

# BrainParser: Deformable Templates Guided Discriminative Models for Robust 3D Brain MRI Segmentation

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**Abstract** Automatically segmenting anatomical structures from 3D brain MRI images is an important task in neuroimaging. One major challenge is to design and learn effective image models accounting for the large variability in anatomy and data acquisition protocols. A deformable template is a type of generative model that attempts to explicitly match an input image with a template (atlas), and thus, they are robust against global intensity changes. On the other hand, discriminative models combine local image features to capture complex image patterns. In this paper, we propose a robust brain image segmentation algorithm that fuses together deformable templates and informative features. It takes advantage of the adaptation capability of the generative model and the classification power of the discriminative models. The proposed algorithm achieves both robustness and efficiency, and can be used to segment brain MRI images with large anatomical variations. We perform an extensive experimental study on four datasets of T1-weighted brain MRI data from different sources (1,082 MRI scans in total) and observe consistent improvement over the state-of-the-art systems.

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## 1 Introduction

In neuroimaging studies, brain MRI segmentation is often a critical preprocessing step. Automated segmentation enables morphometric analysis of cortical and subcortical structures in large datasets [19], a scenario in which manual labeling is impractical. The automatically segmented regions can be used to extract informative characteristics of structures, such as volumes and shape. In the clinic, these features have the potential to be used to evaluate the condition of a subject. Moreover, the identified boundaries between cortical and subcortical structures can aid the planning of brain surgery [76]. In neuroscience research, statistics derived from the segmentations of control and experimental groups can be used to identify structural differences between them. In the context of disease studies, such differences can lead to the identification of new pathological biomarkers. For instance, the atrophy and morphological change of hippocampus have been identified as important markers for Alzheimer’s disease [31].

Several practical segmentation methods are widely used in neuroimaging studies, e.g. Caret [15], FreeSurfer [19], FSL [65], ITK-SNAP [88], SPM [21], and the segmentation utilities in 3D Slicer [51]. However, robustness against variations in the input imaging data remains an open problem. The main difficulties are due to: (1) variations in image intensities due to differences in MRI acquisition (hardware, pulse sequence, imaging parameters); (2) anatomical variations within and across populations. Intensity normalization and image registration [29,35] can be used to standardize the images prior to segmentation, but only to some extent, since many of the variations are intrinsic.

In this paper, we aim to build a robust system that automatically segments T1-weighted brain MRI volumes into anatomical sub-cortical and cortical structures. We approach the 3D brain image segmentation problem from a statistical modeling perspective, combining a generative and a discriminative model with feature augmentation and adaptation. Generative and discriminative models were first explored and compared in the machine learning and computer vision literatures [49,67]. It has been shown that, while generative models outperform discriminative models when the size of the training dataset is small, the latter often have a better asymptotic behavior [44]. Works that attempt to combine both types of model include [32,56,37,28], which show that integrating the two types of models can be beneficial.

Specifically, the goal of brain MRI segmentation is to separate the voxels of an input scan into a number of classes. In some studies, these classes correspond to the three basic tissue types in the brain: gray matter (GM), white matter (WM), and cerebrospinal fluid (CSF) [75,50,39,61,81,6,43]. Other works have attempted to produce labels at the level of brain structures (e.g.,

hippocampus, pallidum, putamen, etc.) [19,58,34,7], which is a more difficult problem but yields a richer description of the data.

To produce these labels, generative models typically rely on two components: a *prior* term that summarizes the statistical frequency and spatial distribution of labels and a *likelihood* term that models how these labels translate into intensities. Then, Bayesian inference can be used to estimate which labels (i.e., which segmentation) are the most likely given the observed image intensities. The prior term is usually in the form of a statistical atlas endowed with deformation model [22,39,19]. Other priors include the use of principal component analysis (PCA) to model whole shape variations [53,85] or Markov random fields (MRF) to impose local shape constraints [19,80,59,9]. For the likelihood term, Gaussian distributions or mixtures thereof have been predominant in the literature, due to their ease of inference [18,85,52,13]. Because the Gaussian parameters (means and variances) are estimated during the optimization, these algorithms are robust against changes in MRI contrast. Moreover, they can also explicitly model image artifacts such as the MRI bias field, making them robust against them as well (see for instance [39]).

Recently proposed multi-atlas methods such as [26,82,35,3,27,78,7,57] can also be seen as generative models. These methods are based on deforming a number of labeled templates to a test scan, and then fusing the deformed labels into a single, enhanced estimate of the segmentation. As explained in [57], these algorithms can be interpreted as generative models in which the intensity and label of each voxel are taken from one of the deformed templates as indexed by a latent, discrete field.

Discriminative models attempt to directly estimate the label of each voxel given the local appearance of the image around it. To do so, a number of features are computed for each voxel and fed to a classifier that attempts to infer the corresponding label. Popular choices of features include image intensities, gradients, textures and other derived local measures [69]. Choices of classifier range from simple rule-based classifiers [41] to more complicated frameworks such as support vector machines (SVM [38,36,1,89]), AdaBoost [55,48] and the increasingly popular random forests [8,87,25,86]. These techniques show promising results when segmenting tissues, structures, and even tumors ([5,42]) if the variations of the test data with respect to the training data are relatively small. Unfortunately, changes in MRI contrast due to differences in imaging hardware or acquisition protocol considerably reduce the performance of these methods, limiting their applicability to MRI scans that have not been acquired the same way as the training dataset.

Comparing generative and discriminative models, we observe that deformable templates [17,52] guided by generative models can efficiently model anatomical structures, thanks to their flexibility and adaptability. However, their simplified assumptions on underlying image statistics (e.g., Gaussian distributions) limit their ability to deal with complex intensity patterns. On the other hand, such patterns can be efficiently captured by discriminative models thanks to their capacity of fusing together a large number of features.

However, as described above, they are sensitive to global intensity changes and have difficulties to include spatial information.

In this paper, we propose combining the strengths of the two approaches by fusing deformable templates (generative) and informative features (discriminative). The presented approach is based on using the estimated segmentation and the parameters of the generative model to normalize the image intensities and extract robust, invariant local features. This creates an augmented feature space that can be used in a discriminative framework, effectively fusing the two types of model.

The rest of this paper is organized as follows. First, Section 2 surveys other attempts of combining generative and discriminative models in brain MRI segmentation. Section 3 describes the proposed segmentation framework. Section 4 describes how to train the model and use it to segment previously unseen test cases. A set of experiments and their corresponding results are described in Section 5. Finally, Section 6 concludes the paper.

## 2 Further related work

Here we discuss other works in the literature that are similar to the proposed approach, highlighting the differences between them:

- Verma et al. [73] use Bayesian and SVM models to classify the intra- and inter-patient tissue distributions. However, the two families of methods are separately used for the different sub-problems, rather than in an integrated fashion.
- In [69], a discriminative classifier is used for appearance, and its classification results are regularized by generative models (PCA) capturing the shape prior. The two types of models are represented as separate modules and not jointly considered. In this paper, we utilize the same discriminative model, whereas the generative models are totally different. As shown in the experiments later on in this paper, the proposed method outperforms this algorithm, which is not robust across datasets due to the inability of the discriminative model to adapt to the new data.
- Wels et al. [76] cast the subcortical segmentation as a posterior maximizing problem with shape parameters. The whole problem is decomposed into four sub-problems in sequence and each is solved by a discriminative model. Their segmentation then locates, rotates, scales, and refines the boundaries of structures in order, which performs a rough generative process. Nevertheless, the four underlying classifiers are purely discriminative so their weaknesses remain. If some stage fails, wrong information is propagated without any adaptation or correction. On the contrary, our discriminative model can benefit from the adaptation capability of generative models so the robustness is achieved.
- Wels et al. [77] also propose a hybrid method for tissue segmentation. However, their approach is sequential: the discriminative model serves for initializing and constraining the subsequent fitting of the generative model,

which is carried out with an expectation maximization algorithm. The approach is validated on 38 scans from the same dataset (Internet Brain Segmentation Repository) using only three classes (WM/GM/CSF). It is unclear how this method would generalize to a higher number of brain structures, since the application of discriminative models at the structure level is more challenging than at tissue class level (i.e., just three classes).

- Fan et al. [16] use a deformable template (generative) to estimate a rough region of each tissue, and a discriminative classifier (SVM) is trained on all tissue volumetric measures.
- Wang et al. [74] propose a two-stage classifier wrapper for adapting different labeling protocols. They use the second stage classifier to learn the systematic errors from the first stage. As in Fan et al., the classifier independently refines an initial solution provided by the generative model.

Compared with these approaches, the key aspect that differentiates our algorithm is the use of an adaptive, augmented feature space that allows us to effectively fuse generative and discriminative models (as described in Section 3.3), rather than simply cascading them.

### 3 Model description

In this section, a general formulation of the segmentation problem is provided. We first compare the generative and discriminative approach to the segmentation problem. The comparison reveals their complementary properties, and inspires our hybrid method. The idea is to first use a deformable template (the generative component, described in Section 3.2) that produces a first estimate of the segmentation for an input volume. This rough estimate is then used (as described in in Section 3.3) to: (1) calculate the regional statistics of the input volume for local (discriminative) feature normalization, and (2) derive local features such as region boundaries and label priors. These features, in combination with local appearance-based, discriminative features, give an augmented set that yields a rich description of the data by integrating information from the generative and discriminative sides. This augmented feature set will be used to train a classifier as described in Section 4.

#### 3.1 Problem formulation

Our goal is to segment a given 3D volume/image  $\mathbf{V}$  into  $K$  anatomical structures, where  $K$  is a fixed number. A training set of  $N$  volumes with their corresponding annotations (no less than  $K$  labels) is assumed to be available, and we denote this set as  $S = \{(\mathbf{V}_1^{tr}, A_1^{tr}), \dots, (\mathbf{V}_N^{tr}, A_N^{tr})\}$ , where  $\mathbf{V}_i^{tr}$  and  $A_i^{tr}$  are the  $i$ 'th training volume and its corresponding annotation. We represent each structure by a region  $R_i$ . A segmentation is denoted as

$$W = \{(R_i, \Theta_i), i = 0 \dots K\}, \quad (1)$$

where  $R_0$  refers to the background region, and each  $R_i$  consists of all the voxels of the  $i^{\text{th}}$  anatomical structure.

These regions are disjoint and they cover the entire volume:  $\bigcup_{i=0}^K R_i = \Lambda$ , where  $\Lambda$  defines the 3D lattice of the input  $\mathbf{V}$ , and  $R_i \cap R_j = \emptyset, \forall i \neq j$ .  $\Theta_i$  is a vector that includes the model parameters for the appearance and shape of region  $i$ .

If we define  $p(\mathbf{V}(R_i)|R_i, \Theta_i)$  as the likelihood of the volume confined in region  $R_i$  under model parameters  $\Theta_i$ , the optimal solution in a Bayesian framework can be obtained as:

$$\begin{aligned} W^* &= \underset{W}{\operatorname{argmax}} p(W|\mathbf{V}) = \underset{W}{\operatorname{argmax}} p(\mathbf{V}|W)p(W) \\ &= \underset{W}{\operatorname{argmax}} \prod_{i=0}^K p(\mathbf{V}(R_i)|R_i, \Theta_i)p(R_i)p(\Theta_i), \end{aligned} \quad (2)$$

where  $p(R_i)$  is the probability of the shape prior, whereas  $p(\Theta_i)$  puts a prior on the parameters and is usually assumed to be flat i.e.,  $p(\Theta_i) \propto 1$ . Moreover, we have assumed that the shapes of the different regions are independent, which allows us to write  $p(W) = \prod p(R_i)$ . Whereas a more faithful model would consider dependencies between the shapes of the structures, this common assumption greatly simplifies both the training of the model and the inference in the Bayesian framework. We further assume that the intensity inhomogeneity of a region is smooth and small enough [40].

In general, independent identical distribution (i.i.d.) assumptions are made in the likelihood function, and the appearance of each structure is approximated by a Gaussian model [18, 53]. Let  $G(\cdot; \Theta_i)$  denote a Gaussian distribution parameterized by  $\Theta_i$  and let  $v_j$  be the intensity value of voxel  $j$ . The likelihood then can be represented as:

$$p(\mathbf{V}(R_i)|R_i, \Theta_i) = \prod_{\forall j \in R_i} G(v_j; \Theta_i),$$

where  $\Theta_i$  contains the mean and standard deviation of region  $i$ :  $\Theta_i = \{\bar{v}_i, \sigma_i\}$ . As argued in [69], using only generative models with i.i.d. assumptions is often too simplistic to fully account for the realistic textures of MRI data. This insufficiency, especially between structural boundaries, will be addressed by the discriminative models in our method.

For a discriminative approach, there is no explicit data parameter estimated for each input volume  $\mathbf{V}$ ; the model is instead learned in the form of a classifier derived from a training dataset. Thus, the solution vector by a discriminative model becomes

$$W_R = \{R_i, i = 0 \dots K\}. \quad (3)$$

We use  $|\mathbf{V}|$  to represent the total number of voxels in  $\mathbf{V}$  and  $l_j$  to denote the label assigned to voxel  $j$ .  $R_i$  is therefore the set of all  $l_j = i$ . A discriminative classifier directly computes the class label at a voxel  $j$  which has the maximum

class posterior, which is based on the local features computed on the sub-volume  $\mathbf{V}(N_j)$  centered at  $j$ :

$$W_R^* \equiv (l_j^*, j = 1..|\mathbf{V}|) = \underset{W_R}{\operatorname{argmax}} \prod_{j=1}^{|\mathbf{V}|} p(l_j | \mathbf{V}(N_j)), \quad (4)$$

where the label of voxel  $j$  maximizing the equation is denoted as  $l_j^*$ .

If we compare the solution vectors  $W$  and  $W_R$  in Equations 2 (generative) and 4 (discriminative), we make two observations. First, generative models explicitly estimate the data parameters and thus are adaptive to the input. Second, discriminative models can efficiently capture complex local image statistics by combining many low- and mid-level features. As discussed above, generative models make simplistic assumptions for the likelihood term modeling the local appearance, whereas discriminative models struggle capturing the information from larger regions. Therefore, in this study we will use deformable templates guided by generative models (as described in Section 3.2) and seek to fuse them with the discriminative model in Equation 4 (as described in Section 3.3).

### 3.2 Deformable templates guided by generative models

For the generative model of image intensities we adopt a Gaussian mixture model due to its modeling capability [19, 85, 52] and relatively low computational complexity. Let  $\Theta = \{\Theta_i, i = 0, \dots, K\}$ , we introduce the weights of Gaussian components so the parameters for each region are given by:

$$\Theta_i = \{(\bar{v}_i^{(m)}, \sigma_i^{(m)}, \beta_i^{(m)}), m = 1, 2\},$$

where  $\bar{v}_i^{(m)}$ ,  $\sigma_i^{(m)}$ , and  $\beta_i^{(m)}$  are respectively the mean, standard deviation and weight of Gaussian component  $m$  of region  $R_i$ . In this paper, we assume two components  $m = 1, 2$  for each model, which is empirically sufficient to describe the appearance of the anatomical regions. The parameters of these Gaussian components are obtained with a expectation maximization (EM) algorithm [10]. According to this model, we have the following likelihood function of voxel  $j$  in region  $R_i$ :

$$p(v_j; \Theta_i) = \sum_{m=1}^2 \beta_i^{(m)} G(v_j; \bar{v}_i^{(m)}, \sigma_i^{(m)}), \quad (5)$$

Given a volume  $\mathbf{V}$ , we seek to minimize an energy function under the Bayesian formulation of Equation 2 with flat  $p(\Theta_i)$ :

$$\begin{aligned}
& E(W_R, \Theta, \mathbf{V}) \\
&= \sum_{i=0}^K -\log p(\mathbf{V}(R_i)|R_i, \Theta_i) - \log p(R_i) - \log p(\Theta_i) \\
&\approx \sum_{i=0}^K \sum_{j \in R_i} \left\{ -\log p(v_j; \Theta_i) + \kappa \sum_{j' \in N_j} \delta(l(j') \neq l(j)) \right\},
\end{aligned} \tag{6}$$

where the first term assumes the mixture model in Equation 5 and the second term is a Markov Random Field prior for  $p(R_i)$  that encourages smooth region boundaries/surfaces; using other priors is also possible.  $N_j$  is the set of neighboring voxels of  $j$  and  $l(j')$  and  $l(j)$  are respectively the region labels of  $j'$  and  $j$ ;  $\delta(\cdot)$  is Kronecker's delta and  $\kappa$  is a constant that balances the weight of the smoothness prior versus the likelihood of the intensities. We assume  $p(\Theta_i) \propto 1$  so this term can be omitted. Such a flat prior implies that, a priori, we do not prefer any values of the Gaussian parameters over others. In other words, we assume no prior knowledge on the intensities of the image.

An estimate of  $W_R$  and  $\Theta$  can be obtained by minimizing  $E(W_R, \Theta, \mathbf{V})$ :

$$\hat{W} = (\hat{W}_R, \hat{\Theta}) = \underset{W_R}{\operatorname{argmin}} E(W_R, \Theta, \mathbf{V}). \tag{7}$$

Starting from an initial solution (a deformed template containing a volume  $V_a$  and its label annotation  $A_a$ ), we minimize the energy  $E(W_R, \Theta, \mathbf{V})$  in Equation 6 using a region competition algorithm [69, 91]. Henceforth, we denote this algorithm as *gmDT* (generative model based on a deformable template).

To avoid the initial solution is biased, we use the set of training volumes with their corresponding labels,  $S = \{(\mathbf{V}_1^{tr}, A_1^{tr}), \dots, (\mathbf{V}_N^{tr}, A_N^{tr})\}$ , to generate the template denoted as  $(V_a, A_a)$ . We use  $D(\mathbf{V}, \mathbf{V}_n)$  to denote the dissimilarity between  $\mathbf{V}$  and  $\mathbf{V}_n^{tr}$  after applying a linear transformation (in our case, computed with AIR [79]). The learned template volume  $\mathbf{V}_a$  minimizes the total dissimilarity with all other volumes in the training set:

$$\mathbf{V}_a = \arg \min_{\mathbf{V} \in S} \sum_{n=1}^N D(\mathbf{V}, \mathbf{V}_n^{tr}), \tag{8}$$

The corresponding manual annotation of  $V_a, A_a$ , is used as the initial  $W_R$  in Equation 7. Then, the obtained segmentation estimated by *gmDT*,  $\hat{W}_R$ , is an approximation to the optimal solution. A byproduct of *gmDT* is  $\hat{\Theta}$ , the parameter estimates.



### 3.3 Model fusion

Next, we discuss the discriminative model in our method, which is used to incorporate the adapted information from *gmDT*. For a purely discriminative model, we denote the total number of candidate features by  $B$  and the  $k^{th}$  feature for voxel  $j$  computed on volume  $\mathbf{V}(N_j)$  by  $f_{d,k}(j)$ . The discriminative feature vector  $\mathbf{F}_d(j)$  of each voxel  $j$  can be written as:

$$\mathbf{F}_d(j) \equiv [f_{d,1}(\mathbf{V}(N_j)), \dots, f_{d,B}(\mathbf{V}(N_j))] \quad (9)$$

A discriminative classifier, e.g., boosting [23], selects a number of informative features (typically a couple of hundred) from  $\mathbf{F}_d(\cdot)$  and fuses them with appropriate weights. The training process is driven by the minimization of the classification error in the labeled training data and the generalization power of the classifier (VC dimension, [71]). Hence, the quality of a trained discriminative classifier is greatly determined by the effectiveness of its feature set.

To achieve enhanced robustness, the basic idea here is to integrate the adaptiveness and the fusion capability respectively from generative and discriminative models. This is done by augmenting the feature vector  $\mathbf{F}_d$  with  $(\hat{W}_R, \hat{\Theta})$  from the deformable template. This way, we achieve robustness against intensity variations while we equip  $\mathbf{F}_d$  with structure-adapted features.

#### Using $\hat{W}_R$ for intensity normalization

Features computed directly from  $\mathbf{V}(N_j)$  are often sensitive to geometrical and intensity variations, but  $\hat{\Theta}$  from Equation 7 can then be used to normalize  $\mathbf{V}$ . We denote the normalized volume as  $\mathbf{V}_{\hat{\Theta}}$ . The new, augmented feature vector  $\mathbf{F}(j)$  is:

$$\mathbf{F}(j) \equiv [F_{\hat{W}_R}(j), f_{d,1}(\mathbf{V}_{\hat{\Theta}}(N_j)), \dots, f_{d,B}(\mathbf{V}_{\hat{\Theta}}(N_j))], \quad (10)$$

Comparing Equation 10 with Equation 9,  $f_{d,k}(\mathbf{V}_{\hat{\Theta}}(N_j))$  is computed on the normalized volume  $\mathbf{V}_{\hat{\Theta}}$ , instead of  $f_{d,k}(\mathbf{V}(N_j))$ , for a voxel  $j$ .  $F_{\hat{W}_R}(j)$  represents the augmented features based on  $\hat{W}_R$  that will be discussed below.

Normalization is achieved by intensity correction based on matching the intensity of the regions to those from the template volume [29]. Specifically, we search for the linear transform that best matches the intensities in a least squares sense, a problem that can be solved with standard techniques [66].

#### Augmenting atlas features from $\hat{W}_R$

From  $\hat{W}_R$  (given by *gmDT*), we have an estimated region label for each voxel  $j$ . To differentiate this estimated label from  $l_j$  in Equation 4, we use  $\hat{l}_j$  to denote it. From  $\hat{W}_R$ , the displacement vector of a voxel  $j$  to the centroid of structure  $k$ ,  $d_{\hat{R}_k}(j)$ , can be calculated. This displacement vector is a spatial feature which is more adaptive than the absolute coordinates. Similarly, we

can compute the signed distance function of each voxel  $j$  with respect to the estimated region boundary of each anatomical structure. The signed distance  $s_{\hat{R}_i}(j)$  of voxel  $j$  to the boundary of region  $\hat{R}_i$  is:

$$s_{\hat{R}_i}(j) = \begin{cases} + \min_{j' \in \hat{C}_i} d(j, j') & \text{if } \hat{l}(j) = i \\ - \min_{j' \in \hat{C}_i} d(j, j') & \text{otherwise} \end{cases}, \quad (11)$$

where  $d(j, j')$  is the distance between  $j$  to any point  $j'$  on the region boundary  $\hat{C}_i$ . The positive/negative sign indicates that  $j$  is inside/outside  $\hat{R}_i$ . Now our augmented feature vector becomes:

$$F_{\hat{W}_R}(j) = (d_{\hat{R}_0}(j), \dots, d_{\hat{R}_K}(j), s_{\hat{R}_0}(j), \dots, s_{\hat{R}_K}(j)), \quad (12)$$

Henceforth, we denote these features derived from  $\hat{W}_R$  as ‘‘atlas features’’. They correspond to  $F_{\hat{W}_R}(j)$  in Equation 10.

#### 4 Learning and using fusion models

Here we aggregate all the components in Section 3 to define the training and classification (testing) stages of our method. An atlas (template) will first be selected among the training data as the template. As described in Section 3.2, the template will be guided by *gmDT* to adapt to the input volumes. Though we know the true labels of the training data, *gmDT* is applied (normalization and atlas features) in training the discriminative models so the resultant classifiers can model the estimates from *gmDT* in the test stage.

##### 4.1 Atlas selection and feature constitution

In the training stage, we have the set of training volumes with their corresponding labels  $S = \{(\mathbf{V}_n^{tr}, A_n^{tr}), n = 1..N\}$ , which have been normalized to the same scale and properly preprocessed. A template,  $(\mathbf{V}_a, A_a)$ , can be learned based on Equation 8.

For our model, we will need to learn the uncertainty from *gmDT* before training the discriminative models. Using  $A_a$  as the initial labeling, we perform *gmDT* on the rest of training volumes in  $S$ . For each training volume  $\mathbf{V}_n^{tr}$ , an estimated segmentation  $\hat{W}_n^{tr}$  is obtained from *gmDT*. On the other hand, we define the feature set for the discriminative classifiers,  $\mathbf{F}_d(\cdot)$ , as local features such as gradients, curvatures, and Haar-like responses at various spatial scales (approximately 5,000 in total in this paper, but our method is not restricted to the specific  $\mathbf{F}_d(\cdot)$ ). Since the training volumes are in the same size, these features can be computed directly on a pre-defined sub-window (of size  $11 \times 11 \times 11$ ) centered at the target voxel. A detailed discussion of how these features are computed can be found in [69].

We use the estimated  $\hat{W}_n^{tr}$  to: (1) normalize the intensity of  $\mathbf{V}_n^{tr}$  to  $\mathbf{V}_a$  as in Section 3.3 and then compute  $[f_{d,1}(\mathbf{V}_{\hat{\Theta}}(N_j)), \dots, f_{d,B \approx 5000}(\mathbf{V}_{\hat{\Theta}}(N_j))]$  on

the normalized intensity volume; and (2) compute the atlas features  $F_{\hat{W}_R}(j)$  as in Equation 12. Combining the two sets yields the augmented feature set (about 6,000 features),  $F(j)$ , for each voxel  $j$  in the training volume  $\mathbf{V}_n^{tr}$ .

In short, *gmDT* is also applied to the training volumes. The role of the ground truth labels in the training stage is to train the discriminative, supervised classifiers; they do not participate before computing the augmented features.

#### 4.2 Integration within a discriminative framework

Once all the features  $\mathbf{F}(j)$  are computed, we train a classifier upon the training set

$$\mathfrak{N} = \{(l(j), \mathbf{F}(j)), j = 1..T\},$$

where  $l(j)$  is the true label for voxel  $j$ , and  $T$  is the total number of voxels in all the training volumes.

A learning algorithm either directly combines all the features in  $\mathbf{F}(j)$  like SVM [72], or selects a set of features out of  $\mathbf{F}(j)$  such as boosting [23] and random forests [8]. Either way, features are combined into the classifier in order to minimize the training error. When training our model, no preference was given to features derived from the atlas over those computed from image intensities (e.g., Haar-like), and vice versa. Here we adopt the auto-context algorithm using a cascade of PBT (Probabilistic Boosting Tree) classifiers [68] as the discriminative model, which performs feature selection and fusion by exploring a high-dimensional feature space. Note that, in addition to the features in  $\mathbf{F}(j)$ , auto-context [68] itself is an iterative method that incorporates contextual information into the classification by augmenting the feature space as follows. In a first iteration, the classifier is trained on the available set of features. In subsequent iterations, the label posteriors (as estimated by the current classifier) at a fixed set of shifted locations are added to the feature

- 1: **Given:** A set of training volumes together with their image features  $F(j)$  after applying *gmDT*,  $S = \{(\mathbf{V}_n^{tr}, A_n^{tr}), n = 1..N\}$ . The size of each volume is  $|V|$ .
- 2: For each volume  $V_n^{tr}$ , initialize probability maps  $P_n^{(0)}$  with uniform distribution on all the labels.
- 3: Iteratively train  $T$  classifiers:
- 4: **for**  $t = 1$  to  $T$  **do**
- 5:   Make a training set of voxel samples from  $S$ :  $\{(l_{ni}, (F(ni), P_n^{(t-1)}(i))), n = 1..N, i = 1..|V|\}$ , where  $i$  is a voxel of volume  $V_n^{tr}$  and  $l_{ni}$  is from  $A_n^{tr}$ .
- 6:   Train a classifier (in our implementation, a PBT classifier) on both image and context features from  $F(ni)$  and  $P_n^{(t-1)}(i)$  respectively. Output the classifier.
- 7:   Use the trained classifier to compute new probability maps  $P_n^{(t)}$  on all the labels for each training volume  $V_n^{tr}$ .
- 8: **end for**
- 9: The algorithm outputs a sequence of  $T$  trained classifiers for  $p^{(T)}(l_i|F(i), P^{(T-1)}(i))$

**Fig. 1** Training process of auto-context algorithm for our brain segmentation method.

- 1: **Given:**  $N$  training volumes with their corresponding annotations  $(V_1^{tr}, A_1^{tr}), \dots, (V_N^{tr}, A_N^{tr})$
- 2: Learn  $(V_a^{tr}, A_a^{tr})$  from the  $N$  training volumes as the atlas.
- 3: **for**  $i = 1$  to  $N$  **do**
- 4:   Obtain  $\hat{W}_i^{tr} = \{\hat{W}_R, \hat{\Theta}\}$  by minimizing Equation 6. This step is called *gmDT*.
- 5:   **for**  $j = 1$  to  $|A|$  **do**
- 6:     Obtain normalized feature set  $\mathbf{F}_d(i, j)$  based on  $\hat{\Theta}$ :  

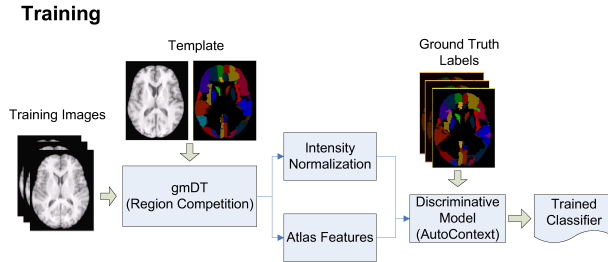
$$\begin{aligned} & [f_{d,1}(\mathbf{V}_i(N_j)), \dots, f_{d,B}(\mathbf{V}_i(N_j))] \\ & \rightarrow [f_{d,1}(\mathbf{V}_{i,\hat{\Theta}}(N_j)), \dots, f_{d,B}(\mathbf{V}_{i,\hat{\Theta}}(N_j))] \end{aligned}$$
- 7:     Merge  $\mathbf{F}_d(i, j)$  with the atlas features based on  $\hat{W}_R$ :
- 8:      $\mathbf{F}(i, j) \equiv [F_{\hat{W}_R}, f_{d,1}(\mathbf{V}_{i,\hat{\Theta}}(N_j)), \dots, f_{d,B}(\mathbf{V}_{i,\hat{\Theta}}(N_j))]$
- 9:   **end for**
- 10: **end for**
- 11: Take samples from  $\{A_i^{tr}(j), \mathbf{F}(i, j) : i = 1..n, j = 1..|A|\}$  to train a multi-class classifier  $p(l|\mathbf{F})$ .

**Fig. 2** Training process. All the training volumes are assumed to be skull-stripped and aligned.

space, implicitly capturing the shape of the structures to segment. A summary of the algorithm is described in Figure 1; the reader is referred to the original paper [68] for further details.

Nevertheless, the key of our proposed method is the augmentation/normalization of features, which implicitly fuses the generative and discriminate aspects of the model. It is not tied to any specific choice of classifier so one could also use boosting, random forests, or any probabilistic classifiers as the discriminative classifier.

Once a classifier has been trained on the training set  $\aleph = \{(l(j), \mathbf{F}(j)), j = 1..T\}$ , we can use it to estimate  $p(l|\mathbf{F}(i))$  for a given test voxel  $i$ . A test volume is required to have the same preprocessing steps and resized such that all features can be correctly computed. Each test voxel will be assigned to the label that maximizes the probability:  $l_i^* = \arg \max_l p(l|\mathbf{F}(i))$ .



**Fig. 3** Training procedures of the proposed fusion method. A template of the volume/label pair is selected and guided by *gmDT* using other training volumes as the input. The augmented feature set  $\mathbf{F}$  is then extracted to train the discriminative model giving the manual labels of the training volumes.

We summarize the training algorithm in Figure 2 and illustrate it in Figure 3. In Figure 3, the identical template,  $(\mathbf{V}_a, A_a)$ , will be used in both training and testing. The testing procedure consists of three stages analogous to those in the training procedure, except the last stage: performing classification using the trained classifier.

## 5 Experiments

To examine the effectiveness of the proposed algorithm, we perform a thorough empirical study using four MRI T1-weighted brain datasets and compare our method against the state-of-the-art systems. We focus on sub-cortical structures due to their popularity in the literature. Besides comparing our method with the two components in our method, *gmDT* and *DM*, we also include the methods developed by other researchers to show the integration can achieve better performance in most scenarios.

We will demonstrate the performance of the integrated method by four parts: (1) Increased importance of the adapted atlas features using the fusion mechanism; (2) Comparison using the same dataset for training and testing (intra-dataset). This scenario is very suit for *DM* and the proposed method is comparable to it; (3) Comparison using different datasets for training and testing (inter-dataset). This scenario is favored by *gmDT* and our method achieved better results; (4) Performance on longitudinal data, which shows the potential of our method to capture the morphological changes by the same subject. Though the fact that *DM* tends to fail in the last two parts is known, we still show its quantitative results for completeness.

### 5.1 Experimental setup

In this section, we elaborate the datasets used in our four parts of experiments, the three main algorithms to compare, the pre-processing steps we applied to the heterogeneous datasets, and the measures we used for our comparisons.

#### 5.1.1 T1 MRI datasets

All the datasets used in our experiments are the following (the suffix indicates the number of volumes in the dataset):

(1) *IBSR18*: We use the 18 scans with 84 manually annotated structures from the Internet Brain Segmentation Repository (IBSR) <sup>1</sup> (we do not include the 120 cortical/subcortical parcellations as 84 structures are sufficient in our experiment). All volumes were scanned at 1.5T. This dataset has been extensively utilized as a benchmark for evaluation in various papers <sup>2</sup>.

<sup>1</sup> <http://www.cma.mgh.harvard.edu/ibsr>

<sup>2</sup> <http://www.cma.mgh.harvard.edu/ibsr/PubsUsingIBSR.html>

(2) *LPBA40*: The LONI Probabilistic Brain Atlas dataset (LPBA) [62] contains volumes from 40 healthy subjects with 56 anatomical structures manually annotated. These structures include both cortical and subcortical regions. All subjects were scanned with a GE Signa 1.5T systems with a SPGR sequence.

(3) *LONI28*: 28 scans from normal subjects were acquired on a GE Signa 1.5T scanner with a SPGR sequence. Eight subcortical structures (left and right hippocampus, putamen, caudate nucleus, and lateral ventricle) were manually delineated by neuroanatomists.

(4) *ADNI980*: The Alzheimer’s Disease Neuroimaging Initiative (ADNI) [31] was launched in 2003 by government, private pharmaceutical companies and non-profit organizations. The goal of ADNI has been to develop measures for the progression of mild cognitive impairment (MCI) and early Alzheimer’s disease (AD). ADNI is the result of efforts of many co-investigators from a broad range of academic institutions and private corporations, and the data keeps growing by its follow-up projects. Readers may see [www.adni-info.org](http://adni-info.org) for more information. In this study, 490 pairs of brain scans (980 volumes in total) were selected from ADNI. They correspond to 490 subjects who are known to be one of the three groups: elderly controls, mild cognitive impairment (MCI), or Alzheimer’s Disease (AD). For each subject, two volumes were acquired 12 months apart for longitudinal analysis. This dataset is particularly challenging because the subjects were scanned at different sites with different scanners. Therefore, these volumes display high variability stemming from intrinsic longitudinal changes of subjects and scanning configurations. More details of the various scanning protocols can be found in <http://adni.loni.ucla.edu/research/protocols/mri-protocols/>.

Note that not all the datasets have manual delineations of all subcortical structures. We summarize the main characteristics of these four datasets, as well as the role they play in our experiments, in Table 1.

Dataset	Number of labels	Role in the study
<i>IBSR18</i>	84 (cortical+subcortical)	Training, Testing
<i>LPBA40</i>	56 (cortical+subcortical)	Training, Testing
<i>LONI28</i>	8 (subcortical)	Testing
<i>ADNI980</i>	0	Testing

**Table 1** Datasets used in this study. Note that we only use 14 subcortical structures in IBSR18 out of the 84 that are available.

### 5.1.2 Algorithms to compare

We adopt region competition [69, 91] as the *gmDT* process which iteratively minimizes the energy  $E(W_R, \Theta, \mathbf{V})$  in Equation 6. We choose region competition due to its simplicity and effectiveness. The template  $(V_a, A_a)$  is learned according to Equation 8, using Mattes mutual information as the dissimilarity function. The region competition process takes about 5 ~ 15 minutes to reach

a steady state (no more change of labels, or fluctuation of labels) of surface evolution. It runs typically for less than 15 iterations. Other approaches such as the level set methods [11,85] could also have been used to perform energy minimization in a similar manner.

On the other hand, we use the auto-context algorithm [68] with PBT as the baseline discriminative classifier. Henceforth, this discriminative model is denoted as *DM*. Its running time is 10 ~ 20 minutes, depending on the number of structures for segmentation.

Comparisons between our method, *gmDT* and *DM* demonstrates the advantages due to the integration. We also list the measures from literatures for the same T1 MRI dataset and structures if available.

### 5.1.3 Pre-processing

Before applying the three algorithms for comparison, these heterogeneous datasets several preprocessing steps. We eliminate the dominant spatial disparity between an input volume and an atlas by the following sequence of preprocessing steps: (1) re-orientation using AIR 2.5 [79] (if the two volumes were at different orientations); (2) skull stripping using BET in FSL 4.0.3 [65]; (3) a 12-parameter global affine registration using AIR 2.5; and (4) a diffeomorphic registration, SyN, from ANTS 1.9 [4].

We chose SyN as our non-linear method stage due to its speed and high accuracy [35]. We use the following settings: three resolution levels (30x50x5 iterations), step-length 0.15, probability mapping (PR) with 4mm radius as cost function, and regularization with a Gaussian filter with standard deviation 3mm. These parameters are obtained empirically and they provide sufficient spatial alignments for *gmDT* across the four test datasets. Under these settings, the SyN registration between an atlas and an image can be done under 30 minutes. Steps prior to SyN can be done in 5 minutes. Once the segmentation result is obtained, each preprocessing step is reverted to map the result back to the original space.

### 5.1.4 Measures for comparison

The main evaluation measure used here is the Dice overlap,  $Dice = \frac{2*|L \cap S|}{(|L|+|S|)}$ , where  $L$  and  $S$  are the sets of voxels manually annotated and those automatically segmented. Precision and recall rates are also used in Table 6, where  $Precision = |L \cap S|/|S|$  and  $Recall = |L \cap S|/|L|$ . Another popular measure in literature, Jaccard coefficient, can be directly calculated from Dice:  $Jaccard^{-1} = 2Dice^{-1} - 1$ . In the inter-dataset tests, as our method and *gmDT* are adaptive, we further compare them in surface consistency. Hausdorff distances [45] and the Mean distances [85] between two sets of surface voxels are measured:

$$H(A, B) = \max_{a \in A} \min_{b \in B} D(a, b), \quad M(A, B) = \sum_{a \in C_A} \min_{b \in C_B} \frac{D(a, b)}{|C_A|},$$

where  $A$  and  $B$  are sets of voxels.  $H(\cdot)$  and  $M(\cdot)$  are respectively the Hausdorff distance and the Mean distance.  $D$  is the underlying distance metric, which is

usually the Euclidean distance or the Manhattan distance.  $C_A$  is the surface of segment  $A$ . Note that both the directed Hausdorff distance and the Mean distances are not symmetric. We will use  $H(A,B)$ ,  $H(B,A)$ , and one direction (segmented to ground truth) of the Mean distances in our evaluation.

## 5.2 Importance of atlas features

To demonstrate the importance of using atlas features based on deformable templates, we compare the features used in two models trained to segment the 56 brain structures of LPBA, one with a fixed annotation  $A_a$  (no adaptation to the input volume) and another with an adapted atlas based on  $gmDT$ . The feature pool is dominated by Haar-like responses due to their effectiveness to describe the appearance at different spatial scales. Derivative features perform similarly to Haar, but are limited to eighteen local derivatives in the x, y, and z directions. Features in the column ‘‘Others’’ include intensities, gradients, and curvatures. As mentioned in Section 4.2, we perform feature selection when training the Probability Boosting Tree (PBT). For each node in PBT, a limited number of features are chosen to form a decision criterion (a boosting classifier) such that the classification error is minimized for the training data arriving at this node. Giving the same complexity of the trained classifiers (the same tree depth and the same number of features used at each tree node), we observe in the table that more atlas features are selected when using an adapted atlas rather than a fixed one; the percentage of the selected atlas features increases from <3% to 6%.

	Atlas	Haar	Location	Derivative	Others
<i>Proportion</i>	0.02%	99.4%	0.2%	0.3%	0.04%
fixed atlas	2.9%	48.1%	31%	18%	0%
adapted atlas	6%	46%	29%	18%	1%

**Table 2** Distributions of the first 120 selected (most informative) features for a fixed annotation  $A_a$  and an adapted  $\hat{W}_R$ . The first row shows the proportion of the corresponding type of features in the whole candidate feature pool for reference. The column ‘‘Others’’ includes the features of intensities, gradients, and curvatures.

## 5.3 Intra-dataset evaluation

In this section, we evaluate our algorithm using training and test data from the same dataset, which is common in brain image segmentation. IBSR18 and LPBA40 are included in this experiment.

### (1) *IBSR18*:

A total of 14 subcortical structures including the left and right lateral-ventricles are evaluated. Due to the relatively few number of volumes in IBSR18, we perform a three-fold cross-validation, i.e., six volumes in each



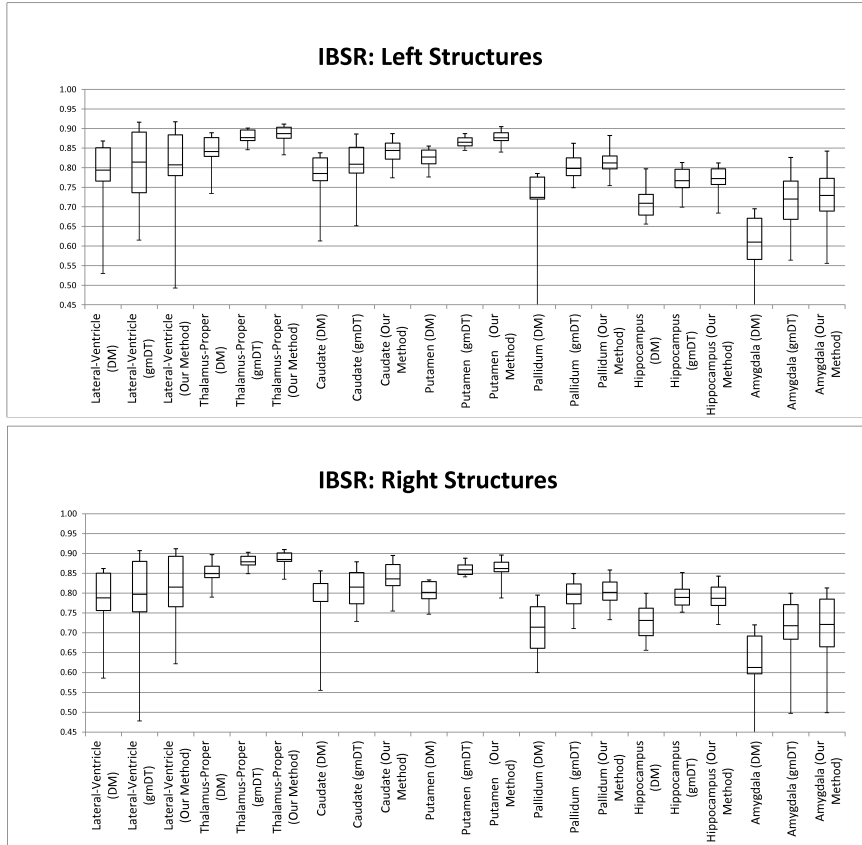
	LHippo.	LCaud.	LPut.	LVent.	LPal.	LThal.	LAmyg.
<i>DM</i>	0.709 (0.036)	0.785 (0.056)	0.827 (0.021)	0.794 (0.082)	0.724 (0.085)	0.841 (0.036)	0.610 (0.065)
<i>SyN</i>	0.708 (0.021)	0.734 (0.045)	0.822 (0.008)	NA	0.732 (0.021)	0.838 (0.012)	0.650 (0.039)
<i>gmDT</i>	0.767 (0.030)	0.809 (0.057)	0.865 (0.011)	<b>0.814</b> (0.093)	0.798 (0.031)	0.877 (0.015)	0.720 (0.086)
Our Method	<b>0.772</b> (0.029)	<b>0.844</b> (0.029)	<b>0.876</b> (0.016)	0.807 (0.108)	<b>0.812</b> (0.027)	<b>0.887</b> (0.015)	<b>0.729</b> (0.062)
	RHippo.	RCaud.	RPut.	RVent.	RPal.	RThal.	RAmyg.
<i>DM</i>	0.731 (0.040)	0.777 (0.072)	0.802 (0.028)	0.788 (0.073)	0.714 (0.057)	0.849 (0.024)	0.613 (0.104)
<i>SyN</i>	0.714 (0.021)	0.712 (0.053)	0.818 (0.012)	NA	0.732 (0.016)	0.840 (0.014)	0.647 (0.043)
<i>gmDT</i>	<b>0.789</b> (0.024)	0.815 (0.042)	0.859 (0.013)	0.797 (0.104)	0.797 (0.036)	0.879 (0.013)	0.718 (0.071)
Our Method	0.787 (0.029)	<b>0.836</b> (0.034)	<b>0.862</b> (0.025)	<b>0.815</b> (0.082)	<b>0.802</b> (0.035)	<b>0.885</b> (0.019)	<b>0.721</b> (0.083)

(a) Dice coefficients of *gmDT* and our method on the IBSR18 dataset.

Method	Hippo.	Caud.	Put.	Lateral-Vent.	Pal.	Thal.	Amyg.
Freesurfer	0.75	0.82	0.81	0.78	0.71	0.86	0.68
FSL	0.62	0.68	0.81	NA	0.73	0.85	0.59
[33]	0.76	0.83	0.87	0.85	0.72	<b>0.89</b>	0.66
[83]	NA	0.80	0.82	NA	NA	0.85	NA
[26]	0.69	0.76	0.78	0.84	0.72	NA	0.64
[2]	0.69	0.80	0.79	NA	0.74	0.84	0.63
[90]	0.70	0.80	0.81	NA	NA	0.84	0.64
[76]	0.73	0.80	0.82	NA	NA	NA	NA
[46]	NA	0.78	0.80	NA	NA	0.84	NA
[14]	0.72	0.80	0.83	<b>0.88</b>	0.77	0.83	<b>0.74</b>
Our Method	<b>0.78</b>	<b>0.84</b>	<b>0.87</b>	0.81	<b>0.81</b>	<b>0.89</b>	<b>0.73</b>

(b) Related reports on IBSR subcortical structures [33].

**Table 3** A comparison of our method and other methods on the IBSR18 dataset. We perform a three-fold cross-validation test on IBSR, each fold used 12 volumes for training and 6 volumes for testing. (a) The detailed Dice coefficients of *DM*, *gmDT*, and our method of the eight subcortical structures, where *DM* is the discriminative method and *gmDT* is the generative model based deformable template approach. The abbreviations are: L/RHippo.=Left/Right Hippocampus, L/RCaud.=Left/Right Caudate Nucleus, L/RPut.=Left/Right Putamen, L/RVent.=Left/Right Lateral Ventricle, L/RPal.=Left/Right Pallidum, L/RThal.=Left/Right Thalamus, L/RAmyg.=Left/Right Amygdala. The standard deviations of the Dice coefficients are shown in parentheses. (b) Other representative, subcortical segmentation results on the IBSR18 dataset [33]. Values are the averages of the left and right structures if they were separately reported. The best values are marked in bold.

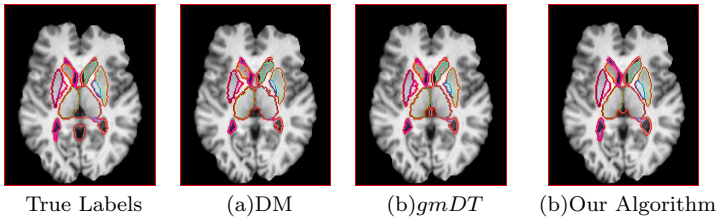


**Fig. 4** Box plots of the *Dice coefficients* of the discriminative model, the generative model based deformable template approach (*gmDT*), and our method on IBSR18. The top and bottom ends of a vertical line are the maximum and minimum values; The upper and lower edges of a box are the quartiles (25% and 75% data). The line inside a box indicates the average value, which is the same as Table 4.

fold are used as the test set, and the other twelve volumes are used to train our model.

We use the box plots in Figure 4 to compare the three methods and Table 3a to lists the Dice overlaps produced by DM, pure *SyN*, *gmDT*, and our method. The measures of *SyN* were included to clarify the improvements between it and our generative process.

In Table 3a, the row *SyN* lists six of the intermediate DICE overlaps before using *gmDT*. *SyN* performs a diffeomorphic transform mainly adapted to larger spatial variations of the whole brain so it can reduce location errors. Based on its rough results, *gmDT* utilized the relatively reliable statistics of each structure so the DICE values can be boosted at least 0.039(Thalami). Therefore, the use of *SyN* in our system is to reduce the risk of failure in



**Fig. 5** Segmentation results on a typical slice view by (a) discriminative method (DM), (b) the generative model based deformable template (*gmDT*), and (c) our algorithm. The left-most picture shows the ground truth labeling for comparison. Only partial of the extracted structures are shown in this 2D view. The example image is IBSR\_09 from the IBSR18 dataset.

*gmDT*; it is still a preprocessing step instead of a main factor in the whole our method.

The data in IBSR18 display larger variations in appearance and shape, and therefore, DM trained upon just 12 images in each fold could not generate satisfactory results. In this scenario, *gmDT* showcases its superior adaptiveness. From Figure 4, we observe that the advantage of *gmDT* is inherited by our fusion method but the similar and better measures.

A visual example of the extracted structures by DM, *gmDT* and our method is shown in Figure 5. Although not all of the 18 structures can be shown in a single 2D slice, better matched caudate nuclei, thalami, and putamens by our algorithm due to the fused information can still be observed. Table 3b lists several published reports on IBSR18, and our measures are at the top on the seven types of gray matter structures, with particularly significant improvements for the pallidum and amygdala. The results are close to [33] which is based on a computationally expensive multi-atlas approach. Their method outperforms ours for the ventricles (0.85 versus 0.80), but is inferior for the pallidum (0.72 versus 0.81) and amygdala (0.66 versus 0.73).

(2) *LPBA40*: In the LPBA40 dataset [62], 40 subjects with fifty-six anatomical structures, both cortical and sub-cortical, were manually delineated for each T1-weighted MRI volumes. We randomly choose 25 volumes for training and use the remaining 15 for testing.

To compare our algorithm against *gmDT* and *DM*, we use Figure 6 and the detailed measures in Table 4 for a comprehensive comparison. From Figure 6, we observe that fusing the two models in our method achieves higher average Dice coefficients than the two baseline methods. Our method gives better measures on L/R hippocampus ( $p$ -value=0.017/0.003) than both the discriminative and the generative models, but the measures of caudate nuclei are worse than DM due to the considerably degraded performance of the generative model. Reasons for this could be (1) the intra-dataset test of LPBA40 containing smaller data variance is favored by DM, and (2) the boundaries between the caudate nuclei and the ventricles in LPBA40 are obscured so *gmDT* could be misled. The weaknesses of *gmDT* in this scenario are the inferior DICE scores and quartile positions. These are all improved by our fusion

	LHippo.	LCaud.	LPut.	RHippo.	RCaud.	RPut.	Total 56 structures
DM	0.796	<b>0.813</b>	0.831	0.786	<b>0.828</b>	0.807	0.767(0.06)
<i>gmDT</i>	0.804	0.726	0.817	0.782	0.795	0.830	0.812(0.04)
Our Method	<b>0.835</b>	0.791	<b>0.838</b>	<b>0.819</b>	0.822	<b>0.845</b>	<b>0.822(0.04)</b>
p-value	0.017	0.136	0.053	0.003	0.051	0.063	2.5E-04

**Table 4** Accuracy measures (*Dice coefficients*) on the LPBA40 test volumes for extracting the 56 structures. We show the average Dice of the 56 structures and the results of six subcortical structures which are highly referred in the later tests. The abbreviations are: L/RHippo.=Left/Right Hippocampus, L/RCaud.=Left/Right Caudate Nucleus, L/RPut.=Left/Right Putamen. Our model is trained on 25 volumes from LPBA40. Values in parentheses are the standard deviations and we mark the best measures in bold. DM is the discriminative method and *gmDT* is our generative model based deformable template approach. The two-sided p-values between our method and *gmDT* of the six structures are also shown in the last row.

method due to the corrections in the discriminative framework, especially in the left/right caudates and the left putamen. The difference in the average Dice overlap between our method and *gmDT* is relatively small ( $\sim 1\%$ ), but still very statistically significant by the p-values shown in Table 4.

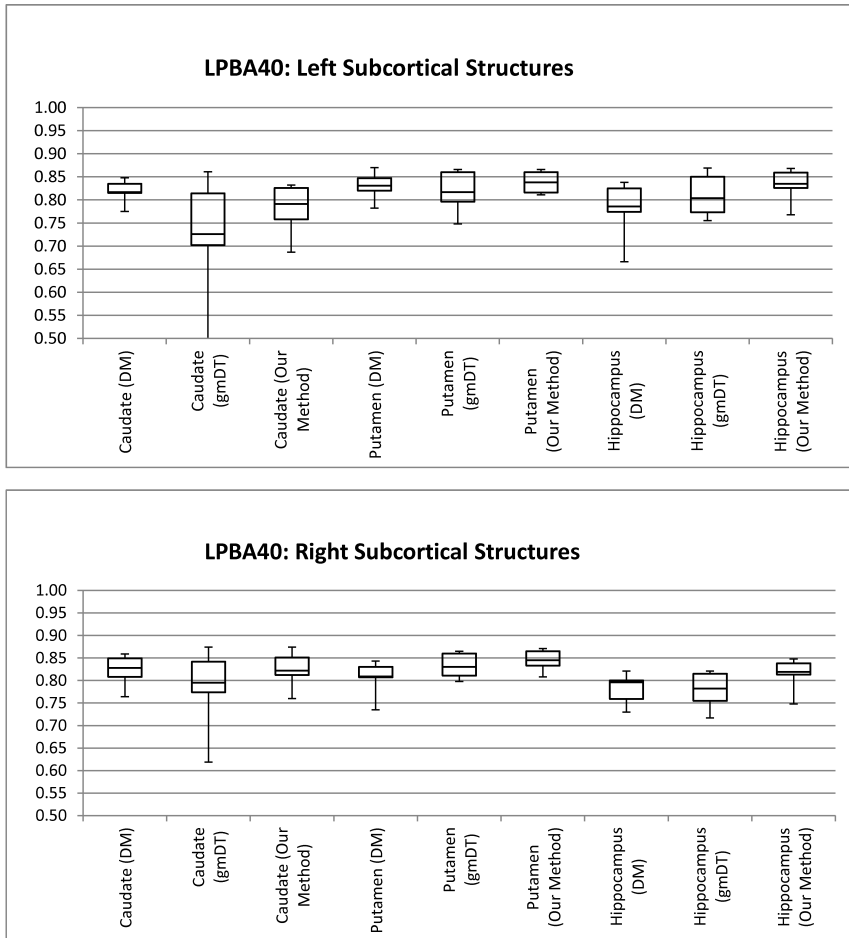
In summary, the two intra-dataset tests show the proposed fusion method successfully combine the advantages of *DM* and *gmDT*, and its performance is at least comparable to *DM*.

#### 5.4 Robustness across different datasets (Inter-dataset)

	Caudate	Putamen	Hippocampus
FreeSurfer	0.65(0.040)	0.64(0.105)	0.57(0.029)
FSL	0.63(0.063)	0.79(0.042)	0.53(0.043)
DM	0.73(0.058)	0.69(0.083)	0.58(0.037)
<i>gmDT</i>	0.74(0.064)	0.79(0.035)	0.57(0.029)
Our Method	<b>0.77(0.045)</b>	<b>0.80(0.035)</b>	<b>0.61(0.029)</b>

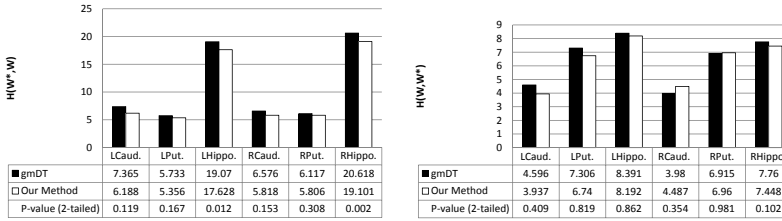
**Table 5** Inter-dataset measures (*Dice coefficients*) on 40 LPBA40 volumes for extracting the six subcortical structures. All values are the averages of the corresponding measures of the left and right structures, and the standard deviations are shown in parentheses. The model was trained from IBSR18. DM is the discriminative method and *gmDT* is our generative model based deformable template approach. The p-values between our method and *gmDT* are  $1.58 \times 10^{-5}$  (Caudate), 0.0135 (Putamen), and  $4.87 \times 10^{-14}$  (Hippocampus).

In the previous experiments, we trained and tested our method on volumes from the same dataset. To build a practical system dealing with clinical data, it is important to test its robustness on a large number of volumes from various sources.



**Fig. 6** Box plots of the *Dice coefficients* of the discriminative model, the generative model based deformable template approach (*gmDT*), and our method on LPBA40. The top and bottom ends of a vertical line are the maximum and minimum values; The upper and lower edges of a box are the quartiles (25% and 75% data). The line inside a box indicates the average value, which is the same as Table 4. All models were trained by the 25 images from LPBA40.

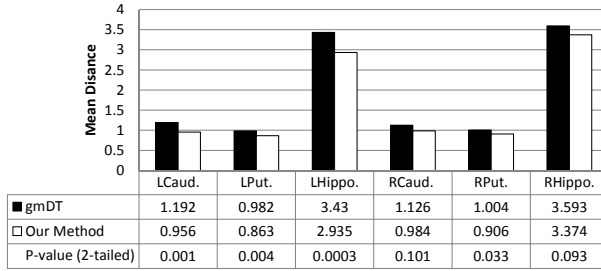
We use the same model trained from IBSR18 as in Section 5.3(1). The test datasets include LPBA40 and LONI28. Both of them have manual annotations of subcortical structures (caudate nuclei, putamens, and hippocampi for both sets, lateral ventricles only for LONI28). These structures are all covered by the annotations of IBSR (see Section 5.3(1)). We choose these subcortical structures because: (1) they are frequently used to evaluate automatic segmentation methods; and (2) they are very relevant in neuro-image studies of diseases such as Alzheimer’s and Parkinson’s. To keep the results comparable with other literatures on the same test datasets, we chose not to re-annotate the structures



**Fig. 7** Inter-dataset *Hausdorff distance* measures in *mm*, on 40 LPBA40 volumes for extracting the three types of subcortical structures (smaller is better). We denote the automated segmented result as  $W^*$  and the ground truth as  $W$ .

but keep the annotations as they were. In the following experiments,  $DM$ ,  $gmDT$ , and our algorithm are trained from the same training data. The other methods (FreeSurfer and FSL) are off-the-shelf systems without any parameter adjustments. Although the difference among the three protocols introduce intrinsic Dice errors, they still share a high proportion of common structure regions and the relative performance between the three models trained by the same data can be observed.

In addition to the quantitative analyses on the two test datasets, we also test the model trained on LPBA40 - Section 5.3(2) - and show visual examples on ADNI980, LPBA40 and LONI28 at the end of this section.

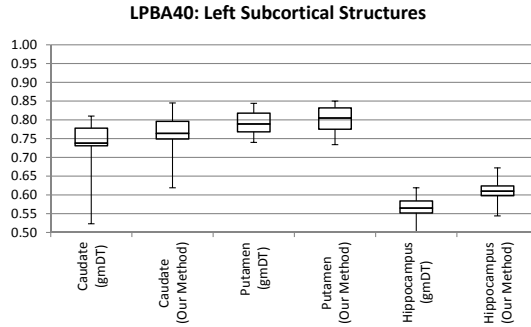


**Fig. 8** Inter-dataset *Mean distance* measures in *mm*, on 40 LPBA40 volumes for extracting the three types of subcortical structures (smaller is better). The standard deviations are shown in parentheses.

(1) *LPBA40*: Table 5 gives the Dice measures of three types of subcortical structures in LPBA40. All values are the averages of the corresponding measures of the left and right structures. Our method demonstrates the best performance among all five competing algorithms. Compared with the intra-dataset results in Table 4, noticeable degradation of the Dice overlaps is found in hippocampus among all methods in Table 5. This is because the subiculum region of hippocampus in LPBA40 was annotated differently from IBSR and many other datasets.

The improvement in surface distance measures given by our method is shown in Figures 7 and 8. We denote the automated segmented result as

$W^*$  and the ground truth as  $W$ . Our method achieves consistent improvement over  $gmDT$ . According to the p-values, the mean distance measures show especially significant improvement on the left caudate ( $p$ -value=0.0010), left putamen ( $p$ -value=0.0041), left hippocampus ( $p$ -value=0.0003), and right putamen ( $p$ -value=0.0326). The box plots in Figures 9 confirm the overall better performance of our method in terms of Dice overlap.



**Fig. 9** Box plots of the *Dice coefficients* of the generative model based deformable template approach ( $gmDT$ ) and our method on LPBA40. The top and bottom ends of a vertical line are the maximum and minimum values; The upper and lower edges of a box are the quartiles (25% and 75% data). The line inside a box indicates the average value, which is the same as Table 5. The model was trained by IBSR18.

(2) *LONI28*: The segmentation accuracy in eight subcortical structures on LONI28 (Left/Right Hippocampi, Left/Right Caudate Nuclei, Left/Right Putamens, and Left/Right lateral ventricles) by the model trained from IBSR18 is shown in Table 6. The results from Freesurfer and Hybrid are from [69]. The Hybrid model is trained by 14 LONI28 volumes and the measures are calculated from the other 14 test volumes. Although our model is trained by IBSR18, our method provides the best F-values in five structures as well as the averages in the last column. This shows the robustness of our method when segmenting the subcortical, gray matter structures in a different dataset.

The Dice overlaps in Table 6 show similar performances by  $gmDT$  and our method. Surface distance measures are showed in Figures 10 and 11. Our method gives smaller Mean and Hausdorff distances, while  $gmDT$  provides smaller  $H(W, W^*)$ . We notice that  $H(W^*, W)$  yields a significant improvement on most structures (LH, RH, LC, RC, LP, RP). From the four tables, we observe that  $W^*$  produced by our method has more consistent surface with  $W$  than  $gmDT$  (better  $H(W^*, W)$  and smaller Mean distance).

The improvement produced by our method can also be observed in the box plots in Figure 12. Our method not only increases the average Dice coefficient in all the tested structures (higher average values), but the worst cases have all improved.

(3) *56 structures on ADNI980, LONI28, and IBSR18*: In addition to the two subcortical tests on our IBSR18 subcortical model, we used the fifty-six

	LH	RH	LC	RC	LP	RP	LV	RV	Av
Precision	47%	54%	78%	78%	71%	77%	81%	72%	70%
Recall	66%	84%	77%	77%	83%	83%	76%	76%	78%
F-value	0.55	0.66	0.77	0.77	0.77	0.80	0.78	0.74	0.74

(a) FreeSurfer

	LH	RH	LC	RC	LP	RP	LV	RV	Av
Precision	77%	64%	81%	83%	68%	72%	81%	81%	76%
Recall	69%	62%	84%	81%	75%	75%	90%	90%	78%
F-value	<b>0.73</b>	0.63	<b>0.82</b>	0.82	0.71	0.73	<b>0.85</b>	<b>0.85</b>	0.77

(b) Hybrid [69](discriminative+shape)

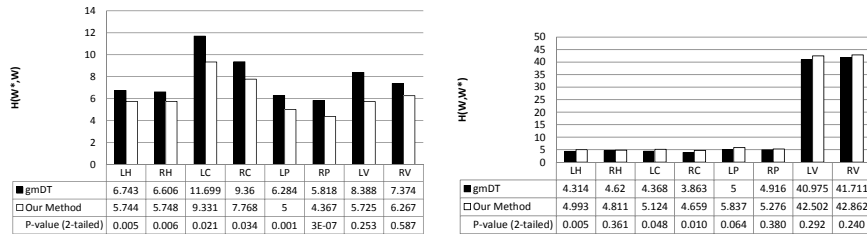
	LH	RH	LC	RC	LP	RP	LV	RV	Av
Precision	61%	56%	83%	83%	78%	78%	78%	78%	74%
Recall	81%	85%	78%	82%	81%	85%	77%	77%	81%
F-value	0.69	0.68	0.80	0.82	<b>0.79</b>	0.81	0.77	0.78	0.77

(c) *gmDT*

	LH	RH	LC	RC	LP	RP	LV	RV	Av
Precision	63%	58%	86%	86%	78%	83%	82%	82%	77%
Recall	83%	86%	79%	80%	81%	82%	75%	77%	80%
F-value	0.72	<b>0.69</b>	<b>0.82</b>	<b>0.83</b>	<b>0.79</b>	<b>0.83</b>	0.78	0.79	<b>0.78</b>

(d) Our Method

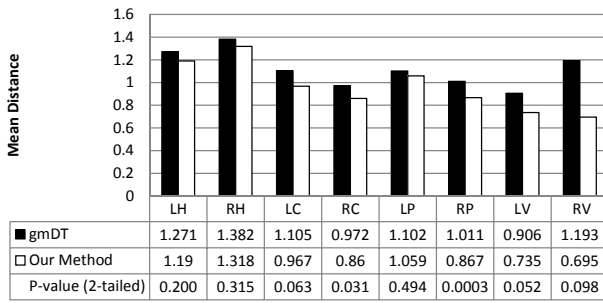
**Table 6** Inter-dataset measures (*Dice coefficients*) on 28 LONI28 test volumes for extracting the eight subcortical structures. The measures of FreeSurfer and Hybrid tested on 14 volumes of LONI28 were from [69], so we use the same measures (precision and recall rates) and F-values for comparison. [69] can be considered as a combination of DM and a shape prior. *gmDT* is the generative model based deformable template approach. The abbreviations are: L/RH=Left/Right Hippocampus, L/RC=Left/Right Caudate Nucleus, L/RP=Left/Right Putamen, L/RV=Left/Right Lateral Ventricle, Av=Average. In [69], the Hybrid model is trained by another 14 volumes of LONI28. Our model is trained from IBSR18 and still showed the best average values among the four methods. The best F-value of each structure is marked in bold.



**Fig. 10** Inter-dataset *Hausdorff distance* measures in *mm*, on 28 LONI28 volumes for extracting the eight subcortical structures (smaller is better). We denote the automated segmented result as  $W^*$  and the ground truth as  $W$ .

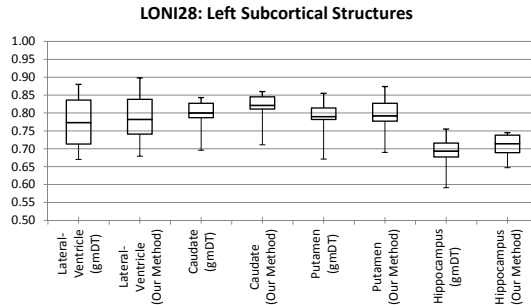
structure model trained upon LPBA40 in Section 5.3(2) to perform cortical and subcortical segmentation on three datasets: ADNI980, LONI28, and IBSR18. Figure 13 shows a number of 2D MRI slices together with their segmentation results. We see from the figure that the intensity patterns and textures are quite different for these datasets. Even within the same dataset, ADNI980, the MRI slices show large variation since not all of them were ac-





**Fig. 11** Inter-dataset *Mean distance* measures in *mm*, on 28 LONI28 volumes for extracting the eight subcortical structures (smaller is better). The standard deviations are shown in parentheses.

quired with the same scanner. However, despite such a high degree of variability in the data, the segmentation results are mostly satisfactory.

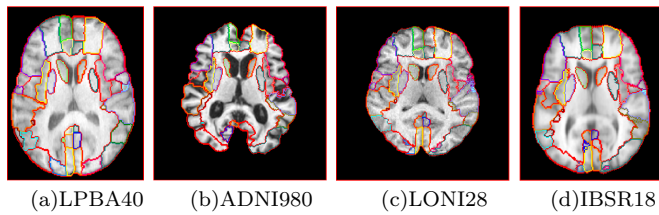


**Fig. 12** Box plots of the *Dice coefficients* of the generative model based deformable template approach (*gmDT*) and our method on LONI28. The top and bottom ends of a vertical line are the maximum and minimum values; the upper and lower edges of a box are the quartiles (25% and 75% ranked data). The line inside a box indicates the average value.

By the first two parts of tests containing disparate training and test datasets, we see the proposed fusion method can achieve the highest compatibility between different protocols, all by experts, than *DM* and *gmDT* given the same training set. The third test further shows its robustness to large data variation. Both properties are important when applying the proposed method as a common brain segmentation tool.

## 5.5 Performance in longitudinal studies

Section 5.4 demonstrated the effectiveness of the proposed algorithm for segmenting structures from several datasets with high variability in anatomy and image intensities. In this section, we show the results of our algorithm on data in a longitudinal study where the main source of variation is the temporal



**Fig. 13** Typical examples of the proposed brain segmentation method on different datasets. We use 2D slices of similar brain locations for comparison. Slice (a) is from LPBA40 to show the original imaging conditions of the training dataset. For clear comparison, these slices shown were skull-stripped and scaled to similar size. These four sets contain totally more than 1,000 volumes, and all the results are obtained by the identical system without parameter tuning.

changes within the same subject. Since the results from [30] indicate the significant atrophy of hippocampi in both MCI and AD cases, and the scans from ADNI980 provide the longitudinal (12 months) data of 490 subjects from AD, MCI, and normal control groups, we use this dataset as our testbed. ADNI980 volumes also display large variations between subjects due to different scanning settings; nevertheless, volumes from the same subject still share the same acquisition configurations. Although our method may need more specific design for high precision hippocampus segmentation, this experiment still shows our advantage with respect to the baseline methods in the longitudinal study.

The tested segmentation methods include DT, *gmDT*, and our algorithm. The training data contains volumes from 7 AD, 7 MCI, and 7 control subjects in ADNI with manual annotations of hippocampi by an expert (these subjects are not part of the 490). The measures listed in Table 7 include the average volume of baseline, the average volume after 12 months, the volume difference between the two times, and the percentage of volume loss.

All the measures obtained by a discriminative approach [68] are listed in Table 7a and those by *gmDT* are listed in Table 7b. As DM is not adaptive, the coverage of the segmented region by DM is conservative if the test hippocampus has a drifted position, a shape distortion like that typically produced by AD, or a different statistical distribution of appearance. DM also fails to identify the hippocampal regions of 13 subjects, so we needed exclude their volumes from Table 7a. For the rest of the scans, the averages of volume change and percentage of loss show noticeable differences between groups. However, their standard deviations are relatively large, indicating that the measures from the direct discriminative model are not stable for the three groups. In Table 7b, the longitudinal differences between AD and normal are not fully demonstrated because *gmDT* (with a simple appearance model) could include more non-hippocampal regions than DM and our method. Instead, our approach gives apparent differences in all columns among the three groups; see Table 7c. The smaller standard deviations provide a better separation of the three groups compared to DM. In addition, the balanced performance among the left hippocampus, the right hippocampus and the average is evident.

	baseline average volume ( $mm^3$ )			12-month average volume ( $mm^3$ )		
	Left	Right	Average	Left	Right	Average
AD	2301.22	2372.67	2336.95	2009.52	2158.16	2083.84
MCI	2527.80	2717.52	2622.66	2342.51	2546.46	2444.48
Normal	3039.01	3082.53	3060.77	2927.31	2980.78	2954.05
	volume change ( $mm^3$ )			percentage loss (%)		
	Left	Right	Average	Left	Right	Average
AD	291.71(544)	214.52(506)	253.11(504)	10.23%(35%)	8.27%(30%)	9.44%(30%)
MCI	185.29(449)	171.06(504)	178.18(453)	7.30%(22%)	6.64%(24%)	7.12%(21%)
Normal	111.70(520)	101.75(592)	106.72(539)	1.65%(21%)	1.40%(24%)	1.67%(22%)

(a) DM : DM fails to process the volumes from 13 subjects so the statistics here are based on 477 subjects. The difference between the three groups as well as the longitudinal change inside each group are demonstrated. However, the standard deviations of all measures are big.

	baseline average volume ( $mm^3$ )			12-month average volume ( $mm^3$ )		
	Left	Right	Average	Left	Right	Average
AD	3545.03	3884.74	3714.89	3507.99	3778.29	3643.14
MCI	3607.94	3870.49	3739.21	3579.45	3846.53	3712.99
Normal	3751.34	4008.77	3880.05	3706.05	3962.01	3834.03
	volume change ( $mm^3$ )			percentage loss (%)		
	Left	Right	Average	Left	Right	Average
AD	37.04(413)	106.45(433)	71.75(382)	0.53%(13%)	2.18%(13%)	1.50%(11%)
MCI	28.49(408)	23.96(462)	26.22(401)	0.19%(11%)	0.21%(12%)	0.29%(11%)
Normal	45.28(359)	46.76(353)	46.02(345)	0.62%(9%)	0.64%(10%)	0.72%(9%)

(b) *gmDT* : The extracted volumes are consistent due to the adaptation power of generative model, but it fails to represent the longitudinal changes of the three groups.

	baseline average volume ( $mm^3$ )			12-month average volume ( $mm^3$ )		
	Left	Right	Average	Left	Right	Average
AD	2999.04	3115.78	3057.41	2837.53	2970.87	2904.20
MCI	3238.77	3415.67	3327.22	3352.16	3150.12	3251.14
Control	3788.27	3938.24	3863.26	3751.32	3897.77	3824.54
	volume change ( $mm^3$ )			percentage loss (%)		
	Left	Right	Average	Left	Right	Average
AD	161.52(161)	144.92(176)	153.22(141)	5.56%(6%)	4.96%(6%)	5.25%(5%)
MCI	88.65(168)	63.51(177)	76.08(149)	2.70%(6%)	1.79%(6%)	2.27%(5%)
Normal	36.95(153)	40.47(153)	38.71(136)	0.98%(4%)	1.06%(4%)	1.03%(4%)

(c) Our Algorithm: The results faithfully reflect both the absolute volumes and longitudinal changes of the three groups compared with DM (a) and *gmDT* (b).

**Table 7** 12-month longitudinal analysis of the extracted hippocampi from the 490 subjects. DM is the discriminative model [68] and *gmDT* is the generative model based deformable template approach. We demonstrate the changes in left hippocampus, right hippocampus, and the average of both sides. Individual volume change is obtained by the baseline volume subtracting the volume of 12-month after. Values listed are the averages of the 490 subjects, and those in parentheses are the standard deviations.

We further compare the results by our method with the estimated hippocampal volume changing rates reported by [60]. Their estimated rates are modeled as nonlinear curves based on manual segmentations of ADNI data. As the average ages of our three groups are 76.82(standard deviation(SD)=6.63 ) for Normal, 76.05(SD=6.67) for MCI, and 76.82(SD=6.44) for AD, our data are mostly located in the range [70-80], where their estimated curves are still

close to linear. Using the average hippocampal volume of 75 years old normal subjects as the standard level, their chart shows the estimated average volume losses as  $400 \text{ mm}^3$  (sMCI, the subjects do not convert to AD),  $650 \text{ mm}^3$  (cMCI, the subjects would progress to AD), and  $750 \text{ mm}^3$  (AD). Our average volume of the whole MCI group is  $573.40 \text{ mm}^3$  less than the Normal group and consistent with this estimation. However, the number of our AD group is  $920.34 \text{ mm}^3$  which is  $170 \text{ mm}^3$  larger than their estimation. This is a hint of what direction to work in if we want to specifically adjust our work in order to obtain precise hippocampal segmentations. Table 7 shows that our method achieves higher robustness when identifying inter- and intra- individual differences, and its potential to help indicate different pathological stages.

## 6 Conclusion and future work

In this paper, we have proposed a system for brain MRI image segmentation by fusing together deformable templates (generative) and informative features (discriminative). It takes the advantages of the generative model for being adaptive and the discriminative classifier for achieving classification power on high dimensional data. This approach uses a new way of combining generative and discriminative models and the complementary properties between them can be efficiently exerted; the information extracted from generative models is considered as an additional channel of features for training the discriminative models. The trained models are improved in two ways: (1) The original features can be normalized according to intrinsic structural statistics. Typical methods to accommodate large variation, such as performing histogram matching [64] or extracting features invariant to intensity change [70], are only based on the statistics of whole volume data. (2) The feature set is augmented with features derived from the estimation by *gmDT*.

A thorough experimental study on T1-weighted datasets demonstrates the robustness of our algorithm. Although discriminative models can perform well if the training and the test data share the same condition of variances, better performance can still be observed by our method as our discriminative models augment the informative feature set with generative features. This advantage leads to improvement over several state-of-the-art algorithms. The inter-dataset and the longitudinal tests show the deficiency of discriminative models in practice and the necessity of introducing the generative information. Our method demonstrates both adaptiveness and precision in these challenging tests and outperforms the two direct models in region overlaps (Dice) and surface fitness. This is different from [74] that the first stage classifier is considered as a black box approach. These advantages also lead to the improvement over several state-of-the-art algorithms on standard datasets such as IBSR18, LPBA40, and ADNI.

An important aspect of the proposed method is its running time, which is approximately one hour. Whether the system is practical depends on the application. In neuroimaging studies, in which research labs often spend months

collecting the data, slow running times are not a problem. For instance, our method is much faster than the widely used FreeSurfer, which requires on average 12 hours to segment a single brain scan. In clinical practice, one hour might not be sufficiently fast. The bottleneck of the algorithm is, as for many other brain MRI segmentation methods, the nonlinear registration. However, the registration can be dramatically sped-up through parallelization.

An aspect of the framework that was not evaluated was its performance on multispectral data. In this scenario, the different data channels represent images acquired with different MRI contrast [54,47,84,24] or even different modalities [20,12]. Segmentations on multispectral data have the potential to be more accurate thanks to the larger amount of information present in the different channels. In our framework, generalization to multispectral scenarios is immediate: the intensities of the additional channels are just extra dimensions of the feature vectors. Another possible line of research would be to analyze the performance of the framework using other generative or discriminative models. For instance, it would be interesting to assess whether introducing explicit shape, regional, or context information in the generative prior has a positive impact of the segmentation. Exploring all these directions remains as future work.

### Information Sharing Statement

An implementation of the method is publicly available for download at the LONI<sup>3</sup> and NITRC<sup>4</sup> websites. We provide a Windows®, a Linux®, and a LONI pipeline version. The software can be used freely in research provided this paper is cited in any material using the results of their application. For other usage, contact the authors.

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<sup>3</sup> <http://www.loni.ucla.edu>

<sup>4</sup> <http://www.nitrc.org/projects/brainparser>

Alzheimer Immunotherapy Research & Development, LLC, Johnson & Johnson Pharmaceutical Research & Development LLC, Medpace, Inc., Merck & Co., Inc., Meso Scale Diagnostics, LLC, Novartis Pharmaceuticals Corp., Pfizer Inc., Servier, Synarc Inc., and Takeda Pharmaceutical Co.. The Canadian Institutes of Health Research is providing funds to support ADNI clinical sites in Canada. Private sector contributions are facilitated by the Foundation for the National Institutes of Health ([www.fnih.org](http://www.fnih.org)). The grantee organization is the Northern California Institute for Research and Education, and the study is coordinated by the Alzheimer's Disease Cooperative Study at the University of California, San Diego. ADNI data are disseminated by the Laboratory for Neuro Imaging at the University of California, Los Angeles. This research was also supported by NIH grants P30 AG010129 and K01 AG030514.

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