Magnetic resonance imaging datasets with anatomical fiducials for quality control and registration

Alaa Taha1,2, Greydon Gilmore1,2,3, Mohamad Abbass1,3,4, Jason Kaj1,4, Tristan Kuehn1,2, John Demarco1,5, Geetika Gupta1,4, Chris Zajner1,3, Daniel Cao1,2,7, Ryan Chevalier3, Abrar Ahmed1,3, Ali Hadi1,3, Bradley G. Karat1,4, Olivia W. Stanley1,5, Patrick J. Park1,2, Kayla M. Ferko1,4, Dimuthu Hemachandra1,2, Reid Vassallo8, Magdalena Jach3, Arun Thurairajah1,6, Sandy Wong1,2, Mauricio C. Tenorio1,2, Feyi Ogunsanya1,5, Ali R. Khan1,2,3,4,9 & Jonathan C. Lau1,2,3,4,9

Tools available for reproducible, quantitative assessment of brain correspondence have been limited. We previously validated the anatomical fiducial (AFID) placement protocol for point-based assessment of image registration with millimetric (mm) accuracy. In this data descriptor, we release curated AFID placements for some of the most commonly used structural magnetic resonance imaging datasets and templates. The release of our accurate placements allows for rapid quality control of image registration, teaching neuroanatomy, and clinical applications such as disease diagnosis and surgical targeting. We release placements on individual subjects from four datasets (N = 132 subjects for a total of 15,232 fiducials) and 14 brain templates (4,288 fiducials), totalling more than 300 human rater hours of annotation. We also validate human rater accuracy of released placements to be within 1 – 2 mm (using more than 45,000 Euclidean distances), consistent with prior studies. Our data is compliant with the Brain Imaging Data Structure allowing for facile incorporation into neuroimaging analysis pipelines.

Background & Summary

Open resources available for reproducible, quantitative assessment of brain correspondence have been limited1. The most common metrics employed for the purpose of examining the quality of image registration, including the Jaccard similarity and Dice kappa coefficients, compute the voxel overlap between regions of interest (ROIs), which have been shown to be insufficiently sensitive when used in isolation or in combination for validating image registration strategies1. The ROIs used in voxel overlap are often larger subcortical structures that are readily visible on magnetic resonance imaging (MRI) scans (e.g., the thalamus, globus pallidus, and striatum), and thus lack the ability to detect subtle misregistration between images which may be crucial where millimetric differences in variability should be accounted for1–5.

Inspired by classic stereotactic methods, our group created, curated, and validated a protocol for the placement of anatomical fiducials (AFIDs) on structural MRI scans of the human brain2. The protocol involves the placement of 32 AFIDs found to have salient features that allow for accurate localization. The AFIDs are described using three-dimensional (x, y, and z) Cartesian coordinates and thus correspondence between points can be computed using Euclidean distances across a variety of applications. After a brief tutorial, AFIDs have been shown to be highly reproducible even when performed by individuals with no prior knowledge of medical imaging.
images, neuroanatomy, or neuroimaging software. This was shown in separate studies where placements were performed on publicly available templates and datasets and a clinical neuroimaging dataset.

The AFID protocol provides a metric that is independent of the registration itself while offering sensitivity to registration errors at the scale of millimeters (mm). This margin is crucial in neuroimaging applications (including morphometric analysis and surgical neuromodulation), where a few mm may represent the difference between optimal and suboptimal therapy.

The aim of this data descriptor is to provide the community with curated AFID placements and their associated MRI images. We release annotations on four datasets (N = 132; 15,232 fiducials) including healthy subjects and patients with neurological disorders, and 14 commonly used MRI templates (4,288 fiducials), totalling more than 300 human rater hours of manual annotation of neuroanatomical structures. Descriptions of the datasets and templates are provided in subsequent sections. We highlight current and prospective applications of our released data in Fig. 1.

**Current applications.** *Registration assessment.* We share our curated AFID annotations for a wide variety of datasets and templates of varying field strengths. This diversity of datasets will facilitate the testing and validation of image registration algorithms that can be used in many contexts. The user can select the datasets and templates that are in line with their neuroimaging application, then use the curated annotations to assess image registration quantitatively. For instance, AFIDs have been used to evaluate the process of iterative deformable template creation, showing that error metrics generated from AFIDs converged differently as a function of template iterations and registration method (i.e., linear vs non-linear). Sharing the AFID placements and their associated images in the Brain Imaging Data Structure (BIDS) format aids in the convenience we strive to provide for the end-user and neuroimaging application developer.

**Prospective applications.** *Registration optimization and quality control.* The released imaging and AFID placement data may be useful in a few ways for improving neuroimaging pipelines: (1) providing centralized and quality controlled neuroimaging data (from 4 different neuroimaging datasets) allowing for a more accurate and generalizable head-to-head comparison among existing software for image registration, and (2) establishing a new registration metric that can be incorporated into neuroimaging software development workflows to optimize registration algorithm performance and ensure quality control.

**Automatic and accurate landmark placement.** Our curated AFIDs can serve as ground truth placements when training machine learning algorithms to automate brain landmark localization. Among the 32 AFIDs we release are the anterior and posterior commissures (AC and PC, respectively). Downstream applications of automatic localization include automatically computing AC-PC transformation (a common process in neuroimaging studies) and aspects of neurosurgical planning that involve the placement of these anatomical landmarks. The diversity of the released data (both hardware and disease status) will be crucial to the generalizability of such tools.

**Surgical targeting.** We release locally curated ultra-high field (7-Tesla; 7-T) MRI data where small structures like the subthalamic nucleus (STN) and zona incerta within the posterior subthalamic area are clearly visible. Ground truth locations of surgical targets (x, y, and z) can be related to AFID locations via predictive models. This approach mitigates the lack of access to best-case neuroimaging in clinical settings due to limited access to high-field MRI or motion degradation.

**Brain anatomy abstraction and anonymization.** AFIDs and the distances between them represent an abstraction of brain anatomy in an anonymized way while still allowing for the accurate pooling of data. Other significant anatomical landmarks (representing lesions, tumors, or other structures) can be described in reference to the AFID “coordinate system” we establish using these curated placements.

**Methods**

**Rationale for fiducial selection and placement assessments.** The current version of the AFID protocol involves the placement of 32 landmarks. They were selected to be easily identified on structural MRI scans across varying field strengths (1.5-T, 3-T, 7-T) and were validated in previous studies. During the selection process, regions that were prone to geometric inhomogeneity and distortion were avoided to enhance the accuracy of fiducial placement. There are 10 fiducials that fall on the midline and 11 located laterally on both hemispheres (see Table 1). The AFID protocol includes landmarks representing salient neuroanatomical features mostly located in the subcortex (see Fig. 2).
Fiducial localization error (FLE) is a term described by Fitzpatrick and colleagues that represents the distance between a fiducial position from its intended location. This term is used when operating image-guidance systems during surgical procedures. In the context of the AFID protocol, and inspired by this extant terminology, we have defined the term anatomical fiducial localization error (AFLE). This value, in mm, can be thought of as the error arising from the placement (i.e., localization) of each fiducial. When used to communicate the accuracy of all AFIDs together, we term it global AFLE. There are three contexts for applying AFLEs: (1) mean AFLE: rater localization error relative to the intended location defined as the mean placement of all raters for a specific fiducial (termed ground truth AFID in subsequent sections). (2) inter-rater AFLE: rater localization error calculated as the pairwise distances between different rater placements. If a single rater performed the AFID protocol more than once, then their mean placement coordinates were used for the pairwise distance calculations. (3) intra-rater AFLE: rater localization error evaluating the precision of multiple placements by a single rater computed as the average pairwise distance between the same rater's placements.

Fig. 1 Current and prospective applications of curated anatomical fiducial (AFID) placements. Top Panel: Current applications in neuroanatomy education and image registration. Middle Panel: Released healthy and pathologic datasets and templates (detailed descriptions can be found in text). Bottom Panel: Prospective applications of AFIDs in stereotactic targeting and as a disease biomarker.
We also adopt the term *fiducial registration error (FRE)* in the context of the AFID protocol and term it the anatomical fiducial registration error (AFRE). It is important to note that AFRE in our context diverges from the original usage by Fitzpatrick and colleagues\(^\text{10}\) which was restricted to describe registration error at fiducials used to drive image registration (i.e., during landmark-based registration). Computed in mm, AFRE is defined as errors arising from the registration protocol performed between two images (often, but not limited to, subject and template). AFRE is the distance after co-registration between each of the 32 AFIDs placed on a moving image and their counterparts placed on the fixed image (i.e., homologous points). The average AFRE of all fiducials is termed the global AFRE. We also establish nomenclature to differentiate various use cases for AFRE. If an individual rater placement is chosen for subsequent analysis, then we term the resulting AFRE to be the real-world AFRE as it is more representative of what would happen in a clinical setting where one rater would apply the AFID protocol. If a ground truth AFID placement is used, then the resulting error is termed consensus AFRE as it represents the average placement among a group of raters prior to the image registration step. In this data descriptor, our focus is on releasing the curated AFID placements and not an assessment of registration, so no AFRE metrics are produced. We still felt it would be useful to introduce AFRE as its computation constitutes

Fig. 2 Curated AFID locations within the brain and usage of the AFIDs validator website. Top Panel: Distribution of AFIDs overlayed on one of the released templates. We also show major subcortical structures with AFIDs (black points) in various anatomical views. Bottom Panel: Uses and outputs from the AFIDs validator website (https://validator.afids.io)\(^\text{8}\). The user decides whether to upload their placements to our database and will receive summary metrics regarding their placements in an interactive 3D coordinate system and tabular format (not shown).
one of the main applications of AFIDs and our shared datasets for quality control (i.e., in the context of image registration).

**Hardware and software used to curate data.** All manually curated AFIDs were placed using the Markups Module of 3DSlicer (an open-source imaging software)\(^1\). The datasets were curated at different times so a reference to the exact version of 3DSlicer and associated modules will be made under each dataset. 3DSlicer was chosen because it offers a variety of modules, particularly markups and registration modules used for fiducial placement and AC-PC transformation. 3DSlicer stores points placed within its 3D coordinate system overlaid on the image giving the possibility of more accurate localization without the need to interpolate to the nearest voxel. The AFID placements released here for templates and datasets were performed on structural T1w MRI images.

**Performing the AFID protocol.** All raters underwent extensive training before being involved in any AFID related studies\(^2\,^3\). More specifically, they (1) attended a synchronous session about 3DSlicer and placed all the AFIDs under the supervision of expert raters, (2) were asked to refer to resources found on our AFID protocol website (https://afids.github.io/afids-protocol)\(^12\) to supplement their learning asynchronously, and (3) uploaded their annotations to the AFIDs validator tool for feedback with further review with an expert to ensure that their annotations were of sufficient quality. We collected demographic data (neuroanatomy, imaging, and 3DSlicer exposure) for raters involved in data curation (see Data Records).

For manual rater placements, the AFID protocol generally began with the placement of the anterior commissure (AC) and posterior commissure (PC) points (AFID01 and 02, respectively), which are defined to be at the center of each commissure. This was then followed by the identification of one or two more midline points (often the pontomesencephalic junction, AFID04, and the genu of corpus callosum, AFID19, are used). After that, an AC-PC transformation is performed, and the rest of the anatomical fiducials are placed. Rater placements deviating from a ground truth fiducial by greater than 10 mm were removed and considered outliers, as these errors are likely to be due to mislabelling and not reflective of true localization accuracy. In addition to subsequent sections, Table 2 provides brief descriptions of the released datasets and templates, information about raters, and AFID placements.

**AFIDs-HCP30 dataset.** *Subject demographics and imaging protocol.* This subset consists of 30 unrelated healthy subjects (age: 21 – 52 years; 15 female) chosen from the Human Connectome Project dataset (HCP). All scans were T1-weighted MR volumes with 1 mm voxels acquired on a 3-T scanner\(^13\).

**Rater demographics and AFID placements.** The AFID protocol was performed a total of three times on this dataset (2,880 fiducials). Five expert raters were involved with annotations (3DSlicer 4.10.0). Each scan within this dataset was assigned for annotation by three expert raters.

**AFIDs-OASIS30 dataset.** *Subject demographics and imaging protocol.* This subset consists of 30 subjects (age: 58.0 ± 17.9 years; range: 25–91; 17 female) selected from the publicly available Open Access Series

### Table 1. Description of curated anatomical fiducials (AFIDs) and related metadata.

| Number | Anatomical Fiducial | Acronym | Side |
|--------|---------------------|---------|------|
| 1      | anterior commissure  | AC      | Midline |
| 2      | posterior commissure | PC      | Midline |
| 3      | infracollicular sulcus | ICS  | Midline |
| 4      | pontomesencephalic junction | PMJ  | Midline |
| 5      | superior interpeduncular fossa | SIPF | Midline |
| 6,7    | superior lateral mesencephalic sulcus | R/L SLMS | Lateral |
| 8,9    | inferior lateral mesencephalic sulcus | R/L ILMS | Lateral |
| 10     | culmen               | CUL     | Midline |
| 11     | intermammillary sulcus | IMS  | Midline |
| 12,13  | mammillary body      | R/L MB  | Lateral |
| 14     | pineal gland         | PG      | Midline |
| 15,16  | lateral aspect of frontal horn at AC | R/L LVAC | Lateral |
| 17,18  | lateral aspect of frontal horn at PC | R/L LVPC | Lateral |
| 19     | genu of corpus callosum | GENU | Midline |
| 20     | splenium of the corpus callosum | SPLE | Midline |
| 21,22  | anterolateral temporal horn | R/L ALTH | Lateral |
| 23,24  | superior anteromedial temporal horn | R/L SAMTH | Lateral |
| 25,26  | inferior anteromedial temporal horn | R/L IAMTH | Lateral |
| 27,28  | indusium griseum origin | R/L IGO | Lateral |
| 29,30  | ventral occipital horn | R/L VOH | Lateral |
| 31,32  | olfactory sulcal fundus | R/L OSF | Lateral |
| Template or Dataset | Brief Description | Field Strength (T) | Raters (N) | Total number of AFID annotations | References |
|---------------------|-------------------|-------------------|-----------|-------------------------------|------------|
| MN1209bAsym         | A population group template consisting of 152 individuals commonly used in the neuroimaging literature | 1.5          | 8 novices | 8 x 4 (1,024 individual points) | Fonov et al.16 |
| MNIColin27          | A template of a single healthy control subject (N = 1) scanned 27 times and averaged together | 1.5          | 2 experts | 2 x 1 (64 individual points) | Lau et al.17 |
| Agile12v2016        | An ultra-high field template consisting of 12 healthy control averaged subjects created at the Centre for Functional and Metabolic Mapping at Western University | 1.5          | 3 experts | 3 x 1 (128 individual points) | Lau et al.2 |
| BigBrainSym         | Ultra-high resolution histological 3D model of the brain (BigBrain) registered to MNI2009bSym space | N/A; histological | 9 total: 3 expert and 6 novices | 9 total: 3 expert and 6 novices | References |
| MNI2009bSym         | The symmetric version of the MNI2009bAsym template | 1.5          | 9 total: 3 expert and 6 novices | 3 x 2 (3,072 individual points) | Lau et al.2 |
| PD-25               | A multi-contrast MNI template of a Parkinson's disease (PD) cohort | 3            | N/A       | N/A                           | References |
| TemplateFlow        | A centralized resource of open-access templates for neuroimaging studies (tpl-MNI152-Lin, NLin2009cAsym, NLin2009cSym, NLin6Asym, NLin6Sym, tpl-MNI305, tpl-OASIS30ANTS, tpl-fsaverage) | 3+          | 4 total: 1 expert and 3 novices | 4 x 1 (128 individual points) | Citic et al.32 |
| AFIDs-HCP30         | A subset of N = 30 healthy control subject images from the Human Connectome Project (HCP) dataset | 3            | 5 experts | 3 x 1 (2,880 individual points) | Van Essen et al.3 |
| AFIDs-OASIS30       | A subset of N = 30 cognitively intact control subject images from the OASIS-1 database selected to exhibit a wide range of normal anatomical variability | 3            | 9 total: 1 expert and 8 novices | 3 x 1 (2,880 individual points) | Marcus et al.14 |
| LHSCPD              | A set of N = 40 PD patient images acquired at University Hospital (Western University, Canada) | 1.5          | 5 total: 2 expert and 3 novices | 5 x 40 (6,400 individual points) | Abbass et al.1 |
| SNSX                | A set of N = 32 control subject images acquired at the Centre for Functional and Metabolic Mapping at Western University | 7            | 9 total: 3 expert and 6 novices | 3 x 2 (3,072 individual points) | Lau et al.2 |

Table 2. Summary of templates and datasets released, raters, and anatomical fiducial (AFID) protocol performances.

of Imaging Studies (OASIS-1) database14 and imaged at 3-T. The subjects were cognitively intact (Mini-Mental State Examination = 30), and the MRI scans were specifically chosen to be challenging (areas with more complex anatomy and asymmetries) to ensure the protocol could work over a broad spectrum of structural variability. More details on the selected subjects can be found in a previous study2. It is worth noting that this subset of the OASIS-1 dataset includes different subjects from other currently existing subsets in the neuroimaging literature (for instance, the one used in the Mindboggle project15).

Rater demographics and AFID placements. The AFID protocol was performed a total of three times on this dataset (2,880 fiducials). Nine raters (1 expert and 8 novices) were involved with annotations (3DSlicer 4.8.1). Each scan within this dataset was randomly assigned for annotation by one expert and two novice raters.

LHSCPD dataset. Subject demographics and imaging protocol. The London Health Sciences Center Parkinson's disease (LHSCPD) dataset currently consists of 40 subjects diagnosed with Parkinson's Disease (age: 60.2 ± 6.8, range: 38–70; 13 female) with images acquired at University Hospital in London, ON, Canada on a 1.5-T scanner (Signa, General Electric, Milwaukee, Wisconsin, USA). The detailed imaging protocol was described in a previous study7. Due to the heterogenous nature of clinical imaging, MRI scans across patients in this dataset are not always consistent in all three dimensions. Ethics approval was received for the anonymized release of patient scans by the Human Subject Research Ethics Board (HSREB) office at Western University (REB# 109045). Patients signed written consent forms for undergoing clinical imaging and open release of this data.

Rater demographics and AFID placements. The AFID protocol was performed a total of five times on this dataset (6,400 fiducials). Five raters (2 experts and 3 novices) were involved with annotations (3DSlicer 4.10.0). Each scan within this dataset was annotated by all five raters.

SNSX dataset. Subject demographics and imaging protocol. The Stereotactic Neurosurgery (SNSX) dataset currently consists of 32 healthy participants (age: 46.2 ± 13.5 years; range: 20–70 years; 12 female) with images acquired at the Centre for Functional and Metabolic Mapping (Western University, Canada) on a 7-T head-only scanner (Siemens Magnetom; Siemens Healthineers, Erlangen, Germany). An 8-channel parallel transmit/32-receive channel coil was used. The detailed imaging protocol and pre-processing steps were documented in a previous study7. Ethics approval was received for open release of patient scans by the HSREB office at Western University (REB# R-17–156). Patients signed written consent forms for participating and open release of this data.
**Rater demographics and AFID placements.** The AFID protocol was performed a total of three times (3,072 fiducials) on this dataset.

Nine raters (3 experts and 6 novices) were involved with annotations (3DSlicer 4.8.1). Each scan within this dataset was randomly assigned for annotation by one expert rater and two novice raters.

**MNI2009bAsym & Agile12v2016 & MNIColin27 templates.** 

**Template details and imaging protocol.** A group of commonly used public templates were annotated. The MNI2009bAsym is a population group template consisting of 152 individuals (age: 18.5–43.5 years) used commonly in the literature. The images were acquired on a Philips 1.5-T Gyroscan (Best, Netherlands) scanner at the Montreal Neurological Institute.

The Agile12v2016 is an ultra-high field template created locally at our institution. It consists of 12 healthy control subjects (age: 27.6 ± 4.4 years; 6 female). Scans were acquired on a 7-T scanner (Agilent, Santa Clara, California, USA/Siemens, Erlangen, Germany) via a 24-channel transmit-receive head coil array.

The MNIColin27 is a template created from one subject scanned 27 times on a Phillips 1.5-T MR unit.

**Rater demographics and AFID placements.** The AFID protocol was performed a total of 32 times (1,024 fiducials/template). The same raters who annotated the AFIDs-OASIS30 dataset also annotated all the templates via 3DSlicer 4.8.1. Each template was annotated by eight raters four times.

**BigBrainSym & MNI2009bSym & PD-25 templates.** 

**Template details and imaging protocol.** BigBrain is an ultra-high resolution histological 3D model of the brain created using a large-scale microtome to cut a complete paraffin-embedded brain (65-year-old male) coronally at 20 µm thickness. The BigBrainSym template refers to the BigBrain registered to MNI2009bSym space, defined in previous studies. The MNI2009bSym is a symmetric version of the MNI2009bAsym.

The PD-25 template is a multi-contrast MNI template of a PD cohort with 3-T field strength. We used the PD25-T1MPRAGE for the AFID placements.

**Rater demographics and AFID placements.** The AFID protocol was performed a total of two times (64 fiducials/template) by two expert raters via 3DSlicer 4.8.1.

**TemplateFlow templates.** 

**Template details and imaging protocol.** All adult human structural MRI templates that were available on TemplateFlow (see Table 2) at the time of manuscript preparation, which had not been previously annotated, were included (n = 8).

**Rater demographics and AFID placements.** The AFID protocol was performed a total of four times (128 fiducials/template). Four raters (1 expert and 3 novices) annotated each template once via 3DSlicer 4.8.1.

**AFLE calculation for all datasets and templates.** All placements for a given scan and fiducial were averaged to achieve the ground truth fiducial placement per participant or template as shown in Fig. 3a. For datasets, ground truth fiducial placements were computed for each subject in a dataset as shown in Fig. 3b.

To compute the mean AFLE, Euclidean distances from the ground truth fiducial location to each of the individual rater placements were averaged for each fiducial. The result is termed the subject or template mean AFLE per fiducial. This process was independently repeated for all subjects. All subject mean AFLEs were averaged to achieve a dataset mean AFLE per fiducial as shown in Fig. 4a. Finally, the dataset mean AFLE per fiducial was averaged across all fiducials to produce the global dataset mean AFLE. In a similar fashion, global inter-rater AFLE was computed for one subject across fiducials and then averaged across all subjects to produce a global dataset inter-rater AFLE shown in Fig. 4b.

**Data Records**

In total, we release the curated AFID placements and associated imaging of 4 datasets and 14 openly available human brain templates (a total of 19,520 manually placed anatomical landmarks — more than 300 human rater annotation hours).

All datasets have been deposited at DOI-issuing repositories separately (i.e., Zenodo or OpenNeuro) and follow the BIDS directory hierarchy (see Table 3 for licensing and metadata). Each dataset's main directory contains 1) files about licensing and metadata, 2) imaging data directories organized per subject, and 3) a "derivatives" directory which includes AFID coordinates also organized per subject. We release both imaging and AFID annotation data. However, since some of the imaging datasets are protected under Data Usage Agreements (DUAs), the user needs to view and accept terms under DUAs before the imaging data becomes available for download.

For ease of acquiring the data, we merge released datasets and templates in a “super” dataset (https://github.com/afids/afids-data) which serves as a centralized repository for this data descriptor and can be used to install all of the imaging and AFID annotation data we release.

In brief, the centralized repository (i.e., super dataset) has three main directories. 1) data: raw and curated coordinate files for datasets and templates alongside the MRI scans on which coordinates have been applied in BIDS format, 2) notebooks: code used for data curation and quality control, and 3) other: rater demographics, interactive glass brain showing applied coordinates, and a list of our curated brain landmarks.

Raw rater annotations were released. These files have a “rater” label under the “desc” BIDS entity with session also encoded (if applicable). Additionally, curated mean placements (have a “groundtruth” tag in “desc” BIDS entity) are made available. We make individual rater placements available so users have the opportunity to select the subset of rater placements for their intended application. However, we believe that our “ground truth” placements provide the best estimate as to where the true AFIDs are located.
The AFIDs coordinates are described using the Markups comma-separated values file (i.e., *.fcsv), which is generated after the raters save their placements on 3DSlicer. The *.fcsv file contains coordinate data organized in rows for each of the 32 landmarks of interest, with columns describing the AFID and corresponding x, y, and z coordinates in native subject or template space. As for the imaging data, all dataset images used for annotations were BIDS compatible and made available in a compressed NIfTI-1 format (i.e., *.nii.gz).

We release our anatomical landmark annotations under the Attribution 4.0 International (CC BY 4.0) license, available in "DERIVATIVE_DATA_USE_AGREEMENT.txt" file at the level of each dataset. Meanwhile, imaging

---

Fig. 3 Ground truth anatomical fiducial (AFID) placement on templates and datasets. (a,b) show the process of computing the intended AFID placement on a neuroimaging template or dataset, respectively. It is the mean of the rater point cloud at each AFID, referred to as "ground truth" in the text.
data are protected by a DUAs which we make available in the "IMAGING_DATA_USE_AGREEMENT.txt" file, also at the level of each dataset.

**Technical Validation**

As mentioned in the methods, raters typically go through the AFID protocol by referring to the detailed documentation and resources we have made available online (https://afids.github.io/afids-protocol)\(^1\). To ensure the placements we share are accurate and reproducible amongst expert and novice raters, we computed the AFLE metrics and validated that they are generally within 1–2 mm. Table 4 summarizes the AFLE metrics computed for each of the templates and datasets. Across all AFID protocol performances, the global mean AFLE metric was 0.99 ± 0.32 mm.

**Usage Notes**

We recommend loading the shared AFID annotation files (*.fcsv) in 3DSlicer alongside their associated images all of which are in BIDS format for ease of navigating. The local neuroimaging datasets we release here (namely, the LHSCPD and SNSX) will be quality controlled and expanded as more participants are recruited. Additionally, new brain landmarks can be added to future versions of the data descriptor once they have met validation standards set by prior related studies\(^2,\)\(^3\). Access to imaging data is granted after users accept the DUAs. Directions on how to gain access have been added for each of the datasets\(^22–25\). For the AFIDs-HCP dataset, users will need to create an account on the HCP website (https://db.humanconnectome.org) and accept the DUA via the portal which will subsequently provide them with an access key to use when cloning our repository. For the AFIDs-OASIS dataset, users will need to accept the DUA on the website (https://www.oasis-brains.org), but no user credentials are required for data.
Table 4. Summary of anatomical fiducial localization errors (AFLE) and Euclidean distances (ED) used for their calculation across released data.

| Template or Dataset | EDs utilized for AFLE metrics | AFLE ± Error |
|---------------------|------------------------------|--------------|
|                     |                              | Mean         | Inter-rater |
| MN152Lin2009bAsym   |                              | 0.99 ± 1.11  | 1.07 ± 0.46 |
| MNIcolin27          | Mean: 1.024 and inter-rater: 896 | 1.71 ± 2.78  | 1.36 ± 0.88 |
| Agi12v2016          |                              | 1.10 ± 1.59  | 1.14 ± 0.48 |
| BigBrainSym         |                              | 0.63 ± 0.50  | 1.25 ± 1.02 |
| MN152Lin2009bSym    | Mean: 64 and inter-rater: 32 | 0.55 ± 0.26  | 1.09 ± 0.52 |
| PD-25               |                              | 0.42 ± 0.24  | 0.83 ± 0.47 |
| MN152Lin            |                              | 1.07 ± 0.45  | 1.74 ± 0.74 |
| MN152Lin2009cAsym   |                              | 1.03 ± 0.40  | 1.67 ± 0.63 |
| MN152Lin2009cSym    |                              | 1.06 ± 0.47  | 1.67 ± 0.63 |
| MNI152Lin6Asym      | Mean: 128 and inter-rater: 192 | 1.16 ± 0.51  | 1.90 ± 0.86 |
| MNI152Lin6Sym       |                              | 1.08 ± 0.54  | 1.73 ± 0.84 |
| MN30i               |                              | 1.14 ± 0.41  | 1.85 ± 0.52 |
| OASIS30ANTs         |                              | 0.78 ± 0.33  | 1.25 ± 0.51 |
| fsaverage           |                              | 1.00 ± 0.44  | 1.65 ± 0.73 |
| AFIds-HCP30         | Mean: 2,880 and inter-rater: 2,880 | 0.66 ± 0.22  | 1.15 ± 0.70 |
| AFIds-OASIS30       |                              | 0.94 ± 0.73  | 1.58 ± 1.02 |
| LHSCPD              | Mean: 6,400 and inter-rater: 12,800 | 1.57 ± 1.16  | 2.01 ± 1.49 |
| SNSX                | Mean: 3,072 and inter-rater: 3,072 | 0.96 ± 0.33  | 1.64 ± 1.37 |
| Total or Average    | Mean: 19,520 and inter-rater: 25,952 | 0.99 ± 0.32  | 1.48 ± 0.34 |

access. Permission to reshare imaging data from OASIS and HCP has been acquired. We share imaging data from our locally curated datasets, SNSX and LHSCPD, under a Creative Commons license so no further user intervention is needed when acquiring those data.

Users can download all the datasets used in this study, after installing DataLad (https://www.datalad.org/#install) and accepting DUAs, with two lines at the command prompt: “datalad install -r https://github.com/afids/afids-data.git” and then “datalad get -r .” after migrating to the directory containing installed data. Alternatively, users can download individual datasets or images by calling “datalad get -r .” at the directory of interest within the installed dataset. Before images download, the user will be prompted to input relevant information (e.g., user credentials) to ensure the DUAs have been accepted. More granular details on downloading data can be found on our main repository. Although users can acquire the data from the individual dataset directories, we highly recommend using the centralized repository.

Code availability

The code used for technical validation as well as prior AFID studies can be found on the GitHub repository page (https://github.com/afids), including the validator tool (https://github.com/afids/afids-validator).

Received: 15 February 2023; Accepted: 26 June 2023; Published online: 12 July 2023

References

1. Rohlfing, T. Image similarity and tissue overlaps as surrogates for image registration accuracy: widely used but unreliable. *IEEE Trans. Med. Imaging* 31, 153–163 (2012).
2. Lau, J. C. et al. A framework for evaluating correspondence between brain images using anatomical fiducials. *Hum. Brain Mapp.* 40, 4163–4179 (2019).
3. Abbass, M. et al. Application of the anatomical fiducials framework to a clinical dataset of patients with Parkinson’s disease. *Brain Struct. Funct.* 227, 393–405 (2022).
4. Chakravarty, M. M. et al. Comparison of piece-wise linear, linear, and nonlinear atlas-to-patient warping techniques: analysis of the labeling of subcortical nuclei for functional neurosurgical applications. *Hum. Brain Mapp.* 30, 3574–3595 (2009).
5. Chakravarty, M. M., Sadikot, A. F., Germann, J., Bertrand, G. & Collins, D. L. Toward a validation of atlas warping techniques. *Med. Image Anal.* 12, 713–726 (2008).
6. Xiao, Y. et al. An accurate registration of the BigBrain dataset with the MNI PD25 and ICBM152 atlases. *Sci. Data* 6, 210 (2019).
7. Lau, J. C. et al. Direct visualization and characterization of the human zona incerta and surrounding structures. *Hum. Brain Mapp.* 41, 4500–4517 (2020).
8. Kai, J. et al. Source code for: Anatomical Fiducial Placement Validator Tool. Zenodo https://doi.org/10.5281/zenodo.7871820 (2023).
9. Keukens, M. C. et al. Ultra-high 7T MRI of structural age-related changes of the subthalamic nucleus. *J. Neurosci.* 33, 4896–4900 (2013).
10. Fitzpatrick, J. M. & West, J. B. The distribution of target registration error in rigid-body point-based registration. *IEEE Trans. Med. Imaging* 20, 917–927 (2001).
11. Fedorov, A. et al. 3D Slicer as an image computing platform for the Quantitative Imaging Network. *Magn. Reson. Imaging* 30, 1323–1341 (2012).
12. Lau, J. C. et al. Source code for: Anatomical Fiducial Placement Protocol. Zenodo https://doi.org/10.5281/zenodo.7871601 (2023).
13. Van Essen, D. C. et al. The WU-Minn Human Connectome Project: an overview. *Neuroimage* 80, 62–79 (2013).
The authors would like to acknowledge the efforts of all the students and trainees who created, maintained, and participated in neuroanatomy workshops and tutorials which established the foundations of the AFID framework, including staff involved in study recruitment, imaging, and maintaining locally curated datasets. Data was provided in part by Open Access Series of Imaging Studies (Principal Investigators: D. Marcus, R. Buckner, J. Csernansky J. Morris; P50 AG05681, P01 AG03991, P01 AG026276, R01 AG021910, P20 MH071616, U24 RR021382). Data was also provided in part by the Human Connectome Project, WU-Minn Consortium (Principal Investigators: David Van Essen and Kamil Ugurbil; 1U54MH091657) funded by the 16 NIH Institutes and Centers that support the NIH Blueprint for Neuroscience Research; and by the McDonnell Center for Systems Neuroscience at Washington University. A.T. was supported by the Canada Research Chairs program (#950-231964), NSERC Scholarship and Parkinson’s Southwestern Society Graduate Student Scholarship in partnership with the Mitacs Center for Systems Neuroscience at Washington University. A.T. was supported by the Ontario Graduate Institute and Centers that support the NIH Blueprint for Neuroscience Research; and by the McDonnell Consortium (Principal Investigators: David Van Essen and Kamil Ugurbil; 1U54MH091657) funded by the 16 NIH Institutes and Centers that support the NIH Blueprint for Neuroscience Research; and by the McDonnell Center for Systems Neuroscience at Washington University. A.T. was supported by the Ontario Graduate Institute and Centers that support the NIH Blueprint for Neuroscience Research; and by the McDonnell Center for Systems Neuroscience at Washington University. A.T. was supported by the Canada Research Chairs program (#950-231964), NSERC Discovery Grant (RGPIN-2023-05568), and Canada Foundation for Innovation (CFI) John R. Evans Leaders Fund Project (#37427), the Canada First Research Excellence Fund, and Brain Canada. J.L. was supported by research start-up funding from the Department of Clinical Neurological Sciences at Western University and an NSERC Discovery Grant (RGPIN-2023-05558), and Canada Foundation for Innovation (CFI) John R. Evans Leaders Fund Project (#37427), the Canada First Research Excellence Fund, and Brain Canada. J.L. was supported by research start-up funding from the Department of Clinical Neurological Sciences at Western University and an NSERC Discovery Grant (RGPIN-2023-05562).

Author contributions
J.L., A.K., A.T., G.G. conceptualized the idea of the data release and obtained ethics approval. J.L., A.K., A.T., G.G., M.A., J.D., G.G., K.F. overlooked studies related to data release. A.T., J.L., and A.K. contributed to writing and preparing the manuscript. J.L., A.K., A.T., G.G., S.W., T.K., J.K., O.S. and M.A. provided and organized code for analysis and dataset organization. All authors contributed to data curation, material preparation, and reviewed the final version of the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Correspondence and requests for materials should be addressed to J.L.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2023