Description
This educational activity is designed to enhance awareness of the risk of progressive multifocal leukoencephalopathy (PML) as a complication of autoimmune and inflammatory disease and treatments and to provide vital information to clinicians regarding its prompt recognition and diagnosis.

Objectives
Upon completing this activity, participants will be able to:
• Describe the epidemiology of PML in the setting of autoimmune and inflammatory disease and the risks associated with individual biologic therapies
• Explain the microbiology of JC virus and the pathogenesis of PML
• Recognize the early signs and symptoms of PML and conduct appropriate diagnostic evaluations
• Engage patients in meaningful shared and informed decision-making that considers PML when choosing immunosuppressive regimens for treatment of their disease.

Target Audience
This activity is intended for rheumatologists, oncologists, neurologists, infectious disease specialists, and other interested health care professionals.

Accreditation
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Disclosure
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Leonard Calabrese, DO, reported consulting relationships with Abbott Laboratories, Amgen Inc., Centocor, Elan Corporation, Genentech, Roche, and Wyeth; and teaching and speaking relationships with Abbott Laboratories, Amgen Inc., Centocor, and Genentech.

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Charles L. Bennett, MD, PhD, MPP
Joseph R. Berger, MD
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PROGRESSIVE MULTIFOCAL LEUKOENCEPHALOPATHY IN THE BIOLOGIC ERA: IMPLICATIONS FOR PRACTICE

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This supplement was developed from the audio transcript of a roundtable discussion convened at Cleveland Clinic on January 31, 2011. The transcript was formatted and edited by the Cleveland Clinic Journal of Medicine staff for clarity and conciseness, and was then reviewed, revised, and approved by each of the authors.
The epidemiology of progressive multifocal leukoencephalopathy (PML) has evolved in recent years. Until the availability of natalizumab, PML was seen primarily by physicians who treated patients with human immunodeficiency virus (HIV) infection. This situation changed in 2005 when PML was first associated with natalizumab therapy in patients with multiple sclerosis (MS). This discovery focused attention and investigative resources on the relationship between PML and natalizumab, and subsequently on other biologically based therapies. In the years since, we have learned about the pathogenesis of PML and developed concepts of risk mitigation. Increasingly selective, potent, and innovative biologically based immune-based therapeutics have led to enhanced therapeutic options, but also to more surprises, with PML being observed in unexpected patient populations. The MS community and clinicians who routinely use biologic therapies are alert to this disease and its implications for patients. PML remains rare enough, however, that community-based clinicians may be less attuned to the impact and risks of these therapies.

This Cleveland Clinic Journal of Medicine supplement addresses the issues of awareness, recognition, and management of PML for clinicians whose patients may be at risk, including those in the fields of infectious disease, neurology, oncology, and rheumatology. The supplement provides an overview of the pathogenesis and clinical picture of PML, its evolving epidemiology, and the current approaches to its management.

The articles and their accompanying discussions are based on a roundtable held at Cleveland Clinic on January 31, 2011. The roundtable’s expert faculty contributed insights from several different perspectives, including laboratory research on JC virus–induced demyelinization in PML; front-line experience starting in the early 1980s when PML was understood to be an HIV-related disease and continuing to its current status as a potential complication of biologic therapy; and management of a large pharmaceutical safety program that monitors and facilitates reporting of potential fatal drug side effects.

Readers of these articles will acquire understanding of the history and clinical picture of PML, appreciation of the influence of biologic therapies in several specialties, and enhanced awareness of when to consider PML and what actions to take when the diagnosis is a possibility.

Leonard Calabrese, DO
Supplement Editor
History and current concepts in the pathogenesis of PML

ABSTRACT

The JC virus (JCV), first described in 1971, is responsible for initiation of progressive multifocal leukoencephalopathy (PML), a disease characterized by demyelinating plaques and a classic triad of symptoms consisting of cognitive impairment, visual deficits, and motor dysfunction. To establish a diagnosis of PML, evidence of the presence of JCV DNA in pathologic tissue is necessary. The host range for productive infection of JCV is controlled by factors in the cell nucleus that bind to the viral promoter, initiating transcription of mRNA for the coordinated synthesis of viral proteins. Oligodendrocytes, astrocytes, and CD34+ and CD19+ cells of the immune system have the necessary binding proteins in sufficient concentration to allow lytic infection to occur. A strong link between JCV infection in cells of the immune system and cells of the nervous system points to the importance of the tissue origin of JCV latency, the bone marrow that harbors CD34+ cells. The emergence of PML in patients treated with natalizumab and other immune-altering agents supports this observation and provides new insights into the pathogenic mechanisms of JCV infection.

The neuropathology of progressive multifocal leukoencephalopathy (PML) was first reported in 1958 following examination of brain tissue from two cases of chronic lymphocytic leukemia and one case of Hodgkin lymphoma. The classic triad of symptoms of PML—cognitive impairment, visual deficits, and motor dysfunction—had been observed previously but had not been formally described. Until PML was discovered in patients with autoimmune diseases treated with biologic therapies that do not directly suppress immunity, PML had been considered a very rare, virus-induced demyelinating disease of the white matter that occurred in immune-compromised patients. The incidence of PML rose sharply in the mid-1980s with the pandemic of human immunodeficiency virus (HIV)-1 infection and continues as an acquired immunodeficiency syndrome–defining illness at a rate of approximately 1% to 3% of HIV-1 seropositive individuals; more recently, it has been seen in approximately 1 in 850 natalizumab-treated individuals who have multiple sclerosis (MS). The incidence of PML in natalizumab-treated MS patients increases with dosing; among those who receive 24 or more doses, the incidence is 1 in 400.

The cause of PML was unknown until 1971, when viral particles were observed by electron microscopy in PML brain lesions and subsequently isolated at the University of Wisconsin, Madison, in cultures of human fetal brain tissue. The designation of JC virus (JCV) was derived from the initials of the patient whose brain tissue was used for culture and isolation. Variants in the noncoding region of the genome were then serially identified as Mad 1, Mad 2, and so on, representing the geographic location, Madison, Wisconsin, where the virus was identified.

The JCV, a polyomavirus, is a nonenveloped DNA virus withicosahedral structure containing double-stranded DNA genomes. The circular genome of JCV contains early and late transcription units, the latter of which encodes three virion structural proteins—Vp1, Vp2, and Vp3. Humans generate antibodies directed against the amino terminal end of Vp1 and perhaps Vp2 and Vp3.

JC VIRUS PATHOGENESIS

JCV pathogenesis is studied in cell cultures derived from human fetal brain tissue. In vitro, JCV robustly infects astrocytes, making it important to identify the culture’s cellular phenotypes. A cell line was developed that allows multiplication of JCV and, more recently, human multipotential progenitor cells were isolated and are being grown from the human developing brain at various gestational stages. The lineage pathways of these cells can be differentiated into astrocytes, oligodendrocytes, and neurons. Initiating infection in progenitor cells with JCV virions made it possible to determine which cells were
susceptible to infection. JCV susceptibility is evident in progenitor-derived astrocytes and glial cells, which reflects the pathologic process in PML brain tissue. Neuronal cells, by contrast, are not susceptible to infection.

**JC VIRUS CHARACTERISTICS: GLOBAL DISTRIBUTION, TRIAD OF SYMPTOMS**

Subcortical multifocal white matter lesions are the classic feature of PML on neuroimaging. Seroepidemiology of JCV has revealed ubiquitous distribution, with 50% to 60% of adults aged 20 to 50 years demonstrating antibody to JCV. The percentage of the population with antibody increases with age, but may vary among geographic regions. Prevalence is lower among remote populations.

Although the initial site of JCV infection is not well characterized, we know that the primary infection is not in the brain. The JCV has a selective tropism for replication in glial cells in the human brain, but the absence of an animal model for PML has hindered our understanding of the JCV migration to the brain and the initiation and development of central nervous system infection.

Although humans carry JCV-specific antibodies, the clinical significance of these antibodies is unknown. Antibody levels rise during active infection, at times to very high titers, but offer no protection. T-cell–mediated immune responses directed to structural and nonstructural proteins are important in controlling infection.

A high index of suspicion for PML is warranted in individuals who demonstrate the classic triad of symptoms (cognitive impairment, visual deficits, and motor dysfunction) and in whom magnetic resonance imaging shows evidence of demyelinated plaque lesions; however, evidence of the presence of JCV DNA in pathologic tissue is necessary to confirm a diagnosis of PML.

The development of an in situ DNA hybridization assay using a biotinylated probe has facilitated identification of JCV DNA in the infected nuclei of the pathologic tissue. The presence of JCV DNA in cerebrospinal fluid (CSF) samples can be detected using a quantitative polymerase chain reaction assay, targeting the viral genome in the amino terminal end of the viral T protein. This T protein coding region was targeted because it does not crossreact, even with other human polyomaviruses, and it is intolerant of mutations. This assay is certified by the Clinical Laboratory Improvement Amendments, licensed by the National Institutes of Health, and it is the most sensitive (to levels of 10 copies/mL sample) assay available.

**JC VIRUS SUSCEPTIBILITY FACTORS**

Despite the high prevalence of JCV infection, PML is rare, suggesting important barriers to its development. Although the receptor for JCV has been identified as alpha 2,6-linked sialic acid, the host range for productive infection is controlled by factors within the cell nucleus that bind to the viral promoter; this process initiates transcription of mRNA for the coordinated synthesis of viral proteins. Only certain cells have the necessary DNA binding proteins in high enough concentrations to allow lytic infection to take place, spreading by cell-to-cell contact. These cells include oligodendrocytes, the primary target for JCV, whose destruction leads to PML; astrocytes; and the CD34+ and CD19+ cells of the immune system. JCV can also be found in urine, at times in very high concentrations. It is present in the uroepithelial cells and multiplies without apparent pathologic consequences. Virus isolated from the urine has not been grown in cell culture systems in the laboratory setting.

Bone marrow CD34+ hematopoietic progenitor cells represent a potential pathway of JCV pathogenesis: in six people with PML, latent JCV DNA was demonstrated in pathologic tissue from lymph, spleen, or bone marrow biopsies taken months to years before the patient developed neurologic disease.

The initial site of JCV infection is not known, but it is believed that the virus infects stromal cells within the tonsils and then disseminates through peripheral blood (Figure 1). The CD34+ hematopoietic stem cells carry the virus in bone marrow, and the uroepithelial cell is the carrier in the kidney. The nucleotide arrangement of the regulatory region that drives infection is markedly different in the kidney than in the bone marrow or brain tissue of patients with PML. JCV isolated from the kidney is referred to as the archetype sequence, which, unlike PML in the brain, has no tandem repeats. No pathology is associated with JCV kidney infection, although about 30% of the population excrete JCV in the urine.

Upon immunosuppression, reactivation of the virus occurs, with evidence of the virus found in CD10 and CD19/20 lymphocytes in the peripheral blood of some individuals. Blood-to-brain viral dissemination results in infection of oligodendrocytes, astrocytes, and progenitor cells.

**Susceptibility is related to nucleotide sequences**

Susceptibility to PML is associated with promoter/enhancer nucleotide sequences. The tandem repeat nucleotide structure has been found in the peripheral blood leukocytes and the CSF of patients with PML. Although the arrangement of nucleotide sequences in the viral regulatory region is highly variable among patients with PML, there are no alterations in the sequence within the origin of DNA replication. These highly conserved sequences contain regions for DNA-binding proteins that drive transcription, initiating the life cycle of the virus.
The nuclear transcription factor NF-1 is a cell-specific regulator of JCV promoter/enhancer activity. In humans, the NF-1 family of DNA-binding proteins is encoded by four discrete genes, one of which is NF-1 class X (NF-1X), a critical transcription factor that affects JCV cells. The human brain makes NF-1X in concentrations greater than the concentrations of other NF-1 transcription family members of DNA-binding proteins. NF-1X is located adjacent to and interacts with another family of transcription factors, activator protein-1, which has also been associated with JC viral activity.

Another transcription factor, Spi-B, binds to sequences present in the JCV promoter/enhancer. Spi-B is a regulator of JCV gene expression in susceptible cells and appears to play an important role in JCV activity. The expression of Spi-B is upregulated in patients with MS who are treated with the monoclonal antibody natalizumab, a population of patients in whom PML has been recently described.11–15

Natalizumab binds to the alpha-4 integrin molecule, preventing hematopoietic stem cells and developing B cells from attaching to a vascular-cell adhesion molecule and forcing them to migrate from the bone marrow (Figure 2).16 An ideal environment is created for JCV when the natalizumab-induced increase in CD34+ cells in the circulation is combined with upregulation of gene cells involved in B-cell maturation. JCV can reside in the bone marrow in a latent state and can use B cells and their DNA-binding proteins to initiate viral multiplication, eventually gaining entry into the brain to cause PML.

In addition to natalizumab, PML has been described in patients treated with efalizumab, another biologic agent that binds alpha-4 integrin molecules on the surface of T and B cells, preventing their entry into the brain, gut, and skin, and forcing migration of bone marrow CD34+ into peripheral circulation for long periods.9,17,18 Rituximab, another monoclonal antibody, binds the CD20 surface molecule on B cells, causing their depletion from the peripheral circulation through complement-mediated cytolysis.7

### Risk factors for development of PML

Measurable risk factors for PML include:

- Rising antibody titers
- Evidence of viremia, especially persistent viremia associated with repeat sequences in the regulatory region of the viral genome
- Ineffective T-cell (CD4 and CD8) responses
- Molecular host factors (ie, Spi-B expression in B cells) that support JCV infection in potentially susceptible cells.

The presence of more than one of these risk factors is necessary for development of PML.

### VIRAL LATENCY IN B LYMPHOCYTES IN BONE MARROW

A strong link between JCV infection in cells of the immune system and those of the nervous system points to the importance of the tissue origin of JCV latency. Bone marrow harbors CD34+ cells that migrate into the peripheral circulation and undergo differentiation to pre-B and mature B cells, augmenting JCV growth. The emergence of PML in patients treated with natalizumab, rituximab, efalizumab, and other immune-altering drugs underscores this observation.

As noted, the incidence of PML in natalizumab-treated patients with MS and Crohn disease rises as the number of doses increases. Analysis of blood samples
collected from patients treated with natalizumab at baseline and again during treatment at months 1 to 12 and beyond 24 months demonstrates that the frequency of CD34+ cells in the peripheral circulation increases with the duration of therapy, adding credence to the theory that CD34+ cells act as a reservoir for latent virus. A higher frequency of CD34+ cells is associated with viremia. A higher frequency of CD34+ cells is associated with viremia.

The role of Spi-B in JC virus latency
Understanding the role of Spi-B during JCV latency and reactivation is increasingly important as the number of patients treated with immunomodulatory agents that can develop PML continues to rise. Spi-B is highly represented in the B cell and CD34+ cell fractions. Spi-B expression in B cells correlates with reactivation of JCV in immune cells in natalizumab-treated patients. In a sample of four patients with MS treated with natalizumab who developed PML, T-cell responses have been ineffective (absent or aberrant). Two patients had no detectable T-cell response to JCV; the other two demonstrated response, but their CD4 T-cell responses were dominated by interleukin-10–producing cells.

Longitudinal examination of CSF samples from 13 MS patients who were treated with natalizumab and subsequently developed PML revealed persistence of viral load even though all patients experienced immune reconstitution inflammatory syndrome and most had high levels of anti-JCV antibodies.19

Dr. Rudick: Have you found the virus in B cells in healthy people?

Dr. Major: We never tested this directly. From human brain cultures, we know that Spi-B is made in glial cells, not in neurons. We are considering the idea that wherever JCV binds, it takes advantage of certain types of DNA-binding proteins in the molecular regulation. If the binding takes place in an immune system cell, for example, Spi-B plays an important role.

Dr. Berger: Koralnik et al demonstrated JCV excretion in urine in MS patients after 12 months of treatment with natalizumab, and at 18 months, viremia in 60% of the patients.20 Yet, repeated studies of patients taking natalizumab have failed to demonstrate viremia or conversion of virus in the archetype. How do these findings correlate with your thoughts on the action of natalizumab in the pathogenesis of PML?

Dr. Major: We certainly know that natalizumab forces migration of hematopoietic stem cells and pre-B cells out of the marrow, but our findings have differed somewhat from those of Koralnik’s laboratory. For example, in the several hundred nucleotide sequences we have looked at in PML brain tissue, we have found the Mad 1 genotype once. We consider Mad 1 to be a potential laboratory contamination, so if we find Mad 1 we resequence the sample. We never clone because cloning can introduce alterations; we sequence directly from the clinical tissue. We can identify Mad 1 because our assay is very sensitive. In normal individuals, CD34+ cells compose approximately 0.01% of the peripheral circulation; in individuals treated with natalizumab, however, their composition is 0.1% to 0.3%. So if there is a potential for latent infection, we have an opportunity to find it in those cells. Its presence does not necessarily mean that the individual is going to develop PML, however; there are other controlling factors.

Dr. Berger: Does natalizumab upregulate Spi-B in glial cells?

Dr. Major: Does natalizumab force the migration of hematopoietic stem cells and pre-B cells from the bone marrow by preventing their attachment to vascular-cell adhesion molecule (VCAM). The JC virus (JCV) can reside in the bone marrow in a latent state for extended periods and migrate to the peripheral circulation. JCV uses B cells and their DNA-binding proteins to initiate viral replication.


Dr. Berger: Has Spi-B been detected in B cells in developing brain?

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Dr. Rudick: Do you support a role for Spi-B in controlling late phase JCV in latency?

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Dr. Rudick: Have you found the virus in B cells in healthy people?
Dr. Major: Yes we have, in about one-third. It is higher than what we would expect to see in the normal population.

Dr. Rudick: How can that finding be turned into something that’s clinically useful?

Dr. Major: If you’re trying to identify persons who are more susceptible to PML given underlying risk factors—treatment with natalizumab or rituximab, presence of HIV infection, or some other immune-altering condition—looking at one parameter isn’t going to help. Based on the available data, rising antibody titers signals an active infection, and viremia of any kind means probable latent infection. Because this is a small event in very few cells, you will not have the numbers of cells needed to identify susceptibility in a normal population. For now, we monitor patients at risk and, if we find viremia, we assess the cell population to determine whether a molecular factor like Spi-B is upregulated. We hope to develop an assay in which we can obtain one test tube of blood and report T-cell responses, molecular factors, antibody titer, and presence or absence of viremia. Such an assay would provide the data necessary to make a clinical decision.

REFERENCES


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The clinical features of PML

ABSTRACT

The symptoms associated with progressive multifocal leukoencephalopathy (PML) reflect the location of pathologic brain lesions. These symptoms include visual deficits, cognitive impairment, and motor weakness; in patients with acquired immunodeficiency syndrome (AIDS), presenting signs can also include gait disturbance, dysarthria, dysphasia, and ocular palsy. Recently, PML has been observed in patients treated with biologic agents; natalizumab recipients currently represent the second largest group of patients with PML (behind patients with AIDS). Although brain biopsy is the most accurate and reliable method for diagnosing PML, it is rarely used today. Diagnosis is usually based on detection of JC virus in the cerebrospinal fluid by polymerase chain reaction, the clinical presentation, and demonstration of PML brain lesions on magnetic resonance imaging. With immune reconstitution, the prognosis of PML has improved markedly.

Progressive multifocal leukoencephalopathy (PML) was a rare disease until the era of human immunodeficiency virus (HIV) infection, when the number of cases of PML markedly increased. We are now entering a new era in which PML is being observed in patients treated with biologic agents for diseases not associated with development of PML.

This article reviews the epidemiology and symptoms that characterize PML, the identification of lesions on radiographic imaging that support the diagnosis, the value of laboratory studies and immunocytochemistry in the diagnosis, and clinical outcomes.

CHANGING EPIDEMIOLOGY OF PML

The presentation and epidemiology of PML have evolved over the last several decades. Prior to the acquired immunodeficiency syndrome (AIDS) era, nearly one-half of patients with PML were affected by visual deficits. Among patients with AIDS-related PML, motor weakness is a feature for more than one-half, while cognitive impairment is the most common clinical presentation in natalizumab-related PML, affecting nearly one-half of patients (Figure).

The pre-AIDS era

Lesions of subcortical white matter characterize PML and the patient’s clinical manifestations reflect their location. Brooks and Walker reviewed 69 pathologically confirmed and 40 virologically and pathologically confirmed cases of PML in the era before AIDS, and categorized the neurologic signs and symptoms at onset and during disease progression; the clinical picture had three significant findings:

- Impaired vision: Defective vision, most commonly homonymous hemianopsia, was the most frequent presenting sign, present in 35% to 45% of cases. At the time of diagnosis, 6% to 8% of the patients were cortically blind because of bioccipital pathology.
- Motor weakness: Motor weakness was the initial sign in 25% to 33% of patients. At the time of diagnosis, hemiparesis or hemiplegia was present in nearly all patients.
- Changes in mentation: A change in mentation, including personality change, difficulty with memory, emotional lability, and frank dementia, was the presenting sign in approximately one-third of cases and eventually involved most patients.

AIDS-related PML

The epidemiology of PML changed with the AIDS pandemic. From 1958 to 1984, Brooks and Walker identified 230 cases of PML; in the period from 1981 to 1994, Berger and colleagues described 154 cases of AIDS-related PML that had been identified by the University of Miami Medical Center and the Broward County medical examiner’s office. The frequency of PML from 1991 through 1994 was 12-fold greater than the frequency 10 years earlier, from 1981 through 1984. Among the patients with AIDS-related PML, the most common initial symptoms were weakness (42%), speech abnormalities (40%), cognitive abnormalities (36%), gait abnormalities (29%), sensory loss (19%), and visual impairment (19%), followed by seizures, diplopia, and...
limb incoordination. The most common findings at the time of initial physical examination were weakness (54%), followed by gait abnormalities (20%), cognitive abnormalities (20%), dysarthria (24%), aphasia (19%), sensory loss (19%), visual impairment (17%), and oculomotor palsy (6%). For about 5% of patients with PML, it is the heralding manifestation of AIDS.

Although clinical features consistent with cerebral hemisphere lesions are most common, brainstem and cerebellar findings are also observed. Among these are ataxia, dysmetria, dysarthria, and oculomotor nerve palsies.2–4 Other signs and symptoms associated with PML include headache, vertigo, seizures, sensory deficits, parkinsonism,5 aphasia, and neglect syndromes.1–4 In some cases, the coexistence of encephalitis with HIV infection could have accounted for some of the symptoms.

PML associated with monoclonal antibody therapy
Natalizumab is an alpha-4-beta-1 integrin inhibitor approved for the treatment of relapsing-remitting multiple sclerosis (MS); patients taking natalizumab represent the second largest group with PML (the largest group is patients with AIDS). Natalizumab-associated PML has some noteworthy features. The most common clinical presentations are cognitive disorders (48%), motor abnormalities (37%), language disturbances (31%), and visual defects (26%). Lesions are often monofocal rather than multifocal and the most common site of involvement is the frontal lobe.6 Among MS patients with natalizumab-associated PML, 30% to 40% have gadolinium-enhancing lesions on magnetic resonance imaging (MRI) at the time of diagnosis.

IMMUNE RECONSTITUTION INFLAMMATORY SYNDROME

The immune reconstitution inflammatory syndrome (IRIS) is a paradoxical clinical deterioration that typically occurs in the face of immunologic recovery (Table).7,8 Although not unique to PML or opportunistic infections, it is frequently observed in the setting of HIV infection and natalizumab-associated PML, and it may be concurrent with the diagnosis of PML.6,9

Among patients with HIV, predictors for the development of IRIS include antiretroviral naive, profoundly low CD4 lymphocyte counts (< 50 cells/mm3), a rapid decrease in HIV load, and the presence of active or subclinical opportunistic infections at the time of initiation of combined antiretroviral therapy. Tan and colleagues8 have reported the largest series to date. Of the 54 patients in their series, 36 developed PML and IRIS simultaneously, and 18 had worsening of preexisting PML. Although some investigators have recommended corticosteroid therapy for PML-IRIS, no controlled trials have been conducted and caution has been advised, particularly in patients without contrast enhancement on MRI or mass effect.

DIAGNOSTIC TESTING: NEUROIMAGING, CEREBROSPINAL FLUID ANALYSIS

Neuroimaging, including computed tomography (CT) and MRI, is a useful diagnostic tool for investigating a patient with PML. Cerebrospinal fluid (CSF) analysis for the presence of JC virus (JCV) may play a significant role, but it primarily serves to rule out other illnesses.

Computed tomography: lesion size may not reflect clinical status
On CT, demyelinating lesions appear as subcortical hypodensities, often with a propensity for parieto-occipital areas that are confined to the white matter at the junction interface of the gray-white junction of the cortex.9–11 Lesions may be seen in the corpus callosum, thalamus, and basal ganglia,9 but changes in the size of lesions observed on CT do not necessarily reflect clinical progression.12 Prior to the availability of highly active antiretroviral therapy (HAART) for the treatment of AIDS, mass effect was exceptionally rare.
However, the development of IRIS with PML, typically in AIDS patients following the use of HAART, may be associated with edema. Single-dose intravenous contrast and delayed, double-dose contrast CT scanning enhancement is observed in a minority of patients, typically fewer than 10%. This enhancement is generally faint and peripherally located.

Magnetic resonance imaging may show lesions before clinical disease

MRI is vastly more sensitive than CT in detecting the demyelinating lesions of PML. On rare occasions, MRI will clearly demonstrate pathology when CT is normal. In fact, MRI may show lesions in advance of clinically apparent disease. The characteristics of these lesions are hyperintensity on T2-weighted imaging, fluid-attenuated inversion recovery sequences, and hypointensity on T1-weighted image. Apparent diffusion coefficients (ADC) on MRI are typically normal to low in new lesions and at the advancing edge of lesions; the ADC was typically higher in the center of lesions.

As observed on CT, approximately 10% of patients exhibit a faint rim of gadolinium enhancement. Enhancement is more common with PML-IRIS, and the distribution of lesions parallels what is seen pathologically. Enhancement PML lesions have altered signal characteristics compared with the surrounding white matter. In contrast, 15% of HIV-associated PML showed gadolinium enhancement on MRI at the time of diagnosis.

Cerebrospinal fluid analysis

With the exception of polymerase chain reaction (PCR) for JCV, the primary utility of lumbar puncture in the setting of possible PML is to exclude the presence of other illnesses, including treatable infections.

CSF findings in patients with PML are nonspecific, with most patients demonstrating a normal profile. A mild lymphocytic pleocytosis, which is rarely (if ever) more than 25 leukocytes/mL, occurs in 15% of patients. Total protein level is mildly elevated in approximately 20% to 30% of patients.

The CSF examination in HIV-infected patients with PML may reflect changes associated with HIV: low-grade lymphocytic pleocytosis (< 20 cells/mm³), mildly elevated protein (< 65 mg/dL), and elevated immunoglobulin G and oligoclonal bands. These abnormalities should not be attributed to PML.

**TABLE**

**Features of PML-IRIS**

- Paradoxical worsening of clinical or radiographic finding with recovery of the immune system
- New or increased neurologic deficits
- Increase in the number or size of lesions on neuroimaging
- Contrast enhancement of brain lesions
- Brain edema
- Concurrent with diagnosis of PML

IRIS = immune reconstitution inflammatory syndrome; PML = progressive multifocal leukoencephalopathy

DIAGNOSIS

The most reliable and accurate method for the diagnosis of PML remains brain biopsy that demonstrates the characteristic triad of histopathologic findings (demyelination, bizarre astrocytes, and enlarged oligodendrocyte nuclei) coupled with evidence of JCV infection. With respect to the latter, in situ hybridization or immunocytochemistry can be employed. In situ DNA-DNA hybridization is a method of annealing JCV DNA to complementary strands either in paraffin-embedded tissue or in frozen sections from biopsy samples.

In immunocytochemistry, antibodies to both T antigen and the common polyomavirus capsid antigen are used to detect cells undergoing productive viral infection. Cells that are positive by in situ hybridization are in a stage of active viral replication. Cells positive by immunocytochemistry that are expressing viral capsid antigens are in a stage of viral transcription and translation (ie, undergoing productive infection). In addition to their utility in confirming a diagnosis of PML, these techniques have demonstrated the presence of JCV in perivascular locations and at sites distant from foci of demyelination. Alternatively, PCR may be used to demonstrate JCV in brain tissue.

In the absence of biopsy, which few deem necessary today, a widely employed approach to diagnosis requires the demonstration of:

- JCV in the CSF by PCR
- Compatible clinical presentation
- An MRI finding consistent with PML
- No other alternative diagnosis.

With an ultrasensitive PCR technique, sensitivities should approach or may exceed 95%, but PCR sensitivity remains at 75% in some laboratories. Because the viral copy numbers in the CSF may be low, particularly in a patient treated with a monoclonal antibody such as natalizumab, the CSF PCR may be falsely negative.

If clinical suspicion of the disease remains high in the face of an initially negative CSF JCV, the CSF analysis should be repeated. CSF analysis for JCV is approximately 99% specific, but recent studies demonstrating low copy numbers of JCV in the CSF of patients with MS have raised concerns about potential pitfalls of this assay.
PROGNOSIS

Until recently, PML was regarded as virtually universally fatal. The mean survival in the pre-AIDS era was approximately 6 months, and mortality was 80% within 9 months of disease onset. Rarely, patients had long survivals that ranged from 5 years to 19 years.

In the early years of the AIDS era, survival with PML did not appear to differ significantly from that observed in the pre-AIDS years. In the largest study of HIV-associated PML in the era prior to HAART, the median survival was 183 days. However, the majority of individuals were dead within 3 months of diagnosis. Only 8% to 10% of patients survived longer than 12 months, which has been regarded as “prolonged survival.” This long survival skewed the mean and median survival rates in this population.

Several factors have since been identified that correlate with prolonged survival in HIV-associated PML, including PML as the heralding manifestation of HIV, CD4 counts exceeding 300 cells per mm3, contrast enhancement of the lesions on radiographic imaging, low copy number or decreasing JCV titers in CSF21–24 and the presence of JCV-specific cytotoxic T cells.25 A better prognosis has also been postulated for higher CSF levels of macrophage chemotactic protein-126 and PML associated with JCV VP1 loop-specific polymorphisms.27

Prognosis of HIV-associated PML improves with immune system restoration

In the era of HAART, not only has the incidence of HIV-associated PML declined, but the prognosis of affected patients has improved as well. This development highlights the importance of restoration of the immune system in both disease prevention and survival. Some estimate that as many as 50% of HAART-treated patients with PML exhibit prolonged survival. In one study of 25 patients, the median survival was more than 46 weeks.28

Nonetheless, PML continues to have the worst prognosis of any AIDS-related cerebral disorder, with those having advanced immunosuppression being most susceptible to the disorder. For AIDS patients with PML, those who were HAART-naive at the time of diagnosis appear to have better survival than treatment-experienced patients.29 Survival also correlates with reduced JCV load in the CSF30 and improved CD4 lymphocyte counts (CD4 counts > 100 cells/mm3).31

Prognosis of natalizumab-associated PML is different

The prognosis of natalizumab-associated PML differs from that of HIV-associated PML. In a series of 35 patients, 25 (71%) patients were alive on average 6 months after diagnosis.32 Prognosis was worse with a longer time to diagnosis and the presence of widespread disease.

Most deaths in patients taking natalizumab who developed PML have occurred during IRIS. Steroid treatment of IRIS appears to improve prognosis,3 but no scientifically rigorous study has been undertaken to demonstrate this recommendation. Among the survivors, neurologic deficit was mild in one-third, moderate in one-third, and severe in one-third of patients.

CONCLUSION: DISPELLING SOME MYTHS

Several assumptions about PML are not necessarily true. For example, although PML implies the presence of multifocal lesions as a characteristic of the disease, the lesions may be monofocal, especially with natalizumab-associated PML. The lesions of PML may show early gadolinium enhancement on neuroimaging. Although lesions typically are seen in subcortical white matter, cortical involvement also may be observed. Cerebellar granular cell degeneration may occur in association with PML or in isolation. Disease progression and death are not inevitable, even in the absence of treatment. The most important determinant for survival is restoration of the immune system.

DISCUSSION

Dr. Calabrese: Why are sensory deficits so common?

Dr. Berger: We don’t know. Because we see involvement in the parietal lobe, we would anticipate observing sensory deficits. I think that a lot of sensation occurs deep in the thalamic area, which is not often involved in PML. Also, we often don’t test for some of the deficits that may occur.

Dr. Rudick: Do you know of any cases of natalizumab-associated PML detected as an incidental finding on MRI, making a case for screening MRI in patients without clinical symptoms?

Dr. Berger: There have been a handful of cases, including one of the seminal cases of natalizumab-associated PML, in which MRI abnormalities were observed in advance of clinically recognized symptomatology.

Dr. Calabrese: The correlate question is, if a patient with a risk factor—be it HIV or treatment with a biologic agent—has a common neurocognitive sign or perhaps some subtle motor findings, does a normal MRI have 100% negative predictive value?

Dr. Berger: I have yet to see somebody with PML who has a normal MRI.

Dr. Simpson: What you may see are lesions that are not typical MRI lesions of white matter hypointensity. In some cases, as Dr. Berger mentioned in his summary, we’ll see cerebellar degeneration—atrophy—but not necessarily white matter lesions.
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Pharmacovigilance and PML in the oncology setting

ABSTRACT

Methods developed by the Southern Network on Adverse Reactions project, the only state-funded pharmacovigilance program in the nation, are invaluable in identifying rare and serious drug events and in disseminating related safety reports quickly throughout the medical community. An important discovery was identification and reporting of an association of rituximab and progressive multifocal leukoencephalopathy (PML) in patients without human immunodeficiency virus (HIV). A recent investigation identified 57 patients with rituximab-associated PML, including bone marrow samples, brain biopsies, and autopsy materials from patients with lymphoma and PML who tested positive for JC virus. The investigation identified an association of rituximab-chemotherapy administration and PML, although a causal relationship remains an area of active investigation. Additional investigations evaluated the epidemiology of PML in the oncology setting before and after the introduction of rituximab for lymphoma treatment. Focused analyses investigated risk factors for development of this rare complication. Further studies are needed to investigate the pathophysiology, epidemiology, and risk factors for PML developing among HIV-negative cancer patients who receive rituximab and chemotherapy.

Organizations such as the Southern Network on Adverse Reactions (SONAR), which has become the only state-funded pharmacovigilance initiative in the nation, have identified potentially fatal and previously unreported side effects associated with 43 drugs—with the majority of these drugs involving the hematology and oncology disciplines.

Dr. Bennett reported that he has no financial relationships that pose a potential conflict of interest with this article.

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time for a “black box” warning to appear on a package insert following FDA approval is 7 to 10 years.3

**Timely and thorough data collection**

Academic pharmacovigilance organizations such as SONAR operate differently from the FDA and pharmaceutical manufacturers in their search for adverse drug events (Figure).4 SONAR collects reports from investigators, clinicians, attorneys, patients, and family members on suspected treatment-related adverse events and investigates these reports carefully. Direct calls to hospitals and large centers can be useful in searching for cases, using information obtained from Internal Review Boards and medical records.

SONAR investigators perform extensive literature reviews, may request more data from authors, and request and review additional FDA case reports. Unfortunately, obtaining data from the FDA can be difficult and time-consuming. Data can be requested through the Freedom of Information Act (FOIA), but receiving it may take more than a year, and the information in the public record may be redacted. SONAR obtains laboratory tests and imaging records and works with scientists to better understand the pathophysiology of potential treatment-related rare adverse events, investigate epidemiologic estimates of the side effect rate, and evaluate risk factors for development of toxicity.

Adverse events are usually identified by SONAR within 2 years post–drug approval—a 5-year improvement over the FDA on this important metric. Once an adverse event is positively identified, the information is disseminated throughout the worldwide medical community via journal articles and presentations at medical conferences. Funding is grant-based from sources such as the National Institutes of Health (NIH), the state of South Carolina, and the University of South Carolina.

**FDA, manufacturer reports may be incomplete and delayed**

In contrast with SONAR, the FDA relies heavily on MedWatch to detect cases of adverse events. The safety record compiled by MedWatch is often incomplete because the program relies on voluntary submissions of adverse events; further, the inordinate amount of followup required of physicians discourages many from participating. The time to identify an adverse event can be several years, and the FDA disseminates adverse event reports via package inserts. The network that evaluates the safety information and identifies initial safety signals is mainly internal to FDA employees, as is the funding.

Pharmaceutical manufacturers frequently compile data from their own proprietary databases. Although they attempt to follow up on reports of rare adverse events, it is often difficult or impossible for the company to obtain followup information from busy clinicians. Identifica-
tion of an adverse event typically takes 7 to 12 years for most pharmaceutical manufacturers—reflecting the barriers experienced in obtaining detailed information from clinicians about potential new serious adverse drug reactions. Findings are frequently disseminated through “Dear Doctor” letters. Manufacturers’ investigative networks, like those of the FDA, are largely internal and the amount of funding of they allocate to drug safety investigations is unknown.

## RARE EVENTS MAY INVOLVE FEW CASES

Of our major publications,5-14 many findings are based on a small number of cases—for example, only 13 cases for clopidogrel-associated thrombotic thrombocytopenic purpura (TTP)13 and 9 for pure red cell aplasia caused by epoetin alfa.12 Important findings also come from meta-analyses,8,10 although this avenue in our pharmacovigilance approach is less typical.

The 2008 study on mortality and venous thromboembolism associated with erythropoiesis-stimulating agents highlights the importance of basic scientific investigation in identifying rare events. Administration of epoetin alfa to raise hemoglobin levels had been approved by the FDA in 1989 for use in patients undergoing dialysis and in 1993 for supportive use in patients with some types of cancers. We discovered that epoetin alfa promoted cancer growth based on analysis of published data and reports in conjunction with basic scientific studies of erythropoietin and erythropoietin receptors in solid cancers.

## RITUXIMAB AND VIRAL REACTIVATION

In the case of viral reactivation associated with the use of rituximab, a warning about hepatitis B reactivation was added to the package insert in 2004.15 In 2006, a warning about other viral infections was added to the package insert.16 In late 2006, a letter was sent to health care professionals from the manufacturer and the FDA with the warning that PML had been observed in two patients with systemic lupus erythematosus (SLE) who were treated with rituximab (an off-label use), both of whom were negative for human immunodeficiency virus (HIV).16 A few months later, a black box warning to this effect was added to the package insert.16

After we identified PML as an adverse event from rituximab in HIV-negative patients,14 we obtained case reports from clinicians at 12 cancer centers or academic hospitals (22 cases). We also reviewed FDA reports (11 cases), the manufacturer’s database (30 cases), and publications (18 cases) using the search terms “leukencephalopathy,” “rituximab,” “immunosuppressed,” “lymphoma,” and “leukemia.” The unique data sources included clinical observations, the medical literature, FDA MedWatch, and the manufacturer.17

Of rituximab-treated patients who developed PML, the mean age was 61 years (range, 30 to 89 years), 56% of patients were women, and the mean number of rituximab doses was six (range, 1 to 28). Six patients had undergone stem cell transplants (four autologous), and 26 were also taking a purine analogue.17

Among 57 patients, a median of 16 months elapsed between first taking rituximab to development of PML (range, 1.0 to 90.0 months), and 5.5 months from the last dose of rituximab to development of PML (range, 0.3 to 66.0 months). The median time from diagnosis of PML to death was only 2.0 months (range, 0.4 to 122 months). Reported survival rates for patients with rituximab-associated PML who did not undergo stem cell transplantation was less than 10%.17

The symptoms of PML are easily confused with those that might be expected in an older patient with lymphoma, making early detection especially difficult. More than one-half (54.4%) had confusion or disorientation, and many had focal motor weakness (33.3%), loss of coordination (24.6%), difficulty speaking (21.2%), and vision changes (17.5%).

### Effects on T and B cells and role of JC virus

At the time of PML diagnosis, 90% of patients had either a severely low CD4+ count or a low CD4+:CD8+ ratio. Based on clinical trial data, cytotoxic chemotherapy and not rituximab appears responsible for the abnormal CD4+ count and the low CD4+:CD8+ ratio in rituximab-treated patients.

Little is known about how T cells function after rituximab administration. In idiopathic thrombocytopenic purpura, the response to B-cell depletion induced by rituximab is associated with significant changes in the T-cell compartment.18 In a study of patients with either SLE or Evans syndrome, rituximab therapy was found to modify T-cell phenotype and cytokine profiles.19 The rapid effect of rituximab in multiple sclerosis suggests that it targets a process thought to be T-cell mediated.20

Our early hypothesis was that rituximab contributes to viral reactivation and PML through inhibiting T- and B-lymphocyte interactions. We now believe that the bone marrow plays an important role, which may explain the process by which natalizumab can cause PML. Five of five bone marrow samples from patients with lymphoma and PML tested positive for JC virus (JCV) compared with only two of 86 bone marrow samples from patients without PML. The JCV is latent in CD34+ hematopoietic cells and probably in early B lymphocytes. Chemotherapy mobilizes the stem cells from bone marrow and causes quantitative T-cell depletion. Rituximab reduces the qualitative T-cell response, and B-cell depletion results in expansion of progenitor cells containing the latent JCV. The hypothesis is limited in that it is based on a retrospective case series and
is not verified in a laboratory model.

Of the 57 cases of PML identified in 2009, two patients were given rituximab for hematologic disorders and had no chemotherapy other than steroids. These data suggest that rituximab confers risk on its own.27

Quantifying risk of developing PML from rituximab
Calculating the odds of developing PML from rituximab therapy is difficult. The background rate of PML is an important consideration. One population-based study estimated the incidence of PML in patients with hematologic malignancies at 0.07%. This estimate was based on three cases of PML observed in patients with hematologic malignancies over a period of 11 years in a single Canadian province.21 Another study found a higher incidence of 0.52% in patients with chronic lymphocytic leukemia, although all of these patients were also treated with fludarabine.22 Accurately calculating the risk of PML attributable to the underlying malignancy as opposed to immune suppression from treatment is complicated by the rarity of the disease. Fludarabine is the chemotherapeutic agent most closely associated with PML. However, its well known side effects of T-lymphocytic depletion and complicating opportunistic infections similar to those seen in acquired immunodeficiency syndrome (AIDS) make such an association intuitive.23

Kavenaugh and Matteson reported that about 8,000 SLE patients had received rituximab treatment and two of these patients had developed PML.24 PML has been reported previously among 30 SLE patients who had not received rituximab, suggesting that SLE is a predisposing disorder.25,26

In the setting of hematologic malignancy, rituximab-associated PML incidence estimates are complicated by a low basal risk of PML seen among persons with the disease state prompting rituximab therapy and an inability to determine risk attributable to rituximab. A recent study demonstrated an association between rituximab and PML in patients with non-Hodgkin lymphoma (NHL). The retrospective, monocentric cohort study assessed data from 976 NHL patients diagnosed in Italy from 1994 to 2008, including 517 patients who received at least one dose of rituximab. Inclusion of rituximab into standard chemotherapy regimens for NHL caused a significantly higher incidence of PML cases (rate difference, 2.2 every 1,000 patient-years; 95% confidence interval, 0.1–4.3).27 More such studies of viral reactivation syndromes are obviously needed.

Ideally, randomized clinical trials of the use of rituximab in patients with lymphoma would serve as guidance, but because the drug as the standard of care for treatment of lymphoma, is so widely used, randomization would be impractical.

Future planned studies include a case-control study of T-cell markers after chemotherapy administration with or without exposure to rituximab, a case-control study of bone marrow specimens from disease-matched and treatment-matched controls, and a cohort study using a large electronic medical records database or a government database.

■ CONCLUSION

The methods developed in the SONAR project will permit exploration of important hypotheses regarding the detection and prevention of rare adverse events in oncology, forming a basis for subsequent investigations. Based on our recent findings, rituximab may be associated with multiple viral reactivation syndromes; screening and early detection can potentially be helpful in preventing these complications.

■ DISCUSSION

Dr. Calabrese: Your approach to identifying rare adverse events is novel and aggressive, but the seeming limitations in a disease such as lymphoma are (1) rituximab is now a standard of care so everybody with lymphoma gets it, and (2) going back to the earliest descriptions of PML, lymphoma has always been represented as a predisposing factor. Moving ahead, how then can you calculate an effect size for a drug like rituximab?

Dr. Bennett: There’s no way to do it; we’re sort of stuck. Of the 57 cases with PML that we reported in Blood,17 two patients received rituximab for hematologic disorders and received no chemotherapy besides steroids. In those two patients, we could not blame the development of PML on lymphoma. Those types of patients suggest that rituximab may be implicated, but examining this question with a case-control or even a cohort study is an expensive proposition.

Dr. Simpson: My experience in terms of collaborating with the FDA has been distinctly unrewarding. Some years ago I had been looking into an adverse effect related to the nucleoside analog reverse transcriptase inhibitor d4T, in which there was a rapidly progressive neuromuscular weakness syndrome that looked like Guillain-Barré and lactic acidosis. The FDA itself reported 12 cases at an international AIDS conference and did not have any answers. I was charged by the AIDS Clinical Trials Group and other branches at the NIH to try to figure it out. When I requested access to FDA data, I ran into an unbelievable bureaucratic morass. Ultimately, we had to go through the FOIA to get them to release anything.

Dr. Bennett: The FOIA is the only way to get anything from the FDA. It takes about a year and a half and much information is redacted.
Dr. Bennett: The Risk Evaluation and Mitigation Strategies program authorizes the FDA to require post-marketing surveillance of all adverse events from manufacturers. We published 11 cases of TTP in association with clopidogrel, obtained from surveillance of directors of plasmapheresis centers in the United States. Not one of them had been reported to the FDA directly. However, we had an article 6 weeks after clopidogrel received FDA approval. Now, 10 years later, there are about 120 clopidogrel-associated TTP cases in the FDA database. Its estimated incidence is still one in a million, although we hear about the side effect every night on TV during commercials for the drug on the evening news.

Dr. Major: The FDA is more open now than in the past to trying to get a handle on what’s going on with biologic therapies. We need to do a little more homework up front on biologic agents in order to anticipate some adverse events. For example, the migratory nature of CD34+ cells through the circulation following natalizumab therapy was not appreciated, even though data in the literature already supported this phenomenon when integrin receptors are blocked.

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ABSTRACT

The treatment benefits of natalizumab in patients with multiple sclerosis (MS) appear to exceed those of other disease-modifying drugs, but progressive multifocal leukoencephalopathy (PML) has been identified as a risk in patients receiving natalizumab. As of August 2011, a total of 150 cases of natalizumab-associated PML had been reported worldwide. The overall risk is estimated at approximately 1.66 in 1,000 patients. Independent risk factors for natalizumab-associated PML are number of infusions beyond 36 and prior use of immunosuppressive drugs. Classifying JC virus antibody status appears to be useful in treatment decision-making for individual MS patients. Patient tolerance for risk plays an important role in the selection of therapy, and the treating physician’s perception and tolerance of risk may differ markedly from the patient’s. Physicians can help patients make individual informed decisions regarding the use of natalizumab, given the known risk of PML.

Natalizumab, and more recently, fingolimod, are generally used as second-line agents. Fingolimod, the first oral agent to receive FDA approval for the treatment of relapsing-remitting MS, is a functional antagonist of sphingosine-1-phosphate receptors. The reductions in annualized relapse rates in two phase 3 controlled trials of fingolimod were approximately 55% compared with placebo or intramuscular interferon beta-1a. Because of its more convenient oral route of administration and its documented efficacy, widespread use of fingolimod is anticipated. However, adverse reactions affecting more than 10% of patients include headache, influenza, diarrhea, back pain, liver transaminase elevations, and cough. Because sphingosine-1-phosphate receptors are widespread in many body tissues, off-target effects of fingolimod may be problematic and long-term toxicity is unknown.

In addition to natalizumab and fingolimod, which are currently available for use as second-line agents, several other MS therapies are showing promise. Oral cladribine, teriflunomide, and laquinimod have reported positive phase 3 results in publication or at national meetings, and several other drugs are in late stages of development (alemtuzumab, BG-12, ocrelizumab) based on encouraging phase 2 results. Thus, the options for MS patients are expanding, but drugs with higher efficacy also may pose greater risk.

NATALIZUMAB: ROBUST BENEFITS BUT ASSOCIATED RISK

Natalizumab is a humanized monoclonal antibody that binds to alpha-4 integrin on leukocytes. By inhibiting alpha-4 integrin, natalizumab, the first of a new class of selective adhesion-molecule inhibitors, impedes migration of activated mononuclear leukocytes into the brain and gut.

Significant efficacy

Two phase 3 studies demonstrated more robust efficacy of natalizumab in patients with relapsing-remitting MS than had been observed in prior studies with other agents. In the Natalizumab Safety and Efficacy in Relapsing-Remitting Multiple Sclerosis (AFFIRM) study, which followed patients over 2 years of treatment, natalizumab was
associated with a 68% reduction in the annualized relapse rate\(^1\), a 92% reduction in gadolinium-enhanced lesions on magnetic resonance imaging (MRI), which indicate new, active lesions\(^6\); and an 83% reduction in the mean number of new or enlarging T2 lesions\(^6\) compared with placebo. The likelihood of confirmed worsening on the Kurtzke Expanded Disability Status Score, which is the standard measure for MS-related disability, was also 42% lower in patients assigned to natalizumab compared with placebo.\(^{14}\)

Other reported benefits from natalizumab therapy include a significantly increased probability of maintaining disease-free status\(^7\) and clinically significant improvements on patient-reported quality-of-life measures.\(^{18}\)

Although there have been no head-to-head studies of natalizumab with interferon beta, glatiramer acetate, or fingolimod, there is a widespread view that treatment benefits of natalizumab exceed those of other disease-modifying drugs. In clinical practice, patients with MS who experience breakthrough disease on standard disease-modifying drugs are routinely observed to achieve disease control after switching to natalizumab. Thus, based purely on efficacy, patient-reported outcomes, and the convenience of once-monthly intravenous infusion, natalizumab represents an extremely attractive treatment option for patients with relapsing-remitting MS.

**Use discontinued in 2005**

Natalizumab was approved for treatment of relapsing-remitting MS in November 2004, using the FDA accelerated review pathway. The approval was based on the first-year results of the AFFIRM\(^{14}\) and the Natalizumab plus Interferon Beta-1a for Relapsing Remitting Multiple Sclerosis (SENTINEL)\(^{19}\) studies, both of which were completed in February 2005. In the 3 to 4 months between the drug’s approval and completion of the AFFIRM and SENTINEL studies, approximately 7,000 patients with relapsing-remitting MS received treatment with natalizumab. In February 2005, shortly after the release of the 2-year data, three cases of progressive multifocal leukoencephalopathy (PML) were identified in natalizumab-treated patients (one with Crohn disease, the other two with MS). Clinical and research use of natalizumab was abruptly suspended that month, pending a comprehensive safety review.

A safety study evaluated 3,116 patients who had received natalizumab over a mean exposure of 17.9 monthly doses.\(^{20}\) The study failed to identify any additional cases of PML and concluded that the risk of PML was approximately 1 in 1,000 patients. Abrupt discontinuation of natalizumab also allowed systematic assessment of disease behavior following treatment interruption. In 1,866 patients who had received natalizumab during clinical trials but who discontinued natalizumab after PML was recognized, MS relapses and gadolinium-enhancing lesions returned approximately to baseline levels within 4 to 7 months of natalizumab suspension. Reactivation of MS disease activity was observed even in patients who instituted one of the first-line disease-modifying drugs as substitute therapy.\(^{21}\)

Based on the strong efficacy data and the extensive safety review, an FDA advisory committee recommended reintroduction, and natalizumab was returned to the market in June 2006. Natalizumab may be administered only in accredited infusion centers that agree to a monthly reporting regimen designed to identify all cases of PML. In the United States, natalizumab is available to patients only through the Tysabri Outreach Unified Commitment to Health (TOUCH) Prescribing Program, a restricted distribution program. Aggressive monitoring and reporting is also required in other regions of the world, so that ascertainment of PML associated with natalizumab is thought to be relatively complete.

**Risk related to duration of therapy**

As of June 30, 2011, some 88,100 patients had been dosed with natalizumab in the postmarketing setting.\(^{22}\) The overall risk of PML in these patients was originally estimated to be 1 in 1,000 individuals taking the drug; the risk now is estimated to be 1.66 in 1,000 patients. Postmarketing surveillance has revealed differences in risk when duration of therapy is considered (Figure 1).\(^{22}\)

The most recent data indicate that, for those receiving between 1 and 24 infusions, the risk of PML is exceedingly low: 0.3 per 1,000 patients. For patients receiving between 25 and 36 infusions, the PML risk increases to approximately 1.5 per 1,000 patients. Although the number of cases receiving natalizumab for more than 36 infusions is more limited, the risk does not appear to increase further—the most recent estimated risk of PML for patients receiving 37 to 48 infusions is 0.9 per 1,000 patients.\(^{22}\) The use of immunosuppressive drugs, including antimetabolites (eg, azathioprine), alkylating agents (eg, cyclophosphamide), or anthracyclines (eg, mitoxantrone) prior to natalizumab exposure appears to be an independent risk factor for natalizumab-associated PML, raising risk by approximately two- to fourfold.

Of the first 35 cases of natalizumab-associated PML, 10 cases (29%) were fatal. Among surviving patients, the level of disability was found to be severe in 48%; moderate in 36%; and mild in 16%.\(^{23}\) Improved survival was associated with younger age, less MS-related disability prior to PML, more localized disease on brain MRI at diagnosis, and shorter time from symptom onset to PML diagnosis.\(^{23}\) As of August 4, 2011, there were 150 confirmed cases of natalizumab-associated PML (58 in the United States, 85 in Europe, and 7 from the rest of the world); of these, 29 (19%) have died.\(^{22}\)

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**VARIATION IN PATIENT RISK TOLERANCE**

The postmarketing surveillance of natalizumab clearly demonstrates that risk is associated with administration
of the drug, but risk tolerance varies considerably among individuals with MS. Some patients elect to use natalizumab despite the risk of PML, even when they have relatively mild MS. Other patients decline use of natalizumab even when their MS is severe and has responded poorly to other disease-modifying drugs.

In most cases, based on my experience, patients accept the risk of natalizumab-associated PML if MS disease is their primary consideration. Another major factor is the patient’s prior experiences with disease-modifying drugs; patients who have experienced breakthrough disease activity despite treatment with first-line drugs commonly opt for natalizumab regardless of the risk of PML.

Interestingly, the treating neurologist’s perception of MS severity and risk of PML may differ from the patient’s perception. In a study of 69 natalizumab-treated MS patients and 66 neurologists, Heesen et al found that patients had a significantly worse perception of their disease and were more willing to assume treatment risks and continue natalizumab therapy than their neurologists were.24 About one-half of the neurologists said that they would discontinue natalizumab at a risk level of 1 in 5,000 or lower, whereas only 17% of the patients would stop at this risk level. This finding has significant implications for clinical practice and implies that the neurologist should discuss concerns about MS and risk of treatment with the patient in order to tailor the decision to the patient’s concerns.

Interest in identifying biomarkers to aid in quantifying risk is ongoing. Chen et al found that subclinical reactivation of the JC virus (JCV) occurred frequently in 19 natalizumab-treated MS patients.25 Another study of 24 natalizumab-treated MS patients found no JCV DNA in the blood, although JCV DNA was found in the urine in 25% of patients.26 A large survey of blood and urine from natalizumab-treated MS patients found low sensitivity and specificity for JCV DNA as a predictor for subsequent PML.27 In this study of more than 1,000 natalizumab-treated patients, JCV DNA was detected in 0.3% of patients’ plasma and in 26% of patients’ urine, but PML did not develop in any patient who was JCV-positive. Among five natalizumab-treated patients who developed PML, JCV DNA was detected in none before the advent of symptoms. The presence of JCV DNA in bodily fluids is important for the diagnosis of PML, but it currently holds no predictive clinical value. At present, measuring JCV DNA in blood, cells, or urine as

![Figure 1. Progressive multifocal leukoencephalopathy (PML) incidence estimates by treatment duration (A) and treatment epoch (B), calculated based on natalizumab exposure through January 31, 2011, and 95 confirmed cases as of February 2, 2011. (A) The incidence for each time period is calculated as the number of PML cases divided by the number of patients exposed to natalizumab (eg, for ≥ 24 infusions, the number of PML cases diagnosed with exposure of 24 infusions or more is divided by the total number of patients exposed to at least 24 infusions). (B) The incidence for each epoch is calculated as the number of PML cases divided by the number of patients exposed to natalizumab (eg, for 25 to 36 infusions, the number of PML cases diagnosed during this period is divided by the total number of patients ever exposed to at least 25 infusions and therefore having risk of developing PML during this time). Source: Data on file. Biogen Idec; 2011.](image)
a predictive biomarker for natalizumab-associated PML does not appear to be clinically useful.

Stratifying risk by measuring JCV serology, however, does appear to be a useful strategy. Investigators using a two-step assay to detect and quantify JCV antibodies found 53.6% of MS patients to be seropositive, with a false-negative rate of 2.5%. Of most interest, all 17 natalizumab-associated PML patients who had available blood samples taken an average of 2 years before onset of PML tested positive for JCV antibodies.28 Although studies are ongoing, classification according to JCV-antibody status may be helpful in advising patients. Patients who are JCV-antibody seronegative (about one-half of patients) appear to be at extraordinarily low risk for PML. In these patients, use of natalizumab could be liberalized and continued as long as the JCV-antibody status remains negative. In patients who are seropositive for JCV antibodies, caution is recommended, particularly for patients who had prior immunosuppressive drug therapy and for patients who have received treatment for more than 24 months. Even in JCV-antibody–seropositive patients, use of natalizumab may be advisable depending on disease severity, available options, and the patient’s risk tolerance. JCV-antibody testing is a rare example of a clinically useful biomarker that can guide specific treatment decisions in the field of neurology.

**CURRENT PRACTICE: A PERSONAL MANAGEMENT ALGORITHM**

Based on current evidence, the following opinion on the use of natalizumab for the treatment of MS is offered as a supplement to approved prescribing information. The neurologist must individualize the treatment decision for each patient and recognize that these general comments represent a personal opinion. Several factors affect decisions about the use of disease-modifying drugs in MS, and specifically use of natalizumab: How severe is the disease, and what is the prognosis for future disease progression from the neurologist’s perspective? How concerned is the patient about current or future MS symptoms and disability? What is the patient’s tolerance for medication side effects? For risk taking? Has there been prior immunosuppressive therapy? What is the JCV antibody status? What other options are available for disease management?

These issues require discussion among the neurologist, the patient, and the patient’s family. The neurologist should provide input on disease status, an opinion about prognosis, and a description of appropriate options for disease management. Many patients also want a global recommendation (ie, “Tell me what you think I should do”). The neurologist must tailor that global recommendation to the patient’s perceptions of his or her MS, its treatment, and preferences regarding treatment options.

My current, evolving view of the management of relapsing-remitting MS is outlined in Figure 2. For treatment-naïve patients with active MS who are seronegative for JCV antibodies, I recommend that natalizumab be considered, with JCV serology repeated yearly (Figure 2A). For patients who are JCV-antibody–seronegative and have breakthrough disease, I also recommend natalizumab with JCV serology repeated yearly. If the patient is treatment-naïve and seropositive, I recommend a first-line drug or fingolimod. If the seropositive patient develops breakthrough disease, is risk tolerant, and has not been treated with prior immunosuppressive drugs, I advise switching to natalizumab for 1 to 2 years to determine response and then reassess.

For patients who are already receiving treatment with a first-line drug and whose disease is well controlled, I make no changes in treatment until a breakthrough occurs, at which time I recommend switching to natalizumab with JCV antibody status assessed yearly (Figure 2B). If a patient has been taking natalizumab for more than 2 years and is seronegative, I advise continuing natalizumab (Figure 2C). If seropositive after 2 or more years of natalizumab therapy, I recommend switching to fingolimod and monitoring for disease reactivation.

**DISCUSSION**

**Dr. Calabrese:** Have you perceived growing concern over PML among the MS patient population over the past 2 to 3 years?

**Dr. Rudick:** I would say that it’s pretty stable. Patients who are risk intolerant select out of natalizumab. Some patients would just as soon take their chances with MS rather than deal with additional risk. Since we talk about the risk of PML with patients prior to treatment, the patients who choose natalizumab are able to deal with the risk. The difficult cases are those patients who will be severely disabled before long but choose not to go on natalizumab because they’re very risk averse. It gets even more complicated when the closest family member (parents or spouse) want their relative to use natalizumab but the patient is risk averse. This can become quite complicated—for example, I’ve seen situations where the patient is a minor, and one parent wants their child to use natalizumab but the other is risk-averse. Spouses often see risk differently, and this has led to interesting and difficult discussions. In the case of a child with MS, I listen to preferences from family members, but otherwise I empower the patient to drive the decision.

**Dr. Major:** Our data seem to suggest a higher percentage of individuals who are seropositive—about 56%. The issue, however, is a lack of a standard to define seropositive and seronegative. I suggested that the natalizumab manufacturer collect a bank of samples and
allocate them to laboratories with no vested interest for polymerase chain reaction (PCR) assays, from which consensus definitions of seropositive and seronegative could be developed.

**Dr. Calabrese:** Why is there no confirmatory immunoassay for this virus? We don’t have false positives for hepatitis B or human immunodeficiency virus. We still seem to be relying on older technologies.

**Dr. Major:** To determine the level of antibody, an enzyme-linked immunosorbent assay is just fine.

**Dr. Calabrese:** That’s for sensitivity, but what about specificity?

**Dr. Major:** Specificity is quite good. Everybody now uses the same antigen, the polyoma capsid antigen, VP1, to detect productive viral infection, but there are no set standards for the cutpoints to classify as seropositive and seronegative. Certainly, there’s high sensitivity with PCR to detect JCV DNA in cerebrospinal fluid (CSF), because we’re able to detect very low copy levels of JCV DNA in the CSF.

**Dr. Simpson:** I would like to see a quantitative measure of risk versus benefit for all of the drugs used in MS, not just natalizumab. When you look at any clinical trial, you see a table of adverse events and you see efficacy measures, but you don’t see the two combined. This really is necessary to compare drug A with drug B. Instead, we end up making decisions based on risk tolerance and rather soft criteria. One could argue that we don’t want to be so algorithmic that we take the art out of medicine, but the criteria we use to make decisions are quite soft. I wonder whether you have any suggestions on a more quantitative approach.

**Dr. Rudick:** This is an important point, but a difficult problem. We have much more information about PML associated with natalizumab than we do about many seri-

![Diagram](image-url)

**FIGURE 2.** (A) Decision-making algorithm for patients with active multiple sclerosis (MS) who are treatment-naïve, (B) for patients with MS who are already taking a first-line drug, and (C) for patients who have been receiving natalizumab for more than 2 years. JCV = JC virus.
ous adverse events. For example, there are rare adverse events with interferon beta—severe depression, liver injury, and so forth. But we don’t have precise quantitative data on most adverse drug effects, and in general adverse events are underreported in clinical practice.

The natalizumab-PML situation is somewhat unique. PML is a dramatic, often fatal, disease that is virtually never observed spontaneously in MS, and the strict reporting requirements have resulted in near-complete ascertainment and more precise risk estimates. This situation doesn’t apply to most adverse events associated with other therapies—even for some severe adverse events. But you are correct—focusing exclusively on the risk of PML seems somewhat simplistic because there are clear risks with other drugs, and these need to be factored into treatment decisions.

Dr. Mollov: Do you have good tools that predict how a patient with MS will do over time?

Dr. Fox: We have fair tools, not great tools.

Dr. Rudick: We’re diagnosing MS earlier, sometimes at the first symptom. We’re even beginning to recognize it in patients without symptoms who have MS observed as an incidental MRI finding. It is difficult at the earliest stage of MS to predict severity with any confidence. The best predictor we have is the severity of the disease by MRI criteria. This can provide a general guide to treatment decisions, but it is an imprecise predictor.

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HIV-associated PML: Changing epidemiology and clinical approach

ABSTRACT

Despite the availability of highly active antiretroviral therapy (HAART) for the treatment of human immunodeficiency virus (HIV) infection, there has not been a dramatic decrease in the frequency of progressive multifocal leukoencephalopathy (PML) in the HIV-infected population. Usually a multifocal progressive disease of nonenhancing lesions in white matter, PML can have distinct characteristics in HIV-infected patients, including unifocal static lesions of faint contrast enhancement on imaging and involvement of gray matter. A syndrome of cerebellar degeneration has been described in association with HIV infection in patients positive for JC virus, the papovavirus responsible for PML. The standard of care for HIV-associated PML is HAART to achieve immunologic recovery and optimal HIV virologic control. The prognosis of PML has improved greatly since the advent of HAART.

The appearance of progressive multifocal leukoencephalopathy (PML) as a complication of human immunodeficiency virus (HIV) infection dates to shortly after the first description of acquired immunodeficiency syndrome (AIDS). The advent of highly active antiretroviral therapy (HAART) dramatically altered the nature of HIV infection, resulting in a substantial decline in mortality and, in essence, turning AIDS into a chronic disease. As patients lived longer with HIV infection, one consequence was an increased incidence of neurologic complications. By the early 1980s, AIDS was well recognized as an underlying disorder that predisposed to PML.

As many as 70% of HIV patients will eventually have involvement of either the peripheral or central nervous system (CNS). Most patients with HIV are managed by primary care clinicians, including those in the fields of family practice, internal medicine, or infectious disease, and the complexity of the neurologic disorders associated with HIV often results in either delayed diagnosis or misdiagnosis. For example, the evolution of HIV in the plasma, where most clinicians measure it, may differ from its evolution in the spinal fluid and brain. An emerging issue is that of hepatitis C coinfection, which may itself be associated with central and peripheral neurologic complications.

Treatment of HIV with antiretroviral agents has numerous neurologic implications. These include the potential ability of these agents to penetrate the blood-brain barrier, their efficacy in both treating and preventing cognitive impairment and other CNS disorders, and their toxic effects in the CNS and peripheral nervous system.

NEUROLOGIC COMPLICATIONS OF AIDS

Neurologic disease in AIDS patients can be classified in several ways. One of the most logical, particularly for primary care clinicians, is the separation of primary from secondary neurologic disorders:

- Primary neurologic disorders are enigmatic and difficult to characterize; they include HIV-associated neurocognitive disorders in adults, encephalopathy in children, myelopathy or spinal cord disease, and peripheral neuropathy.
- Secondary complications are related to progressive immunosuppression. These include opportunistic infections such as cytomegalovirus, toxoplasmosis, or cryptococcal meningitis; and neoplasms such as primary CNS lymphoma. Opportunistic infections and neoplasms have declined in incidence in the HAART era.

The prevalence of neuropathy as a neurologic complication of HIV/AIDS is as high as 57%, while PML affects 5% or fewer of HIV-infected patients (Table). PML in these patients is often associated with advanced disease, as reflected by low CD4+ cell counts and increased plasma HIV viral loads.

AT-RISK POOLS FOR PML

The AIDS epidemic significantly changed the epidemiology of PML, turning a formerly rare disease into a much more common one. In South Florida, the incidence of PML in patients with AIDS increased by 12 times from...
the 5-year period 1981 to 1984 compared with 1991 to 1994. Only two non-AIDS cases of PML were reported in South Florida during this 15-year period. At present, nonimmunosuppressed, healthy individuals account for fewer than 1% of all cases of PML. Non-HIV–related PML represents 10% to 20% of all PML cases. Cancer survivors and patients with rheumatoid arthritis who are treated with immunotherapy constitute the largest at-risk pools among this group. PML related to HIV represents 80% to 90% of PML cases, drawing from a pool of 1.2 million HIV-infected individuals in the United States.

## UNIQUE PRESENTATION OF HIV-ASSOCIATED PML

The brain lesion in PML is classically a nonenhancing focal lesion, preferentially in white matter, but lesion characteristics often depart from this characteristic picture. For example, relatively faint contrast enhancement of lesions on magnetic resonance imaging has been observed, as well as involvement of white matter and gray matter. The distribution and character of brain lesions in PML may also differ from the classic picture. For example, the lesion may not be focal, particularly when PML is combined with the symmetric white matter abnormalities that are seen in HIV encephalopathy; this nonclassic presentation can cause difficulty in radiologic differentiation of PML and HIV encephalopathy.

### Cerebellar degeneration

A unique presentation of PML is possible in HIV-infected patients. In 1998, Tagliati et al2 described a syndrome of degeneration of the cerebellum in 10 HIV-infected patients. One patient had JC virus (JCV) detected by polymerase chain reaction (PCR) in cerebellar biopsy tissue. The authors proposed the possibility of latent JCV infection of cerebellar granular cells in HIV-infected patients with cerebellar atrophy, lacking further evidence of other features of PML.

In 2005, Koralnik et al3 described a case of JCV granule neuronopathy in a patient with JCV-associated cerebellar degeneration. Clinical symptoms included gait and limb ataxia, dysmetria, dysarthria, and nystagmus. Magnetic resonance imaging demonstrated diffuse cerebellar atrophy and occasional subtle white matter abnormalities within the cerebellum (Figure). Immunohistochemistry showed preservation of Purkinje cells, and in situ PCR revealed selective depletion of cerebellar granule cells and JCV infection of granule cell neurons.

## MANAGEMENT OF HIV-ASSOCIATED PML

### Optimize HAART

A suppressed plasma HIV viral load is the strongest prognostic factor for an improved course in PML. In the pre-HAART era, the mean survival of HIV-associated PML was 3 to 6 months, with long-term survival estimated at 10%. The use of HAART has achieved a dramatic improvement in long-term survival, to upwards of 50%. Neurologic deficits are often irreversible even with HAART, but most HAART recipients show stability in neurologic status for years.

Other key characteristics associated with improved survival in HIV-associated PML appear to be younger age, PML as the heralding manifestation of AIDS, initiation of HAART upon diagnosis of PML, higher CD4 count, and absence of severe neurologic impairment.

### Investigational therapies

Specific antiviral drug regimens targeting JCV have been tested empirically in case studies and in clinical trials in patients with AIDS- and non–AIDS-related PML. Cytosine arabinoside (Ara-C). Ara-C is a nucleoside analog used as an antineoplastic agent; it terminates chain elongation and inhibits DNA polymerase to confer antiviral activity. Ara-C decreased JCV replication in vitro. Based on anecdotal reports of efficacy in cancer-related cases of PML, Ara-C was tested in a multicenter trial of 57 patients with HIV and biopsy-confirmed PML. Neither intravenous nor intrathecal Ara-C combined with established antiviral therapy for AIDS improved the prognosis of these patients, and Ara-C has since been abandoned as a strategy to treat HIV-related PML.

Cidofovir. The noncyclic nucleoside phosphonate cidofovir garnered attention as a potential treatment for
PML ASSOCIATED WITH HIV

PML based on case reports of efficacy in HIV as well as non-HIV patients. Subsequently, a large multicenter study failed to detect any significant added benefit with cidofovir beyond that of HAART.\(^\text{11}\) Retrospective European studies confirmed the lack of clinical benefit with cidofovir.\(^\text{6,7,12}\)

**Interferon alfa.** Case reports with interferon alfa-2a and -2b for the treatment of PML show conflicting results with respect to clinical response, symptomatic improvement, and survival, but toxicity has been substantial. In a series of 97 patients with AIDS-related PML, Geschwind et al determined that interferon alfa had no effect on survival beyond that of HAART.\(^\text{13}\)

**Mirtazapine.** Serotonin receptor antagonists such as mirtazapine can block JCV entry into glial cells via serotonin 5-hydroxytryptamine receptors, providing a rationale for their use as a potential treatment for PML. Verma et al describe a case of clinical improvement (stable neurologic deficit) and PML lesion regression in a 63-year-old bedbound woman with polycythemia vera with biopsy-proven non–HIV-related PML that had progressed to quadriparesis.\(^\text{14}\)

**Mefloquine.** The antimalarial drug mefloquine inhibits viral replication in cultured human glial cells and astrocytes, inhibits JC viral DNA replication, and showed efficacy against two JCV strains in cell culture.\(^\text{15}\) A randomized study to assess the effectiveness of mefloquine for treatment of PML has been completed and its results await publication.

**SUMMARY**
The incidence of PML has remained unchanged from the pre-HAART to the HAART era, but the prognosis is greatly improved. The clinical presentation of PML in AIDS patients may deviate from the classic triad of progressive, multifocal, white matter disease. It may be static and unifocal, and it may involve gray matter and neurons as well as white matter. The number of neurologic manifestations is vast and can include the cerebellar syndrome. Lumbar puncture with a PCR negative for JCV does not confirm the absence of PML.

The standard of care for HIV-associated PML is HAART, with the goal of achieving immunologic recovery and optimal virologic control. Whether therapeutic results obtained in patients with HIV-associated PML can be translated to the setting of non–HIV-associated PML is unclear.

**DISCUSSION**

Dr. Simpson: As a followup to the Ara-C trial that was published,\(^\text{10}\) PML confirmed by brain biopsy was one of the enrolling criteria, and the planned study population was 65 patients. Longitudinal examination of viral load in cerebrospinal fluid (CSF) was a part of the study, and we found that the lower the viral load, the better the prognosis. Fifty-two patients were enrolled before the trial was stopped because it was clear that Ara-C was not producing a benefit. The patients had multifocal disease but, because Ara-C does not effectively cross the blood-brain barrier, penetration in the brain was minimal even with the use of an intrathecal shunt in this study.

Dr. Major: Do you think viral load in CSF is a predictor of disease severity and outcome in PML?

Dr. Rudick: Generally speaking, that’s probably true. We have found, as have many of our colleagues who run a lot of CSF samples, that high viral loads are not a good thing.

Dr. Bennett: How is it that the incidence of PML has not changed from the pre-HAART to the post-HAART era? How do you account for this in terms of the change in patients’ T-cell function from pre- to post-HAART?

Dr. Simpson: I don’t know. Intuitively, why do patients treated with HAART, who are relatively immune reconstituted, develop PML? The problem is that not everyone is immune reconstituted. HAART fails in some patients. Further, PML remains a disease that is
more common in late-stage HIV among patients with low CD4 counts and high viral loads, meaning that a large population of patients is available to develop this disease. With that said, it is perplexing that the incidence has not gone down more than it has.

**Dr. Major:** There’s a phenomenon called “unmasking PML with HAART,” in which individuals have no signs of PML upon initiation of HAART, but then very shortly after, PML is diagnosed.

**Dr. Berger:** You’re talking about PML immune reconstitution inflammatory syndrome (IRIS).

**Dr. Major:** IRIS can occur before PML, or PML and IRIS can be concurrent. In some patients, once the infection starts, it persists; this suggests that the virus is carried to the brain through the infected lymphocyte populations and may explain why the incidence of PML has not changed from the pre-HAART to the HAART era.

**Dr. Calabrese:** In patients with HIV who develop PML within the first 6 months of HAART, are we seeing the IRIS phenomenon or is it a presenting sign of advanced HIV?

**Dr. Simpson:** It’s well known that a number of opportunistic infections can develop in the setting of HAART. In fact, whether one should delay HAART when initiating therapy for opportunistic infections has been debated for just this reason. Most people presume IRIS to be a massive immunologic hit to all organ systems, as CD4 counts rise dramatically to produce hyperimmune-mediated phenomena such as Guillain-Barré syndrome. To what extent immunologic recovery is or is not linked to PML and why it happens are fascinating questions.

**Dr. Berger:** Opportunistic infections, PML among them, that occur following the initiation of HAART and recovery of the immune system are almost always an IRIS-mediated phenomenon in which the disease has been smoldering and then surfaces because of the release of an inflammatory response.

**Dr. Calabrese:** In patients with cerebellar degeneration, do you typically detect JCV in PCR in the spinal fluid?

**Dr. Simpson:** Not in the early stages, but in some patients with later-stage disease, the answer is yes. Certainly, PCR of CSF samples to look for JCV is the diagnostic test of choice. But in the early days, when we had no idea what caused this cerebellar syndrome, we were doing cerebellar biopsies.

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PML and rheumatology: The contribution of disease and drugs

ABSTRACT
Progressive multifocal leukoencephalopathy (PML), a rare, typically fatal, opportunistic infection caused by the JC virus, is becoming relevant to physicians in multiple specialties, including those who prescribe biologic agents for the treatment of autoimmune disorders. Reports of PML have led to US Food and Drug Administration alerts and warning letters regarding four immunosuppressive agents in recent years (natalizumab, rituximab, efalizumab, and mycophenolate mofetil). Consequently, informed clinical decision-making requires understanding the risk of PML associated with these therapies. An estimate of the relative frequency of PML associated with specific rheumatic conditions has been generated. Systemic lupus erythematosus appears to be associated with susceptibility to PML that cannot be fully explained by the intensity of immunosuppressive therapy. Further, the use of rituximab in patients with rheumatic disease has raised concerns. However, definitive attribution of cause is precluded by the limitations of the currently available data. All patients with rheumatic disease, regardless of the intensity of their current immunosuppressive therapy, should be considered potentially at risk of PML. With an evolving understanding of a greater clinical heterogeneity of PML, advances in diagnostic methods, and significant implications for therapy, PML should be considered in the differential diagnosis of neurologic manifestations of rheumatic diseases.

PML and rheumatology: The contribution of disease and drugs

Progressive multifocal leukoencephalopathy (PML) is a rare, typically fatal, opportunistic infection caused by the JC virus (JCV). Formerly an example of neurologic arcana, PML became an important clinical concern when it developed in patients with human immunodeficiency virus (HIV) infection. More recently, PML has attracted the attention of rheumatologists following reports of its being associated with the use of targeted therapies such as natalizumab and rituximab. A recent survey of rheumatologists’ knowledge of and attitudes towards PML revealed that concerns over PML affect decisions on the use of biologic agents. Further, rheumatologists have important real and perceived learning gaps regarding PML; for example, 41% of those surveyed could not identify the diagnostic test of choice for PML.

PML IN RHEUMATIC DISEASES
The US Food and Drug Administration (FDA) issued an alert in December 2006 following documentation of two fatal cases of PML in patients with systemic lupus erythematosus (SLE), both of whom had been treated with rituximab. We subsequently performed a literature search to identify cases of PML associated with rheumatic diseases. Patients were included only if the information provided was sufficient to substantiate the diagnosis of PML and the rheumatic disease in question; patients were excluded if they had HIV or cancer or had undergone organ transplantation. The search revealed 50 patients with rheumatic diseases who had PML (Table 1); SLE was overrepresented (n = 32) despite a much lower population prevalence of SLE compared with rheumatoid arthritis.

Examination of the immunosuppressive therapies prescribed to these patients within 6 months of the onset of neurologic symptoms attributed to PML revealed that low-dose (≤ 15 mg/d) prednisone, with or without an antimalarial agent, was the only immunosuppressive therapy in 31% of patients with SLE and in 11% of patients with rheumatic diseases other than SLE. Three patients had no documented immunosuppressive therapy in the 6 months prior to the onset of PML. Two patients with SLE were prescribed rituximab; no cases were reported in association with biologic therapies other than rituximab.

In order to circumvent reporting bias, a nationwide hospital discharge database representing nearly 300
million patient discharges was used to determine the relative frequency of PML in patients with rheumatic diseases. Because of the reliance on diagnostic coding, rheumatic diseases were likely underreported in this sample; information on therapies was unavailable. After excluding patients who had HIV or cancer or were organ transplant recipients, four cases of PML were identified per 100,000 SLE discharges. This rate was 10-fold higher than the rate associated with rheumatoid arthritis and 20-fold higher than that of the background population.

These data show that PML is a rare occurrence in patients with rheumatic diseases, and SLE appears to be associated with a predisposition to PML. This predisposition in patients with SLE does not appear to be proportional to the degree of iatrogenic immunosuppression, emphasizing the role of host factors.

**DISEASE-MODIFYING DRUGS AND PML RISK**

In addition to certain disease states, disease-modifying biologic drugs have recently been associated with rare instances of PML.

**Rituximab**

The first case of rituximab-associated PML in the setting of rheumatoid arthritis was recorded in September 2008. The patient had longstanding rheumatoid arthritis and Sjögren syndrome. She received four courses of rituximab and was diagnosed with PML 18 months after the last dose; she died 1 month later. Her therapy for rheumatoid arthritis included a tumor necrosis factor (TNF) antagonist prior to rituximab initiation and treatment with methotrexate and steroids before, during, and after rituximab therapy. Oropharyngeal cancer developed in this patient 9 months prior to the onset of PML and was treated with chemotherapy and radiation therapy.

Another case of PML in a patient with rheumatic disease who had been treated with rituximab was notable because it was the first in which the patient had not previously been treated with an anti-TNF agent.

Ascertaining cause of PML in patients treated with rituximab is difficult because the potential pathogenic mechanism remains unknown. Humoral immunity is not protective against PML, leading to speculation that the loss of other B-cell functions, such as those of antigen-presenting cells or cytokine production, may lead to a defect in cell-mediated immunity. Another theory posits that reconstitution of naïve B cells with latent JCV infection following B-cell depletion from rituximab therapy may somehow facilitate the development of PML.

**Efalizumab**

Efalizumab is a monoclonal antibody that targets CD11a, the alpha subunit of lymphocyte function-associated antigen 1. Efalizumab blocks binding to intercellular adhesion molecule 1, and thereby blocks T-cell adhesion and migration. CD11a is also expressed on a variety of other leukocytes and lymphocytes such as B cells, monocytes, and natural killer cells.

Efalizumab was approved in 2003 by the FDA for the treatment of moderate to severe plaque psoriasis. It is estimated that 46,000 patients have been treated with efalizumab worldwide since its approval. In 2008, a black box warning was added to the efalizumab prescribing information following the occurrence of serious infections, including pulmonary tuberculosis, necrotizing fasciitis, and invasive fungal infections. Subsequently, four cases of PML, three of which were fatal, were reported in psoriasis patients treated with efalizumab. Of note, these were the first cases of PML reported in patients with psoriasis. Of more concern, the affected patients were among a group of approximately 1,100 patients who had been treated with efalizumab for more than 3 years. In February 2009, a public health advisory was issued by the FDA, and efalizumab was voluntarily withdrawn by its manufacturer 2 months later.

**Belatacept**

Belatacept is a recombinant soluble fusion protein of the extracellular domain of human cytotoxic T-lymphocyte antigen-4 with a fragment of a modified Fc domain of immunoglobulin G1. Recently approved by the FDA for prophylaxis of renal transplant rejection, it is a second-generation, higher-avidity version of abatacept. Abatacept is licensed for the treatment of rheumatoid arthritis and is under investigation for the treatment of vasculitis and SLE. Belatacept differs from abatacept by only two amino acids.

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**TABLE 1**

PML associated with rheumatic disease

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<thead>
<tr>
<th>Rheumatic disease</th>
<th>No. (%) patients with PML</th>
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<tr>
<td>Systemic lupus erythematosus</td>
<td>32 (64)</td>
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<tr>
<td>Idiopathic inflammatory myositis</td>
<td>6 (12)</td>
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<td>(5 dermatomyositis/1 polymyositis)</td>
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<td>5 (10)</td>
</tr>
<tr>
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<td></td>
</tr>
<tr>
<td>Wegener granulomatosis</td>
<td>4 (8)</td>
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<tr>
<td>Other†</td>
<td>3 (6)</td>
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</table>

*Confirmed number of diagnoses of PML in patients with rheumatic disease reported in the English-language medical literature through April 2009.
†Sjögren’s syndrome and CD4 lymphopenia (n = 1), localized scleroderma and amyloidosis (n = 1), and destructive polyarthritis (+ANA, Jo1) with CD4 lymphopenia (n = 1)

PML = progressive multifocal leukoencephalopathy
Two cases of PML have been reported in association with belatacept, one in a patient following renal transplantation and the other in a patient following liver transplantation. Both patients had been treated with other standard immunosuppressive therapies for prophylaxis of organ transplant rejection, including mycophenolate mofetil.

Mycophenolate mofetil
Mycophenolate mofetil is the prodrug of mycophenolic acid. Both have been the subjects of FDA alerts regarding PML, based on a January 2008 report of 10 definite and 7 possible cases of PML occurring with mycophenolate mofetil. The patients affected included four with SLE, none of whom underwent a renal transplant.

In a retrospective cohort study of 32,757 renal transplant patients, Neff et al found 14 cases of PML per 100,000 person-years among patients treated with mycophenolate mofetil following kidney transplant compared with none in patients who did not receive mycophenolate mofetil. It is difficult to ascertain risk with mycophenolate mofetil because it is standard therapy among renal transplant patients, leaving few patients in these groups unexposed.

Given the FDA alert with respect to mycophenolate mofetil and PML, the frequent use of mycophenolate mofetil in the setting of SLE, and the concerns about possible predisposition to PML among patients with SLE, it will be important to clarify the level of risk in patients with SLE who are treated with mycophenolate mofetil.

## AGGREGATE EXPERIENCE: REVIEW OF FEDERAL DATABASE

We examined the aggregate experience of PML in association with autoimmune disorders and biologic disease-modifying antirheumatic drug (DMARD) exposures reported in the FDA Adverse Event Reporting System (AERS) database. A total of 19 confirmed cases of PML in patients with rheumatic diseases were uncovered: 10 in patients with SLE, 5 in patients with rheumatoid arthritis, 3 in patients with vasculitis, and 1 in a patient with dermatomyositis. The patients with PML included six who received rituximab for the management of rheumatic diseases (Table 2). In all six patients, rituximab was the most recently prescribed DMARD. Four cases were identified in patients treated with anti-TNF therapy, but three of these had received anti-TNF therapy prior to rituximab, and the other was receiving concomitant cyclophosphamide for rheumatoid vasculitis.

Ten cases of PML were confirmed with cyclophosphamide treatment, and cyclophosphamide was the most recent DMARD prescribed in two of these cases. Five cases were confirmed with mycophenolate mofetil (in four of which it was the most recently prescribed DMARD) and six with azathioprine (in three of which it was the most recently prescribed DMARD).

### Risk of PML with DMARD therapy

**Rituximab.** The confirmation of six cases of PML among rituximab-treated rheumatoid arthritis patients is a source of concern. Nevertheless, PML is a rare adverse event. It occurs in fewer than 1 in 10,000 rituximab-treated patients who have rheumatoid arthritis, among a total of approximately 130,000 such patients. A better understanding of the potential mechanism responsible for the increased risk of developing PML may help in risk prediction and to guide patient selection for this agent.

**Anti-TNF therapy.** A paucity of confirmed cases in patients treated with anti-TNF therapy argues against a significant risk of PML associated with this therapy, especially considering the estimated 2 to 3 million rheumatoid arthritis patients who are receiving treatment with anti-TNF agents. A note of caution is sounded by a recent case report of PML in a rheumatoid arthritis patient. The patient had been treated with infliximab, with the only background therapy being methotrexate. Ongoing vigilance is therefore necessary.

**Mycophenolate mofetil.** All five confirmed cases of PML in mycophenolate mofetil-treated patients had earlier received treatment with cyclophosphamide. These data indicate no clear signal of excess risk with mycophenolate mofetil above that seen with other non-biologic immunosuppressive agents, such as cyclophosphamide or azathioprine.

### CONCLUSION AND RECOMMENDATIONS

PML has been reported in association with a variety of disease states, although a predisposition in patients with SLE has become apparent. Synthetic and biologic immunosuppressive therapies have also been impli-
DISCUSSION

Dr. Simpson: To what extent are these lesions in the brain being attributed to the underlying vasculitis, particularly in SLE, as opposed to pursuing a PML diagnosis, and how might this result in dramatic underreporting of the complication?

Dr. Molloy: We found that PML is almost certainly underdiagnosed, particularly in SLE patients. If a patient succumbs to assumed neuropsychiatric SLE, how often is an autopsy undertaken? One telling paper from Sweden documented four cases of PML in SLE patients. In one of these, the diagnosis was made retrospectively from autopsy tissue that had been banked 20 years previously. It undoubtedly is underdiagnosed.

Dr. Calabrese: Even in the most recent rituximab-associated cases of PML, several patients were empirically given additional immunosuppressive therapy because it was presumed that they had a comorbid neuropsychiatric rheumatic complication. The presence of neuropsychiatric complications ascribed to an autoimmune inflammatory disease generally warrants escalation of immunosuppressive therapy. It has always been standard practice for us to rule out opportunistic infection, but JCV infection has not been on the radar screen until very recently.

Dr. Molloy: I’d like to emphasize that, in our literature review, 50% of the rheumatic disease patients diagnosed with PML had been treated with more intensive immunosuppressive therapy. It was only after they continued to deteriorate that JCV infection was suspected and PML ultimately diagnosed.

Dr. Berger: Is it fair to say that the incidence of PML in SLE is about 10 times that in rheumatoid arthritis?

Dr. Molloy: In the hospital discharge database, it was 10-fold higher in SLE than in rheumatoid arthritis, but we can’t draw a conclusion from the AERS database because we don’t have a denominator. The database consists of voluntary submission of cases.

Dr. Calabrese: The information that we can expect to glean from the database is profoundly limited, for all the reasons that you enumerated. Despite the flaws, we’re obligated to continuously examine it because sometimes a case or two may provide some special insight.

Dr. Simpson: As neurologists, we often lag behind rheumatologists in the use of new treatments, including intravenous immune globulin (IVIG) and now rituximab. Rituximab is becoming the go-to drug for a number of neurologic diseases. I’m using it quite a bit and have observed some dramatic responses in patients with chronic inflammatory demyelinating polyneuropathy, for example, in whom IVIG or plasmapheresis was failing. Anecdotally, some of the turnarounds in polymyositis and even myasthenia gravis are remarkable as well. I’m not sure to what extent neurologists—particularly peripheral neurologists—who use rituximab are recognizing PML.

Dr. Fox: The index of suspicion is probably vastly different among multiple sclerosis (MS) specialists and general neurologists. Neurologists who treat MS will be acutely aware of PML because of its association with natalizumab.

Dr. Berger: Yes, but you’re talking about possibly two orders of magnitude difference between natalizumab and rituximab. In fact, PML is rarely reported in the setting of neurologic disease. It’s mostly reported in the setting of rheumatologic disease.

Dr. Rudick: I don’t necessarily agree with you. Ascertaining the true incidence of PML with agents other than natalizumab is difficult. One is unlikely to miss a case of PML in an MS patient treated with natalizumab, but most cases stemming from the use of these other disease-modifying drugs are probably being missed.
Dr. Calabrese: I get two messages out of this body of work. Number one is that while PML is rare, it is seen across the spectrum of immunosuppressive agents, including biologic and nonbiologic drugs. Number two is that PML is seriously underreported and underrecognized, which is probably leading to suboptimal patient care. Rituximab was recently approved for treatment of Wegener granulomatosis, and this disease is heavily pretreated with cyclophosphamide. You would expect that PML is on the radar among clinicians caring for patients whose diseases warrant the use of increasingly complex, potent, and novel immunosuppressives.

Dr. Berger: There is one other biologic agent you left out—alemtuzumab. It wipes out all of the B cells and T cells; the B cells repopulate but the T cells remain suppressed for a long period. If ever there was a drug whose action mirrors what happens in HIV, alemtuzumab is that drug. Yet, PML is rarely seen with alemtuzumab. Alemtuzumab-associated PML has not been reported in the MS population, and it has only been seen in two transplantations that I’m aware of. I’m not saying that it doesn’t occur, but we’re not seeing it with the same frequency that one would predict given its profile.

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ABSTRACT
Progressive multifocal leukoencephalopathy (PML), a rare opportunistic infection of the central nervous system, occurs mainly in the setting of broad-based and selective immunosuppression. The immunomodulatory agent most often implicated in the development of PML is the monoclonal antibody natalizumab. Management of PML begins with risk stratification. Factors that predict the risk of PML are JC virus (JCV) antibody status, history of chemotherapy use, and cumulative exposure to natalizumab. The risk of natalizumab-related PML increases up to a duration of 36 months of therapy, after which the risk appears to level off. If suspicious for PML, the use of a sensitive JCV polymerase chain reaction assay permits early diagnosis. Immune reconstitution represents the mainstay of treatment for PML. With rapid reversal of immunosuppression followed by immunologic recovery, almost all patients suffer clinical deterioration termed immune reconstitution inflammatory syndrome (IRIS). High-dose corticosteroids are often recommended if a clinical and imaging syndrome resembling IRIS develops after immune restoration.

APPROACH TO PML TREATMENT
The ideal approach to PML treatment is generally two-pronged: antiviral therapies to directly reduce viral replication and immune reconstitution that empowers the immune system to attack the JC virus (JCV). Challenges to treatment are the difficulty in culturing JCV for in vitro studies, lack of an animal model of PML, and infrequency of PML cases.

Antiviral therapies
At present, no antiviral agent has confirmed efficacy in PML. Nucleoside analogues, serotonin 5-hydroxytryptamine receptor antagonists (to block the JCV receptor), and several cytokines provided exciting prospects in preclinical studies for treatment of PML in humans. Unfortunately, subsequent clinical studies of cytarabine, cidofovir, and interferon alfa all yielded disappointing results. A derivative of cidofovir, CMX001, is also being evaluated for efficacy in PML. Mefloquine was identified through a broad pharmaceutical screening study to have strong antiviral effects in vitro, but a clinical trial to assess its effects was stopped. It remains unclear whether the failure of clinical studies after successful in vitro studies is secondary to low drug penetration into the CNS, treatment initiation too late in the course of PML, or other differences not yet fully understood.

Immune reconstitution
Given the widespread failure of antiviral regimens, the mainstay of PML treatment is immune reconstitution. When immunosuppression is secondary to a medical disorder, efforts are pursued to reverse the primary disorder. For example, highly active antiretroviral therapy significantly prolongs survival in antiretroviral-naive acquired immunodeficiency syndrome patients. Decreasing the

Progressive multifocal leukoencephalopathy (PML) is a rare opportunistic infection of the central nervous system (CNS). Although originally associated with broad-based immunosuppression (human immunodeficiency virus infection, lymphoproliferative disorders, and immunosuppressive medications), recognition of PML in patients with selective immunosuppression is growing. This restricted immunodeprivation can arise from autoimmune disorders such as systemic lupus erythematosus, selective immunosuppressive therapies (eg, rituximab, leflunomide, and efalizumab), or immunosuppression limited to the CNS (eg, treatment with natalizumab).

This article reviews approaches to the management of PML, with specific recommendations regarding PML associated with natalizumab therapy.

Dr. Fox reported that he has received consulting fees from Avanir, Biogen Idec, EMD Serono, and Novartis. In addition, research support and consultant and advisory committee fees from Biogen Idec were paid to Cleveland Clinic.

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intensity of immunosuppressive therapy in solid organ transplant may improve survival with PML. When PML is associated with biologic therapies for autoimmune diseases, early diagnosis and immediate suspension of therapy is thought to improve outcomes.

**EXPERIENCE WITH NATALIZUMAB**

PML in the setting of natalizumab therapy is related to cumulative exposure to natalizumab. As of August 4, 2011, there had been 150 cases of natalizumab-related PML documented in more than 88,000 patients exposed to natalizumab worldwide (see page S18, “Multiple sclerosis, natalizumab, and PML: Helping patients decide”). The incidence of PML in natalizumab-treated patients varies according to the number of infusions received, but the incidence of PML by each epoch of treatment exposure (1 to 24 infusions, 25 to 36 infusions, 37 to 48 infusions) appears to have remained stable over time.

The mortality associated with natalizumab-related PML was 19% (29 deaths among the 150 confirmed cases) as of August 4, 2011. In cases with at least at least 6 months of follow-up, mortality has remained at about 20%. Many who survived were left with serious morbidity and permanent disability, although interpretation of disability is difficult because functional impairment is a hallmark of multiple sclerosis (MS) irrespective of PML. Survival in patients with natalizumab-associated PML appears to be better than with PML associated with other conditions, possibly because of early diagnosis achieved through clinical vigilance and swift immune reconstitution through natalizumab discontinuation and either plasmapheresis or immunoabsorption. Predictors of survival include younger age at diagnosis, less disability prior to onset of PML, more localized disease on magnetic resonance imaging (MRI) of the brain, and shorter time from symptom onset to PML diagnosis.

**Clinical characteristics of natalizumab-associated PML**

Several clinical observations should increase suspicion of natalizumab-associated PML. For example, the most common presenting symptoms are cognitive, motor, language, and visual impairment. Gadolinium-enhancing lesions are observed at presentation in about one-half of patients. Seizures and paroxysmal events can occur at presentation, which helps to differentiate PML from an MS relapse.

Approximately one-half of patients with natalizumab-associated PML have an initial viral load of less than 500 copies/mL, underscoring the need for ultrasensitive polymerase chain reaction (PCR) testing. An ultrasensitive JCV assay (Focus Diagnostics, Cypress, California) is available that can detect less than 50 copies/mL of JCV DNA. Because the viral copy numbers in the cerebrospinal fluid (CSF) may be low in patients treated with natalizumab, the CSF PCR may be falsely negative. In several cases of PML, JCV was undetectable in the CSF by PCR, identified only later by repeat PCR or brain biopsy. Serum JCV PCR is not useful in the screening or diagnosis of PML.

Natalizumab-associated PML has not been observed with therapy of 6 months’ or less duration. After 6 months of natalizumab therapy, new MRI lesions are rare in patients who are negative for neutralizing antibodies. A new MRI lesion in such a patient should be considered suspicious for PML. Our standard protocol is to check for neutralizing antibodies at 6 months in all patients treated with natalizumab. Symptoms of PML develop in affected patients whose duration of therapy ranges from 6 to 81 infusions. Symptoms often develop well before PML is diagnosed.

Forty-six percent of patients treated with natalizumab who develop PML have received previous autologous bone marrow transplantation or chemotherapy, including mitoxantrone, azathioprine, methotrexate, and mycophenolate mofetil. In comparison, up to 25% of MS patients who were treated with natalizumab (13% in the United States, 24% in Europe) have had prior chemotherapy treatment. Prior immunosuppressive therapy increases the risk of PML by two- to fourfold, which may explain the higher rate of PML in Europe compared with that of the United States.

**Testing for immune response to JCV**

A JCV enzyme-linked immunosorbent assay (ELISA) test has been developed that identifies patients with an immune response to JCV. Among MS patients, 55% test positive for JCV through this assay. The false-negative rate of the test is 5%, and the overall annual seroconversion rate is estimated to be about 2%, necessitating repeat testing.

Based on results of this assay, the estimated risk of PML in seropositive patients is about 1 in 500. The test was positive in 28 of 28 patients who developed PML. The probability of this relationship occurring by chance is 0.5528, which suggests that this assay is useful to stratify risk for development of PML. Although the rate of false negatives makes the test an imperfect predictor, it is still useful in clinical practice. The test became available clinically in late summer 2011. Further longitudinal observation studies (STRATIFY-1 and STRATIFY-2) on the use of the JCV ELISA to detect anti-JCV antibodies in the blood of natalizumab-treated patients with MS are under way.

**Stratifying risk for natalizumab-related PML**

Three factors may predict the risk of PML: JCV antibody status, history of chemotherapy use, and duration of natalizumab treatment. Estimates of risk of PML have been derived from these factors, with differences in patient profiles producing risk estimates that range from approximately 1 in 40,000 to 1 in 100. Overall, the estimated risk of a JCV-negative person who is chemotherapy-naïve is approximately 1 in 40,000. With prior chemotherapy, this risk increases to approximately 1 in 15,000.
patients who are JCV antibody-positive, the overall risk of PML is 1 in 500 for chemotherapy-naïve patients and 1 in 200 for those previously exposed to chemotherapy. To give these ratios some perspective, the lifetime risk of dying in a car accident is 1 in 100 (Table).6,7

Natalizumab holidays and PML risk
The possibility of reducing the risk of PML in natalizumab-treated patients through natalizumab holidays is attractive. When exploring this option, one must consider whether the risk of recurrent disease activity with treatment interruption outweighs the potentially decreased risk of PML.8 A randomized controlled multicenter clinical trial of natalizumab interruption is ongoing, with the recruitment phase complete after enrollment of 175 patients. Patients taking natalizumab at study entry have been randomized to one of three arms: continuation of monthly natalizumab for 6 months, placebo for 6 months, or an alternate treatment (interferon beta-1a, glatiramer acetate, or monthly intravenous steroids) for 6 months administered open-label by clinician and patient choice. The primary outcome measures are markers of immune function and overall disease activity during treatment interruption and after resumption. Patients are monitored monthly using MRI to measure disease activity. Those who experience relapse will have the option of returning to natalizumab therapy or switching to an alternate treatment. The results of this prospective, randomized, controlled trial will provide a greater understanding of the safety issues surrounding natalizumab holidays.

Management of natalizumab-related PML
Management of patients taking natalizumab starts with risk stratification in an attempt to prevent the development of PML. If suspicion for PML is raised based on symptoms, early diagnosis can be accomplished through the use of a sensitive JCV PCR assay, with a repeat PCR if negative. Natalizumab treatment should be withheld during the workup for PML.

In the setting of natalizumab therapy, where the immunosuppression is compartmentalized to the CNS, functional leukocytes are only millimeters away from where they are needed to fight JCV infection. Plasmapheresis has been shown to accelerate removal of natalizumab, accelerate desaturation of the targeted alpha-4-integrin receptor, and restore leukocyte transmigration in vivo (Figure).9 Desaturation of the integrin receptor occurs at natalizumab serum drug levels less than 1 μg/mL. Statistical modeling from pharmacokinetic measurements during a plasmapheresis study projected that clinically relevant integrin receptor desaturation is accelerated by 82 days through the use of five plasmapheresis treatments.9 Accordingly, plasmapheresis (or immunoadsorption) is recommended in natalizumab-treated patients who develop PML. Putative antiviral therapies can be considered but have so far yielded disappointing results in clinical trials. Additional trials are under way.

When immunosuppression is rapidly reversed in cases of natalizumab-associated PML, an overly exuberant immune response targeting JCV in the CNS is observed 2 to 6 weeks later. The response, termed immune reconstitution inflammatory syndrome (IRIS), is not always easy to differentiate from progression of PML. Nonetheless, most clinicians recommend high-dose corticosteroids if a clinical and imaging syndrome resembling IRIS develops several weeks after immune restoration.10 The objective is to achieve the immune reconstitution needed to control JCV infection while limiting the collateral damage of inflammation on the remaining brain tissue.

<table>
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*Estimate; no cases reported

JCV = JC virus

**SUMMARY**
Risk factors for natalizumab-associated PML include duration of treatment with natalizumab, previous chemotherapy, and JCV antibody serology. Early diagnosis requires the use of an ultrasensitive JCV PCR assay. Treatment is focused on early diagnosis, immediate cessation of pharmacologic causes of immunosuppression, and active efforts to accelerate immune restoration.
DISCUSSION

Dr. Calabrese: What are your thoughts about plasmapheresis for rituximab-related cases of PML?

Dr. Fox: It’s probably not going to be as helpful as with natalizumab. Rituximab has pharmacokinetics that are similar to those of other monoclonal antibodies, with a half-life in the range of 14 to 20 days. So it’s pretty much absent from the body within 1 to 2 months of infusion. The enduring benefit from rituximab comes not from the persistent presence of the monoclonal antibody, but the persistent absence of CD19 B cells. Plasmapheresis is unlikely to be effective because it won’t accelerate return of CD19 B cells to the peripheral circulation. In rituximab-related PML, stimulating the bone marrow to produce more B cells in order to restore the immune system is more likely to be effective. In contrast, I did recommend plasmapheresis in a case of efalizumab-related PML. Because efalizumab is a binding antibody to the CD11a receptor, we wanted to accelerate its removal.

Dr. Molloy: In an MS patient who responds well to natalizumab, do you ever explore a strategy of dose reduction or extending the dosing interval of natalizumab?

Dr. Fox: Let me put that into a clinical context. A 35-year old man has had relapsing-remitting MS for 3 years. Two years ago, after disease activity occurred while he was using an injectable therapy, he started natalizumab and has been clinically and radiologically stable on natalizumab. Then, he gets the JCV assay, it’s positive, and he asks if it’s time to get off natalizumab “because of the risk of that brain virus.” What do I tell him? Should I change the dosing interval? At this point, we are not doing either. One reason is the unpredictable pharmacokinetics of the drug. The dose and dosing regimen were chosen to have 85% or greater receptor saturation in 95% or more of patients over the course of the recommended 4-week dosing interval. If you increase the interval to 6 weeks or 8 weeks, you can’t predict in individual patients whether or not meaningful desaturation occurs and thus allows some immune cells to enter the brain to protect against PML (but not too many, or MS disease activity will return).

Dr. Simpson: Do you have an algorithm for working up patients?

Dr. Fox: It depends on the level of suspicion given the patient’s symptoms. It’s difficult to find a single MS...
patient who does not have some fluctuation of symptoms over time and some worsening of symptoms such as stiffness, fatigue, and cognitive difficulties. They all have changes in mood, so if one took any symptom change—any change in their report of mood and cognition—as the cutoff for a workup, we wouldn’t be giving natalizumab at all. But if a patient or family says, “I am worried,” then we need to work it up. Also, if there are unclear new or worsening neurologic symptoms, we pursue a workup. Often, the change in symptoms is revealed when the patient comes in for his or her monthly infusion and the nurse asks the four questions from the preinfusion checklist (as part of the mandatory Tysabri Outreach: Unified Commitment to Health [TOUCH] prescribing program for natalizumab).11

If there are new symptoms, we hold infusions and do a two-stage evaluation. The first stage is a brain MRI to evaluate for change from baseline (the US Food and Drug Administration requires a brain MRI at baseline before starting natalizumab therapy). Most patients undergo a brain MRI every 6 to 12 months while on natalizumab therapy, with instructions to the neuroradiologist to evaluate carefully for new lesions. In our institution, the PML MRI evaluation is a fine-tooth-comb assessment of lesions from the most recent MRI compared with the current MRI. Depending on the results of the current MRI and on our level of suspicion, we may proceed to a spinal tap, even if the MRI findings are stable. We have done 8 to 10 spinal taps in patients taking natalizumab when we were suspicious enough to evaluate for PML. Occasional patients continue to have active disease, relapses, and new lesions even without developing antibodies while taking natalizumab.

**Dr. Rudick:** We need a quick, quantitative analysis method to compare one MRI with another. It is easy to say, “Consider PML if there are new lesions.” It’s not so easy to know if the lesions are new. We are participating in a National Institutes of Health study regarding identification of biomarkers of interferon’s effects, and the study requires obtaining MRI scans at baseline and 6 months. We have state-of-the-art subtraction MRI to quantify new lesions on the followup MRI. However, there is significant disagreement on the number of new lesions determined by clinical raters, and disagreement between the clinical raters and the numbers generated by the computer program.

**Dr. Major:** Is the incidence of natalizumab-related PML based on the number of months or on the number of infusions?

**Dr. Fox:** It is based on the number of infusions. You bring up a good point because these patients may interrupt treatment when they go on vacation, for example, or have a lapse in insurance coverage. Most patients follow the every-4-weeks protocol and receive 13 infusions in a year. Perhaps 10% to 15% do not follow it precisely.

**Dr. Molloy:** Is everyone who takes natalizumab being followed for PML even if they discontinue natalizumab? Have any differences emerged in the factors that predispose to PML among those who continue therapy compared with those who discontinue? I ask because I’m wondering why the incidence appears to stabilize, or even go down, after 36 infusions.

**Dr. Fox:** PML has not been reported beyond several months after stopping natalizumab; concern about PML can decrease fairly quickly after stopping the drug. Many of us expected the risk of PML to continue rising with cumulative treatment, so were pleasantly surprised to see a plateau in the risk of PML after about 36 months. We don’t understand what leads to this plateau.

**REFERENCES**


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A rational approach to PML for the clinician

ABSTRACT

The first step in the management of progressive multifocal leukoencephalopathy (PML) is awareness of the disease. Patients vulnerable to PML are those with immunosuppression, either through their disease or use of immune-modulating therapy. In patients susceptible to PML who exhibit focal neurologic signs and symptoms, brain magnetic resonance imaging can detect the telltale PML brain lesions—subcortical white matter hyperintense areas on T2-weighted images and fluid-attenuated inversion recovery sequences and hypointensity on T1-weighted images, typically without enhancement. Demonstration of JC virus DNA by ultrasensitive polymerase chain reaction in cerebrospinal fluid is diagnostic for PML. Immune restoration whenever possible is the cornerstone of treatment. Highly active antiretroviral therapy has dramatically improved the prognosis for patients infected with human immunodeficiency virus. Alternatively, restoration of immunity is frequently attended by the immune reconstitution inflammatory syndrome which can be clinically devastating or even fatal. In the case of natalizumab-associated PML, withdrawal of therapy and prompt institution of plasmapheresis to desaturate target receptors provides the best chance for long-term survival.

Our remarkable progress in understanding progressive multifocal leukoencephalopathy (PML) since its discovery more than 50 years ago has evolved in three stages, concurrent with the changing epidemiology of PML: the pre–human immunodeficiency virus (HIV) era; the HIV era, with highly active antiretroviral therapy (HAART) bringing further change; and the biologic therapy era. Before the appearance of HIV, PML developed mostly in patients who had lymphoma, other malignancies, and rare forms of immunosuppression. The development of HIV changed the nature of PML, with more than 75% of cases now reported in HIV-infected patients. Within the HIV population, the epidemiology and prognosis of PML have undergone additional changes since the late 1990s. The introduction of HAART transformed PML from an almost uniformly fatal and inexorably progressive disease to one in which long-term survival is expected, particularly in the setting of robust immune reconstitution.1

The third and most recent stage in the evolution of PML and our understanding of it has coincided with the introduction and use of increasingly potent immunosuppressive regimens and novel biologic immunologic therapies that target various aspects of the integrated immune response. These agents are being applied not only in the field of autoimmune and autoinflammatory disease but also in transplantation and oncology. Collectively, vulnerable populations (ie, patients with lymphoreticular malignancies and autoinflammatory diseases) are now being subjected to therapies that singly or in combination have unknown effects on the immune system. As a byproduct, practitioners who were only vaguely aware of PML in the past now must consider PML in their differential diagnosis, develop a rational plan for evaluating such patients, and recognize when referral to a specialist is indicated. Recent descriptions of atypical forms of PML2,3 add to the challenge for clinicians, as do reports of cases of PML in patients with minimal immunosuppression, in the absence of immunosuppressive therapy, and in patients who appear to have “normal” immune systems but in fact have diseases such as sarcoidosis.4 Rare cases are also being reported in patients with advanced liver disease.5

This article offers recommendations for enhanced awareness of PML, suggestions for improved evaluation of predisposed patients, and a summary of currently accepted treatment strategies.

WHEN TO SUSPECT PML

Consideration of PML in the differential diagnosis is based on the patient’s vulnerability and the signs and symptoms of the disease. In an otherwise immunologically healthy individual, PML rarely accounts for focal
neurologic deficits. Clinicians should therefore focus their suspicions on individuals who are predisposed to PML (Table 1). Predisposed individuals are not always obvious; for example, HIV-positive individuals who have not been diagnosed with HIV infection may present with PML as the heralding manifestation of their disease.

Patients being treated with immunosuppressive biologic agents represent a significant group that is predisposed to PML. At one time, focal neurologic deficits were required to consider the possibility of PML, but cognitive/behavioral abnormalities rather than focal neurologic findings are often the presenting sign in individuals treated with immune-modulating biologic agents. This phenomenon is most strikingly observed in recipients of natalizumab. Any central nervous system (CNS) dysfunction in a patient taking an immunosuppressive biologic agent should arouse suspicion for PML.

Peripheral neuropathy is not caused by PML but can coexist with it. Accordingly, in patients with rheumatologic disease who are receiving immune modifiers, neuromuscular symptoms in the absence of brain abnormalities on magnetic resonance imaging (MRI) argue against consideration of PML but do not rule it out—especially in patients with connective tissue diseases.

Systemic lupus erythematosus (SLE) represents a special challenge for several reasons. First, SLE appears to be a predisposing factor among other connective tissue diseases. In addition, SLE is associated with a variety of CNS complications, including a spectrum of focal and diffuse signs and symptoms that can mimic PML and lead to underdiagnosis.

Underrecognition is a risk in the HIV population as well, where cognitive impairment is common. Irrespective of immune or virologic status, 57% of HIV patients demonstrate impairment on neuropsychiatric testing. Often, mild to moderate cognitive impairment in HIV is attributed to HIV encephalopathy with no further workup, resulting in a missed or late diagnosis of PML.

### Imaging Considerations

In the rheumatologic disease population, especially those with SLE, and the HIV population, neuroimaging is indicated in any patient who presents with cognitive impairment. Typical radiographic characteristics of PML on MRI are subcortical white matter hyperintense areas on T2-weighted images and fluid-attenuated inversion recovery. T1-weighted images will reveal hypointense lesions that usually do not enhance, but may do so in fewer than 10% of patients with PML. Typically, no mass effect is seen.

In addition to rare faint gadolinium enhancement of lesions, other lesion characteristics may depart from the classic picture—for example, white matter and gray matter involvement, and monofocal instead of multifocal lesions. In HIV-positive patients, MRI can demonstrate diffuse cerebellar atrophy and subtle white matter abnormalities within the cerebellum.

Unfortunately, nonspecific white matter lesions occur in HIV infection as well as connective tissue diseases, compromising diagnostic specificity of a single imaging study. Nevertheless, progression of clinical signs and symptoms and progressive MRI changes should prompt a more vigorous diagnostic evaluation for PML. Alternatively, a normal MRI in a patient in whom PML is suspected has strong negative predictive value. In either situation, baseline neuroimaging is not recommended.

### Diagnosis and Referral

A neurology consult is advised when a patient has a predisposing condition for PML or suspicious neurologic signs or symptoms, whether focal or diffuse, and in whom an MRI demonstrates white matter changes.

#### Evaluation for JC virus DNA

When the neurology consult has been scheduled, and before the actual visit to the neurologist, a cerebrospinal fluid (CSF) sample should be obtained and evaluated for JC virus (JCV) DNA using a highly sensitive polymerase chain reaction (PCR) assay. Lumbar puncture in the setting of possible PML is critical to exclude the presence of other opportunistic infections.

The importance of using ultrasensitive PCR assays for diagnosing PML cannot be overstated, as falsely negative CSF PCR has been observed for JCV DNA despite high levels of JCV DNA in spinal fluid when utilizing less sensitive assays. The most sensitive commercial assays can detect as few as 50 copies of JCV DNA per mL of CSF fluid.

The risk of PML imparted by biologic agents other than natalizumab and nonbiologic immunosuppressive agents has been difficult to quantify, but no immune-modifying drug or combination of drugs appears entirely

### Table 1

<table>
<thead>
<tr>
<th>Situations that arouse suspicion of PML</th>
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<tr>
<td>• Cognitive/behavioral abnormalities in patients receiving immune-modulating biologic agents, especially natalizumab</td>
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<tr>
<td>• Any central nervous system dysfunction in the presence of immunosuppressive biologic therapy</td>
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<tr>
<td>• Neurocognitive dysfunction in patients with systemic lupus erythematosus or human immunodeficiency virus infection</td>
</tr>
<tr>
<td>• In patients with predisposing conditions, the finding of white matter changes on magnetic resonance imaging in the presence of suspicious neurologic signs or symptoms</td>
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PML = progressive multifocal leukoencephalopathy
Any patient who has had significant or prolonged immunosuppression should be considered vulnerable, and any patient with suspicion of PML based on unexplained neurologic symptoms warrants CSF examination for JCV DNA.

### Brain biopsy

In patients with progressive clinical and MRI findings that suggest PML, but whose CSF PCR for JCV DNA is repeatedly negative, a brain biopsy is appropriate regardless of background immunosuppression. In the patient with rheumatologic disease, for example, suspicion of PML should be heightened if there is neurologic deterioration in the face of escalating antiinflammatory or immunosuppressive therapy for immune-driven inflammatory disease. Diagnostic urgency is particularly warranted in those disorders where the possibility for immune reconstitution is highest (ie, those receiving immunosuppressive regimens).

### PML MANAGEMENT DEPENDS ON CLINICAL SETTING

Management of PML starts with risk stratification to identify those patients most prone to developing PML based on their immune status; the presence of autoimmune disease; the subtype of disease (in the case of SLE); and the nature, intensity, and duration of their immunosuppression. If a high-risk patient develops signs and symptoms of PML, the diagnosis should be anticipated and serious consideration given to withholding immunosuppressive therapies while the patient is being worked up.

When PML is diagnosed, whether by demonstration of JCV in CSF or documented by brain biopsy in a patient with a suggestive clinical picture, the appropriate management depends on the clinical setting (Table 2). In the HIV-infected patient who is not receiving antiviral therapy, initiation of HAART is the core of treatment. For natalizumab-treated patients, the standard approach is to discontinue natalizumab, institute plasmapheresis to accelerate clearance of therapeutic levels of natalizumab and increase the number and function of leukocytes entering the CNS, and monitor for immune reconstitution.

### Accelerating immune reconstitution

Once a diagnosis of PML is confirmed, immune reconstitution should be accelerated whenever possible. This can include temporary or permanent withdrawal of immunosuppressive therapy and initiation of plasmapheresis. Evidence supports continuing plasma exchange until natalizumab serum drug levels decline to less than 1 μg/mL to achieve desaturation of the alpha-4 integrin receptor. Typically, desaturation of the targeted integrin receptor occurs after five plasmapheresis sessions.

Immune reconstitution may also precipitate a syndrome known as the immune reconstitution inflammatory syndrome (IRIS), characterized by enlargement and contrast enhancement of PML lesions, appearance of new brain lesions, and worsening of neurologic deficits. The infiltration of the brain with inflammatory multinucleated cells and lymphocytes following abrupt immune reconstitution requires treatment. Opinion suggests that judicious use of corticosteroids may control the immune response in the brain in patients with PML-IRIS, although further studies are needed.

### Involving the patient in treatment decisions

Because risk tolerance varies considerably among individuals, patients should be informed of the risks of PML on the basis of their disease and the agents used to treat it. They should also be given information about the effects of individual treatments on the course of their disease, and they should be encouraged to participate in the selection of therapy.

### SUMMARY

The approach to PML in the biologic era starts with an increased awareness of the disease followed by recognition of vulnerable populations and factors that contribute to the development of PML, such as biologic and nonbiologic immunosuppressive therapy. Optimal management includes a low threshold for investigating neurologic signs and symptoms and new-onset signs and symptoms in vulnerable populations, the use of MRI to detect typical PML brain lesions and other atypical brain features (ie, cerebellar atrophy), lumbar puncture and spinal CSF analysis to detect JCV DNA, and timely neurologic consultation for further evaluation. Much still needs to be learned about PML and the risks imparted by background diseases and individual drugs used in rheumatologic, neurologic, and oncologic disease.

### REFERENCES


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