**Multi-atlas and label fusion approach for patient-specific MRI based skull estimation**

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A Multi-Atlas and Label Fusion Approach for Patient-Specific MRI Based Skull Segmentation

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Multi-Atlas and Label Fusion Approach for Patient-Specific MRI Based Skull Estimation

Angel Torrado-Carvajal,1,2* Joaquin L. Herraiz,2,3 Juan A. Hernandez-Tamames,1,2 Raul San Jose-Estepar,2,4 Yigitcan Eryaman,2,3,5 Yves Rozenholc,6,7 Elfar Adalsteinsson,2,8,9,10 Lawrence L. Wald,5,9 and Norberto Malpica1,2

Purpose: MRI-based skull segmentation is a useful procedure for many imaging applications. This study describes a methodology for automatic segmentation of the complete skull from a single T1-weighted volume.

Methods: The skull is estimated using a multi-atlas segmentation approach. Using a whole head computed tomography (CT) scan database, the skull in a new MRI volume is detected by nonrigid image registration of the volume to every CT, and combination of the individual segmentations by label-fusion. We have compared Majority Voting, Simultaneous Truth and Performance Level Estimation (STAPLE), Shape Based Averaging (SBA), and the Selective and Iterative Method for Performance Level Estimation (SIMPLE) algorithms.

Results: The pipeline has been evaluated quantitatively using images from the Retrospective Image Registration Evaluation database (achieving an overlap of \(72.46 \pm 6.99\%\)), a clinical CT-MR dataset (maximum overlap of \(78.31 \pm 6.97\%\)), and a whole head CT-MRI pair (maximum overlap 78.68\%). A qualitative evaluation has also been performed on MRI acquisition of volunteers.

Conclusion: It is possible to automatically segment the complete skull from MRI data using a multi-atlas and label fusion approach. This will allow the creation of complete MRI-based tissue models that can be used in electromagnetic dosimetry applications and attenuation correction in PET/MR.

Key words: atlas-based; label fusion; MRI; skull segmentation; tissue models

INTRODUCTION

Skull segmentation from MRI data is receiving a lot of attention, as there are many applications in which a precise delineation of the skull is needed, in addition to soft tissues. Accurate construction of patient-specific tissue models for dosimetry applications in electromagnetics (EM) (1,2), medical radiation physics (3), or the use of tissue information for attenuation correction in positron emission tomography (PET)/MR (4–6) are three of the most important examples. Treating bone as soft tissue or ignoring it in those applications is known to cause a distorted and biased distribution in the final estimation maps: B1+ field, specific absorption rate (SAR), standardized uptake value (SUV) distribution.

Most common approaches to patient-specific model creation are based on a combination of MRI and computed tomography (CT) images of the subject (7). Figure 1 shows how the MRI allows to better differentiate between soft tissues and to establish their boundaries, while CT provides the bone tissue information.

The use of only MRI instead of MRI+CT has the advantage of reducing radiation dose to subjects, decreasing costs and acquisition time, while allowing detailed information of soft tissues. It will also allow more complete (bigger field of view with respect to CT) and repeated skull imaging of the patient. Nevertheless, MRI-based bone segmentation, specifically automatic segmentation of the skull, is a challenging task. On one hand, bone tissue and air both present low signal intensity on MR images, making it difficult to accurately delimit the bone boundaries. On the other hand, the high complexity of the skull anatomy, its fuzzy boundaries and missing edge features hinders the application of general-purpose segmentation methods.

Background

Several skull segmentation or estimation approaches from MRI images have been presented in the recent literature, based on image postprocessing techniques or on the acquisition of multiple or specific MRI sequences.

*Correspondence to: Angel Torrado-Carvajal, M.Sc. in Biomedical Engineering, Medical Image Analysis and Biometry Lab, Universidad Rey Juan Carlos, Mostoles, Madrid, Spain.
E-mail: angel.torrado@urjc.es

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Image Postprocessing

Mathematical morphology analyses and processes geometrical structures in a binary image by using a structuring element; the operations over the structuring element include intersection, union, inclusion, and complementation. These morphological operators can be used on graphs, surface meshes, solids, and many other spatial structures.

Several approaches have also used deformable models based on an initial surface of the skull, deformed subject to artificial forces derived from the desired features of the segmentation. The strength of these models arises from their ability to include high-level information with low-level local features.

In Akahn et al (8), the authors use T1-weighted and proton density images to segment the skull and other tissues with a hybrid algorithm that uses snakes, region growing, morphological operations, and thresholding. The method described in Dogdas et al (9) finds the inner and outer skull boundaries in T1-weighted images using thresholding and morphological operations, and mask the results with the scalp and brain segmentations to ensure closed and nonintersecting skull boundaries.

Rifai et al (10) proposed a three-dimensional (3D) method for segmenting bone regions in MRI volumes using deformable models and taking the partial volume effect into account. In Wang et al (11) the authors proposed the use of a CT database to create a reliable shape model used to locate the skull shape in MRI.

Most of these works are focused on the upper part of the head, as they only have to deal with tissues surrounding the brain. However, the inclusion of the jaw in the complete head models complicates the application of these methods due to its different signal in MRI. Only the methods based on deformable models have dealt with the complete skull segmentation approach.

Multiple MRI Sequences

The detection of signals from tissues and tissue components with very short T2s is now possible due to the development and implementation of the ultrashort echo time (UTE) sequences (12). These sequences allow detecting signal from previously unobservable tissues such as cortical bone, tendons, ligaments, and menisci. However, classification of the skull in MRI has often been a by-product of classification techniques designed to categorize brain tissue. Several approaches are now focused on these techniques, but need multiple samples for each voxel to determine the corresponding output.

Keereman et al (13) proposed the use of the transverse relaxation rate derived from UTE images to classify the voxels into skull, soft tissue, and air. In Berk et al (14), the authors used UTE triple-echo (UTILE) MRI sequence for bone detection and gradient echoes for Dixon water and fat separation.

These methods show a good accuracy for the skull segmentation task, but they do not guarantee continuous bounding contours which can be a problem for meshing and EM simulation. Additionally, they show limited accuracy of bone segmentation in the interface between bone and air. As each radial projection passes through the center of k-space, the signals are indeed heavily averaged resulting in a considerable reduction of motion artifacts. However, the radial approach translates into an increased blurring, hindering edge detection.

In Belardinelli et al (15), the authors used an approach based on neural networks to segment the skull and the brain from successive T1-weighted 2D images. Hsu et al (16) proposed a method consisting of acquiring T1-weighted, T2-weighted, two echoes from a UTE sequence, and fat and water images using a Dixon method, to classify tissues using fuzzy c-means clustering.

The use of these approaches increase the overall acquisition time. The oversampling of the center of k-space in radial imaging for the UTE sequence makes the acquisition last $\frac{3}{2}$ times longer than a conventional Cartesian acquisition for the same matrix size. The use of several MRI sequences increase even more the complete acquisition time, depending on the nature and the number of sequences to acquire. For this reason, skull segmentation using a unique T1-weighted MRI sequence is desirable, as they are routinely acquired in clinical settings.
Contribution

In this work, we propose a new approach for complete skull estimation based only on T1-weighted images of the human head. This work is the development and assessment of the idea presented in Torrado-Carvajal et al (17), as part of a complete pipeline for tissue segmentation based on MRI only (18).

We use the patient’s MRI T1-weighted images and register the volumes from a multi-atlas CT database. The final segmentation is then statistically estimated from the individual segmentations. This approach deals with the limitations of the previous approaches presented above, and offers a general approach to accurately segment the skull.

We have compared the results provided by four different label fusion methods: majority voting (MV) (19), simultaneous truth and performance level estimation (STAPLE) (20), shape-based averaging (SBA) (21), and selective and iterative method for performance level estimation (SIMPLE) (22).

In this study, we demonstrate that a statistical combination of the multi-atlas data provides highly accurate segmentations. The study is structured as follows: the implementation of the skull segmentation algorithm is detailed, including the description of the multi-atlas CT database, the registration process of the multi-atlas to the patient-specific MRI standard space, a description of the different label fusion techniques used in this work and an introduction of the MRI data sets used to obtain the quantitative measures and the experimental results in Section 3; Section 4 establishes a discussion about our work including its limitations; in Section 5 we draw conclusions and comment on future work.

METHODS

An overview of our multi-Atlas and label fusion skull segmentation pipeline is shown in Figure 2. In this multi-atlas based segmentation approach, the unknown ground-truth segmentation mask $L_G$ of the MRI volume $V_{MRI}$ is estimated as $L_{MRI}$ by registration of a set of $N$ CT volumes $V_n$ and propagating their corresponding segmentations $L_n$. The major steps of the pipeline are: (i) Multi-Atlas CT database. Generation of the multi-atlas CT database $A_n = (V_n, L_n)$, where $A_n$ denotes an element in the atlas, $V_n$ is a CT volume, and $L_n$ its corresponding segmentation label map. (ii) CT-MR intermodality registration. Registration of each atlas volume $V_n$ to the patient-specific target volume $V_{MRI}$, and propagation of each atlas label $L_n$ to the target volume standard space, obtaining a new $L'$ label set containing $N$ label maps $L'_n$. (iii) Label fusion. Combination of the propagated segmentations $L'_n$ to create an estimation $L_{MRI}$ of the ground truth $L_G$.

Step 1: Multi-Atlas CT Database

We use the CT volumes $V_n$ from the whole head CT-scan database for craniofacial reconstruction developed by Tilotta et al (23). The images were acquired from healthy volunteers using whole head Somatom Sensation 16 CT
scanners (Siemens, Erlangen) at the Ouest Parisien (Val d’Or, Saint-Cloud, France) Medical Imaging Center. Subjects were positioned supine. For the current work, we have used a subset of the database, consisting of 19 subjects with ages ranging from 20 to 65 years old.

The segmentation of the CT volumes \( L_n \) is performed by thresholding of the CT data. The Hounsfield scale allows easily differentiating bone from the rest of the tissues. Additionally, an expert radiologist corrected the segmentations to delete segmentation errors due to dental restorations artifacts. These facts ensure that the segmentations are close to the actual ground truth, making the possible intra-rater reliability of the CT segmentation close to zero.

**Step 2: CT-MRI Intermodality Registration**

Every CT volume \( V_n \) from the atlas \( A \) is registered to the MRI volume \( V_{MRI} \) and each atlas segmentation \( L_n \) is propagated to the unseen data standard space (MRI space). To do so, we first need to pre-align the atlas volumes with the MRI volume using an affine registration, and then refine the registration using a nonrigid transformation.

**Mutual Information**

As we are dealing with intersubject CT-MR intermodality registration, we need a measure of the mutual dependence between the images. Mutual information (MI) is a distance measure described in the field of information theory by Collignon et al (24) and Viola and Wells (25). MI is a measure of the mutual dependence of two random variables computed from the gray level joint density of the volumes. The MI of two volumes \( V_A \) and \( V_B \) can be defined as:

\[
MI(V_A, V_B) = \sum_{a \in A} \sum_{b \in B} p(a,b) \log \frac{p(a,b)}{p(a)p(b)}
\]

where \( p(a,b) \) is the joint density function of \( V_A \) and \( V_B \), and \( p(a) \) and \( p(b) \) are the marginal probability distribution functions of \( V_A \) and \( V_B \), respectively. MI is considered as one of the most accurate and robust distance metrics used in registration (26,27).

**Affine Registration**

As the new MRI volume \( V_{MRI} \) is acquired in a different modality than the volumes in the atlas \( A \), the affine registration step transforms the CT volumes \( V_n \) in the atlas to roughly align to \( V_{MRI} \) and prepare them for the nonrigid registration step. This registration can be performed by directly registering each CT volume \( V_n \) to the MRI volume \( V_{MRI} \), obtaining the corresponding transformation matrix \( T_n^{A \to C} \). However, an efficient way to make the pre-alignment of the atlas volumes can be performed as follows:

1. Create an affine registered atlas, \( A^{A \to C} \), where each CT volume \( V_m \ \forall m = 2, \ldots, N \) is affine registered to the volume \( V_1 \) used as reference. In this case \( A^{A \to C} = (V_n^{A \to C}, T_n^{A \to C}) \), where \( V_n^{A \to C} \) is the volume \( V_n \) transformed to the \( V_1 \) space, and \( T_n^{A \to C} \) the corresponding transformation matrix used to obtain this correspondence.

2. Affinely register the MRI volume \( V_{MRI} \) to the CT reference volume \( V_1 \), obtaining the corresponding transformation matrix \( T_1^{C \to MRI} \).

3. Concatenate the inverse transformation matrix \( T_1^{-1} \) with each individual transformation matrix \( T_n^{A \to C} \) of each atlas volume \( V_n^{A \to C} \) producing a final transformation matrix \( T_n^{A \to MRI} \) which is applied to obtain the transformed volumes \( V_n^{A \to MRI} \).

The transformation matrices \( T_n^{A} \) and the transformation matrices \( T_n^{A \to MRI} \) may differ slightly; however, they seem to be good enough to serve as initialization for the nonrigid registration step. This efficient pre-alignment reduces in a factor of \( N \) the computational time for the affine registration.

**Nonrigid Registration**

Once the Atlas volumes \( V_n \) are pre-aligned with the MRI volume \( V_{MRI} \), we need to refine the registration to maximize the similarity between each volume \( V_n \) and the volume \( V_{MRI} \). This step can be performed by directly nonrigid registering each CT volume \( V_n^{A \to MRI} \) to \( V_{MRI} \). In this step we obtain the corresponding transformation \( T_n^{A \to MRI} \) for each volume in the atlas \( A^{A \to MRI} \).

**Label Map Propagation**

The transformation matrices \( T_n^{A \to MRI} \) and the transformations \( T_n^{A \to MRI} \) are then concatenated to obtain the complete transformations \( T_n \). These \( T_n \) are used to directly propagate the segmentation label maps \( L_n \) to the MRI standard space as \( L_n^{MRI} \).

**Step 3: Label Fusion**

Once we have registered the \( N \) CT volumes \( V_n \) and propagated their corresponding segmentations \( L_n \) to the MRI standard space \( V_{MRI} \) of the new data, the final patient-specific skull segmentation \( L_{MRI}^{MRI} \) is estimated as a combination of all the segmentations, \( L_{MRI}^{MRI} \), by using label fusion techniques. Label fusion exploits multi-atlas segmentation as an inference algorithm based on a nonparametric probabilistic model (28); thus, this technique decides how to fuse the information from several registered label maps. In this work, we have compared four label fusion procedures: MV (19), STAPLE (20), SBA (21), and SIMPLE (22) algorithms. Details of every method can be found in the online Supporting Information.

**DATA SETS**

To evaluate the complete segmentation pipeline, and the result of applying the different label fusion methods, we need \( T1 \) input volumes with a known ground truth. We have used data from different sources.

**RIRE**

The Retrospective Image Registration Evaluation (RIRE) project was designed to compare retrospective CT-MR and PET-MR registration techniques (29). The database
contains several intra-patient brain CT, MR, and PET volumes acquired at Vanderbilt University Medical Center. These datasets are now available as open-access data (30). Volumes present several voxel sizes in x and y, and a space between slices of 3 or 4 mm.

As the RIRE project aims at assessing different registration methods, images from the dataset are not aligned (the “truth”) transforms were defined using a prospective, marker-based technique, but they remain unpublished). Thus, we have used the same registration methods as in our pipeline to register the CT images to their corresponding MR images.

Clinical CT-MRI Pairs
A retrospective dataset consisting of six patients of Ruber International Hospital in Madrid that had undergone neurosurgery were analyzed. Every patient had an MR and a CT volume. MR images had been acquired on a General Electric Signa HDxt 3.0 Tesla (T) MR scanner using the body coil for excitation and an eight-channel quadrature brain coil for reception. Subjects were positioned supine. Imaging was performed using an isotropic 3DT1w SPGR sequence with a repetition time (TR) = 10.8 ms, TE = 4.2 ms, inversion time (TI) = 0 ms, number of excitations (NEX) = 1, acquisition matrix = 256 × 192, resolution = 1 mm × 1 mm × 1 mm, flip angle = 20.

Low-dose CT images were acquired on a General Electric Lightspeed VCT scanner with matrix = 512 × 512, resolution = 0.56 × 0.56 mm, slice thickness = 1.25 mm, PITCH = 0.53 mm, acquisition angle = 0°, voltage = 120 kV, radiation intensity = 200 mA. The examination is performed with the subject in the dorsal decubitus position.

Head MR Images
The present study was approved by the Instituto Carlos III Ethics Board and informed consent was obtained from all subjects before recruitment. Our dataset includes a total of 12 healthy subjects (4 males/8 females) aged 22–57 participating in this study. We also acquired an MRI volume for one of the subjects in the original CT database.

Images of the head were acquired on a General Electric Signa HDxt 3.0T MR scanner using the body coil for excitation and an eight-channel quadrature brain coil for reception. Subjects were positioned supine. Imaging was performed using an isotropic 3DT1w SPGR sequence with a repetition time (TR) = 10.024 ms, TE = 4.56 ms, TI = 600 ms, NEX = 1, acquisition matrix = 288 × 288, resolution = 1 × 1 × 1 mm, flip angle = 12. All image datasets were pre-processed using 3D Slicer built-in modules. The preprocessing step included MRI bias correction (N4 ITK MRI bias correction).

IMPLEMENTATION DETAILS
The skull segmentation pipeline has been implemented as an extension of 3D Slicer (31), as shown in Figure 3. The 3D Slicer is a free and open source software platform for visualization and image analysis of medical data. The 3D Slicer platform leverages the benefits of different open-source libraries such as the Insight Registration and Segmentation Toolkit (ITK) and the Visualization Toolkit (VTK), and allows rapid prototyping and development of medical imaging tools and applications (32,33).

The registration steps of the pipeline are performed with the built-in registration module (BRAINSFit) (34), using Mattes Mutual Information with a b-spline transformation model. This step allows registering the atlas dataset to the new volume to apply on of the different label fusion techniques.

MV and SIMPLE have been implemented as embedded Python code in the extension, while we have used the open source implementations of STAPLE – available as part of the Computational Radiology Kit (http://www.nitrc.org/projects/cmrk/) (CRKit)– and SBA – available as part of the Computational Morphometry Toolkit (http://nitrc.org/projects/cmrk/) (CMTK) – for this first development. All the experiments presented in this study were performed over Ubuntu Precise (12.04.3 LTS) on an Intel(R) Core(TM) i7-2600 CPU @ 3.40GHz with 8GB RAM.

RESULTS
In this section, we describe the main quantitative and qualitative results obtained on the datasets.

Quantitative Results
We have used the RIRE and the clinical CT-MRI pairs as input datasets to quantitatively evaluate the performance of the skull estimation pipeline. These datasets contained 16 and 6 subjects, respectively, where we measured the overlap between the ground truth and our estimation by using the Dice coefficient (35).

The Dice coefficient quantifies the similarity between regions by quantifying the spatial overlap. The Dice coefficient is computed as shown in Eq. [2]. Compared with Euclidean distance, the Dice coefficient gives more weight to voxels where the two images classification agree, and retains sensitivity in more heterogeneous data sets by giving less weight to outliers. Its values range between 0 (no overlap) and 1 (perfect agreement).

\[
D = \frac{2(A \cap B)}{(A \cap B) + (A \cup B)}
\]  

[2]

Figure 4A shows the performance of the different label fusion methods on the RIRE dataset. The Dice coefficient between the ground truth and the automated segmentation presents an overlap of 37.66 ± 7.54% for MV and SIMPLE, 71.79 ± 7.28% for STAPLE, and 72.46 ± 6.99% for SBA. Due to the low resolution in the z axis of the volumes in the RIRE database, the registration of some of the volumes in the database is not correct. Majority voting and SIMPLE are not able to discard these results, thus providing very bad segmentation results. STAPLE and SBA iteratively select the better registrations in the database, thus allowing to obtain a good final segmentation.
Figure 4B shows the performance of the different label fusion methods over the clinical CT-MRI dataset. The Dice coefficient between the ground truth and the automated segmentation presents a value of 48.79 ± 20.39% for MV, 78.31 ± 6.97% for STAPLE, 73.70 ± 7.24% for SBA, and 58.06 ± 7.80% for SIMPLE.

Figure 4C shows the performance of the different label fusion methods on the whole head CT-MRI pair. The Dice coefficient between the ground truth and the automated segmentation presents a value of 77.67% for MV, 78.68% for STAPLE, 77.22% for SBA, and 77.67% for SIMPLE.

FIG. 3. Graphical user interface of the skull segmenter extension. It allows loading a new MRI volume and estimating the skull by one of the four label fusion methods used in this study.

FIG. 4. Box and whisker plots of the Dice coefficient for the four estimation methods over the RIRE dataset (A), the clinical CT-MRI dataset (B), and the whole head CT-MRI pair (C). On each box, the central mark is the median, the edges of the box are the 25th and 75th percentiles, the whiskers extend to the most extreme data points [1.5 × inter-quartile range (IQR)], and outliers are plotted individually.
The DICE coefficient is a global measurement of segmentation quality. We have analyzed the spatial distribution of the fractional error by aligning all individual error masks to a template and computing the average error distribution, which is shown in Figure 5. Of the voxels showing some misclassification, only 2% showed a fractional error over 0.5 (the voxel was misclassified in more than half of the patients).

We have also measured the execution time for the three datasets. Table 1 shows the time taken by a complete execution of the registration and each label fusion approach on all datasets. The mean execution time and the 95% confidence interval are presented for each of the methods.

Qualitative Results

Visual inspection of the different approaches compared with the ground truth provides further assessment. Figure 6 shows the estimation result in five slices of a complete head MRI volume of a healthy subject. The method approaches the shape of the skull generally well. However, it can be seen that in some places, such as the frontal sinuses or cervical vertebrae, the estimation deviates from the ground truth skull contour.

Figure 7 shows the 3D rendering for 10 subjects. In this representation we can appreciate the level of detail of the approach.

The evaluation of our method on the RIRE dataset provides more information regarding the estimation of the skull on unhealthy subjects. Figure 8 shows the resulting mask in six pathological subjects from this dataset. The method is able to estimate the boundaries of the skull avoiding the pathological structures, even when they are close to the skull boundaries.

DISCUSSION

MRI-based bone segmentation, and particularly automatic segmentation of the skull, is a challenging task. Due to the nature of the tissue properties, bone and air present low signal intensity on MR images, making it difficult to accurately delimit the bone boundaries, while the high complexity of the skull anatomy, its fuzzy boundaries and missing edge features make it difficult to apply general purpose segmentation methods.

The estimation of the skull based only on the patient-specific MRI is feasible with a previous CT atlas dataset and label fusion techniques. The results show how the estimation of the skull adjusts to the bone boundary limits while differentiating the air. The strategy of this method is to leverage the benefits of “a priori” knowledge from the CT atlas dataset and label fusion techniques.

The proposed method is able to approximate the skull contours and differentiate air from bone with similar accuracy as the results presented in Wang et al (11), that reported a Dice coefficient of 0.75, even though they only showed results on the upper half of the skull. Additionally, their method requires previous annotation of the database by an expert, while our method requires no previous manual intervention.

As a result of the label fusion volume averaging, the contours of the skull are smoothed in several places. For example, our method fails to capture the details of the bone spikes in the area inside the sinuses, as seen in Figure 5. However, these errors are not crucial in most of the potential applications of our method. The skull estimation presented in this study has been successfully used to create complete patient-specific tissue models.

Table 1

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Label fusion technique</th>
<th>Registration time (minutes)</th>
<th>Label fusion time (seconds)</th>
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<tr>
<td>RIRE</td>
<td>MV</td>
<td>19.50 ± 4.57</td>
<td>0.47 ± 0.09</td>
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<td></td>
<td>STAPLE</td>
<td>11.86 ± 2.17</td>
<td>35.64 ± 5.16</td>
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<tr>
<td></td>
<td>SBA</td>
<td>15.50 ± 2.69</td>
<td>5.16</td>
</tr>
<tr>
<td>Clinical CT-MRI pairs</td>
<td>STAPLE</td>
<td>22.89 ± 1.58</td>
<td>53.73 ± 3.47</td>
</tr>
<tr>
<td></td>
<td>SBA</td>
<td>23.40 ± 1.47</td>
<td>21.56 ± 2.16</td>
</tr>
<tr>
<td></td>
<td>SIMPLE</td>
<td>21.71 ± 2.22</td>
<td>38.88 ± 3.16</td>
</tr>
<tr>
<td>Head</td>
<td>MV</td>
<td>21.71 ± 2.22</td>
<td>39.73 ± 1.29</td>
</tr>
<tr>
<td>CT-MRI pair</td>
<td>STAPLE</td>
<td>38.88 ± 3.16</td>
<td>39.73 ± 1.29</td>
</tr>
<tr>
<td></td>
<td>SBA</td>
<td>84.30 ± 5.15</td>
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<tr>
<td></td>
<td>SIMPLE</td>
<td>84.30 ± 5.15</td>
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*Registration time refers to the time taken in registering the whole dataset to the input image. We performed 20 executions for the CT-MRI pair to obtain the mean execution time, as this dataset only contained one subject.*
for EM modeling in EEG/MEG forward problem solving (36) and local SAR management (37,38), and PET/MR attenuation correction (39).

Our pipeline requires cross-modality registration, so that similarity based approaches to label-fusion are not directly applicable. Better results could be achieved...
using MRI-to-MRI registration, should we have MR-CT pairs for all subjects in the database.

Visual inspection of the segmentation results shows the high quality of the segmentations, and the robustness of the method in the presence of tumors and other brain pathologies. The use of this approach may lead to a decrease in patient ionization by removing the need of patient-specific CT acquisitions while obtaining a good estimation of the ground truth. These results are promising and may be included in several protocols such as brain studies in PET-MR scanners.

As can be seen in Table 1, the main drawback of the proposed method is the computational burden introduced by the multiple registrations and information fusion from the entire training data. Our approach provides good results, but computing many non-rigid
registrations is very time consuming. However, the gain to a patient by not performing a CT scan fully justifies the computational cost. Parallelization of the pipeline could reduce the overall time taken for skull estimation. Increasing the size and variety of the atlas could improve the accuracy of the results.

CONCLUSIONS

In this study, we have presented an approach for the estimation of the human skull by using a multi-atlas and label fusion approach based on acquired MR images only. The algorithm presented in this work removes the need of a patient-specific CT acquisition. The approach performs successfully on a wide range of data and could be useful for tasks where the skull estimation is needed such as PET/MI attenuation correction, and SAR calculations.

ACKNOWLEDGMENTS

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REFERENCES


**SUPPORTING INFORMATION**

Additional Supporting Information may be found in the online version of this article.