Magnetization Transfer Imaging and Proton MR Spectroscopy in the Evaluation of Axonal Injury: Correlation with Clinical Outcome after Traumatic Brain Injury

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BACKGROUND AND PURPOSE: Current imaging does not permit quantification of neural injury after traumatic brain injury (TBI) and therefore limits both the development of new treatments and the appropriate counseling of patients concerning prognosis. We evaluated the utility of magnetization transfer ratio (MTR) and proton MR spectroscopy in identifying patients with neuronal injury after TBI.

METHODS: Thirty patients with TBI (21–77 years old; mean age, 42 years; admission Glasgow Coma Scale (GOS) scores 3–15; mean score, 11) were studied on a 1.5-T system with magnetization transfer imaging and MR spectroscopy of the splenium. Magnetization transfer imaging was also performed in the brain stem in all patients, and other areas of the brain were sampled in one patient. The splenium of the corpus callosum and brain stem were studied because these are often affected by diffuse axonal injury. Scans were obtained 2 to 1129 days after injury (median, 41 days). MTR was considered abnormal if it was more than 2 SD below normal. Proton MR spectroscopy was used to calculate the N-acetylaspartate (NAA)/creatine (Cr) ratio. GOS was determined at least 3 months after injury.

RESULTS: In 10 patients with a GOS of 1 to 4, the mean NAA/Cr was 1.24 ± 0.28; two of these patients had abnormal MTR in normal-appearing white matter (NAWM). In 20 patients with a GOS of 5, the mean NAA/Cr was 1.53 ± 0.37 (P < .05); four of these patients had abnormal MTR in NAWM. MTR abnormalities in NAWM were identified in six patients, but these changes did not correlate with GOS or MR spectroscopy changes.

CONCLUSION: MTR and MR spectroscopy can quantify damage after TBI, and NAA levels may be a sensitive indicator of the neuronal damage that results in a worse clinical outcome.
nonanatomic studies as outcome predictors. Single-photon emission CT depicts abnormalities in regional blood flow and lesions not visible on CT or MR studies, and these global blood flow measurements correlate with outcome (19). Holshouser et al (20) used proton MR spectroscopy to examine neonates, infants, and children with brain injury caused by a variety of mechanisms. In this patient population, changes in MR spectroscopy measurements in occipital gray matter correlated with outcome. Choe et al (21) found similar MR spectroscopy changes in adults after TBI when the frontoparietal white matter was studied. In patients with multiple sclerosis (MS), a positive correlation has been identified between the magnetization transfer ratio (MTR) and N-acetylaspartate (NAA) measured by MR spectroscopy in white matter plaques (22, 23). In the present study, we sought to evaluate whether a similar association exists between MTR and proton MR spectroscopy in patients with TBI. We hypothesized that MR spectroscopy and MTR provide complementary functional information that correlates with neurologic outcome in adults with TBI.

**Methods**

**Patients**

After obtaining signed consent, as approved by our institutional review board, 30 adult patients (15 men and 15 women; 21–77 years old; mean age, 42 years) with closed TBI underwent MR imaging and MR spectroscopy. Admission GCS scores (24) and mechanisms of injury were obtained from initial hospitalization records. Admission GCS scores ranged from 3 to 15 (mean score, 11). Injury mechanisms included motor vehicle accidents in 16 patients (53%), falls in seven (23%), assaults in four (13%), and bicycle accidents in three (10%). MR imaging was performed 2 to 1129 days after the injury (median, 41 days). Patients’ outcomes were assessed at least 3 months after the injury by using the Glasgow Outcome Scale (GOS): 1 = death, 2 = persistent vegetative state, 3 = severe disability (conscious but disabled), 4 = moderate disability (disabled but independent), 5 = good recovery (25). GOS scores ranged from 1 to 5 (median, 5).

**MR Studies**

All MR imaging and MR spectroscopy studies were performed on a 1.5-T MR scanner equipped with a spectroscopy software package. Conventional MR imaging included sagittal short-TR/TE sequences (650/11), axial spin-echo (3000/30,90) or fast spin-echo (2700/18,85) sequences, and axial spin-echo (3000/30,90) or gradient-recalled echo (750/40; 10° flip angle) sequence. One 19-millisecond sinc-shaped off-resonance saturation pulse was used per TR: control images were obtained without the saturation pulse. Imaging parameters were chosen to minimize T1 and T2 weighting. The MTR is defined by the equation

\[
MTR = (M_0 - M_3)/M_0 \times 100\%
\]

where \(M_0\) equals the average signal intensity in the absence of the saturation pulse and \(M_3\) equals the average signal intensity in its presence. Images were processed on a Sun Sparc station 330 (Sun Microsystems, Palo Alto, CA) using Interactive Data Language software (Interactive Data Language Systems, Boulder, CO). MTRs were calculated in normal-appearing white matter (NAWM) in areas frequently subject to diffuse axonal injury (the pons within the brain stem and the splenium of the corpus callosum) (Fig 1). Owing to differences in MTRs and NAA levels in gray and white matter, evaluation of the splenium minimizes the effect of partial volume averaging that occurs if a subcortical voxel placed at the gray/white junction is used. MTRs were also obtained in the same areas in healthy control subjects (26), consisting of six men and nine women ranging in age from 26 to 61 years (mean age, 36 years). Regions with MTRs more than 2 SD below MTRs obtained in corresponding areas in the control group were classified as abnormal.

Spectroscopy was initially performed with axial T2-weighted sequences to prescribe rectangular voxels of the white matter of the splenium that minimized gray matter and CSF contamination. The voxel dimensions routinely used were 9 mm (anterior/posterior) \(\times\) 25 mm (right/left) \(\times\) 10 mm (superior/inferior). Voxel locations were chosen by a single investigator to ensure the same location in each subject (Fig 1). The splenium was studied with a stimulated-echo acquisition mode (STEAM) localization technique to minimize contamination. Careful shimmeding was necessary to minimize line widths of the spectra from this region. Water suppression was achieved by using three chemical shift-selective radio-frequency pulses followed by a dephasing gradient applied to each of the three axes. The sequence parameters were as follows: 2000/31 (TR/TE), 2500 Hz spectral bandwidth, 13.7 milliseconds mixing time, 2048 complex points, eight-step phase cycling, and 128 acquisitions. Gradient shimming on the voxel and optimization of the solvent suppression were performed before the start of the acquisition. The spectral processing was performed with ProNMR (Softpulse Software, Guelph, Ontario, Canada) by using zero-filling to 8K data points, 2 Hz line broadening applied in the time domain, one-dimensional Fourier transformation, and zero-order phase correction. Areas under the peaks were estimated using a Marquardt fitting routine to lorentzian lineshapes in the frequency domain. Peak ratios were then calculated for NAA/creatinine (Cr) (Fig 2). NAA/Cr was chosen as the end point because it has been shown to be a more reliable indicator of injury after TBI than NAA/choline (27).
Figure 2. Proton MR spectrum obtained from normal splenium. The NAA/Cr ratio is 1.75. All spectra were acquired using a STEAM sequence (2000/31).

**Statistical Analysis**

Abnormal MTRs and metabolic ratios in subjects with good outcomes (GOS = 5) were compared with those obtained in subjects with less than a good outcome (GOS = 1–4) using Student’s t-test. GOS and GCS were also correlated with the NAA/Cr ratio by using Spearman’s rank-order correlation. A P value of less than .05 was considered statistically significant. NAA/Cr ratios were also correlated with the time interval between injury and MR examination for both outcome groups.

**Results**

Abnormal MTRs in NAWM of the brain stem or splenium were detected in five patients (Fig 3). An additional patient had abnormally reduced MTR values in the NAWM of the temporal and occipital lobes and the internal capsule. Clinical data pertaining to these patients are summarized in the Table. There was no significant correlation between abnormal MTR and clinical outcomes in this group.

The NAA/Cr ratio in the splenium ranged from 0.80 to 2.33. In the 20 patients with a GOS of 5, the mean (± SD) NAA/Cr was 1.53 ± 0.37. For those with a GOS of 4, the mean NAA/Cr was 1.45 (three patients); for a GOS of 3, the mean NAA/Cr was 1.24 (four patients); and for a GOS of 2, the mean NAA/Cr was 0.98 (two patients). The one patient with a GOS of 1 had an NAA/Cr of 1.15 in the splenium. For the 10 patients with less than good outcomes (GOS = 1–4), the mean (± SD) NAA/Cr was 1.24 ± 0.28, which was significantly lower than that for patients with good outcomes (GOS = 5, P < .05) (Fig 4). Overall, we noted a significant correlation between GOS and NAA/Cr levels in the splenium (Y = 0.142X + 0.819, r_s = 0.4378, P < .01, Fig 5). No significant correlation was found between the presence of abnormal MTRs in NAWM and NAA/Cr values nor between GCS and NAA/Cr levels or GCS and MTR changes. In addition, no significant correlation was found when NAA/Cr ratios for both outcome groups were correlated with the time interval between injury and MR examination (Fig 6).

**Discussion**

Important treatment decisions for TBI patients are based on estimates of patient outcome. The Traumatic Coma Data Bank showed that 29% of patients with a postresuscitation GCS of 6 had a good outcome (based on the GOS), 21% died, and the remainder had some degree of disability (28). That a single category of the GCS has such a range of outcomes underscores the rather crude nature of this technique of categorizing TBI. The clinical course of a patient with multiple contusions and a GCS of 6 evolves by different pathophysiological mechanisms than that of a patient with a subdural hematoma, skull fracture, and GCS of 6. In most epidemiologic studies, these two types of injury are grouped together. Only a few studies have compiled data in such a way as to permit a more detailed characterization of this heterogeneous patient population (2, 29). Difficulties predicting outcome also occur in cases of less severe TBI. Rimel et al
Patients with abnormal MTR in NAWM

<table>
<thead>
<tr>
<th>Age</th>
<th>GCS Score</th>
<th>Mechanism of Injury</th>
<th>NAA/Cr in Splenium</th>
<th>GOS Score</th>
<th>Location of MTR Abnormality</th>
</tr>
</thead>
<tbody>
<tr>
<td>77</td>
<td>14</td>
<td>Fall</td>
<td>1.47</td>
<td>3</td>
<td>Temporal, occipital, internal capsule</td>
</tr>
<tr>
<td>51</td>
<td>15</td>
<td>Fall</td>
<td>1.60</td>
<td>5</td>
<td>Pons and splenium</td>
</tr>
<tr>
<td>39</td>
<td>15</td>
<td>MVA</td>
<td>1.12</td>
<td>5</td>
<td>Splenium</td>
</tr>
<tr>
<td>34</td>
<td>15</td>
<td>Fall</td>
<td>1.33</td>
<td>5</td>
<td>Pons</td>
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<tr>
<td>28</td>
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<td>5</td>
<td>Pons</td>
</tr>
<tr>
<td>71</td>
<td>9</td>
<td>Fall</td>
<td>1.21</td>
<td>4</td>
<td>Pons and splenium</td>
</tr>
</tbody>
</table>

Note.—MVA indicates motor vehicle accident.

(30) studied patients with “mild” TBI (GCS = 13–15) and reported that two thirds of these patients had persisting neurobehavioral deficits 3 months after injury and one third of the group was unemployed. Recent epidemiologic studies have also found an association between mild TBI and the development of Alzheimer disease (31, 32), suggesting that neural damage with functional consequences often occurs after mild TBI. Most victims of mild TBI are discharged home with minimal, if any, medical follow-up because their neurologic examinations are normal. Those patients with a persisting cognitive deficit frequently have problems returning to work and reentering into normal family life. Early identification of patients at risk for either cognitive/behavioral deficits after mild TBI or poor/vegetative outcome after more severe TBI will allow an appropriate modification of how these patients are managed.

Numerous pathophysiological mechanisms have been associated with secondary or delayed injury in patients with TBI. Raised intracranial pressure, arterial hypotension, hypoxemia, and pyrexia have all been shown to be independent variables that can cause additional neural injury in the hours and days after injury (33, 34). Current animal studies are being directed at preventing the secondary neuronal damage that occurs from such known mechanisms of injury as excitatory amino acids, apoptosis, ischemia, cytoskeleton destruction, calcium toxicity, free radicals, and mitochondrial dysfunction (35). Determining which patients may benefit from a specific treatment or combination of treatments also depends on early and accurate identification of neuronal damage.

**Magnetization Transfer Imaging**

Magnetization transfer imaging relies on the principle that protons bound in structures exhibit T1 relaxation coupling with protons in the aqueous phase. When an off-resonance saturation pulse is applied, it selectively saturates those protons that are bound in macromolecules. These protons sub-
changes and occurred before changes seen on T2-weighted images. In the rotational acceleration model of axonal injury in the minipig, Kimura et al. (39) and McGowan et al. (40) both showed magnetization transfer imaging to be more sensitive than T2-weighted imaging for detecting histologically identified axonal damage. In a study of human TBI, Bagley et al. (26) reported abnormal MTR in otherwise NAWM only in patients with persistent neurologic deficits. Although we did not identify a similar trend in the current study, the patient populations are somewhat different. Bagley et al. studied a lower percentage of patients with good outcome and did not find any abnormal MTRs in the NAWM of these patients. A GOS of 5 can incorporate a range of functional outcomes that may not be sensitive to lesser amounts of neuronal injury (41). Consequently, it is possible that the MTR abnormalities seen in patients with a GOS of 5 may be characteristic of a subset of patients who have sustained neuronal injury that they have been able to compensate for as they reintegrate into their preinjury activities (42). An additional example of this problem occurs in MS, where there is weak or no correlation between Kurtzke Expanded Disability Status scores and T2 lesion volume or T1-enhanced lesion volume or other such metrics (43, 44). To test functional status better, more detailed neuropsychological tests may be needed. Perhaps the MR measures are more sensitive and specific for TBI than are the current clinical assessments.

Proton MR Spectroscopy

Whereas magnetization transfer imaging provides structural information, MR spectroscopy permits the detection of in vivo neurochemical alterations, such as Cr, choline, myo-inositol, and NAA, in a selected tissue volume. Although glutamate is the only amino acid in the CNS that is more abundant than NAA, the actual role of NAA in the brain is not understood. One proposed function for NAA is that of a brain “osmolyte” (45). Osmolytes are small, nonmetabolizable molecules that would protect the brain-cell volume during recovery from metabolic disturbances of extracellular osmotic pressure (46). NAA can also act as an acetyl group donor to initiate neuronal protein synthesis (47) or lipid synthesis during myelination (48). Regardless of its function, NAA is made exclusively in the mitochondria of neurons and has been shown to be a neuronal marker that decreases in numerous disease processes that have little in common except neuronal loss (49). Examples include spinal cord injury (50), amyotrophic lateral sclerosis (51), Parkinson disease (52, 53), Huntington disease (54), ischemic stroke (55, 56), progressive multifocal leukoencephalopathy (57), epilepsy (58), and MS (23, 59–62). The widespread loss of NAA in patients with MS correlates with the recent documentation of axonal damage throughout the brains of MS patients (63). The dementia associated with Al-
zheimers disease (64–75), Pick disease (76), and AIDS-related dementia (77) has been correlated with decreases in brain NAA. Cheng et al (76) were able to directly correlate loss in NAA as measured by MR spectroscopy with histologic neuronal loss seen in patients with Pick disease. Direct histochemical measurements of NAA in patients with Alzheimer disease have confirmed the lower levels of NAA that have been observed with MR spectroscopy (78, 79). Despite the more widespread use of NAA as a marker of neuronal loss, there are some reports of patients who have shown reversible decreases in NAA after various brain insults (80, 81). DeStefano et al (80) used their control data from the same group of patients with the systemic diseases evaluated in the study; therefore, the NAA changes noted may be relative rather than absolute alterations.

In models of brain injury in the rat, Rubin et al (82) showed a loss of NAA in the cortex of animals 1 hour after injury, and Signoretti et al (83) showed a decrease within minutes of injury. In a swine model of rotational acceleration injury, Cecil et al (84) also found decreased NAA 1 hour after injury and this change remained stable 1 week after injury. Choe et al (21) used MR spectroscopy to examine 10 patients with head injury and found decreased NAA/Cr. The lower NAA levels were statistically significant relative to those in healthy control subjects, and there was a trend for lower NAA/Cr ratios to correlate with poor clinical outcome (21). Neonates, infants, and children with brain injuries from various mechanisms (infection, hypoxia, trauma, and metabolic disorders) were examined with proton MR spectroscopy by Holshouser et al (20), who found that NAA/Cr ratios in occipital gray matter were significantly lower in infants and children with poor outcomes but not in neonates with poor outcomes. Unfortunately, direct comparison of the data from Choe et al (21) and Holshouser et al (20) with our findings cannot be performed, since those studies sampled NAA/Cr in the left frontoparietal white matter and occipital gray matter, respectively (21). Cecil et al (27) found that in uninjured control subjects, NAA/Cr levels in the splenium are higher than those in hemispheric white matter. In the present study, the splenium was chosen to measure NAA/Cr because of the high prevalence of splenium injuries in patients with TBI (85). Given these differences, the work of Choe et al (21) and Holshouser et al (20) still supports the conclusion that NAA/Cr may be a useful indicator of outcome after TBI. Finally, Cecil et al (27) also showed decreased NAA/Cr in the splenium of head-injured patients relative to that in control subjects. Their data support the ability of MR spectroscopy to identify neuronal injury after TBI; however, no correlation between GCS and NAA/Cr was noted (27).

**MR Studies after TBI**

A source of error in studying this patient population is the inevitably heterogeneous nature of patient characteristics and injury types. Age has not been controlled for in this study, yet some decrease in NAA has been shown in elderly patients with chronic ischemia (86) and even in normal aging (87, 88). The spectrum of injury mechanisms present in any TBI population dilutes the power of any observations. The pathobiology and outcome after diffuse axonal injuries and focal contusions may contribute to the variability in MTR and MR spectroscopy seen in this and previous studies (89). In addition, the imaging examinations in this study were performed at a wide range of time points after injury. While DeStefano et al (80) suggest that in some disease processes declines in NAA may be reversible, Cecil et al (27) identified no such change in head-injured patients. Although we did not restudy individual patients over time after their injury, evaluation of the distribution of NAA/Cr ratios relative to the timing of the study suggests that no large changes in NAA levels occur during the time intervals in which these patients were examined (Fig 6). Recent evaluation of whole-brain NAA levels in healthy patients has shown that NAA per cubic centimeter of brain is unchanged with aging (90). Thus, it does not seem reasonable that within the interval of this study (up to 3 years) the decrease in NAA can be attributed to anything other than the traumatic incident.

Despite these concerns, our data support a theory that decreases in NAA/Cr as detected by noninvasive MR spectroscopy may be an indicator of neuronal injury and prognosis after TBI. One explanation for this observation is that MR spectroscopy findings can indicate which patients with a poor initial GCS will make a significant recovery. Two patients in our study with an initial GCS of 7 and 10, respectively, both made a moderate recovery, to a GOS of 4. The NAA/Cr in these patients was above the mean of those with a GOS of 5 (ie, 1.57 and 1.58, respectively). Further studies with such patients are indicated to evaluate this phenomenon.

The strength of the correlation between GOS and NAA/Cr does not permit accurate prognostication using only the NAA/Cr ratio in the splenium. The findings of decreased NAA/Cr in the lobar white matter after TBI by Choe et al (21) and Holshouser et al (20), combined with the changes noted in the splenium in the current study, suggest that more detailed measurements of whole-brain NAA may increase the prognostic power of MR spectroscopy in determining outcome. Additional studies may permit the stratification of patients on the basis of age. Finally, the range of NAA/Cr levels noted in patients with good outcomes (GOS = 5) raises the possibility that MR spectroscopy may be an indicator of neuronal injury that is capable of identifying a subset of patients with “good” outcomes who have persistent cognitive deficits that negatively impact their ability to return to their preinjury level of activity. Conceptually, we view NAA as a specific marker of irreversible neuronal loss, whereas MTR may provide a measure of the entire
spectrum of histopathology, one end of which is neuronal loss. Evaluation of this hypothesis by further exploration of the ability of quantitative MR studies to obtain early prognostic information non-invasively in patients who have sustained a TBI is warranted.

Conclusion

This study shows that quantitative MR imaging and MR spectroscopy can be reliably obtained in patients with TBI. Proton MR spectroscopy measurement of the NAA/Cr ratio in the splenium may provide additional information that is useful in predicting clinical outcome in these patients. In the 10 patients with less than good outcomes, the NAA/Cr ratio was 1.24±0.28, compared with 1.53±0.37 in the 20 patients with a good neurologic outcome. An abnormal MTR in NAWM was identified in 20% of these patients and did not correlate with outcome or posttraumatic alterations in the NAA/Cr ratio.

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