

Bootstrapping Multi-atlas Segmentation Using Multiple Automatically Generated Templates for the Segmentation of the Whole Hippocampus and Subfields

Jon Pipitone¹, Min Tae M. Park¹, Julie Winterburn¹, Tristram A. Lett^{1,9}, Jason P. Lerch^{2,3}, Jens C. Pruessner⁴, Martin Lepage^{4,5}, Aristotle N. Voineskos^{1,6,9}, M. Mallar Chakravarty^{1,6,7,8} and the Alzheimer’s Disease Neuroimaging Initiative*

¹*Kimel Family Translational Imaging-Genetics Lab, Centre for Addiction and Mental Health, Toronto, ON, Canada*

²*Neurosciences and Mental Health Laboratory, Hospital for Sick Children, Toronto, ON, Canada*

³*Department of Medical Biophysics, University of Toronto, Toronto, ON, Canada*

⁴*Douglas Mental Health University Institute, Verdun, QC, Canada*

⁵*Department of Psychiatry, McGill University, Montreal, QC, Canada*

⁶*Department of Psychiatry, University of Toronto, Toronto, ON, Canada*

⁷*Institute of Biomaterials and Biomedical Engineering, University of Toronto, Toronto, ON, Canada*

⁸*Rotman Research Institute, Baycrest, Toronto, ON, Canada*

⁹*Institute of Medical Science, University of Toronto, Toronto, ON, Canada*

Abstract

Introduction: Advances in image segmentation of magnetic resonance images (MRI) have demonstrated that multi-atlas approaches improve segmentation accuracy and precision over regular atlas-based approaches. These approaches often rely on a large number of such manually segmented atlases (e.g. 30-80) that take significant time and expertise to produce. We present an algorithm, MAGeT-Brain (Multiple Automatically Generated Templates), for the automatic segmentation of the hippocampus that minimizes the number of atlases needed while still achieving similar agreement to multi-atlas approaches. Thus, our method acts as an accurate multi-atlas approach when using special, hard-to-define atlases that are laborious to construct.

Method: MAGeT-Brain works by propagating atlas segmentations to a template library, formed from a subset of target images, via transformations estimated by nonlinear image registration. The resulting segmentations are then propagated to each target image and fused using a label fusion method.

We conduct two separate Monte Carlo cross-validation experiments comparing MAGeT-Brain and multi-atlas whole hippocampal segmentation using differing atlas and template library sizes, and registration and label fusion methods. The first experiment is a 10-fold validation (per parameter setting) over

*Data used in preparation of this article were obtained from the Alzheimer’s Disease Neuroimaging Initiative (ADNI) database (adni.loni.ucla.edu). As such, the investigators within the ADNI contributed to the design and implementation of ADNI and/or provided data but did not participate in analysis or writing of this report. A complete listing of ADNI investigators can be found at: http://adni.loni.ucla.edu/wp-content/uploads/how_to_apply/ADNI_Acknowledgement_List.pdf

60 subjects taken from the ADNI Alzheimer’s disease imaging database, and the second is a 5-fold validation over 81 subjects having had a first episode of psychosis. In both cases, automated segmentations are compared with manual segmentations following the Pruessner-protocol. Using the best settings found from these experiments, we segment 246 images of the ADNI1:Complete 1Yr 1.5T dataset and compare these with segmentations from existing automated methods: FSL FIRST, FreeSurfer, MAPER, and SNT. Finally, we conduct a leave-one-out cross-validation (LOOCV) of hippocampal subfield segmentation using five high-resolution manually segmented atlases (Winterburn et al., 2013).

Results: [TODO: update with FEP-XVAL and subfield XVAL results.] Using 9 atlases and 19 template images, MAGeT-Brain achieves a mean Dice’s Similarity Coefficient (DSC) of 0.869 (to Pruessner-protocol manual segmentations) over 10-folds of Monte Carlo significantly lower variability in DSC than multi-atlas segmentation. In a LOOCV, MAGeT-Brain accurately reproduces the subfields...

MAGeT-Brain produces hippocampal volumes in a first episode psychosis patient population that are highly correlated with expert manual segmentation volumes (Pearson $r = 0.877$, $t = 16.244$, $p < 0.001$). Compared to FSL FIRST and FreeSurfer, MAGeT-Brain shows much smaller fixed volume bias (within $250mm^3$ on average) to semi-automated (SNT) segmentations available from ADNI, as well as a conservative, rather than exaggerated, proportional volume bias.

Conclusion: We demonstrate that MAGeT-Brain produces accurate hippocampal segmentations using only 5 atlases over different hippocampal definitions, disease populations, and acquisition types, as well as showing that accurate identification of the hippocampal subfields is possible.

Contact:

Jon Pipitone and M. Mallar Chakravarty
 Kimel Family Translation Imaging-Genetics Research Laboratory
 Research Imaging Centre
 Centre for Addiction and Mental Health
 250 College St.
 Toronto, Canada M5T 1R8
 jon.pipitone@camh.ca; mallar.chakravarty@camh.ca

1 Introduction

The hippocampus is a brain structure situated in the medial temporal lobe, and has long been associated with learning and memory (den Heijer et al., 2012; Jeneson and Squire, 2012; Wixted and Squire, 2011; Scoville and Milner, 2000). The hippocampus is of interest to clinical neuroscientists because it is implicated in many forms of brain dysfunction, including Alzheimer’s disease (Sabuncu et al., 2011) and schizophrenia (Narr et al., 2004; Karnik-Henry et al., 2012). In neuroimaging studies, structural magnetic resonance images (MRI) are often used for the volumetric assessment of the hippocampus. As such, accurate segmentation of the hippocampus and its subfields in MRI is a necessary first step to better understand the inter-individual variability of subject neuroanatomy.

The gold standard for neuroanatomical segmentation is manual delineation by an expert human rater. However, with the availability of increasingly large MRI datasets the time and expertise required for manual segmentation becomes prohibitive (Mazziotta et al., 1995, 2001; Mazziotta et al.; Pausova et al., 2007). This effort is complicated by the fact that there is significant variation between segmentation protocols with respect to specific anatomical boundaries of the hippocampus (Geuze et al., 2004) and this has led to

efforts to create an unified hippocampal segmentation protocol (Jack et al., 2011; Boccardi et al., 2013b,a). In addition, there is controversy over the appropriate manual segmentation protocol to use in a particular imaging study (Nestor et al., 2012). Thus, a segmentation algorithm that can easily adapt to different manual segmentation definitions would be of significant benefit to the neuroimaging community.

Automated segmentation techniques that are reliable, objective, and reproducible can be considered complementary to manual segmentation. In the case of classical model-based segmentation methods (Haller et al., 1997; Csernansky et al., 1998), an MRI atlas that was previously manually labelled by an expert rater is matched to target images using nonlinear registration methods. The resulting nonlinear transformation is applied to the manual labels (i.e. *label propagation*) to warp them into the target image space. While this methodology has been used successfully in several contexts (Chakravarty et al., 2008, 2009; Collins et al., 1995; Haller et al., 1997), it is limited in accuracy due to error in the estimated nonlinear transformation itself, partial volume effects in label resampling, and irreconcilable differences between the neuroanatomy represented within the atlas and target images.

One methodology that can be used to mitigate these sources of error involves the use of multiple manually segmented atlases and probabilistic segmentation techniques, such as those found in the FreeSurfer package (Fischl et al., 2002). FreeSurfer uses a probabilistic atlas of anatomical and tissue classes along with spatial constraints for class labels encoded using a Markov random field model to segment the entire brain.

More recently, many groups have used multiple atlases to improve overall segmentation accuracy (i.e. multi-atlas segmentation) over model-based approaches (Heckemann et al., 2006a, 2011; Collins and Pruessner, 2010; Lötjönen et al., 2010; Aljabar et al., 2009; Leung et al., 2010; Wolz et al., 2010). Each atlas image is registered to a target image, and label propagation is performed to produce several labellings of the target image (one from each atlas). A *label fusion* technique, such as voxel-wise voting, is used to merge these labels into the definitive segmentation for the target. In addition, weighted voting procedures that use *atlas selection* techniques are often used to exclude atlases from label fusion that are dissimilar to a target image in order to reduce error from unrepresentative anatomy (Aljabar et al., 2009). This involves the selection of a subset of atlases using a similarity metric such as cross-correlation (Aljabar et al., 2009) or normalized mutual information. Such selection has the added benefit of significantly reducing the number of nonlinear registrations. For example Collins and Pruessner (2010) demonstrated that only 14 atlases, selected based on highest similarity between medial temporal lobe neuroanatomy as evaluated by normalized mutual information (Studholme et al., 1999) from a library of 80 atlases, were required to achieve accurate segmentations of the hippocampus. Also, several methods have been explored for label fusion. For example, the STAPLE algorithm (Simultaneous Truth And Performance Level Estimation; Warfield et al. (2004)) uses an expectation-maximization framework to compute a probabilistic segmentation from a set of competing segmentations, or the work of Coupé et al. (2012) who show that a subset of segmentations can be estimated using metrics, such as the sum of squared differences in the regions of interest to be segmented.

However, many of these methods require significant investment of time and resources for the creation of the atlas library ranging between 30 (Heckemann et al., 2006a) and 80 (Collins and Pruessner, 2010) manually segmented atlases. This strategy has the main drawback of being inflexible as it does not easily accommodate varying the definition of the hippocampal anatomy (such as the commonly used heuristic of subdividing the hippocampus into head, body, and tail (Poppenk and Moscovitch, 2011; Pruessner et al., 2000)). Furthermore, none of these methods have demonstrated sufficient flexibility to accommodate atlases that are somehow exceptional such as those derived from serial histological data (Chakravarty et al., 2006;

Yelnik et al., 2007) or high-resolution MRI data that enables robust identification of hippocampal subfields (Winterburn et al., 2013; Yushkevich et al., 2009; Mueller and Weiner, 2009; Van Leemput et al., 2009; Wisse et al., 2012). Due to the recent availability of the latter, there has been increased interest in the use of probabilistic methods for the identification of the hippocampal subfields on standard T1-weighted images. Our group recently demonstrated that through use of an intermediary automated segmentation stage, robust and accurate segmentation of the striatum, pallidum, and thalamus using a single atlas derived from serial histological data is possible (Chakravarty et al., 2013). The novelty of this manuscript is the extension of our multi-atlas methodology to the segmentation of hippocampus. Additionally, in this paper we rigorously explore the effects of using multiple input atlases, of varying the size of the template library constructed, and registration and label fusion methods. As a result, we aim to demonstrate that it is indeed possible to reliably apply the segmentation represented in a very small set of segmented input atlases to an unlabelled target image set.

Of particular relevance to the present work is the LEAP algorithm (Learning Embeddings for Atlas Propagation; Wolz et al. (2010)) because of its focus on performing multi-atlas segmentation with a limited number of input atlases. The LEAP algorithm is a clever modification to the basic multi-atlas strategy in which an atlas library is grown, beginning with a set of manually labelled atlases, by successively incorporating unlabelled target images once they themselves have been labelled using multi-atlas techniques. The sequence in which target images are labelled is chosen so that the similarity between the atlas images and the target images is minimised at each step, effectively allowing for deformations between very dissimilar images to be broken up into sequences of smaller deformations. Although Wolz et al. (2010) begin with an atlas library of 30 MR images, this method could theoretically work using a much smaller atlas library. In their validation, LEAP was used to segment the whole hippocampus in the ADNI1 baseline dataset, achieving a mean Dice score of 0.85 against semi-automated segmentations.

Also of interest to this manuscript are the methods that attempt to define hippocampal subfields using standard T1- or T2-weighted data, of which there are few. Van Leemput et al. (2009) demonstrate that the applicability of hippocampal subfield segmentation in T1-weighted images by Bayesian techniques using Markov random field shape priors learned from 10 manual segmentations. This work, available as part of the FreeSurfer package, is limited as the segmentation omits the tail of the hippocampus and the protocol has yet to be fully validated. Yushkevich et al. (2009) manually segment hippocampal subfields on high-resolution (either $0.2mm^3$ isotropic or $0.2mm \times 0.3mm \times 0.2mm$ resolution voxels) T2-weighted MR images acquired from five post-mortem medial temporal lobe samples. Then, using nonlinear registration guided by shape-based models of the subfield segmentations, and manually derived hippocampus masks of the target images, the authors demonstrate accurate parcellation of hippocampal subfields in clinical 3T T1-weighted MRI volumes. Using multi-atlas with bias correction techniques, Yushkevich et al. (2010) demonstrate a semi-automated method of subfield segmentation on in vivo focal T2-weighted MR acquisitions of the temporal lobe. Manual input is only needed to mark divisions between the head, body and tail of the hippocampus on target images.

In this paper we describe a thorough validation of the MAGeT-Brain algorithm for the fully automatic segmentation of the hippocampus and its subfields. First, we address the very idea of bootstrapping a template library from a limited number of input atlases (Chakravarty et al., 2013) for whole hippocampus segmentation by conducting a multi-fold validation experiment over a range of atlas and template library sizes, registration and label fusion methods. This type of validation is done first on a subset of the Alzheimer’s

Disease Neuroimaging Initiative (ADNI) dataset with manual segmentations Pruessner-protocol, and then replicated on a first episode psychosis patient dataset to determine the behaviour of MAGeT-Brain when segmenting younger and differently diseased subjects. Next, we compare MAGeT-Brain with other popular segmentation algorithms (FreeSurfer, FSL FIRST, MAPER, and SNT) on all the images available in the ADNI1:Complete 1Yr 1.5T sample. Lastly, using the optimal parameter settings for MAGeT-Brain found from the previous experiments, we investigate hippocampal subfield segmentation by conducting a leave-one-out validation using the Winterburn et al. (2013) manually segmented high-resolution MR atlases.

2 The MAGeT-Brain Algorithm

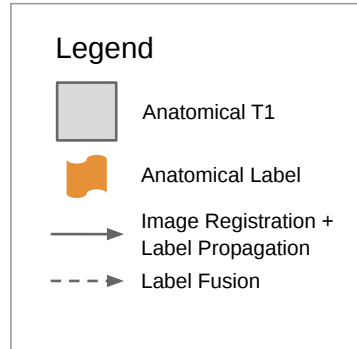
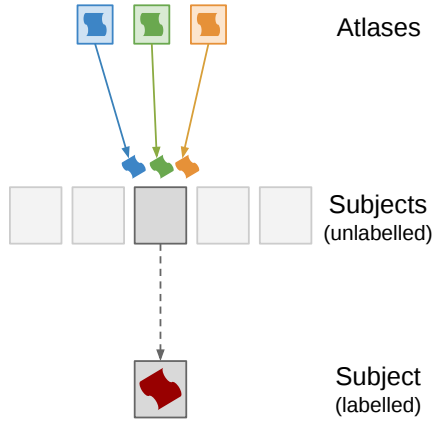
In this paper, we use the term *label* to mean any segmentation (manual or derived) of an MR image. *Label propagation* is the process by which two images are registered and the resulting transformation is applied to the labels from one image to bring them into alignment with the other image. We use the term *atlas* to mean a manually segmented image, and the term *template* to mean an automatically segmented image (i.e. via label propagation). The terms *atlas library* and *template library* describe any set of such images. Additionally, we use the term *target* to refer to an unlabelled image that is undergoing segmentation.

The simplest form of multi-atlas segmentation, which we call *basic multi-atlas segmentation*, involves three steps. First, each labelled input image (i.e. atlas or template) is registered to an unlabelled target image. Second, the labels from each image are propagated to the target image space. Third, the labels are combined into a single label by label fusion (Heckemann et al., 2006a, 2011). The basic multi-atlas segmentation method is described in detail in other publications (Collins and Pruessner, 2010; Heckemann et al., 2011; Aljabar et al., 2009). When only a single atlas is used, basic multi-atlas segmentation degenerates into model-based segmentation: labels are propagated from the atlas to a target, and no label fusion is needed.

MAGeT-Brain (**M**ultiple **A**utomatically **G**enerated **T**emplates) bootstraps the creation of a large template library given a limited input atlas library, and then uses the template library in basic multi-atlas segmentation. Images for the template library are selected from a set of input target images, either arbitrarily or so as to reflect the neuroanatomy or demographics of the target set as a whole (for instance, by sampling equally from cases and controls). The template library images are automatically labelled by each of the atlases via label propagation. Effectively, basic multi-atlas segmentation is then conducted using the template library to segment the entire set of target images (including the target images used in the construction of the template library). Since each template library image has multiple labels (one from each atlas), the final number of labels to be fused for each target may be quite large (i.e. $\# \text{ of atlas} \times \# \text{ of templates}$).

Figure 1 illustrates the MAGeT-Brain algorithm graphically. Source code for MAGeT-Brain can be found at <http://github.com/pipitone/MAGeTbrain>.

Multi-Atlas Segmentation



MAGeT Brain Segmentation

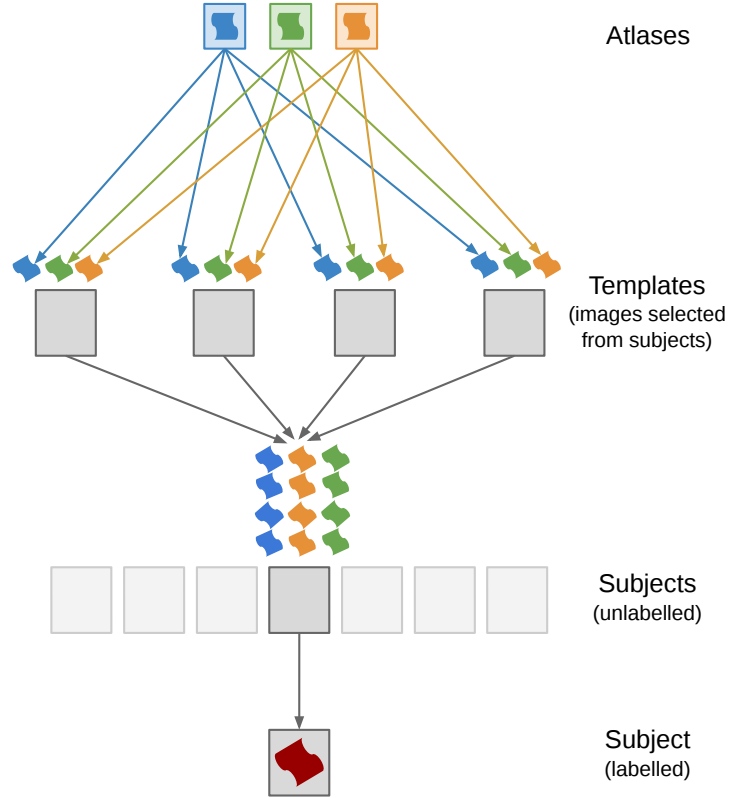


Figure 1: A schematic illustration of basic multi-atlas segmentation and MAGeT-Brain segmentation. In multi-atlas segmentation, manual labels from atlas images are warped (propagated) into subject space by applying the transformations estimated from nonlinear image registration. The resulting candidate labels from all atlas images are then fused to create a final segmentation. In MAGeT-Brain segmentation, a template library is created by sampling (either randomly or representatively) from the subject images. Atlas labels are propagated to all template images and then to each subject image (including those used in the template library). The candidate labels for a subject are then fused into a final segmentation.

3 Experiments

The following section describes experiments conducted to assess the segmentation quality of the MAgE-T-Brain algorithm:

- Experiment 1 investigates MAgE-T-Brain whole hippocampus segmentation of aging and Alzheimer’s diseased subjects over a wide range of parameter settings using a Monte Carlo cross-validation design. The results of this experiment enable us to choose the parameter settings offering the best performance for use in subsequent experiments.
- Experiment 2 consists of a similar cross-validation to explore MAgE-T-Brain segmentations on the brain images of young, first episode psychosis patients, and to establish parameter settings that do not overfit on a single dataset.
- Experiment 3 bridges MAgE-T-Brain with the existing segmentation literature by comparing MAgE-T-Brain whole hippocampus segmentations with those of well-known automated methods (FreeSurfer, FSL FIRST, MAPER, SNT) on the entire ADNI1:Complete 1Yr 1.5T image dataset consisting of 246 brain images of subjects diagnosed as cognitively normal, having mild cognitive impairment, or Alzheimer’s disease.
- Experiment 4 assesses hippocampal subfield segmentation quality in a leave-one-out cross-validation on the five high-resolution manually segmented Winterburn MR atlases (Winterburn et al., 2013).

3.1 Experiment 1: Whole Hippocampus Segmentation Cross-Validation — Alzheimer’s Disease

In this experiment we explore the very idea of bootstrapping a template library for multi-atlas-based segmentation from a small number of input atlases. To do so, we conduct repeated cross-validations of MAgE-T-Brain whilst varying the composition and sizes of the atlas and template libraries used, as well as varying the registration algorithm and label fusion method. The dataset used in this experiment is images from the ADNI dataset (Jack et al., 2008) along with whole hippocampus labels manually segmented following the Pruessner-protocol (Pruessner et al., 2000).

Note, in the Supplementary Materials we have replicated this experiment using the SNT semi-automated segmentations included as part of the ADNI dataset.

3.1.1 Experiment 1: Materials and Methods

ADNI1:Complete 1Yr 1.5T dataset Data used in the preparation of this article were obtained from the Alzheimer’s Disease Neuroimaging Initiative (ADNI) database (adni.loni.usc.edu). The ADNI was launched in 2003 by the National Institute on Aging (NIA), the National Institute of Biomedical Imaging and Bioengineering (NIBIB), the Food and Drug Administration (FDA), private pharmaceutical companies and non-profit organizations, as a \$60 million, 5-year public-private partnership. The primary goal of ADNI has been to test whether serial magnetic resonance imaging (MRI), positron emission tomography (PET), other biological markers, and clinical and neuropsychological assessment can be combined to measure the progression of mild cognitive impairment (MCI) and early Alzheimer’s disease (AD). Determination of sensitive and

Table 1: **ADNI1 cross-validation subset demographics.** CN - Cognitively Normal. LMCI - Late-onset Mild Cognitive Impairment. AD - Alzheimer’s Disease. CDR-SB - Clinical Dementia Rating-Sum of Boxes. ADAS - Alzheimer’s Disease Assessment Scale. MMSE - Mini-Mental State Examination.

	CN			LMCI			AD			Combined		
	<i>N</i> = 20			<i>N</i> = 20			<i>N</i> = 20			<i>N</i> = 60		
Age at baseline Years	72.2	75.5	80.3	70.9	75.6	80.4	69.4	74.9	80.1	70.9	75.2	80.2
Sex : Female	50% (10)			50% (10)			50% (10)			50% (30)		
Education	14.0	16.0	18.0	13.8	16.0	16.5	12.0	15.5	18.0	13.0	16.0	18.0
CDR-SB	0.00	0.00	0.00	1.00	2.00	2.50	3.50	4.00	5.00	0.00	1.75	3.62
ADAS 13	6.00	7.67	11.00	14.92	20.50	25.75	24.33	27.00	32.09	9.50	18.84	26.25
MMSE	28.8	29.5	30.0	26.0	27.5	28.2	22.8	23.0	24.0	24.0	27.0	29.0

a b c represent the lower quartile *a*, the median *b*, and the upper quartile *c* for continuous variables.

Numbers after percents are frequencies.

specific markers of very early AD progression is intended to aid researchers and clinicians to develop new treatments and monitor their effectiveness, as well as lessen the time and cost of clinical trials.

The Principal Investigator of this initiative is Michael W. Weiner, MD, VA Medical Center and University of California San Francisco. ADNI is the result of efforts of many co-investigators from a broad range of academic institutions and private corporations, and subjects have been recruited from over 50 sites across the U.S. and Canada. The initial goal of ADNI was to recruit 800 subjects but ADNI has been followed by ADNI-GO and ADNI-2. To date these three protocols have recruited over 1500 adults, ages 55 to 90, to participate in the research, consisting of cognitively normal (CN) older individuals, people with early or late MCI, and people with early AD. The follow up duration of each group is specified in the protocols for ADNI-1, ADNI-2 and ADNI-GO. Subjects originally recruited for ADNI-1 and ADNI-GO had the option to be followed in ADNI-2. For up-to-date information, see www.adni-info.org.

Sixty 1.5T images were arbitrarily selected from the baseline scans in the *ADNI1:Complete 1Yr 1.5T* standardized dataset. Twenty subjects were chosen from each disease category: cognitively normal (CN), mild cognitive impairment (MCI) and Alzheimer’s disease (AD). Demographics for this subset are shown in Table 1. Fully manual segmentations of the left and right whole hippocampi in these images were provided by one author (JCP) according to the segmentation protocol specified in Pruessner et al. (2000).

Clinical, demographic and pre-processed T1-weighted MRI were downloaded by the authors from the ADNI database (adni.loni.ucla.edu) between March 2012 and August 2012. The image dataset used was the ADNI1:Complete 1Yr 1.5T standardized dataset available from ADNI ¹ (Wyman et al., 2012). This image collection contains uniformly pre-processed images which have been designated to be the “best” after quality control. All images were acquired using 1.5T scanners (General Electric Healthcare, Philips Medical Systems or Siemens Medical Solutions) at multiple sites using the protocol described in Jack et al. (2008). Representative 1.5T imaging parameters were TR = 2400ms, TI = 1000ms, TE = 3.5ms, flip angle = 8°, field of view = 240 x 240mm, a 192 × 192 × 166 matrix (*x*, *y*, and *z* directions) yielding voxel dimensions of 1.25mm × 1.25mm × 1.2mm.

¹<http://adni.loni.usc.edu/methods/mri-analysis/adni-standardized-data/>

Table 2: **ANIMAL registration parameters.**

Parameters	Stage 1	Stage 2	Stage 3
Model Blur (FWHM)	8	8	4
Input Blur (FWHM)	8	8	4
Iterations	30	30	10
Step	8x8x8	4x4x4	2x2x2
Sub-Lattice	6	6	6
Lattice Diameter	24x24x24	12x12x12	6x6x6

Experiment details Monte Carlo Cross-Validation (MCCV), also known as repeated random sub-sampling cross-validation, consists of repeated rounds of validation conducted on a fixed dataset (Shao, 1993). In each round, the dataset is randomly partitioned into a training set and a validation set. The method to be validated is then given the training data, and its output is compared with the validation set.

In this experiment, our dataset consists of 60 1.5T images and corresponding Pruessner-protocol manual segmentations. In each validation round, the dataset is partitioned into a training set consisting of images and manual segmentations used as an atlas library, and a validation set consisting of the remaining images to be segmented by both MAgE-T-Brain and multi-atlas. The computed segmentations are compared to the manual segmentations (see Evaluation below).

A total of ten validation rounds were performed on each subject in the dataset, over each combination of parameter settings. The parameter settings explored are: atlas library size (1-9), template library size (1-20), registration method (ANTS or ANIMAL, described below), and label fusion method (majority vote, cross-correlation weighted majority vote, and normalized mutual information weighted majority vote, described below). In each validation round, both a MAgE-T-Brain and multi-atlas segmentation is produced. A total of $10 \times 60 \times 9 \times 20 \times 2 \times 3 = 6.48 \times 10^5$ validation rounds were conducted and resulting segmentations analysed.

Before registration, all images underwent preprocessing with the N3 algorithm (Sled et al., 1998) to minimize intensity nonuniformity. In this experiment we compared two nonlinear image registration methods:

Automatic Normalization and Image Matching and Anatomical Labeling (ANIMAL) The ANIMAL algorithm carries out image registration in two phases. In the first, a 12-parameter linear transformation (3 translations, rotations, scales, shears) is estimated between images using an algorithm that maximizes the correlation between blurred MR intensities and gradient magnitude over the whole brain (Collins et al., 1994). In the second phase, nonlinear registration is completed using the ANIMAL algorithm (Collins et al., 1995): an iterative procedure that estimates a 3D deformation field between two MR images. At first, large deformations are estimated using a blurred version of the input data. These larger deformations are then input to subsequent steps where the fit is refined by estimating smaller deformations on data blurred with a Gaussian kernel with a smaller full width at half maximum (FWHM). The final transformation is a set of local translations defined on a bed of equally spaced nodes that were estimated through the optimization of the correlation coefficient. For the purposes of this work we used the regularization parameters optimized in Robbins et al. (2004), displayed in Table 2.

Automatic Normalization Tools (ANTs) ANTS is a diffeomorphic registration algorithm which provides great flexibility over the choice of transformation model, objective function, and the consistency of the final transformation (Avants et al., 2008). The transformation is estimated in a hierarchical fashion where the MRI data is subsampled, allowing large deformations to be estimated and successively refined at later hierarchical stages (where the data is subsampled to a finer grid). The deformation field and the objective function are regularized with a Gaussian kernel at each level of the hierarchy. The ANTS algorithm is freely available <http://www.picsl.upenn.edu/ANTS/>. We used an implementation of the ANTS algorithm compatible with the MINC data format, mincANTS <https://github.com/vfonov/mincANTS>.

We used the following command line when running ANTS:

```
mincANTS 3 -m PR[target_file.mnc,source_file.mnc,1,4]
--number-of-affine-iterations 10000x10000x10000x10000x10000
--affine-gradient-descent-option 0.5x0.95x1.e-4x1.e-4
--use-Histogram-Matching --MI-option 32x16000
-r Gauss[3,0] -t SyN[0.5] -i 100x100x100x20
-o transformation.xfm
```

These settings were adapted from the "reasonable starting point" given in the ANTS manual ².

Label fusion methods Label fusion is a term given to the process of combining the information from several candidate labels for an image into a single labelling. In this experiment we explore three fusion methods:

Voxel-wise Majority Vote Labels are propagated from all template library images to a target. Each output voxel is given the most frequent label at that voxel location amongst all candidate labels.

Cross-correlation Weighted Majority Vote An optimal combination of targets from the template library has previously been shown to improve segmentation accuracy (Aljabar et al., 2009; Collins and Pruessner, 2010). In this method, each template library image is ranked in similarity to each unlabelled image by the normalized cross-correlation (CC) of image intensities after linear registration, over a region of interest (ROI) generously encompassing the hippocampus. Only the top ranked template library image labels are used in a voxel-wise majority vote. The ROI is heuristically defined as the extent of all atlas labels after linear registration to the template, dilated by three voxels (Chakravarty et al., 2013). The number of top ranked template library image labels is a configurable parameter and displayed as the size of the template library in the rest of the paper.

The `xcorr_vol` utility from the ANIMAL toolkit is used to calculate the cross-correlation similarity measure.

Normalised Mutual Information Weighted Majority Vote This method is similar to cross-correlation weighted voting except that image similarity is calculated by the normalised mutual information score over the region of interest (Studholme et al., 2001). The `itk_similarity` utility from the EZMinc toolkit³ is used to calculate the normalised mutual information measure between two images.

²<https://sourceforge.net/projects/advants/files/Documentation/>

³<https://github.com/vfonov/EZminc>

Evaluation method The Dice similarity coefficient (DSC), also known as Dice's Kappa, assesses the agreement between two segmentations. It is one of the most widely used measures of segmentation agreement, and we use it as the basis of comparison in this experiment.

$$\text{Dice's coefficient (DSC)} = \frac{2|A \cap B|}{|A| + |B|}$$

where A and B are the regions being compared, and the cardinality is the volume measured in voxels. The labels produced by MAGeT-Brain and multi-atlas segmentation are compared to the manual labels using the Dice similarity coefficient, and the recorded value for each subject at each parameter setting explored in this experiment is the average over ten validation rounds.

Additionally, the sensitivity of MAGeT-Brain and multi-atlas to atlas and template library composition is evaluated by comparing the variability in Dice scores over all validation rounds at fixed parameter settings. This is achieved by first computing the variance of DSC scores in each block of ten validation rounds per subject. The distribution of these statistics across all subjects is then compared between MAGeT-Brain and multi-atlas using a Student's t-test. A significant difference between distributions is taken to show either a larger or smaller level of variability between methods.

3.1.2 Experiment 1: Results

We find that for MAGeT-Brain segmentations, similarity score increases as atlas and template library size is increased, although with diminishing returns and an eventual trend towards a plateau (Figure 2a). For instance, with 9 atlases and using ANTS for registration and majority vote fusion, the mean DSC scores for 1, 5, 9 and 17 templates are 0.844, 0.865, 0.867, 0.868, respectively. A maximum similarity score of 0.869 is found when using 9 atlases, 19 templates, ANTS registration, and majority vote label fusion.

The ANTS registration method consistently outperforms ANIMAL registration over all variable settings we tested (mean increase in DSC is 0.078). Pearson correlations of MAGeT-Brain DSC scores when using weighted voting and when using non-weighted majority vote label fusion (with ANTS registration) for all combinations of atlases and templates are $r > 0.899$, $p < 0.001$, with a mean difference in DSC score of 0.002. This result suggests that using a weighted voting strategy does not significantly improve MAGeT-Brain segmentation agreement, contrary to the findings of Aljabar et al. (2009) for basic multi-atlas segmentation. Thus, in the remainder of our experiments only results using the ANTS registration algorithm and majority vote fusion will be shown.

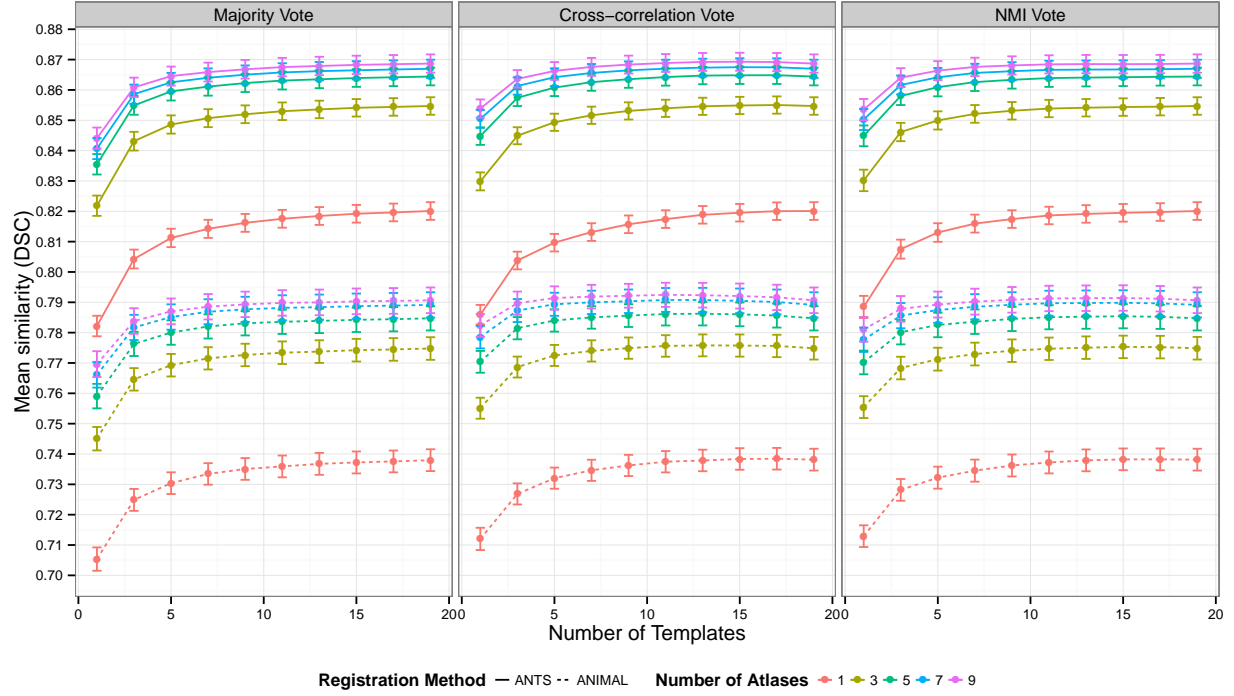
With at least five templates, MAGeT-Brain consistently shows a higher DSC score than multi-atlas segmentation with the same number of atlases: $r = 0.94$, $p < 0.001$, mean difference = 0.008 (Figure 2b). The magnitude of DSC increase grows with template library size but shows diminishing returns with larger atlas libraries. Peak increase (+0.025 DSC) is found with a single atlas and template library of 19 images.

In addition to a mean increase in similarity score over multi-atlas-based segmentation, MAGeT-Brain also shows more consistency in similarity scores across all subjects and validation folds (Figure 2c). A template library of at least 13 images is sufficient to show significant ($p < 0.05$) decrease in variance for all sizes of atlas library tested (1-9 images).

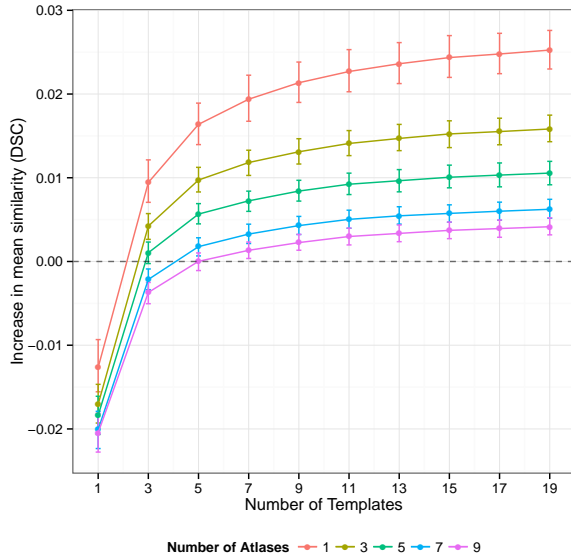
We find similar behaviour with respect to optimal parameter settings and increased consistency of MAGeT-Brain segmentations in the replication of this experiment (Experiment 5, Supplementary Materials) where a different hippocampal definition is used (SNT labels available with the ADNI datasets). This

strongly suggests that these results are independent of the segmentation protocol used and are, instead, features of the MAgE-T-Brain algorithm.

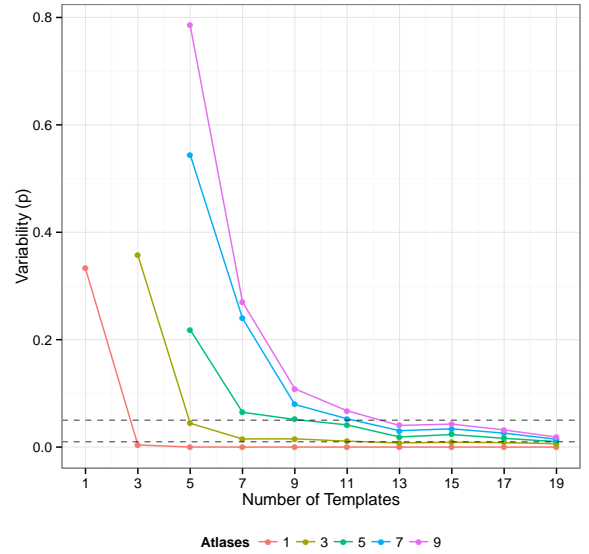
We have omitted results obtained when using an even number of atlases or templates since with these configurations we found significantly decreased performance. We believe this results from an inherent bias in the majority vote fusion method used (see Discussion).



(a) DSC vs. atlas and template library size



(b) Increase in similarity score over multi-atlas



(c) Difference in variability with multi-atlas

Figure 2: **Whole hippocampus segmentation cross-validation on ADNI subjects with Pruessner-protocol manual segmentations.** (2a) Average DSC score of MAGeT-Brain with manual segmentations for 60 ADNI subjects taken over 10 folds of cross-validation at each parameter setting. Error bars indicate standard error. (2b) Increase in DSC of MAGeT-Brain over multi-atlas segmentations. (2c) shows the significance of t-tests comparing the variability in DSC scores of MAGeT-Brain and multi-atlas across validation folds. Only points where MAGeT-Brain mean variability is lower than multi-atlas are shown. Dashed lines indicate p-values of 0.05 and 0.01.

3.2 Experiment 2: Whole Hippocampus Segmentation Cross-Validation — First Episode of Psychosis

To validate that the MAGE-T-Brain works effectively in the context of other neurological disorders, in this experiment we replicate the cross-validation done in Experiment 1 with a dataset of patients having had a single episode of psychosis. We also compare MAGE-T-Brain segmentations with those of two well-known automated segmentation methods, FSL FIRST and FreeSurfer.

3.2.1 Experiment 2: Materials and Methods

First Episode Psychosis (FEP) Dataset All patients were recruited and treated through the Prevention and Early Intervention Program for Psychoses (PEPP-Montreal), a specialized early intervention service at the Douglas Mental Health University Institute in Montreal, Canada. People aged 14 to 35 years from the local catchment area suffering from either affective or non-affective psychosis who had not taken antipsychotic medication for more than one month with an IQ above 70 were consecutively admitted as either in- or out-patients. Of those treated at PEPP, only patients aged 18 to 30 years with no previous history of neurological disease or head trauma causing loss of consciousness were eligible for the neuroimaging study; only those suffering from schizophrenia spectrum disorders were considered for this analysis. For complete program details see Malla et al. (2003).

Scanning of 81 subjects was carried out at the Montreal Neurological Institute on a 1.5-T Siemens whole body MRI system. Structural T1 volumes were acquired for each participant using a three-dimensional (3D) gradient echo pulse sequence with sagittal volume excitation (repetition time=22ms, echo time=9.2ms, flip angle=30°, 180 1mm contiguous sagittal slices). The rectangular field-of-view for the images was 256mm (SI)×204mm (AP). Subject demographics are shown in Table 3.

Expert whole hippocampal manual segmentation of each subject is produced following a validated segmentation protocol (Pruessner et al., 2000).

Winterburn Atlases The Winterburn atlases (Winterburn et al., 2013) are digital hippocampal segmentations of five in-vivo 0.3mm-isotropic T1-weighted MR images. The segmentations include subfield segmentations for the cornu ammonis (CA) 1; CA2 and CA3; CA4 and dentate gyrus; subiculum; and strata radiatum (SR), strata lacunosum (SL), and strata moleculare (SM). Subjects in the Winterburn atlases range in age from 29-57 years (mean age of 37), and include two males and three females.

Experiment details The same overall design as Experiment 1 is followed in this experiment: a Monte Carlo cross-validation (MCCV) is conducted using the pool of 81 first episode psychosis subject brain images and corresponding Pruessner-protocol manual segmentations. Five rounds of validation are conducted for each subject, and each atlas and template library size combination (1-9 atlases, 1-19 templates). In each round, images and their manual labels are randomly selected from the pool, and the remaining images are segmented using MAGE-T-Brain with a random subset of the unlabelled images also serving as template images. Majority vote fusion, and the ANTS registration algorithm are used, as these have shown to behave favourably in previous experiments.

In addition to the MCCV, we segment the entire first episode psychosis dataset using MAGE-T-Brain and two other popular segmentation packages, FSL FIRST and FreeSurfer. Firstly, MAGE-T-Brain is used

Table 3: **First Episode Psychosis Subject Demographics.** ambi - ambidextrous. SES - Socioeconomic Status score. FSIQ - Full Scale IQ.

	N	FEP
		<i>N</i> = 81
Age	80	21 23 26
Gender : M	81	63% (51)
Handedness : ambi	81	6% (5)
left		5% (4)
right		89% (72)
Education	81	11 13 15
SES : lower	81	31% (25)
middle		54% (44)
upper		15% (12)
FSIQ	79	88 102 109

a b c represent the lower quartile *a*, the median *b*, and the upper quartile *c* for continuous variables.

N is the number of non-missing values.

Numbers after percents are frequencies.

with the five Winterburn atlas images and labels as atlases, and a randomly selected subset of 19 psychosis dataset images as templates. Secondly, MAGeT-Brain is run using the same template images, but instead as atlases we use five additional first episode psychosis subjects and corresponding manual segmentations (not included above). Finally, FSL FIRST and FreeSurfer are run with their default settings to segment the entire dataset. The FSL FIRST `run_first_all` script was used according to the FIRST user guide ⁴, and FreeSurfer was run using `recon-all -all`.

Evaluation method Manual and automated segmentations are directly compared using Dice’s similarity coefficient (DSC). In the MCCV, the per-subject DSC value is computed as the average value over the five rounds of validation for a given atlas and template library size. The reported average DSC value per given atlas and template library size is the average DSC value over all subjects segmented.

The Pruessner segmentation protocol differs slightly from the Winterburn protocol, and those used by FreeSurfer and FSL FIRST, in the inclusion of neuroanatomical features and the manner they are delineated (see Winterburn et al. (2013), and Table 9 in the Discussion below). This variation in protocol poses a problem if an overlap measure is used for evaluation: since different protocols will necessarily produce segmentations that do not perfectly overlap, the degree of overlap cannot be solely used to compare segmentation methods using different protocols. In place of an overlap metric, we assess the degree of (Pearson) correlation in average bilateral hippocampal volume produced by each method. Additionally, we evaluate the volume-related fixed and proportional biases in all segmentation methods using Bland-Altman plots (Bland and Altman, 1986).

3.2.2 Experiment 2: Results

As in Experiment 1, we find that similarity score increases with a greater number of atlases or templates but quickly plateaus (Figure 3a). A maximum similarity score of 0.841 is found when using 9 atlases, 19

⁴<http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FIRST/UserGuide>

templates, ANTS registration, and majority vote label fusion.

We found a close relationship in average hippocampal volume between the manual label volumes and MAgE-T-Brain when using the Winterburn atlases, or manually segmented FEP subjects as atlases (Figure 3b). Both sets of volumes are correlated with Pearson $r > 0.88$. FreeSurfer and FSL FIRST volumes are both correlated with manual volumes at Pearson $r > 0.7$.

As Bland and Altman (1986) noted, high correlation amongst measures of the same quantity does not necessarily imply agreement (as correlation can be driven by a large range in true values, for instance). Figure 3c shows Bland-Altman plots illustrating the level of agreement of each method with manual volumes. All methods show an obvious proportional bias: FreeSurfer and FSL FIRST markedly underestimate smaller hippocampi and over-estimate large hippocampi (the limits of agreement are between -2482^3mm and -784^3mm , and between -1653^3mm and 79^3mm , respectively), whereas both MAgE-T-Brain methods show a much less exaggerated, but conservative bias (limits of agreement between -67^3mm and 766^3mm when using FEP atlases, and between -333^3mm and 504^3mm when using Winterburn atlases). On average, FreeSurfer and FSL FIRST overestimate hippocampal volume by about $1600mm^3$ and $800mm^3$, respectively. In contrast, on average MAgE-T-Brain underestimates volumes by about $300mm^3$ when using FEP atlases, and by about $80mm^3$ when using Winterburn atlases.

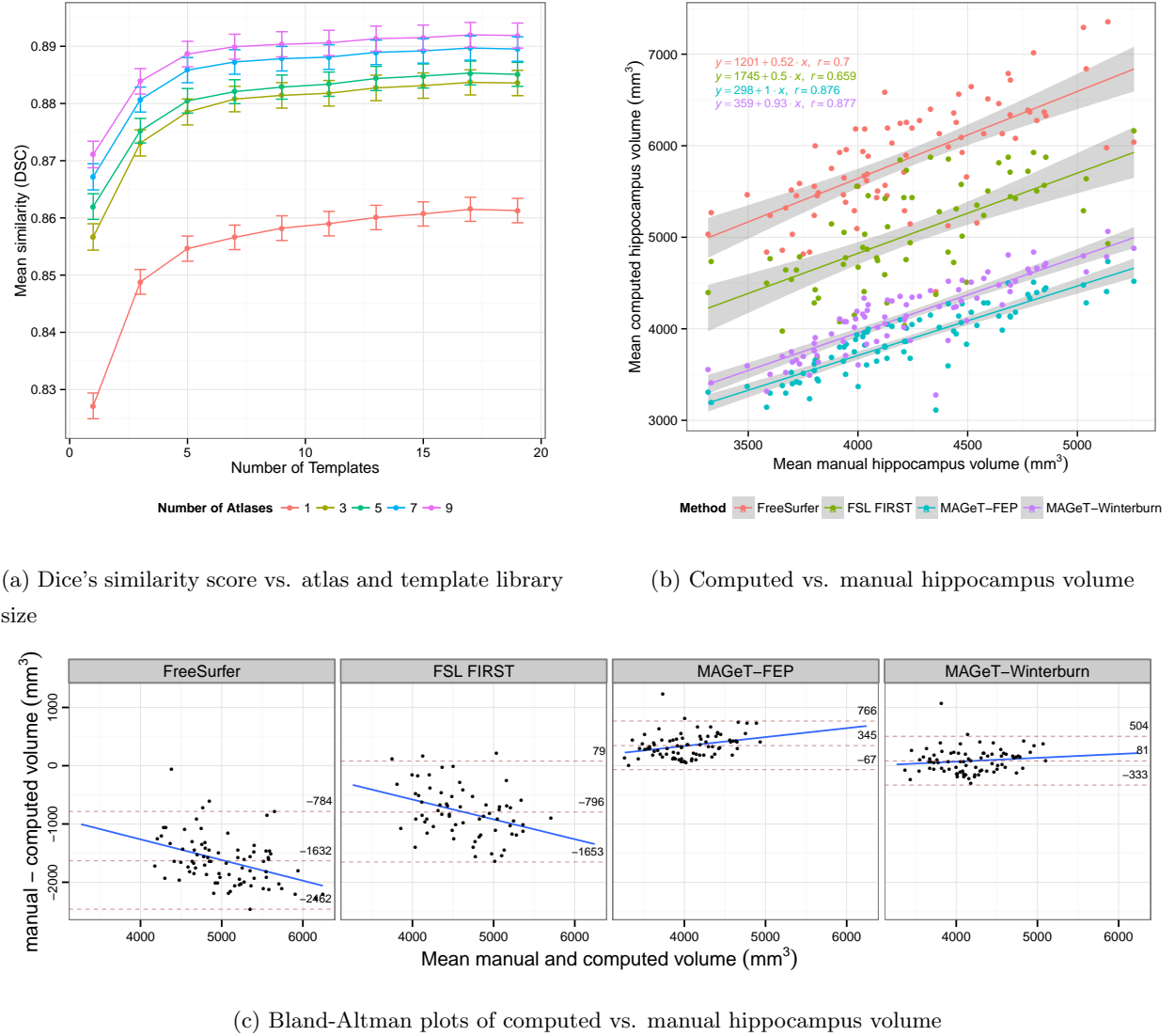


Figure 3: **First Episode Patient dataset validation.** Manual segmentation of all 81 subjects is completed using the Pruessner-protocol. MAGeT-Brain is configured using ANTS registration algorithm and majority vote label fusion. (3a) shows mean DSC overlap similarity score of MAGeT-Brain segmentations as they vary with atlas and template library size over 5-fold validation. Error bars indicate standard error. (3b) shows the volumes of the 81 subjects as determined by FSL FIRST, FreeSurfer, and MAGeT-Brain configured in two ways: with an atlas library of the five Winterburn atlases (MAGeT-Winterburn), or with an atlas library of five manually segmented FEP subjects as atlases (MAGeT-FEP). Linear fit lines are shown, with the shaded region showing standard error. (3c) shows the agreement between computed volumes and manually segmented volume. The overall mean difference in volume, and limits of agreement ($\pm 1.96SD$) are shown by dashed horizontal lines. Linear fit lines are shown for each diagnosis group. Note, points below the mean difference indicate overestimation of the volume with respect to the manual volume, and vice versa.

3.3 Experiment 3: Whole Hippocampus Segmentation Comparison — ADNI1 Complete 1Yr

To validate MAgE-T-Brain segmentation quality with respect to other established automated hippocampal segmentation methods, MAgE-T-Brain was applied to a large dataset from the ADNI project and the resulting segmentations were compared to those produced by FreeSurfer, FSL FIRST, MAPER, as well as semi-automated whole hippocampal segmentations (SNT) provided by ADNI.

3.3.1 Experiment 3: Materials and Methods

ADNI1:Complete 1Yr 1.5T dataset The *ADNI1:Complete 1Yr 1.5T* standardized dataset contains 1919 images in total. SNT, MAPER, and FreeSurfer hippocampal volumes for a subset of images were provided by ADNI, along with quality control data for each FreeSurfer segmentation (guidelines described in (Hartig et al., 2010)).

See Section 3.1.1 for study details, inclusion criteria and imaging characteristics.

Semi-automated segmentations of the left and right whole hippocampi are made available with a subset of ADNI images (Hsu et al., 2002). These labels have been generated using the SNT tool from Medtronic Surgical Navigation Technologies, Louisville, CO (see Supplementary Materials for detailed discussion of the segmentation process).

Winterburn Atlases The Winterburn atlases (Winterburn et al., 2013) are digital hippocampal segmentations of five in-vivo 0.3mm-isotropic T1-weighted MR images, described more fully in Experiment 2, section 3.2.

Experiment details MAgE-T-Brain was configured with an atlas library composed of the five Winterburn atlas images and segmentations. A template library of 19 images were randomly selected from the target dataset of ADNI subjects, and ANTS registration and majority vote label fusion were used as these were found to perform favourably in earlier experiments.

FSL FIRST segmentation was performed using the `run_first_all` script according to the FIRST user guide ⁵. All images in the ADNI1:Complete 1Yr 1.5T dataset were segmented by both methods.

One author (MP) performed visual quality inspection for MAgE-T-Brain and FSL FIRST segmentations using similar quality control guidelines (if either hippocampus was under or over segmented by 10mm or greater in three or more slices then the segmentation did not pass). Only images meeting the conditions of having segmentations from all methods (SNT, MAPER, FreeSurfer, FSL FIRST, and MAgE-T-Brain) and also passing similar quality control inspection were included in the analysis.

Evaluation method As in previous experiments, the Winterburn hippocampal segmentation protocol differs in the delineated neuroanatomical features (Winterburn et al. (2013), and Table 9, Discussion) and so we assess MAgE-T-Brain by the degree of (Pearson) correlation of average hippocampal volume across subjects. We also computed the correlation in hippocampal volume between existing, established automated segmentation methods – FSL FIRST, FreeSurfer, and MAPER, and SNT semi-automated segmentations.

⁵<http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FIRST/UserGuide>

Table 4: **ADNI1 1.5T Complete 1Yr dataset demographics.** CN - Cognitively Normal. LMCI - Late-onset Mild Cognitive Impairment. AD - Alzheimer’s Disease. CDR-SB - Clinical Dementia Rating-Sum of Boxes. ADAS - Alzheimer’s Disease Assessment Scale. MMSE - Mini-Mental State Examination.

	N	CN <i>N</i> = 584			LMCI <i>N</i> = 931			AD <i>N</i> = 404			Combined <i>N</i> = 1919		
Age at baseline Years	1919	72.4	75.8	78.5	70.5	75.1	80.4	70.1	75.3	80.2	71.1	75.3	79.8
Sex : Female	1919	48% (278)			35% (327)			49% (198)			42% (803)		
Education	1919	14 16 18			14 16 18			12 15 17			13 16 18		
CDR-SB	1911	0.0 0.0 0.0			1.0 1.5 2.5			3.5 4.5 6.0			0.0 1.5 3.0		
ADAS 13	1895	5.67	8.67	12.33	14.67	19.33	24.33	24.67	30.00	35.33	10.67	18.00	25.33
MMSE	1917	29 29 30			25 27 29			20 23 25			25 27 29		

a b c represent the lower quartile *a*, the median *b*, and the upper quartile *c* for continuous variables.

N is the number of non-missing values.

Numbers after percents are frequencies.

Table 5: **Number of segmented images and quality control failures of ADNI1:Complete 1Yr 1.5T dataset by method.**

X	SNT	MAGeT	MAPER	FSL	FS
Images	368	368	368	368	368
Failures	n/a	30	n/a	20	88

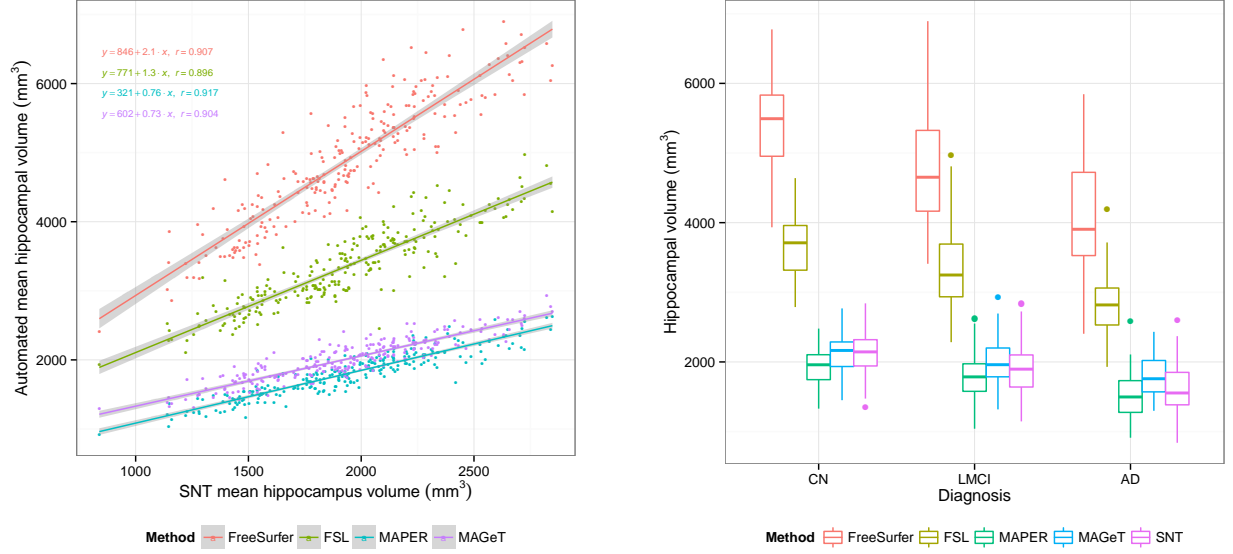
Additionally, we evaluate the volume-related fixed and proportional biases in all segmentation methods using Bland-Altman plots (Bland and Altman, 1986).

3.3.2 Experiment 3: Results

We found a close relationship in total bilateral hippocampal volume between all methods and the SNT semi-automated label volumes (Figure 4a). Volumes are correlated with Pearson $r > 0.78$ for all methods across disease categories. Within disease categories (Figure 4b), MAGeT-Brain is consistently well correlated to manual volumes (Pearson $r > 0.85$), but appears to slightly over-estimate the volume of the AD hippocampus.

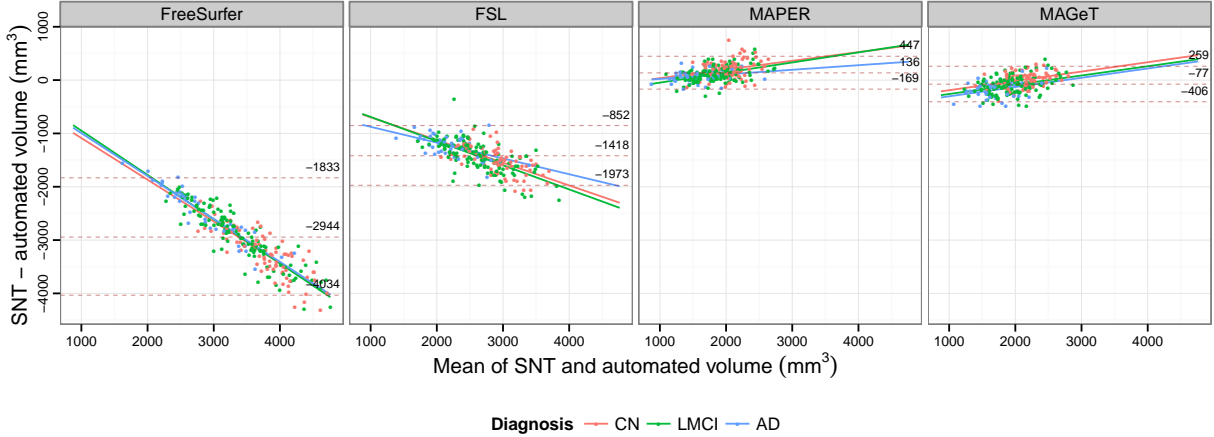
Bland-Altman plots illustrate the level of agreement of each method with SNT segmentation hippocampal volumes (Figure 4c). All methods show an obvious proportional bias: FreeSurfer and FSL FIRST markedly under-estimate smaller hippocampi and over-estimate large hippocampi, whereas MAPER and MAGeT-Brain show a reverse, conservative bias (Figure 4c). Additionally, all methods show a fixed volume bias, with FreeSurfer and FSL FIRST most dramatically over-estimating hippocampal volume by $2600mm^3$ and $2800mm^3$ on average, respectively, and MAPER and MAGeT-Brain within $250mm^3$ on average.

Figure 5 shows a qualitative comparison of MAGeT-Brain and SNT hippocampal segmentations for 10 randomly selected subjects in each disease category, and illustrates some of the common errors found during visual inspection. Mostly frequently, we found MAGeT-Brain improperly includes the vestigial hippocampal sulcus and, although not anatomically incorrect, MAGeT-Brain under-estimates the hippocampal body in comparison to the SNT segmentation.



(a) Computed vs. semi-automated (SNT) segmentation volume

(b) Hippocampal volume by diagnosis group and segmentation method



(c) Bland-Altman plots of computed vs. SNT hippocampus volume

Figure 4: **ADNI1:Complete 1Yr 1.5T dataset segmentation.** (4a) Subject mean hippocampal volume as measured by each of the four automated methods (FreeSurfer (FS), FSL FIRST, MAPER, MAGeT-Brain) versus the semi-automated SNT segmentation volumes. Linear fit lines and Pearson correlations with SNT labels are shown for each method. (4b) Mean hippocampal volume by method and disease category. AD = Alzheimer's disease, LMCI = late-onset mild cognitive impairment, and CN = cognitively normal. (4c) Bland-Altman plots comparing subject mean hippocampal volume as measured in the ADNI1:Complete 1Yr 1.5T dataset by SNT segmentation and each automated methods. The overall mean difference in volume, and limits of agreement ($\pm 1.96SD$) are shown by dashed horizontal lines. Linear fit lines are shown for each diagnosis group. Note, points below the mean difference indicate overestimation of the volume with respect to the SNT volume, and vice versa.

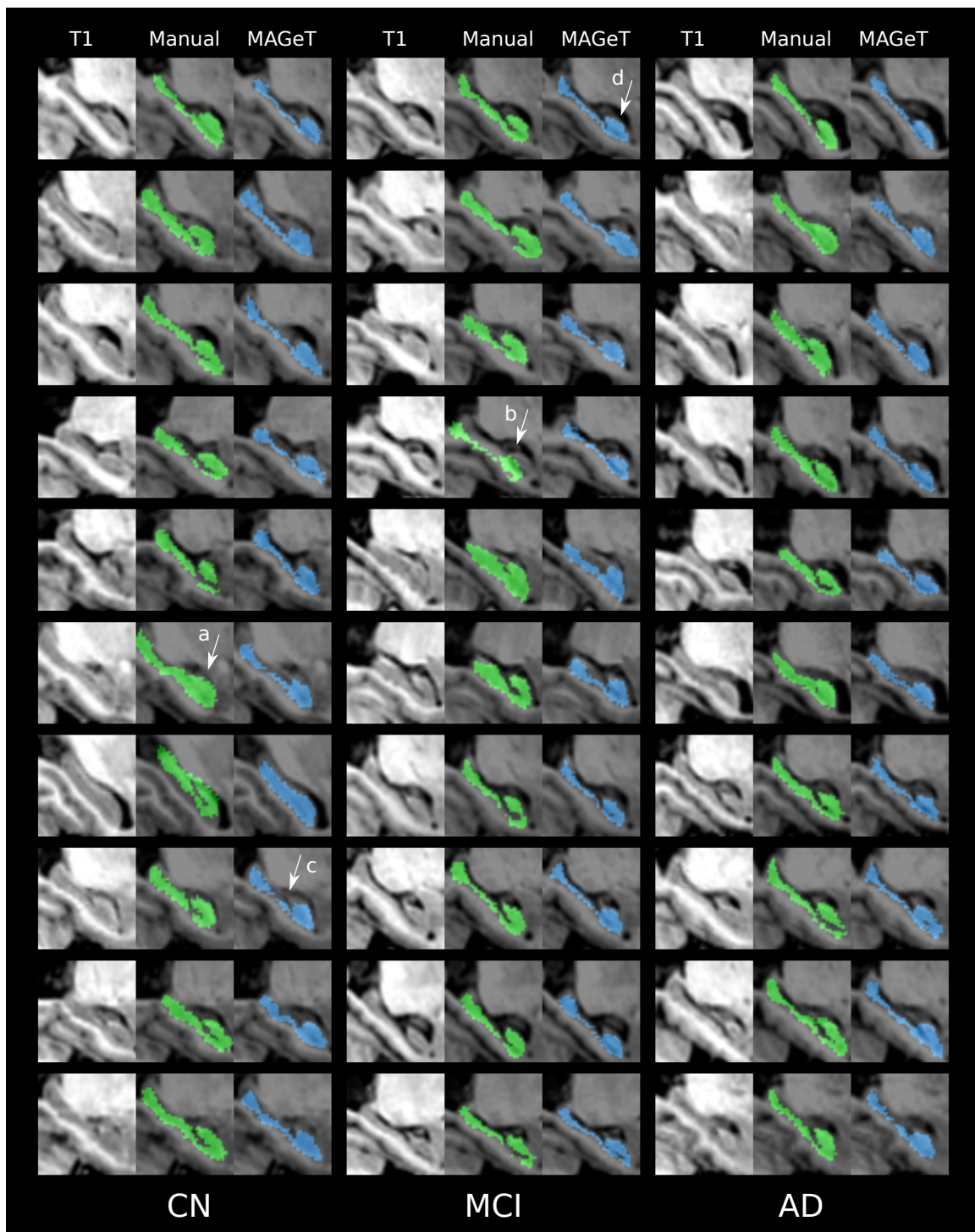


Figure 5: SNT and MAGeT-Brain segmentations for 30 ADNI subjects — 10 subjects randomly selected from each disease category in the subject pool used in Experiment 1 (Section 3.1). Sagittal slices are shown for each unlabelled T1-weighted anatomical image. SNT labels appear in green, and MAGeT-Brain labels appear in blue. Noted are examples of common segmentation idiosyncrasies: (a) over-estimation of hippocampal head and (b) translated segmentation (seen in SNT segmentations only); (c) under-estimation of hippocampal body and (d) improper inclusion of the vestigial hippocampal sulcus by MAGeT-Brain.

3.4 Experiment 4: Hippocampal Subfield Segmentation Cross-Validation

The previous experiment assesses MAgE-T-Brain performance on whole hippocampus segmentation. In this experiment, we evaluate MAgE-T-Brain hippocampal subfield segmentation of standard 3T T1-weighted images at $0.9mm^3$ -isotropic voxels. We use a modified leave-one-out cross-validation (LOOCV) design.

3.4.1 Experiment 4: Materials and Methods

Winterburn Atlases The Winterburn atlases (Winterburn et al., 2013) are digital hippocampal subfield segmentations of five in-vivo $0.3mm$ -isotropic T1-weighted MR images, described more fully in Experiment 2, section 3.2.

Healthy Control Dataset T1 MR images of 14 subjects were acquired as a part of an ongoing study at the Centre for Addiction and Mental Health (Table 6). Subjects were known to be free of neuropsychiatric disorders and gave informed consent. These images were acquired on a 3T GE Discovery MR 750 system (General Electric, Milwaukee, WI) using an 8-channel head coil with the enhanced fast gradient recalled echo 3-dimensional acquisition protocol, FGRE-BRAVO, with the following parameters: $TE/TR/TI = 3.0ms/6.7ms/650ms$, flip angle= 8° , $FOV = 15.3cm$, slice thickness= $0.9mm$, 170 in-plane steps for an approximate $0.9mm$ isotropic voxel resolution.

Experiment details Leave-one-out cross-validation (LOOCV) is a validation approach in which an algorithm is given all but one item in a dataset as training data (in our case, atlas images and labels) and then the algorithm is applied to the left-out item. This is done, in turn, for each item in the dataset and the output across all items is evaluated together.

In this experiment, the Winterburn atlases are resampled to $0.9mm$ -isotropic voxel resolution to simulate standard 3T T1-weighted resolution images. Image subsampling is performed using trilinear subsampling techniques. In each round of LOOCV, a single atlas image is selected and treated as a target image to be segmented by MAgE-T-Brain. So as to have an odd-sized atlas library, atlas image is segmented once using each possible triple of atlas images, and corresponding manual segmentations, from the remaining four unselected atlases. Thus, for each of the five atlases, a total of $\binom{3}{4} = 4$ segmentations are evaluated, resulting in a combined total of $5 \times 4 = 20$ segmentations evaluated overall. We chose an atlas library with an odd number of images so as to ensure unbiased label fusion when using majority voting (see Discussion).

The template library used has a total of 19 images composed of all five resampled atlas images plus the additional 14 images from the healthy control dataset. The ANTS registration algorithm was used for image registration, and majority voting was used for label fusion, as these methods proved most favourable in the previous whole hippocampal validation experiments.

Evaluation method Evaluating the agreement of automated hippocampal subfield segmentations with manual segmentations for T1 images at $0.9mm^3$ -isotropic voxels is inherently ill-defined since there are no manual protocols for segmentation at this resolution. Instead, we must evaluate the reliability, or *precision*, with which MAgE-T-Brain produces hippocampal subfields segmentations at this resolution that correspond in form to the segmentation protocol used by the given high-resolution atlas library images.

Table 6: **Demographics for the hippocampal subfield cross-validation healthy control subject sample used in the template library (excluding the Winterburn atlas subjects).** Education is shown in years.

	N	Control $N = 14$
Age	14	34.5 53.0 62.0
Sex : Male	14	43% (6)
Education : 12	13	15% (2)
13		8% (1)
14		23% (3)
16		15% (2)
18		38% (5)
Handedness : R	14	93% (13)

a b c represent the lower quartile a , the median b , and the upper quartile c for continuous variables.

N is the number of non-missing values.

Numbers after percents are frequencies.

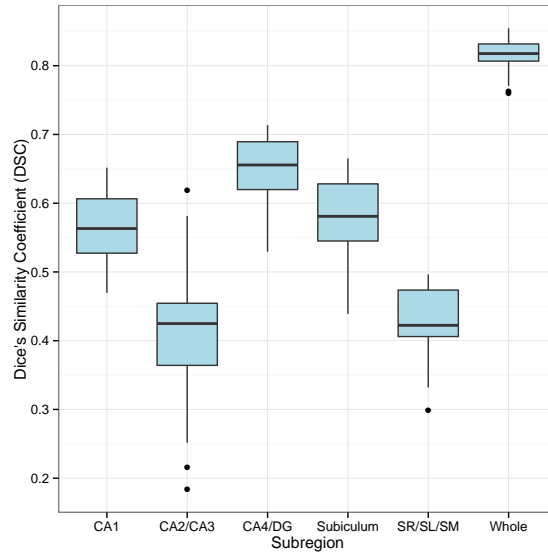
By directly resampling the Winterburn atlas segmentations to $0.9mm^3$ voxels (using standard nearest-neighbour image resampling techniques) we obtain a subsampled version of the labels which preserve the original segmentation protocol within the limits of error from rounding and interpolation. Therefore, using the resampled Winterburn segmentations as definitive for the $0.9mm^3$ resolution we evaluate reliability of MAgE-T-Brain segmentations using DSC overlap scores and evaluate consistency across the range of hippocampal sizes using Bland-Altman plots of subfield volumes.

3.4.2 Experiment 4: Results

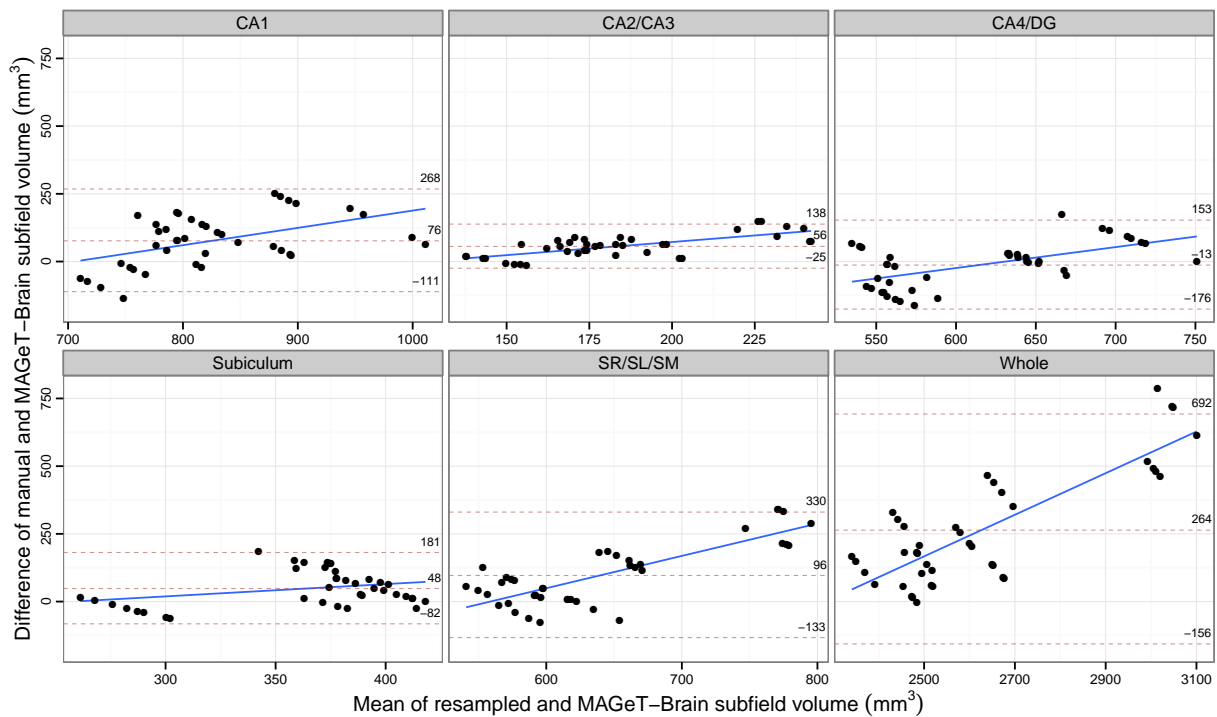
Figure 6a shows the overlap similarity scores between the MAgE-T-Brain segmentations and the resampled Winterburn atlases for each hippocampal subfield across all subjects and folds of the validation. Mean and standard deviation DSC scores of the subfields are shown in Table 7, along with DSC scores for the resampled atlas segmentations when perturbed slightly and compared to the originals. We find that the CA4/DG subfield shows the highest mean DSC score of 0.647 ± 0.051 , followed by the Subiculum and CA1 subfields having scores of 0.563 ± 0.046 and 0.58 ± 0.057 , respectively. Both the CA4/DG and molecular regions score below 0.5. These scores may seem low but not when taken in context and compared to existing (semi-)automated methods (see Discussion). The whole hippocampus is segmented with a mean DSC score of 0.816 ± 0.023 .

Figure 6b contains Bland-Altman plots comparing MAgE-T-Brain volumes with manual volumes across all validation folds. MAgE-T-Brain displays a conservative proportional bias — small hippocampi are overestimated in volume, and larger hippocampi are underestimated (a mean maximum difference of approximately $200mm^3$ across all subfields). MAgE-T-Brain display a slight conservative fixed bias, tending to underestimate all subfields except CA4/DG (mean underestimation: $CA1 = 76mm^3$, $CA2/3 = 56mm^3$, $CA4/DG = -16mm^3$, $Subiculum = 48mm^3$, $SR/SL/SM = 96mm^3$).

Figure 7 shows slices subfield segmentations for a single subject for qualitative inspection.



(a) DSC score by subfield



(b) Bland-Altman plots of computed vs. manual subfield volumes

Figure 6: **Hippocampal subfield cross-validation.** (6a) Similarity of MAGEt-Brain segmentation of subfields and the resampled Winterburn atlas segmentations at $0.9mm^3$ voxel resolution, over all validation folds. Overlap score for each hemisphere is measured separately. (6b) shows the agreement, by subfield, of computed and manual volumes across all validation folds. The overall mean difference in volume, and limits of agreement ($\pm 1.96SD$) are shown by dashed horizontal lines. Linear fit lines are shown. Note, points below the mean difference indication overestimation of the volume with respect to the resampled volume, and vice versa.

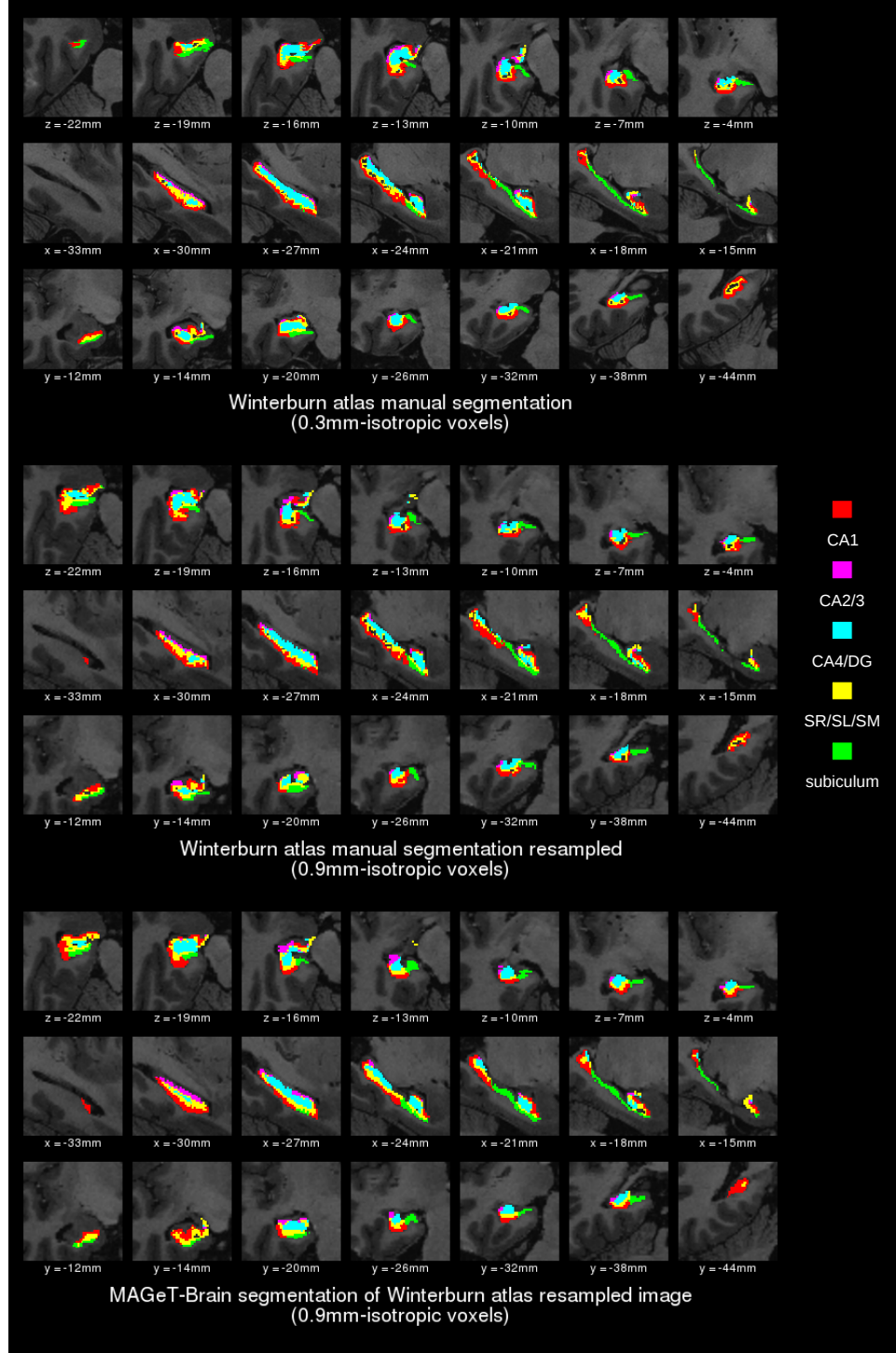


Figure 7: **Detailed subfield segmentation results for a single subject.** In the upper left corner is the original high-resolution Winterburn atlas manual subfield segmentation; in the upper right corner is the Winterburn atlas segmentation subsampled from 0.3mm- to 0.9mm-isotropic voxels; in the lower left corner is the MAgE-T-Brain segmentation of the subsampled Winterburn atlas image from a single fold of the cross-validation. In each segmentation, slices from the left hemisphere are shown in Talairach-like ICBM152 space: the first row shows axial slices from inferior to superior; the second row shows sagittal slices from lateral to medial; the third row shows coronal slices from anterior to posterior.

Table 7: Overlap similarity results for the each of the subfields of the hippocampus. Simulated overlap similarity results are also given for resampled labels that were translated by one voxel (i.e.: $0.9mm$) in all directions. Values are given as mean Dice’s Similarity Coefficient (DSC) \pm standard deviation

Subfield	MAGeT	0.9mm translation
CA1	0.56 ± 0.05	0.27 ± 0.03
CA2/CA3	0.41 ± 0.10	0.12 ± 0.05
CA4/DG	0.65 ± 0.05	0.42 ± 0.05
SR/SL/SM	0.43 ± 0.05	0.19 ± 0.04
Subiculum	0.58 ± 0.06	0.14 ± 0.04

4 Discussion

In this manuscript we have presented the implementation and validation of the MAGeT-Brain framework – a methodology that requires very few input atlases in order to provide accurate and reliable segmentations. Both Experiment 1 (Section 3.1) and Experiment 2(Section ??) compare MAGeT-Brain to basic-multi-atlas segmentation by characterising the change in segmentation quality with varying parameter settings (atlas and template library sizes, registration method, and label fusion method) and differing age and neuropsychiatric populations. Together, these experiments allow us to choose optimal MAGeT-Brain parameter settings for use in subsequent experiments. Experiment 3 (Section 3.3) demonstrates that across 246 images from the ADNI1:Complete 1Yr 1.5T dataset, MAGeT-Brain performs as well as, or better, than other established and popular methods. Finally, Experiment 4 (Section 3.4) demonstrates the reliability of MAGeT-Brain in producing subfield segmentations which match the segmentation protocol of the input atlases despite contrast and resolution limitations in standard T1-weighted image volumes. All of these experiments together demonstrate that MAGeT-Brain’s algorithmic performance is not dependent on a single definition of the hippocampus but is effective with differing hippocampal definitions (Winterburn et al., 2013; Pruessner et al., 2000; Hsu et al., 2002), across image types, and subject populations.

The core claim the MAGeT-Brain method is based on – that we can meaningfully bootstrap a template library from a small set of labelled atlas images – is validated in the cross validation conducted in Experiment 1 (and the replication in in Experiment 2 and Experiment 5, Supplementary Materials). We find that both increasing the number of atlases and the number of templates used improves MAGeT-Brain segmentation over and above basic-multi-atlas segmentations using the same number of atlas images. That is, by taking the extra step of generating a template library using target images, MAGeT-Brain is able to improve the overlap between the automatically generated segmentations and manually generated “gold standard” segmentations. The magnitude of this improvement is greatest with a small number of atlases, but even with larger atlas libraries we have found that generating a template library reduces the variability in segmentation precision (i.e. MAGeT-Brain more consistently produces high quality segmentations than does basic-multi-atlas segmentation). These effects do not appear dependant on the hippocampal segmentation protocol used.

Interestingly, previous work on multi-atlas segmentation methods (Aljabar et al., 2009; Collins and Pruessner, 2010) has found that cross-correlation and normalized mutual information-based weighted label fusion improves segmentation accuracy over simple majority vote label fusion, and yet we did not see

Table 8: Automated segmentation accuracy of the Hippocampus This table shows the best published Dice’s overlap measure between automated and ground truth segmentations. Unless otherwise specified, validation datasets are composed equally of cases and control subjects, and use manual segmentation labels as ground truth in computing DSC scores. AD = Alzheimer’s Disease; MCI = Mild Cognitive Impairment; CN = Cognitively Normal (CN); FEP = First Episode of Psychosis; LOOCV = Leave-one-out cross-validation; MCCV = Monte Carlo cross-validation; SNT = Surgical Medtronic Navigation Technologies semi-automated labels. Some studies of automated segmentation of ADNI images are excluded because they do not provide overlap measures for the hippocampus (Heckemann et al., 2011; Chupin et al., 2009).

Method	Atlases	DSC mean (AD; MCI; CN)	Reference	Validation	Dataset (Truth)
MAGeT-Brain	9	0.841		10-fold MCCV on 69 subjects	ADNI (SNT)
Patch-based label fusion	16	0.861 (0.838; —; 0.882)	Coupe et al. (2011)	LOOCV	ADNI (SNT)
Multi-atlas	20	0.848 (—; 0.798, 0.898)	Wang et al. (2011)	10-fold MCCV on 20 of 139 subjects	ADNI (SNT)
ACM (AdaBoost-based)	21	0.862	Morra et al. (2008)	LOOCV	ADNI (SNT)
LEAP	30	0.848	Wolz et al. (2010)	Segmentation of 182 subjects	ADNI (SNT)
Multi-atlas	30	0.885	Lötjönen et al. (2010)	Segmentation of 60 subjects	ADNI (SNT)
Multi-atlas (MAPS)	55	0.890	Leung et al. (2010)	Segmentation of 30 subjects (10 AD, MCI, and CN)	ADNI (SNT)
MAGeT-Brain	9	0.869		10-fold MCCV on 60 subjects	ADNI (Pruessner)
MAGeT-Brain	9	0.892		5-fold MCCV on 81 subjects	FEP subjects
Neural nets	10	0.740	Powell et al. (2008)	Segmentation of 5 subjects	controls
Probabilistic atlas	11	0.852	van der Lijn et al. (2008)	11 atlases used in 100 rounds of LOOCV on 20 elderly subjects	elderly controls
Probabilistic Atlas	16	0.860	Chupin et al. (2009)	LOOCV	AD subjects
Anatomically-guided EM	17	0.812	Pohl et al. (2007)	LOOCV on 17 controls, segmentation of 33 mixed subjects	mixed diagnosis
Multi-atlas	30	0.820	Heckemann et al. (2006a)	LOOCV	controls
Multi-atlas	30	0.880	Gousias et al. (2008)	30 adult atlas used, segmentation of 33 2yr old subjects	2yr old controls
Multi-atlas	80	0.890	Collins and Pruessner (2010)	LOOCV	controls
Multi-atlas	55	0.860	Barnes et al. (2008)	LOOCV	controls and AD
Multi-atlas	275	0.835	Aljabar et al. (2009)	LOOCV	controls

a significant indication of this effect in the MAgE-T-Brain segmentations. Selectively filtering out atlases with lower image similarity is believed to reduce sources of error from estimating deformations via non-linear registration, partial volume effects from nearest neighbour image resampling, and neuroanatomical mismatch between atlases and subjects. That MAgE-T-Brain does not see the same boost in performance from weighted voting may suggest that the neuroanatomical variability of a template library constructed from study subjects more closely matches any particular subject and thereby leaving less error to filter. From our previous work on the MAgE-T-Brain algorithm we have shown that the reduction in error is not simply a smoothing or averaging effect (Chakravarty et al., 2013).

Although, the goal of this manuscript was not to exhaustively test or validate multiple different voting strategies in the context of our segmentation algorithm, it is important to note that other strategies for voting are available. For example, other groups have used the STAPLE algorithm (Warfield et al., 2004) (or variants of the STAPLE algorithm (Robitaille and Duchesne, 2012)) which weights each segmentation based upon its estimated performance level with respect to the other available candidate segmentations. Further, the sensitivity and specificity parameters can also be tuned to potentially improve segmentation accuracy and reliability. It is likely that using more sophisticated voting methods would have a positive effect on the overall segmentation performance, as demonstrated by the STAPLE algorithm. However, it is also important to note that even in the absence of a more sophisticated label fusion algorithm, MAgE-T Brain performs reasonably well in comparison to other groups that have tested new segmentation algorithm with Alzheimer disease, mild cognitive impairment, and cognitively normal data from the ADNI database (Table 8). In addition, our validation in Experiment 2 (with the first episode psychosis subjects) yields DSC's that are amongst the highest reported. Thus, more work is required to determine the extent to which label-fusion will improve the accuracy of our algorithm.

More work is required to determine the source of the slight decrease in segmentation performance when the number of templates are set to an even number. Our initial concern was that this dip in performance was a by-product of the MAgE-T-Brain algorithm itself. However, this pattern is also found in the results of the multi-atlas segmentations we used in our experiments. We believe that our majority voting methodology is biased towards labels with the lowest numeric values when breaking ties (by way of the implementation of the `mode` function used to determine majority), thus causing the slight bias observed when using an even number of templates. This is another area where the voting scheme could be used to improve performance. However, it is worth noting that this limitation was previously identified by Heckemann et al. (2006b) and, subsequently, other groups have not even considered the potential pitfalls of an even number of candidate labels (e.g. Leung et al. (2010)).

Another concern is the moderate improvement observed in MAgE-T-Brain in comparison to multi-atlas segmentation when using the same number of atlases. The actual benefit in using MAgE-T-Brain is consistency of the labelling regardless of atlas or template choice, as mentioned above. This is an important consideration that few have touched on previously. The 10-fold Monte Carlo cross-validation that we present in Experiment 1 and Experiment 2 is amongst one of the most stringent performed in the multi-atlas segmentation literature. To the best of our knowledge, with the exception of (Wang et al., 2011), other groups using ADNI data for validation do at most a single round of leave-one-out-validation (Table 8). The thoroughness of our validation suggests that our results are reflective of a true average over the choice of parameter settings and are independent of atlas or template choice (provided the input atlases are consistently segmented).

On that note, one author (JW), an expert manual rater (Winterburn et al., 2013), identified regular in-

consistencies in the SNT segmentations: occurrences of over- and under-estimation, as well as misalignments of the entire segmentation volume (Figure 5). Although the SNT segmentations are used as benchmarks for validation in many other studies (Table 8), these segmentation inconsistencies present the possibility that a more accurate and consistent benchmark segmentation protocol ought to be used in order to truly understand the results of such validations. Indeed, our replication of the 10-fold cross-validation using SNT segmentations (Experiment 5, Supplementary Materials) shows noticeably poorer mean similarity scores for both MAGeT-Brain and multi-atlas.

Thus, in comparison to other methodologies in the field MAGeT-Brain performs favourably. Table 8 surveys some of the most recent reported DSC values reported on ADNI dataset, using SNT segmentations for the atlas library and as gold standards for evaluation. While it is difficult to compare segmentation results across studies, gold standards, evaluation metrics, and algorithms it is worth noting that the methods summarized require more atlases (between 16-55) than our MAGeT-Brain implementation with the Winterburn atlases (Winterburn et al., 2013).

There are some important differences between our method and these specific methods. Others have reported the difficulty with mis-registrations in candidate segmentation (i.e. segmentations generated that are then input in the voxel-voting procedure (Collins and Pruessner, 2010)). The work of Leung et al. (2010) tackles this problem by using an intensity threshold that is estimated heuristically at the time of segmentation (this work also reports some of the highest DSC scores for the segmentation of ADNI data). While this method is effective for the ADNI dataset (which is partially homogenized with respect to image acquisition and pre-processing), it is unclear if this type of heuristic is applicable to other datasets. In all cases, these methods require more atlases than our implementation with the Winterburn atlases. Lötjönen et al. (2010) achieve highly accurate segmentation but correct their segmentations using classifications derived using an expectation maximization framework. In their initial work, Chupin et al. (2009) develop their probabilistic methodology using a cohort of 8 healthy controls and 15 epilepsy patients, and then use this method to segment an ADNI sample, with a hierarchical experimentation protocol. These methods suggest that some post-processing of the final segmentations would improve accuracy of the segmentation. While that may be true, there is little consensus regarding how to achieve this.

To the best of our knowledge, no other groups have validated their work using multiple atlas segmentation protocols, different acquisitions, and disease populations in order to demonstrate the robustness of their technique. This is one of the clear strengths of this work. Furthermore, unlike some of the algorithms mentioned, our implementation does not require retuning for new populations or datasets as it inherently models the variability of the dataset through the template library. However it should be noted that the increased accuracy that follows increasing the number of atlases and templates comes at an increased computational cost ($O(\log(n))$), as previously mentioned in other work (Heckemann et al., 2006a).

Among the automated segmentation methods we compared in this paper (FreeSurfer, MAPER, FSL FIRST), we find extremely variable performance of all methods. With the exception of FSL FIRST all methods correlate well with the SNT volumes provided in the ADNI database. However, FreeSurfer and FSL FIRST provide radically different definitions of the size of the hippocampus in comparison to the other methods. Further, when estimating bias of these methods relative to SNT hippocampal volumes we see that large hippocampi are over estimated while small hippocampi are under estimated. By comparison, MAGeT-Brain and MAPER are far more conservative in volume estimation, suggesting these methods may be better suited for estimating true-positives, especially in neurodegenerative disease subjects featuring smaller overall

Table 9: **Summary of labelled subfields of the Hippocampus from recent MRI segmentation protocols.**

Protocol	Labelled Subfields
Winterburn et al. (2013)	CA1, CA2/CA3, CA4/dentate gyrus, strata radiatum/lacunosum/moleculare, subiculum
Wisse et al. (2012)	CA1, CA2, CA3, CA4/dentate gyrus, subiculum, entorhinal cortex
Van Leemput et al. (2009)	CA1, CA2/CA3, CA4/dentate gyrus, presubiculum, subiculum, hippocampal fissure, fimbria, hippocampal tail, inferior lateral ventricle, choroid plexus
Yushkevich et al. (2009)	CA1, CA2/CA3, dentate gyrus (hilus), dentate gyrus (stratum moleculare), strata radiatum/lacunosum/moleculare/vestigial hippocampal sulcus
Mueller et al. (2007)	CA1, CA2, CA3/CA4 & dentate gyrus, Subliculum, entorhinal cortex

hippocampi. However, in this analysis we have only compared methods by total hippocampal volume, and so more work is needed to understand the full extent to which these methods differ.

Finally, we have provided evidence that using the Winterburn high-resolution hippocampal subfield atlases (Winterburn et al., 2013) our algorithmic framework is appropriate for the segmentation of hippocampal subfields in standard T1-weighted data. Subfield segmentation is a burgeoning topic in the literature although very few automated methods are available for the segmentation of 3T data (Yushkevich et al., 2009, 2010; Van Leemput et al., 2009). Table 10 compares segmentation accuracy from some of these methods and MAgE-T-Brain. The overlap DSC scores for MAgE-T-Brain subfields are notably lower but a direct comparison of overlap values must be done cautiously. In the present work, our overlap scores are computed on $0.9mm$ -isotropic voxel resolution images, whereas Yushkevich et al. (2010) uses focal $0.4 \times 0.5 \times 2.0mm$ voxel resolution images, and Van Leemput et al. (2009) use supersampled $0.380mm$ -isotropic voxel resolution images. The larger voxel images we use necessarily entail a greater change in DSC for each incorrectly labelled voxel. In addition, our automated segmentations are compared to manual segmentations resampled from $0.3mm$ -isotropic voxel labels; the resampling process inevitably introduces noise which may lower overlap scores. Lastly, as our method is aimed specifically at situations when manually produced atlases are scarce, in our cross validation we are forced to use three rather than all five of the Winterburn atlases (which, based on our findings with whole hippocampal segmentation, would lead to improved overlap similarity). Although having more atlases would be ideal in this context, these atlases are incredibly time consuming to generate (Winterburn et al., 2013). Nevertheless, the advantage of evaluating MAgE-T-Brain on standard 3T T1-weighted resolution MR images with a publically available atlas library is that our results reflect typical usage scenarios of researchers and clinicians.

One further complication common to all subfield segmentation evaluation is that, by its nature, the Dice’s Similarity Coefficient score penalizes structures with high surface area to volume ratios. Therefore subfield DSC scores will generally be lower than whole hippocampal segmentations. We attempted to put this effect into perspective by comparing MAgE-T-Brain subfield segmentation agreement with the agreement of voxel-shifted manual segmentations (Table 7). The results of this exercise show conclusively, despite the very limited number of atlases we had to work with, that MAgE-T-Brain subfield segmentations are well within the bounds of error of a $0.3mm$ shift.

Our overlap DSC values demonstrates that we can reliably reproduce segmentations for the CA1, subiculum, and CA4/dentate subfields (DSC ≥ 0.5). That the CA2/CA3 and molecular layers are less well repro-

Table 10: **A comparison of subfield segmentation overlap similarity with manual raters.**

Subfield	MAGeT-Brain	Van Leemput et al. (2009)	Yushkevich et al. (2010)
CA1	0.563	0.62	0.875
CA2/3	0.412	0.74	$CA2 = 0.538, CA3 = 0.618$
CA4/DG	0.647	0.68	$DG = 0.873$
presubiculum	—	0.68	—
subiculum	0.58	0.74	0.770
hippocampal fissure	—	0.53	—
SR/SL/SM	0.428	—	—
fimbria	—	0.51	—
head	—	—	0.902
tail	—	—	0.863

duced ($DSC \leq 0.5$) should not be surprising as these are extremely thin and spatially convoluted regions that originally required high-resolution MRI for identification and so it is likely that the extents of these regions are well below the resolution and contrast offered by standard T1-weighted images.

This points to a larger issue of how to truly validate subfield segmentations, both in high resolution images and in standard T1-weighted images. There are several manual subfield segmentation methodologies, and they do not agree on which regions can be differentiated, even on high-resolution scans. See Table 9 for a comparison of manual subfield segmentation methodologies. A further complication is that different researchers have differing operational definitions for the subfields and how they ought to be parcellated. The disagreement in the community has led to an international working group devoted to normalizing the ontology and segmentation rules for the hippocampal subfields (<http://www.hippocampalsubfields.com/>). Both of these disagreements suggest that direct comparison of automated methods using “ground truth”-based overlap similarity metrics, such as Dice’s Similarity Coefficient, are not possible without carefully taking into account the differences in underlying segmentation protocols and image characteristics.

In conclusion, we have demonstrated the viability of leveraging a small number of input atlases to bootstrap a large template library and thereby improve segmentation reliability when using multi-atlas methods. We demonstrated that this method works robustly over hippocampal definitions, different disease populations, and different acquisition types. Finally, we also demonstrate that accurate identification of the hippocampal subfields in standard 3T T1-weighted images is possible.

5 Acknowledgements

We wish acknowledge support from the CAMH Foundation, thanks to Michael and Sonja Koerner, the Kimel Family, and the Paul E. Garfinkel New Investigator Catalyst Award. MMC is funded by the W. Garfield Weston Foundation and ANV is funded by the Canadian Institutes of Health Research, Ontario Mental Health Foundation, NARSAD, and the National Institute of Mental Health (R01MH099167).

Computations were performed on the gpc supercomputer at the SciNet HPC Consortium Loken et al. (2010). SciNet is funded by: the Canada Foundation for Innovation under the auspices of Compute Canada; the Government of Ontario; Ontario Research Fund - Research Excellence; and the University of Toronto.

In addition, computations were performed on the CAMH Specialized Computing Cluster. The SCC is funded by: The Canada Foundation for Innovation, Research Hospital Fund.

ADNI Acknowledgements: Data collection and sharing for this project was funded by the Alzheimer’s Disease Neuroimaging Initiative (ADNI) (National Institutes of Health Grant U01 AG024904). ADNI is funded by the National Institute on Aging, the National Institute of Biomedical Imaging and Bioengineering, and through generous contributions from the following: Abbott; Alzheimer’s Association; Alzheimer’s Drug Discovery Foundation; Amorphix Life Sciences Ltd.; AstraZeneca; Bayer HealthCare; BioClinica, Inc.; Biogen Idec Inc.; Bristol-Myers Squibb Company; Eisai Inc.; Elan Pharmaceuticals Inc.; Eli Lilly and Company; F. Hoffmann-La Roche Ltd and its affiliated company Genentech, Inc.; GE Healthcare; Innogenetics, N.V.; IXICO Ltd.; Janssen Alzheimer Immunotherapy Research Development, LLC.; Johnson & Johnson Pharmaceutical Research Development LLC.; Medpace, Inc.; Merck & Co., Inc.; Meso Scale Diagnostics, LLC.; Novartis Pharmaceuticals Corporation; Pfizer Inc.; Servier; Synarc Inc.; and Takeda Pharmaceutical Company. 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). The grantee organization is the Northern California Institute for Research and Education, and the study is Rev March 26, 2012 coordinated by the Alzheimer’s disease Cooperative Study at the University of California, San Diego. ADNI data are disseminated by the Laboratory for NeuroImaging at the University of California, Los Angeles. This research was also supported by NIH grants P30 AG010129 and K01 AG030514.

We would also like to thank G. Clinton, E. Hazel, and B. Worrell for inspiring this work.

6 Supplementary Materials

6.1 SNT Hippocampal Labels

Semi-automated hippocampal volumetry was carried out using a commercially available high dimensional brain mapping tool (Medtronic Surgical Navigation Technologies, Louisville, CO), that has previously been validated and compared to manual tracing of the hippocampus (Hsu et al., 2002). Measurement of hippocampal volume is achieved first by placing manually 22 control points as local landmarks for the hippocampus on the individual brain MRI data: one landmark at the hippocampal head, one at the tail, and four per image (i.e., at the superior, inferior, medial and lateral boundaries) on five equally spaced images perpendicular to the long axis of the hippocampus. Second, fluid image transformation is used to match the individual brains to a template brain (Christensen et al., 1997). The pixels corresponding to the hippocampus are then labeled and counted to obtain volumes. This method of hippocampal voluming has a documented reliability of an intraclass coefficient better than .94 (Hsu et al., 2002).

6.2 Experiment 5: Whole Hippocampus Segmentation Cross-Validation — Alzheimer’s Disease, SNT Segmentations

This experiment is a replication of Experiment 1 using a pool of 69 images and SNT semi-automated segmentations from the ADNI dataset (Hsu et al., 2002). See Experiment 1 for full details on the ADNI dataset, and validation process.

Table 11: **ADNI1:Complete 1Yr 1.5T SNT cross-validation subset demographics.** CN - Cognitively Normal. LMCI - Late-onset Mild Cognitive Impairment. AD - Alzheimer’s Disease. Hisp - Hispanic. CDR-SB - Clinical Dementia Rating-Sum of Boxes. ADAS - Alzheimer’s Disease Assessment Scale. MMSE - Mini-Mental State Examination.

	CN			LMCI			AD			Combined		
	<i>N</i> = 23			<i>N</i> = 23			<i>N</i> = 23			<i>N</i> = 69		
Age at baseline Years	72.2	75.5	78.5	71.0	77.1	81.4	71.7	77.8	81.8	71.5	76.6	81.3
Sex : Female	43% (10)			43% (10)			43% (10)			43% (30)		
Education	16.0	16.0	18.0	15.0	16.0	18.0	12.0	16.0	16.5	14.0	16.0	18.0
CDR-SB	0.00	0.00	0.00	0.75	1.50	1.50	4.00	4.50	5.00	0.00	1.50	4.00
ADAS 13	4.67	5.67	12.34	14.34	16.00	20.50	23.83	29.00	31.66	10.00	16.00	25.33
MMSE	28.5	29.0	30.0	25.0	27.0	28.0	21.0	23.0	24.0	24.0	27.0	29.0

a b c represent the lower quartile *a*, the median *b*, and the upper quartile *c* for continuous variables.

Numbers after percents are frequencies.

6.2.1 Experiment 5: Materials and Methods

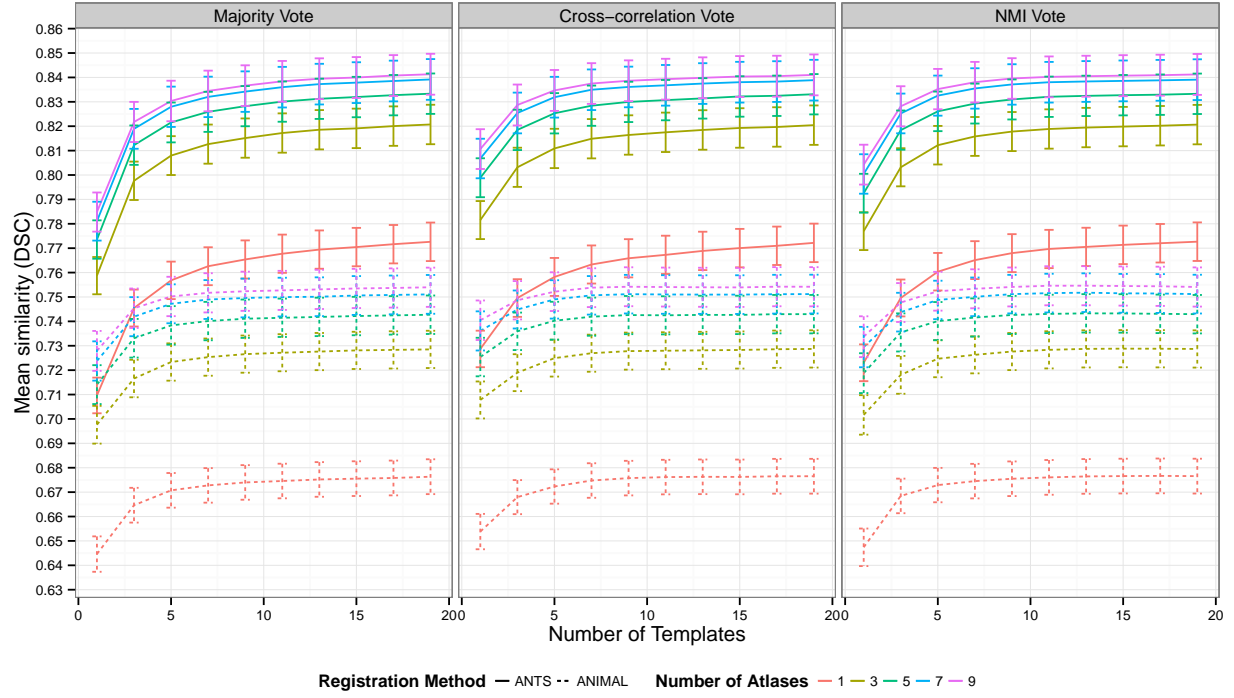
Dataset 69 1.5T images were arbitrarily selected from the baseline scans in the *ADNI1:Complete 1Yr 1.5T* standardized dataset. 23 subjects were chosen from each disease category: cognitively normal (CN), mild cognitive impairment (MCI) and Alzheimer’s disease (AD). Demographics for this subset are shown in Table 1. Each image has a corresponding semi-automated segmentation of the left and right whole hippocampus made available with ADNI images (SNT; see Supplementary Materials).

Experiment details A total of ten validation rounds were performed on each subject in the dataset, for each combination of parameter settings: atlas library size (1-9), template library size (1-20), registration method (ANTS or ANIMAL), and label fusion method (majority vote, cross-correlation weighted majority vote, and normalized mutual information weighted majority vote). A total of $10 \times 69 \times 9 \times 20 \times 2 \times 3 = 7.452 \times 10^5$ validation rounds are conducted. The computed segmentations for a subject are compared to the SNT labels provided by ADNI using Dice’s Similarity Coefficient and the score is averaged over the validation rounds.

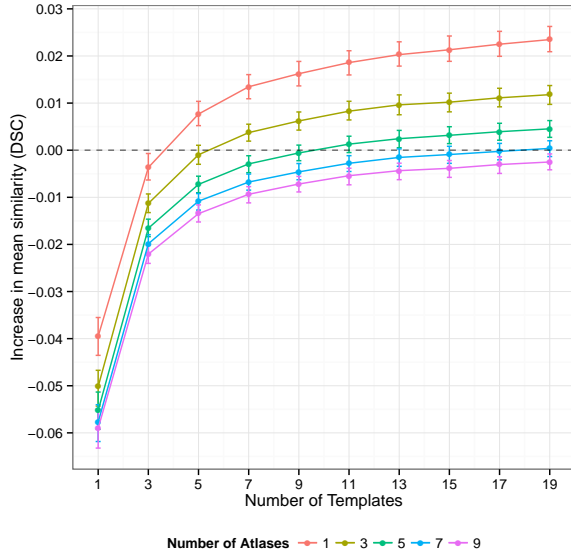
6.2.2 Experiment 5: Results

As when comparing against manual labels in Experiment 1, we find similar behaviour when comparing MAGeT-Brain segmentations to SNT labels: similarity scores increase with increasing numbers of atlases and templates, with diminishing increases in improvement trending towards a plateau (Figure 2a). As in Experiment 1, using ANTS registration leads to significantly increased similarity scores, and there is no significant difference in scores from any of the label fusion methods. Mean DSC score peaks at 0.841 when using 9 atlases, 19 templates, ANTS registration, and majority vote label fusion. Compared to multi-atlas segmentations, we find MAGeT-Brain segmentations show increasing improvement with larger atlas and template libraries when using more than 9 templates and 5 or fewer atlases (Figure 8b). Peak improvement (+0.023 DSC) is found with a single atlas and template library of 19 images. In addition to a mean increase in similarity score over multi-atlas-based segmentation, MAGeT-Brain also shows more consistency in similarity scores across all subjects and validation folds (Figure ??) with a large enough template library.

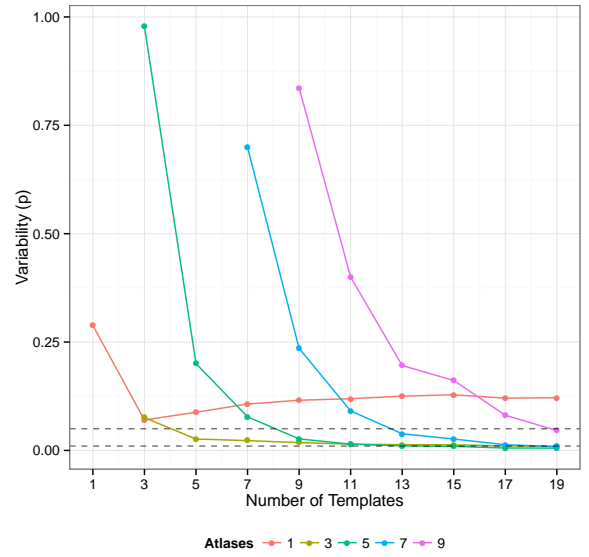
s



(a) DSC vs. atlas and template library size



(b) Increase in similarity score over multi-atlas



(c) Difference in variability with multi-atlas

Figure 8: **Whole hippocampus segmentation cross-validation on ADNI subjects with SNT segmentations.** (8a) Average DSC score of MAGeT-Brain with SNT segmentations for 69 ADNI subjects taken over 10 folds of cross-validation at each parameter setting. Error bars indicate standard error. (8b) Increase in DSC of MAGeT-Brain over multi-atlas segmentations. (8c) shows the significance of t-tests comparing the variability in DSC scores of MAGeT-Brain and multi-atlas across validation folds. Only points where MAGeT-Brain mean variability is lower than multi-atlas are shown. Dashed lines indicate p-values of 0.05 and 0.01.

References

- P. Aljabar, R. a. Heckemann, a. Hammers, J. V. Hajnal, and D. Rueckert. Multi-atlas based segmentation of brain images: atlas selection and its effect on accuracy. *NeuroImage*, 46(3):726–38, July 2009. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2009.02.018.
- B. B. Avants, C. L. Epstein, M. Grossman, and J. C. Gee. Symmetric diffeomorphic image registration with cross-correlation: evaluating automated labeling of elderly and neurodegenerative brain. *Medical image analysis*, 12(1):26–41, Feb. 2008. ISSN 1361-8423. doi: 10.1016/j.media.2007.06.004.
- J. Barnes, J. Foster, R. G. Boyes, T. Pepple, E. K. Moore, J. M. Schott, C. Frost, R. I. Scahill, and N. C. Fox. A comparison of methods for the automated calculation of volumes and atrophy rates in the hippocampus. *NeuroImage*, 40(4):1655–71, May 2008. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2008.01.012.
- J. M. Bland and D. Altman. Statistical methods for assessing agreement between two methods of clinical measurement. *The lancet*, pages 307–310, 1986.
- M. Boccardi, M. Bocchetta, L. G. Apostolova, G. Preboske, N. Robitaille, P. Pasqualetti, L. D. Collins, S. Duchesne, C. R. Jack, and G. B. Frisoni. Establishing Magnetic Resonance Images Orientation for the EADC-ADNI Manual Hippocampal Segmentation Protocol. *Journal of neuroimaging : official journal of the American Society of Neuroimaging*, pages 1–6, Nov. 2013a. ISSN 1552-6569. doi: 10.1111/jon.12065.
- M. Boccardi, M. Bocchetta, R. Ganzola, N. Robitaille, A. Redolfi, S. Duchesne, C. R. Jack, and G. B. Frisoni. Operationalizing protocol differences for EADC-ADNI manual hippocampal segmentation. *Alzheimer’s & dementia : the journal of the Alzheimer’s Association*, pages 1–11, May 2013b. ISSN 1552-5279. doi: 10.1016/j.jalz.2013.03.001.
- M. M. Chakravarty, A. F. Sadikot, S. Mongia, G. Bertrand, and D. L. Collins. Towards a multi-modal atlas for neurosurgical planning. *Medical image computing and computer-assisted intervention : MICCAI ... International Conference on Medical Image Computing and Computer-Assisted Intervention*, 9(Pt 2): 389–96, Jan. 2006.
- M. M. Chakravarty, A. F. Sadikot, J. Germann, G. Bertrand, and D. L. Collins. Towards a validation of atlas warping techniques. *Medical image analysis*, 12(6):713–26, Dec. 2008. ISSN 1361-8423. doi: 10.1016/j.media.2008.04.003.
- M. M. Chakravarty, A. F. Sadikot, J. Germann, P. Hellier, G. Bertrand, and D. L. Collins. Comparison of piece-wise linear, linear, and nonlinear atlas-to-patient warping techniques: analysis of the labeling of subcortical nuclei for functional neurosurgical applications. *Human brain mapping*, 30(11):3574–95, Nov. 2009. ISSN 1097-0193. doi: 10.1002/hbm.20780.
- M. M. Chakravarty, P. Steadman, M. C. van Eede, R. D. Calcott, V. Gu, P. Shaw, A. Raznahan, D. L. Collins, and J. P. Lerch. Performing label-fusion-based segmentation using multiple automatically generated templates. *Human brain mapping*, 34(10):2635–54, Oct. 2013. ISSN 1097-0193. doi: 10.1002/hbm.22092.
- G. E. Christensen, S. C. Joshi, and M. I. Miller. Volumetric transformation of brain anatomy. *IEEE transactions on medical imaging*, 16(6):864–77, Dec. 1997. ISSN 0278-0062. doi: 10.1109/42.650882.

- M. Chupin, E. Gérardin, R. Cuingnet, C. Boutet, L. Lemieux, S. Lehericy, H. Benali, L. Garnero, and O. Colliot. Fully automatic hippocampus segmentation and classification in Alzheimer’s disease and mild cognitive impairment applied on data from ADNI. *Hippocampus*, 19(6):579–87, June 2009. ISSN 1098-1063. doi: 10.1002/hipo.20626.
- D. L. Collins and J. C. Pruessner. Towards accurate, automatic segmentation of the hippocampus and amygdala from MRI by augmenting ANIMAL with a template library and label fusion. *NeuroImage*, 52(4):1355–66, Oct. 2010. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2010.04.193.
- D. L. Collins, P. Neelin, T. M. Peters, and A. C. Evans. Automatic 3D intersubject registration of MR volumetric data in standardized Talairach space. *Journal of computer assisted tomography*, 18(2):192–205, 1994. ISSN 0363-8715.
- D. L. Collins, C. J. Holmes, T. M. Peters, and A. C. Evans. Automatic 3-D model-based neuroanatomical segmentation. *Human Brain Mapping*, 3(3):190–208, Oct. 1995. ISSN 10659471. doi: 10.1002/hbm.460030304.
- P. Coupe, V. Fonov, S. Eskildsen, J. Manjón, D. Arnold, and L. Collins. Influence of the training library composition on a patch-based label fusion method: Application to hippocampus segmentation on the ADNI dataset. *Alzheimer’s & Dementia*, 7(4):S316, July 2011. ISSN 15525260. doi: 10.1016/j.jalz.2011.05.918.
- P. Coupé, S. F. Eskildsen, J. V. Manjón, V. S. Fonov, and D. L. Collins. Simultaneous segmentation and grading of anatomical structures for patient’s classification: application to Alzheimer’s disease. *NeuroImage*, 59(4):3736–47, Feb. 2012. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2011.10.080.
- J. G. Csernansky, S. Joshi, L. Wang, J. W. Haller, M. Gado, J. P. Miller, U. Grenander, and M. I. Miller. Hippocampal morphometry in schizophrenia by high dimensional brain mapping. *Proceedings of the National Academy of Sciences of the United States of America*, 95(19):11406–11411, 1998.
- T. den Heijer, F. V. der Lijn, M. W. Vernooij, M. de Groot, P. J. Koudstaal, a. V. der Lugt, G. P. Krestin, a. Hofman, W. J. Niessen, and M. M. B. Breteler. Structural and diffusion MRI measures of the hippocampus and memory performance. *NeuroImage*, 63(4):1782–9, Dec. 2012. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2012.08.067.
- B. Fischl, D. H. Salat, E. Busa, M. Albert, M. Dieterich, C. Haselgrove, A. van der Kouwe, R. Killiany, D. Kennedy, S. Klaveness, A. Montillo, N. Makris, B. Rosen, and A. M. Dale. Whole brain segmentation: automated labeling of neuroanatomical structures in the human brain. *Neuron*, 33(3):341–55, Jan. 2002. ISSN 0896-6273.
- E. Geuze, E. Vermetten, and J. D. Bremner. MR-based in vivo hippocampal volumetrics: 2. Findings in neuropsychiatric disorders. *Molecular Psychiatry*, 10(2):160, Sept. 2004. doi: 10.1038/sj.mp.4001579.
- I. S. Gousias, D. Rueckert, R. a. Heckemann, L. E. Dyet, J. P. Boardman, a. D. Edwards, and A. Hammers. Automatic segmentation of brain MRIs of 2-year-olds into 83 regions of interest. *NeuroImage*, 40(2):672–84, Apr. 2008. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2007.11.034.

- J. W. Haller, A. Banerjee, G. E. Christensen, M. Gado, S. Joshi, M. I. Miller, Y. Sheline, M. W. Vanier, and J. G. Csernansky. Three-dimensional hippocampal MR morphometry with high-dimensional transformation of a neuroanatomic atlas. *Radiology*, 202(2):504–510, 1997.
- M. Hartig, D. Truran-sacrey, S. Raptentsetsang, N. Schuff, and M. Weiner. USCF FreeSurfer Overview and QC Ratings. 2010.
- R. A. Heckemann, J. V. Hajnal, P. Aljabar, D. Rueckert, and A. Hammers. Automatic anatomical brain MRI segmentation combining label propagation and decision fusion. *NeuroImage*, 46(3):726–38, July 2006a. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2009.02.018.
- R. A. Heckemann, J. V. Hajnal, P. Aljabar, D. Rueckert, and A. Hammers. Automatic anatomical brain MRI segmentation combining label propagation and decision fusion. *NeuroImage*, 33(1):115–26, Oct. 2006b. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2006.05.061.
- R. A. Heckemann, S. Keihaninejad, P. Aljabar, K. R. Gray, C. Nielsen, D. Rueckert, J. V. Hajnal, and A. Hammers. Automatic morphometry in Alzheimer’s disease and mild cognitive impairment. *NeuroImage*, 56(4):2024–37, July 2011. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2011.03.014.
- Y.-Y. Hsu, N. Schuff, A.-T. Du, K. Mark, X. Zhu, D. Hardin, and M. W. Weiner. Comparison of automated and manual MRI volumetry of hippocampus in normal aging and dementia. *Journal of magnetic resonance imaging : JMRI*, 16(3):305–10, Sept. 2002. ISSN 1053-1807. doi: 10.1002/jmri.10163.
- C. R. Jack, M. A. Bernstein, N. C. Fox, P. Thompson, G. Alexander, D. Harvey, B. Borowski, P. J. Britson, J. L. Whitwell, C. Ward, A. M. Dale, J. P. Felmlee, J. L. Gunter, D. L. G. Hill, R. Killiany, N. Schuff, S. Fox-Bosetti, C. Lin, C. Studholme, C. S. DeCarli, G. Krueger, H. A. Ward, G. J. Metzger, K. T. Scott, R. Mallozzi, D. Blezek, J. Levy, J. P. Debbins, A. S. Fleisher, M. Albert, R. Green, G. Bartzokis, G. Glover, J. Mugler, and M. W. Weiner. The Alzheimer’s Disease Neuroimaging Initiative (ADNI): MRI methods. *Journal of magnetic resonance imaging : JMRI*, 27(4):685–91, Apr. 2008. ISSN 1053-1807. doi: 10.1002/jmri.21049.
- C. R. Jack, F. Barkhof, M. A. Bernstein, M. Cantillon, P. E. Cole, C. Decarli, B. Dubois, S. Duchesne, N. C. Fox, G. B. Frisoni, H. Hampel, D. L. G. Hill, K. Johnson, J.-F. Mangin, P. Scheltens, A. J. Schwarz, R. Sperling, J. Suhy, P. M. Thompson, M. Weiner, and N. L. Foster. Steps to standardization and validation of hippocampal volumetry as a biomarker in clinical trials and diagnostic criterion for Alzheimer’s disease. *Alzheimer’s & dementia : the journal of the Alzheimer’s Association*, 7(4):474–485.e4, July 2011. ISSN 1552-5279. doi: 10.1016/j.jalz.2011.04.007.
- A. Jeneson and L. Squire. Working memory, long-term memory, and medial temporal lobe function. *Learning & Memory*, 19(1):15–25, 2012. doi: 10.1101/lm.024018.111.
- M. S. Karnik-Henry, L. Wang, D. M. Barch, M. P. Harms, C. Campanella, and J. G. Csernansky. Medial temporal lobe structure and cognition in individuals with schizophrenia and in their non-psychotic siblings. *Schizophrenia research*, 138(2-3):128–35, July 2012. ISSN 1573-2509. doi: 10.1016/j.schres.2012.03.015.
- K. K. Leung, J. Barnes, G. R. Ridgway, J. W. Bartlett, M. J. Clarkson, K. Macdonald, N. Schuff, N. C. Fox, and S. Ourselin. Automated cross-sectional and longitudinal hippocampal volume measurement in mild

- cognitive impairment and Alzheimer's disease. *NeuroImage*, 51(4):1345–59, July 2010. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2010.03.018.
- C. Loken, D. Gruner, L. Groer, R. Peltier, N. Bunn, M. Craig, T. Henriques, J. Dempsey, C.-H. Yu, J. Chen, L. J. Dursi, J. Chong, S. Northrup, J. Pinto, N. Knecht, and R. V. Zon. SciNet: Lessons Learned from Building a Power-efficient Top-20 System and Data Centre. *Journal of Physics: Conference Series*, 256: 012026, Nov. 2010. ISSN 1742-6596. doi: 10.1088/1742-6596/256/1/012026.
- J. M. Lötjönen, R. Wolz, J. R. Koikkalainen, L. Thurfjell, G. Waldemar, H. Soininen, and D. Rueckert. Fast and robust multi-atlas segmentation of brain magnetic resonance images. *NeuroImage*, 49(3):2352–65, Mar. 2010. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2009.10.026.
- A. Malla, R. Norman, T. McLean, D. Scholten, and L. Townsend. A Canadian programme for early intervention in non-affective psychotic disorders. *The Australian and New Zealand journal of psychiatry*, 37(4):407–13, Aug. 2003. ISSN 0004-8674.
- J. Mazziotta, A. Toga, A. Evans, P. Fox, J. Lancaster, K. Zilles, R. Woods, T. Paus, G. Simpson, B. Pike, C. Holmes, L. Collins, P. Thompson, D. MacDonald, M. Iacoboni, T. Schormann, K. Amunts, N. Palomero-Gallagher, S. Geyer, L. Parsons, K. Narr, N. Kabani, G. Le Goualher, J. Feidler, K. Smith, D. Boomsma, H. Hulshoff Pol, T. Cannon, R. Kawashima, and B. Mazoyer. A four-dimensional probabilistic atlas of the human brain. *Journal of the American Medical Informatics Association : JAMIA*, 8(5):401–30. ISSN 1067-5027.
- J. Mazziotta, A. Toga, A. Evans, P. Fox, J. Lancaster, K. Zilles, R. Woods, T. Paus, G. Simpson, B. Pike, C. Holmes, L. Collins, P. Thompson, D. MacDonald, M. Iacoboni, T. Schormann, K. Amunts, N. Palomero-Gallagher, S. Geyer, L. Parsons, K. Narr, N. Kabani, G. Le Goualher, D. Boomsma, T. Cannon, R. Kawashima, and B. Mazoyer. A probabilistic atlas and reference system for the human brain: International Consortium for Brain Mapping (ICBM). *Philosophical transactions of the Royal Society of London. Series B, Biological sciences*, 356(1412):1293–322, Aug. 2001. ISSN 0962-8436. doi: 10.1098/rstb.2001.0915.
- J. C. Mazziotta, A. W. Toga, A. Evans, P. Fox, and J. Lancaster. A probabilistic atlas of the human brain: theory and rationale for its development. The International Consortium for Brain Mapping (ICBM). *NeuroImage*, 2(2):89–101, June 1995. ISSN 1053-8119.
- J. H. Morra, Z. Tu, L. G. Apostolova, A. E. Green, C. Avedissian, S. K. Madsen, N. Parikshak, X. Hua, A. W. Toga, C. R. Jack, M. W. Weiner, and P. M. Thompson. Validation of a fully automated 3D hippocampal segmentation method using subjects with Alzheimer's disease mild cognitive impairment, and elderly controls. *NeuroImage*, 43(1):59–68, Oct. 2008. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2008.07.003.
- S. Mueller, L. Stables, A. Du, and N. Schuff. Measurement of hippocampal subfields and age-related changes with high resolution MRI at 4T. *Neurobiology of ...*, 1(5):719–726, 2007. doi: 10.1016/j.neurobiolaging.2006.03.007.
- S. G. Mueller and M. W. Weiner. Selective effect of age, Apo e4, and Alzheimer's disease on hippocampal subfields. *Hippocampus*, 19(6):558–64, June 2009. ISSN 1098-1063. doi: 10.1002/hipo.20614.

- K. L. Narr, P. M. Thompson, P. Szeszko, D. Robinson, S. Jang, R. P. Woods, S. Kim, K. M. Hayashi, D. Asuncion, A. W. Toga, and R. M. Bilder. Regional specificity of hippocampal volume reductions in first-episode schizophrenia. *NeuroImage*, 21(4):1563–75, Apr. 2004. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2003.11.011.
- S. M. Nestor, E. Gibson, F.-Q. Gao, A. Kiss, and S. E. Black. A Direct Morphometric Comparison of Five Labeling Protocols for Multi-Atlas Driven Automatic Segmentation of the Hippocampus in Alzheimer’s Disease. *NeuroImage*, Nov. 2012. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2012.10.081.
- Z. Pausova, T. Paus, M. Abrahamowicz, J. Almerigi, N. Arbour, M. Bernard, D. Gaudet, P. Hanzalek, P. Hamet, A. C. Evans, M. Kramer, L. Laberge, S. M. Leal, G. Leonard, J. Lerner, R. M. Lerner, J. Mathieu, M. Perron, B. Pike, A. Pitiot, L. Richer, J. R. Séguin, C. Syme, R. Toro, R. E. Tremblay, S. Veillette, and K. Watkins. Genes, maternal smoking, and the offspring brain and body during adolescence: design of the Saguenay Youth Study. *Human brain mapping*, 28(6):502–18, June 2007. ISSN 1065-9471. doi: 10.1002/hbm.20402.
- K. M. Pohl, S. Bouix, M. Nakamura, T. Rohlfing, R. W. McCarley, R. Kikinis, W. E. L. Grimson, M. E. Shenton, and W. M. Wells. A hierarchical algorithm for MR brain image parcellation. *IEEE transactions on medical imaging*, 26(9):1201–12, Sept. 2007. ISSN 0278-0062. doi: 10.1109/TMI.2007.901433.
- J. Poppenk and M. Moscovitch. A Hippocampal Marker of Recollection Memory Ability among Healthy Young Adults: Contributions of Posterior and Anterior Segments. *Neuron*, 72(6):931–937, Dec. 2011. ISSN 0896-6273. doi: 10.1016/j.neuron.2011.10.014.
- S. Powell, V. A. Magnotta, H. Johnson, V. K. Jammalamadaka, R. Pierson, and N. C. Andreasen. Registration and machine learning-based automated segmentation of subcortical and cerebellar brain structures. *NeuroImage*, 39(1):238–47, Jan. 2008. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2007.05.063.
- J. C. Pruessner, L. M. Li, W. Serles, M. Pruessner, D. L. Collins, N. Kabani, S. Lupien, and A. C. Evans. Volumetry of hippocampus and amygdala with high-resolution MRI and three-dimensional analysis software: minimizing the discrepancies between laboratories. *Cerebral cortex (New York, N.Y. : 1991)*, 10(4):433–42, Apr. 2000. ISSN 1047-3211.
- S. Robbins, A. C. Evans, D. L. Collins, and S. Whitesides. Tuning and comparing spatial normalization methods. *Medical image analysis*, 8(3):311–23, Sept. 2004. ISSN 1361-8415. doi: 10.1016/j.media.2004.06.009.
- N. Robitaille and S. Duchesne. Label fusion strategy selection. *International journal of biomedical imaging*, 2012:431095, Jan. 2012. ISSN 1687-4196. doi: 10.1155/2012/431095.
- M. R. Sabuncu, R. S. Desikan, J. Sepulcre, B. T. T. Yeo, H. Liu, N. J. Schmansky, M. Reuter, M. W. Weiner, R. L. Buckner, R. a. Sperling, and B. Fischl. The dynamics of cortical and hippocampal atrophy in Alzheimer disease. *Archives of neurology*, 68(8):1040–8, Aug. 2011. ISSN 1538-3687. doi: 10.1001/archneurol.2011.167.
- W. B. Scoville and B. Milner. Loss of recent memory after bilateral hippocampal lesions. 1957. *The Journal of neuropsychiatry and clinical neurosciences*, 12(1):103–113, 2000.

- J. Shao. Linear Model Selection by Cross-validation. *Journal of the American Statistical Association*, 88 (422):486–494, June 1993. ISSN 0162-1459. doi: 10.1080/01621459.1993.10476299.
- J. G. Sled, a. P. Zijdenbos, and a. C. Evans. A nonparametric method for automatic correction of intensity nonuniformity in MRI data. *IEEE transactions on medical imaging*, 17(1):87–97, Feb. 1998. ISSN 0278-0062. doi: 10.1109/42.668698.
- C. Studholme, D. Hill, and D. Hawkes. An overlap invariant entropy measure of 3D medical image alignment. *Pattern Recognition*, 32(1):71–86, Jan. 1999. ISSN 00313203. doi: 10.1016/S0031-3203(98)00091-0.
- C. Studholme, E. Novotny, I. G. Zubal, and J. S. Duncan. Estimating tissue deformation between functional images induced by intracranial electrode implantation using anatomical MRI. *NeuroImage*, 13(4):561–76, Apr. 2001. ISSN 1053-8119. doi: 10.1006/nimg.2000.0692.
- F. van der Lijn, T. den Heijer, M. M. B. Breteler, and W. J. Niessen. Hippocampus segmentation in MR images using atlas registration, voxel classification, and graph cuts. *NeuroImage*, 43(4):708–20, Dec. 2008. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2008.07.058.
- K. Van Leemput, A. Bakkour, T. Benner, G. Wiggins, L. L. Wald, J. Augustinack, B. C. Dickerson, P. Goland, and B. Fischl. Automated segmentation of hippocampal subfields from ultra-high resolution in vivo MRI. *Hippocampus*, 19(6):549–57, June 2009. ISSN 1098-1063. doi: 10.1002/hipo.20615.
- H. Wang, J. W. Suh, J. Pluta, M. Altinay, and P. Yushkevich. Optimal weights for multi-atlas label fusion. *Information processing in medical imaging : proceedings of the ... conference*, 22:73–84, Jan. 2011. ISSN 1011-2499.
- S. K. Warfield, K. H. Zou, and W. M. Wells. Simultaneous truth and performance level estimation (STAPLE): an algorithm for the validation of image segmentation. *IEEE transactions on medical imaging*, 23(7):903–21, July 2004. ISSN 0278-0062. doi: 10.1109/TMI.2004.828354.
- J. L. Winterburn, J. C. Pruessner, S. Chavez, M. M. Schira, N. J. Lobaugh, A. N. Voineskos, and M. M. Chakravarty. A novel in vivo atlas of human hippocampal subfields using high-resolution 3 T magnetic resonance imaging. *NeuroImage*, 74:254–65, July 2013. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2013.02.003.
- L. E. M. Wisse, L. Gerritsen, J. J. M. Zwanenburg, H. J. Kuijf, P. R. Luijten, G. J. Biessels, and M. I. Geerlings. Subfields of the hippocampal formation at 7 T MRI: in vivo volumetric assessment. *NeuroImage*, 61(4):1043–9, July 2012. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2012.03.023.
- J. Wixted and L. Squire. The medial temporal lobe and the attributes of memory. *Trends in cognitive sciences*, 15(5):210–217, 2011. doi: 10.1016/j.tics.2011.03.005.
- R. Wolz, P. Aljabar, J. V. Hajnal, A. Hammers, and D. Rueckert. LEAP: learning embeddings for atlas propagation. *NeuroImage*, 49(2):1316–25, Jan. 2010. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2009.09.069.
- B. T. Wyman, D. J. Harvey, K. Crawford, M. A. Bernstein, O. Carmichael, P. E. Cole, P. K. Crane, C. Decarli, N. C. Fox, J. L. Gunter, D. Hill, R. J. Killiany, C. Pachai, A. J. Schwarz, N. Schuff, M. L.

- Senjem, J. Suhy, P. M. Thompson, M. Weiner, and C. R. Jack. Standardization of analysis sets for reporting results from ADNI MRI data. *Alzheimer's & dementia : the journal of the Alzheimer's Association*, Oct. 2012. ISSN 1552-5279. doi: 10.1016/j.jalz.2012.06.004.
- J. Yelnik, E. Bardinet, D. Dormont, G. Malandain, S. Ourselin, D. Tandé, C. Karachi, N. Ayache, P. Cornu, and Y. Agid. A three-dimensional, histological and deformable atlas of the human basal ganglia. I. Atlas construction based on immunohistochemical and MRI data. *NeuroImage*, 34(2):618–38, Jan. 2007. ISSN 1053-8119. doi: 10.1016/j.neuroimage.2006.09.026.
- P. A. Yushkevich, B. B. Avants, J. Pluta, S. Das, D. Minkoff, D. Mechanic-Hamilton, S. Glynn, S. Pickup, W. Liu, J. C. Gee, M. Grossman, and J. A. Detre. A high-resolution computational atlas of the human hippocampus from postmortem magnetic resonance imaging at 9.4 T. *NeuroImage*, 44(2):385–98, Jan. 2009. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2008.08.042.
- P. A. Yushkevich, H. Wang, J. Pluta, S. R. Das, C. Craige, B. B. Avants, M. W. Weiner, and S. Mueller. Nearly automatic segmentation of hippocampal subfields in in vivo focal T2-weighted MRI. *NeuroImage*, 53(4):1208–24, Dec. 2010. ISSN 1095-9572. doi: 10.1016/j.neuroimage.2010.06.040.