CASE REPORT

Adult T-cell Leukemia/Lymphoma as a Methotrexate-associated Lymphoproliferative Disorder in a Patient with Rheumatoid Arthritis

Ichiro Takajo¹,², Kunihiko Umekita¹, Yoshihiko Ikee¹, Koichi Oshima³ and Akihiko Okayama¹

Abstract: The patient was a 74-year-old Japanese woman with rheumatoid arthritis (RA) who developed generalized lymphadenopathy with elevated levels of lactase dehydrogenase (LD), and soluble IL-2 receptor (sIL-2R). She was found to be positive for anti-human T-cell leukemia virus type 1 (HTLV-1) antibodies. Her symptoms and laboratory abnormalities spontaneously regressed after the cessation of methotrexate (MTX), suggesting that she had an MTX-associated lymphoproliferative disorder; however, her lymphadenopathy appeared again approximately 14 months later with LD and sIL-2R elevation. A histopathological analysis and Southern blotting of a lymph node biopsy specimen for HTLV-1 provirus supported the diagnosis of adult T-cell leukemia/lymphoma (ATL) (lymphoma type). These data confirmed that an HTLV-1 positive RA patient may develop ATL.

Key words: rheumatoid arthritis, human T-lymphotropic virus type 1, adult T-cell leukemia/lymphoma, methotrexate-associated lymphoproliferative disorder


Introduction

Patients with rheumatoid arthritis (RA) may present with a wide spectrum of lymphoid proliferation and lymphomas, which together are named lymphoproliferative disorders (LPD). The standardized incidence ratio (SIR) for lymphoma in Japanese RA patients is reported to be significantly high (1, 2). B-cell phenotype has been reported to occur in as much as 80% of LPDs (3). The most common histology of methotrexate (MTX)-LPD found in Japanese RA patients was diffuse large B-cell lymphoma; its frequency was as high as 60% of reported cases (3). The risk of developing LPD was reported to be increased when RA patients were treated with MTX (4). Epstein-Barr virus (EBV) infection has been considered to be involved in the pathogenesis of LPD. A systematic review showed that EBV-induced LPD in RA patients is commonly seen in Asian individuals (5). EBV was detected in 60% of Japanese RA patients with MTX-associated LPD (6). Discontinuation of MTX treatment may result in the spontaneous regression of MTX-associated LPD (7, 8). Thus, immunosuppression caused by MTX has been suspected to be associated with the development of EBV-related B-cell LPD in RA patients.

Human T-cell leukemia virus type 1 (HTLV-1) is the causative agent of the T-cell malignancy adult T-cell leukemia/lymphoma (ATL) (9). The acute or lymphoma types of ATL are aggressive diseases and the prognosis has been reported to be poor, even when they were treated with advanced chemotherapy (10). HTLV-1 is endemic in Japan, where there are an estimated 800,000 HTLV-1 carriers (11). Because women have a higher prevalence of HTLV-1 infection (11), it is reasonable to assume that a relatively high number of Japanese RA patients are infected with HTLV-1. Thus far, only a small number of patients with MTX-LPD have been reported to have the T-cell phenotype and no cases of ATL have been reported as MTX-LPD (3, 8). Thus, whether or not HTLV-1-positive RA patients are at greater risk of developing ATL needs to be investigated.

1Department of Rheumatology, Infectious Diseases and Laboratory Medicine, Internal Medicine, Faculty of Medicine, University of Miyazaki, Japan, ²Youkikai Ikee Hospital, Japan and ³Department of Pathology, Kurume University, Japan

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Correspondence to Dr. Kunihiko Umekita, kunihiko_umekita@med.miyazaki-u.ac.jp
risk of ATL - especially when they receive anti-rheumatic treatment including MTX - is an important question.

We report the case of an RA patient, who developed generalized lymphadenopathy that showed spontaneous regression after the withdrawal of MTX, suggesting that she had an MTX-associated LPD. She was found to be positive for HTLV-I and developed ATL (lymphoma type) 14 months later. It was not clear whether the first episode of MTX-associated LPD occurred due to ATL because no biopsy was performed at the time; however, this case raised the question as to whether T-cell LPD can occur in HTLV-I-positive RA patients who are treated with MTX.

Case Report

A 74-year-old Japanese woman presented to our hospital with poly-arthritis. She tested positive for anti-citrullinated protein antibodies and was diagnosed with seropositive RA based on the 2010 American College of Rheumatology/European League Against Rheumatism criteria. Her disease activity score 28 joints-C-reactive protein (DAS28-CRP) was 4.58, suggesting moderate disease activity. We treated her moderate RA with MTX (4 mg per week) and prednisolone (PSL; 5 mg per day).

Her condition remained stable for 5 years; however, she complained of anorexia and lymph node enlargement was recognized in the left cervical region. Her serum levels of lactase dehydrogenase (LD, 433 IU/L) and CRP (2.14 mg/dL) were elevated. The serum level of soluble IL-2 receptor (sIL-2R) was markedly elevated (34,881 U/mL) and a small number of atypical lymphocytes were present in the peripheral blood (Fig. 1). Although typical ATL cells were not found, HTLV-I antibodies were detected. Computed tomography revealed generalized lymphadenopathy including the mediastinum and abdomen (Fig. 2). MTX treatment was immediately discontinued, and her lymph node enlargement improved. Her serum LD and sIL-2R levels were found to have decreased to normal levels (161 IU/L and 663 U/mL, respectively) at four months after MTX withdrawal. Because MTX withdrawal induced spontaneous regression, she was diagnosed with MTX-associated LPD; however, lymph node biopsy was not performed. Thereafter, her RA was well controlled with a low dose of PSL (3 mg/day).

At fourteen months after the initial event of LPD, she again complained of anorexia and exhibited lymph node enlargement in the left cervical region. On CT, generalised lymphadenopathy, which included the mediastinum and abdomen, was found to be worse than that observed during the first episode of LPD (data not shown). The patient’s serum LD (729 IU/L) and sIL-2R levels (73,208 U/mL) were markedly elevated. The peripheral blood again showed a small number of atypical lymphocytes without typical ATL cells. We performed a cervical lymph node biopsy. A pathological analysis revealed the diffuse proliferation of medium-sized abnormal lymphocytes, suggesting malignant lymphoma (Fig. 3). On immunostaining, the surface marker profile of the lymphocytes with atypia in the lymph node specimen was CD20(-), CD3(+), CD4(+), CD8(-), and CD30(-). The Epstein-Barr encoding region was negative upon in situ hybridization; this was compatible with a T-cell phenotype.
without EBV infection. Southern blotting of the lymph node specimen for HTLV-1 provirus showed a monoclonal proliferation of HTLV-1-infected cells (Fig. 4). She was referred to a hematologist and was diagnosed with ATL (lymphoma type) based on Shimoyama’s criteria (12). Her general condition did not allow us to perform chemotherapy for ATL and she died within a month.

Discussion

It has been realized that RA patients have a higher risk of developing Hodgkin lymphoma and non-Hodgkin lymphoma (13). In addition, immunosuppressive treatments, including MTX, have been reported to increase the risk of LPD. Spontaneous regression after the withdrawal of MTX treatment has been reported, especially in RA patients with EBV-associated LPD (3, 4, 8). Thus, the reactivation of EBV in RA patients with LPD may be induced by immunosuppressive treatment, including MTX (8).

In contrast, ATL is a malignancy of mature T-cells caused by HTLV-1 infection (9). The incidence of ATL among HTLV-1 carriers has been reported to be one in 1,000 person-years (10). There have been several case reports of ATL in patients with rheumatic diseases. Nakamura et al. reported a case of ATL developing in an RA patient who was treated with MTX and tocilizumab for 7 years (14). The complete remission of ATL was achieved with a standard therapeutic regimen, which suggested a preferable clinical course. Moreover, a case was reported in which a spondyloarthritis patient treated with the anti-TNF adalimumab developed ATL (15). This patient was reported to have had a history of oral immunosuppressant treatment; however, it was not clear whether MTX was prescribed. We recently experienced a case in which ATL (chronic type) developed in an HTLV-1 positive RA patient (16). The patient was treated with MTX and infliximab when she developed ATL. After the discontinuation of MTX and infliximab, her ATL showed spontaneous remission. These cases raised the question as to whether ATL occurs as T-cell LPD in HTLV-1-positive patients with rheumatic diseases-especially RA—when they are treated with immunosuppressive agents, including MTX and/or biologic agents.

In the present study, we experienced a case of MTX-LPD followed by the occurrence of overt ATL in an HTLV-1-positive RA patient. The question was whether or not the first episode of LPD in this case was ATL. The first episode might have been EBV-associated B-cell MTX-LPD, because this type of LPD is known to frequently regress upon the withdrawal of MTX (8). If this was the case, two different types of LPD occurred in the same patient within a rela-
The abnormal lymphocytes were CD3 (+) CD4 (+) CD8 (+) CD20 (-) CD30 (-) and EBER (-). The histopathological diagnosis was compatible with adult T-cell leukemia lymphoma. H&E: Hematoxylin and Eosin staining, CD: cluster of differentiation, EBER: Epstein-Barr encoding region.

Figure 4. Southern blotting of the lymph node biopsy specimen to detect HTLV-1 proviral DNA. Southern blotting revealed the monoclonal integration of HTLV-1 proviral DNA (Special Reference Laboratories). 1, positive control; 2, negative control; 3, patient specimen; M, size marker (λDNA/HindIII). Restriction enzyme (E: Eco RI, P: Pst I).

relatively short period. However, the median sIL-2R level in patients with diffuse large B-cell lymphoma is reported to be 1,473 U/mL, while that of ATL (lymphoma type) is reported to be as high as 34,620 U/mL (17, 18). The marked elevation of sIL-2R in the present case not only in the second episode (ATL), but also in the first episode of LPD, supported the possibility that the first episode was also ATL.

A high HTLV-1 proviral DNA load and the clonal expansion of HTLV-1-infected cells in asymptomatic HTLV-1 carriers have been considered major risk factors for the development ATL (19, 20). The incidence of ATL has been reported to increase upon immunosuppressive treatment. HTLV-1-positive patients who received liver transplantation followed by immunosuppressive treatment were reported to have a high risk of developing ATL (21). A neurological disease, HTLV-1-associated myelopathy, was also reported to have developed in a patient undergoing living-donor liver transplantation (22). It is possible that the active expansion of HTLV-1-infected T-cells may occur in HTLV-1-positive patients who are treated with immunosuppressive agents. If the first episode of the present case was less aggressive ATL, it is possible that treatment with MTX caused immunosuppression and was involved in the development of ATL. The withdrawal of MTX may have induced the recovery of the immune system, resulting in the spontaneous re-
gression of ATL; however, the patient in the present case developed aggressive ATL within one year. A similar situation, in which aggressive lymphoma develops after the initial regression of LPD, is commonly observed in patients with EBV-associated MTX-LPD (3).

One of the limitations of the present report is that no biopsy was performed during the first episode of lymphadenopathy. Thus, the nature of MTX-LPD, including the pathology, surface phenotype, and association with EBV or HTLV-1, was not determined in the first episode of MTX-LPD.

In conclusion, the present report confirmed that an HTLV-1-positive RA patient may develop ATL. In addition, the current case indicated the possibility that less aggressive ATL occurs as MTX-LPD, which only regresses upon the withdrawal of MTX. If this is true, then HTLV-1-positive RA patients treated with MTX should be carefully followed to allow for the early diagnosis of ATL. Further studies, with large numbers of HTLV-1-positive RA patients, will be necessary to resolve the clinical question as to whether HTLV-1 infection causes ATL as a T-cell LPD in RA patients and whether MTX treatment is a risk factor for ATL in such patients.

Written informed consent for the publication of this report was obtained from the patient’s son by the author.

The authors state that they have no Conflict of Interest (COI).

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