A New Algorithm for Fully Automatic Brain Tumor Segmentation with 3-D Convolutional Neural Networks

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Abstract

Glioblastoma (GBM) is the most common primary tumor of the central nervous system in the United States, with 3 in 100,000 people diagnosed every year [1]. In this paper, we present a new algorithm for fully automatic brain tumor segmentation based on 3-D Convolutional Neural Networks. Segmentation of the GBM region of the surrounding brain makes it easier to access the image data within MR scans, and consequently can help us better understand GBMs. Most methods for volumetric image data use 2-D convolutions; we present a true generalization of CNNs to 3-D filters to preserve spatial information and increase robustness. In addition, we present a relatively high bias CNN architecture that enables us to both expand the effective data size and reduce the variance of our model. Our median Dice score accuracy is around 89 percent in the whole tumor segmentation. This result represents a significant improvement over past algorithms and demonstrates the power of our approach in generalizing low-bias high-variance methods like CNNs to learn from medium-size data sets.

1. Introduction

Glioblastoma (GBM) is the most prevalent and most aggressive type of brain cancer [2]. As a result of its resistance to existing therapeutic methods, the most common length of survival following diagnosis is 12 to 15 months, and less than 3-5% of patients survive more than five years [1][3]. Much is still being done to understand GBM and its two subtypes, high grade glioma (HGG) and low grade glioma (LGG), but there is still a gold mine of untapped data. Chief amongst these is imaging data in the form of magnetic resonance (MR) scans. Most methods of analyzing and extracting quantitative information from this imaging data requires some form of segmentation of the GBM, and better yet, classification of the tumor into four sub-categories: necrosis, edema, non-enhancing tumor, and enhancing tumor. To date, the gold standard of segmentation is still human radiologist segmentation. However, with the pure size of imaging data being accrued, manual segmentation of all images is no longer a sustainable system.

Nonetheless, existing automated segmentation algorithms are based on high-bias learning algorithms and have failed to give substantial improvement in classification accuracy as described by Subbanna et al., Menze et al., and Bauer et al. [4]. As a result, these high-bias systems are increasingly being replaced with data-driven low-bias models. The development of low-bias systems such as convolutional neural networks for medical purposes is, nevertheless, challenging due to their high-variance, as many medical datasets are limited in size.

In this paper, we propose a novel approach to automatic brain tumor segmentation. We take as input a 3-D MR image and output one of five labels (normal or one of 4 tumor subtypes) for each voxel (the 3-D analog of pixel). Our new approach surpasses current methods by reducing the variance of CNNs applied to medium datasets. First, where most methods use 2-D convolutions for 3-D data, we introduce a CNN that uses 3-D filters, made computationally efficient by the transformation to Fourier space. Consequently, our system is more robust and minimizes loss of spatial information. Second, we present a novel CNN architecture which differs from those traditionally used in brain segmentation and computer vision [4][5]; our CNN architecture uses a higher-bias system than CNNs using pixel-wise neural networks that enables us to both expand the effective data size from patient number to total pixel and reduce the variance of CNNs. Our algorithm for GBM segmentation and characterization exploits the CNNs framework and the structure of medical imaging data to train a model that not only is highly robust but also generalizes data-driven low-bias systems to medium data while minimizing variance.
2. Problem Statement

Our problem amounts to efficiently classifying brain voxels into five categories: non-tumor, necrosis, edema, non-enhancing, and enhancing. The input to our system consists of a preprocessed 3-D image of a brain. Each 3-D image is of size 240 × 240 × 155 × 4, where the depth 4 represents the four modalities of our input images. The goal is to for each voxel correctly and efficiently output its category (non-tumor, necrosis, edema, non-enhancing, and enhancing).

2.1. Dataset

We use the data provided by the MICCAI Multimodal Brain Tumor Image Segmentation Challenge (BRATS) [4]. The 2015 BRATS dataset contains 274 patients: 220 patients with high grade glioblastomas and 54 patients with low grade glioblastomas. For each patient our data contains four MR modalities: T1 post-contrast, T1 pre-contrast, T2 weighted, and FLAIR. In addition, the data contains an expert segmentation for each patient that we will treat as the ground truth. The expert segmentation was made by pooling together the segmentation of eleven different radiologists and includes pixel-wise labeling as one of five categories: non-tumor, necrosis, edema, non-enhancing, enhancing.

All the data has been preprocessed, involving skull stripping, co-registration of the images, and interpolation of the images so that all images are of the same size (240 × 240 × 155 pixels). We were not given any information about the spatial scale of the pixels; and thus, all units for our current method will be in the given pixel-space.

3. Technical Approach

3.1. Convolutional Neural Network Motivation

Convolutional neural networks (CNNs) are powerful low-bias systems in deep learning with remarkable performance in complex tasks ranging from facial recognition to image classification [7][8]. CNNs use 2D convolutions as filters for feature learning and feed the results of these convolution layers into a fully connected neural network. Our algorithm builds on the idea of a CNN as a fully connected neural network trained on sequential convolution layers to improve accuracy and computational efficiency in learning from 3D imaging data. Specifically, we generalize the idea of 2D convolutions to 3D filters that retain the full spatial information of the original stack of images. We then decouple the pixels when providing the training data for a fully connected neural network, thus increasing the size of our training set and providing a computational speedup.

3.2. 3-D Convolutions

We can reduce the bias of our system by considering the input image as a 3-dimensional space of pixels. Expanding our convolution from our usual 2-dimensional convolutions into 3-dimensional convolutions, we get the following equation:

\[
(I * f)(x, y, z) = \sum_{\tau_x=1}^{240} \sum_{\tau_y=1}^{240} \sum_{\tau_z=1}^{155} I(\tau_x, \tau_y, \tau_z)f(x - \tau_x, y - \tau_y, z - \tau_z)
\]

where I is the 3-dimensional image and f is the 3-dimensional filter. Given an image of size \( n \times n \times n \) and a filter of size \( m \times m \times m \) the time complexity of the convolution operation is \( O(m^3n^3) \). We can reduce the time complexity of the convolution operation to \( O(n^3\log(n)) \) by implementing our convolution as an element-wise multiplication in Fourier space. We use a 3-dimensional Difference of Gaussian filter:

\[
D_{\sigma G}(\tau_x, \tau_y, \tau_z) = \frac{1}{(2\pi\sigma^2)^{3/2}} \exp \left( -\frac{\tau_x^2 + \tau_y^2 + \tau_z^2}{2\sigma^2} \right) - \frac{1}{(\sqrt{2}\sigma)^{3/2}} \exp \left( -\frac{\tau_x^2 + \tau_y^2 + \tau_z^2}{\sigma^2} \right)
\]

Each filter is a difference of two 3-dimensional Gaussians (normalized) with scales \( \sqrt{2}\sigma \) and \( \sigma \). We create 8 of these filters with the scales \([\sqrt{2}, 2, 2\sqrt{2}, ..., 16]\). Difference of
Figure 2: Visualization of 3-D Convolution. Similar to the 2-dimensional equivalent, in a 3-dimensional convolution, we create an output image from the dot-product of our 3-dimensional filter and a sub-image centered at every pixel. We represent the filter with the red-box, our image with blue box, and the convolutional product with our white box.

Gaussian is an effective blob detector since it is rotationally symmetric.

To create our final convolutional products, we will consider all four modalities, and the magnitude of the gradient of those images. In addition to the eight filter products, we will include the original pixel intensity and the magnitude of the gradient value. Thus, for each of the four modalities, we will get an expansion into 18 feature images, giving us a total feature space of 72-dimensions for each of our pixels. We choose a filter of size $33 \times 33 \times 33$. Basically, we get for each voxel $[x,y,z]$ the following set of features:

- 1 voxel intensity $V[x,y,z]$
- 1 voxel intensity gradient $\nabla V[x,y,z]$
- 8 DoG convolutions $(V * DoG)[x,y,z]$
- 8 DoG convolutions in gradient space $(\nabla V * DoG)[x,y,z]$

3.3. 3-D Convolutional Neural Network Architecture

Our input consists of a 3-dimensional image with four channels (one for each modality). Our first layer is a non-trained convolutional layer with $72 \times 240 \times 240 \times 155 \times 4$ filters, 3-dimensionally convolved over all channels with our input image. In this layer, our filters are the hand-selected Difference of Gaussian filters for a specified modality with all other values in the other three channels being zero. Our training does not go through this layer.

The following convolutional layers use convolutions of filters sized $1 \times 1 \times 1$ over the number of channels in the preceding layer (either 72 for the first layer or 100 for all other layers). The $1 \times 1 \times 1$ filter is a scalar that we will constantly multiply over the whole image. In a sense, by only wiring the convolution layers onward with these $1 \times 1 \times 1$ filters, we are decoupling all of the pixels in our image. Because the first convolution layer couples neighborhood information via the $33 \times 33 \times 33$ convolutions, all subsequent layers pixels will maintain some level of information of its original neighborhood. Thus, we are able to decrease the number of weights we would have to train while drastically increasing our effective training data size.

On the last layer, we will get five channels, each channel giving us a probability score for every pixel in our image along the five sub-regions (0 = non-tumor, 1 = necrosis, 2 = edema, 3 = non-enhancing, and 4 = enhancing). We use the softmax function as the loss function. To train the convolutional neural network we use stochastic gradient descent by patient. During test time, we classify using a voting metric that assigns a voxel to one of the five categories based on its classification scores (see algorithm 1). This method is equivalent to how the expert segmentations were pooled from 11 different radiologist’s segmentations.

Essentially, our classification priority in descending order goes: enhancing, necrosis, non-enhancing, and edema. If our neural nets returns positive classifications for multiple tumor subtypes, we classify to the positive subtype with the
highest priority. This hierarchical design is based off of the hierarchical majority vote used to combine several different algorithmic results [4].

This seemingly arbitrary methodology makes perfect sense in the context of our classification problem. Tumor segmentations are judged generally in terms of three accuracies: whole tumor accuracy, tumor core accuracy, and enhancing tumor accuracy. Thus, because they have their own accuracy scores, we must prioritize classification of the core over the non-core (edema), and then also the enhancing core over the other core. The enhancing core generally covers a smaller area of the brain, which lends even more reason to be more sensitive to its detection.

Results are reported as the standard Dice score calculated via 10-fold cross validation (see: beginning of next section). We do not use cross validation to select parameters, deciding to keep our neural net parameters set to default values. This is both because the additional computation time would be prohibitive, and also because our Dice scores (which are also calculated from the cross validation) would become biased upwards.

Algorithm 1 Hierarchical majority decision
Our neural network outputs a number between 0 and 1 per voxel for each of the four tumor structures (edema, non-enhancing, necrosis, enhancing), respectively indicated by \( p_{\text{edema}} \), \( p_{\text{nonenh}} \), \( p_{\text{necrosis}} \), \( p_{\text{enh}} \).

```plaintext
1: procedure HIERARCHICALDECISION(Score)
2:   label ← NormalTissue
3:   if \( p_{\text{edema}} \geq 0.5 \) then
4:     label ← Edema
5:   end if
6:   if \( p_{\text{nonenh}} \geq 0.5 \) then
7:     label ← Non − Enhancing
8:   end if
9:   if \( p_{\text{necrosis}} \geq 0.5 \) then
10:    label ← Necrosis
11:   end if
12:   if \( p_{\text{enh}} \geq 0.5 \) then
13:    label ← Enhancing
14:   end if
15: end procedure
```

4. Results
We use Dice Coefficient to compute our results (also known as the Sørensen-Dice Index) [9],

\[
\text{DiceScore} = \frac{2|P_\text{red} \cap R_\text{ef}|}{|P_\text{red}| + |R_\text{ef}|}
\]

where \( P_\text{red} \) is the set of voxels designated as the region of interest in the prediction and \( R_\text{ef} \) is the set of voxels designated as the region of interest in the reference for comparison (in our case, the expert segmentation).

The median Dice performance on the whole tumor detection is 89%! To wit, the inter-radiologist repeatability is only 85%, so our accuracy has saturated with respect to the ground truth. Figure 4 shows a comparison amongst the leading algorithms in the literature. The first row shows the performance of human radiologists. The full distribution of accuracy of the algorithm among the 274 patient is in figure 5. One particularly successful segmentation can be seen in figure 6.

5. Discussion
Our median performance on the whole tumor detection revolves around a dice score of 89, which is right before losing clinical significance. This is very competitive with previous methods. One clear issue with our algorithm is dealing with outliers as can be seen in figure 5. More than half of our segmentations are wildly successful, but some segmentations return sub-50% scores, which you would not typically see with a radiologist.
Figure 6: **Slices-by-Slices Result** A representative slice view of a test case giving 98% Dice score accuracy. Each of the 12 images has three parts. On the left is the T1 post-contrast image at the given slice, the middle image is the expert segmentation labeling, and the right-hand side image is the labeling at the slice by our algorithm. The label color scheme follows that which was introduced in figure 1.

Figure 4 shows a comparison amongst the top results in the literature. Some notable ones in 2013 include ones by Zhao and Subbanna which incorporated Markov Random Fields (MRF), achieving a Dice accuracy of 82, respectively [4]. Festa from 2013 used random forests to achieve a Dice of 62 [4]. In 2014, groups used deep learning and convolution neural nets (CNNs) to achieve accuracies of 85 (Davy) and 88 (Urban) [10]. Our method improves upon the existing vanilla convolutional neural network architecture by making some high biased assumptions that allow us to drastically reduce our variance.

Basically, one of our assumption is that voxels are independent, only coupled by information tied to their feature vectors. Certainly, this is a high-bias assumption, but it allows us to use \( n = 4.5 \) billion training examples rather than only \( n = 274 \) patient samples. Neural nets are ultimately low-bias systems, but our hope is that the improved variance caused by the enlarged sample space will overcompensate for our high-bias assumptions. Furthermore, we contrast our method with standard deep learning algorithms like CNN, which learn the features using convolution kernels on the inputs. Vanilla CNNs use each patient as a single training sample. We, on the other hand, select Difference of Gaussian (DoG) convolution filters to relate voxels to their neighborhoods. Although we are highly biasing our system compared to usual deep learning framework, our features may be more adequate since our subsequent feed-forward neural net can then learn higher level features from each pixel’s lower level features.

### 6. Conclusion

The higher-biased CNN 3-D architecture performs remarkably well on the glioblastoma segmentation problem. The segmentation results are competitive with those using much more complex methods, and we argue our success is due to our smart choice of features along with a greatly enlarged sample space and flexible training method (neural nets). Our algorithm is powerful despite its relatively high bias, and we hope that it may serve the medical community in their work.

The natural next step of this project is a thorough analysis of our models asymptotes. We have claimed that our large data set has significantly reduced model variance, but it is unknown whether we can further reduce variance with more data. Given that our segmentation algorithm is already on par with our reference expert segmentations, we suspect but would like to confirm that our model has already reached its large-data asymptotic performance.

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### References


