# Ultra-High Field Magnetic Resonance Imaging for Stereotactic Neurosurgery 

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#### Abstract

Stereotactic neurosurgery is a subspecialty within neurosurgery concerned with accurate targeting of brain structures. Deep brain stimulation (DBS) is a specific type of stereotaxy in which electrodes are implanted in deep brain structures. It has proven therapeutic efficacy in Parkinson's disease and Essential Tremor, but with an expanding number of indications under evaluation including Alzheimer's disease, depression, epilepsy, and obesity, many more Canadians with chronic health conditions may benefit. Accurate surgical targeting is crucial with millimeter deviations resulting in unwanted side effects including muscle contractions, or worse, vessel injury. Lack of adequate visualization of surgical targets with conventional lower field strengths (1.5/3 Tesla) has meant that standard-of-care surgical treatment has relied on indirect targeting using standardized landmarks to find a correspondence with a histological "template" of the brain. For this reason, these procedures routinely require awake testing and microelectrode recording, which increases operating room time, patient discomfort, and risk of complications. Advances in ultra-high field ( $\geq 7$ Tesla or 7 T ) imaging have important potential implications for targeting structures enabling better visualization as a result of its increased (sub-millimeter) spatial resolution, tissue contrast, and signal-to-noise ratio. The work in this thesis explores ways in which ultra-high field magnetic resonance imaging can be integrated into the practice of stereotactic neurosurgery. In Chapter 2, an ultra-high field MRI template is integrated into the surgical workflow to assist with planning for deep brain stimulation surgery cases. Chapter 3 describes a novel anatomical fiducial placement protocol that is developed, validated, and used prospectively to quantify the limits of template-assisted surgical planning. In Chapter 4, geometric distortions at 7T that may impede the ability to perform accurate surgical targeting are characterized in participant data, and generally noted to be away from areas of interest for stereotactic targeting. Finally, Chapter 5 discusses a number of important stereotactic targets that are directly visualized and described for the first time in vivo, paving the way for patient-specific surgical planning using ultra-high field MRI.


Keywords: accuracy, deep brain stimulation, magnetic resonance imaging, neuromodulation, stereotactic neurosurgery, surgical planning.

Tools used by the surgeon must be adapted to the task and where the human brain is concerned, no tool can be too refined. - LARS LEKSELL


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## Contents

Abstract ..... ii
Acknowledgments ..... v
Co-Authorship Statement ..... vi
List of Figures ..... xi
List of Tables ..... xvii
List of Abbreviations, Symbols, and Nomenclature ..... xviii
1 Introduction ..... 1
1.1 Stereotactic Neurosurgery ..... 2
1.1.1 Early Indications and Treatment Modalities ..... 4
1.1.2 Brain Atlases ..... 6
1.1.3 The Importance of Imaging ..... 8
1.1.4 Deep Brain Stimulation ..... 8
1.1.5 Conventional Approaches for Stereotactic Targeting ..... 10
1.2 Magnetic Resonance Imaging ..... 12
1.2.1 Fundamentals of Magnetic Resonance Imaging ..... 13
1.2.2 Ultra-High Field MRI ..... 15
1.2.3 Quantitative MRI ..... 18
1.3 Thesis Outline ..... 20
2 Ultra-high field template-assisted deep brain stimulation surgery ..... 22
2.1 Introduction ..... 23
2.2 Materials and Methods ..... 23
2.2.1 Deep Brain Stimulation Surgery Workflow ..... 23
2.2.2 7T Group Template Creation ..... 25
2.2.3 Template-to-Patient Registration Workflow ..... 25
2.3 Results: Two Cases ..... 27
2.3.1 Right GPi Implantation for Dystonia ..... 27
2.3.2 Bilateral STN Implantation for Parkinson's Disease ..... 28
2.4 Discussion ..... 30
2.5 Conclusions ..... 32
3 A framework for evaluating correspondence between brain images using anatom- ical fiducials ..... 33
3.1 Introduction ..... 34
3.2 Methods ..... 35
3.2.1 Protocol development ..... 35
3.2.2 Phase 1: Protocol validation for brain templates ..... 37
3.2.3 Phase 2: Protocol validation for individual subjects ..... 37
Region-of-interest segmentation ..... 39
3.2.4 Phase 3: Evaluating subject-to-template registration ..... 39
3.2.5 Phase 4: Evaluating template-to-template registration ..... 40
3.2.6 Source code and data availability ..... 41
3.3 Results ..... 41
3.3.1 Phase 1: Protocol validation for brain templates ..... 41
3.3.2 Phase 2: Protocol validation for individual subjects ..... 43
3.3.3 Phase 3: Evaluating subject-to-template registration ..... 43
3.3.4 Phase 4: Evaluating template-to-template registration ..... 47
3.4 Discussion ..... 49
3.4.1 Protocol development and validation ..... 52
3.4.2 Point-based versus ROI-based metrics ..... 53
3.4.3 Subject-to-template registration ..... 53
3.4.4 Template-to-template registration ..... 54
3.4.5 Teaching neuroanatomy ..... 55
3.4.6 Limitations and future work ..... 55
3.5 Conclusions ..... 56
4 Quantification of local geometric distortion in structural magnetic resonance images: Application to ultra-high fields ..... 57
4.1 Introduction ..... 58
4.2 Materials and Methods ..... 60
4.2.1 Participants and MRI Acquisition Protocol ..... 60
4.2.2 Data Processing ..... 60
4.2.3 Voxel-Level Metrics ..... 61
4.2.4 Region-of-Interest Analysis ..... 62
4.2.5 Validation ..... 62
4.2.6 Effect of Gradient Distortion Correction ..... 63
4.3 Results ..... 64
4.3.1 General Findings ..... 64
4.3.2 Identification and Characterization of Local Distortion ..... 65
4.3.3 Distortion Increases with Isocenter Distance ..... 66
4.3.4 Region-of-Interest Analysis ..... 66
4.3.5 Validation ..... 71
4.3.6 Effect of Gradient Distortion Correction ..... 71
4.4 Discussion ..... 73
4.5 Conclusions ..... 77
5 Direct visualization and characterization of the human zona incerta and sur- rounding regions ..... 79
5.1 Introduction ..... 80
5.2 Materials and Methods ..... 82
5.2.1 Participant and image acquisition details ..... 82
5.2.2 Image pre-processing and template creation ..... 82
5.2.3 Pre-processing: MP2RAGE ..... 83
5.2.4 Pre-processing: T2SPACE ..... 83
5.2.5 Template creation ..... 84
5.2.6 Region-of-interest segmentation ..... 84
5.2.7 Stereotactic target localization ..... 85
5.3 Results ..... 85
5.3.1 Template Creation ..... 85
5.3.2 Direct visualization and segmentation of the zona incerta region ..... 86
5.3.3 Tissue properties of the zona incerta region ..... 90
5.3.4 Deep brain stimulation of the caudal zona incerta ..... 90
5.4 Discussion ..... 90
5.5 Conclusions ..... 96
6 Conclusions and Future Directions ..... 97
6.1 The Limits of Image-Based Targeting ..... 98
6.2 Multiparametric Imaging for Stereotactic Neurosurgery ..... 99
6.3 "Asleep" Deep Brain Stimulation Surgery ..... 100
6.4 Innovations in Stereotaxy ..... 101
6.5 Conclusions ..... 101
Appendices ..... 103
Appendix A Chapter 3 Supplementary Material ..... 104
A. 1 Phase 1: Supplementary Material ..... 104
A. 2 Phase 2: Supplementary Material ..... 116
A. 3 Phase 3: Supplementary Material ..... 120
A. 4 Phase 4: Supplementary Material ..... 130
Appendix B Chapter 4 Supplementary Material ..... 134
B. 1 Supplementary Tables ..... 134
Appendix C Ethics Approvals ..... 139
C. 1 Retrospective images for deep brain stimulation surgery. ..... 140
C. 2 7T images for geometric distortion study. ..... 141
C. 3 Prospective 7T stereotaxy study. ..... 142
Appendix D Copyright Transfers and Reprint Permissions ..... 143
D. 1 Copyright for [Rooney et al., 2007] ..... 144
Bibliography ..... 150
Curriculum Vitae ..... 174

## List of Figures

1.1 The pioneering work of Victor Horsley and Robert Clarke who coined the term "stereotaxy" and proposed a system involving the combination of (a) a mechanical frame with (b) a histological reference space using frozen sections for (c) electrolytic study of the deep cerebellar nuclei [Horsley and Clarke, 1908].
1.2 Stereotactic equipment in humans. (a) The Model V instrument of Spiegel and Wycis introduced in 1947 requiring plaster casting as a means to fix the frame to the head. (b) The first "centre-of-arc" system invented by Lars Leksell in 1949
1.3 Spiral drawing under normal conditions and for a patient with essential tremor.
1.4 Schematic of the cerebellothalamocortical and pallidothalamocortical circuits based on tract-tracing studies in primates [Gallay et al., 2008]. All major cerebellar, basal ganglia, and cortical regions are represented. $\mathrm{M} 1=$ primary motor cortex; $\mathrm{PMc}=$ caudal premotor $; \mathrm{SMA}=$ supplementary motor area; $\mathrm{PMr}=$ rostral premotor; PFC = prefrontal cortex; VLp = ventrolateral posterior thalamus; $\mathrm{VM}=$ ventral medial nucleus; $\mathrm{VLa}=$ ventrolateral anterior; $\mathrm{VApc}=$ ventral anterior parvocellular division; fct = fasciculus cerebellothalamicus; ft = fasciculus thalamicus; $\mathrm{fl}=$ fasciculus lenticularis; $\mathrm{al}=$ ansa lenticularis; STh $=$ subthalamic nucleus; $\mathrm{SNc}=$ substantia nigra pars compacta; $\mathrm{GPi}=$ globus pallidus internus; $\mathrm{GPe}=$ globus pallidus externus11
1.5 Demonstration of the intersubject variability in the location of anatomical structures of the deep brain and specifically the subthalamic region. The upper subfigures represent sagittal sections of the brain from two different cadavers ( Hb 1 is shown with red outlines and filled shapes while Hb 2 is shown with black outlines and light gray filling). The lower subfigure demonstrates variability in the locations for four specimens ( $\mathrm{Hb} 1, \mathrm{Hb} 2, \mathrm{Hb} 3, \mathrm{Hb5}$ ). $\mathrm{PTT}=$ pallidothalamic tractotomy. CTT $=$ cerebellothalamic tractotomy. Figure taken from the open access article by [Gallay et al., 2008].
1.6 (a) Protons spinning in free space, (b) protons spinning under the influence of $B_{0}$, (c) A $B_{1}$ radiofrequency pulse tips the magnetization vector, $\vec{M}$ into the transverse plane and over time relaxes back to equilibrium in alignment with $B_{0}$ (rotating frame). The arrows in (a) and (b) refer to the magnetic moment of the spinning nuclei.
1.7 $\quad T_{1}$ (a) and $T_{2}$ (b) relaxation can be modeled using exponential functions as demonstrated in Equations 1.4 and 1.5. The $T_{1}$ and $T_{2}$ relaxation times were set at 1500 ms and 50 ms respectively in these simulated examples (dashed lines). (c) Demonstration of how for a given echo time (marked by the dashed line) for recording, a mix of both $T_{1}$ and $T_{2}$-based signal is recorded, as is typical with conventional imaging methods.
1.8 One way to increase sensitivity to magnetization (i.e. signal) is to increase $B_{0}$, which scales at least linearly with magnetic field strength. Selected T2weighted coronal images taken at standard clinical field strength (1.5T) and at 7T demonstrating the improvement in image contrast in deep brain structures with an increase in $B_{0}$. The zoomed in regions demonstrate the increased detail in structures of the basal ganglia, specifically the pallidum, subthalamic nucleus, and substantia nigra.
1.9 Comparison of conventional $T_{1}$-weighted imaging where the units are arbitrary with quantitative $T_{1}$ mapping where the values reflect inherent local tissue properties. Individual subject scan at 7-Tesla using the MP2RAGE sequence [Marques et al., 2010].
1.10 Figures from Rooney et al. demonstrating the field-dependent increase in $T_{1}$ values and dispersion in different tissue types [Rooney et al., 2007]. The increase in dispersion of T1 manifests as improved contrast that can be exploited for better delineating boundaries between brain structures. This will be the subject of Chapter 5.
2.1 Side-by-side visualization of standard clinical 1.5T T1 magnetic resonance imaging (MRI) next to the fused ultra-high field MRI T1 and T2 templates in the axial plane. The red dot marks the location of the globus pallidus internus.26
2.2 Right internal pallidum implantation assisted by ultra-high field (UHF) fusion to standard clinical images (1.5T magnetic resonance imaging [MRI] and computed tomography $[\mathrm{CT}])$. The red dot marks the same location fused between modalities for (A) 1.5T MRI with gadolinium, (B) with the SchaltenbrandWahren atlas overlay provided in the clinical neuronavigation software, (C) CT in Leksell frame, (D) UHF T1 average, and (E) UHF T2 average. Also included is the best corresponding coronal section in the post-insertion MRI (F).
2.3 Example screenshot demonstrating the integration of the 7T T2 average into the commercial surgical planning software (Case 1: right internal pallidum insertion). The red dot marks the location of the globus pallidus internus.29
2.4 Bilateral subthalamic nucleus implantation assisted by ultra-high field (UHF) fusion to standard clinical images. The left and right trajectories (yellow and green, respectively) are fused across modalities for (A) 1.5 T magnetic resonance imaging (MRI) with gadolinium, (B) with the Schaltenbrand-Wahren atlas overlay, (C) UHF T1 average, and (D) UHF T2 average. Bilateral subthalamic nuclei were implanted successfully, as demonstrated with select (E) coronal and (F) axial views on postoperative MRI.
3.1 Metrics for evaluating spatial correspondence between brain images include voxel overlap (i.e. ROI-based) metrics as well as point-based distance metrics. The proposed framework involves the identification of point-based anatomical fiducials (AFIDs) in a series of brain images, which provide an intuitive millimetric estimate of correspondence error between images and is also a useful tool for teaching neuroanatomy.
3.2 Each anatomical fiducial in the full AFID32 protocol is demonstrated with crosshairs at the representative location in MNI2009bAsym space using the standard cardinal planes. AC = anterior commissure; $\mathrm{PC}=$ posterior commissure; $\mathrm{AL}=$ anterolateral; $\mathrm{AM}=$ anteromedial; $\mathrm{IG}=$ indusium griseum; IPF = interpeduncular fossa; $\mathrm{LMS}=$ lateral mesencephalic sulcus; $\mathrm{LV}=$ lateral ventricle; $\mathrm{PMJ}=$ pontomesenphalic junction.
3.3 Metrics used for validating AFID placements are shown here in schematic form. Mean, intra-rater, and inter-rater AFLE can be computed for an image that has been rated by multiple raters multiple times.
3.4 K-means clustering of point clouds relative to the mean fiducial location for each of the 32 AFIDs (left). Principle components analysis (bottom right) revealed three different general patterns were identified ranging from highly isotropic (Cluster 1: red) to moderately anisotropic (Cluster 2: blue) to anisotropic (Cluster 3: green). Results are shown for the MNI2009bAsym template. See Appendix Section A. 1 for similar plots for Agile12v2016, Colin27, and the templates combined42
3.5 A comparison of voxel overlap and distance metrics for establishing spatial correspondence between brain regions as evaluated on fMRIPrep output. (A) Multiple views showing the location of AFIDs (black dots) relative to three commonly used ROIs used in voxel overlap measures (the pallidum, striatum, and thalamus). (B,C) The histograms for voxel overlap (Jaccard index) and AFRE, respectively. The distribution for AFRE is more unimodal with a more interpretable dynamic range (in mm ) compared to voxel overlap. Trellis plots demonstrate evidence of focal misregistrations identified by AFRE not apparent when looking at ROI-based voxel overlap alone (D).45
3.6 Investigating relationships between voxel overlap of the striatum and AFRE for each AFID. Focal misregistrations are identified using AFRE for the following AFIDs: 8-10, 14-18, 21-30. The most commonly misregistered regions include the inferior mesencephalon, superior vermis, pineal gland, indusium griseum, and ventricular regions. Horizontal lines are used to demarcate tiers of AFLE error above which AFRE values are beyond a threshold of localization error alone, i.e. the top horizontal line at 3 mm represents more than 2 standard deviations beyond the mean AFLE. Separate plots for the pallidum and thalamus ROIs are provided in Appendix Section A.3.
3.7 Select views demonstrating registration errors between BigBrainSym and MNI2009bSym. The green dots represent the optimal AFID coordinates in MNI2009bSym space superimposed in both templates to provide a basis for comparing registration differences. While many of the midline AFIDs are stable across both templates, the infracollicular sulcus, pineal gland, splenium, and culmen are misregistered in BigBrainSym (red arrows). The AFIDs draw attention to registration differences in the BigBrainSym space in the tectal plate, pineal gland, and superior vermis (blue arrows).
4.1 Workflow for the quantification of local geometric distortion in a single subject at ultra-high field. The leftmost images represent 3T and 7T images of the same subject in native space at the best equivalent sagittal slice. Qualitatively, the 7 T image is more block-shaped than the equivalent 3T image. Checkerboard visualization reveals areas of registration mismatch after rigid and nonlinear registration stages (affine stage omitted from figure). Finally, we quantify local displacement in millimeters overlaid on the 3T image space using the nonlinear deformation field.
4.2 The histogram of mean automated displacements within the masked brain is shown in (a). Since automated displacement is derived from the Euclidean distance between points, all are greater than 0 mm . The histogram has a right skew deviation. The vertical line shows the mean of 0.94 mm . Local voxel displacement is demonstrated to increase with distance from the 7T image isocenter (b). Voxels across all subjects in the study were binned in a 2D histogram (heatmap) according to displacement and distance from isocenter (log-scaled). Mean (thick line) and standard deviation (thin lines) are shown. Mean automated displacement is demonstrated to increase beyond 1 mm at 80 mm .
4.3 Selected images from displacement maps computed for group analysis are overlaid on the MNI152 template with slice references in world coordinates. All voxels shown are significant after controlling for multiple comparisons using FDR ( $q<0.025$ ). Each displacement map has a corresponding image showing an overlay of the maximum $\mathrm{x}, \mathrm{y}$ or z component shown in red, green or blue, respectively, for all significant regions. The main component found was in the z -direction in $56.84 \%$, followed by the x -direction in $22.51 \%$, and finally the $y$-direction in $20.64 \%$,
4.4 The effect of 7T isocenter position on local distortion. Correlation maps of isocenter position against displacement are overlaid on selected sagittal images on the MNI152 template with corresponding world coordinates. All voxels shown are significant after controlling for multiple comparisons using FDR (q $<0.025$ ). The top and bottom images show regions with statistically significant correlation with the $y$ and $z$ isocenter positions, respectively. Red arrows point to corresponding voxel locations where local automated displacement is plotted against isocenter position. The perpendicular black arrows mark the location of the in-plane isocenter in MNI152 space. Positive correlation is observed with a more anterior (increasing y) and superior (increasing z) position. Negative correlation is observed with a more inferior (decreasing $z$ ) isocenter position. Note that there is no significant correlation in a region of the orbitofrontal cortex shown to have high displacement in Figure 4.3, a region known to be prone
to susceptibility effects.

68
4.5 (a) Plot of automated displacement against gold-standard manual displacement demonstrated good correlation ( $\mathrm{R}=0.8755$, p -value $<0.00$ ). (b) Bland-Altman plotting reveals that automated displacement slightly underestimates manual displacement ( mean $=-0.193 \mathrm{~mm}$ ) with good agreement between manual and automated measures. Gray error bands represent $95 \%$ confidence intervals.
4.6 The effect of vendor-provided gradient distortion correction on local displacement in a single subject. A density map of automated displacements within the masked brain with (solid line) and without (dashed line) distortion correction applied (a). The mean displacement is decreased from 0.548 mm to 0.472 mm with distortion correction (a percent reduction of $13.8 \%$ ). Automated displacement maps are overlaid on the subject's own 3T structural scan (b). Decreased displacement is visualized throughout the scan particularly in the suboccipital region. Distortions identified near air-filled sinuses (floor of the middle fossa and orbitofrontal cortex) remain.74
5.1 Visual inspection of AFIDs revealed good convergence with successive template generation steps showing the baseline correspondence between images followed by the linear only template and finally the combined (linear and nonlinear) template after 10 iterations of template building. The information here is corroborated in Tables 5.2 and 5.386
5.2 Study data demonstrating that the ZI (blue) and surrounding structures can be robustly delineated in vivo using 7T MP2RAGE data (a). Actual image values relate to inherent tissue properties; thus thresholding to a set window (10002000 ms ) in MP2RAGE is actually meaningful in comparison to traditional weighted images. The striking similarity between a histological atlas [Schaltenbrand and Wahren, 1977] and in vivo study data is demonstrated in several select slices ( +2.0 and -7.0 mm relative to the anterior commissure in the coronal plane; -3.5 mm relative to the axial plane) (b). Visualization of the models of the RN, STN, and ZI in group average space (c).89
5.3 Study data demonstrates that the optimal target for essential tremor patients is likely the fasciculus thalamicus (fct) rather than the caudal zona incerta. Representative axial slices of the upper mesencephalon in the study group average space depicting T2 and MP2RAGE contrasts (a). The stereotactic target for essential tremor as described by Nowacki [8] is shown as a red dot within the posterior subthalamic area based on the relative location of the STN and RN in T2 images. The same target is superimposed onto the MP2RAGE average with outlines of the fct and cZI in yellow and blue respectively (a). Note that while no contrast can separate the fct from the cZI in the T2 image, the two can be visualized separately in the corresponding MP2RAGE map with the target lying within the fct region. The optimal stimulation electrode for a patient with essential tremor is shown in pink (b) as well as the relative location of the zona incerta and fct (yellow).
5.4 The crosshair is placed on the location identified by [Kerl et al., 2013] as the rostral ZI but overlaid on our joint MP2RAGE and T2SPACE templates. The corresponding location on the the MP2RAGE (T1 map) sequence demonstrates that this feature is actually hypointense on T1 map suggestive of white matter and thus represents the fasciculus lenticularis rather than the rZI. See Figure 5.2 b for the corresponding labels in a histological reference space. . . . . . . . 92

## List of Tables

1.1 $T_{1}$ values at 7-Tesla as acquired from the literature [Marques and Norris, 2017] ..... 20
3.1 Summary of fiducial localization error across brain templates ..... 41
3.2 Mean and inter-rater fiducial localization error pre- and post-QC for the in- cluded OASIS-1 subjects for all AFIDs. ..... 44
3.3 Voxel overlap (Jaccard and Kappa) of the pallidum, striatum, and thalamus after linear registration only and combined linear /nonlinear registration. ..... 46
3.4 AFRE after linear registration alone and combined linear/nonlinear registration. ..... 48
3.5 AFIDs demonstrating evidence of template-to-template misregistration for Big- BrainSym with MNI2009bSym and BigBrainSym with MNI2009bAsym as well as correspondence differences between MNI2009bAsym and MNI2009bSym. 50
4.1 Mean and maximum displacements in millimeters for ROIs part of the lobar and ATAG subcortical atlases. ..... 67
4.2 Mean and maximum displacements in millimeters for ROIs from the Harvard- Oxford atlas meeting thresholds for statistical significance. ..... 69
4.3 Correlation of displacements with change in position of image isocenter for regions meeting thresholds for statistical significance. ..... 70
4.4 Fiducial registration error (in mm ) with successive registration steps in different categories. ..... 72
5.1 MRI sequence details. ..... 82
5.2 Improvement in linear and nonlinear AFRE with multiple iterations of template creation. ..... 87
5.3 AFRE summarized for the final template used in this study (10th iteration). ..... 88
5.4 Summary of T1 values, volume, centroids of key structures of the ZI region. ..... 90
5.5 Comparison between MP2RAGE sequence used in this study and that used by [Forstmann et al., 2014]. ..... 95

## List of Abbreviations and Nomenclature

| General Terminology |  |
| :--- | :--- |
| 3D | three-dimensional |
| 7T | seven Tesla |
| AC | anterior commissure |
| AC-PC | anterior-posterior commissure |
| AChA | anterior choroidal artery |
| al | ansa lenticularis |
| cZI | caudal zona incerta |
| CC | corpus callosum |
| CNR | contrast-to-noise ratio |
| CSF | cereberospinal fluid |
| DBS | deep brain stimulation |
| DRTT | dentatorubrothalamic tract |
| ET | essential tremor |
| fct | fasciculus cerebellothalamicus |
| fl | fasciculus lenticularis |
| FDR | false discovery rate |
| FLE | fiducial localization error |
| FRE | fiducial registration error |
| ft | fasciculus thalamicus |
| GM | gray matter |
| GPe | globus pallidus externus |
| GPi | globus pallidus internus |
| ICBM | International Consortium for Brain Mapping |
| IG | indusium griseum |
| IPF | interpeduncular fossa |
| lin | linear |
| LMS | lateral mesencephalic sulcus |
| LV | lateral ventricle |
| MB | mamillary bodies |
| MER | microelectrode recording |
| MMSE | mini-mental state examination |
| MNI | Montreal Neurological Institute |
| MPTP | 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine |
| nlin | nonlinear |
| OR | operating room |
| PC | posterior commissure |
| PD | Parkinson's disease |
| PMJ | pontomesencephalic junction |
| PSA | posterior subthalamic area |
|  |  |

## General Terminology (Continued)

| raprl | prelemniscal radiations |
| :--- | :--- |
| RN | red nucleus |
| ROI | region-of-interest |
| rZI | rostral zona incerta |
| SNpc | substantia nigra pars compacta |
| SNpr | substantia nigra pars reticulata |
| SNR | signal-to-noise ratio |
| STN | subthalamic nucleus |
| UHF | ultra-high field |
| VIM | ventral intermediate nucleus of the thalamus |
| VLp | ventrolateral posterior nucleus of the thalamus |
| VPL | ventral posterolateral nucleus of the thalamus |
| WM | white matter |
| ZI | zona incerta |


| Medical Imaging Terminology |  |
| :---: | :---: |
| BIDS | Brain Imaging Data Structure |
| CT | computed tomography |
| DICOM | Digital Imaging and Communications in Medicine |
| DTI | diffusion tensor imaging |
| FA | fractional anisotropy |
| HCP | Human Connectome Project |
| MD | mean diffusivity |
| MPRAGE | magnetization prepared rapid acquisition gradient echoes |
| MP2RAGE | magnetization prepared 2 rapid acquisition gradient echoes |
| MRI | magnetic resonance imaging |
| NIfTI | Neuroimaging Informatics Technology Initiative |
| NMR | nuclear magnetic resonance |
| PAT | parallel acceleration technique |
| PD | proton density |
| RF | radiofrequency |
| SA2RAGE | saturation- prepared with 2 rapid gradient echoes |
| SPACE | Sampling Perfection with Application-optimized Contrasts by using different flip angle |
| QMRI | quantitative MRI |
| QSM | quantitative susceptibility mapping |
| T1w | T1-weighted |
| T2w | T2-weighted |
| TE | echo time |
| TI | inversion time |
| TR | repetition time |
| TSE | turbo spin echo |

Terms Introduced by this Thesis
AFID anatomical fiducial
AFLE anatomical fiducial localization error
AFRE anatomical fiducial registration error

## Chapter 1

## Introduction

Neurological disorders represent the predominant contributor to the global burden of disease [Feigin et al., 2017]. For the vast majority of these diseases, effective treatments have unfortunately remained elusive. Neuromodulatory therapy is broadly aimed at providing treatments that improve upon quality of life by either overriding or compensating for pathological activity. The need for improved treatments is paramount particularly with an aging population.

Stereotactic neurosurgery is a subspecialty within the practice of neurosurgery concerned with accurate targeting of brain structures for diagnostic and therapeutic purposes. The word stereotaxy is derived from the Greek words "stereos-", for three-dimensional (3D), and "-taxy", for arrangement ${ }^{1}$ first used by Horsley and Clarke in 1908 [Horsley and Clarke, 1908]. The pioneering work of Leksell and others has resulted in focal surgical treatment options for previously intractable disorders, such as essential tremor and Parkinson's disease [Spiegel et al., 1947, Leksell, 1949, Peters, 2006]. While the mechanical techniques used to accurately target regions with millimetric accuracy have improved relatively modestly in the intervening half century, a variety of neuromodulatory technologies such as biocompatible implanted electrodes and drug delivery systems have since been developed and are available as part of the stereotactic surgeon's armamentarium. Concurrent developments in noninvasive imaging are leading to increasingly robust biomarkers of anatomical and functional substrates of disease that may represent putative therapeutic targets for a wide range of neurodegenerative and chronic illnesses.

This dissertation explores a gap in stereotaxy, namely the continued reliance by neurosurgeons on histological spatial references rather than patient-specific imaging, and presents novel techniques using magnetic resonance imaging (MRI), specifically in the use of highfield images, validation of the correspondence between images, and high-resolution methods to visualize structures not previously seen in vivo.

In this introductory chapter, stereotactic neurosurgery is reviewed from the initial stages of development to present day. The impact of the invention of novel non-invasive imaging methods, i.e. magnetic resonance imaging (MRI), on the practice of stereotaxy is reviewed. Recent MRI advancements, specifically ultra-high field imaging and quantitative MRI, are explored in detail as well as implications of these developments on improving therapeutic targeting of brain structures.

### 1.1 Stereotactic Neurosurgery

Invention and innovation have been central to the evolution of the field of stereotactic neurosurgery, leading to increasingly safer and accurate surgical targeting of brain regions. The history of stereotaxy began in the late 19th century (see [Peters, 2001, Grunert et al., 2003, Peters,

[^0]

Figure 1.1: The pioneering work of Victor Horsley and Robert Clarke who coined the term "stereotaxy" and proposed a system involving the combination of (a) a mechanical frame with (b) a histological reference space using frozen sections for (c) electrolytic study of the deep cerebellar nuclei [Horsley and Clarke, 1908].

2006, Gildenberg and Krauss, 2009] for reviews on this topic). The first attempts at mechanical localization were reported by Russian physician, D. N. Zernov, who invented a device called an "encephalometer" for localizing brain structures based on skull landmarks [Kandel and Shchavinskii, 1973, Blomstedt et al., 2007, Gildenberg and Krauss, 2009]. This work was further extended by his student N. V. Altukhov for targeting deeper structures including the basal ganglia [Grunert et al., 2003, Gildenberg and Krauss, 2009]. However, it was Victor Horsley and Robert Clarke who were the true pioneers and coined the term "stereotaxy" in 1908, reporting the development of a surgical apparatus for targeting of the deep cerebellar nuclei in experimental animals [Horsley and Clarke, 1908] (Figure 1.1). The frame of this instrument was designed to be rigidly fixed to the cranium allowing for the description of points within the frame based on 3D Cartesian ( $\mathrm{x}, \mathrm{y}, \mathrm{z}$ ) coordinates. Concurrent with this physical apparatus, Clarke and Horsley devised a brain atlas from histological sections, providing a mapping between the subject under investigation and a prepared cadaveric specimen (Figure 1.1; covered in Section 1.1.2). While Clarke and Horsley alluded to the use of the apparatus in humans, it was not until 1947 that Ernest Spiegel and Henry Wycis reported the first human device, which
they called the Model V [Spiegel et al., 1947]. After visiting Spiegel, Lars Leksell developed a significant modification by introducing the first "centre-of-arc" based system (Figure 1.2) [Leksell, 1949, Meyerson and Linderoth, 2009]. The stereotactic methodology was further refined in Paris by Jean Talairach [Talairach et al., 1957] into a meticulous system integrated with collinear x-ray pneumoencephalography and cerebral angiography, allowing precise réperage direct (direct investigation) [Gildenberg and Krauss, 2009, Bancaud et al., 1965].


Figure 1.2: Stereotactic equipment in humans. (a) The Model V instrument of Spiegel and Wycis introduced in 1947 requiring plaster casting as a means to fix the frame to the head. (b) The first "centre-of-arc" system invented by Lars Leksell in 1949.

### 1.1.1 Early Indications and Treatment Modalities

While the indications for stereotaxy are growing, here we focus on two classic neurological disorders: Parkinson's disease (PD) and essential tremor (ET). Parkinson's disease, which was first described by James Parkinson [Parkinson, 2002] as the "shaking palsy" or paralysis agitans in 1817 , is characterised by a number of involuntary clinical manifestations including resting tremor, stiffness of motor movements (also called rigidity), slowed movements (bradykinesia), and postural imbalance (or instability). Essential tremor is a common disorder affecting up to $5 \%$ of patients older than 60 characterized by appendicular (arms or legs) or axial tremor typically action-related and occurring in the $6-8 \mathrm{~Hz}$ range. Despite best medical management, this disorder can remain debilitating in around half of patients (Figure 1.3) [Hallett, 2014].

Preceding the development of good pharmacological treatments, clinicians explored surgical therapy as an option for their patients. However, as Gildenberg notes in his historical review [Gildenberg and Krauss, 2009], the consideration of surgical therapy of the basal ganglia was

## Normal



## Essential Tremor



Figure 1.3: Spiral drawing under normal conditions and for a patient with essential tremor.
initially stalled due to the influential opinion of pioneering neurosurgeon, Walter Dandy, who contended that "basal ganglia resections were universally lethal" and who considered the caudate nucleus the "seat of consciousness" [Dandy, 1946, Gildenberg and Krauss, 2009].

Eventually, these views were challenged by a few surgeons who developed open neurosurgical approaches to the deep brain. Russell Meyers began performing surgical procedures in the caudate nucleus with modest benefit [Meyers, 1951a] without the mortality previously reported by Dandy. He went on to develop surgical approaches to the basal ganglia facilitating interruptions of connections, including open approaches to resection of the fiber connections in the deep nuclei [Meyers, 1951b]. Irving Cooper was credited with accidentally discovering that ligation of the anterior choroidal artery (AChA) [Cooper, 1953] resulted in relief of Parkinsonism in a few patients, and went on to produce a larger series where he intentionally ligated the AChA and while some good effect was achieved, observed that unfortunately a subset of patients became either transiently or permanently hemiplegic [Cooper, 1960]. Further investigation revealed that the arterial distribution of the AChA was non-specific to Parkinsonism and not uncommonly supplied blood to important regions including the posterior limb of the internal capsule along the corticospinal tract. These findings resulted in a shift in interest from vascular to parenchymal targeting, with a specific focus on investigating the specific role of deep structures perfused by the AChA. Overall these reports resulted in an understanding that the interruption and disruption of specific deep brain regions could provide effective treatment for movement disorders.

These discoveries using open neurosurgical techniques enabled Spiegel and Wycis, with
their innovations in stereotactic instrumentation, to propose the use of more targeted approaches to the region. In this context, the first reported stereotactic case, a thalamotomy of the medial nucleus of the thalamus was successfully performed by Spiegel and Wycis [Spiegel et al., 1947, Grunert et al., 2003]. The first successful treatments of Parkinson's disease via stereotactic surgery were performed in the 1950's [Narabayashi and Okuma, 1953, Guiot and Brion, 1953, Spiegel and Wycis, 1954, Svennilson et al., 1960, Laitinen et al., 1992]. Cooper began to use more focal treatments, experimenting with different methods for ablation, and played an important role in the development of chemoablation and cryoablation [Cooper, 1960]. Multiple different targets were being explored for ablative (radiofrequency) treatment of Parkinson's disease and debilitating tremor including the posterior ventrolateral area of the thalamus (VLp), globus pallidus internus (GPi), and subthalamic nucleus [Andy et al., 1963, Mundinger, 1965, Houdart et al., 1966, Velasco et al., 1972].

With two exceptions, further technical developments in physical devices for stereotactic treatment were relatively modest until recently. First, Lars Leksell developed an arc-based frame enabling a physical system whereby Cartesian and polar coordinates could be combined (??b). The arc frame simplified the process of determining the trajectory. Once a target location was determined, the trajectory location could be optimized by manipulating the ring and arc angles with the target remaining centered. Second, the invention of frameless computeraided navigation systems [Kelly et al., 1986] revolutionized standard cranial surgery for the localization of intracranial lesions (e.g. tumours), which are now routinely used in clinical practice. Please see [Peters, 2001, Grunert et al., 2003, Peters, 2006] for more details on the development of stereotaxy.

### 1.1.2 Brain Atlases

Brain atlases are a fundamental aspect of stereotactic neurosurgery, and central to the focus of this thesis. The practice of stereotaxy can be considered analogous to geographical exploration and the reliance on a topographic atlas or map. One key distinction must be made between what constitutes a template versus an atlas. A template refers to the brain section, or raw image data prepared in a way that can be described in terms of spatial coordinates, while an atlas refers to an overlay or annotation of the structures in the template. The choice of "reference space" can impact the accuracy of surgical targeting. However, while taking a wrong turn due an inaccurate map can often be compensated for, a deviation from the expected path in a stereotactic procedure could represent the difference between optimal therapy and a devastating complication for the patient. In the following section, we discuss the development of brain atlases leading to the present day.

When neurosurgeons first began considering therapy for deep brain structures, they relied on knowledge from cadaveric specimens and sections [Horsley and Clarke, 1908, Spiegel et al., 1947, Talairach et al., 1957]. A template brain specimen was developed in parallel with the Horsley and Clarke instrument, consisting of glass-mounted frozen sections from a Macaca mulatta (rhesus macaque) specimen [Horsley and Clarke, 1908] (Figure 1.1). Several decades later, these same principles were used to devise a brain template for humans [Spiegel et al., 1947]. These post-mortem brain atlases continued to evolve with two of the most commonly referenced stereotactic atlases being created by Jean Talairach [Talairach et al., 1957] and Schaltenbrand and Wahren [Schaltenbrand and Wahren, 1977].

Computing the correspondence between an external template and the patient or subject is central to the process of atlas-assisted sterotaxy. Without access to other resources, these decisions were originally made by inferring the location of structures based on external cranial landmarks [Grunert et al., 2003]. Talairach appreciated that external cranial landmarks were unreliable for describing the location of structures within the cranial vault employing intra-operative adjuncts such as x-ray and more specifically ventriculographic and angiographic techniques to provide internal references for correspondence, a process which he referred to as a réperage indirect [Talairach et al., 1957]. Ventriculographic studies allowed for alignment based on observed features in both the template and patient images. To establish correspondence between the postmortem brain and the patient under investigation, Talairach observed that several salient features could be visible on ventriculographic studies, specifically, the anterior commissure ( $\mathrm{AC)}$ and posterior commissure (PC), two dense white matter tracts connecting the hemispheres. Talairach's réperage procedure was refined over time into a more comprehensive system for alignment of individual patient datasets with a template space through a series of axes-specific scaling steps, since referred to as the Talairach proportional grid normalization [Talairach et al., 1957, Talairach and Tournoux, 1988, Brett et al., 2002]. Under this process, each subject was brought into alignment using first an AC-PC transformation followed by scaling in the standard cardinal ( $\mathrm{x}, \mathrm{y}, \mathrm{z}$ ) directions relative to AC-PC. This would be considered a 9 degree-of-freedom transformation (consisting of 3 degrees for translation, 3 for rotation, and 3 for scaling). It became possible to thus relate different anatomical labels onto the Talairach template.

Atlas representations have continued to evolve and the process, increasingly refined. Talairach's initial template had several limitations: it was devised from a single subject (a 60 year-old female) and also the left and right hemispheres had to be sectioned in two different orientations (one axial and the other coronal). Symmetry between hemispheres was thus a necessary assumption. Another commonly used atlas developed by Schaltenbrand and Wahren [Schaltenbrand and Wahren, 1977] used three separate brains cut in each of the standard cardi-
nal planes. To prepare the transparent atlas overlays, 10 brains were used in the preparation of the coronal overlays, 13 sagittal, and 7 transverse [Schaltenbrand and Wahren, 1977]. The spatial correspondence between the same structures (particularly thalamic nuclei) in the different orientations has been reported to be poor [Niemann and Van Nieuwenhofen, 1999] as a result of inter-subject variability which is a confound in atlas-based studies. To complicate matters further, tears and processing artifacts (e.g. tissue distortions) were encountered and required substitution with additional brain sections, which in total amounted to 111 brains being used in the preparation of the atlas [Schaltenbrand and Wahren, 1977]. Finally despite high detail in the plane of the sections, individual sections were variably spaced from between 0.5-1.5 mm presenting some problems to their use for accurate stereotactic targeting. The process of improving the correspondence between the atlas and subject or template continues to be refined [Brett et al., 2002, Amunts et al., 2014]. How the correspondence between templates and subject datasets can be quantified, is the subject of study in Chapter 3 of this thesis.

### 1.1.3 The Importance of Imaging

The development of stereotaxy coincided with a number of important imaging innovations crucial to improving targeting accuracy. The discovery of x-rays and the development of x-ray imaging by Wilhelm Roentgen [Roentgen, 1895] followed by ventriculography [Dandy, 1918], and cerebral angiography by Egaz Moniz [Doby, 1992] enabled visualization of the patient's cranial anatomy, ventricles, and cerebral vasculature respectively. These innovations allowed for more accurate correspondence with histological atlases by permitting linear alignment of the atlas to these anatomical features [Spiegel et al., 1947]. The ability to see the cerebral vessels also enabled Talairach's réperage [Talairach et al., 1957], and to directly avoid vessels seen on angiograms [Talairach et al., 1957, Bancaud et al., 1965]. Calcification of the pineal gland could also be used as an internal reference frame for anatomy, and was used, as such, for the first successful thalamotomy [Spiegel et al., 1947, Grunert et al., 2003]. Targeting of specific parenchymal regions remained elusive with x-ray technology since, compared to bone or contrast-enhanced regions, $x$-ray contrast between different soft tissues is very low, and thus these tissues still had to be targeted using indirect methods based on atlas correspondence.

### 1.1.4 Deep Brain Stimulation

Over the last several decades, deep brain stimulation (DBS) has become established as an effective and reversible means of providing focal surgical therapy for patients with movement disorders [The Deep-Brain Stimulation for Parkinson's Disease Study Group, 2001, Hariz, 2017]. These efforts were driven in part by increased recognition of side effects of systemic
pharmacotherapy for the treatment of movement disorders, namely levodopa for Parkinson's disease [Shoulson et al., 1975, Marsden and Parkes, 1976, Lees et al., 1977, Krack et al., 1999, Poewe, 2009, Williams et al., 2010], and thus renewed interest in other treatment options including surgery. DBS was pioneered by Alim Benabid [Benabid et al., 1988, Benabid et al., 1996], who demonstrated that high-frequency ( $>100 \mathrm{~Hz}$ ) electrical stimulation applied to specific deep brain targets could be effective for treating motor symptoms, coinciding with new insights into the basal ganglia circuitry (Figure 1.4) derived from microelectrode recordings of primate PD models by Mahlon Delong and colleagues [Bergman et al., 1990, DeLong, 1990]. Neuromodulatory therapy to specific basal ganglia targets such as the globus pallidus and subthalamic nucleus (STN) were being explored as candidate regions for focal electrical stimulation [Poewe, 2009].

Benabid reported on the efficacy of DBS of the STN region in 1993 [Pollak et al., 1993, Benabid et al., 1994, Limousin et al., 1995]. For Parkinson's disease, DBS allows reduction of dopaminergic therapy [Deuschl et al., 2006], thus decreasing drug-induced motor symptoms like dyskinesia, while enabling more consistent efficacy and minimization of on/off fluctuations [Tomlinson et al., 2010, Okun, 2012]. A recent cost-analysis reported that DBS may be associated with lower medical costs at follow-up due to reductions in the long-term need of polypharmacy ( $\$ 65 \mathrm{~K}$ over 10 years in the United States) [Hacker et al., 2016]. These findings have also motivated earlier intervention in PD patients with demonstration of superiority to medical therapy with respect to motor symptoms, quality of life, and levodopa-related dyskinesias [Schuepbach et al., 2013], as well as evidence for decreased medication costs [Hacker et al., 2016].

Benabid also proposed the use of DBS for the treatment of medically refractory essential tremor (ET), specifically suggesting the ventralis intermedius (VIM) nucleus as the target, which is also known as the ventrolateral posterior (VLp) nucleus ${ }^{2}$ [Benabid et al., 1988, Benabid et al., 1991]. Microelectrode recordings (MER) studies have revealed that neurons in the VIM are synchronous with tremor in the contralateral extremity, and as such believed to play a crucial role in tremor modulation [Narabayashi, 1986]. Other groups have found success with stimulation of other regions including the caudal zona incerta (cZI) within the posterior subthalamic area [Plaha et al., 2006, Blomstedt et al., 2007, Fiechter et al., 2017, Nowacki et al., 2018], based on previous lesional work from Spiegel [Spiegel et al., 1964] and Mundinger [Mundinger, 1965, Mohadjer et al., 1990]. Studies have suggested that the cZI may require comparatively reduced amplitude of stimulation to achieve a therapeutic benefit [Blomstedt et al., 2010], which may result in longer life of the stimulator battery, although this has not been evaluated as a randomized controlled trial. Nowadays, DBS is considered the first-line

[^1]surgical treatment option for patients with essential tremor, given it is a reversible treatment [Tasker et al., 1997, Pahwa et al., 1999, Schuurman et al., 2000, Pahwa et al., 2001].

The exact mechanism by which electrical stimulation works as therapy remains poorly understood. Early on, Benabid and colleagues posited that DBS induced "functional inhibition" of the target region [Benabid et al., 1988, Benabid et al., 1996]. Studies employing neuronal recording methods demonstrate evidence of neuronal suppression in the vicinity of the stimulation target [Kiss et al., 2002, Hamani et al., 2004], which may be mediated by alterations of the extracellular milieu, particularly increased potassium concentrations [Shin et al., 2007, Florence et al., 2016]. Beyond local suppression, evidence suggests more far-reaching interactions with projected regions [Anderson et al., 2004, Miocinovic et al., 2006, Shimamoto et al., 2013, Horn et al., 2017c]. At the level of the local field, these findings may manifest as changes in the oscillatory background activity and suppressed with stimulation, which have been observed in the beta $(12-30 \mathrm{~Hz})$ range in patients with PD [Kuhn et al., 2008] and at the tremor frequency $(6-8 \mathrm{~Hz})$, a concept referred to as thalamocortical dysrhythmia [Gallay et al., 2008], in patients with ET [Hua and Lenz, 2005, Raethjen and Deuschl, 2012, Hallett, 2014, Hariz and Blomstedt, 2017]. Overall, high-frequency DBS likely involves a dynamic interplay between inhibitory and excitatory mechanisms at both the local and network levels, and also involving a range of time scales of clinical effect from immediate (for tremor) to long-term (weeks to months) for obsessive compulsive disorder and refractory depression (see [Herrington et al., 2016] for a recent review).

### 1.1.5 Conventional Approaches for Stereotactic Targeting

The work by Delong and colleagues elucidated clear functional roles in different pathways of the basal ganglia and how dysfunction within this circuit could produce symptoms like Parkinson's disease [Bergman et al., 1990, DeLong, 1990, Poewe, 2009]. Delong studied experimental models of PD (1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine; MPTP) investigating neuronal activity in these deep circuits using microelectrode recordings. Coupled with the routine use of awake MER by Benabid and colleagues, with few exceptions, MER has remained a central component of DBS procedures. Awake MER surgery has been central to DBS surgery and the origins of how the technique was discovered. However, use of MER results in increased operating room time, patient discomfort, and risk of complications [Zrinzo et al., 2012]. Systematic reviews have identified that MER results in a higher hemorrhage rate [Zrinzo et al., 2012].

Side effects include pyramidal effects (stimulation-locked clonic movements), dysarthrophonia, cognitive, behavioural side effects [Krack et al., 2002, Lambert et al., 2012], eye devi-


Figure 1.4: Schematic of the cerebellothalamocortical and pallidothalamocortical circuits based on tract-tracing studies in primates [Gallay et al., 2008]. All major cerebellar, basal ganglia, and cortical regions are represented. M1 = primary motor cortex; PMc = caudal premotor ; SMA = supplementary motor area; $\mathrm{PMr}=$ rostral premotor; $\mathrm{PFC}=$ prefrontal cortex; VLp = ventrolateral posterior thalamus; $\mathrm{VM}=$ ventral medial nucleus; $\mathrm{VLa}=$ ventrolateral anterior; VApc $=$ ventral anterior parvocellular division; $\mathrm{fct}=$ fasciculus cerebellothalamicus; $\mathrm{ft}=$ fasciculus thalamicus; $\mathrm{fl}=$ fasciculus lenticularis; al $=$ ansa lenticularis; STh = subthalamic nucleus; $\mathrm{SNc}=$ substantia nigra pars compacta; $\mathrm{GPi}=$ globus pallidus internus; $\mathrm{GPe}=$ globus pallidus externus.
ation [Shields et al., 2007], and speech effects [Tripoliti et al., 2008, Åström et al., 2010]. In 2000, Hariz first reported an increased risk of complications in those with pre-existing speech and cognitive decline [Hariz et al., 2000, Hariz, 2017]. Beyond complication avoidance, all these factors motivate the need for better ways of optimizing the target location [Delong and Wichmann, 2012], for which improving the quality of pre-operative patient imaging has been crucial.


Figure 1.5: Demonstration of the intersubject variability in the location of anatomical structures of the deep brain and specifically the subthalamic region. The upper subfigures represent sagittal sections of the brain from two different cadavers ( Hb 1 is shown with red outlines and filled shapes while Hb 2 is shown with black outlines and light gray filling). The lower subfigure demonstrates variability in the locations for four specimens $(\mathrm{Hb} 1, \mathrm{Hb} 2, \mathrm{Hb} 3, \mathrm{Hb} 5) . \mathrm{PTT}=$ pallidothalamic tractotomy. $\mathrm{CTT}=$ cerebellothalamic tractotomy. Figure taken from the open access article by [Gallay et al., 2008].

### 1.2 Magnetic Resonance Imaging

The development of MRI began with nuclear magnetic resonance (NMR) when two researchers Felix Bloch and Edward Purcell independently discovered that they could "listen" to atonic nuclei [Bloch, 1946, Purcell et al., 1946]. Further developments were related to the idea that the magnetic field of a sample could be manipulated by applying an external electromagnetic field [Ernst et al., 1987], termed "pulsed NMR" and eventually the suggestion by Richard Ernst that encoding of space within a sample was possible using frequency and phase [Ernst and Anderson, 1966]. These discoveries formed a foundation by which Paul Lauterbur and Peter Mansfield could independently describe magnetic resonance imaging (MRI) [Mansfield and Grannell, 1973, Lauterbur, 1973, Mansfield, 1977]. Here we outline some of the important concepts necessary to provide context for the application of MRI to stereotactic neurosurgery.

### 1.2.1 Fundamentals of Magnetic Resonance Imaging

a)

b)



Figure 1.6: (a) Protons spinning in free space, (b) protons spinning under the influence of $B_{0}$, (c) A $B_{1}$ radiofrequency pulse tips the magnetization vector, $\vec{M}$ into the transverse plane and over time relaxes back to equilibrium in alignment with $B_{0}$ (rotating frame). The arrows in (a) and (b) refer to the magnetic moment of the spinning nuclei.

Nuclear magnetic resonance (NMR) is based on the absorption and emission of energy in the radiofrequency $(\mathrm{MHz})$ range by atomic nuclei. Atoms with an odd number of protons or neutrons have a net spin or magnetic moment, $\mu$, behaving, at a macroscopic level, like tiny magnets. The strength of the magnetic moment quantifies the strength of this magnetism. When placed under the influence of a static magnetic field ( $B_{0}$ ), atoms with net spin align either parallel with or antiparallel to the field producing a net magnetization, $\vec{M}$ (Figure 1.6). This net magnetization is in the direction of $B_{0}$, which is conventionally known as the longitudinal, or $\vec{z}$, direction. There is no net magnetization in the transverse plane, i.e. in the $\vec{x}$ and $\vec{y}$ components, since the orientation of the spins is random in this plane and thus cancel out. Different nuclei precess at a predefined frequency (i.e. Hertz $=\mathrm{Hz}$ ) for a given $B_{0}$, also known as the Larmor frequency:

$$
\begin{equation*}
f_{0}=B_{0} \times(\gamma / 2 \pi) \tag{1.1}
\end{equation*}
$$

where $\gamma / 2 \pi$ is the gyromagnetic ratio, which is constant for a given atomic nucleus and magnetic field strength. For hydrogen nuclei, which in water make up the most abundant source measured in biological tissues, $\gamma / 2 \pi$ is $42.575 \mathrm{MHz} /$ Tesla. The net magnetization at equilibrium, $\vec{M}_{0}$, under the influence of $B_{0}$ can be described by the following equation:

$$
\begin{equation*}
\overrightarrow{M_{0}}=\frac{N_{s} \gamma^{2} \hbar^{2} \overrightarrow{B_{0}}}{4 k_{B} T} \tag{1.2}
\end{equation*}
$$

where $N_{s}$ is the number of spinning atomic nuclei, $\hbar$ is Planck's constant, $k_{B}$ is Boltzmann's constant, and $T$ is the temperature (in degrees K ). Overall, this equation demonstrates that sensitivity to detection of NMR signal is dependent on having an abundance of detectable nuclei
(i.e. protons in water) and a higher main magnetic field strength (see Section 1.2.2). Decreasing the temperature also has the effect of increasing net magnetization but is not feasible in living subjects.

When a radiofrequency (transmit) pulse ( $B_{1}$ ) is applied at an angle relative to $B_{0}$ (classically perpendicular), the aligned nuclei are excited into a higher energy state (Figure 1.6c). Upon cessation of the pulse, the stimulated nuclei return to the equilibrium state $\left(M_{0}\right)$ in alignment with the main magnet, a phenomenon known as relaxation, which is described by the Bloch equations:

$$
\begin{equation*}
\frac{d \vec{M}}{d t}=\vec{M} \times \gamma \vec{B}+\frac{M_{x} \vec{x}+M_{y} \vec{y}}{T_{2}}-\frac{\left(M_{z}-M_{0}\right) \vec{z}}{T_{1}} \tag{1.3}
\end{equation*}
$$

where $T_{1}$ and $T_{2}$ relaxation times represent two independent processes that describe how the spins recover after the application of a radiofrequency $\left(B_{1}\right)$ pulse to return to equilibrium (i.e. in alignment with $B_{0}$ ). More specifically, $T_{1}$ represents the longitudinal relaxation time constant for nuclei to recover to equilibrium and $T_{2}$ represents the transverse relaxation time constant for nuclei to dephase or decay to equilibrium. In an NMR experiment, this process of relaxation back to equilibrium can be recorded using radiofrequency receiver coils. $T_{1}$ and $T_{2}$ relaxation can be modeled as exponential curves with net magnetization in the component longitudinal ( $M_{z}$ ) and transverse ( $M_{x y}$ ) directions described using the following:

$$
\begin{gather*}
M_{z}(t)=M_{z}(0)\left(1-e^{\frac{-t}{T_{1}}}\right)  \tag{1.4}\\
M_{x y}(t)=M_{x y}(0)\left(e^{\frac{-t}{T_{2}}}\right) \tag{1.5}
\end{gather*}
$$

noting that $M_{z}(0)$ is equivalent to $M_{0}$. Specifically, $T_{1}$ is defined as the length of time for $M_{0}$ to recover by a factor of $(1-1 / e)$, where $e$ is Euler's number, that is to $63.2 \%$ of its original value. $T_{2}$ is defined as the length of time for $M_{x y}(0)$ to decay to $1 / e$, or $36.8 \%$ of its maximal value. Example relaxation curves are demonstrated in Figure 1.7.

A basic NMR experiment requires a static ( $B_{0}$ ) field, transmit RF ( $B_{1}+$ ) field, and receiver RF ( $B_{1^{-}}$) coil for detecting magnetic parameters in a test tube with a homogeneous sample. To produce images, gradient coils are a crucial additional element that allows encoding of information about two key properties of spinning atomic nuclei: frequency and phase. Coils are constructed that produce a spatially varying magnetic field by manipulating the frequency and phase of the atomic spins to be unique at every point in the scanned region by using frequency encoding $\left(G_{f}\right)$ and phase encoding $\left(G_{\phi}\right)$ gradients, respectively. These magnetic manipulations occur at a much smaller scale than $B_{0}$ and are in the range of milli-Tesla per meter ( $m T / m$ ). For a gradient $G_{x}$, the resonance frequency at each location can thus be expressed using a linear


Figure 1.7: $T_{1}$ (a) and $T_{2}$ (b) relaxation can be modeled using exponential functions as demonstrated in Equations 1.4 and 1.5. The $T_{1}$ and $T_{2}$ relaxation times were set at 1500 ms and 50 ms respectively in these simulated examples (dashed lines). (c) Demonstration of how for a given echo time (marked by the dashed line) for recording, a mix of both $T_{1}$ and $T_{2}$-based signal is recorded, as is typical with conventional imaging methods.
equation:

$$
\begin{equation*}
\omega(x)=\omega_{0}+\gamma G_{x} x \tag{1.6}
\end{equation*}
$$

The faster and slower precession of atomic nuclei along this gradient is used to spatially encode information about local net magnetization and stored in Fourier space, also known as kspace. In $k$-space, information across the entire region of interest being scanned is summed together and represented in spatial frequency space. The frequency differences can be recovered from the raw signals recorded from the receiver RF using Fourier transformation to produce the images in magnetic resonance imaging (MRI). The interested reader is referred to the following references for a more thorough description of the theoretical and practical foundations of MRI and Fourier transformation [Nishimura, 1996, McRobbie et al., 2003]. Finally, there are inherent inhomogeneities introduced during the construction of gradient coils that can result in geometric distortion that is important to understand in the context of stereotactic neurosurgery. These issues are further explored in Chapter 4 as are other sources of MRI distortion.

### 1.2.2 Ultra-High Field MRI

Measuring signal from precessing nuclei is an inherently noisy process requiring sensors capable of detecting signal differences on the order of parts per million. As demonstrated in Equation 1.2, one way to increase sensitivity to magnetization is to increase $B_{0}$, which scales at least linearly with magnetic field strength. The increased signal can be exploited in differ-
ent ways, permitting investigation of small structures in the deep brain to be imaged at higher resolution. Ultimately, going to higher main magnetic field strength presents an opportunity to improve in vivo visualization of putative stereotactic targets for neuromodulation.

As an exercise, we specifically consider the STN and the impact of resolution. The STN is one of the smaller deep brain nuclei and one of the key targets for Parkinson's disease, that has been better characterized over the the last few years. The volume of the STN as estimated from histological studies at $240 \mathrm{~mm}^{3}$ with a maximal extent of $\sim 10 \mathrm{~mm}$ in the dorsolateral to ventromedial direction and containing approximately 560000 neurons, thus containing approximately 2300-2400 neurons within a $1 \mathrm{~mm}^{3}$ voxel [Hardman et al., 2002, Hamani et al., 2004], the standard clinical resolution acquired for surgical planning. Increasing the resolution by $40 \%$ isotropically (i.e. reducing the voxel dimensions to $0.7 \mathrm{~mm}^{3}$ ) results in an improvement in the overall number of voxels from 240 with 10 along the maximal extent to 700 voxels (almost 3-fold) with 15 voxels along the maximal extent. Estimates within a voxel also improve from 2300-2400 neurons to $\sim 800$ neurons per voxel. The numbers are more drastic with improvements to 0.5 mm isotropic voxels, representing an 8 -fold increase in number of STN voxels to 1920 with a 2 -fold increase in a single dimension to 20 voxels along the maximal extent. Increasing $B_{0}$ permits an increase in signal that can be used to offset the increased noise associated with higher resolution imaging.

There are also smaller structures in the vicinity of the STN such as the fields of Forel and zona incerta that have not been reliably visualized in vivo, although they have historically been considered potential therapeutic targets. Spiegel and Wycis initially had proposed ansotomy, that is lesioning of the ansa lenticularis, and campotomy, that is lesioning of the nuclei campi perizonalis or Fields of Forel as a means of treating Parkinson's disease [Spiegel and Wycis, 1954, Spiegel et al., 1962, Spiegel et al., 1964]. Later, Mundinger and colleagues described lesioning of the cZI [Mundinger, 1965]. However, it is unclear how well the original pioneers of stereotaxy were able to accurately localize the location of their lesions. To our knowledge, no clear estimates of the size of the cZI or fields of Forel have been made using histology or MRI. Direct visualization of these structures at high-fields will be the subject of Chapter 5.

Moving to higher fields introduces many new challenges (see [Uğurbil, 2017] for review). Most directly, $B_{0}$ inhomogeneities increase with the static magnetic field strength inducing "phase accruals" that lead to distortion and signal loss particularly at tissue interfaces. Of particular concern, these spatial distortions could render the use of 7 T impractical for surgical targeting purposes and is explored in detail in Chapter 4 . An increase in $B_{0}$ also results in an increase in the resonance (Larmor) frequency and smaller wavelengths. The increased resonance frequency leads to higher attenuation and thus the need for higher $B_{1}$ intensity; that is, the RF transmit pulses need to be stronger to excite the atoms in an MR experiment. The excitation


Figure 1.8: One way to increase sensitivity to magnetization (i.e. signal) is to increase $B_{0}$, which scales at least linearly with magnetic field strength. Selected T2-weighted coronal images taken at standard clinical field strength (1.5T) and at 7 T demonstrating the improvement in image contrast in deep brain structures with an increase in $B_{0}$. The zoomed in regions demonstrate the increased detail in structures of the basal ganglia, specifically the pallidum, subthalamic nucleus, and substantia nigra.
angles transmitted to the entire volume, which in ideal circumstances are stable throughout, are inhomogeneous due to both the increased attenuation and standing wave effects. This results in variable intensities throughout the sample in the reconstructed image. Second, the strong RF pulse implies that there is an increase in the power deposited into the subject, which is regulated by the FDA by a measurement called the specific absorption rate. Finally, other features that can be perceived as advantages or disadvantages are the changes in the inherent relaxation parameters of different tissues. $T_{1}$ relaxation times increase with field strength, while T2 relaxation times shorten.

To date, all ultra-high field MRI systems are attached to academic institutions and require highly complex and specialized hardware in order to achieve optimal performance. The process to achieve this performance, the delays in imaging and complex decisions regarding hardware design are well-documented by Kamil Ugurbil in his recent commentary [Uğurbil, 2017]. While not standard with ultra-high field MRI systems, radiofrequency coil design innovations have been crucial for mitigating problems with $B_{1}$ inhomogeneity and SAR. Specifically, the development of parallel transmit ( pTx ) technology, that is the use of multichannel transmit coil
elements rather than a single element, has presented an elegant means of appropriately distributing power across the object being imaged, all while maintaining homogeneous excitation and limiting power deposition.

### 1.2.3 Quantitative MRI

Standard anatomical MRI images are "weighted" by a combination of magnetic parameters ( $T_{1}, T_{2}, \mathrm{PD}$ ) and scanner-related idiosyncracies (e.g. $B_{1}$ field inhomogeneities) limiting interpretability. While a radiologist may be able to qualitatively identify any relative differences in intensity for a given image, high variability exists not only between scanners and subjects, but also for single subjects across different sessions. Post-processing using intensity normalization techniques enable some degree of between participant comparison; however, the local tissue value is devoid of meaning (Figure 1.9). Quantitative MRI (QMRI) involves a shift in paradigm from using MRI as a tool strictly for producing images to one that is also used for measurement.

QMRI sequences measure individual MRI contrast-generating parameters in isolation (see [Weiskopf et al., 2015] for a review). QMRI-based methods typically require the acquisition of multiple scans in order to estimate specific magnetic resonance properties (e.g. $T_{1}$ or $T_{2}$ relaxation). These local parameters are inherently more robust than weighted images, which involve a mix of parameters, and thus better reflect intrinsic tissue properties and can be used to identify specific brain structures and also pathology on the basis of a local measurement. These measurements are also more directly comparable between scanners allowing better standardization compared with conventional weighted images [Deoni et al., 2008, Weiskopf et al., 2013].

Chapter 5 of this thesis focusses on examining the utility of longitudinal relaxometry ( $T_{1}$ $=1 / R_{1}$ ) at ultra-high fields for delineating deep brain structures relevant to stereotactic neurosurgery. $T_{1}$ relaxation is also referred to as spin-lattice relaxation as it involves the transfer of energy between local nuclear spins and the surrounding environment, or lattice. Thus, the structure of the environment heavily impacts the local $T_{1}$ values measured. On the one hand, areas of relatively restricted water mobility (i.e. in the axons of white matter), collisions and interactions with surrounding lipid molecules in myelin result in more rapid relaxation, and thus shorter, $T_{1}$ values. On the other hand, areas of high relative water mobility (e.g. cerebrospinal fluid) have longer relaxation times, relying on random interactions with other water molecules rather than any sort of inherently structured lattice. Generally, as it relates to brain tissue, $T_{1}$ values are highest in cerebrospinal fluid, intermediately high in gray matter, and lowest in white matter (see Table 1.1).

T1-weighted Image


Figure 1.9: Comparison of conventional $T_{1}$-weighted imaging where the units are arbitrary with quantitative $T_{1}$ mapping where the values reflect inherent local tissue properties. Individual subject scan at 7-Tesla using the MP2RAGE sequence [Marques et al., 2010].

Many groups have focussed on the inherent contrast-related advantages of T2-based protocols at high field, due to the rich iron content of many basal ganglia nuclei. However, the advantages are not limited to this contrast type, with $T_{1}$ values not only increasing in a fielddependent manner, but also the dispersion between different tissue types; thus increasing the contrast and thus salience between neighbouring structures with different properties [Rooney et al., 2007, Tourdias et al., 2014].

Table 1.1: $T_{1}$ values at 7-Tesla as acquired from the literature [Marques and Norris, 2017]

| region | $T_{1}(\mathrm{~ms})$ | $T_{2}(\mathrm{~ms})$ |
| :--- | :---: | :---: |
| White Matter | $1100-1400$ | 55 |
| Cortical Gray Matter | $1900-2100$ | 50 |
| Cerebrospinal Fluid | 4400 | NA |
| Blood | 2600 | 7 |
| Putamen | $1520-1700$ | NA |
| Caudate Nucleus | $1630-1700$ | NA |
| Globus Pallidus | $1180-1200$ | NA |
| Red Nucleus / Substantia Nigra | $\sim$ White Matter | NA |



Figure 1.10: Figures from Rooney et al. demonstrating the field-dependent increase in $T_{1}$ values and dispersion in different tissue types [Rooney et al., 2007]. The increase in dispersion of T1 manifests as improved contrast that can be exploited for better delineating boundaries between brain structures. This will be the subject of Chapter 5 .

### 1.3 Thesis Outline

Despite significant advances in stereotactic targeting, a recent analysis of multiple national databases in North America (over 28000 procedures) revealed a surprisingly high rate of hardware revision and removal for deep brain stimulation procedures at a rate of up to $48.5 \%$ related to improper targeting or lack of therapeutic effect [Rolston et al., 2016]. This suggests that further work is necessary in both patient and target selection in patients being considered
for neuromodulation. Lack of adequate visualization with conventional lower field strengths (1.5/3 Tesla) has meant that standard-of-care surgical treatment has relied on indirect targeting using standardized landmarks as described in Section 1.1.2. For this reason, these procedures routinely require awake testing and microelectrode recording, which increases operating room (OR) time, patient discomfort, and risk of complications.

In this manuscript-based thesis, the feasibility of using ultra-high field MRI for stereotactic surgery is explored. The fundamental motivation behind this line of work is that the ability to see structures better will allow for more focal, targeted therapy. To answer this question, this work begins in Chapter 2 with a practical clinical example of integrating a 7T template into the clinical workflow for surgical planning. This technical report is followed with a study examining the practical limits of accuracy using traditional template-assisted stereotaxy in Chapter 3. In Chapter 4, the impact of geometric distortion on our ability to use 7T for surgical targeting is investigated. Finally, in Chapter 5, we demonstrate how quantitative MRI acquired at 7T can be used to visualize stereotactic targets never before seen in vivo. The implications of the thesis and future considerations are summarized in Chapter 6.

## Chapter 2

## Ultra-high field template-assisted deep brain stimulation surgery

This chapter is based on the following manuscript:

- Lau, J. C., MacDougall, K. W., Arango, M. F., Peters, T. M., Parrent, A. G., \& Khan, A. R. (2017). Ultra-High Field Template-Assisted Target Selection for Deep Brain Stimulation Surgery. World Neurosurgery, 103, 531-537.


### 2.1 Introduction

Surgical planning for deep brain stimulation (DBS) surgery is variable from centre to centre as a result of a number of factors including availability of imaging modalities, access to stereotactic equipment, and the institutional and neurosurgical experience. Results of a recent international survey highlighted important procedural differences in DBS workflow steps, identifying five distinct procedural clusters among respondents [Abosch et al., 2013]. Many centres continue to depend on lower field magnetic resonance imaging (MRI) at 1.5 Tesla as part of the clinical workflow where direct visualization of traditional DBS targets can be challenging. Even where more high field scanners are available, acquired images may be suboptimal due to patient movement, poorly optimized MRI protocols, and patient-specific considerations (e.g. oblique image acquisitions as a result of torticollis).

Template or atlas guidance has been a fundamental part of stereotactic neurosurgery since Jean Talairach described the réperage radiologique to describe the process of establishing the stereotactic positions of neuroanatomical structures by aligning an anatomical atlas, derived from histological sectioning of a single individual, with patient-specific fluoroscopic images [Bancaud et al., 1965, Talairach and Tournoux, 1988]. Semantically, template assistance involves planning using the unprocessed underlying dataset, while atlas guidance involves using the labeled version of the template. While modern imaging has enabled subcortical visualization not possible with classical x-ray based studies, growing evidence points to the need for even more precise targeting of subcortical substructures [Accolla et al., 2016, Lambert et al., 2012, Vanegas-Arroyave et al., 2016]. In this technical report, we describe our initial experiences with using an ultra-high field (UHF) template to assist neurosurgeons with stereotactic planning for conventional deep brain stimulation targets.

### 2.2 Materials and Methods

### 2.2.1 Deep Brain Stimulation Surgery Workflow

In the typical DBS workflow at our centre, the patient is referred to Neurosurgery for DBS implantation by a movement disorders neurologist. After consenting to the procedure, the patient
undergoes clinical (1.5T) MRI for surgical planning in the weeks leading up to the surgery with an 8-channel head/neck/spine (GE Healthcare, Milwaukee, Wisconsin). Sequences acquired include a three dimensional (3D) T1-weighted image using an axial inversion recovery spoiled gradient recalled echo sequence: echo time $(\mathrm{TE})=4.1 \mathrm{~ms}$, inversion time $(\mathrm{TI})=300$ ms , flip angle $=20$ degrees, resolution $=1.25 \times 1.25 \times 1.50 \mathrm{~mm}$, receiver bandwidth $=22.73$ kHz , field of view $(\mathrm{FOV})=26 \times 26 \mathrm{~cm}^{2}$, matrix size $=256 \times 256$. Selected two dimensional (2D) T2-weighted fast spin echo (FSE) sequences in axial and coronal sections are also acquired to better visualize the target region ( $\mathrm{TE}=110 \mathrm{~ms}, \mathrm{TR}=2800 \mathrm{~ms}$, receiver bandwidth $=20.83 \mathrm{kHz}, \mathrm{FOV}=26 \times 26 \mathrm{~cm}^{2}$, matrix size $=256 \times 224$, slice thickness $=1.5 \mathrm{~mm}$, resolution $=1.25 \times 1.25 \times 1.50 \mathrm{~mm})$. After importing the relevant pre-operative images, the anterior and posterior commissures (AC-PC) are identified using the surgical navigation system (StealthStation S7 Framelink software version 5.4.1, Medtronic Inc., Minneapolis, Minnesota). Indirect targeting based on AC-PC coordinates is performed by the neurosurgeon, aided by direct visualization when feasible. However, it is our experience that the clinical MRI protocols at our centre often do not clearly delineate conventional DBS targets. The total surgical planning process of image import, fusion, and planning requires approximately $60-90$ minutes.

The morning of surgery, a Leksell stereotactic frame is secured to the patient's head after a bilateral field block with a $50: 50$ mixture of $0.5 \%$ bupivacaine and $2 \%$ lidocaine, both with epinephrine. A stereotactic computed tomography (CT) scan is acquired with double-dose contrast while the patient is in the headframe. The CT is fused with the 1.5 T MRI using the image guidance system bringing the surgical plan into the reference frame of the patient. The Leksell target and entry point coordinates are extracted from the surgical navigation workstation requiring roughly 15-30 minutes for CT-MRI fusion and final coordinate calculations. The patient is brought to the operating room where the frame is secured to the operating table in the typical manner. The patient is attached to American Society of Anesthesiologists (ASA) standard monitoring, a non-central IV line is inserted, and IV sedation is initiated (dexmedetomidine continuous infusion at $0.4-0.6 \mathrm{mcg} / \mathrm{kg} / \mathrm{hr}$ ). In our experience, dexmedetomidine, contrary to other sedative medications, has almost no respiratory depression effects and minimal interaction with the microelectrode recording.

The surgical field is prepped and draped in the usual manner. The Leksell frame is set to the specified target coordinates, local anaesthetic is infiltrated, the incision opened, and the cortex exposed via a cranial burr hole. A lead fixation device is attached to the skull surrounding the burr hole (Medtronic Stimloc, Medtronic Inc., Minneapolis, Minnesota). The dura is opened and coagulated, as is the overlying brain. Fibrin sealant is introduced to limit brain shift from pneumocephaly. The microdriver system is attached to the frame and, using Ben's gun [Limousin et al., 1995], up to five microelectrodes are slowly advanced towards the target by
a dedicated neurophysiologist while the electrical activity is recorded (Medtronic Leadpoint, Medtronic Inc., Minneapolis, Minnesota). Once the target region has been mapped physiologically, the patient is awoken to test for effects from stimulation. The macroelectrode is inserted along the course of the best candidate microelectrode. The same procedure is repeated on the opposite side, if indicated. Regarding sedation, the dexmedetomidine infusion is stopped 30 minutes before stimulation, and re-instituted once the surgeon and neurophysiologist are satisfied, continuing until the end of the procedure. The lead or leads are tunnelled to the parietal scalp. The frame is removed and the patient is placed in the supine position and general anaesthetic is administered. The DBS electrodes are tunnelled and connected to the implantable pulse generator (IPG). A post-operative 1.5T MRI is completed the day following surgery to confirm placement. Once the appropriate position is confirmed and the patient is mobilizing well, they are discharged home. The patient is followed by a neurologist for optimizing stimulation parameters with stimulation commencing as early as two weeks post-implantation.

### 2.2.2 7T Group Template Creation

While a number of different ultra-high field templates have been proposed in the literature, for the current report, we have elected to use an unbiased group average created at our institution [Wang et al., 2016]. In brief, 12 healthy control subjects (6 female; age: $27.6+/-$ 4.4 years) were scanned on a 7T scanner (Agilent, Santa Clara, CA, USA/Siemens, Erlangen, Germany) using a 24 -channel transmit-receive head coil array constructed in-house with a receiver bandwidth of 50 kHz . A T1-weighted (T1w) MPRAGE sequence was acquired $\left(\mathrm{TR}=8.1 \mathrm{~ms}, \mathrm{TE}=2.8 \mathrm{~ms}, \mathrm{TI}=650 \mathrm{~ms}\right.$, flip angle $=11^{\circ}, 256 \times 512,230$ slices, resolution $=0.59 \times 0.43 \times 0.75 \mathrm{~mm}^{3}$ ). A T2-weighted (T2w) turbo spin-echo (TSE) $3 \mathrm{D}(\mathrm{TR}=3 \mathrm{D}$ sagittal, matrix: $260 \times 366$, 266 slices, resolution $=0.6 \mathrm{~mm}^{3}, 4$ averages). High-resolution in vivo templates were created by performing group-wise linear and nonlinear registration of 12 normal subjects scanned on a human 7T scanner using both T1w and T2w contrasts (available for download at http://www.nitrc.org/projects/deepbrain7t/) resulting in an unbiased group nonlinear T1w average and T2w averages at submillimeter resolution. These templates demonstrate improved visualization of subcortical nuclei compared to lower field templates.

### 2.2.3 Template-to-Patient Registration Workflow

We propose to assist with target selection by fusing the ultra-high field template to the patient reference (T1w) space (Figure 2.1). The procedures are performed using standard-ofcare clinical imaging as described above, but the planning is augmented by the integration of patient-aligned high resolution templates. The patient $1.5 \mathrm{~T} \mathrm{T1w}$ volume, the reference image,


Figure 2.1: Side-by-side visualization of standard clinical 1.5T T1 magnetic resonance imaging (MRI) next to the fused ultra-high field MRI T1 and T2 templates in the axial plane. The red dot marks the location of the globus pallidus internus.
is exported from PACS and converted to NiFTI file format using dcm2niix [Li et al., 2016]. The reference image is first corrected for intensity inhomogeneities [Sled et al., 1998, Tustison et al., 2010], and subsequently, brain masking is performed using the Brain Extraction Toolkit from the FSL package at a fractional intensity threshold of 0.4 [Smith, 2002]. Through a series of successive image registration steps using NiftyReg [Modat et al., 2010] (default settings in version 1.3.9), the UHF T1w and T2w templates are warped into the patient T1w image space. Registration is initiated with a rigid body ( 6 degrees-of-freedom) registration, and is followed by affine ( 12 degrees-of-freedom). Finally, the template is nonlinearly registered to the reference space using the NiftyReg block-matching deformation technique. The duration of image fusion and registration is 20 minutes. All processing was performed on a modern workstation (Intel Core i5-6400 CPU @ 2.70 GHz x 4; 64-bit; 32 Gb of RAM; Ubuntu 16.04 Long Term Support version), and automated using a bash shell script.

The quality of registration is assessed by the treating neurosurgeon via visual inspection of the spatial correspondence between key neuroanatomical features on both images. Once satisfactory, the transformed templates are exported back to DICOM format using NifTI2Dicom. The DICOM IDs of the newly created DICOM files are unified with the patient pre-operative T1w dataset with one modification to the Study Description header using OsiriX. The aligned UHF templates are imported to the StealthStation as separate image datasets. At several stages throughout processing, the images are quality controlled by the clinical team for salient features that correctly identify the sidedness of each hemisphere in comparison with unprocessed
clinical images (i.e. based on cortical and vascular landmarks). The fusion of the CT with the pre-operative MRI was performed on the day of surgery using the commercial neuronavigation software. This was independent of the template fusion process, but ultimately permitted overlaying of UHF templates and the stereotactic CT in the image space of the pre-operative MRI for visualization, resulting in a transformation of the planned trajectory into physical Leksell coordinates.

### 2.3 Results: Two Cases

We demonstrate the utility of UHF template-assisted stereotactic targeting in two cases: unilateral globus pallidus internus implantation and bilateral subthalamic nucleus implantation.

### 2.3.1 Right GPi Implantation for Dystonia

The first patient was a 51 year-old otherwise healthy right-handed female with severe progressive left-sided dystonia and supranuclear palsy of unknown etiology. Her painful dystonia progressed to the point of complete loss of left upper extremity function requiring an arm sling and she suffered from recurrent severe left shoulder dislocations, despite trials of systemic and local baclofen therapy and a failed orthopedic intervention to correct her dislocation. She opted to undergo implantation of her right internal pallidum to improve her mostly unilateral (left-sided) symptoms.

She underwent a clinical pre-operative MRI scan several weeks prior to her procedure (Figure 2.2a). The basal ganglia structures were not well visualized, made more challenging by her left-sided torticollis (head rotated in the scanner). We employed template-to-patient registration bringing our high resolution T 1 and T 2 templates into the patient space (Figure 2.2d and 2.2e). Quality of registration was assessed manually by the neurosurgeon but was noticeably improved compared to the built-in Schaltenbrand-Wahren atlas provided as part of the commercial software package (Figure 2.2b). Target selection was performed using a combination of indirect (AC-PC coordinate based) and direct targeting techniques, and furthermore assisted by the inclusion of the aligned 7T templates. The final trajectory was decided after ensuring that the trajectory choice using conventional clinical imaging appeared appropriate. The stereotactic CT was registered to the pre-operative patient MRI on the day of surgery using the commercial neuronavigation system (Figure 2.2c). Three microelectrodes were used for recording. Unit potentials were obtained at the expected depth with the central electrode trajectory being chosen after demonstrating the fewest side effects. A 3387 electrode (Medtronic Inc., Minneapolis, Minnesota) was then implanted at the target site (Figure 2.2f). Figure 2.3


Figure 2.2: Right internal pallidum implantation assisted by ultra-high field (UHF) fusion to standard clinical images ( 1.5 T magnetic resonance imaging [MRI] and computed tomography [CT]). The red dot marks the same location fused between modalities for (A) 1.5T MRI with gadolinium, (B) with the Schaltenbrand-Wahren atlas overlay provided in the clinical neuronavigation software, (C) CT in Leksell frame, (D) UHF T1 average, and (E) UHF T2 average. Also included is the best corresponding coronal section in the post-insertion MRI (F).
shows a screenshot of the UHF T2 average integrated into the commercial neuronavigation software.

The patient noted an immediate improvement in her dystonia after surgery. Post-op imaging revealed that the electrode was in the appropriate position with no complications. At four months follow-up, she no longer required an arm sling, and demonstrated significant improvement in her pain and her dystonia with the following stimulation settings: effective contact $=$ 2 , amplitude $=3.5 \mathrm{~V}$, pulse width $=60$ us, frequency $=130 \mathrm{~Hz}$.

### 2.3.2 Bilateral STN Implantation for Parkinson's Disease

The second patient was a 66 year-old left-handed male with a 14-year history of Parkinson's disease. He was referred as a surgical candidate due to significant bradykinesia, dyskinesias, and on/off fluctuations. He consented to bilateral subthalamic nucleus implantation.


Figure 2.3: Example screenshot demonstrating the integration of the 7T T2 average into the commercial surgical planning software (Case 1: right internal pallidum insertion). The red dot marks the location of the globus pallidus internus.

Preoperative imaging again revealed poor visualization of subcortical structures including the STN (Figure 2.4a). Template-to-patient registration was performed (Figure 2.4c and d), and compared against the standard atlas integrated into the neuronavigation software (Figure 2.4 b ). Target selection was performed using a combination of indirect (AC-PC coordinate based) and direct targeting techniques with assistance from the 7T templates. For each side, all five microelectrodes were used for recording. Unit recordings were best identified along lateral and anterior trajectories. The left anterior trajectory was chosen due to no side effects except for mild right hemi-body symptoms at high threshold. On the right, good unit activity was observed in the central, anterior, and medial recordings. The medial electrode demonstrated the best side effect profile. Bilateral Medtronic 3389 leads were implanted (Figure 2.4e and 2.4f).

Postoperatively, this patient noted immediate improvement, likely due to an insertional effect. However, this effect dissipated within two weeks, at which point stimulation was initi-


Figure 2.4: Bilateral subthalamic nucleus implantation assisted by ultra-high field (UHF) fusion to standard clinical images. The left and right trajectories (yellow and green, respectively) are fused across modalities for (A) 1.5 T magnetic resonance imaging (MRI) with gadolinium, (B) with the Schaltenbrand-Wahren atlas overlay, (C) UHF T1 average, and (D) UHF T2 average. Bilateral subthalamic nuclei were implanted successfully, as demonstrated with select ( E ) coronal and ( F ) axial views on postoperative MRI.
ated. At three months post-op, he has halved his levocarbidopa dosage and no longer has any off periods. He is now independent in all activities of daily living. His stimulation settings are the following: L STN bipolar stimulation with effective contacts $=1$ positive and 2 negative, amplitude $=3 \mathrm{~V}$, pulse width $=90$ us, frequency $=130 \mathrm{~Hz} ;$ R STN bipolar stimulation with effective contacts $=9$ positive and 10 negative, amplitude $=3 \mathrm{~V}$, pulse width $=90$ us, frequency $=130 \mathrm{~Hz}$.

### 2.4 Discussion

In this technical report, we have described a workflow for the integration of high-resolution in vivo ultra-high field templates into the surgical navigation system to assist with DBS planning. We have demonstrated that UHF assistance can be helpful for internal pallidum and subthalamic nucleus implantations.

Ultra-high field MRI delivers an over two-fold improvement in signal-to-noise (SNR) compared with standard clinical scanners, allowing in vivo visualization at submillimeter spatial resolution. The improved resolution has enabled clear delineation of small structures like the subthalamic nucleus and pallidum, overall enriching 3D context in the subcortex invaluable for surgical targeting. To date, patient-specific UHF imaging has been limited to research protocols and has not been used directly for surgical planning. While interest in UHF imaging is growing, availability remains limited. By using an unbiased UHF template, our workflow does not require the candidate DBS patient to undergo any additional scans beyond what is conventionally performed at a given institution, and could be used by centers without access to UHF imaging.

Numerous alternative templates exist both with and without atlas labeling. Histological atlases, while capable of providing microscopic detail, may be more difficult to accurately register with clinical images due to differences in scale and contrast, as well as the challenge of spatial correspondences between the histological atlas (derived from a single individual) with the patient. Atlases created at low field are abundant and typically involve larger datasets (e.g. the MNI152 atlas which has 152 subjects). However, the poorer spatial resolution limits visualization of subcortical targets like the STN. Our initial experiences suggest that UHF in vivo templates may serve as an alternative option enabling improved signal and resolution but also improved registration with patient datasets since context regarding expected location of subcortical structures is maintained at a group level. This report has focused on the use of T1 and T2 UHF templates created from a dataset of young healthy volunteers at our centre. This could be considered a limitation of the current work since they may not best represent our patient groups, both in terms of age and pathology. Certain subcortical structures are displaced and atrophic in Parkinson's compared to normal age-matched controls [Xiao et al., 2014], suggesting that a disease-specific atlas could be ideal for DBS planning. One practical problem with such a choice is that it remains unclear how optimal a Parkinson's specific atlas would be for other patient populations encountered in a DBS practice, for example in dystonia or essential tremor. Finally, there is growing evidence that there are separate subtypes of Parkinson's, some of which could be more responsive to DBS than others, suggesting that ultimately subtype specific templates may be more representative. Overall, the choice of atlas is a complex decision, and is an area of further study. Eventually, patient-specific ultra-high field imaging may be feasible. However, calibration and correction of distortion remain challenges [Duchin et al., 2012, Lau et al., 2018a] (also see Chapter 4).

As is routine in clinical practice, the neurosurgeon must be rigorous in establishing the correspondence between any fused images (e.g. CT and MRI), which is performed by manual inspection on the surgical navigation system. When using the proposed workflow, the same level
of rigor is necessary for assuring adequate alignment of the template to the patient dataset. Design and implementation requires a close working relationship with biomedical engineers and medical physicists to debug any software problems and develop safeguards to minimize errors of spatial alignment and data conversion, including DICOM import and export. Our 7T template-assisted workflow serves as an enhancement of the conventional workflow of MRI to CT fusion. The surgeon still performs the target planning using clinical images, but can choose to augment or confirm their targeting with template-assistance by windowing between the template and clinical scan within the neuronavigation software. It should be emphasized that the UHF template is helpful for target placement but obviously not for entry point localization since patient-specific gyral patterns and cortical vessel anatomy would not be accounted for.

Our proposed image fusion workflow could be better optimized through more systematic evaluation of the impact of different pipeline modifications. For example, in our experience, a coarse-to-fine registration approach results in better registration of our template to a patient target image due to better initialization of global image correspondence prior to nonlinear registration. While we have not assessed this systematically, the additional time required for each of the linear (rigid-body and affine) registration components is minimal (less that 1 minute). Ongoing goals in this collaborative project with biomedical engineers include more robust evaluation of the impact of brain extraction and nonlinear fusion methods on speed and accuracy. Overall, we feel that the time required for our current workflow ( 20 minutes) is reasonable given that DBS cases are performed on an elective basis.

### 2.5 Conclusions

We have described a technique for integrating an ultra-high field MRI template into the surgical planning workflow that may serve as a valuable adjunct to standard clinical imaging for stereotactic target selection. The method does not require any additional cost or time to the patient. Prospective studies would help to identify ideal template selection and how this information is best used along with high-quality patient-specific pre-operative imaging.

## Chapter 3

## A framework for evaluating correspondence between brain images using anatomical fiducials

This chapter is based on the following manuscript:

- Lau, J. C., Parrent, A. G., Demarco, J., Gupta, G., Park, P. J., Ferko, K., Khan, A. R., \& Peters, T. M. (2019). A framework for evaluating correspondence between brain images using anatomical fiducials. Human Brain Mapping, Under Review.


### 3.1 Introduction

Establishing spatial correspondence between images is a crucial step in neuroimaging studies enabling fusion of multimodal information, analysis of focal morphological differences, and comparison of within- and between-study data in a common coordinate space. Stereotaxy arose as a result of questions raised by scientists and surgeons interested in the physiology and treatment of focal brain structures [Evans et al., 2012, Horsley and Clarke, 1908, Peters, 2006]. Jean Talairach played a crucial role, observing consistent anatomical features on lateral pneumoencephalograms [Dandy, 1918], or "air studies", that could be consistently localized, specifically the anterior commissure (AC) and posterior commissure (PC) [Schaltenbrand and Wahren, 1977, Talairach et al., 1957], and could thus be mapped to prepared post-mortem brain sections in a 3D coordinate system. The AC-PC line has remained important in the era since magnetic resonance imaging (MRI) has risen to prominence for aligning brain images to create population atlases [Collins et al., 1994, Evans et al., 1992, Talairach and Tournoux, 1988] as well as to project data from structural and functional investigations. Further optimizations enabled by deformable registration have led to atlas enhancements [Fonov et al., 2011] where many more structural features are preserved. The adoption of standard templates has allowed researchers to compile cytoarchitectonic, functional, and structural data across studies via image-based meta-analysis of peak coordinates and statistical maps [Eickhoff et al., 2009, Gorgolewski et al., 2015, Yarkoni et al., 2011].

Ever since the first linearly aligned population templates [Evans et al., 1992, Talairach and Tournoux, 1988], there have been a number of advances in the development of robust higher order nonlinear registration tools. As the options became more numerous, several studies investigated the performance of the different nonlinear registration algorithms [Chakravarty et al., 2009, Evans et al., 2012, Hellier et al., 2003, Klein et al., 2009]. Over the past decade, the most common metrics used to evaluate spatial correspondence are related to voxel overlap between regions-of-interest (ROIs) segmented in both reference and target images. Typically, large subcortical structures well-visualized on standard structural MRIs such as the globus pallidus (pallidum), striatum, and thalamus are used [Chakravarty et al., 2009, Chakravarty et al., 2008, Klein et al., 2009]. While these measures are effective for evaluating spatial correspondence on the macroscale, here we argue that they remain relatively coarse measures of
registration quality and are insensitive to focal misregistration between images. In addition, they do not permit facile identification or description of where these local biases are occurring. These issues are particularly critical as technical advancements in both imaging and stereotaxy are enabling more accurate therapeutic modulation of brain regions where several millimeters could represent the difference between optimal therapy and complications.

In this paper, we sought inspiration from classical stereotactic methods [Schaltenbrand and Wahren, 1977, Talairach et al., 1957], and propose that point-based distances provide a more sensitive metric by which brain image correspondence can be evaluated. Anatomical points have been referred to in the literature using a variety of terms including fiducials, landmarks, markups (sometimes used in combination) but ultimately involve representing an anatomical feature by a three-dimensional ( $\mathrm{x}, \mathrm{y}, \mathrm{z}$ ) Cartesian coordinate. For this manuscript, we have chosen to use the term AFIDs, short for anatomical fiducials, "fiducia" being Latin for trust or confidence. We argue that the advent of automatic segmentation-based methods has led to a relative underemphasis of point correspondence between brain structures. We first sought to determine whether we could define a set of AFIDs that were both consistently identifiable across multiple datasets while also providing a distributed sampling about the brain. Following this, we demonstrate how AFIDs are complementary to segmentation-based metrics for providing a quantitative report of spatial correspondence between structural magnetic resonance images of the brain using more intuitive distance-based measures of alignment. Central to this work was the development of our protocol using an open source framework, enabling reproducibility across sites and centers. The overall study organization is shown schematically in Figure 3.1.

### 3.2 Methods

### 3.2.1 Protocol development

A series of anatomical fiducials (AFIDs) were identified by the lead author (JCL; 10 years experience in neuroanatomy) in consultation with an experienced neurosurgeon (AGP; 20+ years experience practicing stereotactic and functional neurosurgery) with consensus achieved on a set of 32 points, which we refer to as AFID32 (see Figure 3.2; RRID:SCR_016623). AFIDs could generally be classified as midline $(10 / 32=31.25 \%)$ or lateral $(22 / 32$; i.e. 11 structures that could be placed on each of the left and right sides). Regions prone to geometric distortion were avoided [Lau et al., 2018b]. We limited our initial set of AFID locations to deep brain regions where less inter-subject variability exists (millimeter scale) compared to the cortical sulci and gyri (centimeter scale) [Thompson et al., 1996].

## Evaluating correspondence between brain images



Figure 3.1: Metrics for evaluating spatial correspondence between brain images include voxel overlap (i.e. ROI-based) metrics as well as point-based distance metrics. The proposed framework involves the identification of point-based anatomical fiducials (AFIDs) in a series of brain images, which provide an intuitive millimetric estimate of correspondence error between images and is also a useful tool for teaching neuroanatomy.

The AFID points were placed using the Markups Module of 3D Slicer version 4.6.2 [Fedorov et al., 2012] (RRID:SCR_005619). One key feature of 3D Slicer is that it allows markup points to be placed in the 3D coordinate system of the software as opposed to the voxel coordinate system of the image being annotated permitting more refined (sub-voxel) localization. Images are automatically linearly interpolated by the software on zoom. After importing the structural MRI scan to be annotated into 3D Slicer, the anterior commissure (AC) and posterior commissure (PC) points were placed, specifically the center of each commissure rather than the intraventricular edge. After defining an additional midline point (typically the pontomesencephalic junction or intermamillary sulcus), an AC-PC transformation was performed using the built-in Slicer module (AC-PC Transform). For all subsequent AFID placements, the AC-PC aligned image was used. The AFID32 protocol is shown in MNI2009bAsym space in Figure 3.2.

The rest of the methods are organized into four separate phases. Phase 1 involved AFID32 placement in three open access brain templates. Phase 2 involved further placement of the AFIDs in individual subject scans. In Phase 3, AFIDs were used to evaluate subject-to-template registration; and finally, in Phase 4, they were used to assess template-to-template registration quality.

For validation and assessment, we adopted the terminology of Fitzpatrick and colleagues [Fitzpatrick and West, 2001, Fitzpatrick et al., 1998] who defined fiducial localization error (FLE) and fiducial registration error (FRE) as metrics used to evaluate the real-world accuracy of image-guidance systems used in neurosurgery. FLE is defined as error related to the
placement (i.e. localization) of fiducials, while FRE is defined as error related to registration. This body of work has been most concerned with describing the correspondence between preoperative images of a patient and the physical location of the patient and surgical site in the operating room. Here, we use these terms to describe (virtual, image-based) anatomical fiducials (AFIDs) annotated in structural T1-weighted MRI scans.

### 3.2.2 Phase 1: Protocol validation for brain templates

Novice participants $(\mathrm{N}=8)$ were trained over a series of neuroanatomy tutorials to place AFIDs on a number of publicly available brain images: Agile12v2016 [Lau et al., 2017, Wang et al., 2016], Colin 27 [Holmes et al., 1998], MNI2009bAsym (nonlinear asymmetric; version 2009b; RRID:SCR_008796) [Fonov et al., 2011]. Each participant then performed 4 rating sessions independently for each template, for a total of 12 point sets resulting in a total of 96 AFID32 protocols. We computed several different metrics for describing the accuracy (and reliability) of our proposed protocol, all of which are variations of anatomical fiducial localization error (AFLE): mean AFLE, intra-rater AFLE, and inter-rater AFLE as shown in Figure 3.3.

To compute the mean AFLE, the mean AFID coordinate for each brain image was used as an approximation of the ideal coordinate location. Mean AFLE was calculated as the Euclidean distance between the individual position and the group mean. We furthermore calculated intrarater AFLE as the mean pairwise distance between AFIDs placed by the same rater. The individual measures were averaged across all raters as a summary metric. To calculate interrater AFLE, a mean coordinate was computed by averaging the coordinates for each rater as an estimate of the ideal coordinate location for the rater; the mean pairwise distance between AFIDs placed across raters was then calculated as a summary metric. We summarized global and location-specific mean AFLE according to a number of variables: template (group versus individual), rating session (1-4), rater, and AFID.

Time required to complete AFID32 placement for a single MRI was documented by each rater. Outliers were defined as any fiducials deviating from the mean fiducial point by greater than 10 mm . Furthermore, patterns of variability in AFID placement were assessed using Kmeans clustering of fiducial locations (point clouds) relative to the mean fiducial location.

### 3.2.3 Phase 2: Protocol validation for individual subjects

The same participants and the lead author (total $\mathrm{N}=9$ ) performed additional AFID placement on a series of 30 independent brain images from the OASIS-1 database [Marcus et al., 2010] (RRID:SCR_007385). Subjects from the OASIS-1 database were selected from the broad range of ages encountered in the database, restricted to cognitively intact (MMSE 30) participants.


Figure 3.2: Each anatomical fiducial in the full AFID32 protocol is demonstrated with crosshairs at the representative location in MNI2009bAsym space using the standard cardinal planes. AC = anterior commissure; $\mathrm{PC}=$ posterior commissure; AL = anterolateral; $\mathrm{AM}=$ anteromedial; $\mathrm{IG}=$ indusium griseum; $\mathrm{IPF}=$ interpeduncular fossa; $\mathrm{LMS}=$ lateral mesencephalic sulcus; $\mathrm{LV}=$ lateral ventricle; PMJ $=$ pontomesenphalic junction.


Figure 3.3: Metrics used for validating AFID placements are shown here in schematic form. Mean, intra-rater, and inter-rater AFLE can be computed for an image that has been rated by multiple raters multiple times.

Although we controlled for normal cognition by MMSE, we selected for qualitatively challenging images with more complex anatomy (asymmetric anatomy and/or variably-sized ventricles). Details on the 30 scans are provided in Appendix Section A. 2 and organized into the Brain Imaging Data Structure (BIDS) format [Gorgolewski et al., 2016] (RRID:SCR_016124).

Each of the 9 participants placed 10 independent AFID32 protocols for a total of 90 AFID32 protocols and 2880 individual points. Each of the 30 MRI scans from the OASIS-1 database had AFIDs placed by 3 raters to establish inter-rater AFLE (as described in Methods Section Phase 1: Protocol Validation for Brain Templates). Intra-rater AFLE was not evaluated in Phase 2. Quality of rigid registration was visually inspected by an experienced rater (JL).

## Region-of-interest segmentation

BIDS formatting permitted automatic processing of each included OASIS-1 subject using fMRIPrep version 1.1.1 [Esteban et al., 2018, Gorgolewski et al., 2016] (RRID:SCR_016216) with anatomical image processing only. Briefly, the fMRIPrep pipeline involves linear and deformable registration to the MNI2009cAsym template [Avants et al., 2008, Fonov et al., 2011] then processing of the structural MRI through Freesurfer for cortical surface and subcortical volumetric labeling [Dale et al., 1999, Fischl, 2012] (RRID:SCR_001847). We focused on using ROIs commonly used in the literature to evaluate quality of registration in the subcortex [Chakravarty et al., 2009, Hellier et al., 2003, Klein et al., 2009], i.e. the pallidum, striatum, and thalamus provided as part of the fMRIPrep output run through FreeSurfer. The striatum label required combining the ipsilateral caudate nucleus, accumbens, and putamen labels.

### 3.2.4 Phase 3: Evaluating subject-to-template registration

We evaluated the quality of subject-to-template registration using the output provided as part of fMRIPrep version 1.1.1 using conventional ROI-based metrics (i.e. voxel overlap) as well
as distance metrics derived from our manual AFID32 annotations from Phases 1 and 2. The default template for fMRIPrep 1.1.1 was the MNI2009cAsym template. We started by visually inspecting the images qualitatively from the output fMRIPrep html pages. For each individual subject scan, we used the mean fiducial location as the optimal location calculated in Phase 2. The distance between the individual subject AFID location and the corresponding mean AFID location in the template was computed and defined as the anatomical fiducial registration error (AFRE) and computed for linear transformation alone (lin) and combined linear and nonlinear transformation (nlin). Our definition of AFRE differs from the FRE used by Fitzpatrick whose framework for neuronavigation was necessarily limited to rigid-body transformations [Fitzpatrick et al., 1998]. This was compared with ROI-based measures of spatial correspondence, specifically, the Jaccard similarity coefficient $((A \cap B) /(A \cup B))$ and the Dice kappa coefficient $((2 \times A \cap B) /(A+B))$, where A and B are the number of voxels in the source and reference images, respectively.

We were able to use the AFID32 points placed in Phase 1 for the MNI2009bAsym template since the only difference between the MNI2009bAsym and MNI2009cAsym templates was the resampling from 0.5 mm to 1 mm isotropic resolution. AFRE was computed for each AFID location and OASIS-1 subject, along with voxel overlap for the pallidum, striatum, and thalamus. Comparisons between AFRE and voxel overlap were made using Kendall's tau.

### 3.2.5 Phase 4: Evaluating template-to-template registration

BigBrain is a publicly available ultrahigh-resolution ( 20 micron) human brain model that has enabled bridging of macroscale anatomy with near cellular anatomy [Amunts et al., 2013] (RRID:SCR_001593). A deformable mapping provided by the MNI group has permitted the exploration of high-resolution BigBrain neuroanatomy in MNI2009bSym space (BigBrainRelease.2015; Last modified August 21, 2016; accessed August 2, 2018; Available at: ftp:// bigbrain.loris.ca/BigBrainRelease.2015/3D_Volumes/MNI-ICBM152_Space/). In this manuscript, we refer to the registered BigBrain image as BigBrainSym. We quantify the spatial correspondence between BigBrainSym and MNI2009bSym as well as BigBrainSym and MNI2009bAsym templates using the AFID32 protocol to determine whether any significant AFRE could be identified. For MNI2009bAsym, we used mean coordinates for each AFID using rater data from Phase 1. BigBrainSym and MNI2009bSym templates were annotated de novo by three experienced raters (GG, JL, KF). The mean AFID coordinate was used as an approximation of the ideal coordinate location for each template. Spatial correspondence was estimated as the AFRE (i.e. Euclidean distance between points) for each AFID. Correlation between AFLE and AFRE were assessed using Kendall's tau.

Table 3.1: Summary of fiducial localization error across brain templates.

|  | Before QC |  |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Template | mean AFLE (mm) | \# of outliers (\%) | mean AFLE (mm) | \# of outliers (\%) | After QC |  |
| intra-rater AFLE (mm) | inter-rater AFLE (mm) |  |  |  |  |  |
| Agile12v2016 | $1.10 \pm 1.59$ | $3 / 1024(0.29 \%)$ | $1.01 \pm 0.93$ | $0 / 1021(0.00 \%)$ | $1.13 \pm 0.86$ | $1.14 \pm 0.48$ |
| Colin27 | $1.71 \pm 2.78$ | $20 / 1024(1.95 \%)$ | $1.11 \pm 1.05$ | $1 / 1004(0.10 \%)$ | $1.14 \pm 0.92$ | $1.36 \pm 0.88$ |
| MNI2009bAsym | $0.99 \pm 1.11$ | $1 / 1024(0.10 \%)$ | $0.97 \pm 0.80$ | $0 / 1023(0.00 \%)$ | $1.03 \pm 0.78$ | $1.07 \pm 0.46$ |
| Total | $1.27 \pm 1.98$ | $24 / 3072(0.78 \%)$ | $1.03 \pm 0.94$ | $1 / 3048(0.03 \%)$ | $1.10 \pm 0.86$ | $1.19 \pm 0.64$ |

### 3.2.6 Source code and data availability

All data analysis was performed using R-project version 3.5.1. The AFIDs protocol, raw and processed data, processing scripts, and scripts used in this manuscript are available at: https: //github.com/afids.

### 3.3 Results

### 3.3.1 Phase 1: Protocol validation for brain templates

The 8 raters had a mean experience of $11.5 \pm 11.2$ months in medical imaging (range: 0-24 months), $14.3 \pm 17.0$ months in neuroanatomy (range: $0-48$ months), and $7.0 \pm 8.8$ months in 3D Slicer (range: 0-24 months). During the template validation phase, the raters placed a total of 3072 individual points (number of sessions $=4$; templates $=3$; points $=32$ ). Average AFID32 placement time was estimated at between 20-40 minutes. Thus, a total of 1920-3840 minutes (or 32-64 hours) were logged in this phase of the study. The mean, intra-rater, and inter-rater AFLE metrics are summarized in Table 3.1.

For the raw data, the mean AFLE was $1.27 \pm 1.98 \mathrm{~mm}(1.10 \pm 1.59 \mathrm{~mm}$ for Agile12v2016; $1.71 \pm 2.78 \mathrm{~mm}$ for Colin27; $0.99 \pm 1.11 \mathrm{~mm}$ for MNI2009bAsym). Using a threshold of mean AFLE greater than 10 mm from the group mean, we identified 24 outliers out of 3072 independent points ( $0.78 \%$ ). 20/24 ( $83.33 \%$ ) of outliers were the result of variable placement in the bilateral ventral occipital horns (i.e. AFID29 and AFID30) of the Colin27 template. One pair ( $2 / 24 ; 8.33 \%$ ) of outliers was due to left-right mislabeling (indusium griseum; AFID27 and AFID28). One additional point was mislabeled; i.e. the left anterolateral temporal horn point (AFID22) was placed at the left inferior anteromedial horn location (AFID26). After quality control (QC) and filtering outliers, mean AFLE improved to $1.03 \pm 0.94 \mathrm{~mm}(1.01 \pm 0.93 \mathrm{~mm}$ for Agile12v2016; $1.11 \pm 1.05 \mathrm{~mm}$ for Colin27; $0.97 \pm 0.80 \mathrm{~mm}$ for MNI2009bAsym).

Intra-rater AFLE was $1.10 \pm 0.86 \mathrm{~mm}(1.13 \pm 0.86 \mathrm{~mm}$ for Agile12v2016; $1.14 \pm 0.92 \mathrm{~mm}$ for Colin27; $1.03 \pm 0.78 \mathrm{~mm}$ ); and inter-rater AFLE was $1.19 \pm 0.65 \mathrm{~mm}(1.15 \pm 0.49 \mathrm{~mm}$ for Agile12v2016; $1.36 \pm 0.88 \mathrm{~mm}$ for Colin27; $1.07 \pm 0.46 \mathrm{~mm}$ for MNI2009bAsym). Mean, intra-rater, and inter-rater AFLE for each AFID post-QC are summarized in Appendix Section

## A.1.

All subsequent analyses were performed using the mean AFLE metric. We performed a one-way analysis of variance observing evidence of statistically different variance between templates ( F -value $=7.88 ; \mathrm{p}$-value $<0.001$ ). Differences in mean AFLE between templates were identified on subgroup analysis for the right superior lateral mesencephalic sulcus (AFID06), culmen (AFID10), genu of the corpus callosum (AFID19), and left superior anteromedial temporal horn (AFID24), suggesting differences between templates that may contribute to errors in placement. The results for each AFID are also summarized in the Appendix Section A.1.

Furthermore, we observed several distinct patterns of AFID placement using K-means clustering of fiducial locations (point clouds) relative to the mean fiducial location (see Figure 3.4). We identified three different general patterns of point cloud distributions ranging from highly anisotropic to moderately anisotropic to isotropic.

K-means clustering


17


Cluster 2



Principal Componen

Figure 3.4: K-means clustering of point clouds relative to the mean fiducial location for each of the 32 AFIDs (left). Principle components analysis (bottom right) revealed three different general patterns were identified ranging from highly isotropic (Cluster 1: red) to moderately anisotropic (Cluster 2: blue) to anisotropic (Cluster 3: green). Results are shown for the MNI2009bAsym template. See Appendix Section A. 1 for similar plots for Agile12v2016, Colin27, and the templates combined.

As a secondary analysis, we explored whether any evidence of learning over the 4 independent rating sessions could be identified (Appendix Section A.1). Using linear modeling, we identified a general decrease in mean AFLE with increasing session number although this did not meet thresholds of statistical significance (estimate $=-0.02 \mathrm{~mm} / \mathrm{session} ; \mathrm{p}$-value $=0.11$ ).

These trends were explored on the individual rater level. For two out of 8 raters, AFLE varied with session number. Rater04 demonstrated a general linear improvement of $-0.17 \mathrm{~mm} /$ session from an initial mean AFLE of 1.64 mm (i.e. the worst performing initial session); however Rater02 worsened at a rate of $0.12 \mathrm{~mm} /$ session from an initial mean AFLE of 0.59 mm (i.e. the best performing initial session). No significant effect with individual AFIDs was identified. All subgroup analyses were multiple comparisons corrected using FDR ( $q$-value $<0.05$ ).

### 3.3.2 Phase 2: Protocol validation for individual subjects

During the individual subject validation phase, 9 participants completed 10 AFID protocols (= 90 total protocols) and a total of 2880 individual points distributed equally among 30 OASIS-1 datasets. We identified 28 outliers $(0.97 \%)$, defined as individual point placements greater than $1 \mathrm{~cm}(10 \mathrm{~mm})$ away from the group mean. 8/28 outliers ( $28.57 \%$ ) were the result of mislabeled points: three pairs of lateral (non-midline) AFIDs and only one pair due to gross mislabeling of the target AFID structure (placement in bilateral frontal ventricular horns rather than occipital horns). Beyond left-right swapping, the AFIDs most susceptible to outliers were the following points: bilateral ventral occipital horns (AFID29-30) and bilateral indusium griseum origins (AFID27-28). Mean AFLE across the 30 scans and points was $1.28 \pm 3.03 \mathrm{~mm}$ improving to $0.94 \pm 0.73$ after filtering out the outliers. Inter-rater AFLE was $1.58 \pm 1.02 \mathrm{~mm}$ across all AFIDs. Mean AFLE and inter-rater AFLE are summarized for each AFID in Table 3.2 and subject in Appendix Section A.2. FMRIPrep ran successfully on 30/30 datasets (100.0\%).

### 3.3.3 Phase 3: Evaluating subject-to-template registration

The following section uses the AFIDs to evaluate the quality of spatial correspondence between the Phase 2 subject data with the MNI2009cAsym template as processed through fMRIPrep. Visual inspection of the fMRIPrep generated reports revealed no gross misregistrations between MNI2009c and the individual subject scans although a pattern of worse deformable registration in subjects with enlarged ventricles was observed. The rest of this section is concerned with examining the comparative utility of conventional voxel overlap (ROI-based) metrics against the point-based (AFRE) metric proposed in this study (see Figure 3.5a).

Improvements in overlap were identified when going from linear to combined (linear and nonlinear) transformations (Table 3.3). Some heterogeneity in values was noted between ROIs with voxel overlap measures observed to be lowest for the pallidum (the smallest structure evaluated). All Jaccard values after nonlinear transformation were greater than 0.7 (greater than 0.8 for Dice kappa), generally considered to represent good correspondence between two registered images. For simplicity, we report the Jaccard coefficient as our measure of voxel

Table 3.2: Mean and inter-rater fiducial localization error pre- and post-QC for the included OASIS-1 subjects for all AFIDs.

| AFID | Description | Before QC | After QC |  |
| :---: | :---: | :---: | :---: | :---: |
|  |  | Mean AFLE | Mean AFLE | Inter-Rater AFLE |
| 01 | AC | $0.36 \pm 0.21$ (1.29) | $0.36 \pm 0.21$ (1.29) | $0.60 \pm 0.25$ (1.38) |
| 02 | PC | $0.34 \pm 0.16$ (0.88) | $0.34 \pm 0.16$ (0.88) | $0.57 \pm 0.21$ (1.22) |
| 03 | infracollicular sulcus | $0.78 \pm 0.48$ (3.07) | $0.78 \pm 0.48$ (3.07) | $1.34 \pm 0.64$ (3.84) |
| 04 | PMJ | $0.83 \pm 0.49$ (2.44) | $0.83 \pm 0.49$ (2.44) | $1.41 \pm 0.55$ (2.55) |
| 05 | superior interpeduncular fossa | $1.20 \pm 0.75$ (3.50) | $1.20 \pm 0.75$ (3.50) | $2.04 \pm 0.90$ (4.25) |
| 06 | R superior LMS | $1.30 \pm 1.74$ (14.25) | $1.01 \pm 0.55$ (2.85) | $1.70 \pm 0.68$ (3.13) |
| 07 | L superior LMS | $1.36 \pm 1.71$ (13.99) | $1.06 \pm 0.61$ (3.45) | $1.72 \pm 0.71$ |
| 08 | R inferior LMS | $1.13 \pm 0.75$ (5.13) | $1.03 \pm 0.57$ (2.99) | $1.77 \pm 0.74$ |
| 09 | L inferior LMS | $1.10 \pm 0.80$ (5.31) | $1.01 \pm 0.62$ (2.72) | $1.71 \pm 0.86$ |
| 10 | culmen | $0.99 \pm 0.99$ (5.66) | $0.83 \pm 0.62$ (3.07) | $1.35 \pm 0.82$ |
| 11 | intermammillary sulcus | $0.60 \pm 0.31$ (1.62) | $0.60 \pm 0.31$ (1.62) | $1.02 \pm 0.41$ (1.86) |
| 12 | R MB | $0.40 \pm 0.23$ (1.11) | $0.40 \pm 0.23$ (1.11) | $0.69 \pm 0.32$ (1.52) |
| 13 | L MB | $0.36 \pm 0.20$ (1.20) | $0.36 \pm 0.20$ (1.20) | $0.62 \pm 0.29$ (1.62) |
| 14 | pineal gland | $0.68 \pm 0.47$ (1.98) | $0.68 \pm 0.47$ (1.98) | $1.16 \pm 0.69$ (2.63) |
| 15 | R LV at AC | $1.00 \pm 0.90$ (5.28) | $0.91 \pm 0.72$ (4.45) | $1.55 \pm 1.08$ (5.86) |
| 16 | L LV at AC | $1.01 \pm 0.80$ (4.53) | $0.94 \pm 0.70$ (4.53) | $1.60 \pm 1.08$ (5.47) |
| 17 | R LV at PC | $0.92 \pm 0.54$ (3.42) | $0.92 \pm 0.54$ (3.42) | $1.54 \pm 0.77$ (3.84) |
| 18 | L LV at PC | $0.87 \pm 0.42$ (2.20) | $0.87 \pm 0.42$ (2.20) | $1.46 \pm 0.55$ (2.80) |
| 19 | genu of CC | $0.97 \pm 0.81$ (5.16) | $0.89 \pm 0.63$ (3.69) | $1.50 \pm 0.89$ (4.30) |
| 20 | splenium | $0.54 \pm 0.25$ (1.24) | $0.54 \pm 0.25$ (1.24) | $0.91 \pm 0.35$ (1.66) |
| 21 | R AL temporal horn | $1.44 \pm 1.09$ (7.01) | $1.30 \pm 0.86$ (4.45) | $2.21 \pm 1.13$ (5.92) |
| 22 | L AL temporal horn | $1.22 \pm 0.77$ (4.11) | $1.22 \pm 0.77$ (4.11) | $2.04 \pm 1.01$ (4.47) |
| 23 | R superior AM temporal horn | $1.28 \pm 1.27$ (8.22) | $1.12 \pm 0.88$ (4.69) | $1.86 \pm 1.19$ (4.97) |
| 24 | L superior AM temporal horn | $1.09 \pm 1.22$ (7.54) | $0.83 \pm 0.61$ (3.66) | $1.39 \pm 0.85$ (4.60) |
| 25 | R inferior AM temporal horn | $1.69 \pm 1.43$ (9.03) | $1.44 \pm 0.91$ (4.72) | $2.39 \pm 1.23$ (5.07) |
| 26 | L inferior AM temporal horn | $1.99 \pm 1.75$ (8.79) | $1.49 \pm 1.09$ (4.70) | $2.42 \pm 1.47$ (6.64) |
| 27 | R indusium griseum origin | $3.13 \pm 4.19$ (23.44) | $1.77 \pm 0.99$ (4.77) | $2.95 \pm 1.20$ (5.75) |
| 28 | L indusium griseum origin | $2.99 \pm 4.30$ (24.30) | $1.68 \pm 1.00$ (5.00) | $2.75 \pm 1.29$ (5.78) |
| 29 | R ventral occipital horn | $3.64 \pm 10.36$ (78.74) | $0.69 \pm 0.39$ (2.11) | $1.14 \pm 0.54$ (2.53) |
| 30 | L ventral occipital horn | $3.43 \pm 10.38$ (80.42) | $0.86 \pm 0.67$ (4.94) | $1.39 \pm 0.98$ (5.72) |
| 31 | R olfactory sulcal fundus | $0.99 \pm 0.53$ (2.29) | $0.99 \pm 0.53$ (2.29) | $1.71 \pm 0.60$ (2.84) |
| 32 | L olfactory sulcal fundus | $1.21 \pm 0.74$ (4.53) | $1.21 \pm 0.74$ (4.53) | $2.11 \pm 0.92$ (5.81) |

$\overline{\text { AFLE values summarized as: mean } \pm \text { standard deviation (max value) }}$


Figure 3.5: A comparison of voxel overlap and distance metrics for establishing spatial correspondence between brain regions as evaluated on fMRIPrep output. (A) Multiple views showing the location of AFIDs (black dots) relative to three commonly used ROIs used in voxel overlap measures (the pallidum, striatum, and thalamus). (B,C) The histograms for voxel overlap (Jaccard index) and AFRE, respectively. The distribution for AFRE is more unimodal with a more interpretable dynamic range (in mm ) compared to voxel overlap. Trellis plots demonstrate evidence of focal misregistrations identified by AFRE not apparent when looking at ROI-based voxel overlap alone (D).
overlap for all subsequent analyses.
Mean AFRE improved from $3.40 \pm 2.55 \mathrm{~mm}$ with linear transformation alone to $1.80 \pm$ 2.09 with combined linear/nonlinear transformation (p-value $<0.001$ ). AFRE was significantly decreased with nonlinear registration for all AFIDs except the pineal gland (AFID14). AFRE was observed to be higher than mean AFLE measures (see Phase 2: $0.93 \pm 0.73 \mathrm{~mm}$ ) across the same subjects providing evidence that registration error is detectable beyond the

Table 3.3: Voxel overlap (Jaccard and Kappa) of the pallidum, striatum, and thalamus after linear registration only and combined linear /nonlinear registration.

|  |  | Jaccard |  |  | Kappa |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| roi | side | lin | nlin | lin | nlin |  |  |
| pallidum | left | $0.54 \pm 0.13$ | $0.80 \pm 0.03$ | $*$ | $0.69 \pm 0.11$ | $0.89 \pm 0.02$ | $*$ |
|  | right | $0.55 \pm 0.12$ | $0.79 \pm 0.05$ | $*$ | $0.70 \pm 0.11$ | $0.88 \pm 0.03$ | $*$ |
| striatum | left | $0.53 \pm 0.14$ | $0.83 \pm 0.03$ | $*$ | $0.68 \pm 0.13$ | $0.91 \pm 0.02$ | $*$ |
|  | right | $0.55 \pm 0.15$ | $0.82 \pm 0.05$ | $*$ | $0.70 \pm 0.13$ | $0.90 \pm 0.03$ | $*$ |
| thalamus | left | $0.70 \pm 0.11$ | $0.86 \pm 0.03$ | $*$ | $0.82 \pm 0.08$ | $0.93 \pm 0.02$ | $*$ |
|  | right | $0.69 \pm 0.11$ | $0.87 \pm 0.03$ | $*$ | $0.81 \pm 0.08$ | $0.93 \pm 0.02$ | $*$ |

* significant after FDR corrected ( q -value $<0.05$ )
limits of localization error. The number of outlier AFIDs with AFRE $>3 \mathrm{~mm}$ (more than 2 standard deviations above the mean AFLE found in Phase 2 for the same subjects) was 135/960 ( $14.06 \%$ ), representing 22/32 (68.75\%) unique AFIDs identified as misregistered. Each independent OASIS-1 subject had at least one AFID with AFRE $>3 \mathrm{~mm}$ with a mean maximum AFRE of 7.5 mm (Range: $3.16-32.78 \mathrm{~mm}$ ). Although AFLE and AFRE were statistically correlated, the effect size was small (Kendall tau $=0.15$; p-value $<0.001$; Appendix Section A.3).

Subgroup analysis for each AFID is summarized in Table 3.4. AC and PC had the lowest mean AFRE at $0.36 \pm 0.21$ and $0.57 \pm 0.29 \mathrm{~mm}$, respectively. However, registration errors as high as 1.64 mm were observed for PC. The ventricles appeared particularly difficult to align on subgroup analysis of the AFIDs. The highest AFRE among all 32 AFIDs was observed for the right and left ventral occipital horns (AFID29-30) at $3.44 \pm 5.77$ and $4.51 \pm 6.28 \mathrm{~mm}$ respectively with errors in certain cases over 20 mm (OAS1_0109 and OAS1_0203; Appendix Section A.3). Similarly, the lateral ventricle features (AFID15-18) also demonstrated high AFRE ranging from 2.11-3.01 mm on average and up to 7 mm or more. Finally, the alignment of the temporal horn features (AFID21-26) also support this observation with mean errors of $1.67-2.41 \mathrm{~mm}$ with observed errors over 5 mm .

AFRE was negatively correlated with voxel overlap but the estimates were small (tau = $-0.02 ; p=0.03$ ). Subgroup analysis demonstrated the same negative trends for the right pallidum and striatum but these results did not survive multiple comparisons correction (Figure 3.5d). No correlation between voxel overlap measures and individual AFID AFREs survived multiple comparisons correction. Comparing histograms, AFRE demonstrated a more unimodal distribution peaking between 1-2 mm (Figure 3.5b) while voxel overlap exhibited two peaks within the 0.8-0.9 range (Figure 3.5 c ). The AFRE plot also demonstrated a longer tail up to 10 mm , thus permitting a broader dynamic range in which to judge the quality of regis-
tration. In contrast, voxel overlap metrics were sparse in the lower range making interpretation more difficult. Finally, we observed that even where voxel overlap was high, suggesting good spatial correspondence, high AFRE values were also observed for certain AFIDs (see Figure 3.5d). These represent focal AFID locations where two images are misregistered despite stable voxel overlap results (Figure 3.6).


Figure 3.6: Investigating relationships between voxel overlap of the striatum and AFRE for each AFID. Focal misregistrations are identified using AFRE for the following AFIDs: 8-10, 14-18, 21-30. The most commonly misregistered regions include the inferior mesencephalon, superior vermis, pineal gland, indusium griseum, and ventricular regions. Horizontal lines are used to demarcate tiers of AFLE error above which AFRE values are beyond a threshold of localization error alone, i.e. the top horizontal line at 3 mm represents more than 2 standard deviations beyond the mean AFLE. Separate plots for the pallidum and thalamus ROIs are provided in Appendix Section A.3.

### 3.3.4 Phase 4: Evaluating template-to-template registration

Mean AFLE for BigBrainSym and MNI2009bSym was $0.59 \pm 0.40 \mathrm{~mm}$ combined with no outliers (BigBrainSym: $0.63 \pm 0.50 \mathrm{~mm}$; MNI2009bSym: $0.55 \pm 0.26 \mathrm{~mm}$ ). We highlighted

Table 3.4: AFRE after linear registration alone and combined linear/nonlinear registration.

|  |  | Mean AFRE |  |  |
| :---: | :---: | :---: | :---: | :---: |
| AFID | Description | lin | nlin |  |
| 01 | AC | $2.15 \pm 0.97$ (4.96) | $0.36 \pm 0.21$ (0.99) | * |
| 02 | PC | $1.83 \pm 0.96$ (4.58) | $0.57 \pm 0.29$ (1.64) | * |
| 03 | infracollicular sulcus | $2.20 \pm 1.23$ (5.71) | $0.93 \pm 0.53$ (2.11) |  |
| 04 | PMJ | $2.50 \pm 1.36$ (6.06) | $0.68 \pm 0.43$ (2.13) | * |
| 05 | superior interpeduncular fossa | $2.35 \pm 1.06$ (4.75) | $0.76 \pm 0.37$ (1.69) |  |
| 06 | R superior LMS | $2.07 \pm 0.95$ (4.32) | $1.17 \pm 0.74$ (3.52) |  |
| 07 | L superior LMS | $2.03 \pm 0.85$ (4.22) | $1.43 \pm 0.77$ (2.88) |  |
| 08 | R inferior LMS | $2.45 \pm 1.37$ (7.50) | $1.78 \pm 1.11$ (5.41) | * |
| 09 | L inferior LMS | $2.54 \pm 1.26$ (6.63) | $1.83 \pm 0.96$ (3.99) |  |
| 10 | culmen | $4.50 \pm 2.93$ (12.72) | $2.73 \pm 2.81$ (10.12) |  |
| 11 | intermammillary sulcus | $2.81 \pm 1.62$ (6.30) | $1.44 \pm 0.60$ (2.73) |  |
| 12 | R MB | $2.72 \pm 1.67$ (6.90) | $0.93 \pm 0.48$ (1.90) |  |
| 13 | L MB | $2.84 \pm 1.70$ (6.14) | $1.01 \pm 0.62$ (2.93) | * |
| 14 | pineal gland | $2.53 \pm 1.39$ (5.70) | $2.01 \pm 1.24$ (6.16) |  |
| 15 | R LV at AC | $4.44 \pm 1.84$ (7.90) | $2.70 \pm 1.59$ (7.85) | * |
| 16 | L LV at AC | $4.50 \pm 1.95$ (8.40) | $2.11 \pm 1.72$ (7.92) |  |
| 17 | RLV at PC | $4.81 \pm 2.54$ (10.07) | $2.96 \pm 2.42$ (9.46) | * |
| 18 | L LV at PC | $4.80 \pm 2.64$ (10.34) | $3.01 \pm 2.22$ (8.13) |  |
| 19 | genu of CC | $3.73 \pm 1.82$ (7.88) | $1.56 \pm 0.76$ (3.32) | * |
| 20 | splenium | $2.96 \pm 1.88$ (7.57) | $0.97 \pm 0.60$ (2.93) |  |
| 21 | R AL temporal horn | $3.79 \pm 1.71$ (7.50) | $1.70 \pm 1.09$ (5.23) | * |
| 22 | L AL temporal horn | $3.62 \pm 1.45$ (6.98) | $1.67 \pm 0.98$ (4.31) |  |
| 23 | R superior AM temporal horn | $3.34 \pm 1.63$ (7.25) | $1.93 \pm 1.34$ (6.85) | * |
| 24 | L superior AM temporal horn | $3.44 \pm 1.80$ (8.20) | $1.67 \pm 1.25$ (5.80) | * |
| 25 | R inferior AM temporal horn | $4.02 \pm 1.97$ (8.32) | $2.41 \pm 1.16$ (5.61) | * |
| 26 | L inferior AM temporal horn | $4.13 \pm 1.70$ (8.20) | $2.21 \pm 1.09$ (4.84) | * |
| 27 | R indusium griseum origin | $3.36 \pm 2.07$ (8.46) | $2.06 \pm 1.49$ (6.40) | * |
| 28 | L indusium griseum origin | $3.60 \pm 1.68$ (8.83) | $2.05 \pm 1.37$ (5.00) |  |
| 29 | R ventral occipital horn | $5.86 \pm 6.32$ (36.26) | $3.44 \pm 5.77$ (32.78) | * |
| 30 | L ventral occipital horn | $6.99 \pm 6.72$ (33.74) | $4.51 \pm 6.28$ (29.76) | * |
| 31 | R olfactory sulcal fundus | $2.83 \pm 1.36$ (7.50) | $1.37 \pm 0.95$ (3.44) | * |
| 32 | L olfactory sulcal fundus | $2.94 \pm 1.28$ (6.49) | $1.57 \pm 0.84$ (3.41) | * |

AFRE values summarized as: mean $\pm$ standard deviation (max value)

* significant after FDR corrected ( q -value $<0.05$ )

AFRE values beyond a threshold of 2 mm given this represents more than 2 standard deviations beyond the mean AFLE in the templates being studied. AFRE values beyond this minimum were flagged as highlighting focal misregistrations between templates.

The mean AFRE between BigBrainSym and MNI2009bSym was $2.16 \pm 1.99 \mathrm{~mm}$ and between BigBrainSym and MNI2009bAsym was $2.30 \pm 1.83 \mathrm{~mm}$, both above threshold. The largest error was 9.27 mm (MNI2009bSym) and 9.38 mm (MNI2009bAsym), found at the culmen (AFID10). Out of the 32 AFIDs defined, 11 (34.4\%) were above threshold for the symmetric template and 12 ( $37.5 \%$ ) for the asymmetric template. The most prominent misregistrations tended to occur in the posterior brainstem with the infracollicular sulcus (AFID03) and pineal gland (AFID14) quantified as 6.36 mm and 4.42 mm AFRE, respectively. These registration errors can be seen in Figure 3.7 and are summarized by AFID in Table 3.5. In addition, AFRE up to 2.78 mm were observed for AFIDs placed along the lateral mesencephalic sulcus (AFID06-09) and at the superior interpeduncular fossa (AFID05), which represent features demarcating the lateral and superior bounds of midbrain registration. Registration differences between these templates was also above threshold for the left lateral ventricle at the anterior commissure (AFID16), splenium (AFID20), left anterolateral temporal horn (AFID22), bilateral ventral occipital horns (AFID29-30), and bilateral olfactory sulcal fundi (AFID31-32). No correlation between AFRE and AFLE was found using BigBrainSym AFLE (tau $=0.071$; pvalue $=0.57$ ) or MNI2009bSym AFLE (tau $=-0.046$; p -value $=0.71$ ). Interestingly, AFRE was somewhat lower with MNI2009bAsym in many midline AFIDs but higher for certain lateral landmarks, i.e. the left inferior anteromedial temporal horn and bilateral origin of the indusium griseum (AFID26-28).

Finally, we explored the differences in correspondence between the MNI2009bSym and MNI2009bAsym. Note that these differences are not registration errors per se, as the two are not meant to be in the exact same coordinate space. The differences were generally more subtle $(0.88 \pm 0.68 \mathrm{~mm})$ but 4 AFIDs ( $12.5 \%$ ) were found to be above threshold. As expected, correspondence differences greater than 2 mm occurred in lateral rather than midline AFIDs, specifically at the left anterolateral temporal horn (AFID22), bilateral origins of the indusium griseum (AFID27-28), and left lateral ventral occipital horn (AFID30). No correlations between correspondence and AFLE were found ( $\mathrm{tau}=0.210 ; \mathrm{p}$-value $=0.09$ ).

### 3.4 Discussion

The present findings demonstrate that a series of anatomical fiducials, referred to here as AFIDs, can be consistently placed on standard structural MR images and can be used to quantify the degree of spatial alignment between brain images in millimeters. We found that AFIDs
Table 3.5: AFIDs demonstrating evidence of template-to-template misregistration for BigBrainSym with MNI2009bSym and BigBrainSym with MNI2009bAsym as well as correspondence differences between MNI2009bAsym and MNI2009bSym.

| AFID | Description | AFRE (mm) |  |  | Distance** (mm) |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | BigBrainSym vs MNI2009bSym |  | BigBrainSym vs MNI2009bAsym |  | MNI2009bAsym vs MNI2009bSym |  |
| 3 | infracollicular sulcus | 6.36 | * | 5.48 | * | 0.98 |  |
| 9 | L inferior LMS | 2.78 | * | 2.48 | * | 0.68 |  |
| 10 | culmen | 9.27 | * | 9.39 | * | 0.21 |  |
| 14 | pineal gland | 4.42 | * | 4.16 | * | 0.41 |  |
| 16 | L LV at AC | 2.05 | * | 1.22 |  | 0.86 |  |
| 20 | splenium | 2.23 | * | 2.2 | * | 0.1 |  |
| 22 | L AL temporal horn | 4.69 | * | 3.44 | * | 2.45 | * |
| 26 | L inferior AM temporal horn | 1.88 |  | 2.58 | * | 0.98 |  |
| 27 | R indusium griseum origin | 1.21 |  | 3.6 | * | 2.81 | * |
| 28 | L indusium griseum origin | 0.74 |  | 2.88 | * | 2.29 | * |
| 29 | R ventral occipital horn | 2.54 | * | 3.99 | * | 1.63 |  |
| 30 | L ventral occipital horn | 5.88 | * | 4.22 | * | 2 | * |
| 31 | R olfactory sulcal fundus | 2.62 | * | 1.84 |  | 1.1 |  |
| 32 | L olfactory sulcal fundus | 3.06 | * | 4.21 | * | 1.24 |  |



Figure 3.7: Select views demonstrating registration errors between BigBrainSym and MNI2009bSym. The green dots represent the optimal AFID coordinates in MNI2009bSym space superimposed in both templates to provide a basis for comparing registration differences. While many of the midline AFIDs are stable across both templates, the infracollicular sulcus, pineal gland, splenium, and culmen are misregistered in BigBrainSym (red arrows). The AFIDs draw attention to registration differences in the BigBrainSym space in the tectal plate, pineal gland, and superior vermis (blue arrows).
are reproducible, not overtly manually intensive (20-40 minutes once trained), and more sensitive to local registration errors than standard voxel overlap measures. Our entire protocol and study framework leverages open resources and tools, and has been developed with full transparency in mind so that others may freely use, adopt, and modify.

The work presented here is inspired heavily by classical stereotactic methods [Talairach et al., 1957], where point-based correspondence has been used to align brain templates with patient anatomy to enable atlas-based surgical targeting. The anterior and posterior commissure were originally identified as prominent intraventricular features based on air studies, prior to the invention of computed tomography or MRI. The AC and PC have proven to be reliable features on MRI and were adopted by neuroscientists for the alignment of brain images to templates, in what is referred to as the Talairach grid normalization procedure [Brett et al., 2002, Talairach and Tournoux, 1988]. The advent of robust and openly available software for automatic or semi-automatic labeling of regions-of-interest in brain images has led to a relative underemphasis of point-based alignment. We demonstrate here that point-based metrics are more sensitive to focal misregistrations than voxel overlap measures and quantified in millimeters.

Tolerance to focal misregistration in images undoubtedly will depend on the application; but there is no doubt that poor image correspondence can result in inaccurate (and possibly erroneous) predictions and conclusions in neuroimaging studies. Our results evaluating cor-
respondence error in an fMRI preprocessing pipeline revealed local template misregistrations of $1.80 \pm 2.09 \mathrm{~mm}$. For many fMRI or diffusion-based applications, this mean error is about the size of a voxel; and thus may be within an acceptable tolerance. However, mean maximum errors of over 7 mm were also observed and may begin to impact the sensitivity to discovery as well as the accuracy of localization of affected brain regions in a task or connectivity analyses. These misregistrations also may affect the interpretation of voxel-based and deformation-based morphometry studies that seek to investigate subtle shape differences between study populations. Finally, minimizing registration error becomes particularly critical for analyses pertaining to stereotactic interventions like deep brain stimulation (DBS) where millimeters can represent the difference between optimal therapy and side effects.

### 3.4.1 Protocol development and validation

After a single training session, novice raters could place AFIDs at a mean AFLE of approximately $1-1.5 \mathrm{~mm}$ across all AFID32 points. Placement error varied from one template to another and among AFIDs (Appendix Section A.1). Raters had the least amount of error with placements for the MNI2009bAsym and Agile12v2016 templates. In contrast, fiducial placement errors were higher when raters were asked to place AFIDs for individual subjects, i.e. Colin27 as well as the OASIS-1 database. Repeatability was assessed using measures of intrarater and inter-rater AFLE. Intra-rater AFLE was lowest for the MNI2009bAsym and highest in Colin27 (Table 3.1). Inter-rater AFLE was again lowest for MNI2009bAsym and highest in Colin27 and the OASIS-1 datasets. This demonstrates how AFIDs are more difficult to place due to individual variability versus in population templates where the individual nuances of these features may be effectively blurred out. Overall, the placement error remains acceptable (1-2 mm) among all annotated images.

The AC and PC were the most reliably identifiable AFIDs with mean AFLE of less than 0.5 mm and inter-rater AFLE of $0.5-1 \pm 0.3 \mathrm{~mm}$ observed. These results compared favorably to an analysis of experienced neurosurgeons by Pallaravam and colleagues placing the same AC-PC points where they observed a point placement error (equivalent to the inter-rater AFLE metric used here) that was surprisingly higher at $1-2 \mathrm{~mm} \pm 1.5 \mathrm{~mm}$ [Pallavaram et al., 2008]. We speculate that the higher variability in the referenced study was the lack of restriction on how the AC-PC landmarks were placed; that is, some stereotactic neurosurgeons continue to use the intraventricular edge of each commissure, which was the classical technique used by Talairach during air studies, while others used the center of each commissure [Horn et al., 2017a]. The distance from the center to the ventricular edge can be several millimeters likely accounting for this difference. Overall, our findings demonstrate that enforcing certain practices such as using
the center of each commissure play an important role in the consistency and standardization of fiducial placement.

In contrast, certain fiducial points contributed substantially to worse overall estimates of fiducial localization error. In particular, the bilateral ventral occipital horns (AFID29-30) had higher placement errors. Placement was particularly inaccurate for individual subjects where the ventricular atrium tapered completely in many individual subject studies (including Colin27), and thus the posterior continuation into the occipital horn was sometimes difficult to visualize or resolve at all. The bilateral origins of the indusium griseum (AFID27-28) were also difficult for raters to place consistently.

### 3.4.2 Point-based versus ROI-based metrics

Previous work has shown that nonlinear registration improves alignment between structures [Chakravarty et al., 2009, Hellier et al., 2003, Klein et al., 2009], and that the choice of parameters matters. These existing studies have mostly used voxel overlap measures to support their findings. Our results are also in-line with prior work but also demonstrate how AFIDs are complementary and more sensitive than ROI-based metrics for evaluating both local and global spatial correspondence of brain images (see Figure 3.5).

We were able to compare the relative efficacy of AFRE and voxel overlap for subjects from the OASIS-1 database and several commonly used templates. AFRE had a more unimodal distribution and a longer tail facilitating identification of focal misregistrations between images (Figure 3.5). On the other hand, the Jaccard histogram was more sparse towards the tail of the distribution suggesting a poorer ability to discriminate. One key advantage of AFRE is its interpretability, representing the distance in millimeters between aligned neuroanatomical structures in two images, compared to voxel overlap, which is a relative measure and unitless. It is commonly perceived in segmentation studies that voxel overlap measures greater than 0.7 represent accurate correspondence between regions. However, our analysis demonstrates that even with generally high overlap after nonlinear registration, focal misregistrations of AFIDs above 7 mm may be identified (Figure 3.6 and Table 3.4).

### 3.4.3 Subject-to-template registration

We chose to evaluate the subject-to-template registrations computed as part of an fMRI processing pipeline, fMRIPrep [Esteban et al., 2018], as a use case for our AFIDs protocol. Functional MRI studies may not represent the optimal use case due to the relatively coarse spatial resolution relative to the size of misregistration effects we can detect with AFIDs, and because most fMRI researchers are focused on cortical activation while our protocol emphasizes and de-
tects misregistrations in the deep brain regions. Our choice to investigate fMRIPrep registration performance was motivated by their transparent approach to the development of preprocessing software for neuroimaging and BIDS integration [Gorgolewski et al., 2016, Gorgolewski and Poldrack, 2016]. The active developer and support base, as well as growing adoption by many end-users were other contributing factors. Our analysis revealed misregistrations on the order of $1.80 \pm 2.09 \mathrm{~mm}$ and as high as over 30 mm that would be more difficult to identify by qualitative evaluation or ROI-based analysis alone.

While this points to potential caution with the use of standardized pipelines like fMRIPrep for template registration, it should be noted that fMRIPrep was designed with a focus on robustness, rather than accuracy. The underlying parameters and processing steps used in fMRIPrep are fully transparent. In addition, the underlying deformable registration software used [Avants et al., 2008] has been demonstrated to achieve high performance in studies using traditional voxel overlap measures [Klein et al., 2009]. The focal template misregistrations we have identified in fMRIPrep with AFIDs are meant to serve as a baseline for refinement in future versions that can be compared transparently and potentially incorporated for testing new versions as part of a continuous integration workflow. Using additional image contrasts [Xiao et al., 2017] or subcortical tissue priors [Ewert et al., 2019] to drive template registration have been demonstrated using conventional voxel overlap techniques to result in more optimal registrations that can also be tested using the AFIDs framework.

### 3.4.4 Template-to-template registration

We recommend that imaging scientists exercise caution when displaying statistical maps using a template other than the one to which the original deformations were performed. For example, it has become increasingly common to project statistical maps and subject data registered to MNI space using BigBrain for visualization purposes. In this study, we identified clear evidence of registration differences between several templates commonly assumed to be in the same coordinate space: BigBrainSym and MNI2009bSym, and even greater between BigBrainSym and MNI2009bAsym because of the differences in AFID locations in MNI2009bSym and MNI2009bAsym. Specifically, misregistrations as high as over 9 mm have been identified. Many of these errors occur in the midbrain region (Table 3.5), which would have implications in particular if using BigBrainSym to project locations of electrode implantations. In support of other recent work (Horn et al., 2017), this study highlights the importance of understanding which exact template one is using for processing and analysis: that multiple "MNI" templates exist (with different version dates, types, and symmetry), as do registration differences between these templates.

### 3.4.5 Teaching neuroanatomy

Our AFID32 protocol may also hold particular value for teaching neuroanatomy. In fact, evidence from our study suggests that even relative novices can be trained to place AFIDs accurately, including the AC and PC, with comparable accuracy and variability to trained neurosurgeons (Table 3.2). By releasing the data acquired in this study, we provide a normative distribution of AFID placements that can be used to quantify how accurately new trainees can place points. These measures can be used to gauge the comprehension of students regarding the specific location of neuroanatomical structures in a quantitative (millimetric) manner and focus efforts on consolidating understanding based on where localization errors were higher. To date, over a series of locally-held workshops and tutorials, over 60 students have been trained to complete the AFID32 protocol.

### 3.4.6 Limitations and future work

While we have found the AFIDs proposed to be quite reliable, there is clearly location-related heterogeneity in placement error. We make no claims that this set of anatomical fiducials is optimal and in the future, other locations may prove to be more effective than others. Also, for this first proposed set of AFIDs, we limited our locations to deep structures where less inter-subject variability exists compared to cortical features (Thompson et al., 1996); future extensions could include linking our workflow with cortical surface-based [Fischl, 2004] and sulcal-based [Hellier et al., 2003, Perrot et al., 2011, Mangin et al., 2015] methods of spatial correspondence. Development of similar protocols for other neuroimaging modalities such as T2-weighted or diffusion-based contrasts may also be of value. In addition, fiducial localization error may be biased by how the raters were taught to place the fiducials; in our case, we organized an initial interactive tutorial session, and provided text and picture-based resources of how to place the AFIDs. It is also possible that AFLE would be lower if performed by a more experienced group of raters. Also, how AFID placement behaves in the presence of lesional pathology remains an open question. We have made the annotations and images available to allow other groups to propose other AFID locations and descriptions that could be similarly validated. We plan to post any modifications to the protocol as separate versions at the linked repository.

The AFIDs protocol requires correct placement of the anterior commissure (AFID01) and posterior commissure (AFID02) points. We made this decision as it helps to align the brain images into a more standard orientation for subsequent placement of bilateral fiducials. In particular, 4 of the AFIDs are dependent on AC-PC alignment (the lateral ventricles at AC and PC in the coronal plane). It is possible that error in AFID placements could be compounded by ini-
tial error in placement of AC and PC. Fortunately, AC and PC can be placed with high trueness and precision ( $<1 \mathrm{~mm}$ ) (Table 3.2), consistent with prior studies [Liu et al., 2015]. We made the decision to perform AC-PC alignment to permit more accurate placement of lateral AFIDs, which may otherwise have appeared quite oblique from each other if the individual's head was tilted in the scanner. Thus, on balance, AC-PC alignment probably mitigates placement error in lateral AFIDs compared to placing fiducials in the native MRI space. Further research can examine these potential spatial biases more systematically.

Beyond evaluating correspondence, AFIDs could be used for point-based inter-subject or subject-to-template registration. AFIDs used in combination with classic rigid registration algorithms such as Iterative Closest Point [Besl et al., 1992] may result in more optimal initial linear registration between images. In addition, point-based deformable registration using (Bsplines) may produce more efficient, lower order deformable registrations between two images [Bookstein, 1997]. To prevent circular reasoning, we thought this would be best evaluated as independent studies. Finally one compelling extension of this work would be to automate or semi-automate AFID placement, which would enable inclusion of AFID-based metrics in standardized workflows involving template or intersubject registration.

### 3.5 Conclusions

Our proposed framework consists of the identification of anatomical fiducials, AFIDs, in structural magnetic resonance images of the human brain. Validity has been established using several openly available brain templates and datasets. We found that novice users could be trained to reliably place these points over a series of interactive training sessions to within millimeters of placement accuracy. As an example of different use cases, we examined the utility of our proposed protocol for evaluating subject-to-template and template-to-template registration revealing that AFIDs are sensitive to focal misregistrations that may be missed using other commonly used evaluation methods. This protocol holds value for a broad number of applications including intersubject alignment and teaching neuroanatomy.

## Chapter 4

Quantification of local geometric distortion in structural magnetic resonance images: Application to ultra-high fields

This chapter is based on the following manuscript:

- Lau, J. C., Khan, A. R., Zeng, T. Y., MacDougall, K. W., Parrent, A. G., \& Peters, T. M. (2018). Quantification of local geometric distortion in structural magnetic resonance images: Application to ultra-high fields. NeuroImage, 168, 141-151.


### 4.1 Introduction

Ultra-high field ( $\geq 7$ Tesla) magnetic resonance imaging (MRI) allows for improved in vivo visualization of neuroanatomical structures, facilitating detailed morphological and functional study. The superiority of 7T neuroimaging has been demonstrated for visualizing subcortical anatomy due to increased spatial resolution, tissue contrast, and improved signal-to-noise ratio [Abosch et al., 2010, Cho et al., 2010, Kerl et al., 2012]. The subthalamic nucleus (STN) and globus pallidus internus ( GPi ) are more clearly and reliably visualized at 7T, suggesting ultrahigh field MRI can facilitate better understanding of these structures [Kerl et al., 2012, Keuken et al., 2013].

However, the increase in field strength can result in significant geometric artifacts that must be considered, not only in neuroscientific study, but also when considering the use of 7T for image-guided interventions where accuracy is critical [Sankar and Lozano, 2011, Sumanaweera et al., 1994a, Sumanaweera et al., 1994b, Wang et al., 2005]. Stereotactic neurosurgical procedures rely on submillimeter-to-millimeter ranges of accuracy for patient safety, and the presence of geometric distortion in MRI results in loss of spatial confidence in patient images. Geometric inhomogeneity in MR images is the result of multiple scanner and patient-related factors including gradient coil nonlinearity and magnetic susceptibility. Gradient coil nonlinearities are geometric errors introduced through the coil design process with regions closest to the magnet and coil isocenter being the most geometrically accurate. Magnetic susceptibility artifacts occur as a result of local magnetic field inhomogeneities at tissue interfaces, and since they increase linearly with field strength, can become increasingly problematic at ultra-high fields [Littmann et al., 2006].

Several studies have evaluated the problem of geometric distortion at ultra-high field [Cho et al., 2010, Duchin et al., 2012, Neumann et al., 2015]. Results using MRI phantoms have been inconsistent with one study reporting submillimeter accuracy with appropriate higherorder active shimming [Cho et al., 2010] and another claiming deviations in the 2-3 mm range [Neumann et al., 2015]. Furthermore, the extent to which a phantom approximates geometric distortion in in vivo patient data remains unclear. Phantom studies can characterize gradient nonlinearities but do not account for anatomical and physiological distortions resulting from magnetic susceptibilities at tissue interfaces. Another approach involves characterization
of distortion in subjects scanned on both 7T and lower-field MRI. By using a combination of qualitative evaluation of registration, manual fiducial placement error, and analysis in 9 block-shaped regions, Duchin et al. [Duchin et al., 2012] showed that subcortical targets are minimally effected by geometric distortion and most significant in the orbital frontal block. Unfortunately, the results of existing studies are difficult to generalize to other sites as the distribution and magnitude of geometric distortion is known to vary from scanner to scanner [Caramanos et al., 2010]. Replicating these studies at each site with fiducial placement would be manually intensive, and prone to intra-rater and inter-rater variability.

In this work, we present a computational method for quantifying geometric distortion in 7T structural images. Our technique estimates distortion using deformation fields automatically derived from nonlinear registration of 7 T data to corresponding subject data at lower fields. Voxel-wise statistical analysis in a common stereotactic space permits the identification of focal brain regions systematically affected by geometric distortion that should be accounted for in neuroimaging analysis and planning stereotactic interventions. Finally, for validation, we compared our automated measures against manual fiducial placement.


Figure 4.1: Workflow for the quantification of local geometric distortion in a single subject at ultra-high field. The leftmost images represent 3 T and 7T images of the same subject in native space at the best equivalent sagittal slice. Qualitatively, the 7T image is more block-shaped than the equivalent 3T image. Checkerboard visualization reveals areas of registration mismatch after rigid and nonlinear registration stages (affine stage omitted from figure). Finally, we quantify local displacement in millimeters overlaid on the 3T image space using the nonlinear deformation field.

### 4.2 Materials and Methods

### 4.2.1 Participants and MRI Acquisition Protocol

22 participants (mean age $30.8 \pm 9.2$ years; 14 female) were scanned on both 3 T and 7 T imagers at Robarts Research Institute (London, ON, Canada) between 2010 and 2013 (time between scans: $78.2 \pm 217.8$ days; range: 0-633 days; 12 of 22 participants with scans on the same day). Informed consent was obtained.

All subjects were scanned using a 3T MRI scanner (GE Discovery MR750). A T1-weighted (T1w) 3T image was acquired as part of a DESPOT1-HIFI protocol using a 32-channel head coil [Deoni et al., 2007, Deoni et al., 2005, Deoni et al., 2003] with a receiver bandwidth of 19.23 kHz . The DESPOT1-HIFI technique permits rapid high-resolution quantitative T1 mapping extracted from spoiled gradient recall (SPGR) echo images acquired at two different flip angles $\left(4^{\circ} / 18^{\circ}\right)$ with other parameters remaining the same: $\mathrm{TR}=8.36 \mathrm{~ms}, \mathrm{TE}=3.71 \mathrm{~ms}$, matrix $=220 \times 220$, slice thickness $=1 \mathrm{~mm}$, resolution $=1 \mathrm{~mm}$ isotropic. The T1w image acquired with a flip angle of $18^{\circ}$ was used for all further morphometric comparisons between 7 T and 3T. Ultra-high field data were acquired on a 7T imager (Agilent, Santa Clara, CA, USA/Siemens, Erlangen, Germany) using a 16 -channel transmit-receive head coil array with a receiver bandwidth of 50 kHz . A T1-weighted MPRAGE sequence was acquired (TR $=8.1$ $\mathrm{ms}, \mathrm{TE}=2.8 \mathrm{~ms}, \mathrm{TI}=650 \mathrm{~ms}$, flip angle $=11^{\circ}$, matrix $=256 \times 512 \times 172$, resolution $=$ 1 mm isotropic, scan time $=5: 42 \mathrm{~min}$ ). Both 3 T and 7 T datasets were acquired as sagittal source 3D with the readout direction being superior-inferior and the phase-encode direction being anterior-posterior with respect to the brain anatomy.

### 4.2.2 Data Processing

Intensity non-uniformities of both 7T and 3T T1 images were corrected using retrospective bias field correction by N4 [Sled et al., 1998, Tustison et al., 2010]. The Brain Extraction Tool (BET) as implemented in FSL was used for brain masking (fractional intensity threshold $=0.4$ ). At 7T, bias field correction was performed a second time due to evidence of residual intensity non-uniformity [Boyes et al., 2008, Seiger et al., 2015, Tardif et al., 2010].

An image-processing pipeline was developed to iteratively align a source to a target T 1 w image. The 3T T1w image was considered the target for registration. The source volume (7T) was registered iteratively to the target volume (3T) first using rigid body (6 degrees-of-freedom or DOF) and affine (12 DOF) registration via block-based matching [Ourselin et al., 2001], followed by nonlinear registration using a free-form deformation algorithm implemented in
the NiftyReg package using default parameters ${ }^{1}$ [Modat et al., 2010]. The combined linear and nonlinear transformations were concatenated and applied to the source volume, with the result being a 7T T1 image resampled to 3 T space at 1 mm isotropic resolution. For group analysis, transformations were computed from 3T to MNI152 space using the same iterative process of rigid to affine to nonlinear registration.

### 4.2.3 Voxel-Level Metrics

The computed deformation field was converted to a 1 mm isotropic displacement field for each subject providing an estimate of the displacement of a voxel at 7 T to the best corresponding local voxel at 3 T after nonlinear registration. Euclidean distance was computed from the displacement vector at each voxel location as a quantitative estimate of local geometric distortion, which we have called automated displacement for consistency. Voxel-wise scalar displacements in the standard Cartesian directions were also stored so that we could determine which $\mathrm{x}, \mathrm{y}$ or z contributed to the local displacement value. In order to evaluate the principal $\mathrm{x}, \mathrm{y}$ or $z$ component of displacement at each voxel, we used two different metrics: maximum component and relative index. The maximum component was computed by calculating the maximum $\mathrm{x}, \mathrm{y}$ or z component. The relative index was calculated by normalizing the displacement in each component by the sum of displacements $(x+y+z)$.

The image isocenter coordinates were propagated from the 7 T and 3 T origins (world coordinates: $0,0,0$ ) to the MNI152 reference space using the previously computed transformations. The relationship between distance from isocenter and local displacement was plotted on a bivariate histogram across all voxels and subjects.

For group analysis, the scalar automated displacement and component maps were transformed into MNI152 space using the previously computed transformations from 3T to standard space. Mean automated displacement was calculated as a summary measure for all included subjects. To evaluate the effect of the position of the 7T image isocenter on distortion, we computed correlation maps between the $\mathrm{x}, \mathrm{y}$, and z components of the isocenter with corresponding voxel estimates of automated displacement.

Scalar displacement and correlation maps for each subject were imported into R (version 3.2.4) for statistical processing. At each voxel within the reference brain mask, we considered any statistically significant displacement more than 1 voxel ( 1 mm at 3 T ) to be clinically significant, and thus tested the null hypothesis that voxel-wise displacement was less than 1 mm . Non-parametric Wilcoxon rank sum testing was used for statistical analysis. Multiple comparisons correction was performed using the false-discovery rate (FDR) controlling for an adjusted

[^2]p-value ( $q$-value $<0.025$ ). The FDR-corrected q maps were converted to binary masks, which were then applied to the effect size maps and overlaid on the reference atlas for visualization.

### 4.2.4 Region-of-Interest Analysis

Region-of-interest (ROI) analysis was performed using the Harvard-Oxford (Desikan et al., 2006; 113 regions) and ATAG subcortical [Keuken et al., 2013] atlases in MNI space. Since the ATAG atlas is probabilistic, we used binarized ROIs after thresholding for voxels with $\geq$ $10 \%$ likelihood of the corresponding label. In addition, a lobar atlas was derived from the Harvard-Oxford atlas by classifying each of the 113 regions into frontal, parietal, occipital, temporal, insular, and subcortical regions ( 6 per hemisphere; 12 regions). Regional differences in geometric distortion were summarized using two measures: mean displacement (average automated displacement among all voxels in the ROI) and maximum displacement (average displacement after taking maximum automated displacement at each voxel in the ROI). As in the voxel-wise analysis, the null hypothesis of less than 1 mm ( 1 voxel) of displacement was tested non-parametrically. Multiple comparisons correction was performed for all ROIs (137 total) separate from the voxel-wise analysis using an FDR of $q<0.025$. The regional effect of isocenter position was summarized using mean x , y , and z correlation in each ROI (137 x $3=$ 411 total comparisons) also corrected using the same FDR.

### 4.2.5 Validation

Quality of image preprocessing and registration were evaluated by visual inspection (JCL, TYZ) of all subject data by examining correspondence between anatomical features in images as overlays in 3D Slicer [Fedorov et al., 2012]. Quantitative validation was performed by manual fiducial placement on both 7 T and 3T datasets, also in 3D Slicer. Twenty subjects had five midline fiducials using a previously published protocol [Duchin et al., 2012], and five subjects had 75 fiducials. The midline fiducial locations were the following:

1. Anterior commissure
2. Posterior commissure
3. Midpoint of the cerebral aqueduct
4. Midpoint of the optic chiasm
5. Midpoint of the infundibulum

Five randomly chosen subjects had a total of 75 fiducials placed ( 5 midline, 60 whole brain, 10 in highly displaced areas). For the whole brain fiducials, 30 were placed in each hemisphere with the rater choosing salient 3D neuroanatomical locations that could be identified on the cortical surface and sulci. An additional 10 fiducials were chosen based on regions identified as highly displaced, in other words, voxels located in areas of increased displacement identified by overlaying our automated method on the subject's 3 T image.

Each fiducial was placed by two raters (JCL, TYZ) on the 7T image after initial placement at 3 T for each subject. After several weeks, the fiducials were independently placed a second time at 7 T by a single rater (TYZ). The Euclidean distance was used to estimate landmark placement error between corresponding landmarks at 7T by the same rater to estimate intrarater reliability, and between corresponding landmarks on the average fiducial location for TYZ and the corresponding landmark placed by JCL to estimate inter-rater reliability. All fiducials were also classified into categories: midline ( 5 landmarks previously described), lobar (temporal, occipital, parietal, frontal, insular), subcortical, and highly displaced.

The mean 7T fiducial location between all three manual ratings (TYZ x 2, JCL x 1) was used for all subsequent evaluations of registration quality between 7 T and 3 T images. The mean 7T fiducial locations were spatially aligned into 3T space iteratively using the transformations calculated in Section 4.2.2 on Data Processing. The Euclidean distance between the aligned 7 T fiducial location and the corresponding 3 T fiducial estimated registration error for each stage. The fiducial registration error after rigid body and affine registration steps provided an estimate of geometric distortion as calculated manually, which we call manual displacement. We evaluated our candidate automated displacement measure against manual displacement using Pearson correlation and a Bland-Altman plot for agreement [Bland and Altman, 1986]. To determine if there was a significant difference with registration steps, we performed a two-tailed Wilcoxon rank sum test between all combinations of fiducial categories and registration steps ( 14 fiducial categories, $3 \times$ registration steps $=42$ comparisons). These results were adjusted using the false-discovery rate (FDR) controlling for an adjusted p-value ( $q$-value $<0.025$ ) [Genovese et al., 2002].

### 4.2.6 Effect of Gradient Distortion Correction

Data analyzed as part of the study were acquired prior to the scanner upgrade from an Agilent 7 T to a Siemens 7 T . As a result, post hoc gradient distortion correction was not available for evaluation. We prospectively scanned a single subject at 7 T and 3 T using best equivalent MP2RAGE sequences at 0.8 mm resolution [Marques et al., 2010]. At 7T, the MP2RAGE protocol was as follows: $\mathrm{TR}=6.0 \mathrm{~s}, \mathrm{TE}=2.19 \mathrm{~ms}$, matrix $=208 \times 320 \times 320$, resolution $=0.8$
mm isotropic, scan time $=9: 36 \mathrm{~min}$ ). At 3 T , the protocol was: $\mathrm{TR}=5.0 \mathrm{~s}, \mathrm{TE}=3.51 \mathrm{~ms}$, matrix $=208 \times 320 \times 320$, resolution $=0.8 \mathrm{~mm}$ isotropic, scan time $=8: 27 \mathrm{~min}$ ). Both uncorrected and distortion corrected datasets were acquired and processed through our automated framework as described in Section 4.2.2. The density maps of automated displacements within the masked brain were plotted and compared. The corresponding displacement maps with and without correction were visually assessed. Finally, ROI analysis was performed and percent change due to correction was calculated as the difference between corrected and uncorrected displacement divided by uncorrected displacement.


Figure 4.2: The histogram of mean automated displacements within the masked brain is shown in (a). Since automated displacement is derived from the Euclidean distance between points, all are greater than 0 mm . The histogram has a right skew deviation. The vertical line shows the mean of 0.94 mm . Local voxel displacement is demonstrated to increase with distance from the 7T image isocenter (b). Voxels across all subjects in the study were binned in a 2D histogram (heatmap) according to displacement and distance from isocenter (log-scaled). Mean (thick line) and standard deviation (thin lines) are shown. Mean automated displacement is demonstrated to increase beyond 1 mm at 80 mm .

### 4.3 Results

### 4.3.1 General Findings

Nonlinear registration, but not linear registration alone, qualitatively improved our ability to structurally align 7T and 3T images for the same subject (Figure 4.1). As shown in Figure 4.2a, automated displacement was always greater than zero, showed right skew deviation, and
did not meet statistical criteria for normality (Anderson-Darling test, p-value $<0.00$ ). Given these observations, the non-parametric Wilcoxon rank sum test was used for statistical analysis. We performed one-tailed hypothesis testing with the null hypothesis of displacement $<1 \mathrm{~mm}$, as only locations with more than 1 mm (or voxel) of displacement were considered clinically significant. Statistically significant displacements in the opposite direction (i.e., close to zero mm ) were not considered clinically meaningful and thus the lower tail was excluded from analysis.


Figure 4.3: Selected images from displacement maps computed for group analysis are overlaid on the MNI152 template with slice references in world coordinates. All voxels shown are significant after controlling for multiple comparisons using FDR ( $q<0.025$ ). Each displacement map has a corresponding image showing an overlay of the maximum $x$, $y$ or $z$ component shown in red, green or blue, respectively, for all significant regions. The main component found was in the z-direction in $56.84 \%$, followed by the $x$-direction in $22.51 \%$, and finally the y-direction in $20.64 \%$.

### 4.3.2 Identification and Characterization of Local Distortion

By extracting local metrics of displacement from the deformation field, we were able to identify local voxels systematically affected by geometric distortion at ultra-high field, quantified in millimeters. Selected images are shown for a single subject (Figure 4.1) and on group analysis
(Figure 4.3). Overall, $13.2 \%$ of voxels within the brain mask showed statistically significant displacement greater than 1 mm . Voxel-wise analysis identified significant distortions bilaterally on the floor of the anterior and middle fossae in the orbitofrontal cortex, mesial temporal poles, fusiform and temporal-occipital gyri, posterior inferior temporal gyri, and in the occipital/suboccipital regions. The maximum component of displacement was in the $x$ direction at $42.60 \%$, followed by the longitudinal direction ( z ) at $37.31 \%$, and finally in the y direction at $20.01 \%$. Using the relative index, the contribution in the x component was $35.00 \%$, z component was $34.87 \%$, and y component was $30.12 \%$. When analyzing only those voxels identified as significantly displaced after adjustment, the maximum component was predominantly in the longitudinal direction (z component) at $56.84 \%$, followed by the x component at $22.51 \%$, and finally the $y$ component at $20.65 \%$ (Figure 4.3). The relative index also shifted to be first in the $z$ component at $40.31 \%$, followed by the $y$ component at $31.18 \%$, and finally the x component $28.52 \%$. We correlated voxel-wise displacement with the $\mathrm{x}, \mathrm{y}, \mathrm{z}$ positions of the 7 T image isocenter and found that the superior parietal region was negatively correlated and the suboccipital was positively correlated with z position (Figure 4.4). Regions close to air-filled sinuses were not significantly correlated with isocenter position.

### 4.3.3 Distortion Increases with Isocenter Distance

Geometric distortions increased with distance from the image isocenter to over 1 mm at 80 mm distance from the isocenter (Figure 4.2b). The 7T isocenter location in our dataset was $\mathrm{x}=-1.91 \pm 2.67 \mathrm{~mm}, \mathrm{y}=2.27 \pm 7.41 \mathrm{~mm}$, and $\mathrm{z}=3.41 \pm 12.98 \mathrm{~mm}$, and the 3 T isocenter location was $\mathrm{x}=3.17 \pm 3.80 \mathrm{~mm}, \mathrm{y}=-21.11 \pm 5.72 \mathrm{~mm}, \mathrm{z}=-16.04 \pm 9.53 \mathrm{~mm}$ in MNI152 world coordinates. The mean distance between 7 T and 3 T isocenters was $33.18 \mathrm{~mm} \pm 12.54$ mm (differences in each component: $\mathrm{x}=-5.08 \pm 4.26, \mathrm{y}=23.38 \pm 8.47, \mathrm{z}=19.45 \pm 14.99$ ).

### 4.3.4 Region-of-Interest Analysis

Regional effects of from geometric distortion are summarized in Tables 4.1-4.3. After adjustment for multiple comparisons, no lobar ROIs were statistically significant, although there was a trend towards significance in the average voxel-wise displacement in the left occipital and temporal lobes (Table 4.1). Maximum displacements were noted to be highest in bilateral frontal, temporal, and occipital lobes, observed to be $7-8 \mathrm{~mm}, 4-6 \mathrm{~mm}$, and $4-5 \mathrm{~mm}$ respectively, on average across subjects. Using the full Harvard-Oxford atlas, 17 of 113 regions, or $15.0 \%$, met thresholds of statistical significance after FDR correction ( $q<0.025$ ) (Table 4.2; full table available in Appendix Section B.1). Regions identified were: bilateral temporal poles, left anterior middle temporal gyrus, bilateral posterior inferior temporal gyri, bilateral

Table 4.1: Mean and maximum displacements in millimeters for ROIs part of the lobar and ATAG subcortical atlases.

| region | side | atlas | mean | stdev | V | p-value | q-value | max | max stdev |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Left Red Nucleus | left | ATAG | 1.02 | 0.53 | 111 | 0.69 | 1.00 | 1.69 | 0.98 |
| Left Substantia Nigra | left | ATAG | 0.86 | 0.39 | 76 | 0.95 | 1.00 | 1.52 | 0.67 |
| Left Subthalamic Nucleus | left | ATAG | 1.05 | 0.59 | 108 | 0.73 | 1.00 | 1.48 | 0.79 |
| Left Striatum | left | ATAG | 0.59 | 0.14 | 0 | 1.00 | 1.00 | 2.41 | 0.99 |
| Left Globus Pallidus Externus | left | ATAG | 0.48 | 0.15 | 0 | 1.00 | 1.00 | 1.33 | 0.66 |
| Left Globus Pallidus Internus | left | ATAG | 0.55 | 0.21 | 2 | 1.00 | 1.00 | 1.19 | 0.55 |
| Right Red Nucleus | right | ATAG | 1.03 | 0.40 | 131 | 0.45 | 1.00 | 1.71 | 0.72 |
| Right Substantia Nigra | right | ATAG | 0.78 | 0.30 | 31 | 1.00 | 1.00 | 1.48 | 0.80 |
| Right Subthalamic Nucleus | right | ATAG | 0.98 | 0.53 | 99 | 0.81 | 1.00 | 1.44 | 0.70 |
| Right Striatum | right | ATAG | 0.60 | 0.19 | 3 | 1.00 | 1.00 | 2.38 | 0.91 |
| Right Globus Pallidus Externus | right | ATAG | 0.50 | 0.22 | 2 | 1.00 | 1.00 | 1.31 | 0.81 |
| Right Globus Pallidus Internus | right | ATAG | 0.50 | 0.24 | 6 | 1.00 | 1.00 | 1.10 | 0.72 |
| Left Frontal | left | lobar | 0.91 | 0.12 | 30 | 1.00 | 1.00 | 7.69 | 1.19 |
| Left Parietal | left | lobar | 0.81 | 0.11 | 4 | 1.00 | 1.00 | 3.46 | 0.73 |
| Left Occipital | left | lobar | 1.15 | 0.26 | 197 | 0.01 | 0.08 | 4.62 | 1.83 |
| Left Temporal | left | lobar | 1.11 | 0.21 | 195 | 0.01 | 0.09 | 5.99 | 1.53 |
| Left Insular | left | lobar | 0.61 | 0.10 | 0 | 1.00 | 1.00 | 1.80 | 0.30 |
| Left Subcortical | left | lobar | 0.70 | 0.22 | 11 | 1.00 | 1.00 | 2.61 | 1.10 |
| Right Frontal | right | lobar | 0.89 | 0.11 | 24 | 1.00 | 1.00 | 8.07 | 1.71 |
| Right Parietal | right | lobar | 0.76 | 0.11 | 1 | 1.00 | 1.00 | 3.56 | 0.79 |
| Right Occipital | right | lobar | 1.07 | 0.17 | 166 | 0.10 | 0.50 | 4.24 | 0.94 |
| Right Temporal | right | lobar | 1.04 | 0.12 | 169 | 0.09 | 0.43 | 5.75 | 0.96 |
| Right Insular | right | lobar | 0.66 | 0.13 | 1 | 1.00 | 1.00 | 2.07 | 0.62 |
| Right Subcortical | right | lobar | 0.75 | 0.24 | 25 | 1.00 | 1.00 | 2.94 | 0.98 |
| V represents the Wilcoxon | rank sum statistic p-value | is | udiusted | and q-value is FDR ad- |  |  |  |  |  |

$\overline{\mathrm{V}}$ represents the Wilcoxon rank sum statistic, p-value is unadjusted, and q-value is FDR adjusted. None of the regions met thresholds for statistical significance after FDR correction at a rate of $q$-value $<0.025$.


Figure 4.4: The effect of 7 T isocenter position on local distortion. Correlation maps of isocenter position against displacement are overlaid on selected sagittal images on the MNI152 template with corresponding world coordinates. All voxels shown are significant after controlling for multiple comparisons using FDR ( $q<0.025$ ). The top and bottom images show regions with statistically significant correlation with the y and z isocenter positions, respectively. Red arrows point to corresponding voxel locations where local automated displacement is plotted against isocenter position. The perpendicular black arrows mark the location of the in-plane isocenter in MNI152 space. Positive correlation is observed with a more anterior (increasing y) and superior (increasing z) position. Negative correlation is observed with a more inferior (decreasing $z$ ) isocenter position. Note that there is no significant correlation in a region of the orbitofrontal cortex shown to have high displacement in Figure 4.3, a region known to be prone to susceptibility effects.
inferior lateral occipital cortex, bilateral frontal medial cortex, bilateral subcallosal cortex, bilateral anterior parahippocampal gyri, bilateral anterior temporal fusiform cortex, and bilateral occipital poles. None of the subcortical regions defined on the ATAG subcortical atlas were significantly displaced. Maximum displacements around common stereotactic targets, including the substantia nigra and subthalamic nucleus, were $1-2 \mathrm{~mm}$ on average across subjects. The bilateral striatum ROIs, which were the largest subcortical structures in the ATAG atlas, showed the highest maximum displacement in the 2-3 mm range.

Twenty-two regions were identified that demonstrated significant correlations of isocenter position with local displacement as shown in Table 4.3 (for the complete table, see Table S2 in Appendix Section B.1). Most significant correlations were related to the y or z components

Table 4.2: Mean and maximum displacements in millimeters for ROIs from the Harvard-Oxford atlas meeting thresholds for statistical significance.

| region | side | mean | stdev | V | max | max stdev |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Left Temporal Pole | left | 1.18 | 0.28 | 217 | 5.16 | 1.57 | $*$ |
| Right Temporal Pole | right | 1.17 | 0.24 | 213 | 4.87 | 1.50 | $*$ |
| Left Middle Temporal Gyrus Anterior | left | 1.54 | 0.67 | 228 | 3.58 | 1.34 | $*$ |
| Left Inferior Temporal Gyrus Posterior | left | 1.73 | 0.36 | 252 | 5.34 | 0.84 | $*$ |
| Right Inferior Temporal Gyrus Posterior | right | 1.56 | 0.34 | 253 | 5.05 | 0.82 | $*$ |
| Left Lateral Occipital Cortex Inferior | left | 1.31 | 0.26 | 248 | 3.62 | 1.32 | $*$ |
| Right Lateral Occipital Cortex Inferior | right | 1.28 | 0.18 | 253 | 3.45 | 0.57 | $*$ |
| Left Frontal Medial Cortex | left | 1.75 | 0.59 | 251 | 5.68 | 1.12 | $*$ |
| Right Frontal Medial Cortex | right | 1.84 | 0.56 | 253 | 5.77 | 0.94 | $*$ |
| Left Subcallosal Cortex | left | 1.72 | 0.33 | 252 | 6.87 | 0.67 | $*$ |
| Right Subcallosal Cortex | right | 1.80 | 0.38 | 253 | 6.87 | 0.62 | $*$ |
| Left Parahippocampal Gyrus Anterior | left | 1.51 | 0.88 | 244 | 3.84 | 1.89 | $*$ |
| Right Parahippocampal Gyrus Anterior | right | 1.40 | 0.35 | 249 | 4.00 | 1.27 | $*$ |
| Left Temporal Fusiform Cortex Anterior | left | 1.62 | 0.85 | 247 | 3.77 | 1.85 | $*$ |
| Right Temporal Fusiform Cortex Anterior | right | 1.59 | 0.57 | 247 | 3.52 | 1.13 | $*$ |
| Left Occipital Pole | left | 1.64 | 0.65 | 248 | 4.33 | 1.95 | $*$ |
| Right Occipital Pole | right | 1.55 | 0.46 | 252 | 4.01 | 1.08 | $*$ |
| V |  |  |  |  |  |  |  |

$\overline{\mathrm{V}}$ represents the Wilcoxon rank sum statistic, p-value is unadjusted, and q-value is FDR adjusted. * All 13 regions shown met thresholds for statistical significance after FDR correction at a rate of $q<0.025$. For the full table, please see Appendix Section B.1.

Table 4.3: Correlation of displacements with change in position of image isocenter for regions meeting thresholds for statistical significance.

| region | side | atlas | component | corr |  |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Right Subcallosal Cortex | right | HarvardOxford | x | -0.675 | $*$ |
| Left Occipital | left | lobar | y | 0.848 | $*$ |
| Right Occipital | right | lobar | y | 0.767 | $*$ |
| Left Inferior Frontal Gyrus Pars Opercularis | left | HarvardOxford | y | 0.71 | $*$ |
| Left Intracalcarine Cortex | left | HarvardOxford | y | 0.696 | $*$ |
| Right Intracalcarine Cortex | right | HarvardOxford | y | 0.716 | $*$ |
| Left Lingual Gyrus | left | HarvardOxford | y | 0.832 | $*$ |
| Right Lingual Gyrus | right | HarvardOxford | y | 0.802 | $*$ |
| Left Occipital Fusiform Gyrus | left | HarvardOxford | y | 0.772 | $*$ |
| Right Occipital Fusiform Gyrus | right | HarvardOxford | y | 0.795 | $*$ |
| Left Supracalcarine Cortex | left | HarvardOxford | y | 0.646 | $*$ |
| Right Supracalcarine Cortex | right | HarvardOxford | y | 0.795 | $*$ |
| Left Occipital Pole | left | HarvardOxford | y | 0.837 | $*$ |
| Right Occipital Pole | right | HarvardOxford | y | 0.85 | $*$ |
| Right Parietal | right | lobar | z | -0.724 | $*$ |
| Right Postcentral Gyrus | right | HarvardOxford | z | -0.79 | $*$ |
| Right Superior Parietal Lobule | right | HarvardOxford | z | -0.869 | $*$ |
| Left Intracalcarine Cortex | left | HarvardOxford | z | 0.68 | $*$ |
| Right Parahippocampal Gyrus Posterior | right | HarvardOxford | z | 0.669 | $*$ |
| Left Lingual Gyrus | left | HarvardOxford | z | 0.7 | $*$ |
| Right Lingual Gyrus | right | HarvardOxford | z | 0.688 | $*$ |
| Right Occipital Fusiform Gyrus | right | HarvardOxford | z | 0.695 | $*$ |

The corr values represent the correlation of the corresponding $x, y$, or $z$ component of the 7 T image isocenter with mean automated displacement in that region-of-interest, p -value is unadjusted, and q -value is FDR adjusted at a rate of $\mathrm{q}<0.025$. * 22 regions met thresholds for statistical significance after FDR correction. For the full table, please see Appendix Section B.1.
of the isocenter, except for the right subcallosal cortex which correlated with x. At the lobar level, bilateral occipital and right parietal regions were correlated with isocenter position. At the sub-lobar level, regions identified as correlating with the y position of isocenter were: left pars opercularis, bilateral intracalcarine cortex, bilateral lingual gyri, bilateral occipital fusiform gyri, bilateral supracalcarine cortex, and bilateral occipital poles. With z position, the following regions were identified: right postcentral gyrus, right superior parietal lobule, left intracalcarine cortex, right parahippocampal gyrus posterior, bilateral lingual gyrus, and right occipital fusiform gyrus.

### 4.3.5 Validation

Our results were validated using manual fiducial placement on 5 subjects. Successive stages of registration from rigid body to affine to nonlinear registration resulted in mean fiducial error changes from $1.09 \pm 0.77$ to $0.98 \pm 0.69$ to $0.98 \pm 0.67 \mathrm{~mm}$, respectively. Trends toward improvements in fiducial registration error were generally observed with successive stages of registration in Table 4.4. However, after adjusting p-values for multiple comparisons, only the fiducial registration error in the optic chiasm landmark reached statistical significance comparing rigid to nonlinear registration ( $1.61 \pm 0.59$ to $0.88 \pm 0.44 ; q$-value $<0.01$ ) .

Intra-rater reliability was $0.86 \pm 0.35 \mathrm{~mm}$. Inter-rater reliability was $1.14 \pm 0.77 \mathrm{~mm}$. Our automated method demonstrated excellent correlation ( $\mathrm{R}=0.8755$, p -value $<0.00$ ) and agreement on Bland-Altman plotting (Figure 4.5) with manual displacement.

### 4.3.6 Effect of Gradient Distortion Correction

We compared the results using our automated framework with and without vendor-provided gradient distortion correction in a single subject (Figure 4.6 and Table S3 in Appendix Section B.1). The percentage of voxels with displacement greater than 1 mm dropped from $31.9 \%$ without correction to $20.4 \%$ with correction. The mean whole brain displacement decreased from 0.548 mm to $0.472 \mathrm{~mm}-\mathrm{a} 13.8 \%$ reduction in distortion. The density plot was leftshifted with a narrower right-sided tail suggesting fewer geometric outliers. We have included Table S3 (Appendix Section B.1) summarizing the trend of improvements across ROIs, with particularly large reductions in the bilateral parietal and occipital lobes (up to almost $60 \%$ reduction in displacements).

Table 4.4: Fiducial registration error (in mm ) with successive registration steps in different categories.
$\left.\begin{array}{lllllll}\hline \hline \text { region } & \text { rigid } & \text { stdev } & \text { affine } & \text { stdev } & \text { nlin } & \text { stdev } \\ \hline \hline \text { total } & 1.09 & 0.77 & 0.98 & 0.69 & 0.98 & 0.67 \\ \text { temporal } & 0.96 & 0.61 & 0.85 & 0.62 & 0.94 & 0.71 \\ \text { occipital } & 1.26 & 1.13 & 1.02 & 0.83 & 0.78 & 0.42 \\ \text { parietal } & 1.22 & 0.77 & 1.02 & 0.61 & 0.96 & 0.62 \\ \text { frontal } & 0.91 & 0.45 & 0.76 & 0.37 & 0.86 & 0.52 \\ \text { insular } & 0.91 & 0.58 & 0.80 & 0.48 & 0.67 & 0.43 \\ \text { subcortical } & 0.91 & 0.53 & 0.84 & 0.52 & 0.88 & 0.51 \\ \text { midline total } & 0.99 & 0.55 & 0.88 & 0.43 & 0.86 & 0.47 \\ \text { AC } & 0.80 & 0.45 & 0.66 & 0.38 & 0.66 & 0.39 \\ \text { PC } & 0.55 & 0.22 & 0.59 & 0.25 & 0.60 & 0.24 \\ \text { midpoint of aqueduct } & 0.89 & 0.39 & 1.03 & 0.46 & 1.07 & 0.47 \\ \text { optic chiasm } & 1.61 & 0.59 & 1.19 & 0.46 & 0.88 & 0.44\end{array} *\right\}$

Fiducial registration error was assessed for rigid body, affine (12-parameter), and nonlinear registration steps. Fiducials classified as displacement represent selected locations that were chosen retrospectively at areas identified as displaced using the automated method. All interactions were assessed using Wilcoxon rank sum statistics (i.e. two-tailed test for fiducial error with rigid versus affine, rigid versus nonlinear, and affine versus nonlinear registration). * Only one test (rigid versus nonlinear registration for the optic chiasm) met thresholds for statistical significance after FDR correction (q-value $<0.01$ ).


Figure 4.5: (a) Plot of automated displacement against gold-standard manual displacement demonstrated good correlation ( $\mathrm{R}=0.8755$, p -value $<0.00$ ). (b) Bland-Altman plotting reveals that automated displacement slightly underestimates manual displacement (mean $=-0.193 \mathrm{~mm}$ ) with good agreement between manual and automated measures. Gray error bands represent $95 \%$ confidence intervals.

### 4.4 Discussion

The principal results of this study can be summarized as follows. Focal areas of geometric distortion in higher fields can be identified automatically from the deformation field used in nonlinear registration to images acquired at lower fields. We evaluated our automated method on a T1-weighted MRI dataset, the MRI sequence predominantly used as an anatomical reference for structural and functional studies. Distortions were increased at tissue interfaces, particularly the floor of the middle and anterior fossae, but also in the occipital/suboccipital region (Figure 4.1 and 4.3; Table 4.2). Distortions were also increased with distance from the image isocenter to over 1 mm at 80 mm distance from the isocenter (Figure 4.2b). Distortions were correlated with the image isocenter location, particularly in the parietal, occipital, and suboccipital regions, but generally not at tissue interfaces (Figure 4.4; Table 4.3). Traditional deep brain surgery targets including the STN and GPi were minimally displaced (Table 4.1). Geometric distortion correction resulted in a mean reduction in distortion of $13.8 \%$. Finally, our automated framework compared favorably to manual fiducial placement (Figure 4.5). Moving forward, our methodology can be used to evaluate other 7T protocols, including T2weighted [Kwon et al., 2012], susceptibility-weighted [Abosch et al., 2010, Chandran et al., 2016, Schäfer et al., 2012], and T1 inversion recovery images [Sudhyadhom et al., 2009, Tour-


Figure 4.6: The effect of vendor-provided gradient distortion correction on local displacement in a single subject. A density map of automated displacements within the masked brain with (solid line) and without (dashed line) distortion correction applied (a). The mean displacement is decreased from 0.548 mm to 0.472 mm with distortion correction (a percent reduction of $13.8 \%$ ). Automated displacement maps are overlaid on the subject's own 3 T structural scan (b). Decreased displacement is visualized throughout the scan particularly in the suboccipital region. Distortions identified near air-filled sinuses (floor of the middle fossa and orbitofrontal cortex) remain.
dias et al., 2014].
Phantom validation represents an important paradigm for the calibration of MRI scanners and the assessment of geometric distortion. In a phantom comprised of regularly spaced acrylic rods mounted on an epoxy-based skull base, Cho and colleagues showed sub-millimeter accuracy between manually placed fiducials at 7T and on CT [Cho et al., 2010]. Using a proprietary phantom, Dammann et al. [Dammann et al., 2011] found hardware-related geometric distortion generally smaller than 1 mm within a range of 80 mm from the magnet isocenter, consistent with our own observations (Figure 4.2b), except in a T2-weighted sequence. However, results have been inconsistent with other studies suggesting more severe distortions in the several millimeter range [Littmann et al., 2006, Neumann et al., 2015, Watanabe et al., 2006]. Another practical point is that some of the commonly used MRI phantoms [Fonov et al., 2010] do not fit within the field-of-view in head-specific gradient coils. Finally, phantom validation captures gradient nonlinearities [Fonov et al., 2010], but would not be able to capture physiological distortion resulting from magnetic susceptibility at tissue interfaces that are subject-specific.

In vivo studies are better able to evaluate physiological distortion not captured using MRI phantoms. One of the early studies looking at the feasibility of 7T for stereotaxy analyzed subjects scanned at both 1.5 T and 7 T [Duchin et al., 2012]. Using manual segmentation, the
authors observed that fiducials placed in central brain regions were minimally distorted, except for the optic chiasm and infundibulum landmarks abutting the anterior fossa, as we have also confirmed (Figure 4.1, Table 4.3). By dividing the MRI volume into 9 block-shaped regions arranged about the image isocenter, they demonstrated larger distortions proximal to air-filled cavities (inferior frontally). Our methodology expands on the original work by Duchin and colleagues by providing an automated framework that compared favorably with gold-standard manual measures in terms of both correlation and agreement (Figure 4.5). Our own observations are compatible with their work (Figure 4.1 and 4.3; Table 4.2), while providing additional neuroanatomical detail about focal regions of spatial uncertainty at the voxel-wise and regional scales that can be examined prospectively in patient-specific datasets (Figure 4.3; Table 4.14.2). Given the known linear increase in magnetic susceptibility with increasing $B_{0}$, we would expect over 2 x increase in susceptibility artifact from 3 T to 7 T for an equivalent MRI sequence. When isolating the contribution of distortion in the main Cartesian directions, we found that voxels identified as significantly distorted had a predominant z -axis deviation in $56.84 \%$ of voxels, which may result from $B_{0}$-related distortions, consistent with prior work [Neumann et al., 2015, Watanabe et al., 2006].

Geometric distortion can be corrected using a number of different methods. One common technique is to apply spherical harmonics correction based on gradient coil geometry, derived from the design parameters of the coil itself. Langlois et al. demonstrated that applying spherical harmonics correction alone decreased error to the sub-millimeter range in a phantom study at 1.5T [Langlois et al., 1999]. It remains unclear how much spherical harmonics correction improves geometric inhomogeneity at ultra-high field. A recent study observed that applying vendor-provided spherical harmonics correction resulted in higher stereotactic error than without, raising the potential concern that inappropriate use of these algorithms can worsen geometric accuracy [Neumann et al., 2015]. When the spherical harmonics information is not directly available, post hoc phantom-based correction has been found to improve geometric accuracy and reproducibility in multi-site datasets [Jovicich et al., 2006]. However, the estimates on the percent variability accounted for by phantom-based correction are broad, ranging from $17-90 \%$ depending on the study [Caramanos et al., 2010, Fonov et al., 2010, Jovicich et al., 2006]. Altogether, there remains no gold-standard geometric distortion correction algorithm. In a single dataset with and without gradient correction, we demonstrated a promising mean reduction of $13.8 \%$ in our distortion estimate with reductions up to $60 \%$ in the occipital lobes (Figure 4.6 and Table S3). The only exceptions were in regions close to air-filled cavities-where the main cause of distortion is likely related to $B_{0}$ susceptibility rather than gradient inhomogeneity. A complementary approach for $B_{0}$-related distortion correction has recently gained attention as a preprocessing step in the Human Connectome Project (HCP)
[Glasser et al., 2013, van der Kouwe et al., 2008]. The acquired $B_{0}$ field map is scaled by the readout dwell time, and used as a distortion field. This methodology, called field map (or readout) correction has been traditionally applied to echo planar images [Jezzard and Balaban, 1995], but more recently in the HCP, has been used to correct high-resolution structural MR images [Glasser et al., 2013] with the presumption that correction with a low-resolution map is still accurate because $B_{0}$ inhomogeneity is smoothly varying. Qualitative improvements with $B_{0}$ field map correction have been demonstrated, but one possible application of our pipeline is to better quantify the impact of field map correction on residual distortion in gradient-corrected images. Overall, the framework described in this manuscript can be applied generally to evaluate the effect of different correction methods more systematically.

No significant local displacement in the ATAG subcortical atlas was observed, which included STN and GPi regions (Table 4.1). Most of the midline fiducials were minimally displaced, except for the optic chiasm, which was displaced at $1.61 \pm 0.59 \mathrm{~mm}$ with rigid registration alone, improving significantly with nonlinear registration (Table 4.4). Consistent with prior literature, this landmark is likely prone to distortion from magnetic susceptibility at the floor of the anterior fossa proximal to similarly affected inferior frontal regions in our automated analysis (Figure 4.3, Table 4.2), and supported by previous work [Duchin et al., 2012].

Our results demonstrate that local displacement increased with distance from the image isocenter with greater than 1 mm of displacement on average beyond 80 mm (Figure 4.2b). Furthermore, the data are not normally distributed, and researchers and clinicians should be aware of geometric inaccuracies in these outlier regions, particularly with critical applications like stereotactic surgery. In particular, we found that inferior frontal, parasagittal, and occipital cortices were distorted. While these regions are outside traditional zones for neurosurgical intervention, the number of putative neuromodulation targets continues to grow, and one of the more well-studied targets for refractory depression, the subgenual or subcallosal cingulate cortex [Johansen-Berg et al., 2008, Lozano et al., 2008], lies within a region of high distortion (Figure 4.3, Table 4.2). Previous studies have shown that variability in the location of the image isocenter can have an effect on study-related morphometric measures [Caramanos et al., 2010, Jovicich et al., 2006]. While the variability in isocenter placement at both 3T and 7T is quite high in our study, particularly in the z-direction, these results are similar to findings in multi-site structural MRI studies [Jovicich et al., 2006]. By correlating the position of the isocenter with local displacement, we identified the superior parietal region and suboccipital regions as being particularly prone to isocenter displacement. Regions close to air-filled sinuses were not affected in the same way thus suggesting $B_{0}$ may be the likely reason for distortion in these regions. Finally, isocenter discrepancies also highlight important differences between 7T and 3T imagers that result in large differences in isocenter location. At 3T, the isocenter
is typically lower in the z direction with a lower standard deviation since externally, we are aiming for the nasion. At 7T, the magnet and coil are at the extreme end of the table, and thus, we are more limited in our ability to use external landmarks like the nasion for more consistent isocenter placement. Overall, in order to limit large geometric inaccuracies including gradient distortions, our results suggest that a researcher interested in a particular region should place it as close to the 7 T isocenter as possible.

There are several limitations to our work. Our workflow requires nonlinear registration of participant data acquired from a target image at ultra-high field to a reference image at lowerfield (in this case, 3T). We have assumed that the 3T dataset is relatively free of geometric inhomogeneity, even though distortion is known to exist at lower fields [Caramanos et al., 2010, Jovicich et al., 2006]. Despite this, recent clinical studies have suggested a comparable safety profile using 3 T MR-based images for surgical interventions compared with 1.5 T [Cheng et al., 2014, Houshmand et al., 2014]. Certain special precautions including the use of high receiver bandwidth and vendor-provided gradient distortion correction techniques can help to ensure 3T images are geometrically more accurate. Computed tomography (CT) data, which were unavailable for this cohort of subjects, can be considered a gold-standard for geometric evaluation. However, the lack of tissue contrast could result in difficulties with nonlinear registration with MRI. A recent systematic review established that CT/MRI fusion is not trivial with most studies reporting over 1 mm of registration error [Geevarghese et al., 2016, Thani et al., 2011]. Variability in the image isocenter may also be an important confound in our dataset, although our results are similar to data from multi-center anatomical trials [Jovicich et al., 2006]. It is possible that distortion near air-filled sinuses may be an artifact of degraded image quality in these regions [Tardif et al., 2015] rather than true distortion, but nevertheless, the algorithm captures the potential spatial uncertainty in these regions which should be accounted for in neuroimaging studies and surgical planning. Finally, while our project describes focal estimates of geometric distortion at a voxel-level, these results are specific to the scanner and sequence assessed, as well as the gradient coil used.

### 4.5 Conclusions

Our automated framework quantifies, in millimeters, the extent of geometric distortion resulting from ultra-high field structural MRI at the group and subject levels. These results have important implications for evaluating patient-specific local spatial uncertainty. We have demonstrated that our method quantifies voxel-wise distortion and in addition, can help characterize the source of inhomogeneity by observing interactions of our measure with isocenter location and gradient distortion correction methods. Our results point to the need for caution
if using ultra-high field MRI for purposes where morphological accuracy is critical, including stereotactic interventions, as well as the need for scanner and sequence-specific calibration of geometric inhomogeneity. We confirmed that our automated method compared favorably with manual fiducial displacement, thus permitting prospective evaluation of the effect of MRI sequences, putative correction algorithms, and scanner modifications on geometric distortion.

## Chapter 5

Direct visualization and characterization of the human zona incerta and surrounding regions

This chapter is work being prepared for submission.

- Lau, J. C., Parrent, A. G., Xiao, Y., Gilmore, G. G., Demarco, J., MacDougall, K. W., Currie, C., Peters, T. M., \& Khan, A. R. (2019). Direct visualization and characterization of the human zona incerta and surrounding regions.


### 5.1 Introduction

The zona incerta $(\mathrm{ZI})$ is a small but diffuse structure in the deep brain first identified by Auguste Forel in 1877, famously described as "an immensely confusing area about which nothing can be said" [Forel, 1877]. Forel appreciated that the ZI consisted of gray matter located between the external medullary lamina of the thalamus and the corpus Luysi (subthalamic nucleus; STN) of otherwise "indefinite" description. It is particularly telling that Forel found the ZI so difficult to describe given his crucial role in the careful delineation of surrounding fibre tracts still often referred to eponymously as the fields of Forel [Gallay et al., 2008]. Since its original description, much has been learned about the ZI and its surrounds although robust in vivo visualization has remained elusive.

The anatomical boundaries of the ZI have generally been described in the context of its more discrete neighbours rather than based on any consistent feature of the region itself. Packed in a small area between the ventral thalamus, STN, and lateral red nucleus (RN), the ZI is situated at a complex junction of major white matter pathways including the cerebellothalamic, pallidothalamic, medial lemniscal, and corticospinal tracts. Along its dorsal, ventral, and medial borders, the ZI is surrounded by the fasciculus thalamicus ( ft ; also known as the H 1 field of Forel), the fasciculus lenticularis ( fl ; also known as the H 2 field of Forel), and the H field, which is a convergence of the fl and the ansa lenticularis (al), respectively [Nieuwenhuys et al., 2007, Gallay et al., 2008]. The rostral ZI (rZI) is continuous with the reticular nucleus of the thalamus laterally and with the lateral hypothalamus anteromedially. The caudal ZI (cZI) is laterally bounded by the STN and posterior limb of the internal capsule. To date, most of the details regarding the region are the result of meticulous study of carefully prepared postmortem specimens [Schaltenbrand and Wahren, 1977, Morel, 2007, Gallay et al., 2008].

Cytoarchitectonic and myeloarchitectonic studies have identified the ZI as a nuclear complex consisting of loosely arranged neurons of heterogeneous morphology with a diverse immunohistochemical profile [Nieuwenhuys et al., 2007]. In Golgi preparations of the ZI, two main neuronal classes have been identified: principal cells and interneurons [Ma et al., 1997]. Gene expression studies have revealed a common embryological origin along with the reticular nucleus of the thalamus and pregeniculate nucleus of the ventral diencephalon, specifically the prethalamic (prosomere 3) segment, which predominantly contains gabaergic neurons [Wat-
son et al., 2014]. Through immunohistochemical analysis in experimental animals, a general pattern of at least four component ZI sectors has emerged in the rostral, dorsal, ventral, and caudal directions [Mitrofanis, 2005]. Tract-tracing studies have identified extensive and often bilateral connections between the ZI and the cortex, subcortex, and spinal cord [Mitrofanis, 2005, Watson et al., 2014]. At least five functional subsectors within the ZI have been suggested: auditory, limbic, motor, somatosensory, and visual. However, unlike other nearby structures like the STN, no robust immunohistochemical biomarker has been described for the ZI proper.

The diversity of chemical expression and widespread connections suggest an important modulatory role of the zona incerta in regulating brain function. The zona incerta forms extensive inhibitory connections with spinothalamic relay nuclei in experimental animals, and thus may play an important role in modulating neuropathic pain and the somatosensory system [Masri et al., 2009, Truini et al., 2013]. In a perhaps related manner, the rostral ZI is believed to provide inhibitory control over the thalamus during sleep [Llinás and Jahnsen, 1982, Watson et al., 2014], which may also relate to its perceived role in modulating consciousness [Power et al., 1999, Power and Mitrofanis, 1999, Mitrofanis, 2005, Giacino et al., 2014].

In humans, the most well-studied role of the zona incerta is as a putative target for neuromodulatory therapy transmitted either within the caudal zona incerta (cZI) or its vicinity, which has been observed to be highly effective for the treatment of essential tremor [Hariz and Blomstedt, 2017]. These investigations began in the 1960s with leucotomy (selective white matter ablation) treatments [Spiegel and Wycis, 1954, Wertheimer et al., 1960, Spiegel et al., 1962, Spiegel et al., 1964, Mundinger, 1965, Bertrand et al., 1969, Velasco et al., 1975], but as technologies improved, electrical stimulation to these regions has also been demonstrated to be effective [Mohadjer et al., 1990, Velasco et al., 2001, Nowacki et al., 2018, Velasco et al., 1972, Plaha et al., 2006, Blomstedt et al., 2010, Fiechter et al., 2017]. Yet because of poor direct visualization, controversy has remained as to whether the therapeutic effect is derived from modulation of the cell bodies in the cZI, wayward connections such as the cerebellothalamic tracts (also known as the prelemniscal radiations or raprl) [Velasco et al., 1972, Castro et al., 2015] or some combination of both [Blomstedt et al., 2010]. Given the ambiguity and high functional density of the region, some prefer to consider the stereotactic target more broadly as the posterior subthalamic area (PSA) [Hariz and Blomstedt, 2017, Blomstedt et al., 2018, Nowacki et al., 2018]. Targeting of the region relies on identification of the PSA indirectly relative to the adjacent subthalamic nucleus (STN) and red nucleus (RN), which are visible on T2-weighted scans [Blomstedt et al., 2010] (see Section 5.2.6 for more details).

The increased inherent signal resulting from increasing magnetic field strength has presented an opportunity to visualize brain structures that have not been seen at lower field

Table 5.1: MRI sequence details.

| Sequence |  | TE $(\mathrm{ms})$ | TR $(\mathrm{ms})$ | TI | Flip Angle $\left({ }^{\circ}\right)$ | Matrix Size | PAT* $^{*}$ | Averages | Resolution | Acquisition Time (min:sec) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| MP2RAGE | 3D | 2.73 | 6000 | $800 / 2700$ | $4 / 5$ | $342 \times 342 \times 224$ | 3 | 1 | $0.7 \times 0.7 \times 0.7$ | $10: 14$ |
| SA2RAGE | 3D | 0.81 | 2400 | $45 / 1800$ | $4 / 11$ | $128 \times 128 \times 64$ | 2 | 1 | $1.9 \times 1.9 \times 2.1$ | $2: 28$ |
| SPACE | 3D | 398 | 4000 | NA | variable | $320 \times 320 \times 224$ | 3 | 1.6 | $0.7 \times 0.7 \times 0.7$ | $10: 28$ |

* PAT = parallel acquisition technique (acceleration factor)
strengths [Marques and Norris, 2017]. By exploiting T2-related tissue properties, ultra-high magnetic field (7 Tesla; 7T) MRI has enabled direct visualization of many deep brain nuclei with increased resolution and signal-to-noise ratio (SNR) including the red nucleus, substantia nigra, and subthalamic nucleus [Schäfer et al., 2012, Keuken et al., 2013, Plantinga et al., 2018], known to be rich in iron [Zecca et al., 2004, Haacke et al., 2005]. Paralleling these successes, previous attempts at direct visualization of the zona incerta have focussed on the use of T2-based contrast, with purported identification of the rZI, but not the cZI [Kerl et al., 2013]. In this study, we report that by employing high-resolution quantitative T1 mapping at 7-Tesla, robust visualization of the zona incerta is now possible in vivo along the entire rostrocaudal axis, enabling comprehensive anatomical characterization of this previously obscure deep brain region.


### 5.2 Materials and Methods

### 5.2.1 Participant and image acquisition details

We recruited 32 cognitively intact, healthy participants ( $46.2 \pm 13.5$ years; median: 48 years; range: 20-70 years; 12 female and 20 male; right-handed). This study was approved by the Western University Health Sciences Research Ethics Board. All subjects signed a written consent form to participate. The imaging studies were performed in a 7-Tesla head-only scanner (Siemens Magnetom; Siemens Healthcare GmbH, Erlangen, Germany) at the Western University Centre for Functional and Metabolic Mapping (CFMM). An 8-channel parallel transmit/32-receive channel coil was used [Gilbert et al., 2011]. After localization and preparatory sequences, each subject underwent a 3D MP2RAGE [Marques et al., 2010], 3D SA2RAGE [Eggenschwiler et al., 2012], and 3D optimized fast-spin echo (T2 SPACE) acquisitions (see Table 5.1).

### 5.2.2 Image pre-processing and template creation

Upon completion of an MRI scan session, the images were pushed to a DICOM server (dcm4che; https://www.dcm4che.org) with automatic data standardization and conversion to the Brain

Imaging Data Structure (BIDS) [Gorgolewski et al., 2016] using the autobids platform (https : //github.com/khanlab/autobids) deployed on a high-performance compute cluster. Autobids utilizes scanner-specific heuristics enabled by heudiconv (https://github.com/nipy/ heudiconv) preconfigured and validated on multiparametric 7T MRI sequences for DICOM to nifti conversion using dem2niix [Li et al., 2016] and organization into BIDS.

All individual MRI sequences were corrected for gradient nonlinearities using 3D distortion correction [Glasser et al., 2013, Lau et al., 2018a] prior to further processing. The objective of individual preprocessing steps was to adequately prepare the individual MRI sequences for quantitative image analysis and also linear alignment with the subject's T1-weighted structural MRI scan containerized as BIDS apps [Gorgolewski et al., 2017]. The outputs of the preprocessing steps were visually assessed for quality (JL).

### 5.2.3 Pre-processing: MP2RAGE

As part of the MP2RAGE acquisition, two different images were created at separate inversions. Using a lookup table, these inversion images were used to create synthetic quantitative T1 maps devoid of proton density contrast, reception field bias, and first order transmit field inhomogeneity. Minimal pre-processing was necessary except for using the $B_{1}$ field map (SA2RAGE) sequence to correct for intensity inhomogeneity [Eggenschwiler et al., 2012]; specifically, no post hoc intensity nonuniformity correction was employed. This SA2RAGE-corrected T1 map was used for quantitative analysis. The T1w image was used as a reference image for rigidbody alignment of the T2SPACE scan.

### 5.2.4 Pre-processing: T2SPACE

Raw images from the scanner were observed to have significant intensity inhomogeneities. The bias fields were corrected using an initial nonuniformity correction step with N 4 [Sled et al., 1998, Tustison et al., 2010] enabling more accurate registration of the T1w image (and associated brain mask) to T2w. A synthetic T1-T2w fusion image was created by multiplying the T1w by the T2w image [Xiao et al., 2014] and re-estimating the intensity inhomogeneity again with N4. The original T2w image was denoised using the adaptive non-local means method [Manjón et al., 2010] and the obtained inhomogeneity estimation was applied to the denoised image resulting in a final preprocessed T2w image in the scanner space. Rigid registration to the T1w scan was re-estimated using the preprocessed image. Final preprocessed images included both a T 2 w volume in the original scanner space as well as one resampled into the T1w structural space. The process was bootstrapped once after creating an initial T2w template (see Section 5.2.5 Template Creation) and using the template for histogram-
based intensity normalization. Note that because of the combination of post hoc bias field correction and intensity normalization necessary to produce more homogeneous images, the per voxel values of the T2SPACE images are not directly comparable between scans in a quantitative manner. This processing pipeline has been released and containerized as a BIDS app (https://github.com/khanlab/prepT2space/).

### 5.2.5 Template creation

The antsMultivariateTemplateCreation2 pipeline was used for multimodal (T1,T2) template creation [Avants et al., 2011]. A corresponding T2w template (in T1w space) was created after propagating the participant T 2 w images to T 1 w template space using the relevant transformations produced using prepT2space. An initial template was created using rigid body alignment of each participant's T1w scan to the MNI2009bAsym template ( 0.5 mm isotropic resolution) [Fonov et al., 2009]. Over a series of 10 subsequent bootstrapped iterations, the deformable registration was refined (shrink factors: $12 \times 6 \times 4 \times 2 \times 1$; smoothing factors: $6 \times 3 \times 2 \times 1 \times 0 \mathrm{vox}$; max iterations: 100x100x70x50x10; transformation model: Greedy SyN; similarity metric: crosscorrelation). Using the derived affine and nonlinear transforms, the individual sequences (T1map and T2) were transformed and resampled using trilinear interpolation into the template space. Mean intensity images were generated for each parametric sequence. The log Jacobian was computed, providing an estimate of local deformation required to transform each participant into the template space. The scripts for template creation have been archived for reference (https://github.com/jclauneuro/snsx32/). Spatial correspondence was quantified using a recently described anatomical fiducial (AFID) placement protocol by three trained raters who placed 32 AFIDs on the original scanner space for the T1w images using 3D Slicer [Fedorov et al., 2012], so that anatomical fiducial localization error (AFLE; Euclidean distance between point placements) could be calculated. This permitted the computation of residual AFID registration error (AFRE) to be calculated across these same features [Lau et al., 2018b] (RRID:SCR_016623) to quantify the accuracy of template creation, which is the subject of Chapter 3.

### 5.2.6 Region-of-interest segmentation

The zona incerta, subthalamic nucleus, and substantia nigra were segmented using the 10th iteration T1 and T2 combined template using ITK-SNAP version 3.6.0. Each rater segmented the regions twice, with sessions spaced more than two weeks apart. A representative template segmentation was derived by averaging all segmented ROIs and thresholding by majority voting (>50\%) - this was considered the "gold" standard. Three raters segmented the RN
and STN twice using the T2 image (JD, JL, YX). The pZI, rZI, cZI, raprl, and the ft were segmented twice by two raters (GG, JL) using the T1 map image. To our knowledge, the ZI has not been previously segmented from in vivo images. As such, two stereotactic neurosurgeons (AP, KM) were consulted throughout the ZI segmentation process: first, after the initial segmentations by the lead author (JL); second, after identifying critical boundaries of the ZI particularly rostrally; and finally, to review the final consensus segmentation. Several histological human brain atlases were used as references [Schaltenbrand and Wahren, 1977, Morel, 2007, Hawrylycz et al., 2012].

The rostral ZI presented some challenges to accurate identification, not for lack of contrast, but due to difficulty with determining its relationship with the fl and ft. On closer review, we speculate that the fl actually runs through the rostral portion of the ZI. We provide labels for the ZI as a whole, and provide separate labels for the dorsal rZI, vZI, interposed fl, and cZI. The lateral aspect of the central portion of the ZI (between rostral and caudal ends) was too thin to segment along its entire length even at 7 T .

### 5.2.7 Stereotactic target localization

Target locations in the bilateral posterior subthalamic area were placed according to the placement scheme of Nowacki and colleagues [Nowacki et al., 2018] (Figure 5.3). This scheme relies on anatomical targeting based on axial T2-weighted images after performing an initial AC-PC transformation with placement at the center of each commissure. This target involves the identification of three different lines: a horizontal line drawn along the equator of the RN identified on the axial slice of maximal diameter, an oblique line drawn along the long-axis of the STN, and finally, an oblique line perpendicular to the long-axis of the STN intersecting the lateral border of the RN at its equator. The placements were reviewed via consensus with two neurosurgeons who practice stereotactic neurosurgery (AP, KM). Optimal target location was also investigated prospectively in a patient with essential tremor who had undergone cZI implantation with adequate long-term follow-up and Essential Tremor Rating Scale (ETRS) recorded.

### 5.3 Results

### 5.3.1 Template Creation

Mean AFLE after quality control was $0.91 \pm 0.69 \mathrm{~mm}$. Visual inspection of AFIDs revealed good spatial correspondence as well as evidence of good anatomical detail in subcortical struc-


Figure 5.1: Visual inspection of AFIDs revealed good convergence with successive template generation steps showing the baseline correspondence between images followed by the linear only template and finally the combined (linear and nonlinear) template after 10 iterations of template building. The information here is corroborated in Tables 5.2 and 5.3.
tures of the final template (Figure 5.1). Mean AFRE decreased globally across all AFID points with registration complexity improving from $29.62 \pm 11.58 \mathrm{~mm}$ at baseline to $3.47 \pm 1.62 \mathrm{~mm}$ after linear registration and to $2.80 \pm 0.90 \mathrm{~mm}$ after deformable registration (one iteration). Further increases were noted with successive iterations of template generation to $2.06 \pm 0.92$ mm after 4 iterations to $1.27 \pm 1.02 \mathrm{~mm}$ after 10 iterations. Mean AFRE improved to a limit of $2.82 \pm 1.47 \mathrm{~mm}$ with linear registration alone with little improvement beyond 6 iterations, while improvements were noted up to 10 iterations with deformable registration (see Table 5.3 and Figure 5.1).

Several AFID features were particularly difficult to register, with registration errors higher than the expected manual anatomical fiducial localization error. In particular, we identified several features with more than 6 mm of registration error: the lateral mesencephalic sulci, temporal horns, and origins of the indusium griseum bilaterally (Figure 5.1 and Table 5.3).

### 5.3.2 Direct visualization and segmentation of the zona incerta region

We pooled submillimetric ( 0.7 mm isotropic) in vivo 7T MRI data from cognitively intact participants to characterize the human ZI in relation to surrounding structures of the subthalamic region with a specific focus on longitudinal (T1) relaxometry. After windowing to a threshold of between $1000-2000 \mathrm{~ms}$, the contrasts from T1 mapping within the ventral thalamus and subthalamic region were strikingly similar to classical Nissl staining (Figure 5.1) with white matter generally darker stained relative to gray matter. This permitted facile identification of

Table 5.2: Improvement in linear and nonlinear AFRE with multiple iterations of template creation.

| iteration | lin AFRE | nlin AFRE |
| :---: | :---: | :---: |
| 0 | $29.88 \pm 11.75(61.50)$ | $29.88 \pm 11.75(61.50)$ |
| 1 | $3.47 \pm 1.62(10.01)$ | $2.87 \pm 0.94(8.58)$ |
| 2 | $3.31 \pm 1.57(9.95)$ | $2.61 \pm 0.95(8.59)$ |
| 3 | $3.15 \pm 1.53(9.79)$ | $2.37 \pm 0.95(8.44)$ |
| 4 | $3.04 \pm 1.50(9.66)$ | $2.12 \pm 0.96(8.39)$ |
| 5 | $2.94 \pm 1.47(9.52)$ | $1.90 \pm 0.97(8.11)$ |
| 6 | $2.86 \pm 1.45(9.38)$ | $1.71 \pm 1.00(8.05)$ |
| 7 | $2.80 \pm 1.43(9.33)$ | $1.53 \pm 1.04(8.05)$ |
| 8 | $2.79 \pm 1.44(9.18)$ | $1.41 \pm 1.05(7.88)$ |
| 9 | $2.73 \pm 1.37(9.19)$ | $1.34 \pm 1.08(7.81)$ |
| 10 | $2.82 \pm 1.47(9.23)$ | $1.33 \pm 1.09(7.84)$ |

AFRE values summarized as: mean $\pm$ standard deviation (max value)
the zona incerta and surrounding structures in reference to a number of classic and modern atlases [Schaltenbrand and Wahren, 1977, Morel, 2007, Hawrylycz et al., 2012, Mai et al., 2015].

The ZI could be visualized along its entire rostrocaudal axis and was distinct from the surrounding white matter tracts, specifically the external medullary lamina of the ventral thalamus, fasciculus thalamicus (ft), and fasciculus lenticularis (fl). Regions of high T1 signal were identified both superior and inferior to the fl which we believe represent distinct dorsal and ventral components of the rZI. Interestingly, the ventral component which we observed has been obscurely named on the classic Schaltenbrand atlas, where it is unnamed on select coronal sections (Figure 2.1 b ; Coronal: +2.0 mm ), and ambiguously named on relevant sagittal sections. The labelling is no clearer in modern atlases with some atlases preferring to incorporate this label as a protrusion of the lateral hypothalamus. Caudally, the cZI was clearly distinct from nearby gray matter nuclei including the STN and RN (Figure 2.1b; Coronal: -7.0 mm). Furthermore, we identified a distinct hypointense region within the posterior subthalamic area, anterior to the cZI and anterolateral to the RN coinciding with the fct (aka raprl), previously only identified on histological sections (Figure 2.1b; Coronal: -7.0 mm ; Axial: -3.5 mm ).

Segmentation of the ZI and nearby RN and STN enabled visualization of the spatial relationship between these structures as three-dimensional models (Figure 5.2). We also provided segmentations of the cZI separately as well as the relevant portions of the ft and fct. Based on our analysis, we estimate that the ZI proper spans $\sim 15 \mathrm{~mm}$ along its main axis, $\sim 5 \mathrm{~mm}$ in the medial-to-lateral direction, and varying in height from thinner than 0.5 mm along its lateral boundary to as thick as $\sim 5 \mathrm{~mm}$ in the cZI. The volume of the ZI proper was $219.5 \mathrm{~mm}^{3}$ on the left and $211.5 \mathrm{~mm}^{3}$ on the right. The cZI volume was $63.8 \mathrm{~mm}^{3}$ on the left and $65.9 \mathrm{~mm}^{3}$

Table 5.3: AFRE summarized for the final template used in this study (10th iteration).

| AFID | Description | Mean AFRE |
| :---: | :---: | :---: |
| 01 | AC | $0.41 \pm 0.20(1.05)$ |
| 02 | PC | $0.47 \pm 0.29(1.56)$ |
| 03 | infracollicular sulcus | $0.69 \pm 0.21(1.19)$ |
| 04 | PMJ | $0.93 \pm 0.63(2.86)$ |
| 05 | superior interpeduncular fossa | $0.70 \pm 0.29(1.26)$ |
| 06 | R superior LMS | $1.11 \pm 0.67(2.93)$ |
| 07 | L superior LMS | $1.00 \pm 0.54(2.41)$ |
| 08 | R inferior LMS | $1.35 \pm 1.23(6.81)$ |
| 09 | L inferior LMS | $1.56 \pm 1.30(6.13)$ |
| 10 | culmen | $2.16 \pm 1.08(5.40)$ |
| 11 | intermammillary sulcus | $0.80 \pm 0.44(1.99)$ |
| 12 | R MB | $0.64 \pm 0.36(1.60)$ |
| 13 | L MB | $0.59 \pm 0.38(1.81)$ |
| 14 | pineal gland | $1.66 \pm 0.93(4.19)$ |
| 15 | R LV at AC | $2.11 \pm 1.42(5.28)$ |
| 16 | L LV at AC | $1.90 \pm 1.37(5.48)$ |
| 17 | R LV at PC | $2.84 \pm 1.90(8.15)$ |
| 18 | L LV at PC | $2.37 \pm 1.57(6.53)$ |
| 19 | genu of CC | $1.28 \pm 0.74(3.52)$ |
| 20 | splenium | $0.94 \pm 0.53(2.47)$ |
| 21 | R AL temporal horn | $1.98 \pm 1.31(5.25)$ |
| 22 | L AL temporal horn | $1.92 \pm 1.06(4.97)$ |
| 23 | R superior AM temporal horn | $1.72 \pm 1.36(6.41)$ |
| 24 | L superior AM temporal horn | $1.41 \pm 0.85(3.92)$ |
| 25 | R inferior AM temporal horn | $2.29 \pm 1.26(5.17)$ |
| 26 | L inferior AM temporal horn | $1.86 \pm 0.99(5.44)$ |
| 27 | R indusium griseum origin | $2.07 \pm 1.44(4.70)$ |
| 28 | L indusium griseum origin | $2.04 \pm 1.67(7.84)$ |
| 29 | R ventral occipital horn | $2.44 \pm 4.11(24.49)$ |
| 30 | L ventral occipital horn | $3.85 \pm 5.05(22.42)$ |
| 31 | R olfactory sulcal fundus | $1.43 \pm 1.02(4.05)$ |
| 32 | L olfactory sulcal fundus | $1.60 \pm 1.10(6.10)$ |

$\mathrm{AF} \overline{\overline{\mathrm{RE}} \text { values summarized as: mean } \pm \text { standard deviation (max value) }}$

■


## A



Table 5.4: Summary of T1 values, volume, centroids of key structures of the ZI region.

|  |  | ZI | cZI | fct | RN | STN |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\mathrm{T} 1(\mathrm{~ms})$ | left | $1329.1+/-111.4$ | $1361.5+/-63.9$ | $1192.2+/-39.6$ | $1169.2+/-36.5$ | $1115.6+/-65.4$ |
|  | right | $1315.3+/-107.2$ | $1339.5+/-59.9$ | $1184.4+/-36.6$ | $1163.7+/-34.8$ | $1128.6+/-47.5$ |
| volume $\left(\mathrm{mm}^{3}\right)$ | left | 219.5 | 63.75 | 162.5 | 324.2 | 168.5 |
|  | right | 211.5 | 65.88 | 161.5 | 337.5 | 163.6 |
| AC-PC Coordinates | left | $(9.1,2.5,2.6)$ | $(12.3,8.7,4.9)$ | $(10.4,5.1,1.5)$ | $(4.4,6.5,5.9)$ | $(9.7,0.3,3.4)$ |
|  | right | $(-9.7,2.6,2.3)$ | $(-13.0,9.0,4.0)$ | $(-10.2,5.2,1.7)$ | $(-4.6,6.4,5.8)$ | $(-10.3,0.4,3.2)$ |

on the right. Regarding relevant nearby white matter regions, the fct was $\sim 4-5 \mathrm{~mm}$ along its longest axis, representing 1-3 voxels if relying on DTI alone compared to 5-7 voxels with the MP2RAGE protocol with a total volume of $162.5 \mathrm{~mm}^{3}$ on the left and $162.5 \mathrm{~mm}^{3}$ on the right. Note that this is similar in size to the STN in most reported studies (e.g. [Xiao et al., 2014]).

### 5.3.3 Tissue properties of the zona incerta region

The mean T1 relaxometry values and volumes for the bilateral red nucleus, subthalamic nucleus, and zona incerta structures are provided in Table 5.4. The zona incerta appears prominent as a hyperintense region located in the subthalamic region with T 1 values generally in the $1300-1500 \mathrm{~ms}$ range with similar relaxometry parameters to cortex (see Table 1.1), providing in vivo support that the ZI represents a distinct gray matter region.

### 5.3.4 Deep brain stimulation of the caudal zona incerta

T1 values, volumes, and centroid locations are summarized in Table 5.4. We found sufficient contrast (contrast-to-noise ratio $=2.7-4.3$ ) to allow separation of the cZI and fct, which has important implications for more precise targeting within the PSA. We discovered that the T2based indirect method for targeting described was closer to the fct than the cZI centroid (3.6 vs 5.4 mm ) in both our template and also the active contact location in a patient with excellent post-operative outcome for essential tremor (Essential Tremor Rating Scale improvement by $60 \%$ total and by $80 \%$ for hand tremor and function).

### 5.4 Discussion

The present study demonstrates that robust visualization of the zona incerta region is possible using high-resolution quantitative T1 mapping. We report the first precise delineation of the zona incerta region in vivo providing estimates of the morphology (volume, dimensions) and


Figure 5.3: Study data demonstrates that the optimal target for essential tremor patients is likely the fasciculus thalamicus (fct) rather than the caudal zona incerta. Representative axial slices of the upper mesencephalon in the study group average space depicting T2 and MP2RAGE contrasts (a). The stereotactic target for essential tremor as described by Nowacki [8] is shown as a red dot within the posterior subthalamic area based on the relative location of the STN and RN in T2 images. The same target is superimposed onto the MP2RAGE average with outlines of the fct and cZI in yellow and blue respectively (a). Note that while no contrast can separate the fct from the cZI in the T2 image, the two can be visualized separately in the corresponding MP2RAGE map with the target lying within the fct region. The optimal stimulation electrode for a patient with essential tremor is shown in pink (b) as well as the relative location of the zona incerta and fct (yellow).
longitudinal relaxation-based properties of the region. Furthermore, we have identified a region of the rostral ZI inferior to the thalamic fasciculus, which to our knowledge has not previously been labeled on histological atlases of the human brain (Figure 5.2) we believe this represents the ventral rZI. Due to the striking similarity in tissue contrast with classic post-mortem Nissl staining, we were able to segment the fct (raprl) as a substructure within the posterior subthalamic area separate from the cZI. By exploiting the signal advantages of ultra-high field MRI and template averaging, we demonstrated that the zona incerta is visible in vivo, determined that this nuclear region can be decoupled from surrounding fibre pathways, and were able to employ this methodology for prospective identification of the active stimulation location for deep brain stimulation.

Efforts at visualizing small structures of the deep brain using high-field MRI have mostly focussed on transverse relaxation properties (i.e. T2- and T2* shortening) due to the welldocumented iron-rich, and thus paramagnetic contrast produced by many subcortical nuclei [Zecca et al., 2004, Haacke et al., 2005]. Increasing the strength of the main magnetic field $\left(B_{0}\right)$ results in an at least linear increase in SNR, a three- to four- fold increase compared to conventional clinical 1.5 Tesla MRI, along with significantly higher resolution. Visualization at high fields has led to more robust imaging of small structures including the STN and SN using T2-based contrast mechanisms [Keuken et al., 2013]. Others have attempted to visualize the ZI using T2 contrast, yet the region has remained elusive except for one study purporting to show that the rZI is visible, but which we show here is actually the fasciculus lenticularis (Figure 5.4). Instead, protocols for stereotactic targeting of the cZI have relied on the relative visibility of the surrounding red nucleus and STN, which can be used to infer the location of the stereotactic target within the PSA. Overall the poor visualization of ZI on T2 contrast suggests that this region is not a strong generator of T 2 contrast.


Figure 5.4: The crosshair is placed on the location identified by [Kerl et al., 2013] as the rostral ZI but overlaid on our joint MP2RAGE and T2SPACE templates. The corresponding location on the the MP2RAGE (T1 map) sequence demonstrates that this feature is actually hypointense on T1 map suggestive of white matter and thus represents the fasciculus lenticularis rather than the rZI. See Figure 5.2b for the corresponding labels in a histological reference space.

In the present study, we focus on longitudinal (T1) rather than T 2 relaxation properties of the ZI, to demonstrate its salience, and propose its use as a robust in vivo biomarker for delineating structures in the region. T1 relaxation times increase in a field-dependent manner, as does the dispersion between brain tissue types [Rooney et al., 2007], which have the effect of improving contrast between tissue types at 7T. This advantage has been exploited to parcellate thalamic nuclei [Tourdias et al., 2014] and investigate cortical laminae [Trampel et al., 2017].

Surgical planning and in vivo histology have been considered as important potential applications of the MP2RAGE sequence since its initial development [Marques et al., 2010, Marques and Gruetter, 2013]. Using this method, we demonstrate that the ZI can also be visualized along its entire rostrocaudal axis using this MRI-based tissue property (Figure 5.2). Furthermore, we found sufficient difference in T1-related tissue parameters to allows separation of the cZI from nearby white matter tracts, including the fct of the posterior subthalamic area and the ft and other components of the fields of Forel in the rZI. Rostrally, this permitted more complete characterization of the relationship between the fl and rZI, which we believe divides the rZI into dorsal and ventral components, which have been described in immunohistochemical analysis of experimental animals [Mitrofanis, 2005, Watson et al., 2014] and in at least one human brain atlas [Mai et al., 2015]. Although the increase in T1 tissue values with field strength has been perceived as a disadvantage due to increased scan time, our experience demonstrates that sufficient resolution and contrast can be attained with a scan time of just over 10 minutes (Figure 5.2; Table 5.1) with an additional 2 minute sequence for on scanner $B_{1}$ inhomogeneity correction [Eggenschwiler et al., 2012].

Since the boundaries of the ZI have not previously been well-defined in three dimensions, we elected to perform consensus segmentations using group averaging as a strategy to further boost the SNR for accurate delineation of these structures from our 7T data. Our interpretation of the boundaries of the zona incerta using in vivo sequences was based on meticulous comparison with annotations of the zona incerta from classical and modern histological atlases [Schaltenbrand and Wahren, 1977, Morel, 2007, Hawrylycz et al., 2012, Mai et al., 2015]. Our technique compares favorably with classic post-mortem atlases for mesoscale morphometric analysis due to our ability to pool structural information from a set of 32 participants and also circumvent the challenges of histological processing that can result in tissue shrinkage, deformation, and tears. While errors in registration may result in errors in segmentation of the individual subjects, we were able to confirm adequate spatial correspondence by using visual quality control and also by determining that fiducial registration error was in the millimetric range (Table 5.2 and Figure 5.1). Our high-resolution template creation approach allowed for the pooling of data from multiple participants $(\mathrm{N}=32)$ into a single reference space allowing us to better account for intersubject variability compared to histological atlas creation, which is also prone to tissue deformations [Morel, 2007]. Due to the complexity of this structure, future efforts will involve further clarifying the exact boundaries of the rostral ZI through a combination of detailed histological analysis and population studies of the region.

Based on our analysis, T1 maps appear to represent an optimal quantitative contrast by which to visualize the zona incerta region. As already mentioned, there is very close correspondence in intensity between our in vivo T1 maps with Nissl staining [Schaltenbrand and

Wahren, 1977, Weiskopf et al., 2015]. However, the correspondence with histology is not strictly one-to-one, noting that the RN and STN appear hypointense on T1 maps despite being largely gray matter regions in the current study (Table 5.4; Figure 5.2) as well as in work from other groups (e.g. [Keuken et al., 2017]). We believe that the relatively low T1 map contrast in these regions is related to the density of myelinated tracts travelling within these nuclear regions. In the ZI region, which is relatively sparsely nucleated, the traversing white matter tracts seem to occur at specific spatial clusters (ft and raprl). Multiparametric approaches may help further characterize underlying tissue properties. We have found the multicontrast approach (both T1 and T2) helpful for establishing that a region previously identified as the rZI [Kerl et al., 2013] actually represents the fl (Figure 5.4). Unfortunately, due to limitations in our T2 protocol, we could not investigate the T 2 values quantitatively (see Methods).

Our analysis demonstrates that there is sufficient signal and contrast within the PSA region to allow separation of the cZI from the fct and ml (see Table 5.4). We discovered that commonly used T2-based indirect anatomical target and optimal stimulation locations were closer to the fct than the cZI centroid. These findings are in line with other work suggesting that a proportion of benefit is derived from stimulation of wayward white matter tracts in the fct (raprl) [Spiegel et al., 1964, Mundinger, 1965, Velasco et al., 1972, Mohadjer et al., 1990, Blomstedt et al., 2010, Blomstedt et al., 2018], and also concordant with recent studies employing diffusion tensor imaging (DTI) [Fiechter et al., 2017, Dallapiazza et al., 2018, Velasco et al., 2018]. Compared to DTI-based measures, high-field T1 mapping has higher SNR, is less prone to image distortions, and is acquired at inherently higher resolution ( 0.7 mm compared to $2-3 \mathrm{~mm}$ isotropic). We have determined that the dimensions of the fct within the PSA is $\sim 4-5 \mathrm{~mm}$ along its longest axis, representing 1-3 voxels if relying on DTI alone compared to 5-7 voxels using our MP2RAGE protocol. Nowacki and colleagues have previously suggested that the DRTT (dentatorubrothalamic tract within raprl) is the key area within the PSA for targeting [Fiechter et al., 2017]. However, their analysis of the active contact location suggests that the most efficient stimulation location was further away $(1.84 \pm 1.24 \mathrm{~mm})$ from the DRTT than the least efficient stimulation location $(0.92 \pm 1.21 \mathrm{~mm})$ - although this did not meet thresholds of statistical significance [Nowacki et al., 2018].

Radiofrequency inhomogeneity is a well-known problem with increasing magnetic field strength biasing the interpretation of T1 maps since stronger RF pulses are required for excitation and detection leading to more pronounced intensity bias in the images [Rooney et al., 2007] (Fig1). The MP2RAGE sequence used in this study was designed to be a self bias-field correcting sequence free of PD and T2* weighting, RF receiver field bias, and low-order RF transmit field bias [Marques et al., 2010]. In this study, we also acquired an additional lowresolution $B_{1}$ mapping protocol, SA2RAGE [Eggenschwiler et al., 2012], as a means to detect

Table 5.5: Comparison between MP2RAGE sequence used in this study and that used by [Forstmann et al., 2014].

| Sequence |  | TE(ms) | TR (ms) | TI | Flip Angle (\%) | Matrix | PAT* | Averages | Resolutio | Acquisition Time (min:sec) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Current Stud | 3D | 2.73 | 6000 | 0/700 | 4/5 | $342 \times 342 \times 224$ | 3 | 1 | $0.7 \times 0.7 \times 0.7$ | 10:14 |
| Forstmann 2014 | 3D | 2.45 | 5000 | 0/2750 | 4/11 | $320 \times 32 \times 240$ | 2 | 1 | $0.7 \times 0.7 \times 0.7$ | 10:57 |

* PAT = parallel acquisition technique (acceleration factor)
and subsequently correct the transmit inhomogeneity. Finally, and perhaps most importantly, our acquisitions benefited from the use of parallel transmission of RF pulses using a multiple channel approach [Gilbert et al., 2011], a hardware-based approach for achieving better intensity homogeneity. This combination of hardware and software calibrations has helped to improve the quality and homogeneity of our 7T T1 map acquisitions enabling more robust identification and characterization of the ZI and surrounding region.

Some discrepancy was noted when comparing values reported by others who have investigated the subthalamic region using MP2RAGE-based T1 mapping [Forstmann et al., 2014, Keuken et al., 2017]. In particular, our values tended to be $\sim 100-200 \mathrm{~ms}$ shorter within the STN and SN. More perplexing was that we also noted some difference in age-related trends using our ROIs. Several factors may account for the discrepancies between our studies. Beyond inter-scanner variability in hardware (e.g. our use of parallel transmission) and our use of SA2RAGE-based correct (see the paragraph above), inter-protocol variability has been reported [Stikov et al., 2015]. In addition, there were some differences in acquisition parameters (Table 5.5), which unfortunately due to assumptions about mono-exponential longitudinal relaxation in the MP2RAGE implementation can introduce additional sequence-dependent measurement variability [Rioux et al., 2016]. These factors all lead the T1 value measurements to tend to be more precise than accurate, contributing to inter-site differences. For ROI segmentation, Forstmann et al. performed meticulous labelling of the STN and SN for every subject by two raters, a highly manually intensive process [Forstmann et al., 2014]. Our approach, instead, relied on careful delineation of the ROIs in reference space, with propagation error quantified to $\sim 1 \mathrm{~mm}$ using a recently described metric for estimating registration accuracy between template and subject scans [Lau et al., 2019] (see Chapter 3). Perhaps the most important difference relates to the Forstmann group's choice to use "conjunction masks" as the choice of ROI (that is the intersection between the labels of two raters as the choice of ROI). This approach ensures that summary measures are obtained for voxel regions confidently within the structure of interest, thus preventing boundary effects such as the impact of changes in surrounding structures; but on the other hand, underestimates the volume of the structures (e.g. the STN sizes reported by Forstmann and colleagues tended to be lower $\sim 50 \mathrm{~mm}^{3}$ than those reported by other groups). In the future, these data can be pooled into larger multisite analyses
with standardized workflows that will better clarify our findings on a larger scale.
Our findings add to the growing body of knowledge that the optimal DBS target within the posterior subthalamic area is at the anterior boundary of the cZI abutting or directly within the fasciculus cerebellothalamicus [Herrington et al., 2016]. This suggests that direct targeting of the white matter, in other words connection-based targeting, may be central to efficacy, which is increasingly being recognized for the treatment of tremor [Akram et al., 2018] and other disorders [Horn et al., 2017b, Horn et al., 2017c]. Our approach using T1 mapping for visualizing relevant WM tracts ( $\mathrm{fct}, \mathrm{al}, \mathrm{ft}, \mathrm{fl}$ ) is in contrast to more common methods using diffusion-based imaging. In regards to human in vivo studies, DTI studies have mostly focussed on connections between larger cortical and subcortical structures since achieving high resolution (submillimetric) images in clinically feasible timeframes for DTI remains a challenge. There is also increasing acknowledgement that connectivity-based methods are prone to producing false-positive tracts [Maier-Hein et al., 2017]. An additional advantage of using T1 mapping is that the images can simultaneously be used as a baseline structural scan and furthermore employed to identify the target, eliminating the need for an image fusion step, which can introduce error.

### 5.5 Conclusions

We demonstrate using ultra-high field MRI and template averaging that the zona incerta is visible in vivo in control and patient datasets. In addition, we determined that this nuclear region can be decoupled from surrounding fibre pathways. Due to the striking similarity in tissue contrast with classic post-mortem Nissl staining, we were able to segment the fasciculus cerebellothalamicus as a substructure within the posterior subthalamic area separate from the caudal ZI. Furthermore, we identified a region of the rostral ZI inferior to the thalamic fasciculus, which to our knowledge has not previously been labeled on histological atlases of the human brain, and which we believe represents the ventral rZI described in experimental animals [Mitrofanis, 2005]. This work enables high-resolution in vivo visualization of a region previously only seen on histology, paving the way for patient-specific optimization and characterization of stereotactic targets.

## Chapter 6

## Conclusions and Future Directions

The work in this thesis has explored ways in which ultra-high field magnetic resonance imaging can be integrated into the practice of stereotactic neurosurgery. The unique contributions can be summarized as follows:

- Chapter 2: For the first time, an ultra-high field MRI template is integrated into the surgical workflow to assist with surgical planning for deep brain stimulation surgery cases.
- Chapter 3: A novel anatomical fiducial placement protocol is developed, validated, and used prospectively to quantify the limits of template-assisted surgical planning.
- Chapter 4: A novel morphometry workflow is employed to characterize local geometric distortion in ultra-high field MR images identifying systematic regions with higher geometric uncertainty, which should be taken into account if using high-field imaging for surgical planning.
- Chapter 5: A number of important stereotactic targets (i.e. the zona incerta and fasciculus thalamicus) are directly visualized and characterized for the first time in vivo at high resolution using T1 mapping methods at ultra-high field.

By identifying the limits of atlas- and template-assisted stereotactic planning, characterizing regional biases of spatial uncertainty, and identifying anatomical targets not prreviously visible in vivo, this work paves the way for patient-specific surgical planning using ultra-high field MRI.

### 6.1 The Limits of Image-Based Targeting

Proof-of-principle evidence for template-assisted surgical planning for deep brain stimulation surgery is provided in Chapter 2. By employing deformable (nonlinear) registration methods, this represents a reasonable technique where patient imaging is inadequate, i.e. when only standard field MRI is available and where clinical imaging is subpar (motion artifact, etc.). In Section 3.3.3, the geometric limits of template-assisted registration with patient datasets were explored, demonstrating that with current state-of-the-art deformable registration methods, errors in point-based registration were present even beyond the limits of localization error. This analysis revealed misregistrations on the order of $1.80 \pm 2.09 \mathrm{~mm}$ and as high as over 30 mm . These results provide evidence that standard deformable registration methods remain insufficiently accurate for the purposes of stereotactic neurosurgery, and also suggest that high-quality patient-specific imaging may be a welcome alternative.

While using patient-specific imaging avoids the problem encountered with template registration where two images with different anatomical features are matched, geometric distortion in images, particularly with an increase to high-field may present some issues. Several sources for geometric distortion issues have been highlighted in Chapter 4. In a dataset of subjects scanned at ultra-high field and standard field, we were able to characterize the location of these spatial biases. These $B_{0}$-related issues tended to occur at air-tissue boundaries such as the floor of the middle fossa. Fortunately, the work in this thesis has determined that the areas of the deep brain which are the putative targets for the majority of stereotactic procedures are relatively protected from these effects paving the way for the use of patient specific 7T imaging. These results are not necessarily generalizable to other scanners given the unique combination of hardware. However, the framework used that employs voxel-wise morphometric analysis can be used by other groups. Also in the future, other specific sources of distortion that are sequence-specific, i.e. Maxwell distortions, should be explored.

Despite advances in resolution for patient imaging and well-optimized distortion correction methods, other factors, that may be limiting the ability to accurately target these small brain structures, become more important. In the neurosurgical literature, the term application accuracy is used to describe the multifactorial nature of this problem in the surgical environment. That is, targeting error is influenced by many different variables from the quality of preoperative imaging, image fusion, the quality of fixation of the frame or pins to the head, to the limits of physical accuracy of the devices being employed to perform targeting [Maciunas et al., 1994, Grunert et al., 2003, Henderson et al., 2004, Shamir et al., 2011]. It has been perceived that the physical or mechanical accuracy of current physical frames is on the order of 1-2 mm. Recent evidence suggests some modest improvement in targeting accuracy (to $1-1.5 \mathrm{~mm}$ error) with modern robotic devices for stereotactic targeting [Cardinale et al., 2013]. While worth mentioning, this thesis has focussed on optimizing the image-based aspects of the application accuracy problem.

### 6.2 Multiparametric Imaging for Stereotactic Neurosurgery

The latter part of this thesis has focussed on using T1 mapping for visualizing small structures in the deep brain. Integration with other quantitative MRI-based parameters would provide richer multi-contrast information that could be used to better delineate between different local tissue types. For example, other acquisition types that characterize local magnetic properties using multi-echo acquisitions and associated post-processing methods (R2* and quantitative susceptibility mapping), and local diffusion-related properties such as mean diffusivity (MD) and fractional anisotropy (FA) have been demonstrated to be beneficial for surgical planning.

These different imaging methods reflect to different extents underlying biological features related to molecular-level content (water, iron, myelin, gray matter) and organization (anistropy, etc.).

Over the past few years, connectomic approaches that focus on the connections between regions to optimizing neuromodulation sites has become increasingly popular [Fox et al., 2014]. These methods focus on using diffusion-based or functional MRI-based sequences to determine coherence or connectivity based on DBS lead locations. For Parkinson's disease, these studies have focussed on large tract connections to cortex and their association with lead location [Vanegas-Arroyave et al., 2016, Akram et al., 2017, Horn et al., 2017b, Horn et al., 2017c]. However, despite convergence of evidence from multiple groups, these methods have recently been discovered to be prone to false-positive results [Maier-Hein et al., 2017]. Ultimately, the approach taken in this thesis, particularly with increasingly higher resolution imaging, should be considered complementary to these endeavours enabling accurate localization of smaller tracts and structures for integration into connectivity analyses.

## 6.3 "Asleep" Deep Brain Stimulation Surgery

Over the last several years, a number of groups have shifted to "asleep" DBS surgery-that is, advocating for direct image-based directing using pre-operative imaging with intraoperative verification with the patient under general anaesthesia [Hyam et al., 2015, Chen et al., 2016, Brodsky et al., 2017]. To date, all the existing asleep DBS protocols have relied on imaging at standard magnetic field strengths. Overall, good success has been documented, although a recent meta-analysis suggests that patients who were awake during surgery suffered from fewer stimulation-related side effects than those performed asleep [Ho et al., 2017]. This has been corroborated in a recent single center study demonstrating improved outcomes in awake versus asleep patients [Blasberg et al., 2018]. Another recent retrospective study observed that while image-based targeting was appropriate in the vast majority of cases, in $20 \%$ of cases the imaging-based target proved to be suboptimal [Lozano et al., 2018]. Still, others continue to advocate for MER recordings with recent evidence demonstrating that multiple MER tracks are more beneficial than the use of a single track [Bjerknes et al., 2018]. These recent findings all point to the need for caution and that other adjuncts, whether microelectrode recording or better imaging (see Chapter 5), may help to facilitate improved clinical benefit in patients undergoing stereotactic neurosurgery. In the future, 7T imaging for in vivo visualization and direct targeting of brain structures for DBS may eliminate the need for awake surgery enabling personalized therapy where DBS target selection is guided by high-quality, patient-specific imaging.

Microelectrode recording during surgery has been central to the origins of stereotaxy and the discovery of targets for DBS implantation, and yet there has been a clear shift with improving imaging towards making this technique obsolete. With improving imaging, the number of trajectories to be explored may be decreased, which should help mitigate any risk of complications to the patient and decrease overall operating room time. So what becomes of MER in the future? There is no doubt about the role of MER for neuroscientific study, but will it remain important in specific circumstances for clinical decision-making? These discussions are mostly beyond the limits of the presented work. However, my belief is that MER will not disappear and will continue to be an important component of functional neurosurgery. However, the importance of single neuron activity in clinical practice will shift from its use for localization to processing of these signals for decoding more complex behaviours and activities, including its use in brain-computer interfaces.

### 6.4 Innovations in Stereotaxy

Functional neurosurgery is an exciting subspecialty within neurosurgery with many new technological developments over the past several decades. Directional leads and the possibility of current steering allow for some tolerance to targeting error by allowing "sculpting" of the field of stimulation. MRI-guided laser ablation has become a valid treatment option for temporal lobe epilepsy [Willie et al., 2014, Gross et al., 2018]. Closed loop deep brain stimulation devices now allow for responsive treatment based on specific brain signals [Morrell, 2011, Herron et al., 2016]. Optogenetic [Gradinaru et al., 2010] and chemogenetic [Roth, 2016] modulation have come to the forefront as tools in neuroscience. Initial safety trials have at least demonstrated short-term safety of gene therapy in human subjects [LeWitt et al., 2011]. Robotic devices are enabling more and more accurate stereotactic placement [Cardinale et al., 2013]. All these developments can be aided by more accurate image-based targeting.

### 6.5 Conclusions

To conclude, the question of target selection not only for classic indications like Parkinson's disease and Essential Tremor but the growing number of indications remains an unsolved problem. Determining the exact location of the most effective target that maximizes benefit while minimizing side effects or complications will be greatly assisted by having access to higherresolution in vivo brain imaging, facilitated by higher field imaging. With the recent approval of specific ultra-high field scanner configurations by the FDA, more groups are investing in this technology, and accelerations in developments in this growing field are to be anticipated over
the next decade. The opportunity to pool ultra-high field data across multiple centers will also enable generalizability of the findings in this thesis.

## Appendices

## Appendix A

# Chapter 3 Supplementary Material 

A. 1 Phase 1: Supplementary Material

## Phase 1：Protocol Validation for Brain Templates

This notebook contains results validating the AFID protocol on three openly available templates（Agi1e 12 v 2016, colin27，and ICBM2009bAsym）．
The first step is to initialize the variables，define useful functions，and load all the raw fosv data into df＿raters．

## Template Averages

For each template，we calculate the mean value for each AFID32 point and store it in a separate ．fcsv file so that it can be loaded back into 3D Slicer．
Deviation of the values by $>10 \mathrm{~mm}$ will be classified as an outtier．

Phase 1：Raw Data Analysis
Also classify extreme outliers，defined as $>=10 \mathrm{~mm}$ from the group mean
＇Total： $1.27+/-1.98 \mathrm{~mm}$ ；Outiers： $24 / 3072$（ $0.78 \%$ ）＇
＇Agile12v2016： $1.10+/-1.59 \mathrm{~mm}$ ；Outtiers： $3 / 1024$（ $0.29 \%$ ）
＇Colin27： $1.71+/-2.78$ mm；Outilers：20／1024（1．95\％）＇
＇MN1152NLin2009bAsym： $0.99+/-1.11 \mathrm{~mm}$ ；Outiers： $1 / 1024$（ $0.10 \%$ ）＇

Template Averages：Post－QC
Template averages were recreated after quality control and filtering of outiers．
＇Total： $1.03+/-0.94 \mathrm{~mm}$ ；Outiers： $1 / 3048$（ $0.03 \%$ ）${ }^{\text {＇}}$
＇Agile12v2016： $1.01+/-0.93 \mathrm{~mm}$ ；Outiers：0／1021（0．00\％）＇
＇Colin27： $1.11+/-1.05 \mathrm{~mm}$ ；Outliers： $1 / 1004$（ $0.10 \%$ ）＇
＇MN152NLin2009bAsym： $0.97+/-0.80 \mathrm{~mm}$ ；Outiers： $0 / 1023(0.00 \%)$＇

| AFID | Description | Agile 12v2016 Pre－QC | Agile 12v2016 Post－QC | Colin27 Pre－QC | Colin27 Post－Qc | MNI2009bAsym Pre－QC | MNI2009bAsym Post－QC | Total Pre－QC | Total Post－QC |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 01 | AC | 0．33＋0．16（0） | $0.33 \pm 0.16$（0） | 0．34土0．29（0） | 0．34土0．29（0） | 0．35 0.20 （0） | $0.35 \pm 0.20$（0） | 0．34土0．22（0） | 0．34土0．22（0） |
| 02 | PC | 0．340． 19 （0） | 0．340．19（0） | 0．35＋0．18（0） | 0．35＋0．18（0） | 0．33＋0．14（0） | 0．33＋0．14（0） | 0．34＊0．17（0） | 0．34＊0．17（0） |
| 03 | infracollicular sulcus | $1.25 \pm 0.47$（0） | $1.25 \pm 0.47$（0） | $1.22 \pm 0.48$（0） | $1.22 \pm 0.48$（0） | 1．08＋0．46（0） | 1．08＋0．46（0） | 1．17 10.47 （0） | 1．17 17.47 （0） |
| 04 | PMJ | 0．83＋0．47（0） | 0．83＋0．47（0） | 0．97 0.65 （0） | 0．97＋0．65（0） | 0．84土0．52（0） | 0．84＋0．52（0） | 0．87 $0.544^{(0)}$ | 0．87 1.17 .54 （0） |
| 05 | superior interpeduncular fossa | 1．15＋0．61（0） | 1．15＋0．61（0） | 0．96 0.600 （0） | 0．96＋0．60（0） | 1．12＋0．50（0） | $1.12 \pm 0.50$（0） | $1.08 \pm 0.57$（0） | 1．08＋0．57（0） |
| 06 | R superior LMS | 0．75 $\pm 0.48$（0） | 0．75 $\pm 0.48$（0） | 1．16 00.69 （0） | 1．16 00.69 （0） | 0．68＋0．50（0） | 0．68＋0．50（0） | 0．85 0.59 （0） | 0．85 00.59 （0） |
| 07 | L superior LMS | 0．93＋0．59（0） | 0．93 10.59 （0） | 1．05 0.0 .57 （0） | 1．05＋0．57（0） | 0．9110．90（0） | $0.91 \pm 0.90$（0） | $0.96 \pm 0.71$（0） | 0．96 +0.71 （0） |
| 08 | R inferior LMS | 1．55＋1．14（0） | 1．55＋1．14（0） | 1．61＋1．07（0） | 1．61＋1．07（0） | 1．47＋0．96（0） | $1.47 \pm 0.96$（0） | 1．54＊1．05（0） | 1．54＋1．05（0） |
| 09 | L inferior LMS | 1．39＋1．11（0） | 1．39＋1．11（0） | 1．79＋1．32（0） | 1．79＊1．32（0） | 1．63＋1．19（0） | 1．63＋1．19（0） | 1．60土1．21（0） | 1．60 1.21 （0） |
| 10 | culmen | 1．03 $\pm 0.73$（0） | 1．03＋0．73（0） | 0．68 +0.24 （0） | 0．68 00.24 （0） | 0．61 00.32 （0） | $0.61 \pm 0.32$（0） | $0.77 \pm 0.50$（0） | 0．77 0.0 .50 （0） |
| 11 | intermammillary sulcus | 0．73 0.34 （0） | 0．73 00.34 （0） | 0．68＋0．34（0） | 0．68 00.34 （0） | 0．70 00.38 （0） | 0．70 00.38 （0） | 0．70 0.3 .35 （0） | 0．70 00.35 （0） |
| 12 | R MB | $0.37 \pm 0.28$（0） | 0．37 0.28 （0） | 0．4440．32（0） | 0．44＊0．32（0） | 0．48土0．34（0） | 0．48 +0.34 （0） | 0．44土0．31（0） | 0．44＊0．31（0） |
| 13 | LMB | 0．43 $\pm 0.27$（0） | 0．43 $\pm 0.27$（0） | $0.53 \pm 0.32$（0） | 0．53 0.3 .32 （0） | 0．5000．31（0） | 0．5000．31（0） | 0．49 0.300 （0） | 0．49＋0．30（0） |
| 14 | pineal gland | 0．7000．33（0） | 0．7000．33（0） | 0．9440．33（0） | 0．94 00.33 （0） | 0．68＊0．51（0） | 0．68＋0．51（0） | 0．77 $\pm 0.42$（0） | 0．77 70.42 （0） |
| 15 | R LV at AC | 0．99＋1．48（0） | 0．99＋1．48（0） | 0．68＋0．42（0） | 0．68＋0．42（0） | 0．62 0.50 （0） | 0．62＋0．50（0） | 0．75 0.0 .92 （0） | 0．75 $\pm 0.92$（0） |
| 16 | Llv at AC | 1．06＋1．60（0） | 1．06＋1．60（0） | 0．73 +0.42 （0） | 0．73 $\pm 0.42$（0） | 0．62＋0．51（0） | 0．62＋0．51（0） | $0.79 \pm 0.98$（0） | 0．79 $\mathrm{T}^{0.98 \text {（0）}}$ |
| 17 | RLV at PC | 1．13 +1.35 （0） | 1．13＋1．35（0） | 1．12＋1．01（0） | 1．12＋1．01（0） | 1．0000．60（0） | $1.00 \pm 0.60$（0） | $1.08 \pm 1.00$（0） | 1．08＋1．00（0） |
| 18 | L LV at PC | $1.23 \pm 1.46$（0） | $1.23 \pm 1.46$（0） | $1.32 \pm 1.02$（0） | $1.32 \pm 1.02$（0） | 1．03＋0．58（0） | $1.03 \pm 0.58$（0） | 1．18土1．05（0） | 1．18＋1．05（0） |
| 19 | genu of CC | 1．0000．46（0） | $1.00 \pm 0.46$（0） | $0.63 \pm 0.24$（0） | 0．63 0.24 （0） | 0．78＋0．48（0） | 0．78 70.48 （0） | 0．80 0.44 （0） | 0．800．44（0） |
| 20 | splenium | 0．71 10.39 （0） | 0．71 1 0．39（0） | 0．52 0.2 .27 （0） | 0．52＋0．27（0） | 0．80 11.10 （0） | 0．80⼟1．10（0） | 0．68土0．73（0） | 0．68 0.0 .73 （0） |
| 21 | R AL temporal horn | 1．44＋1．20（0） | $1.44 \pm 1.20$（0） | 1．52＋0．79（0） | 1．52＋0．79（0） | $1.15 \pm 0.89$（0） | $1.15 \pm 0.89$（0） | $1.36 \pm 0.98$（0） | 1．36＋0．98（0） |
| 22 | LAL temporal horn | $1.64 \pm 1.92$（1） | 1．32＋0．91（0） | 1．1000．56（0） | 1．1000．56（0） | 1．16 00.94 （0） | 1．16 00.94 （0） | $1.29 \pm 1.27$（1） | 1．19＋0．82（0） |
| 23 | R superior AM temporal horn | 0．62 0.3 .38 （0） | 0．62 0.38 （0） | 1．31 11.71 （0） | 1．3111．71（0） | 0．83 0.9 .91 （0） | 0．83 00.91 （0） | 0．91土1．15（0） | 0．91土1．15（0） |
| 24 | L superior AM temporal horn | 0．59＋0．39（0） | 0．59＋0．39（0） | $2.02+1.90$（0） | 2．02＋1．90（0） | 0．95 0.0 .98 （0） | 0．95 00.98 （0） | 1．17 11.36 （0） | 1．17 1.1 .36 （0） |
| 25 | R inferior AM temporal hom | 1．31＋1．20（0） | 1．31＋1．20（0） | 1．49＋0．94（0） | 1．49＋0．94（0） | 1．40 11.00 （0） | 1．40＋1．00（0） | 1．40＾1．04（0） | 1．40 11.04 （0） |
| 26 | Linferior AM temporal horn | 1．36＋1．16（0） | $1.36 \pm 1.16$（0） | $1.41 \pm 1.06$（0） | 1．41 1.1 .06 （0） | 1．39 0.7 .76 （0） | 1．39 0.7 .76 （0） | $1.38 \pm 0.98$（0） | 1．38＋0．98（0） |
| 27 | R indusium griseum origin | 2．58 4.99 （1） | 1．38＋0．75（0） | 1．70 $\pm 1.08$（0） | $1.70 \pm 1.08$（0） | $1.26 \pm 0.82$（0） | 1．26＋0．82（0） | $1.81 \pm 2.92$（1） | $1.43 \pm 0.90$（0） |
| 28 | L indusium griseum origin | 2．57 $\pm 4.91$（1） | 1．52＋1．14（0） | $2.11 \pm 1.44$（0） | 2．1111．44（0） | 1．4000．98（0） | 1．4000．98（0） | 1．99＊2．94（1） | $1.66 \pm 1.22$（0） |
| 29 | R ventral occipital hom | 1．59＋1．07（0） | 1．59 1.07 （0） | $11.38 \pm 4.82$（10） | 0．8000．45（0） | 2．07 $\pm 4.11$（1） | 1．34＊1．25（0） | 4．84土5．78（11） | $1.30 \pm 1.08$（0） |
| 30 | L ventral occipital horn | 1．09＋1．13（0） | 1．09＋1．13（0） | 10．04土4．78（10） | 1．63＋2．94（1） | 1．25＋1．32（0） | $1.25 \pm 1.32$（0） | $3.96 \pm 5.02$（10） | $1.28 \pm 1.78$（1） |
| 31 | R olfactory sulcal fundus | 1．17 1 0．68（0） | 1．17 17.68 （0） | 1．41 0.95 （0） | 1．41 1.0 .95 （0） | 1．14＊0．59（0） | 1．14＋0．59（0） | $1.23 \pm 0.75$（0） | 1．23＋0．75（0） |
| 32 | L olfactory sulcal fundus | $1.23 \pm 0.54$（0） | $1.23 \pm 0.54$（0） | $1.41 \pm 1.00$（0） | $1.41 \pm 1.00$（0） | $1.13 \pm 0.63$（0） | $1.13 \pm 0.63$（0） | $1.25 \pm 0.74$（0） | 1．25 20.74 （0） |

[^3]
## Demographics of Raters

## Details regarding experience, etc.

| rater_id | imaging_exp | neuro_exp | slicer_exp | description |
| :---: | :---: | :---: | :---: | :---: |
| Rater01 | 24 | 24 | 24 | undergrad_student |
| Rater02 | 0 | 0 | 0 | medical_student |
| Rater03 | 8 | 0 | 8 | undergrad_student |
| Rater04 | 24 | 6 | 0 | grad_student |
| Rater05 | 0 | 24 | 0 | grad_student |
| Rater06 | 24 | 12 | 12 | grad_student |
| Rater07 | 12 | 48 | 12 | grad_student |
| Rater08 | 0 | 0 | 0 | undergrad_student |

## Secondary Analyses

## We first evaluated whether there was any evidence of learning across sessions (excluding session 0 which was completed as part of a group tutoria). There were negative trends in the mean AFLE with increasing session number but these did not meet thresholds of statistical analysis. The first column is the effect. second column is the associated $p$-value. Wese did not meet thresholds of statistical analysis. The first column is the effect, second column is the associated $p$-value. <br> | (Intercept) | 1.090 | 0.0000 |
| :--- | :--- | :--- |
| session | -0.024 | 0.1141 |

## Did specific raters demonstrate any learning?

Because of the trends, we explored further to determine whether any specific raters demonstrated any learning. After multiple comparisons correction, only two raters demonstrated statistically significant change with session number. Rater \#4 was observed to start at a baseline increased rating error in the first session ( 1.64 mm ) but demonstrated a decrease in AFLL with session number improving by $0.1-0.2 \mathrm{~mm}$ per session based on the linear model (statistically significant improvement). On the contrary, Rater \#2 who started with an intercept of 0.59 mm (better than the average) showed worsening of rater error with time.

| rater | (Intercept) | pval_(Intercept) | session | pval_session | pval_session_adjusted | pval_session_significant |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| 1 | 0.78 | 0 | 0.02 | 0.5644 | 0.6450 | FALSE |
| 2 | 0.59 | 0 | 0.12 | 0.0001 | 0.0009 | TRUE |
| 3 | 1.30 | 0 | -0.06 | 0.1624 | 0.2783 | FALSE |
| 4 | 1.64 | 0 | -0.17 | 0.0002 | 0.0009 | TRUE |
| 5 | 1.05 | 0 | 0.04 | 0.2881 | 0.3841 | FALSE |
| 6 | 1.08 | 0 | -0.04 | 0.1739 | 0.2783 | FALSE |
| 7 | 0.84 | 0 | 0.02 | 0.7086 | 0.7086 | FALSE |
| 8 | 1.45 | 0 | -0.12 | 0.0210 | 0.0560 | FALSE |

## Did AFLE improve for specific AFIDs?

We wanted to see if specific AFIDs tended to improve with more training (i.e. more sessions). This analysis did not survive multiple comparisons analysis.

| fid | (Intercept) | pval_(Intercept) | session | pval_session | pval_session_adjusted | pval_session_significant |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 0.36 | 0.0000 | -0.01 | 0.7009 | 0.8307 | FALSE |
| 2 | 0.38 | 0.0000 | -0.01 | 0.3531 | 0.7955 | FALSE |
| 3 | 1.28 | 0.0000 | -0.03 | 0.4534 | 0.7955 | FALSE |
| 4 | 0.65 | 0.0000 | 0.09 | 0.0588 | 0.4391 | FALSE |
| 5 | 1.08 | 0.0000 | 0.00 | 0.9802 | 0.9802 | FALSE |
| 6 | 0.71 | 0.0000 | 0.06 | 0.2590 | 0.7296 | FALSE |
| 7 | 0.92 | 0.0000 | 0.01 | 0.8661 | 0.8941 | FALSE |
| 8 | 1.76 | 0.0000 | -0.08 | 0.4408 | 0.7955 | FALSE |
| 9 | 1.82 | 0.0000 | -0.07 | 0.5221 | 0.7955 | FALSE |
| 10 | 0.79 | 0.0000 | -0.01 | 0.8035 | 0.8866 | FALSE |
| 11 | 0.85 | 0.0000 | -0.06 | 0.0686 | 0.4391 | FALSE |
| 12 | 0.47 | 0.0000 | -0.01 | 0.6396 | 0.8307 | FALSE |
| 13 | 0.41 | 0.0000 | 0.03 | 0.2885 | 0.7296 | FALSE |
| 14 | 1.02 | 0.0000 | -0.10 | 0.0076 | 0.2431 | FALSE |
| ${ }^{15}$ | 0.68 | 0.0051 | 0.03 | 0.6932 | 0.8307 | FALSE |
| 16 | 0.74 | 0.0044 | 0.03 | 0.7880 | 0.8866 | FALSE |
| 17 | 1.00 | 0.0002 | 0.04 | 0.6840 | 0.8307 | FALSE |
| 18 | 1.25 | 0.0000 | -0.02 | 0.8506 | 0.8941 | FALSE |
| 19 | 0.87 | 0.0000 | -0.02 | 0.5664 | 0.8239 | FALSE |
| 20 | 0.86 | 0.0000 | -0.09 | 0.0188 | 0.3005 | FALSE |
| 21 | 1.22 | 0.0000 | 0.07 | 0.4391 | 0.7955 | FALSE |
| 22 | 1.05 | 0.0000 | 0.07 | 0.3945 | 0.7955 | FALSE |
| 23 | 1.21 | 0.0000 | -0.13 | 0.1677 | 0.7296 | FALSE |
| 24 | 1.34 | 0.0001 | -0.08 | 0.5017 | 0.7955 | FALSE |
| 25 | 1.76 | 0.0000 | -0.13 | 0.1829 | 0.7296 | FALSE |
| 26 | 1.64 | 0.0000 | -0.10 | 0.2964 | 0.7296 | FALSE |
| 27 | 1.04 | 0.0000 | 0.15 | 0.0629 | 0.4391 | FALSE |
| 28 | 1.47 | 0.0000 | 0.08 | 0.4954 | 0.7955 | FALSE |
| 29 | 1.79 | 0.0000 | -0.18 | 0.0984 | 0.5246 | FALSE |
| 30 | 1.82 | 0.0004 | -0.20 | 0.2627 | 0.7296 | FALSE |
| 31 | 1.46 | 0.0000 | -0.09 | 0.2158 | 0.7296 | FALSE |
| 32 | 1.33 | 0.0000 | -0.03 | 0.6341 | 0.8307 | FALSE |

Intra-Rater AFLE

| template | mean | sd |
| :--- | :--- | :--- |
| Agile12v2016 | 1.13 | 0.86 |
| Colin27 | 1.14 | 0.92 |
| MN1152NLLin2009bAsym | 1.03 | 0.78 |

## Inter-Rater AFLE



## Summary of Validation Results (Post-QC)

Mean AFLE, Intra-Rater AFLE, Inter-Rater AFLE

2／24／2019
PHASE1＿template＿validation

| AFID | Description | Agile12v2016 Mean AFLE Mean AFLE | Colin27 Mean AFLE | MNI2009bAsym | $\begin{array}{\|l\|l\|} \hline \text { Total } \\ \text { Mean } \\ \text { AFLE } \end{array}$ | Agile 12v2016 Intra－Rater | Colin27 Intra－Rater | MNI2009bAsym Intra－Rater | $\begin{array}{\|l\|l} \hline \text { Total } \\ \text { Intra- } \\ \text { Rater } \end{array}$ | Agile 12 v 2016 Inter－Rater | Colin27 Inter－Rater | MNI2009bAsym Inter－Rater | $\begin{array}{\|l\|} \hline \text { Total } \\ \text { Inter- } \\ \text { Rater } \end{array}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 01 | AC | $0.33 \pm 0.16$ | $0.34 \pm 0.29$ | 0．35 $\pm 0.20$ | $0.34 \pm 0.22$ | $0.41 \pm 0.15$ | 0．49 0.3 .35 | 0．44＊0．23 | 0．45 $\pm 0.24$ | 0．31 +0.16 | 0．30 0.12 | $0.37 \pm 0.16$ | 0．33 $\pm 0.04$ |
| 02 | PC | 0．34 +0.19 | 0．35＋0．18 | 0．33 $\pm 0.14$ | $0.34 \pm 0.17$ | 0．43 $\pm 0.22$ | 0．42 $\pm .13$ | 0．39 +0.17 | $0.41 \pm 0.17$ | 0．33 $\pm .10$ | 0．39 0.13 | $0.34 \pm 0.13$ | $0.35 \pm 0.03$ |
| 03 | infracollicular sulcus | $1.25 \pm 0.47$ | $1.22 \pm 0.48$ | 1．08さ0．46 | $1.17 \pm 0.47$ | 0．93 $\pm 0.36$ | $0.70 \pm 0.32$ | 0．70 $\pm 0.40$ | $0.78 \pm 0.36$ | 1．600．72 | 1．67 $\pm 0.76$ | 1．47 $\pm 0.75$ | 1．58＋0． |
| 04 | PMJ | 0．83土0．47 | 0．97 $\pm 0.65$ | 0．84土0．52 | 0．87 $\pm 0.54$ | 0．80＾0．23 | 0．89 0.49 | 0．76 0.28 | $0.81 \pm 0.34$ | 1．06ะ0．50 | 1．23 20.74 | 1．17 ${ }^{\text {a }}$ ．55 | $1.15 \pm 0.08$ |
| 05 | superior interpeduncular fossa | $1.15 \pm 0.61$ | $0.96 \pm 0.60$ | $1.12 \pm 0.50$ | $1.08 \pm 0.57$ | $1.04 \pm 0.37$ | $1.02 \pm 0.61$ | $1.00 \pm 0.51$ | $1.02 \pm 0.48$ | $1.38 \pm 0.83$ | $1.07 \pm 0.59$ | $1.20 \pm 0.63$ | $1.22 \pm 0.1$ |
| 06 | R superior LMS | 0．75 0.4 .48 | $1.16 \pm 0.69$ | 0．68＊0．50 | $0.85 \pm 0.59$ | 1．07 $\pm 0.38$ | $1.44 \pm 0.50$ | 0．91 0.46 | 1．14土0．48 | $0.63 \pm 0.28$ | $1.17 \pm 0.59$ | 0．55 $\pm 0.34$ | $0.78 \pm 0.34$ |
| 07 | L superior LMS | 0．93＋0．59 | 1．05＋0．57 | 0．91 0.90 | 0．96 0.71 | 1．18さ0．32 | $1.04 \pm 0.44$ | $1.27 \pm 0.87$ | $1.16 \pm 0.58$ | 1．03＋0．46 | $1.25 \pm 0.68$ | 0．62 0.31 | 0．97 $\pm 0.32$ |
| 08 | R inferior LMS | 1．55 $\pm 1.14$ | $1.61 \pm 1.07$ | $1.47 \pm 0.96$ | 1．54土1．05 | 1．81土1．07 | $1.49 \pm 0.77$ | $1.35 \pm 0.81$ | 1．55 0.88 | 1．65 1.08 | 2．06 1.116 | 1．87 1.28 | $1.86 \pm 0.21$ |
| 09 | L inferior LMS | 1．39 +1.11 | $1.79 \pm 1.32$ | $1.63 \pm 1.19$ | $1.60 \pm 1.21$ | 1．66＋0．91 | $1.88 \pm 1.37$ | 1．60ı1．22 | $1.71 \pm 1.14$ | 1．53 1.07 | $2.06 \pm 1.29$ | 2．05 1．45 | $1.88 \pm 0.31$ |
| 10 | culmen | 1．03＋0．73 | 0．68＋0．24 | 0．61 0.32 | $0.77 \pm 0.50$ | $1.16 \pm 0.70$ | 0．72 $\pm .22$ | 0．61 $\pm .25$ | 0．83 0.49 | 1．100．41 | 0．77 $\pm 0.25$ | 0．70 $\pm .29$ | 0．85 $\pm .21$ |
| 11 | intermammillary sulcus | 0．73 $\pm 0.34$ | 0．68＋0．34 | $0.70 \pm 0.38$ | 0．70 0.35 | 0．69き0．41 | 0．74 $\pm 0.41$ | 0．83 $\pm 0.47$ | 0．76 0.42 | $0.82 \pm 0.39$ | 0．72 $\pm 0.31$ | 0．68 $\pm 0.36$ | 0．74 $\pm 0.07$ |
| 12 | R mb | $0.37 \pm 0.28$ | $0.44 \pm 0.32$ | $0.48 \pm 0.34$ | 0．44土0．31 | 0．51 $\pm 0.31$ | $0.47 \pm 0.14$ | 0．54 $\pm 0.34$ | $0.51 \pm 0.27$ | 0．32 +0.18 | $0.51 \pm 0.42$ | 0．51 $\pm 0.39$ | 0．45＋0．11 |
| 13 | LMB | 0．43＊0．27 | 0．53 $\pm 0.32$ | 0．50 $\pm 0.31$ | 0．49 5.30 | $0.52 \pm 0.29$ | 0．50 0.15 | 0．58 $\pm .28$ | 0．53 $\pm 0.24$ | 0．400．19 | $0.63 \pm 0.45$ | 0．52 $\pm 0.33$ | 0．52 $\pm .12$ |
| 14 | pineal gland | 0．7000．33 | 0．9440．33 | 0．68土0．51 | $0.77 \pm 0.42$ | $0.91 \pm 0.24$ | $1.16 \pm 0.37$ | 0．83 $\pm 0.57$ | 0．97 $\pm .42$ | $0.63 \pm 0.25$ | 0．70 0.41 | $0.68 \pm 0.35$ | $0.67 \pm 0.04$ |
| 15 | R LV at AC | 0．99 1.4 .48 | 0．68＊0．42 | 0．62 0.50 | $0.75 \pm 0.92$ | 1．29土1．50 | 0．74 $\pm 0.41$ | 0．74 +0.45 | 0．92 $\pm 0.93$ | 1．1000．66 | 0．81 $\pm 0.35$ | 0．73 $\pm 0.30$ | $0.88 \pm 0.20$ |
| 16 | Llv at AC | ${ }^{1.06+1.60}$ | 0．73 +0.42 | $0.62 \pm 0.51$ | 0．79 0.98 | 1．3441．50 | $0.76 \pm 0.33$ | 0．78 0.441 | 0．96 0.92 | $1.31 \pm 1.09$ | 0．91 0.35 | 0．77 $\pm 0.32$ | 0．99＋0．28 |
| 17 | R LV at PC | 1．13＋1．35 | 1．12＋1．01 | $1.00 \pm 0.60$ | $1.08 \pm 1.00$ | 1．35 1.35 | 1．19＊1．04 | 0．90 0.60 | 1．14＊1．01 | $1.29 \pm 0.65$ | $1.38 \pm 0.59$ | $1.32 \pm 0.58$ | $1.33 \pm 0.05$ |
| 18 | LLV at PC | $1.23 \pm 1.46$ | $1.32+1.02$ | 1．03 0.58 | 1．18 $\times 1.05$ | 1．48土1．29 | $1.18 \pm 1.07$ | $0.91 \pm 0.54$ | $1.19 \pm 1.00$ | 1．50⒈04 | $1.65 \pm 0.72$ | $1.40 \pm 0.61$ | $1.52 \pm 0.13$ |
| 19 | genu of CC | 1．00＾0．46 | $0.63 \pm 0.24$ | 0．78 0.48 | 0．80 0.44 | 1．100．66 | 0．62 $\pm .21$ | 0．84＊0．42 | 0．85 0.49 | 0．99＋0．46 | $0.75 \pm 0.28$ | 0．99 0.48 | 0．91 0.14 |
| 20 | splenium | 0．71 $\pm 0.39$ | 0．52＋0．27 | 0．80 1.10 | $0.68 \pm 0.73$ | 0．90 0.40 | 0．69 0.21 | $0.86 \pm 0.52$ | 0．81 0.3 .39 | $0.67 \pm 0.32$ | $0.47 \pm 0.15$ | 0．67 $\pm 0.29$ | 0．60 0.11 |
| 21 | R AL temporal horn | ${ }^{1.44 \pm 1.20}$ | 1．52＋0．79 | $1.15 \pm 0.89$ | $1.36 \pm 0.98$ | $1.55 \pm 1.26$ | $1.71 \pm 0.59$ | 1．33土1．12 | 1．53 $\pm 1.00$ | $1.72 \pm 1.00$ | $1.65 \pm 0.80$ | $1.32 \pm 0.67$ | $1.56 \pm 0.21$ |
| 22 | $\begin{array}{\|l} \hline \begin{array}{l} \text { LAL temporal } \\ \text { horn } \end{array} \\ \hline \end{array}$ | $1.32 \pm 0.91$ | $1.10 \pm 0.56$ | 1．16さ0．94 | $1.19 \pm 0.82$ | $1.32 \pm 1.07$ | $1.29 \pm 0.39$ | $1.49 \pm 0.93$ | $1.36 \pm 0.82$ | 1．46 $\pm 0.91$ | $1.15 \pm 0.52$ | $1.31 \pm 0.62$ | $1.30 \pm 0.16$ |
| 23 | R superior AM temporal horn | ${ }^{0.62 \pm 0.38}$ | $1.31 \pm 1.71$ | 0．83土0．91 | 0．91 11.15 | 0．70 0.36 | 1．73 11.90 | $0.72 \pm 0.33$ | 1．05 1.19 | $0.69 \pm 0.37$ | $1.35 \pm 0.74$ | 0．80 0.34 | $0.95 \pm 0.35$ |
| 24 | L superior AM temporal horn | 0．59＋0．39 | 2．02＋1．90 | $0.95 \pm 0.98$ | 1．17 11.36 | $0.66 \pm 0.31$ | $2.42 \pm 2.12$ | $0.80 \pm 0.26$ | $1.29 \pm 1.44$ | $0.71 \pm 0.36$ | $2.08 \pm 1.07$ | $0.89 \pm 0.32$ | $1.23 \pm 0.7$ |
| 25 | R inferior AM temporal horn | $1.31 \pm 1.20$ | 1．49＋0．94 | $1.40 \pm 1.00$ | 1．40＋1．04 | $1.65 \pm 0.97$ | $1.36 \pm 0.81$ | $1.44 \pm 0.68$ | $1.48 \pm 0.80$ | $1.55 \pm 0.86$ | 1．80さ1．14 | 1．87 1.25 | $1.74 \pm 0.17$ |
| 26 | L inferior AM temporal horn | ${ }^{1.36 \pm \pm 1.16}$ | $1.41 \pm 1.06$ | $1.39 \pm 0.76$ | $1.38 \pm 0.98$ | $1.56 \pm 0.94$ | $1.37 \pm 0.99$ | $1.40 \pm 0.88$ | $1.44 \pm 0.90$ | 1．67 $\pm 0.88$ | $1.70 \pm 1.21$ | $1.67 \pm 0.78$ | 1．68＋0．0 |
| 27 | R indusium griseum origin | $1.38 \pm 0.75$ | 1．70さ1．08 | $1.26 \pm 0.82$ | $1.43 \pm 0.90$ | $1.10 \pm 0.54$ | $1.55 \pm 1.00$ | ${ }^{1.18 \pm 0.86}$ | $1.28 \pm 0.81$ | $1.71 \pm 1.08$ | 2．00 1.17 | $1.35 \pm 0.78$ | 1．69＋0．3 |
| 28 | L indusium griseum origin | 1．52＋1．14 | $2.11 \pm 1.44$ | $1.40 \pm 0.98$ | $1.66 \pm 1.22$ | $1.37 \pm 0.74$ | 1．61 1.32 | $1.45 \pm 0.75$ | $1.47 \pm 0.94$ | $2.12 \pm 1.35$ | $2.74 \pm 1.76$ | $1.59 \pm 0.90$ | $2.15 \pm 0.57$ |
| 29 | $\begin{aligned} & \mathrm{R} \text { ventral occipital } \\ & \text { horn } \end{aligned}$ | $1.59 \pm 1.07$ | 0．80さ0．45 | $1.34 \pm 1.25$ | $1.30 \pm 1.08$ | $1.85 \pm 1.28$ | $0.95 \pm 0.24$ | 2．08さ1．63 | 1．69＋1．29 | $1.58 \pm 0.85$ | $0.93 \pm 0.63$ | $1.13 \pm 0.59$ | $1.21 \pm 0.33$ |
| 30 | L ventral occipital horn | 1．09＋1．13 | $1.63 \pm 2.94$ | $1.25 \pm 1.32$ | $1.28 \pm 1.78$ | 1．22＋1．15 | $0.86 \pm 0.31$ | $1.90 \pm 1.45$ | $1.37 \pm 1.16$ | $1.36 \pm 0.88$ | $4.98 \pm 6.77$ | $1.12 \pm 0.60$ | 2．49＋2．16 |
| 31 | R olfactory sulcal fundus | $1.17 \pm 0.68$ | $1.41 \pm 0.95$ | $1.14 \pm 0.59$ | $1.23 \pm 0.75$ | $1.44 \pm 0.75$ | $1.91 \pm 0.65$ | $1.23 \pm 0.56$ | $1.53 \pm 0.69$ | 0．96 0.51 | $1.35 \pm 0.68$ | $1.20 \pm 0.69$ | $1.17 \pm 0.19$ |
| 32 | L olfactory sulcal fundus | $1.23 \pm 0.54$ | $1.41 \pm 1.00$ | 1．13土0．63 | 1．25 $\times 0.74$ | $1.24 \pm 0.57$ | 1．56さ1．13 | 1．07 $\pm 0.43$ | 1．29 20.77 | 1．29＋0．71 | 1．41 1.00 | $1.27 \pm 0.66$ | $1.32 \pm 0.08$ |

ANOVA for Templates
A difference in placement error between templates was identified by ANOVA．

2/24/2019
PHASE1_template_validation

| 'F-value: 7.88; p-value: 0.0004 |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| fid | Fval | pval | adjusted | significant |
| 1 | 0.03 | 0.9695 | 0.9767 | FALSE |
| 2 | 0.13 | 0.8760 | 0.9490 | False |
| 3 | 1.45 | 0.2406 | 0.5918 | FALSE |
| 4 | 0.72 | 0.4900 | 0.7215 | FALSE |
| 5 | 1.01 | 0.3696 | 0.6571 | FALSE |
| 6 | 7.28 | 0.0011 | 0.0119 | TRUE |
| 7 | 0.38 | 0.6840 | 0.8755 | FALSE |
| 8 | 0.16 | 0.8535 | 0.9490 | FALSE |
| 9 | 0.90 | 0.4118 | 0.6935 | FALSE |
| 10 | 7.61 | 0.0008 | 0.0119 | true |
| 11 | 0.12 | 0.8897 | 0.9490 | FALSE |
| 12 | 1.05 | 0.3546 | 0.6571 | FALSE |
| 13 | 0.84 | 0.4362 | 0.6979 | FALSE |
| 14 | 4.04 | 0.0206 | 0.1319 | FALSE |
| 15 | 1.57 | 0.2124 | 0.5918 | FALSE |
| 16 | 1.83 | 0.1659 | 0.5310 | FALSE |
| 17 | 0.17 | 0.8398 | 0.9490 | FALSE |
| 18 | 0.71 | 0.4960 | 0.7215 | FALSE |
| 19 | 6.38 | 0.0025 | 0.0198 | TRUE |
| 20 | 1.30 | 0.2779 | 0.5918 | FALSE |
| 21 | 1.39 | 0.2530 | 0.5918 | FALSE |
| 22 | 0.61 | 0.5467 | 0.7289 | FALSE |
| 23 | 3.19 | 0.0456 | 0.1826 | FALSE |
| 24 | 11.61 | 0.0000 | 0.0009 | true |
| 25 | 0.24 | 0.7905 | 0.9490 | FALSE |
| 26 | 0.02 | 0.9767 | 0.9767 | FALSE |
| 27 | 2.22 | 0.1142 | 0.4060 | FALSE |
| 28 | 3.32 | 0.0401 | 0.1826 | FALSE |
| 29 | 3.76 | 0.0271 | 0.1443 | FALSE |
| 30 | 0.61 | 0.5433 | 0.7289 | FALSE |
| 31 | 1.28 | 0.2819 | 0.5918 | FALSE |
| 32 | 1.23 | 0.2959 | 0.5918 | FALSE |

## K-means clustering of point cloud distributions

Across all templates; and template specific


Agile12v2016 only


Colin27 only



All templates combined


```
2/24/2019
R version 3.5.1 (2018-07-02)
Matrix products: default
LAPACK: /System/Library/Frameworks/Accelerate.framework/versions/A/Frameworks/vecLib.framework/versions/A/libLAPACK.dylib
locale: 
l
M
```



```
10aded via a namespace (and not attached):
```



```
llll
llll
lal
lll
lll
```


## A. 2 Phase 2: Supplementary Material

Phase 2: Protocol Validation for Individual Subjects
This notebook contains results validating the AFID32 protocol on individual subjects from the OASIS-1 databank.

OAS1 Subset: Demographics
Demographics here.
'Total: $58.0+/$ - 17.9 years; Range: $25-91$
'Female: 17/30 (56.7\%)'
'Total: $1.28+$ +- $\mathbf{3} .03 \mathrm{~mm}$; Outliers: $28 / 2880(0.97 \%)^{\prime}$

|  | fid | subject | mri_session | name | description | ean_A |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 137 | 29 | OAS1_0203 | MR1 | 29 | R ventral occipital horn | 16.19882 |
| 1501 | 29 | OAS1_0216 | MR1 | 29 | R ventral occipital horn | 17.77257 |
| 1502 | 30 | OAS1_0216 | MR1 | 30 | L ventral occipital horn | 11.61197 |
| 2043 | 27 | OAS1_0256 | MR1 | 28 | Lindusium griseum origin | 15.80488 |
| 2044 | 28 | OAS1_0256 | MR1 | 27 | $R$ indusium griseum origin | 15.35560 |
| 2141 | 29 | OAS1_0263 | MR1 | 29 | R ventral occipital horn | 39.35092 |
| 2142 | 30 | OAS1_0263 | MR1 | 30 | L ventral occipital horn | .6437 |
| 2173 | 29 | OAS1_0263 | MR1 | 29 | R ventral occipital horn | 78.744 |
| 2174 | 30 | OAS1_0263 | MR1 | 30 | L ventral occipital horn | 80.42163 |
| 2205 | 29 | OAS1_0263 | MR1 | 29 | R ventral occipital horn | 39.39868 |
| 2206 | 30 | OAS1_0263 | MR1 | 30 | L ventral occipital horn | 39.79291 |
| 2235 | 27 | OAS1_0266 | MR1 | 27 | R indusium griseum origin | 23.44415 |
| 2236 | 28 | OAS1_0266 | MR1 | 28 | L indusium griseum origin | 24.30401 |
| 2267 | 27 | OAS1_0266 | MR1 | 27 | R indusium griseum origin | 10.56158 |
| 2268 | 28 | OAS1_0266 | MR1 | 28 | L indusium griseum origin | 12.044 |
| 2299 | 27 | OAS1_0266 | MR1 | 27 | $R$ indusium griseum origin | 12.987 |
| 2300 | 28 | OAS1_0266 | MR1 | 28 | Lindusium griseum orig | 12.35749 |
| 2534 | 6 | OAS1_0303 | MR1 | 6 | R superior LMS | 2487 |
| 2535 | 7 | OAS1_0303 | MR1 | 7 | L superior LMS | 13.9873 |
| 2653 | 29 | OAS1_0343 | MR1 | 29 | R ventral occipital horn | 15.8310 |
| 2942 | 30 | OAS1_0365 | MR1 | 30 | L ventral occipital horn | . 929 |
| 3387 | 27 | OAS1_0456 | MR1 | 27 | R indusium griseum origin | 23.38522 |
| 3388 | 28 | OAS1_0456 | MR1 | 28 | L indusium griseum origin | 23.76189 |
| 3390 | 30 | OAS1_0456 | MR1 | 30 | L ventral occipital horn | 17.74944 |
| 3419 | 27 | OAS1_0456 | MR1 | 27 | $R$ induseum griseum origin | 10.64591 |
| 20 | 28 | OAS1_0456 | MR1 | 28 | L induseum griseum origin | 10.43077 |
| 3451 | 27 | OAS1_0456 | MR1 | 27 | R indusium griseum origin | 12.883 |
| 3452 | 28 | OAS1_0 | M | 28 | L indusium griseum origin | 13.5399 |

Individual Subject Results: Post-QC
Re-analysis after quality control and filtering of outiers.
'Total: $0.94+/-0.73 \mathrm{~mm}$; Outiers: $0 / 2872$ ( $0.00 \%$ ) ${ }^{\prime}$

Inter-Rater AFLE
'Total: $1.58+/-1.02 \mathrm{~mm}$

| AFID | Description | Mean AFLE Pre-QC | Mean AFLE Post-QC | Inter-Rater AFLE Post-QC |
| :---: | :---: | :---: | :---: | :---: |
| 01 | AC | $0.36 \pm 0.21$ (1.29) | $0.36 \pm 0.21$ (1.29) | $0.60 \pm 0.25$ (1.38) |
| 02 | PC | $0.34 \pm 0.16$ (0.88) | $0.34 \pm 0.16$ (0.88) | 0.57 $\pm .21$ (1.22) |
| 03 | infracollicular sulcus | $0.78 \pm 0.48$ (3.07) | $0.78 \pm 0.48$ (3.07) | $1.34 \pm 0.64$ (3.84) |
| 04 | PMJ | 0.83+0.49 (2.44) | 0.83+0.49 (2.44) | $1.41 \pm 0.55$ (2.55) |
| 05 | superior interpeduncular fossa | $1.20 \pm 0.75$ (3.50) | $1.20 \pm 0.75$ (3.50) | $2.04 \pm 0.90$ (4.25) |
| 06 | R superior LMS | $1.30 \pm 1.74$ (14.25) | $1.01 \pm 0.55$ (2.85) | $1.70 \pm 0.68$ (3.13) |
| 07 | L superior LMS | 1.36+1.71 (13.99) | $1.06 \pm 0.61$ (3.45) | $1.72 \pm 0.71$ (3.89) |
| 08 | R inferior LMS | 1.13 1.0 .75 (5.13) | $1.03 \pm 0.57$ (2.99) | $1.77 \pm 0.74$ (3.43) |
| 09 | L inferior LMS | $1.10 \pm 0.80$ (5.31) | $1.01 \pm 0.62$ (2.72) | $1.71 \pm 0.86$ (3.71) |
| 10 | culmen | 0.99+0.99 (5.66) | $0.83 \pm 0.62$ (3.07) | $1.35 \pm 0.82$ (3.42) |
| 11 | intermammillary sulcus | 0.600.31 (1.62) | 0.600.31 (1.62) | $1.02 \pm 0.41$ (1.86) |
| 12 | R MB | $0.40 \pm 0.23$ (1.11) | $0.40 \pm 0.23$ (1.11) | $0.69 \pm 0.32$ (1.52) |
| 13 | Lmb | $0.36 \pm 0.20$ (1.20) | $0.36 \pm 0.20$ (1.20) | $0.62 \pm 0.29$ (1.62) |
| 14 | pineal gland | $0.68 \pm 0.47$ (1.98) | $0.68 \pm 0.47$ (1.98) | $1.16 \pm 0.69$ (2.63) |
| 15 | R LV at AC | $1.00 \pm 0.90$ (5.28) | $0.91 \pm 0.72$ (4.45) | $1.55 \pm 1.08$ (5.86) |
| 16 | Llvat AC | $1.01 \pm 0.80$ (4.53) | 0.94+0.70 (4.53) | $1.60 \pm 1.08$ (5.47) |
| 17 | R LV at PC | 0.92+0.54 (3.42) | 0.92+0.54 (3.42) | $1.54 \pm 0.77$ (3.84) |
| 18 | LLVat PC | $0.87 \pm 0.42$ (2.20) | $0.87 \pm 0.42$ (2.20) | $1.46 \pm 0.55$ (2.80) |
| 19 | genu of CC | 0.97+0.81 (5.16) | 0.89+0.63 (3.69) | $1.50 \pm 0.89$ (4.30) |
| 20 | splenium | 0.54 40.25 (1.24) | 0.54 00.25 (1.24) | 0.91 0.35 (1.66) |
| 21 | R AL temporal horn | $1.44 \pm 1.09$ (7.01) | $1.30 \pm 0.86$ (4.45) | $2.21 \pm 1.13$ (5.92) |
| 22 | LAL temporal horn | 1.22+0.77 (4.11) | $1.22 \pm 0.77$ (4.11) | $2.04 \pm 1.01$ (4.47) |
| 23 | R superior AM temporal horn | $1.28 \pm 1.27$ (8.22) | 1.12+0.88 (4.69) | 1.86 $\pm 1.19$ (4.97) |
| 24 | L superior AM temporal horn | 1.09+1.22 (7.54) | $0.83 \pm 0.61$ (3.66) | $1.39 \pm 0.85$ (4.60) |
| 25 | R inferior AM temporal horn | $1.69 \pm 1.43$ (9.03) | $1.44 \pm 0.91$ (4.72) | $2.39 \pm 1.23$ (5.07) |
| 26 | Linferior AM temporal horn | $1.99 \pm 1.75$ (8.79) | $1.49 \pm 1.09$ (4.70) | $2.42 \pm 1.47$ (6.64) |
| 27 | R indusium griseum origin | $3.13 \pm 4.19$ (23.44) | $1.77 \pm 0.99$ (4.77) | 2.95+1.20 (5.75) |
| 28 | L indusium griseum origin | $2.99 \pm 4.30$ (24.30) | $1.68 \pm 1.00$ (5.00) | 2.75 1.2 .29 (5.78) |
| 29 | R ventral occipital horn | $3.64 \pm 10.36$ (78.74) | 0.69+0.39 (2.11) | $1.14 \pm 0.54$ (2.53) |
| 30 | L ventral occipitial horn | $3.43 \pm 10.38$ (80.42) | 0.86 0.0 .67 (4.94) | $1.39 \pm 0.98$ (5.72) |
| 31 | R olfactory sulcal fundus | $0.99 \pm 0.53$ (2.29) | 0.99+0.53 (2.29) | $1.71 \pm 0.60$ (2.84) |
| 32 | L olfactory sulcal fundus | $1.21 \pm 0.74$ (4.53) | $1.21 \pm 0.74$ (4.53) | $2.11 \pm 0.92$ (5.81) |

## Secondary Analyses

We evaluated whether there was any evidence of an effect of demographics on AFLE.

| (Intercept) | 0.7694 | $0+00$ |
| :--- | :--- | :--- |
| age |  |  |


| age | 0.0030 | $1 e-04$ |
| :--- | :--- | :--- |

Did AFLE worsen with the age of the subject for specific AFIDs?
We wanted to see if specific AFIDs tended to worsen with age of the OAS1 participant scan. Worsened for AFID17-18: bilateral LV at PC.

| fid | (Intercept) | pval_(Intercept) | age | pval_session | pval_session_adjusted | pval_session_significant |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 0.19 | 0.0102 | 0.00 | 0.0133 | 0.1422 | FALSE |
| 2 | 0.24 | 0.0001 | 0.00 | 0.0964 | 0.3426 | FALSE |
| 3 | 0.93 | 0.0000 | 0.00 | 0.3885 | 0.5920 | FALSE |
| 4 | 0.86 | 0.0000 | 0.00 | 0.8868 | 0.9063 | FALSE |
| 5 | 0.81 | 0.0033 | 0.01 | 0.1364 | 0.4095 | FALSE |
| 6 | 1.24 | 0.0000 | 0.00 | 0.2292 | 0.4584 | FALSE |
| 7 | 0.66 | 0.0035 | 0.01 | 0.0572 | 0.2466 | FALSE |
| 8 | 0.79 | 0.0003 | 0.00 | 0.2276 | 0.4584 | FALSE |
| 9 | 0.60 | 0.0074 | 0.01 | 0.0557 | 0.2466 | FALSE |
| 10 | 10.61 | 0.0075 | 0.00 | 0.3133 | 0.5321 | FALSE |
| 11 | 0.67 | 0.0000 | 0.00 | 0.5306 | 0.7075 | FALSE |
| 12 | 12 0.52 | 0.0000 | 0.00 | 0.1408 | 0.4095 | FALSE |
| 13 | 13 0.42 | 0.0000 | 0.00 | 0.4399 | 0.6120 | FALSE |
| 14 | 14.73 | 0.0001 | 0.00 | 0.7578 | 0.8362 | FALSE |
| 15 |  | 0.0025 | 0.00 | 0.7391 | 0.8362 | FALSE |
| 16 | 6 0.88 | 0.0008 | 0.00 | 0.8194 | 0.8741 | FALSE |
| 17 | 170.18 | 0.3163 | 0.01 | 0.0000 | 0.0013 | true |
| 18 | 8.44 | 0.0030 | 0.01 | 0.0029 | 0.0461 | true |
| 19 | 0.92 | 0.0002 | 0.00 | 0.9063 | 0.9063 | FALSE |
| 20 | 0.44 | 0.0000 | 0.00 | 0.2772 | 0.5217 | FALSE |
| 21 | 10.92 | 0.0043 | 0.01 | 0.2049 | 0.4584 | FALSE |
| 22 | 12.14 | 0.0001 | 0.00 | 0.7487 | 0.8362 | FALSE |
| 23 | 0.99 | 0.0029 | 0.00 | 0.6622 | 0.8150 | FALSE |
| 24 | 2 0.62 | 0.0057 | 0.00 | 0.3294 | 0.5321 | FALSE |
| 25 | 1927 | 0.0002 | 0.00 | 0.5861 | 0.7502 | FALSE |
| 26 | 1.86 | 0.0000 | -0.01 | 0.3325 | 0.5321 | FALSE |
| 27 | 1.13 | 0.0021 | 0.01 | 0.0616 | 0.2466 | FALSE |
| 28 | 1.25 | 0.0008 | 0.01 | 0.2183 | 0.4584 | FALSE |
| 29 | 0.41 | 0.0039 | 0.00 | 0.0404 | 0.2466 | FALSE |
| 30 | 0.36 | 0.1322 | 0.01 | 0.0345 | 0.2466 | FALSE |
| 31 | 10.84 | 0.0000 | 0.00 | 0.4201 | 0.6111 | FALSE |
| 32 | 0.84 | 0.0022 | 0.01 | 0.1563 | 0.4167 | FALSE |
| R version 3.5.1 (2018-07-02) <br> Platform: x86_64-apple-darwin14.5.0 (64-bit) |  |  |  |  |  |  |
|  |  |  |  |  |  |  |
| Running under: macos High Sierra 10.13 .2 |  |  |  |  |  |  |
| Matrix products: default ${ }_{\text {BLAS }}$ /System/Library/Frameworks/Accelerate.framework/Versions/A/Frameworks/vecLi |  |  |  |  |  |  |
|  |  |  |  |  |  |  |
| LAPACK: /System/Library/Frameworks/Accelerate.framework/versions/A/Frameworks/vecLib.framework/versions/A/libLAPACK.dylib |  |  |  |  |  |  |
| 1ocale: |  |  |  |  |  |  |
| [1] en_CA.UTF-8/en_CA.UTF-8/en_CA.UTF-8/C/en_CA.UTF-8/en_CA.UTF-8 |  |  |  |  |  |  |
| attached base packages: |  |  |  |  |  |  |
|  | ] stats | graphics gr | Devic | ces utils | datasets methods | base |
| other attached packages: |  |  |  |  |  |  |
|  |  |  |  |  |  |  |
| loaded via a namespace (and not attached): |  |  |  |  |  |  |
|  | 1] Rcpp_0. | 12.17 comp | compiler_3.5.1 pi |  | lar_1.3.0 bindr_ | bindr_0.1.1 |
|  | 51) base64en | nc-0.1-3 tools |  |  | $\mathrm{d}^{-0.1-2}$ json1i ${ }^{\text {a }}$ | ${ }^{\text {jsonlite_1.5 }}$ |
|  | 9) evaluate |  | IRdisplay 0 0.5.0 IR |  |  | bindrcpp_0.2.2 |
|  | ${ }^{7} 1$ repr_0 0 | 15.0 withr |  |  | ingr_1.3.1 dplyr | ${ }_{\text {dplyr_0 }}$ |
|  | 11 grid_3. | 5.1 tidy | tidyselect_0.2.4 glle |  | e_1.3.0 R6_2.2. | ${ }_{\text {R6_2 }}$ 2.2.2 2 |
|  | $5]$ pbdzMe_ | 0.3-3 purr | r_0.2 | 2.5 magr | rittr_1.5 scales | colorspace_1.3-2 |
|  | 9] htmltoo | 1s_0.3.6 misc | misc3d_0.8-4 ass |  | ertthat_0.2.0 colors |  |
|  | 3] stringi_ | -1.2.4 lazy |  |  | sell_0.5.0 crayon | n_1.3.4 |

## A. 3 Phase 3: Supplementary Material

```
2/24/2019

Phase 3: Subject-to-Template Evaluation
```

This notebook compares voxel overlap measures against AFID-based metrics for evaluating spatial correspondence. The OASIS-1 dataset from from PHASE2 was processed using the Ants-based T1-to-MN
Attaching package: 'dplyr'
The following objects are masked from 'package:plyr':
arrange, count, desc, failwith, id, mutate, rename, summarise,
summarize
The following objects are masked from 'package:stats':
filter, lag
The following objects are masked from 'package:base':
intersect, setdiff, setequal, union
Loading required package: magrittr
Attaching package: 'ggpubr'
The following object is masked from 'package:plyr':
mutate
ROI Overlap
Values for pallidum, striatum, and thalamus.

| roi | side | jaccard_lin | jaccard_nlin | jaccard_lin_vs_nlin | kappa _lin | kappa_nlin | kappa_lin_vs_nlin |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| pallidum | left | 0.54*0.13 | 0.800.03 | * | 0.69+0.11 | 0.89+0.02 | . |
| pallidum | right | 0.55 0.12 | 0.79+0.05 | * | 0.7000.11 | 0.88+0.03 | - |
| striatum | left | 0.53*0.14 | 0.83 0.03 | . | 0.68+0.13 | 0.91 0.02 | - |
| striatum | right | 0.55 $\pm 0.15$ | 0.82 $\pm 0.05$ | * | 0.7000.13 | 0.90 0.03 | . |
| thalamus | left | 0.7000.11 | 0.86 0.003 | * | 0.82+0.08 | 0.93 0.0 .02 | . |
| thalamus | right | 0.69 0.11 | 0.87 $\pm 0.03$ | . | $0.81 \pm 0.08$ | 0.93 0.02 | . |

```


AFRE
 OAS1_0109 which accounts for the maximally error observed in this analysis of \(>30 \mathrm{~mm}\) AFRE

\section*{Nonlinear Transform Results}
\[
\begin{aligned}
& \text { 'Total: } 1.80+\text { +/ } 2.09 \mathrm{~mm} \text {; Range: } 0.07-32.78 \text { ' } \\
& \text { 'Mean Max: } 7.55 \mathrm{~mm} \text { ' }
\end{aligned}
\]

\section*{Linear Transform Results}
'Total: \(3.40+/-2.55 \mathrm{~mm}\); Range: \(0.28-36.26\); Mean Max: '
'Mean Max: 10.25 mm'
Wilcoxon rank sum test with continuity correction

\(\mathrm{W}=716930, \mathrm{p}\)-value \(<2.2 \mathrm{e}-16\)
alternative hypothesis: true 10 ocation shift is not equal to
\begin{tabular}{|c|c|c|c|c|}
\hline AFID & Description & Mean AFRE lin & Mean AFRE lin & lin vs nlin \\
\hline 01 & \({ }^{\text {AC }}\) & \(2.15 \pm 0.97\) (4.96) & 0.36 \(\pm 0.21\) (0.99) & . \\
\hline 02 & PC & \(1.83 \pm 0.96\) (4.58) & 0.57 0.29 (1.64) & . \\
\hline 03 & infracollicular sulcus & \({ }^{2.20 \pm 1.23(5.71)}\) & 0.93+0.53 (2.11) & . \\
\hline 04 & PMJ & 2.50⒈36 (6.06) & 0.68+0.43 (2.13) & . \\
\hline 05 & superior interpeduncular fossa & \(2.35 \pm 1.06\) (4.75) & 0.76 00.37 (1.69) & . \\
\hline 06 & R superior LMS & 2.07 \(\pm 0.95\) (4.32) & 1.17+0.74 (3.52) & . \\
\hline 07 & L superior LMS & 2.03+0.85 (4.22) & 1.43+0.77 (2.88) & * \\
\hline 08 & R inferior LMS & \(2.45 \pm 1.37\) (7.50) & 1.78+1.11 (5.41) & . \\
\hline 09 & L inferior LMS & \(2.54 \pm 1.26\) (6.63) & 1.83+0.96 (3.99) & - \\
\hline 10 & culmen & \(4.50 \pm 2.93\) (12.72) & \(2.73 \pm 2.81\) (10.12) & - \\
\hline 11 & intermammillary sulcus & \(2.81 \pm 1.62\) (6.30) & 1.44+0.60 (2.73) & * \\
\hline 12 & RMB & \(2.72 \pm 1.67\) (6.90) & 0.93+0.48 (1.90) & - \\
\hline 13 & LMB & \(2.84 \pm 1.70\) (6.14) & 1.01 0.62 (2.93) & . \\
\hline 14 & pineal gland & \(2.53 \pm 1.39\) (5.70) & 2.01+1.24 (6.16) & \\
\hline 15 & R LV at AC & \(4.44 \pm 1.84\) (7.90) & 2.70 11.59 (7.85) & - \\
\hline 16 & LLV at AC & \({ }^{4.50 \pm 1.95}\) (8.40) & 2.11 11.72 (7.92) & - \\
\hline 17 & R LV at PC & 4.81 2.54 (10.07) & 2.96+2.42 (9.46) & * \\
\hline 18 & LLV at PC & \(4.80 \pm 2.64\) (10.34) & \(3.01 \pm 2.22\) (8.13) & * \\
\hline 19 & genu of CC & \({ }^{3.73 \pm 1.82}\) (7.88) & 1.56+0.76 (3.32) & * \\
\hline 20 & splenium & \(2.96 \pm 1.88\) (7.57) & 0.97+0.60 (2.93) & * \\
\hline 21 & R AL temporal horn & \({ }^{3.79 \pm 1.71 ~(7.50) ~}\) & 1.70+1.09 (5.23) & * \\
\hline 22 & L AL temporal horn & \({ }^{3.62 \pm 1.45}\) (6.98) & 1.67 1.0 .98 (4.31) & * \\
\hline 23 & R superior AM temporal horn & \(3.34 \pm 1.63\) (7.25) & 1.93+1.34 (6.85) & - \\
\hline 24 & L superior AM temporal horn & \(3.44 \pm 1.80\) (8.20) & 1.67+1.25 (5.80) & * \\
\hline 25 & R inferior AM temporal horn & \(4.02 \pm 1.97\) (8.32) & \(2.41 \pm 1.16\) (5.61) & - \\
\hline 26 & Linferior AM temporal horn & 4.13 \(\pm 1.70\) (8.20) & \(2.21 \pm 1.09\) (4.84) & . \\
\hline 27 & R indusium griseum origin & \(3.36 \pm 2.07\) (8.46) & 2.06+1.49 (6.40) & * \\
\hline 28 & L indusium griseum origin & \(3.60 \pm 1.68\) (8.83) & \(2.05 \pm 1.37\) (5.00) & * \\
\hline 29 & R ventral occipital horn & \({ }^{5.86 \pm 6.32 ~(36.26) ~}\) & 3.44+5.77 (32.78) & - \\
\hline 30 & L ventral occipital horn & 6.99+6.72 (33.74) & 4.51 1 6.28 (29.76) & * \\
\hline 31 & R olfactory sulcal fundus & \(2.83 \pm 1.36\) (7.50) & 1.37+0.95 (3.44) & . \\
\hline 32 & L olfactory sulcal fundus & \(2.94 \pm 1.28\) (6.49) & 1.57 \(\pm 0.84\) (3.41) & * \\
\hline
\end{tabular}


Subject level analysis of lin versus nlin
Revealed 3 subjects where mean AFRE was not statistically different. However, individual afids demonstrated high AFRE.
One subject appeared to be well registered with linear registration alone. The other two had extreme registration errors (over 8 mm AFRE).
\begin{tabular}{|c|c|c|c|}
\hline subject & AFRE_lin & AFRE_nlin & AFRE_qval_significant \\
\hline OAS1_0010 & 4.44土2.06 (11.19) & \(1.82 \pm 1.25\) (5.61) & - \\
\hline OAS1_0086 & \(3.19 \pm 1.41\) (6.40) & \(1.33 \pm 1.14\) (5.66) & - \\
\hline OAS1_0101 & \(3.10 \pm 2.22\) (8.83) & 2.46+2.14 (8.36) & \\
\hline OAS1_0109 & 4.86+8.12 (36.26) & 3.89+7.34 (32.78) & \\
\hline OAS1_0114 & 3.31 \(\pm 2.03\) (8.32) & 1.52 11.21 (5.88) & - \\
\hline OAS1_0117 & \(4.08 \pm 2.15\) (9.76) & 1.74+1.81 (10.12) & . \\
\hline OAS1_0145 & 2.33+1.69 (7.57) & 1.27+1.45 (6.85) & - \\
\hline OAS1_0177 & \(2.84 \pm 1.78\) (7.06) & \(1.54 \pm 0.82\) (2.83) & - \\
\hline OAS1_0180 & 4.08+2.20 (10.13) & 2.52 1.69 (6.45) & - \\
\hline OAS1_0188 & 3.35+1.80 (9.08) & 1.65 +1.21 (5.08) & - \\
\hline OAS1_0200 & \(2.56 \pm 1.45\) (7.88) & \(1.47 \pm 0.96\) (4.84) & - \\
\hline OAS1_0203 & 3.78+3.89 (23.96) & 2.4633.89 (22.40) & \\
\hline OAS1_0216 & \(2.19 \pm 1.61\) (7.58) & 1.73+1.02 (4.48) & \\
\hline OAS1_0239 & 2.89+2.03 (11.68) & 1.56+1.63 (8.55) & * \\
\hline OAS1_0249 & 3.34+1.65 (8.53) & 1.63+1.08 (4.66) & - \\
\hline OAS1_0255 & 3.48+1.77 (6.68) & \(1.29+0.75\) (3.16) & * \\
\hline OAS1_0256 & 4.16 +2.00 (7.90) & 1.52+0.97 (3.56) & * \\
\hline OAS1_0263 & \(3.97 \pm 2.36\) (10.34) & 1.29+0.96 (3.96) & . \\
\hline OAS1_0266 & 3.67 \(\pm 1.18\) (7.50) & \(1.60 \pm 1.22\) (6.16) & * \\
\hline OAS1_0274 & \(2.90 \pm 1.87\) (7.73) & \(1.99+2.34\) (8.13) & * \\
\hline OAS1_0284 & 3.90 \(\pm 2.59\) (13.41) & \(1.84 \pm 1.85\) (8.39) & * \\
\hline OAS1_0303 & 2.70+1.23 (5.41) & \(1.47 \pm 0.85\) (4.03) & . \\
\hline OAS1_0343 & \(3.32 \pm 1.69\) (7.95) & \(2.43 \pm 1.95\) (9.46) & * \\
\hline OAS1_0345 & \(2.31 \pm 1.30\) (6.16) & \(1.57 \pm 1.10\) (4.11) & * \\
\hline OAS1_0357 & 2.58 \(\pm 1.61\) (7.43) & \(1.47 \pm 1.35\) (5.30) & - \\
\hline OAS1_0365 & 4.18+2.07 (9.61) & \(1.65 \pm 1.35\) (6.50) & * \\
\hline OAS1_0371 & 2.64+1.31 (6.92) & \(1.33 \pm 0.82\) (3.81) & - \\
\hline OAS1_0395 & \(3.27 \pm 1.96\) (11.14) & \(1.68 \pm 1.28\) (6.68) & - \\
\hline OAS1_0398 & 3.85 3.04 (12.40) & \(1.81 \pm 1.80\) (9.09) & - \\
\hline OAS1_0456 & 4.64 \(\pm 2.73\) (12.72) & 2.42+2.01 (9.59) & * \\
\hline
\end{tabular}

\footnotetext{
https://mc.manuscriptcentral.com/LongRequest/hbm?DOWNLOAD=TRUE\&PARAMS=xik_R17sEqXQgrEiZcJ3Fi4yQ5vW9sAZQN8zLziy8onBLAmxR1Qvq3mN... 4/9
}
```

2/24/2019
PHASE3_subject_to_template

```

\section*{Comparison of AFRE and Voxel Overlap}

\section*{}
A

D

side \(\circ\) left \(\Delta\) right

\section*{Correlation between AFRE and voxel overlap}

A weak negative correlation was found between AFRE and standard voxel overlap measures at the global dataset level and for each specific ROI in isolation.
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|}
\hline \multicolumn{10}{|l|}{```
data: compare_overlap_AFRE_nlin$jaccard and compare_overlap_AFRE_nlin$AFRE
z=-2.1686, p-value = 0.03011
alternative hypothesis: true tau is not equal to 0
sample estimates:
    tau
-0.01911451
```} \\
\hline \multicolumn{10}{|c|}{Kendall's rank correlation t} \\
\hline \multicolumn{10}{|l|}{\begin{tabular}{l}
data: compare_overlap_AFRE_nlinskappa and compare_overlap_AFRE_nlinsAFRE \\
\(z=-2.1686, p\)-value \(=0.03011\) \\
alternative hypothesis: true tau is not equal to 0 \\
sample estimates: \\
tau \\
-0.01911451
\end{tabular}} \\
\hline \multicolumn{10}{|c|}{Kendal1's rank correlation tau} \\
\hline \multicolumn{10}{|l|}{```
data: compare_overlap_AFRE_nlin$jaccard and compare_overlap_AFRE_nlin$kappa
z=113.2, p-value < 2.2e-16
alternative hypothesis: true tau is not equal to 0
sample estimates:
tau
    1
```} \\
\hline roi & side & AFRE jaccard & AFRE jaccard_pval & AFRE_kappa & AFRE_kappa_pval & AFRE _accard_pval_adjusted & AFRE jaccard_pval_significant & AFRE_kappa_pval_adjusted & AFRE_kappa_pval_signif \\
\hline pallidum & left & -0.021863046 & 0.31813186 & -0.021863046 & 0.31813186 & 0.3817582 & FALSE & 0.3817582 & FAL \\
\hline pallidum & right & -0.045837311 & 0.03634823 & -0.045837311 & 0.03634823 & 0.1090447 & FALSE & 0.1090447 & FALSE \\
\hline striatum & left & -0.034786535 & 0.11219299 & -0.034786535 & 0.11219299 & 0.2243860 & FALSE & 0.2243860 & FALSE \\
\hline striatum & right & -0.049644573 & 0.02339900 & -0.049644573 & 0.02339900 & 0.1090447 & FALSE & 0.1090447 & FALSE \\
\hline thalamus & left & 0.006607498 & 0.76287336 & 0.006607498 & 0.76287336 & 0.7628734 & FALSE & 0.7628734 & FALSE \\
\hline thalamus & right & -0.029857412 & 0.17277515 & -0.029857412 & 0.17277515 & 0.2591627 & FALSE & 0.2591627 & FALSE \\
\hline
\end{tabular}

\section*{For each AFID and ROI}
```

AFID plots along the y-axis.
'Number of significant correlations (individual AFIDs vs voxel overlap): 0/192 (0.0%)'

```

Pallidum

side \(\circ\) left \(\Delta\) right

Striatum

side \(\circ\) left \(\Delta\) right

Thalamus
https://mc.manuscriptcentral.com/LongRequest/hbm?DOWNLOAD=TRUE\&PARAMS=xik_R17sEqXQgrEiZcJ3Fi4yQ5vW9sAZQN8zLziy8onBLAmxR1Qvq3mN... 6/9


\section*{AFLE versus AFRE}

In this section, we examine whe
A positive correlation between AFLE and AFRE was found to be statistically significant although the actual effect size of the correlation was small.
'Num Outilie AFIDs (> 2 s.d. above mean AFLE): 135/960 (14.06\%)'
'Num Unique Outtier AFIDs (>2 s.d. above mean AFLE): \(22 / 32(68.75 \%)^{\prime}\) Kendall's rank correlation tau
data: AFLE_vs_AFRESAFRE and APLE_Vs_APRESmean_AFLE
\(z=6.9796\), p-value \(=2.959 e-12\)
alternative hypothesis: true tau is not equal to 0
alternative hypoth
sample estimates:
\(\underset{0.1504519}{\text { tau }}\)


\section*{Secondary Analyses}

We evaluated whether there was any evidence of an effiect of demographics (e.g. age of the participants being registered) on AFRE. Age resulted in a global AFRE change of \(0.0075 \mathrm{~mm} /\) year (i.e. a small but statistically significant We evaluated wherher there was any evidence of an effece of demographics (e.g. age of the participants being registered) on
effect). No specific AFIDs were found to contribute to this age-related AFRE change after multiple comparisons correction.
\begin{tabular}{|l|l|l|}
\hline (Intercept) & 2.1635 & 0.0000 \\
\hline age & 0.0075 & 0.0191 \\
\hline
\end{tabular}
\begin{tabular}{|l|l|l|l|l|l|l|}
\hline fid & (Intercept) & pval_(lintercept) & age & pval_age & pval_age_adjusted & pval_age_significant \\
\hline 1 & 1.15 & 0.03 & 0.00 & 0.83 & 0.86 & FALSE \\
\hline 2 & 1.22 & 0.01 & 0.00 & 0.96 & 0.96 & FALSE \\
\hline 3 & 1.96 & 0.00 & -0.01 & 0.42 & 0.75 & FALSE \\
\hline 4 & 0.72 & 0.23 & 0.01 & 0.13 & 0.55 & FALSE \\
\hline 5 & 1.77 & 0.00 & 0.00 & 0.66 & 0.83 & FALSE \\
\hline 6 & 1.74 & 0.00 & 0.00 & 0.78 & 0.83 & FALSE \\
\hline 7 & 1.03 & 0.01 & 0.01 & 0.05 & 0.43 & FALSE \\
\hline 8 & 0.62 & 0.25 & 0.03 & 0.01 & 0.16 & FALSE \\
\hline 9 & 1.54 & 0.00 & 0.01 & 0.20 & 0.55 & FALSE \\
\hline 10 & 6.53 & 0.00 & -0.05 & 0.02 & 0.32 & FALSE \\
\hline 11 & 1.41 & 0.03 & 0.01 & 0.23 & 0.56 & FALSE \\
\hline 12 & 0.90 & 0.18 & 0.02 & 0.15 & 0.55 & FALSE \\
\hline 13 & 0.75 & 0.28 & 0.02 & 0.08 & 0.50 & FALSE \\
\hline 14 & 2.07 & 0.00 & 0.00 & 0.72 & 0.83 & FALSE \\
\hline 15 & 2.72 & 0.00 & 0.01 & 0.30 & 0.64 & FALSE \\
\hline 16 & 2.12 & 0.03 & 0.02 & 0.20 & 0.55 & FALSE \\
\hline 17 & 2.32 & 0.05 & 0.03 & 0.16 & 0.55 & FALSE \\
\hline 18 & 1.65 & 0.14 & 0.04 & 0.04 & 0.40 & FALSE \\
\hline 19 & 2.30 & 0.01 & 0.01 & 0.65 & 0.83 & FALSE \\
\hline 20 & 2.34 & 0.00 & -0.01 & 0.61 & 0.83 & FALSE \\
\hline 21 & 2.52 & 0.00 & 0.00 & 0.77 & 0.83 & FALSE \\
\hline 22 & 2.87 & 0.00 & 0.00 & 0.74 & 0.83 & FALSE \\
\hline 23 & 3.42 & 0.00 & -0.01 & 0.26 & 0.60 & FALSE \\
\hline 24 & 2.88 & 0.00 & -0.01 & 0.67 & 0.83 & FALSE \\
\hline 25 & 2.51 & 0.00 & 0.01 & 0.36 & 0.71 & FALSE \\
\hline 26 & 2.21 & 0.01 & 0.02 & 0.19 & 0.55 & FALSE \\
\hline 27 & 3.19 & 0.00 & -0.01 & 0.56 & 0.83 & FALSE \\
\hline 28 & 3.20 & 0.00 & -0.01 & 0.61 & 0.83 & FALSE \\
\hline 29 & 2.43 & 0.38 & 0.04 & 0.40 & 0.75 & FALSE \\
\hline 30 & 2.19 & 0.45 & 0.06 & 0.21 & 0.55 & FALSE \\
\hline 31 & 2.48 & 0.00 & -0.01 & 0.52 & 0.83 & FALSE \\
\hline 32 & 2.47 & 0.00 & 0.00 & 0.70 & 0.83 & FALSE \\
\hline & & 5.03 \\
\hline
\end{tabular}

R version 3.5.1 (2018-07-02)
Platform: x86_64-apple-darwin14.5.0 (64-bit
Running under: macos High Sierra 10. 13.2 ,
Matrix products: default
BLAS: /System/Library/Frameworks/Accelerate.framework/Versions/A/Frameworks/vecLib.framework/Versions/A//ibBLAS.dylib
LAPACK: /System/Library/Frameworks/Accelerate.framework/Versions/A/Frameworks/vecLib.framework/Versions/A/liblAPACK.dylib
locale:
11] en_CA.UTF-8/en_CA.UTF-8/en_CA.UTF-8/C/en_CA.UTF-8/en_CA.UTF-8
attached base packages
[1] stats
11
\({ }^{[11}\) bindrcpp_0.2.2 ggpubr_0.1.8 \(\quad\) magrittr_1.5 \(\quad\) ggplot2_3.0.0 \(\quad\) reshape2_1.4.3
(6) digest_0.6.16 dplyr_0.7.6 plyr_1.

1oaded via a namespace (and not attached)







\section*{A. 4 Phase 4: Supplementary Material}
```

2/24/2019

Phase 4: Template-to-Template Evaluation


PHASE4_template_to_template


BigBrainSym versus ICBM2009b Asym
Here we evaluated the spatial correspondence between BigBrainSym and MNI2009bAsym (asymmetric) knowing that BigBrainSym was registered to MNI2009bSym rather than MNI2009bAsym. AFRE should be higher than for
MNI2009bSym.
NIL2009bSym
'Total: $2.30+1-1.83 \mathrm{~mm}$

## ICBM2009b: Sym versus Asym

samated the distance between AFIDs for ICBM2009b sym and asym templates. Note that calling the difference AFRE is not technically correct as the two templates are not aligned to one another. However, the syntax was
kept the same for simplicity.
'Total: $0.88+/-0.68 \mathrm{~mm}{ }^{\prime}$

Is there any correlation of the errors reported with FLE?
Here we take our computed AFLE values for ICBM2009b-Asym and ICBM2009b-Sym and find that there is no correlation with the AFRE found.

```
Warning message in cor.test.default(sumnary_asym_vs_symSAFRE, summary_sym_df&mean,
    kendal1's rank correlation tau
    data: summary_asym_vs_symSAPRE and summary_sym_df$mear
    alternative hypothesis: true tau is not equal to 0
    sample estimates
    H: tau
```

2/24/2019
PHASE4_template_to_template


|  | AFID | Description | AFRE for BigBrainSym vs MNI2009bSym | Star: BigBrain and Sym | AFRE for BigBrainSym vs MNI2009bAsym | Star: BigBrain and Asym | AFRE for MNI2009b: Asym vs Sym | Star: Asym vs Sym |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 3 | 03 | infracollicular sulcus | 6.36 | * | 5.48 | * | 0.98 |  |
| 9 | 09 | L inferior LMS | 2.78 | * | 2.48 | . | 0.68 |  |
| 10 | 10 | cuimen | 9.27 | . | 9.39 |  | 0.21 |  |
| 14 | 14 | pineal gland | 4.42 | * | 4.16 |  | 0.41 |  |
| 16 | 16 | Llvat AC | 2.05 | . | 1.22 |  | 0.86 |  |
| 20 | 20 | splenium | 2.23 | . | 2.20 | . | 0.10 |  |
| 22 | 22 | L AL temporal horn | 4.69 | * | 3.44 |  | 2.45 | - |
| 6 | 26 | L inferior AM temporal horn | 1.88 |  | 2.58 | - | 0.98 |  |
| 27 | 27 | R indusium griseum origin | 1.21 |  | 3.60 | * | 2.81 | - |
| 28 | 28 | L indusium griseum origin | 0.74 |  | 2.88 | - | 2.29 | - |
| 29 | 29 | R ventral occipital horn | 2.54 | * | 3.99 | * | 1.63 |  |
| 30 | 30 | L ventral occipital horn | 5.88 | * | 4.22 | . | 2.00 | - |
| 31 | 31 | R olfactory sulcal fundus | 2.62 | * | 1.84 |  | 1.10 |  |
| 32 | 32 | L olfactory sulcal fundus | 3.06 | * | 4.21 | - | 1.24 |  |

R version 3.5.1 (2018-07-02)
Platform: x8666-apple-darwin14.5.0 (64-bit)
Running under: macos High Sierra 10.13.2
Matrix products: default
BLAS: /System/Library/Frameworks/Accelerate.framework/Versions/A/Frameworks/vecLib.framework/versions/A/1ibBLAS.dylib
LAPACK: /System/Library/Frameworks/Accelerate.framework/Versions/A/Frameworks/vecLLb.framework/versions/A/libLAPACK.dylib
locale:
_len_CA.UTF-8/en_CA.UTF-8/C/en_CA.UTF-8/en_CA.UTF-
attached base packages:
[1] stats
graphics grDevices utils datasets methods base
other attached packages:
$[1]$ ggplot2_3.0.0 reshape2_1.4.3 digest_0.6.16 plyr_1.8.4

https://mc.manuscriptcentral.com/LongRequest/hbm?DOWNLOAD=TRUE\&PARAMS=xik_3LHrwSCjMZ5brtcpoyhcwx.Jnh574PQKDjHos9smgMeCJF32WKfhM. .

## Appendix B

# Chapter 4 Supplementary Material 

B. 1 Supplementary Tables

| region | side | atlas | mean | stdev | v | $p$-value | $q$-value | max | max stdev |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Left Red Nucleus | left | ATAG | 1.02 | 0.53 | 111 | 0.69492 | 1.00000 | 1.69 | 0.98 |
| Left Substantia Nigra | left | ATAG | 0.86 | 0.39 | 76 | 0.95079 | 1.00000 | 1.52 | 0.67 |
| Left Subthalamic Nucleus | left | ATAG | 1.05 | 0.59 | 108 | 0.72770 | 1.00000 | 1.48 | 0.79 |
| Left Striatum | left | ATAG | 0.59 | 0.14 | 0 | 1.00000 | 1.00000 | 2.41 | 0.99 |
| Left Globus Pallidus Externus | left | ATAG | 0.48 | 0.15 | 0 | 1.00000 | 1.00000 | 1.33 | 0.66 |
| Left Globus Pallidus Internus | left | ATAG | 0.55 | 0.21 | 2 | 1.00000 | 1.00000 | 1.19 | 0.55 |
| Right Red Nucleus | right | ATAG | 1.03 | 0.40 | 131 | 0.44937 | 1.00000 | 1.71 | 0.72 |
| Right Substantia Nigra | right | ATAG | 0.78 | 0.30 | 31 | 0.99953 | 1.00000 | 1.48 | 0.80 |
| Right Subthalamic Nucleus | right | ATAG | 0.98 | 0.53 | 99 | 0.81473 | 1.00000 | 1.44 | 0.70 |
| Right Striatum | right | ATAG | 0.60 | 0.19 | 3 | 1.00000 | 1.00000 | 2.38 | 0.91 |
| Right Globus Pallidus Externus | right | ATAG | 0.50 | 0.22 | 2 | 1.00000 | 1.00000 | 1.31 | 0.81 |
| Right Globus Pallidus Internus | right | ATAG | 0.50 | 0.24 | 6 | 1.00000 | 1.00000 | 1.10 | 0.72 |
| Left Frontal | left | lobar | 0.91 | 0.12 | 30 | 0.99960 | 1.00000 | 7.69 | 1.19 |
| Left Parietal | left | lobar | 0.81 | 0.11 | 4 | 1.00000 | 1.00000 | 3.46 | 0.73 |
| Left Occipital | left | lobar | 1.15 | 0.26 | 197 | 0.01043 | 0.07938 | 4.62 | 1.83 |
| Left Temporal | left | lobar | 1.11 | 0.21 | 195 | 0.01257 | 0.08610 | 5.99 | 1.53 |
| Left Insular | left | lobar | 0.61 | 0.10 | 0 | 1.00000 | 1.00000 | 1.80 | 0.30 |
| Left Subcortical | left | lobar | 0.70 | 0.22 | 11 | 0.99999 | 1.00000 | 2.61 | 1.10 |
| Right Frontal | right | lobar | 0.89 | 0.11 | 24 | 0.99985 | 1.00000 | 8.07 | 1.71 |
| Right Parietal | right | lobar | 0.76 | 0.11 | 1 | 1.00000 | 1.00000 | 3.56 | 0.79 |
| Right Occipital | right | lobar | 1.07 | 0.17 | 166 | 0.10496 | 0.49586 | 4.24 | 0.94 |
| Right Temporal | right | lobar | 1.04 | 0.12 | 169 | 0.08810 | 0.43104 | 5.75 | 0.96 |
| Right Insular | right | lobar | 0.66 | 0.13 | 1 | 1.00000 | 1.00000 | 2.07 | 0.62 |
| Right Subcortical | right | lobar | 0.75 | 0.24 | 25 | 0.99982 | 1.00000 | 2.94 | 0.98 |
| Left Lateral Ventricle | left | HarvardOxford | 0.37 | 0.05 | 0 | 1.00000 | 1.00000 | 1.33 | 0.42 |
| Left Thalamus | left | HarvardOxford | 0.86 | 0.41 | 63 | 0.98206 | 1.00000 | 2.45 | 1.11 |
| Left Caudate | left | HarvardOxford | 0.64 | 0.25 | 8 | 1.00000 | 1.00000 | 1.81 | 0.58 |
| Left Putamen | left | HarvardOxford | 0.57 | 0.12 | 0 | 1.00000 | 1.00000 | 1.95 | 0.72 |
| Left Pallidum | left | HarvardOxford | 0.51 | 0.18 | 1 | 1.00000 | 1.00000 | 1.44 | 0.71 |
| Brainstem | NA | HarvardOxford | 1.31 | 0.64 | 179 | 0.04587 | 0.25135 | 7.50 | 1.93 |
| Left Hippocampus | left | HarvardOxford | 0.72 | 0.20 | 11 | 0.99999 | 1.00000 | 3.09 | 1.78 |
| Left Amygdala | left | HarvardOxford | 1.14 | 0.49 | 131 | 0.44937 | 1.00000 | 3.23 | 1.86 |
| Left Accumbens | left | HarvardOxford | 0.93 | 0.40 | 79 | 0.93966 | 1.00000 | 1.95 | 1.04 |
| Right Lateral Ventricle | right | HarvardOxford | 0.39 | 0.08 | 0 | 1.00000 | 1.00000 | 1.33 | 0.46 |
| Right Thalamus | right | HarvardOxford | 0.99 | 0.41 | 113 | 0.67218 | 1.00000 | 2.74 | 1.10 |
| Right Caudate | right | HarvardOxford | 0.60 | 0.18 | 1 | 1.00000 | 1.00000 | 1.63 | 0.53 |
| Right Putamen | right | HarvardOxford | 0.59 | 0.26 | 13 | 0.99998 | 1.00000 | 1.96 | 0.95 |


| Right Pallidum | right | HarvardOxford | 0.51 | 0.25 | 8 | 1.00000 | 1.00000 | 1.35 | 0.82 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Right Hippocampus | right | HarvardOxford | 0.67 | 0.16 | 1 | 1.00000 | 1.00000 | 2.50 | 0.89 |
| Right Amygdala | right | HarvardOxford | 1.07 | 0.54 | 107 | 0.73824 | 1.00000 | 2.72 | 1.23 |
| Right Accumbens | right | HarvardOxford | 0.82 | 0.39 | 51 | 0.99426 | 1.00000 | 1.79 | 0.94 |
| Left Frontal Pole | left | HarvardOxford | 0.91 | 0.24 | 43 | 0.99766 | 1.00000 | 6.92 | 1.62 |
| Right Frontal Pole | right | HarvardOxford | 0.91 | 0.12 | 41 | 0.99816 | 1.00000 | 7.49 | 2.02 |
| Left Insular Cortex | left | HarvardOxford | 0.61 | 0.10 | 0 | 1.00000 | 1.00000 | 1.80 | 0.30 |
| Right Insular Cortex | right | HarvardOxford | 0.66 | 0.13 | 1 | 1.00000 | 1.00000 | 2.07 | 0.62 |
| Left Superior Frontal Gyrus | left | HarvardOxford | 1.06 | 0.32 | 143 | 0.30508 | 1.00000 | 4.38 | 1.25 |
| Right Superior Frontal Gyrus | right | HarvardOxford | 1.01 | 0.36 | 114 | 0.66057 | 1.00000 | 4.36 | 1.29 |
| Left Middle Frontal Gyrus | left | HarvardOxford | 0.73 | 0.12 | 0 | 1.00000 | 1.00000 | 2.59 | 0.52 |
| Right Middle Frontal Gyrus | right | HarvardOxford | 0.67 | 0.13 | 0 | 1.00000 | 1.00000 | 2.31 | 0.49 |
| Left Inferior Frontal Gyrus Pars Triangularis | left | HarvardOxford | 0.85 | 0.25 | 51 | 0.99426 | 1.00000 | 2.30 | 0.65 |
| Right Inferior Frontal Gyrus Pars Triangularis | right | HarvardOxford | 0.81 | 0.24 | 34 | 0.99927 | 1.00000 | 2.29 | 0.75 |
| Left Inferior Frontal Gyrus Pars Opercularis | left | HarvardOxford | 0.81 | 0.28 | 43 | 0.99766 | 1.00000 | 2.32 | 0.69 |
| Right Inferior Frontal Gyrus Pars Opercularis | right | HarvardOxford | 0.71 | 0.17 | 9 | 0.99999 | 1.00000 | 1.95 | 0.48 |
| Left Precentral Gyrus | left | HarvardOxford | 0.87 | 0.12 | 15 | 0.99997 | 1.00000 | 3.95 | 1.04 |
| Right Precentral Gyrus | right | HarvardOxford | 0.83 | 0.12 | 8 | 1.00000 | 1.00000 | 3.85 | 0.87 |
| Left Temporal Pole | left | HarvardOxford | 1.18 | 0.28 | 217 | 0.00110 | 0.00940 | 5.16 | 1.57 |
| Right Temporal Pole | right | HarvardOxford | 1.17 | 0.24 | 213 | 0.00184 | 0.01479 | 4.87 | 1.50 |
| Left Superior Temporal Gyrus Anterior | left | HarvardOxford | 0.89 | 0.42 | 57 | 0.98957 | 1.00000 | 2.41 | 0.97 |
| Right Superior Temporal Gyrus Anterior | right | HarvardOxford | 0.87 | 0.22 | 56 | 0.99052 | 1.00000 | 2.26 | 0.55 |
| Left Superior Temporal Gyrus Posterior | left | HarvardOxford | 0.88 | 0.23 | 66 | 0.97692 | 1.00000 | 2.13 | 0.60 |
| Right Superior Temporal Gyrus Posterior | right | HarvardOxford | 0.87 | 0.27 | 57 | 0.98957 | 1.00000 | 2.22 | 0.78 |
| Left Middle Temporal Gyrus Anterior | left | HarvardOxford | 1.54 | 0.67 | 228 | 0.00021 | 0.00196 | 3.58 | 1.34 |
| Right Middle Temporal Gyrus Anterior | right | HarvardOxford | 1.13 | 0.42 | 155 | 0.18527 | 0.79317 | 2.83 | 0.84 |
| Left Middle Temporal Gyrus Posterior | left | HarvardOxford | 1.16 | 0.29 | 196 | 0.01146 | 0.08262 | 3.73 | 0.94 |
| Right Middle Temporal Gyrus Posterior | right | HarvardOxford | 1.02 | 0.28 | 138 | 0.36309 | 1.00000 | 3.17 | 0.84 |
| Left Middle Temporal Gyrus TemporoOccipital | left | HarvardOxford | 1.11 | 0.27 | 174 | 0.06444 | 0.32697 | 2.99 | 0.53 |
| Right Middle Temporal Gyrus TemporoOccipital | right | HarvardOxford | 1.05 | 0.31 | 140 | 0.33943 | 1.00000 | 3.02 | 0.83 |
| Left Inferior Temporal Gyrus Anterior | left | HarvardOxford | 1.30 | 0.53 | 190 | 0.01954 | 0.12166 | 3.29 | 1.35 |
| Right Inferior Temporal Gyrus Anterior | right | HarvardOxford | 1.08 | 0.29 | 161 | 0.13781 | 0.60905 | 2.79 | 0.62 |
| Left Inferior Temporal Gyrus Posterior | left | HarvardOxford | 1.73 | 0.36 | 252 | 0.00000 | 0.00001 | 5.34 | 0.84 |
| Right Inferior Temporal Gyrus Posterior | right | HarvardOxford | 1.56 | 0.34 | 253 | 0.00000 | 0.00001 | 5.05 | 0.82 |
| Left Inferior Temporal Gyrus TemporoOccipital | left | HarvardOxford | 1.09 | 0.32 | 136 | 0.38726 | 1.00000 | 3.50 | 1.15 |
| Right Inferior Temporal Gyrus TemporoOccipital | right | HarvardOxford | 1.08 | 0.17 | 186 | 0.02712 | 0.15483 | 3.82 | 0.85 |
| Left Postcentral Gyrus | left | HarvardOxford | 0.90 | 0.15 | 44 | 0.99736 | 1.00000 | 3.03 | 0.67 |
| Right Postcentral Gyrus | right | HarvardOxford | 0.87 | 0.18 | 41 | 0.99816 | 1.00000 | 3.30 | 0.88 |


| Left Superior Parietal Lobule | left | HarvardOxford | 0.88 | 0.26 | 72 | 0.96311 | 1.00000 | 2.32 | 0.54 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Right Superior Parietal Lobule | right | HarvardOxford | 0.91 | 0.32 | 86 | 0.90651 | 1.00000 | 2.55 | 0.67 |
| Left Supramarginal Gyrus Anterior | left | HarvardOxford | 0.89 | 0.18 | 51 | 0.99426 | 1.00000 | 2.41 | 0.54 |
| Right Supramarginal Gyrus Anterior | right | HarvardOxford | 0.82 | 0.17 | 19 | 0.99994 | 1.00000 | 2.31 | 0.56 |
| Left Supramarginal Gyrus Posterior | left | HarvardOxford | 1.00 | 0.21 | 122 | 0.56320 | 1.00000 | 2.70 | 0.54 |
| Right Supramarginal Gyrus Posterior | right | HarvardOxford | 0.87 | 0.14 | 20 | 0.99993 | 1.00000 | 2.55 | 0.48 |
| Left Angular Gyrus | left | HarvardOxford | 1.01 | 0.23 | 136 | 0.38726 | 1.00000 | 2.75 | 0.62 |
| Right Angular Gyrus | right | HarvardOxford | 0.96 | 0.23 | 96 | 0.83956 | 1.00000 | 2.84 | 0.51 |
| Left Lateral Occipital Cortex Superior | left | HarvardOxford | 1.10 | 0.23 | 186 | 0.02712 | 0.15483 | 3.37 | 0.57 |
| Right Lateral Occipital Cortex Superior | right | HarvardOxford | 1.07 | 0.23 | 174 | 0.06444 | 0.32697 | 3.28 | 0.61 |
| Left Lateral Occipital Cortex Inferior | left | HarvardOxford | 1.31 | 0.26 | 248 | 0.00000 | 0.00003 | 3.62 | 1.32 |
| Right Lateral Occipital Cortex Inferior | right | HarvardOxford | 1.28 | 0.18 | 253 | 0.00000 | 0.00001 | 3.45 | 0.57 |
| Left Intracalcarine Cortex | left | HarvardOxford | 0.65 | 0.27 | 13 | 0.99998 | 1.00000 | 1.77 | 0.61 |
| Right Intracalcarine Cortex | right | HarvardOxford | 0.53 | 0.23 | 3 | 1.00000 | 1.00000 | 1.51 | 0.59 |
| Left Frontal Medial Cortex | left | HarvardOxford | 1.75 | 0.59 | 251 | 0.00000 | 0.00001 | 5.68 | 1.12 |
| Right Frontal Medial Cortex | right | HarvardOxford | 1.84 | 0.56 | 253 | 0.00000 | 0.00001 | 5.77 | 0.94 |
| Left Juxtapositional Lobule Cortex (formerly SMA) | left | HarvardOxford | 1.05 | 0.43 | 128 | 0.48731 | 1.00000 | 3.97 | 1.40 |
| Right Juxtapositional Lobule Cortex (formerly SMA) | right | HarvardOxford | 1.05 | 0.53 | 116 | 0.63691 | 1.00000 | 3.82 | 1.49 |
| Left Subcallosal Cortex | left | HarvardOxford | 1.72 | 0.33 | 252 | 0.00000 | 0.00001 | 6.87 | 0.67 |
| Right Subcallosal Cortex | right | HarvardOxford | 1.80 | 0.38 | 253 | 0.00000 | 0.00001 | 6.87 | 0.62 |
| Left Paracingulate Gyrus | left | HarvardOxford | 0.73 | 0.27 | 27 | 0.99975 | 1.00000 | 2.63 | 1.09 |
| Right Paracingulate Gyrus | right | HarvardOxford | 0.73 | 0.22 | 17 | 0.99996 | 1.00000 | 2.54 | 0.95 |
| Left Cingulate Gyrus Anterior | left | HarvardOxford | 0.65 | 0.15 | 1 | 1.00000 | 1.00000 | 2.12 | 0.79 |
| Right Cingulate Gyrus Anterior | right | HarvardOxford | 0.70 | 0.20 | 11 | 0.99999 | 1.00000 | 2.41 | 0.93 |
| Left Cingulate Gyrus Posterior | left | HarvardOxford | 0.52 | 0.11 | 0 | 1.00000 | 1.00000 | 1.75 | 0.32 |
| Right Cingulate Gyrus Posterior | right | HarvardOxford | 0.54 | 0.19 | 1 | 1.00000 | 1.00000 | 1.83 | 0.51 |
| Left Precuneus Cortex | left | HarvardOxford | 0.63 | 0.12 | 0 | 1.00000 | 1.00000 | 2.80 | 0.92 |
| Right Precuneus Cortex | right | HarvardOxford | 0.54 | 0.09 | 0 | 1.00000 | 1.00000 | 2.53 | 0.55 |
| Left Cuneal Cortex | left | HarvardOxford | 0.84 | 0.23 | 38 | 0.99875 | 1.00000 | 2.51 | 0.82 |
| Right Cuneal Cortex | right | HarvardOxford | 0.71 | 0.18 | 2 | 1.00000 | 1.00000 | 2.29 | 0.80 |
| Left Frontal Orbital Cortex | left | HarvardOxford | 1.10 | 0.22 | 191 | 0.01794 | 0.11702 | 6.20 | 1.50 |
| Right Frontal Orbital Cortex | right | HarvardOxford | 1.08 | 0.27 | 150 | 0.23139 | 0.96061 | 6.74 | 2.16 |
| Left Parahippocampal Gyrus Anterior | left | HarvardOxford | 1.51 | 0.88 | 244 | 0.00001 | 0.00008 | 3.84 | 1.89 |
| Right Parahippocampal Gyrus Anterior | right | HarvardOxford | 1.40 | 0.35 | 249 | 0.00000 | 0.00003 | 4.00 | 1.27 |
| Left Parahippocampal Gyrus Posterior | left | HarvardOxford | 0.80 | 0.23 | 33 | 0.99937 | 1.00000 | 2.24 | 0.52 |
| Right Parahippocampal Gyrus Posterior | right | HarvardOxford | 0.76 | 0.13 | 4 | 1.00000 | 1.00000 | 2.10 | 0.31 |
| Left Lingual Gyrus | left | HarvardOxford | 0.70 | 0.26 | 17 | 0.99996 | 1.00000 | 2.63 | 1.38 |
| Right Lingual Gyrus | right | HarvardOxford | 0.63 | 0.20 | 4 | 1.00000 | 1.00000 | 2.28 | 0.90 |


| Left Temporal Fusiform Cortex Anterior | left | HarvardOxford | 1.62 | 0.85 | 247 | 0.00000 | 0.00004 | 3.77 | 1.85 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Right Temporal Fusiform Cortex Anterior | right | HarvardOxford | 1.59 | 0.57 | 247 | 0.00000 | 0.00004 | 3.52 | 1.13 |
| Left Temporal Fusiform Cortex Posterior | left | HarvardOxford | 1.04 | 0.31 | 125 | 0.52537 | 1.00000 | 3.58 | 1.15 |
| Right Temporal Fusiform Cortex Posterior | right | HarvardOxford | 1.08 | 0.25 | 164 | 0.11738 | 0.53604 | 3.49 | 0.60 |
| Left Temporal Fusiform Cortex TemporoOccipital | left | HarvardOxford | 0.64 | 0.23 | 8 | 1.00000 | 1.00000 | 2.04 | 0.89 |
| Right Temporal Fusiform Cortex TemporoOccipital | right | HarvardOxford | 0.65 | 0.21 | 6 | 1.00000 | 1.00000 | 2.53 | 0.77 |
| Left Occipital Fusiform Gyrus | left | HarvardOxford | 1.03 | 0.57 | 104 | 0.76861 | 1.00000 | 2.92 | 1.71 |
| Right Occipital Fusiform Gyrus | right | HarvardOxford | 0.83 | 0.42 | 68 | 0.97288 | 1.00000 | 2.56 | 1.07 |
| Left Frontal Operculum Cortex | left | HarvardOxford | 0.61 | 0.14 | 0 | 1.00000 | 1.00000 | 1.71 | 0.54 |
| Right Frontal Operculum Cortex | right | HarvardOxford | 0.55 | 0.15 | 0 | 1.00000 | 1.00000 | 1.46 | 0.41 |
| Left Central Opercular Cortex | left | HarvardOxford | 0.61 | 0.15 | 0 | 1.00000 | 1.00000 | 1.91 | 0.48 |
| Right Central Opercular Cortex | right | HarvardOxford | 0.57 | 0.13 | 0 | 1.00000 | 1.00000 | 1.71 | 0.45 |
| Left Parietal Operculum Cortex | left | HarvardOxford | 0.55 | 0.12 | 0 | 1.00000 | 1.00000 | 1.64 | 0.37 |
| Right Parietal Operculum Cortex | right | HarvardOxford | 0.53 | 0.11 | 0 | 1.00000 | 1.00000 | 1.70 | 0.48 |
| Left Planum Polare | left | HarvardOxford | 0.65 | 0.14 | 0 | 1.00000 | 1.00000 | 1.67 | 0.49 |
| Right Planum Polare | right | HarvardOxford | 0.79 | 0.25 | 31 | 0.99953 | 1.00000 | 1.78 | 0.44 |
| Left Heschl Gyrus | left | HarvardOxford | 0.62 | 0.26 | 21 | 0.99991 | 1.00000 | 1.53 | 0.42 |
| Right Heschl Gyrus | right | HarvardOxford | 0.68 | 0.21 | 6 | 1.00000 | 1.00000 | 1.64 | 0.49 |
| Left Planum Temporale | left | HarvardOxford | 0.65 | 0.16 | 1 | 1.00000 | 1.00000 | 1.87 | 0.43 |
| Right Planum Temporale | right | HarvardOxford | 0.63 | 0.13 | 0 | 1.00000 | 1.00000 | 1.72 | 0.46 |
| Left Supracalcarine Cortex | left | HarvardOxford | 0.59 | 0.21 | 3 | 1.00000 | 1.00000 | 1.53 | 0.52 |
| Right Supracalcarine Cortex | right | HarvardOxford | 0.49 | 0.17 | 1 | 1.00000 | 1.00000 | 1.15 | 0.36 |
| Left Occipital Pole | left | HarvardOxford | 1.64 | 0.65 | 248 | 0.00000 | 0.00003 | 4.33 | 1.95 |
| Right Occipital Pole | right | HarvardOxford | 1.55 | 0.46 | 252 | 0.00000 | 0.00001 | 4.01 | 1.08 |

V represents the Wilcoxon rank sum statistic, $p$-value is unadjusted, and $q$-value is FDR adjusted. * 13 regions met thresholds for statistical significance after FDR correction at a rate of $q<0.025$.

## Appendix C

## Ethics Approvals

## C. 1 Retrospective images for deep brain stimulation surgery.

## Western <br> Resear@lastern University Health Science Research Ethics Board HSREB Delegated Initial Approval Notice

Principal Investigator: Dr. Mandar Jog
Department \& Institution: Schulich School of Medicine and DentistrylClinical Neurological Sciences,London Health Sciences Centre

Review Type: Delegated
HSREB File Number: 109045
Study Title: Retrospective medical image review of patients who have undergone implantation of deep brain stimulation electrodes for the purpose of electrode localization using advanced medical image reconstruction techniques.
HSREB Initial Approval Date: March 07, 2017
HSREB Expiry Date: March 07,2018

|  |  |  |
| :--- | :--- | :--- |
| Documents Approved and/or Received for Information: |  |  |
| Document Name | Comments | Version Date |
| Western University Protocol | Received February 16, 2017 |  |
| Data Collection Form/Case Report Form | Version 1.0 | $2017 / 02 / 15$ |


#### Abstract

The Western University Health Science Research Ethics Board (HSREB) has reviewed and approved the above named study, as of the HSREB Initial Approval Date noted above.

HSREB approval for this study remains valid until the HSREB Expiry Date noted above, conditional to timely submission and acceptance of HSREB Continuing Ethics Review.

The Western University HSREB operates in compliance with the Tri-Council Policy Statement Ethical Conduct for Research Involving Humans (TCPS2), the International Conference on Harmonization of Technical Requirements for Registration of Pharmaceuticals for Human Use Guideline for Good Clinical Practice Practices (ICH E6 R1), the Ontario Personal Health Information Protection Act (PHIPA, 2004), Part 4 of the Natural Health Product Regulations, Health Canada Medical Device Regulations and Part C, Division 5, of the Food and Drug Regulations of Health Canada.


Members of the HSREB who are named as Investigators in research studies do not participate in discussions related to, nor vote on such studies when they are presented to the REB.

The HSREB is registered with the U.S. Department of Health \& Human Services under the IRB registration number IRB 00000940 .

## C. 2 7T images for geometric distortion study.



## Office of Research Ethics

The University of Western Ontario
Room 4180 Support Services Building, London, ON, Canada N6A 5C1
Telephone: (519) 661-3036 Fax: (519) 850-2466 Email: ethics@uwo.ca Website: www.uwo.ca/research/ethics
Use of Human Subjects - Ethics Approval Notice
Principal Investigator: Dr. T.M. Peters
Review Number: 16189
Review Level: Full Board
Review Date: May 19, 2009
Protocol Title: Structural and Functional MR imaging in Frontal and Temporal Lobe Epilepsy at 1.5T, 3T, and 7T
Department and Institution: Imaging, Robarts Research Institute Sponsor: CIHR-CANADIAN INSTITUTE OF HEALTH RESEARCH

$$
\text { Ethics Approval Date: October 7, } 2009 \quad \text { Expiry Date: July 31, } 2015
$$

Documents Reviewed and Approved: UWO Protocol, Letter of information \& consent form dated Aug. 31/09 \& Advertisement dated Aug. 31/09
Documents Received for Information:

This is to notify you that The University of Western Ontario Research Ethics Board for Health Sciences Research Involving Human Subjects (HSREB) which is organized and operates according to the Tri-Council Policy Statement: Ethical Conduct of Research Involving Humans and the Health Canada/ICH Good Clinical Practice Practices: Consolidated Guidelines; and the applicable laws and regulations of Ontario has reviewed and granted approval to the above referenced study on the approval date noted above. The membership of this REB also complies with the membership requirements for REB's as defined in Division 5 of the Food and Drugg Regulations.

The ethics approval for this study shall remain valid until the expiry date noted above assuming timely and acceptable responses to the HSREB's periodic requests for surveillance and monitoring information. If you require an updated approval notice prior to that time you must request it using the UWO Updated Approval Request Form.

During the course of the research, no deviations from, or changes to, the protocol or consent form may be initiated without prior written approval from the HSREB except when necessary to eliminate immediate hazards to the subject or when the change(s) involve only logistical or administrative aspects of the study (e.g. change of monitor, telephone number). Expedited review of minor change(s) in ongoing studies will be considered. Subjects must receive a copy of the signed information/consent documentation.
Investigators must promptly also report to the HSREB:
a) changes increasing the risk to the participant(s) and/or affecting significantly the conduct of the study;
b) all adverse and unexpected experiences or events that are both serious and unexpected;
c) new information that may adversely affect the safety of the subjects or the conduct of the study.

If these changes/adverse events require a change to the information/consent documentation, and/or recruitment advertisement, the newly revised information/consent documentation, and/or advertisement, must be submitted to this office for approval.

Members of the HSREB who are named as investigators in research studies, or declare a conflict of interest, do not participate in discussion related to, nor vote on, such studies when they are presented to the HSREB.

## C. 3 Prospective 7T stereotaxy study.

Research Ethics

Western University Health Science Research Ethics Board HSREB Full Board Initial Approval Notice

Department \& Institution: Schulich School of Medicine and Dentistry\Medical Biophysics, Western University
Review Type: Full Board
HSREB File Number: 108952
Study Title: Ultra-high field magnetic resonance imaging for stereotactic neurosurgery
Sponsor:
HSREB Initial Approval Date: March 19, 2017
HSREB Expiry Date: March 19, 2018
Documents Approved and/or Received for Information:

| Document Name | Comments | Version Date |
| :--- | :--- | :--- |
| Letter of Information \& Consent | LOI patients clean | $2017 / 03 / 08$ |
| Data Collection Form/Case Report Form | Data collection form | $2017 / 01 / 22$ |
| Advertisement | poster clean | $2017 / 02 / 17$ |
| Instruments | Surgeon Questionnaire | $2017 / 02 / 17$ |
| Western University Protocol | Protocol 108952 Clean | $2017 / 02 / 28$ |
| Letter of Information \& Consent | LOI clinician clean | $2017 / 02 / 28$ |
| Letter of Information \& Consent | LOI controls clean | $2017 / 02 / 28$ |
| Letter of Information \& Consent | LOI patients clean | $2017 / 02 / 28$ |

The Western University Health Science Research Ethics Board (HSREB) has reviewed and approved the above named study, as of the HSREB Initial Approval Date noted above.

HSREB approval for this study remains valid until the HSREB Expiry Date noted above, conditional to timely submission and acceptance of HSREB Continuing Ethics Review.

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- Chapter 4: Lau, J. C., Khan, A. R., Zeng, T. Y., MacDougall, K. W., Parrent, A. G., \& Peters, T. M. (2018). Quantification of local geometric distortion in structural magnetic resonance images: Application to ultra-high fields. NeuroImage, 168, 141-151.

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# Curriculum Vitae 

Name

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## Education and Degrees

Western University<br>London, Ontario, Canada<br>2017 - Present, PhD Candidate in Biomedical Engineering<br>2011 - Present, Resident in Neurosurgery<br>Queen's University<br>Kingston, Ontario, Canada<br>2007-2011, MD

## Awards

Canadian Institutes of Health Research (CIHR) Frederick Banting and Charles Best Canada Graduate Scholarships Doctoral Award (2017-Present)

Canadian League Against Epilepsy, Mary Ann Lee Resident Research Prize (2018)
Department of Clinical Neurological Sciences, Resident Research Prize (2018)
Canadian League Against Epilepsy, Top Trainee Abstract Award (2017)
Department of Clinical Neurological Sciences, Resident Research Prize (2017)
Canadian Neurological Sciences Federation Congress, Best Resident Poster (2016)

## Recent Manuscripts

(* - co-directed with Dr. Joseph Megyesi)

1. Lau JC, Parrent AG, Xiao Y, Demarco J, MacDougall KW, Cruckley C, Peters TM, Khan AR. Direct visualization and characterization of the human zona incerta and surrounding fiber tracts. In preparation.
2. Lau JC, Parrent AG, Demarco J, Ferko K, Gupta G, Khan AR, Peters TM. A framework for evaluating correspondence between brain images using anatomical fiducials. Human Brain Mapping. Under review.
3. DeKraker J, Ferko K, Lau JC, Kohler S, Khan AR. Hippocampal morphology and cytoarchitecture in 3D BigBrain histology. Cerebral Cortex. Submitted.
4. Gui C, Lau JC*, Kosteniuk SE, Lee DH, Megyesi JF. Radiology reporting of low-grade glioma growth underestimates tumor expansion. Acta Neurochirurgica. Accepted.
5. Lau JC, Kosteniuk SE, Walker T, Iansavitchene A, Macdonald DR, Megyesi JF. Operative complications with and without image-guidance: systematic review and metaanalysis of the Ommaya reservoir literature. World Neurosurgery. 2019. 122:404-414.
6. Santyr BG, Lau JC, Mirsattari SM, Burneo JG, de Ribaupierre S, Steven DA, Parrent AG, MacDougall KW, Khan AR. Novel connectivity map normalization procedure for improved quantitative investigation of structural thalamic connectivity in temporal lobe epilepsy patients. Journal of Magnetic Resonance Imaging. 2018. 48:1529-1539.
7. Kosteniuk SE, Lau JC*, Megyesi JF. The Impact of Functional Magnetic Resonance Imaging on Clinical Outcomes in a Propensity-Matched Low-Grade Glioma Cohort. World Neurosurgery. 2018. 120:1143-1148.
8. Lau JC, Khan AR, MacDougall KW, Parrent AG, Peters TM. Quantification of local geometric distortion in structural magnetic resonance images: application to ultra-high fields. NeuroImage. Special Issue: Neuroimaging with Ultra-High Field MRI: Present and Future. 2018. 168:141-151.
9. Gui C, Kosteniuk SE, Lau JC*, Megyesi JF. Tumor growth dynamics in serially-imaged low-grade glioma patients. Journal of Neuro-Oncology. 2018. 139:167-175.
10. Lau JC, Kosteniuk SE, Macdonald DR, Megyesi JF. Image-guided Ommaya reservoir insertion for intraventricular chemotherapy. Acta Neurochirurgica. 2018. 160:539-544.
11. DeKraker J, Ferko K, Lau JC, Kohler S, Khan AR. Unfolding the hippocampus: applications in subfield segmentation and standardization. NeuroImage. 2018. 167:408-418.
12. Lau JC, MacDougall K, Peters TM, Parrent AG, Khan AR. Ultra-high field MRI templateassisted target selection for deep brain stimulation surgery. World Neurosurgery. 2017. 103:531-537.
13. Santyr BG, Goubran M, Lau JC, Kwan B, Salehi F, Lee DH, Mirsattari SM, Burneo JG, Steven DA, Parrent AG, de Ribaupierre S, Hammond RR, Peters TM, Khan AR. Investigation of hippocampal sub-structures in focal temporal lobe epilepsy with and without hippocampal sclerosis at 7T. Journal of Magnetic Resonance Imaging. 2017. 45(5):1359-1370.
14. Lau JC, Kosteniuk SE, Bihari F, Megyesi JF. Functional magnetic resonance imaging for preoperative planning in brain tumour surgery. Canadian Journal of Neurological Sciences. 2017. 44:59-68.

[^0]:    ${ }^{1}$ Contemporary use of the word "stereotactic" may actually have mixed origins, "tactus" being Latin for touch.

[^1]:    ${ }^{2}$ Not to be confused with the ventral posterolateral (VPL) or main somatosensory nucleus of the thalamus

[^2]:    ${ }^{1}$ https://sourceforge.net/projects/niftyreg/; git version hash key 83d8d1182ed4c227ce4764f1fdab3b1797eecd8d downloaded and compiled January 13, 2016

[^3]:    https：／／mc．manuscriptcentral．com／LongRequest／hbm？DOWNLOAD＝TRUE\＆PARAMS＝xik＿Mbi415k82xNsH2RCenqEHsLXvXH89mphe4STkVJRJzBEVCn4Q1．．．

