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Susceptibility-weighted imaging study in male and female ice hockey players over a single season

Clinical article

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Object. Concussion, or mild traumatic brain injury (mTBI), is a commonly occurring sports-related injury, especially in contact sports such as hockey. Cerebral microbleeds (CMBs), which appear as small, hypointense lesions on T2*-weighted images, can result from TBI. The authors use susceptibility-weighted imaging (SWI) to automatically detect small hypointensities that may be subtle signs of chronic and acute damage due to both subconcussive and concussive injury. The goal was to investigate how the burden of these hypointensities changes over time, over a playing season, and postconcussion, in comparison with subjects who did not suffer a medically observed and diagnosed concussion.

Methods. Images were obtained in 45 university-level adult male and female ice hockey players before and after a single Canadian Interuniversity Sports season. In addition, 11 subjects (5 men and 6 women) underwent imaging at 72 hours, 2 weeks, and 2 months after concussion. To identify subtle changes in brain tissue and potential CMBs, nonvessel clusters of hypointensities on SW1 were automatically identified, and a hypointensity burden index was calculated for all subjects at the beginning of the season (BOS), the end of the season (EOS), and at postconcussion time points (where applicable).

Results. A statistically significant increase in the hypointensity burden, relative to the BOS, was observed for male subjects with concussions at the 2-week postconcussion time point. A smaller, nonsignificant rise in the burden for female subjects with concussions was also observed within the same time period. There were no significant changes in burden for nonconcussed subjects of either sex between the BOS and EOS time points. However, there was a statistically significant difference in the burden between male and female subjects in the nonconcussed group at both the BOS and EOS time points, with males having a higher burden.

Conclusions. This method extends the utility of SWI from the enhancement and detection of larger (> 5 mm) CMBs, which are often observed in more severe cases of TBI, to cases involving smaller lesions in which visual detection of injury is difficult. The hypointensity burden metric proposed here shows statistically significant changes over time in the male subjects. A smaller, nonsignificant increase in the burden metric was observed in the female subjects.

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Key Words
- concussion
- susceptibility-weighted imaging
- mild traumatic brain injury
- ice hockey
- sex-based difference

Abbreviations used in this paper: BOS = beginning of the season; CMB = cerebral microbleed; DWI = diffusion-weighted imaging; EOS = end of the season; GRE = gradient recalled echo; HIB = hypointensity burden; ImPACT = Immediate Postconcussion Assessment and Cognitive Test; mTBI = mild traumatic brain injury; SCAT2 = Sport Concussion Assessment Tool–2; SWI = susceptibility-weighted imaging.

* Drs. Shenton and Echlin share senior authorship of this work.

It is estimated that there are 1.7 million traumatic brain injury (TBI) occurrences from all causes per year and that emergency department visits and hospitalizations due to TBI are increasing.18 This number does not include instances of TBI that are either treated by a primary care physician or are left untreated, so the actual number is expected to be higher. Sports are increasingly recognized as a significant source of TBIs in the young adult popu-
Susceptibility-weighted imaging study in ice hockey players

Cerebral microbleeds (CMBs) have been observed to accumulate, it has also been shown that the outcome for TBI may be normal for most patients. In noninvasive imaging modalities such as CT scanning or conventional MRI, concussions are difficult to measure and the results from measuring the net motion of the tissue water one can infer the integrity of the structures. Given this, DWI has been used to search for evidence of diffuse axonal injury in concussion, and we can also investigate the initial burden of low-intensity clusters that may occur in response to concussion. Through this method we can detect changes in the number of low-intensity clusters that may occur in response to concussion, and we can also investigate the initial burden of these clusters in each sex, how the clusters correlate with numbers of previous concussions, and how they change over time when no further concussion is reported.

Methods

Study Subjects

A cohort of 45 adult hockey players (25 men and 20 women) underwent imaging at the beginning of the season (BOS), the end of the season (EOS), and at 72 hours, 2 weeks, and 2 months postconcussion using an SWI protocol. Data from these subjects were processed and analyzed. However, not all of the participants who suffered a concussion during the season underwent imaging at each time point, due to subject withdrawal from the study or scheduling conflicts. For example, 4 of the male subjects were not included in the study either due to claustrophobia at imaging, self-withdrawal from the study, or noncompletion of the BOS imaging. The mean age of the patients at the BOS was 23 ± 2 years for the men and 21 ± 4 years for the women. From these 2 groups there were 5 confirmed concussions for the men and 6 for the women during the season. The internal review boards of the participating universities approved this study.

Previous concussions were self-reported by these subjects. Of the 16 men who did not have concussions during the season, 3 had 2 previous concussions and 5 had 1 previous concussion. Of the 14 women who did not have T2*-weighted gradient recalled echo (GRE) scans, which are sensitive to hemorrhagic regions due to the paramagnetic nature of blood breakdown products, for example hemosiderin. However, standard MRI methods have shown mixed results when investigators have tried to correlate imaging results with clinical outcomes.

In addition to structural measurements, one can use MRI to detect CMBs that often accompany TBI over a range of severity. Susceptibility-weighted imaging (SWI) is a relatively new MRI modality that involves further processing of a GRE data set. The contrast found in SWI is generated through the sensitivity of the phase measurement to disruptions in the magnetic field. The phase is sensitive to differences in magnetic susceptibility that, in the case of TBI, are provided by the iron present in pools of static blood resulting from (micro)hemorrhages. The SWI modality takes these phase disruptions and uses them to generate contrast that can greatly enhance the detection of small pockets of blood. The sensitivity of SWI is greater than that of a standard GRE sequence in which the contrast is related to the T2* distribution in the tissue. The current study is the first to use SWI to assess the effects of sports concussion in a prospective manner.

In the case of concussion, it is not expected that there will be the type of large CMBs (> 5.7 mm has been suggested for visual inspection studies) that have been observed to accompany stroke or severe TBI. Instead, we use SWI to detect smaller, more subtle intensity changes in the sensitized images. Our method, described below, is to interrogate the time course of clusters of low-intensity voxels. Through this method we can detect changes in the number of low-intensity clusters that may occur in response to concussion, and we can also investigate the initial burden of these clusters in each sex, how the clusters correlate with numbers of previous concussions, and how they change over time when no further concussion is reported.

Methods

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concussions during the season, only 1 had had a previous concussion. Several of the subjects who suffered a concussion during the season of interest also had previous concussions. Of the 5 men who had at least 1 concussion during the season (1 of the injured men had 2 concussions during the season between the 2-week and 2-month time points, and hence did not complete the 2-month and EOS time points), 2 of them had 1 previous concussion, 2 of them had 2 concussions, and a single subject reported 3 previous concussions. Of the 6 women who had a concussion during the season, 3 of them had no previous concussions, 1 had 1 previous concussion, 1 had 3 previous concussions, and a single subject reported 4 previous concussions.

Imaging Protocol

Subjects underwent imaging on a 3-T Philips Achieva equipped with an 8-channel head coil. The SWI data set was acquired using a multiecho gradient-echo pulse sequence consisting of 5 echoes, with an initial TE of 6 msec; the interval between echoes was 6 msec. To generate the SWI volumes analyzed here, the TE 30 msec data were used. The TR was 34 msec. Slices were acquired in the transverse plane and the acquired FOV was 220 × 128 mm with a voxel volume of 0.5 × 0.5 × 2 mm³. This was then reconstructed to a voxel volume of 0.491 × 0.491 × 1 mm³. A SENSE factor of 1.2 was used in the right-left direction, and the partial Fourier k-space acquisition percentage factor was 67%. The flip angle was 17° and the total acquisition time was 6 minutes and 39 seconds. The magnitude and phase images were saved for each volume in the PAR/REC format of the Philips system and were used in the SWI processing. It is also to be noted that a gradient coil change occurred during the collection of these data. Its effect on the imaging data and the final SWI volume voxel intensities was investigated (see below). However, because our goal here was to detect nonspecific “hypointensity” rather than to compare calculated parameter values (such as the apparent diffusion coefficient or the spin-spin relaxation time), it was not expected that the coil change would significantly affect these results. Nonetheless, analysis of the image intensity range and statistics was undertaken to ensure that this was in fact the case.

Image Processing

The SWIs were processed using in-house code written in C++ and incorporated functions from the FreeSurfer libraries. First, background pixels were removed from the magnitude volumes using the Brain Extraction Tool (BET) from the Functional MRI of the Brain (FMRIB) Software Library (FSL). A mask was then generated from the resulting brain-only volume and applied to the phase volume before the phase was unwrapped using the PRELUDE method, also from Functional MRI of the Brain (FMRIB) Software Library (FSL). Seven initial phase splits were chosen as an input to PRELUDE, and that value was selected as a compromise between speed of processing and quality of the resulting phase unwrapping. To create the SWI volume, the following steps were performed: 1) a gaussian smoothing filter of 2-mm kernel width was applied to the unwrapped phase volume; 2) the resulting smoothed volume was subtracted from the original phase volume, resulting in an effectively high-pass filtered phase volume; 3) a negative phase mask of the form for \( q > 0 \rightarrow q = 1 \); for \( q \leq 0 \rightarrow q = 1 + (q/\pi) \), where \( q \) is the phase, was multiplied by itself 4 times and applied to the high-pass filtered unwrapped phase mask; and 4) this result was then multiplied by the masked magnitude volume, resulting in the final SWI volume. In addition, a minimum-intensity projection over 4 slices in the imaging-volume slice direction was performed and used to identify vessels, and hence the maximum-intensity threshold described in the following section.

Image Analysis

Analysis of the SWI volumes was performed using in-house code written in Matlab (The Mathworks, Inc.). Our hypothesis was that there would be few or no large hypointense regions that could be identified through visual inspection, and therefore our analysis focused on detecting smaller hypointensities that would be difficult to detect through visual inspection. In this analysis, the intensity that corresponded to maximum value of the lowest 10% of pixel intensities was used as a threshold value, and each voxel whose intensity value was at or below that level was recorded. In addition, the code also calculated the minimum/maximum/mean/median of the SWI voxel intensities for each subject to determine whether different image intensity ranges existed in the data for different cohorts or across the coil change date.

The algorithm then identified connected clusters of these marked voxels by using the 26-neighbor criterion for 3D space. This not only includes the closest neighboring voxels, but also neighboring voxels on the diagonal. To reduce the possibility of any vessels being included in these clusters, an upper limit on the cluster size included in the analysis was set at 35 voxels. The exact number chosen for either the cluster-size threshold or the image-intensity threshold was not found to be critical to the results presented here. Various cluster and intensity thresholds were chosen and similar results were found for all (data not shown), as long as the cluster threshold was low enough not to include obvious vessels that were identified as such by following their path manually. In addition, this cluster threshold corresponds roughly to a 3-voxel × 4-voxel hypointense region that is connected across 3 slices (~ 1.5 × 2 × 3 mm³), and it was determined to have eliminated almost completely the observable vessels in these data from the cluster set. No minimum cluster size was set because single-voxel “clusters” were not observed to contribute significantly to the final volume of interest.

From the total number of voxels that met the criterion that the cluster size be less than 35 voxels, we calculated the hypointensity burden (HIB), which we define as follows: \( \text{HIB} = (\text{total number of voxels in accepted clusters/total number of brain voxels}) \times (\text{volume in mm}^3 \text{ of 1 voxel}) \). The HIB gives the scaled percentage of brain volume that contains nonvessel hypointensities. The measure is useful because it allows us to detect both potential microbleeds as well as other more subtle forms of damage that could present as hypointensity, such as neural microstructural changes examined in the accompanying articles in this current series (Pasternak et al., and Sasaki et al.).
Advantages of this measure include the fact that it is completely automatic, the level of HIB can be investigated over time, and it can be compared at baseline to potentially characterize levels of previous damage. In addition, hypointensities that are not due to recent damage are not expected to change over time, and therefore add only a constant offset and would not affect the postconcussion HIB time course.

Results

Figure 1 shows orthogonal views from a representative subject; it depicts the SWI with an overlay of the discovered clusters. The exact shapes of the clusters are difficult to visualize in a single plane because clusters can be in more than a single slice. Note that obvious in-plane vessels are not discovered by the algorithm, nor are large vessels perpendicular to the plane of the image. Note also that, due to the small size of these clusters, they would not generally be classified as “traditional” CMBs (diameters larger than ~5 mm).

In Fig. 2, we show the HIB level of the 13 female subjects (Fig. 2 upper) and the 8 male subjects (Fig. 2 lower) in the nonconcussed group as a function of the number of self-reported previous concussions. There was no statistical difference in HIB level for the male subjects, regardless of the number of previous concussions. Note, however, that the HIB level was higher for the male than for the female subjects, even for the participants with no reported concussions, and this difference was statistically significant (p < 0.003, 2-tailed test, unequal variance) for that group. Note that there were not the same numbers of subjects at each number of previous concussions (see above in the discussion of the subjects for the multiplicity at each number). There was only a single female subject who had 1 previous concussion, and who also did not have an additional concussion during the season (nonconcussed group).

Figure 3 shows the HIB levels for all nonconcussed subjects (female subjects designated by circles and male subjects by squares) irrespective of previous concussions at each of the 2 imaging time points. Between sexes, the BOS and EOS data were significantly different from each other (BOS, p < 0.0005, EOS, p < 0.05; unpaired t-test, 2-sided, unequal variance). The BOS and EOS data for the nonconcussed group did not, however, differ significantly from the BOS to the EOS within each sex.

In Fig. 4 we show the HIB versus time point curves for the concussion groups. Figure 4A shows the individual data for the 6 female subjects and Fig. 4B shows the individual data for the 5 male subjects. Note that for the male subjects, the last time point only includes data from 2 individuals; the others in this group did not complete the final imaging time point. The cohorts for both sexes consisted of participants who had HIB levels that changed over time and those for whom the HIB level remained constant (2 of 5 men, 2 of 6 women). The 1 female subject who completed only the BOS imaging time point, but later suffered a concussion, was not included in these data. For 2 of the subjects (1 male and 1 female), 1 of the postconcussion time points was coincident with the EOS time point for the other subjects. The data for these subjects were included in the calculations for, and
displayed with, the appropriate postconcussion time point rather than at EOS.

In Fig. 4C we show the mean ± SD of the HIB for both sexes on a single plot to compare the HIB time course for each sex. The difference in mean HIB values between the sexes was statistically significant for the 2-week time point. Note that the overall change from BOS to HIB maximum was greater for the male than for the female subjects. We measured this increase by calculating the mean change in HIB at 2 weeks postconcussion compared to the BOS: $\Delta HIB = \Sigma (HIB (2 \text{ weeks}) - HIB (BOS))/N$, where N is the number of subjects, and found that $\Delta HIB$ (female) = 0.0003 ± 0.0004, whereas $\Delta HIB$ (male) = 0.0006 ± 0.0007. Note that although the mean HIB level had its peak at the 72-hour time point in women in this study, this may be due to the fact that SWI could be obtained in only half of the female subjects at this time point. It should also be observed that there were subjects for each sex whose HIB value generally did not change with time.

Figure 5 displays the mean ± SD for the concussion group and all nonconcussed subjects, regardless of previous concussion status, for female subjects (Fig. 5 upper) and male subjects (Fig. 5 lower). The concussed subjects are represented by the diamonds and the nonconcussed subjects by the triangles. Note that the BOS and EOS data were almost identical for female subjects in both cohorts, but that this was not the case for male subjects. In male subjects, the EOS and BOS levels in the nonconcussed group were higher than those for the concussed cohort.

The HIB data at each time point were tested for statistical significance, versus that at the BOS time point for both sexes. Only the 2-week time point for the concussed male subjects’ data showed a statistically significant increase over the BOS data (asterisk in Fig. 5 lower).

**Discussion**

*Study Overview*

Although most studies use visual inspection to detect CMBs, this strategy is less useful for cases of milder injury that may not produce lesions with enough reduced image intensity (or volume) to be detected by the eye alone. One must also be careful not to interpret the discovered...
areas depicted in Fig. 1 as traditionally defined CMBs (for example, > 5.7 mm in patients with cerebral amyloid angiopathy). Here we tailor our analysis method to the expectation that there will be few if any CMBs of that size in our cohort and that the damage may be occurring at a much smaller-length scale. We note that there have been recent observations of submillimeter pericapillary hemosiderin, which may be contributing to the clusters discovered in this work.

We also note that we make no claim that this simple method specifically detects microbleeds, and there will inevitably be a background of false-positive detections. For example, there are a few small clusters present in the aqueduct in Fig. 1. However, we believe that, given the time course of the HIB metric postconcussion, we are also detecting changes that are specific to the concussion. Although it is true that there will be a baseline of clusters detected that are not due to the concussion, this baseline will be manifest as a common offset for each group. This common offset does not, however, explain the sex-based difference in the BOS HIB, which will be discussed below.

Given that each voxel may produce a signal that arises from tissue in multiple states of health or injury, the voxel intensity may not be reduced to the same extent as it is in larger or multiple lesions resulting from stroke or more severe trauma. Although we observed few visually obvious (≥ 5 mm) signs of damage in these data, we did find, by characterizing and following clusters of the lowest-intensity voxels, time-dependent behavior of our HIB metric after concussion. In addition we note that, in the nonconcussed group data, the BOS and EOS HIB levels were the same, implying that the observed time dependence of HIB in the concussed group data was not due to the hardware or threshold, but rather was related to the concussion. It is also important to note that we are only discovering a population of connected voxels of a certain size. This was done to reduce the possibility of detecting vessels with the algorithm, but note that large CMBs would also not be detected, and therefore the algorithm is tuned to our current application. This method is also fully automatic and less computationally intensive than the method of Barnes et al., which uses support vector machines to separate CMBs from other hypointensities.

**Hardware Upgrade**

One complication in the analysis of these data was that the gradient coil was changed during the season. However, it is not likely that this change would affect the selection of the lowest 10% of image intensities because it is unlikely that the gradients would decrease higher image intensities into the lowest decile in significant numbers sufficient to affect the results presented here. We also found no evidence of regional heterogeneity in the final clusters selected for analysis. In addition, we found no statistically significant difference in image intensity range, median, or maximum value from the BOS to EOS time points (within the sexes). In addition, because the within-group HIB in all groups is statistically the same for BOS and EOS, this strongly suggests that the coil change was not a significant factor in the results of this analysis.

**Sex-Based Differences in the HIB Metric**

To our knowledge, this is the first study to examine sex-based differences in patients with sports-related concussions that included prospective incidence rates and evidence of organic brain injury. The first striking feature of Figs. 2 and 3 is the difference in HIB level between the male and female subjects. This difference occurred at all time points, and for both the nonconcussed and concussed groups. One possible explanation for the difference in the BOS HIB level between the sexes is that the HIB lowest 10% threshold is different between the sexes, and therefore the difference in threshold allows more clusters to be discovered. There was, in fact, a statistically significant difference in the HIB lowest 10% threshold between the male and female subjects in the nonconcussed group data (using that as the data set with the largest number of individuals). This was true for both the BOS (p < 0.002, 2-tailed, unequal variance) and the EOS (p < 0.05, 2-tailed, unequal variance).

One would expect that the data with the largest threshold would have the largest range of values (assuming that...
the intensity-value histograms have similar shapes), but this was not the case. We find that although the female subjects had the largest intensity threshold value—mean and median—the ranges of intensity values were larger for the male subjects. Therefore, the SWI intensity-value distributions were statistically different between the sexes, but not in a way that explains the sex-based difference in HIB values; that is, for a higher threshold one would expect more clusters to be detected and therefore a higher detected HIB. In fact, the HIB level for the female subjects was less than that for the male subjects in both groups. In addition, there was no statistically significant difference in the threshold between time points within each sex.

One hypothesis for the shift of HIB to lower levels is that the male subjects may have accumulated more subconcussive hits resulting in hypointensities over their careers and that this resulted in cumulative damage and/or HIB and higher BOS HIB. In addition, whereas the total number of previous concussions was approximately the same for each sex in the concussed group, the distribution among subjects was not the same. More specifically, all of the male subjects in the concussed group had had 1 or more previous concussions, whereas this was not the case for the female subjects (see more detailed discussion below). It should be noted that although the number of previous concussions was recorded in this study, these were self-reported, and no other information regarding the concussions was available.

The time courses of mean HIB for the male and female subjects were roughly equivalent, but there is obviously a range of behavior in the individual subjects. In addition, the maximum increase in HIB was greater for the male subjects. This greater increase, although not statistically significant due to the small number of subjects in each group (and also taking into account that there was little change over time for 2 subjects in each group), was nonetheless suggestive of a larger effect for the male than for the female subjects. This may be due to differences in degree of trauma for these particular individuals, but should be investigated with a larger cohort. It is also interesting that the HIB level renormalized by the EOS time point for both sexes.

Previous Concussions

Figure 5 allows the direct comparison of the nonconcussed and concussed group data. Note that the BOS and EOS HIB levels were identical for both groups of female subjects, but not for the male subjects. This may be due to several factors. First, the number of previous concussions in the concussed female subjects was \{4,3,0,0,0,1\}; given in order of highest to lowest BOS HIB level, whereas for the concussed male subjects the number of previous concussions was \{1,2,3,1,2\}. It is interesting that the number of previous concussions correlated well with the observed BOS HIB level in the female subjects. The picture from the concussed group data of the male subjects is more complicated, and points to the importance of a method for estimating cerebral damage postconcussion.

As noted earlier in this paper, the number of previous concussions was self-reported, and there is a tendency for the male subjects to underreport the number and severity of concussions. In addition, the nonconcussed cohort of male subjects reported more previous concussions (11 total in 16 males) than did the female cohort (1 total in 14 females), and this could account for the higher BOS and EOS HIB levels in the nonconcussed group of male subjects. Assuming that the HIB level reflects damage due to concussion, the smaller number of previous concussions in the female subjects is consistent with the lower overall HIB level in the nonconcussed group, and with the BOS and EOS levels matching at those time points for both groups of female subjects. In addition, the sole statistically significant increase in postconcussion HIB versus BOS HIB was observed in the cohort of male subjects. This, along with the fact that each individual in the concussed group of male subjects had previous concussions, is consistent with the idea that damage due to concussions may be cumulative.27 Hockey players are a unique group, both because of the observed rate of concussion in the sport for both sexes, and because the nature of routine play is such that it is unlikely that a player can avoid any sort of trauma to the head (even when concussions are neither identified nor self-reported). Given what is known about the occurrence of multiple subconcussive blows and later concussions, this method may be useful in the detection of cumulative effects of concussion.

Participation Rates

We also note that there was a lower participation rate among the male subjects than the female subjects—the male subjects failed to undergo imaging at the 2-month and EOS time points at a higher rate than the female subjects. The documented participation rate demonstrates the difficulty inherent in these studies. The participation goals of the athlete and the researcher conflict, because the identification of concussion results in the injured athlete being restricted from competition.

Study Limitations

There are several limitations in this study. The strength of this study was that independent specialist medical observers were involved in direct concussion identification and diagnosis. Therefore it was more likely that a concussion would be identified and recorded. This independent direct observation and diagnosis maximizes the number of concussions that would be identified for each group. Subjects were also broken into groups differentiated by sex and diagnosis (concussion/no concussion). However, not all subjects who suffered a concussion during the season completed all of the scheduled imaging time points. This was primarily due to player perception that participation in the study had the potential to curtail his or her ability to return to play after a concussion. Interestingly, overall participation was greater among the female than among the male subjects. Given the variability in subject outcome and response to concussion, the low numbers in each group mean that large effects in HIB are needed to achieve statistical significance.

Another limitation of this study was that the Immediate Postconcussion Assessment and Cognitive Test (ImPACT) and Sport Concussion Assessment Tool–2
(SCAT2) scores collected were also highly variable. Echlin and colleagues,16 in an earlier publication that presented the clinical outcomes data, attributed this to one or more of the following: the small sample size, the inherent variability of the tests, or the possibility of malingering of subjects in the baseline tests. Therefore, it is not surprising that no correlation was found between the HIB metric and the ImPACT or SCAT2 scores (data not shown). However, in that earlier study we did find that a statistically significant effect was present for the SCAT2 total symptom score for concussed subjects, which increased (the SCAT2 total symptom score increases as the number of self-perceived symptoms and severity of symptoms increase) at postconcussion and renormalized by the end of the season, similar to the time course of the HIB. Nonetheless, given the factors discussed above, further study of this metric in other populations, and in comparison with pathological data, will be needed to make further inferences regarding a link between the HIB and clinical outcomes.

Another limitation is that imaging was not performed at one or more points during the season for subjects not suffering a concussion. These data would have aided in characterizing the reliability of the HIB metric and would have helped to put the magnitude of changes observed in the concussed subjects in context.

Conclusions

The SWI modality allows us to detect hypointense brain regions that have been shown to be associated with the presence of blood-breakdown products originating from damaged vessels. This method extends the utility of SWI from the enhancement and detection of larger (> 5 mm) CMBs that are often observed in more severe TBI to concussion in which visual detection of injury is difficult. The HIB metric proposed here shows statistically significant changes over time in the male subjects. A smaller nonsignificant increase in the burden metric was observed in the female subjects. Data acquired using this method could be used for the monitoring of players throughout their careers and could lead to improved diagnoses and return-to-play guidelines.

Disclosure

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