Systems/Circuits

Echoes from Intrinsic Connectivity Networks in the Subcortex

¹⁰Josephine M. Groot,^{1,2} Steven Miletic,² Scott J. S. Isherwood,² Desmond H. Y. Tse,³ Sarah Habli,⁴ ¹⁰Asta K. Håberg,^{5,6} Birte U. Forstmann,² ¹⁰Pierre-Louis Bazin,^{1,7*} and ¹⁰Matthias Mittner^{1*}

¹Department of Psychology, UiT–Arctic University of Norway, Tromsø, 9037, Norway, ²Integrative Model-based Cognitive Neuroscience research unit, University of Amsterdam, Amsterdam, 1001 NK, The Netherlands, ³Department of Neuropsychology and Psychopharmacology, Maastricht University, Maastricht, 6200 MD, The Netherlands, ⁴Department of Psychology, Norwegian University of Science and Technology, Trondheim, 8900, Norway, ⁵Department of Neuromedicine and Movement Science, Norwegian University of Science and Technology, Trondheim, 8900, Norway, ⁶Department of Radiology and Nuclear Medicine, St. Olavs Hospital, Trondheim, 7006, Norway, and ⁷Departments of Neurophysics and Neurology, Max Planck Institute for Human Cognitive and Brain Sciences, Leipzig, 04303, Germany

Decades of research have greatly improved our understanding of intrinsic human brain organization in terms of functional networks and the transmodal hubs within the cortex at which they converge. However, substrates of multinetwork integration in the human subcortex are relatively uncharted. Here, we leveraged recent advances in subcortical atlasing and ultra-high field (7 T) imaging optimized for the subcortex to investigate the functional architecture of 14 individual structures in healthy adult males and females with a fully data-driven approach. We revealed that spontaneous neural activity in subcortical regions can be decomposed into multiple independent subsignals that correlate with, or "echo," the activity in functional networks across the cortex. Distinct subregions of the thalamus, striatum, claustrum, and hippocampus showed a varied pattern of echoes from attention, control, visual, somatomotor, and default mode networks, demonstrating evidence for a heterogeneous organization supportive of functional integration. Multiple network activity furthermore converged within the globus pallidus externa, substantia nigra, and ventral tegmental area but was specific to one subregion, while the amygdala and pedunculopontine nucleus preferentially affiliated with a single network, showing a more homogeneous topography. Subregional connectivity of the globus pallidus interna, subthalamic nucleus, red nucleus, periaqueductal gray, and locus coeruleus did not resemble patterns of cortical network activity. Together, these finding describe potential mechanisms through which the subcortex participates in integrated and segregated information processing and shapes the spontaneous cognitive dynamics during rest.

Key words: 7 Tesla; dual regression; functional connectivity; network integration; resting-state

Significance Statement

Despite the impact of subcortical dysfunction on brain health and cognition, large-scale functional mapping of subcortical structures severely lags behind that of the cortex. Recent developments in subcortical atlasing and imaging at ultra-high field provide new avenues for studying the intricate functional architecture of the human subcortex. With a fully data-driven analysis, we reveal subregional connectivity profiles of a large set of noncortical structures, including those rarely studied in fMRI research. The results have implications for understanding how the functional organization of the subcortex facilitates integrative processing through cross-network information convergence, paving the way for future work aimed at improving our knowledge of subcortical contributions to intrinsic brain dynamics and spontaneous cognition.

Received June 1, 2023; revised July 11, 2023; accepted July 28, 2023.

*P.-L.B. and M.M. contributed equally to this work.

The authors declare no competing financial interests.

Correspondence should be addressed to Matthias Mittner at matthias.mittner@uit.no. https://doi.org/10.1523/JNEUROSCI.1020-23.2023

Copyright © 2023 the authors

Introduction

A large body of research in the past decades has focused on descriptions of the macroscopic organization of the human brain in terms of intrinsic functional connectivity (FC) and its role in orchestrating cognition and behavior (Damoiseaux et al., 2006; W. H. Lee et al., 2019; Liégeois et al., 2019). The integration of distributed, functionally specialized brain networks is thought to be essential, especially for higher-level cognition and consciousness (Senden et al., 2014; Bell and Shine, 2016). With a variety of

Author contributions: A.K.H. and B.U.F. designed research; S.M., S.J.S.I., D.H.Y.T., and S.H. performed research; J.M.G., P.-L.B., and M.M. analyzed data; J.M.G. wrote the first draft of the paper; J.M.G., S.M., S.J.S.I., D.H.Y.T., S.H., A.K.H., B.U.F., P.-L.B., and M.M. edited the paper; J.M.G. wrote the paper.

This work was supported by The Netherlands Organization of Scientific Research Grant 016.Vici.185.052 to B.U.F.

methods, specific sites for network convergence have been identified in the posterior cingulate cortex (PCC), anterior cingulate cortex, and the posterior parietal cortices (Tomasi and Volkow, 2011; Bell and Shine, 2015; Lyu et al., 2021), revealing an ensemble of transmodal regions in the cortex that enable efficient global communication (Van der Heuvel and Sporns, 2011; Grayson et al., 2014). With a novel multivariate approach, it was revealed that subtle signals from functionally specialized subdivisions within these regions have connectivity profiles that mirror, or "echo," the activity of different networks, potentially indicating a mechanism through which they facilitate cross-network information integration (Leech et al., 2012; Braga et al., 2013; Braga and Leech, 2015).

Although this work has provided important insights, the dominating corticocentric view overlooks potential contributions from the highly diverse and interconnected structures in the subcortex (Bell and Shine, 2016; Forstmann et al., 2017; Tian et al., 2020). This knowledge gap is likely related to the challenges associated with visualizing the subcortex using conventional MRI because of the varied magnetic tissue properties and generally weaker signal-to-noise ratio (SNR) compared with the cortex (De Hollander et al., 2017; Keuken et al., 2018). Nonetheless, many subcortical structures are part of extensive cortico-subcortical circuitry and demonstrate widespread FC to networks, including the default mode network (Haber, 2003; Bär et al., 2016; T. W. Lee and Xue, 2018; Ji et al., 2019; Li et al., 2021). Compared with the smaller subcortical nuclei in the deep brain, larger structures, such as the thalamus (Tha) and striatum (Str), have received a relatively high amount of attention, establishing their hub-like properties and roles in integrative processing (Choi et al., 2012; Jarbo and Verstynen, 2015; Hwang et al., 2017; Greene et al., 2020; Seitzman et al., 2020; Cheng and Liu, 2021). However, most of the subcortex remains underrepresented in human fMRI studies, and the majority of available evidence is based on lower field strength (3 Tesla), often combined with extensive spatial smoothing, both of which limit the spatial resolution needed to resolve smaller nuclei and increase the risk for signal blurring (De Hollander et al., 2015; Forstmann et al., 2017).

Because of these shortcomings, the functional architecture of the subcortex and its role in integrative processing remain poorly understood. Given that subcortical dysfunction is heavily implicated in a wide range of neuropsychiatric diseases, advancing this knowledge may be vital for our understanding of healthy cognitive functioning as well as improving disease models. Charting the topography of network echoes within the subcortex provides a compelling approach to accomplish new insights into the subcortical contributions to whole-brain communication and higher-level cognition. Following previous work (Leech et al., 2012; Braga et al., 2013), we define an echo as a unique subregional connectivity profile that traces the activity pattern of a functional network. By leveraging recent advances in automated parcellation algorithms and sensitive fMRI protocols for the subcortex at ultra-high field (Bazin et al., 2020; Miletic et al., 2020), we aim to extend the previously established multivariate echo analysis to a large set of subcortical structures, including those rarely studied with human fMRI: the Tha, Str, globus pallidus externa (GPe), globus pallidus interna (GPi), subthalamic nucleus (STN), claustrum (Cl), hippocampus (HPC), amygdala (Amg), substantia nigra (SN), red nucleus (RN), ventral tegmental area (VTA), locus coeruleus (LC), periaqueductal gray (PAG), and pedunculopontine nucleus (PPN). Similar to findings for the cortex, we expect that subcortical structures organized to facilitate multinetwork integration demonstrate a heterogeneous subregional topography of intrinsic echoes from separate functional networks, which are likely hidden with previous univariate connectivity analyses.

Materials and Methods

Participants. The study was approved by the Ethics Review Board of the University of Amsterdam and the Regional Committees for Medical and Health Research Ethics in Norway. Forty healthy adults between 19 and 39 years old (21 female, mean age = 26.5 years, SD = 5.5 years) were recruited from the general population in Norway and screened for MRI compatibility. Exclusion criteria were self-reported (history of) neurologic or psychiatric disease, impaired vision, or any contraindications for MRI, such as metal implants. Written informed consent was obtained from all participants before data collection. All materials, code, and unthresholded group-level statistical maps from multivariate as well as (supplementary) univariate connectivity analyses are publicly available in an Open Science Framework repository at https://osf.io/wt3uc.

fMRI acquisition and preprocessing. Neuroimaging data were collected with a Siemens MAGNETOM Terra 7 Tesla (7 T) system with a 32-channel phased-array head coil. Structural images were obtained with a MP2RAGE sequence (Marques et al., 2010) in 224 sagittal slices at 0.75 mm isotropic voxel resolution (TR = 4300 ms; TI_{1,2} = 840, 2370 ms; flip angles_{1,2} = 5, 6°; TE = 1.99 ms; FOV = 240 × 240 × 168 mm). Functional images were acquired using a gradient EPI sequence with a voxel resolution of 1.5 mm isotropic (82 transverse slices per volume; TR = 1380 ms; TE = 14 ms; flip angle = 60°; in-plane acceleration factor (GRAPPA) = 3; multiband acceleration factor = 2; partial Fourier = 6/8). An additional EPI sequence with opposite phaseencoding direction was performed for susceptibility distortion correction purposes. Heart rate and respiratory data were acquired with a fingerclip and waistband, respectively, to correct for physiological noise, which is especially prominent in the subcortex.

MR images were preprocessed with fMRIPrep (version 20.2.6) (Esteban et al., 2019) in the Nipype framework (Gorgolewski et al., 2011). The structural (T1-weighted) scan was corrected for intensity nonuniformity with N4BiasFieldCorrection (ANTs version 2.3.3) (Tustison et al., 2010) and skull-stripped with antsBrainExtraction using the OASIS30ANTs target template. Brain tissue segmentation of CSF, white matter, and gray matter was performed with FAST (FSL version 5.0.9) (Zhang et al., 2001). For each of the two resting-state runs, a reference volume and its skull-stripped version were generated. A fieldmap based on the EPI references with opposing phaseencoding directions was calculated with 3dQwarp (AFNI) (Cox, 1996), and susceptibility distortion correction was applied to the EPI reference before coregistration to the T1-weighted reference using the boundary-based registration cost-function in bbregister with 6 degrees of freedom (FreeSurfer) (Greve and Fischl, 2009). Headmotion parameters (rotation and translation) were estimated with MCFLIRT (FSL version 5.0.9) (Jenkinson et al., 2002), and slice-time correction to half of the acquisition range (0.674 s) was performed with AFNI's 3dTshift. Following fMRIPrep, data were spatially smoothed with a FWHM Gaussian kernel of 1.5 mm using SUSAN (S. M. Smith and Brady, 1997) and denoised with a first-level GLM in FEAT (Woolrich et al., 2001) that included fMRIPrep-derived confound regressors, including the following: mean signal in CSF and white matter, framewise displacement, six rotation and translation parameters, and discrete-cosine transform basis functions to model low-frequency scanner drifts. In addition, cardiac and respiratory sources of nuisance were based on acquired physiological data and modeled with RETROICOR (Glover et al., 2000) using the MATLAB PhysIO toolbox (Kasper et al., 2017) in TAPAS (Frässle et al., 2021). For one subject with missing physiological data, the same number of fMRIPRrep's anatomic component-based noise correction (aCompCor; Behzadi et al., 2007) regressors were entered in the model instead. The modeled data were obtained via linear regression and normalized. Finally, the two residual runs were concatenated and registered to the ICBM 152 Nonlinear Assymetrical template version 2009c (MNI152Nlin2009cAsym) (Fonov et



Figure 1. Overview of the data analysis.

al., 2009) using the nonlinear registration tool in antsRegistration (Avants et al., 2008) with the transformation parameters provided by fMRIPrep.

Experimental design and ROIs. Two runs of 15 min eyes-open wakeful rest (fixation on centered cross) were collected together with structural (or anatomical) scans during the first of four sessions that were part of a larger multisession 7 T study. The anatomical and experimental data acquired during the other sessions are not part of this study. Figure 1 provides an overview of the analysis, extending the data-driven echo approach (Leech et al., 2012; Braga et al., 2013) to the subcortex. With this multivariate technique, unique FC patterns are estimated while controlling for other subsignals within a region, revealing a more subtle subregional functional organization beyond a region's global connectivity profile that remains concealed with univariate analyses (Leech et al., 2012).

Fourteen subcortical ROIs were defined based on open-source parcellations (Table 1; Fig. 2a). Binary ROI masks were computed from the Multi-contrast Anatomical Subcortical Structure Parcellation algorithm (MASSP) (Bazin et al., 2020) that is based on quantitative MRI data (N = 105, ages 18-80) from the 7 T Amsterdam ultra-high field adult lifespan database (Alkemade et al., 2020) in high-resolution MNI space (MNI152Nlin2009bAsym) (Fonov et al., 2009). The MASSP parcellations include the Tha, Str, Cl, GPe, GPi, SN, STN, VTA, RN, Amg, PAG, and PPN. The LC was defined with the 7 T Probabilistic LC Atlas based on 53 healthy adults aged 52-84 years (Ye et al., 2021). In addition, the 17-network cortical parcellation (Yeo et al., 2011) was used for extracting a mask of the HPC, which was taken from the Default C network. To validate the results for noncortical structures, we also assessed whether we could reproduce the pattern of echoes within various cortical regions, including the PCC, medial prefrontal cortex (mPFC), and visual cortex (Braga et al., 2013). We used the same cortical network parcellation to derive masks for the striate and extrastriate cortex (Visual Central network) and the PCC and mPFC (Default A network). For bilateral ROIs, left and right hemispheres were combined into a single binary mask, and all masks were resampled to the resolution of the functional data

Table 1. Parcellation details for ROIs

			Mean (SD)	
	Abbreviation	N voxels ^{a}	tSNR	Source ^b
Forebrain				
Thalamus	Tha	6130	47.94 (6.31)	MASSP
Striatum	Str	8552	52.17 (8.11)	MASSP
Globus pallidus externa	GPe	1241	35.44 (5.58)	MASSP
Globus pallidus interna	GPi	453	34.18 (4.39)	MASSP
Subthalamic nucleus	STN	93	32.30 (4.25)	MASSP
Claustrum	Cl	683	59.12 (4.71)	MASSP
HPC	HPC	2894	37.84 (10.44)	17-network cortical
				parcellation
Amygdala	Amg	1063	39.89 (7.22)	MASSP
Midbrain				
Substantia nigra	SN	481	31.51 (5.32)	MASSP
Red nucleus	RN	232	33.75 (3.05)	MASSP
Ventral tegmental area	VTA	220	37.68 (3.05)	MASSP
Periaqueductal gray	PAG	198	32.37 (10.56)	MASSP
Brainstem				
Locus coeruleus	LC	98	39.01 (7.31)	7 T Probabilistic
				LC Atlas
Pedunculopontine nucleus	PPN	135	40.00 (3.29)	MASSP

 ${}^{a}\!Number of voxels$ (N voxels) in functional space (1.5 mm isotropic voxel size) and mean (SD) of ROI-wise tSNR values.

^bSource: MASSP (Bazin et al., 2020); 17-network cortical parcellation (Yeo et al., 2011); 7 T Probabilistic LC Atlas (Ye et al., 2021).

with FLIRT using nearest-neighbor interpolation (version 6.0) (Jenkinson and Smith, 2001). The probabilistic LC mask was thresholded liberally so that voxels that overlapped 1% or more were included in the resampled mask.

Statistical analysis. The individual preprocessed resting-state timeseries were masked with each of the binary ROIs and decomposed into 10



Figure 2. Parcellations of subcortical ROIs and reference networks. *a*, Subcortical ROIs defined with open-source atlases and *b* data-driven reference networks from a whole-brain canonical ICA on the resting-state timeseries, labeled according to their maximum spatial correlation with a 17-network cortical parcellation. Corresponding whole-brain tSNR maps are shown in Extended Data Figure 2-1. SomA/B, Somatomotor A/B; ConA/B/C, Control A/B/C; TemPar, Temporal Parietal; DefA/B, Default A/B; VisC, Visual Central; VisP, Visual Peripheral; LimA/B, Limbic A/B; SalA/B, Salience/Ventral Attention A/B.

spatiotemporal independent subregions with a spatially restricted group canonical independent component analysis (canICA) as implemented in Nilearn. Although the temporal concatenation ICA approach is a popular technique in combination with dual regression, biases in the estimation of group-level networks may arise with varying degrees of interindividual variability (Hu and Yang, 2021). Instead, canICA applies a hierarchical approach in which individual data are decomposed before canonical correlation analysis to identify group commonalities (Varoquaux et al., 2010). The ROI-wise canICAs were restricted to find 10 independent components. Model order selection constitutes a main challenge in ICA, and the exact number of underlying signals in the diverse subcortical structures remains unknown. While prior analyses on the PCC demonstrated qualitatively similar outcomes for various model orders (Leech et al., 2012), conducting such comprehensive comparisons for all included structures was beyond the scope of this study. Instead, we opted to follow previous approaches and fix the number of components, addressing interregional differences in network echoes rather than precise dimensionality of individual structures.

Following spatiotemporal decomposition, the unique whole-brain FC of each independent component (subregion) was then investigated with dual regression (Beckmann et al., 2009; Zuo et al., 2010). First, the 10 spatial maps from the canICA were regressed onto every individual's whole-brain resting-state data to estimate the subject-specific time course for each subregion. By simultaneously entering all 10 spatial maps as design matrix, the time course for each subregion was estimated while statistically controlling for the variance in the other subregions' time courses. Second, the 10 subject-specific independent time courses were regressed onto the subject's resting-state data to obtain spatial maps corresponding to the whole-brain, voxel-wise unique FC of each subregion. These subject-level FC maps were then combined in a nonparametric group-level analysis using random permutation testing (5000 permutations) with threshold-free cluster enhancement. This resulted in one group-level t-statistical map for each of the 10 subregions within each individual ROI that was thresholded with family-wise error correction at *p* < 0.05.

To quantify the presence of echoes from canonical resting-state networks within subcortical regions, the thresholded group-level FC maps were spatially correlated with data-driven reference networks obtained from a canICA on the whole-brain timeseries restricted to find 20 independent components. Based on visual inspection and low spatial Pearson product-moment correlation coefficients with an established 17-network cortical parcellation (Yeo et al., 2011), four independent components (r = 0.05, r = 0.04, r = 0.13, r = 0.04) were identified as artifactual and removed from further analysis. The resulting 16 reference networks were masked with the cortical network parcellation to remove any voxels located outside cortical gray matter (e.g., cerebral white matter, subcortex, CSF).

The extent of the spatial correlation between the FC map for each subregion and the reference networks was used to identify whether patterns of cortical network activity were mirrored, or echoed, in the unique subregional time courses.

Results

Data-driven networks correspond to existing cortical network parcellations

The 16 data-driven reference networks were labeled automatically according to their maximum spatial correlation with the well-established 17-network cortical parcellation (Yeo et al., 2011; Fig. 2b), which is based on rs-fMRI data from 1000 individuals. Despite large differences in field strength, data resolution, and parcellation method, we found correlation coefficients ranging from 0.21 to 0.67 (mean r = 0.44, SD = 0.14), generally indicating moderate to good spatial overlap with their reference network counterparts (Fig. 2b, bottom right): Somatomotor A (r = 0.66), Somatomotor B (r = 0.30), Control A (r = 0.46), Control B (r = 0.51), Control C (r = 0.57), Salience/Ventral Attention A (r = 0.34), Salience/Ventral Attention B (r = 0.54), Temporal Parietal (r = 0.21), Dorsal Attention A (DorA, r = 0.46), Dorsal Attention B (DorB, r = 0.25), Default A (r = 0.42), Default B (r = 0.47), Limbic A (r = 0.30), Limbic B (r = 0.41), Visual Central (r = 0.49), and Visual Peripheral (r=0.67). The data-driven Temporal Parietal network also partially overlapped with the Control A network parcellation (r = 0.15).

Together, the reference networks covered 66% of cortical gray matter defined in the parcellation by Yeo et al. (2011). The strongest deviation was observed in the anterior temporal cortex, which was not remedied by increasing model order (40 or 100 independent components) or a cortically restricted canICA. To assess corresponding variations in temporal SNR (tSNR), we calculated voxel-wise tSNR values as the ratio of the mean and SD of the resting-state timeseries after temporal high-pass filtering (1/128 s). Individual tSNR maps were registered to standard MNI space and averaged (voxel-wise) across subjects and runs. Compared with other cortical areas, reduced tSNR in the temporal lobe was



Figure 3. Echoes of intrinsic connectivity networks in the subcortex. *a*, The number of distinct subregions within an ROI with an FC profile that resembled a reference network (Subregions) and the number of different reference networks that were echoed within a region (Networks) both defined by counting above-threshold spatial correlations. *b*, The maximum spatial correlation between each ROI and each reference network, independent of subregion, for nine ROIs that demonstrated at least one above-threshold spatial correlation to any reference network. Subregional FC profiles for a subset of structures and their spatial correlation with reference networks are illustrated in Extended Data Figure 3-1. The same analysis was repeated with reference networks taken from the 17-network cortical parcellation (Yeo et al., 2011) shown in Extended Data Figure 3-2 as well as for three cortical ROIs (Extended Data Fig. 3-3). SomA/B, Somatomotor A/B; ConA/B/C, Control A/B/C; TemPar, Temporal Parietal; DefA/B, Default A/B; VisC, Visual Central; VisP, Visual Peripheral; LimA/B, Limbic A/B; SalA/B, Salience/Ventral Attention A/B.

observed; and as a consequence, temporal networks were underrepresented in the analysis (Extended Data Fig. 2-1).

Subcortical structures echo signals from different restingstate networks

The 10 thresholded FC maps for each ROI, representing the unique whole-brain FC of each subregion at group level, were spatially correlated with the 16 unthresholded spatial maps of the data-driven reference networks. Figure 3a summarizes the degree of network echoes for the nine ROIs that demonstrated at least one spatial correlation with any reference network above a threshold that was arbitrarily set at the 97th percentile of all spatial correlations (r = 0.16). Echoes were summarized by counting above-threshold spatial correlations in terms of (1) the number of reference networks represented in each ROI and (2) the number of subregions that echoed a reference network. For example, six distinct striatal subregions displayed FC profiles that spatially correlated above-threshold with in total 10 different resting-state networks. Figure 3b presents the actual maximum spatial correlations between each ROI and each reference network, independent of subregion. The reference network that was represented most often was the Salience B network, correlating above-threshold with seven ROIs, followed by Default A, Control C, and Visual Peripheral, each with at least one above-threshold spatial correlation with six different ROIs.

Seven subcortical ROIs echoed signals from more than one network, including the following: Tha, Str, HPC, Cl, GPe, SN, and VTA. The former four ROIs furthermore showed that the echoes from different reference networks were distributed among multiple subregions, indicating evidence for a heterogeneous functional organization. In contrast, both the Amg and PPN showed medium and small spatial correlations, respectively, with only one reference network (Amg: r = 0.37 [DefA]; PPN: r = 0.19 [SalB]). The GPi, STN, RN, PAG, and LC failed to show evidence of echoes as none of their subregions demonstrated a connectivity pattern that resembled the pattern of an intrinsic connectivity network. In some cases, a subregion's FC profile was widespread and shared spatial similarity with more than one reference network. Extended Data Figure 3-1 presents a few FC maps to illustrate the diversity and similarity in connectivity profiles to different reference networks across a subset of subcortical structures.

The FC maps of each subregion were also spatially correlated with the 17-network cortical parcellation (Yeo et al., 2011), which yielded generally lower spatial correlations but a qualitatively similar pattern of results (Extended Data Fig. 3-2). To validate these novel results for the subcortex, we repeated the analyses for three cortical regions that were previously investigated. Results for the PCC, mPFC, and visual cortex are presented in Extended Data Figure 3-3 and are largely consistent with previous findings (Leech et al., 2012; Braga et al., 2013).

Topographic organization of functionally heterogeneous subcortical structures

Figure 4 shows the topographic pattern of network echoes in the subregions of the seven ROIs with more than one above-threshold spatial correlation. Subregions are color-coded according to the reference network they echoed most strongly, whereas



Figure 4. Topography of network echoes within heteromodal subcortical structures. Spatiotemporal decomposition of subcortical structures into independent subregions, color-coded according to their strongest network echo or made translucent if their maximum spatial correlation with any reference network did not reach threshold. SomA/B, Somatomotor A/B; ConA/B/C, Control A/B/C; TemPar, Temporal Parietal; DefA/B, Default A/B; VisC, Visual Central; VisP, Visual Peripheral; LimA/B, Limbic A/B; SalA/B, Salience/Ventral Attention A/B.

subregions with a maximum spatial correlation below threshold (r < 0.16) are translucent. For every ROI, there were several subregions that did not mirror the activity in any intrinsic connectivity network, because they were predominantly functionally connected to other subcortical structures or because their signal largely reflected noise on visual inspection.

Five thalamic subregions echoed signals from various reference networks, demonstrating a heterogeneous organization that was mostly symmetrically distributed in bilateral subdivisions. Left and right ventromedial subregions were both most strongly correlated to the Somatomotor A network (left: r = 0.26, right: r = 0.20), although the right subregion's connectivity profile also spatially overlapped with Salience B (r = 0.20). A more dorsomedial bilateral subregion displayed a connectivity pattern that correlated with the pattern of multiple reference networks, including Default A (r = 0.38), Default B (r = 0.32), and Control A (r = 0.25). Another bilateral subregion, more dorsolaterally located, correlated most strongly with the DorA network (r = 0.31), although there was also spatial overlap with Somatomotor A (r = 0.29), DorB (r=0.25), and Visual Peripheral (r=0.24) networks. Finally, the Default B network was represented in the posterior part of the left-sided Tha (r = 0.22).

Within the Str, there were six different subregions that echoed one or more reference networks, located mostly within the caudate nucleus. A subregion primarily in the left tail of the caudate nucleus spatially correlated with the Default B network (r = 0.21), whereas a subregion covering more of the right tail of caudate nucleus most strongly echoed Control B (r = 0.26), although its widespread connectivity pattern also overlapped with Temporal Parietal (r = 0.23) and Salience A (r = 0.22) networks. A bilateral subregion covering the NAc correlated most strongly with Default A (r = 0.40), whereas another bilateral subregion in the mediodorsal part of the caudate head was functionally connected with Control A (r=0.26) and Default B (r=0.21) networks. Subregions that most strongly echoed the Salience A network included a division in the posterior parts of the left caudate tail and left putamen (r = 0.20) as well as a bilateral region in the lateral NAc (r = 0.19).

For the HPC, we observed that different intrinsic connectivity networks were echoed within four different subregions. In the left hemisphere, a posterior dorsal subregion correlated most strongly with Default A (r=0.34), whereas a more ventrally located subregion correlated exclusively with the Limbic A network (r=0.24). A bilateral anteromedial subregion was functionally connected to the Visual Central network (r=0.21), whereas a posterior dorsal subregion in the right hemisphere echoed the Visual Peripheral (r = 0.30) as well as the Dorsal Attention networks (DorA: r = 0.28, DorB: r = 0.30).

Five subregions of the Cl showed an FC profile that correlated with different reference networks. A small, bilateral subregion in the ventral Cl had a widespread cortical connectivity that had the strongest spatial similarity with DorA (r=0.23), but also Somatomotor A (r=0.20), Dorsal attention B (r=0.19), and Salience B (r=0.19) networks. Left and right subdivisions in the posterior part both echoed the Salience A network (r=0.26 and r=0.21), respectively). In addition, an exclusive functional connection with the Default B network was observed in an anterior subregion of the left Cl (r=0.32) and with the Somatomotor A network in a more posterior subregion of the right Cl (r=0.35).

The GPe and SN each had one subregion with a widespread connectivity profile comprising seven and three reference networks, respectively (Fig. 3b). In the GPe, a bilateral dorsolateral subdivision most strongly echoed the Somatomotor A network (r=0.26), but its signal also correlated with activity in DorA (r = 0.23) and Control networks A and B (r = 0.20 and r = 0.22,respectively). The most pronounced network echo within the SN was from Default A (r = 0.24) and came from a bilateral subregion in the medial anterior SN. The same subregion also showed traces from Salience B (r = 0.22) and Control C (r = 0.16) networks. For the VTA, a large inferomedial subdivision in the right hemisphere was most strongly connected to Salience B (r = 0.19) and just below threshold to Visual Peripheral (r = 0.15) networks. Echoes from the Default A network were furthermore present in two other subregions of the VTA, but spatial correlations were weaker (r = 0.15 and r = 0.13).

Discussion

Despite accumulating insights into the mechanisms of functional integration within the cortex, subcortical substrates of cross-network convergence are largely unexplored. Nonetheless, the subcortex is embedded within an extensive cortico-subcortical architecture that is thought to serve integrative rather than purely segregated functions (Haber, 2003). Here, we aimed to more closely examine the underlying functional organization of subcortical nuclei and their subregional connectivity to functional networks across the cortex.

Consistent with our expectations, we show that individual subcortical structures contain a composite of neural signals that can be decomposed into activity traces of intrinsic network activity. In their study, Braga et al. (2013) showed that activity in multiple networks converges at specific transmodal zones in the cortex, as reflected in a mixture of signals that partially correlate with different networks. We demonstrate that this property is not limited to cortical regions by revealing potential mechanisms for multinetwork integration in the subcortex. The results provide the strongest evidence for functional heterogeneity within the Tha, Str, Cl, and HPC, for which we observed a complex pattern of subregional whole-brain FC that resembled spontaneous activity in distinct functional networks. Subregions in left and right hemispheres had similar spatiotemporal signatures that echoed the same functional networks, showing a symmetrical bilateral topography that is consistent with prior work (Cheng and Liu, 2021).

The Tha and Str are the most commonly represented noncortical structures in studies of global brain connectivity, providing support for their putative role as hub regions (Van der Heuvel and Sporns, 2011; Bell and Shine, 2015, 2016). Whereas several studies report an amalgamation of primarily sensory information within thalamic subregions consistent with its gating function (Tomasi and Volkow, 2011; Ji et al., 2019), we observed traces of somatomotor as well as default mode and dorsal attention networks. The somatomotor subdivisions also spatially overlapped with cingulo-opercular regions of the salience network, which aligns with findings of a "motor integration zone" within ventral thalamic nuclei (Greene et al., 2020). Additionally, dorsal attention, somatomotor, and visual networks converged in a dorsolateral subregion, similar albeit slightly less posterior to the "visual integration zone" in the pulvinar nucleus reported earlier (Greene et al., 2020). For the Str, we observed signal echoes from default mode, control, and salience networks predominantly within the caudate head and left tail, right tail, and left putamen, respectively. Despite large methodological differences across studies, these findings are consistent with prior evidence for "cognitive" integration within the Str (Choi et al., 2012; Greene et al., 2020; Seitzman et al., 2020) and supports thalamic and striatal roles in information integration and higher-level cognitive functioning (Haber, 2003; Hwang et al., 2017).

Although organizational principles may broadly concur, precise functional boundaries and network connections diverge across studies. For example, the subregional profiles identified here partially deviate from another data-driven co-partitioning (Cheng and Liu, 2021) and a voxel-wise winner-take-all approach (Seitzman et al., 2020) for the Tha, as well as the from the striatal architecture reported by Choi et al. (2012). Additionally, we found interhemispheric differences in the HPC (i.e., visual and dorsal attention network echoes in the right and default mode and limbic in the left side) that are inconsistent with reports of lateralized subdivisions along an anterior-posterior axis, as well as the location along this axis of the preferential connection to the default mode network (Blessing et al., 2016; Cheng et al., 2020; Ezama et al., 2021). Given differences in connectivity with entorhinal and parahippocampal cortex (Qin et al., 2016; Seoane et al., 2022), it is possible that the extent of hippocampal and surrounding voxels included in the analysis explains some of the discrepancies across studies, which might be further exacerbated by the effects of spatial smoothing. Furthermore, high degrees of individual variability in subcortical anatomy and FC may result in distortions of group-level estimations (De Hollander et al., 2015; Greene et al., 2020; Sylvester et al., 2020; Tian et al., 2020; Marek and Greene, 2021).

Similar to previous observations for the cortex (Braga et al., 2013), we demonstrate that functional heterogeneity is not

ubiquitously present throughout the subcortex. Within the GPe, SN, and VTA, only one subregion's connectivity profile resembled patterns of functional network activity. A region in the dorsolateral GPe echoed somatomotor as well as dorsal attention and control networks, indicating an integrative site that may support its known role in voluntary, planned movement. Both the SN and VTA showed a pattern of converging signals from default mode and salience networks, although less evident in the VTA. Whereas this association with the default mode network is more established (Bär et al., 2016; Zhang et al., 2016; Edlow, 2021; Li et al., 2021), connectivity to the salience network is less known and may indicate involvement in attention and spontaneous cognition (O'Callaghan et al., 2021).

No clear evidence for functional integration was observed for the Amg and PPN. Whereas the PPN likely takes part in more specialized subcortical circuitry involved in arousal and locomotion (Martinez-Gonzales et al., 2011; Bennarroch, 2013), the Amg was previously proposed as hub structure (Tomasi and Volkow, 2011) and showed dissociable FC profiles from its separate nuclei (Kerestes et al., 2017). Although we did not find evidence for such heterogeneity when controlling for other subregional time courses, we observed an intact connection with the default mode network, which is supported by other work (Kerestes et al., 2017; Sylvester et al., 2020; Harrison et al., 2021). For the remaining structures (i.e., GPi, STN, RN, PAG, and LC), we failed to find network echoes. Although previous univariate FC studies have indicated correlations with widespread cortical activity for some of these structures (e.g., Zhang et al., 2016; Anteraper et al., 2018), the multivariate analysis here did not result in a clear group-level pattern of cortical connectivity. Similar to the PPN, these structures may be less involved in integrating spontaneous signals from distributed functional processes across the cortex, but are likely more strongly embedded in local networks to support segregated functional processing (Singh et al., 2022). Recent findings suggest that neuromodulatory nuclei for dopaminergic and noradrenergic systems are driving systems-level integration and cognition (Zhang et al., 2016; De Gee et al., 2017; Liu et al., 2017). However, not all findings converge. For example, Bär et al. (2016) showed that LC connectivity to the default mode network disappeared when controlling for adjacent neural signals and that hub-like features of midbrain nuclei were not supported by a graph theory analysis. The results presented here align with this observation and emphasize that integrative properties of these structures, among which the LC, remain somewhat elusive. Given proposed roles of the LC in mediating the dynamics of cortical connectivity and neural gain (Aston-Jones and Cohen, 2005; Munn et al., 2021), it is perhaps not surprising that no dissociable traces of functional network activity are observed. That is, the LC may drive global states of network integration and segregation rather than serving as a convergence zone in itself.

In conclusion, our results suggest that subcortical structures exhibit varying degrees of functional heterogeneity. This characteristic might be expressed along a gradient, where structures adjacent to the cortex seem more likely to support multinetwork integration compared with deep brain nuclei. However, several factors may confound interpretations of interregional differences in the subcortex. For example, deep brain nuclei are generally smaller in size and have weaker SNR, while subcortex near the cortex is susceptible to signal bleeding from adjacent cortical voxels, to which they are also reciprocally connected (Choi et al., 2012). This issue might be especially prominent in the Cl, which is a thin sheet-like structure situated directly between the Str and insula. In a recent study, Krimmel et al. (2019) used a novel regression technique on similar high-resolution fMRI data (1.5 mm isotropic voxels) to isolate the signal in the Cl from nearby cortical and striatal voxels, which preserved the widespread FC with cortical networks involved in attention and cognitive control. Although we did not correct for potential signal bleeding beyond limiting the amount of spatial smoothing, our finding of functionally heterogeneous network echoes within the Cl's subdivisions coincides with this work and its postulated role in attention and cognition (Bell and Shine, 2015; Krimmel et al., 2019; J. B. Smith and Jackson, 2020).

It should be noted that recent work highlights the difference in FC between eyes-open and eyes-closed resting-state conditions, particularly with regard to internetwork connectivity of visual and sensorimotor networks to default mode and salience networks (Agcaoglu et al., 2019; Costumero et al., 2020; Han et al., 2023). While a large portion of studies on subcortical connectivity cited here are correspondingly based on eyes-open restingstate fMRI (e.g., Choi et al., 2012; Blessing et al., 2016; Hwang et al., 2017; Greene et al., 2020; Seitzman et al., 2020; Sylvester et al., 2020), future efforts could contrast our results to potential reconfigurations during other resting-state and experimental conditions. Investigating changes in the pattern of echoes according to external factors, such as cognitive demand, and internal state are likely necessary to illuminate their functional relevance (e.g., Leech et al., 2012).

Although the precise significance of network echoes for cognition and behavior is not resolved, we strengthen the evidence that the subcortex participates in cross-network integration through echoing intrinsic network activity. These results may ignite new intriguing hypotheses on the mechanisms of spontaneous cognitive processes, such as mind wandering (Mittner et al., 2016; Zuberer et al., 2021). Previous work has shown that mind wandering correlates with activity and connectivity in the default mode and frontoparietal control networks as well as the subcortex (Mittner et al., 2014; Kucyi et al., 2017; Groot et al., 2022). Given that both subtle and pronounced reorganizations in FC occur with changes in task demand (Leech et al., 2012; Braga et al., 2013; Tian et al., 2020), investigations of how the complex pattern of echoes in the subcortex is perturbed by attentional changes may reveal novel insights into the mechanisms that drive mind wandering.

References

- Agcaoglu O, Wilson TW, Wang YP, Stephen J, Calhoun VD (2019) Resting state connectivity differences in eyes open versus eyes closed conditions. Hum Brain Mapp 40:2488–2498.
- Alkemade A, Mulder MJ, Groot JM, Isaacs BR, Van Berendonk N, Lute N, Isherwood SJ, Bazin PL, Forstmann BU (2020) The Amsterdam Ultra-High Field Adult Lifespan Database (AHEAD): a freely available multimodal 7 Tesla submillimeter magnetic resonance imaging database. Neuroimage 221:117200.
- Anteraper SA, Guell X, Whitfield-Gabrieli S, Triantafyllou C, Mattfeld AT, Gabrieli JD, Geddes MR (2018) Resting-state functional connectivity of the subthalamic nucleus to limbic, associative, and motor networks. Brain Connect 8:22–32.
- Aston-Jones G, Cohen JD (2005) An integrative theory of locus coeruleusnorepinephrine function: adaptive gain and optimal performance. Annu Rev Neurosci 28:403–450.
- Avants B, Tustison NJ, Song G (2008) Advanced normalization tools (ANTS). Insight J. 1–35.
- Bär KJ, De la Cruz F, Schumann A, Koehler S, Sauer H, Critchley H, Wagner G (2016) Functional connectivity and network analysis of midbrain and brainstem nuclei. Neuroimage 134:53–63.

- Bazin PL, Alkemade A, Mulder MJ, Henry AG, Forstmann BU (2020) Multicontrast anatomical subcortical structures parcellation. Elife 9:e59430.
- Beckmann CF, Mackay CE, Filippini N, Smith SM (2009) Group comparison of resting-state fMRI data using multi-subject ICA and dual regression. Neuroimage 47:S148.
- Behzadi Y, Restom K, Liau J, Liu TT (2007) A component based noise correction method (CompCor) for BOLD and perfusion based fMRI. Neuroimage 37:90–101.
- Bell PT, Shine JM (2015) Estimating large-scale network convergence in the human functional connectome. Brain Connect 5:565–574.
- Bell PT, Shine JM (2016) Subcortical contributions to large-scale network communication. Neurosci Biobehav Rev 71:313–322.
- Bennarroch EE (2013) Pedunculopontine nucleus: functional organization and clinical implications. Neurology 80:1148–1155.
- Blessing EM, Beissner F, Schumann A, Brünner F, Bär KJ (2016) A datadriven approach to mapping cortical and subcortical intrinsic functional connectivity along the longitudinal hippocampal axis. Hum Brain Mapp 37:462–476.
- Braga RM, Leech R (2015) Echoes of the brain: local-scale representation of whole-brain functional networks within transmodal cortex. Neuroscientist 21:540–551.
- Braga RM, Sharp DJ, Leeson C, Wise RJ, Leech R (2013) Echoes of the brain within default mode, association, and heteromodal cortices. J Neurosci 33:14031–14039.
- Cheng H, Liu J (2021) Concurrent brain parcellation and connectivity estimation via co-clustering of resting state fMRI: a novel approach. Hum Brain Mapp 42:2477–2489.
- Cheng H, Zhu H, Zheng Q, Liu J, He G (2020) Functional parcellation of the hippocampus by semi-supervised clustering of resting state fMRI data. Sci Rep 10:16402.
- Choi EY, Yeo BT, Buckner RL (2012) The organization of the human striatum estimated by intrinsic functional connectivity. J Neurophysiol 108:2242–2263.
- Costumero V, Bueichekú E, Adrián-Ventura J, Ávila C (2020) Opening or closing eues at rest modulates the functional connectivity of V1 with default and salience networks. Sci Rep 10:9137.
- Cox RW (1996) AFNI: software for analysis and visualization of functional magnetic resonance neuroimages. Comput Biomed Res 29:162–173.
- Damoiseaux JS, Rombouts SA, Barkhof F, Scheltens P, Stam CJ, Smith SM, Beckmann CF (2006) Consistent resting-state networks across healthy subjects. Proc Natl Acad Sci USA 103:13848–13853.
- De Gee JW, Colizoli O, Kloosterman NA, Knapen T, Nieuwenhuis S, Donner TH (2017) Dynamic modulation of decision biases by brainstem arousal systems. Elife 6:e23232.
- De Hollander G, Keuken MC, Forstmann BU (2015) The subcortical cocktail problem: mixed signals from the subthalamic nucleus and substantia nigra. PLoS One 10:e0120572.
- De Hollander G, Keuken MC, Van der Zwaag W, Forstmann BU, Trampel R (2017) Comparing functional MRI protocols for small, iron-rich basal ganglia nuclei such as the subthalamic nucleus at 7 T and 3 T. Hum Brain Mapp 38:3226–3248.
- Edlow BL (2021) Dopaminergic modulation of human consciousness via default mode network connectivity. Proc Natl Acad Sci USA 118:e2111268118.
- Esteban O, Markiewicz CJ, Blair RW, Moodie CA, Isik AI, Erramuzpe A, Kent JD, Goncalves M, DuPre E, Snyder M, Oya H, Ghosh SS, Wright J, Durnez J, Poldrack RA, Gorgolewski KJ (2019) fMRIPrep: a robust preprocessing pipeline for functional MRI. Nat Methods 16:111–116.
- Ezama L, Hernández-Cabrera JA, Seoane S, Pereda E, Janssen N (2021) Functional connectivity of the hippocampus and its subfields in restingstate networks. Eur J Neurosci 53:3378–3393.
- Fonov VS, Evans AC, McKinstry RC, Almli CR, Collins DL (2009) Unbiased nonlinear average age-appropriate brain templates from birth to adulthood. Neuroimage 47:S102.
- Forstmann BU, De Hollander G, Van Maanen L, Alkemade A, Keuken MC (2017) Towards a mechanistic understanding of the human subcortex. Nat Rev Neurosci 18:57–65.
- Frässle S, et al. (2021) TAPAS: an open-source software package for Translational Neuromodeling and Computational Psychiatry. Front Psychiatry 12:680811.

- Glover GH, Li TQ, Ress D (2000) Image-based method for retrospective correction of physiological motion effects in fMRI: RETROICOR. Magn Reson Med 44:162–167.
- Gorgolewski K, Burns CD, Madison C, Clark D, Halchenko YO, Waskom ML, Ghosh SS (2011) Nipype: a flexible, lightweight and extensible neuroimaging data processing framework in Python. Front Neuroinform 5:13.
- Grayson DS, Ray S, Carpenter S, Iyer S, Costa Dias TG, Stevens C, Nigg JT, Fair DA (2014) Structural and functional rich club organization of the brain in children and adults. PLoS One 9:e88297.
- Greene DJ, et al. (2020) Integrative and network-specific connectivity of the basal ganglia and thalamus defined in individuals. Neuron 105:742–758.e6.
- Greve DN, Fischl B (2009) Accurate and robust brain image alignment using boundary-based registration. Neuroimage 48:63–72.
- Groot JM, Csifcsák G, Wientjes S, Forstmann BU, Mittner M (2022) Catching wandering minds with tapping fingers: neural and behavioral insights into task-unrelated cognition. Cereb Cortex 32:4447–4463.
- Haber SN (2003) The primate basal ganglia: parallel and integrative networks. J Chem Neuroanat 26:317–330.
- Han J, Zhou L, Wu H, Huang Y, Qiu M, Huang L, Lee C, Lane TJ, Qin P (2023) Eyes-open and eyes-closed resting state network connectivity differences. Brain Sci 13:122.
- Harrison OK, Guell X, Klein-Flügge MC, Barry RL (2021) Structural and resting state functional connectivity beyond the cortex. Neuroimage 240:118379.
- Hu Y, Yang Z (2021) Impact of inter-individual variability on the estimation of default mode network in temporal concatenation group ICA. Neuroimage 237:118114.
- Hwang K, Bertolero MA, Liu WB, D'Esposito M (2017) The human thalamus is an integrative hub for functional brain networks. J Neurosci 37:5594– 5607.
- Jarbo K, Verstynen TD (2015) Converging structural and functional connectivity of orbitofrontal, dorsolateral, prefrontal, and posterior parietal cortex in the human striatum. J Neurosci 35:3865–3878.
- Jenkinson M, Bannister P, Brady JM, Smith SM (2002) Improved optimisation for the robust and accurate linear registration and motion correction of brain images. Neuroimage 17:825–841.
- Jenkinson M, Smith SM (2001) A global optimisation method for robust affine registration of brain images. Med Image Anal 5:143–156.
- Ji JL, Spronk M, Kulkarni K, Repovs G, Anticevic A, Cole MW (2019) Mapping the human brain's cortical-subcortical functional network organization. Neuroimage 185:35–57.
- Kasper L, Bollmann S, Diaconescu AO, Hutton C, Heinzle J, Iglesias S, Hauser TU, Sebold M, Manjaly ZM, Pruessmann KP, Stephan KE (2017) The PhysIO Toolbox for modeling physiological noise in fMRI data. J Neurosci Methods 276:56–72.
- Kerestes R, Chase HW, Philips ML, Ladouceur CD, Eickhoff SB (2017) Multimodal evaluation of the amygdala's functional connectivity. Neuroimage 148:219–229.
- Keuken MC, Isaacs BR, Trampel R, Van der Zwaag W, Forstmann BU (2018) Visualizing the human subcortex using ultra-high field magnetic resonance imaging. Brain Topogr 31:513–545.
- Krimmel SR, White MG, Panicker MH, Barrett FS, Mathur BN, Seminowicz DA (2019) Resting state functional connectivity and cognitive task-related activation of the human claustrum. Neuroimage 196:59–67.
- Kucyi A, Hove MJ, Esterman M, Hutchison RM, Valera EM (2017) Dynamic brain network correlates of spontaneous fluctuations in attention. Cereb Cortex 27:1831–1840.
- Lee TW, Xue SW (2018) Functional connectivity maps based on hippocampal and thalamic dynamics may account for the default-mode network. Eur J Neurosci 47:388–398.
- Lee WH, Moser DA, Ing A, Doucet GE, Frangou S (2019) Behavioral and health correlates of resting-state metastability in the Human Connectome Project. Brain Topogr 32:80–86.
- Leech R, Braga R, Sharp DJ (2012) Echoes of the brain within the posterior cingulate cortex. J Neurosci 32:215–222.
- Li J, Curley WH, Guerin B, Dougherty DD, Dalca AV, Fischl B, Horn A, Edlow BL (2021) Mapping the subcortical connectivity of the human default mode network. Neuroimage 245:118758.

- Liégeois R, Li J, Kong R, Orban C, Van de Ville D, Ge T, Sabuncu MR, Yeo T (2019) Resting brain dynamics at different timescales capture distinct aspects of human behavior. Nat Commun 10:2317.
- Liu KY, Marijatta F, Hämmerer D, Acosta-Cabronero J, Düzel E, Howard RJ (2017) Magnetic resonance imaging of the human locus coeruleus: a systematic review. Neurosci Biobehav Rev 83:325–355.
- Lyu D, Pappas I, Menon DK, Stamatakis EA (2021) A precuneal causal loop mediates external and internal information integration in the human brain. J Neurosci 41:9944–9956.
- Marek S, Greene DJ (2021) Precision functional mapping of the subcortex and cerebellum. Curr Opin Behav Sci 40:12–18.
- Marques JP, Kober T, Krueger G, Van der Zwaag W, Van de Moortele PF, Gruetter R (2010) MP2RAGE, a self bias-field corrected sequence for improved segmentation and T1-mapping at high field. Neuroimage 49:1271–1281.
- Martinez-Gonzales C, Bolam JP, Mena-Segovia J (2011) Topographical organization of the pedunculopontine nucleus. Front Neuroanat 5:22.
- Miletic S, Bazin PL, Weiskopf N, Van der Zwaag W, Forstmann BU, Trampel R (2020) fMRI protocol optimization for simultanously studying small subcortical and cortical areas at 7 T. Neuroimage 219:116992.
- Mittner M, Boekel W, Tucker AM, Turner BM, Heathcote A, Forstmann BU (2014) When the brain takes a break: A model-based analysis of mind wandering. J Neurosci 34:16286–16295.
- Mittner M, Hawkins GE, Boekel W, Forstmann BU (2016) A neural model of mind wandering. Trends Cogn Sci 20:570–578.
- Munn BR, Müller EJ, Wainstein G, Shine JM (2021) The ascending arousal system shapes neural dynamics to mediate awareness of cognitive states. Nat Commun 12:6016.
- O'Callaghan C, Walpola IC, Shine JM (2021) Neuromodulation of the mindwandering brain state: the interaction between neuromodulatory tone, sharp wave-ripples and spontaneous thought. Philos Trans R Soc Lond B Biol Sci 376:20190699.
- Qin S, Duan X, Supekar K, Chen H, Chen T, Menon V (2016) Largescale intrinsic functional network organization along the long axis of the human medial temporal lobe. Brain Struct Funct 221:3237– 3258.
- Seitzman BA, Gratton C, Marek S, Raut RV, Dosenbach NU, Schlagger BL, Petersen SE, Greene DJ (2020) A set of functionally-defined brain regions with improved representation of the subcortex and cerebellum. Neuroimage 206:116290.
- Senden M, Deco G, De Reus MA, Goebel R, Van den Heuvel MP (2014) Rich club organization supports a diverse set of functional network configurations. Neuroimage 96:174–182.
- Seoane S, Modroño C, Gonzáles-Mora J, Janssen N (2022) Medial temporal lobe contributions to resting-state networks. Brain Struct Funct 227:995– 1012.
- Singh K, Cuazzo S, García-Gomar MG, Stauder M, Vanello N, Passino C, Bianciardi M (2022) Functional connectome of arousal and motor brainstem nuclei in living humans by 7 Tesla resting-state fMRI. Neuroimage 249:118865.
- Smith JB, Lee AK, Jackson J (2020) The claustrum. Curr Biol 30:R1401– R1406.
- Smith SM, Brady JM (1997) SUSAN: a new approach to low level image processing. Int J Comput Vis 23:45–78.
- Sylvester CM, et al. (2020) Individual-specific functional connectivity of the amygdala: a substrate for precision psychiatry. Proc Natl Acad Sci USA 117:3808–3818.
- Tian Y, Margulies DS, Breakspear M, Zalesky A (2020) Topographic organization of the human subcortex unveiled with functional connectivity gradients. Nat Neurosci 23:1421–1432.
- Tomasi D, Volkow ND (2011) Association between functional connectivity hubs and brain networks. Cereb Cortex 21:2003–2013.
- Tustison NJ, Avants BB, Cook PA, Zheng Y, Egan A, Yushkevich PA, Gee JC (2010) N4ITK: improved N3 bias correction. IEEE Trans Med Imaging 29:1310–1320.
- Van der Heuvel MP, Sporns O (2011) Rich-club organization of the human connectome. J Neurosci 31:15775–15786.
- Varoquaux G, Sadaghiani S, Pinel P, Kleinschmidt A, Poline JB, Thirion B (2010) A group model for stable multi-subject ICA from fMRI datasets. Neuroimage 51:288–299.

- Woolrich MW, Ripley BD, Brady M, Smith SM (2001) Temporal autocorrelation in univariate linear modeling of FMRI data. Neuroimage 14:1370–1386.
- Ye R, Rua C, O'Callaghan C, Jones PS, Hezemans FH, Kaalund SS, Tsvetanov KA, Rodgers CT, Williams G, Passamonti L, Rowe JB (2021) An in vivo probabilistic atlas of the human locus coeruleus at ultra-high field. Neuroimage 225:117487.
- Yeo BT, Krienen FM, Sepulcre J, Sabuncu MR, Lashkari D, Hollinshead M, Roffman JL, Smoller JW, Zöllei L, Polimeni JR, Fischl B, Liu H, Buckner RL (2011) The organization of the human cerebral cortex estimated by intrinsic functional connectivity. J Neurophysiol 106:1125–1165.
- Zhang S, Hu S, Chao HH, Li CS (2016) Resting-state functional connectivity of the locus coeruleus in humans: in comparison with the ventral

tegmental area/substantia nigra pars compacta and the effects of age. Cereb Cortex 26:3413-3427.

- Zhang Y, Brady M, Smith S (2001) Segmentation of brain MR images through a hidden Markov random field model and the expectation-maximization algorithm. IEEE Trans Med Imaging 20:45–57.
- Zuberer A, Kucyi A, Yamashita A, Wu CM, Walter M, Valera EM, Esterman M (2021) Integration and segregation across large-scale intrinsic brain networks as a marker of sustained attention and task-unrelated thought. Neuroimage 229:117610.
- Zuo XN, 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.