HippUnfold: Automated hippocampal unfolding, morphometry, and subfield segmentation

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Key words: hippocampal subfields, MRI, morphology, deep learning, neuroimaging software

Abstract

Like neocortical structures, the archicortical hippocampus differs in its folding patterns across individuals. Here, we present an automated and robust BIDS-App, HippUnfold, for defining and indexing individual-specific hippocampal folding in MRI, analogous to popular tools used in neocortical reconstruction. Such tailoring is critical for inter-individual alignment, with topology serving as the basis for homology. This topological framework enables qualitatively new analyses of morphological and laminar structure in the hippocampus or its subfields. It is critical for refining current neuroimaging analyses at a meso- as well as micro-scale. HippUnfold uses state-of-the-art deep learning combined with previously developed topological constraints to generate uniquely folded surfaces to fit a given subject's hippocampal conformation. It is designed to work with commonly employed sub-millimetric MRI acquisitions, with possible extension to microscopic resolution. In this paper we describe the power of HippUnfold in feature extraction, and highlight its unique value compared to several extant hippocampal subfield analysis methods.

Introduction

Most neurological or psychiatric diseases with widespread effects on the brain show strong and early impact on the hippocampus (e.g. [1]). This highly plastic grey matter (GM) structure is also critical in the fast formation of episodic and spatial memories (e.g. [2]). Examination of this structure with non-invasive neuroimaging, such as MRI, provides great promise for furthering our understanding, diagnosis, and subtyping of these diseases and cognitive processes in the hippocampus and its component subfields [3].
In current neuroimaging analyses the hippocampus is typically modelled as a subcortical volume, but it is actually made up of a folded archicortical mantle, or ‘ribbon’ [4]. Representing the hippocampus as such can be leveraged to enable qualitatively new analyses, such as registration, despite inter-individual differences in gyrification or folding structure, through topological alignment. Additionally, representation as a ribbon allows the hippocampus to be factorized into surface area and thickness, which can be further subdivided for laminar analyses. These methods are thus critical in advancing MRI research from the macroscopic scale to the subfield, cortical column, and laminar scales. Similar approaches have already yielded a paradigm shift in neocortical analysis methods [5,6].

Denoting the hippocampal archicortex or ribbon is challenging because it is thin (0.5-2mm), its folding pattern varies considerably between individuals [7,8], and this folding may even continue to change from early development through adulthood [9]. We present here a set of tools to overcome these challenges using a highly sensitive and generalizable “U-Net” deep learning architecture [10], combined with previous work that enforces topological constraints on hippocampal tissue [11].

In previous work [11], we developed a method to computationally unfold the hippocampus along its geodesic anterior-posterior (AP) and proximal-distal (PD, i.e., proximal to the neocortex, with the dentate gyrus being most distal) axes. We demonstrated for the first time several qualitative properties using in vivo MRI, such as the contiguity of all subfields along the curvature of the hippocampal head (anterior) and tail (posterior), previously described only in histology. This pioneering work relied heavily on detailed manual tissue segmentations including the high-myelinated stratum radiatum, lacunosum, and moleculaire (SRLM), a commonly used landmark that separates hippocampal folds along the inward ‘curl’ of the hippocampus. In this work we also considered curvature and digitations along the AP axis of the hippocampus, most prominently occurring in the hippocampal head [4,7,8,11]. Each of these features are highly variable between individuals, making them difficult to capture using automated volumetric atlas-based methods and time-consuming to detect manually.

The current work automates the detailed tissue segmentation required for hippocampal unfolding using a state-of-the-art ‘U-Net’ deep convolutional neural network [10]. In particular, we aimed to capture morphological variability between hippocampi which are not seen using existing automated methods which employ either a single atlas or multi-atlas fusion (eg. [12–14]). U-Net architectures have been shown to be generalizable and sensitive to anatomical variations in many medical image processing tasks [15], making them ideal to overcome this challenge.

Estimating hippocampal subfield boundaries in MRI is challenging since their histological hallmarks are not directly available in MRI due to lower spatial resolution and lack of appropriate contrasts, which is an ongoing hurdle in neuroimaging [16,17]. However, post-mortem studies show that the subfields are topologically constrained according to their differentiation from a common flat cortical mantle [4]. Thus a folded representation of hippocampal tissue provides a powerful intermediate between a raw MRI and subfield labels [18], analogous to the reconstruction of a 3D neocortical surface. This surface can then be parcellated into subregions without topological breaks [5], overcoming many limitations of current subfield segmentation methods [17]. Here, we apply surface-based subfield boundary definitions obtained via manual segmentation of BigBrain 3D histology [19] which was additionally supported by a data-driven
parcellation [20]. We additionally demonstrate how labels used in the popular Freesurfer [21] and Automatic Segmentation of Hippocampal Subfields (ASHS) [12] software packages can be applied under our topologically-constrained framework.

Altogether, we combine novel U-Net tissue classification, previously developed hippocampal unfolding [11], and topologically-constrained subfield labelling [20] together into a single pipeline which we refer to as ‘HippUnfold’ hereinafter. We designed this pipeline to employ FAIR principles (findability, accessibility, interoperability, reusability) with support across a wide range of use-cases centered around sub-millimetric MRI.

Results

HippUnfold aligns and visualizes data on folded or unfolded surfaces

HippUnfold is presented here as a fully-automated pipeline with outputs including hippocampal tissue and subfield segmentations, geodesic Laplace coordinates spanning over hippocampal GM voxels, and inner, midthickness and outer hippocampal surfaces. These surfaces have corresponding vertices, providing an implicit topological registration between individuals.

The overall pipeline for HippUnfold is illustrated briefly in Figure 1. A comprehensive breakdown of each step is provided in the Materials and Methods.

Figure 1. Overview of HippUnfold pipeline. First, input MRI images are preprocessed and cropped around the left and right hippocampi. Second, a U-Net neural network architecture (nnUNet [10]) is used to segment hippocampal grey matter (GM), the high-myelinated stratum radiatum, lacunosum, and moleculare (SRLM), and structures surrounding the hippocampus. Segmentations are post-processed via template shape injection. Third, Laplace’s equation is solved across the anterior-posterior (AP), proximal-distal (PD) and inner-outer (IO) extent of hippocampal GM, making up a geodesic coordinate framework. Fourth, scattered interpolants are used to determine equivalent coordinates between native Cartesian space and unfolded space. Fifth, unfolded surfaces with template subfield labels [20] are
transformed to subjects’ native folded hippocampal configurations. Morphological features (e.g., thickness) are extracted using Connectome Workbench [22] on these folded native space surfaces. Sixth, volumetric subfields are generated by filling the voxels between inner and outer surfaces with the corresponding subfield labels. Additional details on this pipeline can be found in the Materials and Methods.

In addition to subfield segmentation, HippUnfold extracts morphological features and can be used to sample quantitative MRI data along a mid-thickness surface to minimize partial voluming with surrounding structures. This is visualized across n=148 test subjects on an unfolded surface and group-averaged folded surface in Figure 2. Note that the group averaging takes place on a surface and so does not break individual subjects' topologies. Quantitative MRI features examined here include T1w/T2w ratio as a proxy measure for intracortical myelin [23], mean diffusivity, and fractional anisotropy [24,25].

Figure 2. Average hippocampal folded and unfolded surfaces showing subfields, morphometric and quantitative MRI measures from the HCP-YA test dataset (see Table 1 of Materials and Methods). The same topologically defined subfields were applied in unfolded space to all subjects (top), which are also overlaid on quantitative MRI plots (black lines). The dentate gyrus (DG) is represented as a distinct surface, reflecting its unique topology, and is mostly occluded in native space. Thickness was not measured across the dentate gyrus surface. Note that many morphological and quantitative MRI measures show clear distinctions across subfield boundaries.
Clear differences in morphological and quantitative MRI features can be seen across the hippocampus, particularly across subfields as defined here from a histologically-derived unfolded reference atlas [20]. This highlights the advantages of the present method. These folded and unfolded representations of hippocampal characteristics are broadly in line with previous work examining differences in such morphological and quantitative MRI features across hippocampal subfields or along the hippocampal AP extent (eg. [26,27]). However, in previous work these features differed between predefined subfields on average, but did not necessarily follow subfield contours as seen here. Some advantages of the current pipeline that likely contribute to this clarity include i) the detail of the hippocampal GM segmentation, ii) sampling along a midthickness surface to minimize partial voluming with surrounding structures, and iii) the fact that subjects are topologically aligned leading to less blurring of features after group-averaging.

Extant methods do not respect the topological continuity of hippocampal subfields

Several automatic methods for labelling hippocampal subfields in MRI exist, of which Freesurfer [21] (FS, v7.1) and Automatic Segmentation of Hippocampal Subfields [12] (ASHS) are among the most widely adopted. These methods rely on volumetric registrations between a target hippocampus and a reference or atlas. Specifically, ASHS makes use of multi-atlas registration, wherein multiple gold standard manual hippocampal subfield segmentations are registered to a target sample. Typically the multi-atlas consists of roughly a dozen samples which are then fused together to generate a reliable yet oftentimes smooth or simplified final product. FS uses a combination of voxel-wise classification and, bijectively, volumetric registration between a target hippocampus and a probabilistic reference atlas, which is generated via combined in vivo MRI and 9.4T ex vivo hippocampal subfield segmentations [21]. When hippocampi take on different folding configurations, such registrations can become ill-posed. HippUnfold overcomes these limitations in two ways: with extensive training (in this case n=590), U-Net can capture detailed inter-individual differences in folding and, secondly, our unfolding technique ensures that subfield labelling is topologically constrained [18].

We applied Freesurfer’s (v7.1) hippocampal subfields pipeline as well as ASHS using a recent manual subfield multi-atlas [28] to the HCP-YA test set (see Table 1 of Materials and Methods). We then compared resulting subfield segmentations to those generated via HippUnfold in native and unfolded space, which is shown in Figure 3 in one representative subject. For comparison, we additionally mapped FS and ASHS subfield boundaries in folded and unfolded space.
Figure 3. Comparison of HippUnfold, ASHS, and Freesurfer subfield segmentations in native and unfolded space. Sagittal and coronal slices and 3D models are shown for one representative subject. Note that for HippUnfold hippocampal subfields are the same for all individuals in unfolded space, but for ASHS and FS we mapped all subjects’ subfield boundaries which are shown in the black lines in column 4 rows 2 and 4. We then took the mode subfield label from ASHS and FS in unfolded space and projected it back to native space, which is shown in rows 3 and 5.

Both ASHS and FS showed subfield discontinuities in unfolded space in at least some subjects, and FS even showed discontinuities in the group-averaged unfolded subfields. That is, some pieces of a given label were separated from the rest of that label. ASHS does not include an SRLM label and the SRLM produced by FS was not consistently aligned with that used in unfolding. Thus, subfields sometimes erroneously crossed the SRLM, breaking topology and explaining why discontinuities were sometimes observed in unfolded space. Ordering of labels was also not consistent in ASHS and FS. For example, sometimes CA1 would border not only CA2 but also CA3, CA4, and/or DG. Additionally, neither ASHS nor FS extends all subfields to the full anterior and posterior extent of the hippocampus. Instead, both methods simplify most of the anterior hippocampus as being CA1 and opt not to label subfields in the posterior hippocampus at all. These qualities are not in line with the anatomical ground truth shown in both classic and contemporary ex-vivo histological studies [4,8], which were indeed well captured by HippUnfold. FS also over-labelled hippocampal tissue, which can be seen reaching
laterally into the ventricles in the coronal view. Similar errors have been documented for FS in other recent work [29,30].

Trained U-Net performance is similar to manual segmentation

From the HCP-YA dataset, a set of 738 (left and right from 369 subjects) gold standard hippocampal tissue (that is, hippocampal GM and surrounding structures) segmentations were generated according to the manual protocol defined in [20]. Automated tissue segmentation was performed using nnUNet, a recent and highly generalizable implementation of a U-Net architecture [10] wrapped into a Snakemake workflow [DOI]. This software was trained on 80% (n=590) of the gold standard segmentation data described above, with the remaining 20% (n=148) making up a test set. Left and right hippocampi from the same participant were never split across training and testing sets due to their high symmetry. Note that all input images were preprocessed, resampled, and cropped (see Figure 1 and Materials and Methods) prior to training. Within the training set, 5-fold cross-validation was performed as implemented in the nnUNet code. Training took place on an NVIDIA T4 Turing GPU over 72 hours. This process was carried out using either T1w or T2w input data with the same training/testing data split. All default nnUNet data augmentation and hyperparameters were used.

![Diagram](image)

Figure 4. Test set performance in Dice overlaps between HippUnfold and manually unfolded subfields. All values are compared to ground truth manually defined tissues followed by unfolded subfield definition (manual unfold). Two models were trained in parallel using the same labels but different input MRI data modalities consisting of T1w or T2w data. Dotted black lines indicate corresponding values from [12], who include SRLM in all labels and combine CA4 and DG into one label.

Dice overlap depends heavily on the size of the label in question, being lower for smaller labels. Typically a score of >0.7 is considered good, and many fully manual protocols show dice scores of >0.8 for the larger subfields like CA1 or the subiculum, and 0.6-0.8 for smaller subfields like CA2 or CA3 (see [17] for overview). Within the HCP-YA test set, performance was similar or better than most fully manual protocols for T1w and T2w data. Performance on T1w images was only marginally poorer than T2w images which typically better show the SRLM and are popular in manual subfield segmentation protocols [17].

Generalizability to unseen datasets and populations

We aimed to determine whether our pipeline would generalize to unseen datasets with different acquisition protocols and sample populations. Hippocampal morphometry, integrity, and
subfields are often of interest in disease states where atrophy or other structural abnormalities are observed [1,31–33]. For this reason, we examined the HCP-A datasets in which we anticipated cases of severe atrophy would be present in some older subjects. Figure 5A shows results from one representative individual (an 80 y.o. female with signs of age-related atrophy but good scan quality). Another common use-case for hippocampal subfield segmentation is on anisotropic T2w data which is considered optimal for performing manual segmentation in most protocols [17], but may impose challenges for our method due to the difference in resolution. We thus applied HippUnfold to 7T-TSE data and also illustrate one representative subfield segmentation result in Figure 5A.

Figure 5. Examination of HippUnfold performance on additional datasets HCP-A (T1w and T2w) and anisotropic 7T-TSE data. A) Sample subjects’ HippUnfold subfield segmentation in native resolution. The first two rows come from the same subjects but using different input data modalities. B) Subjects flagged for Quality Assurance from each dataset based on Dice overlap with a reference mask approximated via deformable registration. C) Failed subject example illustrating missed tissue (red arrows) at the nnUNet pipeline stage.

Gold standard manual segmentations under the protocol used for subsequent unfolding were not available in the generalization datasets. Manually inspecting results from hundreds of subjects is time consuming. We thus streamlined this process by flagging potential segmentation errors by examining Dice overlap with a more conventional segmentation approach: deformable registration. For all datasets described above, we applied deformable fast
B-spline registration [34] to the corresponding T1w or T2w template. Tissue segmentation results (generated at the nnUNet stage) were then propagated to template space and overlap with standard template hippocampal masks were examined, which is shown in Figure 5B. Any subject with a Dice overlap score of less than 0.7 was flagged and manually inspected for quality assurance. This made up 34/2126 (1.6%) samples in the HCP-YA T2w set (including training and testing subsets), 188/1312 (14.3%) samples from the HCP-A T2w set, 37/1312 (2.8%) samples from the HCP-A T1w set, and 3/92 (3.3%) samples from the 7T-TSE set. Closer inspection revealed that the vast majority of flagged cases were due to missed tissue in the nnUNet segmentation, an example of which is shown in Figure 5C. It is interesting to note that the most flagged cases were seen in the HCP-A T2w dataset even though T2w is a popular acquisition protocol for hippocampal subfield segmentation [17], and showed the best performance within the HCP-YA test set (Figure 4). This was likely not due to the age of subjects since few of the HCP-A T1w were flagged as possible errors, but instead may have been due to T2w scan quality, which was observed to be poor in some subjects, causing poor definition of the outer hippocampal boundaries. We recommend that future users carefully inspect results from any flagged subjects, and cases with errors can be either discarded or manually corrected. We cannot determine whether HippUnfold will work as intended on all new datasets, but within the generalization datasets examined here, results were excellent. Some work has already demonstrated it is possible to synthesize or convert between MRI modalities [35], which could be used to alleviate the dependency on any single MR contrast.

FAIR principles in development

We designed this pipeline to employ FAIR principles (findability, accessibility, interoperability, reusability). As such, we have made use of several tools, conventions, and data standards to make HippUnfold extensible and easy to use.

The default file input-output structure of the HippUnfold command line interface was built in compliance with the Brain Imaging Data Standards (BIDS) [36] Applications (BIDS-Apps) guidelines [37], and easily findable amongst the list of available BIDS Apps¹. This is achieved via Snakebids, a tool designed to interface between BIDS datasets and Snakemake [38]. All aspects of HippUnfold use Snakemake [39], a workflow management system based on Python which is reproducible, scalable, and seamlessly combines shell commands, Python code, and external dependencies in a human-readable workflow. There is no need to install these dependencies, which are containerized within the Singularity or Docker versions of HippUnfold.

Altogether, this means that in a single line this pipeline can be applied intelligently to any BIDS-complaint dataset containing a whole-brain T1w image and a T2w image (whole-brain or limited field of view) without having to specify further details. Typical runtimes on a standard desktop are 1 hour per subject, but this is further parallelized for faster processing when multiple subjects and added compute resources (or cloud computing) are available. Additional flags can be used to extend functionality to many other use-cases, including T1w only, T2w only, diffusion-weighted imaging, cases where a manual tissue segmentation is already available, or ex-vivo tissue samples.

¹ https://bids-apps.neuroimaging.io/apps/
Outputs of HippUnfold follow the standards for BIDS derivatives, and include preprocessed input images, volumetric subfield segmentations, inner, midthickness, and outer hippocampal surfaces, vertex-wise morphometric measures of thickness, curvature, and gyration, and a brief quality control (QC) report. All surface-based outputs are combined into a Connectome Workbench [40] specification file for straightforward visualization in alignment with HCP neocortical reconstructions. Outputs can be specified to include images in the original T1w space or in the resampled, cropped space that processing is performed in.

All code, code history, documentation, and support are offered online.²

Discussion

One of the most powerful features of HippUnfold is its ability to provide topological alignment between subjects despite differences in folding (or digitation) structure. This is a critical element of mainstream neocortical analysis methods that, until now, has not been carried out systematically in the archicortex, or hippocampus. The power of this form of topological alignment is evident when mapping morphological or quantitative features across the hippocampus in a large population, which we demonstrate in Figure 2.

We compare HippUnfold to other commonly used tools for hippocampal analysis, Freesurfer v7.1 (FS) and Automated Segmentation of Hippocampal Subfields (ASHS) (Figure 3). Both of these methods rely on smooth deformation of single or multi-atlas references, indicating they do not easily transfer to drastically different hippocampal folding patterns, which are often seen in the hippocampal head and tail across individuals. Both of these methods showed unfolded subfield patterns that were less consistent with ground truth histological literature than the output provided by HippUnfold. Common issues in other methods include introducing breaks in subfield topology, simplifications like the exclusion of the hippocampal tail, or inconsistent ordering of subfields. This highlights some of the advantages of HippUnfold, which was designed to overcome these issues explicitly.

Several factors make surface-based methods difficult to implement in the hippocampus, including its small size, and the difficulty of distinguishing the hippocampal sulcus or SRLM laminae that separate hippocampal folds. Here we have overcome these issues using a highly generalizable and sensitive neural network ‘U-Net’ architecture, combined with our previously developed topological unfolding framework. Together, these methods achieved similar or better Dice overlap scores than what is typically seen between two manual raters on all subfields. We tested performance on new datasets (‘generalization’ datasets with different characteristics than the HCP training set) and saw good performance in nearly all cases. Specifically, we tested other common imaging protocols including different sample age groups (HCP-A) and thick-slice 7T TSE acquisitions often used in targeted hippocampal subfield imaging [17]. Though error rates were low, we do show how and why such errors sometimes occur, highlighting the importance that future users examine the brief quality control reports included for each subject. Thus, while HippUnfold is shown to work well with all datasets examined here, we expect the widespread adoption of higher-resolution acquisition techniques will further improve feasibility at other research institutes.

² https://github.com/khanlab/hippunfold
One important limitation of our method is that HippUnfold did not consistently show clear
digitation in the hippocampal head, body, and tail which was sometimes seen in manual
segmentation in the training set and in other work (see Supplementary Materials). This reflects a
lack of detail compared to histological ground truth materials, and affects downstream
processing. That is, an overly smoothed hippocampal surface will appear thicker and have a
smaller surface area compared to one that captures the full extent of digitations. This smaller
surface area also results in each subfield boundary being proportionally shifted. Future work
could improve this pipeline by training and testing with higher-resolution data where digitations
can more clearly be distinguished both in labelmaps and in the underlying images.

The current work has applications beyond subfield imaging, enabling new investigations
of the hippocampus on a columnar and laminar scale. For example, rather than employing
ROI-based analyses, statistics can be performed on a per-vertex basis for vertices generated at
different depths. This is in line with state-of-the-art neocortical analysis methods [5], and opens
up the possibility of more precise localization of hippocampal properties. Similarly, it is worth
noting that the methods used here are not necessarily restricted to MRI, as we have used the
same surface-based unfolding in combination with manual segmentation to characterize the
hippocampus in 3D BigBrain histology [20].

Altogether, we show that the BIDS App ‘HippUnfold’ that we have developed in this work
(i) respects the different internal hippocampal folding configurations seen between individuals,
(ii) can be applied flexibly to T1w or T2w data, sub-millimetric isotropic or thick-slice anisotropic
data, and (iii) compares favourably to other popular methods including manual segmentation,
ASHS, and Freesurfer. We believe this tool will open up many avenues for future work including
examination of variability in hippocampal morphology which may show developmental
trajectories or be linked to disease, or the examination of hippocampal properties perpendicular
or tangential to its laminar organization with diffusion-weighted imaging. Finally, it is worth noting
that the methods described here stand to improve existing techniques by providing greater
anatomical detail and, critically, greater precision through topological alignment across
individuals who vary in anatomical structure.

Materials and Methods

Data

HippUnfold was designed and trained with the Human Connectome Project (HCP) 1200
young adult subject data release (HCP-YA) [41], and additionally tested on the HCP Aging
dataset (HCP-A) [42], and anisotropic (or thick-slice) 7T data (7T-TSE) from [28] which is
considered optimal by many hippocampal subfield researchers [17]. These data are
summarized briefly in Table 1.

Table 1. MRI datasets used in training, evaluation, and comparison to extant methods. Methods employed
include those proposed here (HippUnfold), the same processing but with manual segmentation (similar to
previous work [20]) (manual unfold), Freesurfer v7.1 [21], and an atlas of manual segmentations [28] used
in ASHS [12].
| Name    | Modalities | Resolution     | Sample size (L+R) | Methods employed                      |
|---------|------------|----------------|-------------------|---------------------------------------|
| HCP-YA  | T1w, T2w  | 0.7x0.7x0.7mm  | n=590 (training) | HippUnfold                            |
|         |            |                |                   | Manual unfold                         |
|         |            |                | n=148 (testing)   | HippUnfold                            |
|         |            |                |                   | Manual unfold                         |
|         |            |                |                   | Freesurfer (v7.1)                     |
| HCP-A   | T1w, T2w  | 0.8x0.8x0.8mm  | n=1312            | HippUnfold                            |
| 7T-TSE  | T2w        | 0.4x0.4x1.0mm  | n=70              | HippUnfold                            |
|         |            |                |                   | Manual subfields                      |
|         |            |                |                   | (ASHS atlas)                          |

nnUNet training

UNet performs classification of each input image voxel, and it is not constrained by smooth displacements used in deformable atlas registration. This is important because smooth deformable registration can be ill-posed for an atlas with a different hippocampal folding configuration than the target. For example, when trying to register a hippocampus with 2 anterior digitations to one with 4 anterior digitations, topological breaks may be seen which leads to loss of detail and disproportionate stretching or compression of some subfields, an issue that is discussed in [43]. Instead, a U-Net classifies voxels individually based on a combination of local low-level and global high-level image features with no explicit smoothness constraints.

In the current work, gold standard training and test comparison segmentations were generated in a semi-automated but heavily supervised manner: a U-Net implementation (NiftyNet [44], which is no longer maintained) was trained on existing data from [11]. This was then applied to new HCP-YA data and results were manually inspected. In many cases, results were poor due to the relatively small training sample size, but good quality segmentations from roughly 50% of subjects were selected and corrected by a manual rater (JD or MY) before being added to the initial training set for a new, de-novo application of U-Net training. This process is typically referred to as incremental learning, and was applied in four iterations until a larger set of high quality, manually inspected and corrected segmentations (738 samples from 369 subjects) was achieved.

Once the gold-standard training data was obtained, we applied a U-Net implementation called nnUNet [10]. nnUNet was built to include many state-of-the art deep learning techniques including sensible hyperparameter selection, built-in 5-fold cross-validation, and other features that have been shown to perform well and minimize possible sources of bias in medical imaging. We thus applied all default parameters in our use of this tool. Training was repeated using the same labelmaps but different underlying images for T1w, T2w, and DWI images. For each of these modalities, training took place on an NVIDIA T4 Turing GPU over 72 hours. Additional new models (or fine-tuned models) can also be trained and supplied within our code framework.
Key image features to recognize

There is a highly complex interaction between image contrast, resolution, signal dropout (which is common to the inferior temporal lobes and can affect the anterior hippocampus), and other features such as the acquisition point-spread function and resulting spatial autocorrelation or other imaging artifacts [30,45]. Thus it is difficult to determine whether HippUnfold or even manual segmentation will be straightforward on a given dataset without careful inspection of images.

One key image feature recognized in many manual protocols is the SRLM (stratum radiatum, lacunosem, and moleculaire), which consists of high-myelinated laminae surrounding the hippocampal sulcus [46]. This is typically most easily observed in T2w images, but it is also possible to see in T1w and DWI images as well [47,48]. This feature should ideally be visible on coronal slices of the hippocampal body, but also in the hippocampal head where it follows the folding of digitations (see example Figure 6). Similarly, digitations should also be visible along the outer boundary of the hippocampus. This can be difficult to distinguish since many parts of the hippocampus are wrapped by the alveus followed by the CSF of the third ventricles. Thus there can sometimes be partial voluming between alveus, CSF, and hippocampal grey matter. As an approximate rule, if these features are easily visible using the human eye, then there is a good chance HippUnfold will also be able to distinguish them. If these features are blurred or show limited visibility, then HippUnfold may fail or else produce smoothed segmentations with limited digitations.
Figure 6. Examples of discernable folding in different image types. Hippocampal folds (highlighted by the red lines) are flanked by the SRLM and/or cysts and/or CSF of the hippocampal sulcus on the inside, and by alveus or CSF of the third ventricles on the outside, making them difficult to distinguish in some image types. The first and second image sets show the same slices from the same individual, but it is considerably more difficult to see hippocampal folds in the T1w image making them less preferable. These features can even be seen out-of-plane with thick slice 7T-TSE images (bottom image set, data from [28]), as in the digitations of the hippocampal tail in the sagittal view.

**HippUnfold detailed pipeline**

1. Preprocessing and resampling. Data is gathered via snakebids [49], which automatically and flexibly queries the specified BIDS directory for T1w and T2w images. Data is loaded and saved using NiBabel [50]. Processing of each image is as follows:
a. **T1w**: N4 bias correction is performed using the Advanced Normalization Toolkit (ANTs) [51] followed by affine registration (NiftyReg [34]) to CITI168 atlas [52]. This transformation is composed (Convert 3D or c3d [53]) with a precomputed transform from CITI168 to oblique to the long-axis of the hippocampus. Images are resampled to 0.3mm$^3$ and cropped to 128x256x128 voxels centered on the CITI168 left and right hippocampi. Left hippocampi are flipped sagittally to resemble right hippocampi. We refer to this as cropped coronal oblique space.

b. **T2w**: N4 bias correction is performed as above, and if multiple T2w images are present then they are rigidly registered (NiftyReg) and then averaged, a rudimentary form of super-resolution sampling (eg. [54]). Rigid registration to the corresponding T1w image is then performed (NiftyReg), and resampled to cropped coronal oblique space as above.

A ‘modality’ flag is used to determine which image modalities should be used if multiple are present in the input BIDS directory. Within the HippUnfold code, optional flags can be used to skip preprocessing and registration. Manually segmented hippocampal tissues can also be specified, which can be useful in ex-vivo MRI or other modalities on which the current nnUNet-based segmentation is not expected to work. In all cases, data are resampled to cropped coronal oblique space to match the nnUNet training setup. It is possible to skip this step only if a manually segmented hippocampal tissue class image is also provided (in which case nnUNet is not applied).

2. **Tissue class segmentation.** If a manually segmented hippocampal tissue image is not supplied, then the input image will be run through nnUNet [10], a state-of-the-art implementation of a deep convolutional neural network (U-Net) designed for image segmentation [55,56]. The output of nnUNet is a segmentation of tissue classes: hippocampal grey matter (GM) and the surrounding tissues which are used in defining unfolded coordinate boundaries: SRLM, medial temporal lobe cortex (MTLc), pial surface, hippocampal-amygdalar transition area (HATA), indusium griseum (IndGris), cysts, and the dentate gyrus granule cell layer (DG) (which also makes up part of hippocampal grey matter but which marks an endpoint of the unfolding coordinate framework and so it was given a distinct label).

3. **Post-processing.** Here we employed template shape injection [57] to correct possible segmentation errors, making labelmaps more amenable to the previously developed hippocampal unfolding methods. The basic principle of template shape injection is to perform highly fluid deformable registration of a template segmentation labelmap to a given subject’s segmentation labelmap. This differs from typical registration-based segmentation methods in that the registration is optimizing labels rather than image feature similarity (i.e. registration is performed with binarized and smoothed labels as multiple contrasts, rather than on MRI intensities). Specifically, we used mean squared error between labels as the cost function, which is minimized when identical labels are overlapping. In our implementation, we apply multi-contrast deformable registration using Greedy [53]. It should be noted that in principle this step is not necessary for our pipeline, but in practice it helps avoid possible errors due to nnUNet segmentation faults (see main text Figure 5).

The reference template that we applied was created using manual segmentations from
an open source ex-vivo dataset [58] that was manually segmented according to our previous manual hippocampal unfolding protocol [11]. Labelmaps from 22 samples were combined using a standard template building ANTs script 'buildtemplateparallel.sh' [59]. This template generation entails averaging all images and then registering each sample to the average, iteratively refining and sharpening the average image. This ex-vivo dataset was selected for template building because we had high confidence in the quality of these segmentation since they contained higher resolution and contrast than other datasets while still including multiple samples.

4. Unfolding. This code is described in [11] and was modified in [20], but we will provide a short summary here. A Laplace field varying from 0 to 1 is generated across hippocampal grey matter, with 0 being at its anterior boundary with the HATA and 1 being at its posterior boundary with the IndGris (anterior-posterior or AP). This provides a scaled, smooth, geodesic way to index points along this axis. Another Laplace field is generated across the proximal-distal (or PD) axis of the hippocampus (MTLc to DG), and together these two fields provide a coordinate system spanning hippocampus grey matter along two dimensions, which we plot as a flat rectangle (with a 2:1 aspect ratio to reflect the fact that the hippocampus is longer than it is wide). A third field is generated across the thickness of hippocampal grey matter (SRLM to outer boundary, or inner to outer, or IO). By default, the IO Laplace field is replaced by an equivolumetric model [6,60], which helps account for the effects of curvature on laminar features (though this replacement can optionally be disabled). We then compute displacement fields for transforming each voxel from native space to the ‘unfolded’ space spanned by these three (AP, PD, and IO) fields, and vice-versa.

Specifically, transformations for going between this unfolded space and native space are defined from Cartesian coordinates (x,y,z) to each Laplace field (AP, PD, and IO) for all hippocampal grey matter voxels. We performed piecewise linear interpolation (griddata from SciPy [61]) to go from each unfolded coordinate (AP, PD, IO) to back to Cartesian coordinates (x,y,z). Rather than map Cartesian coordinates to Laplace coordinates ranging from 0-1 (as in previous work [11]), we scale these gradients to make up a standard rectangular prism with a size of 256x128x16 voxels (dimensions corresponding to AP, PD, and IO, respectively), at a voxel size of 0.15625mm$^3$ isotropic. This reference space is easily changed in the config file if a different unfolded resolution, size, or aspect ratio is desired. Each of these displacement fields is saved as a standard ITK 3D warp file in NIfTI format that can subsequently be applied to NIfTI or GIfTI files.

Unfolding of the dentate gyrus (DG) is introduced in the current work. This is performed with the same methods described above but over the domain of the dentate gyrus rather than all hippocampal grey matter. IO and PD fields are swapped with respect to the rest of the hippocampus reflecting the fact that during its development, the DG breaks from the rest of the cortical mantle and wraps around its terminus (CA4), making it topologically perpendicular to the rest of the hippocampus [62]. Endpoints for the DG are defined within the template shape used in step 3. Due to the thinness of the DG, it is often thinner than one voxel and so Laplace fields cannot easily be generated with the methods used in previous work. Thus, template shape injection is used to define the AP, PD, and IO fields within the DG, meaning that topological alignment between individuals
does not necessarily follow the same Laplacian coordinate framework used in the rest of the hippocampus. Rather, this represents a more traditional volumetric approach to alignment via a template. The unfolded DG was defined by a rectangular prism with a size of 256x32x16, reflecting the fact that it is smaller than the rest of the hippocampus (PD) but still spans the same long (AP) axis.

5. **Subfield definition.** In previous work [20] we performed a highly detailed 3D ground truth segmentation of hippocampal subfields using 3D BigBrain histology [20]. We mapped subfields using our Laplace coordinate framework, which provides implicit, topologically constrained registration between hippocampi. Thus, HippUnfold applies the same subfield boundary definitions to new samples in unfolded space, which are then propagated back to native space. Specifically, reference subfield labels already in unfolded space are warped to each subjects’ native space using the warp files generated in step 4.

6. **GIFTI formatted outputs.** In order to facilitate integration with other popular neuroimaging analysis tools, we have provided outputs in commonly used gifti surface formats in addition to volumetric nifti formats. Standardized unfolded surfaces corresponding to the inner, midthickness and outer surface were generated for one standard unfolded template and propagated to each subjects’ native, folded space using the warp files generated in step 4. Note that unfolded space is mapped to a rectangle rather than a sphere as is typically used in the neocortex, and so surfaces are not fully enclosed. Tessellation of vertices are available in 2K (0.5mm average distance between vertices), 7K (1.0mm between vertices) or 32K versions (32K corresponds to the number of unfolded coordinates used in previous work, or 256x128). Standardized unfolded tessellations were generated by starting with a 512x256 grid with each point connected to its neighbours, making a uniform mesh in unfolded space. Mesh vertices were iteratively removed until vertex distances after transforming to an averaged native space were achieved with the above spacings. In the case of the 32K surfaces, meshes were generated with 256x128 points with no vertices being removed, meaning that vertex distances are uniform in unfolded space but highly variable in folded space.

7. **Morphometry.** Connectome Workbench commands [22,63] are used to extract measures of thickness between inner and outer gifti surfaces, as well as curvature and gyrification along midthickness surfaces. To correct for distortion between native and unfolded space tessellations, gyrification measures were divided by the difference in surface areas between folded and unfolded space. Additional data (for example, fMRI, DWI, or others) can be sampled at each vertex with the code provided in HippUnfold. With the implicit registration provided by unfolded space and the tessellation of these surfaces, such data can readily be compared across hippocampal samples without the need for further registration. These data can be subgrouped according to subfield labels, as in ROI analysis styles, or each vertex can be examined separately as in searchlight or data-driven analysis styles. Alternatively, gradient-based analyses can be applied based on Laplace coordinates and their corresponding surface mesh tessellations (see [64] for example).
Acknowledgements

This work was supported by a Canadian Institutes for Health Research Project Grant (CIHR Grant # 366062) to A.K. and S.K. AK was supported by the Canada Research Chairs program #950-231964, NSERC Discovery Grant #6639, and Canada Foundation for Innovation (CFI)John R. Evans Leaders Fund project #37427, the Canada First Research Excellence Fund, and Brain Canada. J.D. was funded through a Natural Sciences and Engineering Research Council doctoral Canadian Graduate Scholarship (NSERC CGS-D). R.A.M.H was supported by a BrainsCAN postdoctoral fellowship for this work.

Data were 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. Data and/or research tools used in the preparation of this manuscript were obtained from the National Institute of Mental Health (NIMH) Data Archive (NDA). NDA is a collaborative informatics system created by the National Institutes of Health to provide a national resource to support and accelerate research in mental health. Dataset identifier: dx.doi.org/10.15154/1520707. This manuscript reflects the views of the authors and may not reflect the opinions or views of the NIH or of the Submitters submitting original data to NDA. Data was also provided in part by Berron et al., 2017, in their published work “A protocol for manual segmentation of medial temporal lobe subregions in 7 Tesla MRI” [28] which includes MRI images and subfield segmentations.
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