Age-Related Changes in the Pediatric Brain: Quantitative MR Evidence of Maturational Changes during Adolescence

R. Grant Steen, Robert J. Ogg, Wilburn E. Reddick, and Peter B. Kingsley

PURPOSE: To determine whether a quantitative MR imaging method to map spin-lattice relaxation time (T1) can be used to characterize maturational changes in the normal human brain. METHODS: An inversion-recovery technique was used to map T1 transversely at the level of the basal ganglia in a study population of 19 healthy children (4 to 10 years old) and 31 healthy adolescents (10 to 20 years old), and in a normative population of 20 healthy adults (20 to 30 years old). RESULTS: Nonparametric analysis of variance showed that T1 decreases with age in the genu, frontal white matter, caudate, putamen, anterior thalamus, pulvinar nucleus, optic radiation, cortical gray matter (all \(P < .0001\)), and occipital white matter. There was a significant reduction in T1 between childhood (mean age, 7.1 ± 1.4) and adolescence (mean age, 13.5 ± 2.6) in all brain structures, but there was also a significant reduction in T1 between adolescence (mean age, 13.5 ± 2.6) and adulthood (mean age, 26.5 ± 3.4) in all brain structures except occipital white matter. Regression shows that T1 declines to within the range (mean ± 2 SD) of young adult T1 values by about 2 years in the occipital white matter, by about 4 years in the genu, by 11 years in the cortical gray matter, by 11 years in the frontal white matter, and by 13 years in the thalamus. CONCLUSION: Brain structures mature at strikingly different rates, yet the ratio of gray matter T1 to white matter T1 does not change significantly with age. Thus, conventional MR imaging methods based on inherent contrast are insensitive to these changes. Age-related changes tend to reach completion sooner in white matter than in gray matter tracts. Such normative data are essential for studies of specific pediatric disorders and may be useful for assessing brain maturation in cases of developmental delay.

Index terms: Brain, growth and development; Brain, magnetic resonance


Maturation of the human central nervous system begins in utero and continues through adolescence, with dramatic changes apparent in the first few years of life (1–5). Markers of maturation are available in the physical examination of the child, including head diameter, head circumference, general alertness, motor reflexes, muscle strength and tone, motor milestones, and structure and function of the sensory apparatus (3). However, it has been difficult to characterize maturational changes in the normal human brain or to assess objectively the degree of maturation in an individual child. Magnetic resonance (MR) imaging has provided a window into the intact brain for more than a decade. There has been great interest in characterizing patterns of change seen in pediatric patients by means of various T1- and T2-weighted imaging sequences, which form the repertoire of conventional MR imaging (6–12). Yet the complex changes in the pediatric brain have not been fully characterized because of the relative subjectivity of evaluation possible with conventional MR imaging. MR spectroscopy has also been used to characterize age-related changes in the pediatric brain (13, 14), but this
technique has less spatial resolution than conventional MR imaging, is more difficult to implement, and is available at fewer medical centers.

Newer techniques of quantitative MR imaging are more objective and potentially more sensitive to subtle brain changes than are conventional MR imaging techniques (15). Recently, quantitative MR imaging techniques have led to striking success in identifying brain abnormalities associated with fragile-X syndrome (16), schizophrenia (17), multiple sclerosis (18), phenylketonuria (19), epilepsy (20), and human immunodeficiency virus (HIV)-associated dementia (21, 22). These findings suggest that quantitative MR imaging may offer the clinician sufficient sensitivity and accuracy to characterize the schedule of brain maturation in healthy children more objectively, or perhaps to identify with greater surety those children who are developmentally delayed.

Subjects and Methods

We used T1 mapping to assess the changes occurring in a healthy population of children and adolescents, and we compared our findings with a normative population of healthy young adults 20 to 30 years old.

Subjects were either hospital personnel or the children of hospital personnel, all of whom reported good health. The protocol to image subjects was reviewed and approved by our Institutional Review Board. Subjects included 19 children 4 to 10 years old (mean, 7.1 years ± 1.4), 31 adolescents 10 to 20 years old (mean, 13.5 years ± 2.6), and 20 adults 20 to 30 years old (mean, 26.5 years ± 3.4). Parents or guardians of subjects signed an informed consent form after a brief description of the MR protocol or, in the case of volunteers older than 18, the informed consent was signed by the subject. None of the volunteers was sedated. Of the 50 subjects less than age 20, about 54% were female, 46% male, and 56% were white, 44% black; the distribution of adult subjects was roughly comparable.

Quantitative MR Imaging

All MR imaging was performed on a 1.5-T Magnetom SP63 (Siemens Medical Systems, Iselin, NJ) using a standard Siemens head coil. A conventional T1- or T2-weighted image set of the brain was acquired in the transverse plane to enable selection of a section through the brain most comparable to the section level selected in the other volunteers. Conventional T1-weighted images were acquired with parameters as follows: 266/6/1 (repetition time/echo time/excitations), 23-cm field of view, 90° flip angle, 192 × 256 matrix, 19 sections with a thickness of 5 mm, and a total imaging time of 54 seconds. In some cases, conventional T2-weighted images were acquired with parameters as follows: 3500/19,93/1 (three echoes per repetition time for each effective echo time), 23-cm field of view, 192 × 256 matrix, 19 sections with a thickness of 5 mm, and a total imaging time of 3 minutes 57 seconds. A representative T2-weighted image is shown (Fig 1A) at the section level selected for the parametric T1-weighted image.

Quantitative imaging of T1 was done with a precise and accurate inversion-recovery (PAIR) sequence optimized and validated previously (15, 23–25). A 5-mm tissue section was selected and imaged with a phase-sensitive inversion recovery sequence of 2500/20 with four different inversion times: 100, 500, 900, and 2460. To minimize imaging time, a 20-cm field of view was used, with a 72 × 256 matrix, for an effective in-plane resolution of 2.1 × 0.8 mm; total imaging time was 13 minutes 40 seconds. An inversion recovery image is shown (Fig 1B) to indicate the quality of the raw data used to calculate the parametric image.

Measurement of T1

After acquisition, PAIR images were transferred to an off-line Silicon Graphics workstation for calculation of a T1 map. An empirically determined signal intensity (SI) difference threshold of 100 was applied to the difference between the 100 and 2460 T1 images in order to identify pixels within the phase-encoded region that were either noise or tissue with signal indistinguishable from noise (25). Each pixel identified as noise was excluded and the remaining pixels were submitted to a curve-fitting procedure as an array containing SI values derived from each of the four inversion recovery images. Calculations were performed using a Levenberg-Marquardt algorithm to fit SI values to the following equation, modified from the original (26) to include a parameter for effective flip angle of the inversion pulse:

\[
SI = a \left[ 1 - 2ke^{-\frac{TR-TE}{2}/T1} \right] + ke^{-\frac{TR}{T1}} - (1-k)e^{-\frac{TE}{T1}} \]

where \(a\) is the spin-density factor corrected for T2 losses, \(k\) is the cosine of the effective flip angle of the inversion pulse, TR is the repetition time between successive excitations, TE is the echo time, T1 is the inversion time, and T1 is the spin-lattice relaxation time. The fitting procedure returned values for \(a\), \(k\), and T1 in each pixel, and the T1 value was used to produce a parametric T1 map in which each pixel gray-scale value was equivalent to the T1 relaxation time in milliseconds. A parametric T1 map is shown (Fig 1C), with several regions of interest (ROIs) indicated on the image. The T1 of various structures was measured by identifying ROIs on the 500-T1 image, using criteria described in detail previously (15), then applying those ROIs to the parametric T1 map. Error of the T1 fit was determined on a pixel-by-pixel basis (25), and further analysis excluded pixels identified as having an error above a statistical threshold. Output from the computer
included the number of pixels counted within each ROI, mean T1, and the standard deviation (SD) of T1.

Regression of T1 Against Age

Mean T1 was measured in each of nine brain regions in all subjects (Table 1). Mean T1 was regressed against the subject’s age for each brain region separately, using a standard least-squares algorithm in a statistical software package (SPSS 6.1, SPSS Inc, Chicago, Ill) running on a Macintosh Quadra 650. Young adult subjects were also analyzed separately to determine the range of T1 values expected in fully mature subjects (23). The mean T1 (± 2 SD) of the 20 young adults was accepted as the normal range of adult T1 for each brain structure.

Proportion of Pixels with T1 Values Characteristic of Gray and White Matter

Parametric T1 maps were imported to a Macintosh Ilvx with a Raster Ops display, and images were analyzed using Adobe Photoshop 2.5 (Adobe Systems Inc, Mountain View, Calif). Pixels from bone and tissue external to the brain were erased using standard Photoshop tools, and a histogram analysis was performed on the remaining pixels. Pixels were classified as pure white matter, pure gray matter, or neither, according to criteria established previously, which were derived from T1 data averaged from a group of 24 healthy children (24). The range of pixel values defined as pure white matter was 568 milliseconds (mean genu, T1 – 2 SD) to 712 milliseconds (mean occipital white matter, T1 + 2 SD). The range of pixel values accepted as pure gray matter was 1160 milliseconds (mean cortical gray matter, T1 – 1 SD) to 1440 milliseconds (mean cortical gray matter, T1 + about 5 SD). Pixels falling outside these ranges of T1 were assumed to represent volumes containing both white and gray matter, or gray matter contaminated with cerebrospinal fluid partial volume, and were not counted.

Statistical Tests

Nonparametric tests were used exclusively for data analysis because of potential abnormal distribution of age-related T1 values. The Kruskal-Wallis analysis of variance (ANOVA) tested for trends across all subject groups (group 1, healthy children 4 to 10 years old; group 2, healthy adolescents 10 to 20 years old; group 3, healthy adults 20 to 30 years old). The Mann-Whitney U test was used for pairwise comparisons between these groups.

Results

Quantitative MR Imaging

A parametric T1 map is shown (Fig 1C) for the youngest volunteer, with ROIs indicated. Measurements of T1 made in nine regions of the brain are summarized for all subjects by age.
group (Table 1). Nonparametric analysis of variance (Kruskal-Wallis ANOVA) showed a significant trend of decreasing T1 with increasing age in every brain structure examined. The least significant T1 finding reported was found in occipital white matter (P < .004); the significance of all other age regressions was at least P = .0001. However, the ratio of gray matter T1 to white matter T1 (gray/white ratio) did not change significantly with age. Pairwise comparison between different age groups showed a significant difference in T1, even between adolescents and adults, in every brain structure examined (Table 1).

Simple factorial ANOVA, with age as a factor and with race and sex as covariates, showed that age was a significant source of T1 variation in every brain structure except occipital white matter. Neither sex nor race was a significant covariate in any brain structure except frontal white matter. In frontal white matter, the covariates of sex and race were slightly correlated with T1 (P = .019 and P = .022, respectively), although our sample size is too small for this result to be convincing.

Regression of T1 Against Age

Regressions of T1 with age are shown for frontal white matter (Fig 2A), caudate (Fig 2B), anterior thalamus (Fig 2C), and cortical gray matter (Fig 2D). A fitted regression line is shown for all data points (n = 70), together with an indication of the range of T1 values (mean T1 ± 2 SD) found in adults less than 30 years old. Goodness-of-fit data for each regression of T1 with age are summarized (Table 2). Mean T1 of every brain structure decreases significantly with increasing age, and the R² (multiple correlation coefficient) for this correlation averages .64 (± .14). The highest correlation coefficients were found in the pulvinar nucleus (r = .81), the anterior thalamus (r = .80), and the cortical gray matter (r = .76). Correlation coefficients averaged .73 (± .11) for all gray matter structures, but only .55 (± .13) for all white matter structures. The approximate age at which the fitted regression line crosses the upper bound of the normal T1 range for adults is expressed as the age at which the regression “normalizes” (Table 2).

**Pixels with T1 Values Characteristic of Gray and White Matter**

There were significant changes with age in the raw count of gray matter and white matter pixels. Regression analysis shows that the raw number of gray matter pixels declines with age (r = .79), while the raw number of white matter pixels increases with age (r = .84) (Fig 3A). Both these relationships are significant (P < .0001). Furthermore, there is a significant inverse relationship between raw pixel counts of gray matter and white matter in subjects (n = 64; R² = .62; F test = 41.31; P < .0001) (Fig 3B).

Pixel count data are summarized in Table 3. Kruskal-Wallis ANOVA shows a significant trend toward an increase in the proportion of
pixels with T1 characteristic of white matter ($P < .0001$) and a decrease in the proportion of pixels with T1 characteristic of gray matter ($P < .0001$).

**Discussion**

Data presented here show unequivocally that T1 decreases with age throughout the brain and that these changes can continue through adolescence. For example, T1 of cortical gray matter continues to change through adolescence, and the mean T1 of cortical gray matter in adolescents is significantly greater than in young adults (Table 1). Regression analysis shows that cortical gray matter T1 does not normalize to within the normal adult range until about 11 years old, and the regression curve does not approximate the mean adult value until about 20 years of age. Conversely, white matter T1
tends to normalize before 8 years of age, except in the frontal region (Table 2). Therefore, our data suggest that age-related changes tend to reach completion sooner in white matter than in gray matter. Previously, it has been difficult to determine the age at which maturation of cortical gray matter occurs, either by histology or conventional MR imaging, because gray matter does not show as obvious a sign of maturation as does white matter. Our data also show clearly that the ratio of gray matter T1 to white matter T1 does not change significantly with age. Therefore, conventional MR imaging methods based on the inherent gray/white matter contrast would be insensitive to the age-related changes we describe.

**Table 2: Age regression data from a quadratic fit of quantitative MR data from healthy control subjects**

<table>
<thead>
<tr>
<th>Brain Structure</th>
<th>$R^2$</th>
<th>ANOVA F Test</th>
<th>P</th>
<th>Y-Intercept</th>
<th>Age, y, at Which Regression Normalizes*</th>
<th>Age, y, at Which Structure Matures†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genu</td>
<td>.56</td>
<td>11.6</td>
<td>.0001</td>
<td>645.6</td>
<td>~4</td>
<td>2</td>
</tr>
<tr>
<td>Splenium</td>
<td>.41</td>
<td>5.2</td>
<td>.009</td>
<td>697.1</td>
<td>6.4</td>
<td>2</td>
</tr>
<tr>
<td>Frontal white</td>
<td>.73</td>
<td>29.5</td>
<td>&lt;.0001</td>
<td>695.9</td>
<td>11.2</td>
<td>5</td>
</tr>
<tr>
<td>Caudate</td>
<td>.73</td>
<td>29.1</td>
<td>&lt;.0001</td>
<td>1175.4</td>
<td>9.5</td>
<td>3</td>
</tr>
<tr>
<td>Putamen</td>
<td>.55</td>
<td>11.1</td>
<td>.0001</td>
<td>1017.7</td>
<td>5.7</td>
<td>...</td>
</tr>
<tr>
<td>Thalamus</td>
<td>.80</td>
<td>44.9</td>
<td>&lt;.0001</td>
<td>942.0</td>
<td>12.6</td>
<td>...</td>
</tr>
<tr>
<td>Pulvinar nucleus</td>
<td>.81</td>
<td>47.0</td>
<td>&lt;.0001</td>
<td>1083.5</td>
<td>11.4</td>
<td>...</td>
</tr>
<tr>
<td>Optic radiation</td>
<td>.62</td>
<td>16.3</td>
<td>&lt;.0001</td>
<td>716.0</td>
<td>7.9</td>
<td>...</td>
</tr>
<tr>
<td>Occipital white matter</td>
<td>.45</td>
<td>6.5</td>
<td>.004</td>
<td>670.0</td>
<td>~2</td>
<td>3</td>
</tr>
<tr>
<td>Cortical gray matter</td>
<td>.76</td>
<td>34.4</td>
<td>&lt;.0001</td>
<td>1333.5</td>
<td>11.0</td>
<td>1.5</td>
</tr>
<tr>
<td>Gray/white matter ratio</td>
<td>.30</td>
<td>2.5</td>
<td>NS</td>
<td>1.93</td>
<td>Before birth</td>
<td>...</td>
</tr>
</tbody>
</table>

* The age at which regression normalizes is the approximate age at which the fitted regression line (for all 70 subjects) crosses the upper boundary of the normal range of adult T1 (defined as mean T1 for a particular structure + 2 SDs).

† The age at which structures were thought to mature on the basis of conventional MR appearance is abstracted from Holland et al (8) and shown for comparison.

**Fig 3.** Parametric image pixel counts as a function of age in 50 children and 20 adults; all pixels counted were exclusively in brain (excluding bone, scalp, and subcutaneous fat).

A, Raw count of pixels defined as gray matter (solid diamonds) and white matter (open diamonds), uncorrected for any differences in head size. Nonlinear regressions are shown for gray matter ($r = .84; R^2 = 12.60; P < .0001$) and for white matter ($r = .79; R^2 = 10.51; P < .0001$).

B, The relationship between gray matter pixels and white matter pixels in 70 subjects shows an inverse relationship between the two pixel categories ($r = .64; R^2 = 6.86; P < .0001$).
We also report that there is a reduction of mean white matter T1 with increasing age (Table 1), which may correlate with the process of myelination in white matter tracts. Frontal white matter T1 does not normalize to within the normal adult range until about 11 years old, and the regression curve does not approximate the mean adult value until about 25 years of age. Evidence of continued myelination of white matter tracts is shown by the increase in the raw number of white matter pixels with age and the decrease in the raw number of gray matter pixels with age (Fig 3A). Furthermore, there is a significant inverse relationship between the number of gray matter pixels and the number of white matter pixels (Fig 3B). Our findings thus suggest that some brain areas, initially categorized as gray matter in children, can undergo maturational changes that eventually result in pixel categorization as white matter.

We believe these results are important because they give a more detailed picture of maturational changes than was possible in the past. Our results could potentially be extended to young children in order to characterize the full sequence of normal brain development. Current methods of characterizing brain maturation depend on clinical history, physical examination, and individual developmental observations (3). These methods are usually sufficient to identify severe problems in development, but mild developmental abnormalities often cannot be reliably identified on the basis of clinical findings alone. Developmental delay is diagnosed when a child fails to reach developmental milestones at the expected age, with an adequate leeway given to account for the broad range of normal variation. Because developmental delay has a population prevalence estimated to be as high as 10% (3), an objective method for characterizing brain maturation would be very useful. Conventional MR imaging of the brain has therefore been recommended for children with asymmetric findings on the neurologic examination, abnormal head growth, extremity weakness, changes in motor status, blindness, deafness, seizures, or other substantial impairments (3).

Yet current conventional MR imaging methods of assessing structural maturity of the brain during childhood or adolescence have limited accuracy and sensitivity. Conventional T1- and T2-weighted sequences show that brain maturation proceeds sequentially, from central to peripheral, from inferior to superior, and from posterior to anterior areas (10). However, the relative gray/white matter contrast normalizes to an adult appearance by only 3 years of age, and the brain of a young child is thus largely indistinguishable from that of an adult on conventional MR images (10). Part of the problem is that automated algorithms for adjusting image contrast (window and center), which are routinely available at the imaging console, make image contrast somewhat arbitrary. However, it is also possible that conventional MR imaging is simply too insensitive to detect subtle age-related changes in the brain. It would therefore be advantageous to have another method available to the clinician to identify developmental delay, particularly if such a method could be combined with a standard conventional MR imaging examination for suspected developmental delay.

Previous quantitative MR imaging techniques to study brain maturation have been of limited utility in that they appear to be completely insensitive to changes in the adolescent brain.

**TABLE 3:** Proportion (mean percentage ± SD) of white and gray pixel types, calculated by dividing the number of pixels within a range of T1s by the total number of pixels in the image plane within the brain (excluding bone, scalp, and subcutaneous fat)

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Age Group</th>
<th>Mann-Whitney U Test, P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Children (10 Years Old and Younger), n = 19</td>
<td>Kruskal-Wallis 1 x 2</td>
</tr>
<tr>
<td>White matter pixels, % (SD)</td>
<td>Adolescents (10 to 20 Years Old), n = 28</td>
<td>1 x 3</td>
</tr>
<tr>
<td>(T1, 568–712)</td>
<td>Adults (20 to 30 Years Old), n = 19</td>
<td>2 x 3</td>
</tr>
<tr>
<td>Gray pixels, % (SD)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(T1, 1160–1440)</td>
<td>12.1 (2.7)</td>
<td>.0001</td>
</tr>
<tr>
<td></td>
<td>16.4 (3.6)</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td></td>
<td>22.6 (3.4)</td>
<td>.0007</td>
</tr>
</tbody>
</table>

Note.—Nonparametric statistical tests were used for all comparisons because proportional data do not have a normal distribution. The Kruskal-Wallis one-way ANOVA tested for trends across all groups (70 subjects in total). The Mann-Whitney U test was used for pairwise comparisons between groups.
Prior efforts to measure brain T1 suggested that the mean T1 of white matter of 2 year olds was within 1 SD of the mean adolescent or adult white matter T1 (data summarized in Table 2) (8). Similarly, the mean T1 of gray matter approximated the adult value by only 1 year of age, and the T1 of the caudate was indistinguishable from the adult value by 3 years of age. However, prior techniques for measuring T1 were not rigorously quantitative, were generally not well validated, and were not used to examine an adequate normative sample of subjects. A well-validated, precise, and accurate method will be required if quantitative MR imaging is to prove useful in characterizing brain maturation, and this method must be applied to a large sample of healthy subjects.

Careful measurement of T1 by a PAIR technique has already demonstrated significant age-related changes in a healthy population of 55 adults between 18 and 72 years old (23). This PAIR method has now been improved, using a three-parameter fit to correct for a 2% to 3% underestimation of T1 due to flip angle imperfection in the inverting pulse (P. Kingsley, unpublished data). Application of the new technique of T1 mapping to a population of 50 healthy children demonstrates that T1 can be measured with sufficient precision to show age-related changes in T1 through adolescence (Tables 1 and 2) and that various brain tissues mature at a strikingly different rate (Table 2). However, caution must be exercised because there are not yet enough data to determine whether T1 mapping values can be measured with sufficient accuracy to assess maturation of the individual child.

The age range from birth to about 3 years is of greatest interest to neurologists, because this is the time during which developmental delay is usually first noted (3). A quantitative MR imaging method that could corroborate neurologic findings suggestive of developmental delay within this age range would be extremely useful. As yet, we have little data from children under 4 years of age, because our current protocol precludes sedation of healthy children. However, we recently performed a T1 mapping examination on a 10-month-old boy with neuroblastoma, who was imaged to evaluate cord compression. This child’s conventional MR imaging examination of the brain was interpreted as normal, without evidence of metastatic disease. In this case, we found that frontal white matter T1 was 1047 milliseconds, occipital white matter T1 was 989 milliseconds, and cortical gray matter T1 was 1331 milliseconds. While these findings are anecdotal, they suggest that T1 is very high at birth and further suggest that the T1 decline we describe during childhood and adolescence is perhaps even more striking during infancy. We will continue to accrue control data from infants who are imaged for possible disease but whose findings are normal.

Our data showing age-related changes in the pediatric brain are largely consistent with data obtained by histologic analysis of brain tissue samples. Brain myelination is a dynamic process that proceeds at different rates in different neuronal systems; it can be asynchronous within a given neuronal system (1, 2) and has been reported to be nearly complete in infants as young as 10 months old (27). It is possible that the reduction in T1 we report in myelinating nerve tracts is related to an increase in the number of protons associated with myelin and to a decrease in the number of free-water protons (28), so that our T1 measurement is actually an approximation to a biexponential decay. However, we have no evidence of biexponentiality of the T1 decay curve in white matter, even in young children. To determine objectively whether T1 reduction is caused by an increase in the number of protons associated with myelin, it might be possible to image myelin selectively, using a technique that discriminates between the very short T1 of bound water and the longer T1 of free water (29).

It is also possible that the reduction in brain T1 with increasing age we report is simply related to a reduction in brain tissue water content. Brain water content decreases from 88% at birth to 82% at 6 months of age, as a result of a nearly 50% increase in dry weight of the brain (30). Brain water content continues to decrease, albeit at a slower rate, for several more years. Myelin, which is about 30% protein and 40% water, eventually constitutes one half the dry weight of white matter. The degree of myelination is closely correlated with the cholesterol content of brain, which doubles within the first 6 months of life and continues to increase through at least 4 years of age (30). Our data do not yet permit us to discern among water content, degree of myelination, or some other factor as dominant in affecting brain tissue T1.

The greatest limitation of the PAIR method described here is that it is a single-section tech-
nique, making it less useful for examining patients who may have a diffuse pathologic process. However, we believe it is important to maintain a high degree of precision and accuracy when measuring T1; the mean T1 difference between children and adults averaged only 6.9% in white matter and 12.1% in gray matter. Any loss in precision or accuracy may make it impossible to detect such a small difference. Therefore, we are reluctant to use a multisection technique until it has been well validated; none of the techniques that could potentially speed up T1 measurement has been validated as extensively as the PAIR technique, and it is not clear that any would be as accurate and precise. It may soon be possible to obtain four-section coverage of the brain by implementing an interleaved PAIR technique, but this technique has not yet been validated.

The T1 mapping method presented here provides essential normative data for studies of patients with specific brain abnormalities, but it may also be useful for assessing maturation in suspected cases of developmental delay. An advantage of using this method to characterize brain maturation is that it might help to focus clinical attention if a particular neuronal system is delayed. This could be important because careful neurologic assessment of developmentally delayed children was unable to provide an etiologic diagnosis in 37% (22 of 60) of patients with severe delay (31).

Acknowledgments

We gratefully acknowledge the help of Carol Greenwald, who helped to recruit subjects, and of the families of St Jude Children’s Research Hospital who so generously contributed their time and effort.

References


