Development of a tool for automatic segmentation of the cerebellum in MR images of children



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Declaration

I, Priya Lakshmi Narayanan, hereby declare that the above thesis is my own unaided work both in concept and execution, and that apart from the normal guidance from my supervisor, I have received no assistance.

The thesis has been presented by me for examination for the degree of Doctor of Philosophy in Medicine in Biomedical Engineering.



Abstract

Thesis: Development of a tool for automatic segmentation of the cerebellum in MR images of children

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The human cerebellar cortex is a highly foliated structure that supports both motor and complex cognitive functions in humans. Magnetic Resonance Imaging (MRI) is commonly used to explore structural alterations in patients with psychiatric and neurological diseases. The ability to detect regional structural differences in cerebellar lobules may provide valuable insights into disease biology, progression and response to treatment, but has been hampered by the lack of appropriate tools for performing automated structural cerebellar segmentation and morphometry.

In this thesis, time intensive manual tracings by an expert neuroanatomist of 16 cerebellar regions on high-resolution T1-weighted MR images of 18 children aged 9-13 years were used to generate the Cape Town Pediatric Cerebellar Atlas (CAPCA18) in the age-appropriate National Institute of Health Pediatric Database (NIHPD) asymmetric template space. An automated pipeline was developed to process the MR images and generate lobule-wise segmentations, as well as a measure of the uncertainty of the label assignments. Validation in an independent group of children with ages similar to those of the children used in the construction of the atlas, yielded spatial overlaps with manual segmentations greater than 70% in all lobules, except lobules VIIb and X. Average spatial overlap of the whole cerebellar cortex was 86%, compared to 78% using the alternative Spatially Unbiased Infratentorial Template (SUIT), which was developed using adult images.

Since multi atlas fusion methods account for inter subject variability and reduce residual errors that may arise from using an atlas developed from a different population, we were interested in investigating whether our pre-adolescent training data could be used to accurately segment an adult data set. The performance of three different methods to propagate labels from the pre-adolescent training data used in the construction of the CAPCA18 atlas to 35 adult subjects were compared in the next section. These included direct warping of CAPCA18, multi atlas majority voting (MAMV) and multi atlas generative model

(MAGM) based label fusion. Our results demonstrate that multi atlas based label propagation methods achieved good segmentation accuracy and consistently performed better than direct warping, presumably due to the fact that these methods capture more of the variation present in the training data.

Finally, we compared cerebellar segmentation accuracy in the 35 adult test subjects between these two different multi atlas label fusion methods using the manually segmented data of 20 training subjects that were used in the construction of SUIT. Even though the training and test data were collected in two different laboratories on two different scanner platforms, mean Dice coefficients in the test subjects were greater than 0.89 in all regions except lobules VIIb, VIIIa, VIIIb, and X, confirming the robustness and reliability of our multi atlas segmentation pipeline.

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Table of Contents

Declaration	I
Abstract	II
Acknowledgements	IV
Table of Contents	V
List of Tables	VII
List of Figures	VIII
List of Abbreviations	XII
Preface	XIII

1. Introduction	1
1.1 Anatomy of the human brain	3
1.2 Medical Imaging	6
1.3 Medical Imaging Modalities	7
1.3.1 Computed Tomography	7
1.3.2 Positron Emission Tomography	8
1.3.3 Magnetic Resonance Imaging	8
1.4 Image Registration	12
1.5 Brain Template	20
1.6 Brain Atlas	23
1.7 Structural MR Image Analysis	26
1.7.1 MR Image Label Map	27
1.7.2 Automated cerebellar structure segmentation methods	27
1.7.3 Atlas-based segmentation	
1.7.4 Multi-atlas based segmentation	29

2. Improved labelling of cerebellar structures in children using probabilistic atlas-based segmentation	
2.1 Introduction	
2.2 Methods	
2.3 Results	
2.4 Discussion	
2.5 Conclusion	

3. Assessment of different label propagation methods for segmenting cerebellar structures in adults using the CAPCA18 pediatric cerebellar atlas	54
3.1 Introduction	55
3.2 Materials and Methods	59
3.3 Results	66
3.4 Discussion	72
3.5 Conclusion	73

74
75
90

scussion)4
----------	----

nclusions102

List of Tables

Table 3.1 Comparison of average lobular volumes for three randomly selected test subjects

 obtained using manual tracing and the three different label propagation methods.

 67

Table 4.2 Mean Dice coefficients in 35 test subjects of spatial overlap between the STAPLE estimate and automated segmentations in 20 hemispheric cerebellar regions using multi atlas majority voting (MAMV) and multi atlas generative model based (MAGM) label fusion, respectively.

 83

List of Figures

1.9 MR sequential processing of structural T1 images that first extracts the intracranial cavity (ICC) from the background (BG), and then iteratively segments the tissue classes present in the ICC into gray matter (GM), white matter (WM) and cerebrospinal fluid (CSF)......26

1.10 Schematic of atlas based segmentation
--

|--|

3.6 Maximum likelihood label maps and their associated lobular boundaries for two different coronal slices (a and b) superimposed on normalised T1 images of one test subject. Segmentations were generated using direct warping (left), multi atlas majority voting (MAMV; middle), and multi atlas generative model label fusion (MAGM; right). The look up table and colours representing each structure are listed in Figure 2.2. The visualization and the colour overlay was generated using Freeview software (http://www.nmr.mgh.harvard.edu/martinos). Yellow arrows indicate the transverse sinuses that are erroneously included in Crus I using direct warping segmentation. The red arrows indicate areas where MAGM contours follow the cerebellar gray matter boundaries more

List of abbreviations

The following table describes the significance of various abbreviations and acronyms used throughout the thesis.

СТ	Computed Tomography
CAT	Computer Assisted Tomography
PET	Positron Emission Tomography
MRI	Magnetic Resonance Imaging
RF	Radio Frequency
TR	Repetition Time
NMR	Nuclear magnetic resonance
3D	Three dimension
MAMV	Multi Atlas Majority Voting
MAGM	Multi Atlas Generative Model
LOOCV	Leave One Out Cross Validation
CUBIC	Cape Universities Brain Imaging Centre
NIHPD	National Institute of Health Pediatric Database
GM	Gray Matter
WM	White Matter
CSF	Cerebral Spinal Fluid
AC	Anterior Commissure
PC	Posterior Commissure
SNR	Signal to Noise Ratio
AAL	Automatic Anatomical Labelling
LONI	Laboratory of Neuro Imaging
LPBA40	LONI Probabilistic Brain Atlas
AIR	Automated Image Registration
SUIT	Spatially Unbiased Infra tentorial Template
CAPCA18	Cape Town Pediatric Cerebellar Atlas
STAPLE	Simultaneous Truth and Performance Level Estimation
SPM	Statistical Parametric Mapping
FMRIB	Centre for Functional Magnetic Resonance Imaging of the Brain
FSL	FMRIB Software Library
FLIRT	FMRIB's Linear Image Registration Tool
MNI	Montréal Neurological Institute
ICBM	International Consortium for Brain Mapping

Preface

This dissertation proposes a reliable method for automatically parcellating cerebellar structures in magnetic resonance (MR) images of children. This method involves the use of a fully automated pipeline for processing MR images and generating parcellation labels associated with the cerebellum.

A comprehensive introduction provides the necessary background and context to this work. Thereafter, this thesis includes three independent articles that will be submitted for publication. The articles are presented respectively in chapters two, three and four, then a chapter that discusses and expands on findings and a concluding chapter that summarises salient points from the preceding chapters. The organisation of the work into chapters is designed to facilitate direct access to and concise evaluation of different methodologies, and follows the logical progression of the work. As a complete document, however, this thesis contains some necessary repetition. This is because each core chapter is presented as an independent article.

For the purpose of thesis examination the contributions from co-authors are given below.

Chapter one provides the background and rationale for the current study, as well as a discussion of clinical imaging modalities, MRI physics and registration, and an overview of recent methods of atlas-based segmentation. It also provides the background theory for image segmentation methods investigated in the remainder of the thesis.

Chapter two presents the construction of the Cape Town Probabilistic Cerebellar Atlas (CAPCA18) from MR images of 18 healthy children aged 9 to 13 years, and its implementation for atlas-based segmentation of cerebellar structures, as validated on a

different set of children dataset. Further, the output segmentations are validated against manual segmentation using the standard performance metric for image analysis. Ernesta Meintjes supervised and directed the work covered in chapter two. The project design and implementation of the algorithms was that of the candidate, as well as the writing of the manuscript. Lilla Zöllei provided critical input on improving the validation methods. This chapter has been prepared as a manuscript, reviewed by all co-authors and is ready to be submitted for publication.

Chapter three details work conducted by the candidate during a study visit to the Martinos Center for Biomedical Imaging in Boston, in which all the subjects and corresponding parcellations from the CAPCA18 datasets were used to yield the cerebellar segmentation on adult dataset (Buckner et al., 2004). We utilized a group of 35 adult test subjects to investigate three label propagation methods: 1) using a traditional approach with a probabilistic group atlas; 2) individual registration of subjects to each reference atlas, followed by multi atlas majority voting (MAMV) label propagation; and 3) individual registration of subjects to each reference atlas, followed by multi atlas generative model (MAGM) based label propagation. We further validated for each method using 1) segmentations derived from the simultaneous truth and performance level estimate (STAPLE) algorithm; and 2) quantitative comparison with manual tracings of three test subjects. Ernesta Meintjes directed and supported the work. Lilla Zöllei provided expert advice and systematic guidance for conducting complex registration experiments to achieve precise registration results. The candidate designed the study, implemented the algorithms, performed all the tests and analyses, and prepared the manuscript. This chapter has been prepared as a manuscript and will be submitted for publication.

Chapter four presents results of experiments performed using the developed pipeline with datasets collected in two different labs. We present an optimized pipeline to obtain

automated parcellation of 28 cerebellar structures using manually delineated atlases on a set of test subjects using two types of multi-atlas based segmentation. The performance of the methods is evaluated using the STAPLE algorithm obtained in terms of Dice coefficient on test subjects. In addition, the results are evaluated against manual segmentation using the leave-one-out cross-validation (LOOCV) strategy on training subjects.

Chapter five is a critical discussion of the findings of the previous chapters. It identifies the limitations of the methods presented in the previous chapters. It also suggests possible improvements to the current implementation of the label propagation method in cerebellar structures.

Chapter six presents the conclusions of the study, summarising all salient points.

Chapter 1

Introduction

Rationale and Motivation

The different modalities of Magnetic resonance imaging (MRI), along with computational image analysis tools, have enhanced our ability to "see through" the human brain. MRI makes it possible to investigate brain structures and structure-related pathologies that haven't previously been fully understood. It enables the investigation of, amongst others, anatomical structures of the brain across different age groups, helping us better understand the developing brains of children. With automated parcellation, it also has the potential to provide insight into the neurobiology of psychiatric diseases and into responses to treatments.

This thesis aims to contribute to the development of tools for pipelining analysis of the cerebellum, specifically in pre-adolescent children. Many brain image analysis tools leave out the cerebellum and little detail about its sub-structures is available to the research community. Further, most atlases are developed using adult data and often perform poorly in pediatric data as they do not account for subtle differences in the developing brain. Subtle differences between white and gray matter in the cerebellum make segmentation tricky. Further, segmentation targeting smaller regions of the cerebellum is challenging, even for expert neuroanatomists.

To date, neuroscientists rely on manual procedures, which are time consuming and susceptible to risks related to inter- and intra-rater reliability. Due to time and cost constraints, manual procedures also aren't feasible for large samples. Because of the associated difficulties, manual segmentations generated for this study are employed as spatial priors and propagated to test performance in test subjects whose segmentations are unknown. In each chapter, the test subjects constitute a different sample from that used to develop the

algorithms. It is important for the reader to note the details of the test dataset and training dataset in each chapter.

This thesis presents a reliable cerebellar atlas construction procedure, based on structural T1 MRI images acquired at the Cape Universities Brain Imaging Centre (CUBIC) in Cape Town, South Africa. This work originated due to the limitations associated with structural segmentations of the cerebellum in children. The main objective was to obtain precise structural segmentation output using prior knowledge obtained from the manual segmentations performed on 18 subjects. The cerebellar atlas built in this study is designed to be used in routines to obtain cerebellar structure segmentation on any given test subject. The adult test subjects used for evaluation were the part of prior publication (Buckner et al., 2004) and the data were available with Freesurfer software package.

Contributions

In this work, we present the Cape Town Pediatric Cerebellar Atlas (CAPCA18) for segmentation of cerebellar structures.

- 1. Investigation on using CAPCA18 by evaluating on test dataset from children.
- 2. Investigation of using CAPCA18 atlas images using three different label propagation methods by using a dedicated pipeline and analysis methods for achieving cerebellar structure segmentation on adult test subjects. Three methods used for label propagation are direct warping, multi atlas based majority voting (MAMV) and multi atlas based generative model (MAGM).
- Investigation of using spatially unbiased infratentorial template (SUIT) atlas images using multi atlas based segmentation for achieving cerebellar structure segmentation on adult test subjects.

1.1 Anatomy of the human brain

The human brain comprises 100 billion neurons at birth. A child's brain has only one fifth of the volume of an adult brain (Thompson et al., 2005). The cells proliferate as the child continues to learn and to progress through various developmental stages. The dendritic branching of neurons increases with the increase in number of synaptic connections, which in turn is influenced by the learning process, and by environmental and genetic factors. So the shape of a brain depends on different developmental stages. Brain development is also influenced by gene expression (Armstrong et al., 1995; Ashburner 2009; Panizzon et al., 2009).

Growth in the field of MRI has enabled researchers to study the anatomical structures of the brain in more detail than was possible through post-mortem autopsy. Emerging neuroscience studies focus on mapping the structures, physiology, functions and connectivity of brain structures in individuals and across different populations. Central to these tasks is the construction of comprehensive brain atlases for different populations.

Brains can be anatomically divided into different regions. Based on embryonic development, the brain has three divisions– namely forebrain, midbrain and hindbrain. The forebrain, or cerebrum, is the largest part of the brain, where most information processing takes place. The midbrain consists of brainstem and connects the hindbrain and the forebrain. The hindbrain consists of brainstem, the cerebellum and the pons.

As further context for a discussion of the cerebellum, it is important to understand the tissues of the brain. Gray matter (GM) comprises largely of unmyelinated neurons, most of which are interneurons. The gray matter regions are the areas of nerve connections and processing. White matter (WM) is made mostly of myelinated neurons that connect the regions of gray matter to each other and to the rest of the body. The white matter acts as the information highway of the brain, receiving and sending information from and to every other part of the brain.

The cerebellum, which is the focus of this study, is one of the major parts of the central nervous system. It is located at the base of the skull, above the brain stem and beneath the occipital lobe of the cerebral cortex. This region is part of several neuronal networks. It is functionally heterogeneous, or involved in multiple functions, including sensory, motor, cognitive and affective processing (Stoodley and Schmahmann, 2009; Schlerf et al., 2010; Strick et al., 2009; Schmahmann and Sherman; 1998; Makris et al, 2005; Buckner et al., 2011).

Lobar parcellation using the identification of major fissures allowed the cerebellum to be divided into three sections, as in histological images Figure (1.1). These were the anterior lobe, the posterior lobe and the flocculonodular lobe. The anterior lobe comprises lobules I – V. The posterior lobe has superior posterior and inferior posterior regions. The superior posterior lobe comprises lobule VI and Crus I; and the inferior posterior lobe contains the lobules from Crus II to lobule IX. The flocculonodular lobe corresponds to lobe X. Smaller parcellation in an individual cerebellar lobe is called a lobule. In each lobule of the cerebellum, medial structures called vermis are present closer to the midline and hemispheric structures are present more laterally.

The most striking gross morphologic feature of the cerebellum is its highly convoluted cortical mantle (Figure 1.2). The convolutions allow a large cortical surface area to be contained within its volume. This distinctive property adds to the challenge associated with analysing the cerebellar structure and function (Van Essen et al., 2002).

Understanding the functionalities of the cerebellum and its inter-connections with cerebral lobes is a challenging task, one that typically necessitates the integration of several computational tools. Depending on the research problem, it's necessary to choose from a disconnected array of technologies and analysis methods, and to integrate these to develop a streamlined approach. Drawing on multiple tools tends to yield a greater amount of useful information and to improve statistical observation.



Figure 1.1 Somatotopic representation of structure and function of cerebellum (Left). (Images reproduced from Ermanno Manni and Laura Petrosini (2004)). Histological images (Right) representing anterior view (A & B), posterior view (C & D), inferior view (E & F) of the cerebellum.



Figure 1.2 Inferior view of the cerebellum representing the vermis of the inferior posterior lobe (Left A); Inferior view when the region is spread open; one can see the pyramid, uvule and nodule, and the smaller division of vermis (Right B).

1.2 Medical Imaging

Developments in medical imaging have made it possible to store images from living subjects in digital format. In recent years, an unprecedented volume of medical images has become available to researchers, helping them to completely characterise the imaged anatomies.

Information from different imaging modalities, such as computed tomography, positron emission tomography and MRI, makes it possible to study different tissues and their functional characteristics. Use of multiple acquisitions of the same subject and comparison across subjects or groups of subjects from the same population has become increasingly popular in recent years.

Developments in medical imaging technology, acquisition devices and storage facilities have resulted in a need for computational tools for processing medical images. As well as in processing information, the computational tools must allow for the comparison of information across subjects. Ultimately, this type of tool should create a visual representation of the different structures present in the images, along with the overall scan and quantitative results.

1.3 Medical Imaging Modalities

1.3.1 Computed Tomography

Computed tomography (CT), also known as computed axial tomography or computerassisted tomography (CAT), is a medical imaging procedure that uses computer-processed Xrays to produce tomographic images or "slices" of specific areas of the body. These crosssectional images are used for diagnostic purposes in various medical disciplines. Digital geometry processing is used to generate a three-dimensional image of the imaged anatomy from a large series of two-dimensional X-ray images taken around a single axis of rotation. Historically, the images generated were in the axial or transverse plane, perpendicular to the long axis of the body. However, modern scanners allow this volume of data to be reformatted in various planes or even as volumetric three dimensional (3D) representations of structures.

CT scanning of the head is typically used to detect infarction, tumours, calcifications, haemorrhage or bone trauma. Hypodense (dark) structures can indicate infarction. Hyperdense (bright) structures indicate calcifications or haemorrhage, and bone trauma can be seen as disjunction in bone windows. Tumours can be detected by the swelling and anatomical distortion they cause, or by surrounding oedema. Basically CT uses ionized (or X-ray) radiation to image the region of interest. It gives less information about the soft tissues and more information about bone structures and calcification.

1.3.2 Positron Emission Tomography

In the case of positron emission tomography (PET) imaging, short-lived radio isotopes are injected into the bloodstream of a living subject. Changes in regional blood flow in anatomic structures are quantified by the inference of radioactive decay. The output map shows the tissues in which the molecular tracer has become concentrated, and can be interpreted by a physician or radiologist in the context of the patient's diagnosis and treatment plan.

Nuclear medicine imaging in PET is non-invasive, with the exception of intravenous injections, which are usually painless medical tests. Important applications of PET are in clinical oncology, for identifying tumours leading to metastases, and in diagnosing and evaluating brain disease (grading various types of dementia). PET images are usually acquired with low resolution. Better clinical information is obtained by fusing the images from different modalities, like MRI and PET. This makes it necessary to perform automatic registration (Woods et al., 1993) between the different modality images.

1.3.3 Magnetic Resonance Imaging

Magnetic resonance imaging (MRI) is a safe modality that does not use any radioactive isotopes or ionizing radiation to image the anatomy of a structure. Instead, MRI uses the property of nuclear magnetic resonance (NMR) to image the nuclei of atoms inside the body. MRI can create more detailed images of the body than X-rays. It provides excellent contrast between the different soft tissues of the body, is non-invasive and can produce a volumetric, or three-dimensional, image. It can also produce images that are sensitive to the degree of blood flow in vessels, tissue oxygenation, tissue perfusion, diffusion, function and concentration of metabolites in tissues.

When atomic nuclei with magnetic properties are placed in a magnetic field, they can absorb electromagnetic waves of characteristic frequencies. The exact frequency depends on the type of nucleus, the field strength, and the physical and chemical environment of the nucleus. The absorption and re-emission of such radio waves is the basic phenomenon utilised in MRI.

Hydrogen proton (1H) is the most commonly used because of its abundance in bodily tissues, given that they occur in water and fat. Nuclei that have at least one unpaired proton, like the hydrogen proton, possess inherent spins. They have magnetic properties that distinguish them from non-magnetic isotopes.

A mechanical analogy to the magnetic nature of the nucleus is that it has a spinning mass with a small net positive charge. Because of the motion of the electric charge, a small magnetic field is created. The magnetic properties of the atomic nuclei make them precess around the external field. The frequency ω of this precessing motion is given by the following equation, called the Larmor equation:

$$\omega_{\rm L} = \gamma \times B_0, \tag{1.1}$$

in which ω_L is the angular Larmor frequency (unit: MHz), and γ is the gyromagnetic ratio (unit: MHz/T), which describes the ratio of the mechanical and magnetic properties of the nucleus and depends on the type of nucleus. B₀ is the strength of the magnetic field in Tesla (T). When an electromagnetic wave of appropriate frequency (equal to the Larmor frequency) reaches a nuclei in a state of lower energy, the nuclei is transferred to a state of higher energy. This is one of the few fundamental equations to understand in relation to NMR, MRI, its technology and its applications.

In the absence of a magnetic field, hydrogen protons are randomly aligned and the net magnetisation is zero. When a person is placed inside the powerful magnetic field (B_0) of an

MRI scanner, the magnetic moments of many protons become aligned with the direction of the field, creating a net magnetisation parallel to the main magnetic field.

A radiofrequency (RF) pulse applied to a coil perpendicular to B_0 , creates a small rotating magnetic field ($B_1(t)$) at the Larmor precession frequency perpendicular to the main magnetic field. For the duration of the RF pulse, the spins will precess around B_1 , effectively rotating the net magnetisation out of the longitudinal plane into the transverse plane perpendicular to the B_0 field. Upon termination of B_1 , the spins lose energy and return to their equilibrium state. This loss of magnetisation creates the MR signal.

The energy loss occurs in two main ways – spin-lattice relaxation and spin-spin interactions.

Spin-lattice interactions involve an exchange of energy between the spins and their surroundings. This results in the recovery of the longitudinal component of the magnetisation after a time (T1).

Spins also interact with themselves in a more rapid process than spin-lattice interactions, and this leads to a loss of phase coherence among the spins. The time for the resulting loss in the transverse component of the magnetisation is called T2. In general, T2 is less than T1. Different tissues have different T1 and T2 time constants. For example, myelinated white matter has a shorter T1 than gray matter. This means that in T1-weighted MR images of adult brains, white matter recovers faster and therefore appears brighter in the images than gray matter. In T2-weighted images, gray matter will appear brighter than white matter.

A tissue with a long T1 and T2 (like water) is dark in a T1-weighted image and bright in a T2-weighted image. A tissue with a short T1 and a long T2 (like fat) is bright in a T1-weighted image but gray in a T2-weighted image.

T1 contrast may be manipulated in images by varying the repetition time (TR), which defines the time between successive RF pulses. In this thesis, structural T1 weighted images have been used throughout.

An MR pulse sequence diagram is a simple way of showing how the RF pulses and gradients are applied during an acquisition. Enormous numbers of pulse sequences have been developed, each with its own application.

Three mutually orthogonal magnetic field gradients – namely the slice select gradient, phase encoding gradient and frequency encoding gradient – are usually applied in different combinations and at different times during image acquisition to achieve spatial encoding of the MRI signal to construct the MR image.

Arbitrary orientations of the linear gradients can be achieved through the simultaneous application of fixed-orientation gradients in the X, Y, and Z axes. The basis for spatial localisation is the fact that the nuclear precession frequency is directly proportional to the magnetic field (eq. 1.1). The gradients induce linearly varying magnetic fields, resulting in spatially encoded precessional frequencies. For example, if the gradient is applied during excitation, only spins that match the resonance condition will be excited. In this way, an imaging slice is selected. Similarly, by applying the gradients during readout, the location from which a signal with a specific frequency and phase originated can be determined, yielding a "profile" image.

In order to produce a topographic image, it is necessary to acquire multiple profile images, so signal excitation needs to be repeated many times, with allowances for relaxation time between excitation. The program that controls the timings and order that the different components of the imaging sequence are applied is the pulse sequence.

Brain MR images may be affected by various sources of noise, for example, magnetic field inhomogeneities, ambiguous intensity patterns inside and outside the brain and partial volume effects. Partial volume effects occur when the intensities of voxels are influenced by multiple structures, resulting in a blurring of intensity across boundaries. These effects are common in MRI data where the resolution of the images is relatively large compared to the brain microstructure and images are often not isotropic.

Wrap-around artefacts (Figure 1.3) occur when a part of the image folds around and is visible on the opposite side of the image. This usually happens when the field of view is too small or has not been well placed. Such artefacts were found in a few datasets used in chapters 3 and 4 of this thesis (artefacts not shown in said chapters). These artefacts were removed during the pre-processing stage of the pipeline.



Figure 1.3 Wrap-around artefact (yellow arrow) that shows the patient nose in each of the orthogonal planes (sagittal, and axial)

1.4 Image Registration

Image registration is the process of aligning two images of the same scene or slice so that the output provides more information about the underlying anatomy than either of the separate images. In three-dimensional (3D) space, the volumetric image (target) from one coordinate system is aligned to a reference image in a different coordinate system.

Registration can also be applied to multimodal images. For example, the registration of CT and MR images performed during image fusion (Sasikala et al., 2007) is useful to detect

localised atrophy or neuropathology associated with certain diseases, or to aid in the diagnose of brain tumours. The review paper by Maintz and Viergever (1998) provides a comprehensive overview of key concepts related to image registration.

Registration is an important part of MRI data analysis. A typical image registration algorithm consists of three components that are performed in an iterative way: (see Figure 1.4)

- an alignment measure that quantifies the quality of alignment using an objective function
- a class of admissible geometric transformations that can be applied to the image or images; i.e., that can be employed to spatially "warp" the image(s)
- an optimiser, which seeks the transformation that results in the best alignment, as quantified by the alignment measure and finally maximizes similarity.



Figure 1.4 Schematic representing the three components of image registration

When determining the spatial correspondence between any two images that portray the same scene, one image is considered fixed and referred to as the reference and the other image is the target. The registration framework basically consists of transformation, optimisation and interpolation steps. Transformation helps to achieve spatial correspondence by transforming from one coordinate space to another coordinate space. Optimisation is used during a multiresolution approach when using high resolution images in order to reduce computational demands. Multiresolution registration refers to the process where registration is performed iteratively in regions of increasing spatial resolution. Multiresolution based optimization (Jenkinson et al., 2002) is commonly used in most registration methods.

Registration algorithms are classified in terms of the following five features:

- 1. dimensionality,
- 2. the nature of the transformation,
- 3. the basis functions used during registration,
- 4. imaging modality, and
- 5. whether registration is within or between subjects.

Dimensionality depends on the spatial dimensions of the images involved in the registration process, and registration may be between images with either similar or different dimensionalities. For example, intra-operatively multiple 2D projection images often need to be registered to a 3D image acquired previously during surgical planning.

Transformations are categorized into rigid, affine, non-linear, projectile and curved transformations. Linear transformations can be either rigid or affine; rigid allows only for translation and rotation, while affine transformations allow for translation, rotation, scaling and shear between the images. Projectile and curved transformations are not used in our study.

In rigid body registrations, six parameters are considered: three translations and three rotations. Using these parameters one image is transformed to another image. Each point (x_1, x_2) in a 2D image is mapped to co-ordinates (y_1, y_2) of another image by the simple transformation y=Mx.



Figure 1.5 Example of 2D image rigid registration

Affine registration is a type of linear geometric transformation that involves translation, rotation, scaling and shear. For each point (x_1, x_2, x_3) in a 3D image, an affine mapping transforms these co-ordinates into coordinates (y_1, y_2, y_3) in another space. The transformation is given by equations 1.2, 1.3 and 1.4.

$$y_1 = m_{11}x_1 + m_{12}x_2 + m_{13}x_3 + m_{14} \tag{1.2}$$

$$y_2 = m_{21}x_1 + m_{22}x_2 + m_{23}x_3 + m_{24} \tag{1.3}$$

$$y_3 = m_{31}x_1 + m_{32}x_2 + m_{33}x_3 + m_{34} \tag{1.4}$$

In matrix form the affine transformation is given simply by y=Mx. (eq 1.5).

$$\begin{pmatrix} y_1 \\ y_2 \\ y_3 \\ 1 \end{pmatrix} = \begin{pmatrix} m_{11} & m_{12} & m_{13} & m_{14} \\ m_{21} & m_{22} & m_{23} & m_{24} \\ m_{31} & m_{32} & m_{33} & m_{34} \\ 0 & 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \\ x_3 \\ 1 \end{pmatrix}$$
(1.5)

Equation 1.6 defines a translation by p (i.e. y=x+p) and equation 1.7 a rotation q_1 radians about the x axis.

$$\begin{pmatrix} y_1 \\ y_2 \\ y_3 \\ 1 \end{pmatrix} = \begin{pmatrix} 1 & 0 & 0 & p_1 \\ 0 & 1 & 0 & p_2 \\ 0 & 0 & 1 & p_3 \\ 0 & 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \\ x_3 \\ 1 \end{pmatrix}$$
(1.6)

$$\begin{pmatrix} y_1 \\ y_2 \\ y_3 \\ 1 \end{pmatrix} = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & \cos(q_1) & \sin(q_1) & 0 \\ 0 & -\sin(q_1) & \cos(q_1) & 0 \\ 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \\ x_3 \\ 1 \end{pmatrix}$$
(1.7)

Similarly, rotations q_2 and q_3 radians about the y and z axes, respectively, are given by equations (1.8) and (1.9).

$$\begin{pmatrix} y_1 \\ y_2 \\ y_3 \\ 1 \end{pmatrix} = \begin{pmatrix} \cos(q_2) & 0 & \sin(q_2) & 0 \\ 0 & 1 & 0 & 0 \\ -\sin(q_2) & 0 & \cos(q_2) & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \\ x_3 \\ 1 \end{pmatrix}$$
(1.8)
$$\begin{pmatrix} y_1 \\ y_2 \\ y_3 \\ 1 \end{pmatrix} = \begin{pmatrix} \cos(q_3) & 0 & \sin(q_3) & 0 \\ -\sin(q_3) & 1 & \cos(q_3) & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \\ x_3 \\ 1 \end{pmatrix}$$
(1.9)

Scaling is needed to change the size of an image and also to correct for non-isotropic voxels. An example of scaling along the orthogonal axis is shown in eq. 1.10.

$$\begin{pmatrix} y_1 \\ y_2 \\ y_3 \\ 1 \end{pmatrix} = \begin{pmatrix} q_1 & 0 & 0 & 0 \\ 0 & q_2 & 0 & 0 \\ 0 & 0 & q_3 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \\ x_3 \\ 1 \end{pmatrix}$$
(1.10)

Shear is computed using the matrix in eq.1.11.

$$\begin{pmatrix} y_1 \\ y_2 \\ y_3 \\ 1 \end{pmatrix} = \begin{pmatrix} 1 & q_1 & q_2 & 0 \\ 0 & 1 & q_3 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \\ x_3 \\ 1 \end{pmatrix}$$
(1.11)

Registration performed using non-linear transformations uses the image that has already been approximately registered with the template according to a twelve-parameter affine registration. The transformation model for defining nonlinear warps uses deformations consisting of a linear combination of low-frequency periodic basis functions. These are the low-frequency components of a 3D discrete cosine transform (DCT).

Registration using the non-linear volumetric procedure fails to align the cortical folding patterns because of its highly nonconvex nature of the energy functions when initialized with an affine transform. To achieve this finer refinement in the deformation, we can integrate the surface based information into a volumetric registration procedure. The resultant is a 3D displacement field that aligns both cortical folding patterns and subcortical structures of the brain (Postelnicu and Zöllei et al., 2009).

Registration can be performed using either extrinsic or intrinsic features. Extrinsic methods rely on artificial objects attached to the patient. These are objects that are designed to be visible and accurately detectable in all of the pertinent modalities. Intrinsic methods rely only on patient-generated image content. Registration can be based on a limited set of identified salient points known as landmarks, on the alignment of segmented binary structures (most commonly object surfaces), or on measures computed directly from the image gray values.

Landmark-based registration employs a large set of points to represent a structure. It allows complex registration to be achieved accurately faster than volume-based registration. The iterative closest point method (Besl and Mckay 1992) is a widely used landmark-based registration algorithm. In a shape analysis study of the brains of children with fetal alcohol syndrome (FAS), surface models of the hippocampus and caudate nucleus from different subjects were registered using vertices as landmark points (Joseph et al., 2014).

Registration methods are further categorised as either single-modality or multimodality methods. A single-modality method tends to register images acquired by the same scanner or sensor type, whereas a multi-modality registration method tends to register images acquired by different scanners or sensor types. In the case of MR images, however, multimodal registration is carried out on images that were acquired using different pulse sequences and as such have different contrasts.

Finally, registration methods are categorised based on the number of subjects, i.e. intrasubject, inter-subject or atlas registration methods. In intra-subject registration, scans of the same subject taken at different instants are registered. For example, the high-resolution Colin27 MRI brain template was created by performing intra-subject registration of the same subject scanned at 27 different instants.

Inter-subject registration involves registering scans from multiple subjects and is useful for comparing anatomical structures in different brains. The Montreal Neurological Institute (MNI) used this type of registration to develop the MNI305 human brain template. MR images from 305 subjects were mapped into stereotactic space, intensity normalised and averaged on a voxel-by-voxel basis as part of the International Consortium for Brain Mapping (ICBM) (Evans et al., 1993). Later, 152 subjects were affinely registered to the

MNI305 template to create the ICBM152 template (Mazziotta et al., 2001; Mazziotta et al., 1995).

The ICBM152 template is widely used as a registration target during functional analyses in a variety of popular computational neuroimaging tools, including SPM (www.fil.ion.ucl.ac.uk/spm/), FSL (www.fmrib.ox.ac.uk/fsl/) and AIR (bishopw.loni.ucla.edu/air5). Precise registration algorithms have become critical for brain mapping applications because it allows comparisons of experimental findings from different subjects.

In structural imaging, image registration methods are applied sequentially. Linear registration is initially used to obtain global correspondence. This is followed by non-linear parametric transformation yielding smooth deformations across local structures (Figure 1.6). Non-linear deformations range from smooth parametric deformations to high-dimensional, non-parametric deformations. High-dimensional, non-parametric deformations are not driven by intensity variations, but take geometric information, curvature and other image features into account and follow either elastic or optical flow-based registration.

Since human brains are highly variable and also differ depending on age, gender, race, etc., it may be difficult to choose a suitable registration target. Standard brain templates are often used to provide an anatomical reference for individual or population based assessment of brain structure and function.


Figure 1.6 Average anatomical image of the cerebellum before registration (left) ; Average anatomical image after affine linear registration to the national institute of health pediatric database (NIHPD) template (middle); Average anatomical image after affine linear and non-linear registration with the NIHPD template (right).

(For more detail on image registration, reader can refer to Maintz and Viergever (1998), Crum et. al., 2004 and Crum et al., 2014.)

1.5 Brain Template

Brain templates are constructed from one or more representations of the brain (Toga and Thompson 2001). Brain templates have facilitated the increase in the number of investigations focusing on the structural and functional organization of the brain.

In humans and other species, the brain's complexity and variability across subjects is so large that templates are essential to manipulate, analyse and interpret brain data from multiple subjects. Registration helps to establish a common coordinate system for evaluation of different images. Images are typically registered onto a reference space via affine transformations (Mazziota et al., 2001).

The reference space can be defined by a single subject as per the initial coordinate system proposed by Talairach (Talairach and Tournoux, 1988). The Talairach coordinate system is based on the post-mortem brain of a single subject. However, such an atlas template is biased towards the chosen reference subject space and may not represent the average geometry of the population. Several different approaches have been explored for using multiple subjects in the construction of average intensity atlases. The Montreal Neurological Institute (MNI) constructed the standard MNI305 whole-brain template by registering 305 scans using 9-

parameter linear transformation. Their next generation template, popularly known as the ICBM152 template (Evans et al., 1993), was built using 12-parameter linear and non-linear registration in a stereotactic co-ordinate system. It also included the previously excluded posterior regions and reduced the complexities arising from inter-subject structure variability associated with individual brains. The ICBM template in MNI space is representative of the average brain size and shape. Cortical structures are, however, quite blurred due to the effect of low-pass filtering in the averaging process. The initial template covers the whole head with posterior regions excluding the cerebellum. The brain template is the result of an averaging operation after linear and non-linear registration.

The historical evolution of different whole brain templates using linear and non-linear registration is well documented by Evans et al. (2012) in a review of brain templates. The characteristic features of available adult brain templates are listed in Table 1. In addition to these adult templates, the National Institutes of Health Pediatric Database (NIHPD) has constructed a set of age-specific paediatric templates for child populations. These were constructed via iterative non-linear registration (Fonov et al., 2011) and enable mapping of the brain using a standard co-ordinate system.

The stereotactic co-ordinate system is one example of a widely used standard co-ordinate system in the neuroscience community. It uses the anterior and posterior commissure (AC-PC) to determine the origin of the coordinate system (Figure 1.7). The X-axis is defined as the line passing through the AC point and perpendicular to the AC-PC line. The Y-axis is defined by the line connecting the most superior point of the AC and the most inferior point of the PC. The Z-axis is defined by the line on a vertical plane (while the horizontal plane is defined by the X-axis and Y-axis) passing through the inter-hemispheric fissure and the AC point (Chau et al., 2005).

The process of positional normalisation remains a mandatory step in the processing pipeline. Also, all brain templates are always presented in the stereotactic co-ordinate system.



Figure 1.7 The intersection of lines passing through anterior commissure (AC) and posterior commissure (PC) define the origin of the stereotactic co-ordinate system. The left quadrant takes negative values and the right quadrant takes positive values.

Feature	Talairach	MNI-305	Colin-27	MNI-152	ICBM-452
	Tournoux	(1995)	(1998)	(2001)	(2003)
	(1988)				
No. of subjects	of subjects 1		1	152	452
*Gender	*Gender F-1		M-1	NA	NA
Age (years)	60	23.4±4.1	NA	NA	NA
Image type	2D	3D	3D	3D	3D
Population specific	No	Yes	No	Yes	Yes
Type of atlas	Non-digital	Digital	Digital	Digital	Digital
Registration	Manual	Linear	Linear and	Linear and	Linear and
procedure	(Intra-subject)	(Inter-subject)	non-linear	non-linear	non-linear
			(Intra-subject)	(Inter-subject)	(Inter-subject)
Spatial	NA	9-parameter	Affine	9-parameter	12-parameter
transformation		linear		affine	affine
Coverage	Excludes brain	Does not fully	Full head and	Full head and	Full head and
	stem and	cover top of	cerebellum	cerebellum	cerebellum
	cerebenum	cerebellum			
Cortical detail	Lack of cortical	Lack of cortical	Improved	Lack of	Improved
information	detail	detail	cortical detail	cortical detail	cortical detail
Contrast	Less significant	Significant	Significant	Significant	Significant
between gray					
and					
whitematter					
Size (length,	Smaller than	Larger than	Same as	Same as MNI-	Same as
width and	other brain	Talairach	MNI-305	305	MNI-305
height of	templates	Tournoux			
brain)					

Table. 1.1 Features of different adult brain templates

*M-Male; F-Female; NA- Not applicable

1.6 Brain Atlas

The establishment of stereotactic space led to a framework for consolidation of statistical information across subjects (Evans et al., 2012). After achieving mapping to a brain template, individual anatomical brain structures require identification. This can be accomplished by

using an established atlas in which brain structures have been parcellated, often referred to simply as a brain atlas. A brain atlas consists of a spatial map of the relative locations of individual structures. It describes the underlying anatomy of a structure in terms of structural morphology, size and shape. Atlases are typically created from multiple subjects through manual segmentation of the anatomical structures of the brain.

Brain atlases may be extended to include a probabilistic map of individual brain structures. A probabilistic atlas encodes the underlying anatomical variability in the structures across different subjects. A probability map of segmented structures in the standard coordinate system makes it possible to describe image data with significant statistical and visual power. Anatomical variability is determined by using data from multiple subjects during the atlas construction procedure. Volumetric parcellated atlases for identification of brain structures have been developed by various groups around the globe. Automatic Anatomical Labelling (AAL) is an early atlas comprising 45 volumes per hemisphere created from the Colin27 brain, non-linearly warped to MNI152 space (Tzourio-Mazoyer et al., 2002). It included both cerebral and cerebellar parcellations. The Laboratory of Neuro Imaging (LONI) used 40 brain images to generate the LONI Probabilistic Brain Atlas (LPBA40 Shattuck et al., 2008). It is available to the neuroimaging community in a set of different variants in normalized stereotactic space. The atlas construction pipeline used different normalisation techniques to generate different variants of the same atlas. The probability map of LPBA40 was derived from manual labels using the well-established LONI protocols.

Analysis of structural MR images using templates and atlases requires certain standard preprocessing steps, described in the following section.







Figure 1.8 Sagittal, coronal and transaxial slices of 18 control children used in this study for cerebellar atlas construction and structure parcellation.

1.7 Structural MR Image Analysis

Currently almost all neuroimaging software packages recommend the use of a series of computational tools. This is called a processing pipeline. Structural analysis of MR images is aided by writing in-house scripts specifically to ensure compatibility with in-house data. This is greatly facilitated by routines in open-source tools. Briefly, the raw MR images (Figure. 1.8) are skull-stripped, corrected for inhomogeneities and segmented into different tissue classes (Figure. 1.9), namely gray matter (GM), white matter (WM) and cerebrospinal fluid (CSF). These procedures are common to most of the serial processing computational tools, and complex model parameterisation takes care of inhomogeneity and bias correction (Pham et al., 2000).



Figure 1.9 MR sequential processing of structural T1 images that first extracts the intracranial cavity (ICC) from the background (BG), and then iteratively segments the tissue classes present in the ICC into gray matter (GM), white matter (WM) and cerebrospinal fluid (CSF).

1.7.1 MR Image Label

Image segmentation is defined as partitioning of an image into non-overlapping, constituent regions that are distinct and homogeneous with respect to anatomy or a characteristic such as intensity. In this work, a segmentation method ideally finds the set of voxels that correspond to distinct anatomical structures of the cerebellum. Ω represents the image domain and *Si* is the number of segmentations in the image.

$$\Omega = \bigcup_{i=1}^{k} Si \tag{1.12}$$

It's assumed that the value of k is known, based on prior knowledge of the cerebellar anatomy. For whole brain analyses, it is common to use k=3; corresponding to GM, WM and CSF tissue classes. In this study, we used k=16 as this was the number of structures delineated.

1.7.2 Automated cerebellar structure segmentation methods

Segmenting cerebellar structures in MR images is a challenging task because the intensity distributions of cerebellar gray matter differ from that in the rest of the brain, the cerebellum has a complex, convoluted structure with smaller substructures and physiological noise occurs around the structure (Diedrichsen et al., 2010). Different structures of the cerebellum have the same intensity profile, making it challenging to define the border between two structures accurately. Further, not all boundaries between structures are visible on MR images. Therefore cerebellar structure segmentation methods cannot rely on intensity information alone; additional knowledge such as the deformation warp is needed to separate structures of interest from the background.

The current brain structure segmentation literature can be broadly classified into two categories, based on the type of additional knowledge that they incorporate, namely atlasbased segmentation and multi-atlas based segmentation. These are briefly introduced in the following sections. In practice, this categorization is not always clear-cut and a combination of these methods may result in superior performance.

1.7.3 Atlas-based segmentation

Atlas-based segmentation refers to the process of generating segmentation via registration of an atlas and a candidate image. The spatial correspondence established between the intensity images is used to obtain an output label map of the structure on the candidate image (Figure 1.10). By using a non-rigid deformation field in the registration process, it is possible to extract information relating to subtle differences between subjects.

An atlas-guided approach is generally suited for segmentation of structures that are stable over the population of study (Pham et al., 2000). It has numerous biomedical applications, such as quantification of tissue volumes, diagnosis and localisation of pathology, investigation of particular structures, treatment planning and computer-integrated surgery.



Figure 1.10 Schematic of atlas based segmentation. Atlas construction is illustrated to the left of the blue line and atlas based segmentation to the right. Training images are registered to a standard template and the resulting transformation warps are applied to the individual atlases comprising the manually traced data of each subject. The labels in each location of the registered individual atlases are averaged to create the probability map. During segmentation, the test image is first registered to the template. The inverse of this transformation is then applied to the probability map to warp the labels to the test image.

1.7.4 Multi-atlas based segmentation

Multi-atlas based segmentation has proven to be one of the most competitive techniques in medical image segmentation (Heckemann et al., 2006, Asman et al., 2013, Wang et al., 2013). In a multi-atlas based segmentation approach (Figure 1.11), multiple atlas images are registered to a target image and their segmentations are combined, to obtain segmentation of the target image. This approach produces robust segmentation, transferring segmentations from expert-labelled images, or atlases, to a target image using deformable image registration. Errors produced by label transfer are further reduced by label fusion, which

combines the results produced by all atlases into a consensus solution. Regardless of the approach, fusion algorithms are generally dependent on large atlas sets and highly accurate registration.



Figure 1.11 Schematic of multi atlas based segmentation. The segmentation of one test image is determined from the pairing of each training image (18 in total) with the test image. Linear and nonlinear warping is used to transform the test image to each training image to obtain 18 distinct segmentations based on the 18 individual atlases. These 18 deformed segmentations of the test image are transformed back into the test image space using the inverse transformations. Label fusion methods are applied to obtain the final segmentation from the 18 distinct segmentations.

Chapter 2

Improved labelling of cerebellar structures in children using probabilistic atlas-based segmentation

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Abstract

Consistent morphometry and functional localization of cerebellar cortex in a standard coordinate system is important for interpretation of functional data, as well as comparisons of results and anatomy across sites and populations. We present the first pediatric cerebellar atlas for improved labelling of cerebellar structures in children. The Cape Town Pediatric Cerebellar Atlas (CAPCA18) was constructed in the age-appropriate National Institute of Health Pediatric Database (NIHPD) asymmetric template space using manual tracings of 16 cerebellar compartments in 18 healthy children aged 9-13 years from Cape Town, South Africa. The probabilistic CAPCA18 atlas provides both valid assignments of labels to specific spatial coordinate locations, as well as a quantitative measure of the uncertainty of such assignments. Segmentation using this atlas was validated through comparison to manual tracing in an independent set of 14 healthy children in the same age range and from the same community. Results were compared to the performance of the Spatially Unbiased Infratentorial Template (SUIT) on the same dataset. Average spatial overlap of the whole cerebellar cortex for test images with their corresponding manual segmentations using the probabilistic CAPCA18 atlas was 86%, compared to 78% after normalisation to the SUIT template. Spatial overlap between automated and manual segmentations using the CAPCA18 atlas of individual lobules in both hemispheres were 70% or higher for all lobules, except lobules VIIb and X where the overlap was on the order of 50-60%. On average, these values were 8% higher than those obtained using SUIT for this pediatric dataset.

Keywords: Probabilistic atlas, cerebellum, brain mapping

2.1 Introduction

The human brain is a complex structure and mapping its functional organization presents an ongoing challenge. Recent findings suggest that the cerebellum is functionally heterogeneous, with different topological regions subserving sensory, motor, cognitive, and affective processing (Stoodley and Schmahmann, 2009; Schlerf et al., 2010; Strick et al., 2009; Schmahmann and Sherman, 1998; Makris et al., 2005). As such, it has become increasingly important to identify precisely which lobule is activated in functional imaging studies. Efforts to map cerebellar function have, however, been limited by the fact that available cerebellar atlases are generally limited to gross morphologic relationships (Crosby et al., 1962; Carpenter et al., 1976; DeArmond et al., 1976; Waddington et al., 1984; Roberts et al., 1987; Kretshmann and Weinrich, 1992), that the individual cerebellar lobules are generally not labelled, and only limited sections are depicted in either one or two of the cardinal planes with large gaps between these. Furthermore, the terminology used to identify the fissures and lobules in these atlases is not uniform and is often contradictory.

Schmahmann et al. (2000) presented a human cerebellar atlas with sections at 2mm intervals in three cardinal planes based on high-resolution T1-weighted Magnetic Resonance (MR) images of a single human cerebellum that was coregistered to the Montreal Neurological Institute (MNI) template (Evans et al., 1993) and annotated using a revised and simplified nomenclature. Using the above MR Image Atlas of the Human Cerebellum as a basis for identification of landmarks and fissures, Makris et al. (2005) developed a manual method aided by a set of computer-assisted algorithms to facilitate the parcellation of the cerebellar cortex into 32 Parcellation Units (PUs) per hemicerebellum in a manageable period of time. In their implementation, the fissures divide the cortex into lobules, while longitudinal divisions separate the vermis from the hemispheres, and subdivide the hemispheres into medial and lateral zones. The large lateral hemispheric region of Crus I and II is divided into a further two zones. The authors found that intraclass correlation coefficients (McGraw and Wong, 1996; Shrout and Fleiss, 1979) for both intra- and inter-rater reliability were significantly improved by clustering PUs according to lobar divisions, anatomical connectivity, or functional connectivity. Lobar clusters are widely used (Pierson et al., 2002) and divide the cerebellum into anterior, posterior and flocculonodular lobes that are separated by the primary and the posterolateral fissures, respectively. In all these studies only data acquired from adults were used.

Subsequently, Diedrichsen (2006) developed the high-resolution Spatially Unbiased Infratentorial Template (SUIT) of the cerebellum by normalizing individual cerebella of 20 healthy adults non-linearly to each other before averaging, which improved specificity when labelling regions in functional MRI data.

Although atlases are widely used to assign anatomical labels to locations, there is a high risk for error due to high spatial variability of individual cerebellar structures between different populations. Probabilistic atlases allow for the assignment of labels to specific regions, while providing a quantitative measure of the uncertainty of such assignments. Currently, whole brain probabilistic atlases typically treat the cerebellum as a single structure without any lobular divisions (Hammers et al., 2003; Shattuck et al., 2008). In 2009, the first probabilistic cerebellar SUIT atlas was created (Diedrichsen et al., 2009) based on manual tracings of lobules on T1-weighted MRI scans (1mm isotropic resolution) of 20 healthy adult participants (10 male, 10 female, age range 19-27 years). The SUIT atlas defines twenty-eight compartments: lobules I-IV and V divided into left and right hemispheres; lobules VI, Crus I, Crus II, VIIb, VIIIa, VIIIb, IX, and X divided into vermal sections in addition to left and right hemispheres. This atlas aims specifically to improve inter-subject co-registration of

cerebellums to yield improved specificity of cerebellar activations and valid assignments of functional activations to specific cerebellar lobules. Recently, Bogovic et al. (2013) described a method that uses multiple object geometric deformable models to perform cerebellar segmentation and demonstrated superior performance compared to SUIT atlas-based and multi-atlas fusion approaches.

We were interested in examining cerebellar anatomy in children (age 9-13 years) from the Cape Coloured (mixed ancestry) community in Cape Town, South Africa. Since manual tracing is both time intensive and subjective, we wanted to perform automatic cerebellar segmentation. It has been noted previously, however, that a specialized atlas should be created for research in children (Diedrichsen, 2006) as the shape and ratio of gray matter to white matter in the cerebellums of children differ significantly from that of adults (Fonov et al., 2011). To our knowledge, no pediatric cerebellar atlas is currently available.

In developing an atlas, one important decision relates to the registration target (template) to use. It has been shown previously that a template closer to the study population reduces morphometric bias and the amount of non-linear deformation required to establish spatial alignment (Yoon et al., 2009) between the template and subject. Previously, Wilke et al. (2008) developed the 'Template O Matic' toolbox for SPM to create an age specific whole brain template by initially using linear co-registration of subjects and regressing for age and gender in pediatric populations. The resulting template, however, appears smoothed and lacks anatomical detail in the areas of greatest variability.

The unbiased nonlinear National Institutes of Health Pediatric Database (NIHPD) template created by Fonov et al. (2011) provides better spatial resolution and improved contrast compared to the classical International Consortium for Brain Mapping (ICBM152) template. Templates are available for different age ranges for normal brain development. In their work, the pediatric population was grouped into five categories between the ages of 4.5 and 18.5 years. For each category, two templates were constructed, one that preserves asymmetry and another with symmetric hemispheres. Deformation studies using these five different templates have shown that the average magnitude of deformation increases with increasing difference in age between the template used and the subject being studied (Fonov et al., 2011). In the present study we used the NIHPD unbiased nonlinear template that preserves asymmetry closest to the age range of our subject population (7.5 years to 13.5 years).

In this work, we present the probabilistic Cape Town Pediatric Cerebellar Atlas (CAPCA18), for improved labelling of cerebellar structures in children. The atlas was constructed in the already established age-appropriate NIHPD template space from manual tracings of 16 cerebellar compartments in 18 healthy children (age range 9-13 years, 6 male) according to the nomenclature introduced in the MRI atlas of the human cerebellum (Schmahmann et al., 2000). The probabilistic presentations of each compartment provide a quantitative measure of the spatial variability. Our atlas was validated by comparing manually traced segmentations with automated segmentations using the CAPCA18 atlas in an independent set of 14 healthy children (age range 8.9-11.8 years, 10 male) from the same community.

2.2 Methods

High resolution T1-weighted structural images were acquired on a 3T Allegra (Siemens, Erlangen, Germany) MRI scanner using a Magnetization Prepared Rapid Gradient Echo (MPRAGE) sequence (TR 2300ms, TE 3.93ms, TI 1100ms, 160 slices, flip angle 12 degrees, 1.3x1.0x1.0mm³, 6.03 minutes) in 18 healthy children (mean age 11.8±1.2 years) from the Cape Coloured community in Cape Town, South Africa who were recruited as typically developing controls for ongoing studies of Fetal Alcohol Spectrum Disorder (Jacobson et al.,

2008). A further 14 healthy children (mean age 10.5 ± 0.8 years) from the same community were scanned using a volumetric navigated (Tisdall et al., 2012) multiecho (ME) MPRAGE sequence (van der Kouwe et al., 2008) (128 sagittal slices, TR 2530ms, TE 1.53/3.21/4.89/6.57ms, TI 1100ms, flip angle 7 degrees, 1.3x1.0x1.3mm³). All children were scanned according to protocols that had been approved by the Faculty of Health Sciences Human Research Ethics Committee at the University of Cape Town; parents of all children provided written informed consent and children provided oral assent.

Image Pre-processing

The images were reoriented with the horizontal line defined by the anterior posterior commisure (ACPC orientation) and the sagittal planes parallel to the midline. The images were resampled to isotropic 1mm³ voxels using windowed sinc interpolation in Brain Voyager (Goebel et al., 2006). The resulting images were cropped to a fixed bounding box by removing the empty slices from the volume using a customised script written in MATLAB (www.mathworks.com). This process ensured removal of the neck and generated similarly oriented whole brain images.

Manual Tracing

The cerebella of the 18 input subjects were manually traced in native space according to the revised nomenclature defined by Schmahmann et al. (2000) using Multitracer (Woods et al., 2003) software on a tablet PC by an expert neuroanatomist (CW) who was blind to the age and sex of the children. Although tracings were largely performed in the sagittal view, all three planes were used to ensure three dimensional (3D) continuity. Tracings performed in the coronal and axial planes appeared as dots in the sagittal view and were used to define

lateral boundaries of structures. Tracings were performed at four times magnification. The following lobules of the cerebellum were traced on both the left and right hemispheres: lobules I-V, VI, Crus I, Crus II, VIIb, VIII, IX and X.

The right hemispheres of 10 randomly selected subjects were re-traced at a later time by CW in order to compute intra-rater reliabilities. Cerebella of 14 test subjects were traced by a different neuroanatomist (NB) using the same protocol. Inter-rater reliabilities for lobules of eight cerebellar hemispheres traced by both neuroanatomists were compared using intraclass correlation coefficients (ICC).

The midline slice was defined as the sagittal slice in which the cerebral aqueduct was most clearly visible. Left and right hemispheric subregions were traced separately. The anterior lobe comprises lobules I-V; lobule VI, Crus I and II, and lobule VIIb form the superior-posterior lobe; and lobules VIII, IX and X comprise the inferior posterior lobe. For each lobule, the areas demarcated by the drawn contours as belonging to that lobule were masked on all relevant slices and combined to construct a 3D volume of each lobule. The resulting volumes and their surfaces were inspected visually to identify and correct tracing errors (examples shown in Figure 2.1) in an iterative way. The resulting masks define 16 cerebellar compartments, 8 in each hemisphere, each of which was labelled with a unique integer value. Tracings for one hemisphere of one brain are shown in the sagittal plane in the right panel of Figure 2.2, with the corresponding masked areas and their color representations shown in the middle and left panels, respectively. For each subject we also generated a cerebellar mask comprising the sum of the gray matter parcellations and the total cerebellar white matter.



Figure 2.1 Illustration of Crus II volumes for the 18 training subjects with tracing errors in two (indicated by the black arrows), which were manually corrected by the expert neuroanatomist.



Figure 2.2 Cerebellar lobules and their color representations. Left and right hemispheric structures are represented using the same colors and unique labels. The image on the right shows manually traced contours in the sagittal view for one subject, while the middle panel shows the corresponding masked areas and their color representations on the left.

Construction of the Probabilistic Atlas

The T1 weighted images of the 18 training subjects were spatially normalized to the ageappropriate NIHPD asymmetric template (Fonov et al., 2011) using discrete cosine non-linear deformation. We used the SPM5 unified segmentation (Ashburner and Friston, 2005) method that uses a mixture of Gaussian models to simultaneously perform spatial normalization, bias correction, and tissue classification. The tissue probability maps (gray/white/cerebrospinal fluid (CSF)) of the template were used to classify tissues as gray matter, white matter and CSF. Further, all SPM's default settings were used. Following alignment of the T1 images to the template, individual lobular masks and cerebellar masks of each subject were resampled into the template space using nearest neighbour interpolation.

Co-registered cerebellar masks were averaged in the NIHPD template space to generate an average cerebellar mask. In this work, we chose to set probabilities of voxels with values of 0.3 or less equal to zero, as these voxels were part of the cerebellum in only 30% or fewer of the subjects, and values greater than 0.3 equal to unity. Using a threshold helps to reduce the effects of outliers and individual subject variance. The average of the whole brain images of the input subjects normalised to the NIHPD template space was multiplied by the resulting cerebellar mask to construct an average cerebellar image (Figure 2.3A).

In a similar way, resampled lobular masks in normalised space were averaged across subjects to generate probability maps for each structure, where the value in each voxel location denotes the probability of a voxel belonging to said structure. Typically, there is spatial variability of individual cerebellar structures between subjects even after alignment to a reference template. The probability maps indicate the proportion of subjects in whom a specific lobule occupies a location in the reference space and as such also quantifies the spatial variability in different regions. This enables one to visualize both the spatial extent of individual sub-regions and their spatial variability. Figure 2.3B shows the maximum probability map generated by combining the averaged lobular masks in normalised space. Each voxel is assigned a value equal to the maximum probability for that location. Further, voxels for which the maximum probability is 0.3 or less are set to zero as this indicates that

these voxels belonged to a cerebellar lobule in 30% or fewer of the subjects in the initial dataset.



Figure 2.3 A. Average cerebellar image in normalised space, R-Right, L-Left; B. Maximum Probability maps of individual structures; bright colors (white) indicate voxels with high probability and darker (red) colors indicate voxels with lower probability. C.Maximum likelihood labellings superimposed on the average image generated after spatial normalization with the NIHPD (7.5-13.5 years) asymmetric template. The colors represent the lobules of the cerebellum as per the color look up table in Figure 2.2. The coronal slices range from MNI coordinates -75 to -33.

From the maximum probability map, we also constructed an atlas of maximum likelihood labellings, which assigns to every voxel a label indicating the individual cerebellar structure located at that voxel most often. At the boundaries between structures, where a voxel may have equal probability of belonging to two different lobules, the label that occurs most often in a 3x3x3mm³ region surrounding the voxel was assigned to the voxel (Figure 2.3C).

Validation

The images of the 14 test subjects were pre-processed using the same pipeline as the training subjects. To segment the cerebella of the 14 test subjects using our CAPCA18 cerebellar atlas, the unified normalisation algorithm in SPM5 was applied to simultaneously perform tissue classification and normalisation of each subject's whole-brain images to the NIHPD asymmetric template. Labels from our CAPCA18 atlas were resampled to subject space using the inverse of the deformation warps obtained during normalisation.

We computed for each test subject both individual lobular volumes and total cerebellar gray matter volume. We compared the percentage of total gray matter volume that each lobule occupies in children from our study with values previously reported for adults (Makris et al., 2005).

For comparison, labelling was also performed using the SUIT cerebellar atlas that is available in SPM5 and was developed using adult data. Briefly, each subject's images were cropped to the region of the cerebellum after co-registration to the ICBM152 template. Individual cerebella were normalised to the SUIT template to obtain deformation maps for each subject. Manual editing was required around the anterior borders between the cerebellum and the temporal and occipital lobes for 8 of the test subjects. After editing, the cerebellar cortex achieved 78% average overlap with the SUIT template, which is higher than the 75% previously reported (Diedrichsen et al., 2009). Normalisation was repeated after manual editing and the labels from the SUIT atlas were resampled to the cropped test subject space using the inverse of the deformation warps obtained during normalisation. Finally, labels were resampled to native test subject space using the suit_reslice command.

Cerebellar parcellations derived from manual tracings were compared to those generated using our CAPCA18 atlas and the SUIT atlas using both volumetric measurements and the Dice coefficient as a measure of spatial overlap (Dice et al., 1945). CAPCA18 atlas-based labelling was compared to manual tracings in all 16 hemispheric regions. SUIT labelling differs from the conventions followed in this paper, in that the vermis is defined as a single medial entity in SUIT, while it was not traced separately in our work and as such was included in our hemispheric regions. In order to avoid errors due to omission or inclusion of the vermal regions when comparing data from SUIT to manual tracings, the left and right SUIT hemispheric regions and the SUIT vermal region were combined for each lobule to define whole lobules and compared to whole lobules (left plus right) from the manual tracings. As such, SUIT atlas-based labelling was compared to manual tracings in 8 whole lobules, comprising both hemispheric and vermal regions combined.

2.3 Results

Intra-rater reliabilities for 10 right hemispheres traced on two separate occasions by CW yielded ICCs ranging from 0.72 to 0.96 for different structures (Table 2.1). Only in the small lobules IX and X were ICCs below 0.85. Furthermore, inter-rater reliabilities for lobules of eight cerebellar hemispheres traced by two different neuroanatomists show ICCs greater than 0.8 in five of eight lobules traced. ICCs in VIIb and X were unacceptably low, indicating that these regions are difficult to trace reliably.

Table 2.1 Intra-rater reliability for 8 regions in 10 right hemispheres traced on two separate occasions by an expert neuroanatomist, and inter-rater reliability for eight lobules in eight hemispheres traced by two different neuroanatomists.

Cerebellar	ICC	ICC
Lobules	Intra-	Inter-
	rater	rater
I-V	0.91	0.92
VI	0.87	0.83
Crus I	0.85	0.81
Crus II	0.89	0.69
VIIb	0.85	0.43
VIII	0.96	0.88
IX	0.83	0.87
Х	0.72	0.46

Table 2.2 gives the average lobular volumes for the 14 test subjects after automatic segmentation using the CAPCA18 atlas. Volumes are expressed as a percentage of total cerebellar gray matter volume. The average gray matter volume for our test subjects is 113.2 \pm 11.5cm³, which compares favourably with gray matter volumes of 112.9 \pm 18.9cm³ reported by Makris et al. (2005) in adults. Crus I, Crus II and lobule VIIb together occupy 49.4% of the total cerebellar gray matter. For comparison, percentage lobular volumes reported by Makris et al. (2005) for adults are included in Table 2.2. Proportionally, lobules VI, VIIb and X comprise smaller fractions of the total cerebellar gray matter in children, while lobule IX is proportionately about 50% larger than in adults.

Table 2.2 Mean and standard deviation (SD) of lobular volumes generated from automatic segmentation of the cerebella in 14 test subjects using the CAPCA18 atlas. Volumes are expressed as a percentage of total cerebellar gray matter volume.

Lobule	Ri	ght	Le	eft	Total	Total Hemisphere	% Difference
	Hemis	spheric	Hemis	pheric	Hemispheric	Volume (Makris	between relative
	Vol	ume	Vol	ume	Volume	et al., 2005)	lobular sizes in
	(% 0	f total	(% of	ftotal	(% of total	(% of total	pediatric and adult
	cerebel	lar gray	cerebel	lar gray	cerebellar gray	cerebellar gray	cerebella
	matter	volume)	matter v	volume)	matter volume)	matter volume)	
	Mean	SD	Mean	SD	Mean	Mean	
I-V	5.9	0.3	6.7	0.4	12.6	12.1	4.1
VI	6.4	0.4	7.0	0.5	13.4	16.3	-17.8
Crus I	12.5	1.4	12.2	1.5	24.7	23.5	5.1
Crus II	9.3	1.2	8.5	1.2	17.8	16.4	8.5
VIIb	3.3	0.4	3.7	0.5	6.9	9.0	-23.3
VIII	7.2	1.0	6.6	0.8	13.8	14.9	-7.4
IX	4.7	0.5	4.9	0.5	9.6	6.4	50.0
Х	0.6	0.1	0.6	0.0	1.2	1.4	-14.3
Total	49.9	4.8	50.2	5.0	100.0	100.0	

Table 2.3 compares the mean lobular volumes for the 14 test subjects obtained from manual tracing with those generated from automatic segmentation with the CAPCA18 atlas. Volumes of lobule X were larger bilaterally after automatic segmentation compared to manual tracing (paired Student's *T*-test), as were left Crus I, left VIIb, and left IX. Left Crus II is the only region that was significantly smaller following automatic segmentation compared to manual tracing tracing.

In Figure 2.4, we present Dice coefficients comparing spatial overlap of automatic segmentations using the CAPCA18 atlas with manual tracings for the 16 hemispheric regions. The box and whisker plots indicate that, on average, spatial overlap between manual and automatic segmentations are 70% or higher in all regions, except lobules VIIb and X where it is of the order of 50% and 60%, respectively. Notably, these are the two regions where inter-rater reliabilities for manual tracings were unacceptably low.



Figure 2.4 Box-and-whisker plots of Dice coefficients that quantify spatial overlap in 14 test subjects between manual tracing and automatic segmentation using the CAPCA18 atlas. Red represents the right hemisphere and blue the left hemisphere.

Table 2.3 Comparison of mean and standard deviation (SD) of lobular volumes from manual tracing and automatic segmentation with the CAPCA18 atlas in 14 test subjects.

Cerebellar	Right			Left		
Lobules	Volume from Volume from		t	Volume from	Volume from	t
	Manual	Automatic		Manual	Automatic	
	Tracing (cm ³)	Segmentation		Tracing (cm ³)	Segmentation	
		(cm ³)			(cm ³)	
I-V	6.3 (1.1)	6.1 (0.3)	0.9	6.5 (1.2)	6.7 (0.3)	-0.8
VI	6.8 (1.1)	6.5 (0.1)	0.8	6.7 (1.2)	7.0 (0.1)	-1.2
Crus I	12.0 (2.1)	12.5 (0.2)	-1.3	11.0 (1.2)	12.1 (0.2)	- 4.1*
Crus II	9.5 (1.9)	9.2 (0.2)	0.9	10.4 (2.9)	8.4 (0.2)	3.9*
VIIb	3.6 (0.4)	3.3 (0.0)	2.3†	3.0 (0.4)	3.7 (0.1)	-4.3*
VIII	7.5 (1.2)	7.1 (0.2)	1.4	7.2 (2.0)	6.6 (0.0)	1.5
IX	4.2 (0.5)	4.6 (0.1)	-2.1†	4.3 (0.3)	4.9 (0.1)	-3.7*
Х	0.4(0.0)	0.6 (0.0)	-9.4***	0.4(0.0)	0.6 (0.0)	-12.9***

Values are Mean (SD); †p<0.1; *p<0.05; **p<0.001; ***p<0.0001

Figure 2.5 presents a color map of the Dice coefficients in the 16 hemispheric cerebellar regions for all 14 test subjects, where cold colors (cyan) denote poor spatial overlap and warm colors (red) denote high spatial overlap (>0.7). It is evident that spatial overlap drops below 0.6 in most subjects bilaterally in lobules VIIb and X. For the remaining regions, Dice

coefficients are typically above 0.7 in most subjects. Spatial overlap of manual and automatic total cerebellar gray matter was 86%.



Figure 2.5 Colormap of Dice coefficients of spatial overlap between automatic segmentation using CAPCA18 and manual tracing for 14 test subjects (TS) bilaterally in 8 hemispheric cerebellar regions.

Table 2.4 Comparison of mean and standard deviation (SD) of volumes of cerebellar lobules obtained from manual tracing and SUIT segmentation in 14 test subjects.

Cerebellar	Cerebellar hemispheric volumes				
Lobules	Volume from Volume from		t		
	Manual	SUIT			
	Tracing	Segmentation			
	(cm ³)	(cm ³)			
I-V	12.8 (2.1)	14.3 (2.5)	-1.6†		
VI	13.5 (2.0)	18.3 (3.0)	-5.9***		
Crus I	23.0 (1.8)	20.0 (4.4)	2.5*		
Crus II	19.9 (1.7)	15.0 (4.2)	3.8**		
VIIb	6.7 (0.9)	7.5 (1.9)	-1.5†		
VIII	14.7 (2.2)	14.0 (3.2)	0.8		
IX	8.6 (1.2)	5.8 (1.2)	5.3***		
Х	0.8(0.1)	0.9(0.2)	-1.6†		

Values are Mean (SD); †p<0.1; *p<0.05; **p<0.001; ***p<0.0001

Table 2.4 presents a comparison of mean whole lobular volumes (left and right hemispheres and vermis combined) in the 14 test subjects from segmentation using the SUIT atlas with

volumes obtained from manual tracings. Hemispheric volumes from SUIT segmentation were significantly different to those from manual tracing in four of eight lobules (paired Student *T*-test), and tended to be larger, although below conventional levels of significance, in a further three lobules. Volumes were equivalent only in lobule VIII. Notably, one of the regions that SUIT overestimates is lobule VI, which is proportionately smaller in children than in adults, while lobule IX which is underestimated with SUIT is proportionately larger in children.

Figure 2.6 compares Dice coefficients that quantify spatial overlap in the 14 test subjects between manual tracing and automatic segmentation using either the SUIT (red) or CAPCA18 (blue) atlases in 8 whole lobules, when left and right hemispheres (and vermal regions in the case of SUIT) have been combined. CAPCA18 performs significantly better (Student *T*-test) than SUIT in lobules I-V, VI, CrusI, VIII and IX.

Figure 2.7 compares in 14 test subjects for total cerebellar gray matter spatial overlap

Figure 2.6 Box-and-whisker plots of Dice coefficients that quantify spatial overlap in 14 test subjects between manual tracing and automatic segmentation using either SUIT (red) or CAPCA18 (blue) for whole lobules. $\dagger p < 0.1$; $\ast p < 0.05$; $\ast p < 0.01$; $\ast p < 0.01$

between manual tracing and automatic segmentation using SUIT (green) with spatial overlap between manual tracing and automatic segmentation using CAPCA18 (yellow). In all except one subject, spatial overlap of manual tracings with CAPCA18 segmentations are higher than between manual tracings and SUIT segmentation. On average, gray matter spatial overlap using SUIT before and after manual editing were 75% and 78%, respectively, while it was 86% using the fully automated CAPCA18 atlas.

Figure 2.7 Spatial overlap of total cerebellar gray matter from manual tracing with total cerebellar gray matter from automatic segmentation using either SUIT (green) or CAPCA18 (yellow) in 14 test subjects.

2.4 Discussion

We have presented a probabilistic three dimensional cerebellar atlas for children with parcellations into 16 macroanatomic structures and performed validations comparing manual tracing with automatic segmentation using either the SUIT atlas or our new CAPCA18 atlas using both volumetric measurements and Dice coefficients. The probabilistic cerebellar atlas encodes the individual subject variability in different cerebellar structures, while the maximum likelihood labelling assigns to each voxel a unique label according to the label most often assigned to a voxel in that location. Average spatial overlap of the whole cerebellar cortex for pediatric test images with their corresponding manual gray matter segmentations using our probabilistic CAPCA18 atlas was 86%, compared to 78% after

normalisation to the SUIT template. This finding suggests that our pediatric cerebellar atlas helps to reduce bias and segmentation errors that might occur from using an atlas constructed from adult data.

We also computed spatial overlap in individual cerebellar lobules and achieved spatial overlap in both hemispheres of 70% or higher in all lobules, except lobules VIIb and X where it was of the order of 50-60%. Notably, these are the smallest cerebellar lobules, comprising only 1.2% (lobule X) and 6.9% (lobule VIIb) of the total cerebellar gray matter, so that even small differences will result in low Dice coefficients. These are also regions that are difficult to trace reliably as evidenced by the poor inter-rater reliabilities in these regions. The accuracy of structure segmentation depends on its spatial location and tissues surrounding the structure. The inferior lobe VIIb comprises a very thin layer of gray matter and the inclusion of additional white matter in the manual tracing of this structure in the training subjects appear to have resulted in lower accuracies when automatically segmenting this area. Lobe X is the smallest structure in the cerebellum making it very sensitive to errors.

In all regions except lobules VI and IX, Dice coefficients for the children in our study using CAPCA18 compare favourably to values reported in adults by Bogovic et al. (2013) who used an automatic classification of cerebellar lobules algorithm that employs an implicit multi-boundary evolution (ACCLAIM) approach. The ACCLAIM method relies on good contrast between CSF and GM and high spatial resolution (0.828x0.828x1.1mm³) and as such may perform less well in our pediatric data.

Fissures in an average child brain are not as clearly distinguishable as in an average adult brain and the gray matter to white matter ratio also tends to vary in this population (Fonov et al., 2011). As a result, improved co-registration and segmentation results are expected using a cerebellar atlas for children. To our knowledge, this is the first pediatric cerebellar atlas with 16 parcellations that can serve as a better spatial prior to analyze cerebellar morphometry in children.

The average percentage gray matter volume occupied by each lobule in the cerebellar hemispheres were compared with values previously reported by Makris et al. (2005) to examine potential relative size differences in children compared to adults. It was found that lobule IX is proportionately larger in children, while lobules VI, VIIb and X are proportionately smaller. The increased volume in lobule IX in children may be due to increased gray matter around the posterior lobes of the cerebellum. It has been shown previously, when deforming pediatric brains to the adult ICBM152 nonlinear template, that gray matter around the posterior lobe is increased in children and that the shape and ratio of gray and white matter in children is different to that of adults (Fonov et al., 2011), with differences ranging from 4 to 23%. The good spatial overlap achieved between manual tracing and automated parcellation using CAPCA18 in lobule IX provides further support for the fact that this region is relatively larger in children.

We have used publicly available SPM software to construct the atlas, which ensures that our CAPCA18 atlas can be easily used for both segmentation and localisation in morphometric and functional imaging studies, respectively.

For the data used in the current study no manual editing was required when performing segmentation using the newly presented CAPCA18 cerebellar atlas. It is, however, important to check the images generated after initial pre-processing as well as after each intermediate step to ensure that no errors are introduced in the pipeline when computing the deformation between the atlas and test subject.

One limitation of the present study is that the number of boys and girls were not equal in either the training or test data sets. In the training data set there were twice as many girls as

51

boys, while 70% of the test subjects were boys. This could have introduced bias into the atlas, which may have resulted in poorer segmentation performance in the test subjects. We examined gender differences by comparing lobular volumes from manual tracings in boys and girls. Although lobular volumes in both hemispheres were equivalent between boys and girls (all p's > 0.1) in this small sample, we expect that the atlas may be improved if the training data set had equal numbers of boys and girls.

2.5 Conclusion

This paper presents the CAPCA18 pediatric cerebellar atlas which can be used to automatically segment the cerebella of children into 16 lobules. The segmentation was validated in 14 test subjects from the same age group as children whose data were used in the construction of the atlas. We believe that the maximum probability maps and the maximum likelihood labellings will be a useful tool in analysing the cerebellum in structural and functional imaging studies in children. Future work will focus on improving the accuracy of individual parcellations and performing validation in a larger number of subjects. Higher dimensional registration and using multiple atlases for label fusion can potentially improve the segmentation.

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Chapter 3

Assessment of different label propagation methods for segmenting cerebellar structures in adults using the CAPCA18 pediatric cerebellar

atlas

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Abstract

Effective methods for the propagation of cerebellar labels from structural atlases facilitate studies of cerebellar morphometry. The cortical cerebellar structures in our atlas were manually labelled in 18 healthy pre-adolescent children (training data) and comprise 8 labels per hemisphere. Typically, labels are propagated from a single probabilistic atlas or from individual atlases of the training dataset (multi atlas approach). Here we propagate the training data labels generated from a pre-adolescent population to 35 adult test subjects after establishing spatial correspondence between them using efficient and robust linear and nonlinear registration tools. This paper compares the performance of three different label propagation methods for labelling cerebellar structures. First we use a probabilistic atlas constructed from the training dataset and propagate the atlas labels to test subjects by warping the label map directly on to the test image. We compare this direct warping approach to two different multi atlas label propagation methods: multi atlas majority voting (MAMV; Heckemann et al., 2006) and multi atlas generative model (MAGM) based label propagation (Iglesias et al., 2012). Every training image is first registered to each test image in a pairwise manner (Postelnicu-Zöllei et al., 2009), generating 18 registered training atlases for each test

image. Label propagation on each test image is achieved by combining the deformed output segmentations iteratively for all structures. The improved final multi atlas segmentation is thus derived from 18x35 segmentations for the whole test dataset. The resulting segmentations from these three methods were compared with manual tracings available for 3 test subjects using the Dice coefficient, the modified Hausdorff distance, and volumetrics. Further, the performance of the three methods was evaluated using the simultaneous truth and performance level estimate (STAPLE) (Warfield et al., 2004). Results indicate that multi atlas based label propagation methods consistently perform better than direct warping. The multi atlas based segmentation captures all possible variations present in the training data and helps to achieve precise segmentations closer to manual segmentations.

3.1 Introduction

Magnetic resonance imaging (MRI) is a non-invasive method to study neuroanatomical structures of the brain *in vivo* and is useful to advance our understanding of neurological disorders (Thompson et al., 2004) and evaluate longitudinally (Reuter et al., 2012; Bernal-Rusiel et al., 2012) neuropathology and treatment (Thompson et al., 2007; Gering et al., 2001). In general, parcellated brain atlases are generated by manual delineation of structures of interest on MR images acquired from a group of subjects by an expert neuroanatomist adhering to standard protocols. This manual method is extremely time consuming and subject to inter- and intra-rater variations. To eliminate this, automatic parcellation of structures can be performed using already parcellated brain atlases that produce reproducible and consistent results compared to manual segmentation. Only a few whole brain parcellated atlases, such as the automated anatomic labelling atlas (Tzourio-Mazoyer et al., 2002) and LONI probabilistic brain atlas LPBA40 (Shattuck et al., 2008), are publicly available for parcellation/segmentation of brain structures. Popular software packages like FreeSurfer (http://surfer.nmr.mgh.harvard.edu/fswiki) are often used in brain mapping to study cortical
thickness and perform morphometric analysis over the whole brain. However, the automated parcellation is limited to cerebral structures limiting its usefulness for morphometric analyses of cerebellar structures. Using recent advances in registration and label propagation methods, this work aims to improve methods for automatic parcellation of cerebellar structures.

Average intensity images (templates) are usually presented in a stereotactic coordinate system, such as Montreal Neurological Institute (MNI) or Talairach space. The standard whole brain MNI305 template was constructed by registering 305 scans using 9-parameter linear transformation. The next generation template released by MNI, popularly known as the International Consortium of Brain Mapping (ICBM152) template (Evans et al., 1993), was constructed from co-registration of images from 152 subjects. Templates built in stereotactic coordinate systems aid inter-subject comparisons and alleviate the complexities arising from structural variability between individual brains. The historical evolution of different whole brain templates using linear and non-linear registration has been well documented by Evans et al. (2012). The ICBM152 template serves as the registration target in analysis tools such as SPM (Ashburner et al., 1999) and FSL (Smith et al., 2004). The template, however, has poor signal to noise ratio (SNR) and lacks anatomical detail to distinguish the subregions of the cerebellum. Recently, unbiased non-linear age-specific templates for children in various age categories have been made available by Fonov et al. (2011) along with their corresponding tissue priors. These templates have higher SNR and increased resolution compared to the classic ICBM152 template.

Parcellations on an average template image serve as a map of the spatial arrangement of anatomical structures of the brain and is called an atlas. Maximum likelihood label maps indicate the label that occurs in a specific voxel in the largest number of subjects during atlas construction, while probabilistic maps provide a measure of the uncertainty of a specific label assignment. The CAPCA18 atlas used in our study provides labels for 16 distinct cerebellar structures, 8 in each hemisphere, namely lobules I-V, VI, Crus I, Crus II, VIIb, VIII, IX and X.

Label propagation refers to the process of transferring labels from an atlas to a test image following some sort of spatial transformation and is highly dependent on the registration framework that is used to initially establish the spatial correspondence. Registration algorithms (Woods et al., 1993; Ashburner et al., 1999; Fischl et al., 1999; Jenkinson and Smith 2001; Joshi et al., 2004; Postelnicu-Zöllei et al., 2009) differ primarily in their objective function, the optimization method used, the degrees of freedom, and the number of parameters used. Registration methods are primarily classified into volume- or surface-based where the former uses intensity differences and the latter uses topological properties and geometric features of the cortex to guide the registration. A recent study comparing surfaceand volume-based registration (Ghosh et al., 2010) clearly demonstrated the superior performance of surface-based approaches. The spatial transformation generated from registration is then applied to the label map to propagate the labels to the test image. The accuracy of the resulting segmentation decreases if the anatomy of the atlas is very different from that of the test subject. In such cases, one possible solution is to use multi atlas based segmentation methods that better capture the underlying anatomical variation by using multiple registrations. The multiple atlas concept was originally demonstrated for segmenting microscopy images of bee brains (Rohlfing et al., 2004). Label propagation in human brain images using multi atlas majority voting (MAMV; Heckemann et al., 2006) is achieved by registering several atlases to the test image and then combining the resulting segmentations to obtain the final segmentation of the test image. Recently, a multi atlas generative model

(MAGM) approach (Iglesias et al., 2012; Iglesias et al., 2013) was used to apply the final enhanced segmentation of the test subject. This method is independent of the intensities of the training images and exploits the relationship between the volume of the test image and its propagated labels. Previous studies have shown that multi atlas approaches increase the accuracy and robustness of cortical segmentations with label fusion (Wang et al., 2013; Asman et al., 2013).

The cerebellum, which is much smaller than the cerebrum, is functionally heterogeneous with different topological regions responsible for various sensory, motor, cognitive, and affective processing (Stoodley and Schmahmann, 2009; Schlerf et al., 2010; Strick et al., 2009; Makris et al., 2005). The cerebellar atlas in MNI coordinate space developed by Schmahmann et al. (2000) details the gross morphology of the cerebellum and its arrangement of fissures. Diedrichsen et al. (2009) constructed a probabilistic atlas of the cerebellum and brainstem, popularly known as the Spatially Unbiased Infratentorial Template (SUIT), by warping images of 20 healthy adults into MNI space. This probabilistic atlas provides the probability with which labels occur at each spatial location as opposed to segmentation *per se*. Recently, cerebellar segmentation in adults using SUIT was compared with multi object deformable models that were custom built from manual delineations of cerebellar structures (Bogovic et al., 2013). Performance of the latter was shown to be superior compared to both SUIT atlasbased segmentation and multi atlas approaches.

To our knowledge, the performance of different automated label propagation methods has not been established for cerebellar segmentation. The present study aims to compare the accuracy of cerebellar segmentation in 35 adult test subjects using three different label propagation methods applied to the CAPCA18 atlas that was constructed from manual tracings of cerebellar structures in 18 healthy pre-adolescents. Volumes are compared in the 16 cerebellar regions, 8 in each hemisphere. Finally, we evaluate the segmented output using the Dice coefficient and the modified Hausdorff distance metric in 3 representative test subjects compared to manual segmentations by an expert neuroanatomist (CW).

3.2 Materials and Methods

CAPCA18 Atlas

The details of the CAPCA18 atlas construction are provided in the previous chapter. Briefly, high resolution T1-weighted images were acquired on a 3T Allegra MRI scanner (Siemens, Erlangen, Germany) in the sagittal plane using a magnetization-prepared rapid gradient echo (MPRAGE) sequence in 18 healthy pre-adolescent children (mean age 11.8±1.2 years, range 9-13 years, 6 males) who had been recruited as control subjects for on-going studies on fetal alcohol spectrum disorders (Jacobson et al., 2011) in Cape Town, South Africa. The cerebella of the 18 children were segmented into 16 regions, 8 in each hemisphere, by an expert neuroanatomist (CW) using Multitracer software (Woods et al., 2003) according to the nomenclature defined by Schmahmann et al. (2000). Each cerebellar region includes some white matter due to the fine branching structure of the cerebellar white matter making it difficult to exclude during the delineation process. Each child's brain was warped to the ageappropriate NIHPD asymmetric template using the SPM5 unified segmentation approach. Co-registered images were averaged to generate an average cerebellar image in normalised space. Further, the same transformations were applied to the individual lobular masks and the cerebellar masks of each subject, which were then averaged to construct the probabilistic atlas and the maximum likelihood label maps in the NIHPD template space.

MRI protocol for adults (test dataset)

High resolution MRI scans were acquired on a 1.5T Vision system (Siemens, Erlangen, Germany) for 35 adult subjects as part of studies being conducted at the Washington University Alzheimer's Disease Research Center (ADRC) in St. Louis. T1-weighted magnetization prepared rapid gradient echo (MPRAGE) scans were obtained according to the following protocol: two sagittal acquisitions, FOV = 256, matrix = 256 x 256, resolution = 1x1x1.25 mm³, TR 9.7ms, TE 4ms, flip angle 10 degrees, TI 20ms, TD 200ms. The two acquisitions were averaged to increase the contrast-to-noise ratio. Descriptions of these images have appeared in prior publications (e.g., Buckner et al., 2004; Fotenos et al., 2005).

Label propagation using direct warping

Skull stripped intensity normalised images of the test subjects were co-registered to the average image generated from the co-registered training images. The image volume of each test subject was registered to the target image using a two-step volumetric registration. First we applied linear registration using FLIRT (http://fsl.fmrib.ox.ac.uk/fsl/), followed by non-linear volumetric (MRI_NL_ALIGN) registration in Freesurfer (http://surfer.nmr.mgh.harvard.edu/fswiki). The deformation field was geometrically driven and diffused to the volume based on elasticity to align the volumetric images (Postelnicu-Zöllei et al., 2009).

The registration yields a deformation from each test image to the atlas. By applying the inverse transformation, the maximum likelihood label map of the CAPCA18 atlas was warped to the native coordinate system of the test image and could be used directly to propagate individual labels to the test subject. Figure 3.1 illustrates the steps involved in

generating the automated segmentation of the cerebellar structures for a test subject using this direct warping approach.





Label Propagation using multi atlas segmentation

The images of each of the training subjects and their respective label maps generated from the manual tracings are collectively referred to as the multi atlas.

The images of the training subjects were processed using Freesurfer software (http://surfer.nmr.mgh.harvard.edu/). The processing steps included skull stripping (Ségonne et al., 2004), intensity normalisation (Sled et al., 1998), cortical and sub-cortical structure segmentation, surface generation, registration to the spherical atlas, and cortical thickness calculation (Dale and Sereno, 1993; Dale et al., 1999; Fischl et al., 1999). The surfaces were superimposed on the normalised T1 weighted images and visually assessed for correctness of gray-white and gray-CSF boundaries. White matter boundaries were edited manually to

ensure the correct surface generation. Freesurfer quality control procedures are described in detail at the following link <u>http://surfer.nmr.mgh.harvard.edu/fswiki/FsTutorial/TroubleshootingData</u>. The surfaces of the training subjects were edited to aid the surface registration and to achieve better alignment of the folding patterns between the brains of different subjects.

After pre-processing, the surfaces from the training subjects were used as the moving sources and the surfaces of the test subjects as the targets for combined volume- and surface-based (CVS) registration. Every training subject's surface was registered to each test subject's surface (i.e. 18x35 registrations). The robust CVS registration tool (Postelnicu-Zöllei et al., 2009) achieves good correspondence between cortical folding patterns and between the subcortical structures of the brain. The method uses the geometric information from the warp and incorporates volume registration using Navier elasticity. The resulting non-linear displacement fields are used to register both cortical and subcortical structures. We performed the complex registration on a high performance computer cluster facility which has 127 batch nodes, two quad-core Xeon 5472 3.0 GHz CPUs and 32 GB RAM. All nodes were running the 64-bit version CentOS 6. The pair-wise registration between the training and test surfaces yielded 630 (18x35) registered outputs and their associated deformation morphs.

To obtain the final segmentation of each of the 35 test subjects, we combined the multiple registered atlases resulting from the CVS registration (volume resampled output) of the training atlases. Two different label propagation techniques were used: multi atlas majority voting (MAMV) (Heckemann et al., 2006) and multi atlas generative model based label propagation (MAGM; Iglesias et al., 2012). Figure 3.2 shows a flow chart of our processing

pipeline for multi atlas based segmentation.



Figure 3.2 Flow chart showing the processing pipeline for multi atlas based segmentation. Every image in the training set is registered to the individual test subject's image, whereafter the resulting deformation is applied to each training atlas. Two different label fusion strategies are used to propagate labels from the registered training atlases to the image of the test subject.

Majority voting assigns a label to each voxel of the test image by determining the label that occurs most frequently in the corresponding voxel on all the registered training atlases. All training images are weighted equally and it is assumed that each training atlas represents an accurate segmentation. The effects of independent noise associated with a particular label are reduced using this approach.

The generative model approach uses probabilistic modelling of segmentations from a set of segmented images to generate a second set of final segmentations. The model is independent of the intensity of the training images. The intensity of the test image and the respective deformed outputs are used to model the final segmentation of the test image. The mathematical formulation of the model was derived from the implementations by Sabuncu et al. (2010) and Iglesias et al. (2012). The model includes a single model parameter β , where β

= 0 treats each voxel independently, $\beta = \infty$ corresponds to a single label map, and β = finite value yields the combined output. Typically, in our experiments we used β =0 to obtain the fused output from MAMV and β =0.3 for MAGM.

Validation

Spatial overlap

We validated the segmentation in three randomly selected test images using the Dice coefficient as a measure of spatial overlap between automatic segmentations using the three different label propagation approaches and manually traced regions. The Dice coefficient (Dice et al., 1945) is given by

Dice coefficient (L) = $\frac{2*(M \cap T)}{M+T}$ \forall L=1, 2, ... 16,

where M represents the manual segmentation, T represents the automated segmentation of the test image, and L represents the number of labels, i.e. the number of distinct segmented regions. The three test images were manually traced by the same expert neuroanatomist (CW) who performed the manual tracings of the training images. Further, lobular volumes were computed and compared in the three test subjects for segmentations generated using the different label propagation methods and manual tracings. Analysis of variance (ANOVA) was used to examine volumetric differences between the different methods.

Modified Hausdorff Distance

The modified Hausdorff distance computes similarity of shapes (Dubuisson and Jain, 1994) and was used to compare for the same three test subjects automated segmentations achieved using the three different label propagation approaches with manual segmentations. H_d denotes the modified Hausdorff distance between the manual and automated segmentations,

d(M, T) is the distance of the manually traced surface M from the automated segmentation surface T, and d(T,M) is the distance of the surface T from the surface M. The modified Hausdorff distance is given by:

 $H_d(M,T) = \max(d(M,T), d(T,M)), \text{ where}$ $d(M,T) = \frac{1}{Nx} \sum_{x \in My \in T} ||x - y||_2 \text{ and}$

Nx is the number of elements in the set. Typically, the distance is calculated from the boundary voxels of M to its nearest boundary voxel in T and vice versa for individual label volumes. If the boundaries of segmentations match, the distance will be zero and will increase as the differences in the boundary increase.

Validation using STAPLE

The structural segmentations of the 35 test images resulting from the three different label propagation methods were further compared using the Simultaneous Truth and Performance Level Estimation (STAPLE) algorithm (Warfield et al., 2004). The STAPLE algorithm effectively estimates the probabilistic true segmentation and also measures the performance of the algorithm/rater, given the input segmentations from the different rater or segmentation algorithms of an image. It uses expectation maximization for estimating the true segmentation and determines the performance parameters of the results from a collection of segmentations and a prior model. The Dice coefficient was recorded using STAPLE.

3.3 Results

Table 3.1 presents average lobular volumes for three test subjects using each of the label propagation methods compared to manual tracing. Other than lobules left I-V and left VIIb, volumes using the four different methods are similar. In left I-V, direct warping yields inflated volumes, while volumes of VIIb were smaller for all the automated segmentation methods compared to manual tracing.

Spatial overlap and modified Hausdorff distances

Figure 3.3 shows box-and-whisker plots comparing the Dice coefficients and modified Hausdorff distance metrics for automated segmentations using the three different label propagation approaches, each compared to manual tracings, in three test images for right and left cerebellar lobules. Except for lobule VIIb, Dice coefficients are at least as good or better for the two multi atlas approaches compared to direct warping. Except for lobule VIIb, Hausdorff distances are lowest for MAGM.



Figure 3.3 Box and whisker plots of Dice coefficients (top row) and Hausdorff distances (bottom row) in three randomly selected test subjects for 8 structures per cerebellar hemisphere for automated segmentations using three different label propagation strategies, each compared to manual tracings. (1): Direct warping (magenta); (2): multi atlas majority voting (MAMV; green); (3): multi atlas generative model (MAGM; blue).

Label	Structure		-					
ID	names		F	р				
		Maria						
		Tracing	Direct	MAMV Lobular	MAGM Lobular			
		Lobular	Lobular	volumes	volumes			
		volumes	volumes	volumes	volumes			
		(mm^3)	(mm ³)	(mm^3)	(mm^3)			
LOIL	· 1	()	()	()	()			
Left He	emisphere							
1	I-V ^a	4506 (577)	5892 (939)	4410 (188)	4784 (151)	4.4*	0.04	
2	VI	7078 (428)	5785 (1354)	5716 (596)	6411 (537)	1.8	0.2	
3	Crus I	11520 (2732)	13890 (1981)	11353 (1816)	11709 (1886)	0.9	0.4	
4	Crus II	8148 (2464)	9889 (535)	11185 (790)	10596 (1248)	2.4	0.1	
5	VIIb ^b	4322 (404)	3239 (217)	2628 (76)	2908 (261)	23.2**	0.0002	
6	VIII	7357 (830)	7949 (998)	7077 (124)	6601 (622)	1.8	0.2	
7	IX	2892 (809)	4283 (546)	3778 (463)	3883 (625)	2.6	0.1	
8	Х	482 (121)	498 (102)	367 (86)	435 (78)	1.1	0.4	
Right Hemisphere								
9	I-V	6227 (1011)	6599 (978)	5288 (637)	5695 (458)	1.5	0.2	
10	VI	7744 (300)	6512 (1704)	5546 (1299)	6286 (1092)	1.7	0.2	
11	Crus I	11804 (3133)	14408 (971)	11673 (419)	12217 (512)	1.7	0.2	
12	Crus II	8452 (2278)	10071 (323)	11148 (610)	10846 (804)	2.7	0.1	
13	VIIb	3478 (521)	3922 (217)	2762 (887)	2627 (699)	2.8	0.1	
14	VIII	7096 (261)	7350 (776)	6076 (687)	6008 (541)	3.9†	0.05	
15	IX	4236 (787)	5962 (572)	5422 (836)	5604 (770)	2.8	0.1	
16	Х	446 (100)	480 (90)	436 (83)	579 (122)	1.2	0.3	

 Table 3.1 Comparison of average lobular volumes for three randomly selected test subjects obtained using manual tracing and the three different label propagation methods.

Values are Means (standard deviations) †p<0.1; *p<0.05; **p<0.001; ***p<0.0001

^aDirect warping > manual (*p*=0.01), Direct warping > MAMV (*p*=0.01), Direct warping > MAGM (*p*=0.04) ^bManual > Direct warping (*p*=0.001), Manual > MAMV (*p*<0.001), Manual > MAGM (*p*<0.001)

Visual Assessment of lobule I-V boundaries for different segmentation approaches

In Figure 3.4 is shown superimposed on an intensity normalized test image the contours from the three different label propagation methods and manual tracing (shown in blue) for left lobule I-V. The contours are shown on a coronal (top) and mid-sagittal (bottom) slice. Visually, differences between manual tracing (blue) and the multi atlas label propagation (red and white) contours are smaller. Direct warping (yellow) appears to overestimate the area by erroneously including voxels around the boundary of the cerebellum. MAGM produces the best results and is the only one of the automated methods that successfully excludes some of the finer white matter branches evident on the sagittal slice.



Figure 3.4 Boundaries for segmented left lobule I-V superimposed on an intensity normalised structural T1 image on a single coronal (top) and sagittal (bottom) slice. The colour overlays indicate the boundaries obtained using the different segmentation methods: manual tracing (blue), direct warping (yellow), multi atlas majority voting (MAMV; red), and multi atlas generative model label fusion (MAGM; white).

In Figure 3.5 we compare the area of left lobule I-V as per the different segmentation methods on all the sagittal (top) and coronal (bottom) slices on which it appears in a single test subject. We observe that in most of the slices, except near the edges of the structure, the difference in area between manual tracing and direct warping tends to be larger than for the two multi atlas label propagation methods.



Figure 3.5 Bar plot of the area of left lobule I-V for the different segmentation methods on different sagittal (top) and coronal (bottom) slices: Manual segmentation (Blue); Direct warping (Cyan); MAMV (yellow); MAGM (brown).

Figure 3.6 shows for two different coronal slices (a and b) of one test subject the maximum likelihood label maps and their associated boundaries as generated using direct warping (left), MAMV (middle), and MAGM label fusion (right). While all three methods achieved good spatial overlap with the cerebellum as a whole, it is evident that the transverse sinuses (indicated by the yellow arrows) are erroneously included as part of the cerebellum only in the direct warping segmentation. By contrast, the two multi atlas methods that benefit from additional information related to inter-subject variability correctly exclude the sinuses. The red arrows indicate regions where segmentation using MAGM label fusion generates contours that more closely follow the cerebellar gray matter boundary than the other two methods.



Figure 3.6 Maximum likelihood label maps and their associated lobular boundaries for two different coronal slices (a and b) superimposed on normalised T1 images of one test subject. Segmentations were generated using direct warping (left), multi atlas majority voting (MAMV; middle), and multi atlas generative model label fusion (MAGM; right). The look up table and colours representing each structure are listed in Figure 2.2. The visualization and the colour overlay was generated using Freeview software (<u>http://www.nmr.mgh.harvard.edu/martinos</u>). Yellow arrows indicate the transverse sinuses that are erroneously included in Crus I using direct warping segmentation. The red arrows indicate areas where MAGM contours follow the cerebellar gray matter boundaries more closely.

Figure 3.7 presents the Dice coefficient in terms of box and whisker plots between the ground truth estimate from STAPLE and label propagated segmentations using the three different methods for the 35 test subjects. The STAPLE method allows one to assess overall performance of segmented outputs in the absence of manual delineations for all subjects. MAGM consistently produces the highest Dice coefficients across all cerebellar lobules.



Figure 3.7 Box and whisker plots for right (top) and left (bottom) cerebellar lobules of Dice coefficients comparing the STAPLE truth estimate for 35 test subjects to segmentations obtained using 1. direct warping (magenta); 2. MAMV label propagation (green); and 3. MAGM label propagation (blue).

3.4 Discussion

In this work we present results for three different label propagation strategies, namely direct warping, MAMV, and MAGM label fusion, to automatically segment 16 cerebellar lobules in 35 adult test subjects using our CAPCA18 atlas. For three subjects that had been manually traced, Dice coefficients that assess spatial overlap for each cerebellar structure with the manually traced ground truth were typically higher for the two multi atlas approaches, while Hausdorff distances that assess deviations in shape from the ground truth were lowest for MAGM. Further, it was shown for lobule I-V that the segmented areas on nearly all slices from the two multi atlas approaches are more similar to those computed using manual tracing, while direct warping consistently overestimates the area. Dice coefficients comparing segmentation results in all 35 test subjects with the STAPLE truth estimate consistently yielded higher values for the two multi atlas approaches, with the highest values for MAGM.

These results demonstrate the power of multi atlas label propagation strategies that are able to account for the underlying anatomical variability between subjects. The accurate alignment of cortical surfaces that is achieved through the pairwise combined volume and surface based registration helps to capture all possible variations among the training subjects. One advantage of multi atlas methods is that they are less sensitive to age related differences, which might otherwise result in poor segmentation accuracy in situations where the ages of the test subjects are very different to those of the training subjects used in creating the atlas. This is evident in the high Dice coefficients (>0.9 in all lobules except lobule VIIb, >0.8 in lobule VIIb) achieved in the present adult data set when comparing segmentations to the STAPLE truth estimate, considering that the atlas was created from 18 pre-adolescent children.

These results also serve to demonstrate that multi atlas approaches (especially MAGM) work well in situations where the training and test data were acquired using different sequences on different scanners. The method does not depend on the similarity of intensities across the atlases and the test volume, but on the consistency of intensities within regions in the test volume. This quality makes the method well suited to parcellation of cerebellar data acquired using different MR modalities.

In the three test subjects for whom gold standard manual tracings were available, the segmentation accuracy was worse for the two multi atlas approaches than direct warping *only* in lobule VIIb. Anatomical variability in this region and the intersection of white matter fibers posterior to the neighbouring structure poses difficulties in determining the border voxels of this structure and may contribute to inaccuracies in this area. Further, lobule VIIb is a small region sandwiched between two larger lobules, Crus II and lobule VIII. In this region, the surface to surface distance metrics for the two multi atlas approaches are high, as might occur due to differing boundaries across the entire structure, suggesting that the entire region may be displaced.

3.5 Conclusion

In this paper we have presented and compared three different label propagation strategies for automatic cerebellar segmentation using the CAPCA18 atlas. Multi atlas based approaches were shown to consistently perform better than direct warping, both in terms of higher Dice coefficients and reduced Hausdorff distances.

Chapter 4

Multi Atlas based labelling of cerebellar structures

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Abstract

Multi atlas based segmentation has recently become popular in medical image segmentation. Atlas based segmentation framework allow one to incorporate prior segmentation knowledge by exploiting spatial information gathered via registration of images from multiple previously analyzed subjects. However, limited knowledge is available about spatial locations that exhibit large inter-subject variability. In this study, we present an optimized pipeline to obtain automated cerebellar segmentations of 35 test subjects using manual delineations of 28 cerebellar structures in 20 training subjects and two different multi atlas label fusion methods. We assess the accuracy of our segmentation for the 20 hemispheric cerebellar regions in the test data using simultaneous truth and performance level estimates and Dice coefficients of spatial overlap. We also evaluate segmentation accuracy using leave-one-out cross-validation in our training data for which gold standard manual tracings are available. The training and test data were collected in two different laboratories on two different scanner platforms. Mean Dice coefficients in the test subjects were greater than 0.89 in all regions except lobules VIIb, VIIIa, VIIIb, and X.

4.1 Introduction

Magnetic Resonance images (MRI) require a reliable pipeline to generate segmentations of various brain structures for quantitative analysis. Typically, a series of steps are performed before one can analyse the structure of interest (Pierson et al., 2002; Gouttard et al., 2007; Powell et al., 2008; Diedrichsen et al., 2009; Bogovic et al., 2013). These include brain extraction, also known as skull stripping, which is the process of removing non-brain tissue (Shattuck et al., 2001; Smith et al., 2002; Segonne et al., 2004; Sadananthan et al., 2010) and dura from the images, followed by intensity normalisation (Sled et al., 1998), bias correction, spatial normalisation and tissue classification (Ashburner and Friston 2005).

Atlas-based techniques have been used for the segmentation of various structures on images acquired from a range of non-invasive imaging modalities (Gouttard et al., 2007; Shattuck et al., 2008). These approaches are supervised methods where spatial priors are used to guide the segmentation. Conventionally, a single atlas image that comprises different labelled structures is used to generate the segmentation on an unknown (test) image. However, this method allows no control over errors that arise from poor registration when the test image is very different to the atlas image (Aljabar et al., 2009), which may result in incorrect labels being assigned to voxels.

Multi atlas based approaches rely on independent pairwise registrations between every atlas and the individual test image. Deformed segmentation labels are generated by warping every atlas image to the test image. The warped label maps arising from every deformed segmentation act as individual classifiers and the final segmentation of the test subject is obtained after label fusion. One widely used fusion technique is majority voting label fusion. Multi atlas based label fusion was first applied to segmentation of brain structures (Heckemann et al., 2006). A weighted voting approach has been demonstrated for segmentation of cardiac structures (Isgum et al., 2009), while a more sophisticated variational expectation maximisation approach that constructs a generative model (Iglesias et al., 2013) has been used to fuse segmentations from multiple atlases in individual structures. These established approaches have not been evaluated for segmentation of cerebellar structures. To date, there are no cerebellar parcellations available in Freesurfer (http://surfer.nmr.mgh.harvard.edu) which is a popular tool for automatic whole brain segmentation based on probabilistic information. We propose to design a framework using manual labels to introduce cerebellar parcellations in the Freesurfer software.

The cerebellum is located at the rear of the brain, underlying the occipital and temporal lobes of the cerebral cortex (Figure 4.1). Although the cerebellum accounts for approximately 10% of the brain's volume, it contains over 50% of the total number of neurons in the brain. The cerebellum consists of four major lobes, namely the anterior lobe (lobules I-IV and V), the superior posterior lobe (lobules VI and Crus I), the inferior posterior lobe (Crus II and lobules VIIb, VIIIa, VIIIb and IX), and the flocculonodular lobe (lobule X). These regions contribute to various functional domains. The lobular subdivision helps to better understand the structure and function of the cerebellum. Although the majority of the cerebellar outputs are to the motor system of the brain, it is also involved in other higher order brain functions like cognitive, sensory, affective and effective processing (Stoodley and Schmahmann, 2009; Schlerf et al., 2010).

Parcellation of the cerebellum into different sub divisions has proved to be useful to study quantitative morphological characteristics of cerebellar structures (volumetry) and to achieve improved anatomical localization and inter-subject coregistration in functional imaging studies (Makris et al., 2005; Diedrichsen et al., 2009). Automated segmentation of the cerebellum and its structures remains difficult due to its highly convoluted lobules, small volumes of some structures (Buckner et al., 2011), and comparatively different intensity distribution of cerebellar gray and white matter compared to the cerebrum (Datta et al., 2009; Bogovic et al., 2013).

The Schmahmann cerebellar atlas was the first to depict the gross morphology of the cerebellum with both fissures and individual cerebellar lobules. Later a probabilistic representation of the cerebellum comprising 28 parcellations from both hemispheres was constructed from 20 healthy subjects by affine registration to the ICBM152 template, followed by normalisation to the colin27 template (Diedrichsen 2006). The average deformation warp from the colin27 template to individual subjects resulted in the spatial unbiased infratentorial cerebellar template (SUIT). However, the segmented output using the SUIT atlas frequently suffers from inclusion of the transverse sinus and part of the bone marrow (Diedrichsen et al., 2009).

The primary aim of this study was to develop a method to achieve more accurate automated parcellation of cerebellar structures using a multi atlas based approach. More specifically, we wanted to (1) improve the segmentation accuracy of individual cerebellar hemispheric structures compared to a single atlas approach, (2) quantitatively evaluate our output

segmentation accuracy in test subjects using simultaneous truth and performance level estimates (STAPLE), and (3) perform leave one-out cross validation of our training subject segmentation to compare manual and automated segmentations. To obtain feasible functional localisation, the automatic segmentation algorithm should produce accuracy comparable with manual segmentation performed by an expert neuroanatomist. The major contribution of this work is the construction of a multi atlas segmentation pipeline for cerebellar structures and its evaluation using two different datasets. To the best of our knowledge this study is one of only a few studies to evaluate automated cerebellar segmentation using a multi atlas based approach.

4.2 Methods

Training and Test Data Sets

High resolution structural T1-weighted images from two different centers were used in this study. The training data set (N=20) was acquired on a 3T Philips Intera system using an MPRAGE sequence with field of view 256x256x150 mm³ and 1x1x1mm³ resolution. The age range of the subjects was 22 to 45 years. The cerebella of the training data were previously manually delineated into 28 cerebellar structures for the construction of the SUIT atlas (Diedrichsen et al., 2006; Diedrichsen et al., 2009). In this paper we refer to the 28 cerebellar segmentations as label maps. The 28 cerebellar structures and their corresponding colors for visualization are illustrated in Figure 4.1 and Table 4.1, respectively.



Figure 4.1 Three dimensional surface representations of cerebellar cortex in the posterior (left) and anterior (right) views. The middle image shows a two dimensional coronal slice with the manual tracings for one training subject.

Table 4.1 The 28 cerebellar structures and their color representations as used in the current study

Left I_IV	Left Crusl	Vermis VIIb		Right VIIIb
Right I_IV	Vermis Crusl	Right VIIb		Left IX
Left V	Right Crusl	Left VIIIa		Vermis IX
Right V	Left Crusll	Vermis VIIIa		Right IX
Left VI	Vermis Crusll	Right VIIIa	2.13	Left X
Vermis VI	Right CrusII	Left VIIIb		Vermis X
Right VI	Left VIIb	Vermis VIIIb		Right X

In this work, each training subject's label map constitutes one atlas image in our multi atlas. Of the 28 available structures, we used the 20 hemispheric structures, excluding the eight vermal regions, to validate our multi atlas segmentation accuracy. The vermal regions were excluded due to their small volumes.

Our test data set (to be segmented) comprised 35 high resolution T1 structural scans acquired using a magnetization prepared rapid gradient echo (MPRAGE) sequence on a 1.5T Vision system (Siemens, Erlangen, Germany) as part of ongoing studies at the Alzheimer's Disease Research Center in St. Louis and have appeared in prior publications (Buckner et al., 2004; Fotenos et al., 2005). The field of view is 256x256x224mm³ with 1x1x1.25mm³ spatial resolution. The age range of the subjects was 21.5 to 78.2 years.

Image pre-processing

All training and images processed using the Freesurfer test were (http://surfer.nmr.mgh.harvard.edu) software package. The processed surfaces of the test subjects were already available. All except one of the image volumes from the training data set were successfully processed using the Freesurfer recon pipeline and their surfaces constructed. The image volume of the one subject for whom the recon pipeline failed was excluded from further analyses. For six of the training subjects the Freesurfer brain extraction pipeline included portions of dura and exterior regions of the skull that were subsequently corrected by introducing the graph cut (GCUT) algorithm into the pipeline (Sadananthan et al., 2010). Very few voxels remained that required manual correction after skull stripping.

Registration

After computing the gray and white matter surfaces of the training subjects, each training subject was registered to every test subject using non-linear combined volume and surface based registration (Postelnicu-Zöllei et al., 2009). This tool achieves better alignment of both the cortical and subcortical folding patterns between the two datasets. This step was performed using a multi-CPU cluster running Linux. The resulting deformation warps were then used to segment the test images.

Multi Atlas labelling

We used a multi atlas based approach to obtain the most likely segmentation of each test subject from the classifier set which comprised 19 deformed outputs. The intensity of the test image and the respective deformed outputs were used to model the final segmentation of the test subject. Both multi atlas majority voting (MAMV) and multi atlas generative model based (MAGM) label fusion were used to obtain 28 cerebellar labels for each test subject (Iglesias et al., 2012). The parameterization of the model is based on iterative variational expectation maximization (Iglesias et al., 2013) and finally uses β to generate the most likely segmentation label. The parametrization value of β was adapted from a mathematical formulation that appeared in prior literature (Sabancu et al., 2010; Iglesias et al., 2012). In our work we used β =0 to obtain the fused output from MAMV and β =0.3 for MAGM. The model captures all possible prior knowledge about the spatial location, shape and extent of the individual atlas labels to model the posterior probability distribution of the output label on the test subject.

Dice Coefficient

The Dice coefficient (Dice et al., 1945) is a widely used voxel-wise evaluation measure that assesses the spatial overlap between two regions where 1 indicates perfect overlap and 0 indicates no overlap. In our study we only computed Dice coefficients for the 20 hemispheric structures of the cerebellum (excluding the vermal regions) as previous studies have shown lower reliability in finer parcellated structures (Makris et al., 2005). The Dice coefficient is given by

Dice coefficient (DC) =
$$\frac{2*(M \cap T)}{M+T}$$
 \forall Label=1, 2, ... 20

STAPLE evaluation

For all the test subjects, we compared the multi atlas based segmentations for the 20 hemispheric cerebellar structures achieved using MAMV and MAGM label fusion, respectively, with the probabilistic truth estimate derived from the Simultaneous Truth and Performance Level Estimation (STAPLE) algorithm (Warfield et al., 2004). The STAPLE algorithm effectively estimates the probabilistic true segmentation and also measures the performance of the algorithm given a set of input segmentations of an image. It uses expectation maximization to estimate the true segmentations. For each of the 20 hemispheric cerebellar regions we computed the Dice coefficient comparing the automated segmentation using MAMV and MAGM, respectively, with the STAPLE estimate in the 35 test subjects.

Leave-one-out cross validation (LOOCV)

To further validate our multi atlas approach, we used LOOCV on the training dataset for which the ground truth segmentations were available. The LOOCV uses 18 atlases from the training dataset to determine the segmentation of the remaining training image volume. This was done iteratively for all 19 training subjects to yield automated final fused segmentations of each training subject using multi atlas segmentation based on the other 18 training data sets. Dice coefficients were computed to compare manual and automated segmentations for each subject.

In order to assess the effect of varying the number of atlases used in the multi atlas segmentation, we also evaluated the accuracy of our output segmentation in our cross validation experiment when using a subset of only 14 atlases (instead of 18) from the training data set.

4.3 Results

STAPLE evaluation

In Table 4.2 we present for the 20 hemispheric cerebellar regions the mean Dice coefficients of spatial overlap between the STAPLE estimates and the automated segmentations using multi atlas majority voting and generative model based label fusion, respectively, in the 35 test subjects. For both MAMV and MAGM label fusion, the mean Dice coefficient was greater than 0.89 bilaterally in the anterior lobe (I-IV and V), superior posterior lobe (VI and Crus I) and parts of the inferior posterior lobe (Crus II and IX). The remaining lobules of the inferior posterior lobe (VIIb, VIIIa and VIIIb) and the flocculonodular lobe (X) exhibited lower Dice coefficients. These results are presented as box-and-whisker plots in Figures 4.2 and 4.4 for the 10 hemispheric left and right cerebellar structures, respectively. Dice coefficients achieved using the two different label fusion methods were compared using the Wilcoxon signed-rank test. Bilaterally in Crus I, CrusII, VIIIb and IX MAGM label fusion yielded higher Dice coefficients than MAMV. In contrast, the MAMV approach achieved better segmentation overlap with STAPLE bilaterally in I-IV and X, in left V and in right VI.

Table 4.2 Mean Dice coefficients in 35 test subjects of spatial overlap between the STAPLE estimate and automated segmentations in 20 hemispheric cerebellar regions using multi atlas majority voting (MAMV) and multi atlas generative model based (MAGM) label fusion, respectively.

Cerebellar	Mean Dice coefficient					
Structure	MA	MV	MAGM			
Structure	Left	Right	Left	Right		
I-IV	0.93	0.93	0.91	0.92		
V	0.90	0.89	0.89	0.89		
VI	0.92	0.92	0.92	0.91		
Crus I	0.92	0.93	0.93	0.94		
Crus II	0.89	0.91	0.93	0.95		
VIIb	0.78	0.81	0.79	0.84		
VIIIa	0.86	0.87	0.84	0.84		
VIIIb	0.86	0.84	0.87	0.88		
IX	0.89	0.89	0.91	0.92		
Х	0.77	0.64	0.73	0.54		



Figure 4.2 Box-and-whisker plots of the Dice coefficients in 35 test subjects between the STAPLE estimate and segmentations achieved using multi atlas majority voting (MAMV; red) and generative model (MAGM; blue) label fusion for left hemispheric cerebellar structures. Asterisks indicate significant differences between MAMV and MAGM and the color of the asterisks indicate the method that yielded the higher Dice coefficient. *p<0.05; **p<0.01; ***p<0.001

In order to give more insight into the distribution of different Dice coefficients across the test subjects in each of the cerebellar regions, we present in Figures 4.3 and 4.5 clustered plots of the Dice coefficients for the left and right hemispheric regions, respectively. Each color block gives the range of Dice coefficients in a subset of the test subjects. Each subset comprises 20% (i.e. 7) of the test subjects. This allows one to assess the number of test subjects in whom Dice coefficients were either above or below a certain threshold in each of the regions. It is evident from these plots that although mean Dice coefficients were low in lobules VIIIa and VIIIb, Dice coefficients were below 0.8 only in about 20% or fewer of test subjects, compared to lobules VIIb and X where low Dice coefficients were evident in 40% or more of test subjects.



Figure 4.3 Clustered plot representation of the Dice coefficients of spatial overlap with the STAPLE estimate for automated segmentations based on majority voting (1) and generative model based (2) label fusion for the left hemispheric cerebellar structures.



Figure 4.4 Box-and-whisker plots of the Dice coefficients in 35 test subjects between the STAPLE estimate and segmentations achieved using multi atlas majority voting (MAMV; red) and generative model (MAGM; blue) label fusion for right hemispheric cerebellar structures. Asterisks indicate significant differences between MAMV and MAGM and the color of the asterisks indicate the method that yielded the higher Dice coefficient. *p<0.05; **p<0.01; ***p<0.001



Figure 4.5 Clustered plot representation of the Dice coefficients of spatial overlap with the STAPLE estimate for automated segmentations based on majority voting (1) and generative model based (2) label fusion for the right hemispheric cerebellar structures.

Leave-one-out cross validation (LOOCV) - 18 fused segmentations

Table 4.3 presents the mean Dice coefficients for the 19 training subjects of the spatial overlap between manual tracing and automated segmentation achieved with label fusion of the other 18 training subjects using MAMV and MAGM, respectively, in the 20 hemispheric cerebellar regions. Although the same regions as before (lobules I-IV, V, VI, Crus I, Crus II, and IX) achieved higher spatial overlap bilaterally, Dice coefficients of spatial overlap with the ground truth manual tracings were significantly lower than those obtained in the test subjects using the STAPLE validation (Dice coefficients > 0.67 here compared to Dice coefficients > 0.89 in the test subjects using STAPLE). In contrast to the results in our test subjects, where MAGM typically performed better than MAMV, the Wilcoxon signed-rank test revealed higher Dice coefficients in our training subjects using MAMV bilaterally in lobules V, VIIIb and X, as well as left VIIb. Only bilaterally in Crus 1 did MAGM yield higher Dice coefficients than MAMV. These results are illustrated in box-and whisker plots for left and right hemispheric cerebellar regions in Figures 4.6 and 4.7, respectively.

Table 4.3 Mean Dice coefficients in 20 hemispheric cerebellar regions of spatial overlap between manual tracings and automated segmentations using MAMV and MAGM label fusion, respectively, in 19 training subjects.

Cerebellar	Average of Dice coefficient					
Hemisphere	MA	MV	MAGM			
Structures	Left	Right	Left	Right		
I-IV	0.77	0.78	0.76	0.78		
V	0.70	0.69	0.68	0.67		
VI	0.80	0.78	0.79	0.78		
Crus I	0.77	0.78	0.79	0.81		
Crus II	0.73	0.73	0.72	0.73		
VIIb	0.48	0.48	0.37	0.45		
VIIIa	0.56	0.57	0.58	0.52		
VIIIb	0.59	0.61	0.44	0.45		
IX	0.71	0.73	0.69	0.69		
Х	0.62	0.69	0.48	0.58		



Figure 4.6 Box-and-whisker plots of the Dice coefficients in the 19 training subjects of spatial overlap between manual tracings and automated segmentations using multi atlas majority voting (MAMV; red) and generative model based (MAGM; blue) label fusion, respectively, for left hemispheric cerebellar structures. Asterisks indicate significant differences (Wilcoxon signed-rank test) between MAMV and MAGM and the color of the asterisks indicates the method that yielded the higher Dice coefficient. *p<0.01; ***p<0.001.



Figure 4.7 Box-and-whisker plots of the Dice coefficients in the 19 training subjects of spatial overlap between manual tracings and automated segmentations using multi atlas majority voting (MAMV; red) and generative model based (MAGM; blue) label fusion, respectively, for right hemispheric cerebellar structures. Asterisks indicate significant differences (Wilcoxon signed-rank test) between MAMV and MAGM and the color of the asterisks indicates the method that yielded the higher Dice coefficient. *p<0.05; **p<0.01; ***p<0.001.

Figures 4.8 and 4.9 show the distribution of the Dice coefficients for left and right hemispheric cerebellar regions, respectively, in subsets of the training subjects where each subset comprises 20% of the subjects.



Figure 4.8 Clustered plot representation of the distribution of the Dice coefficients of spatial overlap with manual tracings for automated segmentations based on majority voting (1) and generative model based (2) label fusion, respectively, for the left hemispheric cerebellar structures.



Figure 4.9 Clustered plot representation of the distribution of the Dice coefficients of spatial overlap with manual tracings for automated segmentations based on majority voting (1) and generative model based (2) label fusion, respectively, for the right hemispheric cerebellar structures.

Leave-one-out cross validation (LOOCV) - 14 fused segmentations

To investigate the impact of altering the number of fused segmentations used to generate the output segmentation, we compared in the 19 training subjects the Dice coefficient of spatial overlap with manual tracings obtained in each hemispheric cerebellar region for automated segmentation based on 18 atlases with that obtained using 14 atlases. The 14 atlases were selected from the training dataset based on how clearly border pixels between cerebellar lobules could be identified visually. Figure 4.10 presents a comparison of the Dice coefficients in the 20 hemispheric cerebellar regions based on 14 (red) and 18 (blue) fused segmentations for MAMV (top row) and MAGM (bottom row), respectively.

Using 14 or 18 fused segmentations in the 19 training subjects yielded similar values for the Dice coefficient in most regions using both label fusion methods. Only in VIIb and VIIIb were Dice coefficients somewhat higher using 14 atlases than with 18 atlases for MAGM label fusion – this difference was only significant in right VIIIb. Also, the pattern of the data

is consistent with that seen previously with lower Dice coefficients in posterior lobules VIIb, VIIIa, VIIIb and X.



Figure 4.10 Comparison of Dice coefficients of spatial overlap in 19 training subjects of automated segmentations generated using either 14 (red) or 18 (blue) fused label maps with manual tracing in left (left) and right (right) hemispheric cerebellar regions. The top row shows results for MAMV and the bottom row for MAGM. Dagger and asterisks indicate significant differences (Wilcoxon signed-rank test) between using 14 and 18 atlases and the color indicates the method that yielded the higher Dice coefficient. $\dagger p < 0.1$; *p < 0.05.

4.4 Discussion

In this work, we compared automatic segmentation using MAMV and MAGM label fusion and assessed segmentation accuracy using two different approaches in both our training and test data sets, which were acquired at two different sites on different scanner platforms. Although both MAMV and MAGM in both datasets yielded good segmentation accuracy in all regions except lobules VIIb, VIIIa, VIIIb and X, Dice coefficients of spatial overlap with manual tracings in the training data were typically lower than those achieved in the test data with STAPLE. Except for bilateral lobules I-IV and X, left V and right VI, MAGM achieved similar or better spatial overlap with the STAPLE estimate in the test subjects compared to MAMV. In contrast, MAGM performed better than MAMV only bilaterally in Crus 1 in the training data set. The fact that the volume of lobule X is smaller than the other cerebellar structures, penalizes the Dice coefficient and may account for the low spatial overlap achieved in this region.

The cerebellum is a complex structure with fine parcellations. To date, only a few studies have used SUIT or manual delineation to perform cerebellar segmentation (Bogovic et al., 2013; Makris et al., 2005; Pierson et al., 2002). A previous study using manual delineation emphasized the trade-off between achieving fine parcellations and reliability and finally used only 24 parcellation labels excluding the vermal structures of lobule VI, Crus I, Crus II and VIIb. In the present work, we similarly only evaluated our segmentation accuracy in the 20 hemispheric regions, excluding the smaller vermal regions in which reduced accuracy is expected.

It has previously been reported (Diedrichsen et al., 2009) that the subjects used in the SUIT atlas construction exhibited anatomical variability in Crus II, VIIb and IX. Notably, VIIb is one of the structures that consistently yielded low Dice coefficients in both the test and the training data sets in our study. Using both MAMV and MAGM, the structures located posteriorly in the cerebellum (VIIb, VIIIa, VIIIb, IX and X) generally achieved lower segmentation overlap compared to more anterior regions. Amongst these posterior regions, VIIb, VIIIa and VIIIb achieved the lowest Dice coefficients. Increased variability in posterior
lobules VIIb and VIIIb amongst different raters has also been reported by Bogovic et al. (2013).

Poor contrast often makes it difficult to locate borders during manual delineation of cerebellar structures, so that tracings are based on relative spatial position and contours on adjacent slices rather than boundaries per se. We suspect that cerebellar structures VIIb, VIIIa and VIIIb in the training data may have suffered from these problems in the delineation phase resulting in poor segmentation accuracy in these regions. This may explain why MAMV generally performed better in the training data than MAGM, as MAMV exploits knowledge of relative position rather than voxel intensity distributions. Other studies (Artaechevarria et al., 2009) using multi atlas parcellation have also reported better performances for different methods (for example, majority weighting and local weighting) in different data sets. We also observed intensity variations among the training data, which may have contributed to the boundary estimation problem in the posterior cerebellar lobules.

Using fewer atlases appears to have increased the Dice coefficient in the posterior lobules VIIb and VIIIb using MAGM. It may be that excluding from the training data set atlases in which border pixels between different anatomical regions were not clearly visible due to poor contrast, reduced the variability in these regions resulting in better segmentations and increased Dice coefficients.

To date, very little work has been done on automatic cerebellar parcellation. Using available delineated images we developed an optimised pipeline to achieve reliable cerebellar segmentation. Although both the training and test data exhibited better and worse

performance in the same cerebellar regions, Dice coefficients were higher in all regions in the test data than the training data. This may be due to greater similarities between multi atlas based segmentation and the STAPLE estimates, compared to ground truth manual tracings that may vary significantly between subjects. Further, segmentation accuracy was reduced in the posterior cerebellar lobules, presumably due to greater inter-subject variability in these regions and poor contrast resulting in uncertainty in identifying boundaries during manual delineation.

4.5 Conclusion

In this paper we present a reliable method to achieve automated cerebellar segmentation using multiple atlases. The approach was implemented and evaluated using STAPLE in our test dataset and compared to manual tracings using leave-one-out cross validation in our training data. Although performance of MAMV and MAGM label fusion were generally similar, MAGM performed better in most regions in the test data and MAMV mostly performed better in the training data. Automated segmentation accuracy was generally poor in lobules VIIb, VIIIa and VIIIb and further work is needed to improve segmentation accuracy in these areas.

Chapter 5

Discussion

This thesis investigates different techniques and approaches to the development of a robust tool for segmentation of cerebellar structures on magnetic resonance (MR) images and different validation strategies, with the aim to develop a robust pipeline for use in large population imaging studies. The idea of constructing a pediatric cerebellar atlas and combining it with different label propagation methods to achieve cerebellar segmentation was one of the first of its kind and has rarely been addressed in the literature. In the previous chapters we presented the results obtained using these different methods for the segmentation of cerebellar structures in single or sequential images. The methods used were evaluated on various datasets collected from different imaging centres well-known for their work in MRI.

Anatomical delineation of cerebellar structures

Although manual delineation is extremely labour intensive, it is still considered the gold standard for volumetric assessment. Observer bias may, however, affect the results whenever a manual method is used. Prior studies of the cerebellum have emphasized the challenges associated with manual delineation of CrusII, lobules VIIb and VIII (Diedrichsen et al., 2009; Bogovic et al., 2013) and that raters should consider different image features like size, depth of fissures, location of the fissure, starting and ending points of the fissure, and lobule boundaries when deciding where to draw boundaries. Our expert rater completed manual tracing of individual lobules of a complete cerebellum in roughly 30-60 minutes. Intra-class correlation coefficients (ICCs) of individual lobules traced at two different times by the same tracer were computed to assess the variability of manual tracings.

Probabilistic maps of cerebellar structures

Probability maps show the extent, shape and arrangement of regional cerebellar structures. The map also gives an indication of the certainty with which a given voxel can be assigned to a specific structure, based on the frequency with which that voxel was assigned to said structure in a set of images. The spatial distribution of our probability map reveals that individual structures superimpose accurately in medial regions, but that uncertainty increases as one moves laterally in the structures. CAPCA18 yielded 16 maximum likelihood labels derived from the maximum probability atlas.

Quantitative evaluation of the CAPCA18 atlas

Spatial overlap between automatically segmented cerebellar structures of a pediatric data set using the CAPCA18 atlas and manual tracings of the same data set yielded values above 70% bilaterally in lobules I-V, VI, crusI, crusII, VIII and IX, which compares favourably with previous results (Bogovic et al., 2013). Spatial overlap of regions derived from CAPCA18 segmentation and manual tracing were consistently higher than spatial overlap of SUIT segmentation and manual tracing, with significant improvement in lobules I-V, VI, CrusI, VIII and IX cerebellar structures. It has been noted previously that direct comparison of quantitative segmentation results across publications are difficult and not always fair due to inconsistencies in the manual segmentation protocol, the quality of the imaging data, and differences in the patient populations (Collins et al., 2010; Wang et al., 2012). Despite these limitations, the present work demonstrates that cerebellar segmentation using the CAPCA18 atlas est than segmentation using the SUIT atlas.

Since there is no previous literature on cerebellar segmentation in children, our results help to establish baseline values for future studies. Furthermore, it helps to highlight the significant complexities associated with automatic segmentation of certain regions of the cerebellum. In prior studies of brain structure segmentation, Shattuck et al., (2008) suggest that variability in manual tracing data can also be used as a guide in determining the spatial overlap that would be considered acceptable from an automated segmentation. To this end, we have reported in all structures both Dice coefficients and ICCs obtained from inter- and intra-rater reliability tests. As expected, Dice coefficients and inter-rater ICCs were low and high in the same regions.

When comparing the average percentage of gray matter volume occupied by each lobule with previously reported values in adults (Makris et al., 2005), we found that lobule IX occupied a relatively greater volume in children than adults, while lobules VI, VIIb and X were relatively smaller in children. Fonov et al. (2011) compared deformations when normalising to an adult template compared to a pediatric template and reported greater deformation when using an adult template for pediatric data as the shape and ratio of gray matter and white matter near the posterior lobe of the cerebellum differ between children and adults. We also observed in our data that segmentation using CAPCA18 yielded significantly better spatial overlap in lobule IX compared to SUIT, and that the volume of lobule IX obtained from SUIT segmentation (SUIT was developed using adult data) was significantly smaller than that obtained from manual tracing. It is possible in the future to address more directly questions relating to relative differences across cerebellar regions in different populations using Jacobian maps of deformation.

Comparing different label propagation methods to achieve cerebellar segmentation of an adult dataset

The quality of the manual labels forces an upper bound on the quantitative metrics — an automatic procedure can't agree with a manual rater or the segmentation from the same rater at two instants cannot agree leaving subtle discrepancies and limiting the accuracy achievable by any automated method (Shattuck et., 2008; Collins et al., 2010; Wang et al., 2012). Keeping this in mind, we compared segmentation accuracy achieved using our CAPCA18 atlas with three different label propagation strategies to manual segmentation in 8 structures per hemisphere in 3 adult subjects in terms of spatial overlap and modified Hausdorff distances (Figure 3.3). The distance metric indicated that multi atlas based generative model (MAGM) label fusion consistently performed better than the other two methods in all regions except lobule VIIb. Previous studies (Ghosh et al., 2010) in younger children have demonstrated that surface based registration introduce significantly less bias and that cortical alignment is better in surface based methods compared to volume based registration. Overall, combining deformable registration using cortical surfaces and multi atlas based label fusion improved accuracy compared to direct warping in our study.

In chapter 4, we used the cerebellar parcellations that were available for 20 adult subjects that had been used in the construction of the SUIT atlas (Diedrichsen et al., 2009) combined with two different multi atlas based segmentation methods (MAMV and MAGM) to automatically segment the cerebella of 35 test subjects, in order to evaluate the accuracy achievable when using adult training data to achieve segmentation of adult test subjects (compared to the pediatric training data used in the previous chapter). We compared spatial overlap between automated segmentations in the test data and their STAPLE estimates, and performed leave one out cross validation to segment each training subject using the remaining training atlases for comparison with available manual tracings. Lobules I-IV, V, VI, crus I, crus II and IX

achieved spatial overlaps greater than 0.7. Differences in signal intensity in the training images negatively impacted accuracy in the posterior regions of the cerebellum for both MAMV and MAGM resulting in low spatial overlap in these regions.

Furthermore, we examined the impact on segmentation accuracy of varying the number of training atlases (14 versus 18) used to generate the final fused image. In contrast to previous studies (Heckemann et al., 2006; Lötjönen et al., 2009), we found that Dice coefficients of fused outputs from 14 and 18 atlases were similar in most structures using both MAMV and MAGM. Unexpectedly, Dice coefficients were somewhat higher using 14 atlases than 18 atlases in VIIb and VIIIb for MAGM label fusion – a difference that was only significant in right VIIIb. This bias might be due to the intensity difference mentioned previously or intersubject variability in this region and requires further investigation.

Comparison of CAPCA18 and SUIT Atlases

The SUIT atlas (Diedrichsen et al., 2009) developers mention the presence of high intersubject variability in the posterior lobes of the cerebellum, which one would expect to translate into lower segmentation accuracy in these regions. This may explain why in chapter 4, which uses the manual tracings that were used in the construction of the SUIT atlas to segment 35 adult test subjects, spatial overlap in lobules VIIb, VIIIa, VIIIb and X are consistently lower than in other regions. In comparison, spatial overlap was low compared to other regions only in lobule VIIb when using the CAPCA18 atlas to segment the same 35 adult test subjects.

Computational complexity and performance time

Regardless of which multi atlas fusion method was used, the time to perform the combined volume and surface based registration between the atlases and the test image, was approximately 24 hours without parallelization. This time can be reduced by using parallelization. The time to generate the fused images was negligible compared to that required for the multi atlas registrations. It is worth emphasizing, however, that the additional computational investment significantly improved the resultant segmentation accuracy compared to direct warping.

Limitations

Labelling accuracy depends on the quality of the images, the pre-processing steps the images were subjected to, the registration algorithms used, the label propagation pipeline (atlas/multi atlas), and the methods used to evaluate performance. With regards the first three, we assumed that each data set comprised a group of normal individuals whose brain images were acquired, pre-processed, and labelled in a consistent manner.

We are not aware of any comparisons in literature between the registration of interpolated versus non-interpolated (bias-field corrected and uncorrected, intensity normalized and non-normalized, etc.) images. Images from the subjects used in the construction of the CAPCA18 atlas were linearly interpolated using BrainVoyager before pre-processing in SPM. All of the images in this study were linearly interpolated twice, once to linearly register each brain to a template, and a second time to linearly register each source brain to a target brain in the template space, prior to nonlinear registration.

Regarding the registration methods themselves, each one has a similarity measure, transformation model, regularization method, and optimization strategy. A superior transformation model coupled with an unsuitable similarity measure, for example, would most likely lead to suboptimal results. We ensured that the output of individual registrations was precise to carry out the label propagation. In the context of combined volume and surface based registration, we followed the optimal parameter settings as per the advice from the author of the tool.

Finally for the reslicing of the subjects label volumes, we used nearest neighbour interpolation to preserve the label values and ensure more consistent behaviour at the edges of individual structure.

Future work

We used images from healthy Cape Coloured children who were typically developing controls in a different study to construct the CAPCA18 atlas. The accuracy of automated segmentation using CAPCA18 was evaluated in a different group of healthy children from the same ethnic group. It would be interesting to assess the performance of the CAPCA18 atlas both in children from different ethnic groups and in children in whom there may be some underlying pathology, such as children prenatally exposed to alcohol. Longitudinal studies of this nature would allow one to detect pathology-related changes over time rather than differences that may exist between the individual's brain and the template. CAPCA18 cerebellar parcellations can also be useful in accurately labelling cerebellar regions in functional MRI studies.

The demand for tools for robust structural analysis of MR images continues to grow. Given the interest and prevalence of data from different populations from different laboratories, large-scale multi-site experiments are becoming more routine and are producing more data than before. Multi-centre studies and meta-analyses of cerebellar structures using our CAPCA18 atlas has the potential to improve our understanding of the structure and function of the cerebellum and how it may be altered/affected in different conditions.

Chapter 6

Conclusions

This thesis presents the first tool for automated segmentation of the cerebellum on 3D MR images of children. The CAPCA18 atlas has been shown to provide precise and consistent output segmentations. In this work, we used improved registration algorithms and compared different multi atlas label fusion methods to obtain the final segmentations. The data generated reveals differences between adult and child cerebella.

The evaluation of the proposed atlas demonstrates that it can be used with success in both pediatric and adult data sets. Segmentation accuracies are shown to be comparable or better than values previously reported using SUIT.

Factors that may impact the cerebellar segmentation accuracy have been described and limitations highlighted. To date, the cerebellum is rarely studied. Construction of a user-friendly tool using the CAPCA18 atlas combined with multi atlas label fusion techniques will help facilitate research on cerebellar structural abnormalities that accompany various conditions and can greatly advance our understanding of the role of this region, which is still not very well understood.

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