Fingolimod-associated PML with mild IRIS in MS
A clinicopathologic study

ABSTRACT

Objective: To clarify the clinical, neuropathologic, and virologic characteristics of progressive multifocal leukoencephalopathy (PML) and its immune reconstitution inflammatory syndrome (IRIS) in a patient with fingolimod-treated MS.

Methods: A case study.

Results: A 34-year-old patient with MS using fingolimod for 4 years had a gradual progression of right hemiparesis and aphasia with a new subcortical white matter lesion in the precentral gyrus by initial MRI. Blood tests were normal, except for lymphopenia (160 cells/μL). One month after the cessation of fingolimod, brain MRI depicted a diffusely exacerbated hyperintensity on fluid-attenuated inversion recovery and diffusion-weighed imaging in the white matter with punctate gadolinium enhancement, suggesting PML-IRIS. A very low level of JC virus (JCV)-DNA (15 copies/mL) was detected in the CSF as judged by quantitative PCR. Brain tissues were biopsied from the left frontal lesion, which showed some small demyelinated foci with predominant loss of myelin-associated glycoprotein with infiltrations of lymphocytes and macrophages, but clear viral inclusion was not observed with hematoxylin-eosin staining. JCV-DNA was uniquely detectable in an active inflammatory demyelinating lesion by in situ hybridization, possibly suggesting an early phase of PML. DNA extracted from the brain sample was positive for JCV-DNA (151 copies/cell). It took 3 months to normalize the blood lymphocyte count. The patient was treated with 1 g of IV methylprednisolone for 3 days and a weekly oral dose (375 mg) of mefloquine, and her symptoms gradually improved.

Conclusion: Low CSF JCV-DNA and unfound viral inclusions initially made her diagnosis difficult. The clinical course of fingolimod-associated PML may be associated with mild immune reconstitution.

GLOSSARY

DMT = disease-modifying treatment; DWI = diffusion-weighted imaging; FLAIR = fluid-attenuated inversion recovery; IRIS = immune reconstitution inflammatory syndrome; JCV = JC virus; MAG = myelin-associated glycoprotein; MBP = myelin basic protein; PML = progressive multifocal leukoencephalopathy; RRMS = relapsing-remitting MS.

Fingolimod is an oral disease–modifying drug approved for relapsing-remitting MS (RRMS), which downregulates the sphingosine-1-phosphate receptor and subsequently prevents lymphocyte egress from lymph nodes.¹ Until now, 13 cases of fingolimod-associated progressive multifocal leukoencephalopathy (PML) including this case have been reported in patients with RRMS (personal oral communication, Mitsubishi Tanabe Pharma, 2017);²,³ however, there has been no report of immune reconstitution inflammatory syndrome (IRIS). We herein report a case of fingolimod-associated PML accompanied by IRIS with favorable clinical course, which showed early pathologic changes of PML-IRIS in a biopsied specimen.

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Funding information and disclosures are provided at the end of the article. Go to Neurology.org/nn for full disclosure forms. The Article Processing Charge was funded by JSPS KAKENHI.

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CASE A 34-year-old woman was diagnosed as having RRMS with typical MRI lesions, the so-called Dawson fingers, when she had experienced the initial mild symptoms of paresthesia and dysphagia at the age of 14 years. During 11 years treated with interferon-β1b, she had 12 MS relapses. She had no history of immunosuppressive treatment. At the age of 30 with an Expanded Disability Status Scale score of 6.0, her treatment was changed to fingolimod, and she had no further relapse and disability worsening, although the JC virus (JCV)-antibody was positive (Index 1.13).

Early in January, she noticed gradually aggravating muscle weakness of the right hand, dysarthria, and hemiparetic gait. She visited our clinic by late-March, and she discontinued fingolimod treatment because of lymphopenia (160 cells/μL).

She was admitted to our hospital by mid-April. Initial brain MRI on admission showed fluid-attenuated inversion recovery (FLAIR)-high signal lesions sharply delineated toward the cortex and consecutive deep white matter with hyperintense diffusion-weighted imaging (DWI) signals in the left frontoparietal region (figure 1, A and B). There was no mass effect and no gadolinium enhancement in any lesions (figure 1C). All blood tests including biochemistry and blood cell counts were normal, except for lymphopenia (442 cells/μL). CSF examination showed no cells, elevated protein concentration (81 mg/dL), elevated IgG index (0.89), and positive oligoclonal IgG bands.

One month after admission with gradually worsening aphasia and right hemiplegia, secondary brain MRI revealed several enlarged lesions (figure 1, D and E) with gadolinium enhancement in T1-weighted hypointensity (figure 1F), showing a punctate pattern. A very low level of CSF-JCV-DNA was detected in the CSF by quantitative PCR (15 copies/mL) when examined at the National Institute of Infectious Diseases (Tokyo, Japan). In mid-June, the patient was treated with 1 g of IV methylprednisolone for 3 days and oral mefloquine (375 mg/wk) for 6 months. Two months after the treatment, CSF-JCV became negative, and the blood lymphocyte count was normalized. Her aphasia and right hemiplegia gradually improved.

RESULTS OF PATHOLOGICAL STUDY In late-May, stereotactic needle brain biopsy was performed after written informed consent was obtained. Seven tissue fragments (1 mm in width and 2–5 mm in length) were obtained from the left frontal lobe (figure 2A). With hematoxylin-eosin staining, proliferation of small round cells was apparent, including glial cells and inflammatory components such as lymphocytes, plasma cells, and macrophages. Typical JCV-infected oligodendroglia presenting viral inclusions in the markedly enlarged nuclei were hardly seen. In all fragments of biopsied tissue, CD68-positive cells were scattered, and there were some aggregations of CD68-positive cells (figure 2B, circled). In these lesions, there were multifocal subacute (figure 2, C–E) and acute (figure 2, F–L) demyelinating oval lesions with loss of myelin-associated glycoprotein (MAG) (figure 2, C and G). In subacute lesions without viral inclusion (figure 2, C–E), the immunoreactivity against myelin basic protein (MBP) was partially lost (figure 2D), whereas CD8-positive lymphocytes were observed only at the periphery (figure 2E), suggesting PML-like MAG-dominant demyelinated lesions.

In acute demyelinating lesions, oligodendroglialike cells with medium-sized nuclei were present with clustered inflammatory cells (red yellow arrow in figure 2F). Their nuclei were smaller than typical JCV-infected oligodendroglia, known as a diagnostic hallmark of PML, but mildly enlarged over lymphocytic nuclei. These cells were negative for glial fibrillary acidic protein or lymphocytic markers, and many of them showed intranuclear structures in dots, suggesting JCV-infected oligodendroglia in the early stage of viral infection. Moreover, a cell with relatively large irregular nucleus resembling a JCV-infected astrocyte was also present (yellow arrow in figure 2F). This lesion was in the active demyelinating phase with loss of MAG but completely preserved MBP infiltrated with CD3+, CD8+, and CD79a-
Figure 2  Neuropathology of PML lesions in a patient with MS treated with fingolimod

(A) Fragments of biopsied brain tissues (hematoxylin-eosin; HE staining). Affected lesions were obscure with HE staining. A and B are serial sections (scale bar: 500 mm). (B) Although demyelinated lesions were not clear with Klüver-Barrera staining, there were 4 demyelinating areas with high density of CD68⁺ cells lacking myelin-associated glycoprotein (MAG) or myelin basic protein (MBP). Two of them were relatively in subacute (C–E) and acute phases (F–L). (C) In a typical oval demyelinating lesion with massive macrophage infiltration, there was a lesion with dominant loss of MAG compared with partially lacked MBP (D) with CD8⁺ positive cells at the periphery (E), suggesting a subacute demyelinated lesion with distal oligodendroglialopathy. C–E is a serial section (scale bar: 200 μm). (F) In another active inflammatory lesion (HE staining), there were massive infiltrations of macrophages and lymphocytes diffusely in demyelinating lesions. In these lesions, oligodendroglial-like cells with mildly enlarged nuclei were present (green arrows). An astroglia-like cell suspicious for JC virus (JCV) infection was also seen (yellow arrow). Within serial sections of F, loss of MAG was present (G) with completely preservation of MBP (H), suggesting in acute phase. F–H is a serial section (scale bar: 200 μm). (I–L) Properties of inflammatory cells (I: CD3, J: CD8, and K: CD79a). There were numerous CD3⁺ T cells and CD79a⁺ B cells and plasma cells, and both helper and cytotoxic T cells were present in an almost equal ratio, where JCV-infected cells were observed (L: in situ hybridization (ISH) VP1). I–L is a serial section (scale bar: 50 μm). PML = progressive multifocal leukoencephalopathy.
positive cells (figure 2, G–I), suggesting a very active inflammatory lesion, where several JCV-infected cells were uniquely observed (figure 2L).

Oligodendroglia-like cells with dot-shaped inclusions were seen in the lesions (figure 3, A and B), reflecting JCV proliferation in promyelocytic leukemia nuclear bodies.\(^5\) Immunohistological examination of JCV capsid proteins or agnoprotein revealed only a few cells (figure 3C). In situ hybridization for JCV genomic DNA detected at least 7 JCV-positive cells (figure 3D) with definite high-intense nuclear signals, confirming JCV infection. The distribution of JCV-positive cells was associated with active inflammatory lesions showing the infiltration of helper and cytotoxic T cells, suggesting a mild host inflammatory response against JCV.

The quantitative real-time PCR examination targeting the JCV large T gene was repeatedly performed in CSF (figure 3E) and brain samples (figure 3F).\(^6\) The PCR testing of brain tissue revealed the presence of JCV-DNA (151 copies/cell) (figure 3F). These results indicated that the biopsied brain lesions represented PML pathology, but not MS. The viral titer was markedly low, and nuclear features of oligodendroglia indicated an early stage of JCV infection. Small inflammatory foci may suggest mild host responses against JCV infection, consistent with fingolimod-associated PML-IRIS.

**DISCUSSION** The clinical and pathologic diagnosis of PML-IRIS is challenging, particularly when the consequence of JCV reactivation and immune reconstitution is not clear.\(^3\) PML is essentially rare in patients with fingolimod-associated MS without other previous immunosuppressive treatments. The low CSF viral load and the presence of markedly few JCV-positive cells have raised the matter of difficulty in PML diagnosis in the present study. Clinical worsening was accelerated after the cessation of fingolimod with punctate gadolinium enhancement in MRI, suggesting PML-IRIS. Pathologic examination of biopsied brain tissue revealed some JCV-DNA–positive cells only in an active demyelinating lesion dominantly lacking MAG\(^7\) and JCV-infected glial cells with dot-shaped

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Figure 3 Detection of JC virus (JCV) genomic DNA in mildly enlarged nuclei of oligodendroglia-like cells

(A and B) Oligodendroglia-like cells with dot-shaped inclusions (hematoxylin-eosin). Dot-shaped inclusions were seen at the inner nuclear periphery in A (arrows), and large and small punctuated structures were present in B (arrows). (C) Immunohistochemistry against JCV capsid protein (VP2/VP3C antibody). There was only one cell suspicious for immunoreactivity against JCV capsid protein, but apparently positive cells were hardly detectable for agnoprotein. (D) In situ hybridization (ISH) for JCV genomic DNA. Sensitive ISH detected 7 oligodendroglia-like cells with JCV genomic DNA in the mildly enlarged nuclei. A–D is a serial section (scale bar: 10 \(\mu\)m). (E and F) PCR detection of JCV genomic DNA. DNA was extracted from either CSF specimen (E) or biopsied brain tissues (F), and the JCV-DNA copy number in each sample was determined by a quantitative PCR targeting the JCV large T gene. NC = negative control (no PCR template); PC = Positive control (approximately 100 copies/reaction). PCR were performed in duplicate (green arrows in E and red arrows in F).
inclusions surrounded by inflammatory cells, suggesting an early phase of PML in association with immune reconstitution. The pathologic features in fingolimod-associated PML in MS may include only early and weak characteristic features with immunoreaction, differing from previously reported PML by other disease-modifying treatments (DMTs) such as natalizumab, an anti-α4-integrin monoclonal antibody.

The rarity of fingolimod-associated PML-IRIS may be related to its different mechanisms compared with other DMT. In natalizumab, most PML cases may be related to its different mechanisms compared to fingolimod-associated PML-IRIS. The kinetics of immune reconstitution in patients with MS was varied from half to several months, which might have impacts on the occurrence of PML-IRIS. Other possible reasons for rarity in fingolimod-associated PML-IRIS may be the lack of pathologic studies and the relatively mild clinical symptom, causing underestimation.

Because of weak pathologic changes in brain tissue and an extremely low JCV-DNA level in CSF, MRI was the most useful and sensitive technique used to detect PML and PML-IRIS in the present case, showing DWI-high and FLAIR-high signals like previous PML cases.

AUTHOR CONTRIBUTIONS
Shuhei Nishiyama: study conception; acquisition, analysis, and interpretation of data for the work; and writing of the manuscript. Tatsuro Misu: substantial contribution to the study conception and design of the work; writing and revising manuscripts; and supervision of the acquisition, analysis, and interpretation of data for the work. Yukiko Shishido-Hara: substantial contribution to the design of the work; writing of the manuscript; and supervision of the acquisition, analysis, and interpretation of data for the work. Kazuo Nakamichi: study conception and design of the work; supervision of the acquisition, analysis, and interpretation of data for the work. Masayuki Sajo: supervision of the acquisition and interpretation of data for the work. Yoshiki Takai: analysis, acquisition, and interpretation of data for the work. Kentarou Takei: acquisition, analysis, and interpretation of data for the work. Naoki Yamamoto: acquisition and interpretation of data for the work. Teiji Tominaga, Ichiro Nakashima, Kazuo Fujihara, Hiroshi Kuroda, and Ryuta Saito: acquisition and interpretation of data for the work. Yoshiki Takai: analysis, acquisition, and interpretation of data for the work. Masaki Aoki: substantial contribution to the design of the work; writing of the manuscript; and supervision of the acquisition, analysis, and interpretation of data for the work. Kentarou Takei and Naomi Yamamoto report no disclosures. Hiroshi Kuroda, Ryuta Saito, Mika Watanabe, and Teiji Tominaga have received Grants-in-Aid for Scientific Research. Ichiro Nakashima has received Grants-in-Aid for Scientific Research; has received funding for travel; received speaker honoraria from Tanabe Mitsubishi Pharma Corporation; has received research funding from LSI Medience Corporation; and is an editorial board for Multiple Sclerosis International. Kazuo Fujihara has received Grants-in-Aid for Scientific Research; serves on the editorial board for Clinical and Experimental Neuroimmunology (2009 till present) and an advisory board member of Sri Lanka Journal of Neurology; and has received research support from Bayer Schering Pharma, Biogen Idec, Mitsubishi Tanabe Pharma Corporation, Novartis Pharma, Chugai Pharmaceutical, Ono Pharmaceutical, Nihon Pharmaceutical, Merck Serono, Alexion Pharmaceuticals, MedImmune, and Medical Review; has received funding for travel and speaker honoraria from Bayer Schering Pharma, Biogen Idec, Eisai Inc., Mitsubishi Tanabe Pharma Corporation, Novartis Pharma, Astellas Pharma Inc., Takeda Pharmaceutical Company Limited, Asahi Kasei Medical Co., Daiichi Sankyo, and Nihon Pharmaceutical; serves as an editorial board member of Clinical and Experimental Neuroimmunology (2000 till present) and an advisory board member of Sri Lanka Journal of Neurology; and has received research support from Bayer Schering Pharma, Biogen Idec Japan, Asahi Kasei Medical, The Chemo-Sero-Therapeutic Research Institute, Teva Pharmaceutical, Mitsubishi Tanabe Pharma, Teijin Pharma, Chugai Pharmaceutical, Ono Pharmaceutical, Nihon Pharmaceutical, and Genzyme Japan. Masashi Aoki has received Grants-in-Aid for Scientific Research. Go to Neurology.org/nn for full disclosure forms.

Received July 25, 2017. Accepted in final form September 26, 2017.

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Fingolimod-associated PML with mild IRIS in MS: A clinicopathologic study
Shuhei Nishiyama, Tatsuro Misu, Yukiko Shishido-Hara, et al.

*Neurol Neuroimmunol Neuroinflamm* 2018;5;
DOI 10.1212/NXI.0000000000000415

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