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SEMI-AUTOMATIC 3-D SEGMENTATION OF ANATOMICAL STRUCTURES OF BRAIN MRI VOLUMES USING GRAPH CUTS

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ABSTRACT
We present a semi-automatic segmentation technique of the anatomical structures of the brain: cerebrum, cerebellum, and brain stem. The method uses graph cuts segmentation with an anatomic template for initialization. First, a skull stripping procedure is applied to remove non-brain tissues. Then, the segmentation is done hierarchically by first, extracting first the cerebrum from the brain, and then from the remaining volume the cerebellum and the brain stem are separated. This method is fast and can separate different anatomical structures of the brain in spite of weak boundaries. We describe our approach and present experimental results demonstrating its usefulness.

Index Terms— Biomedical image processing, Image segmentation, Magnetic resonance

1. INTRODUCTION

The quantitative analysis of anatomical structures in MRI brain volumes is becoming a cornerstone in the study and detection of cerebral disease. In particular, volumetric quantification of cerebral and cerebellar tissues is essential in image-based assessment of neuroanatomical disorders such as autism and Asperger’s syndrome. The major difficulties of such studies are to overcome lack of boundaries, poor contrast, and noise, due primarily to the acquisition system and the partial volume effect. Therefore algorithms such as active contours [5, 4, 9] or region growing are subject to “leakage”, i.e., propagation beyond the true anatomical borders. Since a manual delineation of the anatomical structures would be too time consuming, several segmentation techniques have been developed to increase robustness: Active contours with shape model prior knowledge [7], atlas registration [8], and interactive graph cuts segmentation [1].

In the first case, a prior shape is incorporated into the active contour evolution in order to further constrain the segmentation. Shape priors can be modeled by a known class of shapes or through statistical training. So the choice of the models for the training determines the accuracy of the segmentation. The second method uses a combination of rigid and non-rigid transformation of an atlas to detect the internal structures in MR images of the brain. Although quite successful, this method requires high computational cost and a good atlas.

Finally, interactive graph cuts represents the volume as a discrete graph, composed of vertices representing the image voxels, as well as edges connecting the vertices, typically using a 6 or 26 neighborhood. The user marks certain voxels as object or background, defining the terminals of the graph, and the optimal segmentation is found using a max-flow / min-cut algorithm. The quality of the object extraction depends on the number of seeds used in the initialization. Indeed, in [1], the segmentation is refined by additional seeds, which the user adds. In this paper, we extend this method to fit into our specific problem, i.e., the segmentation of brain structures.

1.1. Our contribution

Despite their advantages, graph cuts segmentations can lead to erroneous results. In order to obtain a satisfactory segmentation, many seeds must be used to give a strong spatial constraint. In this paper, we propose a solution to avoid this time-consuming step. We use a template that gives, after a simple registration, a rough localization of the different structures (i.e. cerebrum, cerebellum, and brain stem). The registration steps only consist of a centroid alignment and a scaling of the template. The 3D template is shown in Figure 2(b), and once aligned, gives hard spatial constraints on the segmentation. One other way to give a good initialization is to start from an atlas. A similar approach would be to register a highly accurate and detailed atlas to the current volume, and then apply a strong erosion to ensure the seeds will be entirely inside each respective brain structures. Instead, we choose to use a simpler template to demonstrate the robustness of our method. Indeed, the initialization does not need to be very accurate to give a precise segmentation.

The second contribution is the hierarchical graph cuts segmentation. Indeed since the Boykov and Jolly graph cuts method is a binary segmentation, we proceed in three steps to separate each anatomical structure: brain/non-brain, cerebrum/cerebellum and brain stem, and cerebellum/brain stem.
2. SEGMENTATION APPROACH

We begin by briefly reviewing the graph theory that is used to minimize our energy function. We then describe how we build this function using the image data. We then list each step of our algorithm.

2.1. Graph cut theory

Following [2], consider an undirected graph \( G = (V, E) \) that is composed of vertices \( V \) and undirected edges \( E \) that connect the vertices. Each edge \( e \in E \) is assigned a non-negative cost. There are two special vertices (also called terminals) in the graph that are identified as the source \( s \) and the sink \( t \). With the exception of the terminals, the vertices are comprised of voxels \( P \) in the image. An example, displayed in 2D for simplicity, is shown in Figure 1. A cut \( C \) on the graph is a partition of \( V \) into two disjoint sets \( S \) and \( T = V - S \) such that \( s \in S \) and \( t \in T \), as shown in Figure 1(b). The cost of the cut is the sum of the costs of all edges that are severed by the cut. The minimum cut problem is to find the cut with the smallest cost energy:

\[
\text{Cost}(C) = \sum_{e_{i,j} \in C} w_{i,j} \quad i, j \in V \quad (1)
\]

where \( e_{i,j} \) represents the edge connecting the vertices \( i \) and \( j \) and \( w_{i,j} \) denotes the weight associated with the respective edge. There are numerous algorithms that solve this problem in polynomial time, see [2] for more details.

![Fig. 1. A simple 2D graph for a 3x3 image (a) and its minimal cut (b). Figure based on [1].](image)

2.2. Graph Cuts Image Segmentation

In the case of volume segmentation, the vertices will be the voxels \( P \) and two other nodes denoting the "object" \( O \) terminals and the "background" \( B \) terminals. Our goal then is to take a set of voxels \( P \) and compute a labeling that minimizes the cost of the cut. The idea behind the graph cut segmentation is to minimize an energy function defined on a graph, according to the cut of minimum weight. As presented in [1], the energy function can be written as:

\[
E = \sum_{p \in P} \left( D_p(f_p) + \sum_{q \in N(p)} V_{p,q}(f_p, f_q) \right) \quad (2)
\]

where \( E \) is the energy, \( p \) and \( q \) are voxels, \( N \) is the neighborhood formed from the vertex connectivity. \( D_p \) measures the cost of assigning the voxel \( p \) to the set \( f_p \), while \( V_{p,q} \) measures the cost of assigning the adjacent voxels \( p, q \) to the same set. As proved in [1], the segmentation defined by the minimum cost cut in Equation 1 also minimizes Equation 2. In our implementation,

\[
D_p(f_p) = \begin{cases} 
\text{MAX} & p \in O, f_p = S \\
\text{MAX} & p \in B, f_p = T \\
0 & \text{otherwise}
\end{cases}
\]

where \( \text{MAX} \) is a large positive constant, and

\[
V_{p,q} = \begin{cases} 
\exp\left(\frac{-(I_p - I_q)^2}{2\sigma^2}\right)/\text{dist}(p, q) & p, q \in N \\
0 & \text{otherwise}
\end{cases}
\]

where \( \text{dist}(p, q) \) is the Euclidean distance between voxels \( p \) and \( q \). The parameter \( \sigma \) is a constant that denotes the variance of the voxel value inside the object. In our experiments, we empirically find a fixed \( \sigma \) for each of the three different graph cuts (skull-stripping, cerebrum extraction, cerebellum and brain stem segmentation), and apply the same \( \sigma \)s for all data sets.

![Fig. 2. Initialization of the graph cuts segmentation for the non-brain tissue (a), and for the anatomical structures (b). In (b), the template consists of voxels in the cerebrum (red), cerebellum (blue), and brain stem (green).](image)

2.3. Implementation

First, we register the template to the current volume by aligning its centroid to the centroid of the volume we want to segment. We also scale the template by determining the size of the brain bounding box of the current volume. After the seeding steps, we start a three-phase segmentation process. In the first phase, we apply a skull stripping procedure, which isolates the brain from extracranial tissues to facilitate the subsequent segmentation. To achieve this, the user first brushes non-brain tissue as shown in Figure 2(a) as sink seeds, and source seeds are specified for all voxels in the template, shown in (b). Then we compute the graph over the volume data and its optimal cut.
In the second step, we extract the cerebrum from the brain by using the cerebrum seeds as the source of the graph cuts and the cerebellum and brain stem seeds as the sink. Once the cerebrum removed from the previous graph, we apply another graph cuts with only the brain stem terminals as the source and the cerebellum terminals as the sink to segment those two structures. The process is described in the Figure 3.

The complete algorithm then consists of the following steps:

1. Register the template to the MRI brain volume.
2. Brush non-brain tissue seeds, background included.
3. Apply a first graph cuts segmentation for the skull stripping, with all the template seeds as the source and the non-brain tissue seeds as the sink.
4. From the brain, run another graph cuts to extract the cerebrum. The cerebellum and brain stem seeds of the template are the sink of the graph cuts.
5. After extracting the cerebrum from the brain, apply another binary graph cuts to separate the cerebellum from the brain stem.

3. EXPERIMENTAL RESULTS

We apply our algorithm on T2 MRI axial volumes of brain with 256 * 256 slices. Two volumes are displayed in Figure 4. The whole segmentation (skull stripping included) completes in less than 50 sec on a Pentium 4 2.66 GHz processor. The seeds given by the template give a good prior information on the segmentation. In particular, it prevents any leaking even if the boundaries between the structures are not clear, as shown in Figure 5. In this slice, due to partial volume effect, the border between the cerebellum and the cerebrum is unclear, as shown in (a). Many active contour techniques will not produce accurate results at the region denoted with the white arrows.

This algorithm has been tested on six brain T2 datasets. All datasets have been successfully segmented. With the proposed initialization, the graph cut segmentation is able to retrieve the contour of the cerebellum, even when the boundary is unclear, as shown in Figure 5(b). We plan to perform more extensive testing on more data when it becomes available to fully validate the algorithm. However, our initial results demonstrate that the proposed method shows much promise.

4. CONCLUSION

This paper presented a method based on graph cuts to segment the cerebrum, the cerebellum, and the brain stem in a brain volume. We described our approach and experimentally demonstrated its usefulness.

In our experiments, we consider only a Gaussian function as a neighboring weight, with a unique constant parameter $\sigma$ for all the testing volume data. Any further work would include a computation of a specific sigma for each volume, depending on the voxels’ intensity distribution in order to improve the segmentation.

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6. REFERENCES


Fig. 4. Segmentation of two MRI brain volumes in different slices. The red, blue and green contour represent respectively the cerebrum, the cerebellum and the brain stem.

(a) (b)

Fig. 5. MRI Brain segmentation. The contour of the cerebellum is not clearly defined (a) but our algorithm is able to retrieve it (b).


