Contents lists available at ScienceDirect

NeuroImage

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Brain structure-function coupling provides signatures for task decoding and individual fingerprinting



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ARTICLE INFO

Keywords: Graph signal processing Fingerprinting Decoding Task fMRI Functional connectivity

ABSTRACT

Brain signatures of functional activity have shown promising results in both decoding brain states, meaning distinguishing between different tasks, and fingerprinting, that is identifying individuals within a large group. Importantly, these brain signatures do not account for the underlying brain anatomy on which brain function takes place. Structure-function coupling based on graph signal processing (GSP) has recently revealed a meaningful spatial gradient from unimodal to transmodal regions, on average in healthy subjects during resting-state. Here, we explore the specificity of structure-function coupling to distinct brain states (tasks) and to individual subjects. We used multimodal magnetic resonance imaging of 100 unrelated healthy subjects from the Human Connectome Project both during rest and seven different tasks and adopted a support vector machine classification approach for both decoding and fingerprinting, with various cross-validation settings. We found that structurefunction coupling measures allow accurate classifications for both task decoding and fingerprinting. In particular, key information for fingerprinting is found in the more liberal portion of functional signals, with contributions strikingly localized to the fronto-parietal network. Moreover, the liberal portion of functional signals showed a strong correlation with cognitive traits, assessed with partial least square analysis, corroborating its relevance for fingerprinting. By introducing a new perspective on GSP-based signal filtering and FC decomposition, these results show that brain structure-function coupling provides a new class of signatures of cognition and individual brain organization at rest and during tasks. Further, they provide insights on clarifying the role of low and high spatial frequencies of the structural connectome, leading to new understanding of where key structure-function information for characterizing individuals can be found across the structural connectome graph spectrum.

1. Introduction

The existence of brain signatures based on functional magnetic resonance imaging (fMRI), meaning specific features uniquely characterizing either tasks or individuals, has emerged from the development of advanced data analysis methods in the last two decades. On the one hand, the application of pattern recognition techniques to neuroimaging data proved the capability of fMRI to *decode* task-specific brain activity (Gao et al., 2020; Haynes and Rees, 2006; Li and Fan, 2019; Richiardi et al., 2011; Wang et al., 2020). Significant progress in this direction was made by the recent advent of deep learning (Gao et al., 2020; Li and Fan, 2019; Wang et al., 2020), even if it remains nontrivial to interpret the biological meaning of the learned features. On the other hand, similarly to a *fingerprint*, fMRI-based features can accurately identify individuals from a large group (Amico and Goñi, 2018a; Biazoli et al., 2017; Finn et al., 2015; Mansour et al., 2021; Van De Ville et al., 2021). In a seminal paper by Finn et al. (2015), functional connectivity (FC) profiles were used to successfully classify subjects across resting state test-retest sessions, and even between task and rest conditions. The fronto-parietal network emerged as the main contributor to subject discrimination, and was shown to predict individual cognitive behavior (i.e., level of fluid intelligence). In addition to functional activity, brain anatomical features, such as cortical morphology and whitematter structural connectivity, were also proven useful for brain finger-

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https://doi.org/10.1016/j.neuroimage.2022.118970.

Received 14 November 2021; Received in revised form 12 January 2022; Accepted 3 February 2022 Available online 4 February 2022. 1053-8119/© 2022 The Author(s). Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/)



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printing (Kumar et al., 2017; Lin et al., 2020; Valizadeh et al., 2018; Wachinger et al., 2015; Yeh et al., 2016).

In this context, a still unexplored brain feature, which could offer new insights into task decoding, individual fingerprinting and behavioral correlates, is the degree of coupling between function and structure, i.e., how brain functional activity and connectivity align to the underlying structural connectivity architecture as measured with diffusion-weighted (DW) MRI. Early attempts to investigate structurefunction relationships in the brain spanned from simple approaches, such as correlational analyses (Amico and Goñi, 2018b; Goñi et al., 2014; Honey et al., 2009; Mišić et al., 2016; Zhang et al., 2011), to more complex ones, like whole brain computational and communication models (Amico et al., 2021; Avena-Koenigsberger et al., 2018; Deco et al., 2011; Griffa et al., 2017; Mišić et al., 2015; Seguin et al., 2020). More recently, graph signal processing provided a novel framework for a combined structure-function analysis (Huang et al., 2018; Medaglia et al., 2018; Preti and Van De Ville, 2019). Within this setting, Preti and Van De Ville (2019) quantified the degree of structure-function dependency for each brain region, by means of the newly introduced Structural-Decoupling Index (SDI). This nodal metric quantifies the degree of local (dis)alignment between structure and function, and it is obtained by decomposing the structural connectome into harmonics in the graph frequency domain, and projecting the functional signals (fMRI frames at each timepoint) in the space spanned by the structural harmonics. The functional signals are then filtered into low and high structural graph frequencies, giving rise to coupled and decoupled signal components, respectively. The ratio between the energy of these two signal portions yields the SDI of a brain region. During resting state in healthy subjects, local structure-function (de)coupling showed a very characteristic and behaviorally relevant spatial distribution, spanning from lower-order functional areas such as visual and somatosensory cortices, with activity highly constrained by the structure underneath, to higher-order ones, with activity more liberal. However, the extent to which this configuration changes in different task-related states, or in different subjects, still remains unexplored. Moreover, the quantification of the structurefunction coupling at the level of single brain connections may bring new insights into brain organization principles and their uniqueness to brain states and individuals. In particular, do structure-function dependency patterns represent a signature of a particular task-related state? Can they act as a brain fingerprint uniquely identifying individuals? And which structure-function dependency features are more relevant to task decoding, subject fingerprinting, and inter-individual cognitive variability?

To answer these open questions, we analyzed the structural and functional data during resting state and seven different tasks of 100 unrelated healthy subjects from the Human Connectome Project (HCP) (Van Essen et al., 2013), and obtained their structure-function signatures quantified through: (i) the SDI, and (ii) a new GSP-based decomposition of the FC. The latter is obtained by assessing the functional connectivity between fMRI signal components that are more coupled or decoupled to the underlying structure, named coupled-FC (c-FC) and decoupled-FC (d-FC), respectively. These GSP-derived features quantify brain structure-function coupling at the level of either single brain regions (SDI) or single brain connections (c-FC, d-FC) and were used to classify different tasks and individuals. In both cases, the classification showed high accuracy for all the three structure-function coupling measures, across various cross-validation settings. Two specific networks including regions that are key to either task decoding or individual fingerprinting based on structure-function coupling emerged. Results were then compared with the classification performances obtained with conventional nodal (node strength) and edgewise measures of FC, without knowledge from the underlying structure. Finally, nodewise and edgewise structure-function couplings in resting state were shown to correlate with individual cognitive traits including fluid intelligence and sustained attention, particularly in the high-frequency FC components (d-FC) of the structural connectome.

2. Material and methods

2.1. Methods outline

The methodological pipeline is illustrated in Fig. 1. From the fMRI timecourses (Fig. 1A) of 100 individuals during rest and seven tasks, conventional edgewise and nodal FC measures (FC matrix and FC node strength) were computed (Fig. 1B). In parallel to that, the GSP pipeline outlined in (Preti and Van De Ville, 2019) was implemented to decompose functional signals at each timepoint onto the underlying structural bases and filter them in coupled (low-frequency) and decoupled (high-frequency) portions (Fig. 1C). Structure-function coupling was then evaluated at the level of connections and regions, by means of coupled and decoupled FC and structural-decoupling indexes, respectively (Fig. 1D). c-FC and d-FC are FC matrices derived from the coupled and decoupled portions of fMRI timecourses. The SDI quantifies instead the amount of local alignment between brain functional signals and the underlying structural connectivity network at the nodal level. Next, the task decoding and individual fingerprinting accuracy obtained from the nodal and edgewise structure-function coupling features (SDI, c-FC and d-FC, Fig. 1D), as well as from nodal and edgewise measures of FC not taking into account the underlying brain structure (Fig. 1B), were assessed with support vector machine (SVM) classification (Fig. 1E) and compared. Finally, multivariate relationships between the different nodal and edgewise features and individual cognitive traits were assessed with partial least square correlation (PLSC) analyses.

2.2. Data and preprocessing

100 unrelated healthy subjects (number of subjects $N_s = 100$) of the HCP dataset U100 - HCP900 data release (54 females, 64 males, mean age = 29.1 ± 3.7 years) were included in the study. Ethical approval was obtained within the HCP. Analyses were restricted to these 100 subjects to ensure absence of any family relationship which may influence fingerprinting results. fMRI acquired with $N_T = 8$ different task conditions (resting state and 7 tasks: emotion, gambling, language, motor, relation, social, working memory), each recorded with $N_E = 2$ phase encoding directions (right-left and left-right), as well as DW-MRI sequences were pre-processed with state-of-the-art pipelines, in order to obtain regional functional time courses and their structural connections, based on a parcellation with $N_{ROI} = 379$ regions (360 cortical areas (Glasser et al., 2016) and 19 subcortical ones as provided by the HCP release (Fischl et al., 2002; Glasser et al., 2013)). Each brain area was assigned to one of the 7 Yeo et al. (2011) networks through majority voting procedure or to the subcortical network for post hoc analyses. Minimally preprocessed data from the HCP were selected (Glasser et al., 2013; Van Essen et al., 2013) and the following additional pre-processing steps were performed. Nuisance signals were removed from voxel fMRI time courses (linear and quadratic trends, six motion parameters and their first derivatives, average white matter and cerebrospinal fluid signals and their first derivatives) and average time courses were computed in each region of the parcellation, previously resampled to the functional resolution, and z-scored. To remove the effect of the paradigm on task data, only for task classification, paradigms were regressed out trial by trial from functional time courses (a separate regressor for each task trial was included in the model). Functional connectomes were obtained as Pearson's correlation between pairwise time courses and FC nodal strength was computed for each region as the sum of absolute values of all the connections of that region (Fig. 1B).

The same DW-MRI processing pipeline detailed in (Preti and Van De Ville, 2019) was used to reconstruct whole brain tractograms including 2 million fibers, using a spherical deconvolution approach and the Spherical-deconvolution Informed Filtering of Tractograms 2 (SIFT2 (Smith et al., 2015a), https://www.mrtrix.org/). Structural connectomes were then obtained, after resampling of the same parcellation to diffusion space, as the number of streamlines connecting two regions,



Fig. 1. Method workflow. From fMRI nodal signals at each timepoint (A), functional connectivity (FC) is evaluated through conventional edgewise (FC matrix) and nodewise (FC node strength) measures (B). The graph signal processing (GSP) pipeline is applied to decompose functional signals into the structural harmonics obtained from the eigendecomposition of the structural connectome (SC) Laplacian (C). Functional signals are then filtered into two components; i.e., one coupled and one decoupled from structure, by applying ideal low pass (light blue) and high pass (pink) filters in the graph spectral domain (C). Edgewise and nodewise metrics evaluating structure-function coupling are obtained by computing FC matrices from coupled and decoupled signals (coupled and decoupled FC (c-FC and d-FC), respectively), and the structural decoupling index (SDI). Edgewise and nodal measures of both FC (B) and structure-function coupling (D) enter separate support vector machine (SVM) classifications with various cross validation settings to test their task decoding and fingerprinting value, quantified by task and subject identification accuracies (E).

normalized by the sum of the two regions' volumes. An average structural connectivity (SC) matrix, representative of the whole population, was obtained by averaging the structural connectivity values across subjects.

2.3. Structure-function coupling features

The graph signal processing framework detailed in (Preti and Van De Ville, 2019) was adopted to obtain structure-function signatures (the SDI and the newly introduced c-FC and d-FC) for each subject and acquisition. In brief, the average SC across the population is decomposed into structural harmonics u_k by eigendecomposition of the SC Laplacian $L = I - A_{symm}$ (given the identity matrix I and the symmetrically normalized adjacency matrix A_{symm} of the SC):

$LU = U\Lambda,$

where each eigenvalue $[\Lambda]_{k,k} = \lambda_k$ can be interpreted as spatial frequency of the corresponding structural harmonic (eigenvector) u_k . For each subject, functional data at each timepoint s_t is then projected onto the structural harmonics by assessing spectral coefficients $\hat{s}_t = U^T s_t$, and filtered into two components with ideal low- and high-pass filters (Fig. 1C). A fixed value of c = 50 spectral components were chosen, to be common to all acquisitions, and avoid task- or individual- biases that could affect the following classification. In addition, decoding and fingerprinting analyses were repeated using a median split on the observed energy spectral densities (determined for each subject and task independently) for comparison with previous work (Preti and Van De Ville, 2019). The filtering operation yielded a low-frequency functional activity component $s_t^C = U^{(low)} U^T s_t$, which is coupled to the structure, and a high-frequency one $s_t^D = U^{(high)}U^T s_t$, more decoupled from the structure (where $U^{(low)}$ and $U^{(high)}$ are $N_{ROI} \times N_{ROI}$ matrices with the c first eigenvectors complemented by zeros, and with c first columns of zeros followed by the $N_{ROI} - c$ last eigenvectors, respectively). Pairwise Pearson's correlations of s^C and s^D timecourses were computed to obtain c-FC and d-FC matrices, respectively. The L2 norm across time of s^{C} and s^{D} yielded instead a general measure of coupling and decoupling for each node, and the ratio between the two corresponds to the SDI (Fig. 1D).

2.4. Decoding and fingerprinting patterns

A series of two-factor ANOVA analyses with nodal SDI, nodal FC strength, edgewise FC, c-FC, or d-FC as dependent factor, and subject and task as independent factors, were performed to identify brain patterns of task and subject main effects (decoding and fingerprinting patterns, respectively) for the different measures of interest. Each ANOVA analysis delivered two F-values (one for the subject and one for the task effect) and associated p-value for each brain region or connection, and each measure. The edgewise patterns were summarized at the brain regions' level by taking the row-wise average of the F-value matrices. Fvalues were deemed significant for p < .05 accounting for Bonferroni correction across regions or connections. To assess the complementary contributions of SDI and FC nodal strength / c-FC and FC / d-FC and FC to task decoding and individual fingerprinting, additional three-factor ANOVA analyses were performed on concatenated regional SDI and FC strength / edgewise c-FC and FC / d-FC and FC values including subject, task, measure, and their first-order interactions as independent factors. The interaction terms [task*measure] and [subject*measure] indicate whether the effect of task or subject on brain patterns depends on the way such patterns are quantified; i.e., structure-function coupling or functional connectivity alone.

2.5. Task decoding

Prior to task classification, task paradigms were regressed out from functional time courses to minimize confounds from paradigm-imposed timings, aiming at keeping only differences due to the specific task-related states. Five SVM analyses with $N_{BS} = 8$ classes were performed to classify a brain state bs ($bs = 1, ..., N_{BS}$; i.e., resting state or one of the 7 tasks) based on the $N_{ROI} \times N_E \cdot N_{BS} \cdot N_S = 379 \times 1600$ nodal feature matrices of (1) FC nodal strength and (2) SDI patterns, as well as based on the ($N_{ROI} \cdot (N_{ROI} - 1)/2) \times N_E \cdot N_{BS} \cdot N_S = 71631 \times 1600$ edgewise feature matrices of (3) FC, (4) c-FC and (5) d-FC values, from all subjects and acquisitions. Two different cross-validation settings were explored: a 100-fold (*leave-one-subject-out*) cross-validation, where the $N_E \cdot N_{BS} = 16$ acquisitions from one subject were excluded for each training fold and used as test data; a 10-fold (leave-ten-subject-

out) cross-validation, where $N_E \cdot N_{BS} \cdot 10 = 160$ acquisitions from ten subjects were excluded for each training fold and used as test data. The 10-fold cross-validation was repeated 50 times with different trainingtest data partitions. As a sanity check, the 100-fold leave-one-subjectout cross-validation was repeated while randomizing the task labels. In all cross-validation settings and for each training-test loop, a *one-versusone* multiclass linear SVM classifier with error-correcting output codes modeling was trained on standardized training data (i.e., each predictor variable was centered and scaled to unit variance) using the *fitcecoc* MATLAB v.R2019b function and used to predict the task in the test data (Allwein et al., 2000; Furnkranz, 2002).

2.7. Individual fingerprinting

A second set of SVM classifications, with the same five sets of features (see paragraph 2.6), but with $N_S = 100$ classes, was performed to identify individuals based on their functional or structure-function coupling characteristics. Two different classification and cross-validation settings were explored, considering data obtained from matching or discordant tasks: (1) identification of a subject s doing a specific task bs, based on all other tasks and individuals. This was implemented with a 800-fold (leave-one-subject's-task-out) cross-validation, where the N_E entries (two different encoding directions) of subject s doing task bs were excluded for each fold; (2) identification of a subject s doing a specific task bs, from entries related to only one other different task (all pairwise combinations explored). This was implemented with a set of leave-onesubject-and-task-out cross-validation analyses on data subsets including only entries from a specific task and subject in the test fold, and only entries from a specific different task (all subjects, N_S data points) in the training fold. For each training-test loop, a one-versus-all multiclass linear SVM classifier with output codes modeling was trained on standardized training data and used to predict the subject in the test data (Allwein et al., 2000). We chose a linear SVM approach for its simplicity and ability to handle limited numbers of data points per class combined with large numbers of features. No hyperparameter tuning was involved in the classifiers' training. In addition, for comparison with recent work on functional connectivity data (Finn et al., 2015), fingerprinting accuracy was also quantified with the subject identification rate (with no SVM classification and cross-validation). The identification rate was computed as the rate of success in subject identification when matching each subject s doing a specific task bs with the most similar subject doing a different task (all task combinations explored). The similarity between subject pairs was quantified as the Pearson's correlation coefficient between their respective feature vectors (nodal FC strength; SDI; FC; c-FC; d-FC). As a sanity check, the subject identification rate was also computed while randomizing the subject labels.

2.8. Multivariate correlation with cognition

PLSC analyses (Krishnan et al., 2011) were performed to assess the presence of multivariate correlation patterns between the five sets of nodal and edgewise brain features and 10 cognitive scores across subjects. For the cognitive scores, the 10 cognitive subdomains tested in the HCP were considered, namely, episodic memory, executive functions, fluid intelligence, language, processing speed, selfregulation/impulsivity, spatial orientation, sustained visual attention, verbal episodic memory and working memory (Barch et al., 2013). For subdomains for which more than one unadjusted raw score was available, a single score was obtained by data projection onto the first component from a principal component analysis (Supplementary Fig. S1). For each brain feature, PLSC was repeated N_{BS} times, each time considering only brain values (FC nodal strength, SDI, FC, c-FC, or d-FC) obtained during one task. Given the dimensionality of the data, each PLSC analysis outputs 10 pairs of so-called brain-cognitive saliences corresponding to the left and right singular vectors of the data covariance matrix; 10 singular values indicating the amount of explained covariance; and 10 sets of brain and cognitive latent scores corresponding to data projections onto the brain and cognitive saliences, respectively. Statistical significance of multivariate correlation patterns was assessed with permutation testing (1000 permutations) (McIntosh and Lobaugh, 2004; Zöller et al., 2017). Reliability of nonzero salience values was assessed with bootstrapping procedure (1000 random samples) and computing standard scores with respect to the bootstrap distributions (salience values were considered reliable for absolute standard score > 3) (McIntosh and Lobaugh, 2004; Zöller et al., 2017). Moreover, the generalizability of the multivariate correlation patterns obtained with each PLSC analysis was assessed with a 10-fold crossvalidation procedure in the following way: first, 10 test subjects were removed from the dataset; second, brain and cognitive saliences were estimated from the remaining data (i.e., from the training set which includes $N_S - 10 = 90$ subjects); third, test-subject data were projected onto the saliences obtained from the training set to obtain the test latent scores; fourth, the correlation between original and test latent scores was evaluated. In case of generalizable multivariate correlation patterns, one would expect that original and test latent scores align along the identity line (Loukas et al., 2021). Finally, the r-squared (squared Pearson's correlation) between the latent scores was used to quantify the amounts of cognitive traits' variance explained by the five different brain features. For the edgewise brain features (FC, c-FC, d-FC), a cortical summary of the edgewise saliences was obtained by summing the salience weights of all the edges attached to the individual brain regions.

3. Results

3.1. Group-level structure-function coupling patterns are consistent across tasks

The structure-function coupling assessed with SDI at the nodal level, and with c-FC and d-FC at the edge level, yielded brain patterns of regional and edgewise values for each subject and run which were consistent across tasks (resting state and seven tasks: emotion, gambling, language, motor, relational, social, working memory; each acquired with 2 phase encoding directions). Average SDI, c-FC, and d-FC profiles across subjects for each state are reported in Supplementary Figs. S2 and S3. Consistently with previous work (Preti and Van De Ville, 2019), we observed relatively strong structure-function nodal coupling (lower SDI) in sensory and particularly in visual areas, and relatively strong nodal decoupling (higher SDI) in high-level cognitive networks (Supplementary Fig. S2). Functional connectivity information extracted from the low spatial frequencies of the structural connectome (c-FC) was qualitatively similar to classical functional connectivity (FC), with strong connectivity within visual and somatosensory networks, and low functional connectivity between the default mode (DMN) and limbic networks, and the other brain circuits. Conversely, functional connectivity matrices obtained from high spatial frequencies of the structural connectome (d-FC) were sparser and displayed both anti-correlation and positive-correlation patterns within and between resting state networks. Subcortical regions mainly showed d-FC anti-correlation patterns with cortical circuits (Supplementary Fig. S3).

3.2. Task decoding and fingerprinting patterns are spatially distinct

As a first step, we investigated the existence of possibly distinct brain patterns of structure-function coupling or functional connectivity associated with inter-task and inter-individual variability, respectively. To this end, a set of two-factor ANOVA analyses assessing differences of nodal and edgewise measures across subjects and tasks yielded two spatially distinct whole-brain patterns for each brain measure, characterized by a significant effect for either task or subject. These whole-brain patterns are represented in Fig. 2 for structure-function coupling (SDI, c-FC, d-FC) and in Supplementary Fig. S4 for functional connectivity



Fig. 2. Brain patterns of task decoding and subject fingerprinting effects on nodal Structural-Decoupling Index (SDI), edgewise coupled FC (c-FC), and edgewise decoupled FC (d-FC). F-values from two-factor ANOVA analyses are represented on a standard cortical surface independently from their statistical significance, with colormap scaled between the 5th and 95th percentiles across brain regions for each measure. For edgewise measures, the row-wise average of the F-value matrix is represented. The percentage of statistically significant brain regions or connections (p < .05, Bonferroni-corrected) is reported below each cortical plot. The average F-values across regions belonging to 8 resting state networks are represented in the bar plots, with bars colored according to the corresponding resting-state network. The vertical lines in the bar plots represent the whole-brain average F-values.

(FC nodal strength, FC). The ANOVA analyses revealed similar decoding and fingerprinting patterns for all measures except d-FC, although with some differences. Overall, the task decoding pattern (Fig. 2, left column) clearly involved more prominently regions belonging to unimodal brain circuits, in particular parts of the visual, somatomotor, and auditory networks. On the contrary, the fingerprinting pattern (Fig. 2, right column) was spatially more distributed, spreading across the frontal, parietal, occipital, and temporal lobes, including executive-control and transmodal regions (which have been consistently reported to contribute to subject identification from functional connectivity (Finn et al., 2015)) but not the limbic system and anterior DMN. Compared to SDI, FC nodal strength, and FC, the c-FC fingerprinting pattern showed lesser involvement of the primary and secondary visual cortices. These considerations are also reflected by the average F-value within resting state networks, depicted in the bar plots (Fig. 2) (Yeo et al., 2011). The decoding and fingerprinting patterns of d-FC were distinct from the other measures. Only 29% of brain connections showed a significant effect of the task on d-FC (p < .05, Bonferroni corrected for the number of connections), suggesting a weak d-FC variability across different tasks. Accordingly, the d-FC

decoding pattern was spatially more scattered and mainly involved the primary visual and the inferior parietal cortices. On the contrary, 86% of brain connections showed a significant effect of the subject on d-FC. The d-FC fingerprinting pattern was spatially more localized compared to the other measures and presented a striking resemblance to the frontoparietal executive-control network (Fig. 2). Finally, combined ANOVA analyses including both structure-function coupling and functional connectivity measures as dependent factors, and subject, task, measure, and first order interactions as explanatory factors, showed a significant combined effect of both task and measure (task-measure interaction), and subject and measure (subject-measure interaction) for all brain regions when concatenating SDI and FC nodal strength (p < .05, Bonferronicorrected for the number of brain regions), and for 71% (89%) brain connections for task-measure and 97% (94%) for subject-measure interactions when concatenating c-FC and FC (d-FC and FC) edge values (p < .05, Bonferroni-corrected for the number of brain connections).These interaction analyses indicate that structure-function coupling and functional connectivity measures provide complementary contributions to both task and subject identification.

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Table 1

Task decoding, subject fingerprinting, and brain-cognition relationships. First column: task decoding accuracies for nodewise (FC nodal strength; SDI) and edgewise (FC, c-FC, d-FC) functional and structure-function coupling measures estimated with 100-fold leave-one-subject-out crossvalidation and once-versus-one multiclass linear SVM classifiers. Second column: subject fingerprinting accuracies estimated with 800-fold leave-one-subject's-task-out cross-validation and oneversus-all multiclass SVM classifiers. Third column: brain-cognition r-squared (r^2) computed as the squared Pearson's correlation coefficient between the brain and cognition latent scores obtained from significant partial least squares correlation (PLSC) components. The brain-cognition r^2 quantifies the amount of inter-individual cognitive traits' variance explained by the five different brain features, respectively.

	Task Decoding accuracy	Subject Fingerprinting accuracy	Brain-Cognition r^2
FC nodal strength	0.544	0.984	0.211
nodal SDI	0.756	0.997	0.180
FC	0.919	0.964	0.224
c-FC	0.893	0.972	0.209
d-FC	0.873	1.000	0.654

3.3. Structure-function coupling is able to decode task-related brain states

SVM was used to classify different task-related states (resting state and seven tasks) based on nodewise or edgewise values of functional connectivity as well as structure-function coupling, where task paradigms were regressed out from functional time courses.

For the nodewise metrics, task-classification based on nodal structure-function coupling (SDI) reached an accuracy of 0.756 (chance level = 0.125; randomized data = 0.119, Supplementary Table S1), higher than the one based on FC nodal strength (0.544), showing that SDI is able to outperform a nodal measure (i.e., with equal dimensionality) based on functional data only (Table 1, first column: leaveone-subject-out accuracies; Supplementary Table S2: leave-ten-subjectout accuracies). When keeping the full dimensionality of connections (71,631 features), accuracy for GSP-derived FC values reached 0.893 for c-FC and 0.873 for d-FC, comparable to conventional FC (0.919), showing that structure-function dependencies, analogously to FC, are able to well characterize resting state and the different task conditions (Table 1, second column; Supplementary Table S2). Results were comparable when using a median split of the graph spectral density (rather than a fixed threshold c = 50) for the GSP-based measures' computation (Supplementary Table S3).

3.4. Structure-function decoupling represents an individual fingerprint of brain organization

In addition to characterizing different task-related states, structurefunction coupling measures also revealed to be highly specific to different individuals, which was also the case for functional connectivity. Accuracies for the identification of subjects ranged in fact from 0.964 for edgewise FC to about 1 for nodewise SDI and edgewise d-FC (chance level = 0.010) as assessed with 800-fold leave-one-subject's-task-out crossvalidation setting (Table 1). Both nodewise and edgewise structurefunction coupling measures performed slightly better than their counterparts based on functional connectivity alone (Table 1). Next, we attempted to identify individuals based on training the SVM classifier on only one task and testing it on another task (all task combinations explored; leave-one-subject-and-task-out cross validation). Our results show that even in this more challenging classification setting, subject identification was possible for all functional and structure-function coupling measures, with accuracies above chance level (Fig. 3). When considering nodewise measures, fingerprinting accuracies were higher for FC nodal strength compared to SDI in most task combinations (average accuracies = 0.806 / 0.663 for FC strength and SDI, respectively). However, in this same cross-validation scenario, the performance of edgewise metrics was particularly interesting to observe. The best (near-perfect) accuracies, in fact, were reached by the decoupled FC, largely outperforming both conventional FC and, in particular, coupled FC (average accuracies = 0.897 / 0.428 / 0.997 for FC, c-FC and d-FC, respectively). In general, predicting the subject from resting state data (training fold) to task data (test fold), and from any task to resting state data, was slightly more difficult than cross-task prediction, although there was not a particular pairwise task combination consistently outperforming the other task combinations (Fig. 3). Results were comparable when quantifying the fingerprinting accuracies with the identification rate (average identification rate = 0.683 / 0.431 / 0.739 / 0.250 / 0.985 for nodal FC strength, nodal SDI, FC, c-FC and d-FC, respectively) (Supplementary Fig. S5), with identification rates above values obtained from randomized data (0.011 / 0.011 / 0.010 / 0.011 / 0.010, respectively; Supplementary Table S4). Cross-task identification rates for edgewise functional connectivity were consistent with previous reports (minimum, maximum identification rate = 0.450, 0.990) (Amico and Goñi, 2018a; Finn et al., 2015).

3.5. Structure-function decoupling explains cognitive traits

Finally, functional and structure-function coupling measures explained inter-individual variations of cognitive traits, particularly sustained attention and fluid intelligence scores. Multivariate correlations between subject-specific brain measures (FC nodal strength, nodal SDI, FC, c-FC, and d-FC) in the different tasks (resting state and seven tasks) and 10 scores measuring cognitive subdomains were assessed with PLSC analyses (one PLSC per task and per brain measure). PLSC identifies linear combinations of brain measures that maximally covary with linear combinations of cognitive scores. PLSC analyses revealed significant multivariate correlation patterns between cognitive traits and all five functional and structure-function coupling measures mainly during resting state (p < .05; Supplementary Table S5). During tasks, braincognition multivariate correlations were not statistically significant or statistically significant but weaker compared to resting state, as indicated by lower brain-cognition r-squared values (Supplementary Table S5). When comparing the amount of inter-individual cognitive traits' variance explained by the five different brain features, we found that resting state FC nodal strength, nodal SDI, edgewise FC, and c-FC had similar r-squared values, ranging from 0.180 for SDI to 0.224 for FC (i.e., 18 to 22%, Table 1). However, edgewise d-FC explained a larger amount of inter-individual cognitive variance, reaching 65% (Table 1). In particular, stronger resting state d-FC in regions belonging to the frontoparietal network (including the bilateral posterior superior-frontal gyri, dorsolateral frontal cortices, intraparietal sulci, and inferior temporal gyri), and weaker resting state d-FC in somatosensory, limbic and middle temporal regions, were associated with better sustained attention performances, as shown by the d-FC and cognitive saliences that weigh the contribution of individual variables to the overall multivariate pattern (Fig. 4C). Conversely, larger resting state FC nodal strength, SDI, FC, and c-FC specifically related to a cognitive profile characterized by higher



Fig. 3. Cross-task fingerprinting accuracies for functional and structure-function coupling measures. Subject classification accuracies when using only one condition –task or resting state– for training (matrices' rows) and one for testing (matrices' columns), with all pairwise task combinations explored and for all nodewise (FC nodal strength; SDI) and edgewise (FC, c-FC, d-FC) measures. Classification accuracies were assessed with *leave-one-subject-and-task-out* cross-validation and *one-versus-all* multiclass SVM classifiers. RS=resting state; Emo=emotion; Gam=gambling; Lan=language; Mot=motor; Rel=relational; Soc=social; WM=working memory.

fluid intelligence and spatial orientation, and lower, sustained attention and verbal episodic memory scores (Figs. 4A, B and S6). The FC nodal strength, SDI, FC, and c-FC cortical patterns relating to cognition were spatially similar and mainly involved somatosensory, association, and temporo-parietal brain regions. 10-fold cross-validation analyses indicated good brain and cognitive patterns generalizability, with Pearson's correlation values between original and test latent scores ranging from 0.78 to 0.99 (Supplementary Fig. S7).

4. Discussion

Functional neuroimaging data have shown to provide measures of activity and connectivity with the ability to predict brain states in relation to task execution, as well as to identify individual subjects in a group (Finn et al., 2015; Haynes and Rees, 2006; Richiardi et al., 2011; Van De Ville et al., 2021). In parallel, brain morphology (Wachinger et al., 2015) and structural connectivity (Kumar et al., 2017; Yeh et al., 2016) revealed as well the capability of uniquely identifying individuals. However, brain function and structure are conventionally considered separately, and the relevance of structure-function coupling in underlying distinct brain states (tasks) and inter-individual variability (subjects' fingerprint) remains unexplored.

In relation to the first, the way brain function couples to the underlying structure is likely to adapt to the demands of the task. In line with this, task-related functional activity was shown to be well predicted from structure only in selected brain regions, different for each task (Wu et al., 2020). However, how this structure-function relationship depends on external stimulation, cognitive engagement, and affective state, and whether this can be useful to decode different brain states is still an open question (Suárez et al., 2020). Concerning individual fingerprinting, considering the value of both structural and functional brain features in subject identification, we could expect structure-function coupling profiles to also uniquely characterize individuals, providing a new dimension of inter-individual differences in brain organization. In line with this hypothesis, a recent study showed that the extent of alignment between structure and function correlates with individual differences in cognitive flexibility (Medaglia et al., 2018).

With these premises, we expanded here previous research by introducing new measures of structure-function coupling at the level of single brain connections (c-FC, d-FC) and by identifying the task decoding and individual fingerprinting relevance of such structure-function nodal and edgewise patterns. Our work shows that structure-function coupling can predict both brain states and individuals with high accuracy. The Structural-Decoupling Index and the functional connectivity component decoupled from structure (d-FC) revealed able to identify individual subjects in a group with near-perfect accuracy (Table 1), indicating that the pattern of structure-function coupling is an intrinsic feature (or fingerprint) of an individual's brain organization. The idea of a 'deep' functional fingerprint, independent from brain state configuration, is consistent with recent works reporting good cross-task subject identification from FC data (Abbas et al., 2020; Amico and Goñi, 2018b; Finn et al., 2015) and moderate state-dependency (compared to high subject-dependency) of functional networks (Gratton et al., 2018). Here, we demonstrate that the way brain function aligns (or misaligns) with the underlying structural connectivity provides additional clues on this functional fingerprint.

Therefore, while it is true that structure-function dependencies are sufficiently different across tasks to allow a reliable decoding of brain states, a strong structure-function individual fingerprint exists independently from the task during which brain function is measured. In fact, this fingerprint appears robust to brain state changes, since even a stringent cross-validation setting with pairwise cross-task predictions delivers high fingerprinting accuracies, in particular related to functional connectivity patterns decoupled from structure. This shows, notably, that a great deal of information specific to the individual is present in the high spatial frequencies of the structural decomposition, that is in the portion of functional signals which is more liberal with respect to brain structure. Notably, this finding could be particularly useful in the context of clinical studies (Itani and Thanou, 2021): the information contained in the high frequencies of the structural connectome -which is shown here to distinguish very well among individuals- could in this case represent features characterizing individual patients and reflecting their specific pathological traits. This consideration is reinforced by the PLSC results which indicate that the FC components decoupled from



Fig. 4. Multivariate correlation patterns between classical, coupled-, and decoupled functional connectivity during rest and cognitive traits. Significant partial least square correlation (PLSC) patterns between cognitive traits (first column) and resting state FC (A), c-FC (B), and d-FC (C) (second column). First column: bars and single dots represent the cognitive salience average and dispersion over 1000 bootstraps; yellow shading indicates salience weights significantly different from zero. Second column: brain saliences plotted on the cortical surface. For FC and c-FC (A, B) the significant salience weights were positive, as represented by the yellow-to-red colormap. For d-FC (C) the significant salience weights ranged from negative to positive values, as represented by the blue-to-red colormap.

structure explain a significant percentage of inter-individual cognitive traits' variability, at a level that exceeds the performances of other functional and structure-function coupling measures.

As mentioned above, structure-function dependencies also deliver high accuracy (0.756 for SDI, 0.893 for c-FC, and 0.874 for d-FC, against a 0.125 chance-level) when decoding task-related states. It is important to remark here that, having regressed out task paradigms, task decoding can still detect differences due to task, but not "artificially" induced ones, dependent on the paradigm timing, which prevents biases due to task particularities. Recent studies have shown that the cortical macroscale gradient of structure-function coupling found at rest, opposing primary sensory and association cortices (Preti and Van De Ville, 2019; Vázquez-Rodríguez et al., 2019), can be retrieved from task data as well (Baum et al., 2020; Wu et al., 2020), suggesting similar coupling patterns both in intrinsic (rest) and extrinsic brain states. We can indeed observe the same, when comparing average structure-function coupling patterns among task conditions (average SDI, c-FC, and d-FC maps across subjects in Supplementary Figs. S2 and S3). Nonetheless, specific and non-trivial differences across tasks, not clearly visible at the population level, exist and allow accurate task decoding.

Contributions of brain regions to task and subject identification are in fact not uniformly distributed across the cortex: two clearly distinct patterns were highlighted, one for task decoding and one for individual fingerprinting (see Fig. 2). Interestingly, these two maps group brain regions with distinct structure-function coupling properties. The task pattern mainly involved lower-order regions whose functional activity significantly couples with the structural connectome (Supplementary Fig. S2), including somatomotor, visual and auditory cortices (Preti and Van De Ville, 2019). This means that between-task variations of structurefunction coupling as well as of functional connectivity (Supplementary Fig. S4) mainly occur in regions whose functional activity is on average more constrained by the underlying structure. Conversely, the fingerprinting pattern was spatially more spread and extended to frontal, parietal, occipital, and temporal regions including executive-control and transmodal association cortices whose functional activity tends to decouple from structure (Preti and Van De Ville, 2019), but excluding the limbic system. The latter includes several allocortical regions with simpler cortical layers' organization and early neurodevelopment (Baum et al., 2020), which together may explain the weak subject specificity. Moreover, the structure-function coupling in insulo-limbic regions particularly relates to cortical chemo-architecture and inter-regional similarity of neurotransmitter receptors' concentration (Hansen et al., 2021), which could reduce the degrees of freedom 'available' for interindividual variability. Conversely, transmodal and higher-order cortices present higher cytoarchitectonic complexity and less recurrent connections (Wang et al., 2019). Computational studies suggest that these features could underlie the capacity of higher-order brain regions to support autonomous and spontaneous dynamics more liberal with respect to the structural connectivity architecture (Vázquez-Rodríguez et al., 2019; Wang et al., 2019). On the same line, post mortem histological data and in vivo MRI assessment of cortical myelo-architecture indicate a divergence between microstructural and functional cortical gradients in the executive-control network, suggesting that these fronto-parietal areas may be relatively untethered not only from cortico-cortical whitematter connectivity but also from intra-cortical hierarchical constraints (Paquola et al., 2019). These mechanisms may as well relate to the stronger individual specificity of decoupled functional connectivity patterns observed particularly in executive-control regions.

Indeed, The decoupled component of the functional connectivity was characterized by distinct task- and subject-specific brain patterns compared to the other structure-function coupling and functional connectivity measures. On the one side, d-FC presented limited task specificity, with only 29% of brain connections being associated with a significant task-effect in the two-factor ANOVA analyses and mainly involving inferior parietal and primary visual areas. We speculate that different graph-frequency components may be involved in different tasks, with an omnipresent contribution from lower-order areas (tendentially coupled with structure (Preti and Van De Ville, 2019)), so that multiple graph frequencies and particularly lower frequencies are needed to achieve optimal task differentiation. On the other side, d-FC demonstrated nearperfect subject identification accuracy even in a challenging cross-task classification setting (Fig. 3). Compared to the other measures, the d-FC fingerprinting spatial pattern was more localized and strikingly overlapped with the fronto-parietal executive-control system, in agreement with previous reports on vertex-level inter-subject variability of intrinsic functional connectivity in these regions (Mueller et al., 2013). In parallel, primary sensory networks had a much lower fingerprinting power in d-FC. These are the ones that are more related to the underlying struc-

tural connectivity (Liégeois et al., 2020; Preti and Van De Ville, 2019), so that 'removing' its contribution through graph-domain filtering decreases the relative importance of these networks. The differences observed between SDI, c-FC, and d-FC hint at a neurobiological relevance of the way brain activity (tightly or loosely) couples with the anatomical connectivity substrate, both in regard to the mechanisms underlying brain state reconfiguration across tasks, and to how individual uniqueness is expressed in the brain. In addition, joint analyses of structurefunction coupling and functional connectivity (three-factor ANOVAs) indicated that the two classes of brain features provide complementary contributions to task and subject identification. These results suggest that the alignment of function with structure reveals additional information with respect to the functional connectivity alone. Future work should be done to consolidate and extend these considerations, for example by including subject-specific structural connectivity information -a non-trivial operation that would lead to the definition of multiple spectral domains for brain signals, but opens the perspective of incorporating inter-subject structural variability in the analysis of functional brain signatures.

Differently from previous work that mainly focused on fingerprint patterns and single cognitive domains such as fluid intelligence, here we explored multivariate correlations between functional and structurefunction coupling features, and multiple cognitive traits. We show that inter-individual variations of (nodewise and edgewise) functional connectivity and local structure-function coupling (SDI) during rest consistently explain traits of complex cognition (fluid intelligence, spatial orientation), executive function (sustained attention) and episodic memory (Moore et al., 2015), resembling descriptions of a general intelligence gfactor previously associated with functional connectivity of the default mode network (Smith et al., 2015b). In particular, a relatively stronger nodal structure-function coupling (lower SDI) was associated with better complex cognition, in line with previous work demonstrating a link between less liberal structure-function alignment during task switching and concomitant cognitive flexibility performances (Medaglia et al., 2018). Nonetheless, relatively weaker nodal structure-function coupling was associated with better executive and memory abilities. It might be that certain brain functions, such as complex reasoning, may benefit from more reliable and consolidated brain communication pathways, possibly expressed in a stronger structure-function alignment (Finn et al., 2017; Medaglia et al., 2018; Suárez et al., 2020). Other functions, such as verbal learning and retrieval or attention maintenance, may conversely benefit from a less constrained structure-function alignment, a configuration that might predispose the individual to the integration of new information. On this line, functional connectivity components decoupled from structure (d-FC) were strongly associated with sustained attention, a mental function that plays a ubiquitous role across cognitive domains and perception. The d-FC cortical pattern was spatially spread, with large (absolute) PLS weights spanning all brain lobes. This widespread pattern echoes a large-scale network spanning several cortical, subcortical, and cerebellar regions identified as robust predictors of attention performance (Rosenberg et al., 2016). Yet, the pattern of increased d-FC (positive PLS weights) strongly overlapped with the fronto-parietal network, largely implicated in the selection of sensory contents by attention (Ptak, 2012). The pattern of coupled functional connectivity (c-FC) associated with cognition mainly opposed the fronto-parietal network and posterior DMN to sensorimotor, visual, and anterior cingulate regions. This pattern, associated with complex functions, partially mirrored previously reported macroscale cortical gradients that integrate information across multiple sensory domains into abstract representation, in virtue of a progressively increasing distance from somatosensory areas (Buckner and Krienen, 2013; Margulies et al., 2016). While speculative, these considerations and research in this direction, particularly investigating the role of the medium and high frequencies of the structural connectome, may offer a new understanding of cognitive control mechanisms (Lerman-Sinkoff et al., 2017). Furthermore, in our analyses the relationship between brain features and cognitive traits was predominant in the resting condition, suggesting that intrinsic rather than extrinsic brain states might better reflect general cognitive abilities. Meanwhile, this observation does not exclude that temporal fluctuations of structure-function coupling levels during tasks or rest might tap into specific cognitive-behavioral subdomains and hence improve the prediction of task performance or cognitive traits, which is another avenue for future research (Van De Ville et al., 2021).

Finally, the spatial patterns of structure-function coupling relating to cognition presented similarities both with the task decoding maps in lower-order somatomotor and association cortices, intrinsically characterized by stronger structure-function coupling, as well as with the fingerprinting maps in fronto-parietal regions, characterized by more liberal structure-function coupling (Preti and Van De Ville, 2019) (Figs. 2, 4, and S2). Recent work showed that structural and functional connectivity present distinct patterns of inter-individual variance as they relate to cognition (Rasero et al., 2021; Zimmermann et al., 2018). Intriguingly, our results extend these findings identifying in the structurefunction coupling a possible link between divergent structural and functional connectivity patterns in predicting behavior. In this respect, both the nodewise SDI and the edgewise d-FC capture inter-subject cognitive variability, but along two different axes. Compared to the functional connectivity component decoupled from structure, the coupled component (c-FC) preserves task- and subject-specific information, but to a lesser extent, showing lower fingerprinting accuracies in the crosstask classification setting and weaker brain-cognition relationship. The coupled functional connectivity component may contain large-scale patterns common to individuals in a group, as suggested by its similarity with the classic functional connectivity organization into wellestablished resting state networks (Supplementary Fig. S3), while the decoupled component may contain a larger proportion of subject-specific information. A further exploration of the full structural connectome spectrum and of its derived functional connectivity components is warranted.

This study has a number of limitations and possible developments. First, a relatively restricted HCP sample was used. This choice was motivated by the computational burden of the fingerprinting multi-class SVM classifiers' training. Although recent work indicates reliable FC fingerprinting performance across different subgroups of HCP subjects (Waller et al., 2017), future work should better assess the dependency of accuracy estimates and brain-cognition correlations on more noisy data acquired in routine and clinical settings (Grady et al., 2021). Second, the usage of a gray matter parcellation as opposed to a voxel-based analysis impedes a fine-grained characterization of functional territories that can vary across subjects and tasks (Laumann et al., 2015; Salehi et al., 2020; Wang et al., 2015) with possible impact on the quantification of nodewise and edgewise structure-function coupling features. Nevertheless, a parcellation-based approach facilitates inter-subject comparisons, improves the signal-to-noise ratio of the estimated structural and functional measures, and enables a compact representation of brain fingerprints and decoding patterns. Third, group-level structural connectivity information was used for the computation of GSP-derived metrics. While this choice is convenient since it defines a common spectral domain across subjects and tasks, ways to integrate inter-subject structural variability could be explored in the future. Fourth, this study does not consider time-varying aspects of structure-function dependency (Cabral et al., 2017; Fukushima et al., 2018; Van De Ville et al., 2021): their exploration in the future might provide insight particularly in relation to task decoding and cognitive control mechanisms. Fifth, our analyses are limited to slow temporal scales accessible with fMRI. Previous studies had attempted brain fingerprinting using electrophysiological recordings (Fraschini et al., 2015; Marcel and Millan, 2007; Sareen et al., 2021), but the link between faster brain dynamics and structural topology remains poorly understood (Finger et al., 2016; Glomb et al., 2020). Future research may address how the hierarchy of structure-function dependencies vary at faster temporal scales, possibly carrying distinct fingerprinting and decoding information. Finally, our multivariate correlation analyses explore possible brain patterns relating to cognitive traits, including bootstrap and cross-validation procedure for generalizability assessment. Nonetheless, feature importance in multivariate predictive models of cognition remains difficult to reliably estimate and different machine learning approaches are under investigation (Tian and Zalesky, 2021).

In conclusion, this work demonstrates that structure-function dependencies quantified both at the level of single brain regions and connections form prominent signatures of individual brains' organization reflecting cognitive and behavioral correlates, while at the same time preserving task-dependent information. In particular, the high spatial frequencies of the structural connectome may contain relevant subjectspecific information which deserves further attention in the future.

Declaration of Competing Interest

The authors declare no conflict of interest.

Credit authorship contribution statement

Alessandra Griffa: Conceptualization, Formal analysis, Visualization, Writing – original draft, Writing – review & editing. Enrico Amico: Data curation, Writing – review & editing. Raphaël Liégeois: Data curation, Writing – review & editing. Dimitri Van De Ville: Conceptualization, Supervision, Writing – review & editing. Maria Giulia Preti: Conceptualization, Formal analysis, Visualization, Writing – original draft, Writing – review & editing, Supervision, Data curation.

Data and code availability statement

All data used in this study are available through the Human Connectome Project, WU-Minn Consortium. The code to implement the analyses performed in this study is available at https://github.com/agriffa/GSP_brain_decode_fingerprint.git.

Acknowledgements

Data were provided by the Human Connectome Project, WU-Minn Consortium (Principal Investigators: David Van Essen and Kamil Ugurbil; 1U54MH091657) funded by the 16 NIH Institutes and Centers that support the NIH Blueprint for Neuroscience Research; and by the Mc-Donnell Center for Systems Neuroscience at Washington University. AG was supported by the Swiss National Science Foundation (Grant No. 320030_173153). MGP was supported by the CIBM Center for Biomedical Imaging, a Swiss research center of excellence founded and supported by Lausanne University Hospital (CHUV), University of Lausanne (UNIL), Ecole polytechnique fédérale de Lausanne (EPFL), University of Geneva (UNIGE) and Geneva University Hospitals (HUG). EA acknowledges financial support from the SNSF Ambizione project "Fingerprinting the brain: network science to extract features of cognition, behavior and dysfunction" (Grant No. PZ00P2_185716). RL was supported by the National centre of Competence in Research - Evolving Language (Grant No. 51NF40_180888).

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.neuroimage.2022.118970.

References

Abbas, K., Amico, E., Svaldi, D.O., Tipnis, U., Duong-Tran, D.A., Liu, M., Rajapandian, M., Harezlak, J., Ances, B.M., Goñi, J., 2020. GEFF: graph embedding for functional fin-

gerprinting. Neuroimage 221, 117181. doi:10.1016/j.neuroimage.2020.117181.
Allwein, E.L., Schapire, R.E., Singer, Y., 2000. Reducing multiclass to binary: a unifying approach for margin classifiers. J. Mach. Learn. Res. 1, 113–141.

- Amico, E., Abbas, K., Duong-Tran, D.A., Tipnis, U., Rajapandian, M., Chumin, E., Ventresca, M., Harezlak, J., Goñi, J., 2021. Toward an information theoretical description of communication in brain networks. Netw. Neurosci. 1–20. doi:10.1162/netn_a_00185.
- Amico, E., Goñi, J., 2018a. The quest for identifiability in human functional connectomes. Sci. Rep. 8, 1–14. doi:10.1038/s41598-018-25089-1.
- Amico, E., Goñi, J., 2018b. Mapping hybrid functional-structural connectivity traits in the human connectome. Netw. Neurosci. 2, 306–322. doi:10.1162/netn_a_00049.
- Avena-Koenigsberger, A., Misic, B., Sporns, O., 2018. Communication dynamics in complex brain networks. Nat. Rev. Neurosci. 19, 17–33. doi:10.1038/nrn.2017.149.
- Barch, D.M., Burgess, G.C., Harms, M.P., Petersen, S.E., Schlaggar, B.L., Corbetta, M., Glasser, M.F., Curtiss, S., Dixit, S., Feldt, C., Nolan, D., Bryant, E., Hartley, T., Footer, O., Bjork, J.M., Poldrack, R., Smith, S., Johansen-Berg, H., Snyder, A.Z., Van Essen, D.C., 2013. Function in the human connectome: task-fMRI and individual differences in behavior. NeuroImage 80, 169–189. doi:10.1016/j.neuroimage.2013.05.033, Mapping the Connectome.
- Baum, G.L., Cui, Z., Roalf, D.R., Ciric, R., Betzel, R.F., Larsen, B., Cieslak, M., Cook, P.A., Xia, C.H., Moore, T.M., Ruparel, K., Oathes, D.J., Alexander-Bloch, A.F., Shinohara, R.T., Raznahan, A., Gur, R.E., Gur, R.C., Bassett, D.S., Satterthwaite, T.D., 2020. Development of structure-function coupling in human brain networks during youth. Proc. Natl. Acad. Sci. 117, 771–778. doi:10.1073/pnas.1912034117.
- Biazoli, C.E.J., Salum, G.A., Pan, P.M., Zugman, A., Amaro, E.J., Rohde, L.A., Miguel, E.C., Jackowski, A.P., Bressan, R.A., Sato, J.R., 2017. Commentary: functional connectome fingerprint: identifying individuals using patterns of brain connectivity. Front. Hum. Neurosci. 11. doi:10.3389/fnhum.2017.00047.
- Buckner, R.L., Krienen, F.M., 2013. The evolution of distributed association networks in the human brain. Trends Cogn. Sci. 17, 648–665. doi:10.1016/j.tics.2013.09.017, Special Issue: The Connectome.
- Cabral, J., Kringelbach, M.L., Deco, G., 2017. Functional connectivity dynamically evolves on multiple time-scales over a static structural connectome: models and mechanisms. NeuroImage 160, 84–96. doi:10.1016/j.neuroimage.2017.03.045, Functional Architecture of the Brain.
- Deco, G., Jirsa, V.K., McIntosh, A.R., 2011. Emerging concepts for the dynamical organization of resting-state activity in the brain. Nat. Rev. Neurosci. 12, 43–56. doi:10.1038/nrn2961.
- Finger, H., Bönstrup, M., Cheng, B., Messé, A., Hilgetag, C., Thomalla, G., Gerloff, C., König, P., 2016. Modeling of large-scale functional brain networks based on structural connectivity from DTI: comparison with EEG derived phase coupling networks and evaluation of alternative methods along the modeling path. PLOS Comput. Biol. 12, e1005025. doi:10.1371/journal.pcbi.1005025.
- Finn, E.S., Scheinost, D., Finn, D.M., Shen, X., Papademetris, X., Constable, R.T., 2017. Can brain state be manipulated to emphasize individual differences in functional connectivity? NeuroImage 160, 140–151. doi:10.1016/j.neuroimage.2017.03.064, Functional Architecture of the Brain.
- Finn, E.S., Shen, X., Scheinost, D., Rosenberg, M.D., Huang, J., Chun, M.M., Papademetris, X., Constable, R.T., 2015. Functional connectome fingerprinting: identifying individuals using patterns of brain connectivity. Nat. Neurosci. 18, 1664–1671. doi:10.1038/nn.4135.
- Fischl, B., Salat, D.H., Busa, E., Albert, M., Dieterich, M., Haselgrove, C., van der Kouwe, A., Killiany, R., Kennedy, D., Klaveness, S., Montillo, A., Makris, N., Rosen, B., Dale, A.M., 2002. Whole brain segmentation: automated labeling of neuroanatomical structures in the human brain. Neuron 33, 341–355. doi:10.1016/S0896-6273(02)00569-X.
- Fraschini, M., Hillebrand, A., Demuru, M., Didaci, L., Marcialis, G.L., 2015. An EEG-based biometric system using eigenvector centrality in resting state brain networks. IEEE Signal Process. Lett. 22, 666–670. doi:10.1109/LSP.2014.2367091.
- Fukushima, M., Betzel, R.F., He, Y., van den Heuvel, M.P., Zuo, X.N., Sporns, O., 2018. Structure–function relationships during segregated and integrated network states of human brain functional connectivity. Brain Struct. Funct. 223, 1091–1106. doi:10.1007/s00429-017-1539-3.
- Furnkranz, J., 2002. Round robin classification. J. Mach. Learn. Res. 2, 721-747.
- Gao, Y., Zhang, Y., Cao, Z., Guo, X., Zhang, J., 2020. Decoding brain states from fMRI signals by using unsupervised domain adaptation. IEEE J. Biomed. Health Inform. 24, 1677–1685. doi:10.1109/JBHI.2019.2940695.
- Glasser, M.F., Coalson, T.S., Robinson, E.C., Hacker, C.D., Harwell, J., Yacoub, E., Ugurbil, K., Andersson, J., Beckmann, C.F., Jenkinson, M., Smith, S.M., Van Essen, D.C., 2016. A multi-modal parcellation of human cerebral cortex. Nature 536, 171–178. doi:10.1038/nature18933.
- Glasser, M.F., Sotiropoulos, S.N., Wilson, J.A., Coalson, T.S., Fischl, B., Andersson, J.L., Xu, J., Jbabdi, S., Webster, M., Polimeni, J.R., Van Essen, D.C., Jenkinson, M., 2013. The minimal preprocessing pipelines for the human connectome project. NeuroImage 80, 105–124. doi:10.1016/j.neuroimage.2013.04.127, Mapping the Connectome.
- Glomb, K., Rué Queralt, J., Pascucci, D., Defferrard, M., Tourbier, S., Carboni, M., Rubega, M., Vulliémoz, S., Plomp, G., Hagmann, P., 2020. Connectome spectral analysis to track EEG task dynamics on a subsecond scale. Neuroimage 221, 117137. doi:10.1016/j.neuroimage.2020.117137.
- Goñi, J., Van Den Heuvel, M.P., Avena-Koenigsberger, A., De Mendizabal, N.V., Betzel, R.F., Griffa, A., Hagmann, P., Corominas-Murtra, B., Thiran, J.P., Sporns, O., 2014. Resting-brain functional connectivity predicted by analytic measures of network communication. Proc. Natl. Acad. Sci. 111, 833–838. doi:10.1073/pnas.1315529111.
- Grady, C.L., Rieck, J.R., Nichol, D., Rodrigue, K.M., Kennedy, K.M., 2021. Influence of sample size and analytic approach on stability and interpretation of brainbehavior correlations in task-related fMRI data. Hum. Brain Mapp. 42, 204–219. doi:10.1002/hbm.25217.
- Gratton, C., Laumann, T.O., Nielsen, A.N., Greene, D.J., Gordon, E.M., Gilmore, A.W., Nelson, S.M., Coalson, R.S., Snyder, A.Z., Schlaggar, B.L., Dosenbach, N.U.F., Pe-

tersen, S.E., 2018. Functional brain networks are dominated by stable group and individual factors, not cognitive or daily variation. Neuron 98, 439–452. doi:10.1016/j.neuron.2018.03.035, e5.

- Griffa, A., Ricaud, B., Benzi, K., Bresson, X., Daducci, A., Vandergheynst, P., Thiran, J.P., Hagmann, P., 2017. Transient networks of spatio-temporal connectivity map communication pathways in brain functional systems. Neuroimage 155, 490– 502. doi:10.1016/j.neuroimage.2017.04.015.
- Hansen, J.Y., Shafiei, G., Markello, R.D., Smart, K., Cox, S.M.L., Wu, Y., Gallezot, J.D., Aumont, É., Servaes, S., Scala, S.G., DuBois, J.M., Wainstein, G., Bezgin, G., Funck, T., Schmitz, T.W., Spreng, R.N., Soucy, J.P., Baillet, S., Guimond, S., Hietala, J., Bédard, M.A., Leyton, M., Kobayashi, E., Rosa-Neto, P., Palomero-Gallagher, N., Shine, J.M., Carson, R.E., Tuominen, L., Dagher, A., Misic, B., 2021. Mapping neurotransmitter systems to the structural and functional organization of the human neocortex. (preprint). Neuroscience doi:10.1101/2021.10.28.466336.
- Haynes, J.D., Rees, G., 2006. Decoding mental states from brain activity in humans. Nat. Rev. Neurosci. 7, 523–534. doi:10.1038/nrn1931.
- Honey, C.J., Sporns, O., Cammoun, L., Gigandet, X., Thiran, J.P., Meuli, R., Hagmann, P., 2009. Predicting human resting-state functional connectivity from structural connectivity. Proc. Natl. Acad. Sci. 106, 2035–2040. doi:10.1073/pnas.0811168106.
- Huang, W., Bolton, T.A.W., Medaglia, J.D., Bassett, D.S., Ribeiro, A., Ville, D.V.D., 2018. A graph signal processing perspective on functional brain imaging. Proc. IEEE 106, 868–885. doi:10.1109/JPROC.2018.2798928.
- Itani, S., Thanou, D., 2021. Combining anatomical and functional networks for neuropathology identification: a case study on autism spectrum disorder. Med. Image Anal. 69, 101986. doi:10.1016/j.media.2021.101986.
- Krishnan, A., Williams, L.J., McIntosh, A.R., Abdi, H., 2011. Partial least squares (PLS) methods for neuroimaging: a tutorial and review. NeuroImage 56, 455–475. doi:10.1016/j.neuroimage.2010.07.034, Multivariate Decoding and Brain Reading.
- Kumar, K., Desrosiers, C., Siddiqi, K., Colliot, O., Toews, M., 2017. Fiberprint: a subject fingerprint based on sparse code pooling for white matter fiber analysis. Neuroimage 158, 242–259. doi:10.1016/j.neuroimage.2017.06.083.
- Laumann, T.O., Gordon, E.M., Adeyemo, B., Snyder, A.Z., Joo, S.J., Chen, M.Y., Gilmore, A.W., McDermott, K.B., Nelson, S.M., Dosenbach, N.U.F., Schlaggar, B.L., Mumford, J.A., Poldrack, R.A., Petersen, S.E., 2015. Functional system and areal organization of a highly sampled individual human brain. Neuron 87, 657–670. doi:10.1016/j.neuron.2015.06.037.
- Lerman-Sinkoff, D.B., Sui, J., Rachakonda, S., Kandala, S., Calhoun, V.D., Barch, D.M., 2017. Multimodal neural correlates of cognitive control in the human connectome project. Neuroimage 163, 41–54. doi:10.1016/j.neuroimage.2017.08.081.
- Li, H., Fan, Y., 2019. Interpretable, highly accurate brain decoding of subtly distinct brain states from functional MRI using intrinsic functional networks and long short-term memory recurrent neural networks. Neuroimage 202, 116059. doi:10.1016/j.neuroimage.2019.116059.
- Liégeois, R., Santos, A., Matta, V., Van De Ville, D., Sayed, A.H., 2020. Revisiting correlation-based functional connectivity and its relationship with structural connectivity. Netw. Neurosci. 4, 1235–1251. doi:10.1162/netn.a.00166.
- Lin, Y.C., Baete, S.H., Wang, X., Boada, F.E., 2020. Mapping brain-behavior networks using functional and structural connectome fingerprinting in the HCP dataset. Brain Behav. e01647. doi:10.1002/brb3.1647, n/a.
- Loukas, S., Lordier, L., Meskaldji, D.-.E., Filippa, M., Sa de Almeida, J., Van De Ville, D., Hüppi, P.S., 2021. Musical memories in newborns: a resting-state functional connectivity study. Hum. Brain Mapp. doi:10.1002/hbm.25677, in press.
- Mansour, L.S., Tian, Y., Yeo, B.T.T., Cropley, V., Zalesky, A., 2021. High-resolution connectomic fingerprints: mapping neural identity and behavior. Neuroimage 229, 117695. doi:10.1016/j.neuroimage.2020.117695.
- Marcel, S., Millan, J.D.R., 2007. Person authentication using brainwaves (EEG) and maximum a posteriori model adaptation. IEEE Trans. Pattern Anal. Mach. Intell. 29, 743– 752. doi:10.1109/TPAMI.2007.1012.
- Margulies, D.S., Ghosh, S.S., Goulas, A., Falkiewicz, M., Huntenburg, J.M., Langs, G., Bezgin, G., Eickhoff, S.B., Castellanos, F.X., Petrides, M., Jefferies, E., Smallwood, J., 2016. Situating the default-mode network along a principal gradient of macroscale cortical organization. Proc. Natl. Acad. Sci. 113, 12574–12579. doi:10.1073/pnas.1608282113.
- McIntosh, A.R., Lobaugh, N.J., 2004. Partial least squares analysis of neuroimaging data: applications and advances. NeuroImage 23, S250–S263. doi:10.1016/j.neuroimage.2004.07.020, Mathematics in Brain Imaging.
- Medaglia, J.D., Huang, W., Karuza, E.A., Thompson-Schill, S.L., Ribeiro, A., Bassett, D.S., 2018. Functional alignment with anatomical networks is associated with cognitive flexibility. Nat. Hum. Behav. 2, 156–164.
- Mišić, B., Betzel, R.F., de Reus, M.A., van den Heuvel, M.P., Berman, M.G., McIntosh, A.R., Sporns, O., 2016. Network-level structure-function relationships in human neocortex. Cereb. Cortex 26, 3285–3296. doi:10.1093/cercor/bhw089.
- Mišić, B., Betzel, R.F., Nematzadeh, A., Goñi, J., Griffa, A., Hagmann, P., Flammini, A., Ahn, Y.Y., Sporns, O., 2015. Cooperative and competitive spreading dynamics on the human connectome. Neuron 86, 1518–1529. doi:10.1016/j.neuron.2015.05.035.
- Moore, T.M., Reise, S.P., Gur, R.E., Hakonarson, H., Gur, R.C., 2015. Psychometric properties of the Penn computerized neurocognitive battery. Neuropsychology 29, 235–246. doi:10.1037/neu0000093.
- Mueller, S., Wang, D., Fox, M.D., Yeo, B.T.T., Sepulcre, J., Sabuncu, M.R., Shafee, R., Lu, J., Liu, H., 2013. Individual variability in functional connectivity architecture of the human brain. Neuron 77, 586–595. doi:10.1016/j.neuron.2012.12.028.
- Paquola, C., Wael, R.V.D., Wagstyl, K., Bethlehem, R.A.I., Hong, S.J., Seidlitz, J., Bullmore, E.T., Evans, A.C., Misic, B., Margulies, D.S., Smallwood, J., Bernhardt, B.C., 2019. Microstructural and functional gradients are increasingly dissociated in transmodal cortices. PLOS Biol. 17, e3000284. doi:10.1371/journal.pbio.3000284.

- Preti, M.G., Van De Ville, D., 2019. Decoupling of brain function from structure reveals regional behavioral specialization in humans. Nat. Commun. 10, 4747. doi:10.1038/s41467-019-12765-7.
- Ptak, R., 2012. The frontoparietal attention network of the human brain: action, saliency, and a priority map of the environment. Neuroscientist 18, 502–515. doi:10.1177/1073858411409051.
- Rasero, J., Sentis, A.I., Yeh, F.C., Verstynen, T., 2021. Integrating across neuroimaging modalities boosts prediction accuracy of cognitive ability. PLOS Comput. Biol. 17, e1008347. doi:10.1371/journal.pcbi.1008347.
- Richiardi, J., Eryilmaz, H., Schwartz, S., Vuilleumier, P., Van De Ville, D., 2011. Decoding brain states from fMRI connectivity graphs. NeuroImage 56, 616–626. doi:10.1016/j.neuroimage.2010.05.081, Multivariate Decoding and Brain Reading.
- Rosenberg, M.D., Finn, E.S., Scheinost, D., Papademetris, X., Shen, X., Constable, R.T., Chun, M.M., 2016. A neuromarker of sustained attention from whole-brain functional connectivity. Nat. Neurosci. 19, 165–171. doi:10.1038/nn.4179.
- Salehi, M., Greene, A.S., Karbasi, A., Shen, X., Scheinost, D., Constable, R.T., 2020. There is no single functional atlas even for a single individual: functional parcel definitions change with task. Neuroimage 208, 116366. doi:10.1016/j.meuroimage.2019.116366.
- Sareen, E., Zahar, S., Ville, D.V.D., Gupta, A., Griffa, A., Amico, E., 2021. Exploring MEG brain fingerprints: evaluation, pitfalls, and interpretations. Neuroimage 240, 118331. doi:10.1016/j.neuroimage.2021.118331.
- Seguin, C., Tian, Y., Zalesky, A., 2020. Network communication models improve the behavioral and functional predictive utility of the human structural connectome. Netw. Neurosci. 4, 980–1006. doi:10.1162/netn_a_00161.
- Smith, R.E., Tournier, J.D., Calamante, F., Connelly, A., 2015a. SIFT2: enabling dense quantitative assessment of brain white matter connectivity using streamlines tractography. Neuroimage 119, 338–351. doi:10.1016/j.neuroimage.2015.06.092.
- Smith, S.M., Nichols, T.E., Vidaurre, D., Winkler, A.M., Behrens, T.E.J., Glasser, M.F., Ugurbil, K., Barch, D.M., Van Essen, D.C., Miller, K.L., 2015b. A positive-negative mode of population covariation links brain connectivity, demographics and behavior. Nat. Neurosci. 18, 1565–1567. doi:10.1038/nn.4125.
- Suárez, L.E., Markello, R.D., Betzel, R.F., Misic, B., 2020. Linking structure and function in macroscale brain networks. Trends Cogn. Sci. 24, 302–315. doi:10.1016/j.tics.2020.01.008.
- Tian, Y., Zalesky, A., 2021. Machine learning prediction of cognition from functional connectivity: are feature weights reliable? Neuroimage 245, 118648. doi:10.1016/j.neuroimage.2021.118648.
- Valizadeh, S.A., Liem, F., Mérillat, S., Hänggi, J., Jäncke, L., 2018. Identification of individual subjects on the basis of their brain anatomical features. Sci. Rep. 8, 5611. doi:10.1038/s41598-018-23696-6.
- Van De Ville, D., Farouj, Y., Preti, M.G., Liégeois, R., Amico, E., 2021. When makes you unique: temporality of the human brain fingerprint. Sci. Adv. 7. doi:10.1126/sciadv.abj0751, eabj0751.
- Van Essen, D.C., Smith, S.M., Barch, D.M., Behrens, T.E.J., Yacoub, E., Ugurbil, K., 2013. The WU-Minn human connectome project: an overview. NeuroImage 80, 62–79. doi:10.1016/j.neuroimage.2013.05.041, Mapping the Connectome.

- Vázquez-Rodríguez, B., Suárez, L.E., Markello, R.D., Shafiei, G., Paquola, C., Hagmann, P., Van Den Heuvel, M.P., Bernhardt, B.C., Spreng, R.N., Misic, B., 2019. Gradients of structure–function tethering across neocortex. Proc. Natl. Acad. Sci. 116, 21219– 21227. doi:10.1073/pnas.1903403116.
- Wachinger, C., Golland, P., Kremen, W., Fischl, B., Reuter, M., 2015. BrainPrint: a discriminative characterization of brain morphology. Neuroimage 109, 232–248. doi:10.1016/j.neuroimage.2015.01.032.
- Waller, L., Walter, H., Kruschwitz, J.D., Reuter, L., Müller, S., Erk, S., Veer, I.M., 2017. Evaluating the replicability, specificity, and generalizability of connectome fingerprints. Neuroimage 158, 371–377. doi:10.1016/j.neuroimage.2017.07.016.
- Wang, D., Buckner, R.L., Fox, M.D., Holt, D.J., Holmes, A.J., Stoecklein, S., Langs, G., Pan, R., Qian, T., Li, K., Baker, J.T., Stufflebeam, S.M., Wang, K., Wang, X., Hong, B., Liu, H., 2015. Parcellating cortical functional networks in individuals. Nat. Neurosci. 18, 1853–1860. doi:10.1038/nn.4164.
- Wang, P., Kong, R., Kong, X., Liégeois, R., Orban, C., Deco, G., van den Heuvel, M.P., Thomas Yeo, B.T., 2019. Inversion of a large-scale circuit model reveals a cortical hierarchy in the dynamic resting human brain. Sci. Adv. 5, eaat7854. doi:10.1126/sciadv.aat7854.
- Wang, X., Liang, X., Jiang, Z., Nguchu, B.A., Zhou, Y., Wang, Y., Wang, H., Li, Y., Zhu, Y., Wu, F., Gao, J.H., Qiu, B., 2020. Decoding and mapping task states of the human brain via deep learning. Hum. Brain Mapp. 41, 1505–1519. doi:10.1002/hbm.24891.
- Wu, D., Fan, L., Song, M., Wang, H., Chu, C., Yu, S., Jiang, T., 2020. Hierarchy of connectivity-function relationship of the human cortex revealed through predicting activity across functional domains. Cereb. Cortex 30, 4607–4616. doi:10.1093/cercor/bhaa063.
- Yeh, F.C., Vettel, J.M., Singh, A., Poczos, B., Grafton, S.T., Erickson, K.I., Tseng, W.Y.I., Verstynen, T.D., 2016. Quantifying differences and similarities in whole-brain white matter architecture using local connectome fingerprints. PLOS Comput. Biol. 12, e1005203. doi:10.1371/journal.pcbi.1005203.
- Yeo, B.T., Krienen, F.M., Sepulcre, J., Sabuncu, M.R., Lashkari, D., Hollinshead, M., Roffman, J.L., Smoller, J.W., Zöllei, L., Polimeni, J.R., Fischl, B., Liu, H., Buckner, R.L., 2011. The organization of the human cerebral cortex estimated by intrinsic functional connectivity. J. Neurophysiol. 106, 1125–1165. doi:10.1152/jn.00338.2011.
- Zhang, Z., Liao, W., Chen, H., Mantini, D., Ding, J.R., Xu, Q., Wang, Z., Yuan, C., Chen, G., Jiao, Q., Lu, G., 2011. Altered functional-structural coupling of largescale brain networks in idiopathic generalized epilepsy. Brain 134, 2912–2928. doi:10.1093/brain/awr223.
- Zimmermann, J., Griffiths, J.D., McIntosh, A.R., 2018. Unique mapping of structural and functional connectivity on cognition. J. Neurosci. 38, 9658–9667. doi:10.1523/JNEU-ROSCI.0900-18.2018.
- Zöller, D., Schaer, M., Scariati, E., Padula, M.C., Eliez, S., Van De Ville, D., 2017. Disentangling resting-state BOLD variability and PCC functional connectivity in 22q11.2 deletion syndrome. Neuroimage 149, 85–97. doi:10.1016/j.neuroimage.2017.01.064.