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Novel machine learning approaches for improving the reproducibility and reliability of functional and effective connectivity from functional MRI

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Abstract

Objective. New measures of human brain connectivity are needed to address gaps in the existing measures and facilitate the study of brain function, cognitive capacity, and identify early markers of human disease. Traditional approaches to measure functional connectivity (FC) between pairs of brain regions in functional MRI, such as correlation and partial correlation, fail to capture nonlinear aspects in the regional associations. We propose a new machine learning based measure of FC (ML.FC) which efficiently captures linear and nonlinear aspects. Approach. To capture directed information flow between brain regions, effective connectivity (EC) metrics, including dynamic causal modeling and structural equation modeling have been used. However, these methods are impractical to compute across the many regions of the whole brain. Therefore, we propose two new EC measures. The first, a machine learning based measure of effective connectivity (ML.EC), measures nonlinear aspects across the entire brain. The second, Structurally Projected Granger Causality (SP.GC) adapts Granger Causal connectivity to efficiently characterize and regularize the whole brain EC connectome to respect underlying biological structural connectivity. The proposed measures are compared to traditional measures in terms of reproducibility and the ability to predict individual traits in order to demonstrate these measures’ internal validity. We use four repeat scans of the same individuals from the Human Connectome Project and measure the ability of the measures to predict individual subject physiologic and cognitive traits.

Main results. The proposed new FC measure of ML.FC attains high reproducibility (mean intra-subject $R^2$ of 0.44), while the proposed EC measure of SP.GC attains the highest predictive power (mean $R^2$ across prediction tasks of 0.66). Significance. The proposed methods are highly suitable for achieving high reproducibility and predictiveness and demonstrate their strong potential for future neuroimaging studies.

1. Introduction

The connectivity of the human brain is integral to cognitive capacity, can be an early marker for human disease, and underlies the fundamental functioning of the central nervous system (Ashburner et al 2004). However, measuring connectivity in vivo has proven problematic (Rowe 2010, Fiecas et al 2013, Andellini et al 2015). Functional magnetic resonance imaging (fMRI) of the brain measures the blood-oxygen-level-dependent (BOLD) signal and serves as an indirect measure of neural activity. The brain scan
can be parcellated into neuroanatomical regions and the mean regional time series can be computed from the voxels in each region. By measuring temporal relationships between the mean BOLD signal from two or more regions of the brain, the underlying direct and indirect connectivity and communication within the brain can be probed. The connections between regions can then be used to represent the subject-specific connectome as a connectivity graph with each region represented as a node in the graph, while the edges between nodes are assigned an edge strength proportion to the pairwise regional connectivity.

Connectivity measures are calculated from MRI using a measure of similarity or information transfer between the mean regional BOLD timeseries of a pair of regions. Connectivity metrics can be grouped into undirected functional connectivity (FC) metrics and directed effective connectivity (EC) metrics. Functional connectivity is defined as the temporal coincidence of spatially distant neurophysiological events (Ashburner et al 2004) and it has been used to characterize the human connectome in both health and disease (Cohen et al 2017, Smitha et al 2017). FC is traditionally calculated as the correlation or partial correlation between the regional timeseries. Meanwhile, effective connectivity is defined as the influence one neural system exerts over another (Ashburner et al 2004). Broadly, this is a model-dependent measure wherein the information transfer between mean regional timeseries is quantified from the goodness of fit of a model that predicts one of the timeseries from one or more of the other timeseries. Examples of EC measures include Granger causality (Granger 1969, Spencer et al 2018, Abidin et al 2019, Chockanathan et al 2019), dynamic causal modeling (Park et al 2018, Friston et al 2019), and structured equation modeling (SEM) (Rowe 2010), which have been widely deployed for connectome characterization. EC is inherently directional as it captures the direction of information flow over time (Bielczyk et al 2019). EC is model-dependent and requires more computation than FC but suppresses spurious indirect connections and identifies linkages that are potentially causal and not simply correlated.

Traditional FC and EC measures have several limitations, for which we propose new solutions. Common FC measures include Pearson’s r, partial correlation, and spectral Granger causality (Ding et al 2006). Each of these methods measure a degree of linear association between two mean regional timeseries; however, the actual relationship between mean regional brain activity is nonlinear (Friston et al 2019). Therefore, we propose nonlinear machine learning models that measure FC while capturing such nonlinearities. Of the different EC measures, we focus on Granger causal (GC) methods as they are data-driven approaches that can be used when many neuroanatomical regions, \( N \), are to be analyzed (e.g. \( N > 50 \)). In modern fMRI connectivity analysis \( N \) is often a hundred or more. Alternative causal models, including dynamic causal models and SEMs, typically apply an exhaustive search over possible connectivity patterns, making analysis at this ROI granularity intractable for current compute hardware. Limiting the connectivity to a subset of the brain, such as intra-DMN connectivity, is often used as a workaround, but this restricts the portion of the brain under consideration and can miss important interactions (Rowe 2010, Friston et al 2019). Granger causal methods have limitations as well but these, we hypothesize, are surmountable, including: model selection procedures, regularization, scalability (the traditional GC method requires fitting \( O(N^2) \) submodels where \( N \) is the number of regions under analysis), an inability to capture nonlinear interactions, and the absence of the incorporation of prior knowledge of brain architecture (Ashburner et al 2004). To address these limitations, we propose two measures of effective connectivity. The first measure which we call Machine-Learning FC (ML-FC), uses a nonlinear machine learning model to quantify nonlinear pairwise timeseries associations. Our method is more scalable because the number of required models to fit scales as \( O(N) \). Our second measure, which we call Structurally-Projected Granger Causality (SP-GC), reformulates Granger causal connectivity in two ways. First, we regularize the connectivity computation using a structural connectivity (SC) prior derived from diffusion MRI. Streamline tractography is performed on diffusion MRI from the Human Connectome Project (HCP) and a streamline atlas is generated (Yeh et al 2018). The log of the number of streamlines connecting regions is used as a measure of pairwise structural connectivity. This is used to regularize the functional interactions inferred between regional timeseries via a tradeoff between the raw functional data interactivity and fiber bundle connectivity. As actual neural communication occurs through physical connections, this constraint is a natural choice of a prior to guide brain FC (Allen and Weylandt 2019, Huang and Ding 2016, Dillon et al 2017, Manning et al 2018, Maglanoc et al 2020). The second way we reformulate Granger causal connectivity is to perform dimensionality reduction. Calculating the connectivity in a low dimensional space affords several advantages including: simplifying model optimization as there are fewer weights to tune and providing further regularization to stabilize fMRI interpretation and increase reproducibility. This dimensionality reduction is achieved by projecting the mean regional timeseries into a low dimensional space informed by the streamline SC prior. Each of our proposed measures is evaluated for reproducibility and the ability to predict cognitive and physiological traits of the HCP participants in our study.

A connectivity measure should produce a similar connectivity matrix for a given individual across
repeat fMRI scans that are acquired within a short window of time. Therefore, we evaluated the proposed FC and EC measures reproducibility across four repeated fMRI scans of each individual in our HCP-derived dataset. A reproducible measure better characterizes an individual’s connectivity fingerprint and is therefore more useful to capture true differences between individuals (Waller et al 2017, Noble et al 2019). Reproducibility is necessary, but insufficient to show that the proposed measures have validity; therefore we also measure the predictive power of each FC and EC metric in three relevant domains: a purely physiological domain predicting mean arterial pressure, a purely cognitive domain measuring fluid intelligence, and a combined physiologic and cognitive domain measuring stress. These were chosen as representative targets of interest of researchers and clinicians interested in predictions for physiology (e.g. stroke, aging), cognition (e.g. memory, PTSD), or a combination of the two (e.g. stress, neurodegeneration) for diagnoses and treatment. Measures that are both reproducible and have consistently high predictive power across multiple tasks are significantly more useful as candidate biomarkers (Termenon et al 2016, Waller et al 2017, Noble et al 2017a, 2017b). We postulate that a measure that is both more reproducible and predictive is a better representation of true underlying neural patterns than alternative measures. The contributions of this work are: (1) the development of a new functional connectivity metric (MLFC) and a new effective connectivity metric (MLEC) that efficiently capture nonlinear associations between brain regions, (2) the development of a new effective connectivity metric (SPGC) that incorporates a SC prior while efficiently measuring associations across all brain regions in a low dimensional space, (3) a quantitative comparison of the proposed measures to traditional measures of connectivity in terms of reproducibility and the power to predictive traits of individual subjects. Finally, (4) we recommend individual measures that hold the most potential to advance the study of human brain connectivity in health and disease based on the quantitative comparison.

2. Methods and materials

2.1. Methods

2.1.1. Proposed machine learning-based functional connectivity (MLFC measures)

Characterizing brain connectivity to better understand both health and disease is a complex process requiring measuring both linear and nonlinear aspects of information transfer between brain regions. Classical means of performing this characterization include the use of Pearson’s r, partial correlation, and spectral Granger causality. (For definitions of classical measures of FC, see supplemental section 9.1.2.) Central to this premise, we propose the construction of a machine learning model to calculate functional connectivity, an approach we denote as MLFC. This model predicts the activity, a, at a given node j by using the information present at all other nodes (brain regions) at any given time, t. As illustrated in equation (1), we use a nonlinear model M to predict the activity at region j at time t from all other regions under analysis R except region j

\[ a_{j,t} = M(\mathbf{\hat{d}}_{R \setminus j}, t) + \epsilon_{j,t}, \quad a = \text{activity}, \quad \epsilon = \text{error}. \]

This model simultaneously learns the association between all other nodes’ activity and the target node j. The weight assigned to each covariate quantifies the amount of information the model is using from that node to predict the target node j, which is a putative measure of the connectivity between each node i and j.

This draws on the theory of Granger causality which uses the coefficients of a bilinear model to quantify instantaneous information transfer (i.e. the relationship between signals at a fixed single time t) by predicting the activity of node j at time t, from other nodes, i, with a linear model (Ding et al 2006, Luo et al 2013).

For resting state fMRI, we want to derive a measure of functional connectivity between every set of nodes, resulting in a functional connectivity (FC) matrix. Our procedure using the covariate weights from the predictive model populates one row of the FC matrix at a time. If we repeat the process for each region, we fill the entire FC matrix by fitting N models. The choice of model M determines what associations we can detect between regions from the predicted covariate weights, which enables granular modeling control compared to previous attempts that use only one model (Murugesan et al 2020). In this work we allow M to be any of the following models: (1) the extremely random trees model (ERTs), (2) nonlinear radial basis function kernel support-vector machine regressor (SVM), (3) Extreme Gradient Boosting forest models (XGB) (Chen et al 2016). The ERT was chosen because it produces high performance across a wide domain of machine learning applications (Feczko et al 2018, Mellema et al 2022). The SVM was chosen because it is a high-performing machine learning model which has a more directly interpretable and explicit weights than the ERT (Deshpande et al 2010, Arora et al 2018, Mellema et al 2022). The XGB was chosen because it tends to have higher performance than the ERT, and handles multicollinearity from repeated data subsampling, which we hypothesize will better handle correlated regional information than the ERT.

For each proposed model, we use the following model fitting approach. First, the mean timeseries per region is standardized with a mean of 0 and unit variance. Then, a model is fit to predict regional activity at node j at every time t from other all other nodes i at each time t. Then, a measure of feature
Algorithm 1. GC algorithm. This algorithm describes the steps by which one calculates an effective connectivity matrix $E$ from a neural timeseries $X$ using a standard Granger-causal approach. $X_i$ = neural activity matrix of size $i$ by $t$, where $i \in [1, N]$ and where $t \in [1, T]$. $T$ = number of timepoints. $\tau$ = max lag, $f$ = timeseries predictor function. $j$ = secondary index from 1 to $N$. $E_{ij}$ = effective connectivity matrix of size $N \times N$, indexed by $i$ and $j$. $r$ = reduced timeseries predictor function without region $j$. $\sigma$ = standard deviation.

Inputs: $X_i$, ($i \in [1, N], t \in [1, T]$), $\tau$, $f$;
Output: $E$;

for $i$ := 1 to $N$ do Full model fit
   $X_i = f(X_{i-1}, X_{i-2}, ..., X_{i-\tau})$;

for $j$ := 1 to $N$ do Reduced model fit
   $X'_i = X_i \setminus j$;
   $X'_{ij} = r(X'_{i-1}, X'_{i-2}, ..., X'_{i-\tau})$;
   $E_{ij} = \log(\sigma(r_{i, t, \tau})/\sigma(r_{i, t, \tau}))$;
end

for $i$ := 1 to $N$ do Full model fit
   Input timeseries with number of regions $N$, number of timepoints $T$, max lag $\tau$, timeseries predictor $f$;
   Output effective connectivity matrix $E$;
   For the initial regional timeseries $X$ with $N$ regions fit a full model including region $j$;
   Fit $f$ predicting activity $X$ at time $t$ and node $i$ from times $t-1, ..., t-\tau$;
   Fit a reduced model without region $j$;
   Drop column $j$ from timeseries $X$;
   Fit $r$ predicting activity $X$ at time $t$ and node $i$ from times $t-1, ..., t-\tau$ without node $j$;
   The EC score between $i$ and $j$ equals the log of the ratio of the standard deviation of the residuals of the full and reduced model;

weight or importance is extracted from the model for each nodal covariate $i$. We repeat this for each node $j$ to fully populate an asymmetric FC matrix. The asymmetric matrix is then symmetrized by averaging itself with its transpose. Feature importance is calculated from the Gini importance of the ERT, the covariate weight for the SVM, and the Gini importance weighted by number of samples routed through the decision node for the XGBoost model. The XGBoost model was fit with a group-level hyperparameter search. The ERT and SVM models did not benefit from this search; their default parameters were already optimal. The hyperparameter evaluations were done on HCP data NOT used in training, validation, or testing. For additional model fitting details see supplemental section 9.1.3. In order to evaluate the relative benefits of each proposed ML.FC measure, we test each FC measure’s reproducibility and evaluate its predictive power by using it to infer three individual traits of interest (see section 2.4).

2.1.2. Background of effective connectivity
In addition to functional connectivity, brain connectivity can be quantified with measures of time-delayed information transfer, which we denote as effective connectivity measures. Effective connectivity can be quantified in numerous ways: multivariate Granger-causal (GC) scores (Spencer et al 2018, Abidin et al 2019, Chockanathan et al 2019), bilinear GC modeling (Luo et al 2013), and other measures of directed neural influence (Bielczyk et al 2019). This paper builds new measures from the mathematical foundation of Granger causal modeling. GC measures define a directed edge by quantifying how the past history of activity signal $B$ from a particular brain region informs the future activity of signal $A$, from another brain region. In neuroimaging, signal $B$ is said to be Granger causal of signal $A$ if a model to predict the future of $A$ given all past information from all regions’ signals including $B$ is more accurate than a model that does not include $B$. The degree of causality is called the GC score (Granger 1969). To generate a Granger causal effective connectivity matrix, the Granger score between the regional time courses from each pair of regions is calculated using the GC algorithm (algorithm 1). A full model, $f$, is fit to predict activity in region $i$ at time $t$ from the past history of all regions. Then, a reduced model $f'$ is fit to predict the same activity at time $t$ from the past history of all regions except $j$. The EC score is the log of the ratio of the standard deviation of the residuals of the full and reduced models. By using a linear model $f$, a baseline measure of effective connectivity can be calculated. The linear models with which we calculate the GC score include: an unpenalized multivariate autoregressive (MVAR) model denoted $MV.GC$, an elastic MVAR model with a small $L_1$ and $L_2$ penalty ($L_1 = L_2 = \lambda = 0.1$) denoted $MV.GC_{E=0.1}$, and an elastic MVAR model with a large $L_1$ and $L_2$ penalty ($\lambda = 10$) denoted $MV.GC_{E=10}$. These regularization amounts were chosen empirically to be representative of strong and weak regularization. The timeseries is tested for significant autoregression with the Augmented Dickey Fuller test and any significant autoregression is removed prior to model fitting. Lag values of 1-5 times repetition time (TR) were tested and the model using the lag with the lowest Akaikes information criterion was selected independently for each regional model.

Next, we will build upon this classical GC foundation in two ways: by replacing the GC’s linear MVAR model with a nonlinear multivariate7 machine learning model (explained in section 2.1.3), and by fitting the GC models in a lower dimensional space with a dimensionality reduction that also enforces biological constraints from SC (explained in section 2.1.4).

7 We use the term multivariate to indicate a multivariable model with multiple independent variables.
Algorithm 2. MLEC algorithm. This algorithm describes the steps by which one calculates an effective connectivity matrix \( E \) from a neural timeseries \( X \) using a standard machine-learning effective connectivity. \( X_i = \) neural activity matrix of size \( i \) by \( t \), where \( i \in [1,N] \) and where \( t \in [1,T] \), \( T = \) number of timepoints. \( \tau = \) max log. \( f = \) timeseries predictor function. \( G = \) operator which calculates the importance score of a given model, this is the Gini impurity in a tree-based model or weight in an SVM-based model.

\[
G(f) = \sum_{n=1}^{N} P(n) \cdot (1 - P(n));
\]

\[
E_{ij} = \text{effective connectivity matrix of size } N \times N, \text{ indexed by } i \text{ and } j.
\]

**Inputs:** \( X_i \); \( i \in [1,N], t \in [1,T] \); \( \tau, f \);

**Output:** \( E_i \);

for \( i = 1 \) to \( N \) do Full model fit

\[
X_i = f(X_{i-1}, X_{i-2}, \ldots, X_{i-\tau});
\]

if \( f = \) ERT or XGB then

\[
G(f) = \sum_{n=1}^{N} P(n) \cdot (1 - P(n));
\]

else if \( f = \) SVM then

\[
G(f) = w \cdot \left( \min \|w\|^2 + C \sum \eta_i \right) + (1 - \eta_i \cdot \eta_i \geq 0);
\]

end

For an initial regional timeseries \( X \) with \( N \) regions fit a full model including region \( i \);

Fit a machine learning model \( f \) predicting activity \( X \) at region \( i \) at time \( t \) from times \( t-1, \ldots, t-\tau \);

If an XGBoost or extremely random trees predictor;

Importance score between \( i \) and \( j \) equals the Gini impurity for feature \( j \) with probability of the data being routed down a split \( p \), proportion of data routed to that split \( P_i \), set of all nodes that use feature \( j \) of \( G \);

If a support vector predictor;

Importance score is the weight given feature \( j \) by classification vector \( w \) given the support vector optimization with regularization terms \( \eta_i, \theta, b_i \);

The EC score between \( i \) and \( j \) is the feature importance for \( j \) of the model \( f \) predicting \( i \);

2.1.3. Proposed machine learning-based effective connectivity (ML.EC measures)

We propose a novel EC measure using machine learning model coefficients as in section 2.1.1, but with a time-delay lag, \( \tau \), included. The machine learning model predicts future times steps given up to \( \tau \) past time steps and identifies the important learned features of that model, (algorithm 2). Compared to \( GC \), our proposed measure captures directed influence with only one model fit per region analyzed, and thereby scales as \( O(N) \) rather than \( O(N^2) \) where \( N \) is the number of regions. Furthermore, the machine learning approaches capture nonlinear interactions which the standard \( GC \) approaches do not. We denote this Machine learning effective connectivity approach \( ML.EC \). We test both \( ML.EC \) with an extremely random trees internal predictor (denoted \( .EC_{ERT} \)) and a support vector regressor with a radial basis function kernel, denoted \( ML.EC_{SVM} \).

2.1.4. Proposed structurally-projected effective connectivity (SP.GC measures)

We also propose another novel EC measure which projects the Granger causal models into a lower dimensional space informed by a prior from diffusion MRI. This is a soft constraint which regularizes the EC measure to have at least some agreement with known physical pathways of communication (Huang and Ding 2016, Dillon et al 2017, Manning et al 2018, Allen and Weylandt, 2019, Maglanoc et al 2020). We denote this Structurally Projected Granger Causality approach \( SP.GC \), (see algorithm 3). This approach projects the timeseries into a lower dimensional representation and calculates a full and a reduced model in the lower dimensional space before projecting the predicted activity back into the original space and calculating the error in full versus reduced models. This encourages but does not force, low dimensional timeseries components to lie along known structural networks. This approach also incorporates a sparsity constraint from sparse PCA. Sparse PCA minimizes the number of nonzero terms in each principal component while maximizing the variance explained by the components (Zou and Xue 2018). This sparsity prior and the prior from the SC matrix derived from diffusion MRI encourages these components to robustly represent a physically connected sub-network of the brain.

We implement a SC constraint which encourages a more faithful interpretation the underlying brain functional from fMRI as true functional connectivity lies sparsely along physical connections (Huang and Ding 2016, Dillon et al 2017, Manning et al 2018, Allen and Weylandt, 2019, Maglanoc et al 2020). The physical connectivity prior comes from a SC matrix derived from diffusion MRI. The SC matrix is calculated from the average normalized tractogram from all 1065 subjects in the HCP computed in (Yeh et al 2018). We use a population-level prior so that it may be applied even when diffusion MRI is unavailable for every individual in a study. The strength of the SC between each pair of brain regions is computed from the number of tractography streamlines passing through each region of interest in this normalized, ensemble atlas. Then the log of the total number of streamlines between each region is used as the prior. See supplemental section 9.1.4 for further details. There are a plethora of both direct and indirect connections in the brain, and both direct and indirect connections are captured through
the streamline-derived prior. Each streamline can represent multisynaptic or monosynaptic fibers. In formulating a prior from SC, we encourage activity to lie along the streamline populations. This is an intuitive, interpretable, and logical constraint to add to EC. If there is a strong structural connection between two regions or voxels, it is more likely to have a substantive connection between them. Furthermore, physiologically, regions that are not connected should not communicate without traveling through intermediate regions. Imposing the constraint of a prior along which we hypothesize communication should suppress inferring erroneous connections. This prior is relevant for effective connectivity approaches, where we are attempting to untangle directionality and indirect versus direct communication pathways. So, connections mediated mostly by intermediate nodes, such as an indirect connection from region A to region C through intermediate region B, are explicitly modeled rather than a connection from region A to C being inferred directly. While the concept of using SC from tractography to constrain fMRI interpretation has been used to interpret fMRI before (Huang and Ding 2016, Maglanoc et al 2020), the combination with dimensionality reduction is novel to this work.

We use a formulation of prior-constrained sparse PCA to incorporate our SC prior in the timeseries dimensionality reduction. The objective function for this constrained sparse PCA shown in equation (2); which we have adapted from Dhillon et al (2014) to include the tractography prior:

\[
\overrightarrow{v}_i^\tau = \argmax_{v_i, ||v_i|| = 1, v_i, v_j=0, i\neq j, v_i \neq 0} \left( v_i^T(C + \theta \cdot P_i^T P_i) v_i - \lambda \cdot ||v_i||_1 \right), \quad P_{ij} = \log \left( S_{ij} \right). \tag{2}
\]

The first term in the objective function enforced data fidelity. It depends on the covariance \(C\) \((N_{ROI} \times N_{ROI})\) between mean pairwise mean regional timeseries but is regularized with a structural prior \(P\) with an initial belief \(\theta\). The second term, \(\lambda \cdot \|v_i\|_1\), imposes the \(L_1\) sparsity with weight \(\lambda\). The prior \(P\) is a reformulation of the SC matrix into a matrix where the rows correspond to the individual regions and the columns correspond to a larger network each region can be grouped into. See supplemental section 9.1.4 for further details.

This approach has three advantages over the standard Granger causal measure. First, it incorporates prior information to regularize fMRI interpretation. Second, it reduces the number of measure parameters that must be tuned. In SPGC, only the maximum lag needs be selected for the method as the prior belief weight (\(\theta\) in equation (2)) is fixed at 1 for all experiments, giving equal weighting to the calculated
covariance and the prior. GC with an elastic net requires selection of: (1) an $L_1/L_2$ ratio, (2) penalty weight, and (3) testing of multiple lags up to and including the maximum lag. In practice, for a given maximum lag, only 1 model needs to be fit for SP.GC, while fitting the standard GC requires dozens of cross-validated models to be fit to properly optimize the lag and regularization parameters. A sufficiently large lag selected before dimensionality reduction appropriately weights the lag values through the dimensionality reduction itself (DSouza et al 2017). Third, SP.GC model fitting is faster than standard GC, as there are fewer variables as the number of components is much less than the number of regions.

In this study, the Schaefer functional atlas (Schaefer et al 2018) is used with the cerebellum and striatum added from the AAL atlas (Rolls et al 2020). The Schaefer atlas has the advantage that each ROI is assigned a corresponding coarse and fine RSN label facilitating the construction of our RSN SC prior. We generate two SC network priors, a coarse one with 18 regions (7 RSNs, subcortical structures, and the left and right hemisphere cerebellar gray matter), as well as a fine prior with 38 regions (17 RSNs, subcortical structures, and the left and right hemisphere cerebellar gray matter). The coarse SC prior encourages functional connections that capture left/right hemisphere connections at a whole RSN level, while the fine SC prior encourages the projection of whole brain functional activity to smaller subsections of structurally connected sub-elements of the larger RSNs. We denote the SP.GC approach using the coarser 7 RSN prior $SP.GC_{7}$ and the SP.GC approach using the finer 17 RSN prior $SP.GC_{17}$.

As a baseline of comparison, we also choose to use a PCA projection to a number of components preserving 95% of the variance in the timeseries, analogous to previous work (Luo et al 2013, Abidin et al 2019, Chockanathan et al 2019). The baseline measure does not impose the SC prior. This measure can be computed with algorithm $3_{u}$ using a PCA projection rather than a structurally-constrained projection. We denote this PCA-projected low-dimensional Granger scored measure PC.GC.

### 2.2. Materials

This work uses fMRI data from the HCP (van Essen et al 2012) to evaluate the proposed connectivity measures. We use the four scans of each participant, including on one day: (i) a left-to-right phase-encoded fMRI acquisition and (ii) a right-to-left phase-encoded fMRI acquisition, and on a subsequent day: a repeated (iii) left-to-right and (iv) a right-to-left acquisition. The measures are evaluated for their ability to produce a consistent connectivity matrix across the repeat scans for each subject.

From the HCP database, 805 subjects have the full complement of four repeat scans and demographic information. From these we excluded subjects with substance use (including alcohol and tobacco), as these are known to confound the reproducibility of longitudinal connectivity. Of the remaining 517 subjects, we selected 100 subjects which were demographically diverse and had the least head motion defined by mean framewise displacement between fMRI frames. This minimizes motion confounds which can influence connectivity measures with correlated non-neural signal (Noble et al 2019, Satterthwaite et al 2019). Subjects were selected to match the demographics of the 2010 USA census data (subject distribution shown in figure 1(B)) and the CONSORT diagram of data selection is shown in figure 1(A). I.E. as the 2010 census distribution was 61.5% white, 17.6% Hispanic/Latino, 12.3% Black, 8.6% other, 62 white, 18 Hispanic, 12 Black, and 8 other subjects were chosen, with an even male/female split. A plot of the mean framewise displacement of the selected subset of 100 subjects versus the 517 initial subjects (figure 1(B)), shows the motion level of the chosen set compared to the remainder of HCP under consideration. This selected subset of subjects is later further split for cross-validation of reproducibility and for nested cross-validation for predicting

![Figure 1](image-url)
of target values (with multiple train, validation, and test splits in the nested case).

The selected data was processed with the standard HCP minimal preprocessing pipeline (Glasser et al 2013). Mean regional timeseries were then extracted with the Schaefer atlas with 100 anatomical regions with additional subcortical regions included (Schaefer et al 2018). The Schaefer atlas with 100 regions was chosen over the 200 or 400 region parcellations to decrease computation time while remaining sufficiently large to test our approaches. The Schaefer atlas is a functional atlas whose regions are defined through the clustering of functional activity in fMRI (Schaefer et al 2018), and this functional atlas was chosen because: (1) a functional atlas tends to capture better functional variability than a purely anatomical atlas (Mellema et al 2022), and (2) the Schaefer atlas groups regions into resting-state networks (RSNs), which facilitates inter and intra RSN partitioning and analysis. The cerebellum and striatum from the AAL atlas (Rolls et al 2020) were included as well because both structures contain signals of diagnostic importance and are often overlooked in prior analyses (Stoodley et al 2012).

2.3. Experiment 1: comparison of reproducibility

The reproducibility of the connectivity measures was quantified using five different metrics. These reproducibility metrics include linear, nonlinear, and clustering metrics. The linear metrics included: (1) the average root mean squared difference of each element in the connectivity matrix (after z-score normalizing the elements) across runs of the same subject, and (2) the Pearson’s Correlation, r, between the connectivity elements (edges) of any two pairs of scans of the same subject, averaged over all pairs of scans. The nonlinear metrics of reproducibility included: (1) the average cosine similarity between all edges of any two pairs of scans of the same subject, and (2) the two-way random, single score intraclass correlation coefficient (ICC(2,1)) edgewise (Noble et al 2019) between any two scans of the same subject. Additionally, a clustering score for each EC or FC measure was calculated. This clustering score was the Davies-Bouldin (DB) index (Bezdek and Pal 1998). This index quantifies how well each subject is separated from all the subjects after projection to a low dimensional space and a higher value indicates greater separation (Finn et al 2015). A higher DB score indicates that a more subject-specific fingerprint was identified, capturing aspects of connectivity unique to that subject. Finally, as an additional post-hoc analysis we evaluated how well each measure performed when given less and less of the timeseries data duration. This tested each measure’s ability to maintain high reproducibility using a fraction of the timeseries, which could enable shorter acquisitions facilitating future fMRI studies.

2.4. Experiment 2: comparison of predictability of individuals’ traits

In addition to reproducibility metrics, using the subset of EC and FC measures that had the highest reproducibility we tested how well the connectivity measures could predict three categories of targets: a physiological trait, cognitive trait, and a combined physiological and cognitive trait. The physiological trait chosen from the HCP dataset was mean arterial blood pressure, the cognitive trait was fluid intelligence as measured with the Pennsylvania matrix reasoning test, and the combined physiologic and cognitive trait was the stress and adversity inventory. The combination of physiologic, cognitive, and combined traits was chosen to be more representative of possible real-world targets than a random selection of predictable values. Each of these targets is a single scalar value per subject, averaged over all their visits. i.e. for the first subject’s four scans, there is one scalar target to predict from each of the connectivity ‘fingerprints’. Cross-validation partitioning was performed grouped at the subject level so that a subject’s scans are contained in one partition.

2.4.1. Univariate analysis of effect size

To evaluate how predictive the connectivity measures are, first we performed a univariate analysis. The effect size per edge was measured with Cohen’s d and compared across the measures with high reproducibility, as measured by Pearson’s r and measured with the individual subject clustering score in order to compare predictive information present in the calculated connectivity. An ideal measure will have high reproducibility and high effect size across a variety of predictive tasks.

2.4.2. Multivariate analysis

Univariate analysis does not suffice to show that a connectivity measure is apt to yield accurate multivariate predictions. Therefore, we also selected significant edges (univariate significance p ≤ 0.05) as candidate covariates to form multivariate predictive models for each connectivity measure. To reduce colinearity, pairs of such edges with covariance greater than 0.75 were identified and the edge with higher effect size was retained. The surviving set of edges was used to predict the targets as in experiment 1 with an elastic net predictor and the predictive power was measured. A 10 × 5 cross-validation approach was applied. In the 10-fold cross validation, the data was stratified by the target measure and 10% of subjects were set aside to test each iteration. The inner 5-fold cross validation performed hyperparameter optimization where the elastic net’s L1 /L2 ratio was tuned with a grid search using ratios from the set {0.1, 0.5, 0.7, 0.9, 0.95, 0.99, 1.0}. A nested cross-validation approach was chosen because it tends to provide a low bias estimate of real-world performance (Cawley and Talbot 2010). The regularization weight, α, was
optimized through coordinate descent, and the model with the lowest mean squared error across the inner cross validation folds was evaluated on the 10% test set from the outer fold.

The entire procedure was itself repeated ten times using Monte Carlo iteration as this has been shown to increase the stability of the estimated prediction (Wu et al 2020). The coefficient of determination, $R^2$, of the internally fitted model on the held-out data and averaged over outer Monte Carlo iterations was recorded. A paired one-tailed t-test testing the null hypothesis that the highest performing model’s $R^2$ was not greater than other models’ $R^2$ was performed. The Bonferroni corrected p-value was reported. Bonferroni correction was chosen to minimize type 1 error with a criterion more stringent than FDR correction for answering the specific question of which predictor produces the highest $R^2$.

2.4.3. Multi-input, multivariate analysis
An additional post-hoc analysis was performed to further measure the complementarity of the connectivity measures from the trait predictions described in section 2.4.2. This secondary analysis determines which connectivity measures contain complementary information and compares models built with a combination of complimentary measures to the models built on the original separate measures. To determine measure complementarity, a linear mixed effects (LME) model was fit to predict a participants’ trait by combining the predictions from multiple models each using a single separate connectivity measure. For example, the predicted mean arterial pressure predicted from partial correlation was concatenated with the predicted arterial pressure from the other FC connectivity measures into a vector, and that vector of predictions was used to predict mean arterial pressure. If partial correlation contains trait predictive information that $MLFC_{ERT}$ does not and vice-versa, we would expect the LME model to give significant weight to predictions generated from both partial correlation and $MLFC_{ERT}$. If one of the measures contains only a subset of the information contained by another, the measure with greater predictive information will be assigned a large weight, whilst the other measure will be assigned a small weight close to zero. The LME models were fit with a subject-specific intercept and group level slope per equation (3):

$$y_j = \mu_{0,j} + \beta_1 x_{1,j} + \beta_2 x_{2,j} + \ldots + \beta_n x_{n,j} + \epsilon_j$$

where $j$ indexes the subject and $n$ is the number of measures combined from section 2.4.2. The predicted trait for a subject, $y_j$, is a function of the subject-specific intercept ($\mu_{0,j}$) and the weights ($\beta_i$) on the prediction from each of the $n$ measures. Each predictor ($x_{n,j}$) is the predicted value of subject $j$’s trait (e.g. predicted mean arterial pressure, stress, or fluid intelligence) from each of the $n$ elastic models trained from a single measure (e.g. partial correlation). The fitted LME coefficients with $p \leq 0.05$ and magnitude greater than 10% of the maximum coefficient magnitude were considered to contain complimentary information in the predictions. This produced a subset of complimentary connectivity measures that could then be used in another multi-input, multivariate elastic net model. This secondary model with complimentary measures was also fit with $10 \times 5$ cross-validation and compared to the original models, to test the benefit of combining the complimentary features.

3. Results
3.1. Experiment 1: comparison of reproducibility
3.1.1. Comparison of the reproducibility of functional connectivity
The reproducibility of six functional connectivity metrics: (1) Pearson’s correlation based connectivity, (2) partial correlation connectivity, (3) spectral GC functional connectivity (denoted $SG.CC$), and $MLFC$ based connectivity (4) with XGB, (5) with extremely random trees, and (6) with a radial basis function support vector regressor was quantified with five metrics. These metrics included: Pearson’s $r$, root mean square error (RMSE), ICC, cosine similarity, and the ease of separability (DB clustering score) as described in section 2.3. The results from these comparisons are shown in table 1 with the five right most columns showing the mean reproducibility and 95% confidence interval. The best reproducibility in each column is in boldface. The significant differences (FDR corrected p-value $< 0.05$) between the top result and the other entries within a column are denoted with an asterisk,* . FDR Correction was chosen over familywise type 1 error rate control as we have higher tolerance for type 1 errors and lower tolerance for type 2 errors in this relatively broad comparison using multiple measures. The proposed metrics are distinguished with a grey background. Of all the FC metrics, the proposed $MLFC_{XGB}$ connectivity had the highest Pearson’s $r$ and cosine similarity, as well as the lowest RMSE. Partial correlation connectivity had the best (lowest) DB index while, correlation-based connectivity had a higher ICC than any other metric; however, this metric is less sparse than the other metrics which could artificially inflate this measure for correlation. Though RMSE is a suboptimal metric of reproducibility across multiple metric types due to the different distributions and sparsity of the FC metrics, it is shown in order to facilitate comparisons to the literature. Pearson’s $r$ which is less susceptible to scale variations and sparsity, and the DB clustering score, which captures subject identifiability, were selected as measures for further analysis. A comparison of the six FC methods along these two metrics is shown in figure 2. Under the X
axis, the mean connectivity matrix is shown which is computed across all subject scans for each method. The ordering of the anatomy and RSNs across the columns and rows within each matrix is detailed in supplementary figure S1. The proposed ML.FC_{XGB} with XGB connectivity metric (right) had the best reproducibility by most measures (bold in table 1) including Pearson’s r, while partial correlation had the second best for most measures except DB score, for which it had the best. This suggests conditioning the connectivity between two nodes on the activity of all other nodes is critical for reproducibility.

3.1.2. Comparison of the reproducibility of effective connectivity

The effective connectivity measures were evaluated with Pearson’s r, RMSE, ICC, cosine similarity, and the ease of separability via the DB score, as described in section 2.3. The results from these comparisons are shown in table 2. Significant differences are denoted with an asterisk and the most reproducible measure in each column is boldfaced. The proposed metrics are distinguished with a grey background. We note that the ICC metric for measuring the reproducibility of the GC connectivity measures is influenced by the sparsity of the connectivity measure. The proposed ML.FC_{ERT} connectivity with extremely random trees predictor outperformed all other traditional and proposed methods across all metrics except clustering, where it provided a respectable performance close to the median among the tested methods. The proposed ML.FC_{XGB} connectivity with SVM implementation across all reproducibility measures, suggesting superiority of the ERT based predictor for this connectivity measure.
The regularized GC connectivity measures from the elastic MVAR model (MV.GC) performed significantly better than the unregularized GC measure (MV.GC). The higher elastic penalty (λ = 10) increased Pearson’s r relative to the lower elastic penalty (λ = 0.1), and the lower elastic penalty had a superior cluster separability. The proposed structurally projected GC method, SP.GC17, attained greater reproducibility than either the elastic MV.GC or the PG.GC connectivity measures in its cosine similarity, and was also better in other measures of reproducibility including Pearson’s r and RMSE. To further understand the differences between the GC connectivity metrics (MV.GC, PC.GC and SP.GC), we quantified the stability of these measures as a function of the amount of scan time (fMRI timeseries length) used to measure the causal connectivity. Notably, SP.GC17 using 50% of the initial timeseries has a higher reproducibility than the MV.GC or PG.GC methods using 100% of the timeseries.

Full results of the GC comparison are shown in supplementary figure S2. The stability of the SP.GC17 projection using short timeseries has potential for tangible benefit to analyze data from studies with limited fMRI acquisition duration.

The distributions of Pearson’s r and the clustering score for each of the EC metrics are shown in figure 3. This view manifests the clearly superior performance of ML.ECERT connectivity along the Pearson’s r metric as well as the reasonable clustering performance. Below each measure is the mean connectivity matrix across all subject scans for each measure. The ordering of the anatomy and RSNs across the columns and rows within each matrix is detailed in supplementary figure S1.

### Table 2. Comparison of the reproducibility of the six EC methods including proposed (gray shaded) and traditional (nonshaded) functional connectivity measures. The section where the measure is described is indicated in the Section column. The best performing methods in each metric are shown in boldface. Statistically significant differences from the best performing method after FDR correction at 5% are indicated with *.

| EC measure (ours = ours) | Section | Linear | Nonlinear | Cosine similarity | Clustering DB score |
|--------------------------|---------|--------|-----------|-------------------|-------------------|
| MV.GC                    | 2.1.2   | 0.17 ± 0.00* 1.15 ± 0.01* 0.10 ± 0.00* 0.34 ± 0.01* 11.0 ± 1.1* |
| MV.GC_λ=0.1             | 2.1.2   | 0.29 ± 0.00* 0.93 ± 0.01* 0.01 ± 0.00* 0.57 ± 0.01* 3.97 ± 0.39* |
| MV.GC_λ=10              | 2.1.2   | 0.32 ± 0.01* 0.85 ± 0.02* 0.00 ± 0.00* 0.64 ± 0.01* 5.78 ± 0.44* |
| ML.ECERT                 | 2.1.3   | 0.47 ± 0.00* 0.37 ± 0.01* 0.39 ± 0.00* 0.93 ± 0.00* 6.50 ± 0.52* |
| ML.ECVM                  | 2.1.3   | 0.31 ± 0.00* 0.90 ± 0.01* 0.22 ± 0.00* 0.59 ± 0.01* 10.4 ± 1.4* |
| PC.GC                    | 2.1.4   | 0.33 ± 0.01* 0.82 ± 0.02* 0.15 ± 0.00* 0.67 ± 0.01* 6.84 ± 0.36* |
| SP.GC_7                  | 2.1.4   | 0.32 ± 0.00* 0.84 ± 0.01* 0.07 ± 0.00* 0.65 ± 0.01* 3.00 ± 0.36* |
| SP.GC_17                 | 2.1.4   | 0.36 ± 0.00* 0.76 ± 0.01* 0.06 ± 0.00* 0.73 ± 0.01* 4.32 ± 0.39* |

3.2. Experiment 2: comparison of predictability of individuals’ traits

3.2.1. Comparison of trait predictability using the proposed functional connectivity measures

The ability of the connectivity metrics to predict subject-level traits was evaluated in the second experiment. The most reproducible FC proposed measures (ML.FCERT and ML.FCXGB) were compared to traditional methods of FC: correlation and partial correlation-based connectivity (figure 4). We first performed the univariate analysis of effect size per FC edge as described in section 2.4.1. Figure 4(A) shows the effect size of the top 50 edges in predicting mean arterial pressure, figure 4(B) shows the effect size of the top 50 edges in predicting stress, and figure 4(C) shows the effect size of the top 50 edges in predicting fluid intelligence. The results indicate that the effect sizes of the different methods are comparable for the different predictions except for fluid intelligence, where Correlation and ML.FCERT gave superior performance.

Next, a multivariate analysis was performed (section 2.4.2) in which multiple edges were combined into one model to quantify whether the edges contain complementary information and determine which measures contain the largest total information about the prediction targets. Figure 4(D) shows the $R^2$ on the held-out test data. For all three targets, multivariate connectivity from ML.FCXGB gave the top performance, followed closely behind by partial correlation. ML.FCERT and correlation were distant 3rd and 4th place finishers.

Using the elastic net predictions themselves as input to an LME model allowed us to determine which predictions were complementary (table 3). Our
**Figure 3.** Reproducibility of effective connectivity measures across repeats as measured by Pearson’s $r$ and Davies Bouldin index. The distribution of reproducibility as measured by Pearson’s $r$ and DB clustering score for every pair of EC matrices in the four-scan set per subject is shown. Superior reproducibility for both scores is found at the top of the graph’s Y axes. The mean EC matrix for each measure is displayed under the x axis.

**Figure 4.** Predictive ability of FC measures. (A) Shows the effect sizes of the most reproducible FC measures when used to regress a physiologic target: mean arterial blood pressure. (B) Shows the effect sizes of the most reproducible FC measures when used to regress a physiologic and cognitive target: stress. (C) Shows the effect sizes of the most reproducible FC measures when used to regress a cognitive target: fluid intelligence. (D) Shows the performance (as measured by $R^2$) of a model trained using the top edges ± the 95% CI for each FC connectivity method over ten outer random permutations of the data and ten inner cross validation folds. The top edges used were all edges from the univariate analysis of the training set with $p \leq 0.05$. Significant differences from top performer were calculated with a Bonferroni corrected one-sided t test. $p > 0.05$, $p < 0.05$, $p < 0.01$, $p < 1 \times 10^{-3}$, and $p < 1 \times 10^{-4}$ are indicated with ns, *, **, ***, or **** respectively.

The definition of complementarity here stems from the concepts outlined in section 2.4.3, where the ability of the model to predict from combinations of EC and FC data is analyzed. Complementarity then, is defined as having independent predictive power that is not found in other methods though the magnitude of appropriately regularized coefficients. In this case we observed that combining the complimentary features did not provide a statistically significant improvement in prediction accuracy. $ML\_FC_{XGB}$ contained the information present in the other measures in all cases except mean arterial pressure. For that target, $ML\_FC_{XGB}$ and partial correlation contained complimentary information. The results from figure 4 and table 3 suggest that the boosting method of $ML\_FC_{XGB}$ is particularly well-suited to regularize and discover a stable set of connectivity features, perhaps because of...
Table 3. Complementarity of EC and FC measures. For each set of EC and FC measures, the predicted scores from the elastic-net predictions were fed into a secondary LME model with subject-specific intercepts and group level slopes. This tested for complimentary information contained in the predictions. Those predictions using connectivity metrics were considered to contain complimentary information if the fitted coefficient had a p-value \( \leq 0.05 \) and a magnitude greater than or equal to 10% of the maximum coefficient magnitude. The coefficients are displayed below, with p values in parentheses. Complimentary sets of connectivity features are indicated with boldface.

| Prediction target | Functional connectivity measure | Effective connectivity measure |
|-------------------|---------------------------------|-------------------------------|
| Mean arterial pressure | \(-0.02 (0.00)\) | \(0.10 (0.00)\) | \(0.02 (0.07)\) | \(0.92 (0.00)\) |
| Perceived stress | \(-0.02 (0.05)\) | \(0.06 (0.00)\) | \(0.01 (0.14)\) | \(0.97 (0.00)\) |
| Fluid intelligence | \(-0.02 (0.00)\) | \(0.00 (0.94)\) | \(-0.01 (0.01)\) | \(1.03 (0.00)\) |

Figure 5. Predictive ability of EC measures. (A) Shows the effect sizes of the most reproducible EC measures when used to predict a physiologic target: mean arterial blood pressure. (B) Shows the effect sizes of the most reproducible EC measures when used to predict a physiologic and cognitive target: stress. (C) Shows the effect sizes of the most reproducible EC measures when used to predict a cognitive target: fluid intelligence. (D) Shows the performance (as measured by \( R^2 \)) of a model trained using the top edges \pm 95% CI for each EC connectivity method over ten outer random shufflings of the data and ten inner cross validation folds. The top edges used were all edges from the univariate analysis of the training set with \( p \leq 0.05 \). Significant differences from top performer were calculated with a Bonferroni corrected one-sided t test. \( p > 0.05 \), \( p < 0.05 \), \( p < 0.01 \), \( p < 1 \times 10^{-3} \), and \( p < 1 \times 10^{-6} \) are indicated with ns, *, **, ***, or **** respectively.

3.2.2. Comparison of trait predictability using proposed effective connectivity measures

The predictive ability of the different effective connectivity metrics was evaluated by comparing traditional EC measures of GC, regularized GC with an elastic penalty, and PC.GC to the most reproducible proposed metrics, ML.EC \textsubscript{ERT} with the extremely random trees predictor and SP.GC.

Figures 5(A)–(C) shows the results from the univariate analysis (section 2.4.1). We observed that the highest Cohen's \( d \) was attained for connections measured with PC.GC (red) and ML.EC \textsubscript{ERT} (green) followed by SP.GC (purple). Figure 5(D) shows the performance of multivariate predictive models trained on a set of all edges with univariate \( p \)-value \( \leq 0.05 \) (section 2.4.2). The proposed SP.GC method (purple) and PC.GC (red) explained the most variance, followed by ML.EC \textsubscript{ERT} (green).
The multivariate models (figure 5(D)) revealed that the SP.GC set of edges (purple) tended to contain more total information than the connections computed with the remaining connectivity methods and achieved the highest performance predicting arterial pressure and stress. In close second place was the PC.GC connectivity measure (red), which achieved the highest performance predictive fluid intelligence.

A multi-input analysis (section 2.4.3) was performed to test complementarity across the EC measures using an LME model. From the results in table 3 (bottom), we observe that SP.GC, PC.GC, and MV.GCf,A = 0.1 contained complimentary information for all prediction targets. However, combining complimentary features did not provide a statistically significant improvement for target prediction. This suggests that the regularizing causal measures, via a low dimensional projection or an elastic penalty, extract different information. Furthermore, the cross-prediction comparison indicates that the single connectivity feature most apt to make a given prediction is somewhat problem specific, but SP.GC and PC.GC are well-suited to the variety of prediction tasks examined here.

4. Discussion

Among functional connectivity measures, ML.FCXGB had the highest reproducibility across most metrics. The overall predictive power to predict mean arterial pressure, stress, and fluid intelligence using the multivariate models was also highest for ML.FCXGB. Additionally, our separate analysis of complementarity using second level LME models revealed that the ML.FCXGB approach contained most of the information present in the other FC measures. These results suggest that ML.FCXGB be used as the functional connectivity metric of choice on larger datasets. If there is not enough data to effectively fit the ML.FCXGB across subjects, then we recommend the use of partial correlation based functional connectivity, which also performed well and just behind ML.FCXGB.

Among effective connectivity measures, ML.ECERT performed better in reproducibility than other measures, but suboptimally in the predictive power analysis, suggesting the presence of strong intra-edge correlation and redundant (collinear) information. However, SP.GC performed second highest in reproducibility and was also the most predictive connectivity feature set in two of our three prediction models. Our analysis of reproducibility using fractional timeseries further revealed that SP.GCf17 (a specific formulation of SP.GC) achieved high reproducibility even when applied to a small portion of the IMRI timeseries (supplementary figure S2). Our complementarity analysis using LME models, showed that each of the top EC methods contained information complimentary to each other. Collectively these results indicate that: (1) the incorporation of a structural prior to the SP.GCf17 measure appropriately constrained the GC score with higher predictive power and reproducibility than a standard PCA projection, and (2) the SP.GCf17 measure may be recommended as a method for effective connectivity measurement. If appropriate dMRI priors are unavailable, we recommend either (1) ML.ECERT to capture non-linear EC connectivity with high reproducibility, or (2) PC.GC if predictive power is prioritized over reproducibility.

Prior literature focused on the reproducibility of functional connectivity, using Pearson’s r or partial correlation (Guo et al 2012, Fiecas et al 2013, Liao et al 2013, Andellini et al 2015, Termenon et al 2016, Geerligs et al 2017, Panunzi et al 2017, Wang et al 2017, Noble et al 2017a, 2017b, 2019). Measures of reproducibility can be highly confounded (usually inflated) by motion, therefore the selection of subjects was aimed to minimize this confound (Noble et al 2019). Our estimates of reproducibility of FC measures is somewhat lower than prior reports in the literature (Noble et al 2017a) which is likely a result of our strict motion thresholds limiting the inflation of regional correlation from motion. This is supported by previous reviews of connectivity (Noble et al 2017b, 2019). The aforementioned prior research characterizes the reproducibility of correlation and partial correlation well, but used a limited set of reliability metrics, typically ICC or R² (Termenon et al 2016, Waller et al 2017, Noble et al 2017a). This study complements the prior work in three ways. First, we provide a characterize reproducibility using a multitude of metrics (linear, nonlinear, and clustering). Second, we characterize predictive power using several metrics (ICC, R², cosine similarity, DB score, and accuracy across three relevant neurophysiologic targets). Finally, we propose new measures of connectivity, quantify their reproducibility, and compare them to traditional measures. These proposed measures proved to be the most reproducible and contained the greatest predictive power of the FC measures analyzed. The measure proposed by Murugesan et al (2020) is a special case of the generalized framework for ML.FC proposed in this work. Our framework generalizes across model types (SVM, ERT, and XGB), hyperparameter optimization, develops machine learning connectivity measures for both FC and EC, and adds regularization priors, while the measure in (Murugesan et al 2020) only pertains to FC. Additionally, we evaluate the reproducibility of the proposed models, which is absent from such prior literature.

Examinations of the reproducibility of effective connectivity have been limited to bivariate GC (Fiecas et al 2013), which is a Granger causal estimate using only pairs of regional timeseries, rather than
the more comprehensive multivariate estimates using all regional timeseries employed in this study. This research drives the study of the reproducibility of EC measures beyond bivariate GC analysis. Furthermore, although many studies do not characterize performance, reproducibility is necessary but insufficient quality of a desirable neuroimaging predictor (Termenon et al 2016, Waller et al 2017, Noble et al 2017a, 2017b). Our analysis of predictive power over multiple tasks addresses this gap as well.

When comparing EC and FC measures, the best FC measures had slightly higher reproducibility and predictive power than the best EC measures. However, FC and EC capture different connectivity information and should be considered complimentary rather than competing. FC, for example, may capture longer range interactions, while shorter direct interactions may be better captured by EC. If interpretability is the most important criterion, then we recommend the use of \( SP.GC_{17} \). If predictive performance is the most important criterion, then we recommend \( ML.FC_{XGB} \). If both are equally important, we recommend combining the two, using the top \( X \) percent of informative edges (from a univariate analysis) from each measure, where \( X \) is determined based on the number of samples in the study so as to make the ratio of features to samples tractable to train a predictive model without overfitting.

There are several limitations of this research. First, comparison of reproducibility across different connectivity types can be problematic when the distribution of recovered connectivity values is concentrated to a few values. For example, a calculated GC connectivity with an elastic penalty where the \( \lambda \) penalty is extremely high can have 99% values of 0 connectivity, and the remaining nonzero connections can be uniformly distributed on a logarithmic scale within the range \([0, \infty] \). On the other hand, correlational connectivity values are within the range \([-1,1]\) and tend to be more normally distributed. By using multiple metrics of comparison including several metrics of both reproducibility and predictive power, we increase confidence in the relative usefulness of the analyzable connectivity measures. However, additional reproducibility metrics could be explored. For example, summary graph connectivity measures may exhibit useful predictive properties and reliability, but the differing sparsity between methods would need to be addressed to make such a comparison. The second limitation is that metrics of connectivity and reproducibility can be dependent upon preprocessing decisions, number of subjects, and number of replicates per subject. Future studies could explore the dependency of FC and EC measures to denoising, global signal regression, and choice of atlas, as well as look at similarity among the new and classic methods in terms of edge similarity, measures of graph similarity, and how these change with more noise added. When additional large-scale datasets with multiple replicates per subject become available, further work on reproducibility across more replicates and subjects would also be valuable and welcome. Finally, the third limitation is that, in an effort to focus on the ability of measures to quantify biological variability and not merely quantify motion artifacts, we purposefully chose a sample of subjects with exceptionally low motion. As researchers implement better practices which yield lower and lower intrinsic motion and as motion suppression methods improve, our choice becomes more and more reasonable. However we emphasize that our results pertain to studies with low motion, long scan times, and high temporal resolution. Furthermore, we chose to use all available data per subject, including four longer-duration scans per subject to make each prediction, which can increase the estimated \( R^2 \). The real-world ability to predict in these domains we would expect to be roughly uniformly lower for all methods, however, this study quantified the relative ability of the models to make predictions within a dataset with lower motion. A full study estimating the real-world power of these methods is beyond the scope of this work, which proposes and tests the internal validity of novel algorithms. However, we have also successfully made realistic predictions with these measures in a Parkinson’s disease predictive modeling system, and direct the reader to that body of work (Mellema et al 2023).

5. Conclusions

This study proposes a functional connectivity metric (\( ML.FC \)) and an effective connectivity metric (\( ML.EC \)) that efficiently capture nonlinear associations between brain regions. This study also proposes a connectivity metric (\( SP.GC \)) that encourages the connectivity recovered from fMRI to respect underlying biological SC and efficiently measures causal associations across all brain regions, which provides researchers new capabilities for connectivity analysis. This study compared the proposed measures to traditional ones using quantitative reproducibility metrics and by quantifying their capacity to make accurate predictions of traits of individual subjects to show the internal validity of these proposed metrics. This included a physiologic trait, cognitive trait, and combined physiologic and cognitive trait. The proposed measures produced higher measures of reproducibility and were found to be more predictive across the traits. Based on the study results, two of the proposed methods: \( ML.FC_{XGB} \) and \( SP.GC_{17} \) are recommended as the connectivity measures of choice for functional and effective connectivity, respectively. The contributions of this work hold potential to further the development of tools to characterize the human connectome in health and disease and make meaningful individualized predictions of neuropsychological and neurobiological states.
Data availability statement

The data that support the findings of this study are openly available at the following URL/DOI: www.humanconnectomeproject.org/.

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Materials and ethics statement

To facilitate reuse and extension, the authors are pleased to provide full source code for the connectivity measures and manuscript analyses at: (https://git.biohpc.swmed.edu/s169682/CausalMeasures). Datasets used for this study are publicly available at (www.humanconnectomeproject.org/). The data used for this study data was gathered with written informed consent from all participants. Furthermore, this manuscript follows NIH policy on the inclusion of women and members of racial and ethnic minority groups in funded clinical research. Participant data was collected and handled in accordance with NIH guidelines, HIPPA guidelines, local statutory requirements, and handled with the principles embodied in the Declaration of Helsinki.

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