Multiple sclerosis in children: value of serial MR studies to monitor patients.

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Multiple Sclerosis in Children: Value of Serial MR Studies to Monitor Patients

A series of six children with clinical (4) and laboratory (2) evidence of multiple sclerosis is described. The mean age at onset was 12 years and the female-male ratio was 5:1. All patients had white matter abnormalities on initial MR scans. On follow-up MR studies, performed every 3 to 5 months, all children exhibited changing patterns of CNS signal abnormalities. In three cases, clinically silent brain lesions were detected. In four patients with an acute clinical attack, large lesions were present, demonstrating a lamellar structure on T1- and T2-weighted images. The lesions were seen best on long TR/short TE spin-echo sequences. Combined sagittal and axial series with EKG gating and flow-compensation technique were best for MR follow-up studies.

Our results show that MR is useful for monitoring patients with multiple sclerosis.

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Multiple sclerosis (MS) is diagnosed clinically by the demonstration of white matter dysfunction separated in time and anatomic location [1]. MR imaging detects white matter abnormalities of MS and has been used to identify multiple anatomic sites of the disease [2-6]. And serial MR examinations can be used to identify serial episodes of the disease [7-11]. The aim of the present study was to assess CNS focal white matter abnormalities in childhood MS and to relate changes in their appearance over time with the clinical course of the disease.

Subjects and Methods

Between January 1987 and January 1989, 341 children, ages 2–14 years old, were referred for MR of the brain for evaluation of various neurologic deficits. One hundred seventy-nine children were in the 2–9 age group, and 162 children were in the 10–14 age group. There were 187 boys and 154 girls. The clinical work-up of patients with focal CNS white matter abnormalities revealed the presence of MS in six patients, all in the 10–14 age group. No case of MS was observed in the 2–9 age group.

MR was performed on a superconductive 1.5-T magnet system (Gyros can S 15, Philips Company Eindhoven, Netherlands). T1-weighted images were obtained by using inversion recovery (IR) pulse sequences, 1500/450/30/4 (TR/TE/excitations) and partial saturation spin-echo (SE) pulse sequences (450-600/20/4). T2-weighted images were obtained with a conventional spin-echo technique (multislice-multiecho sequence [SE 2500/30,60/2]) and with a gradient-echo technique (reduced flip angle of 15° and the shortest TR/TE). Sagittal long TR/long TE SE sequences were obtained with EKG gating and flow-compensation technique. For data acquisition, a matrix of 128 or 256 × 256 was used. The thickness of the slices ranged from 5.3 to 6 mm. In general, the slice gap was 10% of the slice thickness. In all patients imaging was performed in transverse and sagittal planes and in some cases coronal sections were also obtained. MR investigations of the cervical spine were done using a rectangular surface coil. Surface-coil imaging of the optic nerve was performed in two patients (STIR sequence) [12]. As gadopentetate dimeglumine was not approved for use in children, contrast scans were not done at that time.
Results

The average age of onset of MS in the present series was 12 years; the female-male ratio was 5:1. Table 1 gives the clinical presentation, CSF analysis, visual evoked responses (VER), brainstem auditory evoked responses (BAER), and EEG results. According to the clinical classification of Poser et al. [13], four children had clinically definite MS and two had laboratory supported definite MS.

The patients presented clinically with motor symptoms (4/6), visual disturbances (2/6), urinary dysfunction (2/6), gait disturbances (1/6), vertigo (1/6), and nystagmus (1/6). Of all paraclinical tests (see Table 1) MR proved to be the most useful for establishing the clinical diagnosis of MS. Clinically definite MS was diagnosed in four patients and laboratory supported definite MS in two patients. The time interval between the first clinical attack and the first examination by MR ranged from 2 weeks to 6 months (average, 5 weeks). Initial MR imaging detected an average of 16 lesions per patient (range, 3–50 lesions); follow-up MR, performed after an interval of 3 months (range, 7–13 weeks), found an average of 21 lesions per patient (range, 3–85 lesions). A third MR examination 5 months later (range, 6 weeks–12 months) demonstrated an average of 27 lesions per patient (range, 3–115 lesions). In four of six patients, a fourth MR study 8 months later showed a further increase in the number of lesions. Of six patients with follow-up MR, three had no clinical attack to suggest new MS plaques in the brain. Therefore, in three of six patients clinically silent lesions and/or progression of the demyelinating process was detected by MR. The MS lesions were located most frequently in the centrum semiovale (6/6), optic radiation (3/6), basal ganglia (3/6), brainstem (3/6), cervical medulla (3/6), corpus callosum (2/6), subcortical white matter (2/6), cerebral gray matter (1/6), and cerebellar peduncle (1/6). On initial MR examination all patients had brain white matter abnormalities that fulfilled the criteria of the MR classification of MS developed by Paty et al. [14].

On follow-up examinations all patients exhibited changing patterns of the multifocal CNS signal abnormalities: increase and decrease in number and size of MS foci, disappearance of lesions, lesions found in completely new locations, and confluence of the lesions (Figs. 1–4).

In four patients with acute clinical attack, large lesions (8–21 mm in diameter) were present, demonstrating a lamellar structure on both T1- and T2-weighted images (Figs. 1 and 2A).

The large foci all demonstrated a central spherical core of low signal on T1-weighted images, isointense with CSF. This core was surrounded by a thick rim of high signal on T1- and proton-density-weighted images. In two children with a fresh demyelinating process, extensive perifocal white matter edema was demonstrated on MR (Fig. 1B). On follow-up this edema had resolved after an interval of 11 weeks (case 1) and 6 weeks (case 3). The large lesions in cases 1, 3, 4, and 6 showed a gradual decrease in size but still kept their anatomic characteristics during the observation period of 19 weeks (case 3), 26 weeks (case 4), and 15 months (cases 1 and 6).

All lesions had the highest signal/noise ratio compared with CSF and gray and white matter on proton-density-weighted, long TR/short TE SE sequences.

### TABLE 1: Case Description

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Sex</th>
<th>Age at Onset (years)</th>
<th>First Symptoms</th>
<th>Oligoclonal Bands (OB)</th>
<th>CSF WBC</th>
<th>Protein</th>
<th>Visual Evoked Responses (VER)</th>
<th>Brainstem Auditory Evoked Responses (BAER)</th>
<th>EEG</th>
<th>Diagnostic Criteria (Poser et al. [13])</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>11</td>
<td>Weakness of left hand, vertigo</td>
<td>+</td>
<td>24</td>
<td>33</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Laboratory supported definite MS*</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>13</td>
<td>Blurred vision, staggering gait, urinary dysfunction</td>
<td>-</td>
<td>24</td>
<td>27</td>
<td>Normal&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Normal</td>
<td>Normal</td>
<td>Clinically definite MS&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>13</td>
<td>Spastic hemiplegia, nystagmus</td>
<td>+</td>
<td>27</td>
<td>39</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal</td>
<td>Laboratory supported definite MS&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>13</td>
<td>Divergent strabismus</td>
<td>+</td>
<td>16</td>
<td>23</td>
<td>Abnormal</td>
<td>Normal</td>
<td>Normal</td>
<td>Clinically definite MS&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>11</td>
<td>Enuresis, weakness, paresthesia of lower limbs</td>
<td>-&lt;sup&gt;c&lt;/sup&gt;</td>
<td>39</td>
<td>28</td>
<td>Abnormal</td>
<td>-</td>
<td>Sharp waves</td>
<td>Clinically definite MS&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>13</td>
<td>Weakness of left arm</td>
<td>+&lt;sup&gt;b&lt;/sup&gt;</td>
<td>29</td>
<td>58</td>
<td>Normal</td>
<td>-</td>
<td>Normal</td>
<td>Clinically definite MS&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>7*</td>
<td>F</td>
<td>12</td>
<td>Diplopia, vertigo, staggering gait</td>
<td>+</td>
<td>25</td>
<td>27</td>
<td>Abnormal</td>
<td>Normal</td>
<td>Normal</td>
<td>Clinically definite MS&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> First investigation of OB 12 months after onset.
<sup>b</sup> VER normal but computerized perimetry abnormal.
<sup>c</sup> Elevated lgG but no OB.
* Recent case of MS not included in present study.
Fig. 1.—Case 1: 11-year-old girl with laboratory supported definite MS.
A, Coronal T1-weighted (700/30) spin-echo MR image 4 weeks after first clinical attack shows three separate lesions with lamellar structure. (i = inner zone of the lesion, o = outer zone)
B, Axial T2-weighted (2500/60) spin-echo MR image shows large lesion with surrounding edema. (i = inner zone, o = outer zone, e = edema)
C, Axial proton-density-weighted (2500/30) spin-echo MR image 11 weeks after first clinical attack shows lamellar structure of the lesion in the right centrum semiovale (large arrow). There was complete resorption of the previous edema.
D, Axial proton-density-weighted (2500/30) spin-echo MR image 15 months after first clinical attack shows further decrease in the size of the large lesion (large arrow). There was persistence of the lamellar structure.

Fig. 2.—Case 3: 13-year-old girl with laboratory supported definite MS. MR shows the changing appearance of the acute and chronic MS focus on serial images.
A, Sagittal T1-weighted (600/30) spin-echo MR image shows lamellar structure of acute MS plaque. Diameter of lesion is 21 mm. (i = inner zone, o = outer zone)
B, Sagittal T2-weighted (600/35) gradient-echo MR image (flip angle = 15°) 1 year later shows decreased size and hazy appearance of chronic MS plaque (arrow) without any inner structure.

Combined series in axial and sagittal planes proved best for serial comparison of the distribution and number of focal white matter abnormalities on MR studies. Even small foci (less than 3 mm in diameter) could be reassessed on multiple follow-up examinations.

In case 5, a patient with clinically rapid progression, confluence of the periventricular lesions was observed and significantly worsened during the observation period of 6 months (Fig. 3).

Discussion
The frequency of MS with known onset in childhood ranges from less than 0.4% of all MS cases [15] and 2.7% [16] and
6% [17]. Early onset of MS has been described at 4 years [18] and at 2 years [19], and a patient with autopsy-verified MS, described by Shaw and Alvord [20], reportedly had the first of 11 attacks at the age of 10 months.

The female-male ratio in adults with MS is reported to be about 2:1; but in children it has been reported at 3:1 [16], 4:1 [21], and 5:1 in the present study. The frequency of MS in our population seems high in this 2-year observation period, although exact figures cannot be derived from this selective group of patients referred for MR. Whether this high frequency is due to a previously unrecognized early onset or represents an epidemic cluster of childhood MS cannot be determined at this time.

Initial clinical symptoms of MS in childhood are sensory symptoms (26.4%), optic neuritis (14%), diplopia (11%), pure motor symptoms (11%), gait disturbance (8%), blurred vision (6%), cerebellar ataxia (5%), sensory and motor symptoms (5%), and sphincter problems (0.8%) [16]. Brett [22] called optic neuritis in childhood a potential harbinger of MS, because isolated optic neuritis often is found to be the first clinical manifestation of MS [22–25]. Compston et al. [26] reported a conversion rate from optic neuritis to clinically definite MS of 60% within 8 years, Rizzo and Lessell [27] reported 74% and Francis et al. [28] reported 75% within 15 years. Dementia processes after rapid progression of childhood MS and psychotic symptoms were also described [19, 21]. In a few reports [29–32] the initial symptoms resembled an acute encephalopathy, and sometimes MS presented clinically and radiologically as a brain tumor or abscess. Mild headache, dizziness, nausea, vomiting, and vertigo [15] sometimes accompany the first clinical attack. Focal abnormalities of CNS white matter are encountered frequently in MR of the aging brain and in patients with cerebrovascular or cardiac risk factors. In the pediatric population MR white matter abnormalities are reported in acute disseminated encephalomyelitis (ADEM), progressive multifocal leucoencephalopathy, CNS lymphoma, mitochondrial encephalomyopathy, herpes simplex encephalitis, mucopolysaccharidosis, homo- and heterozygote adrenoleukodystrophy, methotrexate-encephalopathy, neurosarcoidosis, and others [33]. Thus, the MR findings are nonspecific. Sequential MR examinations, however, are capable of assessing changes in number, size, and configuration of white matter abnormalities. Under the prerequisite of careful reassessment, this changing pattern can suffice for the diagnosis of a disseminated disease. Serial MR cannot only prove dissemination in space but also dissemination in time, demonstrating clinically silent MS lesions in the brain.

Serial MR cannot rule out ADEM, since a remitting/relapsing course in some cases of ADEM has been described, but usually a differentiation is not a problem by clinical means [14]. If high standards of consistency are placed on patient positioning within the imaging system, comparative imaging might be easily and reliably accomplished in any desired plane. In our experience the most exact reassessment of previously assessed lesions in repeated MR studies was achieved by combining sagittal and transverse sections using exactly the same pulse parameters, slice thickness, and number of slices. Flow-compensation technique and EKG gating should be applied.

Pathologically, the acute MS plaque comprises perivenous hypercellularity. Around the inflamed vessels, large spaces between myelinated axons suggest edema. Lymphocytes are confined to perivascular spaces or may spread throughout the surrounding parenchyma [34]. In fresh MS plaques myelin breakdown products, largely lipids, are found free and in macrophages [35–37]. At the edges of the lesions numerous fat-filled macrophages are found. This pathoanatomical description matches the MR morphology of acute MS foci, observed in our series. The center of low signal seen on T1-weighted images, of moderate signal seen on mixed sequences, and of high signal seen on T2-weighted images possibly represents the perivascular infiltration with lymphocytes. The surrounding rim of high signal on T1- and proton-density-weighted images probably corresponds to fat-laden macrophages or to free myelin products. Thus, MR could be used to assess the in vivo progression and regression of fresh demyelinating plaques. In two cases of our series the rim of high signal on
T1-weighted images was demonstrable during a period of 15 months (cases 1 and 6).

Anatomically, the chronic MS plaque is characterized by proliferation of glial tissue [34]. The degree of gliosis contributes significantly to the intensity of the MR signal [8]. On MR the chronic MS plaques show up as smaller homogeneous lesions without any inner structure. The best contrast/noise to surrounding white matter is shown on long TR/short T2SE sequences.

Acute plaques in MS can mimic the radiologic signs of a brain tumor or abscess. In two children having their first attack we observed large tumorlike foci on cranial CT and MR. The presence of a tumorlike focus in the brain on CT demands screening for additional, subtle, or invisible smaller lesions with MR in order to avoid unnecessary brain surgery.

In summary, serial MR was capable of detecting silent brain lesions without concomitant clinical attacks and to unfold the disseminating character of the given disease process. In four of six patients in our series, very large, fresh demyelinating plaques were encountered, which demonstrated a lamellated structure persisting over a long time. In all these patients MR was able to monitor the regression of the acute MS plaque.

In acute MS, MR offers a tool to evaluate the patient’s response to drug therapy.

REFERENCES