# The ANTsX Ecosystem for Mapping the Mouse Brain

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# Abstract

Precision mapping techniques coupled with high resolution image acquisition of the mouse brain permit the study of the spatial organization of gene activity and their mutual interac-18 tion for a comprehensive view of salient structural/functional relationships. Such research 19 is facilitated by standardized anatomical coordinate systems, such as the well-known Allen 20 Common Coordinate Framework (AllenCCFv3), and the ability to spatially map to such 21 standardized spaces. The Advanced Normalization Tools Ecosystem (ANTsX) is a comprehensive open-source software toolkit for generalized quantitative imaging, which includes 23 template building and mapping functionality, with applicability to multiple organ systems, modalities, and animal species. Herein, we illustrate the utility of ANTsX for generating precision spatial mappings of the mouse brain. First, we provide ANTsX-based protocols for mapping MERFISH, fMOST, and lightsheet datasets to AllenCCFv3 accounting for com-27 mon artefacts and other confounds. Novel ANTsX contributions include recently developed 28 ANTsX functionality for generating a velocity flow-based mapping spanning the spatiotem-29 poral domain of a longitudinal trajectory which we apply to the Developmental Common 30 Coordinate Framework (DevCCF). Additionally, we present an automated structural mor-31 phological pipeline for determining volumetric and cortical thickness measurements analogous to the well-utilized ANTsX pipeline for human neuroanatomical structural morphology. 33 This latter development also illustrates a more general open-source ANTsX framework for determining tailored brain parcellations using the AllenCCFv3 and DevCCF templates.

# <sub>36</sub> 1 Introduction

ysis of the mouse brain. It is now possible to track single cell neurons in mouse brains, 1 observe whole brain developmental changes on a cellular level.<sup>2</sup> associate brain regions and tissues with their genetic composition,<sup>3</sup> and locally characterize neural connectivity.<sup>4</sup> Much of this scientific achievement has been made possible due to breakthroughs in high resolution imaging techniques that permit submicron, 3-D imaging of whole mouse brains. Associated research techniques such as micro-optical sectioning tomography, 6 tissue clearing, 1,7 spatial transcriptomics<sup>9</sup> are all well-utilized in the course of scientific investigations of mesoscale relationships in the mouse brain. An important component of this research is the ability to map the various image data to anatomical reference frames<sup>11</sup> for inferring spatial relationships between structures, cells, and genetics. This has motivated the development of detailed structural image at lases of the mouse brain. Notable examples include the Allen Brain Atlas and Coordinate Frameworks (AllenCCFv3), 13 the Waxholm Space, 14 and more recently, the Developmental Common Coordinate Framework (DevCCF). 15 Despite the significance of these contributions, challenges 51 still exist in large part due to the wide heterogeneity in associated study-specific image data. For example, variance in the acquisition methods can introduce artifacts such as tissue distortion, holes, bubbles, folding, tears, and missing slices. These severely complicate assumed correspondence for conventional spatial mapping approaches.

Over the past two decades there have been significant advancements in mesoscopic anal-

# <sub>66</sub> 1.1 Mouse-specific brain mapping software

To address such challenges, several software packages have been developed over the years comprising solutions of varying comprehensibility, sophistication, and availability. An early contribution to the community was the Rapid Automatic Tissue Segmentation (RATS) package<sup>16</sup> for brain extraction. More recently, several publicly available packages comprise well-established package dependencies originally developed on human brain data. SPMMouse, <sup>17</sup> for example, is based on the well-known Statistical Parametric Mapping

(SPM) software package. 18 The automated mouse atlas propagation (aMAP) tool is largely a front-end for the NiftyReg image registration package<sup>19</sup> applied to mouse data which is currently available as a Python module.<sup>20</sup> NiftyReg is also used by the Atlas-based Imaging Data Analysis (AIDA) MRI pipeline<sup>21</sup> as well as the Multi Atlas Segmentation and Morphometric Analysis Toolkit (MASMAT). Whereas the former also incorporates the FMRIB Software Library (FSL)<sup>22</sup> for brain extraction and DSIStudio<sup>23</sup> for DTI processing, the latter uses NiftvSeg and multi-consensus labeling tools<sup>24</sup> for brain extraction and parcellation. In addition, MASMAT incorporates N4 bias field correction<sup>25</sup> from the Advanced Normalization Tools Ecosystem (ANTsX)<sup>26</sup> as do the packages Multi-modal Image Registration And Connectivity anaLysis (MIRACL), <sup>27</sup> Sammba-MRI, <sup>28</sup> and Small Animal Magnetic Resonance Imaging (SAMRI).<sup>29</sup> However, whereas Saamba-MRI uses AFNI<sup>30</sup> for image registration; MIRACL, SAMRI, and BrainsMapi<sup>31</sup> all use ANTsX registration tools. Other packages use landmark-based approaches to image registration including SMART—<sup>32</sup>an R package for semi-automated landmark-based registration and segmentation of mouse brain based on WholeBrain.<sup>33</sup> FriendlyClearMap<sup>34</sup> uses the landmark-based registration functionality of Elastix.<sup>35</sup> Finally, the widespread adoption of deep learning techniques has also influenced development in mouse brain imaging methodologies. For example, if tissue deformations are not considered problematic for a particular dataset, DeepSlice can be used to determine affine mappings $^{36}$  with the optimal computational efficiency associated with 81 neural networks.

# 1.2 The ANTsX Ecosystem for mouse brain mapping

As noted previously, many of the existing packages designed for processing mouse brain image data use ANTsX tools for core processing steps in various workflows, particularly its pairwise, intensity-based image registration capabilities and bias field correction. Historically, ANTsX development is originally based on fundamental approaches to image mapping, 37–39 particularly in the human brain, which has resulted in core contributions to the field such as the well-known and highly-vetted Symmetric Normalization (SyN) algorithm. 40 Since its development, various independent platforms have been used to evaluate ANTsX image regis-

- tration capabilities in the context of different application foci which include multi-site brain
  MRI data, 41 pulmonary CT data, 42 and most recently, multi-modal brain registration in the
  presence of tumors. 43
- Apart from its registration capabilities, ANTsX comprises additional functionality such as template generation, <sup>44</sup> point set data approximation, <sup>45</sup> and deep learning networks specifically trained for mouse data (see Table 1). The comprehensive use of the toolkit has demonstrated superb performance in multiple application areas (e.g., consensus labeling, <sup>46</sup> brain tumor segmentation, <sup>47</sup> and cardiac motion estimation <sup>48</sup>). Importantly, ANTs is built on the Insight Toolkit (ITK) <sup>49</sup> deriving benefit from the open-source community of scientists and programmers and providing an open-source venue for algorithmic development, evaluation, and improvement.

## 1.3 ANTsX-based open-source contributions

Consistent with previous ANTsX development, the newly introduced capabilities introduced below are available through ANTsX (specifically, via R and Python ANTsX packages), and illustrated through self-contained examples in the ANTsX tutorial (https://tinyurl.com/antsxtutorial) with a dedicated GitHub repository specific to this work (https://github.com/ntustison/ANTsXMouseBrainMapping).

#### 108 1.3.1 The DevCCF velocity flow model

Recently, the Developmental Common Coordinate Framework (DevCCF) was introduced to the mouse brain research community as a public resource<sup>15</sup> comprising symmetric atlases of multimodal image data and anatomical segmentations defined by developmental ontology. These templates sample the mouse embryonic days (E) 11.5, E13.5, E15.5, E18.5 and postnatal day (P) 4, P14, and P56. Modalities include light sheet flourescence miscroscopy (LSFM) and at least four MRI contrasts per developmental stage. Anatomical parcellations are also available for each time point and were generated from ANTsX-based mappings of gene expression and other cell type data. Additionally, the P56 template was integrated

Table 1: Sampling of ANTsX functionality

	ANTsPy: Preprocessing
bias field correction	n4_bias_field_correction()
image denoising	denoise_image()
	ANTsPy: Registration
image registration	registration()
image transformation	apply_transforms()
template generation	<pre>build_template()</pre>
landmark registration	<pre>fit_transform_to_paired_points()</pre>
time-varying landmark reg.	<pre>fit_time_varying_transform_to_point_sets()</pre>
integrate velocity field	<pre>integrate_velocity_field()</pre>
invert displacement field	<pre>invert_displacement_field()</pre>
	ANTsPy: Segmentation
MRF-based segmentation	atropos()
Joint label fusion	joint_label_fusion()
diffeormorphic thickness	kelly_kapowski()
	ANTsPy: Miscellaneous
Regional intensity statistics	label_stats()
Regional shape measures	label_geometry_measures()
B-spline approximation	<pre>fit_bspline_object_to_scattered_data()</pre>
Visualize images and overlays	plot()
ANTsPyNet: Mouse-specific	
brain extraction	mouse_brain_extraction(modality="t2")
	mouse_brain_extraction(modality="ex5")
brain parcellation	mouse_brain_parcellation()
cortical thickness	mouse_cortical_thickness()
super resolution	<pre>mouse_histology_super_resolution()</pre>

ANTsX provides state-of-the-art functionality for processing biomedical image data. Such tools, including deep learning networks, support a variety of mapping-related tasks. A more comprehensive listing of ANTsX tools with self-contained R and Python examples is provided as a gist page on GitHub (https://tinyurl.com/antsxtutorial).

with the Allen CCFv3 to further increase the practical utility of the DevCCF. These processes, specifically template generation and multi-modal image mapping, were performed using ANTsX functionality in the presence of image mapping difficulties such as missing slices, tissue distortion.

Given the temporal gaps in the discrete set of developmental atlases, we also provide an open-source framework, through ANTsX, for inferring correspondence within the temporally continuous domain sampled by the existing set of embryonic and postnatal atlases of the DevCCF. This recently developed ANTsX functionality permits the generation of a diffeomorphic velocity flow transformation model, influenced by previous work. The resulting time-parameterized velocity field spans the stages of the DevCCF where mappings between any two continuous time points within the span bounded by the E11.5 and P56 atlases is determined by integration of the optimized velocity field.

#### 29 1.3.2 Structural morphology and cortical thickness in the mouse brain

One of the most frequently utilized pipelines in the ANTsX toolkit is that of determining cortical thickness in the human brain originating with the Diffeomorphic Registration-based
Cortical Thickness (DiReCT) algorithm.<sup>52</sup> This development was later expanded to constitute a complete cortical thickness pipeline for the human brain for both cross-sectional<sup>53</sup> and
longitudinal<sup>54</sup> using T1-weighted MR image data. These pipelines were later significantly
refactored using deep learning innovations.<sup>26</sup>

In contrast to the pipeline development in human data,<sup>26</sup> no current ANTsX tools exist to create adequate training data for the mouse brain. In addition, mouse brain data acquisition often has unique issues, such as lower data quality or sampling anisotropy which limits its applicability to high resolution resources (e.g., AllenCCFv3, DevCCF), specifically with respect to the corresponding granular brain parcellations derived from numerous hours of expert annotation leveraging multimodal imaging resources.

Herein, we introduce a mouse brain cortical thickness pipeline comprising two novel deep learning components: two-shot learning brain extraction from data augmentation of two ANTsX templates generated from two open datasets<sup>55,56</sup> and single-shot brian parcellation derived from the AllenCCFv3 labelings mapped to the corresponding DevCCF P56 time point. Although we anticipate that this cortical thickness pipeline will be beneficial to the research community, this work demonstrates more generally how one can leverage ANTsX tools for developing tailored brain parcellation schemes. Evaluation is performed on an independent open data set<sup>57</sup> comprising longitudinal acquisitions of multiple specimens.

# 50 Results

# 2.1 The DevCCF Velocity Flow Model

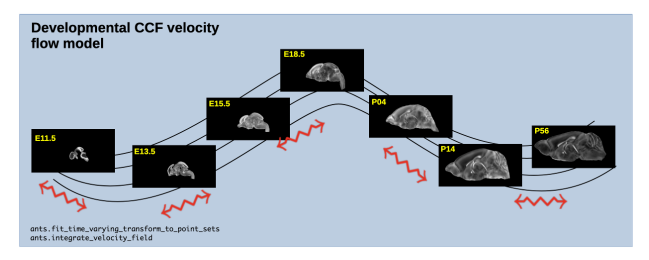


Figure 1: Using a velocity flow model, the transformation between any two temporal time points within the DevCCF is possible.

To continuously interpolate transformations between the different stages of the DevCCF atlases, a velocity flow model was constructed using DevCCF derived data and ANTsX functionality recently introduced into both the ANTsR and ANTsPy packages. Both platforms include a complete suite of functions for determining dense correspondence from sparse 155 landmarks based on a variety of transformation models ranging from standard linear models 156 (i.e., rigid, affine) to deformable diffeomorphic models (e.g, symmetric normalization).<sup>40</sup> The 157 latter set includes transformation models for both the pairwise scenario and for multiple sets, as in the case of the DevCCF. ANTsX, being built on top of ITK, uses an ITK image 159 data structure for the 4-D velocity field where each voxel contains the x, y, z components of 160 the field at that point. 161

#### 2.1.1 Data preparation

Labeled annotations are available as part of the original DevCCF and reside in the space of each developmental template which range in resolution from  $31.5 - 50\mu m$ . Across all atlases, the total number of labeled regions exceeds 2500. From these labels, a common set

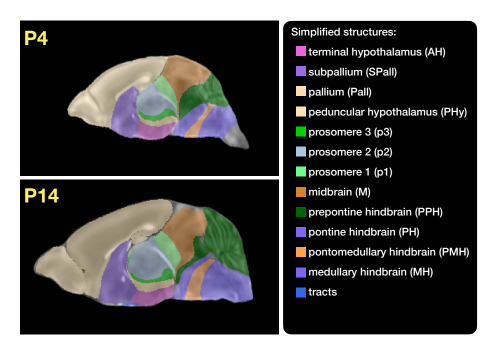


Figure 2: Annotated regions representing common labels across developmental stages which are illustrated for both P4 and P14.

of 26 labels (13 per hemisphere) across all atlases were used for optimization and evaluation.

These regions are illustrated for the P4 and P14 stages in Figure 2.

Prior to velocity field optimization, all data were rigidly transformed to a common space. 168 Using the centroids for the common label set of each DevCCF atlas, each atlas was rigidly 169 aligned to the space of the P56 atlas. In order to determine the landmark correspondence 170 across DevCCF stages, the multi-metric capabilities of ants.registration(...) were used. 171 Instead of performing intensity-based pairwise registration directly on these multi-label im-172 ages, each label was used to construct a separate fixed and moving image pair resulting in a 173 multi-metric registration optimization scenario involving 24 binary image pairs (each label 174 weighted equally) for optimizing diffeomorphic correspondence between neighboring time 175 point at lases using the mean squares metric and the symmetric normalization transform. 176

To generate the set of common point sets across all seven developmental atlases, the label boundaries and whole regions were sampled in the P56 atlas and then propagated to each atlas using the transformations derived from the pairwise registrations. We selected a sampling rate of 10% for the contour points and 1% for the regional points for a total number of points being per atlas being 173303 ( $N_{contour} = 98151$  and  $N_{region} = 75152$ ). Regional

boundary points were weighted twice as those of non-boundary points during optimization.

#### 2.1.2 Optimization

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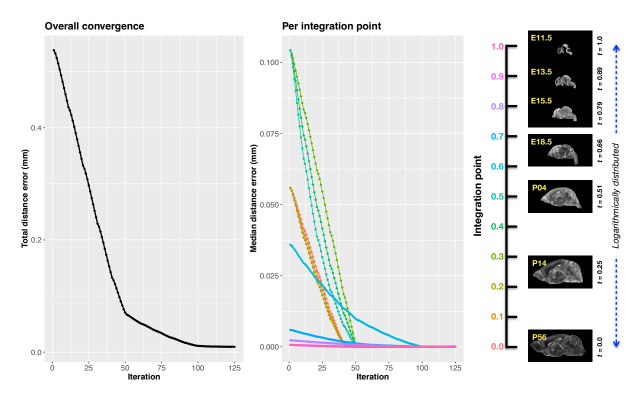


Figure 3: Convergence of the optimization of the velocity field for describing the transformation through the developmental stages from E11.5 through P56.

The velocity field was optimized using the input composed of the seven corresponding point 184 sets and their associated weight values, the selected number of integration points for the 185 velocity field (N = 11), and the parameters defining the geometry of the spatial dimensions 186 of the velocity field. Thus, the optimized velocity field described here is of size [256, 182, 360] 187  $(50\mu \text{m isotropic}) \times 11$  integration points for a total compressed size of a little over 2 GB. 188 This choice represented weighing the trade-off between tractability, portability, and accuracy. 189 However, all data and code to reproduce the results described are available in the dedicated 190 GitHub repository. 191

The normalized time point scalar value for each atlas/point-set in the temporal domains [0, 1] was also defined. Given the increasingly larger gaps in the postnatal timepoint sampling, we made two adjustments. Based on known mouse brain development, we used 28 days for the

P56 data. We then computed the log transform of the adjusted set of time points prior to normalization between 0 and 1 (see the right side of Figure 3). This log transform, as part of the temporal normalization, significantly improved data spacing.

The max number of iterations was set to 200 with each iteration taking six minutes. At each 198 iteration we looped over the 11 integration points. At each integration point, the velocity field 199 estimate was updated by warping the two immediately adjacent point sets to the integration 200 time point and determining the regularized displacement field between the two warped point 201 sets. As with any gradient-based descent algorithm, this field was multiplied by a small step 202 size  $(\delta = 0.2)$  before adding to the current velocity field. Convergence is determined by the 203 average displacement error over each of the integration points. As can be seen in the left 204 panel of Figure 3, convergence occurred around 125 iterations when the average displacement 205 error over all integration points is minimized. The median displacement error at each of the 206 integration points also trends towards zero but at different rates. 207

#### 2.1.3 The transformation model

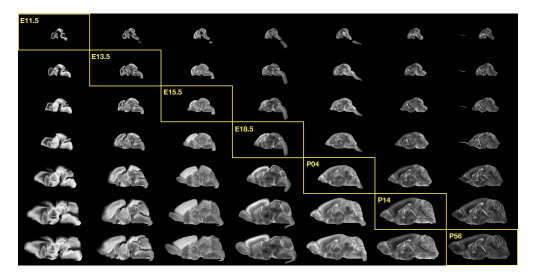


Figure 4: Mid-sagittal visualization of the effects of the transformation model in warping every developmental stage to the time point of every other developmental stage. The original images are located along the diagonal. Columns correspond to the warped original image whereas the rows represent the reference space to which each image is warped.

Once optimized, the resulting velocity field can be used to generate the deformable transform

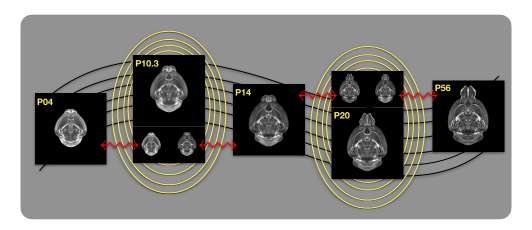


Figure 5: Illustration of the use of the velocity flow model for creating virtual templates at continuous time points not represented in one of the existing DevCCF time points. For example, FA templates at time point P10.3 and P20 can be generated by warping the existing temporally adjacent developmental templates to the target time point and using those images in the ANTsX template building process.

between any two continuous points within the time interval bounded by E11.5 and P56. In 210 Figure 4, we transform each atlas to the space of every other atlas using the DevCCF 211 transform model. Additionally, one can use this transformation model to construct virtual 212 templates in the temporal gaps of the DevCCF. This is illustrated in Figure 5 where we 213 used the optimized velocity field to construct virtual templates at time point P10.3 and 214 P20—arbitrarily chosen simply to demonstrate the concept. After situating these time points 215 within the normalized time point interval, the existing adjacent DevCCF at lases on either 216 chronological side can be warped to the desired time point. A subsequent call to one of the 217 ANTsX template building functions then permits the construction of the template at that 218 time point. Note that both of these usage examples can be found in the GitHub repository 219 previously given. 220

# 2.2 The Mouse Cortical Thickness Pipeline

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One of the most well-utilized pipelines in the ANTsX toolkit is the generation of cortical thickness in the human brain from T1-weighted MRI. Starting with the novel Diffeomor-phic Registration-based Cortical Thickness (DiReCT) algorithm,<sup>52</sup> a complete algorithmic workflow was developed for both cross-sectional<sup>53</sup> and longitudinal<sup>54</sup> using T1-weighted MR

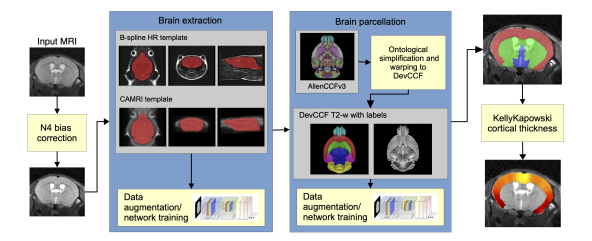


Figure 6: The mouse brain cortical thickness pipeline integrating two deep learning components for brain extraction and brain parcellation prior to estimating cortical thickness. Both brain extraction and parcellation pipelines rely heavily on ANTsX tools for template building and data augmentation as well as open-science data availability.

image data. This contribution was later refactored using deep learning<sup>26</sup> which leveraged the earlier results for training data.

In the case of the mouse brain, the lack of training data and/or tools to generate training
data make a similar developmental trajectory difficult. In addition, mouse data is often
characterized by unique issues such as frequent anisotropic sampling which are in sharp
contrast to high resolution resources available within the community, e.g., AllenCCFv3 and
DevCCF. Using ANTsX and other publicly available data resources, we developed a complete
mouse brain structural morphology pipeline as illustrated in Figure 6 and detailed below.

#### 2.2.1 Two-shot mouse brain extraction network

In order to create a generalized mouse brain extraction network, we built whole-head templates from two publicly available datasets. The Center for Animal MRI (CAMRI) dataset<sup>55</sup>
from UNC consist of 16 T2-weighted MRI of voxel resolution  $0.16 \times 0.16 \times 0.16 \times 0.16mm^3$ . The
second high-resolution data set<sup>56</sup> comprised 88 specimens each with three spatially aligned
canonical views with in-plane resolution of  $0.08 \times 0.08mm^2$  with a slice thickness of 0.5
mm. These three orthogonal views were used to reconstruct a single high-resolution volume per subject using a B-spline fitting algorithm developed in ANTsX.<sup>45</sup> From these two

datasets, two symmetric isotropic ANTsX templates<sup>44</sup> were generated having different "defacing" aesthetics analogous to our publicly available ANTsX human brain templates.<sup>53</sup> Bias field simulation, intensity histogram warping, noise simulation, random translation and warping, and random anisotropic resampling in the three canonical directions was used for data augmentation in creating a T2-weighted brain extraction network.

#### 2.2.2 Single-shot mouse brain parcellation network

To create a brain parcellation network conducive to cortical thickness map generation, we used the AllenCCFv3 and the associated allensdk Python utility. Using allensdk, a gross parcellation labeling was generated which included the cerebral cortex, cerebral nuclei, brain stem, cerebellum, main olfactory bulb, and hippocampal formation. This labeling was mapped to the T2-weighted template component of the P56 DevCCF was used to create a brain parcellation network in combination with the same data augmentation used for the brain extraction network.

#### $_{255}$ 2.2.3 Evaluation

For evaluation, we used the another publicly available dataset<sup>57</sup> which is completely independent from the data used in training the brain extraction and parcellation networks. Data includes 12 specimens each imaged at seven time points (Day 0, Day 3, Week 1, Week 4, Week 8, Week 20) with available brain masks. In-plane resolution is  $0.1 \times 0.1 mm^2$  with a slice thickness of 0.5 mm.

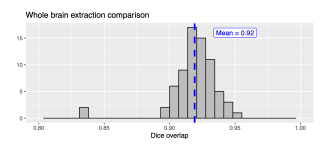


Figure 7

# Discussion

The ANTsX ecosystem is a powerful framework that has demonstrated applicability to multiple species and organ systems, including the mouse brain. This is further evidenced by 263 the many other software packages that use various ANTsX components in their own mouse-264 specific workflows. The extensive functionality of ANTsX per se makes it possible to create 265 complete processing pipelines without requiring the integration of multiple packages. These 266 open-source ANTsX components not only perform well but are available across multiple popular platforms which facilitates the construction of tailored pipelines for individual study 268 solutions. These components are also supported by years of development not only by the 269 ANTsX development team but by the larger ITK community. 270 In the case of the development of the DevCCF, ANTsX was crucial in providing necessary 271 functionality for yielding high quality output. First, for the generation of the individual 272 developmental stage multi-modal, symmetric templates, ANTsX is unique amongst image 273 analysis software packages in providing existing solutions for template generation which have 274 been thoroughly vetted, including being used in several studies over the years, and which 275 continue to be under active refinement. At its core, computationally efficient and quality template generation requires the use of precision pairwise image mapping functionality which, 277 historically, is at the origins of the ANTsX ecosystem. And these mapping capabilities extend 278 beyond template generation to the mapping of other image data (e.g., gene expression maps) 279 to template for providing further insight into the mouse brain. 280 Despite the significant expansion of available developmental age templates beyond what pre-281 viously existed (e.g., Allen CCFv3), there still exist temporal gaps in the DevCCF. However, 282 pioneering work involving diffeomorphic transformations allowed us to continuously situate 283 the existing templates within a time-varying velocity flow model. This allows one to deter-284 mine the diffeomorphic transformation from any one temporal location to any other temporal 285 location within the time span defined by the E11.5 and P56 templates. This functionality 286 is built on multiple components from the Insight Segmentation and Registration Toolkit 287 including the B-spline scattered data approximation technique for field regularization and 288

velocity field integration using fourth order Runge-Kutta. This velocity field model permits

- intra-template comparison and the construction of virtual templates where a template can be
  estimated at any continuous time point within the temporal domain. This novel application
  can potentially enhance our understanding of intermediate developmental stages.
- We also presented a mouse brain pipeline for brain extraction, parcellation, and cortical thickness that did not necessitate the extensive quantity of data required for training our analogous human brain pipeline.
- To increase its impact and reproduce the results shown previously, we have made the data and code publicly available at https://github.com/ntustison/ANTsXMouseBrainMapping.

Field regularization is provided by a B-spline scattered data approximation technique<sup>51</sup> which
permits individual point weighting. Both field regularization and integration of the velocity
field are built on ITK functions contributed from ANTsX development.

# Methods

The following methods are all available as part of the ANTsX ecosystem with analogous elements existing in both ANTsR (ANTs in R) and ANTsPy (ANTs in Python) with and ANTs/ITK C++ core. However, most of the development for the work described below was performed using ANTsPy. For equivalent calls in ANTsR, please see the ANTsX tutorial at https://tinyurl.com/antsxtutorial.

## Preprocessing: bias field correction and denoising

As in human studies, bias field correction and image denoising are standard preprocessing steps in improving overall image quality in mouse brain images. The bias field, a gradual 309 spatial intensity variation in images, can arise from various sources such as magnetic field in-310 homogeneity or acquisition artifacts, leading to distortions that can compromise the quality 311 of brain images. Correcting for bias fields ensures a more uniform and consistent repre-312 sentation of brain structures, enabling accurate quantitative analysis. Additionally, brain 313 images are often susceptible to various forms of noise, which can obscure subtle features 314 and affect the precision of measurements. Denoising techniques help mitigate the impact 315 of noise, enhancing the signal-to-noise ratio and improving the overall image quality. The 316 well-known N4 bias field correction algorithm<sup>25</sup> has its origins in the ANTs toolkit which 317 was implemented and introduced into the ITK toolkit. Similarly, ANTsX contains an im-318 plementation of a well-performing patch-based denoising technique<sup>58</sup> and is also available as 319 an image filter to the ITK community.

# ANTsXNet mouse brain applications

- 322 General notes regarding deep learning training.
- All network-based approaches described below were implemented and organized in the
- ANTsXNet libraries comprising Python (ANTsPyNet) and R (ANTsRNet) analogs using the
- Keras/Tensorflow libraries available as open-source in ANTsX GitHub repositories. For the
- various applications, both share the identically trained weights for mutual reproducibility.
- Training data was provided by manual labeling by various co-authors and expanded using
- both intensity-based and shape-based data augmentation techniques.
- 329 Intensity-based data augmentation consisted of randomly added noise based on
- 330 ITK functionality, simulated bias fields based on N4 bias field modeling, and his-
- togram warping for mimicking well-known MRI intensity nonlinearities. 26,59 These
- augmentation techniques are available in ANTsXNet (only ANTsPyNet versions are
- 333 listed): simulated bias field: antspynet.simulate\_bias\_field(...), image noise:
- antspyhet.add\_noise\_to\_image(...), and MRI intensity nonlinear characteriza-
- sation: antspynet.histogram\_warp\_image\_intensities(...). Shape-based data
- augmentation used both random linear and nonlinear deformations. This func-
- tionality is also instantiated within ANTsXNet in terms of random spatial warping:
- antspynet.randomly\_transform\_image\_data(...).
- For all GPU training, we used Python scripts for creating custom batch generators. As such
- batch generators tend to be application-specific, we store them in a separate GitHub reposi-
- tory for public availability (https://github.com/ntustison/ANTsXNetTraining). In terms of
- GPU hardware, all training was done on a DGX (GPUs: 4X Tesla V100, system memory:
- <sup>343</sup> 256 GB LRDIMM DDR4).
- 344 Brain extraction.
- Similar to human neuroimage processing, brain extraction is a crucial preprocessing step for
- accurate brain mapping. Within ANTsXNet, we have created several deep learning networks
- for brain extraction for several image modalities (e.g., T1, FLAIR, fractional anisotropy).
- Similarly, for the developmental brain atlas work<sup>15</sup> we developed similar functionality for

mouse brains of different modalities and developmental age. All networks use a conventional 2-D U-net architecture<sup>60</sup> and perform prediction in a slice-wise fashion given the 350 limitations of the acquisition protocols (e.g., missing slices, slice thickness). Currently, 351 coronal and sagittal networks are available for both E13.5 and E15.5 data and coronal 352 network for T2-weighted MRI. In ANTsPyNet, this functionality is available in the program antspynet.mouse brain extraction(...). Even when physical brain extraction is 354 performed prior to image acquisition, artifacts, such as bubbles or debris, can complicate 355 subsequent processing. Similar to the brain extraction networks, a 2-D U-net architecture<sup>60</sup> 356 was created to separate the background and foreground. 357

<sup>358</sup> Miscellaneous networks: Super-resolution, cerebellum, and hemispherical masking.

To further enhance the data prior to designing mapping protocols, additional networks were created. A well-performing deep back projection network<sup>61</sup> was ported to ANTsXNet and expanded to 3-D for various super-resolution applications,<sup>62</sup> including mouse data. Finally, features of anatomical significance, namely the cerebellum and hemispherical midline were captured in these data using deep learning networks.

# Intra-slice image registration with missing slice imputation

Volumetric gene expression slice data was collated into 3-D volumes. Prior to mapping 365 this volume to the corresponding structural data and, potentially, to the appropriate template, alignment was improved using deformable registration on contiguous slices. How-367 ever, one of the complications associated with these image data was the unknown num-368 ber of missing slices, the number of consecutive missing slices, and the different locations 369 of these missing slices. To handle this missing data problem, we found that data in-370 terpolation using the B-spline approximation algorithm cited earlier<sup>45</sup> (ANTsPy function: 371 ants.fit bspline object to scattered data(...)). This provided sufficient data in-372 terpolation fidelity to perform continuous slicewise registration. Other possible variants that 373 were considered but deemed unnecessary was performing more than one iteration cycling 374 through data interpolation and slicewise alignment. The other possibility was incorporating 375 the super-resolution technique described earlier. But again, our data did not require these

## 378 Image registration

The ANTs registration toolkit is a complex framework permitting highly tailored solu-379 tions to pairwise image registration scenarios.<sup>63</sup> It includes innovative transformation mod-380 els for biological modeling<sup>40,51</sup> and has proven capable of excellent performance.<sup>41,64</sup> Var-381 ious parameter sets targeting specific applications have been packaged with the different ANTSX platforms, specifically ANTs, ANTsPy, and ANTsR.<sup>26</sup> In ANTsPy, the function ants.registration(...) is used to register a pair of images or a pair of image sets where 384 type of transform is a user-specified option that invokes a specific parameter set. For 385 example type of transform='antsRegistrationSyNQuick[s]' is an oft-used parameter set. Initially, linear optimization is initialized with center of (intensity) mass alignment typically 388

followed by optimization of both rigid and affine transforms using the mutual information similarity metric. This is followed by diffeomorphic deformable alignment using symmetric normalization (SyN) with Gaussian<sup>40</sup> or B-spline regularization<sup>51</sup> where the forward transform is invertible and differentiable. The similarity metric employed at this latter stage is typically either neighborhood cross-correlation or mutual information. Note that these parameter sets are robust to input image type (i.e., LSFM, Nissl staining, and the various MRI modalities) and are adaptable to mousing image geometry scaling. Further details can be found in the various documentation sources for these ANTsX packages.

# Template generation

ANTsX provides functionality for constructing templates from a set (or multi-modal sets) of input images as originally described<sup>44</sup> and recently used to create the DevCCF templates.<sup>15</sup>
An initial template estimate is constructed from an existing subject image or a voxelwise average derived from a rigid pre-alignment of the image population. Pairwise registration between each subject and the current template estimate is performed using the Symmetric

Normalization (SyN) algorithm.<sup>40</sup> The template estimate is updated by warping all subjects to the space of the template, performing a voxelwise average, and then performing a "shape update" of this latter image by warping it by the average inverse deformation, thus yielding a mean image of the population in terms of both intensity and shape.

## 407 Continuous developmental velocity flow transformation model

Given multiple, linearly or non-linearly ordered point sets where individual points across are 408 in one-to-one correspondence, we developed an approach for generating a velocity flow trans-409 formation model to describe a time-varying diffeomorphic mapping as a variant of the inexact 410 landmark matching solution. Integration of the resulting velocity field can then be used to 411 describe the displacement between any two time points within this time-parameterized do-412 main. Regularization of the sparse correspondence between point sets is performed using a 413 generalized B-spline scattered data approximation technique, 45 also developed by the ANTsX 414 developers and contributed to ITK. 415

To apply this methodology to the developmental templates, <sup>15</sup> we coalesced the manual par-416 cellations of the developmental templates into 26 common anatomical regions (13 per hemisphere). We then used these regions to generate invertible transformations between succes-418 sive time points. Specifically each label was used to create a pair of single region images 419 resulting in 26 pairs of "source" and "target" images. The multiple image pairs were used 420 to iteratively estimate a diffeomorphic pairwise transform. Given the seven atlases E11.5, 421 E13.5, E15.5, E18.5, P4, P14, and P56, this resulted in 6 sets of transforms between succes-422 sive time points. Given the relative sizes between atlases, on the order of 10<sup>6</sup> points were 423 randomly sampled labelwise in the P56 template space and propagated to each successive 424 atlas providing the point sets for constructing the velocity flow model. Approximately 125 425 iterations resulted in a steady convergence based on the average Euclidean norm between transformed point sets. Ten integration points were used and point sets were distributed along the temporal dimension using a log transform for a more evenly spaced sampling. 428

# Synthesizing isotropic image volumes from orthogonal views

As we mentioned previously, the multi-plane, high resolution data<sup>56</sup> was used to create single isotropic volumes using the B-spline fitting algorithm.<sup>45</sup> This algorithm is encapsulated in ants.fit\_bspline\_object\_to\_scattered\_data(...) where the input is the set of voxel intensity values and associated physical location. Since each point can be assigned a confidence weight, we use the the normalized gradient value to more heavily weight edge regions.

## Visualization

To complement the well-known visualization capabilities of R and Python, e.g., ggplot2 and matplotlib, respectively, image-specific visualization capabilities are available in the ants.plot(...) (Python) and plot.antsImage(...) (R). These are capable of illustrating multiple slices in different orientations with both other image overlays as well as label images. Data availability. All data and doftware used in this work are publicly available. The DevCCF atlas is available at https://kimlab.io/brain-map/DevCCF/.

ANTsPy, ANTsR, ANTsPyNet, and ANTsRNet are available through GitHub at the ANTsX Ecosystem (https://github.com/ANTsX). A GitHub repository specific to the work discussed in the manuscript was created and is available at https://github.com/ntustison/ANTsXMouseBrainMapping.

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