Radiologically Isolated Syndrome: A Review for Neuroradiologists

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ABSTRACT

SUMMARY: Radiologically isolated syndrome refers to an entity in which white matter lesions fulfilling the criteria for multiple sclerosis occur in individuals without a history of a clinical demyelinating attack or alternative etiology. Since its introduction in 2009, the diagnostic criteria of radiologically isolated syndrome and its clinical relevance have been widely debated by neurologists and radiologists. The aim of the present study was to review the following: 1) historical evolution of radiologically isolated syndrome criteria, 2) clinical and imaging findings in adults and children with radiologically isolated syndrome, 3) imaging features of patients with radiologically isolated syndrome at high risk for conversion to MS, and 4) challenges and controversies for work-up, management, and therapeutic interventions of patients with radiologically isolated syndrome.

ABBREVIATIONS: CIS = clinically isolated syndrome; DIS = dissemination in space; DIT = dissemination in time; MAGNIMS = Magnetic Resonance Imaging in MS; RIS = radiologically isolated syndrome; RRMS = relapsing-remitting multiple sclerosis; CADASIL = cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy; ADEM = acute disseminated encephalomyelitis

Radiologically isolated syndrome (RIS) refers to an entity in which white matter lesions that are characteristic in morphology and location of a demyelinating disease, fulfilling the revised 2017 McDonald Criteria for dissemination in space (DIS) (On-line Table), but without a clinical history of demyelinating attacks or ongoing neurologic deterioration or other alternative causes of the white matter lesions such as those from vascular, infectious, toxic, and drug-related pathology. Some might argue that RIS is not truly a clinical diagnosis and instead is part of a continuum from health to disease that cannot currently be distinguished on the basis of imaging and clinical features.

The widespread use of brain MR imaging in multiple settings has led to frequent reporting of such incidental lesions on T2-weighted and FLAIR pulse sequences. In most cases of RIS, patients are being evaluated for headache, trauma, or nonspecific dizziness, symptoms that are not characteristically attributed to MS, and they have not had prior clinical episodes of neurologic deficits.

While the prevalence of RIS remains unknown, incidentally found white matter lesions resembling demyelination occur in 0.1%–0.7% of the general population. The incidence of RIS has been estimated at 0.8 per 100,000 person-years in a Swedish cohort consisting of 1907 patients, compared with the 10.2 per 100,000 person-year incidence of MS. The prevalence of RIS is known to be increased in healthy relatives of patients with MS. Gable et al found that the prevalence of RIS was 2.9% in the healthy relatives of patients with MS in contrast to the prevalence of 2.4% in nonfamilial healthy controls. Liu et al examined 326 consecutive patients 10–55 years of age presenting with headache and found that the Barkhof and 2010 McDonald criteria for DIS were met in 2.4%–7.1% and 24.4%–34.5%, respectively. These reported figures, however, far exceed the prevalence of MS. Overall, on the basis of postmortem studies, the prevalence of RIS seems to fall within 0.06%–0.7%.

Since RIS was first defined in 2009, the implications of these imaging findings have been widely investigated. The conversion rate of RIS to MS and the debate over whether to start MS disease-modifying therapies (with their potentially serious side effects) in RIS are subjects of intense scrutiny, summarized herein.
Patients for which such data were available. The mean time to the definite MS was noted in the longitudinal follow-up of 10 of 30 Development of clinically isolated syndrome (CIS) or clinically nervous system demyelinating disorder, toxic conditions, or other individuals without a history of symptoms consistent with a central fulfilled at least 3 of 4 Barkhof criteria (Table 1) for DIS in RIS; however, they suggested that identical MR imaging criteria for DIS in RIS is based on the presence of imaging findings of RIS in a radiology-based scientific journal to date.

**Criteria Evolution**

The diagnosis of RIS has relied on the imaging criteria used to make the diagnosis of MS. The 2010 McDonald Criteria simplified the DIS criteria for the diagnosis of MS by focusing on the distribution of the demyelinating lesions (juxtacortical, periventricular, infratentorial, and spinal cord) rather than the total number of lesions. The 2010 and 2017 revised McDonald criteria did not address the RIS entity because of the paucity of longitudinal data surrounding RIS. However, in 2017, the Magnetic Resonance Imaging in MS (MAGNIMS) European group tried to better define the imaging criteria for RIS. The MAGNIMS consensus group did not propose a strict recommendation on diagnosing and treating RIS; however, they suggested that identical MR imaging criteria for DIS and dissemination in time (DIT) should be used for RIS and MS (Table 2).

In other words, instead of using the Barkhof criteria as proposed in Okuda et al, the revised 2017 McDonald and/or MAGNIMS criteria should be applied (On-line Table). On the basis of the new guidelines, DIS in RIS is based on the presence of T2/FLAIR hyperintense lesions in at least 2 of the following topographies: cortical or juxtacortical white matter, periventricular white matter, spinal cord, and the infratentorial (brainstem and/or cerebellum) compartment (Fig 1).

A detailed clinical history and a meticulous neurologic examination are essential to rule out any clinical evidence of MS before labeling the patient as fulfilling the definition of RIS. The criteria used for diagnosing clinically definite MS require the presence of clinical relapses (relapsing-onset MS) or ongoing neurologic deterioration from the start (progressive onset). MAGNIMS opined and the 2017 McDonald Criteria Panel agreed that if a clinical episode occurs in a subject positive for RIS-DIS/DIT then a diagnosis of MS can be made.

### Imaging Findings

In the article of Okuda et al on RIS, 10 of 41 (24%) patients with periventricular, juxtacortical, spinal, or infratentorial white matter lesions had ≥1 enhancing plaque at the time of RIS presentation. Of these 41 patients, MR imaging progression (the presence of new T2 foci, new gadolinium enhancement, or enlargement of pre-existing lesions) on longitudinal MR imaging, fulfilling the DIT criteria, was identified in 59% of the cohort (24/41). In a subsequent multicenter cohort of 456 patients with RIS, the frequency of periventricular, juxtacortical, spinal cord, and infratentorial lesions was 98.7%, 90.1%, 35.2%, and 30.4%, respectively. Gadolinium-enhancing lesions were found in 82.3% of this study cohort (Fig 2). In a study of 19 patients with RIS and 20 individuals with relapsing-remitting MS (RRMS), no significant difference was found between the

<table>
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<th>Inclusion Criteria</th>
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<td>≥1 T2-hyperintense lesions on T2-weighted scans involving at least 2 of the following 4 locations: 1. Periventricular white matter 2. Cortical/juxtacortical 3. Spinal cord 4. Infratentorial</td>
<td>Neurologic dysfunction suggestive of MS based on historical symptoms and/or objective signs MR imaging abnormalities explained by other disease processes, especially considering age, vascular, toxins, or drug-related abnormalities</td>
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groups in T1-weighted and T2-weighted lesion volume or frequency and distribution of lesions. In addition to white matter lesions, cortical lesions have been described in 40% of patients with RIS in a study by Giorgio et al. The authors found that these cortical lesions involved the frontotemporal lobes most commonly and were more frequent in patients with RIS with the following: 1) oligoclonal bands in the CSF, 2) DIT on follow-up, and 3) coexistent cervical cord lesions. The unanswered question remains as to whether patients with RIS with cortical lesions have a higher MS conversion rate.

The perivenular distribution of the lesions is a key feature in distinguishing RIS from other mimickers (eg, migraines, strokes, synovitis-acne-pustulosis-hyperostosis-osteitis syndrome). A high percentage of RIS lesions show the central vein sign, specifically when higher magnetic fields and susceptibility-weighted sequences are used. The paramagnetic rim sign refers to the signal drop (ie, susceptibility artifact) at the edge of the white matter lesions in chronic demyelination. A study of 15 individuals with RIS reported the central vein sign and paramagnetic rim sign in 93% and 73% of the study population, respectively.

**Advanced Imaging**

On 3D quantitative volumetric studies, normalized whole-brain volume, normalized cerebral cortical volume, and normalized cerebellar white matter volume have been found to be significantly lower in patients with RIS compared with healthy individuals. In addition, on MR spectroscopy studies, decreased N-acetyl-aspartate/creatinine levels in lesional regions, normal-appearing white matter, and cortical gray matter have been found in patients with RIS compared with healthy controls, suggestive of early neuronal damage.

Studies using DTI have described microstructural changes in the brain MR imaging of individuals with RIS, which are confined to the lesional area; this finding is in contrast to the altered white matter integrity in both lesions and normal-appearing white matter in RRMS. De Stefano et al examined brain MRIs of 19 individuals with RIS compared with 20 with RRMS and 20 healthy controls. They found lesional magnetization transfer to be significantly lower in those with RIS than in healthy controls, but significantly higher than in those with RRMS. Normal-appearing white matter magnetization transfer values were similar in patients with RIS and healthy controls but significantly higher than normal-appearing white matter in patients with RRMS.

Giorgio et al identified lower fractional anisotropy values in white matter tracts with lesions in patients with RIS compared with healthy controls. In addition, the study compared the functional connectivity in individuals with RRMS and RIS versus healthy controls. There was no difference in resting-state network connectivity among those with RIS, RRMS, and controls; however, patients with RIS had significantly lower functional connectivity in sensorimotor and working memory subnetworks than patients with RRMS.

Similar changes were reported in advanced MR imaging of the spinal cord in RIS. In a study of 3T spinal cord MR imaging in 24 individuals with RIS and 14 healthy controls, the spinal cord

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**FIG 1.** A, Axial T2-weighted image in a 21-year-old patient presenting to the emergency department with the worst headache of her life shows left periventricular posterior fossa lesions (arrow). B, The FLAIR scan demonstrates periventricular (Dawson fingers-like) and juxtacortical lesions (arrows). C, None of the lesions show gadolinium enhancement.

**FIG 2.** A, A 32-year-old woman being evaluated for benign positional vertigo. The FLAIR scans have severe confluent periventricular disease and a few juxtacortical foci (arrows) of high signal. B, On postgadolinium scanning, numerous ring-enhancing demyelinating lesions are present, both periventricular-periatrial on the right and juxtacortical bilaterally.
cord magnetization transfer ratios were lower in patients with RIS. By contrast, there were no significant differences in the DTI metrics (fractional anisotropy and mean, perpendicular, and parallel diffusivity) between individuals with RIS and healthy controls.

Evolution to MS
A number of clinical and paraclinical features have been studied in RIS to help identify whether markers exist that portend early clinical conversion to MS. These predictors may help identify patients with high-risk RIS who might benefit from early treatment with MS disease-modifying therapies, though treatment is controversial at this juncture in time.

In a multicenter study of 451 individuals with RIS, approximately two-thirds of individuals developed new lesions on longitudinal MR imaging, while one-third of them developed CIS within 5 years of the index MR imaging. Since this publication in 2014, it has been accepted that RIS with DIS and DIT plus CIS implies a diagnosis of MS. Approximately 10% of the entire cohort eventually developed primary-progressive MS.

Patients with spinal cord lesions appear to be at particular risk of conversion, with 56% converting to MS. In 1 study of patients with RIS, 84% of individuals who had cervical cord lesions converted to MS, while only 7% of those without cervical cord lesions converted. Similarly, thoracic cord lesions, though occurring more rarely, place patients with RIS at higher risk of conversion to MS (hazard ratio, 2.23 versus 2.02 for cervical cord lesions).

In contrast to spinal lesions, periventricular and juxtacortical lesions were not found to be predictive of clinical conversion. Additionally, the number of lesions and their location in the brain are not a strong predictor of clinical conversion. A few studies have found the presence of infratentorial lesions to be predictive of clinical conversion, while others have not.

Lebrun et al suggested that the criteria for high-risk RIS should include younger age, the presence of gadolinium-enhancing lesions, and abnormal visual-evoked potentials. According to the multinational study mentioned above, which included 451 individuals with RIS (78.5% females; mean age, 37.2 years), younger age (hazard ratio, 0.98; 95% CI, 0.96–0.99), male sex (hazard ratio, 1.93; 95% CI, 1.24–2.99), and the presence of lesions within the cervical or thoracic spinal cord (hazard ratio, 3.08; 95% CI, 2.06–4.62) were identified in the multivariate analysis as the most significant predictors for the development of MS. The presence of contrast enhancement on baseline MR imaging, the location of the lesions in the brain, a family history of MS, and the CSF profile were not strong predictors of clinical conversion in this model. Older men with spinal cord lesions and RIS had a higher rate of conversion to primary-progressive MS.

CSF biomarkers have also been studied in patients with RIS. Oligoclonal bands have been the most important CSF biomarker to predict clinical conversion to MS. In the 2017 McDonald MS criteria, oligoclonal bands can replace DIT. Smaller cohorts have shown a higher risk if patients have elevated interleukin 8, neurofilament light chain, and immunoglobulin G indexes in the spinal fluid.

Lebrun et al also found a higher rate of clinical conversion to MS in pregnant patients with a significantly shorter mean conversion time.

Overall, the presence of spinal cord lesions, oligoclonal bands in CSF, and younger age is the most important predictor for conversion. Whether the presence of contrast-enhancing lesions can be a predictor of clinical conversion is controversial.

Relationship to Clinically Isolated Syndrome
Patients are considered to have CIS when they present with their first clinical symptom suggestive of CNS demyelination and do not fulfill the McDonald criteria for clinically definite MS. The neurologic event might involve the optic nerve, spinal cord, brain stem, cerebellum, or cerebral hemispheres. The clinical event needs to persist in a constant fashion for at least 24 hours and occur in the absence of fever or infection to be called a demyelinating attack. In a study of 156 individuals with CIS with a median follow-up of 7 years, 42% converted to clinically definite MS. The rate of conversion of CIS to MS appears to be higher than that in individuals with RIS. MR imaging is the most helpful tool to predict conversion to MS in patients with CIS. MR imaging abnormalities suggestive of demyelinating disease are seen in 50%–70% of individuals with CIS. The risk of conversion to clinically definite MS is approximately 60%–82% in those with abnormal MR imaging findings, in contrast to 8%–25% in those with a normal MR imaging findings. In individuals with RIS who develop CIS, the presence of DIT (eg, the presence of both gadolinium-enhancing and nonenhancing lesions) on MR imaging will fulfill the criteria for a diagnosis of MS. Several MR imaging features have been associated with a higher risk of conversion of CIS to MS, including the involvement of white matter tracts attributed to motor function, the involvement of tracts near the corpus callosum, the presence of spinal cord lesions, the development of new brain lesions, and lateral ventricle enlargement.

Differential Diagnosis
As stated in the original article of Okuda et al, the label of RIS can be applied only after excluding other causes of white matter lesions. Okuda et al added, as part of these criteria (see Table 1), “The MR imaging anomalies are not due to the direct physiologic effects of substances (recreational drug abuse, toxic exposure) or a medical condition” and “Exclusion of individuals with MR imaging phenotypes suggestive of leukoaraiosis or extensive white matter pathology lacking involvement of the corpus callosum.” Similarly, the MAGNIMS 2018 statement lists as exclusion criteria “MR imaging abnormalities explained by any other disease process, with particular attention to aging or vascular-related abnormalities, and those due to exposure to toxins or drugs.” These statements are included because an age limit has not been defined for RIS. Therefore, an elderly patient with small-vessel ischemic disease with lesions in the posterior fossa and periventricular region who develops a new ischemic focus may fulfill the McDonald 2017 and the RIS imaging criteria of Okuda et al.

When one considers the causes of white matter lesions that may mimic the RIS pattern, the list spans several entities
Table 3: Selected entities often mistaken for RIS

<table>
<thead>
<tr>
<th>Entity</th>
<th>Distinguishing Clinical Features</th>
<th>Distinguishing Imaging Features</th>
<th>Important Keys to Diagnosis</th>
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<tbody>
<tr>
<td>Toxins/drug-related</td>
<td>Altered mental status changes during intoxication, history of substance abuse</td>
<td>Deep gray matter frequently involved, symmetric lesions</td>
<td>Urine and serum toxicology tests</td>
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<td>Age-related leukoaraiosis</td>
<td>Older individual, cardiovascular risk factors</td>
<td>Small (&lt;3 mm), nonenhancing lesions in periventricular and deep white matter, coexistent</td>
<td>Lacunar disease and atypical white matter lesions for RIS in an older individual</td>
</tr>
<tr>
<td>Migraines</td>
<td>Headache or aura predominates</td>
<td>Predominantly subcortical white matter lesions that are small (&lt;3 mm) and do not enhance, few</td>
<td>Headache history and lack of typical imaging features consistent with RIS</td>
</tr>
<tr>
<td>Vasculitis</td>
<td>Episodic neurologic symptoms with superimposed strokes</td>
<td>Gray and white matter lesions coexist, may have enhancing vessel wall and/or leptomeningeal</td>
<td>Systemic symptoms present, elevated erythrocyte sedimentation rate and/or C-reactive protein level, extra-/intracranial vessels abnormal, brain biopsy</td>
</tr>
<tr>
<td>CADASIL</td>
<td>Strokelike episodes, family history of similar clinical syndrome</td>
<td>White matter disease favoring anterior temporal tip subcortical regions, external capsule,</td>
<td>Genetic testing diagnostic</td>
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<tr>
<td>Collagen vascular diseases</td>
<td>Clinical history of arthritis, long-standing chronic disease, episodic</td>
<td>Gray and white matter lesions ± vasculitis, occasional encephalitis</td>
<td>Clinical symptoms of a systemic disorder and presence of serologic autoantibodies/ inflammatory markers</td>
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<tr>
<td>ADEM</td>
<td>Encephalitis, seizures, children &gt; adults, history of viral/vaccine prodrome</td>
<td>Gray matter disease predominates, more diffuse enhancement, may have positive findings on DWI</td>
<td>History of prodromal virus infection or vaccination, encephalopathic</td>
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<tr>
<td>Posttraumatic</td>
<td>History of ≥1 traumatic, sports-related event</td>
<td>Favors gray-white matter junction, hemorrhagic products present, classic tears in splenium-brain stem deep gray matter</td>
<td>Hemorrhage and stereotypical locations of disease at shearing sites</td>
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Note:—ADEM indicates acute disseminated encephalomyelitis.

including vascular (migraine changes, vasculitis, CADASIL, neonatal hypoxic-ischemic injuries, anoxic episodes, and small-vessel ischemic disease), inflammatory/infectious entities (sarcoidosis, lupus, acute disseminated encephalomyelitis, progressive multifocal leukoencephalopathy, synovitis-acne-pustulosis-hyperostosis-ostitis syndrome), posttraumatic changes (remote clinical and subclinical trauma, sports-related, ie, boxers), drug-related (splenial demyelination in seizure medications versus causes of posterior reversible encephalopathy syndrome), and idiopathic (Table 3).39 Two series40,41 of “misdiagnosed MS” reported the most common final diagnoses to be migraine, fibromyalgia, psychiatric/psychogenic disease, nonspecific white matter abnormalities, small-vessel ischemic disease, and neuromyelitis optica spectrum disorders. As opposed to these entities described above, which have overt signs and symptoms, patients with RIS are, by definition, asymptomatic.

Despite this very broad differential diagnosis, subspecialty-trained neuroradiologists have been shown to be very accurate in recognizing the pattern of white matter lesions suggestive of MS and therefore RIS. In a study performed at a large academic center,42 the order in which MS was suggested in a differential diagnosis of the neuroradiologist’s report determined the likelihood of a final diagnosis in patients who did not previously have a diagnosis of MS. When MS was listed as the only diagnosis, the neuroradiologists were correct in the final diagnosis in 22/24 cases (92.3%). When “demyelinating disorder” only was listed, they were correct in 43/51 cases (84.3%), and when MS was listed as the first potential diagnosis in a list of differentials, the final diagnosis was proved to be MS in 38% (3/8) of cases. The final diagnoses in those patients inaccurately reported as “likely MS” included migraines (n = 9), peripheral neuropathy (n = 4), postconcussive lesions (n = 3), substance abuse (n = 2), cerebrovascular disease (n = 2), and no diagnosis (n = 5).42

The imaging findings that are more suggestive of RIS than other entities include the central vein sign of demyelination around a vein and involvement of the corpus callosum (corpus callosum–septum pellucidum interface). The original RIS diagnostic criteria of Okuda et al” excluded patients if they were “lacking involvement of the corpus callosum.” Moreover, spinal involvement also helps to practically exclude diagnoses like migraines, posterior reversible encephalopathy syndrome, progressive multifocal leukoencephalopathy, and so forth. A good review of the accuracy of MR imaging for MS alone and in combination with clinical findings, serologic tests, and CSF markers can be found in the study by Schaffler et al.43
Clinical Approach to RIS

Despite RIS being described more than a decade ago, there are no specific, universally adopted guidelines for the monitoring and treatment of this evolving entity. This issue is mostly secondary to a paucity of available longitudinal studies and lack of expert consensus on how one should approach such patients. Hence, clinicians are left with relying on short studies, review articles, and their anecdotal experiences.

An initial pragmatic approach that many clinicians take after an individual fulfills the criteria for RIS is similar to how clinicians evaluate and monitor patients with suspected CIS and MS. First, a detailed history and physical examination by an experienced neurology clinician are recommended to ensure that a patient truly has RIS, especially because treatment is recommended for most patients with CIS and MS. The emphasis, therefore, is to discover prior or current neurologic signs and symptoms that might fulfill the CIS or MS criteria. At this juncture, most clinicians are hesitant to start patients with RIS on prophylactic disease-modifying therapies because a substantial number of patients with RIS will not develop MS and there are potential serious risks associated with MS therapies.

Obtaining paraclinical testing beyond an MR imaging is also extremely important because this can help stratify patients with RIS into low-risk (eg, no spine lesions, normal lumbar puncture) or high-risk (eg, spine lesions, presence of CSF-restricted oligoclonal bands) for conversion to MS. Hence, most neurology clinicians will recommend a lumbar puncture to assess markers of autoimmunity and inflammation, including CSF pleocytosis, an elevated immunoglobulin G index, and/or CSF-restricted oligoclonal bands. In addition, just like with MS, serologic testing is necessary for ruling out mimickers of demyelinating disorders (systemic/rheumatic disorders, nutritional deficiencies, infectious diseases, and so forth) and for identifying conditions that have a stronger propensity to cause disabling neurologic attacks (eg, seropositive neuromyelitis optica spectrum disorders, anti-myelin oligodendrocyte glycoprotein–associated disorders, and so forth).

Other tests that are commonly performed that may be associated with the prognosis of patients with RIS include visual-evoked potential testing and optical coherence tomography. Electromyography, nerve conduction studies, and other ancillary studies are typically not performed unless there are “red flags” in the history and/or examination suggesting an alternative diagnosis (eg, peripheral neuropathy).

Most clinicians will also obtain a total serum 25-hydroxyvitamin D level on their patients because chronic vitamin D deficiency is associated with a risk of developing MS. If a patient with RIS has low vitamin D, vitamin D supplementation is recommended.

After the initial assessment and work-up for RIS is completed, clinicians are then challenged by how patients with RIS should be monitored across time. Currently, there is a paucity of data that help guide clinicians on when surveillance imaging should be performed in RIS, especially as it relates to assessing evidence of DIT.

The MAGNIMS group and the Consortium of Multiple Sclerosis Centers MR Imaging Task Force have published expert consensus guidelines that can be followed. Future and ongoing prospective longitudinal studies will help refine these guidelines along with the advent of newer imaging techniques (central vein sign, paramagnetic rim sign) and other biomarkers (eg, neurofilament light chain).

At this time, the treatment approach for patients with RIS is controversial. As previously highlighted, a substantial number of patients with RIS will not develop CIS or MS, especially those who are considered to have low-risk RIS. A recent comprehensive review article on RIS discussed the importance of clinical trials in RIS because data are lacking establishing the benefit of starting treatment in RIS. Hence, treatment is not recommended for most upfront, though if new inflammatory activity (ie, gadolinium-enhancing lesions) is seen on follow-up MR imaging, many neurologists would strongly consider starting therapy.

In 1 retrospective study in which a small cohort of patients with RIS received MS disease-modifying therapies, the 5-year risk of developing an MS-defining clinical event was 45% (33/73 patients) in those receiving disease-modifying therapies versus 31% (117/378) in those who did not receive disease-modifying therapies. There are several RIS randomized clinical trials ongoing that are evaluating whether starting dimethyl fumarate or teriflunomide will prevent or delay conversion to clinically definite MS versus a placebo.

RIS in Pediatrics

There are limited data on the prevalence and clinical significance of RIS in children. In a review of brain MRIs in 3966 pediatric patients, only 1 patient (0.03%) was identified as fulfilling the RIS criteria. In a longitudinal multicenter study of 38 pediatric patients with a median age of 15 years who were diagnosed with RIS based on the McDonald 2010 DIS criteria, 42% experienced a first clinical neurologic attack with a median interval of 2 years. In addition, radiologic progression was reported in 61% of the study cohort. These reported numbers are very similar to those of RIS in adult patients; however, the study found that clinical conversion to MS occurs after a shorter interval in pediatric patients compared with adults. Most important, the authors also reported that the presence of ≥2 CSF-related oligoclonal bands or spinal cord lesions was a significant predictor of clinical conversion.

The conventional MR imaging findings in children are not different from those seen in adult patients with RIS. Because there are sparse data available on RIS in children, there is not sufficient evidence to conclude that these patients could potentially benefit from early disease-modifying treatment.

CONCLUSIONS

There are patients without a defined neurologic event who have MR imaging findings that fulfill the imaging criteria for MS. After an extensive work-up to exclude other neurologic, systemic, or iatrogenic etiologies, these patients are labeled as fulfilling the criteria for RIS. At baseline and in follow-up, the presence of imaging factors (spinal lesions, enhancing lesions, corpus callosum lesions), demographic characteristics (men, pregnant women, younger age, relative of a patient with MS), and/or CSF markers (oligoclonal bands) incurs a higher risk of subsequent conversion to MS. Radiologists who identify patients with classic MS-like lesions in the appropriate clinical and patient group settings should suggest...
the RIS entity. The surveillance of patients with RIS, the agent and extent of treatment, and long-term outcomes are subjects of ongoing debate and clinical trials. More studies are also needed with advanced MR imaging techniques to investigate whether novel technologies being developed will help identify which patients with RIS are at greatest risk for conversion to clinically definite MS.


REFERENCES

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