Longitudinal quantification and visualization of intracerebral haemorrhage using multimodal magnetic resonance and diffusion tensor imaging

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Abstract

Objective: To demonstrate a set of approaches using diffusion tensor imaging (DTI) tractography whereby pathology-affected white matter (WM) fibres in patients with intracerebral haemorrhage (ICH) can be selectively visualized.

Methods: Using structural neuroimaging and DTI volumes acquired longitudinally from three representative patients with ICH, the spatial configuration of ICH-related trauma is delineated and the WM fibre bundles intersecting each ICH lesion are identified and visualized. Both the extent of ICH lesions as well as the proportion of WM fibres intersecting the ICH pathology are quantified and compared across subjects.

Results: This method successfully demonstrates longitudinal volumetric differences in ICH lesion load and differences across time in the percentage of fibres which intersect the primary injury.

Conclusions: Because neurological conditions such as intracerebral haemorrhage (ICH) frequently exhibit pathology-related effects which lead to the exertion of mechanical pressure upon surrounding tissues and, thereby, to the deformation and/or displacement of WM fibres, DTI fibre tractography is highly suitable for assessing longitudinal changes in WM fibre integrity and mechanical displacement.

Keywords

Diffusion tensor imaging, intracerebral haemorrhage, longitudinal study, magnetic resonance imaging

Introduction

Intracerebral haemorrhage (ICH) is a neurological condition whose occurrence can have profound implications for patient health and recovery. Spontaneous ICH is typically caused by hypertension, which can produce micro-aneurysms at the bifurcation of arterioles [1]. In clinical settings, ICH is frequently associated with poor patient outcome, with over 70% of patients experiencing residual motor deficits [2, 3]. Approximately one third of patients with ICH who receive a computed tomography (CT) scan within 3 hours of injury onset exhibit haematoma expansion on follow-up CT scans [4], suggesting that neuroimaging is critical to the assessment of injury progression and to the determination of treatment steps necessary for optimal recovery. While some studies based on CT neuroimaging have demonstrated the predictive correlation between haemorrhage volume and patient mortality [5], morbidity in the context of motor outcome is not consistently reflected by CT measurements due to the fact that many neural fibres involved in motor control are confined within relatively small areas and are, thus, susceptible to mechanical pressure [2]. In fact, it has been shown that pressure applied to white matter (WM) may cause fibre deformations and lead to significant loss of neuronal transmission, to demyelination and to axonal shearing [6]. In the representative case of the corticospinal tract (CST), even a small volumetric extent of ICH may have severe debilitating effects [2]. Based on this knowledge, methodologies which can reveal the integrity of peri-lesional WM tracts can be beneficial in establishing a more comprehensive view of injury dynamics.

Whereas conventional structural magnetic resonance imaging (MRI) sequences such as fluid attenuated inversion recovery (FLAIR) and gradient recalled echo (GRE) imaging can reveal important structural information including the presence of oedema and haemorrhage, the integrity of WM tracts is often better observed using diffusion tensor imaging (DTI) [7–13]. Many investigators have utilized DTI to extract quantitative metrics of ICH which are predictively correlated
with motor outcome [2, 14, 15]. In addition, DTI tractography allows for the three-dimensional reconstruction of WM tracts, which greatly enhances the visual and quantitative exploration of fibre integrity. In the case of the CST, Jung and Jang [14] have investigated the relationships between the location and severity of injury along this tract, on the one hand, and motor outcome, on the other hand. The combination of tractography and 3D anatomical models derived from structural imaging volumes can be used to assess longitudinal changes in WM tracts, particularly from the standpoint of how affected fibres are displaced and may subsequently recover over time. By means of freely-available neuroimage analysis software packages such as 3D Slicer (slicer.org), TrackVis (trackvis.org) and the LONI Pipeline (pipeline.loni.usc.edu), this case study evaluation demonstrates the versatility of 3D ICH modelling and DTI fibre tractography in the context of longitudinal visualization and quantification of ICH. The usefulness of the paradigm and its applicability to the clinical assessment of this condition are illustrated in three representative clinical cases of ICH.

Methods

Patients and image acquisition

Multimodal neuroimaging volumes were acquired from three patients with spontaneous ICH at the Ronald Reagan Medical Center of the University of California, LA (UCLA). Informed written consent was provided either by the subjects themselves or by their authorized legal representatives and neuroimage volume acquisition was conducted with the approval of the Institutional Review Board of the School of Medicine at UCLA. Patients 1, 2 and 3 (aged 63, 56 and 44, respectively) were admitted to the neurointensive care unit (NICU) with Glasgow Coma Scores (GCS) of 14, 6 and 7, respectively, and discharged with GCS scores of 11, 7 and 6, respectively. All three patients received an initial MRI scan within 12 hours of admission. Patient 1 underwent thrombolytic therapy and was discharged with GCS scores of 14, 6 and 7, respectively, and was admitted to the neurointensive care unit (NICU) with Glasgow Coma Scores (GCS) of 14, 6 and 7, respectively, and discharged with GCS scores of 11, 7 and 6, respectively. All three patients received an initial MRI scan within 12 hours of admission. Patient 1 underwent thrombolytic therapy and received a follow-up scan 15 days after the injury date. Patient 2 underwent endoscopic evacuation of the haemorrhage and received a follow-up scan 15 days after the injury date. Patient 3 underwent endoscopic evacuation of the haemorrhage and received a follow-up scan 3 days after the injury. Patient 3 underwent endoscopic evacuation and received three additional scans: the first 2 days after injury, the second one 7 days after injury and the third one 12 days after injury. A total of 13 healthy adults (six females, aged 39.77 ± 15.13 years (mean ± standard deviation)) were also included in the study so as to statistically compare the quantitative metrics obtained from patients with ICH to those of a normative sample.

MRI volumes were acquired at 3.0 T using a Trio TIM scanner (1 mm³ voxel size, Siemens Corp., Erlangen, Germany). The acquisition protocol consisted of magnetization prepared rapid acquisition gradient echo (MP-RAGE) T₁-weighted imaging, fluid attenuated inversion recovery (FLAIR), turbo spin echo (TSE) T₂-weighted imaging, gradient recalled echo (GRE) T₂-weighted imaging and DTI. For the latter, volumes with 21 diffusion gradient directions were acquired for Patients 1 and 2 and with 64 directions for Patient 3. A similar acquisition protocol was used for the healthy control subjects as well. For patients with ICH, conventional CT scans were also acquired.

Image processing

Prior to any analysis, all MRI and DTI volumes were co-registered. Image processing was performed using the LONI Pipeline environment (pipeline.loni.usc.edu), including bias field correction, skull stripping and volume co-registration. Haemorrhagic tissues were segmented from GRE volumes and oedematous tissues were segmented from T₂ and FLAIR volumes. The procedure for pathology identification is described in detail elsewhere by Irimia et al. [16]. Briefly, non-haemorrhagic oedema was identified from T₂-weighted GRE imaging and FLAIR, whereas large haemorrhagic lesions and micro-haemorrhages were identified from susceptibility-weighted imaging (SWI). Diffuse axonal injury (DAI) was found to be apparent in DTI volumes. This protocol for identifying TBI-related pathology as well as details on its validation is described more extensively in a previous publication [16]. 3D Slicer software was used to generate 3D models and visualizations of pathology. To co-register WM surface models to pathology models, FreeSurfer was utilized to segment healthy-appearing WM, grey matter (GM) and cerebrospinal fluid (CSF) from T₁-weighted volumes using methodologies described elsewhere [17]. TrackVis and Diffusion Toolkit were used to reconstruct fibre tracts from DTI volume via deterministic tractography. Specifically, a brain mask was first created using FSL [18] to eliminate extracerebral noise. TrackVis was then used to reconstruct and render fibre tracts, which were subsequently loaded and viewed in 3D Slicer. In each subject, fibre tracts which did not intersect pathology-affected regions were discarded.

To reconstruct the CST, seed regions were placed in the brain stem and internal capsule and the WM tracts intersecting these regions were isolated.

The mean fractional anisotropy (FA) of each ICH patient was compared to the distribution of mean FA values in the sample of healthy control subjects as follows. First, for both patients with ICH and healthy subjects, the mean FA over the brain stem portion of the CST was calculated as

$$\text{mean FA}_{\text{CST}} = \frac{\text{sum of FA in CST}}{\text{length of CST}}$$

where

$$\frac{\text{sum of FA in CST}}{\text{length of CST}}$$

and

$$\text{length of CST}$$

are the volumes of the ICH lesion at times i and i + 1, respectively. To quantify the extent to which fibres were affected by pathology, the sum over the lengths of fibres which intersected the pathology was divided by the sum of the lengths of fibres in the whole brain, thus yielding the percentage of fibres in the brain which intersected the primary injury.

To infer whether and to what extent the mean FA in the brain stem portion of the CST were significantly different in
each ICH patient compared to the normative sample of healthy adults, the Z score of each patient’s mean FA with respect to the reference sample was computed at each time point. The statistical significance of the difference in mean FA values between every patient at each time point and the control sample was then quantified under the null hypothesis that no difference in this measure existed between the control sample and each ICH patient. \( p \) values were calculated based on the assumption that the computed Z score followed a standard-normal distribution with zero mean and unit variance. Both Z scores and \( p \) values are reported.

Results

Initial and follow-up MRI scans are displayed in Figures 1 and 2, with the time of the scan indicated at the top of the Figure. Figure 3 displays representative slices acquired from conventional CT to additionally illustrate the extent of the lesions present in each subject. In Figures 4 and 5, respectively, the results of the pathology segmentation at the initial and subsequent time points are displayed. WM models were created for each subject and displayed to provide an anatomical reference. Oedema is shown in cyan and blood is shown in red. Columns A and B display the WM fibres which intersect pathology-affected regions. For each subject, an enlarged, representative view is displayed in column C for closer inspection.

For Patient 1, hyper-intensities in the initial FLAIR scan indicate the presence of oedema in the tissue surrounding the haemorrhage and around the anterior horn of the right lateral ventricle. The GRE and \( T_2 \) sequences reveal the haemorrhagic lesion load better than the \( T_1 \)-weighted scan. The right ventricle appears comparably smaller than the left one, suggesting that the haematoma is displacing surrounding tissue and that it is thereby exerting pressure upon the ventricle. This effect is more obvious in the trigone of the lateral ventricle. Assessment of the CSTs reveal that the fibres in the right CST at the level of the internal capsule are somewhat displaced by the oedema towards the midline, a finding which is consistent with previous descriptions. The 2-week follow-up scans indicate significant resolution of the haemorrhage, with persisting—although reduced—oedema. The left and right ventricles appear more similar in size with respect to the acute scan and the right CST exhibits notable recovery in terms of its displacement.

The initial scans for patient 2 indicate severe haemorrhage in the left hemisphere. The level at which the axial slices are displayed also indicate the presence of a significant mid-line

![Figure 1](image1.png)
shift at the level of the thalamus. As observed in the previous case, the MR volumes indicate a reduction in ventricle volume. Whereas the right CST (Figure 6) appears healthy, the left CST is directly impacted by the haematoma and could not, for this reason, be reconstructed up to the primary somatosensory cortex due to increased diffusion isotropy within the haemorrhagic region. In contrast, DTI tractography based on scans acquired at follow-up demonstrates successful reconstruction of the left CST. Information provided by the GRE sequence indicates a reduction in haematoma size.

Patient 3 suffered a haemorrhage in the insula of the left hemisphere. No significant mid-line shift is observed, in contrast to the previous subjects. The subsequent scans indicate reduction in haematoma volume, with oedema persisting around the lesion. Reconstruction of the CST suggests the presence of mostly healthy-appearing fibres.

Table I displays the results of volumetric and WM fibre-related quantifications for each patient. The time of the scan is indicated at the top of each column. Patient 1 was found to have a total lesion load of 62.4 cm³ at the acute time point. On
Figure 4. The intersection between WM fibre tracts and the ICH pathology at the first time point is displayed in addition to 3D models of the pathology. Columns (A) and (B) display the left and right hemispheres, respectively. Column (C) displays an enlarged, representative view of the intersection between fibres and pathology-affected regions.

Figure 5. Visualizations of fibre tracts intersecting the ICH pathology at the second time point.
follow-up, this volume was reduced to 34.2 cm$^3$, with a decrease in both haemorrhage and oedema. The percentage of haemorrhage resolved was calculated to be 67.68%. The percentage of fibres affected was initially 22.53%, decreasing to 18.05% on follow-up. Patient 2 had an initial total lesion load of 88.51 cm$^3$, which decreased to 73.66 cm$^3$, with 64.7% of haemorrhage resolving between the first and second time points. The percentage of fibres affected was 27.32% initially, but increased to 36.53%. Finally, for Patient 3, lesion load was tracked across four time points. The initial volumes of haemorrhagic and oedematous tissue were 39.83 cm$^3$ and 10.59 cm$^3$, respectively. Whereas the size of the haemorrhage decreased to 16.03 cm$^3$ by the time of the final scan, oedematous tissue increased in volume to 86.04 cm$^3$, resulting in a total lesion load of 102.07 cm$^3$ by the 12th day after injury. The percentage of fibres affected by pathology decreased from 6.4% to 5.57% from the first to the third time point. Because DTI volumes were not collected on either Day 2 or Day 12 for clinically-related reasons, data for these time points are unavailable. Inspection of Table I also indicates that the mean FA in the mid-brain portion of the CST was significantly lower in each ICH patient and at every time point after injury compared to the reference sample of healthy control subjects.

**Discussion**

This study demonstrates an approach in which perilesional fibres can be selectively extracted and visualized based on MRI and DTI scans in patients with ICH. Given that mechanical pressure due to mass effects can alter the anatomical paths of WM fibres, the ability to track pathology-affected fibres over the evolution of injury is relevant to the study of ICH. The modelling of haemorrhagic and oedematous pathology provides useful knowledge on injury location with respect to brain tissue landmarks. This is relevant when considering that extravasated blood components released after ICH impose cytotoxic, pro-oxidative and pro-inflammatory effects on nearby viable brain cells [19]. In the case of oedema, coagulation enzymes such as thrombin, which is produced in response to haemorrhage, have been shown in both animal and human studies to induce a variety of negative effects, among which are oedema formation [1, 20]. Thus, the generation of 3D pathology models is informative when targeting affected regions for preventive treatment.

In all three cases analysed in this study, the amount of haemorrhage was observed to decrease between the initial and final scans. Conversely, cerebral oedema was observed to progressively increase for Patients 2 and 3, which may be a cause for the increase in percentage of affected fibres in the case of Patient 2. This finding, however, is consistent with secondary brain injury mechanisms, which may impose pathophysiological effects leading to increases in cytotoxic or vasogenic oedema [21, 22].

As previously discussed, the CST is frequently studied in ICH cases due to the critical function of the former in voluntary fine motor control [2, 14]. For that reason, this framework includes the ability to model the CST explicitly, which provides further detail on how injury has affected the ICH patient. For example, several authors have extrapolated that rapid and good recovery after stroke is associated with the resolution of factors such as peri-lesional oedema or inflammation, whereas slow but good recovery is associated with brain plasticity [23–25]. Extending upon this finding, Kwon et al. [26] suggested that the former scenario can be attributed to the preservation of the CST over the course of injury, whereas the latter case can be attributed to recovery of the CST. The finding that every ICH patient had lower mean FA in the mid-brain portion of the CST at each time point after injury may suggest appreciable atrophy in the mid-brain section of this functionally-prominent bundle of WM fibres, which is consistent with previous findings of WM degeneration in patients with ICH [27].

The utility of the methodology presented in this study is not restricted to the ability to perform visual observations. By placing fiducial markers in key regions of interest, it is also possible to use the approach proposed here to quantify fibre displacement across various time points. The challenge, however, would be to first devise a method whereby the same fibre tracts located in some given anatomical region can be reliably identified across time points. Since tractography is

Figure 6. The CST is explicitly modelled and displayed simultaneously with 3D models of ICH pathology. Columns (A) and (B) correspond to the acute and chronic scans, respectively.
Table I. Quantification of ICH evolution. The post-injury time of each scan is indicated in days (d) at the top of each column. Volumes are quoted in cm$^3$, with the percentage resolution of haemorrhage also being indicated at the bottom of the table. Blank entries indicate unavailability of the data (see text). The formula for the calculation of the percentage of fibres intersecting the ICH pathology is indicated in the Methods section.

<table>
<thead>
<tr>
<th>Post-injury day</th>
<th>Patient 1</th>
<th>Patient 2</th>
<th>Patient 3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>3</td>
<td>15</td>
<td>1</td>
</tr>
<tr>
<td>Oedema volume (cm$^3$)</td>
<td>29.73</td>
<td>23.64</td>
<td>16.66</td>
</tr>
<tr>
<td>Haemorrhage volume (cm$^3$)</td>
<td>32.67</td>
<td>10.56</td>
<td>71.85</td>
</tr>
<tr>
<td>Lesion load (cm$^3$)</td>
<td>62.4</td>
<td>34.2</td>
<td>88.51</td>
</tr>
<tr>
<td>Midline shift (mm)</td>
<td>4.08</td>
<td>4.72</td>
<td>5.47</td>
</tr>
<tr>
<td>FA of mid-brain CST</td>
<td>$\mu$ 0.3796</td>
<td>0.3823</td>
<td>$\sigma$ 0.3921</td>
</tr>
<tr>
<td></td>
<td>$\sigma$ 0.1485</td>
<td>0.2013</td>
<td>$z$ -7.038</td>
</tr>
<tr>
<td></td>
<td>$p$ &lt;0.0001</td>
<td>&lt;0.0001</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Haemorrhage amount resolved (%)</td>
<td>— 67.68</td>
<td>—</td>
<td>64.7</td>
</tr>
<tr>
<td>WM fibres affected (%)</td>
<td>22.53</td>
<td>18.05</td>
<td>27.32</td>
</tr>
</tbody>
</table>

initiated from seed regions, the reconstructed fibres may vary depending upon scan quality or upon physiological factors such as water anisotropy.

A final consideration is the limitation of DTI metrics such as fractional anisotrophy (FA) to describe WM property dynamics in peri-lesional regions. Particularly, given some location affected by oedema immediately after injury, one might expect FA in such a region to decrease at first [28] and then gradually to increase as haematomata volume decreases. This is because the typical lesion pattern in ICH consists of a haemorrhagic core surrounded by non-haemorrhagic oedema. However, in these case studies, average FA measured in peri-lesional regions indicated insignificant changes between the acute phase and at follow-up. On one hand, this may suggest a lack of recovery. On the other hand, the case studies suggest that, when the amount of cerebral oedema increases subsequent to the acute stage of ICH, the longitudinal comparison of FA values measured at locations within peri-lesional tissue may not be indicative of recovery due to the poor predictability of lesion shape dynamics. Nevertheless, the use of DTI metrics is a potentially useful direction for future applications to the study of ICH.

**Conclusion**

By utilizing DTI in conjunction with conventional structural imaging sequences, it is possible to selectively visualize fibres affected by injury pathology and to quantify the effect of the latter upon the former. Traditional methods of placing seed regions of interest (ROI) in pathology-affected regions can limit the tractography-based reconstruction to fibres which only pass through the selected voxels in the ROI. Thus, one drawback of DTI tractography is that some given reconstructed fibre bundle may cover only a portion of the tract of interest, as in the case of ROIs anatomically located within the CST. On the other hand, the approach described here allows one to reconstruct entire fasciculi such as the CST and then to calculate fibres which intersect pathology-affected regions. This strategy is applicable to any WM structure. Furthermore, the results obtained from 3D ICH analysis can be helpful for assessing longitudinal changes in the locations of WM structures from the acute to the chronic stage.

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**Declaration of interest**

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