The Human Connectome Project's Neuroimaging Approach

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Abstract

Non-invasive human neuroimaging has yielded many exciting discoveries about the brain. Numerous methodological advances have also occurred, though inertia has slowed their adoption. This paper presents an integrated approach to data acquisition, analysis, and sharing that builds upon recent advances, particularly from the Human Connectome Project (HCP). The "HCP-style" paradigm has seven core tenets: (1) collect multimodal imaging data from many subjects; (2) acquire data at high spatial and temporal resolution; (3) preprocess data to minimize distortions, blurring, and temporal artifacts; (4) represent data using the natural geometry of cortical and subcortical structures; (5) accurately align corresponding brain areas across subjects and studies; (6) analyze data using neurobiologically accurate brain parcellations; and (7) share published data via user-friendly databases. We illustrate the HCP-style paradigm using existing HCP datasets and provide guidance for future research. Widespread adoption of this paradigm should accelerate progress in understanding the brain in health and disease.

Introduction

The Human Connectome Project (HCP) began in 2010 with ~\$40M awarded by the National Institutes of Health (NIH) to two consortia to develop improved neuroimaging methods and to acquire a dataset of unprecedented size and quality for mapping the normal human macro-scale connectome ^{1,2} that is, the long-distance connections between all of the brain's areas. Better maps of the brain's areas and their connections will deepen our understanding of healthy brain function and may improve our ability to understand and treat neurological and psychiatric disorders. The "WU-Minn-Ox" HCP consortium, centered at Washington University, University of Minnesota, and Oxford University, spent its first two years developing the state of the art methods discussed here, and then proceeded to acquire, analyze, and share high quality multimodal neuroimaging, behavioral, and genotype data from 1200 healthy young adult twins and non-twin siblings^{3,4}. The HCP focused primarily on MRI-based imaging modalities to measure brain architecture, function, connectivity, and the regular, topographic, organization of some brain areas; however, magnetoencephalography (MEG) data was also acquired in some

subjects (see Box 2). As the HCP's core activities draw to a close in 2016, its contributions include (i) a set of broadly useful MRI acquisition protocols, consisting of thoroughly tested and optimized pulse sequences and image reconstruction algorithms; (ii) an unprecedented collection of exceptionally high quality, freely shared neuroimaging data; (iii) numerous publicly available neuroimaging software and informatics tools; (iv) a growing number of intriguing discoveries emerging from analyses of HCP data; (v) the emergence of an "HCP-style" paradigm for neuroimaging data acquisition, analysis, and sharing; and (vi) a growing number of HCP-style projects that will study different age ranges and brain disorders.

This article is both a position paper and a primer, promoting awareness and adoption of the integrated HCP-style paradigm along with guidance on its use. We provide an accessible conceptual overview of seven core tenets of the paradigm and summarize several recent discoveries made possible through its application. Additional technical information and discussion is provided in the cited publications and in 18 sections of the Supplementary Information (SI) that expand on specific topics raised in the main text. Investigators interested in practical aspects of implementing the HCP paradigm in their own lab can reference the MRI protocols, HCP software, and HCP Course materials listed below.

The seven tenets of the HCP-style paradigm span the domains of acquisition, analysis, and sharing of MRI-based data: (1) acquiring large amounts of high quality, multi-modal data on as many subjects as feasible to measure architectural, functional, connectional, and topographical information (2) acquiring this MRI data with high spatial, temporal, and angular resolution for structural, functional, and diffusion MRI using cutting-edge accelerated acquisition protocols⁵; (3) minimizing blurring introduced by preprocessing and removing distortions, noise, and temporal artifacts as selectively and completely as possible^{6,7}; (4) representing cortical data (surface vertices) and subcortical data (volume voxels) in a common geometrical framework ("CIFTI grayordinates") that is optimal for each⁶; (5) accurately aligning corresponding brain areas across subjects and studies using "areal features" related to connectivity and architecture^{8,9}; (6) using structurally and functionally relevant brain parcellations (preferably based on multiple modalities) to provide a strong neuroanatomical framework for condensing complex neuroimaging data and enhancing statistical sensitivity and power without blurring across areal boundaries⁸; and (7) routine sharing of extensively analyzed results such as statistical maps¹⁰ (plus raw and preprocessed data when feasible¹¹) together with the code used for the analysis, so that other neuroscientists can make precise comparisons across studies, along with replicating and extending findings.

The HCP-style paradigm builds on recent advances in data acquisition, analysis, and sharing from many neuroimaging labs worldwide, both within and outside the HCP consortia. It benefitted greatly from the NIH's emphasis on methodological development prior to high-throughput data acquisition. The HCP-style paradigm differs from common practices in the human neuroimaging community that emerged during the early years of PET and fMRI (the 'traditional paradigm'), which often involve (i) voxel-based ('volumetric') analysis of data acquired at relatively coarse spatial and temporal resolutions on small groups of subjects; (ii) extensive unconstrained spatial smoothing that attempts to compensate for imperfect cross-subject alignment, improve signal-to-noise ratio (SNR), and satisfy oversimplified statistical assumptions¹²; (iii) not using parcellations at all or using parcellations derived from Brodmann's 2D schematic drawing; and (iv) reporting summary results using 3D volumetric coordinates while keeping the extensively analyzed data private. Relative to the new paradigm, the traditional paradigm has limitations in spatial fidelity, neuroanatomical fundamentals, and the robustness of cross-study comparisons, along lines also noted by Turner^{13,14}.

Though most neuroimaging studies still use the traditional paradigm, important exceptions dating as far back as the 1990's include numerous "retinotopy" studies that use functional MRI (fMRI) data to map the arrangement of visual areas on cortical surface meshes of individual subjects^{15,16}. Thanks in part to this emphasis on surface-based analyses, the visual system is one of the better understood functional systems in the human brain. We anticipate accelerated progress in understanding other brain regions as the new paradigm is applied to them more routinely.

A solid conceptual understanding of MRI acquisition, preprocessing, and analysis (tenets 1-5) is important for applying the HCP-style paradigm to one's own studies, and for understanding the methodological choices made by the HCP and its successors (see below). Many of these choices were driven by fundamental properties of human brain anatomy, MR physics, and human physiology.

Tenets 1 and 2: Multi-modal Data Acquisition Informed by Brain Anatomy and Physiology.

Comprehensive non-invasive brain mapping of many subjects using multi-modal MRI. The HCP acquired ~4 hours of structural, functional, and diffusion images (along with auxiliary 'field map' images to help with image preprocessing). The structural images were T1-weighted (T1w) and T2-weighted (T2w) images (see Supplementary Topic #1 for a discussion of the relationship between these terms and MRI contrast). These images allow non-invasive measurement of architectural properties like myelin content within the cerebral cortex and cortical thickness that match well with corresponding invasive measures 17-19, see also Supplementary Topic #1. The structural images also generate geometrical models of brain anatomy (cortical surface meshes and subcortical segmentations 17,20-27) that are critical for all other analyses. The functional images included task-based fMRI, where the subjects carried out behavioral tasks while in the scanner²⁸, and resting-state scans, where subjects fixated on a crosshair. These functional images enable measurement of functional activation within brain areas, and 'functional connectivity' based on correlations of the fMRI signal between brain areas in the absence of a task or controlled external stimuli. They can also map the internal topographic organization within brain areas, such as the ordered representation of visual space within visual areas using task driven^{15,16,29} or resting-state-based methods (visuotopic organization)⁸. The similarity of these detailed maps across species³⁰ and in comparison with invasive studies in non-human primates, 31,32 confirms that fMRI can accurately capture properties of brain organization. Diffusion MRI uses directionally dependent diffusion of water within white matter to encode average axonal fiber orientations, which can be connected by streamlining tractography algorithms to estimate structural brain connectivity, yielding results that are modestly correlated with invasively measured structural connectivity 33,34. Diffusion MRI also allows estimates of microscopic structural properties of brain tissue, such as the integrity and orderliness of axons (e.g. 35-37. Thus, the HCP MRI data enables brain architecture, function, connectivity, and topography to be estimated non-invasively. The large number of subjects (~1100) imaged by the HCP enables statistically precise estimation of these measures in the healthy young adult population and investigation of the nature and extent of their individual variability, while the focus on twins (and their non-twin siblings) enables assessment of their heritability.

How brain anatomy drives MRI spatial resolution. The cerebral cortex, the seat of human cognition, is a folded sheet, 1.6 - 4mm thick (mean ~2.6 mm), with a surface area of ~1,000 cm²/hemisphere 18,38. High quality structural images (both T1w and T2w) are vital for generating accurate geometrical models upon which all other analyses are based (surface meshes for the cerebral cortex and grey matter ROIs for the subcortical structures 6. For the main HCP, we

acquired these images at an isotropic spatial resolution of 0.7mm (voxel cubes 0.7mm on each side), enabling accurate individual subject maps of cortical myelin content and thickness together with precise models of the inner (white matter) and outer (pial) surfaces. In general, we recommend acquiring T1w and T2w images at 0.8mm isotropic or better (half the minimum thickness of the cortex), in contrast to the single 1mm T1w image traditionally acquired. Such high-resolution images benefit from tightly fitting multi-channel MR receive head coils, such as the Siemens 32-channel head coil used by the HCP.

For fMRI, it is important to acquire images of sufficient spatial resolution to distinguish between cortical gray matter, cerebrospinal fluid (CSF), and white matter, and between opposite banks of cortical folds. The HCP acquired high-resolution fMRI data at 2 mm isotropic (with high resolution fMRI being defined as having voxels smaller than the mean cortical thickness of 2.6 mm, see Supplementary Fig. 1) on its customized 3-tesla (3T) scanner (and at 1.6 mm isotropic at 7T). High-resolution fMRI enables more precise localization of functional signals ⁶ in contrast to resolutions as low as 4 mm that have been traditionally used in fMRI.

Diffusion MRI (dMRI) aims to address a fundamentally resolution-starved problem: mapping connections through brain white matter whose individual axons are mostly <1 micron diameter with MRI voxels that exceed 1 mm isotropic, contain hundreds of thousands of axons 40, and typically include multiple populations of fibers crossing in various directions. Both of the HCP consortia used customized MRI scanners and customized MRI pulse sequences to acquire in vivo human diffusion data of unprecedented resolution 41-46. The 3T WU-Minn HCP scanner uses very strong magnetic field gradients (100 mT/m), approximately double that which commercial scanners could achieve when the project began in 2010. Along with new and optimized dMRI pulse sequences, this enabled 1.25mm isotropic resolution in human dMRI *in vivo* (see Supplementary Topic #2 and Supplementary Figs. 2 and 3 for more details) and 1.05mm isotropic at 7T^{45,47}, in contrast to traditional dMRI with 2-2.5 mm resolution. The recently introduced Siemens Prisma MRI scanner (80 mT/m) developed in part because of the HCP enables the community to acquire similar dMRI data.

Faster, longer imaging for cleaner fMRI data and more robust white matter fiber orientation estimates. Most MRI-based functional imaging relies on the Blood Oxygen Level Dependent (BOLD) contrast mechanism in which increased neural activity indirectly alters the regional deoxygenated hemoglobin content over a time course ('hemodynamic response function', or HRF) of many seconds⁴⁸. Accordingly, one might think that it is sufficient to acquire fMRI images at intervals modestly faster than the HRF (2-3 s Repetition Time, TR, has been typical for whole brain imaging). However, fMRI images are corrupted by many sources of timedependent (temporal) artifacts, including subject movement, physiology, and scanner instability. Many artifacts occur more rapidly than the typical fMRI volume TR and thus are aliased into the signal. High temporal resolution fMRI data (i.e., TR < 1 s) facilitates more selective and effective artifact removal⁷, and improves statistical efficiency, particularly for multi-variate statistics⁴⁹. Recent data also suggest that there may be faster components to the BOLD response^{50,51}, which would be better captured with faster scans. The HCP acquired fMRI data at TR = 0.72 s for 3T and 1 s for 7T using MRI pulse sequences that acquire many brain slices at a time ('multi-band' - also known as 'simultaneous multi slice' - sequences, see Supplementary Topic #3^{5,43,52-54} and Supplementary Fig. 4). For diffusion imaging, using these sequences to reduce the TR allows for more images to be acquired with diffusion weightings at different strengths and along a larger number of unique spatial directions in the same amount of time. The resulting higher angular resolution and effective SNR enables multiple fiber populations to be more robustly identified in each imaging voxel, helping to resolve crossing fiber pathways and improve tractography^{5,44}.

The HCP used 1 hour of scan time for each of task fMRI, resting state fMRI, and diffusion imaging. This enabled seven fMRI tasks to be acquired covering multiple cognitive domains and involving most of the brain²⁸ http://protocols.humanconnectome.org/HCP/3T/task-fMRI-protocol-details.html, along with four 15-min resting state fMRI runs (helping to provide more stable connectivity estimates⁵⁵), and 270 different diffusion measures (90 unique diffusion directions for each of three different diffusion weighting strengths (i.e. "shells"), enabling better fiber orientation and tissue microstructure modeling at high resolution^{56,57}. See Supplementary Topic #4 for particulars on various "HCP-Short" protocols optimized for subjects who are less tolerant of long scans than healthy young adults.

Tenets 3 and 4: Optimizing the Precision of Data Analysis

Aligning all MRI modalities within the subject's physical space. A key early task of preprocessing is to align all MRI modalities so that every image volume is in the subject's undistorted physical space. Head movements occur in the scanner, both between acquisition of each image and during acquisition. Because head movement during T1w and T2w images causes uncorrectable blurring, the HCP acquired at least two pairs of T1w and T2w images, and used all scans that were rated good or excellent in overall quality (11; see Supplementary Topic #5 for further discussion of this issue and potential solutions). Spatial effects of movement during fMRI or dMRI acquisitions can be corrected after the fact using image registration to realign the images; however, movement also induces temporal variations of image intensities that must be corrected through other means described below. Movement between images of different modalities is corrected with advanced registration algorithms that use knowledge of brain tissue boundaries and the expected image intensity gradients across those boundaries to align images⁵⁸.

Ideally, image registration within each subject would entail simple translations or rotations, but MRI images have many forms of spatial distortion that must be unwarped before an image faithfully represents the individual brain's real physical dimensions and thus can be precisely aligned. These distortions all derive from imperfections in the scanner's encoding of physical space using perturbations (linear gradients) of the main magnetic field. All MRI images are slightly distorted by small nonlinearities in the space-encoding gradients that are correctible using information measured and specified by the manufacturer (gradient distortion, see Supplementary Topic #6). In addition, placement of the subject's head in the scanner induces idiosyncratic variations in the magnetic field (field inhomogeneities), particularly near the airfilled spaces in the head (sinuses and ear canals), which can be measured using the auxiliary 'field map' scan mentioned above. Correcting these distortions with a field map is critical for accurately aligning data across modalities within each subject, though traditionally this scan is often omitted, leading to poor alignment in some brain regions. Scanner vendors provide a standard field map scan (based on gradient echoes), but the HCP developed an alternative 'spin echo EPI field map', that is faster, at least as accurate, and also enables correction of fMRI image intensity biases (see Supplementary Topic #7). These field map scans should exactly match the geometry and distortion properties of the functional imaging scans. Diffusion imaging scans have additional 'eddy current' distortions caused by switching on and off the strong magnetic field gradients used to generate diffusion weighting. The HCP developed a new algorithm for correcting eddy current distortions that outperforms previous methods and allows quality control and outlier detection (see Supplementary Topic #8, 59-61). With field maps, advanced distortion correction methods, and anatomically informed registration algorithms, the HCP's spatial preprocessing pipelines⁶ precisely align the T1w, T2w, fMRI, and diffusion data to

the subject's physical space while minimizing interpolation-induced blurring by combining all spatial transformations in a single spline interpolation.

Cleaning time-dependent artifacts from MRI timeseries data. In functional imaging, many processes cause time-dependent intensity fluctuations other than the BOLD-related fluctuations of interest. This structured temporal noise is especially problematic for resting-state functional connectivity because it uses correlations of noisy signals in different parts of the brain, rather than correlation of a predefined noiseless task design with noisy data. Temporal noise can either increase or reduce functional connectivity artifactually and lead to incorrect conclusions⁶² ⁶⁴. Major sources of structured temporal noise include subject movement, subject physiology, and the physics of MRI. It is useful to consider the fMRI timeseries as containing fluctuations arising from 1) the BOLD signals of interest, 2) structured temporal noise, 3) and unstructured random 'Gaussian' temporal noise 11. Temporal cleanup aims to remove temporal noise as selectively as possible while minimally impacting the BOLD signals of interest. The HCP devised a method based on Independent Components Analysis (ICA, 65, which splits the fMRI timeseries into structured components and unstructured noise, representing all of the timeseries fluctuations. The structured components are then categorized by a machine learning classifier (FIX) into signal and noise', and the noise components are removed by regressing them out of the data. For HCP data, the ICA+FIX approach removes spatially specific (i.e. affecting only part of the brain in a particular way) structured noise components and retains spatially specific BOLD signal components (i.e. resting state networks, RSNs) with better than 99% accuracy⁷. In contrast, traditional approaches that include bandpass filtering, WM and CSF regression, and censoring of high-motion frames^{63,64}) are not as selectively at targeting structured noise while preserving BOLD signal. However, a controversy remains about "global" fluctuations in fMRI (see Box 1 and also Supplementary Topic #9 and Supplementary Figs. 5 and 6), as the spatial ICA+FIX approach is not designed to separate these 'global' fluctuations in the average fMRI signal into signal and noise. For dMRI, subject motion and physiology can cause signal loss. Such 'outliers' can be detected using a new approach that builds a model of the diffusion signal from all of the data, allowing artifactual measurements to be identified and replaced with estimates of what the signal should have been (see Supplementary Topic #8, ⁵⁹⁻⁶¹).

Box 1: *Global fMRI Temporal Fluctuations*. A hotly debated topic is how best to handle 'global' signals and artifacts present in fMRI data^{62,63,105-107}. The core phenomenon involves slow and spatially widespread fluctuations in the fMRI timeseries. The average timeseries across the brain includes both neural signal (e.g. the average of all the resting state networks) and structured noise (e.g. physiological noise), and the debate centers on whether the average timeseries should be regressed out of the data to remove the effects of global noise⁶² or whether it should be preserved to avoid distorting neural signals by removing their average. Additive global *noise* will by definition artifactually increase 'full' correlations across the brain (especially problematic if it differs across groups), but removal of a global neural *signal* will artifactually decrease correlations across the brain and, more problematically, distort the pattern of correlations for particularly large/strong networks^{62,108,109}.

What is needed to get past the current impasse is a method that selectively separates global signal and global noise. Though no such method is yet available, we offer several observations about global fMRI fluctuations. 1) In HCP data, the global signal (after ICA+FIX cleanup) is primarily a gray matter signal, with a near zero amplitude in white matter voxels and low-level anti-correlation with CSF (Supplementary Fig. 5). 2) Within grey matter, the mean grey time course has much higher amplitude in sensory regions (e.g. visual, auditory, vestibular, and somatosensory cortex, plus the lateral geniculate

nucleus and other thalamic nuclei) and is lower in the cerebellum and many neocortical cognitive regions (Supplementary Fig. 6). Thus, the global signal is not uniform across the brain, or even across grey matter, and these observations constrain the possible sources of the global fluctuations:

- (i) Physiological effects. Substantial fMRI timeseries variance is attributable to physiological variables, including rate and depth of breathing, heart rate, and end tidal pCO2¹¹⁰⁻¹¹²). Indeed, any physiological process that affects global perfusion pressure of the brain (i.e. the above three, or even yawning) may transiently alter the amount of deoxyhemoglobin in the brain and contribute to global, physiologically artifactual fluctuations of the BOLD signal. Different brain areas and tissue types may be differentially sensitive to such changes in perfusion pressure because of differing metabolic rates, which may explain regional differences in global signal amplitude.
- (ii) Head movements (direct, biophysical). Transient head movements are sometimes followed by global fluctuations in the fMRI signal, leading to suggestions that head movements could directly cause these fluctuations 62,64,106. However, head movements are unlikely to cause global fluctuations directly, given the lack of a plausible biophysical mechanism for how head movement might induce tissue-specific and brain-area-specific effects on the fMRI timeseries, let alone globally uniform changes in signal intensity short of complete head coil exit and reentry (See Supplementary Topic #9).
- (iii) Head movements (indirect, neural). Head movements may induce neural responses in somatosensory, vestibular, and/or visual systems that may in turn modulate wider brain activity in a pattern very consistent with that mentioned above.
- (iv) Widespread/High amplitude RSNs, average of the RSNs. As discussed above, a portion of the global fluctuations will represent the average of the RSNs with particular weighting to more widespread or stronger networks.

It will be important to separate global physiological effects from spatially widespread neural effects (using either extrinsic physiological monitoring or intrinsic fMRI-data-driven means yet to be developed). Short of a complete solution to the problem of global fluctuations, we note that some approaches to analysis (such as the use of partial correlation when estimating functional connectivity, and multi-variate measures in general) are much more immune to the effects of global fluctuations than others (e.g., full correlation measures and uni-variate measures in general)¹¹³.

The traditional approach to reducing unstructured Gaussian noise involves spatial and temporal smoothing, which unfortunately affect both signal and noise unselectively. A common problem in traditional neuroimaging is that smoothing mixes the grey matter functional signals of a given brain area with signals from white matter, CSF, and nearby brain areas, markedly reducing the spatial fidelity of neuroimaging data. Parcellation (tenet 6) enables averaging of data without mixing signal across different tissue types or between different brain areas in so far as the parcellation is neurobiologically accurate, and acts as a neuroanatomically informed method of nonlinear smoothing. Another option is to selectively smooth unstructured noise more than structured BOLD signal (see Supplementary Topic #10). This enables reduction of certain artifacts (see Supplementary Fig. 7) and enhancement of SNR without concomitant spatial blurring and without invoking a brain parcellation.

Preserving precision from voxels to CIFTI "grayordinates." It is vital that the preprocessing and analysis techniques used on high resolution HCP-style data maintain this hard-won spatial resolution. Besides removing distortions and accurately aligning data across modalities within each subject, the data must be put into an analysis framework that enables accurate crosssubject and cross-study comparisons (tenet 5 below). The traditional neuroimaging paradigm represents spatial data in a regular 3D array of voxels (an image volume) that reflects the output of the MRI image reconstruction process. Because the cerebral cortex has a sheet-like folded geometry, it is advantageous to visualize and analyze cortical data on surface mesh vertices. instead of volume voxels 15,25,26,38. However, subcortical grey matter structures are mostly globular in shape and remain best represented using volume voxels. To represent the inherently dual geometry of the brain, the HCP developed the CIFTI file format⁶, which integrates surface vertices and subcortical voxels into a single file and represents each major brain gray matter structure using the appropriate geometry (see Figure 1). CIFTI gray matter vertices and voxels are called 'grayordinates.' Analyses focusing only on grayordinates (e.g., fMRI) dramatically reduce file sizes and computational burdens (see Figure 1 legend). For dMRI, surface representations allow more accurate anatomical constraints in tractography (e.g., the white surface can be used as the destination for tractography). The final common outputs of the HCP preprocessing and analysis

preprocessing and analysis pipelines are datasets in the CIFTI standard space⁶. CIFTI is natively supported by Connectome Workbench, and other tools are currently implementing CIFTI compatibility (e.g., FSL, FreeSurfer, FieldTrip, nibabel). In particular, FSL's PALM software enables permutation-based statistical inference on CIFTI datasets (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/PALM ⁶⁶). The CIFTI grayordinates

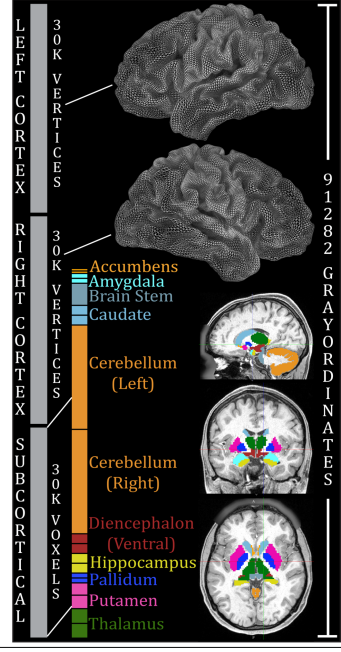


Figure 1. Combined representation of cortical surface vertices and subcortical voxels in the CIFTI grayordinates standard space. Left and right cerebral cortices contribute about 30k surface vertices each (fewer than the 32k vertex standard meshes for each hemisphere, because the non-cortical medial wall is not included). Additionally, 19 subcortical grey matter structures combine to contribute about 30k volume voxels. In total, there are 91,282 grayordinates corresponding to all of the grey matter sampled at a 2 mm average vertex spacing on the surface and as 2 mm voxels subcortically. The HCP's minimal preprocessing pipelines ensure that each subject has 91,282 aligned grayordinates, thereby facilitating cross-subject comparisons of data within this coordinate system. This entails more than a two-fold reduction in file size relative to the >200,000 voxels needed for an equivalent 2 mm isotropic volume representation. For the 1.6mm 7T HCP data, we have developed a 1.6mm standard grayordinates space with 170,494 grayordinates using "59k" surface meshes. Reproduced with permission from Reference 6.

analysis framework allows each brain structure to be registered using the approach best suited to it (see Tenet 5).

Tenet 5: Accurately Aligning Brain Areas Across Subjects and Studies.

Cross-subject comparisons. Analyses that combine neuroimaging data obtained from different individuals benefit greatly from accurately aligning corresponding brain areas, so that like is compared with like ¹⁴, Alignment of subcortical and cortical structures is best achieved through differing means. Subcortical structures are less variable in size, shape, and position across subjects, and reasonable alignment is achieved by nonlinear 3D registration to a standard template, as in the traditional paradigm. The HCP currently uses FSL's FNIRT for nonlinear registration of T1w images to MNI space, but registrations that also incorporate dMRI fiber orientations in the white matter may improve alignment of subcortical white matter fiber tracts (See Supplementary Topic #11^{67,68}).

For human cerebral cortex, the complexity and variability of folding patterns and of areas relative to folds^{38,69-71} poses a profound challenge that is an ill-posed problem for nonlinear 3D registration. Though alternative approaches such as hyperalignment (where alignment is not spatially constrained) are useful in some contexts, many neuroscientific questions are best addressed by aligning subjects to a common space. Registering cortical surfaces rather than brain volumes enables better alignment because accurate cortical segmentation of each subject reduces the dimensionality of the registration problem from 3D to 2D—alignment need only occur tangential to the cortical sheet as opposed to a 3D alignment of the folded sheet itself^{26,38,72-76}.

Though surface-based registration of the cerebral cortex to an atlas is not a part of the traditional paradigm, it has been adopted by a growing number of investigators. Most use surface registration based on cortical folding patterns^{26,38,72-76}, which achieves 'geographic correspondence' in regions where folding patterns are consistent across individuals and 'functional correspondence' in regions where areal boundaries are consistently correlated with folding. Because cortical folding patterns are highly variable in many regions of cortex and their relationships to cortical areas are similarly variable, folding patterns alone are not enough to achieve alignment of (functional) cortical areas across most of cerebral cortex. Registration needs instead to be driven by features more closely tied to cortical areas, such as architecture, function, connectivity, and topography⁷⁷. Such areal-feature-based cortical surface registration can dramatically improve the sharpness of group average maps and enhance group statistics, as it largely compensates for differences across subjects in cortical areal size, shape, and position. Indeed, alignment based on T1w/T2w myelin maps, resting state network maps, and visuotopic connectivity maps vields sharper task fMRI contrast maps with higher statistics, even though the task fMRI data were not used in the registration (Figure 2)8,9. Areal-feature-based aligned data was recently released by the HCP (dubbed 'MSMAll' because it uses the Multimodal Surface Matching algorithm, and all of the useful modalities listed above). We believe that many studies will benefit from using areal-feature-based surface registration as a starting point, even if they choose to make final comparisons across subjects using other means (e.g. hyperalignment⁷⁸, or individual subject parcellation (see below and ref.⁸).

Cross-study comparisons. Traditionally, cross-study comparisons based on 3D stereotaxic coordinates were assumed to align if everyone used a common standard volume space (e.g., MNI). These 3D coordinates were typically derived from thresholded statistical maps of the 'significant' activations (or connectivity seed locations). Though such standard space coordinates allow for gross comparisons across studies at the gyral level (e.g., ref.⁷⁹, questions

of whether a particular cortical area was activated in particular task contrast typically reveal clouds of points whose distribution may be suggestive, but often inconclusive (e.g., ref. 19). Stereotaxic coordinates have high spatial uncertainty (relatively small changes in the thresholded statistical maps used to produce them can have large effects on the resulting coordinate locations, see Supplementary Topic #12 and Supplementary Fig. 8), and they do not contain information about the spatial extent or boundaries of activation. Conclusive cross-study comparisons require assessment of the overlap of activated regions and whether their boundaries align or not. Cross-study comparisons also rely on the absence of any "drift" in alignment between the studies (see Supplementary Topic #13 and Supplementary Fig. 9). When drift is eliminated and boundaries are compared, direct and conclusive assessments of cross-study agreement are possible (see Supplementary Fig. 10, also Figure 6).

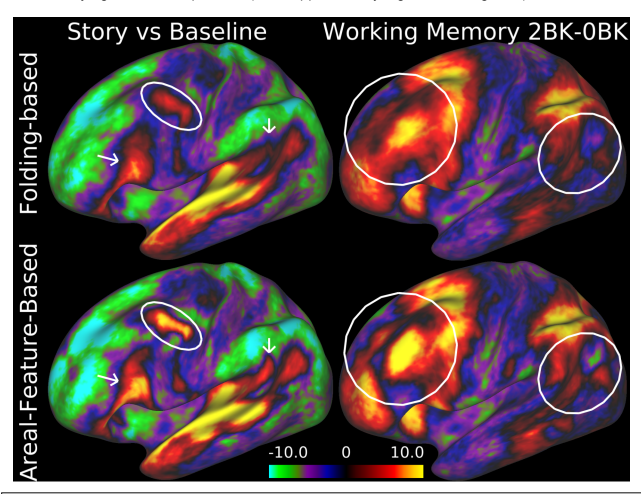


Figure 2. Improved intersubject registration using information based on areal features in addition to cortical folding. The top row shows group task-fMRI z-stat maps ("Story vs Baseline" contrast from the language task on the left and the two-back vs zero-back contrast in the working memory task on the right) from 120 Q1 and Q2 HCP subjects after intersubject registration using the Multimodal Surface Matching (MSM) method constrained only by folding (FreeSurfer's 'sulc' maps). Bottom row shows sharper group task-fMRI maps and higher z-statistics when using resting-state networks (RSN) along with myelin maps to constrain the registration. For the data in this figure and figure 6, subject recruitment procedures and informed consent forms, including consent to share de-identified data, were approved by the Washington University institutional review board. Data at http://balsa.wustl.edu/97V4.

Tenet 6: Neuroanatomically Accurate Maps of Brain Areas.

Accurately subdividing the brain into its constituent parts (areas or parcels) has been a holy grail for brain cartographers and other neuroanatomists for more than a century⁸⁰⁻⁸². The goal is to reliably identify parcels that are distinct from one another in architecture, function, connectivity, and/or topographic organization and are consistently identifiable in most or all individuals. For neuroimaging, parcellation serves to: (i) clarify 'where we are' in the brain when describing results¹⁴; (ii) facilitate comparisons across individuals and studies; and (iii) reduce the dimensionality of complex and noisy datasets while respecting neuroanatomically distinct boundaries. Parcellation can dramatically improve statistical sensitivity and power by averaging across neuroanatomically similar units of the brain, enhancing SNR and reducing the vast number of statistical comparisons often carried out in neuroimaging (circumventing problematic corrections for multiple comparisons that have affected a substantial fraction of the neuroimaging literature ¹²). As noted above, parcellation is an attractive alternative to traditional spatial smoothing for analyses that are interested in effects at the areal level (in practice, most studies that provide a standard space coordinate table).

A cortical areal parcellation should ideally have four key qualities: (i) It should be based on many (hundreds of) well aligned subjects so that it represents the typical areal arrangement in the studied population (e.g., healthy young adults for the HCP). (ii) It should reflect complementary and converging evidence from multiple modalities across the whole cerebral neocortex for completeness and for higher confidence in boundaries. Most parcellation efforts to date have relied instead on information from a single modality, such as architecture^{70,83}, retinotopy^{16,29}, or resting-state fMRI⁸⁴⁻⁸⁶, and often do not cover the whole hemisphere. (iii) It should reflect existing terminology for areas previously reported in the neuroanatomical literature and a rational terminology for newly defined areas. (iv) It should be possible to automatically replicate the parcellation in individual subjects' datasets based on multi-modal areal fingerprints – even in subjects with atypical areal arrangements or those from future studies. The recently reported "HCP_MMP1.0" (HCP MultiModal Parcellation, version 1.0) meets these criteria⁸, but we expect this will not be the final word on cortical parcellation. Also subcortical structures and the cerebellum need analogous multi-modal parcellations.

The HCP_MMP1.0 parcellation (Figure 3) contains 180 distinct areas per hemisphere, symmetrically arranged across the two hemispheres. 83 areas corresponded reasonably well with areas defined in one or more prior studies and were assigned the same names, while 97 were assigned new names. The group average parcellation was used to train a machine-learning classifier (a multi-layer perceptron⁸⁷) to enable a fully automated process for identifying areas in individual subjects based on each area's multi-modal fingerprint, even in individuals having atypical topological arrangements of areas (Supplementary Fig. 11).

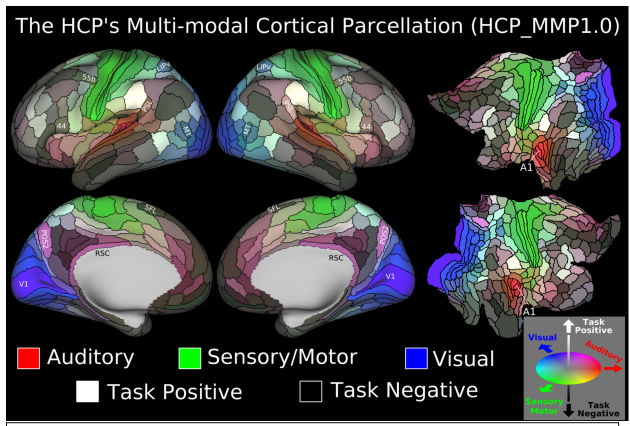


Figure 3. The HCP_MMP1.0 (HCP Multi-Modal Parcellation, v 1.0). Each panel shows 180 cortical areas delineated and identified in the left or right hemisphere, displayed on an inflated or flattened cortical surface. Black outlines indicate areal borders. Colors indicate to what extent the areas are associated in the resting state to auditory (red), somatosensory (green), visual (blue), task positive (towards white), or task negative (towards black) systems. The legend on the bottom right illustrates the 3D color space used in the figure. Reprinted from ref⁸. Data at http://balsa.wustl.edu/WN56.

More generally, the classifier can reliably detect cortical areas in individual subjects without using the HCP's task fMRI data (using only myelin maps, thickness maps, resting state network maps, and resting state visuotopic maps), enabling generalization of the approach to many non-HCP neuroimaging datasets. Individual investigators wishing to apply the HCP Pipelines, areal-feature-based registration, and trained areal classifier to their own individual subjects, need only acquire T1w, T2w, field map, and fMRI data as described above, as even non-resting state fMRI data include similar patterns of connectivity⁸⁸. The population-based parcellation can also be applied directly to registered individual subjects, though it will not be accurate where individual subjects deviate from the typical layout of areas. The HCP's multimodal 'hard' parcellation v1.0 provides a neuroanatomical foundation for future neuroimaging analyses, though other 'weighted' approaches provide valuable complementary information as well (see Supplementary Topic #14). It's worth noting that FreeSurfer uses an algorithm to automatically label gyri and sulci in individual subjects based on manually generated training labels that is similar in spirit to our areal classifier⁸⁹.

Tenet 7: Routine Data Sharing and Advanced Informatics for Data Management

Data sharing for individual laboratories. Data may be publicly shared at many levels, including extensively processed, preprocessed, and unprocessed stages together with the code used for analysis. Data sharing is increasingly mandated by journals and funding agencies. Although the HCP has shared its data using an advanced informatics infrastructure described below, most individual investigators only publish their results in journal articles, as the additional time and informatics skills required for more expansive data sharing have made it impractical. Generating and editing manuscript figures is typically a tedious process involving multiple software applications that neither work efficiently together nor aid in preparing data for post-publication sharing. Routine sharing of extensively processed data would be expedited by a streamlined process for generating publication-ready figures that is integrated with a service for easy post-publication sharing of the associated data.

An integrated approach to data visualization, figure generation, and data sharing is now available. Connectome Workbench (a freely available general-purpose neuroimaging visualization platform) provides a flexible interface for displaying rich multi-modal neuroimaging datasets. "Workbench" enables visualization of various data types overlaid on surfaces or volumes (NIFTI, GIFTI, or CIFTI files). Multi-panel montage figures can be generated by combining web browser-like tabs that can also be annotated (e.g., text, symbols, and imported image panels) directly in Workbench. For example, Figure 4 is one of 42 published figures in one study 8 exported from Workbench without further editing. Importantly, Workbench allows such complex displays to be saved as "scenes" that can be quickly re-opened, edited, re-saved, and exported as publication-ready images. Data from such figures can be shared by uploading the scene file to the BALSA database (http://balsa.wustl.edu; directly from Workbench, together with basic information about the publication. Scene-specific URLs enable one-click linking from a published figure to the corresponding page in BALSA (see Figure legends). Adoption of similar strategies by other neuroimaging software and databases would accelerate sharing of extensively analyzed data and facilitate more accurate and diverse comparisons across studies.

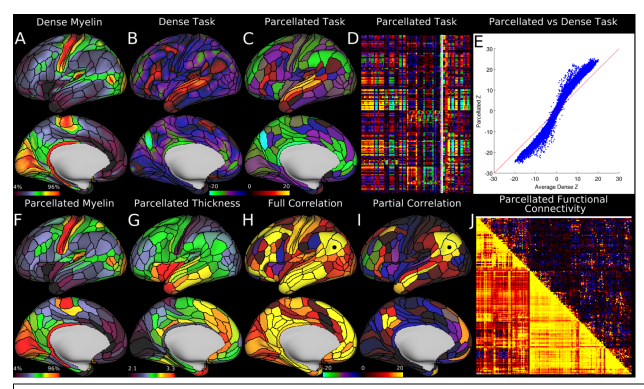


Figure 4. Example parcellated analyses run using HCP data and the HCP MMP1.0 cortical parcellation. Panels A and F, showing dense and parcellated myelin maps respectively, are very similar despite a dramatic dimensionality reduction. Panel G shows a map of parcellated cortical thickness (in millimeters; corrected for folding by regressing out mean surface curvature). Panels B and C show example dense and parcellated task fMRI analysis (LANGUAGE Story vs Baseline). Panel D shows the entire HCP task fMRI battery's Z statistics (86 contrasts; 47 distinct and 39 sign-reversed versions) analyzed in parcellated form and displayed as a matrix (rows are parcels, columns are contrasts, white outline indicates the map displayed in Panel C). Panel E demonstrates a major improvement in Z statistics from fitting a task design on parcellated data instead of fitting it on dense data, and then parcellating afterwards. Panels H and I show parcellated functional connectivity maps on the brain (seeded from area PGi, black dot). In both cases, the task negative (default mode) network is apparent. Panel J shows a parcellated connectome matrix view with the full correlation connectome below the diagonal and the partial correlation connectome above the diagonal (white line shows the displayed partial correlation brain map). The figure matches the Connectome Workbench scene available on the BALSA database (http://balsa.wustl.edu/RG0x). Notably, Panel E was generated in matlab and saved as a PNG and then loaded in to Workbench as its own tab. Reprinted from ref⁸.

Large-Scale Data Sharing and an Advanced Informatics Infrastructure. Well designed data management tools and processes are vital for the successful collection and analysis of high quality HCP-Style data on a large scale. ConnectomeDB, the primary database for HCP data, was designed for user-friendly sharing of unprocessed and preprocessed data from a large number of individual subjects, which can be selectively downloaded using a variety of flexible criteria https://db.humanconnectome.org 11,90. ConnectomeDB is based on the XNAT informatics platform (http://www.xnat.org 91, which manages imaging data through a structured workflow from acquisition through quality control and automated processing 90. It also provides data entry and import tools for task and biometric data collected during image acquisition as well as behavioral and other non-imaging data. ConnectomeDB also serves as the data management platform for the Lifespan Connectome projects, has been deployed by some Connectomes Related to Human Disease projects (see below), and can benefit other groups carrying out

large-scale HCP-style studies (source code at http://www.xnat.org/download/). Given the complexity of the HCP project and the massive size of the HCP dataset, documentation and distribution required a multi-faceted approach (Box 2).

BOX 2 —Accessing and using HCP data, software, and protocols.

Overview of the HCP's Data Sharing. ConnectomeDB (http://db.humanconnectome.org) is the primary repository of HCP imaging, behavioral, and demographic data. 3T structural, functional, and diffusion imaging data for ~1100 subjects is shared in unprocessed, minimally preprocessed (MPP), and more extensively processed forms and staged over 4 major releases (Q1, Q1-Q3, S500, S900) with the final S1100 release slated for summer, 2016. 7T data acquired on 180 subjects has been released in part with the rest to follow shortly. MEG data acquired from 95 subjects gives a window on fast temporal dynamics of the brain and was released in the fall of 2015¹¹⁴. Importantly, the S900 release and the forthcoming S1100 release include the improved intersubject registration provided by "MSMAII". The HCP_MMP1.0 parcellation is available in BALSA (http://balsa.wustl.edu), and individual subject parcellations for each HCP subject will be released on ConnectomeDB in the future (anticipated fall, 2016).

HCP Data Is Much Larger than that of Past Studies. With the high spatial and temporal resolution of HCP data, the dataset is at least an order of magnitude larger than widely used open access neuroimaging data sets like ADNI¹¹⁵. For a single subject, the compressed NIFTI-formatted unprocessed data is around 10 GB, and the preprocessed data is nearly 30GB.

HCP Data Is Widely Used via Multiple Modes of Data Access. To date, >5,200 investigators have agreed to HCP Open Access Data Use Terms (~520 to Restricted Access terms for accessing family structure and other sensitive data). Users can (i) access ConnectomeDB directly to download 'packages' for individual subjects, user-selected subject groups, or HCP-specified groups; (ii) purchase "Connectome-in-a-Box" for the cost of the hard drives, which can be shared by investigators at a given institution; and (iii) access data via the Amazon cloud for processing on the cloud (or for download). Data downloaded directly from ConnectomeDB exceeds 5400 terabytes (TB), with an additional 2000 TB transferred via hard drives (Connectome-in-a-Box) and the Amazon cloud.

Data documentation. The richness and complexity of the HCP data requires extensive documentation for users to understand what is available, how the datasets are organized, and how they were processed, including quality control measures. Available resources include (i) a *Reference Manual* associated with each data release (http://humanconnectome.org/documentation); (ii) *HCP course materials* (lectures, tutorials, and associated data) available at http://humanconnectome.org/courses/); (iii) publications, e.g. for database organization http://humanconnectome.org/courses/); (iii) https://wiki.humanconnectome.org/display/PublicData/Home), which provides additional documentation, FAQs, and updates (including known issues and planned fixes).

Software Sharing. Also important to the replicability of neuroimaging studies is the sharing of the software used for analysis. Scripts for HCP pipelines are available on GitHub (https://github.com/Washington-University/Pipelines/releases), and their usage is described in the HCP course materials. This includes pipelines for MEG as well as MRI data. Connectome Workbench is available as binaries and source code

(http://www.humanconnectome.org/software/get-connectome-workbench.html, on GitHub, and in http://neuro.debian.net) along with tutorials. Additionally many of the labs that make up the HCP consortia share their software on either their own websites or the HCP website.

MRI protocols. The MRI protocols used for the main HCP 3T and 7T scans are available at http://protocols.humanconnectome.org/HCP/; HCP-style pulse sequences have been widely distributed (http://www.cmrr.umn.edu/multiband/index.shtml). The HCP-Short protocols used for the HCP Lifespan (see below) are at http://protocols.humanconnectome.org/lifespan/

HCP-Style Follow Up Studies. The original (healthy young adult) HCP will be succeeded by several large-scale imaging projects that use "HCP-style" data acquisition, analysis, and sharing to study different populations in health and disease. Data sharing for many of these projects will be managed by the Connectome Coordination Facility (www.humanconnectome.org/ccf), an NIH-funded resource devoted to supporting data sharing from the Connectomes Related to Human Disease (CRHD) projects (11 currently funded, with more anticipated in 2017) plus three Lifespan HCP projects (see Figure 5). The CCF is an extension of the ConnectomeDB infrastructure and will provide a common interface for searching within and across projects. In addition, it is developing common acquisition protocols and maintaining the HCP's processing pipelines to maximize cross-study comparisons and analysis, and partnering with other connectomics studies such as the Adolescent Brain Cognitive Development (ABCD) study and UK Developing HCP (dHCP http://www.developingconnectome.org/) to encourage harmonization of acquisition and analysis methods. Versions of the HCP-Short Protocol (see Supplementary Topic #4) have been adapted for a variety of vendors and scanner models, with the goal of producing images of similar spatial and temporal resolution.

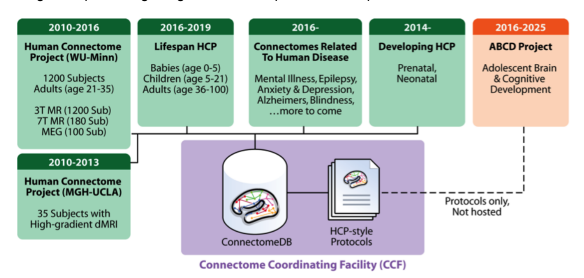


Figure 5. Current and future projects that use HCP-style data acquisition, analysis, and sharing. Projects shown in green will use the Connectome Coordination Facility (CCF, in purple) as their primary mode of data sharing.

Concluding comments

The NIH had two interrelated core goals in launching the HCP: 1) to generate a publicly shared, widely used repository of high quality multi-modal neuroimaging data suitable for mapping the human connectome, and 2) to advance the methods used in neuroimaging research.

More than 140 studies have acknowledged the use of HCP data, and for half of these the authors were largely or entirely outside the HCP consortium (https://wiki.humanconnectome.org/x/YQDJAw). These studies address a wide range of questions using various imaging modalities. Five brief examples highlight some of this progress. Smith et al., 92 identified a 'positive-negative' phenotypic axis that covaries with differences in the strength of functional connectivity in a subset of brain regions that are generally implicated in higher cognitive functions. Yeo et al⁹³ investigated the overlap and segregation of the brain's functional networks, finding that association regions tend to overlap with at least two networks whereas somatosensory and visual regions are more isolated. Wang et al. ⁹⁴ investigated functional network parcellations at the individual level using HCP data. Hawrylycz et al. 95 demonstrated correlations between gene expression patterns in postmortem human cortex and HCP-based in vivo functional connectivity. Tavor et al. 96 showed that a model relating task-independent (rfMRI) measurements to task activity can accurately predict task activation maps for unseen subjects, suggesting a coupling between brain connectivity and function at the level of individual subjects. After release of the final "S1100" imaging and genotyping data (summer, 2016) and the individual-subject cortical parcellations (projected fall, 2016), we anticipate that a growing fraction of studies using HCP data will use the full set of available subjects, better-aligned data, multimodal parcellations of group average and individual subjects, and family structure (e.g., twins). Capitalizing on the full richness of the HCP data offers the best prospects for robust and neurobiologically grounded findings. Even with large amounts of high quality data, it remains extremely important to use caution in interpreting complex neuroimaging datasets and the indirect inferences related to brain connectivity available from neuroimaging. Neither functional connectivity based on rfMRI nor structural connectivity based on dMRI tractography provides good quantitative estimates of the actual strength of direct anatomical connectivity between areas or regions. This reflects a variety of methodological biases and other limitations that have become better understood in recent years ^{2,34,97} see Supplementary Topic #15), but remain serious limitations even with the advances provided by the HCP-style paradigm. Research continues to further validate non-invasive connectivity methods and maximize their accuracy.

The HCP-style paradigm brings together a wide range of methodological advances, many provided by investigators or projects outside the HCP, and others reflecting direct HCP innovations. New MRI scanner technology (Siemens Prisma) and pulse sequence technology (multi-band) were developed in part through the HCP, enabling imaging of finer details faster. Many improvements in image distortion correction, registration, and data-driven image denoising enable selective removal of artifacts while preserving the precision of the signals of interest. The CIFTI grayordinates analysis framework together with areal-feature-based surface registration largely compensates for individual variability across subjects without causing or resorting to blurring of the data. A multi-modal cortical parcellation provides a new neuroanatomical foundation for studies of the human cerebral cortex and is a valuable prerequisite for generating area-to-area connectomes in individuals and group averages. Intuitive, user-friendly data visualization software together with integrated neuroimaging databases and software pipelines enable easier data processing and sharing.

The overarching objectives of the HCP-style paradigm are (i) to acquire and analyze neuroimaging data as accurately as possible in each individual from their original brain imaging voxels to their individually mapped brain areas; (ii) to use precise inter-subject alignment while avoiding smoothing to minimize blurring across brain areal boundaries when comparing results across individuals, groups, and studies; and (iii) to routinely share neuroimaging data in a way that facilitates conclusive cross-study comparisons. Unlike traditional volume-based approaches that often result in statistically significant 'blobs' or 3D coordinates of unclear neuroanatomical identity, the HCP-style paradigm aims to remain as faithful as possible to the underlying neuroanatomy, even in the face of the remarkable individual variability of human brains. Indeed, the gains in spatial sharpness and clarity provided by the HCP-style paradigm (see Figure 6) are qualitatively analogous to those made in astronomy after the introduction of adaptive optics and space telescopes to overcome the atmospheric blurring that plagues conventional groundbased telescopes 98. We provide concrete, generally applicable guidelines for individual labs wishing to embark on a new study using the HCP-style paradigm, though of course not all recommendations will fit every study, and methods will continue to improve (Supplementary Topic #16).

We propose that the diverse technological and conceptual advances in imaging integrated by (and developed within) the HCP meet the criteria of a new paradigm for neuroimaging research⁹⁹. The "HCP-style" paradigm has a differing set of assumptions (e.g., blurriness in brain images is not a 'fact of life' but an avoidable artifact of traditional methods and that neuroanatomical localization is critical to understanding the brain). It invokes new terminology, and provides substantial new capabilities for further research. It is also disruptive to 'business as usual' as practiced by the majority of neuroimaging investigators, and legacy data may not

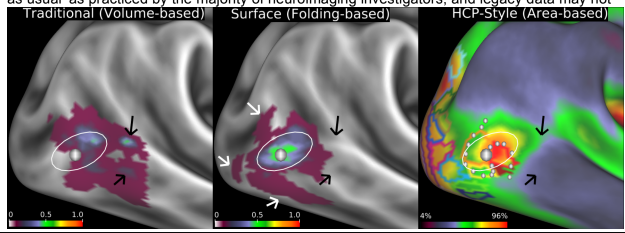


Figure 6. Fidelity of localizing area MT+ (architectonic area hOc5) when mapped to the cortical surface by different methods. Left: volume-based mapping of probabilistic cytoarchitectonic area hOc5¹⁰³ from 10 postmortem subjects mapped to a cortical atlas surface. Black arrows point to locations where the volumebased mapping spreads across gyral and sulcal folds. Center: Surface-based registration of hOc5 from the same 10 subjects mapped to individual surface reconstructions, then to a surface-based atlas using FreeSurfer's folding-based surface registration method⁷³. White arrows identify 'outlier' hOc5 from individual subjects that are not well aligned to the FreeSurfer group average owing to imperfect correspondence between areal boundaries and sulcal folds. Left and center panels adapted from ref. 38. The white oval is in the same location across all panels showing how the volume-based alignment drifts away from the surfacebased alignments, in addition to having substantially lower cross-subject overlap, Right: Group average cortical myelin map (from 196 HCP subjects) with a yellow/orange/red hotspot indicating the MT+ complex and retinotopic areal Maximum Probability Maps (MPMs) (from 12 non-HCP subjects) both registered using arealfeature-based surface registration and 'de-drifted' (see Supplementary Topic #13) (adapted from ref. 104). As shown in Supplementary Fig. 10, the white dots represent a contour along which functional connectivity rapidly changes that aligns with the border between retinotopic areas MT and pMST from a separate study. This is an example of the more conclusive cross-study, cross-modal boundary comparisons made possible by the HCP-style analysis paradigm. Data at http://balsa.wustl.edu/kNpD.

be easily comparable with data from the HCP-style paradigm while maintaining the new paradigm's improved standards of spatial fidelity and neuroanatomical localization (Supplementary Topic #17, 18). Thus, moving to the new paradigm should not be undertaken lightly. That said, we believe the benefits decidedly outweigh the costs, as the new paradigm opens up a variety of novel analysis strategies that will likely accelerate our understanding of human brain structure and function and may even prove useful in a clinical setting. Instead of relying on rough evidence of spatial proximity as in the old paradigm, we aim in the new paradigm to precisely compare the overlap and boundaries of results from differing studies to see if they likely refer to the same brain areas or not. Additionally, we are able to separate differences in brain function or connectivity much more precisely from differences in brain areal size, shape, and position. We can investigate the significance of variations in individual subject areal topologies from the population's typical topology. Finally, the HCP-style paradigm is better positioned to exploit further improvements in brain image acquisition that will enable even finer grained study of the brain at the level of cortical layers or columns, as well as finer subcortical organization 100-102. Nonetheless, the challenges to neuroimaging remain daunting, and we hope that new technologies and ideas will lead to further paradigmatic advances in the future.

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Supplementary Information and Figures for The Human Connectome Project's neuroimaging approach

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The Supplementary Information is divided into three main parts. Part I (Supplementary Notes) includes 18 brief text notes, each expanding on a specific technical or discussion issue in the order that it was raised in the main text. Supplementary References follow in Part II. Part III (Supplementary Figures) includes 12 figures that illustrate important points raised in the main text or supplementary notes.

I. Supplementary Notes

1. What do the terms T1-weighted and T2-weighted mean in terms of MRI and myelin contrast?

Students learning about brain imaging (for research or clinical purposes) are exposed early on to the distinction between "T1-weighted (T1w)" and "T2-weighted (T2w)" images. T1w images have an intensity profile of white matter (WM) > grey matter (GM) > cerebrospinal fluid (CSF). In contrast, T2w images have the opposite intensity profile (CSF > GM > WM). Although these terms were originally based on the types of MR contrast in the images from simple gradient echo or spin echo sequences, they have come to refer more to the direction of contrast in MRI images, given the increasing complexity of the MR pulse sequences currently in use for 3D high resolution brain imaging (and scanner manufacturers, radiologists, and many brain imagers use the terms in this way). Two particular 3D sequences in current widespread use are 3D MPRAGE (magnetization prepared rapid gradient echo)¹ and 3D SPACE (single slab variable flip angle turbo/fast spin echo)². However, none of these terms is completely accurate in describing the MR contrast mechanisms that create each image. The T1w contrast (WM>GM>CSF) in an MPRAGE image is dominated by the T1 relaxation mechanism but also has T2* and proton density (PD) components³. On the other hand, the contrast in a SPACE image can be manipulated via the variable flip angle scheme for the refocusing RF pulse train⁴ to generate T2w⁵, T1w⁶, or proton density weighted (PDw) images. The T2w contrast (CSF>GM>WM), which contains both T1 and T2 relaxation mechanisms⁵, is the most commonly used contrast for 3D SPACE and was the choice in the HCP's T2w SPACE protocol. Given the complexity surrounding the MR contrast mechanisms in 3D structural acquisitions, we refer to the HCP MPRAGE and SPACE acquisitions simply as T1w and T2w images, consistent with

their intensity profiles (albeit with no intention to imply that just a single underlying contrast mechanism is present). We also note that in our experience the best results for T1w and T2w images are achieved using the 3D MPRAGE and SPACE sequences at 0.8 mm isotropic resolution or better, half the thickness of the thinnest cortex.

T1w and T2w images can be used to compute a T1w/T2w ratio in order to map myelin content in cortical grey matter⁷. As mentioned above, T1, T2, T2*, and PD all contribute to some degree to the contrast in the T1w and T2w images. The lipids in myelin affect T1 contrast⁸, and iron, which colocalizes with myelin particularly in cortical grey matter⁹, affects T2* contrast⁸. Proton density also appears to contribute to grey/white contrast and may also contribute to intracortical myelin contrast¹⁰. Thus, many types of MR contrast also contribute to the T1w/T2w ratio and hence to the grey matter myelin maps produced by this ratio. We treat these maps as an in vivo "myelin stain," rather than a quantitative measure of MR properties, and their validation is primarily neuroanatomical, rather than biophysical (i.e. they correlate strongly with known patterns of myelination within the grey matter⁷). Though there has been debate as to whether the T1w/T2w ratio is the best approach for in vivo myelin mapping (as opposed to alternatives such as quantitative T1 images¹⁰⁻¹⁴, quantitative magnetization transfer images, quantitative T2* images¹⁵, T1w/T2*w ratio images¹⁶, myelin water fraction images, etc), we are unaware of an approach that produces as high a contrast-to-noise ratio (CNR) per unit acquisition time and field strength, enabling spatial resolution to be maximized within a fixed scan time (< 13 min combined between the T1w and T2w acquisitions at 3T and 0.8mm isotropic). Also, T1w and T2w images are both very useful for generating accurate white matter and pial surface models and subcortical segmentations¹⁷. Thus, we recommend that most studies acquire both images (and then the T1w/T2w myelin maps come 'for free'), but studies focusing on quantification may elect to use some of their available scan time to acquire quantitative MR parameter maps. A FLAIR type acquisition (also based on the 3D SPACE sequence with an inversion recovery preparation) could be used in place of the T2w SPACE acquisition if FLAIR contrast is important for other reasons. However, such FLAIR images do not have as much myelin contrast or as high SNR as the T2w images mentioned before, so the myelin maps they produce are noisier.

2. Why is the HCP's Diffusion Protocol Special?

The HCP's diffusion protocol differs from traditional diffusion imaging in several important ways. It uses many more diffusion encoding directions (270), enabled by an hour-long scanning session and three-fold multi-band acceleration (MB=3), which incurs little SNR penalty. The WU-Minn-Ox HCP dMRI data were acquired on a customized scanner with stronger diffusion encoding gradients that can operate with 100 mT/m maximal gradient strength (G_{max}). This reduces diffusion encoding times (TE) compared to standard gradients of 40 mT/m (Siemens Trio) for a given diffusion weighting magnitude (b value). Consequently, signal loss due to T2 decay during diffusion encoding is significantly less and the resultant gain in SNR is traded for higher spatial resolution, HCP dMRI data achieved 1.25 mm isotropic resolution with sufficient SNR for b-values as high as 3000 s/mm² and used b shells of 1000, 2000, and 3000 s/mm² 18. (Note that Siemens subsequently introduced the 3T Prisma, developed in part through their collaboration with the HCP with a maximum gradient strength of 80 mT/m, which enables users in the community to use HCP-style diffusion protocols). The 7T data also have high SNR with an even higher spatial resolution (1.05 mm isotropic), 130 directions at b-values of 1000 and 2000 s/mm² 19 (see Supplementary Fig. 2). The MGH/UCLA HCP consortium took a different approach with their even more powerful gradients (300 mT/m G_{max}) by scanning subjects at 1.5mm isotropic resolution and b values up to 10000 s/mm² 20. Both HCP consortia chose to use the original monopolar (i.e., Stejskal-Tanner²¹) diffusion encoding sequence. It has a shorter TE relative to the bipolar (twice-refocused) sequence²² that has been popular for whole brain

diffusion MRI in the last decade. The bipolar sequence reduces eddy currents during acquisition at the expense of an increased TE and thereby reduced SNR. The HCP could tolerate substantially worse eddy current effects with the monopolar diffusion encoding at high b-value because eddy current distortions are instead corrected during post-processing using a newly developed approach²³ that goes beyond traditional affine image registration (see Note #8 below). The HCP also acquired two sets of scans having reversed phase encoding directions. This enables correction of signal that had piled up in one phase encoding direction and spread out in the other without an SNR penalty. Indeed SNR actually increases across the image when both datasets are averaged, producing a relatively unbiased estimate of the signal intensities throughout the final image²⁴). Tractography examples indicative of the data quality are shown in Sotiropoulos et al.¹⁸. Another example (Supplementary Fig. 3) illustrates the ability to precisely map patterns of cortico-striatal connectivity from human HCP tractography data that closely parallel the tracer-based connectivity patterns reported in the macaque.

3. Simultaneous Multi-slice (Multi-Band) Echo Planer Imaging (EPI)

The high spatial and temporal resolution of the HCP data would not have been achievable without the accelerated imaging enabled by Simultaneous Multi-slice (Multi-Band) EPI. This transformative development has become the new standard for EPI imaging, the workhorse sequence for brain fMRI and dMRI. In MR imaging, there is always a tradeoff between resolution and data acquisition times. Higher resolution data simply take longer to acquire. In fMRI, this tension leads to longer TRs (i.e. worse temporal resolution) with increasing spatial resolution. In dMRI, the trade-off is between spatial resolution and angular resolution (in qspace). The approach adopted by the HCP to substantially alleviate this tradeoff is the simultaneous multi-slice (SMS), multi-band (MB) technique that was first employed in brain imaging to reduce the time to acquire very high resolution functional imaging data²⁵. In conventional EPI image acquisition, each 2D slice covering the volume of interest is acquired sequentially in time. In SMS/MB, multiple slices are excited and acquired simultaneously using multi-banded pulses. The resultant image represents the sum of the simultaneously excited slices. These overlapping slices can be unaliased using spatial information inherent in the multi-channel array of coils that are used for receiving the signals (see Supplementary Fig. 4). These coils are sensitive to regions of the brain in close proximity to their physical locations and lose sensitivity with increasing distance. Each coil yields a linear combination of signals from the different slices (weighted by their sensitivity profiles), and a matrix inversion provides a solution to separate the slices from one another. The imaging time is reduced by the number of simultaneously excited slices (MB factor), which receive the same contrast encoding. This is especially important for EPI-based fMRI and dMRI, as contrast encoding time usually outweighs the EPI readout time. The MB factor cannot be increased too high, however, as it is affected by the number of receive coils in the array, their size, proximity to the sample, and layout. There is a penalty to be paid in noise enhancement and cross-slice contamination with increasing MB factors, which must be kept to a negligible level, but these parameters can be measured to determine feasible MB acceleration factors. Displacing slices with respect to each other at the image acquisition stage prior to unaliasing significantly improves the maximum usable MB factors²⁶. In the HCP's 3T fMRI data acquisition, an MB factor of 8 was used with a 32-channel array coil after extensive testing. This allowed 2 mm isotropic whole brain coverage in 0.72s as opposed to the 5.6s that a single band fMRI sequence would take to acquire such data. significantly improving data quality and SNR efficiency²⁷. For HCP diffusion MRI, a MB factor of 3 was employed, making it feasible to obtain high spatial and g-space resolution data. In dMRI. the ultimate limitation in acceleration is the T₁ relaxation of spins, as going faster than approximately 2xT₁ per volume of interest (whole brain for the HCP) becomes SNR-inefficient. However, when high spatial resolution is employed in dMRI, MB factors are restricted by RF

power deposition before reaching this ultimate limit. The dMRI sequence requires more power than the fMRI sequence because, as a spin echo sequence, it requires the use of a 180-degree refocusing pulse in addition to the 90-degree excitation pulse that it shares with a gradient echo fMRI sequence. Further, the MB pulses aggravate the situation because they also increase the power demands. Thus the sequence is limited by the safely achievable peak power and total power deposited in the subject. There are solutions to these limitations developed by the HCP^{19,28,29}, some of which were employed in the 7 Tesla dMRI component of the HCP data acquisition (as higher field strength also leads to higher power requirements).

4. HCP-Short Protocols

The HCP-Short protocols are similar in key aspects to the HCP-Full protocol, but are less than half the total scan duration (~1-2 h total scanning over one or two sessions vs ~4 h over four sessions). HCP-Short is designed for studies in which the subject population may not tolerate intensive multi-session scanning and individual scans of 15 min duration, owing to age and/or brain disorders. The HCP-Short protocols aim to provide sufficient high-quality data to enable HCP-style analyses, including selective removal of artifacts, intersubject registration using arealfeature-based alignment (e.g., MSMAII), and cortical parcellation using the areal classifier. There are multiple variants of the HCP-Short protocols, because some protocols have projectspecific aspects (e.g., related to the particular age range or brain disorder under investigation). The initial HCP-Short protocol was developed for the HCP Lifespan Pilot project, a 1-year effort by both HCP consortia to obtain pilot data for the recently launched HCP Lifespan Development and Lifespan Aging projects (see Figure 5 in main text). We describe below adaptations of the Lifespan HCP-Short protocols proposed for the HCP-Aging and HCP-Development Lifespan projects, which will both use the 3T Prisma platform at each of the four consortium data acquisition sites. Note that these protocols are currently undergoing further piloting, with final protocols and parameters to be available at http://protocols.humanconnectome.org/lifespan/.

Structural scans for HCP-Short are similar to those for HCP-Full, but the spatial resolution is slightly lower (0.8 mm isotropic voxels instead of 0.7 mm), and only a single pair of T1w and T2w scans is collected (albeit with a repeated acquisition if the scanning technician deems them to be only "fair" or worse quality). This increases SNR and shortens scan duration slightly (13 min total for T1w and T2w vs 16 min) while preserving the benefits of high spatial resolution for cortical segmentation and maps of myelin and cortical thickness. Additionally, the Lifespan HCP may use 'volumetric navigators' (vNavs) to record (and possibly correct for) within-scan head motion^{30,31} – see also Note #5 below.

For fMRI scans, the HCP-Short protocol acquires data at the same spatial and temporal resolution as for HCP-Full (2 mm isotropic; TR = 0.72-0.8 s) using a MB factor of 8. Phase encoding is along the A/P axis for the Prisma (vs the L/R axis for the customized WU Connectom scanner). In addition, HCP-Short acquires a spin-echo AP-PA field map pair.

For rfMRI, HCP-Short scans are shorter in duration (usually 5.0-5.5 min/scan, but 3 min/scan for younger children, vs 15 min/scan for HCP-Full), and in total scan time (30-40 min vs 1 h). The reproducibility of resting-state networks (RSNs) increases steeply with total scan time over the first 30 min of rfMRI acquisitions (with additional improvement up to 80-100 min)³². It may be possible to augment information about RSNs using task-fMRI scans (see below), given that functional connectivity is similar when measured during tasks or the resting state³³, and this may add to the useful data for intersubject alignment and individual-subject parcellation.

The task-fMRI aspect of the HCP-Short protocols utilized to date is much briefer (ranging from no task fMRI to ~30 min total, vs 1 h for HCP-Full) and allows for project-specific tasks appropriate for particular age ranges or brain disorders. The HCP Lifespan projects are currently considering options to acquire task data using up to five tasks, each with a notably short (~2 min) scan duration. More generally, the HCP-style paradigm supports diverse task fMRI protocols to reflect project-specific aims without major adverse impacts on intersubject registration ("MSMAII" surface registration does not use task-fMRI activations) or areal classification (which also does not require task-fMRI activations, cf. Ref. ³⁴).

For dMRI, HCP-Short acquires 22 min of dMRI data (four 5.5 min scans) at a spatial resolution of 1.5 mm isotropic voxels (vs. 1.25 mm for HCP-Full), two b values (1,500 and 3,000 s/mm²) and ~75-90 directions/shell. The lower maximal gradient strength for the Prisma (80 mT/m, vs 100 mT/m for the WU customized Connectom scanner) yields a longer TE and slightly lower SNR per image, but HCP-Short uses a simultaneous multi-slice ("multi-band") factor of 4 for dMRI on the Prisma (vs. 3 for main HCP) while keeping within SAR and peak power limits because of additional pulse sequence development that occurred after the main HCP began. Possible refinements are being evaluated that would enable a MB factor of 5 or 6 using the two channel parallel transmit (pTx) capability of the Prisma and the local and global SAR regularized multi-band pTx pulses recently developed at UMinn^{29,35,36}.

The Lifespan HCP-Short protocol will also likely include a ~5 min Arterial Spin Labeling (ASL) perfusion MRI scan, which provides a quantitative measurement of cerebral blood flow (CBF). Since perfusion is normally coupled to metabolism, ASL is a useful surrogate marker of basal brain function, as well as aiding in the discrimination between neuronally-related vs cardiovascular-related effects seen in BOLD imaging. The Lifespan HCP projects are currently evaluating both background suppressed 3D segmented GRASE pseudo-continuous ASL (pCASL)³⁷ and 2D multi-band EPI based pCASLa³⁸.

5. Head Movement During 3D Structural Scans

Head movement during 3D T1w and T2w scans produces currently uncorrectable image blurring and banding and is a major problem for both research and clinical neuroimaging. Even a single translation of a millimeter or so during a scan of 6 – 8 min duration will cause blurring. The main HCP reduced the problem by acquiring two T1w and T2w images. Moreover, if these were not of adequate quality - rated good or excellent based on a systematic manual QC evaluation

(http://humanconnectome.org/documentation/S900/HCP_S900_Release_Appendix_IV.pdf, parts I – III, pp. 17 - 28), subjects were asked to complete an additional scanning session where up to two additional pairs of T1w and T2w images were acquired. All T1w and T2w scans from the same session that were rated good or excellent in overall quality were used in further processing. Though the HCP invested in an external motion tracking camera, this technology was in its very early stages, and problems associated with tracking chip attachment to skin and skin movement independent of the rigid body motion of the head were not solved. As a result this approach was abandoned early in the project. An alterative approach that shows promise involves 'navigator' sequences that acquire low resolution EPI data within the dead time of T1w and T2w sequences. This data can be used to detect subject motion, and, in addition, adjust the imaging gradients to compensate for subject movement, avoiding blurring issues. Also the sequence can reacquire lines in k-space that were acquired during subject movement so as not to use k-space data corrupted by motion in the final Fourier reconstruction. If the robustness of these approaches can be established, future studies may wish to take advantage of such online movement correction of the sequence consistently high quality structural image acquisitions.

6. Gradient Distortion

Spatial localization as well as diffusion encoding is achieved through the gradients of the MR system. However, the gradients are not perfectly linear in space, which leads to image distortions as well as errors in diffusion encoding. For dMRI in particular, such gradient nonlinearities can influence the magnitude and direction of diffusion-sensitizing gradients⁴⁰ in addition to spatial distortions. Spatial distortions due to gradient nonlinearities need to be corrected if images acquired on different scanners are to be compared. Also the correct diffusion weighting magnitude and direction must be used if one is to avoid errors in the estimated diffusion parameters that drive tractography algorithms. In the HCP these corrections were implemented within the HCP's minimal preprocessing pipelines¹⁷.

A second problem exists with respect to calibration of the gradients. For example, if the sequence calls for 40 mT/m, do we actually get 40 mT/m or just 39.2 mT/m? Usually the calibrations are within a small percentage of the targeted value, but they are not exact. This was initially a problem for combining 3T and 7T data on the same subject in the main HCP. These issues were carefully evaluated and calibrated using a grid "phantom" and corrected in postprocessing, by enabling additional degrees of freedom in the boundary-based registration of the HCP's minimal preprocessing pipelines^{17,41} (from 6 DOF to 12 DOF).

7. Why Spin Echo Field Maps?

Maps of susceptibility-induced image distortion can be derived from a pair of spin-echo EPI images with reversed phase encoding directions. We call these "Spin Echo Field Map" scans in the HCP-style protocols. They offer several advantages over the more traditional dual-echo. gradient-echo based field maps, where the magnetic field is computed based on the phase difference of images taken with two differnt echo times. Spin echo field maps are faster to acquire (and thus more robust to subject motion), and can be matched exactly to the gradient echo fMRI data in terms of resolution, geometry, and distortion (via matching the echo spacing). As a result, the distortion is identical between the two images, making image registration of the distorted spin echo and gradient echo fMRI images substantially easier. The HCP tested the two distortion correction approaches and found that they performed similarly, leading to the recommendation that the faster imaging approach be used. Importantly, additional information can be obtained from the spin echo images. Gradient echo fMRI images suffer from signal dropout due to dephasing in areas of high magnetic field inhomogeneity that is absent in the spin echo field map images with similar image distortion. This difference can be used to segment these regions of dropout, and these segmentations can serve as useful information to know where the fMRI signal is likely to be unreliable.

Additionally, the spin echo and gradient echo images can be combined to estimate the difference in the effects of the B1 transmit field the two images, along with an estimate of the B1 receive field on the fMRI image³⁴. This receive field is more accurate than the field generated from T1w and T2w images acquired in a different imaging session (and which also experience somewhat different transmit field effects), and thus can be used to bias-correct the fMRI data without the need to acquire additional sequences or use additional preparatory time on the scanner. For some fMRI analyses (e.g., resting-state correlations and task z-statistics), bias correction of fMRI is not needed. However, for other analyses (e.g., beta maps, variance maps), an fMRI intensity bias field can make the data less biologically interpretable across the whole image. For such analyses, accurate intensity bias correction is important³⁴. The latest version of the HCP's minimal preprocessing pipelines (v3.15.1 or later) now enables this sort of

correction; moreover, as a simple multiplicative bias field, it can be retrospectively applied to prior HCP data as well.

8. Eddy Current Distortion Correction and dMRI Outlier Removal

A major problem for dMRI data arises from eddy currents induced by magnetic field modulations that are distinct for each diffusion direction and are more pronounced in the mono-polar diffusion-encoding scheme chosen by the HCP (for SNR efficiency as mentioned in Note #2). dMRI data also suffer from signal dropouts caused by head movement and other sources. analogous to those that plaque fMRI data. The HCP's 'eddy' software (released in FSL, and not to be confused with the older 'eddy correct' script based on affine registration) corrects for linear and nonlinear eddy current distortions and also has an option to use model-based outlier detection and replacement to correct corrupted individual slices^{24,42,43}. Counter to the typical practice of individually registering each diffusion-weighted volume to a reference, eddy uses all data volumes jointly. A Gaussian-process-based generative data model is used to make predictions about what the 3D dMRI volumes should look like at every point in q-space given knowledge of the different distortions. Inversion of this model enables accurate estimation of the distortion fields, which can then be applied to correct the data. This combination of monopolar acquisition and eddy-based post-processing results in fewer artifacts and less distortion than traditionally achieved, yet with lower TE and therefore higher SNR, which the HCP used to achieve higher spatial resolution (see Note #2).

9. More about the Global Signal and Temporal Cleanup of fMRI Data

One idea that has gained substantial currency in the literature is that global signal fluctuations are often induced by direct effects of subject movement 44-46. Invoking this hypothesis, some argue that it is necessary to regress out the global signal in order to adequately correct for subject movement. However, to our knowledge, no direct biophysical causal mechanism has been proposed to account for subject movement leading to global fluctuations in the fMRI signal (particularly those with the tissue specific and area specific effects shown in Supplementary Figs. 5 and 6). Instead, it is worth considering the kinds of effects seen in fMRI images that are clearly attributable to head movement and which can be removed using ICA+FIX (ICA+FIX consists of detrending, aggressive 24 movement parameter regression, and non-aggressive noise ICA component regression): 1) Largely linear effects of moving the head within an inhomogeneous receive coil. These effects tend to produce "rings" around the perimeter of the brain, reflecting signal intensity increases as the head gets closer to the head coil on one side and decreases as the head gets further from the coil on the other side. Such spatially specific effects are readily identified by ICA+FIX and eliminated from the data. 2) Complex interactions between head motion and the magnetic field inhomogeneities in the fronto-polar, orbito-frontal, anterior temporal, and inferior temporal cortices. These effects are also spatially specific and are readily identified by ICA+FIX and eliminated from the data. 3) "Spin history" effects of movement caused by some portions of the brain being excited sooner than they should and other parts being excited later than they should by the slice selection profiles. In a 'worst case' scenario where the head moved exactly one slice up or down in the slice direction, this would produce a strong banding pattern in the data. Such bands are readily identified by ICA+FIX and eliminated from the data. In summary, no plausible biophysical mechanism has been proposed by which head movement would induce a truly global change in fMRI signal intensity (short of the subject completely exiting the head coil), as opposed to spatially specific changes in fMRI signal intensity that are visible to spatial ICA-based approaches. Further, the global fMRI signals are highly tissue and area specific (Supplementary Figs. 5 and 6), and no plausible biophysical mechanism has been proposed for how subject motion could directly cause such

phenomena. Instead, the hypothesis advanced in the main text Box 1 about global fluctuations being partly due to non-neuronal physiological fluctuations and partly due to neural effects seems much more likely.

Approaches such as ICA+FIX and similar methods^{47,48} are attractive for removing the direct temporal image intensity effects of movement from fMRI data (along with other spatially specific effects like MRI physics related artifacts or arterial and venous pulsation artifacts). Indeed, ICA+FIX results approach that of the more aggressive censoring ('scrubbing') approach when movement becomes severe, but are not forced to make binary 'scrub/not scrub' choices based on thresholds in framewise displacement (FD) and/or framewise image intensity difference (DVARS) measures that may themselves be quite noisy in high temporal resolution data. Instead ICA+FIX derives a weighted solution directly from the data. That said, it remains an open question whether scrubbing (for example, based on those frames where ICA+FIX begins to approach the scrubbing solution by removing substantial amounts of unstructured noise variance from the data) would be useful if we put aside for a moment the effects of global noise. A similar question can be asked regarding the standard bandpass filtering that is often using to clean fMRI data, but has not been demonstrated to be useful above and beyond ICA+FIX and appropriate global noise cleanup for structured noise removal.

Unfortunately, white matter and CSF timeseries regression are unlikely to represent an effective approach to global noise cleanup, as they have relatively little correlation with the global fluctuations as long as they are derived from ROIs that are free from grey matter contamination (which can arise from spatial smoothing or partial volume effects of 'white matter' voxels encroaching into gray matter, particularly in low resolution datasets), see also Supplementary Fig. 5. Indeed approaches that claim to remove global artifacts without global signal regression using white matter or cerebrospinal fluid regressors are likely actually removing the much stronger mean grey matter signal that contaminates voxels near the grey matter. These approaches would likely behave differently if white matter and cerebrospinal fluid tissue masks were eroded by at least 2 imaging voxels or 4mm from the grey matter. Similarly it is not clear that physiological noise modeling approaches, such as that currently implemented in FSL's PMN⁴⁷ will deal with all global noise, though they will likely help. Imperfections in the modeled effects of physiological noise parameters on BOLD fMRI data will lead to incomplete physiological noise clean up. As mentioned in the main text Box 1, a recommendation for the complete approach to fMRI timeseries noise cleanup will have to wait until a selective and effective approach to global noise cleanup becomes available.

Besides ICA+FIX^{48,49} and ICA-AROMA^{50,51}, Multi-Echo-ICA (ME-ICA) is another approach that uses ICA to identify and remove non-BOLD noise components from functional timeseries^{52,53}. In this approach, multiple images are acquired at different echo times (TEs), and each component is scored as to how well it fits a T2* or BOLD-like decay curve. Importantly, although this approach is able to separate BOLD-like from non-BOLD-like components, it does not remove BOLD-like global fluctuations from physiological sources (e.g., the global fluctuations in deoxyhemoglobin from rate and depth of breathing, heart rate, and end tidal pCO2 mentioned in the main text Box 1). Thus, it does not represent a solution to the global signal/global noise problem raised here. Additionally, multi-echo EPI is currently limited in the spatial and temporal resolution that it can achieve (on the order of 4 mm isotropic in published results with a TR of 2.5 s). With the incorporation of multi-band into a multi-echo sequence, the spatial-temporal resolution of this technique should be improved, enabling accurate cortical analysis (i.e., 2.5 mm isotropic or better). At that point, a thorough evaluation of the pros and cons of standard multi-band vs. combined multi-band and multi-echo should be done. Tradeoffs to consider include spatial-temporal resolution and temporal SNR, relative to performance in timeseries de-noising.

Currently, we recommend using one of the other ICA-based techniques with standard multiband fMRI. It is possible that the temporal resolution recommendations of <1 s could be relaxed for multi-echo techniques, though the impact of fewer timepoints on multi-variate methods such as ICA or partial correlation would need to be investigated.

10. Selective Spatio-temporal Smoothing of Unstructured Noise

A major limitation of spatial and temporal smoothing (or filtering) is that they smooth signal and noise indiscriminately. When using smoothing to improve SNR, the data is also being blurred across areal boundaries or removing high frequency temporal information that is known not to be fully noise^{54,55}. For the purposes of estimating dense functional connectivity, an attractive alternative makes use of Principal Components Analysis (PCA) to represent an fMRI 4D timeseries as a series of weighted spatial eigenvectors, ordered according to the size of their eigenvalues. The eigenvalues of the PCA decomposition together form an eigenspectrum (a Wishart distribution in the null case⁵⁶). A white noise timeseries will have this characteristic spectrum of eigenvalues, and structured signal (or noise) present in the data will increase the eigenvalues above and beyond this null eigenspectrum. Even in a dataset containing structured signal (e.g. area-specific resting-state fluctuations), the tail of the eigenspectrum will be largely unstructured noise. A Wishart function can be fit to this tail, making a prediction of the null eigenspectrum for the dataset. This null distribution can be subtracted from the data, forcing most of the eigenvalues representing noise to zero, and the PCA series can then be converted back to a timeseries that has higher SNR (for individual subjects), or correlated as a PCA series with higher SNR and fewer artifacts (for group data). Although reducing the small eigenvalues may remove some very weak signals, the technique avoids the blurring of strong signals that occurs with typical spatial and temporal smoothing. Additionally, this Wishart rolloff approach can be used to remove the effects of spatially autocorrelated noise and ringing from iteratively generated group PCA series⁵⁷ (httpz://www.humanconnectome.org/documentation/mound-andmoat-effect.html). Supplementary Fig. 7 illustrates both effects.

11. Improving Subcortical Alignment Using Diffusion Fiber Orientations

One arena in which the current HCP Pipelines could likely be improved is in the alignment of the subcortical white matter, which is currently based only on T1w images that have relatively little contrast within white matter. Much as areal-feature-based registration improves the alignment of cortical areas over and above using only folding patterns, using fiber orientation information to drive white matter tract alignment will provide the registration algorithm with more information about the identity of the fiber tracts being aligned. Because of their low within white matter contrast, T1w images will only approximately align these tracts. Future versions of the HCP Pipelines may integrate a diffusion fiber orientation-based nonlinear registration (e.g. http://dti-tk.sourceforge.net/pmwiki/pmwiki.php?n=Main.HomePage)^{58,59}.

12. Effects of Thresholding on Reproducibility of Neuroimaging Results

Two interrelated issues arise from the traditional use of statistical thresholding for display of neuroimaging results and identification of regions (which are often then converted to 3D standard space coordinates as the sole means of data sharing):

(1) Statistical significance maps (e.g. Z or p maps) are affected by more than the measure of interest (e.g., %BOLD change or relative myelin content). They also reflect an estimate of the error of the measurement. While such maps are statistically interpretable as the likelihood that a finding differs from chance, and thus have a role for establishing statistically significant effects,

they are less biologically interpretable than maps of the original effect of interest. For example, in data generated using modern multi-channel head coils, regions near the center of the brain have lower Z-statistic values owing to the higher amount of unstructured noise in those regions after correcting for the sensitivity profile of the head coil. This will decrease Z-statistics even if the %BOLD change in a subcortical central region is the same as that in a cortical region. Thus, when searching for biological boundaries in the data, one should arguably use maps of the effect of interest, rather than Z-statistic maps. Further, biologically important boundaries in the data are likely to be identified most reliably by looking for large and rapid changes in the original effect of interest over a small spatial distance. Thus focusing on gradients (measures of spatial change) in modalities can be an effective way to identify biologically important boundaries in one's data, as was done in the HCP's recent multi-modal parcellation³⁴. Using such gradients may aid in comparing across studies to determine the overlap of results (i.e., do two studies activate or connect the same brain area, neighboring brain areas, or more distant brain areas).

(2) Traditionally, statistical significance maps are thresholded (usually at p<0.05 after some method of multiple comparison correction), and the regions that are identified are generally represented by the stereotaxic coordinates of the spatial center of gravity. Often the thresholded statistical maps are all that is provided in the published figures of a study, making it harder to identify the underlying patterns in the data (only the "peak" and "valley" blobs are identified). One problem that could be caused by this approach is unintentionally hiding evidence of an artifactual pattern that is more obvious in the unthresholded data (e.g. after thresholding one might find a few blobs on gyri and sulci, but the unthresholded data might clearly have a gyral vs sulcal bias). Another issue is that the location of the statistical threshold contour can vary much more across studies than the underlying maps of the effect of interest or their gradients (See Supplementary Fig. 8). Relatively uninteresting differences in Z statistical maps can easily turn activations whose significance is right around the threshold value from being above threshold to below threshold and vice-versa as illustrated in the figure. These differences include slight changes in the spatial distribution of the noise, slight errors in the measure of the effect of interest, or even something as simple as adding more subjects or fMRI timepoints. When looking for cortical areal borders, the focus should be on where the effect of interest changes markedly across the cortical surface, not where the variance of non-interest changes. Also, in the limit, given enough data in an individual subject or enough subjects, nearly the entire brain may have statistically significant activation and deactivation in a task fMRI study using even a very conservative statistical threshold, but this says little about the biological significance of this whole brain activation and deactivation 60. Thus, it is advantageous to use gradients in the maps of the effect of interest to delineate areal borders or regions of interest, rather than statistical thresholds. Some of the spatial uncertainty present in the traditional neuroimaging analysis paradigm may reflect the use of statistical thresholding to define clusters from which 3D activation peaks, or centers of mass are computed (also, many of the widely used multiple comparison correction and statistical thresholding approaches have very inflated false positive rates⁶¹). Thus, it would be helpful if the maps of the effect of interest, their gradients, and statistical significance maps were displayed in publications and uploaded to neuroimaging databases such as BALSA (see main text), OpenfMRI (https://openfmri.org), and NeuroVault (http://neurovault.org). Finally, as noted in the main text tenet 6, parcellated analyses provide an excellent alternative approach that has greater sensitivity and power while avoiding many of these issues.

13. Group Average Registration Drift

Group average registration drift is a phenomenon that occurs when registration templates are being generated with non-rigid registration algorithms or when different registration algorithms or templates are used to drive alignment to a common standard space. A particularly striking example of drift is the ~37% increase in brain volume that occurs when one registers HCP study brains to the MNI152 template (see Supplementary Fig. 9), owing to a gradual expansion in the non-rigid components of the registration during the iterative process of template generation that was used to create the original MNI space^{62,63}. Drift can be removed by computing the group average registration effect and concatenating its inverse onto each individual subject's registration. It is both desirable and feasible to avoid such drifts in future brain templates. If such drifts are eliminated⁶⁴, very precise cross-study, cross-modal boundary comparisons can be made. This is true even when using different areal features in different studies after a gentle folding-based registration to a common geographic template (see Supplementary Fig. 10), so long as the two studies' samples are drawn from the same population. Importantly, the current grayordinates space is still based on MNI space for subcortical structures and thus still has this drift embedded in it. A future version of the CIFTI grayordinates space may eliminate this drift.

14. 'Weighted' Approaches to Parcellation

Parcellation of the brain is often carried out by clustering together neighboring grayordinates on the basis of the similarity of their timeseries. This typically yields a large number of nonoverlapping parcels, with a single contiguous group of grayordinates in each parcel (or network node). This can be referred to as a "hard parcellation" 65,66. However the clustering does not have to be "hard", but may instead be "fuzzy" or even probabilistic. One approach to generating a "fuzzy parcellation" involves high-dimensional spatial independent-component analysis (ICA)⁶⁷. Using ICA, each network "node" is described by a spatial map of varying weights. Notably each map may overlap with other nodes' maps and may include more than one set of contiguously neighbouring grayordinates. Network edges (connections between nodes) are estimated by comparing the fMRI time series associated with the nodes (usually derived using multiple regression). With any of these parcellation approaches, brain connectivity can be represented as a 'parcellated connectome', which can be visualized simply as an N_{nodes} x N_{nodes} network matrix or a graph (explicitly showing nodes and the strongest edges⁶⁸) or using more sophisticated visualization approaches that embed nodes and edges into spatial representations of the brain⁶⁹. Hard parcellations have the advantage of being simpler representations and are therefore a more parsimonious yet accurate representation, if they are a good biological model of the data. They are also easier to interpret as distinct neuroanatomical entities. On the other hand, soft parcellations may be useful where the biology does not fit the model of nonoverlapping spatial segregation or there is a premium on maximizing the homogeneity of signals within parcels. Multiple regression using a weighted parcellation may be able to produce "purer," more homogeneous timecourses when multiple neural signals are spatially superimposed (e.g. the distinct signals related to the eccentricity and polar angle axes of visuotopic organization in the visual cortex³⁴). Further, an ICA-based weighted parcellation by definition will not allow rank-deficient node-timeseries matrices to occur, whereas this condition can be a problem for partial correlation of hard parcellations (i.e. especially if a parcellation includes homotopically related parcels across the two hemispheres).

15. Current Limitations to Generating Human Connectomes Using Non-invasive Approaches

The Human Connectome Project's core goal is to create area-to-area connectivity matrices using diffusion tractography and functional connectivity. While much progress has been made in this direction (in particular in defining the cortical areas to be connected³⁴), much work remains to assess and maximize the accuracy of the non-invasive measures of brain connectivity by comparing them to invasive tracers in non-human primates.

Despite its well documented capabilities for localizing major fiber tracts in the white matter (see below), using diffusion tractography to map connections between distant gray matter regions has several limitations and is a field of active research⁷⁰⁻⁷². Some of the limitations are fundamental, whereas others are amenable to methodological advances. One fundamental limitation of diffusion tractography is that it is unable to provide accurate directionality information about connections (it is impossible to know if an estimated connection is $A \rightarrow B$, A \leftarrow B, or A \leftarrow \rightarrow B) because water diffusion in axons is bidirectional. In contrast, invasive retrograde and anterograde tracers reveal the directionality of connections^{71,73}. Thus, directionality information would have to be inferred through some other means (indeed, this prospect was one reason that MEG was included as an imaging modality in the HCP). Another problem is the indirect nature of the measurements: a mapping is necessary from measurements of bulk water diffusion displacements to axons and fiber patterns. Due to the limited resolution of MRI (~mm³), hundreds of thousands of axons coexist in the same brain imaging voxel⁷⁴, giving rise to different complex geometries (crossing, "kissing", branching, fanning, converging). Distinguishing these configurations from the measured diffusion signal is an ill-posed problem in general. For instance some axons in white matter branch by bifurcation at quasi-right angles⁷⁵, making it difficult for tractography to distinguish axonal branching from crossing of independent fiber bundles. These issues can cause errors when tracking either deep in the white matter or close to the cortex.

Current tractography paradigms are also challenged in reaching terminations within gray matter and in accurately tracking in the vicinity of the WM/GM boundary^{74,76,77}. There is a pronounced bias in tractography streamlines towards being higher on gyral crowns and much lower in sulci. When streamline density is normalized relative to flat regions (zero surface curvature in sulcal banks), it becomes clear that the bias is actually more of an anti-sulcal bias rather than as strong an increase in gyral connectivity. Such biases are strongest when tractography streamlines return to the cortex from the deep white matter, and so they are most prevalent when seeding bidirectionally within the white matter (e.g. 'matrix3' in FSL's probtrackx2 tractography algorithm). Pushing to higher resolution does help (as shown when fusing 3T and 7T HCP dMRI data⁷⁸), but does not fully address this issue (as shown with 250um isotropic resolution in post-mortem macaques⁷⁷). The problems arise because (i) near sulcal fundi, superficial white matter and deep grey matter are dominated by tangentially running "U-Fibers," which direct streamlines around the sulci, rather than allowing them to penetrate the grey matter; and (ii) in 'gyral blades' (white matter lying between sulcal banks), fiber bundles are predominantly oriented towards gyral crowns. It is possible to predict components of a neuroanatomically reasonable gyral bias in the tractography using simple geometric models^{74,76,77} that assume that each cubic mm (or square mm of the cortical midthickness surface) should, to first order, send out and receive a comparable number of axons. Thus gyral crowns should have somewhat more axons crossing their white matter surfaces and sulcal fundi somewhat less; however, the bias observed in tractography is frequently much higher ^{74,76}. This geometric model can serve as a reference upon which tractographic model improvements can be objectively evaluated.

Concerns about the performance of tractography near the grey-white interface and the influence of folding patterns on the locations of gradients in tractographic connectivity led to tractography not being used as a modality in the HCP's multi-modal parcellation of the cerebral cortex³⁴. That said, it will be interesting to compare tractographic connectivity gradients to the multi-modal consensus areal borders once large amounts of HCP tractographic data have been generated and shared. Should tractography produce gradients that overlap substantially with the multi-modally defined cortical areal borders, tractography features could aid in predicting the locations of cortical areas in individual subjects using the multi-modal areal classifier³⁴. If tractography gradients do not have strong overlap with the multi-modal consensus areal borders, future improvements in tractographic models could be objectively evaluated with respect to these reference areal borders to aim for better agreement.

Collectively, these issues give rise to significant challenges in keeping track of area-to-area connections as they travel from cortex through the deep white matter within fascicles and then fan out again to reach distant regions of cortex. To the extent that axons do not remain clustered together relative to the voxel size of the diffusion scan, these fascicles—which themselves have complex cross-sectional shapes—form bottlenecks where the accuracy of information about the strengths of inter-areal connections will be eroded. If, on the one hand, fibers from one area tend to cluster together within the deep white matter and then fan out again when they reach cortex, this will relax the spatial resolution requirements needed to accurately reconstruct area-to-area pathways. On the other hand, if axons relatively randomly distribute themselves within major fascicles, accurate reconstructions of area-to-area pathways would require dramatic improvements over current diffusion MRI spatial resolution. Invasive tracer studies in non-human primates compared with tractography may help to resolve which reality predominates within the brain; though the plethora of studies that show agreement between tractography results and other modalities and in various contexts (see Ref. 71 for a review of such evidence) suggest that the geometrical realities may be somewhat on the favorable side.

The above limitations in estimating direct connection strengths using tractography are difficult to fully address. However, better data and better methods can help to reduce errors. Work within the HCP consortium has aimed towards these goals 18,19. The HCP pushed the spatial resolution of dMRI to reduce partial volume effects, while keeping high SNR and high angular resolution. At the same time, data at multiple angular contrasts (b values) have been sampled, which can further help in accurately mapping of fiber orientations^{79,80}. Up to three fiber crossings within the deep white matter, as well as various microstructural metrics, 81-84 can be routinely modeled using the current HCP data releases. New approaches for mapping connections are also being explored, including incorporation of fiber dispersion⁸⁵ in tractography and neighbourhood-wise tracking⁸⁶. Better, neuroanatomically informed models of axonal trajectories near the cortical ribbon incorporated into tractography may help to reduce gyral and sulcal biases. In Ref. 87, the relationship of fiber orientations with cortical features is explored using post-mortem high-resolution histology and used to inform generative models for in-vivo diffusion MRI. This work hopes to use mesoscale information to better estimate the paths of tractography at the transition between white and gray matter. As improved tractography models aimed at measuring accurate area-to-area connection strengths emerge, results can be compared directly with invasive tracer studies in the macague to see whether the modifications to the tractography models have improved the agreement between tracers and tractography⁷⁶.

Aside from the question of accurate area-to-area connections and connectomes, tractography can still provide useful information about the major fascicles themselves, for understanding how large territories of cortex are connected (e.g., refs. ^{88,89}, for understanding how large fascicles have changed through evolution ^{90,91}, for understanding differences in fascicles in health and

disease, or for the purpose of surgical planning. Thus, even without yet having a fully accurate and validated area-to-area structural connectome, investigators are able to see how large ensembles of brain regions are connected to one another and to investigate white matter microstructural changes along these fascicles.

A different set of limitations affect functional connectivity measures. Obtaining connectivity estimates using statistical dependences of the time-varying BOLD signal is also a very indirect method and has its own caveats. Functional connectivity can be strong between parts of cortex that are known not to be directly connected (e.g., the horizontal meridian representations in V1 across hemispheres⁹²). Thus, functional connectivity reflects both direct and indirect connections, likely including common inputs. One approach that attempts to make functional connectivity more direct is partial correlation functional connectivity. This multi-variate approach attempts to identify the unique pairwise statistical influences between cortical areas. Such analyses are very demanding in terms of the number of timepoints and data quality required to produce meaningful data, and might not behave optimally in the context of neuroanatomical parcellations (as opposed to the ICA-based parcellations mentioned in Note #14). Estimating reliable directionality information from rfMRI timeseries is also problematic, partly because of the temporal smoothing of the BOLD response function, although causal inference methods not based on temporal lag are being developed and evaluated for rfMRI data (e.g., see references and discussions in Ref. 93). Another conceptual limitation, raised in Ref. 71 involves an area that participates in many resting state networks and as a result integrates many competing signals. Such an area might actually appear less connected than other areas, and a graph theoretic analysis searching for hubs might overlook it⁹⁴.

Significant progress had been made, in part by the HCP consortium, on how to correct for the temporal fluctuations induced by subject movement (main text and Note #9). That said, as discussed in Box 1 and Note #9, standard full correlation-based functional connectivity has a global signal/global noise problem (as opposed to partial correlation, which is largely immune) that has been much debated in the literature. This problem will need to be solved before full correlation functional connectivity provides an unbiased measure for assessing differences between groups with differing amounts of global noise (fortunately, global noise, by virtue of its global nature, does not affect the locations of gradients). All these functional connectivity approaches (full correlation, partial correlation, attempts at directional correlation) would benefit from comparison with invasive tracer data from non-human primates, just as tractography has done, for validation and to improve their accuracy at representing brain connectivity. Such a framework, just like for tractography, could aid in making decisions about processing and modeling strategies for the generation of more accurate non-invasive human functional connectomes.

16. The 'Standard' Individual Lab's Study in the HCP-Style Paradigm

The HCP-Style neuroimaging paradigm includes a set of standard, concrete recommendations that are applicable to many if not most human neuroimaging studies (i.e., what is desirable for investigators to do in the absence of a good reason to do something different):

Acquisition: A standard study should acquire 3D T1w and T2w structural images on a 3T scanner with a multi-channel head coil (ideally 32 or more receive channels) with 0.8 mm isotropic resolution or better (half the minimum thickness of the cerebral cortex), which can be acquired in ~13 minutes. (3T is currently the most mature platform that produces higher SNR than 1.5T, and the Siemens Prisma scanner is farthest along with the necessary technology and stability). Functional MRI data should be acquired using task and/or resting state paradigms

according to the purpose of the study. The functional MRI data are acquired with whole-brain isotropic spatial resolution of 2.5 mm or better (less than the 2.6mm mean thickness of the cerebral cortex) and a TR of 1 second or less. As much fMRI data is acquired as is feasible (ideally 1 hour or more, and at least 30 min is strongly recommended³²). A phase-reversed pair of spin echo images is acquired with the same geometry and echo spacing as the fMRI data in about 1 minute (for mapping the b0 distortion field and the fMRI receive intensity bias field). If diffusion imaging is a part of the study, it should be acquired using a monopolar diffusion encoding scheme with whole brain coverage and with as high spatial resolution (i.e. less than 2 mm isotropic), as many directions (more than 60 for high angular resolution), and as high a bvalue (ideally more than b=1000 s/mm², and multiple shells are also generally helpful), as is feasible for the scanner being used. At a minimum, phase reversed b0s should be acquired and as many unique directions as possible (rather than repeating directions for averaging). Ideally phase-reversed averages are obtained if there is time to acquire hundreds of directions as in the main HCP. Both the fMRI and diffusion MRI data must be acquired using a multi-band EPI sequence to maximize SNR efficiency and to reach the targets for spatial resolution, temporal resolution, and angular resolution.

Preprocessing: The HCP's minimal preprocessing pipelines, or alternatives with demonstrably superior capabilities, should be used to remove all image distortions, remove the spatial effects of subject motion, align data across modalities, and generate individual subject cortical surfaces, subcortical segmentations, and myelin and thickness maps. Structured temporal noise is removed from fMRI data using ICA+FIX or a similar approach and it is removed from DWI data using the outlier modeling available in FSL's 'eddy' or a similar approach. Cortical data are registered on the surface using MSMAll or a similar areal-feature-based approach to maximize alignment of cortical areas across subjects and, if the study is creating its own arealfeature-based registration template, group registration drift relative to a gentle folding-based geographic registration is removed to enable comparison with other studies that use differing areal features. Volume data are registered using a nonlinear volume registration approach (perhaps in the future based on fiber orientation information if diffusion data has been acquired. but currently based on the T1w image). CIFTI files are created as the final outputs of these registrations in the 2mm standard grayordinates space. Spatial and temporal smoothing are avoided as a rule (unless specific study goals require it, e.g. matching the spatial resolution between fMRI and a grid of ECOG electrodes), and unstructured noise is reduced using parcellation or an approach like the Wishart rolloff method in Note #10. Parcellation is done using a neuroanatomical parcellation that reflects the structural and functional organization of the brain (e.g. the HCP's multi-modal parcellation) and is based on accurately aligned data of hundreds of subjects. If greater accuracy is desired, individual subject parcellations can also be created using a previously trained cortical areal classifier (so far this has been tested on 1 hour of resting state fMRI data, but in principle other kinds of fMRI data should also work, though the minimum amount of data needed has not yet been established).

Analysis: Analyses of fine spatial patterns are carried out on the dense (grayordinate-wise) CIFTI files, however statistical analyses of whole brain, area-level patterns are carried out on parcellated CIFTI files to benefit from gains in sensitivity and power. Statistical comparisons across the whole brain are thus carried out on parcellated data, whereas finer comparisons might be carried out only within predefined brain areas. Obviously there are a very large number of analyses that could arise from HCP-Style studies, including those that wish to go beyond areal-feature-based registration to establish non-spatial correspondences across subjects such as those in hyperalignment⁹⁵. The critical features of an HCP-Style analysis relate to the neuroanatomical fidelity of the spatial localization at the areal level and the use of neuroanatomy in addition to statistics to validate and describe the results.

<u>Data Sharing:</u> Results are visualized on surfaces for the cerebral cortex and in volume slices for subcortical nuclei. If thresholded statistical maps are generated, unthresholded versions of the effect of interest are also made available. When appropriate, figures of brain images are created directly in Connectome Workbench or another neuroimaging software with equivalent features for figure creation, and these figures are saved as "scenes" which link the data to the figure along with any annotations of the figure. These scenes are linked within the resulting publication using unique identifying URLs and the scenes are uploaded to a neuroimaging database so that they and their data are available for download by other investigators. This will allow users access to the brain maps of published figures to facilitate direct cross-study comparisons. Any customized analysis code is uploaded to a code sharing service (e.g., GitHub) and also shared with readers.

17. Can't I Just Map between the Group Average Surface and Group Average Volume MNI Space?

A frequent question on the HCP-Users mailing list concerns how to compare group average or atlas data in MNI space to data within the standard CIFTI grayordinates space (91282) grayordinates, 2mm spacing). Although it is possible to compare group average or atlas data in the standard grayordinate space to an individual subject's physical volume space (by using the MSMAll registered individual surfaces to form a cortical ribbon in the subject's physical volume space and projecting the data into this ribbon), there exists no method to bring data from a group average surface into group average MNI volume space that maintains the spatial localization accuracy of the HCP-style neuroimaging paradigm. Insofar as a particular cortical region has a consistent folding pattern across subjects and these folds have a consistent relationship with cortical areas, the decrease in accuracy is smaller, but this condition holds only for a minority of cortical regions (e.g., the central sulcus, the calcarine sulcus, and the insula). Even in these regions, there is a substantial amount of degradation in spatial localization precision in the volume (e.g., V1 in Ref. 96), and inaccuracy in mapping to a group average cortical surface (e.g., area 2 in Ref. 97). This makes neuroanatomically valid comparisons between group average data in MNI volume space and group average data on the surface inherently challenging'.

The problem manifests itself most strikingly when comparing data mapped to group average cortical surfaces (where matching between vertices is derived from areal feature-based registration) to data mapped to average structural images registered in the volume using nonlinear volume registration (in this case FNIRT, as in Ref. 17). Supplementary Fig. 12 shows the issue in a comparison between group average and individual data. In most cortical regions, group average surfaces and curvature maps are much blurrier than individual subject surfaces and curvature maps. The only exceptions are regions where there is strong consistency in folding patterns across subjects and between cortical areas and folds⁹⁶, such as the central sulcus, the calcarine sulcus, and the insula. This has an important impact on the group average surface contour, which shrinks from the geometric effects of averaging. This shrinkage can be approximately corrected for on the surface when performing computations on group average surfaces (e.g., computing gradients or resampling data) by also making use of group average vertex areas across the individual subjects' midthickness surfaces in their native physical volume spaces. This surface shrinkage makes mapping to or from group average surfaces into group average volume space (e.g., MNI standard space) highly problematic, as the surfaces will shrink and miss large portions of grey matter. To compare group average data from two studies while maintaining the neuroanatomical fidelity of the HCP-style paradigm, it is necessary to process both studies using the HCP-style neuroimaging analysis approach. This means cortical atlases should be surface-based and any averaging across subjects should occur on the surface after areal features have been aligned. Another approach, generating a surface from a group average volume template and registering it to a surface template using folding patterns, may avoid the shrinkage issue^{98,99}; however, surfaces previously produced using this approach are of poorer quality than individual subject surfaces owing to the blurring from tissue partial voluming effects on the volume template from inadequate cross-subject cortical alignment. There is also no guarantee that cortical areas will be aligned across templates, and the volume-based cortical data of interest will still be poorly aligned.

18. How can I compare surface-based data generated on other platforms to the HCP_MMP1.0 parcellation and other group average surface data?

Many investigators have carried out surface-based analyses using atlases and platforms other than fs_LR⁹⁷ and Connectome Workbench, but may be interested in comparing their results to HCP group average data, particularly the HCP_MMP1.0 cortical parcellation. This requires a mapping between the surfaces (individual or group average) on which the investigator's analysis has been carried out and the HCP group average surface mesh. The HCP_MMP1.0 parcellation and most other HCP group average datasets are represented on the '32k_fs_LR' mesh (32,492 vertices per hemisphere, with a 2 mm average vertex spacing on the midthickness surface). This mesh has good geographic correspondence between the left and right hemisphere surfaces as a result of landmark-constrained interhemispheric registration between the left and right hemispheres (fs_L and fs_R) of FreeSurfer's fsaverage atlas surface⁹⁷. For datasets analyzed in FreeSurfer, data from the fsaverage atlas surfaces can be mapped to the 32k_fs_LR mesh by resampling from the registered fsaverage atlas spherical surface to the 32k_fs_LR atlas surface (see https://wiki.humanconnectome.org/display/PublicData/HCP+Users+FAQ; entry #9 – "How do I map data between FreeSurfer (native or fsaverage mesh) and HCP's fs_LR mesh?")⁹⁷. Conversely, data on the 32k_fs_LR mesh can be mapped onto FreeSurfer surfaces by reversing the order of files called in the resampling command. In general, we encourage

Conversely, data on the 32k_fs_LR mesh can be mapped onto FreeSurfer surfaces by reversing the order of files called in the resampling command. In general, we encourage investigators to map data onto the 32k_fs_LR mesh (the correspondence between left and right hemispheres is useful in a variety of contexts), but recognize there may be project-specific reasons for mapping in the reverse direction. For investigators using other surface-based analysis platforms (e.g., SUMA, BrainVoyager, or BrainVisa), analogous inter-atlas transformations have yet to be generated but such methods will be posted to the HCP-Users FAQ if and when they are implemented.

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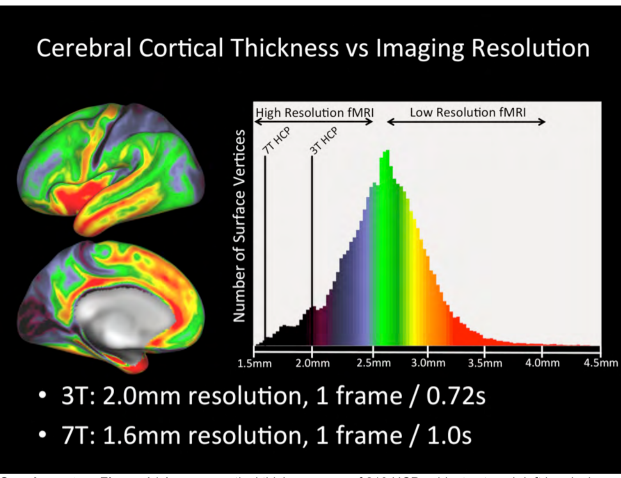
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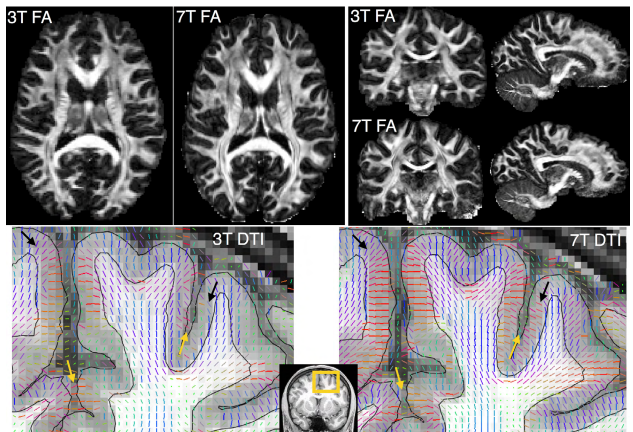
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III. Supplementary Figures

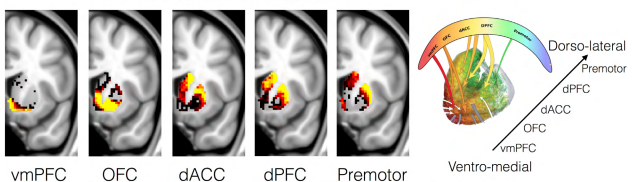


Supplementary Figure 1 | Average cortical thickness map of 210 HCP subjects at each left hemisphere vertex and the associated colorized histogram. The mean cortical thickness is around 2.6 mm, and this roughly divides fMRI data into high resolution (less than the mean cortical thickness) and low resolution (more than the mean cortical thickness). The HCP's 3T and 7T chosen resolutions are also plotted. For the data in this figure and other Supplementary Figures, subject recruitment procedures and informed consent forms, including consent to share de-identified data, were approved by the Washington University institutional review board.

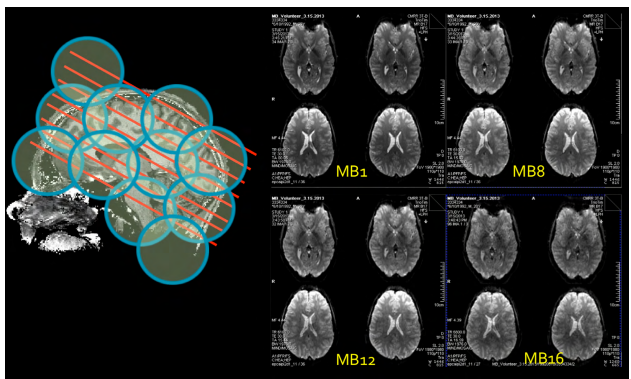


Supplementary Figure 2 | dMRI data for 3T and 7T scans of the same subject (HCP Subject 158035). Top row: Fractional anisotropy (FA) maps (axial, coronal and sagittal views). Notice that B1 inhomogeneities at 7T lead to poor SNR and noisy FA estimates in the inferior temporal regions (evident in the coronal views), but efforts have been taken to minimize them ¹⁹. Bottom row: DTI principal fiber orientations (coronal zoomed view of the area delineated by the yellow box). The orientations are RGB color-coded (Red: Left–Right, Green: Anterior–Posterior, Blue: Superior–Inferior) and superimposed on the structural T1w image. The pial surface and the WM/GM boundary surface are also shown. Reproduced, with permission, from Ref. 78.

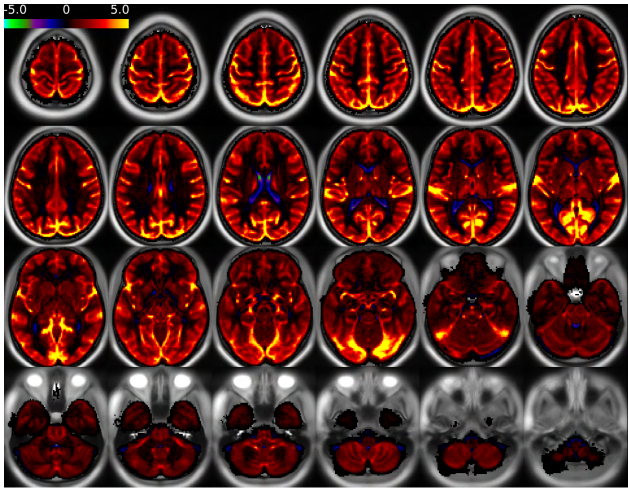
Cortico-Striatal Connections



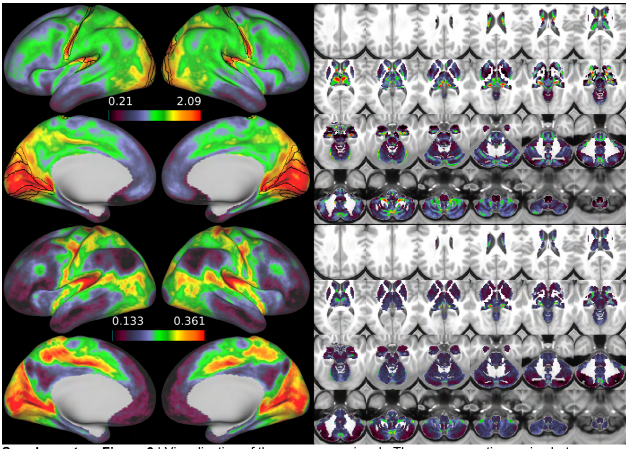
Supplementary Figure 3 | Patterns of cortico-striatal connectivity revealed by tractography. Seed locations were in different cortical regions, including vmPFC (ventromedial prefrontal cortex), OFC (orbitofrontal cortex), dACC (dorsal anterior cingulate cortex), dPFC (dorsal prefrontal cortex) and Premotor cortex. Path probabilities (yellow: high, red: intermediate, black: low) are obtained using probabilistic tractography (FSL's probtracx2, http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/FDT/UserGuide) and the Matrix 3 (bidirectional white matter voxel to gray-matter terminations) algorithm to compute dense connectomes. N=150 HCP subjects were analyzed and averaged. Note the strong similarity to patterns of tracer-based connectivity reported in the macaque monkey, shown in the sketch on the right (reproduced with permission from Ref. 100).



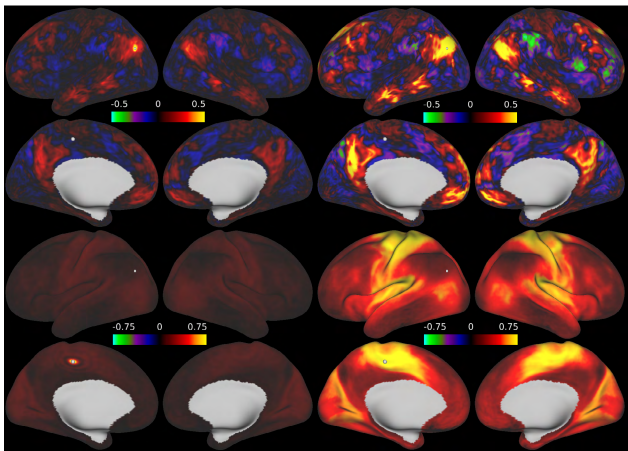
Supplementary Figure 4 | Multi-band imaging schematic and exemplar results. Left: A schematic representation of the array coil elements over a human head image and a multi-band (MB) excitation (8 slices, in red). Each coil detects a linear combination of signals from each slice weighted by the sensitivity profile of that coil. Right. Four slices from a whole brain data set with standard acquisition (upper left, MB1) and with SMS/MB acquisition with MB=8, MB=12, and MB=16, all obtained using a 32-channel coil on a 3T scanner. Visual inspection of the M=16 image reveals discernible artifacts (e.g., ghosting), whereas the MB=12 and MB=8 images appear much cleaner. Quantitative estimates of cross slice contamination and reconstruction noise can be made¹⁰¹. The HCP used MB=8 for fMRI and MB=3 for dMRI (now MB=4 or more for HCP Short).



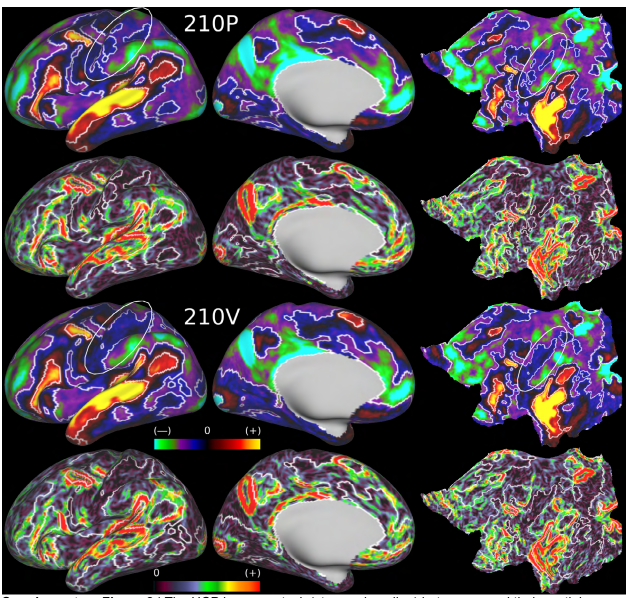
Supplementary Figure 5 | Beta map of the mean fMRI timeseries. The timecourses were averaged over the whole brain (including gray matter, white matter, and CSF), regressed into the data of each subject, and then averaged across (n = 210 HCP subjects). One particularly striking characteristic of this map is how tissue-specific the global signal is (after ICA+FIX data cleanup), being generally most positive in grey matter, close to zero in white matter, and slightly negative in the ventricles. The tissue specificity of this signal argues against a non-physiologic, non-BOLD contrast-based cause of the signal (e.g., direct biophysical effects of subject head motion). This map by itself does not tell us to what extent the global signal is physiological noise vs. neural signal. Although the data are averaged in the volume across subjects, they still appear relatively sharp because they are not smoothed. Data at http://balsa.wustl.edu/0L1m.



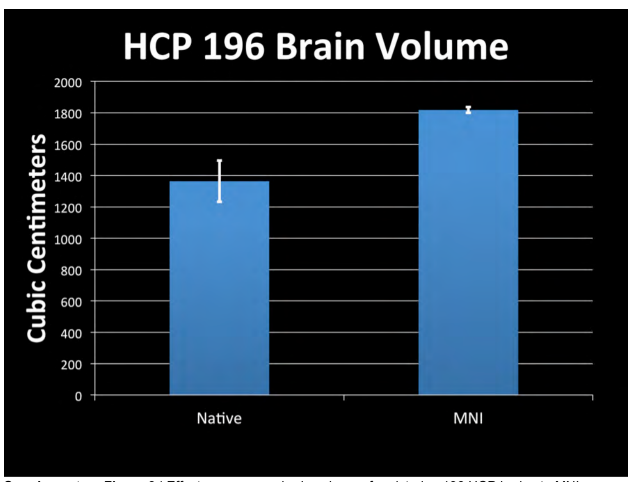
Supplementary Figure 6 | Visualization of the mean grey signal. The mean grey timeseries beta map (top rows) and the ratio of the variance of the mean grey timeseries to the total BOLD variance (i.e., variance classified as signal by ICA+FIX; bottom rows). The absolute magnitude of the mean grey signal is highest in sensory regions including visual cortex, early somatosensory cortex (particularly of the face), early auditory cortex, and several thalamic nuclei including the LGN/MGN. Visual, somatosensory, auditory, and likely vestibular cortical areas are highlighted with black outlines. Data based on averaging across 210 HCP subjects, aligned using MSMAll for the cortical surface. When the global signal is strong in individual subjects, it closely matches the group average pattern, however when it is weaker, it may or may not match the group pattern. In these cases, the global signal often looks like one or another widespread RSN (e.g., the task positive or task negative (default mode) network). This is further evidence that removing the global signal as a preprocessing step may distort resting state functional connectivity and hence that we need a more selective way to clean global noise out of the data. The bottom row is a relative measure of how much the fMRI timeseries will be altered by regressing out the mean grey timeseries (on average across subjects), as it is a measure of the proportion of the total BOLD signal represented by the mean grey timeseries. Regressing out the mean grey signal will also tend to cause resting state gradient boundaries to move somewhat in the regions where this map has sharp gradients³⁴. Data at http://balsa.wustl.edu/2VnN.



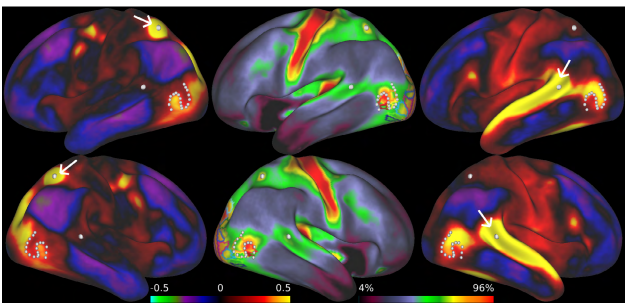
Supplementary Figure 7 | Effects of the Wishart rolloff on dense functional connectivity maps of both an individual subject and group data (210 HCP subjects; MSMAll surface registration). Top rows show an individual subject, before (column 1) and after (column 2) Wishart rolloff for a seed location in lateral parietal cortex (white dot in upper left panels). The correlation increases dramatically as unstructured spatio-temporal noise is reduced, however the map is not substantially "smoothed" as it would be with typical smoothing algorithms. Bottom rows show a group dataset before and after Wishart rolloff for a seed location in the posterior cingulate sulcus (white dot in lower left panels). The dataset has been created using the MIGP⁵⁷ algorithm to generate a group PCA series (d=4500) that represents the group concatenated timeseries. Because of the hard cutoff at PCA component number 4500, there is a 'ringing' pattern resulting from spatial autocorrelation in the spatio-temporal noise that is represented by the PCA components with the lowest eigenvalues. If a Gaussian filter had been applied, this pattern of "local connectivity" would be a blob instead of rings. The Wishart rolloff eliminates these rings and again dramatically increases the SNR of the data. Data at http://balsa.wustl.edu/rrpl.



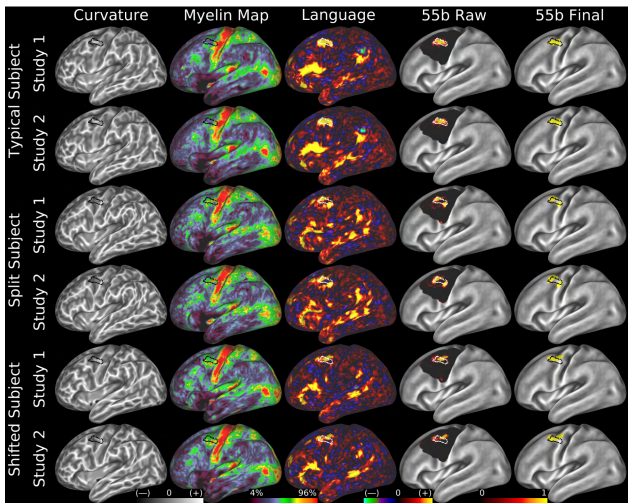
Supplementary Figure 8 | The HCP language task (story vs baseline) beta maps and their spatial gradients. Beta maps (rows 1 and 3) and gradient maps (rows 2 and 4) are from two independent groups of 210 HCP subjects, "210P" (rows 1 and 2) and "210V" (rows 3 and 4). Because of the large number of high quality HCP subjects, the beta maps are very similar across the two groups, and the strong gradients in the beta maps are also very similar. Also shown are white contours of a Bonferroni corrected significance threshold across all 91282 grayordinates (z+/-~5). Two things are apparent: 1) Because of the large amount of high quality data, most of the brain is either significantly activated or deactivated (an issue that has been discussed before, see Ref. 60). Thus the statistical threshold is not particularly biologically meaningful (a point about statistical thresholds that generally applies). At the same time, the statistical threshold is also not strongly reproducible, in spite of the large amount of high quality data (highlighted ellipses show large differences in the area of activation classified as "significant" that are not particularly impressive when viewing the unthresholded beta maps). In contrast, the strong gradients in the beta maps are much more reproducible, are likely also more biologically meaningful, and hence provide a better substrate for defining regions of activation or comparing across studies. Data at http://balsa.wustl.edu/PrmK.



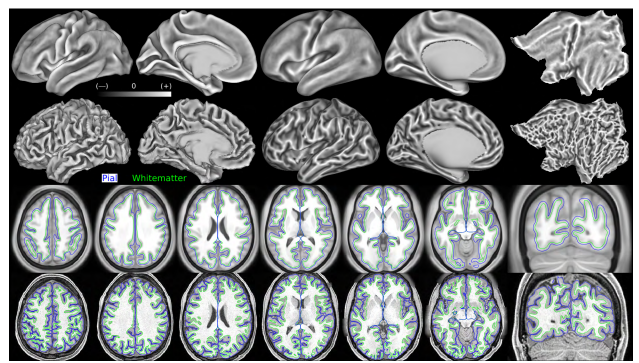
Supplementary Figure 9 | Effects on average brain volume of registering 196 HCP brains to MNI space. The total brain volume (minimal preprocessing pipelines' whole brain mask) measured in the subject's native physical space is around ~1350 cc; however, after registration to MNI space it is ~1800 cc, though the variability in brain volumes goes down as indicated by the narrower standard deviation error bars. This reduction in variability is the intended effect of registration, but the increase in brain volume is group average registration drift that was built into MNI standard space during the non-rigid registrations of the template generation process ⁶²⁻⁶⁴.



Supplementary Figure 10 | A comparison between the HCP data and published retinotopic parcellation data. Data from Ref. 64 and from 120 HCP subjects (from Q1-2) were registered using MSM areal-feature-based registration ¹⁰² and group average registration drift was removed from both. Because of this, a contour in functional connectivity in the MT+ region distinguishing strong connectivity to the heavily myelinated IPS hotspot (LIPv, column 1) and to the STS (column 3) lines up with the border between MT+V4t (orange and yellow) and MST+FST (red and maroon, middle column). This illustrates the kind of precise cross-modal, cross-study boundary comparisons possible using the HCP-Style paradigm. Data at http://balsa.wustl.edu/x2Lz.



Supplementary Figure 11 | Classification of area 55b in individual subjects by the areal classifier. The typical location of area 55b is shown in black or white outlines on the inflated left hemisphere surface. The top two rows show a subject having an area 55b in the typical location found in the population. Rows 1 and 2 are entirely separate 'test/retest' runs of this subject through the full HCP MRI acquisition and analysis protocol. Column 1 is the subject's individual curvature map, column 2 is the subject's myelin map, column 3 is a d=40 RSN map that shows strong connectivity between 55b and other areas of the language network, column 4 is the raw probabilistic output of the areal classifier, column 5 is the final output of the areal classifier. Rows 3 and 4 show a different subject whose 55b is atypically split (heavy myelination running through the population average location of 55b and a concomitant lack of resting state connectivity). In both runs through the protocol, this subject shows a split 55b that is accurately detected by the areal classifier, showing that the classifier can accurately classify atypical subjects and that these atypical patterns are stable across time. Rows 5 and 6 show a third subject whose 55b is atypically shifted relative to its neighbors. Again the pattern is stable across time and the areal classifier is able to accurately detect the area. Reprinted from Ref. 34. Data at http://balsa.wustl.edu/WPPn.



Supplementary Figure 12 | Effects of averaging surface coordinates and folding maps after areal feature-based registration (MSMAII). The first row shows the group average midthickness surface (left), and the group average curvature map also displayed on inflated and flat surfaces (center and right). The group consisted of 210 subjects. The second row shows an individual subject's midthickness surface (left), and the individual's curvature map displayed also on inflated and flat surfaces (center and right). Note how much less detailed the group average surfaces and curvature maps are in most regions of cortex. However, in regions with consistent folding patterns across subjects and consistent relationships between cortical areas and folds. the group average patterns remain sharp (e.g. the central and calcarine sulci and the insula). The third row shows the group average T1w volume (after FNIRT nonlinear volume registration to MNI152 space of each subject in the group) together with the group average white (green) and pial (blue) surface contours. The fourth row shows the same individual subject's T1w volume together with the individual's white and pial surfaces (after aligning both the T1w volume and the surfaces to the group average in MNI space using FNIRT nonlinear volume registration). Despite the high precision of the white and pial surfaces in following the grey matter ribbon in the individual, the group average surfaces do not follow the group average volume particularly well, except in the regions where there are consistent folding patterns across subjects and consistent relationships between cortical areas and folds (as mentioned above). These issues also occur with folding-based surface registration (not shown), though they are less severe, because for folding-based registration the dominant factor is inconsistency in folding patterns across subjects (as no attempt is made to directly align cortical areas). The midthickness surfaces are the average of the white and pial surfaces (this average is performed on each individual, and the group midthickness surface is the average of the individual midthickness surfaces). Data at http://balsa.wustl.edu/7gP3.