A Patient-Specific Segmentation Framework for Longitudinal MR Images of Traumatic Brain Injury

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ABSTRACT

Traumatic brain injury (TBI) is a major cause of death and disability worldwide. Robust, reproducible segmentations of MR images with TBI are crucial for quantitative analysis of recovery and treatment efficacy. However, this is a significant challenge due to severe changes caused by edema (swelling), bleeding, tissue deformation, skull fracture, and other effects related to head injury. In this paper, we introduce a multi-modal image segmentation framework for longitudinal TBI images. The framework is initialized through manual input of primary lesion sites at each time point, which are then refined by a joint approach composed of Bayesian segmentation and construction of a personalized atlas. The personalized atlas construction estimates the average of the posteriors of the Bayesian segmentation at each time point and warps the average back to each time point to provide the updated priors for Bayesian segmentation. The difference between our approach and segmenting longitudinal images independently is that we use the information from all time points to improve the segmentations. Given a manual initialization, our framework automatically segments healthy structures (white matter, grey matter, cerebrospinal fluid) as well as different lesions such as hemorrhagic lesions and edema. Our framework can handle different sets of modalities at each time point, which provides flexibility in analyzing clinical scans. We show results on three subjects with acute baseline scans and chronic follow-up scans. The results demonstrate that joint analysis of all the points yields improved segmentation compared to independent analysis of the two time points.

Keywords: Image segmentation, Atlas formation, Longitudinal analysis

1. INTRODUCTION

Traumatic brain injury (TBI) occurs when an external force traumatically injures the brain, typically due to car accidents, accidental falls, and wartime injuries. It is a major cause of death and disability worldwide, especially in children and young adults, and it affects 1.7 million Americans annually.\textsuperscript{1,2} Robust, reproducible segmentations of MR images with TBI are crucial for quantitative analysis of recovery and treatment efficacy. However, this is a significant challenge due to severe changes such as edema (swelling), bleeding, tissue deformation, skull fracture, and other effects related to head injury. Acute and chronic images of Subject I is shown in Fig. 1 where non-hemorrhagic lesions (edema / swelling) are shown as hyperintense regions in FLAIR while hemorrhagic lesions (bleeding) are shown as hypointense regions in T2 and GRE. Despite the clinical importance of quantifying changes in TBI patient images, few research has been done on segmentation of MR images of TBI patients. A previous work by Thatcher et al.\textsuperscript{3} used fuzzy C-means and/or k-nearest-neighbor (kNN) algorithms and manual classification to segment 3D MR images of TBI patients without a longitudinal component.

Many others have proposed automatic segmentation methods for 3D MR images. For segmenting normal brain MR images, van Leemput et al.\textsuperscript{4} and Zhang et al.\textsuperscript{5} proposed atlas based methods, while Tu et al. proposed
Brain tumor is an example of a pathology that have similar properties to TBI. For segmenting brain MR images with tumor, Prastawa et al. proposed methods based on outlier detection and subject specific modification of atlas priors. Clark et al. proposed a automatic tumor segmentation using knowledge-based techniques. Ho et al. proposed a level-set based tumor segmentation method. Menze et al. presented a generative model for brain tumor segmentation using multi-modal MR images. For brain MR images with TBI, automatic segmentation is difficult due to the variability of lesion types, shapes, and appearances. In this paper, we introduce a novel user-initialized multi-modal image segmentation framework for longitudinal (4D) MR images with TBI.

This paper is organized as follows: Section 2 describes a patient-specific segmentation framework for longitudinal MR images of TBI patients. The framework is composed of Bayesian segmentation with user initialization, and personalized atlas construction. Section 3 shows the results of the proposed method. We compare the result of temporally independent segmentation with our result. We present conclusions and potential future work related to our method in Section 4.

2. METHOD

2.1 Bayesian segmentation with user initialization

The segmentation framework for multi-modal MR images is depicted in Fig. 2. The framework is initialized through manual input of primary lesion sites and affine-registered atlas at each time point, which are then refined by a joint approach composed of Bayesian segmentation and construction of a personalized atlas.

Suppose the multi-modal images at time point $t$ are $X_t = x_{1,t}, \ldots, x_{N,t}$ with $D_t$ the number of channels, where we use mixtures of Gaussians to model the data following van Leemput et al. We segment the images by maximizing the log likelihood function:

$$\ln p(X_t | \alpha, \mu, \Sigma) = \sum_{i=1}^{N} \ln \left( \sum_{j=1}^{K_t} \alpha_{i,j,t} N(x_{i,t} | \mu_{j,t}, \Sigma_{j,t}) \right)$$

where $t \in \{1 \ldots T\}$ is the number of time points, $K_t$ is the number of classes at time point $t$, $N(x_{i,t} | \mu_{j}, \Sigma_{j})$ is the multivariate Gaussian distribution, $\alpha_{i,j,t} = p(z_{i,t} = j)$ is the prior, and $z_{i,t} \in \{1 \ldots K\}$ is the tissue class label at position $i$ and time point $t$. We use the Expectation-Maximization (EM) algorithm to maximize the log likelihood function.
Figure 2. Our semi-automatic segmentation framework for MR images with TBI.

- E step: update the expectation weights $p(z_{i,t} = j | x_{i,t}, \mu_{j,t}, \Sigma_{j,t})$ at each location $i$ of each time point $t$.
- M step: update the Gaussian parameters $\mu_{j,t}, \Sigma_{j,t}$ at time point $t$.

We use user input and an affine-registered atlas to initialize the parameters $\mu_{j,t}, \Sigma_{j,t},$ and $\alpha_{j,t}$. The user input are spheres $S_t$ indicating lesions and the number of lesion classes $L_t$ at each time point $t$. We use the K-means algorithm\textsuperscript{14} to get initial estimates $\mu_{j,t}$ and $\Sigma_{j,t}$ for each lesion class. For other tissue classes (white matter (WM): $k = 1$, gray matter (GM): $k = 2$, cerebrospinal fluid (CSF): $k = 3$ and background (BG): $k = 4$), we use the affine-registered atlas (masked by user input $S_t$) to estimate $\mu_{j,t}$ and $\Sigma_{j,t}$. The initial priors $\alpha_{j,t}$ of each class are obtained by modifying the standard atlas using $S_t$, following Prastawa et al.\textsuperscript{7} We assume that lesions are found in WM and GM regions, so the initial prior for each lesion class becomes $\alpha_{j,t} = w(\alpha_{1,t} + \alpha_{2,t}) + S_t$, for $j \geq 5$, where $w$ is a uniform weight for lesions chosen to be 0.001 in our calculation. The $\alpha_{j,t}$ of other classes are linearly transformed to ensure that $\sum_{j=1}^{K} \alpha_{j,t} = 1$ at each location. The initial priors of one subject are shown in Fig. 3. The results of the EM algorithm are the posteriors $P_t$ which are $p(z_{i,t} = j | x_{i,t}, \mu_{j,t}, \Sigma_{j,t})$ at each location $i$ of each time point $t$.

One advantage of our method is that the number of channels $D_t$ at different time point $t$ can vary, which allows us to use different sets of modalities at each time point, a practice which can occur in clinical scanning, and thus provides flexibility for handling clinical diagnosis.

2.2 Construction of a personalized atlas

We use the segmentation results (the posteriors) to create personalized atlases using the unbiased diffeomorphic atlas construction method.\textsuperscript{15} Here we assume that there is no topological changes between the images of different time points. The flowchart of construction of personalized atlas is depicted in Fig. 4. In atlas construction, we estimate an average $\bar{P}$ (set of probability density functions / PDF) that requires the minimum amount of deformation $h_t$ to transform into the posteriors $P_t$ at every time point $t$, specifically:

$$\{h_t, \bar{P}\} = \arg\min_{h_t \in \mathcal{G}, \bar{P}} \sum_t \|P_t \circ h_t - \bar{P}\|^2 + D(e, h_t)^2$$  \hspace{1cm} (2)$$

where $D(e, h_t)$ is the distance of deformations $h_t$ to the identity transform $e$. We use these average PDF $\bar{P}$ as tissue prior probability maps in subsequent segmentations to get more consistent results by combining information from all time points.
The number of classes $K_t$ at different time points in longitudinal TBI images typically varies because bleeding or edema can disappear in follow-up scans. Following, we address this problem by combining the posteriors of lesions ($j \geq 5$) at a specific time point $t$. The combined lesion class formed by $p(z_{i,t} = j | x_{i,t}, \mu_j, \Sigma_j) = \sum_{j=5}^{K_t} p(z_{i,t} = j | x_{i,t}, \mu_j, \Sigma_j)$ follows our observation that both hemorrhagic and non-hemorrhagic lesions in preceding scans can still be observed in subsequent scans, though they may change appearance due to recovery. Compared to deformable atlas building using the MR image intensities, the benefit of this approach is that intensity calibration is not needed.

Fig. 5 shows an example of a constructed personalized atlas.

### 3. RESULTS

We apply our framework to data sets of three patients with TBI, each with two time points (acute and chronic). The image data of each subject include T1, T2, FLAIR, and GRE modalities. We use manual segmentations by a human expert as ground truth. Also, we compare our result to supervised segmentation (i.e., independent segmentations at each time point). We compare our results and the supervised segmentation results to the ground truth using the Dice coefficient, which is a standard similarity index in the range of 0 to 1 to measure the volumetric overlap of two binary segmentations. Dice coefficient values comparing segmentation results to the ground truth are shown in Table 1. Please note that the Dice coefficients are relatively low due to the complex boundary shape and relatively random spatial distribution of lesions. The volume of lesions for each subject in the manual segmentation is shown in Table 2. Our framework generally performs better than independent segmentations, with the primary exception of subject III where there is almost no lesion in the chronic scans and the lesion volumes are very small. The visualization of deformation field via determinant of Jacobian and vector...
Figure 5. Constructed personalized atlas for Subject II, where the average PDF $\bar{P}$ is deformed to the space at each time point and functions as tissue prior maps.

<table>
<thead>
<tr>
<th>Lesion types</th>
<th>Dice values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ANHL</td>
</tr>
<tr>
<td>Independent analysis</td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>0.5080</td>
</tr>
<tr>
<td>II</td>
<td>0.2165</td>
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<tr>
<td>III</td>
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<tr>
<td>Joint analysis</td>
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<tr>
<td>II</td>
<td>0.4398</td>
</tr>
<tr>
<td>III</td>
<td>0.4768</td>
</tr>
</tbody>
</table>

Table 1. Dice values comparing automatic segmentation results against ground truth, using temporally independent segmentations and our approach. ANHL is acute non-hemorrhagic lesion, AHL is acute hemorrhagic lesion, CL is chronic lesion.

magnitude is shown in Fig. 6. Axial view of acute images of subject I and of the associated segmentation using our framework are presented in Fig. 7.

Figure 6. Visualization of the deformation field of Subject I via Jacobian determinant (a) and vector magnitude (b).
Table 2. Lesion volumes (in voxels) for each subject in the manual segmentations. ANHL is acute non-hemorrhagic lesion, AHL is acute hemorrhagic lesion, CL is chronic lesion.

<table>
<thead>
<tr>
<th>Subject</th>
<th>ANHL</th>
<th>AHL</th>
<th>CL</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>57297</td>
<td>24486</td>
<td>9678</td>
</tr>
<tr>
<td>II</td>
<td>57526</td>
<td>36820</td>
<td>8847</td>
</tr>
<tr>
<td>III</td>
<td>50151</td>
<td>18136</td>
<td>1060</td>
</tr>
</tbody>
</table>

Figure 7. Results of our method for the acute images of Subject I. NHL is non-hemorrhagic lesion, HL is hemorrhagic lesion.

4. CONCLUSIONS

In this paper, we presented a segmentation framework for longitudinal TBI images. The framework is initialized through manual input describing primary lesion sites at each time point, which are then refined by a joint approach composed of Bayesian segmentation and construction of a personalized atlas. Our proposed framework has the advantage of being able to deal with different sets of modalities at each time point. The proposed joint analysis of different time points yields improved results compared to independent analysis.

There are several limitations in our proposed method. One limitation of the proposed method is that we assume there are no topological changes in the images of different time points so that we can use diffeomorphic atlas construction method to build personalized atlas. However, for longitudinal images of TBI patients this assumption is not always true. One possible solution is to mask the lesion area before registration. Alternatively, registration methods which are robust to missing correspondence could be used. Moreover, the approach will have limited applicability when lesion volumes are very small. In this case, the proposed joint approach may not be able to segment and capture these structures.

In the future, we would like to model the topological changes and thus make the atlas construction robust to topological changes. We will explore potential application of our method to aid registration of longitudinal MR images, with TBI, such as the geometric metamorphosis method proposed by Niethammer et al. We intend to apply our method to quantify recovery from longitudinal brain MR images with TBI under different treatments, with the potential of determining effective treatment strategies in the future.

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REFERENCES


