MRI Evidence of Altered Callosal Sodium in Mild Traumatic Brain Injury

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ABSTRACT

BACKGROUND AND PURPOSE: Mild traumatic brain injury is a leading cause of death and disability worldwide with 42 million cases reported annually, increasing the need to understand the underlying pathophysiology because this could help guide the development of targeted therapy. White matter, particularly the corpus callosum, is susceptible to injury. Animal models suggest stretch-induced mechanical stretch of the axonal membrane resulting in ionic shifts and altered sodium ion distribution. The purpose of this study was to compare the distribution of total sodium concentration in the corpus callosum between patients with mild traumatic brain injury and controls using sodium ($^{23}$Na) MR imaging.

MATERIALS AND METHODS: Eleven patients with a history of mild traumatic brain injury and 10 age- and sex-matched controls underwent sodium ($^{23}$Na) MR imaging using a 3T scanner. Total sodium concentration was measured in the genu, body, and splenium of the corpus callosum with 5-mm ROIs; total sodium concentration of the genu-to-splenium ratio was calculated and compared between patients and controls.

RESULTS: Higher total sodium concentration in the genu (49.28 versus 43.29 mmol/L, $P = .01$) and lower total sodium concentration in the splenium (which was not statistically significant; 38.35 versus 44.06 mmol/L, $P = .08$) was seen in patients with mild traumatic brain injury compared with controls. The ratio of genu total sodium concentration to splenium total sodium concentration was also higher in patients with mild traumatic brain injury (1.3 versus 1.01, $P = .001$).

CONCLUSIONS: Complex differences are seen in callosal total sodium concentration in symptomatic patients with mild traumatic brain injury, supporting the notion of ionic dysfunction in the pathogenesis of mild traumatic brain injury. The total sodium concentration appears to be altered beyond the immediate postinjury phase, and further work is needed to understand the relationship to persistent symptoms and outcome.

ABBREVIATIONS: CC = corpus callosum; mTBI = mild traumatic brain injury; TSC = total sodium concentration

Mild traumatic brain injury (mTBI) is the leading cause of death and disability in the United States and worldwide, with approximately 42 million cases annually. Patients may have a complex array of symptoms, including cognitive disturbance, headache, and visual impairment, and there is a critical need to gain further insight into the pathophysiology underlying the injury. It is known that sodium is critical to cellular homeostasis, which maintains fluid volume in the intracellular and extracellular compartments, maintains resting potential across membranes, and triggers action potential. Mild TBI causes mechanical injury to axons, resulting in widespread membrane depolarizations and activation of cellular ionic cascades, thereby causing disruption of sodium homeostasis. There is a resultant increase in intracellular sodium, which lowers the threshold for membrane depolarization.

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rate, and von Mises stress are highest in and around the CC. Because the CC is a group of anatomic tracts that integrate information across cerebral hemispheres, patients with mTBI tend to experience deficits in integrative functions (cognitive slowing, confusion, difficulty with complex tasks) rather than focal neurologic deficits.

Noninvasive imaging of brain sodium on clinical MR imaging scanners is very challenging due to low signal-to-noise ratio. Recent advances in technologies achieved at our site, such as coil design, data acquisition, and sodium quantification, now allow us to study ionic changes noninvasively at clinical field strengths of 3T or higher. The purpose of this pilot study was to measure total sodium concentration (TSC) in the CC in patients with mTBI using sodium ($^{23}$Na) MRI and to compare TSC and its spatial distribution across the CC with that in healthy controls.

MATERIALS AND METHODS
The study was performed under approval by the institutional review board. Informed consent was obtained from each of the subjects studied.

Human Subjects and Clinical Assessments
Eleven patients (5 men and 6 women; age range, 19–70 years) with a history of mTBI (as defined by the American Congress of Rehabilitation Medicine) and 10 age- and sex-matched healthy controls were prospectively recruited for sodium ($^{23}$Na) MRI. Review of clinical charts was performed for pertinent clinical history and assessment, including postconcussive symptoms, neurologic examination, and scores on the Standardized Assessment of Concussion.

MR Imaging Acquisition
Sodium ($^{23}$Na) MRI scans were performed on a clinical 3T scanner (Magnetom Prisma; Siemens, Erlangen, Germany) with a custom-built 8-channel dual-tuned ($^1$H-$^{23}$Na) transmit/receive head array coil. The twisted projection imaging pulse sequence was applied to a 3D volume covering the whole head (FOV = 220 mm, matrix size = 64, 3D isotropic, nominal resolution = 3.44 mm, rectangular radiofrequency pulse duration = 0.5 ms, TE/TR = 0.3/100 ms, flip angle = 90°, rings = 28, P [key parameter] = 0.4, projections = 1595, averages = 4, TA [time of acquisition] = 10.6 minutes). This scheme of data acquisition produced a typically high SNR of 55 in gray matter, 35 in white matter, and 57 in CSF in the square-root of the sum-of-squares sodium image of healthy controls before the correction for coil sensitivities. A magnetization-prepared rapid acquisition of gradient echo ($^1$H-MR imaging pulse sequence was performed for structural imaging of the brain (FOV = 256 x 216 mm$^2$, matrix size = 384 x 324, slice thickness = 1 mm at 144 slices, TE/TR = 3.56/2.220 ms, acceleration factor = 3, TA = 4.6 minutes).

Image Preprocessing
Sodium images were corrected for intensity inhomogeneity relating to the array coil by dividing by a low-resolution version of the images reconstructed from the $k$-space center of an optimally selected diameter of 9.0/FOV. Normalization was then accomplished with conversion of image intensity into sodium concent-

RESULTS
Five men and six women with an age range 19–70 years and a history of mTBI were studied, with an average time since injury of 16 weeks. All patients were symptomatic at assessment. Ten of 11 subjects with mTBI had formal clinical assessments at our institution. One subject declined this assessment.

All of the 10 patients who underwent clinical assessment were symptomatic at time of imaging (detailed in the On-line Table).

FIG 1. The TSC of the genu was higher in patients with mTBI compared with controls ($P = .01$).

FIG 2. The TSC in the splenium was lower in patients with mTBI compared with controls ($P = .08$).
Of note, most patients reported headaches, sleep disturbances, dizziness, decreased concentration, and an average Standardized Assessment of Concussion score of 27.5 (highest Standardized Assessment of Concussion score = 30, normal score ≥ 25).21

The mean TSC in the genu, body, and splenium of the CC in patients with mTBI was 49.28 ± 0.51 mmol/L, 46.04 ± 0.43 mmol/L, and 38.35 ± 0.36 mmol/L, respectively, compared with 43.29 ± 0.46 mmol/L, 45.25 ± 0.38 mmol/L, and 44.06 ± 0.46 mmol/L in controls. There were statistically significant differences between patients with mTBI and controls with respect to the callosal TSC in the genu (49.28 versus 43.29 mmol/L, P < .01) (Fig 1). No significant differences in the average TSC were found between patients and controls in the body (P = .32) and splenium of the corpus callosum (P = .08) (Fig 2).

The average ratio (genu/splenium) was 1.3 in patients and 1.01 in controls (P = .001) (Fig 3).

DISCUSSION

The results of this investigation show differences in the callosal TSC between a small group of symptomatic patients with mTBI and age- and sex-matched healthy controls. Differences in the TSC between study groups varied depending on the location within the CC, with higher genu TSC and a trend that did not reach statistical significance of lower splenial TSC in patients with mTBI (Fig 4).

The findings corroborate a growing body of literature that underscores the importance of cytosolic sodium as a marker of tissue injury after trauma. In mTBI, twisting and stretching of axons results in mechanical disruption of membranes, altered function of voltage-gated sodium channels and sodium-potassium adenosine triphosphatase,4,5,22 and changed regulation of the expression of sodium channels,6,7 with potential persistent sodium abnormality.

It is not fully clear what caused the change of TSC in the CC. TSC has contributions from both intracellular and extracellular compartments, and derangements in either of these compart-

![FIG 3. The TSC of the genu/splenium ratio was higher in patients compared with controls (P = .001).](image)

![FIG 4. The TSC heat color map of the corpus callosum superimposed on a midline MPRAGE image in a control subject (top) and a patient with mTBI (bottom) shows that the TSC is higher in the genu and lower in the splenium in the patient with mTBI.](image)
ments could contribute to the altered TSC. Because the extracellular compartment rapidly equilibrates with a larger plasma sodium pool, changes in intracellular sodium concentration could certainly affect TSC measures. Alterations in the relative size of the compartments would also be expected to affect TSC measurements. Work is currently underway to attempt to estimate intracellular and extracellular contributions to the TSC signal.

Why injury may affect sodium differently across various parts of the CC is unclear. There is variable myelination across the CC, and expression of some specific voltage-gated sodium channels is known to track with myelination. In addition, von Reyn et al demonstrated anatomic redistribution of voltage-gated sodium channels in the CC in an animal model of mTBI. These factors may contribute to anatomic differences in TSC across the CC after injury.

Many medications may affect sodium homeostasis. The most relevant ones were specifically screened and included in the chart review. Patient 5 was on the antidepressant escitalopram (selective serotonin reuptake inhibitor). There are a few recent reports of escitalopram causing a syndrome of inappropriate antidiuretic hormone secretion and hyponatremia. Our patient exhibited no signs of the syndrome of inappropriate antidiuretic hormone secretion and had normal serum chemistry values. Patient 3 was treated with lamotrigine, a central nervous system voltage-gated sodium channel blocker, for posttraumatic dystonia. In this subject, the TSC genu/splenium ratio among the mTBI group was the closest to that of the control cohort (Fig 5).

Limitations of this study include the small sample size; however, here we show proof of concept that TSC can be measured in vivo, noninvasively, on a clinical 3T scanner in subjects with mTBI after injury. This represents the first report we are aware of suggesting sodium homeostasis abnormality in human subjects with mTBI using a noninvasive method. Already discussed is the need to estimate cellular compartmental contributions to the TSC to further elucidate ionic abnormalities in mTBI. This pilot work included a heterogeneous population of patients with respect to time since injury, history of prior mTBI, and medications that may affect the sodium balance. In this preliminary study, no T2WI/FLAIR was performed in these subjects.

CONCLUSIONS

This study shows complex differences in the TSC in the CC in symptomatic patients with mTBI compared with age- and sex-matched healthy controls. Specifically, the TSC in the genu of the CC was elevated. Further work is needed to understand the relationship between the TSC change and symptom resolution or outcome prediction.

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