Ultra-high field (10.5 T) resting state fMRI in the macaque

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Abstract

Resting state functional connectivity refers to the temporal correlations between spontaneous hemodynamic signals obtained using functional magnetic resonance imaging. This technique has demonstrated that the structure and dynamics of identifiable networks are altered in psychiatric and neurological disease states. Thus, resting state network organizations can be used as a diagnostic, or prognostic recovery indicator. However, much about the physiological basis of this technique is unknown. Current approaches to resting state measurements in macaques face unique challenges associated with signal-to-noise, the need for contrast agents limiting translatability, and within-subject designs. These limitations can, in principle, be overcome through ultra-high magnetic fields. However, imaging at magnetic fields above 7T has yet to be adapted for fMRI in macaques. Here, we demonstrate that the combination of high channel count transmitter and receiver arrays, optimized pulse sequences, and careful anesthesia regimens, allows for detailed single-subject resting state analysis at high resolutions using a 10.5 Tesla scanner. In this study, we uncover thirty spatially
detailed resting state components that are highly robust across individual macaques and closely resemble the quality and findings of connectomes from large human datasets. This detailed map of the rsfMRI ‘macaque connectome’ will be the basis for future neurobiological circuit manipulation work, providing valuable biological insights into human connectomics.

Keywords
Functional connectivity; Rhesus macaque; Resting-state; Spontaneous activity; Functional MRI (fMRI)

1. Introduction

Resting state functional connectivity refers to the temporal correlations between spontaneous blood oxygenation level dependent (BOLD) signals obtained using functional magnetic resonance imaging (fMRI). These fluctuations were first noted in motor cortex (Biswal et al., 1995), but since then many other large-scale networks of correlated temporal patterns in the resting brain have been identified (Biswal et al., 2010; Heuvel and Pol, 2010; Smith et al., 2013; Wang et al., 2009). It was clear even before the advent of fMRI that functional networks at multiple spatial and temporal scales are embedded in the mammalian brain (Essen and Maunsell, 1983; Felleman and Essen, 1991); many years of fMRI neuroimaging have catalogued the spatial and, to some degree, dynamic organization (Allen et al., 2014; Hindriks et al., 2016; Ju and Bassett, 2020; Hutchison et al., 2013; Hutchison et al., 2013) of these networks. These different networks have distinct temporal properties (~0.01 - ~0.1 Hz fluctuations) and persist through different states such as sleep or anesthesia. Moreover, they are generally consistent across subjects (Smith et al., 2013, 2009) and to some degree generalize across species (Li et al., 2013; Jonckers et al., 2011).

As a method, characterization of resting state networks has proven invaluable for psychiatry (Floris et al., 2018; Konova et al., 2015; Hulvershorn et al., 2013), basic neuroscience (Fox and Raichle, 2007; Glasser et al., 2016; Smith et al., 2013), and psychology (Laird et al., 2011). In particular, it offers the ability to simultaneously characterize networks across the whole brain within a short period of time, and without the need to engage the subject in order to collect the data, as opposed to classical task-based experimental designs. These brain networks (and any individual variations) can then be studied across development, aging or disease and provide a high-throughput approach to both basic and clinical human neuroscience.

For several reasons, the resting state technique is a particularly important one to implement in the macaque monkey model. First, macaques offer a critical intermediary between humans and rodent models and have gross neuroanatomy that is highly homologous to that of humans (Hutchison and Everling, 2012; Mars et al., 2016; Petrides and Pandya, 2002; Watson and Platt, 2012; Heilbronner et al., 2016). Second, macaques are a valuable model for many complex behaviors and cognitive processes, and thus potentially for psychiatric diseases. Third, macaques offer the opportunity to perform types of invasive manipulations, including microstimulation, chemogenetics (Galvan et al., 2019; Nagai et al., 2016; Raper et al., 2019; Upright et al., 2018; Cushnie et al., 2020) and optogenetics (Galvan et al., 2017;
Khateeb et al., 2019), which are impossible or of limited applicability in humans. Fourth, additional measurement techniques that are rare or impossible in humans, like single-unit electrophysiological recordings and neuroanatomical tract-tracing using invasive tracers, have allowed for careful validation of non-invasive neuroimaging tools such as fMRI and the BOLD response (Logothetis, 2003; Logothetis et al., 2001) or diffusion-weighted-imaging for tractography (dMRI) (Bakker et al., 2012; Markov et al., 2012). Thus, findings in nonhuman animal models constrain theory and experimentation in human neuroscience. On the other hand, investigations using non-invasive tools in humans can guide us in designing and conducting experiments to elucidate the basic mechanism of those findings using invasive tools in animals.

Macaques, however, do have a critical disadvantage as a model organism for resting state fMRI (rsfMRI). That is, until now, our ability to collect high resolution, high signal-to-noise ratio (SNR) whole brain data in macaques has been extremely difficult. This is a multifaceted problem resulting mainly from the macaques’ small head size (~6% of the human brain volume), the use of standard field strength magnets, the challenges in doing alert studies (especially at high resolutions) and the unavailability of commercial high channel count surface coils arrays. Moreover, work on macaques is costly and labor intensive. Thus, sample sizes in classical macaque experiments are generally low (usually 2-4 subjects per experiment), which contradicts the notion of generalization and out-of-sample prediction in human studies. That, in turn, means the utility of neuroimaging in macaques would be greatly enhanced if we could develop ways to improve SNR and contrast-to-noise ratio (CNR) dramatically.

One tool to improve SNR is using exogenous blood-pool contrast agents that induce strong susceptibility gradients in their vicinity. Such agents include monocrystalline iron oxide nanoparticles (MION), or similar more modern agents (Ferumoxytol, Senirem, Feraheme) as a contrast agent (Moeller et al., 2009; Tsao et al., 2003; Vanduffel et al., 2001). However, this approach has several disadvantages: (1) the signal then reflects blood volume changes (Leite et al., 2002) rather than the typical blood oxygenation level dependent (BOLD) response, complicating its translatability to typical human studies, (2) it has a slightly slower hemodynamic response (Leite et al., 2002; Pelekanos et al., 2020), and (3) it can accumulate in the brain and causes potentially long-lasting health disadvantages (Li et al., 2012; Mustafa and Mohammed-Rasheed, 2019). The SNR problem can to some degree also be overcome with larger datasets (i.e. many hours of data collection) but this is expensive and impractical and is accompanied by unavoidable risks to the subjects if general anesthetics are utilized.

One way to overcome the sensitivity problem of acquiring high resolution fMRI is to use ultra-high field strengths in combination with high channel count receiver arrays (Lagore et al., 2020; Uğurbil et al., 2019), an approach which has proven extremely productive, to this end, in human studies (Martino et al., 2018, 2011; Ugurbil, 2014; Uğurbil et al., 2019, 2019, 2003; Yacoub et al., 2008, 2001; Zimmermann et al., 2011). Increasing magnetic fields not only provide enhanced SNR but also lead to higher BOLD contrast, especially for the microvascular BOLD contributions that produce mapping signals with higher fidelity to sites of neuronal activity (Uğurbil, 2018; Ugurbil, 2016). Thus, increasing the magnetic field produces multiplicative gains in BOLD based fMRI studies. Because of these advantages
and an increasing number of solutions introduced to tackle challenges confronted in imaging at such magnetic fields, the ultrahigh magnetic field of 7 Tesla has emerged as the preferred platform for non-clinical fMRI in humans (Uğurbil, 2018). Following this trend, recently, efforts have been undertaken to significantly surpass 7 Tesla in magnetic field strength and introduce magnetic fields greater than 10 Tesla.

While some macaque studies focused on resting state functional magnetic resonance imaging (rsfMRI) (Gilbert et al., 2016; Goulas et al., 2017; Hutchison et al., 2011; Hutchison et al., 2013; Schaeffer et al., 2018; Vincent et al., 2009, 2007) to date, to our knowledge, resolutions in macaque rsfMRI continue to be greater than 1 mm³ as well as longer than 1 second.

Here we explore the advantages of 10.5T neuroimaging for rsfMRI in the macaque. combined with a specialized 32 Rx and 8 Tx channel coil (Lagore et al., 2020), implemented on a 10.5T human scanner with a commercial console (Siemens). We demonstrate robust, within subject and group ‘connectome’ style BOLD-based rsfMRI data in 6 lightly anesthetized macaques (for a similar approach at 3T see Autio et al., 2020). Our approach is extremely robust and allows us to repeatedly and reliably measure intrinsic BOLD signals in macaques at high spatial and temporal resolution and with high SNR. Our findings, extending previous approaches, show robust resting state networks similar to human findings that are consistent between macaques and, importantly, identifiable robustly in the individual. Equally important, the components we identify adhere extremely well to anatomical architecture, as well as showing homogeneous sensitivity to cortex and subcortical regions. Our findings demonstrate the general utility that 10.5T imaging has for in vivo intrinsic BOLD investigations in macaques and non human primates in general.

2. Materials and methods

2.1. Animal preparation

We obtained data from 6 adult macaque monkeys (Macaca fascicularis: 4 female, and Macaca mulatta, 1 male and 1 female). Weights ranged from 3.0 kg to 9.0 kg. Experimental procedures were carried out in accordance with University of Minnesota Institutional Animal Care and Use Committee approval and in accord with the National Institute of Health standards for the care and use of non human primates. All subjects were fed ad libitum and pair housed within a light and temperature controlled colony room. Animals were not water restricted. None of the subjects had any prior implant or cranial surgery. Animals were fasted for 14-16 hours prior to imaging.

On scanning days, anesthesia was first induced by intramuscular injection of atropine (0.5 mg/kg) and ketamine hydrochloride (7.5 mg/kg). Subjects were transported to the scanner anteroom and intubated using an endotracheal tube. Initial anesthesia was maintained using 1.5-2% isoflurane mixed with oxygen (1L/m during intubation and 2L/m during scanning to compensate for the 12m length of the tubing used).

Subjects were placed onto a custom-built coil bed with integrated head fixation by placing stereotactic ear bars into the ear canals. The position of the animal corresponds to the sphinx
position. For functional imaging, the isoflurane level was lowered to 1%. Experiments were performed with animals freely breathing. Additionally, for Macaca fascicularis subjects only, an initial bolus injection of 1.5ug/kg fentanyl was administered IV followed by a continuous administration of 3ug/kg/hr using a syringe pump. Rectal temperature (~99.6°F), respiration (10-15 breaths/min), end-tidal CO2 (25-40), electrocardiogram (70-150 bpm) and SpO2 (>90%) were monitored using an MRI compatible monitor (IRADIMED 3880 MRI Monitor, USA). Temperature was maintained using a circulating water bath as well as chemical heating pads and padding for thermal insulation. Anesthesia was used to eliminate motion effects and physiological stress as well as allow the imaging of surgically naïve subjects. Isoflurane is a vasodilator (Farber et al., 1997) and modulates cerebrovascular activity (Vincent et al., 2007). However, previous studies have shown successful synchronous BOLD fluctuations and resting state activity in macaques as well as mice and rats under isoflurane (Hutchison et al., 2011; Hutchison et al., 2013; Okada et al., 2018; Vincent et al., 2009, 2007).

2.2. Data acquisition

All data were acquired on a passively shielded 10.5 Tesla, 88 cm diameter clear bore magnet coupled to Siemens gradients (“SC72” body gradients operating at a slew rate of 200 mT/m/s, and 70 mT/m maximal strength) and electronics (Magnetom 10.5T Plus) (Siemens, Erlangen, Germany). Within the gradient set and the bore-liner, the space available for subject insertion had a 60 cm diameter.

The 10.5T system operates on the E-line (E12U) platform which is directly comparable to clinical platforms (3T Prisma/Skyra, 7T Terra). As such, the user interface and pulse sequences were identical to those running on clinical platforms. A custom in-house built and designed RF coil with an 8-channel transmit/receive end-loaded dipole array of 18 cm length (individually) combined with a close-fitting 16-channel loop receive array head cap, and an 8-channel loop receive array of 50 × 100mm size located under the chin (Lagore et al., 2020). The size of 14 individual receive loops of the head cap was 37 mm with 2 larger ear loops of 80 mm - all receiver loops were arranged in an overlapping configuration for nearest neighbor decoupling. The resulting combined 32 receive channels were used for all experiments and supported 3-fold acceleration in phase encoding direction. The coil holder was designed to be a semi-stereotaxic instrument holding the head of the animal in a centered sphynx position via customized ear bars. The receive elements were modelled to adhere as close to the surface of the animal’s skulls as possible. Transmit phases, for the individual transmit channels were fine-tuned for excitation uniformity for one representative mid-sized animal and the calculated phases were then used for all subsequent acquisitions. Magnetic field homogenization (B0 shimming) was performed using a customized field of view with the Siemens internal 3D mapping routines. Multiple iterations of the shims (using the adjusted FOV shim parameters) were performed and further fine adjustment performed manually on each animal. Third order shim elements were ignored for these procedures.

In all animals a B1 + (transmit B1) fieldmap was acquired using a vendor provided flip angle mapping sequence (example result in Sup. Fig. 9) and then power calibrated for each individual. Following B1 + transmit calibration, 3-5 averages (23 minutes) of a T1 weighted
magnetization prepared rapid acquisition gradient echo protocol (3D MP-RAGE) were acquired for anatomical processing (TR = 3300 ms, TE = 3.56 ms, TI = 1140 ms, flip angle = 5°, slices = 256, matrix = 320 × 260, acquisition voxel size = 0.5 × 0.5 × 0.5 mm³). Images were acquired using in plane acceleration GRAPPA = 2. A resolution and FOV matched T₂ weighted 3D turbo spin echo sequence (variable flip angle) was run to facilitate B₁ inhomogeneity correction.

Before the start of the functional data acquisition, five images were acquired in both phase encoding directions (R/L, L/R or F/H, H/F as appropriate) for offline EPI distortion correction. For four monkeys, six runs of 700 continuous 2D multiband (MB) EPI (Moeller et al., 2010; Setsompop et al., 2012; Uğurbil et al., 2013) functional volumes (TR = 1110 ms; TE = 17.6 ms; flip angle = 60°, slices = 58, matrix = 108 × 154; FOV = 81 × 115.5 mm²; acquisition voxel size = 0.75 × 0.75 × 0.75 mm³) were acquired. Images were acquired with a left-right phase encoding direction using in plane acceleration factor GRAPPA = 3, partial Fourier = 7/8th, and MB or simultaneous multislice factor = 2. For the two other monkeys, five runs of 700 continuous 2D MB EPI functional volumes were acquired (TR = 1200 ms; TE = 17.6 ms; flip angle = 60°, slices = 58, matrix = 106 × 212; FOV = 79.5 × 159 mm²; acquisition voxel size = 0.75 × 0.75 × 0.75 mm³). Images were scanned in foot-head phase encoding direction using in-plane acceleration factor GRAPPA = 3, partial Fourier = 7/8th and MB = 2. The change in sequence parameters was made after upgrading to the 10.5 Plus (E12U) platform which brought with it software changes that made the left right phase encoding more favorable. Since macaques were scanned in sphinx positions, the orientations noted here are what is consistent with a (head first supine) typical human brain study (in terms of gradients) but translate differently to the actual macaque orientation. While left / right orientation is the same, foot-head is actually anterior-posterior in the monkey.

2.3. Image preprocessing

Image processing was performed using a custom pipeline relying on FSL (Jenkinson et al., 2012), ANT's (Avants et al., 2014, 2011), AFNI (Cox, 1996) and a heavily modified CONN (Whitfield-Gabrieli and Nieto-Castanon, 2012) toolbox. Images were first motion corrected using mcflirt (registration to the first image). Motion parameters never exceeded 0.5 mm or 0.4° rotation except in one run of one animal, which was discarded (the animal experienced anesthesia induced emesis during the last run). Images were slice time corrected and EPI distortion corrected using topup. High magnetic fields and large matrix sizes are commonly associated with severe EPI distortions. Supplementary Fig 1 shows an overview of the extent of the distortion as well as the result of the corrected warps. Anatomical images were nonlinearly warped into the NMT (Seidlitz et al., 2018) template using ANT's and 3DQwarp in AFNI. The distortion correction, motion correction and normalization were performed using a single sinc interpolation. Images were spatially smoothed (FWHM = 2 mm), linear detrended, denoised using a linear regression approach including heart rate and respiration, as well as a 5 component nuisance regressor of the masked white matter and cerebrospinal fluid and band-pass filtering (0.008 - 0.09 Hz) (Hallquist et al., 2013). While we chose here to present the data as is typically done in resting state applications, to demonstrate the raw
SNR and overall BOLD sensitivity we show an unsmoothed example on the individual macaques brain in Fig. 1. A raw tSNR map of one animal can be found in Sup. Fig. 10.

2.4. Independent component analysis

We used group ICA to uncover ordered reproducible components across subjects. This approach uncovers a set of interpretable group components. Algorithms were used as implemented in the GIFT (Calhoun et al., 2001) software package. Our preprocessing pipeline produced one nuisance corrected concatenated time course of all runs per subject. The temporal dimension of the concatenated time course was then reduced using principal component analysis (PCA). Spatial resting state components were then estimated using the Infomax algorithm approach with the standard GIFT parameters. Time-courses and spatial maps associated with each component were then back projected as implemented in the group-ICA tools in GIFT. Since the process of dimensionality reduction and model selection are somewhat arbitrary (one needs to specify the number of components to estimate), there is no real consensus on how many components to extract. It has been noted that there is no best dimensionality for the underlying neurophysiology of multiple distributed systems (Cole et al., 2010). There are always multiple valid solutions, but we argue that a higher model order (Abou-Elseoud et al., 2010; Kiviniemi et al., 2009; Smith et al., 2009) than previously used (Hutchison et al., 2011; Hutchison et al., 2013; Vincent et al., 2007) is beneficial in macaques. While one can compartmentalize the brain into networks with associations to visual, motor, sensory, and auditory processing, the goal in macaque work is usually shifted towards finer-grain coding to allow for better cross-method comparison. Previous papers have not ignored this but have rather been limited by data quality (e.g., a high dimensional ICA decomposition cannot be obtained from a short rsfMRI run at standard field strength using high resolution). Multiple criteria (Jafri et al., 2008; Li et al., 2007; Zuo et al., 2010) currently exist for optimally selecting the number of independent components for a given dataset. The minimum description length criterion, however, yielded a mean estimation of 703 (SD = 69) independent components for our dataset. To obtain a manageable number of components and to allow for easier comparison with human and previous macaque datasets, 40 components were extracted. To test for the reliability of the decomposition the ICASSO (Li et al., 2007) toolbox was used and ICA iterated 20 times. Lastly, mean group independent components were scaled to empirically derived z-scores. These z-scores are an approximation of the temporal correlation between each voxel and the associated components time-course. A threshold of +/-2 was used as a lower limit of functional connectivity for visualization.

2.5. Identification of resting state networks and visualization

Components resulting from the independent component analysis were manually inspected and labeled according to anatomical and functional locations. Since ICA can be extremely noise sensitive, some extracted components are typically associated with motion, scanner noise artifacts or physiological confounds. In our dataset, 6 out of 40 components were discarded as they showed noisy, nonspecific, low correlation activation patterns or correspondence to large veins. Data was visualized in volume space (custom code written in c++ and openGL) or projected to the surface reconstruction of the NMT template (Seidlitz Yacoub et al. Neuroimage. Author manuscript; available in PMC 2020 December 17.
et al., 2018) using AFNI (Cox, 1996) and SUMA (Saad et al., 2004; Saad and Reynolds, 2012).

2.6. Single subject ICA

Since studies in macaque usually have low numbers of subjects, and because subtle individual features of resting-state connectivity can be lost in group analysis, we also performed single subject ICA (example of ICA components extracted from data not subjected to spatial smoothing in Fig. 1) Parameters were matched to those of the group ICA.

2.7. Independent component network connectivity

Temporal correlations between components resulting from spatial ICA analysis as performed here can be high (Calhoun et al., 2003), since our approach to ICA maximizes the statistical independence in space while reducing the dimensionality of the temporal domain. To explore the relationship between the independent components from a temporal perspective we performed functional network connectivity analysis (FNC) (Jafri et al., 2008) between our resulting ICs. Our ICs were further clustered using a purely data driven hierarchical clustering procedure (nonspatial complete-linkage clustering). FNC then analyzes the entire set of connections between pairs of networks in terms of the within- and between-network connectivity sets. Resulting F-statistics were used to test statistically significant connections (connection threshold p<0.05 p-uncorrected, cluster threshold p<0.05 p-FDR corrected using MVPA omnibus test).

3. Results

3.1. Quantitative image metrics

Since this study presents results from a novel instrument (10.5T) we report simple image metrics. Supplementary Fig. 10 presents representative raw tSNR metrics from one functional run of one subject. Next, we evaluated the mean fraction of outliers per fMRI volume for all subjects (AFNI 3dToutcount with alpha = 0.001). A mean of $4.4851 \times 10^{-4}$ with a standard deviation of $9.8672 \times 10^{-4}$ voxels were selected as outliers. Additionally, we investigated the physiological noise regime of our data by computing functional network connectivity on the unsmoothed data while including 1-5 acquired rsfMRI runs incrementally. Supplementary Fig. 11 demonstrates that z-scored connectivity values increase as a function of runs included in the analysis indicating that our data is not noise dominated by physiological noise.

3.2. Group resting state networks

Group-ICA decomposed the dataset of 6 monkeys into 40 independent components as specified. ICASSO returned a stability index of 0.96 (SD = 0.04) demonstrating that the components are close to orthogonal clusters and highly consistent across multiple ICA iterations. Out of the 40 extracted components, 30 were deemed physiologically relevant, containing more than one anatomical area, adhering to the gray matter as well as displaying a reasonable heavy-tailed frequency distribution. The spatial maps of the resting state networks (RSNs) obtained via ICA analysis are shown in Figs. 2 and 3. The order of
components is non informative. The computed components account for 56.39% of the data’s variance. An overview of the cortical coverage, as well as the overlap of the components can be found in supplementary Fig. 2. The resulting 30 RSNs are described below following anatomical classification from the Saleem and Logothetis atlas (Saleem and Logothetis, 2012):

- **Network 1 (lateral occipital):** This RSN is mainly in the lateral occipital cortex. It covers portions of early visual areas V1, V2, and V4. The network also includes anticorrelations to regions within the intraparietal sulcus.

- **Network 2 (dorsal superior temporal sulcus):** This RSN includes the areas surrounding the dorsal portion of the STS, including areas 7a, FST, MT, and MST.

- **Network 3 (somatosensory):** This RSN is primarily composed of somatosensory cortical regions, extending somewhat into the primary motor cortex as well. Anticorrelations with insula and lateral intraparietal area are observed.

- **Network 5 (pulvinar-pons-cerebellum):** This RSN is primarily subcortical and includes portions of the dorsal thalamus, particularly the medial and lateral pulvinar, much of the pons, and part of the cerebellum.

- **Network 6 (medial occipital):** This RSN covers medial areas of the occipital lobe. It is primarily composed of area V2, but extends into aspects of areas V6 and, more rostrally, 7m.

- **Network 7 (lateral and cingulate sulci):** This RSN follows the lateral sulcus closely, and thus covers primary auditory cortex, some auditory association areas, area 7op, and large portions of the insula. Although poorly visible on the surface, this network has substantial connectivity with cingulate areas 23c and 24c.

- **Network 8 (ventromedial subcortex):** This RSN covers a large swath of the ventromedial subcortical surface of the brain, including amygdala, basal forebrain, and hippocampus. Cortically, correlations in anterior cingulate cortex 24a/b as well as anticorrelations in nearby 23a/b are observed.

- **Network 9 (middle STS-dlPFC-cingulate):** This RSN covers the middle portions (dorsal-ventrally) of the superior temporal sulcus. This covers many areas associated with multisensory processing, including TPO, TAA, TEO, PGA, and some of the insula. The network also includes portions of the anterior and posterior cingulate gyrus, the outer edges of the intraparietal sulcus, and the dorsolateral prefrontal cortex (specifically areas 46 and 8a). Anticorrelations are seen with visual areas V2 and V6.

- **Network 11 (striatal-medial cortical):** This RSN covers the entire striatum, as well as some medial cortical areas, including precuneus, V6, and dorsomedial frontal areas 6M and 8M. Additional connectivity to orbitofrontal cortex (area 13) and anticorrelations with frontal pole and midbrain are present.
Network 12 (cerebellar-frontal pole): This RSN covers the cerebellum and ventromedial frontal pole.

Network 13 (dorsomedial cortex): This RSN covers a large portion of the dorsomedial cortex, including areas of the frontal lobe (medial areas 9, 8, 6, 4, and 24) and parietal lobe (somatosensory cortices, precuneus, and are 23).

Network 14 (intraparietal sulcus): This RSN covers much of the intraparietal sulcus, including LIP, VIP, and MIP. Rostrally, it extends into the insula.

Network 15 (insula): This RSN is mostly situated in the insula and adjacent operculum. Dorsally, it bleeds into the somatosensory and premotor cortices; ventrally and rostrally, it bleeds into the ventrolateral prefrontal cortex.

Network 16 (caudal dorsomedial cortex): This RSN is in the dorsal parietal and occipital lobes. It includes portions of V1, V3, V4, V6, precuneus, Opt, LIP, and PEA.

Network 19 (orbitofrontal cortex): This RSN covers nearly all of the orbitofrontal cortex, spanning from orbitofrontal regions of the frontal pole to ventral insula regions of the orbitofrontal cortex. The network extends somewhat dorsally into the medial prefrontal cortex (mainly area 32).

Network 20 (midbrain): This RSN is centered in the midbrain, but extends into portions of the thalamus, hypothalamus, cerebellum, basal forebrain, and pons.

Network 21 (posteromedial cortex): This RSN covers a large portion of the posteromedial cortex, including posterior cingulate areas 23 and 31, parietal area precuneus, and, caudally, some of V6 and V2. There are smaller lateral regions in this network as well: portions of TPO, 7a, and LIP are present. Finally, much of the intraparietal sulcus shows anticorrelations to this RSN.

Network 23 (sensorimotor): This RSN is situated centrally and dorsomedially, spanning the border between the frontal lobe and parietal lobe. Thus, it includes primary motor cortex, primary somatosensory cortex, supplementary motor area, premotor cortex, anterior and posterior cingulate cortices, and PE.

Network 24 (lower brainstem): This RSN is situated in the pons and medulla.

Network 25 (mid-dorsal cingulate): This RSN is mainly situated in the caudal anterior cingulate cortex (area 24’, sometimes called the midcingulate cortex) and posterior cingulate cortex (areas 23 and 31).

Network 26 (dorsal thalamus): This RSN is situated in the dorsal portion of the thalamus.

Network 30 (dorsal parietal cortex): This RSN is located in the dorsal parietal cortex, and covers the intraparietal sulcus, including areas LIPv, LIPd, area 5 and MIP. It extends towards the medial wall, covering areas V6, precuneus, and parts of the posterior cingulate cortex.
Network 31 (ventral thalamus): This RSN is located in the ventral thalamus, with some extension into the midbrain.

Network 32 (anterior cingulate-insula): This RSN covers the entire anterior cingulate cortex (area 24), much of the insula, and a small portion of the lateral parietal cortex (7op and PFG).

Network 33 (posterior cingulate-lateral parietal cortices): This RSN covers the posterior cingulate cortex (areas 23 and 31) as well as lateral parietal areas 7op and Tpt. There is also a small area of correlation in the temporal pole.

Network 35 (dorsolateral-cingulate-thalamus): This RSN includes areas in the dorsolateral prefrontal cortex such as area 8 as well as the anterior cingulate cortex area 24 and some rostral thalamus. Anticorrelations to posterior cingulate cortex and precuneus regions are observed.

Network 36 (lateral prefrontal right): This RSN covers the right lateral prefrontal cortex around the principle and arcuate sulcus. Anticorrelations with the medial prefrontal cortex, such as area 9 and 25, as well as part of the anterior cingulate cortex are observed.

Network 38 (lateral prefrontal left): This RSN covers the lateral left prefrontal cortex around the principle and arcuate sulcus.

Network 39 (sensorimotor 2): This RSN covers both sides of the central sulcus, and thus involves both motor and somatosensory cortices. The network also includes anticorrelations of posterior cingulate cortex and more caudal somatosensory areas.

Network 40 (frontal pole): This RSN is mainly in the frontal pole. Caudally, it extends into the dorsal prefrontal cortex and anterior cingulate cortex.

3.3. Individual resting state networks

Since group analysis can obscure individual anatomical and functional details, we present all individual ICA results in supplementary Figs. 3-8. Fig. 4 shows the consistency of our single subject independent component results in all subjects for four representative components. Note the consistency in spatial location at the same threshold. This finding demonstrates the consistency and data quality of our imaging approach and shows that our group level findings are highly representative of the individual monkey.

Projection to surface space can additionally obscure results and is generally ill-suited for use in individual animal physiological use. Fig. 5. shows 7 of the 30 components displayed from one representative single subject volume space projected onto the NMT template. The results demonstrate the close correspondence of the components with anatomical and functional landmarks as well as specificity to both cortical and subcortical regions.

3.4. Independent component network connectivity

Data driven hierarchical clustering of the independent component time courses revealed 9 clusters of components, with strong positive functional connectivity within cluster and
between adjacent clusters, and strong negative functional connectivity between non-adjacent clusters. Even though clustering was not based on spatial information, the resulting adjacencies reveal anatomical neighboring relationships (Fig. 6). Clusters fall into the following categories: 1. Subcortical components, 2. Prefrontal components, 3. Visual components, 4. Parietal components, 5. Superior-temporal components and 6. Motor components.

4. Discussion

Here we present the first demonstration of connectome style quality resting state fMRI data in the macaque at ultra-high magnetic fields significantly beyond 7T. The delineated RSNs closely resemble the quality and findings of connectomes from large human datasets, providing a detailed map of the rsfMRI ‘macaque connectome’ as a basis for future interventional work.

Resting state functional connectivity patterns are an extremely important tool used in current neuroscientific research into the origin of mental disease states (Bai et al., 2011; Lawrie et al., 2002; Lustig et al., 2003; Quigley et al., 2001; Xie et al., 2012; Zilverstand et al., 2018). Yet, much of this technique’s physiological basis is not well understood. Since macaques offer an extremely promising translational model (Heilbronner and Chafee, 2019; Krubitzer and Huffman, 2000; Consortium et al., 2020; Hutchison et al., 2011; Hutchison and Everling, 2012; Milham et al., 2018), a detailed map of the rsfMRI ‘macaque connectome’ is needed to build the basis for future interventional work. While human rsfMRI studies have made significant advances in the ability to acquire highly sensitive and informative images of functional connectivity (Essen et al., 2012; Glasser et al., 2016; Harms et al., 2018), including at relatively high spatial and temporal resolutions (Moeller et al., 2010; Vu et al., 2016), similar non-invasive neuroimaging advances in the macaque have not been as widespread or productive.

Several barriers have previously prevented such translations from being successful. Since invasive work, such as single unit recording, anatomical dissection, or chemo- and optogenetic manipulations in the macaque is usually done on the individual subject level at high spatial and temporal scales, extremely robust within individual rsfMRI maps are needed. In vivo experiments using very high magnetic fields facilitate such high spatial-temporal resolutions and overall sensitivity across the entire brain. Next, high channel count coils with sufficient parallel imaging performance, specifically designed for macaque setups (Autio et al. 2020; Autio et al. this issue), are also essential for achieving the translational sensitivity and spatial-temporal resolutions. The use of parallel imaging is essential for reducing the volume acquisition times and hence increasing the temporal sampling rate of the fMRI time series. However, this results in a spatially non-uniform increase in the noise of the measurement through the so called g-factor noise (Pruessmann et al., 1999). The g-factor noise is mitigated with higher channel counts and independently with increasing magnetic fields (Ohliger et al., 2003; Wiesinger et al., 2006, 2004). Thus, the two approaches employed together, as in this 10.5T study, work synergistically to provide multiplicative gains, enabling high accelerations with suppressed g-factor noise (Uğurbil et al., 2019).
Further, even though human studies are typically completed within a couple of hours and macaque studies have the luxury of extending well beyond this to increase overall sensitivity, extending session duration can still be problematic in terms of physiological (and therefore fMRI) stability. For this reason, shorter scans with higher SNR are still desirable.

Another approach to achieving sensitivity with shorter scan durations is the use of contrast agents, which greatly improves lower field studies. However, the use of contrast agents has significant limitations as well, such as: 1) It does not measure intrinsic BOLD signals (which is what the vast majority of human studies use), 2) studying temporal dynamics or paradigms other than block designs is much more difficult, and 3) there are risks associated with its use and it is hence not used in non-clinical human studies.

While implementation on a human-based scanner is not absolutely necessary, it does provide a convenient avenue by which imaging methods and sequences routinely used in the human can be applied to the macaques (Autio et al. this issue). Finally, while obtaining high resolution functional images of the alert monkey would be more ideal than the low anesthesia strategy we employed here, this is still an enormous technical challenge that we and others are working to overcome. Despite this, our data demonstrate 10.5T coupled with high channel count arrays can achieve submillimeter acquisitions using the advanced accelerated sequences developed for the Human Connectome Project (Uğurbil et al., 2013). Further, the setup described here could be used for other applications, such as task fMRI.

The macaque connectome has been studied previously (Hutchison et al., 2011) using the ultrahigh magnetic field of 7 Tesla with 1.3 × 1.3 × 1.5 mm\(^3\) resolution and 3 mm FWHM spatial smoothing, revealing 11 macaque RSNs using group ICA analysis. Here, following this earlier work (Hutchison et al., 2011) we report the first whole brain (0.75 mm\(^3\) isotropic at ~1.2s TR, smoothed at 2mm) comprehensive group independent component analysis in macaques at 10.5T. Although our data was of sufficient quality to produce RSNs without spatial smoothing, in this first 10.5T study, we employed spatial smoothing following coming practices employed in the resting state fMRI literature. Our approach identified 30 reproducible RSNs covering the entire macaque cortex as well as subcortical contributions throughout the thalamus, brainstem and cerebellum. The RSNs we were able to identify closely resemble higher model order RSNs identified in large sample sizes of human experiments (Abou-Elseoud et al., 2010; Kiviniemi et al., 2009; Li et al., 2007; Smith et al., 2013, 2012, 2009). In these findings, high model order RSNs are a tradeoff between the number of components extracted and the detectability of unique networks observed. Extremely high model order in ICA produces a larger number of networks but fails to uncover additional brain regions involved in a network. Our networks represent an intermediary between extremely high model orders (Moeller et al., 2009) and more conservative approaches (Hutchison et al., 2011), covering nearly the entire brain (see Fig. 7 and supplementary Fig. 2) while uncovering networks that include long range functional connections. Using this approach, we were able to demonstrate remarkable reproducibility of the RSN decomposition even in the individual animal without the use of contrast agents. This achievement is of utmost importance when trying to understand and model the relationship between real anatomical neural connections only obtainable invasively and functional connectivity based on neuroimaging. Our RSN findings reveal very similar...
network organization between humans and macaques at the mesoscopic level (see Fig. 7 for a qualitative comparison). We observe RSNs in all of the networks that are commonly identified in human experiments (Beckmann et al., 2005; Jafri et al., 2008; Smith et al., 2013, 2009). For example, we identify multiple networks that resemble the commonly described fronto-temporal, default-mode, fronto-parietal, cerebellar, visual, somato-motor, intraparietal, lateral prefrontal, basal ganglia, insular, and cingular networks (Fig. 7).

Previous work found 20 stimulus evoked components (Moeller et al., 2009) using contrast agents or 11 high quality intrinsic components (Hutchison et al., 2011). Extending this work, we were able to resolve previously not observed (Hutchison et al., 2011; Moeller et al., 2009; Vincent et al., 2009, 2007) lateralized dorsolateral prefrontal components (RSN 36,38) that have readily been described in the human literature and attributed to language, memory and cognitive attentional processes (Beckmann et al., 2005; Jafri et al., 2008; Smith et al., 2009). Our lateralized network does not include the parietal contribution present in the human literature, however. Another set of components that have previously remained elusive (Vincent et al., 2009, 2007) in the macaques are the dorso medial and orbital contributions of the prefrontal cortex. In our findings (RSN 19, 40) these were readily present. It is difficult to assess why these findings have previously not been reported, but SNR limitations as well as susceptibility artefacts close to the orbits and sinuses have likely contributed to the problem. One difference in our findings with the human literature is the absence of a large component covering the primary visual cortex’s caudal extent. Our visual cortex related RSNs (RSN 1,6) cover both medial and lateral portions of V1, V2 but miss the occipital pole. This is likely the result of our coil having the lowest sensitivity in visual cortex due to the macaque’s sphinx position. Another difference in our findings compared to previous macaque rsfMRI studies (Hutchison et al., 2011; Moeller et al., 2009; Vincent et al., 2009, 2007), but in agreement with high model order RSN studies (Abou-Elseoud et al., 2010; Kiviniemi et al., 2009) as well as clustering approaches (Leech et al., 2012, 2011), is the fractioning of RSNs that are associated with the default mode network. In our findings multiple RSNs (21, 26, 30, 31, 33, 39) show at least partial overlap with posteromedial cortex. Another finding that has previously not been reported is the detailed delineation of the cingulate cortex that can be observed in our RSNs. RSNs (5,8,9,11,13,21,24,25,32,33,35,39) all exhibit involvement of the cingulate cortex partitioning the anatomical areas 23a,b,c and 24a,b,c. While some of these networks demonstrate overlap in their cingulate cortex involvement, unique partitioning is directly observable in others (RSNs 5,24,25,26,35). Again, it is unclear why these effects were not previously found but inhomogeneous excitation from local transmitters or low SNR in medial regions further away from the scalp (especially true if headposts are used), and, compared to our study, lower BOLD contrast, could contribute to this omission.

To address one of our motivations, the robust identification of RSNs within the individual subject, single subject ICA using the same model order of 40 was performed and assessed for reproducibility and compared to the group results. Compared to previous studies (Hutchison et al., 2011) we found remarkable reproducibility for most of the 30 resulting RSNs. First, using the same thresholds as in the group-ICA resulted in more widespread diffuse activity, yet the pattern and anatomical correspondence of the main group-ICA findings were largely preserved in the individual. It is difficult to compare our current
approach to previous findings since we used diffeomorphic transformations for normalization that likely resulted in better anatomical correspondence and acquired significantly more data in an individual subject.

Taken together, we have achieved a highly detailed, individually robust delineation of RSNs in the macaque that closely resembles the quality and findings of connectomes from large human datasets, further demonstrating the benefits of using extremely-high magnetic fields in vivo.

One countering recent argument and effort (Consortium et al., 2020; Milham et al., 2018) that has seen high translational success has been the cross laboratory pooling of macaque fMRI datasets. We argue that our rationale of advocating high field macaque neuroimaging does not negate or change the need for such databases, but rather aims at providing data and solutions for a different problem. The big data consortium approach allows for the study of general principles derived from rsfMRI data in macaques such as physiological states (Xu et al., 2019) (anesthetized vs. awake) or macroscale area organizations (Xu et al., 2018) within the species. These are incredibly important efforts that can also serve an important translational role. What these approaches cannot provide, and where we believe extremely-high field comes into play, is measuring the effects of within subject manipulation using invasive tools. If we are interested in figuring out how structural connections and their electrical and chemical changes enable functional connectivity in rsfMRI and behavior, disease, or developmental processes, resting state neuroimaging at high magnetic fields coupled with within-subject invasive manipulations will pave the way.

5. Conclusions

To our knowledge, this is the first detailed demonstration of a macaque connectome (or HCP) style resting state dataset, which has been available for human applications for around 10 years. While previous macaque resting state studies have demonstrated typical RSNs, none have achieved whole brain sub-millimeter acquisitions with an approximate 1 sec temporal resolution, let alone doing it using only intrinsic BOLD signals and with a scan duration of around an hour. Data sets such as the current one allow for more straightforward translations of macaque data to the human for the purposes of understanding brain connectivity in individuals and how it is altered in disease.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Bibliography

Abou-Elseoud A, Starck T, Remes J, Nikkinen J, Tervonen O, Kiviniemi V, 2010 The effect of model order selection in group PICA. Hum. Brain Mapp 31. doi: 10.1002/hbm.20929, NA-NA.

Allen EA, Damaraju E, Plis SM, Erhardt EB, Eichele T, Calhoun VD, 2014 Tracking whole-brain connectivity dynamics in the resting state. Cereb. Cortex 24, 663–676. doi: 10.1093/cercor/bhs352. [PubMed: 23146964]

Autio JA, Glasser MF, Ose T, Donahue CJ, Bastiani M, Ohno M, Kawabata Y, Urushibata Y, Murata K, Nishigori K, Yamaguchi M, Hori Y, Yoshida A, Go Y, Coalson TS, Jbabdi S, Sotiropoulos SN, Kennedy H, Smith S, Essen DCV, Hayashi T, 2020 Towards HCP-Style macaque connectomes: 24-Channel 3T multi-array coil, MRI sequences and preprocessing. Neuroimage 215, 116800. doi: 10.1016/j.neuroimage.2020.116800. [PubMed: 32276072]

Avants BB, Tustison NJ, Stauffer M, Song G, Wu B, Gee JC, 2014 The Insight ToolKit image registration framework. Front. Neuroinform 8 (44). doi: 10.3389/fninf.2014.00044.

Avants BB, Tustison NJ, Wu J, Cook PA, Gee JC, 2011 An Open Source Multivariate Framework for n-Tissue Segmentation with Evaluation on Public Data. Neuroinformatcs 9, 381–400. doi: 10.1007/s12021-011-9109-y. [PubMed: 21373993]

Bai F, Xie C, Watson DR, Shi Y, Yuan Y, Wang Y, Yue C, Teng Y, Wu D, Zhang Z, 2011 Aberrant hippocampal subregion networks associated with the classifications of aMCI subjects: a longitudinal resting-state study. Plos One 6, e29288. doi: 10.1371/journal.pone.0029288. [PubMed: 22216234]

Bakker R, Wachtler T, Diesmann M, 2012 CoCoMac 2.0 and the future of tract-tracing databases. Front. Neuroinform 6 (30). doi: 10.3389/fninf.2012.00030.

Beckmann CF, DeLuca M, Devlin JT, Smith SM, 2005 Investigations into resting-state connectivity using independent component analysis. Philos. Trans. R. Soc. B Biol. Sci 360, 1001–1013. doi: 10.1098/rstb.2005.1634.

Biswal B, Yetkin FZ, Haughton VM, Hyde JS, 1995 Functional connectivity in the motor cortex of resting human brain using echo-planar MRI. Magn. Reson. Med 34, 537–541. doi: 10.1002/mrm.1910340409. [PubMed: 8524021]

Biswal BB, Mennes M, Zuo X-N, Gohel S, Kelly C, Smith SM, Beckmann CF, Adelstein JS, Buckner RL, Colcombe S, Doganowski A-M, Ernst M, Fair D, Hampson M, Hoptman MJ, Hyde JS, Kiviniemi VJ, Köttér R, Li S-J, Lin C-P, Lowe MJ, Mackay C, Madden DJ, Madsen KH, Margulies DS, Mayberg HS, McMahon K, Monk CS, Mostofsky SH, Nagel BJ, Pekar JJ, Peltier SJ, Petersen SE, Riedl V, Rombouts SARB, Rypma B, Schlaggar BL, Schmidt S, Seidler RD, Siegel GJ, Sorg C, Teng G-J, Veijola J, Villringer A, Walter M, Wang L, Weng X-C, Whitfield-Gabrieli S, Williamson P, Windischberger C, Zang Y-F, Zhang H-Y, Castellanos FX, Milham MP, 2010 Toward discovery science of human brain function. Proc. Natl. Acad. Sci. USA 107, 4734–4739. doi: 10.1073/pnas.0911855107. [PubMed: 20176931]

Calhoun VD, Adali T, Pearlson GD, Pekar JJ, 2001 A method for making group inferences from functional MRI data using independent component analysis. Hum. Brain Mapp 14, 140–151. doi: 10.1002/hbm.1048. [PubMed: 11559959]

Cushnie AK, El-Nahal HG, Bohlen MO, May PJ, Basso MA, Grimaldi P, Wang MZ, Velasco E.M.F., de, Sommer MA, Heilbronner SR, 2020 Using rAAV2-retro in rhesus macaques: Promise and caveats for circuit manipulation. J. Neurosci. Methods 345, 108859. doi: 10.1016/j.jneumeth.2020.108859. [PubMed: 32668316]

Cole DM, Smith SM, Beckmann CF, 2010 Advances and pitfalls in the analysis and interpretation of resting-state fMRI data. Front. Syst. Neurosci 4 (8). doi: 10.3389/fnsys.2010.00008.

Consortium, T., Prim DE, PRIME-D GCW, Milham M, Petkov CI, Margulies DS, Schroeder CE, Basso MA, Belin P, Fair DA, Fox A, Kastner S, Mars RB, Messinger A, Poirier C, Vanduffel W,
Essen DCV, Alvand A, Becker Y, Hamed SB, Benn A, Bodin C, Boretius S, Cagna B, Coulon O, El-Gohary SH, Evrard H, Forkel SJ, Friedrich P, Froudist-Walsh S, Garza-Villarreal EA, Gao Y, Gozzi A, Grigis A, Hartig R, Hayashi T, Heuer K, Howells H, Ardesch Jarraya B, Jarrett W, Jedema HP, Kagan I, Kelly C, Kennedy H, Klink PC, Kwok SC, Leech R, Liu X, Madan C, Madushanka W, Majka P, Mallon A-M, Marche K, Meguerditchian A, Menon RS, Merchant H, Mitchell A, Nenning K-H, Nikolaidis A, Ortiz-Rios M, Pagani M, Pareek V, Prescott M, Procyk E, Rajimehr R, Rautu I-S, Raz A, Roe AW, Rossi-Pool R, Roumazeilles L, Sakai T, Sallet J, Garcia-Saldivar P, Sato C, Sawiak S, Schiffer M, Schwiedrzik CM, Seidlitz J, Sein J, Shen Z, Shmuel A, Silva AC, Simone L, Sirmipilatze N, Sliwa J, Smallwood J, Tasserie J, Schotten M.T., de, Toro R, Trapeau R, Ungerleider L, Vezoli J, Wang Z, Wells S, Williams B, Xu T, Xu AG, Yacoub E, Zhan M, Ai L, Amiez C, Balezeau F, Baxter MG, Blezer ELA, Brochier T, Chen A, Croxson PL, Damatac CG, Dehaene S, Everling S, Fleischer J, Freiwald W, Guedj C, Hadj-Bouziane F, Harel N, Hiba B, Jung B, Koo B, Laland KN, Leopold DA, Lindenfors P, Meunier M, Mok K, Morrison JH, Nacef J, Nagy J, Pink M, Reader SM, Roelfsema PR, Rushworth MFS, Russ BE, Schmid MC, Sullivan EL, Thiele A, Tudorov OS, Todorov BS, Wilson CRE, Ye FQ, Zarco W, Zhou Y. 2020 Accelerating the evolution of nonhuman primate neuroimaging. Neuron 105, 600–603. doi: 10.1016/j.neuron.2019.12.023. [PubMed: 32078795]

Cox RW, 1996 AFNI: Software for analysis and visualization of functional magnetic resonance neuroimages. Comput. Biomed. Res 29, 162–173. doi: 10.1006/cbmr.1996.0014. [PubMed: 8812068]

Essen DCV, Maunsell JHR, 1983 Hierarchical organization and functional streams in the visual cortex. Trends Neurosci. 6, 370–375. doi: 10.1016/0166-2236(83)90167-4.

Essen DCV, Ugurbil K, Auerbach E, Barch D, Behrens TEJ, Bucholz R, Chang A, Chen L, Corbetta M, Curtiss SW, Penna SD, Feinberg D, Glasser MF, Harel N, Heath AC, Larson-Prior L, Marcus D, McAllarareas G, Moeller S, Oostenveld R, Petersen SE, Prior F, Schlaggar BL, Smith SM, Snyder AZ, Xu J, Yacoub E, Consortium, W.-M.H., 2012 The Human Connectome Project: A data acquisition perspective. Neuroimage 62, 2222–2231. doi: 10.1016/j.neuroimage.2012.02.018. [PubMed: 22366334]

Farber NE, Harkin CP, Niedfeldt J, Hudetz AG, Kampine JP, Schmeling WT, 1997 Region-specific and agent-specific dilation of intracerebral microvessels by volatile anesthetics in rat brain slices. Anesthesiology 87, 1191–1198. doi: 10.1097/00000542-199711000-00024. [PubMed: 9366472]

Felleman DJ, Essen DCV, 1991 Distributed hierarchical processing in the primate cerebral cortex. Cereb. Cortex 1, 1–47. doi: 10.1093/cercor/1.1.1. [PubMed: 1822724]

Floris DL, Lai M-C, Nath T, Millham MP, Martino AD, 2018 Network-specific sex differentiation of intrinsic brain function in males with autism. Mol. Autism 9 (17). doi: 10.1186/s13229-018-0192-x.

Fox MD, Raichle ME, 2007 Spontaneous fluctuations in brain activity observed with functional magnetic resonance imaging. Nat. Rev. Neurosci. 8, 700–711. doi: 10.1038/nrn2201. [PubMed: 17704812]

Galvan A, Raper J, Hu X, Paré J, Bonaventura J, Richie CT, Michaelides M, Mueller SAL, Roseboom PH, Oler JA, Kalin NH, Hall RA, Smith Y. 2019 Ultrastructural localization of DREADDs in monkeys. Eur. J. Neurosci. 50, 2801–2813. doi: 10.1111/ejn.14429. [PubMed: 31063250]

Galvan A, Stauffer WR, Acker L, El-Shamayleh Y, Inoue K-I, Ohayon S, Schmid MC, 2017 Nonhuman primate optogenetics: recent advances and future directions. J. Neurosci. Official J. Soc. Neurosci. 37, 10894–10903. doi: 10.1523/jneurosci.1839-17.2017.

Gilbert KM, Gati JS, Barker K, Everling S, Menon RS, 2016 Optimized parallel transmit and receive radiofrequency coil for ultrahigh-field MRI of monkeys. NeuroImage 125, 153–161. doi: 10.1016/j.neuroimage.2015.10.049. [PubMed: 26497267]

Glasser MF, Smith SM, Marcus DS, Andersson JLR, Auerbach EJ, Behrens TEJ, Coalson TS, Harms MP, Jenkinson M, Moeller S, Robinson EC, Sotiropoulos SN, Xu J, Yacoub E, Ugurbil K, Essen DCV, 2016 The Human Connectome Project’s neuroimaging approach. Nat. Neurosci 19, 1175–1187. doi: 10.1038/nn.4361. [PubMed: 27571196]

Goulas A, Stiers P, Hutchison RM, Everling S, Petrides M, Margulies DS, 2017 Intrinsic functional architecture of the macaque dorsal and ventral lateral frontal cortex. J. Neurophysiol 117, 1084–1099. doi: 10.1152/jn.00486.2016. [PubMed: 28003408]

Neuroimage. Author manuscript; available in PMC 2020 December 17.
Hallquist MN, Hwang K, Luna B, 2013 The nuisance of nuisance regression: spectral misspecification in a common approach to resting-state fMRI preprocessing reintroduces noise and obscures functional connectivity. Neuroimage 82, 208–225. doi: 10.1016/j.neuroimage.2013.05.116. [PubMed: 23747457]

Harms MP, Somerville LH, Ances BM, Andersson J, Barch DM, Bastiani M, Bookheimer SY, Brown TB, Buckner RL, Burgess GC, Coalson TS, Chappell MA, Dapretto M, Douaud G, Fischl B, Glasser MF, Greve DN, Hodge C, Jamison KW, Jbabdi S, Kandala S, Li X, Mair RW, Mangia S, Marcus D, Maselli D, Moeller S, Nichols TE, Robinson EC, Salat DH, Smith SM, Sotiroopoulos SN, Terpstra M, Thomas KM, Tisdall MD, Ugurbil K, van der Kouwe A, Woods RP, Zöllei L, Essen DCV, Yacoub E, 2018 Extending the human connectome project across ages: imaging protocols for the lifespan development and aging projects. Neuroimage 183, 972–984. doi: 10.1016/j.neuroimage.2018.09.060. [PubMed: 30261308]

Heilbronner SR, Chafee MV, 2019 Learning how neurons fail inside of networks: nonhuman primates provide critical data for psychiatry. Neuron 102, 21–26. doi: 10.1016/j.neuron.2019.02.030. [PubMed: 30946820]

Heilbronner SR, Rodriguez-Romagueras J, Quirk GJ, Groenewegen HJ, Haber SN, 2016 Circuit-Based Corticostriatal Homologies Between Rat and Primate. Biol. Psychiat 80, 509–521. doi: 10.1016/j.biopsych.2016.05.012. [PubMed: 27450032]

van den Heuvel MP, Pol HEH, 2010 Exploring the brain network: A review on resting-state fMRI functional connectivity. Eur. Neuropsychopharm 20, 519–534. doi: 10.1016/j.euroneuro.2010.03.008.

Hindriks R, Adhikari MH, Murayama Y, Ganzetti M, Mantini D, Logothetis NK, Deco G, 2016 Can sliding-window correlations reveal dynamic functional connectivity in resting-state fMRI? Neuroimage 127, 242–256. doi: 10.1016/j.neuroimage.2015.11.055. [PubMed: 26631813]

Hulvershorn LA, Mennes M, Castellanos FX, Martino AD, Milham MP, Hummer TA, Roy AK, 2013 Abnormal amygdala functional connectivity associated with emotional lability in children with attention-deficit/hyperactivity disorder. J. Am. Acad. Child Psy 53. doi: 10.1016/j.jaac.2013.11.012, 351–61.e1.

Hutchison RM, Everling S, 2012 Monkey in the middle: why non-human primates are needed to bridge the gap in resting-state investigations. Front. Neuroanat 6 (29). doi: 10.3389/ fnana.2012.00029.

Hutchison RM, Leung LS, Mirsattari SM, Gati JS, Menon RS, Everling S, 2011 Resting-state networks in the macaque at 7T. Neuroimage 56, 1546–1555. doi: 10.1016/j.neuroimage.2011.02.063. [PubMed: 21356313]

Hutchison RM, Womelsdorf T, Allen EA, Bandettini PA, Calhoun VD, Corbetta M, Penna SD, Duyn JH, Glover GH, Gonzalez-Castillo J, Handwerker DA, Keilholz S, Kiviniemi V, Leopold DA, Pasquale de, F, Sporns O, Walter M, Chang C, 2013 Dynamic functional connectivity: promise, issues, and interpretations. Neuroimage 80, 360–378. doi: 10.1016/j.neuroimage.2013.05.079. [PubMed: 23707587]

Jafri MJ, Pearlson GD, Stevens M, Calhoun VD, 2008 A method for functional network connectivity among spatially independent resting-state components in schizophrenia. Neuroimage 39, 1666–1681. doi: 10.1016/j.neuroimage.2007.11.001. [PubMed: 18082428]

Jenkinson M, Beckmann CF, Behrens TEJ, Woolrich MW, Smith SM, 2012 FSL. Neuroimage 62, 782–790. doi: 10.1016/j.neuroimage.2011.09.015. [PubMed: 21979382]

Jonckers E, Audekerke JV, Visscher GD, de Linden AV, Verhoeve M, 2011 Functional connectivity fMRI of the rodent brain: comparison of functional connectivity networks in rat and mouse. Plos One 6, e18876. doi: 10.1371/journal.pone.0018876. [PubMed: 21533116]

Khateeb K, Griggs DJ, Sabes PN, Yazdan-Shahmorad A, 2019 Convection enhanced delivery of optogenetic adeno-associated viral vector to the cortex of rhesus macaque under guidance of online MRI images. J. Vis. Exp doi: 10.3791/59232.
Kiviniemi V, Starck T, Remes J, Long X, Nikkinen M, Haapea J, Veijola J, Moilanen I, Isohanni M, Zang Y-F, Tervonen O, 2009 Functional segmentation of the brain cortex using high model order group PICA. Hum. Brain Mapp 30, 3865–3886. doi: 10.1002/hbm.20813. [PubMed: 19507160]

Konova AB, Moeller SJ, Tomasi D, Goldstein RZ, 2015 Effects of chronic and acute stimulants on brain functional connectivity hubs. Brain Res. 1628, 147–156. doi: 10.1016/j.brainres.2015.02.002. [PubMed: 25721787]

Krubitzer L, Huffman KJ, 2000 Arealization of the neocortex in mammals: genetic and epigenetic contributions to the phenotype. Brain Behav. Evol 55, 322–335. doi: 10.1159/000006667.

Lagore R, Moeller S, Delabarre L, Grant A, Radder J, Zimmermann J, Ugurbil K, Yacoub E, Harel N, Adriany G, 2020 An 8 channel dipole transceive and 24 loop receive array for non-human primate head imaging at 10.5T. In: Proceedings of the 28th International Social for Magnetic Resonance in Medicine Scientific Meeting ISMRM.

Laird AR, Fox PM, Eickhoff SB, Turner JA, Ray KL, McKay DR, Glahn DC, Beckmann CF, Smith SM, Fox PT, 2011 Behavioral interpretations of intrinsic connectivity networks. J. Cognit. Neurosci 23, 4022–4037. doi: 10.1162/jocn_a_00077. [PubMed: 21671731]

Lawrie SM, Buechel C, Whalley HC, Frith CD, Friston KJ, Johnstone EC, 2002 Reduced frontotemporal functional connectivity in schizophrenia associated with auditory hallucinations. Biol. Psychiat 51, 1008–1011. doi: 10.1016/s0006-3223(02)01316-1. [PubMed: 12062886]

Leech R, Braga R, Sharp DJ, 2012 Echoes of the brain within the posterior cingulate cortex. J. Neurosci 32, 215–222. doi: 10.1523/jneurosci.3689-11.2012. [PubMed: 22219283]

Leech R, Kamourieh S, Beckmann CF, Sharp DJ, 2011 Fractionating the default mode network: distinct contributions of the ventral and dorsal posterior cingulate cortex to cognitive control. J. Neurosci 31, 3217–3224. doi: 10.1523/jneurosci.5626-10.2011. [PubMed: 21368033]

Leite FP, Tsao D, Vanduffel W, Fize D, Sasaki Y, Wald LL, Dale AM, Kwong KK, Orban GA, Rosen BR, Tootell RB, Mandeville JB, 2002 Repeated fMRI using iron oxide contrast agent in awake, behaving macaques at 3 tesla. NeuroImage 16, 283–294. doi: 10.1006/nimg.2002.1110. [PubMed: 12030817]

Li L, Hu X, Preuss TM, Glasser MF, Damen FW, Qiu Y, Rilling J, 2013 Mapping putative hubs in human, chimpanzee and rhesus macaque connectomes via diffusion tractography. Neuroimage 80, 462–474. doi: 10.1016/j.neuroimage.2013.04.024. [PubMed: 23603286]

Li M, Kim HS, Tian L, Yu MK, Jon S, Moon WK, 2012 Comparison of two ultrasmall superparamagnetic iron oxides on cytotoxicity and mr imaging of tumors. Theranostics 2, 76–85. doi: 10.7150/thno.3462. [PubMed: 22272221]

Li Y-O, Adali T, Calhoun VD, 2007 Estimating the number of independent components for functional magnetic resonance imaging data. Hum. Brain Mapp 28, 1251–1266. doi: 10.1002/hbm.20359. [PubMed: 17274023]

Logothetis NK, 2003 The underpinnings of the BOLD functional magnetic resonance imaging signal. J. Neurosci 23, 3963–3971. doi: 10.1523/jneurosci.23-10-03963.2003. [PubMed: 12764080]

Logothetis NK, Pauls J, Augath M, Trinath T, Oeltermann A, 2001 Neuro-physiological investigation of the basis of the fMRI signal. Nature 412 (150). doi: 10.1038/35084005.

Lustig C, Snyder AZ, Bhakta M, O'Brien KC, McAvoy M, Raichle ME, Morris JC, Buckner RL, 2003 Functional deactivations: change with age and dementia of the Alzheimer type. Proc. Natl. Acad. Sci 100, 14504–14509. doi: 10.1073/pnas.2259252100. [PubMed: 14608034]

Markov NT, Ercsey-Ravasz MM, Gomes ARR, Lamy C, Magrou L, Vezoli J, Misery P, Falchier A, Quilodran R, Gariel MA, Sallet J, Gamanut R, Huissoud C, Clavagnier S, Giroud P, Sappey-Mariner D, Barone P, Duhay C, Toroczkai Z, Knoblauch K, Essen DCV, Kennedy H, 2012 A weighted and directed interareal connectivity matrix for macaque cerebral cortex. Cereb. Cortex New York N Y 1991 24, 17–36. doi: 10.1093/cercor/bhs270. [PubMed: 23010748]

Mars RB, Verhagen L, Gladwin TE, Neubert F-X, Sallet J, Rushworth MFS, 2016 Comparing brains by matching connectivity profiles. Neurosci. Biobehav. Rev 60, 90–97. doi: 10.1016/j.neubiorev.2015.10.008. [PubMed: 26627865]

Martino FD, Esposito F, van de Moortele P-F, Harel N, Formisano E, Goebel R, Ugurbil K, Yacoub E, 2011 Whole brain high-resolution functional imaging at ultra high magnetic fields: An application
to the analysis of resting state networks. Neuroimage 57, 1031–1044. doi: 10.1016/j.neuroimage.2011.05.008. [PubMed: 21600293]

Martino FD, Yacoub E, Kemper V, Moerel M, Uludağ K, Weerd PD, Ugurbil K, Goebel R, Formisano E, 2018 The impact of ultra-high field MRI on cognitive and computational neuroimaging. Neuroimage 168, 366–382. doi: 10.1016/j.neuroimage.2017.03.060. [PubMed: 28396293]

Milhan MP, Ai L, Koo B, Xu T, Amiez C, Balezeau F, Baxter MG, Blezer ELA, Brochier T, Chen A, Croxson PL, Damatac CG, Dehaene S, Everling S, Fair DA, Fleysher L, Freiwald W, Froudist-Walsh S, Griffiths TD, Guedj C, Hadj-Bouziane F, Hamed SB, Harel N, Hiba B, Jarraya B, Jung B, Kastner S, Klink PC, Kwok SC, Laland KN, Leopold DA, Lindenfors P, Mars RB, Menon RS, Messinger A, Mennier M, Mok K, Morrison JH, Nacef J, Nagy J, Rios MO, Petkou CI, Pinski M, Poirier C, Procyk E, Rajimehr R, Reader SM, Roelfsema PR, Rudko DA, Rushworth MFS, Russ BE, Sallet J, Schmid MC, Schwiedrzik CM, Seidtitz J, Sein J, Shmuel A, Sullivan EL, Ungereider L, Thiele A, Todorov OS, Tsao D, Wang Z, Wilson CRE, Yacoub E, Ye FQ, Zarco W, Zhou Y, Margoulies DS, Schroeder CE, 2018 An open resource for non-human primate imaging. Neuron 100, 61–74. doi: 10.1016/j.neuron.2018.08.039, e2. [PubMed: 30269990]

Moeller S, Nallasamy N, Tsao DY, Freiwald WA, 2009 Functional connectivity of the macaque brain across stimulus and arousal states. J. Neurosci. Official J. Soc. Neurosci 29, 5897–5909. doi: 10.1523/jneurosci.0220-09.2009.

Moeller S, Yacoub E, Olman CA, Auerbach E, Strupp J, Harel N, Uğurbil K, 2010 Multiband multislice GE-EPI at 7 tesla, with 16-fold acceleration using partial parallel imaging with application to high spatial and temporal whole-brain fMRI. Magn. Reson. Med 63, 1144–1153. doi: 10.1002/mrm.22361. [PubMed: 20432285]

Mustafa TA, Mohammed-Rasheed MA, 2019 Accumulation and cytotoxicity assessment of TAT-IONPs on cancerous mammalian cells. Anim. Biotechnol 1–6. doi: 10.1080/10495398.2019.1658595.

Nagai Y, Kikuchi E, Lerchner W, Inoue K, Ji B, Eldridge MAG, Kaneko H, Kimura Y, Oh-Nishi A, Hori Y, Kato Y, Hirabayashi T, Fujimoto A, Kumata K, Zhang M-R, Aoki I, Suhara T, Higuchi M, Takada M, Richmond BJ, Minamimoto T, 2016 PET imaging-guided chemogenetic silencing reveals a critical role of primate rostromedial caudate in reward evaluation. Nat. Commun 7, 13605. doi: 10.1038/ncomms13605. [PubMed: 27922009]

Ohliger MA, Grant AK, Sodickson DK, 2003 Ultimate intrinsic signal-to-noise ratio for parallel MRI: Electromagnetic field considerations. Magn. Reson. Med 50, 1018–1030. doi: 10.1002/mrm.10597. [PubMed: 14587013]

Okada S, Bartelle BB, Li N, Breton-Provencher V, Lee JJ, Rodriguez E, Melican J, Sur M, Jasanoﬀ A, 2018 Calcium-dependent molecular fMRI using a magnetic nanosensor. Nat. Nanotechnol 13, 473–477. doi: 10.1038/s41565-018-0092-4. [PubMed: 29713073]

Pelekano V, Mok RM, Joly O, Ainsworth M, Kyriazis D, Kelly MG, Bell AH, Kriegeskorte N, 2020 Rapid event-related, BOLD fMRI, non-human primates (NHP): choose two out of three. Sci. Rep. 10, 7485. doi: 10.1038/s41598-020-64376-8.

Petrides M, Pandya DN, 2002 Comparative cytoarchitectonic analysis of the human and the macaque ventrolateral prefrontal cortex and corticocortical connection patterns in the monkey: ventrolateral prefrontal cortex in human and monkey. Eur. J. Neurosci 16, 291–310. doi: 10.1046/j.1460-9586.2001.02090.x. [PubMed: 12169111]

Pruessmann KP, Weiger M, Scheidegger MB, Boesiger P, 1999 SENSE: sensitivity encoding for fast MRI. Magn. Reson. Med 42, 952–962. doi: 10.1002/sici.1522-2594(199911)42:5<952::aid-mrm16>3.0.co;2-s. [PubMed: 10542355]

Quigley M, Cordes D, Wendt G, Turski P, Moritz C, Haughton V, Meyerand ME, 2001 Effect of focal and nonfocal cerebral lesions on functional connectivity studied with MR imaging. Ajnr Am. J. Neuroradiol 22, 294–300. [PubMed: 11156772]

Raper J, Murphy L, Richardson R, Romm Z, Kovacs-Balint Z, Payne C, Galvan A, 2019 Chemogenetic inhibition of the amygdala modulates emotional behavior expression in infant rhesus monkeys. Eneuro 6. doi: 10.1523/eneuro.0360-19.2019, ENEURO.0360-19.2019.

Saad ZS, Reynolds RC, 2012 SUMA. Neuroimage 62, 768–773. doi: 10.1016/j.neuroimage.2011.09.016. [PubMed: 21945692]
Saad ZS, Reynolds RC, Argall B, Japee S, Cox RW, 2004 SUMA: An interface for surface-based intra- and inter-subject analysis with AFNI. In: Proceedings of the 2nd IEEE International Symposium Biomed Imaging Nano Macro Ieee Cat 04ex821, pp. 1510–1513. doi: 10.1109/isbi.2004.1398837.

Saleem KS, Logothetis NK, 2012 A Combined MRI and Histology Atlas of the Rhesus Monkey Brain in Stereotaxic Coordinates. Academic Press.

Schaeffer DJ, Johnston KD, Gilbert KM, Gati JS, Menon RS, Everling S, 2018 In vivo manganese tract tracing of frontal eye fields in rhesus macaques with ultra-high field MRI: Comparison with DWI tractography. Neuroimage 181, 211–218. doi: 10.1016/j.neuroimage.2018.06.072. [PubMed: 29964189]

Seidlitz J, Sponheim C, Glen D, Ye FQ, Saleem KS, Leopold DA, Ungerleider L, Messinger A, 2018 A population MRI brain template and analysis tools for the macaque. NeuroImage 170. doi: 10.1016/j.neuroimage.2017.04.063.

Setsompop K, Gagoski BA, Polimeni JR, Witzel T, Wedeen VJ, Wald LL, 2012 Blipped-controlled aliasing in parallel imaging for simultaneous multislice echo planar imaging with reduced g-factor penalty. Magn. Reson. Med 67. doi: 10.1002/mrm.23097, 1210–24. [PubMed: 21858868]

Smith SM, Fox PT, Miller KL, Glahn DC, Fox PM, Mackay CE, Filippini N, Watkins KE, Toro R, Laird AR, Beckmann CF, 2009 Correspondence of the brain’s functional architecture during activation and rest. Proc. Natl. Acad. Sci. USA 106. doi: 10.1073/pnas.0905267106, 13040–5. [PubMed: 19620724]

Smith SM, Miller KL, Moeller S, Xu J, Auerbach EJ, Woolrich MW, Beckmann CF, Jenkinson M, Andersson J, Glasser MF, Essen DVC, Feinberg DA, Yacoub ES, Uğurbil K, 2012 Temporally-independent functional modes of spontaneous brain activity. Proc. Natl. Acad. Sci. USA 109. doi: 10.1073/pnas.1121329109, 3131–6. [PubMed: 22323591]

Smith SM, Vidaurre D, Beckmann CF, Glasser MF, Jenkinson M, Miller KL, Nichols TE, Robinson EC, Salimi-Khorshidi G, Woolrich MW, Barch DM, Uğurbil K, Essen DVC, 2013 Functional connectomics from resting-state fMRI. Trends Cogn. Sci. 17. doi: 10.1016/j.tics.2013.09.016, 666–82. [PubMed: 24238796]

Tsaö DY, Freiwald WA, Knutsen TA, Mandeville JB, Tootell RBH, 2003 Faces and objects in macaque cerebral cortex. Nat. Neurosci 6, 989–995. doi: 10.1038/nn1111. [PubMed: 12925854]

Uğurbil K, 2018 Imaging at ultrahigh magnetic fields: History, challenges, and solutions. Neuroimage 168, 7–32. doi: 10.1016/j.neuroimage.2017.07.007. [PubMed: 28698108]

Uğurbil K, 2016 What is feasible with imaging human brain function and connectivity using functional magnetic resonance imaging. Philos. Trans. R. Soc. Lond. Ser. B Biol. Sci 371, 20150361. doi: 10.1098/rstb.2015.0361. [PubMed: 27574313]

Uğurbil K, 2014 Magnetic resonance imaging at ultrahigh fields. IEEE Trans. Bio-med. Eng 61, 1364–1379. doi: 10.1109/tbme.2014.2313619.

Uğurbil K, Adriany G, Andersen P, Chen W, Garwood M, Gruetter R, Henry P-G, Kim S-G, Lieu H, Tkac I, Vaughan T, Moortele P-FVD, Yacoub E, Zhu X-H, 2003 Ultrahigh field magnetic resonance imaging and spectroscopy. Magn. Reson. Imaging 21, 1263–1281. doi: 10.1016/j.mri.2003.08.027. [PubMed: 14725934]

Uğurbil K, Auerbach E, Moeller S, Grant A, Wu X, Moortele P-FVD, Olman C, Delabarre L, Schillak S, Radder J, Lagore R, Adriany G, 2019 Brain imaging with improved acceleration and SNR at 7 Tesla obtained with 64-channel receive array. Magn. Reson. Med 82, 495–509. doi: 10.1002/mrm.27695. [PubMed: 30803023]

Uğurbil K, Xu J, Auerbach EJ, Moeller S, Vu AT, Duarte-Carvajalino JM, Lenglet C, Wu X, Schmitter S, Moortele P-FVD, Strupp J, Sapiro G, Martino FD, Wang D, Harel N, Garwood M, Chen L, Feinberg DA, Smith SM, Miller KL, Sotiropoulos SN, Jbabdi S, Andersson JLR, Behrens TEJ, Glasser MF, Essen DVC, Yacoub E, Consortium, W.-M.H., 2013 Pushing spatial and temporal resolution for functional and diffusion MRI in the Human Connectome Project. Neuroimage 80, 80–104. doi: 10.1016/j.neuroimage.2013.05.012. [PubMed: 23702417]

Upright NA, Brookshire SW, Schnebelen W, Damatac CG, Hof PR, Browning PFG, Croxson PL, Rudebeck PH, Baxter MG, 2018 Behavioral effect of chemogenetic inhibition is directly related to receptor transduction levels in rhesus monkeys. J. Neurosci. Official J. Soc. Neurosci 38, 7969–7975. doi: 10.1523/jneurosci.1422-18.2018.
Vanduffel W, Fize D, Mandeville JB, Nelissen K, Hecke PV, Rosen BR, Tootell RBH, Orban GA, 2001 Visual motion processing investigated using contrast agent-enhanced fMRI in awake behaving monkeys. Neuron 32, 565–577. doi: 10.1016/s0896-6273(01)00502-5. [PubMed: 11719199]

Vincent JL, Kahn I, Esson DCV, Buckner RL, 2009 Functional connectivity of the macaque posterior parahippocampal cortex. J. Neurophysiol 103, 793–800. doi: 10.1152/jn.00546.2009. [PubMed: 19955295]

Vincent JL, Patel GH, Fox MD, Snyder AZ, Baker JT, Esson DCV, Zempel JM, Snyder LH, Corbetta M, Raichle ME, 2007 Intrinsic functional architecture in the anaesthetized monkey brain. Nature 447, 83–86. doi: 10.1038/nature05758. [PubMed: 17476267]

Vu AT, Jamison K, Glasser MF, Smith SM, Coalson T, Moeller S, Auerbach EJ, Ugurbil K, Yacoub E, 2016 Tradeoffs in pushing the spatial resolution of fMRI for the 7T human connectome project. Neuroimage 154, 23–32. doi: 10.1016/j.neuroimage.2016.11.049. [PubMed: 27894889]

Wang J, Wang L, Zang Y, Yang H, Tang H, Gong Q, Chen Z, Zhu C, He Y, 2009 Parcellation-dependent small-world brain functional networks: a resting-state fMRI study. Hum. Brain Mapp 30. doi: 10.1002/hbm.20623, 1511–23. [PubMed: 18649353]

Watson KK, Platt ML., 2012 Of mice and monkeys: using non-human primate models to bridge mouse- and human-based investigations of autism spectrum disorders. J. Neurodev. Disord 4 (21). doi: 10.1186/1866-1955-4-21.

Whitfield-Gabrieli S, Nieto-Castanon A, 2012 Conn : a functional connectivity toolbox for correlated and anticorrelated brain networks. Brain Connect. 2, 125–141. doi: 10.1089/brain.2012.0073. [PubMed: 22642651]

Wiesinger F, Moortele PV, de Adriany G, Zanche ND, Ugurbil K, Pruessmann KP, 2006 Potential and feasibility of parallel MRI at high field. NMR Biomed. 19, 368–378. doi: 10.1002/nbm.1050. [PubMed: 16705638]

Wiesinger F, Moortele PV, de Adriany G, Zanche ND, Ugurbil K, Pruessmann KP, 2004 Parallel imaging performance as a function of field strength—An experimental investigation using electrodynamic scaling. Magn. Reson. Med 52, 953–964. doi: 10.1002/mrm.20281. [PubMed: 15508167]

Xie C, Bai F, Yu H, Shi Y, Yuan Y, Chen Gang, Li W, Chen Guangyu, Zhang Z, Li S-J, 2012 Abnormal insula functional network is associated with episodic memory decline in amnestic mild cognitive impairment. Neuroimage 63. doi: 10.1016/j.neuroimage.2012.06.062, 320–7. [PubMed: 22776459]

Xu T, Falchier A, Sullivan EL, Linn G, Ramirez JSB, Ross D, Feczko E, Opitz A, Bagley J, Sturgeon D, Earl E, Miranda-Dominguez O, Perrone A, Craddock RC, Schroeder CE, Colcombe S, Fair DA, Milham MP, 2018 Delineating the macroscale areal organization of the macaque cortex in vivo. Cell Rep. 23, 429–441. doi: 10.1016/j.celrep.2018.03.049. [PubMed: 29642002]

Xu T, Sturgeon D, Ramirez JSB, Froudast-Walsh S, Margulies DS, Schroeder CE, Fair DA, Milham MP, 2019 Inter-individual variability of functional connectivity in awake and anesthetized rhesus monkeys. Biol. Psychiatry Cognit. Neurosci. Neuroimaging doi: 10.1016/j.bpsc.2019.02.005.

Yacoub E, Harel N, Ugurbil K, 2008 High-field fMRI unveils orientation columns in humans. Proc. Natl. Acad. Sci 105, 10607–10612. doi: 10.1073/pnas.0804110105. [PubMed: 18641121]

Yacoub E, Shmuel A, Pfeuffer J, Moortele PVD, Adriany G, Andersen P, Vaughan JT, Merkle H, Ugurbil K, Hu X, 2001 Imaging brain function in humans at 7 Tesla. Magn. Reson. Med 45, 588–594. doi: 10.1002/mrm.1080. [PubMed: 11283986]

Zilverstand A, Huang AS, Alia-Klein N, Goldstein RZ, 2018 Neuroimaging impaired response inhibition and salience attribution in human drug addiction: a systematic review. Neuron 98, 886–903. doi: 10.1016/j.neuron.2018.03.048. [PubMed: 29879391]

Zimmermann J, Goebel R, Martino FD, van de Moortele P-F, Feinberg D, Adriany G, Chaimow D, Shmuel A, Ugurbil K, Yacoub E, 2011 Mapping the organization of axis of motion selective features in human area MT using high-field fMRI. Plos One 6, e28716. doi: 10.1371/ journal.pone.0028716. [PubMed: 22163328]

Zuo X-N, Kelly C, Adelstein JS, Klein DF, Castellanos FX, Milham MP, 2010 Reliable intrinsic connectivity networks: Test–retest evaluation using ICA and dual regression approach. Neuroimage 49, 2163–2177. doi: 10.1016/j.neuroimage.2009.10.080. [PubMed: 19896537]
Fig. 1.
Independent components data in individual subject space. Displayed are 6 RSNs in one unsmoothed macaque dataset processed without spatial smoothing. For this visualization thresholds were set to 4,−4. The results show robust results in the unsmoothed volumes at native resolution demonstrating our high SNR regime. Networks adhere to the cortical ribbon topography without any registration performed.
Fig. 2.
Cortical surface representation of the first 15 resting state networks (RSNs) identified using group independent component analysis (GICA) in 6 macaque monkeys. Overlaid color maps represent thresholded z-scores. Individual subjects were normalized to the NMT template. Each component shows the medial and lateral view of each hemisphere independently as well as the dorsal and ventral view of the hemispheres combined.
Fig. 3.
Cortical surface representation of the second 15 resting state networks (RSNs) identified using group independent component analysis (GICA) in 6 macaque monkeys. Overlayed color maps represent thresholded z-scores. Individual subjects were normalized to the NMT template. Each component shows the medial and lateral view of each hemisphere independently as well as the dorsal and ventral view of the hemispheres combined.
Fig. 4.
Consistency of single subject resting state networks. The figure demonstrates the same four resting state components in all six monkeys. Thresholds were kept consistent with group analysis results.
Fig. 5. Example resting state networks (RSNs) of one macaque following single-subject independent component analysis (ICA). Images are normalized to the NMT space and overlaid onto the NMT-template. Overlaid color maps represent thresholded z-scores. Arbitrary slices were selected to demonstrate the diversity of components from subcortical to cortical components as well as the detailed anatomical correspondence.
Fig. 6. Functional Network Connectivity of the 30 extracted Independent components. Lines portray significant connections between the networks and color indicates positive (red) and negative (blue) functional connectivity. Networks were clustered using hierarchical clustering. Results demonstrate strong positive within cluster network connections and primarily negative between cluster connections.
Fig. 7.
Qualitative comparison between resting state networks identified from the Human Connectome Project database (supplied by the CONN toolbox and Connectome Workbench) and our six subject macaque group resting state networks. Individual regions of interest were smoothed using a 10 mm spherical kernel interpolated in 3D. Gray shading represents a reconstruction of the estimated pial surface.