Collegiate athlete brain data for white matter mapping and network neuroscience

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We describe a dataset of processed data with associated reproducible preprocessing pipeline collected from two collegiate athlete groups and one non-athlete group. The dataset shares minimally processed diffusion-weighted magnetic resonance imaging (dMRI) data, three models of the diffusion signal in the voxel, full-brain tractograms, segmentation of the major white matter tracts as well as structural connectivity matrices. There is currently a paucity of similar datasets openly shared. Furthermore, major challenges are associated with collecting this type of data. The data and derivatives shared here can be used as a reference to study the effects of long-term exposure to collegiate athletics, such as the effects of repetitive head impacts. We use advanced anatomical and dMRI data processing methods publicly available as reproducible web services at brainlife.io.

Background & Summary
Elite athletes are highly motivated individuals with physical and psychological characteristics that make them uniquely fit to compete in a sport. Understanding the impact of sports (positive as well as negative) to the brain is an ongoing and active research need1–26. Advancing understanding of the effects of sports on the brain is currently hindered by a lack of openly shared datasets and the challenge of collecting data from elite athletes. Our datasets address this challenge by providing magnetic resonance imaging data from 33 athletes and 9 matched controls. As of today, this is the first publicly available and openly shared datasets of college-level athletes.

Relatively recently, efforts to generate large-scale neuroimaging datasets to study the effects of sport on the brain have been funded. Take for example, trackTBI27, ENIGMA28, and the CARE consortium29, which are large scale projects involving multiple investigators and institutions. To contribute to the advancement of science, a good percentage of the data in these projects is or is planned-to-be released for sharing. Because of the administrative burden and funding models, these datasets share data with several restrictions spanning all the way from requirements for co-authorship, study participation or research partnership. Our dataset is different, as we collected a smaller sample of subjects (42) from a single institution and shared the dataset openly without administrative overhead. We release this dataset with the intent for it to be used in combination with other datasets, as the size of this dataset alone limits the power for inferences and generalizations.

We share the data utilizing a recently developed unique approach that exploits the free and secure cloud computing platform brainlife.io. The approach integrates, into a single record, both data and reproducible web-services implementing the full processing pipeline30–32. The raw data contains both anatomical (T1 weighted) and diffusion-weighted (dMRI) magnetic resonance imaging data. The processed data contains 14 types...
derivatives across 42 participants, comprising of 84 brain masks (2 per participant), 42 Freesurfer outputs (1 per participant), 168 fiber orientation distribution (FOD) images (4 per participant), 588 diffusion parameter maps (14 per participant), 42 tractograms (1 per participant), 2,562 segmented major tracts (61 per participant), 756 connectivity matrices (18 per participant), and 42 cortex map data types (1 per participant). The total size of the repository is approximately 676.08 GB of data derivatives comprising 1470 datasets.

The processing pipeline implemented to process this dataset utilizes mainstream neuroimaging software libraries including FSL33–35, FreeSurfer36–55, MRTrix 3.056, DIPY57, and connectome_workbench58. The corresponding brainlife.io Apps were developed with a light-weight specification and utilizing modern methods for software containerization making the analyses trackable, reproducible, and reusable on a wide range of computing resources59. The present descriptor describes the repository and pipelines published via brainlife.io mechanisms. These resources will allow the broader research community to explore the effects of long-term sports participation by exploring high-quality preprocessed neuroimaging, replicate previous examinations of the data, and examine a wide variety of hypotheses without the impediment of the aforementioned barriers.

Methods

Data sources. Data is publicly available at https://doi.org/10.25663/brainlife.pub.14.

Neuromaging data sources. Data were collected at the Indiana University Imaging Research Facility (IRF, https://www.indiana.edu/~irf/). Data collection was approved by the Indiana University Institutional Review Board.

Study participants. A total of fifty-one male participants participated in the study. Twenty-one participants were 4th and 5th-year varsity Indiana University (IU) football “starters” (age 21.1 ± 1.5 years). This number accounts for approximately 60% of the total IU football team active players matching our criteria. Potential participants were excluded if they reported a diagnosed concussion within 6 months of the beginning of the study. The only other exclusion criteria was for safety in the MRI environment. One football player did not complete the study, and three football players did not complete the diffusion MRI (dMRI) scans. This left 17 usable datasets from the Football group.

Following scanning, football players received a socioeconomic status survey gathering information regarding estimated family income and the area in which they were raised (i.e. urban, small town, suburbs). Nineteen members of the IU cross-country running team (age 20.2 ± 2.5 years) were included as a non-collision sports group and were matched to the football players based on age and experience level. Three of the players’ anatomical (T1w) images were unusable and thus their data was not included. This left us with 16 eligible members. Our access to the socioeconomic status of non-athlete undergraduates was limited to Psychology and Neuroscience undergraduates who had filled out the socioeconomic survey in an IRB approved subject pool. Eleven controls (non-athletes; age 19.9 ± 3) participants were selected from the limited pool matched on age, sex, estimated family income. None of these individuals were athletes, no additional information on the exercise habits was collected for this group. Two of the controls’ diffusion data contained artifacts that were beyond correction with our processing protocol and thus their data were not included, leaving nine usable datasets. Overall, the released dataset contains usable data from 17 football players, 16 cross-country runners, and 9 non-athletes for analysis (N = 42). In regards to concussion history, two football players had been diagnosed with a concussion approximately 3 years before the study and one player had been diagnosed approximately 2 years before the study. There was no history of concussion in the cross-country runners while at IU. No information was collected regarding the concussion history of the participants before their arrival at IU, however. Although we did not have this information available, we can estimate around 7.25% of football players have been diagnosed with a concussion prior to college given estimates from the literature (Dompier et al., 2015). Participants gave informed written consent that was approved by the Indiana University Institutional Review Board. All participants were recruited through flyers handed out by the athletic trainers of each team or posted around campus. Participants were compensated for participation with a cash payment.

The data detailed here was collected as part of a larger study, which included task-related functional MRI (fMRI) data. The results from the fMRI portion of the study is described in60. Due to this, and the limitations of gathering information via subject pool, focus was placed on collecting neuroimaging data. No other cognitive or behavioral data was collected on the participants, including handedness, IQ, GPAs, or diagnoses of any neuro-cognitive or development disorders.

Neuroimaging parameters. Participants were imaged using a 3-Tesla TIM Trio scanner located in the Imaging Research Facility at Indiana University. A 12-channel head coil was used as the 32-channel coil did not fit the heads of our larger subjects. Diffusion-weighted magnetic resonance imaging (dMRI) data were collected with two phase-encoding schemes, i.e. anterior-posterior (AP) and posterior-anterior (PA). The following parameters were used for the dMRI pulse sequence: TR/TE = 4930/99.6 ms, iPAT acceleration factor = 2; voxel size = 2x2x2 mm isotropic, 143 diffusion-weighting directions. As student athletes have demanding schedules, emphasis was given to minimizing time of participation when designing the study. Because of this, and the additional fMRI component of the larger study, only two diffusion gradient strengths (b-values) were collected. Sixty-four diffusion gradient directions were collected for each gradient strength, b = 1000 s/mm² and b = 2000 s/mm², respectively. Fifteen non-weighted images were also acquired (b = 0). On T1-weighted (T1w) anatomical image was acquired for each participant using the following sequence: TR/TE = 1800/2.67 ms, TI = 900 ms, flip angle = 9°, bandwidth = 150 Hz/pixel, 160 sagittal slices, FOV = 256 mm, matrix = 256 × 256, slice thickness = 1 mm, resulting in 1 mm isotropic voxels.

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Eleven controls (non-athletes; age 19.9 ± 3) participants were selected from the limited pool matched on age, sex, estimated family income. None of these individuals were athletes, no additional information on the exercise habits was collected for this group. Two of the controls’ diffusion data contained artifacts that were beyond correction with our processing protocol and thus their data were not included, leaving nine usable datasets. Overall, the released dataset contains usable data from 17 football players, 16 cross-country runners, and 9 non-athletes for analysis (N = 42). In regards to concussion history, two football players had been diagnosed with a concussion approximately 3 years before the study and one player had been diagnosed approximately 2 years before the study. There was no history of concussion in the cross-country runners while at IU. No information was collected regarding the concussion history of the participants before their arrival at IU, however. Although we did not have this information available, we can estimate around 7.25% of football players have been diagnosed with a concussion prior to college given estimates from the literature (Dompier et al., 2015). Participants gave informed written consent that was approved by the Indiana University Institutional Review Board. All participants were recruited through flyers handed out by the athletic trainers of each team or posted around campus. Participants were compensated for participation with a cash payment.
Anatomical (T1w) preprocessing. Raw anatomical (T1w) images were preprocessed using the fsl_anat functionality provided by the FMRIB Software Library (FSL)33–35 implemented as brainlife.app.273. In brief, the raw anatomical (T1w) images were cropped and reoriented to match the orientation of the MN152 template. Then, the cropped and reoriented images were linearly and non-linearly aligned to the MN152 0.8 mm template using flirt and fslirt respectively61–63. The linearly aligned images will hereafter be referred to as the ‘acpc aligned’ anatomical (T1w) images. The warps generated from the non-linear alignment were subsequently used for mapping diffusion metrics to the cortex64. Following alignment, the ‘acpc aligned’ anatomical (T1w) images were processed via Freesurfer’s recon-all function to generate pial (i.e. cortical) and white matter surfaces and to parcellate the brain into known anatomical atlases36–39 implemented as brainlife.app.0. The Destrieux (aparc.a2009s) atlas was used for subsequent white matter tract segmentation and for mapping of diffusion metrics to the cortical surface65. For network generation, the multi-modal 180 cortical node parcellation66 was mapped to the Freesurfer segmentation of each participant implemented as brainlife.app.239. Finally, the ‘acpc aligned’ anatomical (T1w) image was segmented into different tissue-types using MRTrix 3.0’s 5tigen functionality35,56 implemented as brainlife.app.239. The gray- and white-matter interface mask was subsequently used as a seed mask for white matter tractography.

Preprocessed anatomical images, and their derivatives, were visually QA’d for common artifacts by BC, RS, and FP. Specifically, the ‘acpc aligned’ anatomical (T1w) images were examined for proper alignment and tissue-contrast. Freesurfer surfaces and parcellations were examined for common surface artifacts and improper voxel identification in parcellations. Any identified issues were manually corrected in Freesurfer and reuploaded before further analysis. The gray- and white-matter interface mask was visually examined for proper separation of the gray- and white-matter in the ‘acpc aligned’ anatomical (T1w) issue.

Diffusion (dMRI) preprocessing. Raw dMRI images were first reoriented to match the orientation of the MN152 template using the fsreorient2std command provided by FSL. The gradients orientation were then checked using MRTrix 3.0’s dwigradcheck functionality67. Following gradient checking, PCA denoising was performed using MRTrix 3.0’s dwidenoise functionality68. This was followed by Gibbs deringing using MRTrix 3.0’s mrdqgsbbs functionality69. The pseudo-facing distortions corresponding to each phase encoding direction (i.e. PA and AP) were then combined into a single corrected image in a method similar to the one described in Andersson and colleagues (2003)70,71 (i.e. top-up command) as provided by FSL72,73. Eddy-current and motion correction was then applied via the eddy_cia8.0 with replacement of outlier slices (i.e. repol) command provided by FSL74–77. Following this, dMRI images were debiased using ANTS’s n4 functionality78 and the background noise not associated with the diffusion signal was cleaned using MRTrix 3.0’s dwidenoise functionality79. Finally, the preprocessed dMRI images were registered to the ‘acpc aligned’ anatomical (T1w) image using FSL’s epi_reg functionality80–83 and resliced to 1 mm isotropic voxels. The preceding steps were implemented as brainlife.app.68. In sum, the dMRI data was interpolated 4 times: 1) following top-up, 2) following eddy, 3) following epi_reg, and 4) during reslicing. These steps were implemented as brainlife.app.68. Brainmasks of the preprocessed, acpc-aligned dMRI images were then used for subsequent modelling and tractography using FSL’s bet2 functionality84 implemented as brainlife.app.163.

Quality control was estimated by calculating the Signal to Noise Ratio (SNR) of the diffusing data. To quantify the SNR in the preprocessed, acpc-aligned dMRI data, the workflow provided by Dipy to map SNR in the corpus callosum was used77,77,78 implemented as brainlife.app.120. SNR values reported are generated from this step.

White matter microstructure modeling (DTI). In order to investigate advanced microstructural properties of white matter, the diffusion tensor (DTI) model was fit to the preprocessed, acpc-aligned dMRI data using FSL’s dtifit functionality implemented as brainlife.app.292. For white matter tract profiles, the default parameters of dtifit were used and the b = 1000 shell was chosen for fitting. However, for mapping of the DTI measures to the cortex, both the b = 1000 and b = 2000 shells were used, kurtosis was calculated, and the sum of squared errors was outputted following the parameters used in Fukutomi et al.85.

White matter microstructure modeling (NODDI). In order to investigate advanced microstructural properties of white matter, the Neurite Orientation Dispersion and Density Imaging (NODDI)79 model was fit to the multi-shell (i.e. b = 1000, 2000 s/mm²) acpc-aligned dMRI data via the Accelerated Microstructure Imaging via Convex Optimization (AMICO; https://github.com/daducci/AMICO85) toolbox implemented as brainlife.app.365. The AMICO toolbox was used in order to significantly speed-up the time necessary to fit the NODDI model by reformulating the NODDI model as a linear system, without sacrificing accuracy86. For major white matter tract analysis, the isotropic diffusivity parameter (diso) was set to 3.0 × 10⁻³ m²/s (the rate of unhindered diffusion of water) while the intrinsic free diffusivity parameter (dI) was set to 1.7 × 10⁻³ mm²/s. For cortical white matter parcel analyses, the isotropic diffusivity parameter was also set to 3.0 × 10⁻³ mm²/s while the intrinsic free diffusivity parameter was set to 1.1 × 10⁻³ mm²/s, which is the optimal value of diffusivity found by Fukutomi et al.87.

White matter microstructure modeling (CSD). The CSD model was fit to the preprocessed multi-shell data utilizing MRTrix 3.0 dwi2fod function across 4 maximum spherical harmonic orders (i.e. Lmax’s) parameters (2,4,6,8) implemented as brainlife.app.239. Lmax’s 6 and 8 were chosen for subsequent white matter tractography, however, all Lmax’s are included in the released data.

White matter microstructure modeling (Tractography). Anatomically-constrained probabilistic tractography (ACT)88 via MRTrix3’s tckgen functionality implemented as brainlife.app.297 was used to generate tractograms on preprocessed multi-shell dMRI data for each participant. A total of 1.5 million was tracked over both Lmax6 and Lmax8. The two tractograms were then combined to create a single tractogram of 3 million streamlines
via Vistasoft functionality implemented as brainlife.app.305. The step-size was set to 0.2 mm for both lmax6 and lmax8. The minimum length of streamlines was set to 25 mm, and the maximum length was set to 250 mm. A maximum angle of curvature of 35° was set. The merged tractogram of 3 million streamlines was then used for subsequent white matter tract segmentation and network generation.

**White matter microstructure modeling (Segmentation & Cleaning).** 61 human white matter tracts were segmented using a custom version of the white matter query language67 implemented as brainlife.app.188. These tracts include the following: L/R arcuate, aslant, corticospinal tract (CST), contralateral anterior frontal cerebellar tracts, contralateral motor cerebellar tracts, inferior fronto-occipital fasciculus (IFOF), inferior longitudinal fasciculus (ILF), middle longitudinal fasciculus-angular gyrus (MDLF-ang) and superior parietal lobule (MDLF-spl), motor cerebellar tracts, occipital cerebellar tracts, superior longitudinal fasciculus components 1 & 2 and 3 (SLF-1 & 2, SLF-3), temporal-parietal connection, thalamic cerebellar tracts, uncinate, vertical occipital fasciculus (VOF), Baum's and Meyer's loops, cingulum, frontal thalamic tracts, motor thalamic tracts, parietal arcuate (pArc), parietal thalamic tracts, spinothalamic tracts, and temporal thalamic tracts. The callosal tracts, including anterior frontal, forceps major, forceps minor, middle frontal, and parietal corpus callosum, are also included.

Following tract segmentation, outlier streamlines were removed using mba's mbaComputeFibersOutliers functionality68 implemented as brainlife.app.195. For each tract, the spatial ‘core’ representation of the tract was computed by averaging the streamline coordinates across all streamlines in a tract. Streamlines were removed if their length was 4 standard deviations from the length of the ‘core’ representation and/or were located 4 standard deviations away from the ‘core’ representation of the tract. The cleaned segmentations were then used for all subsequent analyses.

**White matter microstructure modeling (Tract profiles).** Tract profiles37 for each DTI parameter estimate (i.e. AD, FA, MD, RD) and NODDI parameter estimate (i.e. NDI, ODI, ISOVF) were generated by estimating the “core” representation of each tract, resampling and segmenting each streamline into 200 equally-spaced nodes, applying a gaussian weight to each streamline based on the distance away from the “core”, and obtaining the weighted average metric at each node. This was performed using MATLAB code utilizing the Compute_FA_AlongFG command provided by Vistasoft (https://github.com/vistasoft/vistasoft) implemented as brainlife.app.361.

**White matter network modeling (Network generation).** Structural networks were generated using the multi-modal 180 cortical node atlas and the merged tractograms for each participant using MRTRix3’s tck2connectome38 and tck2csift29 functionality implemented as brainlife.app.394. SIFT2 was used to generate a cross-sectional area weight value for each streamline in order to accurately reflect density. Connectomes were then generated by computing the number of streamlines intersecting each ROI pairing in the 180 cortical node parcellation. Multiple adjacency matrices were generated, including: count, density (i.e. count divided by the node volume of the ROI pairs), length, length density (i.e. length divided by the volume of the ROI pairs), and average and average density AD, FA, MD, RD, NDI, ODI, and ISOVF. Density matrices were generated using the -invmodevol option69. For non-count measures (length, AD, FA, MD, RD, NDI, ODI, ISOVF), the average measure across all streamlines connecting and ROI pair was computed using MRTRix3’s tck2scale functionality using the -precise option70 and the -scale_file option in tck2connectome. These matrices can be thought of as the “average measure” adjacency matrices. Before figure generation, nodes in which less than 50% of the participants had a connection were removed.

**Cortical white matter microstructure modeling (Cortex mapping).** DTI and NODDI measures were mapped to each participant’s cortical white matter parcels following methods found in Fukutomi and colleagues44 using functions provided by Connectome Workbench66 implemented as brainlife.app.379. First, mid-thickness surfaces between the cortical pial surface and white matter surface provided by Freesurfer segmentation were computed using the wb_command -surface-cortex-layer function provided by Workbench command. A Gaussian smoothing kernel (FWHM = ~4 mm, σ = 5/3 mm) was applied along the axis normal to the surface, and DTI and NODDI measures were mapped using the wb_command -volume-to-surface-mapping function. Freesurfer was used to map the average NODDI parameter estimates to subcortical white matter parcels.

**Demographics, brain size, body size.** We performed multiple one-way ANOVAS between the groups utilizing the python repository statsmodels’ ols function to test for differences in the following: age, body weight, SMI, average gray-matter cortical thickness, total brain volume, gray-matter cortical volume, and white matter volume. Bonferroni multiple comparisons correction was performed, and all reported p-values were significantly below a corrected p < 0.0083 (0.05/6 measures).

**Data visualization.** A majority of the images generated for this descriptor were generated using a number of brainlife.io applications utilizing functionality from FSL and DIPY. A list of these Apps include: Generate images of NODDI/DTI, Generate figures of whole-brain tractogram (TCK), Generate images of mask overlaid on DWI, Generate an image of ODF, Generate images of DWI overlaid on T1, Generate images of tissue type masks, Generate images of T1/DWI, and Plot response function. The other images were generated using brainlife.io’s visualization functionality.
Data Records

The data outputs on brainlife.io are organized using https://brainlife.io/datatypes. These DataTypes allow applications to exchange and archive data. Data outputs can be conveniently downloaded from brainlife.io using the BIDS standard92. The data outputs described below can be downloaded at https://doi.org/10.25663/brainlife.pub.1493. The standard does not yet provide a specification for processed dMRI, tractograms, white matter tracts, and connectivity matrices. The brainlife.io platform will be updated as soon as the BIDS standard fully describes a specification for the models and tractography, tractometry, and network data. For the time being the specification follows the work previous work30. We also provide two additional online tables reporting input and output specimens as requested by the Scientific Data guidelines (see Online-only Table 1 and Supplemental Table 1).

**T1-weighted anatomical.** T1w image preprocessed and linearly- and nonlinearly- aligned to the MNI152 0.8 MM template using FMRIB Software Library (FSL)'s fsl_anat functionality.

upload/sub-{}/anat/
  sub-{}_tag-preprocessed_tag-acpc-aligned_desc-{}_T1w.json
  sub-{}_tag-preprocessed_tag-acpc-aligned_desc-{}_T1w.nii.gz
  sub-{}_tag-preprocessed_tag-standard_desc-{}_T1w.json
  sub-{}_tag-preprocessed_tag-standard_desc-{}_T1w.nii.gz

**Non-Linear image warping.** Warp files describing the non-linear alignment between the raw anatomical (T1w) image and the template image generated from fsl_anat. No current BIDS structure exist, brainlife.io structure is as follows:

dt-neuro-warp.tag-preprocessed.tag-standard.id-{}
  _info.json
  inverse-warp.nii.gz
  warp.nii.gz

**Freesurfer.** Freesurfer output directory containing all derivatives generated during Freesurfer's recon-all. No existing BIDS structure, brainlife.io structure is as follows:

dt-neuro-freesurfer.tag-preprocessed.tag-acpc_aligned.tag-hippocampal.
tag-thalamic_nuclei.id-{}
  _info.json
  output

**Multi-atlas Transfer Tool (MaTT).** The surface and volumated mapping files of the 180 node multimodal parcellation to individual participant surfaces. No existing BIDS structure, brainlife.io structure is as follows:

dt-neuro-parcellation-surface.id-{}
  _info.json
  lh.parc.annot.gii
  lh.parc.white.gii
  rh.parc.pial.gii
  key.txt
  lh.parc.inflated.gii
  rh.parc.annot.gii
  rh.parc.inflated.gii
  key.txt
  lh.parc.pial.gii
  rh.parc.pial.gii
  lh.parc.inflated.gii
  rh.parc.inflated.gi
  dt-neuro-parcellation-volume.tag-SupraTentorial.id-{}
  _info.json
  key.txt
  label.json
  parc.nii.gz

**Tissue-type masks.** The 5-tissue type images (GM, WM, CSF, GMWMI) used for tracking. No existing BIDS structure, brainlife.io structures is as follows:

dt-neuro-mask.tag-5tt_masks.id-{}
  _info.json
  csf.nii.gz
  gm.nii.gz
  wm.nii.gz
  mask.nii.gz

**Diffusion-weighted imaging (dMRI).** The final preprocessed dMRI data used for all further modeling and analyses following FSL Topup & Eddy - CUDA and MRTRix3 preproc.
The final dMRI brain mask used for all modeling and analyses. No existing BIDS structure, brainlife.io structures are as follows:

dt-neuro-mask.tag-brain.id-{}
  _info.json
  mask.nii.gz

Neurite Orientation Dispersion Density Imaging (NODDI). The neurite density, orientation dispersion, and isotropic volume fraction maps generated from NODDI AMICO. No existing BIDS structure, brainlife.io structure is as follows:

dt-neuro-noddi.id-{}
  _info.json
  dir.nii.gz
  ndi.nii.gz
  odi.nii.gz
  isovf.nii.gz

Diffusion Tensor Imaging (DTI). The fractional anisotropy, mean diffusivity, axial diffusivity, and radial diffusivity maps generated from DTIFIT.

Constrained Spherical Deconvolution (CSD). CSD models fit across $L_{\text{max}} = 2, 4, 6,$ and 8 using Fit Constrained Spherical Deconvolution Model For Tracking. No existing BIDS structure, brainlife.io structure is as follows:

dt-neuro-csd.tag-preprocessed.id-{}
  _info.json
  lmax2.nii.gz
  lmax4.nii.gz
  lmax6.nii.gz
  lmax8.nii.gz
  response.txt

Tractograms. The merged tractograms across $L_{\text{max}} = 6$ and $L_{\text{max}} = 8$, totaling 3 million streamlines generated from the Anatomically-constrained tractography (ACT) app.

White Matter Anatomy (wma) Segmentation. Major track segmentation generated from White Matter Anatomy Segmentation. No existing BIDS structure, brainlife.io structure is as follows:

dt-neuro-wmc.id-{}
  _info.json
  classification.mat
  surfaces
  tracts

Segmentation cleaned. The cleaned tracks from Remove Fiber Outliers. No existing BIDS structure, brainlife.io structure is as follows:

dt-neuro-wmc.tag-cleaned.id-{}
  _info.json
  classification.mat
  output_fibercounts.txt
  tracts
**Tract profiles.** Mapping of DTI and NODDI metrics along the core of the segmented white matter tracts using the Tract Analysis Profiles app. No existing BIDS structure, brainlife.io structure is as follows:

```
dt-neuro-tractprofile.id-{}  
_info.json  
profiles
```

**Network generation.** Network adjacency matrices were generated using the Structural Connectome MRTRix3 (SCMRT) (SIFT2) app. No existing BIDS structure, brainlife.io structure as follows:

```
dt-raw.tag-networkmatrices.id-{}  
_info.json  
output
```

**Cortical DTI and NODDI mapping.** Surfaces and DTI and NODDI measure files mapped to the surface generated from Cortical Tissue Mapping. No existing BIDS structure, brainlife.io structure is as follows:

```
dt-neuro-cortexmap.id-{}  
_info.json  
cortexmap  
Func  
Label  
surf
```

**Technical Validation**

In this section, we provide a qualitative evaluation of the data derivatives made available on brainlife.io. We provide qualitative analysis of the preprocessing of the anatomical (T1w) image, including the seed mask, and pial and white matter surfaces generated from Freesurfer. Qualitative images of the dMRI preprocessing, dMRI modeling (CSD and NODDI), dMRI tractography, network generation, and mapping of diffusion measures to the cortical surface are also provided. We further provide a quantitative analysis of the SNR of the dMRI data following preprocessing.

**Anatomical (T1w) preprocessing.** Anatomical (T1w) images were linearly aligned to the MNI152 0.8 mm template and further segmented into gray-matter, white-matter, CSF, and gray- and white-matter interface masks using brainlife.app.273. See Methods: Anatomical (T1w) preprocessing for more details. Each participant’s aligned anatomical images (T1w), and all derivatives generated from the aligned images, are provided. Figure 1a exemplifies the quality of the linear alignment obtained with brainlife.app.300 in representative participants from each athlete group (i.e. Football: top, Cross-Country: middle, and Non-Athlete: bottom). The gray- and white-matter interface mask (1b) and white matter boundary (1c) are overlaid on the ‘acpc-aligned’ anatomical (T1w) image to further provide quality assurance. These images were generated with brainlife.app.312.

Following alignment and segmentation, Freesurfer was used to generate cortical and white matter surfaces, along with the Destrieux Atlas parcellation using brainlife.app.0. Figure 2a demonstrates the quality of the surface generation representative participants from each athlete group (i.e. Football: top, Cross-Country: middle, and Non-Athlete: bottom). Images of the Destrieux (aparc.a2009s) atlas on the pial surface, along with images of the pial and white matter surface outlines overlaid on the ‘acpc-aligned’ anatomical (T1w) image, are provided as a means of quality assurance. Figure 2b illustrates the mapping of the 180 node multimodal atlas to representative participants from each group mapped using brainlife.app.23. These images were generated using brainlife.io’s Freeview and Connectome Workbench viewers.

**Diffusion (dMRI) preprocessing.** Raw dMRI images were corrected for Gibbs ringing, susceptibility-weighting, eddy currents, motion, biasing, and Rician background noise using a combination of methods. Following preprocessing, the dMRI images were aligned to the ‘acpc-aligned’ anatomical (T1w) image. See Methods: Diffusion (dMRI) preprocessing for more details. The preprocessing was performed using brainlife.app.68. Following preprocessing, the signal-to-noise ratio (SNR) was computed for each subject in the non-diffusion weighted volumes (i.e. b = 0) and the diffusion-weighted volumes (i.e. b = 1000,2000) separately as a means for quality assurance. The SNR was computed using brainlife.app.120.

Figure 3a demonstrates the quality of alignment of the dMRI and ‘acpc-aligned’ anatomical (T1w) image from representative participants from each group. The fractional anisotropy (FA) map (see Methods: White matter microstructure: DTI for more details on DTI fitting) from each subject is overlaid in red-yellow on the ‘acpc-aligned’ anatomical (T1w) images. Overall, the alignments of the dMRI and the anatomical image are anatomically-sound. The fully preprocessed dMRI images from each participant, along with their corrected b-vectors and b-values, are provided. The images were generated using brainlife.app.309. Figure 3b documents the non-diffusion weighted and diffusion-weighted SNRs for each participant. The average SNR for Football players (orange) following preprocessing was 28.354 (±5.772 SD) for non-diffusion weighted volumes. This was slightly lower than the SNR for Cross-country runners (pink) with an average SNR in the non-diffusion weighted volumes of 34.944 (±4.594 SD). Non-athletes overall had the lowest average SNR in the non-diffusion weighted volumes (23.002 ± 7.784 SD).

**White matter microstructure modeling: DTI and NODDI.** Following preprocessing, models of microstructure were fit to the dMRI images. Specifically, the diffusion tensor (DTI) and neurite orientation dispersion
density imaging (NODDI) models were fit to the $b = 1000$ and $b = 1000, 2000$ shells respectively. See Methods: White matter microstructural modeling: DTI & NODDI for more details. DTI was fit using brainlife.app.292, while NODDI was fit using brainlife.app.365. The DTI and NODDI maps for each participant are provided. Figure 4 demonstrates the quality of fit of both the DTI and NODDI models on representative participants from each group. Specifically, mid-axial slices of the fractional anisotropy (FA), mean diffusivity (MD), axial diffusivity (AD), and radial diffusivity (RD) from the DTI model, and the orientation dispersion and neurite density indices (ODI, NDI) and the isotropic volume fraction (ISOVF) from the NODDI model are presented. There is a high anatomical correspondence between the measures and known anatomical properties. For example, the ventricles across all three participants are saturated in the MD maps, as water moves maximally isotropically. In the white matter, FA and NDI are highest in the highest concentrations of white matter, while ODI is lowest. The images were generated using brainlife.app.302 and brainlife.app.367.

White matter microstructural modeling: CSD. In order to map white matter macrostructure via white matter tractography, the constrained spherical deconvolution model (CSD) was fit to each participant across 4 maximum spherical harmonic orders (i.e. $L_{\text{max}}$: 2, 4, 6 and 8). $L_{\text{max}} = 6, 8$ were chosen for tracking. See Methods: White matter microstructural modeling (CSD) for more details. The CSD was fit using brainlife.app.238. Each participant’s CSD fits across all four $L_{\text{max}}$’s are provided. Figure 5 demonstrates the quality of fit of the CSD model on representative participants from each group using $L_{\text{max}} = 8$. In the left column, the response function generated is mapped to a sphere, while the right column corresponds to the fiber orientation distribution function (fODF). The response functions demonstrate a quality fit due to the relatively flat shape and sharp folding in the center. In the fODF maps, clear anatomy is distinguished in regions of the highest white matter concentration. Images were generated using brainlife.app.311 and brainlife.app.317.

Fig. 1 Anatomical (T1w) preprocessing: Alignment and segmentation. Representative mid-plane images from each group of the preprocessed anatomical (T1w) image. (a) Sagittal slice of MNI152 (ACPC) aligned T1 used for all subsequent preprocessing, (b) Axial slice with gray - and white-matter interface outlined (red) used as a seed mask for tractography, and (c) Coronal slice with white-matter boundary outlined (red). Images were generated using brainlife.app.300 and brainlife.app.312.
Fig. 2  Anatomical (T1w) preprocessing: Freesurfer and 180 node multimodal atlas mapping (a) Representative images from each group of the Freesurfer outputs: pial (red) and white (blue) matter surfaces, and the aparc.a2009s + aseg (i.e. Destrieux) parcellation. Images were generated using brainlife.io’s Freeview viewer. (b) Representative images from each group of the 180-node multimodal (hcp-mmp) atlas mapped to an inflated representation of the cortical surface. Images were generated using brainlife.io’s Connectome Workbench viewer.

Fig. 3  Qualitative and quantitative quality-assurance measures following dMRI preprocessing. (a) Representative mid-axial images of the fractional anisotropy (FA) (red-yellow) image overlaid on the ‘acpc aligned’ anatomical (T1w) image. Images were generated using brainlife.app.309. (b) The signal-to-noise ratio for each participant from each group (football: orange, cross-country: pink, non-athlete: blue) following dMRI preprocessing. The average SNR was computed in the Corpus Callosum across the b = 0 shells (crosses) and b = 1000,2000 shells (circles). Standard deviation across the non-b = 0 shells are plotted as error bars for each participant.
White matter microstructure modeling: anatomically-constrained tractography. Following the fitting of the CSD model to each participant, anatomically-constrained tractography (ACT) was performed on $L_{\text{max}} = 6,8$ to generate whole-brain tractograms containing 3 million streamlines. These tractograms were used for subsequent segmentation and network generation. See Methods: White matter microstructure modeling (Tractography) for more details. The tractograms were generated using brainlife.app.297, and merged using brainlife.app.305. Each participant’s whole-brain tractogram containing 3 million streamlines per tractogram is provided. Figure 6a demonstrates the quality of the tractography in representative participants from each group. The whole-brain tractogram of each participant shows high densities of streamlines filling the entire brain volume, as expected. The images were generated using brainlife.app.310.

White matter microstructure modeling: Segmentation and cleaning. Following tractography, each participant’s whole-brain tractogram was segmented using a recently published methodology using anatomical definitions of common white matter tracts. In brief, this segmentation classifies streamlines as belonging to a particular tract based on their cortical terminations and known shape characteristics. See Methods: White matter microstructure modeling (Segmentation & Cleaning) for more details. Tract segmentation was performed using brainlife.app.188. Following segmentation, each tract was cleaned by removing outlier streamlines using brainlife.app.195. Each participant’s segmentation, both cleaned and uncleaned, is provided. Figure 6b provides representative white matter tract segmentation from participants from each group. From a qualitative perspective, each segmentation fills the whole-brain volume and contains a relatively high density of streamlines per tract. Each of the tracts is listed on the right. The images were generated using the brainlife.io tract segmentation viewer.
White matter microstructure modeling: Tract profiles. Following white matter tract segmentation and cleaning, tract profilometry was performed for each participant and each tract. In brief, a central representation (i.e. ‘core’) of each tract was computed by weighted-average of the X,Y, and Z coordinates of all the streamlines in the tract. Streamlines were then resampled to 200 equally spaced nodes, and the average microstructural measures (DTI, NODDI) were computed at each node. The first and last ten nodes were removed, and
then the profiles for each tract were averaged across each group. See **Methods: White matter microstructure modeling (Tract profiles)** for more details. Tract profiles were computed using brainlife.app.361. Tract profilometry data for all participants and tracts are provided. Figure 7a provides a representative Right ILF from a Football player. Figure 7b illustrates the centralized core of the ILF and the ODI values mapped along the tract. Figure 7c provides the group average ODI and FA tract profiles for the Right ILF. These profiles document the ability of this methodology for identifying group differences along a tract. Images were generated using the Matlab Brain Anatomy toolbox https://github.com/francopestilli/mba86 scripts available at https://github.com/bacaron/athlete-brain-study.

**White matter microstructure modeling: Network adjacency matrix generation.** The whole-brain tractograms from each participant were used to generate structural connectomes. Specifically, measures of streamline count and density are computed between each node in the multimodal 180 cortical node parcellation and network matrices are generated94,95. See **Methods: White matter microstructure modeling (Network generation)** for more details. Network adjacency matrices were generated using brainlife.app.394. Figure 8 demonstrates group average connectivity matrices using log streamline count, log density, and average FA across the streamlines connecting nodes from each group and the total dataset. Images were generated using the `imagesc` function in MATLAB.

**Cortical white matter microstructure mapping.** Diffusion-based measures of microstructure were also mapped to a surface representation of the midthickness (i.e., the average coordinates between pial and white matter boundary surface64), hereafter simply referred to as 'cortical.' For more details on the differences between the two mappings, see **Methods: White matter microstructural modeling: DTI & NODDI.** The DTI and NODDI
maps for each participant for the cortical white matter mapping analyses are provided as well. These measures were then mapped to the cortex following the procedures described in Fukutomi et al. 2018. See Methods: Cortical white matter microstructure mapping for more details. Diffusion measures were mapped using brainlife.app. Figure 9 demonstrates the quality of fit of DTI and NODDI measures on the cortical surface. Specifically, the FA and ODI maps mapped to a representative participant’s cortical surface. Anatomic landmarks, including higher FA and lower ODI in motor and somatosensory cortices, are consistent across participants and map well to the results presented in Fukutomi et al. 2018. These images were generated using brainlife.io’s Connectome Workbench viewer.

Mass and brain size. To further reduce the burden to full understanding of the dataset provided, we examined the potential differences between the groups in terms of mass and brain size. We collected data from each participant provided by the Freesurfer segmentations regarding total brain volume, cortical volume, white matter volume, and cortical thickness and computed one-way ANOVAs between our groups. We identified a significant difference in body mass between Football players and the other two groups (F(2,39), p < 0.0001; Fig. 10e). However, we did not observe any significant effects of group on brain volume (Fig. 10a), cortical volume (Fig. 10b), white matter volume (Fig. 10c), or cortical thickness (Fig. 10d).

Usage Notes
The data are publicly available on brainlife.io using the following https://doi.org/10.25663/brainlife.pub.1493. Data can be accessed for visualization and download without requiring a login. The data can be browsed directly using any major web-browser.

Data files can also be downloaded, and some can be organized into BIDS standard. The data derivatives are stored in numerous formats, including NIFTI, TCK, GIFTI, and .mat. Access to the published data is currently supported via (i) web interface and (ii) Command Line Interface (CLI).

The brainlife.io CLI can be installed on most Unix/Linux systems using the following command:

```bash
npm install brainlife.io -g
```

The CLI can be used to query and download partial or full datasets. The following example shows the CLI command to download all T1w datasets from a subject in the publication data:

```bash
bl pub query # this will return the publication IDs
```
The following command downloads the data in the entire project (from Release 2) into BIDS format:

```
bl bids download --pub <insert pub id> --subject 1_001 --datatype \nneuro/anat/t1w–tag “fsl_anat”
```

The following command downloads the data in the entire project (from Release 2) into BIDS format:

```
bl bids download --pub 5f2c3765beafe924c962dd8d
```

Additional information about the brainlife.io CLI commands can be found at https://github.com/brainlife/cli.

Table 1. Description and web-links to the open-source code and open cloud services used in the creation of this dataset.

| Application                         | Github repository                                      | Open Service DOI     | Git branch  |
|-------------------------------------|--------------------------------------------------------|----------------------|-------------|
| Brain Alignment                     | https://github.com/brainlife/app-fsl-anat              | https://doi.org/10.25663/brainlife.app.273 | v1.0        |
| Cortical and Subcortical Segmentation | https://github.com/brain-life/app-freesurfer            | https://doi.org/10.25663/bl.app.0         | 1.11        |
| White matter Tissue Segmentation   | https://github.com/brainlife/app-mrtrix3-3tt            | https://doi.org/10.25663/brainlife.app.239 | binarize-v1.0 |
| Multi-Atlas Transfer Tool (MaTT)   | https://github.com/faskowit/app-multiAtlasTT           | https://doi.org/10.25663/bl.app.23       | 0.5         |
| dMRI preprocessing and model fitting | https://github.com/brain-life/app-mrtrix3-preproc     | https://doi.org/10.25663/bl.app.68            | 1.6         |
| SNR Calculation                     | https://github.com/davhunt/app-snr_in_cc/tree/plot     | https://doi.org/10.25663/bl.app.120        | plot        |
| Brain mask Generation               | https://github.com/brain-life/app-FSLBET                | https://doi.org/10.25663/brainlife.app.163 | dwi         |
| NODDI model fit                     | https://github.com/brainlife/app-noddi-amico           | https://doi.org/10.25663/brainlife.app.365 | 1.3         |
| Diffusion Tensor Fit                | https://github.com/brainlife/app-fdtDtFIFT              | https://doi.org/10.25663/brainlife.app.292 | v1.0        |
| Constrained Spherical Deconvolution Fit | https://github.com/bacaron/app-mrtrix3-act       | https://doi.org/10.25663/brainlife.app.238 | csd_generation-v1.0 |
| Tractography                        | https://github.com/bacaron/app-mrtrix3-act             | https://doi.org/10.25663/brainlife.app.297 | 1.3         |
| Merging Tractography Files          | https://github.com/bacaron/app-mergeTCK                 | https://doi.org/10.25663/brainlife.app.305 | two-tck     |
| White Matter Anatomy Segmentation   | https://github.com/brainlife/app-wmaSeg                  | https://doi.org/10.25663/brainlife.app.188 | 3.7         |
| Remove Tract Outliers               | https://github.com/brainlife/app-removeTractOutliers   | https://doi.org/10.25663/brainlife.app.195 | 1.3         |
| Tract Profiles                      | https://github.com/brain-life/app-tractanalysisprofiles | https://doi.org/10.25663/brainlife.app.361 | 1.8         |
| Cortex Tissue Mappin                | https://github.com/brainlife/app-cortex-tissue-mapping | https://doi.org/10.25663/brainlife.app.379 | v1.1        |
| Structural Connectome               | https://github.com/brainlife/app-sift2-connectome-generation | https://doi.org/10.25663/brainlife.app.394 | sift2_v1.0  |

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References

1. Hillman, C. H., Erickson, K. I. & Kramer, A. F. Be smart, exercise your heart: exercise effects on brain and cognition. *Nat. Rev. Neurosci.* 9, 58–65 (2008).
2. Voss, M. W., Nagamatsu, L. S., Liu-Ambrose, T. & Kramer, A. F. Exercise, brain, and cognition across the life span. *J. Appl. Physiol.* 111, 1505–1513 (2011).
3. Penedo, F. J. & Dahn, J. R. Exercise and well-being: a review of mental and physical health benefits associated with physical activity. *Curr. Opin. Psychiatry* 18, 189–193 (2005).
4. Reimers, C. D., Knapp, G. & Reimers, A. K. Does physical activity increase life expectancy? A review of the literature. J. Aging Res. 2012, 243958 (2012).
5. Esteban-Cornejo, I. et al. Physical Fitness, White Matter Volume and Academic Performance in Children: Findings From the ActiveBrains and FITKids2 Projects. Front. Psychol. 10, 208 (2019).
6. Chadlock-Heyman, L. et al. Physical Activity Increases White Matter Microstructure in Children. Front. Neurosci. 12, 950 (2018).
7. Ruoletsalinen, I. et al. Physical activity, aerobic fitness, and brain white matter: Their role for executive functions in adolescence. Dev. Cogn. Neurosci. 42, 100765 (2020).
8. Sarauli, D., Costanzi, M., Mastroirilli, V. & Farioli-Vecchioli, S. The Long Run: Neuroprotective Effects of Physical Exercise on Adult Neurogenesis from Youth to Old Age. Curr. Neuropharmacol. 15, 519–533 (2017).
9. Hütting, K. & Röder, B. Beneficial effects of physical exercise on neuroplasticity and cognition. Neurosci. Biobehav. Rev. 37, 2243–2257 (2013).
10. El-Swey, I., Harasym, D., Turco, C. V., Locke, M. B. & Nelson, A. J. Exercise-Induced Neuroplasticity: A Mechanistic Model and Prospects for Promoting Plasticity. Neuroscientist 25, 65–85 (2019).
11. Radak, Z., Marton, O., Nagy, E., Koltai, E. & Goto, S. The complex role of physical exercise and reactive oxygen species on brain. J. Sport and Health Sci. 2, 87–93 (2013).
12. Brunetta, H. S., Holwerda, A. M., van Loon, L. J. C. & Holloway, G. P. Mitochondrial ROS and Aging: Understanding Exercise as a Preventive Tool. Journal of Science in Sport and Exercise 2, 15–24 (2019).
13. Sze, D.-Y., Heo, J.-W., Ko, J. R. & Kwak, H.-B. Exercise and Neuroinflammation in Health and Disease. Int. Neurol. J. 23, 582–92 (2019).
14. Jäncke, L., Koeneke, S., Hoppe, A., Rominger, C. & Hägg, J. The architecture of the golfer’s brain. PLoS One 4, e4785 (2009).
15. Hägg, J. et al. Structural brain correlates associated with professional handball playing. PLoS One 10, e0124222 (2015).
16. Hägg, J., Koeneke, S., Bezzola, I. & Jäncke, L. Structural neuroplasticity in the sensorimotor network of professional female ballet dancers. Hum. Brain Mapp. 31, 1196–1206 (2010).
17. Deng, F. et al. Plasticity in deep and superficial white matter: a DTI study in world class gymnasts. Brain Struct. Funct. 223, 1849–1862 (2018).
18. Montenigro, P. H. et al. Cumulative Head Impact Exposure Predicts Later-Life Depression, Apathy, Executive Dysfunction, and Cognitive Impairment in Former High School and College Football Players. J. Neurotrauma 34, 328–340 (2017).
19. Bahrami, N. et al. Subconcussive Head Impact Exposure and White Matter Tract Changes over a Single Season of Youth Football. Radiology 281, 919 (2016).
20. Churchill, N. W., Caverzas, E., Graham, S. J., Hutchison, M. G. & Schweizer, T. A. White matter microstructure in athletes with a history of concussion: Comparing diffusion tensor imaging (DTI) and neurite orientation dispersion and density imaging (NODDI). Hum. Brain Mapp. 38, 4201–4211 (2017).
21. Churchill, N. W. & Hutchison, M. G. Di Battista, A. P., Graham, S. J. & Schweizer, T. A. Structural, Functional, and Metabolic Brain Markers Differentiate Collision versus Contact and Non-Contact Athletes. Front. Neurol. 8, 390 (2017).
22. Davenport, E. M. et al. Subconcussive impacts and imaging findings over a season of contact sports. Concussion 1, C119 (2016).
23. Sobolovov, S. M. et al. The effect of repetitive subconcussive collisions on brain integrity in collegiate football players over a single football season: A multi-modal neuroimaging study. Neuroimage: Clinical 18, 708–718 (2017).
24. Lefebvre, G. et al. Diffusion tensor imaging in contact and non-contact university-level sport athletes. J. Neurotrauma https://doi.org/10.1089/neu.2020.7170 (2020).
25. Mayinger, M. C. et al. White matter alterations in college football players: a longitudinal diffusion tensor imaging study. Brain Imaging Behav. 12, 44–53 (2018).
26. Mayer, A. R. et al. A prospective microstructure imaging study in mixed-martial artists using geometric measures and diffusion tensor imaging: methods and findings. Brain Imaging Behav. 11, 698–711 (2017).
27. Yue, J. K. et al. Transforming research and clinical knowledge in traumatic brain injury pilot: multicenter implementation of the common data elements for traumatic brain injury. J. Neurotrauma 30, 1831–1844 (2013).
28. Koerte, I. K. et al. The ENIGMA sports injury working group: an international collaboration to further our understanding of sport-related brain injury. Brain Imaging Behav. https://doi.org/10.1007/s11868-020-00370-y (2020).
29. Broglio, S. P. et al. A National Study on the Effects of Concussion in Collegiate Athletes and US Military Service Academy Members: The NCAA-DoD Concussion Assessment, Research and Education (CARE) Consortium Structure and Methods. Sports Med. 47, 1437–1451 (2017).
30. Avesani, P. et al. The open diffusion data derivatives, brain data upcycling via integrated publishing of derivatives and reproducible open cloud services. Sci Data 6, 69 (2019).
31. Stewart, C. A. et al. Jetstream: A self-provisioned, scalable science and engineering cloud environment. https://doi.org/10.1145/2792745.2792774 (2015).
32. Towns, J. et al. XSEDE: Accelerating Scientific Discovery. Computing in Science Engineering 16, 62–74 (2014).
33. Woolrich, M. W. et al. Bayesian analysis of neuroimaging data in FSL. Neuroimage 45, 5173–86 (2009).
34. Smith, S. M. et al. Advances in functional and structural MR image analysis and implementation as FSL. Neuroimage 23(Suppl 1), S208–19 (2004).
35. Jenkinson, M., Beckmann, C. F., Behrens, T. E. J., Woolrich, M. W. & Smith, S. M. FSL. Neuroimage 62, 782–790 (2012).
36. Dale, A., Fischl, B. & Sereno, M. I. Cortical Surface-Based Analysis: I. Segmentation and Surface Reconstruction. Neuroimage 9, 179–194 (1999).
37. Dale, A. M. & Sereno, M. I. Improved Localization of Cortical Activity by Combining MRI and MEG with MRI Cortical Surface Reconstruction: A Linear Approach. Journal of Cognitive Neuroscience 5, 162–176 (1993).
38. Desikan, R. S. et al. An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. Neuroimage 31, 968–980 (2006).
39. Fischl, B. & Dale, A. M. Measuring the thickness of the human cerebral cortex from magnetic resonance images. Proc. Natl. Acad. Sci. USA 97, 11050–11055 (2000).
40. Fischl, B., Liu, A. & Dale, A. M. Automated manifold surgery: constructing geometrically accurate and topologically correct models of the human cerebral cortex. IEEE Medical Imaging 20, 70–80 (2001).
41. Fischl, B. et al. Whole brain segmentation: automated labeling of neuroanatomical structures in the human brain. Neuroimage 33, 341–355 (2002).
42. Fischl, B. et al. Sequence-independent segmentation of magnetic resonance images. Neuroimage 23, 569–584 (2004).
43. Fischl, B., Sereno, M. I. & Dale, A. Cortical Surface-Based Analysis: II. Inflation, Flattening, and a Surface-Based Coordinate System. Neuroimage 9, 195–207 (1999).
44. Fischl, B., Sereno, M. I., Tootell, R. B. H. & Dale, A. M. High-resolution intersubject averaging and a coordinate system for the cortical surface. Hum. Brain Mapp. 8, 272–284 (1999).
45. Fischl, B. et al. Automatically Parcellating the Human Cerebral Cortex. Cereb. Cortex 14, 11–22 (2004).
46. Han, X. et al. Reliability of MRI-derived measurements of human cerebral cortical thickness: The effects of field strength, scanner upgrade and manufacturer. Neuroimage 32, 180–194 (2006).
47. Jovicich, J. et al. Reliability in multi-site structural MRI studies: Effects of gradient non-linearity correction on phantom and human data. Neuroimage 30, 436–443 (2006).
48. Kuperberg, G. R. et al. Regionally localized thinning of the cerebral cortex in Schizophrenia. Arch. Gen. Psychiatry 60, 878–888 (2003).
49. Reuter, M., Schmansky, N. J., Rosas, H. D. & Fischl, B. Within-subject template estimation for unbiased longitudinal image analysis. Neuroimage 61, 1402–1418 (2012).
50. Reuter, M. & Fischl, B. Avoiding Asymmetry-Induced Bias in Longitudinal Image Processing. Neuroimage 57, 19–21 (2011).
51. Reuter, M., Rosas, H. D. & Fischl, B. Highly Accurate Inverse Consistent Registration: A Robust Approach. Neuroimage 53, 1181–1196 (2010).
52. Rosas, H. D. et al. Regional and progressive thinning of the cortical ribbon in Huntington's disease. Neurology 58, 695–701 (2002).
53. Salat, D. et al. Thinning of the cerebral cortex in aging. Cereb. Cortex 14, 721–730 (2004).
54. Segonne, F. et al. A hybrid approach to the skull stripping problem in MRI. Neuroimage 22, 1060–1073 (2004).
55. Segonne, F., Pacheco, J. & Fischl, B. Geometrically accurate topology-correction of cortical surfaces using nonseparating loops. IEEE Trans. Med. Imaging 26, 518–529 (2007).
56. Tournier, J.-D. et al. MRtrix3: A fast, flexible and open software framework for medical image processing and visualisation. Neuroimage 202, 116137 (2019).
57. Garyfallidis, E. et al. Dipy, a library for the analysis of diffusion MRI data. Front. Neuroinform. 8, 7 (2014).
58. Van Essen, D. C. et al. The WU-Minn Human Connectome Project: an overview. Neuroimage 80, 62–79 (2013).
59. Merkel, D. Docker: lightweight linux containers for consistent development and deployment. Linux J. 2014, 2 (2014).
60. Kellar, D., Newman, S., Pestilli, F., Cheng, H. & Port, N. L. Comparing fMRI activation during smooth pursuit eye movements among contact sport athletes, non-contact sport athletes, and non-athletes. Neuroimage Clin 18, 413–424 (2018).
61. Jenkinson, M. & Smith, S. A global optimisation method for robust affine registration of brain images. Med. Image Anal. 5, 143–156 (2001).
62. Jenkinson, M., Bannister, P., Brady, M. & Smith, S. Improved optimization for the robust and accurate linear registration and motion correction of brain images. Neuroimage 17, 825–841 (2002).
63. Greve, D. N. & Fischl, B. Accurate and robust brain image alignment using boundary-based registration. Neuroimage 48, 63–72 (2009).
64. Fukutomi, H. et al. Neurite imaging reveals microstructural variations in human cerebral cortical gray matter. Neuroimage https://doi.org/10.1016/j.neuroimage.2018.02.017 (2018).
65. Destrieux, C., Fischl, B., Dale, A. & Halgren, E. Automatic parcellation of human cortical gyri and sulci using standard anatomical nomenclature. Neuroimage 53, 1–15 (2010).
66. Glasser, M. F. et al. A multi-modal parcellation of human cerebral cortex. Nature 536, 171–178 (2016).
67. Jeurissen, B., Leemans, A. & Sijbers, J. Automated correction of improperly rotated diffusion gradient orientations in diffusion weighted MRI. Med. Image Anal. 18, 953–962 (2014).
68. Versart, J. et al. Denoising of diffusion MRI using random matrix theory. Neuroimage 142, 394–406 (2016).
69. Kellner, E., Dhiatal, B., Kiselev, V. G. & Reisert, M. Gibbs-ringing artifact removal based on local subvoxel-shifts. Magn. Reson. Med. 76, 1574–1581 (2016).
70. Andersson, J. L. R., Skare, S. & Ashburner, J. How to correct susceptibility distortions in spin-echo echo-planar images: application to diffusion tensor imaging. Neuroimage 20, 870–888 (2003).
71. Andersson, J. L. R. & Sotiropooulos, S. N. An integrated approach to correction of off-resonance effects and subject movement in diffusion MR imaging. Neuroimage 125, 1063–1078 (2016).
72. Andersson, J. L. R., Graham, M. S., Zsoldos, E. & Sotiropooulos, S. N. Incorporating outlier detection and replacement into a non-parametric framework for movement and distortion correction of diffusion MRI images. Neuroimage 141, 556–572 (2016).
73. Andersson, J. L. R., Graham, M. S., Drobnjak, I., Zhang, H. & Campbell, J. Susceptibility-induced distortion that varies due to motion: Correction in diffusion MR without acquiring additional data. Neuroimage 171, 277–295 (2018).
74. Andersson, J. L. R. et al. Towards a comprehensive framework for movement and distortion correction of diffusion MR images: Within volume movement. Neuroimage 152, 450–466 (2017).
75. Tustison, N. J. et al. Large-scale evaluation of ANTs and FreeSurfer cortical thickness measurements. Neuroimage 99, 166–179 (2014).
76. Smith, S. M. Fast robust automated brain extraction. Hum. Brain Mapp. 17, 143–155 (2002).
77. Descoteaux, M., Deriche, R., Le Bihan, D., Mangin, J.-F. & Poupon, C. Multiple q-shell diffusion propagator imaging. Med. Image Anal. 15, 603–621 (2011).
78. Jones, D. K., Knösche, T. R. & Turner, R. White matter integrity, fiber count, and other fallacies: the don'ts and don'ts of diffusion MRI. Neuroimage 73, 239–254 (2013).
79. Zhang, H., Schneider, T., Wheeler-Kingshott, C. A. & Alexander, D. C. NODDI: practical in vivo neurite orientation dispersion and density imaging of the human brain. Neuroimage 61, 1000–1016 (2012).
80. Daducci, A. et al. Accelerated Microstructure Imaging via Convex Optimization (AMICO) from diffusion MRI data. Neuroimage 105, 32–44 (2015).
81. Jeurissen, B., Tournier, J.-D., Dhollander, T. Connolly, A. & Sijbers, J. Multi-tissue constrained spherical deconvolution for improved analysis of multi-shell diffusion MRI data. Neuroimage 103, 411–426 (2014).
82. Tournier, J.-D., Calamante, F., Gadian, D. G. & Connelly, A. Direct estimation of the fiber orientation density function from diffusion-weighted MRI data using spherical deconvolution. Neuroimage 23, 1176–1185 (2004).
83. Takekura, H., Cañada, C. F. & Wandell, B. A. & Pestilli, F. Ensemble Tractography. PLoS Comput. Biol. 12, e1004692 (2016).
84. Smith, R. E., Tournier, J.-D., Calamante, F. & Connelly, A. Anatomically-constrained tractography: improved diffusion MRI streamlines tractography through effective use of anatomical information. Neuroimage 62, 1924–1938 (2012).
85. Bullock, D. et al. Associative white matter connecting the dorsal and ventral posterior human cortex. Brain Struct. Funct. https://doi.org/10.1007/s00429-019-01907-8 (2019).
86. Pestilli, F., Yeatman, J. D., Rokem, A., Kay, K. N. & Wandell, B. A. Evaluation and statistical inference for human connectomes. Nat. Methods 11, 1058–1063 (2014).
87. Yeatman, J. D., Dougherty, R. F., Myall, N. J., Wandell, B. A. & Feldman, H. M. Tract profiles of white matter properties: automating fiber-tract quantification. PLoS One 7, e49790 (2012).
88. Smith, R. E., Tournier, J.-D., Calamante, F. & Connelly, A. The effects of SIFT on the reproducibility and biological accuracy of the structural connectome. Neuroimage 104, 253–265 (2015).
89. Smith, R. E., Tournier, J.-D., Calamante, F. & Connelly, A. SIFT2: Enabling dense quantitative assessment of brain white matter connectivity using streamlines tractography. Neuroimage 119, 338–351 (2015).
90. Hagmann, P. et al. Mapping the structural core of human cerebral cortex. PLoS Biol. 6, e159 (2008).
91. Smith, R. E., Tournier, J.-D., Calamante, F. & Connelly, A. SIFT: Spherical-deconvolution informed filtering of tractograms. Neuroimage 67, 298–312 (2013).
92. Gorgolewski, K. J. et al. The brain imaging data structure, a format for organizing and describing outputs of neuroimaging experiments. Sci Data 3, 160040 (2016).
93. Carolan, B. & Pestilli, F. Collegiate athlete brain data for white matter mapping and network neuroscience. brainlife.io https://doi.org/10.56563/brainlife.pub14 (2020).
94. Yeh, C.-H., Jones, D. K., Liang, X., Descoteaux, M. & Connelly, A. Mapping Structural Connectivity Using Diffusion MRI: Challenges and Opportunities. *J. Magn. Reson. Imaging* https://doi.org/10.1002/jmri.27188 (2020).

95. Rubinov, M. & Sporns, O. Complex network measures of brain connectivity: uses and interpretations. *Neuroimage* **52**, 1059–1069 (2010).

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**Author contributions**

B.C., R.S. and F.P. wrote the manuscript text, performed analyses, and prepared Figs. 1–10. B.C., D.B., L.K., J.F., B.M., and F.P. contributed to software development. B.C., D.K., H.C., S.N., N.P. and F.P. contributed to data curation. All authors reviewed the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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