Altered Relationship between Working Memory and Brain Microstructure after Mild Traumatic Brain Injury

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ABSTRACT

BACKGROUND AND PURPOSE: Working memory impairment is one of the most troubling and persistent symptoms after mild traumatic brain injury (MTBI). Here we investigate how working memory deficits relate to detectable WM microstructural injuries to discover robust biomarkers that allow early identification of patients with MTBI at the highest risk of working memory impairment.

MATERIALS AND METHODS: Multi-shell diffusion MR imaging was performed on a 3T scanner with 5 b-values. Diffusion metrics of fractional anisotropy, diffusivity and kurtosis (mean, radial, axial), and WM tract integrity were calculated. Auditory-verbal working memory was assessed using the Wechsler Adult Intelligence Scale, 4th ed, subtests: 1) Digit Span including Forward, Backward, and Sequencing; and 2) Letter-Number Sequencing. We studied 19 patients with MTBI within 4 weeks of injury and 20 healthy controls. Tract-Based Spatial Statistics and ROI analyses were performed to reveal possible correlations between diffusion metrics and working memory performance, with age and sex as covariates.

RESULTS: ROI analysis found a significant positive correlation between axial kurtosis and Digit Span Backward in MTBI (Pearson $r = 0.69$, corrected $P = .04$), mainly present in the right superior longitudinal fasciculus, which was not observed in healthy controls. Patients with MTBI also appeared to lose the normal associations typically seen in fractional anisotropy and axonal water fraction with Letter-Number Sequencing. Tract-Based Spatial Statistics results also support our findings.

CONCLUSIONS: Differences between patients with MTBI and healthy controls with regard to the relationship between microstructure measures and working memory performance may relate to known axonal perturbations occurring after injury.

ABBREVIATIONS: AK = axial kurtosis; AWF = axonal water fraction; BCC = body of the corpus callosum; DKI = diffusional kurtosis imaging; DSF = Digit Span Forward; DSB = Digit Span Backward; DSS = Digit Span Sequencing; FA = fractional anisotropy; gCC = genu of the corpus callosum; HC = healthy control; LNS = Letter-Number Sequencing; MTBI = mild traumatic brain injury; pCR = posterior corona radiata; sCC = splenium of the corpus callosum; sCR = superior corona radiata; SLF = superior longitudinal fasciculus; WMTI = white matter tract integrity; WAIS = Wechsler Adult Intelligence Scale; WRAT = Wide Range Achievement Test

Mild traumatic brain injury (MTBI) is an important public health problem with many serious consequences. While most patients with MTBI recover symptomatically within 2–3 weeks after injury, at least 15% of patients report persistent cognitive symptoms that are an important source of distress and disability after injury. Now an important body of work reveals MTBI-related WM injury using DTI and diffusional kurtosis imaging (DKI). More recently, WM tract integrity (WMTI) metrics derived from an advanced compartmental diffusion WM model have been proposed to describe microstructural characteristics in the intra- and extra-axonal environments of WM, including axonal water fraction (AWF), intra-axonal diffusivity, and extra-axonal axial and radial diffusivities.

One of major barriers to applying such findings to clinical cohorts is that the disorder is extremely heterogeneous and most current studies group clinically heterogeneous cohorts of patients with MTBI together, representing a broad spectrum of clinical symptoms. Thus, there is a specific need to understand domain-specific symptoms as they relate to detectable microstructural injuries, to better understand patient-specific injury and recovery.
One of the most common and clinically significant symptoms in patients with MTBI is deficits in working memory, which often negatively affect quality of life. This result comes as no surprise because working memory, which involves the capacity to temporarily store and manipulate information in pursuit of a goal, is at the core of critical cognitive functions such as comprehension, learning, reasoning, and decision-making. Working memory is conceptualized as comprising 3 main components: the central executive, responsible for manipulation of information and allocation of attention and processing resources, and 2 maintenance systems, the phonologic loop (verbal and auditory information) and the visuospatial sketchpad (visual and spatial information). There have been a few studies showing associations of working memory performance with measures of WM microstructure such as fractional anisotropy (FA) and AWF in healthy individuals, believed to relate to differences in axon volume and myelination. However, such associations have not yet been investigated in patients with MTBI.

Here, we hypothesize that WM injury in patients with MTBI can affect the relationship between microstructural changes to the WM and working memory performance. To test this hypothesis, we investigated the relationship between WM microstructural changes assessed using diffusion MR imaging (DTI, DKI, WMTI) and a set of Wechsler Adult Intelligence Scale, 4th ed (WAIS-IV) subtests tapping auditory-verbal working memory functions in patients with MTBI, comparing them against healthy controls (HCs). We also performed subgroup analyses based on working memory performance and time since injury.

**Materials and Methods**

**Study Population**

This study was approved by our institutional review board at New York University Langone Health. All experiments were performed in accordance with relevant guidelines and regulations, and written informed consent was provided by all subjects before the procedure. We prospectively recruited subjects who were seen for clinical care in the Emergency Department or Institutional Concussion Center. Inclusion criteria were the following: 1) adult individuals in the age range of 18–65 years; 2) diagnostic MTBI criteria defined by the American Congress of Rehabilitation Medicine, including either loss of consciousness of <30 minutes or altered consciousness at time of the accident and a Glasgow Outcome Score of 13–15; and 3) injury within 4 weeks. We excluded patients with the following: 1) a history of traumatic brain injury, neurologic illness, or psychiatric disorder; 2) a history of participation in organized contact sports; and 3) any contraindication to MR imaging. We also excluded non-native English speakers and non-right-handed individuals to avoid any potential confounding effects of language and handedness. We studied 19 patients with MTBI (mean age, 30 ± 7 years; age range, 22–45 years; average time since injury, 16 days; 9 men) and 20 HCs (mean age, 33 ± 10 years; age range, 19–65 years; 9 men). For all subjects, formal neurocognitive tests including the WAIS-IV working memory subtests were performed, and MR images were acquired within 1 day of neurocognitive tests. Additionally, to characterize subjects, the Wide Range Achievement Test, 4th ed, Word Reading subtest (WRAT-4) was performed, and the scores were converted to intelligence quotient scores as a brief measure of intelligence. Subgroups of the subjects in this study were previously included in 2 works with nonoverlapping hypotheses.

**MR Imaging Protocol**

MR imaging was performed using a 3T MR imaging scanner (Magnetom Skyra; Siemens, Erlangen, Germany). Diffusion imaging was performed with 5 b-values (250, 1000, 1500, 2000, and 2500 s/mm²) using 5 diffusion-encoding direction schemes (6, 20, 20, 30, and 60, respectively). Three images with b=0 s/mm² were also acquired. Multiband (factor of 2) EPI was used for accelerated acquisitions with an anterior-posterior phase-encoding direction. Other parameters included the following: FOV = 220 × 220 mm, acquisition matrix = 88 × 88, number of slices = 56, image resolution = 2.5 × 2.5 × 2.5 mm³, TR/TE = 4900/95 ms, bandwidth = 2104 Hz/pixel, a generalized autocalibrating partially parallel acquisition factor of 2. An additional image with b=0 s/mm² with a reversed phase-encoding direction was acquired for geometric artifact correction. The total scan time was 12 minutes.

**Working Memory Assessment**

Working memory was assessed with age-appropriate WAIS-IV subtests, which included Digit Span and Letter-Number Sequencing (LNS). In the Digit Span Forward (DSF) task, examinees repeat a sequence of numbers read to them. In the Digit Span Backward (DSB), the same procedure is followed, except that examinees repeat the numbers in reverse order. In the Digit Span Sequencing (DSS), examinees repeat the numbers in ascending order. In the LNS task, examinees separate numbers from letters and state in ascending/alphabetical order a mixed sequence of numbers and letters read to them. Raw scores were converted into standardized age-corrected z scores with a zero mean and a unitary variation, with higher scores indicating better performance.

**Image Analyses**

**Diffusion Image Processing.** The diffusion images underwent the preprocessing steps including Marchenko-Pastur principal component analysis denoising. Gibbs correction, distortion correction with the topup command in FSL (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/topup), eddy current distortion and motion correction with the eddy command in FSL (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/eddy), and outlier detection. In total, 11 diffusion metrics including DTI (FA, mean diffusivity [MD]), axial diffusivity [AD], radial diffusivity [RD]), DKI (mean kurtosis [MK], axial kurtosis [AK], radial kurtosis [RK]), and WMTI (AWF, intra-axonal diffusivity, extra-axonal axial diffusivity, and extra-axonal radial diffusivity) metrics were calculated using in-house software developed in Matlab R2017a (MathWorks, Natick, Massachusetts).

**Tract-Based Spatial Statistics.** We used the standard Tract-Based Spatial Statistics (TBSS; http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/TBSS) to reveal possible correlations between working memory test z scores and diffusion metrics. Briefly, subject FA maps were normalized to the FA template through a nonlinear coregistration, and voxelwise statistical analysis was performed on FA values.
uses projected onto the FA skeleton by looking for local maximum values perpendicular to the skeleton using a permutation-based nonparametric testing randomize command in FSL (https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/Randomise/UserGuide) with the threshold-free cluster enhancement (TFCE) option. All other parametric maps underwent the same transformations and processes. The tract skeleton was thresholded at an FA of 0.2 for DTI and DKI metrics. For WMTI metrics, analysis was restricted to WM regions consisting of single-fiber orientations (FA threshold of 0.4), as recommended.11,33 Age and sex were included as covariates. The number of permutations was set to 5000.

ROI Analysis. ROI analysis was performed on 18 major WM tracts, including genu/body/splenium of the corpus callosum (gCC/bCC/sCC), right and left anterior/posterior limb of the internal capsule, right and left anterior/superior (sCR)/posterior corona radiata (pCR), right and left cingulum, right and left superior longitudinal fasciculus (SLF), and whole WM. ROI regions were generated on the basis of the Johns Hopkins University ICBM-DTI-81 WM labels atlas.32 Briefly, all subjects’ FA maps were nonlinearly registered to the FA template; then, a “reversed warping” procedure was performed to assign the atlas labels to each subject’s space. The ROIs in each subject’s space were manually corrected if necessary. For each ROI, the mean value was obtained only in voxels with FA ≥ 0.2 for DTI and DKI metrics and with FA ≥ 0.4 for WMTI metrics, to restrict analysis to WM regions, as recommended.11,33

Statistical Analysis
The MANCOVA was used to test group differences in the length of education and WRAT-4 intelligence quotient scores, with age and sex as covariates, using SPSS Statistics software, Version 25.0 (IBM, Armonk, New York). Results were considered significant for \( P < .05 \).

For TBSS, a statistical threshold level of \( P < .05 \) was applied after family-wise error correction for multiple comparisons.

For ROI analysis, both the Pearson partial correlation and Spearman rank correlation were performed to measure the associations between diffusion metrics and WAIS-IV subtest scores in each ROI, using SPSS. Age and sex were included as covariates. All \( P \) values were corrected for multiple comparisons using the Benjamini-Hochberg correction. A statistical threshold level of corrected \( P < .05 \) was used. Correlation coefficients were calculated. We also used Fisher R-to-Z transformation18 to test for differences in between-group correlations.

On the basis of the ROI analysis results, specific regions demonstrating significant correlations with working memory performance were further interrogated by dividing subjects into subgroups based on their working memory test \( z \) scores (<1, ≥1) and time since injury (<2 weeks, 2–4 weeks). Subgroup comparisons were performed using MANCOVA, with age as a covariate. Results were considered significant at \( P < .05 \).

RESULTS
The average length of education for patients with MTBI was 15.5 ± 1.7 years, not statistically different from that in healthy controls (16.3 ± 1.8 years). Also, WRAT-4 intelligence quotient scores were not statistically different between MTBI (108.5 ± 12.0) and HC (113.6 ± 14.3) groups. WAIS-IV subtests were not significantly correlated with age and length of education for MTBI and HC groups, except for positive correlations between the length of education and DSS in HCs (\( P = .03 \)).

From TBSS analysis, in the MTBI group, we found a significant positive correlation between AK and DSB, primarily in the right SLF (Fig 1A), which was not present in the HC group (Fig 1B). Most interesting, we found complete loss of relationships between FA and LNS in the MTBI group (Fig 2A), while a significant correlation was shown in the HC group (Fig 2B), most notably in parietal WM, sCR/pCR, bCC/sCC, and SLF. We also found no significant correlation between AWF and LNS in the MTBI group (Fig 3A), while there were multiple areas showing a statistically significant positive correlation between AWF and LNS in the HC group, involving parietal WM, sCR/pCR, and bCC/sCC (Fig 3B). No other diffusion metrics showed areas of significant correlation surviving correction for multiple comparisons.

ROI analysis also found a significant correlation between AK and DSB in the right SLF in the MTBI group (Pearson \( R = 0.69, P = 0.04; \) Spearman \( \rho = 0.75, P = 0.01 \)) that was not present in the HC group. The correlation coefficients observed in the MTBI and HC groups were significantly different (Fisher R-to-Z transformation, \( P = .01 \)). On the other hand, we also observed loss of relationships between FA and LNS in the MTBI group but found a significant positive correlation in the right pCR in the HC group (Pearson \( R = 0.67, P = 0.04; \) Spearman \( \rho = 0.57, P = 0.09 \)). The correlation coefficients observed in the MTBI and HC groups were not significantly different, but there was a trend toward significance (Fisher R-to-Z transformation, \( P = .06 \)). No other significant correlations were found after correction for multiple comparisons. These results are summarized in Table 1.

On the basis of the results of the ROI analysis, subjects were further divided into subgroups according to their working memory test \( z \) scores (<1 or ≥1) and time since injury (<2 weeks or 2–4 weeks). Details of the subgroup characteristics are given in Tables 2 and 3. With regard to subgroup analysis, we found a significant difference in AK between the MTBI subgroup of 2–4 weeks from injury with DSB <1 and the MTBI subgroup
of 2–4 weeks from injury with $\text{DSB} \geq 1$ within the right SLF (Fig 4A). Significant differences in FA were shown between the MTBI subgroup of <2 weeks of injury with $\text{LNS} < 1$ and the HC subgroup with $\text{LNS} < 1$, and between the HC subgroup with $\text{LNS} < 1$ and the HC subgroup with $\text{LNS} \geq 1$, within the right pCR (Fig 4B).

**DISCUSSION**

This study shows significant differences in the relationships between specific WM microstructural markers and auditory-verbal working memory performance between patients with MTBI and healthy controls. Specifically, in the MTBI group, a significant correlation between AK and DSB was present in the right SLF, while the HC group demonstrated no such relationship (Fig 1), a finding observed using both TBSS and ROI analyses. Patients with MTBI also appear to lose the normal associations seen in controls between diffusion metrics (FA, AWF) and LNS (Figs 2 and 3). Furthermore, subgroup analyses also showed a significant difference in AK between the MTBI subgroup of 2–4 weeks of injury with $\text{DSB} < 1$ and the MTBI subgroup of 2–4 weeks of injury with $\text{DSB} \geq 1$. A significant difference was also found between the MTBI subgroup of <2 weeks of injury and the HC subgroup, both having $\text{LNS} < 1$.

Notably, the relationship between AK and DSB in patients with MTBI is mainly present in the right SLF (Fig 1), a structure critical for attention, memory, emotion, and language, linking frontoparietal WM regions critical to working memory. In particular, the right SLF is critical for attention, visuospatial function, and short-term memory. Previously, similar results were reported showing relations between the SLF and working memory deficits assessed by FA and the visual 2-back d-prime index in patients with severe and diffuse traumatic brain injury. In this study, we found decreased AK in the right SLF, reflecting decreased tissue complexity along the long axis of the axon in patients with MTBI with...

Table 1: Results of the ROI analysis showing significant positive correlations between AK and DSB in the MTBI group, and between FA and LNS in the HC group

| Diffusion Metrics vs Working Memory | ROI       | MTBI Pearson | MTBI Spearman | HC Pearson | HC Spearman | Fisher R-to-Z (P) |
|-----------------------------------|-----------|--------------|--------------|------------|-------------|-----------------|
| AK vs DSB                         | Right SLF | 0.69a        | 0.75b        | 0.04       | 0.67b       | 2.32b (0.03)    |
| FA vs LNS                         | Right pCR | 0.25         | 0.75         | 0.11       | 0.04        | -1.59 (0.15)    |

*a Corrected P values after Benjamini-Hochberg correction for multiple comparison.

*b Significant results.
sists of several separate pathways connecting the cerebral cortex to complex bundle of fibers related to working memory because it contributes to axonal injury known to occur after MTBI, because AWF is a measure that indicates disruption in this normal relationship, presumed to relate to WM. However, in patients with MTBI, we found reactive astrogliosis.

Table 2: Subgroup characteristics defined by their working memory test z scores (LNS <1 or ≥1) and time since injury (<2 weeks or 2–4 weeks)*

|                  | LNS <1 |                  | LNS ≥1 |
|------------------|--------|------------------|--------|
|                  | MTBI (<2 Weeks) | MTBI (2–4 Weeks) | HC     | MTBI (<2 Weeks) | MTBI (2–4 Weeks) | HC     |
| No.              | 5      | 10               | 9      | 3                  | 1                  | 11     |
| Age              | 31 ± 6 | 31 ± 9           | 31 ± 8 | 25 ± 3             | 31                 | 35 ± 12 |
| LNS              | −0.13 ± 0.51 | 0.07 ± 0.41     | 0.07 ± 0.32 | 1.55 ± 0.39     | 2                  | 1.82 ± 0.62 |

*Data are means unless otherwise indicated.

Table 3: Subgroup characteristics defined by their working memory test z scores (DSB <1 or ≥1) and time since injury (<2 weeks or 2–4 weeks)*

|                  | DSB <1 |                  | DSB ≥1 |
|------------------|--------|------------------|--------|
|                  | MTBI (<2 Weeks) | MTBI (2–4 Weeks) | HC     | MTBI (<2 Weeks) | MTBI (2–4 Weeks) | HC     |
| No.              | 4      | 12               | 12     | 4                  | 4                  | 8      |
| Age              | 30 ± 6 | 29 ± 8           | 31 ± 7 | 37 ± 8             | 26 ± 4             | 37 ± 14 |
| DSB              | −0.67 ± 0.27 | −0.24 ± 0.25    | 0.08 ± 0.38 | 1.42 ± 0.5       | 1.58 ± 0.32       | 1.79 ± 0.71 |

*Data are means unless otherwise indicated.

FIG 4. Results of subgroup analysis. A. Boxplots of AK show a significant difference between the MTBI subgroup of 2–4 weeks from injury with DSB <1 and the MTBI subgroup of 2–4 weeks from injury with DSB ≥1. B. Boxplots of FA show significant differences between the MTBI subgroup of <2 weeks of injury and LNS <1 and the HC subgroup with LNS <1 and between the HC subgroup with LNS <1 and the HC subgroup with LNS ≥1. The asterisk indicates P < .05; w, weeks.

poorer performance on DSB. AK is known from animal validation studies to be affected by axonal injury46 as well as secondary reactive astrogliosis.47

Also differentiating MTBI from HC groups in TBSS analysis was a loss of diffusely present normal associations that we have seen in healthy controls (Figs 2 and 3); in controls, it has been shown that FA and AWF directly correlate with performance on LNS, a complex working memory task.24 However, in patients with MTBI, we found a disruption in this normal relationship, presumed to relate to WM injuries known to occur after MTBI,14 because AWF is a measure that reflects axonal density and/or myelination.

ROI analyses (Table 1) showed related statistically significant correlations focused in the right SLF in patients with MTBI as well as in the right pCR in healthy controls, compared with the more diffuse TBSS results. TBSS uses maximum values projected onto the WM skeleton, making it more sensitive to maximal deviations in diffusion metrics. Both TBSS and ROI analyses revealed positive correlations in the right SLF and right pCR, suggesting that these regions are strong, potential anatomic landmarks important to working memory performance, and possibly impairment. Along with the SLF, a structure critical to working memory,18–40 the corona radiata is also a complex bundle of fibers related to working memory because it consists of several separate pathways connecting the cerebral cortex to subcortical structures,24,48,49 including fiber tracts in the pCR, which connect to the precuneus, a part of the default mode network implicated in working memory performance.50,51

With regard to the subgroup analysis, within the same time-since-injury period of 2–4 weeks, there was a significant difference in AK between the MTBI subgroups with higher and lower working memory performance in the right SLF (Fig 4A). Decreased AK, believed to relate to axonal injury46 and/or secondary reactive astrogliosis,47 may specifically be a more useful indicator in the MTBI subgroup with lower working memory performance. Also, within the lower LNS range, significantly elevated FA was observed in the MTBI subgroup within 2 weeks of injury compared with the corresponding HC subgroup with the same lower LNS range (Fig 4B). Elevated FA was reported frequently within 2 weeks of injury, believed to reflect injury-related cytotoxic edema52 or reactive astrogliosis.53 We found only a significant difference in FA between the MTBI subgroup within 2 weeks of injury, believed to reflect injury-related cytotoxic edema52 or reactive astrogliosis.53

There are several limitations in the presented study. First, there is a wide age range from 19 to 65 years. Any age effects were minimized using age-corrected WAIS-IV subtest z-scores derived from the published normative sample (n = 2200) divided into 13 age bands, spanning ages 16–90.25 Moreover, age is included as a covariate in all statistical analyses. Second, subgroups were defined on the basis of a somewhat arbitrary statistical threshold value for LNS and DSB (Fig 4). Further work could study various performance groups. Third, this study includes a relatively small
CONCLUSIONS

There are differences between patients with MTBI within 4 weeks of injury and healthy controls in terms of the relationships between brain microstructure and working memory performance. These findings may relate to known WM injury and changes in functional organization occurring after MTBI. Further study on the effect of time since injury on working memory performance may provide insight into the temporal dynamics of working memory deficits in patients with MTBI.

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