Decreased susceptibility of major veins in mild traumatic brain injury is correlated with post-concussive symptoms: A quantitative susceptibility mapping study

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ABSTRACT

Cerebral venous oxygen saturation (SvO2) is an important biomarker of brain function. In this study, we aimed to explore the relative changes of regional cerebral SvO2 among axonal injury (AI) patients, non-AI patients and healthy controls (HCs) using quantitative susceptibility mapping (QSM). 48 patients and 32 HCs were enrolled. The patients were divided into two groups depending on the imaging based evidence of AI. QSM was used to measure the susceptibility of major cerebral veins. Non-parametric testing was performed for susceptibility differences among the non-AI patient group, AI patient group and healthy control group. Correlation was performed between the susceptibility of major cerebral veins, elapsed time post trauma (ETPT) and post-concussive symptom scores. The ROC analysis was performed for the diagnostic efficiency of susceptibility to discriminate mTBI patients from HCs.

The susceptibility of the straight sinus in non-AI and AI patients was significantly lower than that in HCs (P < 0.001, P = 0.004, respectively, Bonferroni corrected), which may indicate an increased regional cerebral SvO2 in patients. The susceptibility of the straight sinus in non-AI patients positively correlated with ETPT (r = 0.573, P = 0.003, FDR corrected) while that in AI patients negatively correlated with the Rivermead Post Concussion Symptoms Questionnaire scores (r = −0.582, P = 0.018, FDR corrected). The sensitivity, specificity and AUC values of susceptibility for the discrimination between mTBI patients and HCs were 88%, 69% and 0.84. In conclusion, the susceptibility of the straight sinus can be used as a biomarker to monitor the progress of mild TBI and to differentiate mTBI patients from healthy controls.

1. Introduction

Traumatic brain injury (TBI) is one of the major health problems worldwide and affects over 1.7 million people each year in the United States alone (Paul et al., 2010). It is estimated that 70–90% of TBI cases is mild TBI (mTBI), which is defined by a Glasgow Coma Scale of 13 to 15. This includes those cases where the period of loss of consciousness is < 30 min and the period of post-traumatic amnesia is < 24 h (Ahman et al., 2013). Mild brain injury can have serious post-concussive symptoms including headache, fatigue, dizziness, poor memory, depression, irritability and concentration difficulties. These symptoms can persist for several months to years and significantly affect the quality of life (Ahman et al., 2013; Doshi et al., 2015). Several serious cerebral changes associated with chronic mTBI have been reported, such as: axonal injuries (AI), decreased cerebral blood flow, cognitive dysfunction, chronic traumatic encephalopathy, Alzheimer’s disease,
Therefore, the early detection of cerebral SvO₂ changes may play a
cerebral tissue oxygenation in severe TBI (McCredie et al., 2017).

treatment of red blood cell transfusion had no measurable impact on
hypothermia treatment (Sun et al., 2016). McCredie et al. found the
sure of oxygen correlated with a poor outcome of severe TBI with mild
e of cerebral tissues, indicating a neuroprotective response following mTBI
creased cerebral venous oxygen saturation (SvO₂) may indicate either a
creased cerebral blood flow in mTBI patients exceeds the oxygen demand of
the cerebral tissues, indicating a neuroprotective response following mTBI
(Doshi et al., 2015). Monitoring SvO₂ may also reflect the therapeutic
effect. For example, Sun et al. found the low brain tissue partial
pressure of oxygen correlated with a poor outcome of severe TBI with mild
hypothermia treatment (Sun et al., 2016). McCredie et al. found the
treatment of red blood cell transfusion had no measurable impact on
cerebral tissue oxygenation in severe TBI (McCredie et al., 2017).
Therefore, the early detection of cerebral SvO₂ changes may play a
significant role in better understanding of the patient's pathologic
changes. Near-infrared spectroscopy (NIRS) has been applied to detect the
cerebral oxygenation in TBI patients. However, uncertainty still
exists regarding the reliability of this technology, and it cannot detect the
deep cerebral regional tissues oxygenation due to superficial cov-
erage in the spectrophotometric methods (Davies et al., 2015; Ferrari
et al., 2004).

Recently, quantitative susceptibility mapping (QSM) has been used to
quantify the cerebral SvO₂ (Haacke et al., 2010; Tang et al., 2013; Xu
et al., 2014; Fan et al., 2014; Liu et al., 2017). Unlike other phase-based
methods, the cerebral SvO₂ from QSM is not dependent on the or-
ientation of the cerebral veins. In previous studies, QSM has been used to
explore the changes of regional cerebral SvO₂ in a rat stroke model, healthy individuals and acute ischemic stroke patients (Busch et al.,
2016; Hsieh et al., 2016; Xia et al., 2014). To date, only one paper has
reported relative changes of cerebral SvO₂ in mTBI patients (Doshi et al.,
2015). In that study, Doshi et al. found the decreased venous
susceptibility, indicating increased cerebral SvO₂ in the left thala-
modriate vein and right basal vein of Rosenthal in 14 mTBI patients.

The purposes of this study include: 1) to use QSM to measure the
difference in the susceptibilities of major veins among the mTBI pa-
tients without AI, patients with AI and HCs; 2) to evaluate the di-
nostic efficiency of susceptibility to discriminate mTBI patients from
HCs; 3) to study the relationships between the susceptibility of major
cerebral veins with ETPT and post-concussive symptom scores in mTBI
patients.

2. Materials and methods

2.1. Subjects profiles

This study was approved by the Institutional Review Board of
Tianjin First Central hospital. Written informed consent was acquired from
all the subjects before the MRI study. Data from 49 mTBI patients
were collected from August 2013 to April 2016. The inclusion criteria
were made based on the definition of mTBI by the American Congress
of Rehabilitation Medicine (1993), as follows: 1) the patients were
right-handed and aged 18 years or older without contraindications for
MR (such as any metal in the head and body, pacemaker, claus-
trophydia); 2) the patients had an initial Glasgow Coma Scale score of
13–15; 3) any period of loss of consciousness was approximately 30
min or less; 4) any period of post-traumatic amnesia in patients was < 24
h; 5) the patients may have changes of mental status, such as being dis-
oriented, confused or dazed; 6) all patients had head CT data, and MRI
data including T₁-weighted and T₂ FLAIR images (T₁WI, T₂ FLAIR),
diffusion weighted imaging (DWI) and susceptibility weighted imaging
(SWI); 7) CT imaging appearance of mTBI was negative; and 8) the
patients did not have any other structural imaging changes except for
AI, such as skull fracture, bleeding, cerebral contusion, hematoma,
vessel malformation. The exclusion criteria were: 1) the age of patient
was under 18 years; 2) the patient was pregnant; 3) the patients had a
history of previous TBI; 4) the patient had another disorders including a
neurological disease, brain tumor, epilepsy, seizure, drug or substance
abuse, or any other systemic disease which might affect the cerebral
oxygen metabolism; and 5) the imaging quality was too poor for ob-
servation and analysis because of motion artifact. According to the in-
clusion and exclusion criteria, one patient with poor imaging quality
was excluded due to the motion artifacts in SWI images.

Finally, 48 mTBI patients (16 females, 32 males; age range 19 to
58 years; mean 35.38 ± 10.60 years) were recruited. The diagnostic
criteria of AI included: 1) linear hypointensity on SWI or hyperintensity
on T₂WI and/or DWI and this abnormal signal was found on the in-
terface between white and gray matter; 2) cerebral microbleeds or AI
were found in the white matter of the cerebral cortex, corpus callosum,
brainstem or cerebellum on SWI, which met with the mild AI classifi-
cation of Adams (Adams et al., 1989; Daveva et al., 2015); and 3) there
was no other brain damage such as: cerebral contusion, epidural or
subdural hematoma or subarachnoid hemorrhage, etc. 16 AI patients
were found in this group (6 females, 10 males, age range 19 to 56
years, mean 32.06 ± 10.00 years). The non-AI group was considered when
no abnormalities were found on any MRI sequence (T₁WI, T₂ FLAIR,
DWI or SWI), this included 32 patients (10 females, 22 males, age range
from 19 to 58 years, mean 37.03 ± 10.65 years). Finally, the hema-
tocrit (Hct) (0.27 to 0.50, mean 0.39 ± 0.04) and the ETPT (12 to
360 h, mean 70.73 ± 73.42 h) was recorded for all the patients. 32
HCs (12 females, 20 males, age range 20 to 59 years, mean 37.78 ± 9.81
years) were enrolled from the local community. They
had no history of previous TBI, contraindications to MRI, neurological
problems, brain tumor, epilepsy, seizure, drug or substance abuse, or
any other systematic diseases.

2.2. Post-concussive symptom scores

To evaluate the post-concussive symptoms after mTBI, the Rivermead Post-Concussion Symptoms Questionnaire (RPQ) (King
et al., 1995) was given to all 48 patients within 24 h of admission by a
neurosurgeon with 10-year experiences (R.G.). The RPQ test consists of
16 scoring items including headache, dizziness, nausea, noise sensi-
tivity, sleep disturbance, fatigue, irritability, depression, frustration,
poor memory, poor concentration, taking longer to think, blurred vi-
sion, light sensitivity, double vision and restlessness. These clinical
symptoms were evaluated with a five-point scale corresponding to 5
response alternatives on an ordinal level: never had any symptoms
(category 0), had symptoms but they have resolved (category 1), had
symptoms with weak problems (category 2), had symptoms with
moderate problems (category 3), and had symptoms with severe prob-
lems (category 4). In this study, the total RPQ score was calculated as the
sum of scores for each of the 16 events. Scores could range from 0, if
the patients never had any symptoms (category 0) to 64 if the subjects
had problems in all 16 elements of category 4.

2.3. SWI parameters and processing

All MR imaging was performed at 3.0 T (MRI Siemens Tim Trio
system, Siemens Medical Systems, Erlangen, Germany) using an 8-
channel phased-array coil. Earplugs were used to reduce noise during
the MR scanning and foam pads were applied to keep the head sta-
tionary. To ease the effects of postural changes, all subjects lied in a
dorsal position for at least 30 min before the MRI scanning under
peaceful circumstances. All transverse images were obtained parallel to
the anterior-posterior commissural line on the middle sagittal plane and included the whole cerebral parenchyma. Conventional axial $T_1$WI and $T_2$FLAIR were collected to exclude subjects with other brain structural abnormalities and congenital diseases. Then a high resolution 3D flow-compensated SWI sequence was used and both magnitude and phase images were reconstructed. The imaging parameters of the $T_1$WI sequence were: TR/TE = 250/2.46 ms; number of slices = 21; voxel resolution = $0.9 \times 0.8 \times 5.0$ mm$^3$; field of view = $240 \times 100$ mm; receiver bandwidth = 330 Hz/pixel; flip angle = 70°; 1 acquisition; total acquisition time = 66 s. The imaging parameters of the $T_2$ FLAIR sequence were: TR/TE = 9290/93 ms; TI = 2500 ms; number of slices = 25; voxel resolution = $0.9 \times 0.9 \times 4.0$ mm$^3$; field of view = $220 \times 90.6$ mm; slice thickness = 4.0 mm; receiver bandwidth = 287 Hz/pixel; flip angle = 130°; 1 acquisition; averages 1; total acquisition time = 169 s. The imaging parameters of the DWI sequence were: TR/TE = 5500/130 ms; b = 0/1000; number of slices = 20; voxel resolution = $1.8 \times 1.8 \times 5.0$ mm$^3$; field of view = $240 \times 100$ mm; slice thickness = 5.0 mm; receiver bandwidth = 968 Hz/pixel; EPI factor = 136; 1 acquisition and total acquisition time = 40 s. The imaging parameters of the SWI sequence were: TR/TE = 27/20 ms; voxel size = $0.5 \times 0.5 \times 2$ mm$^3$; field of view = $230 \times 200$ mm; receiver bandwidth = 120 Hz/pixel; 1 acquisition; flip angle = 15°; number of slices = 56; and acquisition time = 179 s. The total acquisition time of $T_1$, $T_2$ FLAIR, DWI and SWI was under 8 min (454 s).

The QSM images were reconstructed using SMART (Susceptibility Mapping and Phase Artifacts Removal Toolbox, Detroit, Michigan, USA) post-processing software (Haacke et al., 2010, 2015) and included the following steps: first, the unwanted low signal regions outside the brain were eliminated using Brain Extraction Tool (BET) in FMRIB Software Library (FSL) (Smith, 2002); the background phase was reduced using a 96 × 96 homodyne high-pass filter (due to the retrospective nature of this study, the raw unfiltered phase images were not available and the phase images obtained using the SWI sequence were already processed by a 96 × 96 homodyne high-pass filter, and hence, no additional filtering was applied during post-processing); finally, QSM data were generated using truncated k-space division with a regularization threshold of 0.1 (Haacke et al., 2010, 2015). These post-processing steps led to a high resolution QSM showing the cerebral venous structures clearly.

The analysis of the QSM data was performed with SPIN (signal processing in nuclear magnetic resonance, Detroit, Michigan, USA) software. The susceptibility of major cerebral veins was measured on the maximum intensity projection (MIP) over 16 slices. To include major veins and eliminate the artifacts from the sinuses cavities, the coverage extended from the highest level of the cerebrum to the lowest level of the cerebellum. Volumes of interests (VOIs) of major veins were outlined manually on the consecutive slices of MIP QSM images, which showed the cerebral veins clearly. VOIs of major veins were delineated according to the normal anatomic structures and distribution regions of cerebral major veins and the susceptibility of cerebral veins were measured by two neuroradiologists, respectively (C.Z. with five years of experience in neuroradiology; C.C. with eight years of experience in neuroradiology), who was blinded to the epidemiological and clinical information of all the participants. The major cerebral veins included the right and left cortical veins, septal veins, thalamostriate veins, internal cerebral veins, the basal veins and straight sinus. Consecutive slices were drawn to cover the body of major veins except for the topmost and lowermost levels of cerebral veins, to avoid the partial volume effect. The VOIs of cortical veins were drawn to include the whole cerebral hemisphere. The VOIs of other cerebral veins were drawn close to the cerebral veins as much as possible and cover the whole distribution regions of cerebral veins (Fig. 1). Before the measurement of venous susceptibility, a threshold of 90 parts per billion (ppb) was used to set the susceptibility of brain tissues surrounding the cerebral veins in the VOIs to 0, to avoid their influence on the measurement of susceptibility of cerebral major veins in the VOIs. Cerebral microbleeds and other small vessels were also avoided when drawing the major veins. Mean and standard deviation of the susceptibility in regional cerebral veins were recorded. 

2.4. Statistical analysis

The statistical evaluation of data was performed using SPSS 19.0 (SPSS Inc. Chicago, Ill., USA) software. In order to explore the reproducibility of the QSM method, the agreement between the susceptibility of cerebral veins from the HCs in our study and that from normal people studied by Doshi et al. (2015) was calculated using Pearson's correlation and inter-class correlation (ICC) was also calculated in mTBI patients and HCs. The normal distribution of data was explored by the Kolmogorov-Smirnov test. The difference of age and gender among three groups was explored using ANOVA and the Chi-square test. Due to the non-normal distribution of the susceptibility of left septal vein in HCs, the difference in susceptibility of cerebral veins among three groups was evaluated using non-parametric test. Similarly, due to the non-normal distribution of ETPT and RPQ scores between AI and non-AI group was explored using the Mann-Whitney U test and the two independent sample t-test, respectively. The correlations between the susceptibility of cerebral veins in the non-AI group and RPQ scores, between the susceptibility of cerebral veins in the AI group and either ETPT or RPQ scores were calculated using Pearson’s correlation analysis. The correlations between the susceptibility of cerebral veins in non-AI group and ETPT were calculated using Spearman's correlation analysis. The diagnostic efficiency of susceptibility to discriminate mTBI patients and HCs was evaluated using the receiver operating characteristic curve (ROC). For the multiple comparison of susceptibility in cerebral major veins among non-AI patients, AI-patients and HCs, a Bonferroni-corrected $P < 0.0045 (0.05/11)$ was considered as significant. The false discovery rate (FDR) correction was applied for the correlations between the susceptibility of cerebral veins from the HCs in our study and that from normal people studied by Doshi et al.; the correlations between the susceptibility of cerebral veins in AI and non-AI group and either ETPT or RPQ scores, an FDR-corrected $P < 0.05$ was considered significant.

3. Results

3.1. Demographic and clinical characteristic of subjects

The difference in demographic and RPQ scores, etiologies of the AI and non-AI patients and HCs are shown in Table 1. There was no significant difference in age and gender among the three groups ($P = 0.172, P = 0.848$, respectively). The mean ETPT in the AI and non-AI group was 110.13 ± 105.93 h and 51.03 ± 39.26 h, respectively, while there was no significant difference in the ETPT between the two mTBI groups ($P = 0.071$). No significant difference in RPQ scores between the AI (9.06 ± 5.58) and non-AI (10.06 ± 6.69) group was found ($P = 0.588$).

3.2. Reproducibility of the QSM method

In order to explore the reproducibility of the QSM method, we compared our results to those reported in Doshi et al. (2015). We found the susceptibility of major cerebral veins from the HCs in our study was highly consistent with that from normal controls reported in their study (Pearson's correlation; $r = 0.914, P = 0.011$; FDR corrected) (Fig. 2). We also calculated the ICC in mTBI patients and HCs between two measurements. The ICC values for right cerebral cortical vein, left cerebral cortical vein, right thalamostriate vein, left thalamostriate vein, right septal vein, left septal vein, right internal cerebral vein, left internal cerebral vein, right basal vein, left basal vein and straight sinus
in mTBI patients were 0.97, 0.98, 0.98, 0.97, 0.99, 0.99, 0.99, 0.95, 0.96, 0.99, respectively and the ICC values for above these veins in HCs were 0.96, 0.95, 0.97, 0.96, 0.98, 0.99, 0.99, 0.98, 0.98, 0.99, respectively, demonstrating good reproducibility of the analysis method.

3.3. Analyses of the susceptibility among the three groups

The comparison of susceptibility of major cerebral veins among the three groups using non-parametric test is presented in Table 2. There was a significant difference in the susceptibility of the straight sinus among the three groups (P < 0.001, Bonferroni corrected). Specifically, the susceptibility of the straight sinus in non-AI (256 ± 35, P < 0.001, Bonferroni corrected) and AI (265 ± 61, P = 0.004, Bonferroni corrected) group was significantly lower than that in the healthy control group (311 ± 43) (Fig. 3). But there was no significant difference in the susceptibility of straight sinus between the two mTBI groups (P = 1.000).

3.4. Diagnostic efficiency of susceptibility of straight sinus for discrimination between patients and HCs

Due to a significant difference in the susceptibility of the straight sinus among the three groups, the sensitivity, specificity, and area under the curve (AUC) of susceptibility of the straight sinus for the discrimination between mTBI patients, non-AI patients, AI patients and HCs using the receiver operating characteristic curve (ROC) are shown in Fig. 4. The best cut-off value for the discrimination between mTBI patients and HCs was 298 ppb with a sensitivity of 88% (95% CI: 75–95), specificity 69% (95% CI: 50–84), and AUC value 0.84 (95% CI:...
The best cut-off value for the discrimination between non-AI patients and HCs was 273 ppb with the sensitivity 75% (95% CI: 57–89), specificity 84% (95% CI: 67–95), and AUC of 0.86 (95% CI: 0.75–0.93) (Fig. 4b). The best cut-off value for the discrimination between AI patients and HCs was 290 ppb with sensitivity 81% (95% CI: 54–96), specificity 69% (95% CI: 50–84), and AUC of 0.80 (95% CI: 0.66–0.90) (Fig. 4c). We also calculated the sensitivity, specificity, and AUC of susceptibility of the other veins for the discrimination between mTBI patients, non-AI patients, AI patients and HCs using the ROC (Supplementary Table 1, Supplementary Figs. 1–10).

| Table 1 | The difference in demographic and post-concussive symptom scores among non-AI patients, AI patients and HCs. |
|------------------|--------------------------------------------------|
| **Clinical parameters** | **Non-AI patients (n = 32)** | **AI patients (n = 16)** | **HCs (n = 32)** | **P value** |
| Age (year) | 37.03 ± 10.65 | 32.06 ± 10.00 | 37.78 ± 9.81 | 0.172* |
| Gender (male/female) | 22/10 | 10/6 | 20/12 | 0.848 |
| ETPT (h) | 51.03 ± 39.26 | 110.13 ± 105.93 | – | – |
| RPQ scores | 10.06 ± 6.69 | 9.06 ± 5.58 | – | – |
| The etiologies of patients with mTBI (n = 48) | | | | |
| Assault (n = 15) | 9 | 6 | – | – |
| Struck by vehicle (n = 24) | 19 | 5 | – | – |
| Falling (n = 6) | 3 | 3 | – | – |
| Blunt trauma from falling objects (n = 2) | 1 | 1 | – | – |
| Blast injury (n = 1) | 0 | 1 | – | – |

* mTBI = mild traumatic brain injury, AI = axonal injury, RPQ = Rivermead Post-Concussion Symptoms Questionnaire.

** Fig. 2. ** The correlation between susceptibility of cerebral veins in our study and that in the Doshi et al.’s study.

The susceptibility of each major cerebral vein including left septal vein, right septal vein, left thalamostriate vein, right thalamostriate vein, left basal vein and right basal vein from HCs measured in our study were highly consistent with the measured susceptibility values from normal controls reported by the study of Doshi et al.’s by Pearson’s correlation (r = 0.914, P = 0.011, FDR corrected) (the error bars represent standard deviation of measured susceptibility of cerebral veins).

0.74–0.91 (Fig. 4a). The best cut-off value for the discrimination between non-AI patients and HCs was 273 ppb with the sensitivity 75% (95% CI: 57–89), specificity 84% (95% CI: 67–95), and AUC value 0.86 (95% CI: 0.75–0.93) (Fig. 4b). The best cut-off value for the discrimination between AI patients and HCs was 290 ppb with sensitivity 81% (95% CI: 54–96), specificity 69% (95% CI: 50–84), and AUC of 0.80 (95% CI: 0.66–0.90) (Fig. 4c). We also calculated the sensitivity, specificity, and AUC of susceptibility of the other veins for the discrimination between mTBI patients, non-AI patients, AI patients and HCs using the ROC (Supplementary Table 1, Supplementary Figs. 1–10).

| Table 2 | The difference in susceptibility (ppb) of each cerebral vein among non-AI, AI patients and HCs. |
|------------------|--------------------------------------------------|
| **Cerebral veins** | **Non-AI patients (n = 32)** | **AI patients (n = 16)** | **HCs (n = 32)** | **P value** |
| Right cortical vein | 164.782 ± 8.55 | 160.768 ± 7.78 | 163.057 ± 7.39 | 0.257 |
| Left cortical vein | 165.496 ± 10.09 | 162.911 ± 13.11 | 160.367 ± 8.10 | 0.131 |
| Right thalamostriate vein | 163.104 ± 13.01 | 164.702 ± 13.16 | 164.374 ± 13.87 | 0.900 |
| Left thalamostriate vein | 161.797 ± 12.35 | 159.770 ± 13.09 | 164.650 ± 11.60 | 0.391 |
| Right septal vein | 151.096 ± 15.09 | 146.878 ± 22.27 | 151.327 ± 14.98 | 0.65 |
| Left septal vein | 151.199 ± 12.59 | 157.667 ± 37.23 | 154.088 ± 20.42 | 0.636 |
| Right cerebral internal vein | 199.521 ± 28.39 | 195.364 ± 36.10 | 189.564 ± 28.45 | 0.418 |
| Left cerebral internal vein | 199.775 ± 31.45 | 195.383 ± 28.24 | 195.374 ± 28.45 | 0.847 |
| Right basal vein | 165.936 ± 7.42 | 170.132 ± 8.99 | 169.794 ± 8.78 | 0.117 |
| Left basal vein | 169.614 ± 10.93 | 165.559 ± 8.16 | 170.977 ± 8.73 | 0.185 |
| Straight sinus | 255.514 ± 35.25 | 264.840 ± 60.80 | 310.634 ± 43.05 | < 0.001* |

* Stand for P < 0.0045, Bonferroni corrected.
3.5. Correlation analyses with ETPT and post-concussive symptom scores

In the non-AI group, a positive correlation between the susceptibility of the straight sinus and the ETPT was observed ($r = 0.573$, $P = 0.003$, FDR corrected), as shown in Fig. 5. No significant correlation was found in the AI group ($P = 0.992$, FDR corrected). On the other hand, in the AI group, a negative correlation between the susceptibility of the straight sinus and the RPQ scores was observed ($r = -0.582$, $P = 0.018$, FDR corrected), as shown in Fig. 6, but no significant correlation was found in the non-AI group ($P = 0.886$, FDR corrected). There were no correlations between the susceptibility of other cerebral veins (the right and left cortical veins; septal veins; thalamostriate veins; internal cerebral veins; basal veins) and ETPT or RPQ scores in the AI or the non-AI groups ($P > 0.05$).

4. Discussion

There were three important findings in our study. First, the susceptibility of the straight sinus in non-AI and AI patients was significantly lower than that in HCs indicating an increased regional cerebral SvO2 in patients after mTBI. The best cut-off values for the discrimination between mTBI patients, non-AI patients, AI patients and HCs were 298 ppb, 273 ppb and 290 ppb. Second, as the ETPT increased, the susceptibility of straight sinus in non-AI patients was increased, indicating that the regional cerebral SvO2 tended to decrease to normal levels. Third, the RPQ scores increased with the decreasing susceptibility of the straight sinus, suggesting an increase in cerebral SvO2 due to severe post-concussive symptoms. The lower relative susceptibility in the straight sinus suggesting a higher level of oxygenated blood is consistent with the findings in Doshi et al. (2015). In that study, the differences between the subgroups (non-AI patient group and AI patient group) of mTBI patients are not considered. However, the abnormalities such as moderate bleeds, small bleed, nonspecific hyperintensities, arachnoid cyst, and pericallosal lipoma in the mTBI patients of Doshi et al.’s study may affect the susceptibility of the veins. In addition, the healthy controls (HCs) in their study had some abnormalities, such as capillary telangiectasia, pineal gland cyst, and nonspecific hyperintensity, which may also affect their results. Besides, the sample size of their study was relatively small (14 patients). These may explain the difference between the findings obtained in our study and Doshi et al.’s study. Nonetheless, the susceptibility of cerebral veins of HCs measured in our study are in good agreement with those reported by Doshi et al. (2015), indicating good reproducibility for using QSM to...
evaluate the cerebral SvO₂.

One reason for lower relative susceptibility may be from a decrease in cerebral metabolic rate of oxygen consumption in mTBI patients, which has been suggested in some earlier papers (Chen et al., 2004; Harris et al., 2012; Hovda et al., 1991). During the acute and sub-acute stages, the cerebral tissues might use another energy source, such as lactate, instead of glucose in some extreme conditions including trauma and starvation (Prins et al., 2004). One possible mechanism is mitochondrial dysfunction post TBI leading to a lower oxygen consumption.

In our study, we only found significant difference in the susceptibility of the straight sinus among three groups. There are a few possible reasons for this. First, the size of straight sinus was larger than that of the other cerebral veins and the susceptibility quantification of the other cerebral veins may be affected by partial volume effects because of the limited resolution in the through-plane direction. Hence, it was easier to highlight the significant susceptibility difference due to the reduced partial volume effects. In addition, the partial volume effects are also dependent on the orientation of the vessel. However, for veins in similar orientations, the estimate of the relative change in susceptibilities of these veins should not be affected much, as similar level of underestimations in susceptibilities are caused by partial volume effects. Nonetheless, in McDaniel et al. (2017), a promising technique was proposed, in which the partial volume effects can be compensated through a joint reconstruction using both magnitude and phase images for vessels in certain orientations. This method provides more accurate estimation of the oxygen saturation of the vessels. Second, the cerebral region drained by the straight sinus was larger than that drained by other small cerebral veins, and hence, the change of susceptibility due to abnormal metabolism may be more obvious. It has been reported that straight sinus mostly receives the deep cerebral venous system draining deep gray matter, the inferior sagittal sinus and some regions of the upper brainstem through the confluence of the inferior sagittal sinus and great vein of Galen (Jain et al., 2013; Krishnamurthy et al., 2014). The great vein of Galen receives bilateral cerebral internal, basal, thalamostriate and septal veins (Pearl et al., 2011). The straight sinus also drains the deep brain structures including thalamus, cerebral nuclei, midbrain, pontine nuclei and parahippocampal gyrus. These brain structures stand for more conserved parts of the brain and are responsible for basic activities to sustain life such as the consciousness, respiratory and cardiovascular systems (Jain et al., 2013). This may also explain the more significant changes of the susceptibility in the straight sinus than the other veins.

No significant difference in the susceptibility of the straight sinus between the AI and non-AI patient groups was observed in this study. This may be due to the different ETPT between AI and non-AI patients. Even though severe cerebral tissue damage could cause reduction in SvO₂ with an accompanying decrease in arterial blood supply (Shen et al., 2007), the time at which this change is measured is critical. Since the ETPT for the non-AI group was 51 h and was much shorter than the 110 h in the AI group, the different imaging time points of this cross-sectional study may assess different stages of injury and recovery in each group, making a direct comparison of SvO₂ challenging. This recovery is not unexpected (LosoiH et al., 2016). While the susceptibility of the straight sinus correlated with ETPT in the non-AI group, this was not observed in the AI group. This distinction may reflect the different pathological mechanisms in these two patient groups. The cerebral tissue damage in AI patients was more severe than that in non-AI patients (Cunningham et al., 2005; Martin et al., 1997). For the AI patients, the more extensive lesions in white matter tracts over widespread areas, such as brainstem, corpus callosum, basal ganglia and cerebral hemispheres, and the cerebral function may influence the rate of SvO₂ recovery (Cunningham et al., 2005; Martin et al., 1997).

It is also found that the susceptibility of the straight sinus in AI patients negatively correlated with RPQ scores, which may indicate severe post-concussive symptoms. The RPQ is a commonly implemented neurophysiological test used to evaluate the severity of clinical symptoms after mild or moderate TBI (King et al., 1995). The RPQ has been considered as a unitary construct with no further demand to analyze its underlying structures (Potter et al., 2006) and has shown reliability and validity in the studies (King et al., 1999; King, 1996). The increased cerebral SvO₂ may indicate reduced consumption of oxygen due to severe post-concussive symptoms which could be reflected by headache, dizziness, restlessness etc.

However, in non-AI patients, we did not find significant correlation between the susceptibility of straight sinus and RPQ scores, suggesting that the normal oxygen utilization is a key factor in maintaining brain function. There are a few possible reasons for this. First, the AI patients had more severe traumatic brain injury than the non-AI patients, which can cause obvious post-concussive symptoms. Second, the increased cerebral SvO₂ of non-AI patients can return to the normal level with ETPT, indicating that the post-concussive symptoms can be alleviated in non-AI patients. Although we found no significant difference in RPQ scores between AI and non-AI group, the RPQ scores of non-AI patients were slightly higher (10.06 ± 6.69) than that of AI patients (9.06 ± 5.58).

There are a few limitations in our study. First, for generating QSM data, the background field removal was achieved by using homodyne high-pass filtering. It is known that high-pass filtering leads to underestimation of the susceptibilities of structures with relatively low spatial frequency, such as the basal ganglia structures. This is the main reason of the reduction of the gray/white matter contrast in Fig. 1. Using more advanced background field removal algorithms such as Sophisticated Harmonic Artifact Reduction for Phase data (SHARP) (Schwezer et al., 2011) and Projection onto Dipole Fields (PDF) (Liu et al., 2011), the accuracy of susceptibility quantification can be further improved. Second, the number of patients included in this study is relatively small. More patients, especially AI patients, should be evaluated to observe the difference in susceptibility, ETPT and RPQ scores. Third, our study design was cross-sectional. A longitudinal study should be performed to explore the change of relative susceptibility in cerebral major veins and the correlation between the relative susceptibility of cerebral major veins, ETPT and RPQ test. Fourth, we only measured the Hct in patient groups but not in healthy controls. Thus, the quantification and direct comparison of venous oxygen saturation between mTBI patients and HCs was not available. With Hct known, the venous oxygen saturation can be calculated using the following equation (Ahman et al., 2013): $\Delta X_{\text{vein-tissue}} = \Delta X \times \text{Hct} \times (1 - Y_i)$, where $\Delta X_{\text{do}}$ is the difference in susceptibility per unit hematocrit between totally oxygenated blood and totally deoxygenated blood ($\Delta X_{\text{do}} = 4\pi \times 0.27 \text{ ppm}$) (He and Yablonskiy, 2009; Jain et al., 2010). Hematocrit (Hct) is defined as fractional hematocrit value in the major draining veins, which is approximately 40% (Haacke et al., 1997). $Y_i$ is the cerebral SvO₂ and $\Delta X_{\text{vein-tissue}}$ is the susceptibility of the cerebral veins. For the non-AI patients and AI patients, the cerebral SvO₂ of straight sinus were calculated as 81.10 ± 2.8% and 79.58 ± 4.1%, respectively. Assuming Hct = 0.4 for female and Hct = 0.42 for male (Haacke et al., 1997; Fan et al., 2012; Jain et al., 2010), the cerebral SvO₂ of straight sinus in HCs was estimated as 78.66 ± 3.5%. We found there were significant differences in cerebral SvO₂ of straight sinus among the non-AI patients, AI patients and HCs using one-way ANOVA ($P = 0.017$, Bonferroni corrected). Especially for the non-AI patients, the cerebral SvO₂ of straight sinus was significant higher than that in the HCs ($P = 0.014$, Bonferroni corrected). Finally, we did not measure cerebral perfusion. The increase in local perfusion may be another reason for increased cerebral SvO₂. The increase in local perfusion may exceed the oxygen demand of the traumatic cerebral tissues, indicating the neuroprotective response following mTBI (Doshi et al., 2015). In the future, we will detect the changes of cerebral perfusion to decide whether the increased local perfusion was one reason for increased regional cerebral SvO₂ in mTBI patients.
5. Conclusion
The decreased susceptibility of the straight sinus appears to correlate with post-concussive symptoms. In the case of the non-AI group, the decreased susceptibility of the straight sinus correlated with EPTP, indicating a recovery to normal levels of oxygenation over time. The susceptibility of the straight sinus provides an important biomarker for monitoring the progress of mild TBI and to differentiate mTBI patients from healthy controls.

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Conflicts of interest
The authors disclose no conflicts.

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