- 1 Full Title:
- 2 Decoding task-specific cognitive states with slow, directed functional networks in the
- 3 human brain
- 4 Short title: Directed fMRI functional networks for decoding cognitive states

5 Authors:

6 Devarajan Sridharan^{1,2}, Shagun Ajmera¹, Hritik Jain¹ and Mali Sundaresan¹

7 Affiliations:

- 8 ¹Centre for Neuroscience, Indian Institute of Science, Bangalore 560012, India
- 9 ²Department of Computer Science and Automation, Indian Institute of Science, Bangalore 560012, India

10 **Corresponding author:**

- 11 Devarajan Sridharan
- 12 E-mail: sridhar@iisc.ac.in (DS)

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16 Abstract

Flexible functional interactions among brain regions mediate critical cognitive functions. Such interactions 17 can be measured from functional magnetic resonance imaging (fMRI) data with either instantaneous (zero-18 lag) or lag-based (time-lagged) functional connectivity; only the latter approach permits inferring directed 19 functional interactions. Yet. fMRI hemodynamic response 20 the is slow. and sampled at a timescale (seconds) several orders of magnitude slower than the underlying neural dynamics 21 (milliseconds). It is, therefore, widely held that lag-based fMRI functional connectivity, measured with 22 approaches like as Granger-Geweke causality (GC), provides spurious and unreliable estimates 23 24 of underlying neural interactions. Experimental verification of this claim has proven challenging because neural ground truth connectivity is often unavailable concurrently with fMRI recordings. We address this 25 challenge by combining machine learning with GC functional connectivity estimation. We estimated 26 instantaneous and lag-based GC functional connectivity networks using fMRI data from 1000 participants, 27 28 drawn from the Human Connectome Project database. A linear classifier, trained on either instantaneous or 29 lag-based GC, reliably discriminated among seven different task and resting brain states, with over 80% 30 cross-validation accuracy. With network simulations, we demonstrate that instantaneous and lag-based GC exploited interactions at fast and slow timescales, respectively, to achieve robust classification. With 31 32 human fMRI data, instantaneous and lag-based GC identified distinct, cognitive core networks. Finally, 33 variations in GC connectivity explained inter-individual variations in a variety of cognitive scores. Our findings show that instantaneous and lag-based methods reveal complementary aspects of functional 34 35 connectivity in the brain, and suggest that slow, directed functional interactions, estimated with fMRI, provide robust markers of behaviorally relevant cognitive states. 36

38 Author Summary

Functional MRI (fMRI) is a leading, non-invasive technique for mapping networks in the human brain. Yet, 39 fMRI signals are noisy and sluggish, and fMRI scans are acquired at a timescale of seconds, considerably 40 slower than the timescale of neural spiking (milliseconds). Can fMRI, then, be used to infer dynamic 41 processes in the brain such as the direction of information flow among brain networks? We sought to 42 answer this question by applying machine learning to fMRI scans acquired from 1000 participants in the 43 Human Connectome Project (HCP) database. We show that directed brain networks, estimated with a 44 technique known as Granger-Geweke Causality (GC), accurately predicts individual subjects' task-specific 45 46 cognitive states inside the scanner, and also explains variations in a variety of behavioral scores across individuals. We propose that directed functional connectivity, as estimated with fMRI-GC, is relevant 47 48 for understanding human cognitive function.

49 Introduction

Coordinated activity among brain regions underlies a variety of cognitive functions [1]. Mapping functional 50 51 coupling among brain regions is, therefore, key to mapping brain function and for understanding how the brain produces behavior. Human fMRI studies have commonly investigated such functional coupling with 52 correlation-based measures, including the Pearson correlation coefficient [2,3] and partial correlations [4] 53 between pairs of brain regions. These measures frequently incorporate regularization penalties to estimate 54 sparse functional networks [5]. Correlation-based measures are ideal for characterizing "instantaneous" 55 functional coupling, representing functional interactions among brain regions that occur at timescales faster 56 57 than the sampling rate of the measurement [6]. In contrast, comparatively few studies, have examined lagbased measures of functional connectivity to examine time-lagged interactions among brain regions [7,8]. 58

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Measures of linear dependence and feedback, based on Granger-Geweke causality (GC) [9,10] represent 60 61 a powerful approach for estimating both instantaneous and lag-based functional connectivity. These 62 measures are firmly grounded in information theory and statistical inferential frameworks [9-11]. GC 63 measures have been widely applied to estimate functional connectivity in recordings of brain activity made 64 with electroencephalography (EEG: [12]), magnetoencephalography (MEG: [13]) and electrocorticography 65 (ECoG: [14]). However, the application of GC measures to brain recordings made with functional magnetic 66 resonance imaging (fMRI) has provoked significant controversy [15–18]. Because the hemodynamic 67 response is produced and sampled at a timescale (seconds) several orders of magnitude slower than the 68 underlying neural processes (milliseconds), previous studies have argued that lag-based measures, 69 particularly lag-based GC, produce spurious and unreliable estimates of functional connectivity, when 70 applied to fMRI data (fMRI-GC) [18-21].

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Three primary confounds have been identified with inferring connectivity with fMRI-GC. First, systematic differences in hemodynamic lags across regions could yield spurious directionality of GC connections [16,18]. Second, in simulations, measurement noise added to the signal during fMRI acquisition significantly degrades GC functional connectivity estimates [19]. Finally, downsampling recordings to the

typical fMRI sampling rate (seconds), three orders of magnitude slower than the timescale of neural spiking
 (milliseconds), effectively eliminates all traces of functional connectivity inferred by GC [19].

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79 The controversy regarding the application of GC to fMRI data continues to date, primarily because of the 80 lack of access to ground-truth in neural data. On the one hand, claims regarding the efficacy of GC estimates are primarily based on simulations [11,22], and are only as valid as the underlying model of 81 neural activity and hemodynamic responses. Because the precise mechanism by which neural responses 82 83 generate hemodynamic responses is an active area of research, strong conclusions cannot be drawn based on fMRI simulations alone. On the other hand, establishing ground-truth validity for fMRI functional 84 85 connectivity requires invasive neurophysiological recordings across many brain regions, concurrently during fMRI scans, a prohibitive enterprise. 86

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88 We seek to address this controversy by applying machine learning to fMRI-GC networks, which works 89 around these challenges. We estimated instantaneous and lag-based GC connectivity with fMRI data drawn from 1000 human subjects, recorded under seven different task conditions and in the resting state 90 (~8000 functional scans drawn from the Human Connectome Project database; [23]). We trained a linear 91 92 classifier, based on GC connectivity features alone, to discriminate among the different task and resting conditions, and assessed classifier accuracy with cross validation. The results show that instantaneous and 93 lag-based GC connectivity, estimated from fMRI data, can decode task-specific cognitive states with 94 95 superlative accuracies. Next, with simulations, we show that slow, multi-second timescales emerge in 96 sparse, random networks despite individual neurons operating at fast, millisecond timescales – a result that 97 explains why directed functional connectivity can be reliably estimated with GC in slowly sampled fMRI 98 data. Finally, we demonstrate that GC connectivity features can be used as predictors [24] to explain inter-99 individual variations in behavioral scores across a variety of cognitive tests. The results suggest that 100 instantaneous and lag-based GC measures applied to fMRI data permit mapping slow, emergent and behaviorally relevant functional interactions in the human brain. 101

102 Results

103 GC estimated from slowly sampled fMRI data suffices to distinguish task and resting states

We asked if instantaneous GC (iGC) and directed GC (dGC) (SI Mathematical Note Section S1) 104 connectivity would flexibly reconfigure with task demand, by testing if GC connectivity sufficed to accurately 105 classify among seven different task states or the resting state (SI Table S1A; Methods; [9,10]). Data were 106 obtained from 1000 participants from the Human Connectome Project (HCP) database [25] (RRID: 107 SCR 008749). We used connection weights among brain regions in each network (iGC or dGC) as feature 108 vectors in a linear classifier based on Support Vector Machines (SVM) for high dimensional predictor data. 109 Accuracies for classifying resting state from a working memory task (WM task) are described first; 110 accuracies for other tasks are presented subsequently. 111

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Both iGC and dGC connectivity were able to distinguish the working memory task from resting state 113 significantly above chance (Fig. 1B: p<0.001, permutation test), Maximum accuracy (median, [95% CI]) was 114 97.3% [96.3 - 98.0%] with iGC and 92.0% [90.5 - 93.2%] with dGC (SI Fig. S1B Yeo Parcellation, iGC: 115 precision= 97.2, recall= 97.4; dGC: precision= 90.9, recall= 93.2). k-fold (k=10) cross-validation accuracy 116 was comparable (iGC: 97.1% [96.2 - 97.9%], dGC: 91.7% [90.3 - 93.0%]). These numbers correspond to 117 maximum cross validation accuracy across all five parcellations tested (SI Table S3; SI Fig. S1A); 118 accuracies with each parcellation are shown in the Supporting Information (SI Fig. S1B). Non-linear 119 classifiers, such as SVMs based on radial basis function kernels produced similar results, with comparably 120 above chance classification accuracy for both iGC and dGC connectivity (SI Fig. S1C). 121

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Figure 1. Discriminating between task and resting state with instantaneous and directed GC networks.

A. Schematic of task state classification based on instantaneous (iGC) and directed (dGC) Granger Geweke Causality with fMRI data from 1000 subjects (see text for details).

B. Two-way classification accuracies (leave-one-out) for each of seven tasks versus resting state based on
 GC. Red unfilled bars and blue filled bars: accuracies based on dGC and iGC features, respectively (task

key in SI Table S1). Error-bars: Clopper-Pearson binomial confidence intervals. Chance accuracy: 0.5 (notshown).

C. Two-way task versus resting state classification accuracies based on dGC (red dots) and iGC (blue dots), as a function of number of task scan time points (volumes). Dashed lines: linear fits.

D. Two-way task versus resting state classification accuracies based on dGC after averaging dGC matrices

over different numbers of subjects (x-axis). Each task is represented with a different color. Colored dashed
 lines: biexponential fits. Black dashed horizontal and vertical lines: 95% accuracy and n=5 subjects'
 average, respectively.

E. Two-way classification accuracies across each pair of tasks. Cells: classification accuracies for each pair
 of tasks based on dGC (lower triangular matrix) or iGC (upper triangular matrix). Diagonal cells: number of
 task scan timepoints. Highlighted cells: lowest (dashed-line border) and highest (solid-line border)
 accuracies achieved with dGC (red) and iGC (blue).

F. N-way classification accuracies among all seven tasks. Dashed line: chance accuracy (14.3%). Otherconventions are the same as in panel B.

G. Two-way sub-task classification accuracies (SI Table S1B) based on GC. ns.: accuracy not significantlyabove chance. Other conventions are the same as in panel B.

145 H. (Left) Two-way task versus resting state classification accuracies obtained with regional time series subsampled at 2x (filled symbols) and 3x (open symbols) of the TR (720 ms) (y-axis) plotted against 146 accuracies obtained with the original data (1x, x-axis) for each of 7 tasks. Red: dGC, Blue: iGC. Dashed 147 diagonal line: Line of equality (x=y). (Right) N-way classification accuracies among all seven tasks with data 148 sampled at 1x, 2x, 3x of the original TR. Other conventions are the same as in panel F.For panels B.E.F: 149 150 accuracies correspond to highest values, across all parcellations tested, and hyperparameter optimization was done for panel B. For panels C,G,H: accuracies correspond to Shirer et al [26] 14-network parcellation. 151 152 For panel D: accuracies correspond to Shirer et al [26] 90-node parcellation.

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We repeated these analyses by classifying the six other tasks (SI Table S1) versus resting state. iGC and dGC connectivity could accurately classify each task from resting state significantly above chance. For iGC, maximum classification accuracies ranged from 90.1%, for emotion task vs. resting state classification, to 97.1%, for language task vs. resting state classification. Similarly, for dGC, accuracies ranged from 78.1%,

for emotion task vs. resting state classification, to 92.8%, for language task vs. resting state classification (Fig. 1B). In general, classification accuracy increased with more scan timepoints for each task versus resting state classification (Fig. 1C), consistent with GC being an information theoretic measure; we confirmed this result with simulations also (SI Fig. S1D).

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In these analyses, classification accuracies based on dGC were systematically lower than those based on 163 iGC. We asked if dGC accuracies were poorer due to noise corrupting the fit of the autoregressive model, 164 and if a more consistent estimate could be obtained by averaging dGC connectivity features, to remove 165 uncorrelated noise, across subjects. We addressed this guestion by partitioning the data into two groups --166 a training (T) group and a test (S) groups - with 500 subjects each. We trained the classifier on group T 167 and tested the classifier prediction by averaging GC matrices across several folds of S, each fold 168 containing a few (m=2, 4, 5, 10, 25 or 50) subjects; the procedure was repeated by exchanging training and 169 test datasets (see Methods). For the vast majority of tasks (6/7), dGC's classification accuracy was more 170 than 95% with as few as m=5 subjects within each fold of the test set (Fig. 1D). These results suggest that 171 averaging dGC matrices across a few subjects, yielded reliable estimates of dGC connectivity. 172

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174 We considered other factors that, in addition to intrinsic connectivity differences, could have produced these superior classification accuracies. First, GC-based accuracies for classifying task versus resting state 175 scans might arise from differences in brain regions activated during each of these scans. In addition to 176 task-relevant sensory input, overt motor responses always occurred during task scans but were absent 177 during resting state scans [23,27]. Could GC features discriminate among more subtle connectivity 178 variations across/within tasks? Second, scan data from the HCP database was sampled at a TR (repetition 179 time) of 720 ms, considerably faster than the TR for conventional fMRI scans. Would GC accuracies 180 degrade if the data were sampled at much slower sampling rate (~2000 ms), in line with conventional fMRI 181 182 TR?

183

We addressed the first question in two stages. First, we asked if GC connectivity features would be able to classify which of the seven tasks each subject was performing in the scanner. First, we performed a pairwise classification of each task from the other. Maximum classification accuracies for iGC (dGC) ranged

from 87% (67%) for the emotion vs. gambling task classification to 98% (91%) for the language vs. social 187 task classification. Again, the number of timepoints for each task proved to be a strong indicator of 188 classification accuracies (Fig. 1E): average inter-task classification accuracies were highest for the 189 language task (iGC: 97%, dGC:88%, n=316 timepoints) and lowest for the emotion task (iGC: 91%, dGC: 190 77%, n=176 timepoints). Next, we performed an n-way classification analysis across all 7 tasks, again 191 using linear SVM (Methods). Accuracies were significantly above chance (14.3% for 1-in-7 classification) 192 for classifying among the seven tasks (Fig. 1F; maximum accuracy, iGC: 74.4% [73.3%-75.4%]; dGC: 193 47.6% [46.4%-48.7%]; p<0.001, permutation test). These results indicate that functional connectivity was 194 consistently estimated with GC, and reliably different across tasks. 195

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Second, each of the different tasks in the HCP database comprised of blocks of contiguous trials, each 197 corresponding to one of (at least) two different sub-tasks ([27]; SI Table S1B). For example, the motor task 198 comprised of blocks of movements of the right or left hand interleaved with blocks of trials involving 199 movement of the right or left foot. Similarly, the working memory task comprised of interleaved blocks of 0-200 201 back and 2-back tasks. We asked, therefore, if GC connectivity could distinguish among subtler variations in brain states across sub-tasks within each task. We sought to classify across two sub-tasks for each of six 202 tasks (SI Table S1B). In all cases, except one, both iGC and dGC connectivity discriminated between each 203 pair of sub-tasks with higher than chance accuracies (Fig. 1G; maximum accuracy, iGC: 89.2% [87.6% -204 90.7%]; dGC: 80.1% [78.9% - 82.9%]; p<0.05 permutation test). These results indicate that GC functional 205 connectivity could accurately distinguish among sub-tasks within each task as well. 206

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Next, we tested whether GC connectivity estimated from slowly sampled fMRI data could accurately 208 classify task and resting states. We downsampled the data to either one half (2xTR=1440 ms) or one third 209 (3x TR=2160 ms) of its original sampling rate, by decimation, while also concatenating the decimated data 210 211 to the end of the sub-sampled timeseries to preserve the overall number of timepoints (Methods). We repeated both of the previous classification analyses - pairwise task versus resting state classification (Fig. 212 1H left), as well as n-way inter-task classification (Fig. 1H right). Following downsampling, we observed that 213 classification accuracies were marginally higher than accuracies in the original data in the case of dGC (2x: 214 215 p=0.02; 3x: p=0.06; Wilcoxon one-tailed signed rank test), and were even higher than those in the original

data in the case of iGC (2x: p=0.01; 3x: p=0.01), across tasks. These results indicate that the superlative
 sampling rate of the HCP fMRI data was not the primary reason for these high classification accuracies for
 GC-based classification.

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We performed three control analyses to further confirm these results: i) by performing stationarity tests on the data prior to GC estimation and classification; ii) using single full regression to estimate GC [28,29], instead of estimating with separate full and reduced regressions; and iii) incorporating motion scrubbing [30] to ensure that the classification accuracies were not driven by head motion artifacts. These controls are described in SI Results, section on "Control analyses". In every case, we obtained equivalent or superior classification accuracies (SI Fig. S2), confirming that the results were not due to data nonstationarity, biases in GC estimation or head motion artifacts.

227

These results demonstrate that both iGC and dGC yielded task-specific signatures of functional connectivity even with slowly sampled fMRI data (TR~2000 ms): these estimates were consistent across subjects and reliably different across tasks to permit successful classification. Furthermore, these superlative classification accuracies were obtained despite widely held caveats concerning the application of GC to fMRI data [28].

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235 Correlation-purged GC connectivity suffices for accurate task-state classification

236 Correlation-based (zero-lag) connectivity measures (e.g. partial correlations or PC) have been widely 237 applied to estimate functional connectivity from fMRI data [5,31]. In fact, several previous studies [18,19] 238 have argued that correlation-based measures are more reliable and should be preferred to lag-based 239 measures like GC [11], for estimating functional connectivity with fMRI data. We tested this claim here with 240 a three-fold analysis approach.

241

First, we asked how classification accuracies based on PC connectivity would compare with those reported above, based on GC connectivity. Maximum classification accuracies with PC connectivity ranged from 96-99% for task versus resting state classification, and were consistently higher than accuracies with GC

connectivity (Fig. 2A). These results are along expected lines: estimators based on same-time covariance, such as PC, are less susceptible to noise than those based on lagged covariance, such as GC (derived analytically in the Methods, section on *Functional connectivity estimation and classification with partial correlations*). In addition, as mentioned previously, GC is an information theoretic measure: classification accuracy with iGC and dGC increased systematically with more scan time points, asymptotically matching PC accuracies (SI Fig. S1D).

251

252 Figure 2. Classification accuracies with GC purged of instantaneous correlations.

A. Two-way task versus resting state classification accuracies, based on partial correlations (PC; grey unfilled bars). Numbers reported correspond to highest leave-one-out classification accuracies across parcellations, obtained with hyperparameter optimization. Corresponding accuracies for dGC (red dots) and iGC (blue dots) are shown for comparison. Other conventions are as in Fig. 1B.

B. Schematic illustrating procedure for purging data of instantaneous correlations. fMRI regional timeseries were purged of instantaneous correlations by either whitening the data with zero-phase component analysis (ZCA), separately for each task and resting state scan, or by projecting data into a space spanned by the generalized eigenvectors (GEV), common to both task and resting state scans. GC and PC were then estimated with the ZCA or GEV projections of the timeseries data, followed by classification analysis based on GC or PC connection strength as features.

C. (Top) Two-way task versus resting state classification accuracies following ZCA-based decorrelation.
 Gray circles: Classification accuracies based on PC. Other conventions are as in Figure 1B. Dashed line:
 chance accuracy (50%).(Bottom) Same as in top panel, but for classification following GEV-based
 decorrelation.

D. (Top) Schematic showing unweighted directed graph obtained from dGC; this digraph representation encodes only the dominant direction of connectivity, and not its magnitude. (Bottom) Two-way task versus resting state classification accuracies based on dGC digraph representations. Secondary ordinate (y-axis on the right): number of scan timepoints for each task.(Panels C-D). GC features were estimated with the Shirer et al [26] 14-network parcellation.

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Second, we asked if lag-based connectivity could accurately classify task from resting state, once the data 273 were purged of all instantaneous correlations. To accomplish this, we adopted two approaches: i) zero-274 phase component analysis (ZCA) and ii) generalized eigenvalue decomposition (GEV) (Methods). Briefly, 275 ZCA (or the Mahalanobis transformation) produces whitened time series data that is closest, in a least 276 squares sense, to the original regional time series data. As an alternative approach, we decorrelated both 277 task and resting state time series jointly by projecting them onto a single set of generalized eigenvectors 278 (GEV). These approaches provided empirical upper and lower bounds on GC's performance on correlation-279 purged data (Methods). 280

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GC connectivity features sufficed to successfully classify all tasks from resting state, even in correlation-282 purged data. With ZCA, iGC accuracies ranged from 84% to 96% whereas dGC accuracies ranged from 283 82% to 96% across tasks. With GEV, iGC accuracies ranged from 60% to 71% whereas dGC accuracies 284 ranged from 56% to 76% across tasks; in each case, classification accuracies were significantly above 285 chance (p<0.001, permutation test). We confirmed that performance in each case was not an artifact of the 286 decorrelation procedure (ZCA/GEV) by randomly interchanging task and resting state labels for each pair of 287 datasets across subjects (Methods); shuffling labels reduced classification accuracy to chance. Note that in 288 289 every case, classification performance based on PC connectivity was at chance (Fig. 2C), a direct consequence of removing instantaneous correlations from the data. Despite this, classification accuracies 290 based on iGC connectivity were not at chance; in the next section, we discuss potential reasons for these 291 differences between iGC and PC classification accuracies. 292

293

Third, we asked if an unweighted directed graph (digraph) network representation - whose edges indicated 294 the dominant direction, but not magnitude, of connectivity (Fig. 2D) - would suffice to distinguish task from 295 resting brain states (Methods). Again, dGC directed graphs successfully distinguished each task from 296 resting state well above chance. Classification accuracies ranged from 56% for the motor task versus 297 resting state classification to 68% for the relational task versus resting state; for each task, classification 298 accuracies were significantly above chance (p<0.001; permutation test). Interestingly, we did not see a 299 strong influence of the number of data points on classification accuracy in this case (Fig. 2D, purple dots). 300 301 For instance the emotion task (n=176 timepoints) was classified with an accuracy of 62% from resting state,

which was comparable to the classification accuracy of working memory (n=405 timepoints) from resting state (64%). Both iGC and PC, which are symmetric connectivity measures, could provide no directed connectivity information.

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These results demonstrate that lag-based connectivity contained sufficient information to classify task from resting state even when instantaneous correlations were entirely purged from the data. Moreover, unweighted directed connectivity graphs alone, indicating the direction, but not scalar magnitude, of GC connectivity, sufficed to accurately classify task from resting brain states. These findings indicate that directed functional connectivity measures, like dGC, provide connectivity information that is distinct from, and complementary to, what can be obtained with undirected functional connectivity measures, like PC.

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313 Instantaneous and directed GC identify complementary aspects of functional connectivity

What characteristics of functional connectivity are respectively identified by instantaneous and lag-based connectivity? And how can lag-based connectivity be reliably estimated with fMRI data, which is sampled at time scales orders of magnitude slower than neural timescales? We addressed both of these questions, first, with simulations (this section) and, then, with real data (next section).

318

First, we tested the ability of GC to reliably recover functional interactions in simple, two-node feedforward 319 networks operating at different timescales (Fig. 3A). We simulated fMRI data using a two-stage model 320 (Methods): i) a latent variable model that describes the dynamics of the nodes (vector Ornstein-Uhlenbeck 321 process; [32]); ii) a convolution of these neural dynamics with a hemodynamic response function to obtain 322 the simulated fMRI time series [18,19]. Based on this model, we simulated activity in two 2-node networks. 323 In the first network, individual node decay timescales were set to 50 ms, whereas in the second network, 324 these were set to 1000 ms (parameters in SI Table S6A). For convenience, we refer to these two network 325 timescales as "fast" (50 ms) and "slow" (1000 ms). We then varied the sampling interval (Ts) of the 326 simulated data from 50 ms to 1450 ms in steps of 100 ms. Connections at both "fast" and "slow" timescales 327 were generally discovered by iGC regardless of sampling interval, although connections at slow timescales 328 were less robustly detected than those at fast timescales (Fig. 3A). On the other hand, the connection in 329

the "fast" timescale network was not discovered by dGC when the sampling interval was higher than 50 ms, in line with the results of Smith et al [18]. However, the connection in the "slow" timescale network was reliably discovered by dGC across a wide range of sampling intervals, upto, and exceeding 1000 ms. In each case, dGC failed to discover the underlying interaction when the sampling interval was much higher than the slowest timescale in each network, consistent with recent theoretical results [6]. These findings suggest that dGC can detect slow neural processes, which operate at a timescale slower than TR, in fMRI data.

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Figure 3. Robustness of GC estimates depend on network timescales in simulated hemodynamic data.

A. (Top) Two-node networks with fast (50 ms; left) or slow (1000 ms; right) decay timescales of individual 340 nodes. Each subpanel shows ground truth connectivity either as a schematic (left) or connectivity matrix 341 (right). In the matrix, a non-zero entry at cell (i, i) corresponds to a connection from node i (source) to node 342 i (destination).(Bottom) dGC (red), iGC (blue), and PC (black) connection strengths as a function of 343 344 sampling intervals. Filled circles and solid lines: Strengths of true connections and curve (biexponential) fits, respectively. Open circles and dashed lines: Strengths of spurious connections and curve fits, 345 respectively. Dashed vertical line: Sampling interval of 750 ms, mimicking the TR of the fMRI data. Matrices 346 to the right of each plot show GC connection strengths estimated at sampling interval of 750 ms. Black 347 squares surrounding matrix cells denote significant connections (Methods). For iGC and PC (symmetric 348 connectivity), only the lower triangular matrix is shown, for clarity. 349

B. (Top left) Schematic showing a cluster of neurons, each with timescale 50ms, connected with sparse,

random, net excitatory connectivity. Matrix: Connectivity among the 100 neurons in a representative cluster.

Red: excitatory connections; blue: inhibitory connections. Each such cluster forms one of the nine nodes in

the simulated network. (Top right) Connectivity among the nine nodes in the network. (Bottom left)

Eigenspectrum (upper panel) of a representative 100 neuron cluster, showing one slow emergent timescale

355 corresponding to the real-part of one eigenvalue close to zero. Histogram (lower panel) showing timescales

of all eigenmodes, with the slowest eigenmode at >2000ms. (Bottom right) Eigenspectrum (upper panel) of

357 sub-network DEF exhibits multiple slow emergent timescales. Histogram (lower panel) showing timescales

of all eigenmodes, with three slow eigenmodes at ~1000-6000 ms.

C. Same as in A, but for simulated 9-node networks. (Left) Sub-network ABC, (middle) sub-network DEF,
(right) sub-network GHI. Other conventions are as in panel A.

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How might such slow timescales, orders of magnitude slower than spike times and membrane time 362 constants, arise in fMRI data? To answer this question, we availed of established results in random matrix 363 364 theory. Connectivity in randomly connected E-I networks of neurons can produce slow timescales, without fine-tuning of network parameters [32,33]. We modeled sparse, random, net excitatory connectivity in a 365 small network of (N=100) neurons with connection parameters drawn from previous studies (SI Table S6B; 366 367 [32,34,35]). The eigenspectrum of the network revealed that each network exhibited one eigenvalue close to zero corresponding to a slow timescale (~1000 ms or greater, Fig. 3B bottom left); the latter constitutes 368 an emergent timescale associated with the dominant eigenmode that is a property of network connectivity 369 370 (Methods).

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We modeled nine such networks, organized into three, non-interacting, clusters (Fig. 3B top right): a) a 372 cluster with a purely feedforward connection across two networks, b) a cluster with recurrent excitatory (E-373 E) feedback connections among two networks and c) a cluster with recurrent excitatory-inhibitory (E-I) 374 375 feedback connections among two networks. In each case, connectivity across networks was mediated by a 376 small proportion (5%) of neurons in each network (parameters in SI Table S6B). This configuration mimics "small-world" connectivity in brain networks [36], with locally-connected brain regions interacting through 377 sparse, long-range connections [37]. The eigenspectra revealed that dynamics in all clusters operated at 378 timescales of around 6000 ms, comparable to or slower than the individual network timescales (Fig. 3B 379 bottom right). To simulate fMRI data we averaged the activity across all 100 neurons in each network and 380 convolved it with a canonical HRF. As before, these nine timeseries were then sampled at various sampling 381 intervals, including a 750 ms interval mimicking the scan TR, and analyzed with GC to detect significant 382 connections. 383

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iGC and dGC identified complementary aspects of connectivity with these simulated data (Fig. 3C). iGC robustly identified feedforward and excitatory (E-E) feedback connections. dGC also estimated these

connections, albeit with the following differences: First, in the feedforward network dGC occasionally 387 identified a spurious connection, albeit much weaker in magnitude, in the direction opposite to the true 388 connection (Fig. 3C, left column, red dashed line). Second, when the E-E feedback connections were 389 precisely balanced in strength (symmetric), dGC also failed to identify the connection reliably (SI Fig. S3A). 390 Yet, when these connections were of different strengths dGC reliably identified both connections, and their 391 relative strengths (Fig. 3C, middle column, red). In contrast, when the connections were of different signs 392 (E-I feedback) dGC robustly identified both connections, whereas iGC failed to reliably detect this 393 connection (Fig. 3C, right column, blue). Yet, taken together, iGC and dGC identified all three connection 394 types reliably. 395

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Next, we compared the efficacy of connectivity estimation with partial correlations (PC). While PC robustly 397 identified both feedforward and feedback E-E connections (Fig. 3C left and middle columns, black), it, 398 399 surprisingly, failed to estimate feedback E-I connections, particularly when these were balanced in strength (Fig. 3C right column, black). When the E and I connection strengths were not balanced, but were strongly 400 biased in favor of the E or the I connection, PC estimates varied with the sign of the more dominant 401 connection (SI Fig. S3B, right top). These results generalize beyond these particular simulations; in SI 402 403 Mathematical Note, Section S2 and S3, we identify, analytically, network configurations for which PC estimates systematically deviate from ground-truth connectivity (see also SI Results, section on "Analytical 404 relationship between PC and iGC"). 405

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Taken together, these results indicate that instantaneous and lag-based connectivity measures can reveal complementary aspects of brain connectivity. In addition, the results challenge the notion that correlationbased measures, like PC, should be favored over lag-based measures, like dGC for measuring functional connectivity in the brain [18]. Rather, the strengths and weaknesses of each measure (GC and PC) must be recognized when seeking to apply these to brain imaging data.

412

Identifying a cognitive core system and predicting behavioral scores with GC connectivity

Our classification analyses and simulations suggested that iGC and dGC reliably recover task-specific brain networks, the latter when slow-timescale processes occur within the network. We asked whether iGC and

dGC connectivity merely reflected reliable statistical patterns of brain activity, or whether it would be relevant for understanding the nature of information flow in the brain, and its relationship to behavior. To answer this question, we first investigated whether each measure would identify brain networks with consistent outflow and inflow hubs across tasks. Next, we asked whether GC connectivity would be relevant for predicting brain-behavior relationship.

421

First, we sought to identify a common core of "task-generic" connections across cognitive tasks. For this, 422 we applied a feature selection approach - recursive feature elimination (Methods) - a technique that 423 identifies a minimal set of features that provide maximal cross validation accuracy (generalization 424 performance) [38]. Prior to analysis of real data, we validated RFE by applying it to estimate connectivity 425 differences in two simulated networks (Fig. 4A,B). RFE accurately identified connections that differed in 426 simulation ground-truth: specifically, differences in fast timescale connections were reliably identified by 427 iGC, and in slow timescale connections by dGC (Fig. 4B bottom). RFE based on dGC and iGC also 428 accurately identified the relevant connections, but not always their directionality, even when systematic 429 variations in hemodynamic lag occurred across regions; the results are described in SI Results, section on 430 "Effects of regional variations in hemodynamic lag". 431

432

Figure 4. Recursive feature elimination (RFE) identifies task-generic and task-discriminative networks based on GC connectivity.

435 A. Schematic showing two simulated networks each with fast (50 ms; ABC) and slow (1000 ms; DEF) sub-

436 networks, with distinct connectivity patterns. Network activity was simulated for 375 seconds with a

sampling interval of 5 ms, convolved with the hemodynamic response function and sub-sampled at 750 ms

to yield 500 simulated time points.

B. (Top) RFE curves, with classification accuracy as a function of remaining features, for classification
based on dGC (left) and iGC (right). (Bottom) Maximally discriminative features following RFE based on
dGC (left) and iGC (right). Entries denote average beta weights across RFE iterations.

442 C. RFE curve for two-way classification of each of six tasks (all tasks except Motor) versus rest, based on

dGC (top) and iGC (bottom). Color conventions are as in Figure 1D. Data points: RFE accuracies; solid

444 lines: piecewise linear fits. Vertical dashed line: location of the elbow for each RFE curve.

D. Task-generic connections following task-versus-resting RFE, based on dGC (left) and iGC (right)
features, using Shirer et al (2012) 14-network parcellation [26] (SI Table S4); each network is indicated with
a different color and a label. Directed dGC connections are shown as tapered links, broad at the source
node and narrow at the destination node. Undirected iGC connections are shown as bidirectional links
between the respective pair of nodes. Colors of the connections represent the color of the destination node.
E. Same as in panel D, but for n-way classification across the six tasks. Color conventions are as in panel
B.

F. Same as in panel E, but for task-discriminative connections, which maximally discriminated each task from the five others, following n-way RFE, based on dGC features (left) and iGC features (right). Other conventions are the same as in panel C.

455

We applied RFE to classify tasks versus resting state; we chose these six tasks (all tasks except the motor task) as being the most likely to engage common cognitive control mechanisms (Fig. 4C). For these RFE analyses we employed a 14 network functional parcellation [26], as it consistently gave good classification accuracies with both iGC and dGC connectivity (SI Fig. S1B). Following RFE, we applied a binomial test across tasks (Methods) to identify a common core of task-generic connections, separately for iGC and dGC.

462

RFE identified distinct task-generic networks with iGC and dGC, which comprised of connections that 463 distinguished a majority of tasks from resting state. The iGC task-generic network revealed a visuospatial 464 network hub, which connected with the anterior salience, dorsal DMN, higher visual and posterior salience 465 networks (Fig. 4D, right). The dGC task-generic network confirmed the hub-like connectivity of the 466 visuospatial network but, in addition, revealed consistent directed information outflow from the visuospatial 467 network to the other networks (Fig. 4D, left). In addition, dGC revealed consistent inflow into the higher-468 469 visual network across tasks, including from the visuospatial, right executive control, and auditory networks, consistent with the ability of top-down inputs from these networks to strongly modulate sensory encoding in 470 higher visual cortex [39]. Finally, the higher-visual network projected consistently to the sensorimotor 471 network, suggesting a final common pathway, across these tasks, for influencing behavior. Interestingly, 472 473 the only network providing inflow into the visuospatial network hub was the anterior salience network, in line

with a previous study that indicated a role for the salience network in controlling other task positive networks [7].

476

Similarly, we also identified connections that were maximally discriminative across tasks; again iGC and dGC showed distinct sets of these "task-discriminative" connections (Fig. 4E-F; SI Results, section on "Identifying task discriminative networks with GC").

480

To address GC's relevance for understanding brain-behavior relationships we tested whether the strength of functional connections estimated with iGC and dGC could predict inter-individual variations in behavioral scores as measured by a standard cognitive battery (Methods; SI Table S7). We employed a leave-one-out prediction analysis based on multilinear regression followed by robust correlations of predicted and observed scores (Fig. 5A; p<0.05 with Benjamini-Hochberg correction; Methods).

486

487 Figure 5. GC connectivity explains inter-individual variations in behavioral scores.

488 A. (Left) Schematic of behavioral score prediction analysis. GC connectivity strengths for each task were 489 used as independent factors to predict behavioral scores using linear regression with a leave-one-out 490 approach. 51 different behavioral scores were predicted (SI Table S7), compared against observed scores 491 (upper right), and their correlation values plotted as a matrix (lower right).

B. Exemplar score predictions based on dGC (left panels) and iGC (right panels). In order (from left to right): List Sorting score predicted from Working memory task dGC connectivity, Anger Emotion Recognition score from Emotion task dGC connectivity, Endurance score from Motor task iGC connectivity and Reading score from Language task iGC connectivity.

C. (Top) Prediction statistics for selected scores based on dGC connectivity. Correlation coefficients (r values) between the predicted and observed scores are plotted in the top half of each stem plot, and significance (p values) are plotted in the bottom half. Each score is denoted by a different color, and each sub-panel shows predictions based on GC connectivity for a different task; Stems with open symbols represent non-significant correlation coefficients, whose corresponding p-values are not shown. p values are floored at 10-4for ease of visualization. (Bottom) Same as in top panel, but predictions based on iGC connectivity.

D. (Top) Inter-subject correlation matrix of composite behavioral scores. Row and column indices: subjects. (Bottom) Cumulative distributions (solid lines) and density function estimates (filled area) of correlation coefficients between observed and predicted composite scores, for the same subject (yellow) or across different subjects (grey). Predictions were based on GC estimates from the relational and working memory tasks. p-value: Kolmogorov-Smirnov test.

508

Both iGC and dGC predicted key behavioral scores. Several scores were predicted uniformly well by iGC 509 across tasks (Fig. 5B, right; Fig. 5C, bottom; SI Fig S5B). Scores of fluid intelligence (Penn progressive 510 matrices), grip strength, endurance, and language (reading and picture-vocabulary) (Fig. 5B right; r: 0.077 -511 0.363: p<0.02), were all well predicted by iGC, in addition to scores of spatial orientation (Penn line 512 orientation test)and dexterity (SI Fig. S5B, r:0.081 -0.243; p<0.0125). On the other hand, dGC-based 513 predictions were more selective, in that several behavioral scores were best predicted by dGC based on 514 specific tasks alone (Fig. 5B, left; Fig. 5C, top; SI Fig. S5A). For instance, dGC in the emotion task alone 515 predicted positive affect (Fig. 5C, top; r=0.094, p=0.028) and anger emotion recognition (Fig. 5B, left, cyan; 516 r=0.106, p=0.001) scores, dGC in the gambling task alone predicted self-report scores of perceived social 517 support (r=0.101, p=0.002), fear (r=0.139, p<0.001) and sadness (r=0.107, p=0.001) and dGC in the motor 518 task alone predicted median reaction time in the fluid intelligence test (r=0.123, p<0.001). In addition, dGC 519 in the working memory task predicted a range of scores in the "cognition" category including list sorting(Fig. 520 5B, left,pink; r=0.119, p=0.000), fluid intelligence, picture discrimination speed, spatial orientation and self 521 regulation (discounting of delayed reward; Fig. 5C top, SI Fig. S5A). 522

523

A variety of behavioral scores were also successfully predicted based on PC connectivity (SI Fig. S5C); several of these (~60%) overlapped with those predicted with GC connectivity (SI Results, section on "Predicting behavioral scores with PC connectivity"). Yet, PC connection features that led to successful predictions overlapped strongly with iGC connection features than with dGC connection features, as quantified by the mean correlation between their regression weights in the respective prediction models (PC vs. iGC: r=0.38±0.01, mean±std; PC vs. dGC: r=0.03±0.01, p<0.001, ranksum test).

530

Finally, we tested whether GC connectivity could predict a combined set of behavioral scores unique to 531 each subject. For this, we created a vector of all independent behavioral scores (composite score; 532 Methods), and confirmed that this composite behavioral score uniquely identified each subject in the 533 database, as evidenced by the highest values along the main diagonal of the inter-subject correlation 534 matrix (Fig. 5D top). Following this, we performed the leave-one-out prediction, as before, except that we 535 used dGC and iGC connectivity features from two of the tasks alone (working memory and relational; also 536 see SI Fig. S5D). We then tested whether each subject's predicted composite score would correlate best 537 with her/his own observed composite scores. Although we did not observe the highest correlation values 538 consistently along the main diagonal, the distribution of correlation coefficients along the diagonal were 539 significantly different (and higher) than the distribution of off-diagonal correlation coefficients (Fig. 5D 540 bottom: p<10⁻¹⁵. Kolmogorov-Smirnov test). Inter-individual variation GC connectivity, therefore, contained 541 sufficient information to accurately identify subject-specific behavioral scores in this cohort of subjects. 542

543

In summary, the ability to successfully predict subject-specific behavioral scores suggests that GC functional connectivity is relevant for understanding brain-behavior relationships. Moreover, connection features that were relevant for behavioral predictions with PC overlapped highly with iGC, but not with dGC, thereby validating our simulation results regarding the complementarity of iGC and dGC connectivity estimates.

550 Discussion

Neural processes in the brain range from the timescales of microseconds to milliseconds, for extremely 551 rapid processes (e.g. sound localization), to timescales of several seconds to minutes, for processes that 552 require coordination across diverse brain networks (e.g. when having a conversation), and hours to days, 553 for processes that involve large-scale neuroplastic changes (e.g. when learning a new language). 554 Coordinated activity among brain regions that mediate each of these cognitive processes should manifest 555 in the form of functional connectivity among these regions at the corresponding timescales. Our results 556 indicate that applying Granger-Geweke Causality (GC) with fMRI data permits estimating behaviorally 557 relevant functional connectivity at a timescale corresponding to the sampling rate of fMRI data (seconds). 558

559

The application of GC to neuroscience is a contentious topic, for a variety of reasons [15–17,19,28]. One particular challenge stems from the use of the word "causality": the notion of causality in GC is different from the notion of interventional causality [40]. Our use of the term Granger causality, here, purely reflects its application as a marker of information flow among brain networks [19,41], and is not meant to indicate causality in a physical, interventional sense.

565

With this understanding, our results contain three key insights. First, we show that, either iGC or dGC connectivity suffices to reliably classify task-specific cognitive states with superlative accuracies (Fig. 1B). Instantaneous and directed GC – both measures of conditional linear dependence and feedback [9] – were able to robustly estimate task-specific functional interactions even with slowly sampled fMRI data. Our application of machine learning and classification analysis circumvents the lack of access to ground truth connectivity, and our simulations suggest that GC connectivity is relevant for estimating slow, emergent interactions among brain networks [15–19].

573

574 Second, we show that functional connections identified by iGC and dGC carry complementary information, 575 both in simulated and in real fMRI recordings, and we demonstrate key caveats with employing correlation-576 based measures of functional connectivity like partial correlations, despite superior classification accuracies 577 with these latter measures. First, PC fails to correctly infer reciprocal excitatory-inhibitory interactions, which

can be accurately inferred with lag-based methods like dGC. Second, PC may yield incorrect estimates of 578 functional connectivity that do not match ground truth (SI Fig. S3C). In particular, when the data are well 579 described by an autoregressive model framework our results suggest that instantaneous connectivity 580 measures, like iGC, provide more accurate descriptions of functional connectivity than PC. Third, even with 581 data completely purged of partial correlations, dGC connectivity was sufficient to classify task-specific 582 cognitive states (Fig. 2C). In fact, unweighted directed connectivity alone sufficed to produce accurate 583 classification at accuracies significantly above chance (Fig. 2D). These results indicate that information flow 584 mapped by GC connectivity can be complementary to that of PC, and highlights the need for examining 585 diverse measures, both instantaneous and lag-based, to obtain a complete picture of functional connectivity 586 in the brain. 587

588

Third, differences in inter-individual iGC and dGC connectivity were able to successfully explain inter-589 individual variation in behavioral scores on various cognitive tasks, and to identify an individual-specific 590 composite marker of behavioral scores, with high accuracies. Because these behavioral scores were 591 acquired in a separate testing session outside the scanning session [27], the results suggest that GC 592 connectivity was both individual-specific, and stable over timescales exceeding the scan session, to permit 593 accurate prediction. Moreover, in our analysis, each subject's behavioral score was predicted based on 594 her/his GC connectivity, whereas the regression beta weights - representing the relationship between GC 595 connectivity and behavior - were computed from the population of all subjects excluding that subject (Fig. 596 5A). Successful predictions, therefore, indicate a consistent mapping between GC connectivity and 597 behavioral scores across the population of subjects. These findings complement recent results showing 598 that dynamic, resting-state functional connectivity, based on correlations, can explain significant variance in 599 human behavioral data [42], and indicate the relevance of lag-based connectivity measures for 600 understanding brain-behavior relationships. 601

602

Does GC's discriminatory power rely on directed functional connectivity in the underlying neural response or systematic distortions of this connectivity induced by subsampling [19] and hemodynamic filtering [20,21]? While our findings cannot completely rule out the latter hypothesis, we address, next, three key

606 caveats raised by previous studies for estimating functional connectivity with fMRI-GC, and argue why our 607 results support the former hypothesis.

608

First, several studies have shown that sub-sampling of neural time series, at the scale of fMRI TR, renders 609 functional connections undetectable with GC [11,18-20]. In these studies, GC was estimated with 610 simulated fMRI time series, sampled at an interval (TR) of seconds, and failed to recover underlying neural 611 interactions, which occur at millisecond timescales (e.g. [18]). However, these claims depended strongly on 612 the nature and timescale of the connectivity in the networks employed in these simulations. For instance, a 613 widely cited study [18] employed purely feedforward connectivity matrices with a 50 ms neural timescale in 614 their simulations, and argued that functional connections are not reliably inferred with GC applied to 615 simulated fMRI data. In addition to being neurally implausible, such purely feedforward network 616 configurations yield eigenmodes whose slowest timescales are identical with the timescales of individual 617 nodes [43]. Therefore, such a configuration rendered lag-based measures like GC, irrelevant for estimating 618 neural interactions from slowly sampled fMRI data [18,19]. Furthermore, such connectivity precludes the 619 occurrence of slower, behaviorally relevant timescales of seconds, which readily emerge in the presence of 620 feedback connections, both in simulations [32,33] and in the real brain [44,45]. Our simulations show that 621 slow timescale interactions emerge in networks with sparse, random, net excitatory connectivity, mimicking 622 connectivity in the neocortex [32,33,35]. While earlier studies have employed large-scale, biologically 623 plausible models [46,47] to demonstrate the emergence of slow (<0.1 Hz) emergent functional interactions 624 among brain networks, our results build upon these previous findings and show that such emergent 625 functional interactions at slow timescales can be readily inferred from simulated fMRI data with GC. In fact, 626 GC connectivity continued to robustly classify distinct task states even when data were sampled at 2x or 3x 627 the original sampling interval of the fMRI data. Thus, while it is likely that GC applied to fMRI data is unable 628 to detect connections at timescales faster than TR, our results show that sufficient distinguishing 629 630 information occurs in slow-timescale connections to enable accurate inter-task classification.

631

Sub-sampling alone, may also produce spurious GC causality; the precise conditions under which spurious GC arises for continuous time vector autoregressive processes is an area of active research, and must be addressed in future studies [6,20].

635

Second, previous studies have shown that systematic differences in hemodynamic (HRF) lags (e.g. time to 636 onset, or time to peak) among brain regions may produce spurious dGC estimates [19,21]. With simulations 637 we demonstrated that fMRI-GC could identify differences in slow-timescale network connectivity, despite 638 systematic differences and heterogeneity in HRF onset latencies across nodes (SI Fig. S3D-E). In all 639 cases, applying recursive feature elimination with either dGC or iGC features identified the precise subset 640 of connections that distinguished distinct network configurations. In a majority of cases, dGC also correctly 641 identified the directionality of these connections. In our simulations, the only scenario in which dGC 642 features failed to identify the directionality of connections correctly, was when the onset latency in the 643 "destination" nodes were biased to be systematically earlier than those in the "source" nodes. Nevertheless. 644 in the real data it is unlikely that systematic inter-regional HRF differences were responsible for the 645 observed superior classification accuracies. Variations in HRF delays would indeed confound dGC 646 connectivity estimates - if they occurred consistently between brain regions across subjects and tasks (e.g. 647 SI Fig. S3D, red curves). Yet, such a scenario cannot account for the high classification accuracies among 648 tasks and sub-tasks based on dGC connectivity alone. In other words, even if HRF latency differences 649 systematically biased dGC connectivity estimates, these estimates were sufficiently and reliably different 650 across task cognitive states to permit accurate classification among them. Finally, network properties of key 651 regions identified with fMRI-GC were consistent with their known functional properties of these regions. For 652 instance, dGC identified the visuospatial network as an information outflow hub, across all six cognitive 653 tasks (Fig. 4D left). The visuospatial network comprises frontal cortex regions, including the frontal eve 654 field, as well as posterior parietal cortex, which are both widely implicated in visuospatial attention control 655 [48–51]. In addition, the only network that provided task-generic incoming connections to the visuospatial 656 network was the anterior salience network comprising the anterior fronto-insular cortex and the anterior 657 cingulate cortex [52,53], regions implicated in feature-based attention and the suppression of distractors 658 659 [54]. Information outflow from these key networks identified by dGC is consistent with their role in attention and executive control. 660

661

662 Third, simulations and theoretical results indicate that scanner noise can degrade, or even obliterate GC 663 connectivity estimates [19]. On the other hand, our classification accuracies suggest that GC estimates

were sufficiently robust to scanner noise to permit accurate task and sub-task classification in these data. In fact, we show that averaging dGC connectivity across as few as 5 subjects' data improves classification accuracy to over 95%, for nearly all tasks (Fig. 1D). Such superlative classification accuracies are unlikely to have occurred if scanner noise were to significantly degrade GC estimates.

668

In sum, these results strongly indicate that slow functional interactions in the brain can be meaningfully 669 inferred with GC from fMRI data. While the directionality of interactions measured by GC may need to be 670 interpreted with care [11,21], our results suggest that fMRI-GC may be useful for formulating hypothesis 671 about the role of particular brain regions in providing "top-down" control signals, for modulating activity in 672 other brain regions [7.8], as well as for investigating the nature of information flow in cortical microcircuits 673 with slow sampling rate techniques, such as calcium imaging [55]. The causal role of these brain regions in 674 behavior can then be directly tested with interventional approaches such as transcranial magnetic 675 stimulation, optogenetic inactivation or by examining patient populations with lesions in specific brain 676 regions [56]. Such a systematic analysis will pave the way for a mechanistic understanding of how flexible 677 functional interactions among brain regions mediate complex cognitive behaviors. 678

679

681 Materials and Methods

682 Ethics statement.

The scanning protocol for the Human Connectome Project was approved by the Human Research Protection Office at Washington University at St. Louis' (IRB *#* 201204036). Only de-identified, publicly released data were used in this study.

686

687 **Data and code availability statement.**

Data used in the study is available in the public domain at the Human Connectome Project database (https://db.humanconnectome.org/). Code used for analyses are available at the following link:

690 https://figshare.com/s/9d9131a6780fc8197cf1

Data sharing permissions can be found at the HCP website. Code may be shared or re-used upon requesting the corresponding author. These data and code sharing policies comply with the requirements of all funding agencies supporting this research and comply with institutional ethics protocols.

694

695 **fMRI data, parcellation and time-series extraction.**

We analyzed minimally preprocessed brain scans of 1000 subjects, drawn from the Human Connectome 696 Project (HCP) database (S1200 release) (age range: 22-35 years; 527 females); fMRI acquisition and 697 preprocessing details are described elsewhere [23,25]. SI Table S2 shows the identifiers of the subjects 698 from whom data were analyzed. Data were analyzed from resting state and seven other task conditions (SI 699 Table S1): Emotion processing, Gambling, Language, Motor, Relational processing, Social cognition and 700 Working memory; in most figures, these tasks are referred to with their initial letters. fMRI scans for the 701 relational task were not available for 9/1000 subjects; therefore, we analyzed a total of 7991 scans across 702 all tasks and subjects. 703

704

We employed five different brain parcellations based one anatomical atlas and four functional atlases (SI Table S3). For the tasks versus resting-state classification based on GC connectivity (first section of Results), all 5 parcellations were used. Based on the classification performance in this analysis, we picked the three parcellations with the highest accuracies (90 node and 14 network parcellations of [26] and 96

network parcellation of [57]) and these were used for the pairwise classification analysis of each task versus 709 the other as well as the n-way task classification analyses. Analysis with averaging GC features across 710 subjects (Fig. 1D) was performed with a 90 node parcellation [26]. Classification analyses with data purged 711 of instantaneous correlations and unweighted digraph representations (second section of Results) were 712 performed with the Shirer et al [26] 14 network parcellations. Analyses involving identifying task-generic 713 and task-discriminative networks, as well as behavioral score predictions, based on GC features (last 714 section of the Results) were performed with the Shirer et al [26] 14 network parcellation. Voxel time series 715 were extracted using Matlab and SPM 8 [58], and regional and network time series were computed by 716 averaging the time series across all voxels in the respective region or network. 717

We employed parcellations with fewer, more coarse-grained regions, rather than fine-grained parcellations because Granger Causality estimates were more reliable when the number of regions was fewer than the number of timepoints. Both task and resting scans were of sufficient duration (~200-300 volumes) to permit robust GC estimation. Finally, we noticed that in some parcellations, there were overlapping voxels between some of the regions. To avoid mixing of signals, we assigned each overlapping voxel to the region to whose centroid it was closest, based on Euclidean distance.

724

725 Estimating functional connectivity with GC.

We modeled instantaneous and lag-based functional connectivity between brain regions using conditional 726 Granger-Geweke Causality [9]. The linear relationship between two multivariate signals x and y conditioned 727 on a third multivariate signal z can be measured as the sum of linear feedback from x to y (Fx \rightarrow y|z), linear 728 feedback from y to x (Fy \rightarrow x|z), and instantaneous linear feedback (Fx \circ y|z) [9,41]. To quantify these linear 729 relationships, we model the future of each time series in terms of their past values, using multivariate 730 autoregressive (MVAR) modeling (SI Mathematical Note, Section S1, equation 1). MVAR model order was 731 732 determined with the Akaike Information Criterion (AIC) for each subject, and was typically 1. The MVAR model fit was used to estimate both an instantaneous connectivity matrix using iGC (Fxoylz) and a lag-733 based connectivity matrix using dGC ($Fx \rightarrow y|z$). Details are provided in SI Mathematical Note, Section S1. 734

Briefly, $Fx \rightarrow y|z$ is a measure of the improvement in the ability to predict the future values of **y** given the 735 past values of x, over and above what can be predicted from the past values of z and y, itself (and vice 736 versa for Fy \rightarrow x|z). Fx \circ y|z, on the other hand, measures the instantaneous influence between x and y 737 conditioned on z (see SI Mathematical Note, Section S1). We refer to Fx°vlz, as *instantaneous* GC (iGC). 738 and $Fx \rightarrow y|z$ and $Fy \rightarrow x|z$ as lag-based GC or *directed* GC (dGC), with the direction of the influence (x to y 739 or vice versa) being indicated by the arrow. The "full" measure of linear dependence and feedback Fx,y|z is 740 given by: $Fx,y|z = Fx \rightarrow y|z + Fy \rightarrow x|z + Fx \cdot y|z$. Fx,y|z measures the complete conditional linear dependence 741 between two time series. If, at a given instant, no aspect of one time series can be explained by a linear 742 model containing all the values (past and present) of the other, Fx,y|z will evaluate to zero [41]. 743

744

745 Classification with linear SVM based on GC connectivity.

The connection strengths of the estimated GC functional connectivity matrices were used as feature vectors with a linear classifier based on SVM for high dimensional predictor data. For a parcellation with n ROIs, the number of features for iGC-based classification was n(n-1)/2 (upper triangular portion of the symmetric n×n iGC matrix) and for dGC-based classification it was n^2-n (all entries of the n×n dGC matrix, excluding self-connections on the main diagonal).Based on these functional connectivity features, we asked if we could reliably distinguish each task condition from resting state (e.g. language versus resting) or each task condition from the other

753

For pairwise classification of resting state scans versus each task we used Matlab's fitclinear function, 754 optimizing hyperparameters using a 5-fold approach: by estimating hyperparameters with five sets of 200 755 subjects in turn, and measuring classification accuracies with the remaining 800 subjects. Classification 756 performance was assessed with leave-one-out and 10-fold cross-validation. We also assessed the 757 significance of the classification accuracy with permutation testing (see Methods). In simulations, we 758 observed that the magnitude of GC estimates varied based on the number of timepoints used in the 759 estimation. To prevent this difference in number of timepoints from biasing classification performance, each 760 scan was truncated to a common minimum number of time samples across the respective scans being 761 classified (task, resting) before estimating GC. For each subject, GC connectivity was estimated 762 independently for the two scan runs (left-to-right and right-to-left phase encoding runs), and averaged 763

across the runs. Hyperparameters optimized included the regularization parameter, regularization method 764 765 (ridge/lasso) and the learner (linear regression model. svm/logistic) using the OptimizeHyperparameters option to the fitclinear function. Hyperparameter optimization was 766 performed only for task vs. rest classifications, but not for subject feature averaging, task vs. task, or N-way 767 classification analyses. 768

769

For pairwise classification of each task versus the other, default hyperparameters were used in the 770 fitclinear function and classification performance was assessed with leave-one-out cross-validation. 771 For n-way classification, we used MATLAB's fitcecoc function, which is based on error-correcting output 772 codes, and fits multiclass models for SVMs. Briefly, the function implemented a one-vs-all coding design. 773 for which seven (number of classes in multiclass classification) binary learners were trained. For each 774 775 binary learner, one class was assigned a positive label and the rest were assigned negative labels. This design exhausts all combinations of positive class assignments. Classification performance in n-way 776 classification was assessed with leave-one-out cross-validation. For each classification analysis mentioned 777 above, task scans were truncated to the common minimum number of time samples across each set of 778 779 scans, before estimating GC.

780

781 Classification based on GC connectivity across sub-tasks and with sub-sampled data.

Tasks in the HCP data were run as a block design, alternating between various conditions (sub-tasks). We 782 tested whether GC connectivity would be able to classify among sub-tasks within each task (SI Table S1B). 783 784 fMRI time series corresponding to each sub task was obtained by concatenating blocks of fMRI task time series pertaining to the respective sub task; the temporal order across blocks was preserved while 785 concatenating the data. We also ensured that data at the conjunction of two successive blocks, which 786 represented non-contiguous time points, were not used for GC estimation. The two sub tasks to be 787 classified were then truncated to have same number of time points. GC estimation and pair-wise 788 classification across sub-tasks was performed with the procedure described in the previous section. The 789 Shirer et al [26] 14-network parcellation was used for these analyses. For the motor task, time series for the 790 791 left and right finger movement blocks were combined into a "hand" movement sub-task, and left and right toe movement blocks were combined into a "foot" movement sub-task. 792

793

We also tested whether GC on fMRI data sampled at slower rates would suffice to classify among task and 794 resting states. We obtained time series downsampled at 2x the original sampling interval by removing data 795 at even numbered sample points, and retaining data at odd numbered sample points (k=1, 3, 5...). The 796 797 even-sample point data were appended the end of odd-sample data series, thereby retaining the overall number of data points in the original time series. Again, we ensured that data at the conjunction of the odd-798 and even-sampled data series (last odd-sampled point and first even sampled point), which represented 799 non-contiguous data points, were not used for GC estimation. Similarly, we obtained time series 800 downsampled at 3x the original sampling interval by removing every third data point, starting with the 801 second or third data point, and concatenating these timeseries to retain the overall number of data points in 802 the original timeseries. As before, GC estimation and pair-wise classification was performed with the 803 procedure described in the previous section 804

805

806 **Permutation testing of classifier accuracies.**

We performed permutation tests for evaluating the statistical significance of classifier performance, using 807 the method outlined in [59]. The test involved permuting task labels independently for each subject and 808 809 computing a null distribution of 10-fold cross-validation accuracy. We employed 1000 surrogates and assessed significance of each empirically estimated 10-fold cross-validation accuracy values for dGC and 810 iGC, based on the proportion of samples in the null distribution which were greater than the cross-validation 811 accuracy estimated from the data. We conducted these analyses for the tasks versus resting state 812 classifications, n-way task classification, classification analyses after purging instantaneous correlations 813 and those based on digraph features, separately for the two metrics (dGC and iGC). 814

815

816 Testing for data stationarity and goodness of MVAR model fit.

Computing GC based on VAR modeling assumes that the timeseries represent a stationary process. Four different tests were performed to test whether the MVAR model provided a valid and adequate fit to the data (SI Table S5). We performed these tests for parcellated time-series using scripts provided in the Multivariate Granger Causality (MVGC) toolbox [60]. First, we checked for the stability of the MVAR model fit by computing logarithm of the spectral radius using the *var specrad()* function. A negative value was

taken to indicate a stable fit. Second, we assessed consistency of the model fit, which quantifies what 822 proportion of the correlation structure in data is accounted for by the VAR model, using the consistency() 823 function. We adopted a threshold of 80% (or above) for both task and resting timeseries to consider the 824 data to have passed the test for consistency [60]. Third, we evaluated the whiteness of residuals based on 825 the Durbin-Watson test for absence of serial correlation of VAR residuals, using the whiteness() function. 826 Values of the Durbin-Watson statistic less than 1 or greater than 3 signify a strong positive or negative 827 correlation, respectively among the residuals [60]. Subjects for whom the Durbin-Watson statistic lay 828 between 1 and 3 for more than 90% of the regional timeseries, for both task and resting state data, were 829 considered to have passed the test. Fourth, we checked for stationarity based on the augmented Dicky-830 Fuller unit-root test (ADF), using the mvgc adf() function. As in the previous case, subjects for whom the 831 ADF test statistic was less than its critical value for more than 90% of the regional timeseries, for both task 832 and resting state data, were considered to have passed the test. 833

834

835 Control for motion artifacts.

We checked whether systematic differences in motion artifacts could contribute to the superlative 836 classification accuracies observed with GC. For this, we calculated Frame-wise Displacement (FD) [30] as 837 the sum of temporal derivatives of translational and rotational displacement along the three (x,y,z) axes in 838 mm, with the estimated motion parameters provided by HCP. Frames with FD>0.5mm were considered 839 "misaligned" and were discarded ("scrubbed") while estimating GC values. Because dGC is estimated 840 based on lagged correlations, we also discarded one frame before and after every misaligned frame (AR 841 model order was typically 1 for these data). We then repeated the SVM-based two-way classification of 842 resting state from the seven different task states, with GC features estimated on the "motion scrubbed" 843 data; we also repeated n-way classification among the 7 tasks. Comparison of classification (cross-844 validated) accuracies with and without motion scrubbing, across all 1000 subjects, is shown in SI Fig. S2C. 845

846 847

848 Functional connectivity estimation and classification with partial correlations.

We compared the performance of classification based on GC measures with that based on partial correlations (PC). Partial correlations were computed based on the inverse of the covariance matrix as

outlined previously [4,5]. Like iGC, the PC connectivity matrix is undirected and symmetric. Therefore, only the upper triangular portion of the matrix, including $(n^{*}(n-1)/2)$ PC weights, was used as features in the classification analyses. Classification and cross-validation analyses followed the procedures described in the Methods section on *"Classification with linear support vector machines based on GC connectivity"*.

855

PC connectivity performed consistently better than GC connectivity for classifying task from resting state (Fig. 2A). We propose the following analytical explanation for this observation: PC, an estimator based on instantaneous covariance, is less susceptible to noise than GC, which is based on lagged covariance. This is due to the fact that the estimation of lagged-covariance is susceptible to errors from noise at multiple time-points. For illustration, consider a timeseries generated by a VAR(1) model: $\mathbf{x}(t) = A \mathbf{x}(t - 1) + \mathbf{e}(t)$. The lagged (lag-1) covariance matrix (Σ_1) is estimated from the data as:

862 E $[\mathbf{x}(t) \mathbf{x}(t-1)^T] = E [(A\mathbf{x}(t-1) + \mathbf{e}(t)) \mathbf{x}(t-1)^T] = A E[\mathbf{x}(t-1) \mathbf{x}(t-1)^T] + E [\mathbf{e}(t) \mathbf{x}(t-1)^T]$

Thus, when estimating the lagged covariance, the variance of the interaction term $E[e(t) \mathbf{x}(t-1)^T]$ (second term in the right hand side) contributes to the variance of Σ_1 in addition to the variance in computing the instantaneous covariance $E[\mathbf{x}(t-1) \mathbf{x}(t-1)^T]$ (first term on the right hand side).

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867 Classification based on GC connectivity in zero-lag correlation purged data.

To test the complementarity of information conveyed by GC functional connectivity versus functional connectivity based on instantaneous correlations we decorrelated the regional time series data to purge them of instantaneous correlations. We adopted two approaches for this purpose: i) zero-phase component analysis (ZCA) and ii) generalized eigenvalue decomposition (GEV).

872

i) Zero-phase component analysis (ZCA). Consider demeaned *t*×*r* data matrix **X** of regional timeseries with *t* timepoints and *r* regions, with covariance matrix **C**. Decorrelating the data, to remove correlations among the columns of **X**, is achieved with a whitening transformation. A common whitening transformation is based on principal components analysis (PCA): **Y** = $W_{PCA}X$, with $W_{PCA} = D^{-1/2}E^{T}$ where **D** is a diagonal matrix, with the eigenvalues of **C** on its diagonals, and the columns of **E** contain the eigenvectors of **C**.

While the PCA transformation effectively decorrelates regional timeseries, there is no way to ensure one-toone correspondence of the whitened dimensions across subjects, rendering subsequent classification analysis challenging. Consequently, here we chose a different whitening transformation based on zerophase component analysis (ZCA), also known as the Mahalanobis transformation. Based on this transformation, whitening is achieved as: **Y** = **W**_{ZCA}**X**, with **W**_{ZCA} = **ED**^{-1/2}**E**^T = **C**^{-1/2}. A particular advantage

of the ZCA transformation is that it yields whitened data that is as close as possible to the original data, in a 883 884 least-squares sense [61]. Therefore, each subject's data is projected on to a set of dimensions are most closely aligned with the underlying regional timeseries dimensions. Because the regions exhibit spatial 885 correspondence across subjects (due to fMRI spatial normalization), the ZCA dimensions possess a 886 natural, one-to-one correspondence across subjects, permitting subsequent classification. Before 887 888 classification analysis ZCA dimensions were identified for each subject, separately for task and resting datasets. Regional time series for task and resting data were independently decorrelated by projecting onto 889 their respective ZCA dimensions. GC (and PC) functional connectivity was estimated based on the these 890 decorrelated timeseries, followed by classification analysis, as described previously (Methods section on 891 892 "Classification with linear support vector machines based on GC connectivity"). As proof that the ZCA transformation was working effectively, classification accuracy based on PC (an instantaneous correlation 893 measure) computed from ZCA components was at chance across all tasks (Fig. 2C top). 894

895

ii) Generalized Eigenvalue Decomposition (GEV). Although ZCA effectively purged correlations from the 896 data, for the subsequent classification analyses task and resting state data were projected onto different, 897 respective ZCA dimensions. Thus, the above-chance task versus resting state classification accuracy with 898 GC features derived from ZCA components (Fig. 2C top) could perhaps be explained by, for example, 899 900 systematic differences with how reliably ZCA dimensions were estimated across task and resting-state scans. We therefore sought an approach that could project both task and resting data into the same 901 dimension while simultaneously decorrelating both. Such joint decorrelation may be achieved by projecting 902 the data on to the generalized eigenvectors of the covariance matrices of the two datasets [62]. Let C_T and 903 C_R denote the covariance matrices of the task and resting datasets respectively. The generalized 904 eigenvectors of these two symmetric matrices are given by the columns of $G = E_T D_T^{-1/2} E_R$, where, as 905

before D_T is a diagonal matrix, with the eigenvalues of C_T on its diagonals, and the columns of E_R and E_T 906 contain the eigenvectors of C_R and C_T respectively. It can be readily verified that $G^T C_T G$ and $G^T C_R G$ are 907 both diagonal matrices. Therefore, G is a matrix that jointly diagonalizes both C_T and C_R and projecting 908 either task or resting state data into the columns of G decorrelates the respective timeseries. So, for these 909 analyses, the regional time series for the task and resting state conditions for each subject were jointly 910 decorrelated by projecting them onto a single space spanned the generalized eigenvectors. This was 911 followed by classification analysis with GC features obtained from the decorrelated time series. As before, 912 we confirmed the effectiveness of the decorrelation by computing classification accuracy based on PC from 913 GEV components, which was at chance across all tasks (Fig. 2C bottom). 914

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916 Classification based on unweighted digraph representations of GC connectivity.

An unweighted directed graph (digraph) network representation shows the dominant direction (but not 917 magnitude) of functional connectivity among brain regions. Obtaining significant directed connections with 918 dGC is challenging due to number of multiple comparisons required for testing n²-n connections. To identify 919 significant directed connections, overcoming the multiple comparisons problem, we first subtracted the dGC 920 connectivity matrix from its transpose and then applied the following two-stage procedure. In the first stage, 921 the 1000 subjects were divided into five folds. For each two-way task versus resting state classification, 922 recursive feature elimination (RFE, described in a later section titled "GC feature selection based on 923 Recursive Feature Elimination") was performed based on dGC features of subjects from one fold (i.e. with 924 200 subjects). A minimal set of connection features identified by RFE, and their corresponding symmetric 925 counterparts were then employed in the subsequent analyses; we term these connections K; the cardinality 926 of K (the number of significant connections) was typically in the range of 2 - 86 (2.5th - 97.5th percentile). In 927 the second stage, we identified statistically significant connections among these K features alone. For each 928 of the subjects in the four remaining folds (i.e. 800 subjects), a null distribution for the dGC values of the 929 930 features in K was obtained by estimating dGC following phase-scrambling the time series [8]. Next, we identified significant connections based on dGC values that occurred at the tail of the null distribution: the 931 threshold for significant connections was determined based on a p-value of 0.05 with a Bonferroni 932 correction for multiple comparisons. Classification performance based on digraph features was assessed 933 934 with leave-one-out cross-validation.

935

936 GC connectivity in simulated fMRI time series.

To test the ability of GC measures to reliably recover functional interactions at different timescales, we simulated fMRI time series for model networks. Simulated fMRI time series were generated using a twostage model. The first stage involved modeling latent neural dynamics with a stochastic, linear vector differential equation given by:

where **r** is the multivariate neural state variable representing the state of each neuron (or node) in the 942 network (an N×1 vector, with N being the number of neurons), dr/dt is its temporal derivative, W is the 943 neural ("ground truth") connectivity matrix (dimension N×N), τ is the time constant of each neuron (or node) 944 and $\boldsymbol{\varepsilon}$ is i.i.d Gaussian noise (N(0, Σ)), with $\Sigma = I_N$ (N×N identity matrix). Although this model does not 945 explicitly incorporate signal propagation delays, such vector Ornstein-Uhlenbeck models rank, arguably, 946 among the most common models employed for simulating neural and fMRI time series, in many previous 947 studies [6,18,19]. The multivariate time series r(t), sampled at discrete time points $r(k\Delta)$ with a sampling rate 948 of Δ , were generated based on the discrete time (1-lag) connectivity matrix A(Δ) and a residual noise 949 950 intensity $\Sigma(\Delta)$. Here:

951
$$A(\Delta) = e^{\Delta A}; \qquad \Sigma(\Delta) = (1/\Delta) (\Gamma(0) - e^{\Delta A} \Gamma(0) e^{\Delta A'})$$

where $A = (1/\tau) (W - I_N)$, e^A denotes the matrix exponential, A' is the transpose of A, and $\Gamma(0)$ is the zero 952 lag autocovariance which satisfies the continuous time Lyapunov equation $A\Gamma(0)+\Gamma(0)A'+\Sigma=0$ [19]. In the 953 second stage, the latent neural dynamics were convolved with the hemodynamic response function (HRF) 954 to obtain the simulated fMRI time series: $\mathbf{y} = H \otimes \mathbf{x}$, where H is the canonical hemodynamic response 955 function (hrf; simulated with spm hrf in SPM8), \otimes is the convolution operation and y is the simulated fMRI 956 time series. Finally, following convolution with the hrf, the data were downsampled to 750 ms, to mimic the 957 repeat time (TR) of the HCP fMRI scans used in this study. The same model was used for the different 958 959 simulations used in the manuscript (third section of the Results). The parameters for the 2-node 960 simulations, and for the 9-node (100 neurons per node) simulations are described in SI Table S6.

For the 2-node simulations, iGC and dGC values were estimated by simulating the network for 200 962 timepoints, averaged across 25 repetitions. The 9-node simulations were performed with a 900 neuron 963 network, with 100 neurons per node. Each node had sparse, random excitatory/inhibitory connectivity 964 among its neurons (parameters in Table S6), whereas only 5% of neurons in each node were involved in 965 inter-node connections, to mimic sparse, long-range connectivity in the neocortex [63]. The network was 966 simulated for 200 timepoints, and timeseries from all (100) neurons in each node were averaged to 967 generate 9 node timeseries. iGC and dGC values were estimated from the node timeseries and averaged 968 across 10 independent repetitions. Significance was assessed with a bootstrap approach that involved 969 generating 1000 surrogates by phase scrambling the node timeseries to yield a null distribution of GC 970 values [8], followed by a Benjamini-Hochberg correction for multiple comparisons. 971

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Simulations comparing PC and iGC connectivity were performed as follows: We simulated a 7-node 973 network with a 1-lag VAR model of the form: $\mathbf{X}_{k} = A \mathbf{X}_{k-1} + \mathbf{\varepsilon}_{k}$ where \mathbf{X}_{k} is the state of the discrete time 974 process at discrete timestep 'k', A is the connectivity matrix, and ε is Gaussian noise with covariance matrix 975 Σ_d . A was chosen to be a random matrix with spectral radius less than 1 to ensure stability. Σ was chosen 976 such that the covariance between every pair of residuals was zero (independent residuals) except for the 977 first two residuals. The correlation between these residuals, ε^1 and ε^2 , was parametrically varied between -978 1.0 and 1.0 to systematically vary the strength of iGC connectivity. Note that, under this model, iGC 979 between X¹ and X² vanishes only if and only if ε^1 and ε^2 are uncorrelated [9]. 980

981

982 GC feature selection based on Recursive Feature Elimination (RFE).

We performed features selection for analyses reported in Fig. 2D, 4B,C and S4B based on Recursive 983 Feature Elimination (RFE). RFE identifies a minimal set of features, which provide maximal cross-validation 984 accuracy [38]. Here, we implemented a two-level algorithm, described previously [16,64]. First, the data 985 were divided into N₁ (here, 10) folds. Of these, N₁-1 folds were used as "training" data, and one fold was 986 reserved as "test" data for quantifying the generalization performance of the classifier. Training data were 987 pooled and further divided into N₂ (here, 5) folds. The SVM classifier was then trained on N₂-1 folds 988 (leaving out one fold) and discriminative weights were obtained. The above procedure was repeated N₂ 989 times by leaving out each fold, in turn. Average weights were then computed by averaging the absolute 990

values of the discriminative weights across the N₂ runs. Next, 10% of the features (connections) 991 contributing the lowest average weights were discarded, and the classifier was trained again with only the 992 retained set of features. This procedure of feature selection and training was repeated until no more 993 features remained. At this stage, the generalization performance for every set of retained features (each 994 "RFE level") was assessed using the left out "test" data. The entire procedure was repeated N1 times by 995 leaving out each fold of the original data, in turn, as test data. Final generalization performances and 996 discriminative weights of each RFE level were obtained as the average over N1 folds. We selected the set 997 of connections at the RFE level at which the generalization performance reached an "elbow": a minimal set 998 of connections at which generalization performance dipped dramatically below its maximal level. To identify 999 this elbow (e), we used a custom elbow fitting procedure, requiring a piecewise linear fit to the RFE curve, 000 based on two lines, one for "x>e" and another for "x<=e", with the first line required to have a higher slope 001 than the second. The first point in each RFE curve was excluded from the higher slope line fit (Fig. 4C, 4E, 002 003 SI Fig. S4B). RFE was typically repeated 5 times before determining peak accuracy and corresponding features. 004

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006 Simulating hemodynamic lag variations across nodes.

007 We simulated systematic differences in hemodynamic lags across nodes by varying the onset parameter of the spm hrf function (SPM8; [58]). For network configurations A and B described in Figure 4A, we 800 simulated 4 scenarios: a) same mean HRF onset (μ_L = 3s) across nodes; b) source node HRF onset lagging 009 the destination node by 1s ($\mu_{1-src} > \mu_{1-dst}$); c) source node HRF onset leading destination node by 1s ($\mu_{1-src} > \mu_{1-dst}$); 010 µ_{L-dst}); and d) mixed latencies of lead and lag across source and destination nodes (see next). GC was 011 estimated for 100 simulated participants, by sampling onset latencies for each of the 6 nodes (A-F) from 012 normal distributions (truncated to have only positive latency values), over a range of different standard 013 deviations (σ_1 =0-1s, in steps of 0.2s). Onset latencies were sampled independently across participants, but 014 015 were sampled such that the relative latency between each pair of source and destination nodes, across corresponding network configurations, remained the same for each participant. For example, if the onset 016 latency difference between nodes A and B was 0.7s ($\mu_{L-B}-\mu_{L-A}=0.7s$) for a particular subject, the same 017 difference in onset latency was also maintained between nodes B and C ($\mu_{I-C}-\mu_{I-B}=0.7s$). For simulations 018 019 with mixed latencies (case d), 50% of simulated participants had onset latencies drawn from distributions

with the source node lagging the destination node (case b) and the remaining 50% with the source node leading the destination node (case c). GC values were averaged over 5 runs for each simulated participant. Finally, we performed RFE to identify key connections that distinguished the two network configurations (same procedure as in Fig. 4B). Connections weights of the most discriminative connections following RFE are shown in SI Fig. S3E (for $\sigma_L=0.4$ s). Difference of dGC connections strengths as well as iGC connection strengths, for various values of σ_L , are shown in SI Fig. S3D.

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028 Identifying "task-generic" and "task-discriminative" GC connections.

identify a minimal set of connections that occurred consistently across tasks ("task-generic" То 029 connections), we adopted the following approach. We performed RFE analysis for task versus resting state 030 classification for each of the six tasks (all tasks except motor); we expected each of these tasks to recruit 031 032 common cognitive control mechanisms. We then performed a binomial test to identify connections that were consistently activated across tasks. Briefly, the presence or absence of a connection in the set of RFE 033 features for a given task versus resting state classification was considered as a Bernoulli trial, with 034 probability of success (its presence) p being the mean number of RFE features identified across all six 035 classifications. The number of trials n was the number of tasks versus resting state classifications (here 036 n=6). The probability of a randomly picked connection being present in more than k such RFE sets is given 037 by the cumulative distribution function for the binomial distribution F(k; n, p). Significant connections were 038 identified as those that occurred in k or more tasks, with threshold at the p=0.05 level. 039

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To identify a minimal set of connections that maximally differed across tasks ("task-discriminative" connections), we used RFE with an n-way classifier, to classify among all six tasks (again, except the motor task). The n-way classifier is based on training *n* (here, 6) one-vs-all binary learners. At the second level of the RFE procedure described above, average weights were computed for each of these *n* binary learners by averaging the absolute values of the discriminative weights across the N₂ runs. Next, a set of features obtained by taking union of 1% of the features (connections) contributing the lowest average weights in each learner was discarded, and the classifier was trained again with only the retained set of features.

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While guantifying the overlap between task-generic and task-discriminating connections identified 049 separately for dGC, iGC and PC, we converted the dGC matrix to a lower triangular matrix by reflecting all 050 connections about the main diagonal. The degree of overlap between PC and GC connections was 051 quantified as the number of overlapping connections as proportion of the total number of connections 052 identified by PC. We then computed a null distribution of the degree of overlap by randomly permuting the 053 connection identities within each matrix, while preserving the overall number of connections in each matrix, 054 generating 1000 surrogate samples. The significance of the overlap of task-generic or task-055 and discriminating connections between each pair of metrics (PC-dGC or PC-iGC) was quantified as the 056 fraction of overlapping connections in the data that exceeded this null distribution. 057

058

059 Predicting behavioral scores based on GC connectivity

We asked whether inter-individual differences in GC connectivity would be relevant for predicting inter-060 individual differences in behavioral scores. HCP provides a well-validated battery of behavioral scores 061 assessed with a wide range of cognitive tasks. The task battery is based on the NIH Toolbox for 062 Assessment of Neurological and Behavioral function [65], developed to create a uniform set of measures 063 for rapid data collection in large cohorts. The toolbox includes assessments of cognitive, emotional, motor 064 and sensory processing scores in healthy individuals. We pre-selected, based on domain knowledge, a 065 specific subset of 51 scores for these analyses, using age-adjusted scores, wherever available (listed in SI 066 Table S7). Next, we sought to predict subjects' behavioral scores based on GC connectivity with an 067 068 established leave-one-out approach [66]. Briefly, we used linear regression to predict behavioral scores using, as features, GC estimates of functional connectivity, separately for iGC (91 features or connections) 069 and dGC (182 features). The leave-one-out analysis was performed such that the support vector regressor 070 was fit on all but one subject and the learned beta weights were used to obtain predictions of the left-out 071 subject's behavioral score, using that subject's own GC connectivity weights. Predicted scores were 072 073 correlated with the actual scores using robust correlations ("percentage-bend" correlations; [67]).

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Next, we asked if GC connectivity could identify an individual based on a composite marker of her/his behavioral scores. Because 40 subjects did not have a full complement of behavioral scores, data from the remaining 960 subjects was included in this analysis. The 51 behavioral scores were, each, z-scored

across subjects and formatted into a "composite behavioral score" vector. This vector served as an 078 individual specific composite marker of behavioral scores, as revealed the weak off-diagonal values in the 079 covariance matrix of this vector across subjects (Fig. 5D top). dGC and iGC features of individual tasks, as 080 well as combination of tasks (Relational and Working memory), were used to then predict the composite 081 score marker for individual subjects, using the same leave-one-out procedure as described above. The 082 observed and predicted set of composite scores was correlated across subjects. The distribution of 083 observed versus predicted correlation values for each subject (values on main diagonal; Fig. 5D bottom 084 yellow) were compared against between-subject correlation values (off-diagonal values; Fig. 5D bottom 085 grey) using a Kolmogorov-Smirnov test. 086

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234 Supporting Information

- 235 SI Text. Results and SI Figure Legends
- 236 SI Figure S1. GC classification accuracies for different parcellations and alternative classifiers
- 237 SI Figure S2. Stationarity tests and 1-stage versus 2-stage GC
- 238 SI Figure S3. Relationship between network connectivity, GC and partial correlations
- 239 SI Figure S4. Task generic and discriminative connections based on partial correlations (PC)
- 240 SI Figure S5. Behavioral score predictions based on GC connectivity strengths
- 241 SI Table S1A. Task descriptions.
- 242 SI Table S1B. Description of sub-tasks
- 243 SI Table S2. Subject identifiers
- 244 SI Table S3. Parcellations used in the analysis
- 245 SI Table S4. Network labels in the Shirer et al (2012) 14-network parcellation
- 246 SI Table S5. Number of subjects passing stationarity tests
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- 248 SI Table S7. Behavioral scores and descriptions
- 249 SI Text. Mathematical Note













