of occupational radiation workers as well as in patients who have undergone radiation therapy.

Several factors might be related to the absence of reported cases of FIP1L1-PDGFRα(+) CEL. Firstly, CEL in general is a very rare disease. Besides, FIP1L1-PDGFRα fusion gene has been discovered not long ago and cannot be detected by conventional cytogenetics, therefore, the respective abnormality had not been tested routinely in a significant proportion of patients and the true incidence of FIP1L1-PDGFRα-positive cases is still unknown. To our knowledge there is at least one case of CEL, developed 6 years after radiotherapy for thyroid cancer, reported so far. Interestingly, this patient was positive for t(6;11)(q27;q23)/MLL-AF6, which is a typical molecular marker of AML, but at that time FIP1L1-PDGFRα-rearrangement was not tested. On the other hand, the risk of development of secondary malignancies after radiation exposure depends on a number of factors - cumulative radiation dose received, the individual genetic background, exposure to additional mutagenic factors, etc. that might vary from case to case.

In conclusion, this case contributes to the clinical heterogeneity of a rare entity such as FIP1L1-PDGFRα – positive myeloproliferative neoplasms, and for the possible role of occupational and therapeutic radiation, raising the question if one or both of them might be the causative factor.

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