**DynamicBC:**
A MATLAB Toolbox for Dynamic Brain Connectome Analysis

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**Abstract**

The brain connectome collects the complex network architectures, looking at both static and dynamic functional connectivity. The former normally requires stationary signals and connections. However, the human brain activity and connections are most likely time dependent and dynamic, and related to ongoing rhythmic activity. We developed an open-source MATLAB toolbox DynamicBC with user-friendly graphical user interfaces, implementing both dynamic functional and effective connectivity for tracking brain dynamics from functional MRI. We provided two strategies for dynamic analysis: (1) the commonly utilized sliding-window analysis and (2) the flexible least squares based time-varying parameter regression strategy. The toolbox also implements multiple functional measures including seed-to-voxel analysis, region of interest (ROI)-to-ROI analysis, and voxel-to-voxel analysis. We describe the principles of the implemented algorithms, and then present representative results from simulations and empirical data applications. We believe that this toolbox will help neuroscientists and neurologists to easily map dynamic brain connectomics.

**Key words:** brain connectome; dynamic; effective connectivity; functional connectivity; resting-state fMRI

**Introduction**

The brain connectome collects network architectures. At macroscopic scales, the human brain connectomics provide a comprehensive description of the anatomical pathways and functional interactions among distinct brain areas (Sporns, 2013, 2014). The structural connectomics essentially comprises a comprehensive map of the anatomical connections reflecting axonal pathways (Sporns et al., 2005), and the structural covariance connectivity interpreted as the phenotype of brain development and/or plasticity (Alexander-Bloch et al., 2013; He et al., 2007). Additionally, the functional connectomics can be captured as patterns of functional covariance network (Liao et al., 2013b; Zhang et al., 2011), functional connectivity (FC) and effective connectivity (EC) networks (Friston, 2009, 2011; Marinazzo et al., 2011, 2014; Wu et al., 2013b).

FC measures statistical patterns of interactions among remote brain regions; while EC discerns the transfer of information such as directed causal interactions (Friston, 1994; Rubinov and Sporns, 2010). Consequently, the brain can be seen as a large-scale functional integrated network both during cognitive tasks and at resting state (Bressler and Menon, 2010). Resting-state FC represents the synchronization of spontaneous blood-oxygenation level-dependent (BOLD) activity and is typically analyzed in terms of correlation, coherence, and spatial grouping based on temporal similarities (Beckmann et al., 2005; Biswal et al., 1995; van den Heuvel et al., 2008). These FC analyses always assume that the functional connections remain constant during the whole period of data collection.

However, both emerging theoretical ideas and empirical observations suggest that the human brain connectome is most likely to be time dependent and dynamic, and to be related to ongoing rhythmic activity (Sporns, 2011). The functional repertoire of brain connectome is continually revisited and rehearsed in endogenous neural activity (Deco and Corbetta, 2011). Recent empirical human, macaque, and rat studies...
have observed the phenomenon that brain FC can indeed exhibit nonstationary activity, and change over a short time (Allen et al., 2014; Chang and Glover, 2010; de Pasquale et al., 2010; Di and Biswal, 2013; Handwerker et al., 2012; Hutchison et al., 2013b; Kang et al., 2011; Lee et al., 2013), and even across a few spontaneous points (Liu and Duyn, 2013; Tagliazucchi et al., 2012a; Wu et al., 2013a). Hence, dynamic techniques track the variability of the topology of the brain connectome across different cognitive states (Bassett et al., 2011; Fornito et al., 2012) and the evolution of diseased brain networks (Liao et al., 2013a; Zhang et al., 2014).

Several modeling strategies have been developed to meet brain dynamics’ need (Hutchison et al., 2013a). The most commonly utilized approach, called sliding-window analysis, is performed by conducting FC on a set number of data points. Much earlier defined paradigms information or stationary assumptions regarding the signal must be made prior to its calculation. Another widely used data-driven approach, such as Kalman filtering (KF), is capable of assessing rapidly changing connectivity relationships between brain areas (Kang et al., 2011).

Recently, several advanced toolboxes providing indexes of dynamic brain connectivity have been developed, such as GIFT (http://mialab.mrn.org/software/gift/) (Allen et al., 2014), eConnectome (http://econnectome.umn.edu) (He et al., 2011), and BSMART (www.brain-smart.org/) (Cui et al., 2008). However, most of them either focus on a special modality (e.g., functional magnetic resonance imaging [fMRI], electroencephalography [EEG], magnetoencephalography) and/or include only a subset of measures as part of a more general-purpose toolbox. More important, tracking dynamic FC (d-FC) and dynamic EC (d-EC) extends the repertoire of brain connectome. However, it is still desired that a unified toolbox facilitate the neuroscientist and neurologist to easily map dynamic brain connectome.

We hereby developed a publicly available toolbox named DynamicBC (dynamic brain connectome toolbox; www.restfmri.net/forum/DynamicBC). It is an open-source MATLAB toolbox with user-friendly graphical user interfaces (GUI). The toolbox supports NIFTI and ANALYZE images (*.nii and *.img) that would be preprocessed in the REST (Song et al., 2011) and DPARSF (Chao-Gan and Yu-Feng, 2010) toolkit, in addition to ASCII and .mat formats. The DynamicBC implements both d-FC and d-EC for tracking brain dynamics from fMRI. Particularly, a distribution-free time-varying parameter regression strategy was implemented. In addition, multiple region of interest (ROI) setting ways are provided, for example, seed-to-voxel, ROI-to-ROI and voxel-to-voxel. In the following section, the remainder of this work first describes the principles of the algorithms implemented, and then we present representative results from simulations and real data, to illustrate the reliability of all these dynamic connectivity measures.

**Function Module Implemented in DynamicBC**

**Overview of usage of the toolbox**

The DynamicBC was developed by cross platform MATLAB (Mathworks, Inc.) programming language, with a user-friendly GUI, under a 64-bit Windows environment. It is integrated by the modules of connectivity types, dynamic analysis strategies, and ROI set (Fig. 1).

**Connectivity types selection**

Functional connectivity. The toolbox focuses on the Pearson linear correlation to measure the FC between pair of regions:

\[ r_{xy} = \frac{1}{T} \sum_{t=1}^{T} [x(t) - \bar{x}] \cdot [y(t) - \bar{y}] } }{ \bar{S_x} \bar{S_y} } \]

(1)

where \( r_{xy} \) is the Pearson correlation coefficient, \( x(t) \) and \( y(t) \) are the seed and target variables with means \( \bar{x} \) and \( \bar{y} \), and standard deviations \( S_x \) and \( S_y \), respectively. The summation limit, \( T \), corresponds to the total number of time points. The most resting-state fMRI studies used this Pearson linear correlation with full-length time series, referred to as static FC (s-FC) (Biswal et al., 1995; Fox et al., 2005; Fransson, 2005; Zuo et al., 2012).

Effective connectivity. The bivariate Granger causality (GC) to explore EC was employed in the current toolbox, which tested the null hypothesis that region \( x \) does not Granger-cause region \( y \) measured via linear autoregressive model. The GC index from \( x \) to \( y \) is defined as follows:

\[ F_{x \rightarrow y} = \ln \left( \frac{\sum \xi_t^2}{\sum \eta_t^2} \right), \]

(2)

where \( \xi_t \) and \( \eta_t \) are the residuals of the restricted and unrestricted regression models respectively, and \( \sum \) indicates the variance. The static EC (s-EC) was termed to describe GC relationship by full-length time series.

**Dynamic analysis strategies**

To describe the dynamic connectivity among the brain areas, we employed a time-varying parameter regression method, which is briefly described as follows:

\[ y(t) = x(t) \beta(t) + u(t), \]

(3)

where \( x(t) \) and \( y(t) \) are the seed and target variables respectively, and \( u(t) \) is the approximation error, and \( \beta(t) \) is the coefficient to determine whether two variables covary and reflect the dynamic connectivity between \( x \) and \( y \) at time \( t \).

**Sliding-window analysis.** If we treat \( x \) and \( y \) in a short time window as being generated by an underlying (approximately) and stationary stochastic process, then the model has constant parameters during this short period. Furthermore, if the values of \( x \) and \( y \) are normally distributed and homoscedastic, coefficient \( \beta = r_{xy} \frac{S_y}{S_x} \) can be estimated by ordinary least squares estimate in a short time window. Here, we employ \( r_{xy} \) as the strength of d-FC in the short time window, without considering of the scaling in samples. By transforming equation (3) into the vector autoregressive model, and following a similar procedure, the time-varying GC between \( x \) and \( y \) could be evaluated by means of sliding-window analysis.

Flexible least squares. The value of \( \beta(t) \) may continuously change for various reasons. For example, switching in different tasks, or could be the consequence of underlying physiological process. There are two approaches to estimate the continuous changed model parameters at each observation. One popular methodology is the application of
the KF to infer time-varying parameters (Kalman, 1960). Another tool is the flexible least squares (FLS) (Hastie and Tibshirani, 1993; Kalaba and Tesfatsion, 1989). The KF typically builds on the assumption of a certain distribution in the innovations (which is usually set to be normal distribution), while FLS is distribution-free. Here, we only focus on the distribution-free method. The idea of the FLS method is to assign two types of residual error to each possible coefficient sequence estimate. The first one is the sum of squared residual measurement errors:

\[ r^2_M(b, T) = \sum_{t=1}^{T} (y(t) - x(t)b(t))^2 \]  

matching the prior measurement specification: \( y(t) - x(t)b(t) \approx 0 \). The other is the sum of squared residual dynamic error, in which FLS declares that the vector of coefficients evolves slowly over time \( (b(t+1) - b(t)) \approx 0 \), formally:

\[ r^2_D(b, T) = \sum_{t=1}^{T-1} (b(t+1) - b(t))^T (b(t+1) - b(t)) \]  

with a given \( \mu \) weighting parameter, Kalaba and Tesfatsion (Kalaba and Tesfatsion, 1989) define the incompatibility cost assigned to any \( b \) coefficient sequence as

\[ C(b, \mu, T) = \mu \cdot r^2_D(b, T) + r^2_M(b, T) \]

FIG. 1. The framework of the DynamicBC toolbox. A graphical user interface (GUI) can be started by calling the “DynamicBC” function in the command window of the MATLAB. The following procedures include three parts: the selection of connectivity types (the functional connectivity [FC] and effective connectivity [EC]), selection of dynamic analysis strategies (the sliding-window and flexible least squares [FLS]), and selection of connectivity measures (the seed-to-voxel, region of interest [ROI]-to-ROI, and voxel-to-voxel analysis). The subsequent brain connectomes are then visualized. Color images available online at www.liebertpub.com/brain
The incompatibility cost function $C(\beta, \mu, T)$ generalizes the goodness-of-fit criterion function for ordinary least squares estimation by permitting the coefficient vector $\beta(t)$ to vary over time. When $\mu$ approaches was set to zero, $r_M^2$ can generally be brought down close to zero and the corresponding value for $r_D^2$ will be relatively large, resulting in a rather erratic sequence of estimates. As $\mu$ becomes arbitrarily large, the incompatibility cost function assigns all importance to the dynamic specification. This case yields the ordinary least squares solution, $r_M^2$ is minimized subject to the following formula: $r_D^2 = 0$.

**ROI set for d-FC and d-EC**

The implementation includes three ways to set ROI for dynamic brain connectivity analysis (Fig. 1). Seed-to-voxel (voxel wise) analysis calculated the bivariate FC/EC between seed brain region and every voxel in the whole brain. ROI-to-ROI (ROI-wise) analysis computed the bivariate FC/EC between each pair of ROIs, resulting brain connectivity matrix (network) allows users to further perform graph theoretical analysis (Liao et al., 2010; Wu et al., 2013b). Voxel-to-voxel computed the bivariate FC/EC between every pair of voxels without using a priori seed/ROI to mapping whole-brain connectome (Tomasi and Volkow, 2010; Zuo et al., 2012). In addition, the toolbox then provides to calculate the reliability of uncovering d-FC by short sliding-window method is drawn as the pink and green line, respectively. The yellow filled area indicates the 95% confidence intervals for $\beta$ coefficient (red line) estimated by KF method (Fig. 2B, C), and there is no significant connectivity between the two signals at a time point when the enclosed range did not cover zero; the overlapped blue line is $\beta$ estimated by FLS method. These results suggest that both the FLS and sliding window analysis could capture ground truth $\beta$ over time. Under the different parameters, sliding window analysis and FLS may induce different results, experimental outcomes of FLS for $\mu = 0.01, 0.1, 1, 10, 100, 1000$, and 1000 are plotted in Figure 3.

**Illustrations of Dynamic Brain Connectome**

**Simulation**

The following simulated example is explored here to validate the reliability of uncovering d-FC by short sliding-window analysis and FLS, and the KF analysis were included. The generated data of $y$ and $x_i$ follow a normal distribution with mean 0 and variance 0.3. The generated data of $y$ and $x_i$ are shown in Figure 2A. The FC between $y$ and $x_i$ are evaluated by s-FC and d-FC (included sliding-window analysis (window size = 20 time points, step = 1 time point), FLS ($\mu = 20$), and KF with default parameters (Peng and Aston, 2011)); the s-FC and d-FC by sliding widow method is drawn as the pink and green line, respectively. The yellow filled area indicates

$$y = x_1 \beta_1 + x_2 \beta_2 + \varepsilon,$$

$$\begin{align*}
\beta_1(t) &= 0, \quad x_1(t) = e_1(t), \quad \text{if} \quad t \in [1:50], [101:150], [201:250] \\
\beta_1(t) &= \log(10), \quad x_1(t) = \sin(t + 10) + 0.01, \quad \text{if} \quad t \in [51:100] \\
\beta_1(t) &= \cos(t \cdot \frac{\pi}{5}), \quad x_1(t) = \cos(t \cdot \frac{\pi}{5}), \quad \text{if} \quad t \in [151:200] \\
\beta_1(t) &= 2.5, \quad x_1(t) = \sin(t \cdot \frac{\pi}{2}), \quad \text{if} \quad t \in [251:300] \\
\beta_2(t) &= 0, \quad x_2(t) = e_2(t), \quad \text{if} \quad t \in [1:50], [101:150], [201:300] \\
\beta_2(t) &= -\log(4t), \quad x_2(t) = \cos(t + 10) + 0.01, \quad \text{if} \quad t \in [51:100] \\
\beta_2(t) &= 0.2 \sqrt{t}, \quad x_2(t) = -\sin(t \cdot \frac{\pi}{50}), \quad \text{if} \quad t \in [151:200],
\end{align*}$$


**fMRI data**

We selected 32 young healthy subjects (10 females, all right-handed; age: 25.19 ± 6.71 years) from our previous studies (Liao et al., 2013b; Zhang et al., 2011). The subjects had no history of neurological disorder or psychiatric illness and no gross abnormalities in the brain MRI images. Written informed consent was obtained from all subjects. The study was approved by the Local Medical Ethics Committee at Jinling Hospital, Nanjing University School of Medicine.

We performed functional neuroimaging acquisitions using a Siemens Trio 3T scanner at Jinling Hospital. We used foam padding to minimize head motion. We acquired resting-state functional images using a single-shot, gradient-recalled echo planar imaging sequence (250 volumes, repetition time $= 2000$ msec, echo time $= 30$ msec, flip angle $= 90^\circ$, field of view $= 240 \times 240$ mm², inter-slice gap $= 0.4$ mm, voxel size $= 3.75 \times 3.75 \times 4$ mm³, 30 transverse slices aligned along the anterior–posterior commissure). Subjects were instructed simply to rest with their eyes closed, not to think of anything in particular, and not to fall asleep. Subsequently, we acquired 3D T1-weighted anatomical images in sagittal orientation using a magnetization-prepared rapid gradient-echo sequence (repetition time $= 2300$ msec, echo time $= 2.98$ msec, flip angle $= 9^\circ$, field of view $= 256 \times 256$ mm², voxel size $= 0.5 \times 0.5 \times 1$ mm³, 176 slices without inter-slice gap).

**Preprocessing**

Functional images were preprocessed using the REST (Song et al., 2011), DPARSF (www.restfmri.net) (Chao-Gan and Yu-Feng, 2010) and SPM8 (www.fil.ion.ucl.ac.uk/spm) toolkits. We excluded the first 10 images to ensure steady state longitudinal magnetization, and then we corrected the remaining images for temporal differences and head motion. No
translation or rotation parameters in any given data set exceeded ±1 mm or ±1°. The individual 3D T1-weighted anatomical image was coregistered to the functional images. The 3D T1-weighted anatomical images were segmented (gray matter, white matter, and cerebrospinal fluid). A non-linear spatial deformation was then calculated from the gray matter images to a gray matter template in Montreal Neurological Institute (MNI) space using 12 parameters that were defined by affine linear transformation. This transformation was then applied to the functional images. The normalized images were resliced at a resolution of 3×3×3 mm³.

Nine sources of variances including six head motion parameters, averaged signals from cerebrospinal fluid and white matter, and global brain signal were regressed. Next, the data were band-pass filtered (0.01–0.08 Hz).

**Seed-to-voxel-based s-FC and d-FC patterns**

For seed-to-voxel analysis, a sphere (radius = 6 mm) in the posterior cingulate cortex (PCC; MNI coordinates: −2, −48, 28) was defined as the seed according to previous study (Spreng et al., 2013). The averaged BOLD time series was then obtained from the PCC and linear Pearson correlation analysis was performed in a voxel-wise way using full-length time series to generate s-FC map. Next, the s-FC map of each subject was converted into z map by Fisher’s z transformation.
Finally, z maps were combined across subjects using a fixed-effects analysis (sum and divided by the square root of number of subjects) to generation group s-FC map. The s-FC pattern was consistent with previous resting-state fcMRI studies (Fig. 4A) (Fox et al., 2005; Fransson, 2005; Zhang et al., 2011). The PCC showed positive s-FC with medial prefrontal cortex (mPFC), bilateral inferior parietal lobule, middle temporal gyrus, and superior frontal gyrus. These regions are considered part of the default mode network (DMN) (Raichle et al., 2001). In addition, the PCC showed negative s-FC with brain

FIG. 3. The differential β amplitudes as estimated by the FLS method with different penalty weights μ, which increase by powers of ten: 0.01, 0.10, 1, 10, 100, 1000, and 10,000. The gray line indicates the ground truth β. Color images available online at www.liebertpub.com/brain

FIG. 4. Illustration of the seed-to-voxel-wise s-FC and d-FC. (A) Group-averaged s-FC map following the linear Pearson’s correlation analysis using the full-length of the resting-state blood-oxygenation level-dependent (BOLD) functional magnetic resonance imaging (fMRI) signal with a seed placed in the posterior cingulate cortex (PCC; Montreal Neurological Institute [MNI] coordinates: x = –2, y = –48, z = 28, 6 mm radius sphere). (B) d-FC map following FLS analysis with the PCC seed of a representative healthy subject. Two target ROIs with a 6 mm radius sphere were placed in the medial prefrontal cortex (mPFC, MNI coordinates: x = 6, y = 51, z = 9), and right superior parietal lobule (SPL, MNI coordinates: x = 63, y = –33, z = 39). The d-FC (solid line) and s-FC (dashed line) time series of mPFC (red line) and SPL (blue line) varied in a time-dependent manner. Warm and cool colors indicate brain regions with positive and negative temporal correlations with the PCC seed, respectively. Color scales represent the group-averaged correlation coefficient value (Z) of the s-FC map and individual β amplitude values of the d-FC map, respectively. See Supplementary Movie S1; Supplementary Data are available online at www.liebertpub.com/brain. Color images available online at www.liebertpub.com/brain
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regions involving the frontoparietal control network (FPCN), and the dorsal attention network (DAN).

The above-mentioned PCC seed was also applied to d-FC analysis. The FLS analysis strategy was selected for illustrating here. d-FC organizations of one representative subject are shown in Figure 4B (see corresponding Supplementary Movie S1; Supplementary Data are available online at www.liebertpub.com/brain). To display PCC seeded FC dynamics, two target ROIs with 6 mm radius sphere (mPFC, MNI coordinates: [6, 51, 9], and superior parietal lobule [SPL], MNI coordinates: [63, –33, 39]) were identified in s-FC map (Fig. 4A). As seen from d-FC time series, FC between the PCC seed and the mPFC target, which both are belongs to the DMN, is relatively stable (solid red line); while FC between the PCC seed and SPL target involved in the DAN is highly nonstationary (solid blue line), in some cases exhibiting both strongly negative and correlations within the whole scan. These results give an evidence that the FC between brain regions has more variable in distinct networks than within one network (Allen et al., 2014).

For s-EC and d-EC analysis, we also defined the PCC as seed (see Seed-to-voxel-based s-FC and d-FC patterns section). The averaged PCC BOLD time series was used to compute the GCA value of each voxel using full-length time series, resulting in s-EC map. Then, the s-EC map of each subject was combined across subjects using a fixed-effects analysis to generation group s-EC map. The s-EC from the PCC to whole brain ("out" map) and whole brain to PCC ("in" map) were illustrated in Figure 5A.

Subsequently, we evaluated d-EC using the sliding-window GC analysis. We calculated GC maps between the PCC time series and all other brain voxel for a sliding-window of 50 volumes. For each sliding-window, we obtained the GC value in and out maps. The window was then shifted by 5 volumes and a new GC maps was calculated. This analysis strategy permitted to estimate d-EC over time. The d-EC map showing GC from whole brain to the PCC (upper panel) and from the PCC to whole brain (bottom panel) are illustrated as in Figure 5B (see Supplementary Movies S2 and S3). As see from d-EC time series of the target region mPFC, both the in- and out-influence are relatively stable and consistent changes across time.

ROI-to-ROI-based s-FC and d-FC networks

To illustrate the ROI-to-ROI-based d-FC networks, we selected 43 ROIs, which are involved in the DMN, DAN, and FPCN in line with the previous study (Spreng et al., 2013). Detailed brain regions and corresponding MNI coordinates and abbreviations of each ROI are shown in Supplementary Table S1. We extracted the averaged BOLD time series from each ROI (6 mm radius sphere) from each subject. We computed the Pearson linear correlation coefficient between a pair of ROIs using the full length of the time series, and the square $43 \times 43$ s-FC matrix was obtained individually. Then, s-FC matrices after Fisher’s z transformation were combined across subjects using a fixed-effects analysis to generation group s-FC matrix. To visualize the s-FC network, we positioned the regional centroid of each ROI (node) according to its MNI coordinates, and defined the threshold ($p < 0.01$, Bonferroni corrected) to remove spurious connection (edge).

FIG. 5. Illustration for seed-to-voxel-wise static EC (s-EC) and dynamic EC (d-EC). (A) Group-averaged s-EC map following linear residual-based Granger causality analysis (GCA) using full length of resting-state BOLD fMRI signal with a seed placed in the PCC (MNI coordinates: $-2, -48, 28, 6$ mm radius sphere). (B) d-EC map following sliding-window analysis of linear residual-based GCA with the PCC seed of a representative healthy subject. One target ROI with 6 mm radius sphere placed in the mPFC (MNI coordinates: $3, 39, 18$). The d-EC (solid line) and s-EC (dashed line) time series of mPFC from the in map (red line) and out map (blue line) vary across time. Warm and cool colors indicate brain regions with in influence (from whole brain to the seed) and out influence (from seed to whole brain), respectively. Color scales represent group-averaged GC value (F) of s-EC map and individual $\beta$ amplitude values of d-EC map, respectively. See Supplementary Movies S2 and S3. Color images available online at www.liebertpub.com/brain
Node strength was computed as the sum of the weights of all the connections of a given ROI. Finally, s-FC network was visualized using the BrainNet Viewer (www.nitrc.org/projects/bnv/) (Xia et al., 2013) (Fig. 6A). We observed that a high degree of integration within each sub-network and anti-correlation between the DMN and the DAN and FPCN. The PCC and mPFC exhibited the high node strength, considering as core hubs. These findings are consistent with the previous studies (Fox et al., 2005; Spreng et al., 2013).

For d-FC analysis, the FLS analysis strategy was also used. d-FC network organizations of one representative subject are shown in Figure 6B (see Supplementary Movie S4). As seen from d-FC network dynamics, the fundamental organization, that is integration within network and fractionation between networks, is relatively stable across time (Fig. 6B, upper panel). However, the hub with high regional weights changed hands several times (Fig. 6B, bottom panel), suggesting greater FC variability in the hub for the cognitive process switch. To examine the FC dynamics of core hub PCC, we selected three target ROIs. They are left posterior inferior parietal lobule (pIPL.L), right middle temporal motion complex (MT.R), and right dorsolateral prefrontal cortex (dlPFC.R) in the DMN, DAN, and FPCN, respectively. As seen from the d-FC time series, connectivity between the PCC and pIPL.L (solid red line) is less nonstationary; while that between the PCC and MT.R and between the PCC and dlPFC.R is relatively high nonstationary. This d-FC finding would suggest that some brain regions reveal dual-aligned properties (Spreng et al., 2013).

Quantification of disrupted dynamical connectome in diseased brain would better understand the evolution of disorder. In the current work, we aimed to observe the transition of whole brain connectome that account for absence seizure onset and offset. To this end, the voxel-to-voxel-based d-FC network was constructed for a representative patient with absence epilepsy.

The patient underwent simultaneous EEG-fMRI session by an MR-compatible EEG recording system (Brain Products). During fMRI acquisition, EEG data were continuously recorded through a 10/20 systems with 32 Ag/AgCl electrodes attached to the scalp with conductive cream. EEG electrodes were connected to a BrainAmp amplifier, with a sampling rate of 5 kHz. The EEG data were processed offline to filter out MR artifacts and remove ballistocardiogram artifacts (Brain Vision Analyzer 2.0). Onset and end time of epileptic discharges were marked and classified according to both spatial distribution and morphology. For more detail about the EEG dataset, see our previous studies (Liao et al., 2013a; Zhang et al., 2014). The resting-state fMRI data of patient with absence epilepsy were preprocessed in line with the healthy subject, except to re-sliced at a resolution of $6 \times 6 \times 6$ mm$^3$ to minimize storage and computational requirements. Voxel-to-voxel analysis computed the bivariate FC between every pair of voxels without using a priori seed/ROI. We used the FLS analysis strategy here. In this

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case, we obtained the FCS maps for each time points as shown in Figure 7 (see Supplementary Movie S5). According to information from simultaneously collected EEG data, the FCS maps were separated into preictal (time before seizure onset), ictal, and postictal (time after seizure end) time periods (Liao et al., 2013a). As previously suggested, the thalamus and PCC were involved in seizure initiation, maintenance, and termination during absence seizures. We selected these two core brain regions as the ROIs to track the FCS dynamics. As seen from d-FCS network dynamics, there is a higher FCS of the thalamus (THA) during the ictal period relative to the periods before and after seizures (blue line). Conversely, the PCC d-FCS time series are lower during the ictal period (red line). These findings suggest that the total connections of thalamus and the PCC relate to mechanisms of seizure generation and suspension of default mode of brain function is consistent with an inhibitory effect of seizures on the default mode of brain function, respectively.

### Discussion

We have presented the DynamicBC, a new MATLAB toolbox for the analysis of dynamic brain connectome from multiple functional neuroimaging. In contrast to now available toolboxes, such as, GIFT (Allen et al., 2014), eConnectome (He et al., 2011), BSMART (Cui et al., 2008), and Conn (Whitfield-Gabrieli and Nieto-Castanon, 2012), we now understand that the DynamicBC encompasses both d-FC and d-EC indexes. The aim of the toolbox is to develop a user-friendly GUI for accessibly, and to analyze more comfortably the dynamic brain connectome in which a task or stimulus responses and spontaneous brain activity are measured. To our knowledge, the voxel-level whole brain (GC density/strength) index is firstly available in this toolbox, which could be used to identify the hub of incoming and outgoing information transferring (Wu et al., 2013b).

Particularly, in the current work, we employ FLS algorithm for uncovering the time-varying coefficients of a regression model. Comparing to sliding-window strategy in which the window length is hard to priori define to statistical validation (Hutchison et al., 2013a), the FLS algorithm is a data-driven approach to rapidly assess connectivity changes. In addition, the well-known methodology KF will be added to obtain a comparable result of the transient dynamics (Kang et al., 2011). Meanwhile, FLS and KF model based d-EC algorithms will provide new options to capture the dynamic connectomes.

The present toolbox not only illuminate how much d-FC varies over a scan, but also provide whether the range of d-FC time series variability is significantly different between two populations or between particular regions. The formal group comparisons are challenge for dynamic brain connectome, due to the multi temporal-dimensional nature of the outputs (Hutchison et al., 2013a). According to previous studies, we presented two ways for group analysis. The first sophisticated one, clustering method, collapse the temporal dimension of dynamic connectivity maps/matrices into several outputs or a single one (Allen et al., 2014; Liu and Duyn, 2013). Then, we could perform routine statistical tests between one or more conditions. The second simple one, the variance of the d-FC/d-EC time series (as a proxy of how “stable” a connection is) was calculated automatically. This quantification of dynamic brain connectivity would lead to an improved understanding of the physiological processes of the intact brain or neuropathologic mechanism of the diseased brain.

Another issue related with the recovery of EC networks from BOLD fMRI signal is the possibly confounding effect of the hemodynamic response. To decouple the neuronal activity and the hemodynamic responses, we suggest applying a blind deconvolution procedure, based on the detection of pseudo-events, to the BOLD fMRI signal (Wu et al., 2013a). This blind deconvolution codes have been released and freely available at http://software.incf.org/software/blind-hrf-retrieval-and-deconvolution-for-resting-state-bold.

There are some future directions for the toolbox. Only bivariate dynamic connectivity index are implemented. First, the multivariate or blockwise connectivity measures (Wu et al., 2011) will be integrated in future releases of the tool. Second, implementation of point process event-related nonstationary dynamics of spontaneous activity would be desirable (Tagliazucchi et al., 2012a; Wu et al., 2013a). Third, combining the
different EEG rhythms covary with fMRI connectivity over time (Chang et al., 2013; Tagliazucchi et al., 2012b) will be implemented in the future in the toolbox. Finally, we will add a visualization module for dynamic connectome movie.

**Conclusion**

The DynamicBC toolbox offers a user-friendly and integrated framework to tracking brain connectivity by FC and EC. We provide two brain dynamic analysis strategies and three ways to set connections measures. The illustrative results showed the dynamics of FC patters or network in healthy brain and the whole brain connectome evolved with absence seizures. The current version is freely available at www.restfmri.net/forum/DynamicBC. Users would raise questions and give comments by email (dynamicbrainconn@gmail.com) and online forum (www.restfmri.net/forum/). We hope that this toolbox would make the dynamic brain connectome technique easier to develop, and the clinicians will be benefited from the contribution of the toolbox.

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