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Abstract. Executive functions (EFs) associated with the frontal lobe are vital for goal-orientated behavior. To date, limited efforts have been made to examine the relationships among the behavior, brain activation, and topological organization of functional networks in the frontal lobe underlying various EF tasks, including inhibition, working memory, and cognitive flexibility. In this study, functional near-infrared spectroscopy neuroimaging technique was used to systematically inspect the differences in the brain activation and the topological organization of brain networks between various EF tasks in the frontal lobe. In addition, the relationships between brain activation/network properties and task performances and the relationships between brain activation and network properties were, respectively, examined for different EF tasks. Consequently, we have discovered that the nodal and global properties of the resting-state and task-evoked networks, respectively, exhibited significant correlations with the activation of various brain regions during various EF tasks. In particular, the measure that links the neural activation to the topological organization of the brain networks in the frontal lobe can serve as a synergistic indicator to examine the difference between various EF tasks, which paves a way toward a comprehensive understanding of the neural mechanism underlying EFs.

Keywords: functional near-infrared spectroscopy; behavior; network; activation; executive functions; frontal cortex.

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1 Introduction

Executive functions (EFs) refer to a set of cognitive processes that are essential for the cognitive control of behavior. In particular, EFs implicate the top-down control processes, which are also vital for physical and mental health and academic and job performances.1 EFs consist of three basic cognitive processes, such as inhibition, working memory, and cognitive flexibility, in which inhibition involves resisting dominant response and focusing on the present task, and working memory is associated with holding and operating verbal or nonverbal information in mind, whereas cognitive flexibility is related to shifting between multiple tasks and adjusting the behavior appropriately to a changing circumstance.2 It is widely recognized that EFs are regulated by the frontal lobe,1–3 whereas frontal lesions or dysfunctions can cause a wide range of cognitive deficits and mental diseases, including dyslexia,4 dyscalculia,5 obsessive-compulsive disorder,6 attention-deficit hyperactivity,7 and schizophrenia.8

Interestingly, recent neuroimaging and neuroscience studies demonstrate that the different brain regions in the frontal cortex are responsible for the various cognitive processes of EFs. For example, previous reports have shown that the attentional inhibition (measured by the flanker task) elicited the activation in the supplementary motor area (SMA), the right ventrolateral prefrontal cortex, the superior frontal gyrus, and the left superior/anterior parietal lobe9,10 whereas the response inhibition induced the activation in the right inferior frontal gyrus (IFG) and the right pre-SMA11–14 More important, during the response inhibition task, it has been discovered that the superior part of anterior frontal cortex exhibited reduced activation as compared to that at rest.15,16 In addition, working memory is considered to be linked to the functions of the dorsolateral prefrontal cortex (DLPFC).17–19 In addition, a meta-analysis study has revealed that the lateral and medial regions of the frontopolar area (FPA) get activated during the shifting between cognitive rules.20 An additional study has also illustrated that the FPA plays an essential role in monitoring multiple goals in parallel and in switching between them.21

In addition to the study of brain activation patterns, functional connectivity (FC) analyses have become an increasingly important tool for examining EFs.22 In particular, significant progress has been made in the characterization of the human brain networks, in which graph theory-based network analysis is introduced to describe more comprehensive properties of the functional topological organization of brain networks, such as the small-worldness.23,24 More importantly, the brain FC during a task is modulated primarily by the intrinsic resting-state network architectures, and then secondarily by the task-evoked networks.25 In addition, a previous study has demonstrated that more globally integrated networks enable fast and effective performance during an n-back task.26 Further, a recent functional near-infrared spectroscopy (fNIRS) study associated with cognitive flexibility has examined the relationship between...
topological properties of the resting-state network in the prefrontal cortex and EF scores,
which they have discovered that the total EF scores negatively correlated with regional network properties in the right triangular IFG, whereas the EF scores positively correlated with regional properties in the right dorsal superior frontal gyrus. In addition, Zhao et al.’s study illustrated that the working memory is related to the regional network properties in the right middle frontal gyrus and triangular IFG, whereas the cognitive flexibility is associated with the regional properties in the right middle frontal gyrus.

Although the brain activation and FC were, respectively, examined in a single study,28–31 the relationship between these two perspectives has not been extensively explored. For example, Mennes et al.32 inspected the relationship between resting-state FC and brain activation by using the flanker task, in which they discovered that the resting-state FC within default mode networks showed a negative correlation with the task-evoked brain activation, whereas the task-based FC positively correlated with the brain activation. An additional study based on the visual attention tasks also demonstrated that the local FC density, which mainly decreased in weakly activated/deactivated brain regions, was associated with better behavioral performance and activation in the task-related networks.33

However, to the best of our knowledge, no systematic study has been performed to examine the relationship between the brain activation and the topological properties of FC during the EF tasks. As a result, it is hypothesized for the present study that the linking between the brain activation and the task-based/resting-state FC can serve as a neural index, which might identify the significant difference between various cognitive processes of EFs, including inhibition, working memory, and cognitive flexibility.

To test this hypothesis, the brain activation and resting-state/task-based FC are accessed based on four classic EF tasks (go/no-go, flank, n-back, and switch). The cortical hemodynamic changes at rest are first acquired and then task-based recordings are also performed for the investigation of EFs with fNIRS. The fNIRS is an emerging noninvasive optical neuroimaging technique, which can offer unsurpassed temporal resolution (50 Hz for the present system) and characterize the hemodynamic fluctuations more precisely. In addition, fNIRS can be operated in a more comfortable, portable, quiet, and natural way with very few body constraints and low cost.34 In particular, the brain activation and FC, in this study, are first individually explored and compared among various cognitive processes of EFs. Further, we also carefully examine how the activation and FC network are individually associated with the behavior performances of four various EF tasks. More important, we inspect the relationship between the activation and the topological properties of brain functional networks during tasks or at rest. As the linking between the brain activation and the network properties of FC has never been systematically explored for cognitive processes of EFs, this fNIRS work will pave a new avenue toward a better understanding of the cognitive mechanism of EFs.

2 Methods and Materials

2.1 Participants

A total of 30 college students (15 females and 15 males) were recruited from the campus of the University of Macau, China. Participants aged from 18 to 28 years (mean age: 23.2 years) had normal or corrected-to-normal vision without histories of learning disabilities, medical illness, neurological, or psychiatric disorders. Informed consent documents were signed prior to the experiment. This study was approved by the Medical Ethics Committee with the University of Macau.

2.2 Procedures and Tasks

The experimental tests were performed in a quiet room. 11-min resting-state data were acquired before the performance of EF tasks, which were utilized to construct the intrinsic functional networks in the frontal lobe. During the resting-state recordings, participants were instructed to close their eyes, remain still, and avoid falling asleep. Subsequently, participants went through four typical EF tasks related to different EFs components, with a short break between any two tasks with the time period determined by participants. These four tasks consisted of the flanker task, go/no-go task, n-back task, and switch task. In particular, there were two conditions for each of the four tasks, in which the incongruent, no-go, 2-back, and switch condition, respectively, were the experimental conditions, whereas the congruent, go, 0-back, and repeat condition, respectively, were the control conditions. The schematic of experimental procedures of the four tasks is illustrated in Fig. 1.

For the present experimental design, participants performed the flanker task first after 11-min resting-state recordings, followed by the go/no-go task. In contrast, the switch and n-back tasks were arranged as the last sections for data acquisition in order to reduce the possible influence of difficulty in performing the tasks. However, the order of the switch and n-back tasks was not well counterbalanced, whose effect should be further investigated in the future.

2.2.1 Flanker task

The flanker task was adopted to inspect the inhibition at the attention level.38,39 The event-related stimuli were presented trial by trial. Each trial lasted 15 s, which consisted of a 0.5-s stimulus period, a 1.5-s presentation of a blank screen, and a 13-s rest period with a fixation cross displayed in the center of the monitor. The flank task had two congruent stimuli including “< < < < < <” and “> > > > > >,” and two incongruent stimuli including “< < < < < <” and “> > > > > >.” When the central arrow points to the left (“<”), the participants were required to press the left button, whereas they had to press the right button if it points to the right (“>”). Stimuli of each type were randomly repeated 24 times.

2.2.2 Go/no-go task

The go/no-go task was utilized to study the inhibition at the response level.40,41 The procedure of go/no-go task was very similar to the event-related flanker task, in which a series of English letters were presented. Participants were required to press the space key with their right forefinger as soon as the letter appeared (go condition), except the letter “X” (no-go condition), which needed the participants to withhold their response.

2.2.3 N-back task

The n-back task was used to investigate the working memory.37,42,43 A block-design paradigm was adopted, which contained six 0-back blocks and six 2-back blocks. Each block
had 12 trials, and each trial started with a stimulus period of 0.5 s with a letter in the center of the monitor, followed by a 1.5-s blank screen (interstimulus interval). Participants need to respond to the stimuli by pressing a key with their forefinger. The stimuli materials consisted of a bunch of English letters, which were presented randomly. During the 0-back condition, participants responded to the letter “A” (target), whereas during the 2-back condition, participants needed to respond to any letter that was identical to the one presented two trials before (target). During the n-back task, participants were required go through 0-back and 2-back blocks in turns, and between the predecessor and the successor blocks, there was a 30-s rest period.

2.2.4 Switch task

The switch task was used to inspect cognitive flexibility. The procedure of switch task was quite similar to that of the flanker task, except for an additional 50-ms cue presented before the onset of the stimulus, which demonstrated the stimulus conditions (odd/even or odd/even). The stimuli materials consisted of a serial of Arabic numbers ranged from 0 to 9. For an individual trial, a single letter was presented with a background shaped either like a square (odd/even condition) or like a diamond (low/high condition). Participants were instructed to determine whether the number was odd/even or low/high with a left or right key-press. It should be noted that in the low/high condition, the number 0 to 4 was classified as “low,” whereas 5 to 9 was classified as “high.” The trials were presented randomly either as switch trials or as repeat trials. In switch trials, the task was switched from the former one, whereas in repeat trials, the task was the same as the former one.

2.3 Functional Near-Infrared Spectroscopy Data Acquisition and Preprocessing

The fNIRS signals were acquired at a sample rate of 50 Hz by using a continuous wave (CW) system (CW6 fNIRS system; TechEn Inc., Milford, Massachusetts). The optode arrays comprised 8 laser sources at wavelengths 690 and 830 nm and 16 optical detectors, which generated 32 measurement channels. We used a homemade plastic patch to hold the optodes with a fixed interoptode distance of 3 cm and then placed the patch on each participant’s head to cover the frontal lobe, as displayed in Fig. 2(a). After the experiment, a three-dimensional (3-D) digitizer (PATRIOT, Polhemus, Colchester, Vermont) was used to measure the 3-D coordinates of the optodes, which were further imported to NIRS_SPM44 for spatial registration to generate the 3-D coordinates of all channels. In addition, the distributions of the optodes and channels, mean MNI coordinates of the channels, and anatomical labels are provided in Fig. 2 and Supplementary Material in Table S1, respectively.

Before data preprocessing, the signal quality of each channel was accessed by using the signal-to-noise ratio (SNR), which was the ratio of the mean to standard deviation of the signal intensity. According to Rose criterion,45 data from participants with an SNR of <5 for both wavelengths were excluded for further analysis as the low SNR could undermine the corresponding correlations and network properties.46 The SNRs of bad channels from each wavelength are listed in Table S2 in the Supplementary Material for the excluded participants.

The preprocessing of fNIRS data was performed with Homer2 software:47 (1) converting the raw data to the changes in optical density, (2) removing motion artifacts by spline method,48 (3) bandpass filtering (for filtering parameter
selection, resting state: 0.01 to 0.08 Hz, EF tasks: 0.01 to 0.2 Hz), (4) using principal component analysis to remove the first one or two eigenvectors accounting for 80% of the variance in the optical data, which was mainly contributed by heart rate and blood pressure oscillations, (5) normalizing the concentration changes by subtracting the mean channel-wise concentration, which were further processed by dividing the standard deviation (z-scores), and (6) providing baseline correction with the period from \(-2\) to 0 s for a single trial. The adopted frequency band was used to filter out the high-frequency physiological noises, such as heart beat (~0.8 Hz), cardiac cycles (~1 Hz), and breathing (~0.2 Hz), as well as the slow-frequency physiological drifts. In addition, for resting-state data, data from 180 to 600 s were extracted before the preprocessing to ensure stable signals, which were then detrended after the preprocessing. The method for data preprocessing was in line with those from previous studies. Importantly, both the HbO and HbR signals can be used to characterize the activation and functional network properties of the frontal lobe during the rest or EF tasks. However, most of previous fNIRS work only reported the neuroimaging results based on HbO signals.

In this study, both HbR and HbO data were analyzed to generate the brain activation and construct the brain networks.

2.4 Brain Activation

After the preprocessing, the grand-averaged HbO and HbR data were calculated for each condition associated with each task across all participants. Then, the mean z-scores of HbO and HbR signals during the stimuli period (for the n-back task: 0 s to 26 s; for the other EF tasks: 0 to 13 s) were calculated to map the brain activation for each channel. In particular, we discovered that for most of the cases the task-related brain regions exhibited increased HbO and decreased HbR concentrations.

2.5 Network Topological Properties

The functional networks in frontal lobe were constructed, in which resting-state and task-evoked FC matrices were generated by computing the Pearson correlations between pairs of channels for each participant. In addition, the network topological
properties of constructed matrices were analyzed using GRETNA.61

The network topological properties are further characterized by seven global parameters and three nodal parameters.62-64 The global parameters denote the efficiency of the whole network, which consist of the average clustering coefficient ($C_p$), average characteristic path length ($L_p$), normalized clustering coefficient ($\gamma$), normalized characteristic path length ($\lambda$), small-worldness ($\sigma$), local efficiency ($E_{loc}$), and global efficiency ($E_{glob}$). The nodal parameters represent the efficiency of nodes, which include the nodal degree ($K_{nod}$), nodal efficiency ($E_{nod}$), and nodal betweenness centrality ($BC_{nod}$). In brief, $C_p$ and $\gamma$ measure the local interconnectivity of a network, whereas $L_p$ and $\lambda$ quantify the overall routing efficiency of a network. In addition, $E_{loc}$ plays a role similar to the clustering coefficient, whereas $E_{glob}$ is inversely related to the characteristic path length.62-64 Further, networks with higher $\sigma$ are regarded as systems that are both locally and globally efficient. In contrast, for nodal topological properties, $K_{nod}$ denotes the number of edges linked to a node, which measures the association capability of the given node. However, $E_{nod}$ is inversely correlated with the averaged shortest path length from a given node to the other nodes, which measures the nodal efficiency of information transfer. In addition, $BC_{nod}$ denotes the frequency of a given nodal showing up in a shortest path from one node to another, which exhibits the importance of a given node over the information flow. It is also noted for the present study that the threshold is determined by the network density (sparsity) of the brain networks, which is defined by the ratio ($S$) of the existing edges divided by the maximum possible edges in the network.65 In addition, the global properties of brain networks only represent the information transmission pattern in frontal lobe, whereas the nodal properties denote the role of each channel in the information transmission associated with the frontal lobe network.

Interestingly, previous studies have shown that the resting-state brain functional networks have prominent small-world properties as well.25,46,66 In this study, one sample t-tests against 1.1 were performed on the small-worldness of the resting-state or task-based networks constructed from HbR and HbO data, respectively. The maximum threshold $S$ was determined as 44% to ensure the prominent small-worldness of both the networks.67 In addition, the minimum threshold was determined by the average degree over all nodes $K_{average} > 2 \ln(N)$ with $N = 32$, denoting the number of the nodes, which can guarantee that a random graph is connected.64 The threshold range determined by this procedure was $23% \leq S \leq 44%$.

### 3 Results

#### 3.1 Behavioral Results

The mean accuracy of the responses and reaction time for correct responses were, respectively, calculated for each participant. The accuracy and reaction time in each task were, respectively, subjected to paired t-test to reveal the performance difference between the experimental and the control conditions.

The t-test results are summarized in Table 1, which shows that all experimental cases consumed more reaction time than the control cases ($p < 0.01$). Meanwhile, lower accuracy is respectively detected in the flanker ($p = 0.01$), go/no-go ($p < 0.01$), and n-back task (marginally significant) as compared to that from the control case.

#### 3.2 Brain Activation

In addition, we compared the differences in brain activation between the experimental and the control conditions for each task.

The averaged HbO/HbR time courses for the four EF tasks are displayed in the Supplementary Material in Figs. S1–S4. The brain t-map, generated from the activation difference between the experimental and the control conditions for each task, is visualized in the first two rows of Fig. 3 by using xjView toolbox68 and BrainNet Viewer toolbox69,70 and the detailed t-test results are provided in Table 2. In addition, the Pearson correlation is performed to examine the relationship between the HbO and the HbR activation, and the correlation coefficients are visualized in the third row of Fig. 3.

Figure 3 and Supplementary Material in Figs. S1–S4 show that the brain activation differences between the experimental and the control conditions exhibit different tendencies for the four tasks although only the flanker and switch tasks exhibit significant differences. Specifically, the incongruent case in flanker task induces increased HbO in the bilateral prefrontal and supplementary motor cortex (MC), whereas the switch case

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**Table 1** The t-test results of behavioral data.

| Contrast                  | t   | df  | p     | d    | M1  | SE1 | M2  | SE2 |
|---------------------------|-----|-----|-------|------|-----|-----|-----|-----|
| flk_con_acc—flk_incon_acc | 3.58| 29  | 0.001 | 0.92 | 0.99| 0.01| 0.92| 0.02|
| gng_go_acc—gng_nogo_acc   | 4.12| 29  | <0.001| 1.14 | 0.99| <0.01| 0.96| 0.01|
| nbk_b0_acc—nbk_b2_acc     | 1.92| 29  | 0.065 | 0.47 | 0.99| 0.01| 0.96| 0.01|
| swt_rep_acc—swt_SW_acc    | 0.37| 29  | 0.715 | 0.08 | 0.93| 0.01| 0.92| 0.01|
| flk_con_rt—flk_incon_rt  | −8.45| 29  | <0.001| −0.85| 0.57| 0.02| 0.70| 0.03|
| nbk_b0_rt—nbk_b2_rt      | −6.80| 29  | <0.001| −1.61| 0.47| 0.01| 0.60| 0.02|
| swt_rep_rt—swt_SW_rt     | −5.61| 29  | <0.001| −0.50| 0.95| 0.03| 1.03| 0.03|

Note: Abbreviations: acc, accuracy; rt, reaction time; t, t-value of the paired t-test; df, degree of freedom; d, Cohen’s d; M, mean value, SE, standard error, 1, 2 means control and experimental cases, respectively; flk, flanker task; con, congruent; incon, incongruent; gng, go/no-go task; nbk, n-back task; b0, 0-back condition; b2, 2-back condition; swt, switch task; rep, repeat condition; and SW, switch condition.

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in the switch task elicits increased HbO and decreased HbR in the right FPA and decreased HbO in the left frontal eye fields (FEFs). The decreased HbR is still observed even after the false discovery rate (FDR) correction. In addition, although no significant difference between the two conditions is revealed for the go/no-go and n-back tasks, tendencies toward enhanced HbO activation for the 2-back case are observed across the task period in Supplementary Material in Fig. S3 as compared to that from the 0-back in left DLPFC (channels 9, 11, and 21). In particular, Supplementary Material in Fig. S2 shows that the no-go case elicits HbR/HbO activation for most channels in the frontal lobe, which is also comparable to the go case.

**Table 2** Summary of the t-test results with regard to the brain activation differences.

| Task      | #ch | Label | p(fdr) | p   | t    | df | SD  | d   | M1  | SE1 | M2  | SE2 |
|-----------|-----|-------|--------|-----|------|----|-----|-----|-----|-----|-----|-----|
| HbO Flanker (incon–con) | 28  | L-MC  | 0.388  | 0.033 | 2.27 | 24 | 0.24 | 0.40 | 0.18 | 0.05 | 0.07 | 0.05 |
|           | 30  | R-MC  | 0.388  | 0.048 | 2.09 | 24 | 0.31 | 0.45 | 0.22 | 0.05 | 0.09 | 0.06 |
|           | 31  | R-MC  | 0.388  | 0.029 | 2.33 | 24 | 0.31 | 0.44 | 0.19 | 0.06 | 0.04 | 0.07 |
| Switch (swt–rep) | 14  | L-FEF | 0.790  | 0.049 | −2.07 | 24 | 0.22 | −0.33 | 0.07 | 0.05 | 0.16 | 0.06 |
|           | 6   | R-FPA | 0.574  | 0.018 | 2.54 | 24 | 0.23 | 0.47 | 0.33 | 0.05 | 0.21 | 0.05 |
| HbR Switch (swt–rep) | 6   | R-FPA | 0.005  | <0.001 | −4.50 | 24 | 0.22 | −0.84 | −0.21 | 0.04 | −0.01 | 0.05 |
|           | 8   | R-FPA | 0.077  | 0.005 | −3.10 | 24 | 0.20 | −0.53 | −0.07 | 0.04 | 0.05 | 0.05 |

Note: Abbreviations: #Ch, the label of the channel; incon, incongruent; con, congruent; swt, switch; rep, repeat; L, left hemisphere; R, right hemisphere; t, t-value of the paired t-test; df, degree of freedom; d, Cohen’s d; M, mean value; SE, standard error (1, 2 means the experimental and control cases, respectively).
In addition, the negative correlations between HbO and HbR activations are clearly identified by the widespread dark color in the correlation maps in Fig. 3.

3.3 Functional Network Properties

3.3.1 Functional connectivity

The grand-averaged correlation coefficients for each condition of the four tasks and for the resting state are visualized in the matrix form in Supplementary Material in Figs. S5 and S6 for the HbO and the HbR measures, respectively. In addition, the global and nodal topological properties of networks are calculated and compared for various test cases.

3.3.2 Small-world properties

Whether the task-based or resting-state networks exhibited small-world property (namely $\sigma > 1$) is inspected in this section. One sample $t$-tests are performed on the small-worldliness for the five individual test cases with selected thresholds ($23\% \leq S \leq 44\%$). Figure 4 shows that the brain functional networks during all test cases exhibit the prominent small-world property.

3.3.3 Global properties

Subsequently, we calculated the area under property curves (AUC) over the predetermined thresholds for each test case and compared the AUC between the networks during the tasks and at rest, and also between the experimental and the control conditions for the four tasks. In addition, paired $t$-tests were performed on all topological properties, including the global and nodal properties.

First, the global properties of the networks for each individual test case are compared to those at rest, and the comparison results are provided in Table 3. Interestingly, Table 3 shows that the global properties of HbR-based networks are more sensitive to the task tests as compared to those from the HbO-based ones. For example, only the 2-back case in $n$-back task exhibits decreased $C_p$ and $E_{\text{loc}}$ for HbO-based networks when compared to those at rest. However, this is not the case for HbR-based networks, in which all task-based test cases exhibit decreased local interconnection including decreased $C_p$, $\gamma$, and $E_{\text{loc}}$. In addition, the congruent case in flanker task also exhibits increased overall routing efficiency including decreased $L_p$ and $\lambda$, and increased $E_{\text{glob}}$ as compared to the rest. In contrast, both cases in the go/no-go task and experimental cases in then-back and switch tasks exhibit decreased small-worldness ($\sigma$).

In addition, we also examined the differences in global properties of the networks between the experimental and the control conditions associated with the four tasks. Interestingly, we discovered that only HbR-based networks exhibited significant differences in nodal properties. For example, significantly decreased $BC_{\text{nod}}$ was clearly identified in the left MC (channel 29) for HbO-based networks for all task-based tests, although it was not that significant for the go case. In contrast, HbR-based networks showed decreased network efficiency, including the betweenness, nodal degree, and nodal efficiency in the left FEF (channels 12 or 14) for all task-based tests although it was not significant for some nodal properties associated with the flanker task.

In addition, we also compared the differences in network nodal properties between the experimental and the control conditions. The comparison results are given in Table 4. Regarding the HbO-based networks, the incongruent case from the flanker task exhibited decreased $K_{\text{nod}}$ in the right FPA, and increased $K_{\text{nod}}$ and $E_{\text{nod}}$ in the right FEF. In the $n$-back task, the 2-back case showed decreased $K_{\text{nod}}$ and $E_{\text{nod}}$ in the left MC and increased $BC_{\text{nod}}$ in the right FPA, whereas the switch case

![Fig. 4 Small-worldness of the brain network for each test case. Among the selected threshold (sparsity: 23% to 44%), the brain networks that exhibit the small-world property ($\sigma > 1$) are labeled with asterisks with corresponding colors.](https://www.spiedigitallibrary.org/journals/Neurophotonics)
from switch task showed decreased $K_{nod}$ in the left FPA and decreased $E_{nod}$ in the left FPA and right FEF.

Further, it was observed that the HbR-based networks were more sensitive to the EF demands. In the flanker task, the incongruent case exhibited increased $BC_{nod}$, $K_{nod}$, and $E_{nod}$ in the right DLPFC, and decreased $BC_{nod}$ in the left FPA and FEF and right MC, decreased $K_{nod}$ in the right FPA and left FEF, and decreased $E_{nod}$ in the right FPA. In the $n$-back task, the 2-back case exhibited increased $BC_{nod}$ in the left DLPFC and FEF and right FPA. In addition, increased $K_{nod}$ and $E_{nod}$ were identified in the right DLPFC, whereas decreased ones were revealed in the left FPA. In the switch task, we discovered that the anterior part of left DLPFC (channel 10) and its nearby regions exhibited increased $BC_{nod}$, $K_{nod}$, and $E_{nod}$, whereas the posterior part of left DLPFC (channel 11) and its nearby regions exhibited decreased $BC_{nod}$, $K_{nod}$, and $E_{nod}$.

### Table 3  Significant changes in global parameters for each condition from each task.

| Condition | Para. | $p$  | $t$  | $df$ | $d$  | M1  | SE1 | M2  | SE2 |
|-----------|-------|------|------|------|------|-----|-----|-----|-----|
| HbO       |       |      |      |      |      |     |     |     |     |
| 2-Back    | Cp    | <0.01| −3.17| 24   | −0.85| 0.64| 0.01| 0.68| 0.01|
|           | Eloc  | 0.02 | −2.57| 24   | −0.72| 0.75| 0.01| 0.78| 0.01|
| HbR       |       |      |      |      |      |     |     |     |     |
| Congruent | Gamma | 0.03 | −2.37| 23   | −0.77| 1.43| 0.03| 1.58| 0.05|
| Incongruent| Cp    | <0.01| −3.58| 23   | −1.04| 0.61| 0.01| 0.67| 0.01|
|           | Lp    | 0.02 | −2.41| 23   | −0.70| 1.62| 0.02| 1.68| 0.02|
|           | Lambda| 0.01 | −2.80| 23   | −0.77| 1.04| 0.01| 1.07| 0.01|
|           | Gamma | 0.02 | −2.48| 23   | −0.77| 1.42| 0.03| 1.58| 0.05|
|           | Eloc  | 0.02 | −2.55| 23   | −0.74| 0.76| 0.01| 0.79| 0.01|
|           | Eglob | 0.02 | 2.52 | 23   | 0.73 | 0.63| 0.01| 0.61| 0.01|
| Go        | Gamma | 0.01 | −2.93| 24   | −0.87| 1.40| 0.03| 1.58| 0.05|
|           | Sigma | 0.01 | −2.80| 24   | −0.80| 1.32| 0.03| 1.47| 0.04|
| No-go     | Gamma | 0.01 | −2.93| 24   | −0.78| 1.42| 0.03| 1.58| 0.05|
|           | Sigma | 0.01 | −2.65| 24   | −0.74| 1.33| 0.03| 1.47| 0.04|
| 0-Back    | Cp    | <0.01| −3.11| 24   | −0.86| 0.61| 0.01| 0.67| 0.01|
|           | Gamma | 0.03 | −2.27| 24   | −0.70| 1.42| 0.04| 1.58| 0.05|
|           | Eloc  | 0.01 | −2.99| 24   | −0.86| 0.75| 0.01| 0.79| 0.01|
| 2-Back    | Cp    | <0.01| −3.56| 24   | −0.88| 0.62| 0.01| 0.67| 0.01|
|           | Gamma | 0.02 | −2.54| 24   | −0.69| 1.44| 0.03| 1.58| 0.05|
|           | Sigma | 0.05 | −2.10| 24   | −0.54| 1.37| 0.03| 1.47| 0.04|
|           | Eloc  | 0.02 | −2.56| 24   | −0.73| 0.76| 0.01| 0.79| 0.01|
| Repeat    | Cp    | 0.03 | −2.28| 23   | −0.74| 0.63| 0.01| 0.67| 0.01|
|           | Gamma | 0.03 | −2.32| 23   | −0.63| 1.44| 0.03| 1.57| 0.05|
| Switch    | Gamma | 0.02 | −2.54| 23   | −0.66| 1.44| 0.03| 1.57| 0.05|
|           | Sigma | 0.04 | −2.19| 23   | −0.59| 1.35| 0.03| 1.46| 0.04|

### 3.4 Relationship between the Brain Activation/Network Topological Properties and Task Performance

Subsequently, the Pearson correlation between the cognitive cost (measured by the increase in reaction time) and the brain activation/FC was also generated in this study, which can be used to examine the influence of the brain activation/networks on the task performance. As a result, the task-evoked activation and topological properties of FC were used as the measures to generate the correlations with the cognitive costs in each task. In particular, the task-evoked changes were denoted as the differences in HbO or HbR activation/topological properties between the experimental and the control conditions, whereas the cognitive cost was quantified by the relative reaction time between the experimental and the control conditions.
Table 4 Summary of the t-test results with regard to the AUC of nodal parameters for each task.

| Task         | Para. | Label | #Ch | p(FDR) | p | t  | df | d   | M1  | SE1 | M2  | SE2 |
|--------------|-------|-------|-----|--------|---|----|----|-----|-----|-----|-----|-----|
| HbO Flanker (incon-con) | K     | R-FPA | 6   | 0.76   | 0.05 | −2.09 | 24 | −0.25 | 10.35 | 1.00 | 11.60 | 0.98 |
|              | K     | R-FEF | 19  | 0.76   | 0.03 | 2.30  | 24 | 0.17  | 11.05 | 1.18 | 10.02 | 1.19 |
|              | K     | R-FEF | 19  | 0.65   | 0.03 | 2.30  | 24 | 0.18  | 0.56  | 0.04 | 0.52  | 0.04 |
|              | E     | L-MC  | 23  | 0.57   | 0.02 | −2.47 | 25 | −0.31 | 6.57  | 1.13 | 8.41  | 1.17 |
|              | E     | L-MC  | 23  | 0.99   | 0.04 | −2.15 | 25 | −0.36 | 0.43  | 0.05 | 0.51  | 0.04 |
| N-back (b2-b0) | BC    | R-FPA | 6   | 0.81   | 0.03 | 2.38  | 25 | 0.48  | 9.38  | 1.71 | 5.75  | 1.18 |
|              | K     | L-MC  | 23  | 0.57   | 0.02 | −2.47 | 25 | −0.31 | 6.57  | 1.13 | 8.41  | 1.17 |
|              | E     | L-MC  | 23  | 0.99   | 0.04 | −2.15 | 25 | −0.36 | 0.43  | 0.05 | 0.51  | 0.04 |
| Switching (swt-rep) | K     | L-FPA | 3   | 0.66   | 0.04 | −2.20 | 24 | −0.19 | 9.86  | 1.24 | 11.01 | 1.17 |
|              | E     | L-FPA | 3   | 0.59   | 0.04 | −2.21 | 24 | −0.16 | 0.53  | 0.05 | 0.57  | 0.04 |
|              | E     | R-FEF | 25  | 0.43   | 0.01 | −2.67 | 24 | −0.19 | 0.52  | 0.04 | 0.56  | 0.04 |
| HbR Flanker (incon-con) | BC    | L-FPA | 1   | 0.32   | 0.03 | −2.39 | 24 | −0.67 | 8.53  | 1.46 | 16.73 | 3.15 |
|              | BC    | L-FEF | 13  | 0.32   | 0.05 | −2.07 | 24 | −0.28 | 9.01  | 1.72 | 11.83 | 2.26 |
|              | BC    | R-DLPFC | 16 | 0.32  | 0.03 | 2.30  | 24 | 0.59  | 25.75 | 2.73 | 17.15 | 3.10 |
|              | BC    | R-MC  | 31  | 0.25   | 0.01 | −2.90 | 24 | −0.62 | 12.10 | 2.28 | 21.90 | 3.84 |
|              | K     | L-FPA | 6   | 0.29   | 0.02 | −2.53 | 24 | −0.52 | 12.37 | 0.60 | 13.91 | 0.59 |
|              | K     | L-FEF | 13  | 0.46   | 0.04 | −2.14 | 24 | −0.38 | 10.07 | 0.58 | 11.42 | 0.83 |
|              | K     | R-DLPFC | 16 | 0.13  | <0.01 | 3.19  | 24 | 0.66  | 14.02 | 0.57 | 11.93 | 0.70 |
|              | E     | R-FPA | 6   | 0.41   | 0.03 | −2.37 | 24 | −0.48 | 0.67  | 0.01 | 0.70  | 0.01 |
|              | E     | R-DLPFC | 16 | 0.03  | <0.01 | 3.78  | 24 | 0.76  | 0.70  | 0.01 | 0.65  | 0.02 |
| N-back (b2-b0) | BC    | R-FPA | 6   | 0.16   | 0.01 | 2.78  | 25 | 0.81  | 21.47 | 3.20 | 10.62 | 1.87 |
|              | BC    | L-DLFPC | 10 | 0.41  | 0.05 | 2.10  | 25 | 0.54  | 19.94 | 2.76 | 13.49 | 1.78 |
|              | BC    | L-FEF | 14  | 0.16   | 0.01 | 2.79  | 25 | 0.74  | 18.69 | 3.06 | 9.59  | 1.54 |
|              | K     | L-FPA | 1   | 0.08   | <0.01 | −3.36 | 25 | −0.65 | 10.82 | 0.91 | 13.51 | 0.71 |
|              | K     | R-DLPFC | 24 | 0.62  | 0.04 | 2.18  | 25 | 0.51  | 12.71 | 0.63 | 10.89 | 0.78 |
|              | E     | L-FPA | 1   | 0.14   | <0.01 | −3.15 | 25 | −0.69 | 0.63  | 0.02 | 0.69  | 0.01 |
|              | E     | R-DLPFC | 24 | 0.57  | 0.05 | 2.10  | 25 | 0.49  | 0.67  | 0.01 | 0.64  | 0.01 |
| Switching (swt-rep) | BC    | L-DLFPC | 10 | 0.43  | 0.03 | 2.25  | 24 | 0.35  | 17.25 | 2.04 | 13.32 | 2.48 |
|              | BC    | L-DLFPC | 11 | 0.43  | 0.03 | −2.38 | 24 | −0.44 | 9.70  | 1.84 | 14.38 | 2.38 |
|              | BC    | L-MC  | 23  | 0.43   | 0.04 | −2.12 | 24 | −0.50 | 8.19  | 1.33 | 13.16 | 2.48 |
|              | K     | L-DLFPC | 10 | 0.53  | 0.05 | 2.07  | 24 | 0.49  | 13.88 | 0.58 | 12.36 | 0.66 |
|              | K     | L-DLFPC | 11 | 0.53  | 0.03 | −2.36 | 24 | −0.30 | 9.70  | 0.85 | 10.96 | 0.81 |
|              | K     | R-FEF | 17  | 0.53   | 0.04 | 2.18  | 24 | 0.45  | 8.99  | 0.79 | 7.50  | 0.51 |
|              | E     | L-DLFPC | 10 | 0.49  | 0.04 | 2.17  | 24 | 0.50  | 0.70  | 0.01 | 0.67  | 0.01 |
|              | E     | L-DLFPC | 11 | 0.10   | <0.01 | −3.30 | 24 | −0.31 | 0.61  | 0.02 | 0.64  | 0.02 |
|              | E     | L-FEF | 13  | 0.49   | 0.05 | −2.11 | 24 | −0.37 | 0.62  | 0.01 | 0.65  | 0.02 |
Note that the go/no-go task was excluded from this analysis because no reaction time was measured in no-go condition. We discovered that only HbR activation (but not HbO) exhibited correlations with the cognitive cost. In particular, in the flanker task, the cognitive cost showed positive correlation with HbR changes in the left FEF, whereas it showed negative correlation in the right FPA. In the n-back and switch tasks, positive correlations were revealed in the right DLPFC. However, after the FDR correction, no significant correlation was identified.

In contrast, for both HbO-based and HbR-based networks, the nodal (but not global) topological properties changes showed significant relationships with the cognitive cost. For example, for HbO-based networks, the cognitive cost was positively correlated with $K_{nod}$ in the left FPA and DLPFC, and positively correlated with $E_{nod}$ in the left DLPFC for flanker task, although it showed negative correlation with $BC_{nod}$ in the left DLPFC and MC. Interestingly, in the n-back task, negative correlations were identified for $K_{nod}$ in the left FEF and for $BC_{nod}$ in the left DLPFC. However, in the switch task, the cognitive cost was positively correlated with $K_{nod}$ in the right FEF, whereas negative correlations were revealed for $K_{nod}$ in the right DLPFC, for $E_{nod}$ in the left MC, and for $BC_{nod}$ in the left FEF and MC and right FPA. In contrast, after the FDR correction, only negative correlation for $BC_{nod}$ was identified in the left DLPFC for flanker task.

In addition, for HbR-based networks, the cognitive cost showed negative correlation with $BC_{nod}$ in the left DLPFC and right MC for flanker task. In the n-back task, positive correlations were discovered between $K_{nod}$ and cognitive cost in right MC and FEF and negative ones for $E_{nod}$ in the bilateral DLPFC. In the switch task, positive correlations were identified for $K_{nod}$ and $E_{nod}$ in the left FPA, whereas negative correlations were identified for $K_{nod}$ and $E_{nod}$ in the left MC, and for $BC_{nod}$ in the left DLPFC and right MC. However, this is not the case after the FDR correction, in which no significant correlation was identified.

### 3.5 Relationship between Brain Activation and Networks Topological Properties

Pearson correlation analysis was performed to examine the relationship between the brain activation and the networks topological properties. In particular, we correlated the topological properties of resting-state networks, and task-based network topological properties with task-based brain activation.

Interestingly, the brain activation during the EF tasks exhibited significant correlations with the network topological properties for most of the channels. However, after the FDR correction, it was clearly observed that the brain activation during flanker, n-back, and switch tasks only showed correlation with the intrinsic network properties. In contrast, in the go/no-go task, the activation not only exhibited the correlation with the topological properties derived from intrinsic networks but also exhibited the properties derived from task-evoked and HbR-based networks.

In particular, in the flanker task, the activation in bilateral DLPFC and left FEF showed negative correlation with $\lambda$ for HbO-based intrinsic networks, whereas the activation in left MC was positively correlated with $K_{nod}$ for the same resting-state networks.

In the n-back task, the activation in right FEF was positively correlated with $K_{nod}$ for HbO-based intrinsic networks. In addition, the activation in right MC was positively correlated with $BC_{nod}$ for HbR-based intrinsic networks, whereas the activation in left MC was negatively correlated with $BC_{nod}$ from the same network. In the switch task, the activation in left FPA was negatively correlated with $C_p$, $L_p$, and $\lambda$, whereas it was positively correlated with $E_{glob}$ in HbR-based intrinsic networks.

In the go/no-go task, the activation in left FEF was positively correlated with $K_{nod}$ and $E_{nod}$ for HbO-based and task-evoked networks. In addition, the activation in left MC was negatively correlated with $C_p$ and $E_{loc}$ for HbR-based and task-evoked networks. Moreover, the activation in right FEF was positively correlated with $\gamma$ for the HbR-based intrinsic networks, whereas the activation in right FEF and DLPFC was positively correlated with $\sigma$ from the same networks.

### 4 Discussion

To the best of our knowledge, this is the first study that systematically explored the differences in frontal HbO/HbR-based activation and networks topological properties between various EF components by testing the same group of participants with fNIRS. In particular, we inspected the relationship between the activation/networks topological properties and the task performance. In addition, the relationship between the brain activation and the networks topological properties was also carefully examined as a synergistic characteristic, which can serve as an index to inspect the difference between various EFs tasks in frontal cortex.

#### 4.1 Differences in Behavioral Performance and Brain Activation/Networks Topological Properties

Consistent with previous studies,\textsuperscript{1,71} the present behavioral results showed that the experimental condition consumed more cognitive effort than the control condition, which was demonstrated by both the low accuracy and the slow response for the experimental condition.

In addition, the negative relationship between HbO and HbR activation was well documented in previous studies,\textsuperscript{59,60} which was also confirmed in the present study, as indicated by the correlation maps in Fig. 3. Moreover, different activation tendencies were identified for the four EF tasks, indicating that the subcomponents of EFs were associated with different neural substrates. Specifically, the attentional inhibition elicited enhanced activation in MC. In contrast, the no-go case elicited comparable activation with the go case in frontal lobe, which confirmed the hypothesis that the response inhibition was a facet of goal-oriented response selection rather than passive inhibition.\textsuperscript{61,72} In particular, the working memory was associated with the activation in left DLPFC, whereas the cognitive flexibility was correlated more with the activation in right FPA.

More important, the network properties provided more comprehensive information to characterize the differences between various EF cognitive processes. In particular, the topological properties of HbO-based networks showed the differences from those from HbR-based networks. It was discovered that the HbR-based networks were highly clustered in the frontal lobe as compared to HbO-based ones. Moreover, both the HbO-based and HbR-based networks exhibited prominent small-world properties whether at rest or during task.

For global properties of HbR-based networks, the task-evoked networks exhibited predominantly decreased network efficiency as compared to the intrinsic networks, except that of the incongruent case in the flanker task, exhibiting increased
global efficiency. However, for HbO-based networks, only the 2-back case in n-back task exhibited decreased local efficiency. The new results indicated that HbR-based network properties were better indexed to characterize the network difference between various EF components. In addition, decreased network efficiency demonstrated that different brain regions worked independently to support the goal-oriented behaviors, which was also confirmed by the highly clustered FC displayed in the matrix of correlation coefficients in Supplementary Material in Fig. S6. Interestingly, unlike other EF tasks, only the incongruent case in flanker task exhibited decreased local efficiency as compared to the control condition. In addition, only the incongruent case exhibited increased global efficiency, which indicated higher routing efficiency. Our findings also suggested that the attentional inhibition was achieved in a more integrative way.

Regarding nodal properties, both the HbO-based and the HbR-based networks exhibited decreased network efficiency during EF tasks, as compared to those at rest. However, in the HbO-based networks, the left MC decreased its importance over the information flow, whereas in the HbR networks, the left FEF exhibited decreased network efficiency for all three nodal properties. Consistent with the global properties results, the nodal property analysis revealed the reorganization of the functional networks during EF tasks, in which HbO-based and HbR-based networks exhibited decreased network efficiency in different brain regions.

Importantly, when the nodal property differences were inspected between the experimental and the control conditions, multiple regions exhibited the significance, which demonstrated that the nodal properties were better indicators than the activation to characterize the neural basis-supporting EF tasks. In particular, HbO-based networks exhibited increased network efficiency including “importance over the information flow (BCnod),” “association capability (k nod),” and “efficiency of the information transfer (Emod)” in the right DLPFC, whereas they exhibited decreased network efficiency in FPA, MC, and FEF during flanker task. The results revealed that the right DLPFC, although not significantly activated, played an important role in the organization of HbO-based networks supporting the attentional inhibition. In addition, in the n-back task, the right FPA played an essential role over the information flow during the 2-back case. Moreover, the left FPA exhibited decreased association capability and information transfer efficiency for the switch case in switch task.

Likewise, nodal differences in multiple channels between the experimental and the control conditions were also identified for HbR-based networks, which were in line with those of the HbO-based networks. In particular, similar to HbO-based networks, HbR-based networks exhibited increased network efficiency in the right DLPFC during flanker tasks. This finding confirmed the important role of the right DLPFC during attentional inhibition. Moreover, the n-back task also exhibited increased importance in the left DLPFC and FEF associated with HbR-based networks, in addition to the right FPA. These results were consistent with the previous studies emphasizing the role of left DLPFC in working memory. In the switch task, decreased nodal degree and efficiency were also observed in the left FPA and right FEF although no significant difference was identified between the two conditions. Importantly, in addition to the activation of DLPFC associated with cognitive flexibility, we also discovered increased network efficiency in the anterior part of left DLPFC and decreased network efficiency in the posterior part.

Importantly, for the go/no-go task, the no-go condition exhibited identical brain activation and network properties with the go case. Our findings provided strong evidence that the response inhibition was a facet of goal-oriented response selection rather than passive cognitive inhibition.

## 4.2 Correlations between Network Properties/Brain Activation and Behavioral Performance

Further, we correlated the activation and network properties with behavioral performances, which were measured by the cognitive cost. Interestingly, significant correlations were revealed between network topological properties and behavioral performances. However, for activation, only the HbR case showed correlations with the behavioral performance. For example, in the flanker task, decreased HbR activation in the left FEF was associated with fast response, whereas decreased HbR activation in the right FPA was related to slow response. In addition, in the n-back and switch tasks, decreased HbR activation in the right DLPFC was associated with faster response. However, after the FDR correction, no significant correlation was identified.

More important, we discovered that the nodal topological properties, rather than global properties, of HbO-based and HbR-based networks exhibited significant correlations with the cognitive cost, which indicated that the nodal properties were more sensitive to the cognitive cost changes. Our finding was in line with a previous study, which also showed no significant correlation between the global topological properties and the behavioral performance.

Interestingly, although the left MC exhibited decreased HbO-based network efficiency during task when compared with that at rest, its importance over the information flow enhanced the performance in the flanker task, its association capability was correlated with faster response in n-back task, and its efficiency of information transfer and importance over the information flow were associated with better performance in switch task. As a result, the left MC played a key role in the function of frontal networks, which promoted the behavioral performance.

In HbR-based networks, the importance of left DLPFC and right MC over the information flow was associated with better performance during flanker task. In addition, in the n-back task, the association capability in right MC and FEF inhibited the behavioral performance, whereas the information transfer efficiency in bilateral DLPFC promoted the performance. In the switch task, the association capability and nodal efficiency in left FPA were associated with poor performance, whereas these properties in the left MC promoted the performance. In addition, the importance of left DLPFC and right MC was also associated with better performance.

## 4.3 Relationships between Brain Activation and Networks Topological Properties

Finally, we generated the correlations between the activation and the networks topological properties, in which multiple correlations were identified. For example, in the flanker, n-back, and switch tasks, the activation was mainly modulated by the intrinsic network properties. These findings are in line with the previous reports that the brain FC during task is modulated primarily by the intrinsic resting-state network architectures, and the resting-state networks can predict the brain activation...
and behavioral performance associated with working memory. In addition, the activation during the go/no-go task was not only shaped by the intrinsic network but also regulated by the task-evoked and HbR-based networks.

In addition, in the flanker task, the overall routing efficiency of networks was associated with increased HbO activation in bilateral DLPFC and left FEF. In addition, the association capability in left MC also enhanced the HbO activation. In the n-back task, the association capability of right FEF showed the positive relationship with the HbO activation; the importance of the right MC showed the positive correlation with the HbR activation, whereas the importance of the left MC exhibited the negative correlation with HbR activation. In the switch task, the local efficiency was associated with decreased HbR activation in the left FPA, whereas the global efficiency was associated with increased HbR activation. In addition, for the task-evoked networks during go/no-go task, the association capability and nodal efficiency in left FEF showed positive correlation with the HbO activation, whereas the local efficiency decreased the HbR activation in left MC. In addition, for the intrinsic network during go/no-go task, the local efficiency enhanced the HbR activation in right FEF, whereas the small-worldness increased the HbR activation in right FEF and DLPFC.

5 Conclusions
A systematical investigation of brain activation and FC patterns during different EF tasks was performed. Our findings demonstrate that the fNIRS is a sufficient technique to characterize the topological organization of brain networks and its link with the activation. In particular, linking the neural activation to the topological organization in frontal lobe can serve as a synergistic index to examine the difference between various EF tasks, which paves a new way toward a comprehensive understanding of neural basis underlying the EFs.

Disclosures
No conflicts of interest financial or otherwise, are declared by the authors.

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