Multiple sclerosis in adolescents: CT and MR findings.

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Multiple Sclerosis in Adolescents: 
CT and MR Findings

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The MR and CT findings in 12 adolescents with multiple sclerosis were compared with reported findings in adults. The adolescent group showed a more striking female predominance, more severe disease characteristics, and more frequent infratentorial involvement. Cortical atrophy and abnormal iron accumulation in the basal ganglia were uncommon in the adolescents. Neither group demonstrated a correlation between symptom severity and either extent or location of disease as delineated by MR imaging. MR was more sensitive than CT in detecting demyelinating plaques.


Multiple sclerosis (MS) is a rare disorder in children and adolescents, with this age group accounting for less than 1% of all cases [1]. Vague symptoms and an often erratic clinical course in this age group made it difficult to establish a diagnosis prior to the advent of MR imaging. While a few scattered case reports have appeared in the literature [2–4], no series has delineated the appearance of childhood MS on high-resolution CT and MR studies. We present the imaging spectrum based on a series of 12 documented cases.

Subjects and Methods

MR images obtained during the period from January 1985 to April 1989 of all children and adolescents either referred to the University of Utah for examination because of clinically suspected MS or whose scans themselves were considered suggestive of demyelinating disease were reviewed. Fourteen such cases were found; 12 had CT/MR findings that suggested MS. Ten of the 12 patients were female (Table 1), and five were 17 years old or younger at the time of initial imaging. Seven patients had symptom onset in their mid to late teens but were 18 or older at the time of imaging. Only two patients, among the oldest in the series, had a definite clinical diagnosis of MS prior to MR scanning. Two other patients were referred for suspected MS but were found to have nondemyelinating disorders (one had a high cervical disk herniation with cord compression and the other had progressive idiopathic arteriopathy of childhood). The remaining 12 patients either had CSF studies consistent with MS (presence of oligoclonal banding or evidence of increased immunoglobulin synthesis rates), abnormal visual or somatosensory evoked potentials, or a subsequent relapsing-remitting or chronic-progressive course consistent with clinically definite MS.

Eleven patients had both plain and contrast-enhanced high-resolution axial CT scans. A double-dose delayed scan was obtained in one patient. Axial and sagittal T1- and T2-weighted axial MR scans were obtained at 1.5T in all 12 patients. In one patient, contrast-enhanced MR images were also obtained. MR scans were evaluated for foci of abnormal signal in the periventricular regions, both hemispheres, cerebellum, brainstem, and (in four patients) the spinal cord. The thalamus, putamen, and caudate nuclei were assessed for the presence of abnormally decreased signal on T2-weighted images. The MR index developed by Uhlenbrock et al. [5] was modified and used to assess disease severity in each case (Table 2).
TABLE 1: Case Material

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Sex</th>
<th>Age at First Symptoms</th>
<th>Age at First Scan</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>16</td>
<td>17</td>
<td>18-month history of intermittent right-sided numbness, diffuse weakness</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>14</td>
<td>15</td>
<td>Vague neurologic complaints of diffuse numbness, upper extremity weakness</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>17</td>
<td>20</td>
<td>3-year history of bilateral upper extremity tingling</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>16</td>
<td>16</td>
<td>2-week history of right arm and leg numbness and tingling, weakness of left arm and leg (CSF was positive for oligoclonal banding)</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>17</td>
<td>21</td>
<td>Transient left facial numbness for 4 years; 3-week history of ataxia, diplopia</td>
</tr>
<tr>
<td>6</td>
<td>F</td>
<td>16</td>
<td>22</td>
<td>5-year history of progressive neurologic deficits, clinical diagnosis of MS; new onset of temporal visual field defects</td>
</tr>
<tr>
<td>7</td>
<td>F</td>
<td>18</td>
<td>19</td>
<td>Visual symptoms only</td>
</tr>
<tr>
<td>8</td>
<td>F</td>
<td>17</td>
<td>21</td>
<td>4-year history of visual difficulties, extremity numbness, speech difficulties, loss of bowel and bladder control; clinical diagnosis of MS; recent-onset seizures</td>
</tr>
<tr>
<td>9</td>
<td>F</td>
<td>16</td>
<td>22</td>
<td>6-year history of intermittent, vague neurologic complaints</td>
</tr>
<tr>
<td>10</td>
<td>F</td>
<td>17</td>
<td>17</td>
<td>&quot;Flu&quot; followed by lethargy; facial numbness, hand incoordination</td>
</tr>
<tr>
<td>11</td>
<td>M</td>
<td>17</td>
<td>18</td>
<td>Seizures, bilateral Babinski reflexes; CSF positive for oligoclonal bands</td>
</tr>
<tr>
<td>12</td>
<td>M</td>
<td>15</td>
<td>15</td>
<td>10-day history of left upper lip, cheek, face, scalp numbness; CSF positive for oligoclonal bands</td>
</tr>
</tbody>
</table>

Results

CT

Eleven patients had plain and contrast-enhanced brain CT scans (Table 3). When present, lesions were typically hypo- or isodense on plain scans, with variable enhancement after contrast administration. Differing degrees of plaque enhancement were often seen in the same patient varying from no enhancement in some low-density lesions to ring or inhomogeneously increased density in others. Double-dose delayed scanning was performed in one patient and disclosed additional lesions not identified on the immediate postcontrast study (Fig. 1). The three patients with spinal cord lesions who had negative brain MR studies also had normal CT brain scans. In all patients, more lesions were seen on MR than on CT (Table 3).

MR Imaging

Four patients had symptoms of a spinal cord lesion and thus had cord scans; all four had cord lesions (Table 3). Three of these patients had a solitary focus of increased signal in the cervical cord and normal brain studies (Fig. 2); CSF was positive for oligoclonal bands in all three. The fourth patient had both thoracic and cervical cord lesions in addition to multiple lesions in the periventricular deep white matter characteristic of long-standing MS. Two of the patients had mild mass effect associated with their cord lesions.

The remaining eight patients had multiple brain scans, but no spinal cord studies were performed. All but one (case 11) had supra- and infratentorial lesions (Fig. 3) with most having severe disease as evidenced by multiple confluent periventricular plaques and extensive white matter involvement (Tables 2 and 3). Lesions were typically iso- or hypointense on T1-weighted images and hyperintense on T2-weighted studies (Fig. 4). Beveled or ring lesions with mixed signal on T1-weighted scans and various degrees of increased signal on proton-density scans were also common, as were ovoid plaques (Fig. 1D) that have recently been described in MS [6]. In the one patient who received MR contrast material, additional lesions were identified on the postcontrast study that were not seen on proton-density or T2-weighted studies.

One patient with long-standing severe disease had cortical atrophy as well as abnormally decreased signal in the thalamus, putamen, and basal ganglia on T2-weighted images (Fig. 5); none of the other 10 cases had evidence of these associated abnormalities. One patient had a biopsy-proved hypothalamic glioma in addition to florid MS.

TABLE 2: Modified MR Index for Assessing MS Plaques*

A. Periventricular plaques
   A0. No plaques
   A1. No more than five plaques, with single plaque not larger than 8 mm, or one plaque larger than 8 mm and no more than three plaques; no confluence
   A2. More than five plaques, more than three plaques with one plaque larger than 8 mm, or two or more plaques larger than 8 mm; in addition or exclusively, there is periventricular confluence but the bodies of the ventricles are mainly unaffected
   A3. More than 12 plaques and/or periventricular confluence, including the bodies of the ventricles

B. Hemispheric plaques
   B0. No plaques
   B1. No more than five plaques
   B2. More than five plaques

C. Cerebral atrophy
   C1. White matter atrophy
   C2. Focal cortical atrophy
   C3. Diffuse cortical atrophy
   C4. White matter and cortical atrophy

D. Cerebellar plaques
   D0. No plaques
   D1. One or more plaques

E. Brainstem plaques
   E0. No plaques
   E1. One or more plaques

F. Spinal cord plaques
   F0. No plaques
   F1. One or more plaques

G. Abnormal iron deposition in thalamus, putamen, and/or caudate nuclei
   G0. Normal signal
   G1. Decreased signal

* After Uhlenbrock et al. [5]; category added for abnormal nonheme iron deposition.
TABLE 3: Radiologic Findings

<table>
<thead>
<tr>
<th>Case No.</th>
<th>CT Findings</th>
<th>MR Findings</th>
<th>MR Index*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Brain: normal</td>
<td>Cord: normal intramedullary focus of increased signal on T2-weighted image at C6; mild mass effect</td>
<td>A0, B0, C0, D0, E0, F1, G0</td>
</tr>
<tr>
<td>2</td>
<td>Brain: single lesion with mild enhancement</td>
<td>Brain: multiple confluent periventricular lesions with deep white matter and brainstem lesions. Cord: thoracic cord lesions, T2–T4</td>
<td>A3, B2, C0, D0, E1, F1, G0</td>
</tr>
<tr>
<td>3</td>
<td>Brain: normal</td>
<td>Cord: solitary intramedullary focus of increased signal at C3–C4; no mass effect</td>
<td>A0, B0, C0, C0, E0, F1, G0</td>
</tr>
<tr>
<td>4</td>
<td>Brain: normal</td>
<td>Brain: normal Cord: single midcervical intramedullary lesion; mild mass effect</td>
<td>A0, B0, C0, D0, E0, F1, G0</td>
</tr>
<tr>
<td>5</td>
<td>Brain: none</td>
<td>Brain: multiple large brainstem, cerebellar, and confluent periventricular lesions, iso- or hypointense on T1-weighted image, hyperintense on T2-weighted image</td>
<td>A3, B2, C0, D1, E1, G0</td>
</tr>
<tr>
<td>6</td>
<td>Brain: multiple enhancing plaques, deep cerebral white matter; enhancing suprasellar mass</td>
<td>Brain: brainstem, cerebellar, and confluent periventricular lesions, iso- or hypointense on T2-weighted image, hyperintense on T2-weighted image; hypothalamic glioma</td>
<td>A3, B2, C0, D1, E1, G0</td>
</tr>
<tr>
<td>7</td>
<td>Brain: single enhancing pontine lesion</td>
<td>Brain: multiple confluent periventricular and brainstem lesions of variable signal on T1-weighted image, hyperintense on T2-weighted image</td>
<td>A3, B2, C0, D1, E1, G0</td>
</tr>
<tr>
<td>8</td>
<td>Brain: multiple enhancing lesions, centrum semiovale</td>
<td>Brain: severe confluent periventricular, brainstem, cerebellar, and medullary plaques; generalized cortical atrophy; corpus callosum atrophy; low signal in basal ganglia</td>
<td>A3, B2, C0, D1, E1, G1</td>
</tr>
<tr>
<td>9</td>
<td>Brain: multiple periventricular low-attenuation lesions, variable enhancement</td>
<td>Brain: discrete brainstem and periventricular lesions, hypointense on T1-weighted image, hyperintense on T2-weighted image</td>
<td>A2, B2, C0, D1, E1, G0</td>
</tr>
<tr>
<td>10</td>
<td>Brain: multiple iso-or hypointense lesions with variable enhancement on double-dose delayed scans</td>
<td>Brain: mixed-signal lesions, deep white matter, brainstem, and cerebellum</td>
<td>A3, B2, C0, D1, E1, G0</td>
</tr>
<tr>
<td>11</td>
<td>Brain: normal</td>
<td>Brain: scattered foci of increased signal in perivenricular white matter; two to three lesions not seen on T2-weighted image enhanced after contrast; posterior fossa normal</td>
<td>A1, B1, C0, D0, E0, G0</td>
</tr>
<tr>
<td>12</td>
<td>Brain: normal</td>
<td>Brain: scattered supratentorial periventricular nonconfluent foci of increased signal; solitary lesion at left fifth cranial nerve root in posterior fossa</td>
<td>A1, B2, C0, D0, E1, G0</td>
</tr>
</tbody>
</table>

* See Table 2 for descriptions.
Discussion

Several striking differences between MS in adults and in our series of adolescents were readily apparent. The female to male ratio in most adult MS series varies from 1.4:1 to 1.9:1. By contrast, 10 of our 12 patients with disease onset in adolescence were female.

In the large series of adult cases reported by Uhlenbrock et al. [5], lesion severity as rated on their MR index varied widely from normal to severe. While three of our four adolescents with spinal cord lesions had normal brain MR studies, all but two of the nine cases with positive brain scans had severe disease characterized by the presence of either more than 12 plaques or confluent periventricular lesions (both males in our series had mild disease).

While cerebellar and brainstem lesions were seen in only one third and 14% of adults, respectively, in the Uhlenbrock series, seven of the 10 adolescents had cerebellar plaques and eight of 10 had brainstem disease even though symptoms of posterior fossa involvement were often lacking. In contrast, cortical and white matter atrophy were seen in only one of the adolescents, manifested by sulcal enlargement and thinning of the corpus callosum. The relative lack of brain atrophy may reflect comparatively short disease duration in spite of its severity (30–70% of all patients with MS require at least ambulatory assistance 15 years after symptom onset) [7].

Other investigators report the presence of abnormally decreased signal intensity in the thalamus or putamen on T2-weighted images in 90% of adults with MS [7], probably secondary to nonheme iron deposition. Only one of the 12 adolescents had such findings; this patient had the most severe, long-standing disease in our series.

While we did not actually grade clinical impairment by using the Kurtzke score, severity of symptoms in our series did not correlate with disease extent as seen on MR scans (Tables 1 and 3). Some patients (e.g., case 7) with severe disease on MR had minimal symptoms. This lack of correlation between clinical data and imaging findings is consistent with the results reported for adults [8, 9].
Fig. 2.—Case 3: 20-year-old woman with 3-year history of bilateral upper extremity paresthesias; normal brain scan. Sagittal MR image (2500/80/1) shows solitary focus of increased signal in midcervical cord (arrow).

Fig. 3.—Case 6: 22-year-old woman with 5-year history of facial numbness; recent onset of ataxia, diplopia, and visual-field defects. Coronal MR image (2500/20/1) shows extensive confluent periventricular deep white matter and brainstem disease (arrows).

Fig. 4.—Case 9: 22-year-old woman with 6-year history of vague neurologic complaints. A, Axial MR image (600/20/2) shows low signal foci in brainstem (arrows). B, Hypointense lesions seen in A become hyperintense (arrows) on this T2-weighted scan (2800/70/1).

Fig. 5.—Case 8: 21-year-old woman with 4-year history of progressive visual difficulty, upper extremity numbness, and urinary urgency and frequency; recent onset of seizures. Presumptive diagnosis of MS (CSF positive for oligoclonal bands) was questioned because seizures are uncommon in MS. A and B, Axial (A) and coronal (B) MR scans (2500/70/1) show extensive confluent periventricular and deep white matter lesions. Note striking low signal in basal ganglia (outlined arrows in A–C). C, More anterior coronal section shows cortical atrophy with dilated sulci (black arrows). Note marked thinning of the corpus callosum on B and C.
CT was less sensitive than MR in all patients studied with both methods (Table 3). Five of our patients had solitary or multiple low-density foci on CT with variable enhancement of at least one lesion. In the one patient who had a double-dose scan, enhancing foci were identified on the delayed scan that were not apparent on the immediate postcontrast study. Other authors have also noted that acute lesions display enhancement on delayed scans after administration of high-dose contrast material while chronic lesions remain unchanged. In all cases more lesions were seen on MR than on CT.

Of interest are the two other adolescents whom we scanned for clinically suspected MS. One, an 18-year-old woman with a 3-year history of intermittent upper extremity weakness and paresthesia, had a normal MR brain scan. The sagittal T1-weighted series disclosed C3–C4 disk herniation with cord compression. Diskectomy resulted in complete relief of symptoms. A 13-year-old girl with left upper extremity weakness had numerous dilated vascular channels seen on MR. Idiopathic progressive arteriopathy of childhood (Moya-Moya–type pattern) was confirmed at angiography. MR is extremely helpful in distinguishing these and other clinical MS mimics from true demyelinating disease.

Conclusions
On the basis of this small series of 12 adolescents, we conclude that female predominance as well as the frequency of MR-demonstrated lesions in the cerebellum and brainstem appear to be higher in adolescents than that usually encountered in adults. Atrophy and abnormal iron deposition were uncommon in the adolescents.

REFERENCES