

# The Kapowski Chronicles: A Complete, Volumetric Cortical Thickness Open Source Pipeline with Evaluation on Public Data

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## Abstract

Numerous studies have explored the relationship between cortical structure and brain development, cognitive function, and functional connectivity. The highly convoluted cortical topography makes manual measurements arduous and often impractical given the population sizes necessary for sufficient statistical power. Computational techniques have permitted large-scale studies as they provide robust and reliable localized measurements characterizing the cortex with little or no human intervention. Particularly useful to the neuroscience community are publicly available tools, such as the popular surface-based Freesurfer, which facilitate the testing and refinement of hypotheses. In this paper, we introduce the volume-based Advanced Normalization Tools (ANTS) cortical thickness automated pipeline comprised entirely of well-vetted components developed as part of the ANTS toolkit such as multivariate template construction, SyN (image registration), N4 (bias correction), Atropos (*n*-tissue segmentation), and DiReCT (cortical thickness) all developed as part of the ANTS open science effort. Complementing the open source aspect of ANTS we confirm previous findings of gender differences, age correlation, and hemispherical asymmetry in cortical thickness using the publicly available IXI data consisting of multimodal structural MRI of approximately 600 subjects. To further promote open science and use of the proposed tools, all scripts and templates used to produce the results are hosted on the Neuroimaging Informatics Tools and Resources Clearinghouse (NITRC) website.

**Keywords:** advanced normalization tools, bias correction, cortical thickness, open source software, segmentation, skull stripping

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## 1. Introduction

Neuroscientific investigations into cortical morphological changes/differences have illuminated interesting correlations with normal and pathological neurodevelopment in addition to cognitive function. Historically rooted in the meticulous work of von Economo [22], computational methods for modeling the cortical gray matter have resulted in numerous studies exploring such relationships. These analysis techniques have been instrumental in demonstrating the presence of regional cortical abnormalities in such conditions as Huntington's disease [55, 54, 58], schizophrenia [48], bipolar disorder [43], Alzheimer's disease and frontotemporal dementia [21, 19], Parkinson's disease [35], multiple sclerosis [53], autism [12, 32], migraines [17], chronic smoking [39], alcoholism [28], cocaine addiction [46], Tourette syndrome in children [66], scoliosis in female adolescents [70], obsessive compulsive disorder [62], ADHD [1], obesity [52], and heritable [50] and elderly [8] depression. Further study topics exploring correlations with cortical thickness include normal aging [37], gender differences [42], anatomical asymmetries [41, 2], intelligence [61], athletic ability [72], musical abil-

ity [9, 29], white collar criminals [51], and Tetris-playing ability in female adolescents [31]. Additionally, recent studies have demonstrated possible functional connectivity relationships using cortical thickness measures [73, 40, 33]. Although these study findings are subject to debate and interpretation [30], the availability of quantitative computational methods for extracting such information has proven invaluable for developing and refining fundamental neuroscience hypotheses.

Broadly characterized as surface mesh-based or volumetric [56, 13], such techniques have been introduced and extensively developed over the last 10+ years. Representative of the former is the Freesurfer<sup>2</sup> cortical modeling software package [15, 26, 24, 25, 27] which owes its popularity to public availability, excellent documentation, good performance, and integration with other toolkits, such as the extensive FMRIB software library (FSL) [65]. Similar to other surface approaches (e.g. [18, 45, 44, 36]), the pial and white matter surfaces from individual subject MR data are modeled with polygonal meshes which are then used to determine local cortical thickness based on a specified correspondence between the surface models.

Image volumetric (or meshless) techniques are varied both in algorithmic terms as well as the underlying definition of cortical thickness. An early, foundational technique is the method

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<sup>2</sup><http://surfer.nmr.mgh.harvard.edu/>

of [34] in which the inner and outer surface geometry is used to determine the solution to Laplace’s equation in which thickness is measured by integrating along the tangents of the resulting field lines spanning the boundary surfaces. Subsequent contributions improved upon the original formulation. For example, in [74], an Eulerian PDE approach was proposed to facilitate the computation of correspondence paths. Extending the surface-based work of [44], the hybrid approach of [36] uses the discrete Laplacian field to deform the white matter surface mesh towards the pial surface. Although the Laplacian-based approach has several advantages including generally lower computational times and non-crossing correspondence paths, direct correlative assessments with histology are difficult as the quantified distances are not necessarily Euclidean. Other volumetric algorithms have employed coupled level sets [75], model-free intelligent search strategies either normal to the gray-white matter interface [56] or using a min-max rule [69]. Most relevant to this work is the DiReCT (Diffeomorphic Registration-based Cortical Thickness) algorithm proposed by [16] in which a registration-based approach proposed where the derived diffeomorphic mapping between the white and pial matter surfaces is used to propagate thickness values through the cortical gray matter.

Despite the numerous techniques that have been proposed in the literature (of which only a fraction has been cited), several requisite processing components are common to many of them which are prior to quantification of cortical thickness. These include inhomogeneity correction, skull stripping, and  $n$ -tissue segmentation for differentiating the gray and white matters. For post-processing statistical analysis across populations, construction of population-specific unbiased templates is also beneficial. In addition, some of these steps might include a crucial registration component, e.g. propagating template-based tissue priors for improved segmentation. The requisite additional components coupled with the general lack of availability of published algorithms [38] inhibits performing studies by external researchers and makes comparative evaluations difficult. For example, one recent evaluation study [13] compared Freesurfer (a surface-based method) with two volumetric methods [34, 16]. Whereas the entire Freesurfer processing pipeline has been made publicly available, tuned by the original authors in terms of implementation, and described in great detail (specifically in terms of suggested parameters); both volumetric methods were implemented solely by the authors of the evaluation (not the actual software developers) using unspecified parameters making the comparisons less than ideal.

In this work we describe our cortical thickness software pipeline which we have made open source as part of our Advanced Normalization Tools (ANTs) repository. This includes all the necessary preprocessing steps consisting of well-vetted previously published algorithms for bias correction [68], brain extraction [4],  $n$ -tissue segmentation [6], template construction [7], and image normalization [5]. We also describe improvements made to the original DiReCT algorithm [16]. Equally as important, we demonstrate how to coordinate these pipeline components and provide a set of useful parameters which are used to analyze the publicly available IXI data set. This is en-

capsulated in a well-documented shell script which is also available in ANTs. Furthermore, we provide all the derived image data and other processing scripts on a github repository specifically meant for this publication. This permits the set of results described in this work to be fully reproducible thus permitting other researchers to use a complete volumetric pipeline for measuring cortical thickness.

## 2. Methods and Materials

### 2.1. ANTs volumetric-based cortical thickness estimation pipeline

The ANTs-based cortical thickness estimation workflow is illustrated in Figure 1. The steps are as follows:

1. initial N4 bias correction on input anatomical MRI,
2. brain extraction using a hybrid segmentation/template-based strategy,
3. alternating between prior-based segmentation and white matter posterior probability weighted bias correction,
4. DiReCT-based cortical thickness estimation, and
5. optional normalization to specified template.

Each component, including both software and data, is briefly detailed below with the relevant references for additional information.

Additionally, the coordination of all the algorithmic components is encapsulated in the shell script `antsCorticalThickness.sh`. This includes optimal parameters for each of the algorithmic components which has worked well for our processing and which are used to acquire the results described in this work.

#### 2.1.1. Anatomical template construction

Normalizing images to a standard coordinate system reduces intersubject variability in population studies. Various approaches exist for determining the normalized space such as the selection of a pre-existing template based on a single subject, e.g. the Talairach atlas [67], or a publicly available averaged group of subjects, e.g. the MNI [14] or ICBM [47] templates. Additionally, mean templates constructed from labeled data can be used to construct spatial priors for improving segmentation algorithms. The work of [7] explicitly models the geometric component of the normalized space during optimization to produce such mean templates. Coupling the intrinsic symmetry of SyN pairwise registration [5] and an optimized sharpening/averaging of the template appearance, Symmetric Group Normalization (SyGN) is a powerful framework for producing optimal population-specific templates [7] with arbitrary similarity metric choice.

The ANTs implementation of this technique is currently available as a shell script, `buildtemplateparallel.sh`, and a multivariate version, `antsMultivariateTemplateConstruction.sh`, both of which are distributed as part of the ANTs repository. The latter script permits the construction of multimodal templates (e.g. T1-weighted, T2-weighted, and proton density MRI as

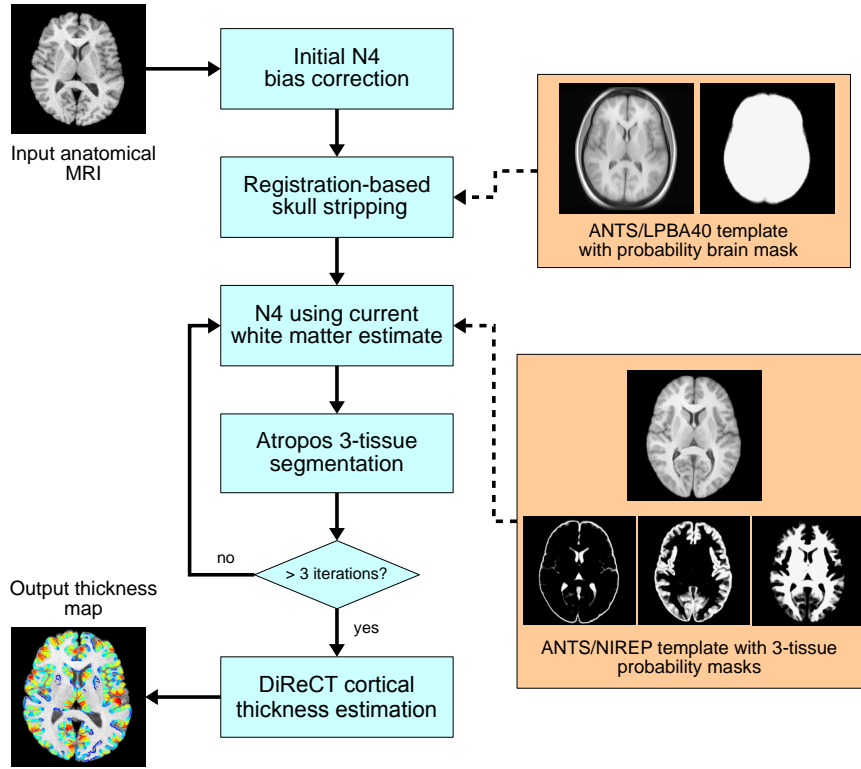


Figure 1: The ANTs T1 processing workflow containing all elements for determining cortical thickness. Not shown is the optional single subject to template registration.

described in the Evaluation section). Both scripts accommodate a variety of computational resources for facilitating template construction. These computational resource possibilities include:

- serial processing on a single workstation,
- parallelized processing on a single workstation with multiple cores using `pexec`<sup>3</sup>,
- parallelized processing using Apple’s XGrid technology<sup>4</sup>,
- parallelized processing using Sun Grid Engine for cluster-based systems<sup>5</sup>, and
- parallelized processing using the Portable Batch System for cluster-based systems<sup>6</sup>.

Within this work multiple templates were created for all stages of image processing and analysis. The creation of these templates are described in the corresponding data section.

### 2.1.2. N4 Bias field correction

Critical to quantitative processing of MRI is the minimization of field inhomogeneity effects which causes artificial low

frequency intensity variation across the image. Large-scale studies, such as the Alzheimer’s Disease Neuroimaging Initiative (ADNI), employ perhaps the most widely used bias correction algorithm, N3 [63], as part of their standard protocol [10].

In [68], we introduced an extension of N3, denoted as N4, which demonstrates improved performance and convergence behavior on a variety of data. This improvement is a result of an enhanced fitting routine (which includes multi-resolution capabilities) and a modified optimization formulation. For our workflow, the additional possibility of specifying a weighted mask in N4 permits the use of the current white matter probability map calculated during the segmentation pipeline for further improvement of bias field estimation. In addition to its public availability through ANTs and the Insight Toolkit, it has also been included in the popular open source Slicer software package for visualization and medical image computing [23].

N4 is used in two places during the individual subject processing (cf. Figure 1). Following conversion of the raw dicom T1-weighted image to Nifti format using our related Neuropipedream set of raw image conversion and organization tools<sup>7</sup>, N4 is used to generate an initial bias corrected image for use in brain extraction. The input mask is created by adaptively thresholding the background from the foreground using Otsu’s algorithm [49]. Following brain extraction, the three-tissue segmentation involves iterating between bias field cor-

<sup>3</sup><http://www.gnu.org/software/pexec/pexec.1.html>

<sup>4</sup>[https://developer.apple.com/hardware/drivers/hpc/xgrid\\_intro.html](https://developer.apple.com/hardware/drivers/hpc/xgrid_intro.html)

<sup>5</sup><http://www.oracle.com/technetwork/oem/grid-engine-166852.html>

<sup>6</sup><http://www.pbsworks.com/>

<sup>7</sup><http://sourceforge.net/projects/neuropipedream/>

rection using the current white matter posterior probability as a weight mask and then using that bias corrected image as input to the Atropos segmentation step (described in subsequent sections).

### 2.1.3. Atropos 3-tissue segmentation

In [6] we presented an open source  $n$ -tissue segmentation software tool (which we denote as “Atropos”) attempting to distill 20+ years of active research in this area particularly some of its most seminal work (e.g. [76, 3]). Specification of prior probabilities includes spatially varying Markov Random Field modeling, prior label maps, and prior probability maps typically derived from our template building process. Additional capabilities include handling of multivariate data, partial volume modeling [60], a memory-minimization mode, label propagation, a plug-n-play architecture for incorporation of novel likelihood models which includes both parametric and non-parametric models for both scalar and tensorial images, and alternative posterior formulations for different segmentation tasks.

### 2.1.4. Brain extraction

Brain extraction using ANTs combines template building, high-performance brain image registration [5], and Atropos with topological refinements. An optimal template for brain extraction is generated offline using labeled brain data. For example, in this work we use the LPBA40 data for generating a brain extraction template and a corresponding brain probability mask which is available on the website associated with this submission.

The warped template probability map is thresholded at 0.5 and the resulting mask is dilated with a radius of 2. Atropos is used to generate an initial 3-tissue segmentation estimate within the mask region. Each of the three tissue masks undergo specific morphological operations which are then combined to create a brain extraction mask for use in the rest of the cortical thickness workflow.

A comparison using open access brain data with publicly available brain extraction algorithms including AFNI’s 3dIntracranial [71], FSL’s BET2 [64], Freesurfer’s `mri_watershed` [57], and BrainSuite [20] demonstrated that our combined registration/segmentation approach [4] performs at the top level alongside BrainSuite (tuned) and FreeSurfer.

### 2.1.5. DiReCT Cortical Thickness Estimation

Although the basic formulation of DiReCT as reported in this work is as it was introduced in [16], we have made several improvements (perhaps the most significant being that this particular ITK-compatible implementation has been significantly multi-threaded, is written in ITK coding style, and has been made publicly available through ANTs complete with a unique user interface design developed specifically for ANTs tools).

## 2.2. Public Data Resources

### 2.2.1. LPBA40 Data for Skull Stripping

For the brain extraction step we used the data from the LPBA40 repository [59]. These data consist of 40 high-

resolution 3D Spoiled Gradient Echo (SPGR) MRI acquisitions which were manually labeled delineating 56 brain structures. Additional post processing included automated brain extraction using FSL’s brain extraction tool [64] which was followed by manual corrections. These 40 brain masks are also included with the database.

The first 20 subjects were used to create a population-specific unbiased average template [7]. The brain masks corresponding to the 20 subjects were warped to the template space using the transforms derived during the template building process. A template probability mask was created by averaging the warped brain masks. For brain extraction of any single individual this ANTS-based LPBA40 template is coarsely registered to the single subject brain. The template probability brain mask is warped to the individual subject and is used as the initial brain mask estimate. As mentioned previously, Atropos and binary morphological operations are used to refine the brain mask estimate.

### 2.2.2. NIREP Data for 3-Tissue Segmentation

The nonrigid image registration evaluation project is an ongoing framework for evaluating image registration algorithms [11]. The initial data set introduced into the project consists of 16 (8 male and 8 female) high resolution skull-stripped brain data with 32 cortical labels manually drawn using published protocol. Given the gray matter labels, the white matter and CSF were identified for each of the 16 subjects using Atropos. Similar to the LPBA40 data set, a NIREP template was created from 8 subjects and each the warped labels were used to create probabilistic estimates of the labeled region boundaries. These probability maps were used as spatial prior probabilities during the 3-tissue segmentation component of the pipeline. Using SyN, the NIREP template is registered to the extracted individual subject brain which is followed by a warping of the NIREP priors to the space of the individual subject. The initial warped white matter probability map is used as the weighted confidence mask in the follow-up bias correction step.

## 2.3. IXI Data for Pipeline Evaluation

The IXI data<sup>8</sup> consists of approximately 600 healthy subjects imaged at three sites using several modalities (T1-weighted, T2-weighted, proton density, magnetic resonance angiography, and diffusion tensor imaging). The database also consists of demographic information such as age, weight, height, ethnicity, occupation category, educational level, and marital status. The number of subjects spanning a range of demographic characteristics makes this a rich data set for validating and exploring correlations with cortical thickness measured using the ANTs pipeline.

<sup>8</sup><http://biomedic.doc.ic.ac.uk/brain-development/>

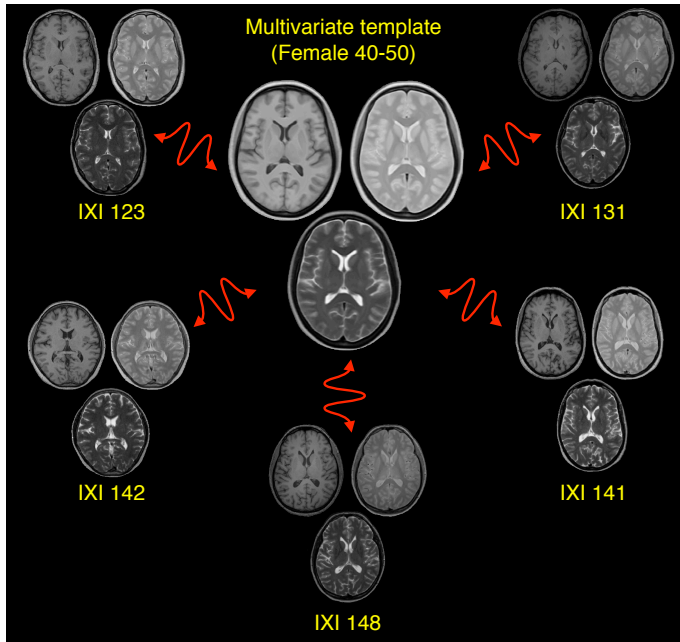


Figure 2: Sample multivariate template constructed from a subset of the IXI data (female, age 40–50). Axial slices of five of the 37 total subjects from this cohort are shown.

### 3. Evaluation

### 4. Discussion

### 5. Conclusions

### Acknowledgments

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