Current and Emerging Therapies in Multiple Sclerosis: Implications for the Radiologist, Part 2—Surveillance for Treatment Complications and Disease Progression

C. McNamara, G. Sugrue, B. Murray, and P.J. MacMahon

ABSTRACT

SUMMARY: An understanding of the new generation of MS drugs in conjunction with the key role MR imaging plays in the detection of disease progression, opportunistic infections, and drug-related adverse effects is of vital importance to the neuroradiologist. Part 1 of this review outlined the current treatment options available for MS and examined the mechanisms of action of the various medications. It also covered specific complications associated with each form of therapy. Part 2, in turn deals with the subject of pharmacovigilance and the optimal frequency of MRI monitoring for each individual patient, depending on his or her unique risk profile. Special attention is given to the diagnosing of progressive multifocal leukoencephalopathy in patients treated with natalizumab as this is a key area in which neuroradiologists can contribute to improved patient outcomes. This article also outlines the aims of treatment and reviews the possibility of “no evidence of disease activity” becoming a treatment goal with the availability of more effective therapies. Potential future areas and technologies including image subtraction, brain volume measurement and advanced imaging techniques such as double inversion recovery are also reviewed. It is anticipated that such advancements in this rapidly developing field will improve the accuracy of monitoring an individual patient’s response to treatment.

ABBREVIATIONS: BV = brain volume; DIR = double inversion recovery; DMT = disease-modifying therapy; IRIS = immune reconstitution inflammatory syndrome; JC virus = John Cunningham virus; MTR = magnetization transfer ratio; NEDA = no evidence of disease activity; NTZ = natalizumab; NTZ PML = natalizumab-associated PML; PML = progressive multifocal leukoencephalopathy; PML IRIS = PML-associated IRIS; PSIR = phase-sensitive inversion recovery

Part 1 of this review covered recent developments in the treatment of MS and discussed the published clinical data on the efficacy and safety of the currently approved and emerging therapies in this condition as they apply to the radiologist. Part 2 will focus on the role of the radiologist in monitoring patients being treated with these medications and will focus on pharmacovigilance, which includes observation for treatment complications and signs of disease progression.

Frequency of MR Imaging for Drug Monitoring

MR imaging is recommended before the initiation or modification of disease-modifying therapy (DMT) and approximately 6 months after a treatment switch to allow sufficient time for the new therapy to reach its therapeutic potential. Therefore, some authors suggest that the reference scan should be obtained no earlier than 3 months after treatment initiation/change to overcome the uncertainty about new lesions occurring before the treatment becomes effective. Continued or worsening disease activity on MR imaging while a patient is on a DMT may prompt a change in therapy. There is evolving evidence that ongoing MR imaging activity can be indicative of a suboptimal therapeutic response; however, due to variable pharmacodynamics, the time until MR imaging activity is suppressed differs among the immunomodulatory agents.

Several studies have attempted to define criteria and strategies for the early identification of suboptimal response in individual patients via a combination of clinical and MR imaging measures during the first 6–12 months after treatment initiation. These criteria are partially or completely based on the detection of disease activity on follow-up brain MR imaging studies, defined as new gadolinium-enhancing lesions or new and/or enlarging T2 hyperintense lesions.

More frequent surveillance may be indicated in clinically aggressive cases or unusual patterns (eg, tumefactive MS) (Table 1). Clinical judgment and experience are critical in such cases. While guidelines on a tolerable threshold for new lesion activity that warrants a change in therapy have been proposed, individual factors will impact the clinician’s decision as to the exact frequency of MR imaging monitoring and time to switch ther-

From the Departments of Radiology (C.M., G.S., P.J.M.) and Neurology (B.M.), Mater Misericordiae University Hospital, Dublin, Ireland.

Please address correspondence to Peter MacMahon, MD, Department of Radiology, Whitty Bldg, North Circular Rd, Mater Misericordiae University Hospital, Dublin 7, Ireland; e-mail: pmacmahon@mater.ie; @McnamaraCillian

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http://dx.doi.org/10.3174/ajnr.A5148
Isolated worsening on the neurologic examination or MR imaging to date, among patients in addition, PML is not the only MR imaging–based monitoring of patients for early progressive multifocal leukoencephalopathy (PML) detection is not recommended for based monitoring of patients for early progressive multifocal leukoencephalopathy (PML) detection is not recommended for brain MR imaging should lead to closer monitoring rather than brain MR imaging should lead to closer monitoring rather than opportunistic infections who are switching DMT opportunistic infections who are switching DMT. MS treatment–related infections include a wide-ranging opportunistic infection that can be observed during MS treat opportunistic infection that can be observed during MS treat opportunistic infection that can be observed during MS treat opportunistic infection that can be observed during MS treat opportunistic infection that can be observed during MS treat opportunistic infection that can be observed during MS treat opportunistic infection that can be observed during MS treat opportunistically with patients identified at the symptomatic stage. The current diagnostic criteria for PML as established by the American Academy of Neurology include the presence of 3 components: a clinical picture suggestive of PML (eg, behavioral and spectrum of pathogens such as varicella zoster virus, as has been described in patients treated with fingolimod.14 There is now growing evidence that drug-related adverse effects can occur when MS medication is discontinued or even several months after a new treatment is started (so-called “carry over opportunistic infections”). Therefore, strict pharmacovigilance, including frequent MR imaging, should be performed in patients who switch therapies, to detect resurgent MS disease activity and adverse effects such as opportunistic infections.15–17 In patients at high risk of developing opportunistic infections (eg, immunocompromised patients, those recently treated with a course of immunosuppressants) who are switching DMTs, brain MR imaging is recommended when the current treatment is discontinued and after the new treatment is started. Thus, heightened pharmacovigilance, including brain MR imaging every 3–4 months for up to 12 months, is required in patients who switch from NTZ to other medications (such as fingolimod, alemtuzumab, and dimethyl fumarate).20 To date, among patients switching to fingolimod treatment after previous natalizumab therapy, there have been 17 suspected cases of PML.18

| Risk Factors for PML | Duration of Therapy ≥24 Months |
|----------------------|-------------------------------|
| Development in Patients Treated with NTZ | Low risk | Prior immunosuppressive therapy | $\geq 24$ mo |
| High risk | Evidence of JCV antibody seropositivity | As indicated |
| | Presence of anti-JCV antibodies | | |
| | Increased treatment duration, especially beyond 2 yr | | |
| | High JCV titers | | |
| | Patients negative for anti-JCV antibody (risk is <0.09/1000) | | |
| | All patients receiving NTZ should be screened for previous JCV infection | | |
| | Serologic testing for patients negative for JCV every 6 mo | | |
| | Patients positive for JCV who have been treated with NTZ for >2 yr should be switched to another second-line therapy | | |
| | Increased frequency of imaging surveillance for high-risk patients (eg, every 3–4 mo) | | |
| | Comparison of surveillance images with previous MR imaging | | |

Note: --JCV indicates John Cunningham virus.

Table 1: Frequency of MRI surveillance

| Clinical Indication | Frequency of Imaging | Imaging Protocol |
|---------------------|----------------------|-----------------|
| RRMS, routine surveillance | Annually for at least the first 2 or 3 years after starting therapy or switching DMT | T2-weighted and contrast-enhanced T1-weighted |
| Higher risk patients (positive for JC virus serum antibodies) with ≥24 mo of NTZ exposure | Every 3–6 months | T2WI, T2 FLAIR, DWI, SWI (if indicated) |
| Low risk of PML (negative for JC virus serum antibodies) | Annually | T2WI, T2 FLAIR, DWI, SWI (if indicated) |
| Patients at high risk of developing opportunistic infections who are switching DMT | MRI when the current treatment is discontinued and 3–6 months after the new treatment is started | T2WI, T2 FLAIR, DWI, SWI (if indicated) |
| Patients who switch from NTZ to other therapeutics (including fingolimod, alemtuzumab, and dimethyl fumarate) | Enhanced pharmacovigilance, including brain MRI every 3–4 mo for up to 12 mo | T2WI, T2 FLAIR, DWI, SWI (if indicated) |
| Patients who require enhanced pharmacovigilance for other reasons | Every 3–6 mo | As indicated |

Note: --RRMS indicates relapsing-remitting MS.

Imaging Surveillance for Opportunistic Infections

Particularly high-risk patients (natalizumab [NTZ] treatment longer than 24 months, serology positive for John Cunningham [JC] virus serology, prior immunosuppression) should be screened every 3–6 months with at least a curtailed MR imaging protocol to include FLAIR, T2-weighted, DWI, and SWI sequences if available.8 Low-risk patients (JC virus seronegative) can be imaged on an annual basis (Table 2).8,11 MR imaging–based monitoring of patients for early progressive multifocal leukoencephalopathy (PML) detection is not recommended for those patients treated with NTZ exclusively, but also for patients being treated with other DMTs, including alemtuzumab, fingolimod, and dimethyl fumarate.12,13 In addition, PML is not the only opportunistic infection that can be observed during MS treatment. MS treatment–related infections include a wide-ranging

Table 2: Risk stratification of natalizumab-associated PML

| Risk stratification of patients Treated with NTZ | Duration of Therapy ≥24 Months |
|-----------------------------------------------|-------------------------------|
| High risk | Prior immunosuppressive therapy | $\geq 24$ mo |
| | Evidence of JCV antibody seropositivity | As indicated |
| | Presence of anti-JCV antibodies | | |
| | Increased treatment duration, especially beyond 2 yr | | |
| | High JCV titers | | |
| | Patients negative for anti-JCV antibody (risk is <0.09/1000) | | |
| | All patients receiving NTZ should be screened for previous JCV infection | | |
| | Serologic testing for patients negative for JCV every 6 mo | | |
| | Patients positive for JCV who have been treated with NTZ for >2 yr should be switched to another second-line therapy | | |
| | Increased frequency of imaging surveillance for high-risk patients (eg, every 3–4 mo) | | |
| | Comparison of surveillance images with previous MR imaging | | |

Note: --JCV indicates John Cunningham virus.

Diagnosing PML

Although differentiating between a new MS lesion and an evolving PML lesion can be challenging, radiologists must be familiar with the key imaging features associated with PML because a timely and accurate diagnosis at the asymptomatic stage is associated with improved survival and functional outcome compared with patients identified at the symptomatic stage.19 MR imaging has been shown to be sensitive in the detection of PML lesions and can identify such lesions several months before the onset of clinical symptoms.20

The current diagnostic criteria for PML as established by the American Academy of Neurology include the presence of 3 components: a clinical picture suggestive of PML (eg, behavioral and
Compared with symptomatic PML, lesions are often large (≥3 cm) when asymptomatic. In symptomatic patients, lesions are more confluent and diffuse. Microscopic lesions, no associated edema or mass effect, are common in asymptomatic patients.

**MRI Features of PML**

| Features                  | Location                  | Size                  | Morphology                     |
|---------------------------|---------------------------|-----------------------|--------------------------------|
| Location                  | Predilection for frontal lobe > occipital lobe > parietal lobe | Small, <3 cm when asymptomatic | Punctate lesions often identified in asymptomatic PML |
| Size                      |                           |                       | Symptomatic PML typically demonstrates more confluent and diffuse lesions |
| Morphology                |                           |                       | Microcysts                     |
| TIW and T2WI              | FLAIR hyperintensity with corresponding T1 hypointense lesions | T1 hypointensity is less pronounced in asymptomatic PML |
| DWI                       | Hyperintensity on DWI     | 30% will demonstrate linear or punctate enhancing lesions; if nodular or rim enhancement is present, active MS plaque or IRIS should be considered |
| SWI                       | Low signal intensity in U-fibers | MS plaque            |
| Differential diagnosis    | Acute disseminated encephalomyelitis | Tumefactive lesion |
|                           | Ischemic changes          | Posterior reversible encephalopathy syndrome |

**FIG 1.** Infratentorial natalizumab-associated PML. Axial FLAIR (A and B) images of a patient demonstrating extensive hyperintense signal involving the cerebellum, middle cerebellar peduncle, and pons. This had progressed from prior imaging. Note minimal mass effect. Postadministration of gadolinium, there is no enhancement demonstrated (C).

**TABLE 3: MRI Features of PML**

| Location               | Features                                      |
|------------------------|-----------------------------------------------|
| Subcortical white matter U-fibers | Predilection for frontal lobe > occipital lobe > parietal lobe |
| Predilection for frontal lobe > occipital lobe > parietal lobe | Small, <3 cm when asymptomatic |
| Punctate lesions often identified in asymptomatic PML | Symptomatic PML typically demonstrates more confluent and diffuse lesions |
| Microcysts | No associated edema or mass effect |
| FLAIR hyperintensity with corresponding T1 hypointense lesions | T1 hypointensity is less pronounced in asymptomatic PML |
| Hyperintensity on DWI | 30% will demonstrate linear or punctate enhancing lesions; if nodular or rim enhancement is present, active MS plaque or IRIS should be considered |
| Low signal intensity in U-fibers | MS plaque |
| Acute disseminated encephalomyelitis | Tumefactive lesion |
| Ischemic changes | Posterior reversible encephalopathy syndrome |

PML-immune reconstitution inflammatory syndrome, particularly during MS follow-up imaging.25,26

The most common area to be affected by NTZ PML is the frontal lobe, followed in frequency by the occipital lobe and then the parietal lobe (Table 3).27,28 Like other aggressive infiltrating lesions, PML can also infiltrate the corpus callosum, though isolated corpus callosal involvement is rare.29 PML lesions occur in the posterior fossa in early PML in approximately 10% of cases at initial presentation (Fig 1).28 These most commonly involve the cerebellum and middle cerebellar peduncles, though the brain stem can be affected.27,30 Crescent-shaped lesions involving the middle cerebellar peduncles and adjacent cerebellar and/or pontine white matter may be specific to PML, rather than MS, because they have, so far, only been reported in patients with PML.30-32 Deep gray matter (including the thalamus and dentate nuclei) PML lesions are unusual (5%) in NTZ PML, particularly in the early disease course.22

A 2016 study published in *Radiology* by Hodel et al27 investigated the radiologic findings of NTZ PML at the asymptomatic stage. They concluded that hyperintensity on DWI and involvement of U-fibers were the most predictive MR imaging features for the diagnosis of asymptomatic NTZ PML. Their results also suggested that punctate lesions may be a highly specific finding for NTZ PML, being exclusively observed in patients with NTZ PML. Most of the lesions observed in their study involved the frontal lobe as has been previously reported in other studies.28

They also found that comparison of the patient’s current MR images with those from previous MR imaging examinations significantly improved the specificity for the detection of NTZ PML lesions and for the diagnosis of PML (from 88% without to 100% with previous MR imaging; P = .05). Recent evidence also suggests that SWI may be of value in the early diagnosis of PML (Fig 2). In particular, low signal intensity observed in U-fibers identified with SWI has been reported as a unique finding in early PML.33,34 Most interesting, the rate of PML in patients treated for <1 year remains virtually zero. Thus, the patient’s length of treatment with natalizumab should be provided to the radiologist when interpreting MR imaging findings.

**PML–Immune Reconstitution Inflammatory Syndrome**

In a bid to halt the progression of PML, plasma exchange or immunoadsorption can be used to remove NTZ from the circulation, which allows lymphocytes to cross the BBB once again.35

cognitive abnormalities, progressive weakness, gait abnormalities, visual field deficits, speech and language disturbances, and/or incoordination), biochemical and/or histologic evidence of the JC virus, and MR imaging findings.21 Compared with symptomatic PML, in which the lesion dissemination typically involves multiple lobes affecting both hemispheres, patients with asymptomatic PML present more frequently with a localized disease in a unilateral and focal distribution.19,20 Most interesting, compared with other PML cohorts, contrast enhancement can be observed quite frequently in natalizumab-associated PML (NTZ PML), occurring in approximately 30% of all patients at diagnosis, and can be the leading radiologic sign in asymptomatic patients.22–24

Substantial evidence indicates that FLAIR is the most sensitive sequence for detecting PML.22 DWI is known to be highly sensitive for detecting acute demyelination but can also aid in the differentiation of acute PML lesions from chronic and subacute demyelinating MS lesions.22 In patients with MR imaging lesions suggestive of PML, the MR imaging protocol should be extended to include pre- and postcontrast T1-weighted imaging to detect inflammatory features and the possible coincidence of PML and
This sudden restoration of cellular immunity can paradoxically result in the worsening of the patient’s neurologic status with the development of immune reconstitution inflammatory syndrome (IRIS).36 The development of IRIS occurs days to weeks following treatment with plasma exchange or immunoadsorption.35,36 PML-associated IRIS (PML IRIS) can, however, also occur following the cessation of NTZ alone without plasma exchange.37 In such cases, immunologic rebound with IRIS takes approximately 3 months postcessation, in keeping with the long half-life of the drug.37 This represents a challenge for the radiologist because neurologic dysfunction and white matter lesions occurring after NTZ interruption may also be caused by MS exacerbations.38 The literature suggests that PML IRIS tends to occur first in such cases followed by recurrences of MS activity at later follow-up imaging, suggesting that timing is key to reaching a diagnosis.17,25,28

Imaging findings that can assist in the diagnosis of PML IRIS include contrast enhancement and rapid progression of existing PML lesions, which begin to exert mass effect through swelling and edema.37,39,40 Current data, though limited, suggest that contrast enhancement with either a patchy or punctate pattern in the border of the PML lesion is the most frequent imaging sign in early PML IRIS (Fig 3).37 Gheuens et al41 have developed a diagnostic model using proton MR spectroscopy to help differentiate PML IRIS lesions from lesions without IRIS on the basis of their metabolic profile and the presence contrast enhancement.

**Assessing Radiologic Response to Therapy**

Several studies have demonstrated that the occurrence of new T2 lesions or gadolinium-enhancing lesions during the first year of a treatment (most studies have investigated treatment with interferon β) correlates with progression of disability.42,43 These 2 measures have relevant differences. While contrast-enhancing lesions are considered a marker of acute inflammation in patients with MS, new T2 hyperintense lesions reflect the permanent footprint from a previous focal inflammatory lesion that developed in the interval between 2 scans.44

Determining ongoing radiologic stability is based on the presence or absence of new lesions (whether T2-weighted or contrast-enhancing lesions).5,42,43,45,46 In a large retrospective study of patients with MS on therapy, the presence of ≥2 contrast-enhancing lesions at 1 year was predictive of poor clinical outcome at 5 years.47 The Barcelona and Cleveland Clinic groups have both shown that the accumulation of a certain number of new T2 lesions, with or without clinical relapse, has a predictive value for disability progression during the next few years.3,43,48 For example, Rio et al49 showed that ≥3 lesions after 1 year of treatment predicted increased disability after 2 years (OR = 8.3, 71% sensitivity, 71% specificity).

A scoring system proposed by Rio et al50 combines commonly used, short-term (ie, 1-year) treatment measures of disease activity, such as clinical relapses, Expanded Disability Status Scale progression, and active MR imaging lesions, to identify patients at risk of developing new clinical activity (ie, relapses) during the ensuing years. The other novelty provided by this proposed score is the use of the number of new T2 hyperintense MR imaging lesions instead of gadolinium-enhancing lesions. Previous approaches have used gadolinium-enhancing lesions to assess MR imaging activity because they are easier to detect as already mentioned.42,43,51 The Rio Criteria or the Modified Rio Criteria are guides to prognosis; but unfortunately, their sensitivity and specificity are not considered accurate enough for routine clinical practice.5,6

Critics have dismissed T2 lesions as a predictive biomarker because of the moderate-to-poor short-term correlation between the accumulation of lesions and outcome. However, long-term studies are much more informative. For example, in the Queen Square 21-year clinically isolated syndromes follow-up study, T2 lesion load was highly predictive of a poor clinical outcome.52 In a
meta-analysis from 23 randomized placebo-controlled clinical trials involving 6591 patients, regression models demonstrated that the treatment effect on T2 hyperintense lesions correlated with the effect on relapses. Most interesting, the development of new T2 lesions in patients with MS on DMT is much more predictive of long-term outcome compared with the development of new T2 lesions in patients on no treatment or a placebo. Such data make a compelling case for suppressing the development of T2 lesions as a treatment target, with the aim of delaying or preventing the onset of secondary-progressive MS. One drawback of such an approach is that the detection of new T2 hyperintense lesions can be challenging, especially if there is a preexisting high load of lesions, and one has to factor in variable positioning on serial scans and 1.5T-versus-3T lesion detection.

Whereas the classification of new T2 lesions can be more difficult and is more time-consuming, T2 signal alterations are a durable “footprint” of plaque formation and thus a relatively reliable means of ascertaining lesion accumulation. Some of the limitations of T2 lesion counting could be overcome by technologic innovations, for example emerging registration and subtraction techniques.

Defining “Nonresponders”
Effective DMT treatment has been shown to be especially beneficial early in the course of MS. Thus, rapid identification of nonresponders is crucial to determine the need for a therapeutic switch. Patients with MS who continue to experience clinical and/or radiologic disease activity despite treatment with DMT are categorized as “nonresponders.” The response of patients with MS to DMT is very heterogeneous; thus, it is especially difficult to determine whether a particular treatment of choice is producing the desired effect. Proposed scoring methods to identify patients with a suboptimal treatment response on the basis of combined clinical and radiologic measures at follow-up have shown considerable variation. Moreover, such criteria have been developed almost exclusively in patients receiving different formulations of interferon β. Few data are available from patients undergoing treatment with the newer DMTs in clinical practice.

Advancing toward “No Evidence of Disease Activity”
With the introduction of new, more effective therapies, there has been a shift in treatment expectations from partial response to potential remission, with no evidence of disease activity (NEDA) being the new treatment target. NEDA is defined as the following: 1) absence of relapses, 2) absence of sustained disability worsening, and 3) absence of radiologic activity, seen as gadolinium-enhancing lesions or new/enlarged T2-hyperintense lesions (NEDA-3). More recently, this composite outcome has been implemented by also considering the absence of brain volume loss (NEDA-4). The clinical significance of NEDA in the long-term, however, is still lacking, and the concept is still evolving because there is debate over additional measures that could or should be included, such as loss of brain volume as a marker. Realistically because DMT is only partially effective, some lesion accumulation is an expected outcome for patients treated with these therapies and neurologists do not routinely switch DMTs with minor asymptomatic relapses. While new T2 lesions during treatment are undesirable, studies are heterogeneous in relation to the number of new lesions considered significant disease activity. In a recent systematic review investigating the role of MR imaging in the assessment of treatment response to interferon β, pooled data of new T2 lesions did not show statistical significance for those patients with only 1 new T2 lesion per year. A 2016 study published by Prosperini et al showed that 34% of patients treated with natalizumab achieved NEDA-3 at 7-year follow-up.

From the clinician’s point of view, there are multiple justifications for switching therapies. First, MS involves diverse pathologic mechanisms that no single DMT addresses. Second, MS is heterogeneous, and one might expect a given agent to work better in one patient than another. Third and perhaps most important, no DMT actually cures MS. Approximately 30% of patients show a suboptimal therapeutic response during the early treatment years. Therefore, NEDA may be an unrealistic long-term expectation, especially when economic factors and adverse effects are taken into account. There are no uniform criteria for choosing the initial DMT to be trialed in an individual patient. This decision is guided by drug, disease, and patient considerations; personal preferences and experiences; and economic and availability factors. Most initial treatment choices would involve a first-line agent before consideration is given to a second-line agent such as NTZ.

Future Areas
Brain Atrophy as a Marker. Numerous studies have used MR imaging–derived methods to assess brain volume (BV) changes in MS, revealing that atrophy can be present even in the early stages of the condition and that it progresses during the disease course. Several MR imaging studies consistently reported associations between whole-brain gray matter loss and increases of lesion load. The clinical relevance of brain atrophy is well-known: Compared with measures of lesion load, brain atrophy shows better correlations with progression of disability and cognitive impairment, with gray matter atrophy being more closely associated with clinical dysfunction than white matter atrophy. The overall effect of treatment on brain atrophy can be reflective of the effect of the therapy on disability. The more recent DMTs have been shown to decrease rates of brain atrophy when compared with a placebo. Thus, recent clinical trials have incorporated BV outcomes as a measure of treatment effect. Current methods for measuring BV in radiology are usually automated software that either uses a segmentation-based approach (such as statistical parametric mapping) for cross-sectional data or a registration-based approach (such as structural image evaluation with normalization of atrophy) for longitudinal analysis. Unfortunately, the clinical interpretation of BV loss in patients with MS can often be difficult with other confounding factors such as hydration status and ongoing inflammation at the time of treatment onset. Moreover, differences in the quality and capabilities of MR imaging hardware and in the software packages used for analysis or processing can generate notable variability in brain atrophy assessment.

In many trials, DMTs have produced only moderate evidence of a reduction in BV loss. This is because anti-inflammatory agents have been shown to increase the rate of BV reduction.
within the first 6 months to 1 year of treatment, followed by stabilization during the second year of treatment. Resolution of this inflammation will lead to an initial accelerated BV loss that has been described as a “pseudoatrophy” effect. Therefore, the most clinically efficacious drugs (with a high impact on inflammation), such as natalizumab, fingolimod or high-dose interferon β, will tend to produce larger-than-placebo BV decreases during the first months of therapy, which, at least in part, may not be related to true tissue damage. The difficulties in clinical interpretation of BV loss outlined above such as the pseudoatrophy effect explain why this novel technique is not currently employed in routine clinical use.

**Image Subtraction to Detect New Lesions.** A second area of growing interest is the accurate detection of active lesions and its achievement. Both new and enlarging lesions, imaged with standard serial nonregistered 2D images, are complicated by repositioning errors and a background of unaltered nonactive lesions. Subtraction images, after image registration, provide an alternative strategy in which the effect of repositioning errors is reduced, cancelling out radiologically stable disease. In both single-center and multicenter settings, 2D subtraction images depicted higher numbers of active lesions with greater interobserver agreement compared with nonregistered 2D T2-weighted spin-echo images. Image subtraction can overcome issues associated with differentiating old T2 lesions from new T2 lesions, thus providing good visualization and quantification of active and resolving lesions. Recent data have shown that automated identification of new or enlarging T2 lesions is robust, accurate, and sensitive, thus supporting its use for evaluating treatment efficacy in clinical trials. Long-interval T2-weighted sequences can be processed with automated subtraction, which has been used in a multicenter trial to provide greater power for assessing treatment efficacy than is possible with monthly contrast-enhanced T1-weighted imaging. However, subtraction requires time-consuming postprocessing steps, and the process can be susceptible to imaging artifacts.

**Advanced Imaging Techniques.** A broadening spectrum of emerging MR imaging techniques allows a more specific evaluation of MS pathology. Included in these new methods are the following: magnetization transfer ratio (MTR), diffusion tensor imaging, restricted proton fraction, and positron-emission tomography techniques. Furthermore, additional pulse sequences such as double inversion recovery (DIR) and phase-sensitive inversion recovery (PSIR) have improved the detection rate of cortical lesions.

Although conventional MR imaging is very sensitive to macroscopic lesions, it lacks sensitivity to the microscopic pathology. Diffusion MR imaging techniques, including DTI and diffusional kurtosis imaging, are sensitive to diffuse microscopic injury in the normal-appearing white matter and gray matter of patients with MS. These techniques provide quantitative parameters such as fractional anisotropy and mean diffusivity, which correlate well with tissue damage that is not visible on T2 and FLAIR imaging. DTI may also assist in evaluating the structural integrity of white matter and can be used for probabilistic or deterministic tractography. In one in vivo study of MS, radial diffusivity appeared to provide the closest correlation with tissue integrity. A multicenter validation study indicated that fractional anisotropy was the most comparable DTI measure across centers and endorsed its use in multicenter clinical trials. Werring et al demonstrated that a steady and moderate increase of the apparent diffusion coefficient can precede the development of new plaques by 6–8 months, and they indicated that structural changes in prelesion normal-appearing white matter occur before inflammation and BBB leakage.

MTR is a technique based on the transfer of magnetization between semisolid and water protons in different structural environments. Longitudinal studies demonstrate decreases in MTR preceding contrast enhancement. The magnitude of the MTR decline during gadolinium enhancement predicts whether a lesion will evolve into a T1 hypointense one. Thus, MTR alteration has been proposed as a marker of overall lesion severity. Partial or complete resolution of the MTR may occur as inflammation is reduced and remyelination ensues. Due to these features, this technique provides a promising primary outcome measure to evaluate remyelinating therapies in clinical trials. Several metrics that have been proposed for further study include intralesional MTR and whole-brain MTR.

DIR sequences simultaneously suppress CSF and white matter signals, thereby increasing the detectability of gray and white matter lesions in MS significantly. Several reports have shown the utility of DIR and PSIR images for detecting MS plaques, especially in intracortical or mixed white matter–gray matter areas compared with FLAIR and T2-weighted images (Fig. 4). Recently showed that such sequences could be recreated by using synthetic MR imaging techniques, which lead to greater diagnostic accuracy than conventional MR images in comparable acquisition times.

Last, there have been studies that have combined the above imaging advances with promising results. Calabrese et al showed that the combination of DIR and DTI constitutes a substantial step forward in the analysis of cortical disease in MS. They
combined these techniques to analyze the cortex of patients with relapsing-remitting MS and found that fractional anisotropy was significantly increased in cortical lesions compared with normal-appearing gray matter. Their study concluded that there is a high correlation between the fractional anisotropy of normal-appearing gray matter and the progression of physical disability during a 3-year follow-up. Another example is the combined use of DIR and PSIR, which allows for more reliable detection of cortical lesions in MS than DIR techniques alone. The addition of PSIR also provides better delineation of lesion boundaries, which results in a more confident classification of lesions as purely intracortical, mixed, or juxtacortical.390

Although the use of quantitative MR imaging measures for disease monitoring shows much promise, their use in routine clinical practice is, in part, hindered by the longer acquisition and postprocessing times and a current lack of standardization.

CONCLUSIONS
The field of MS therapeutics is a rapidly evolving one. For radiologists to provide an informed opinion in discussions of newer MS agents, they must have a working understanding of the strengths and limitations of various novel therapies. The role of imaging in MS has advanced beyond confirmation of disease presence to include the monitoring and surveillance of disease activity and treatment complications. An understanding of the new generation of MS drugs in conjunction with the key role MR imaging plays in the detection of disease progression, opportunistic infections, and drug-related adverse effects is of vital importance to the radiologist and physician alike. Radiologists are in a unique position to detect many of the described complications well in advance of clinical symptoms. This rapidly expanding area will see much change in the years to come, and it is more important than ever for the neurologist and radiologist to work together in the pursuit of improved outcomes for the patient with MS.

ACKNOWLEDGMENTS
The authors thank Drs Seamus Looby and Ronan P. Killeen for providing images for the article.

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