Task-Dependent Individual Differences in Prefrontal Connectivity

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Recent advances in neuroimaging have permitted testing of hypotheses regarding the neural bases of individual differences, but this burgeoning literature has been characterized by inconsistent results. To test the hypothesis that differences in task demands could contribute to between-study variability in brain-behavior relationships, we had participants perform 2 tasks that varied in the extent of cognitive involvement. We examined connectivity between brain regions during a low-demand vigilance task and a higher-demand digit-symbol visual search task using Granger causality analysis (GCA). Our results showed 1) Significant differences in numbers of frontoparietal connections between low- and high-demand tasks 2) that GCA can detect activity changes that correspond with task-demand changes, and 3) faster participants showed more vigilance-related activity than slower participants, but less visual-search activity. These results suggest that relatively low-demand cognitive performance depends on spontaneous bidirectionally fluctuating network activity, whereas high-demand performance depends on a limited, unidirectional network. The nature of brain-behavior relationships may vary depending on the extent of cognitive demand. High-demand network activity may reflect the extent to which individuals require top-down executive guidance of behavior for successful task performance. Low-demand network activity may reflect task- and performance monitoring that minimizes executive requirements for guidance of behavior.

Keywords: connectivity, functional imaging, individual differences, prefrontal cortex, processing speed

Introduction

One aim of cognitive neuroscience has been to identify those aspects of neurophysiology that underlie the consistent individual differences in performance that have long been observed in experimental psychology. Spearman’s (1904) observation that some individuals consistently perform better than others across a broad range of tasks has spawned generations of research investigating the hypothesis that a limited set of resources govern cognitive performance (Spearman 1904; Kahneinan 1973; Norman and Bobrow 1975; Vernon 1983; Baddeley 1986; Just and Carpenter 1992).

One such resource, processing speed, may emerge from individual differences in the efficiency with which cognitive operations can be performed. Cognitive efficiency theories suggest that when these operations can be performed quickly, resource allocation can be minimized and performance maximized. Efficiency theorists have hypothesized that oft-observed correlations between reaction time (RT) and intelligence measures reflect individual differences in “neural efficiency” which, they argue, permit some individuals to overcome cognitive capacity limits more than others (e.g., Jensen 1982, 1998; Vernon 1983). The advent of modern neuroimaging techniques has made it possible to test such hypotheses by permitting more direct observation of brain-behavior relationships than was possible in the past.

Neuroimaging studies in healthy adults support efficiency explanations of individual differences. Results from electroencephalography (EEG) studies have shown differences in amplitude and coherence measures between individuals that correspond to their performance differences (e.g., Gevins and Smith 2000; Grabner et al. 2003; Reiterer et al. 2005). In one study, for instance, Gevins and Smith (2000) required high-ability and low-ability (as measured by WAIS-R performance) participants to perform an n-back working memory (WM) task during EEG recording. The important result was that high-ability participants showed less prefrontal cortex (PFC) and more parietal activity than their low-ability counterparts.

In other EEG studies, reduced “event-related desynchronizations” (ERDs) in alpha frequencies (8-12 Hz), coupled with reduced “event-related synchronizations” (ERSs) in theta frequencies (4-8 Hz), in higher as compared to lower performing individuals have been observed (e.g., Grabner et al. 2003; Babiloni et al. 2009; Del Percio et al. 2009). An ERD is said to occur when the power of some frequency or band of frequencies decreases in response to a stimulus event, whereas an ERS is said to occur when the power of a frequency band increases. ERD in the alpha band, coupled with ERS in the theta band have been interpreted as an index of mental effort (Nunez et al. 2001). Thus, these results suggest reduced mental effort in higher performers compared to lower performers.

Finally, EEG results suggesting reduced neural activity in experts and professional athletes, compared with novices and amateur athletes, suggest support for efficiency explanations of individual differences. In one study, for instance, Babiloni et al. (2009) observed reduced ERDs in the scalp potentials of gymnasts, compared with nongymnasts, while they viewed films of gymnastic performances and judged the artistic and athletic level of the performer. Similar results have been observed when expert performers were compared with nonexperts during actual athletic performance (Del Percio et al. 2009).

Results from positron emission tomography (PET) and functional magnetic resonance imaging (fMRI) studies also show reduced activation in faster than in slower individuals (e.g., Haier et al. 1988, 1992; Larson et al. 1995; Koslyn et al. 1996; Rypma and D’Esposito 1999; Rypma et al. 2002, 2005). In one study for instance, Haier et al. (1992) had 8 participants perform a spatial reasoning task, Raven progressive matrices
that the nature of activation–performance relations may be task
studies reviewed above, different tasks were employed in the
neuroimaging studies may occur for a number of reasons. In the
neural activity.

The between-study variation in brain-behavior relationships
The second observation is that results have been in-
consistent with respect to the direction of brain-behavior
relationships. That is to say, some studies have shown
activation increases with increases in performance, whereas
others have shown activation decreases with increases in
performance. Resolving these discrepancies would have
important implications for, for instance, how we ascribe
cognitive functions to brain regions and whether optimal
performance depends on the amount of activation that
accompanies task performance (e.g., Klimesch 1997, 1999;
Gray et al. 2003) or the speed and efficiency of activation and
communication between brain regions (e.g., Vernon 1983;
Haier et al. 1992; Grabner et al. 2003; Neubauer et al. 2004;
Smith and Jonides 1999; Rypma et al. 2002; Newman et al. 2003;
Maccotta and Buckner 2004; Landau et al. 2007; Bressler et al.
2008), 3 observations can be made about the variation in brain-
behavior relationships. The first observation is that these studies
have been consistent in identifying a frontoparietal network in
which neural activity varies on the basis of individual partici-
pants’ performance, repeated performance of a single task, or
repeated stimulus exposure. These results are important
because they suggest that performance might vary between
individuals on the basis of interactive functions of relatively
distant brain regions whose communication depends on the
integrity of large-scale networks. Bressler (1995, 1996) and
Bressler et al. (2008), for example, have pioneered the concept
of large scale–distributed processing in functionally localized
brain regions. Using local field potentials from up to 15 cortical
sites in 1 hemisphere of functioning adult rhesus monkeys,
Bressler et al. (1993) found task-related multiregional synchroni-
zation over the entire frequency range examined. In subsequent
studies, using local field potential data from extracellular record-
ings, they demonstrated synchrony on physically distant but
functionally related regions during task performance. Thus, it may
be that the efficiency of coordinating such complex systems,
involving the integration of multiple distributed areas differs
between individuals, and is reflected in interactions between
frontal and parietal regions (Bressler 1995; Brovelli et al. 2004,
2008).

Despite the explanatory power of the neural efficiency
hypothesis and suggestive data, neuroimaging findings have not
been replicated consistently across studies. Specifically, some
studies have shown between-subject performance differences
in which greater task-dependent activation was observed in
higher than in lower performing individuals (e.g., Larson et al.
1995; Gray et al. 2003; Newman et al. 2003) and suggest that
neural activity increases with task-related cognitive capacity.
Mixed results in ERD measurements have been observed as
well. For instance, unlike the Grabner et al. (2003) results
reviewed above, Klimesch (1997, 1999) has observed greater
ERD for higher than lower performing participants (see
Jausovec N and Jausovec K 2005, for a review).

Similar discrepant results have been reported in PET and fMRI
studies. Gray et al. (2003), for instance, performed a study
similar to Haier et al.’s (1992) (see above) in which, prior to
fMRI scanning, participants performed the RPM task. During
scanning, participants performed a complex WM task in which
they viewed single letters that appeared sequentially. They were
required to respond each time they observed the appearance of
a letter that had also occurred 3 trials earlier. The difficulty of
the task was varied by the occasional occurrence of “lure” trials
in which a letter presented on a current trial had also appeared
2, 4, or 5 trials previously. Unlike the results of Haier et al.
(1992) described above, they observed, on the lure trials,
greater neural activity within PFC, in participants with higher,
compared with those with lower RPM performance (see also
Brand and Deary 1982; Callicott et al. 2000 see Toffanin et al.
2007, for further review), suggesting that greater visuospatial
WM capacity was associated with greater task-related PFC
neural activity.

Divergent patterns of activation–performance relations across
neuroimaging studies may occur for a number of reasons. In the
studies reviewed above, different tasks were employed in the
different studies. One possibility suggested by the discrepant
results in the Gray et al. (2003) and Haier et al. (1992) studies is
that the nature of activation–performance relations may be task
dependent. It may be that the n-back task used by Gray et al.
(2003) and the tetris task used by Haier et al. (1992) emphasize
different cognitive mechanisms. Other studies have also shown
divergent results (e.g., Tower of London; Newman et al. 2003;
Sternberg-type WM; Rypma et al. 1999; backward digit span;
Larson et al. 1995). Indeed, even subtle variations in task
parameters have been shown to influence activation–perform-
ance relations in both EEG and fMRI studies (Johnson et al.
1997; Rypma, 2006).

In the present study, we sought to examine relationships
between neural activity and performance in this frontoparietal
network, in a single group of participants, using tasks that
varied in the extent of cognitive demand. We employed GCA to
develop a network-based model and to investigate effective
connectivity relationships between brain regions, how they
varied with task demand, and how they varied with individual
differences in participants’ performance.
Previous studies of functional cerebral connectivity have relied on methods that identify sets of brain regions with correlated signal-change patterns (e.g., ICA and PCA; McKeown et al. 1998; Biswal and Ulmer 1999; Allen et al. 2005). These studies have yielded robust delineation of functional connectivity between brain regions by locating discrete temporal structures (e.g., Biswal et al. 1995; Hyde and Biswal 1998; Gusnard et al. 2001; Greicius et al. 2003; Fox et al. 2005). GCA adds important information to that derived from these methods by assessing time-lagged relationships between functionally connected regions, permitting inferences about the directional influences of effective connections. In the present study, we used GCA to gain clues regarding the role of effective connectivity in performance differences between individuals. Brain-behavior relationships have been investigated in the context of fMRI activity (e.g., Rypma and D’Esposito 1999; Rypma et al. 2002, 2006; Grabner et al. 2003; Gray et al. 2003; Beschoner et al. 2008; Perfetti et al. 2009; Rypma and Prabhakaran 2009). How relationships between neural connectivity and behavior vary based on task demands, however, has not yet been systematically investigated.

In this study, a single group of participants performed 2 kinds of tasks during fMRI scanning. The first task was a vigilance task in which participants passively viewed a fixation point and were periodically signaled to press a button. The second task was a digit–symbol substitution task (DSST) in which, on each trial, participants viewed a code table containing digit–symbol pairs, and a single digit–symbol probe that appeared simultaneously below the key (Fig. 1). If the probe pair matched one of those in the table, participants pressed a right-thumb button; otherwise they pressed a left-thumb button. A generation of research on DSST performance indicates that it is maximally sensitive to individual performance differences, that it requires a circumscribed set of cognitive mechanisms (including visual search, pattern matching, and response selection), and that it is minimally sensitive to individual strategy differences (Erber 1976; Grant et al. 1978; Beres and Baron 1981; Wechsler 1981; Joy et al. 2000). Thus, the vigilance task was used to evoke neural activity on the basis of minimal cognitive demand, requiring vigilant attention to stimulus presentation and a simple response. The DSST was used to evoke neural activity that was more cognitively demanding, involving not only attention and simple button-press requirements, but also visual-search and choice-response requirements. Directional influences that were evoked during task performance were assessed using GCA performed on time series data from PFC and parietal regions where neural activity is known to vary between individuals and where DSST-related neural activity has been observed before (Rypma et al. 2006).

Materials and Methods

Participants

Twelve participants (ages 18–27, 7 males and 5 females) were recruited from the Rutgers University and New Jersey Medical School campuses. Participants were excluded if they had any medical, neurological, or psychiatric illness, or if they were taking any type of prescription medication. Participants were screened for depression using the Beck Depression Inventory, which is a 21-item screener for depressive symptoms (BDI; Beck and Steer 1987). Individuals scoring above 14 (i.e., mild depression) were excluded because of the potential for depression to influence brain activity. The study was approved by the University of Medicine and Dentistry of New Jersey and Rutgers University Institutional Review Boards.

Behavioral Tasks

Participants were brought into the behavioral laboratory, signed consent and given a standard battery of tests and questionnaires, and were trained on the computerized DSST by one of the authors (D.A.E.). Participants were then brought to the neuroimaging laboratory. Prior to scanning, they were given brief practice with each of 2 tasks they were to perform during scanning.

Vigilance Task

Each subject performed a task in which he or she stared at a central white fixation cross for 18-s intervals after which the cross changed briefly (500 ms) to a circular checkboard that flickered at 8 Hz for 500 ms, cueing participants to make a bilateral button press. Twenty such events occurred during the 320-s scan (160 images). All scanning parameters were identical to those used for the DSST.

Digit-Symbol Substitution Test

Following performance of the vigilance task, participants performed a task modeled after the DSST from the Wechsler Adult Intelligence Scale (1981). On each fMRI scanning trial, a code table containing digit-symbol pairs and a single digit-symbol probe appeared simultaneously (Fig. 1) for 3.5 s. If the probe pair matched one of those in the table, participants pressed a right-thumb button; otherwise, they pressed a left-thumb button. There were a total of 260 trials in 5 scanning runs (ca. 52 trials per run); the trials for each run were randomly intermixed (jittered) with 23-s rest periods. On half the trials, the probe pair matched one of the digit–symbol pairs in the code table; on the other half, the probe pair did not match one of the pairs in the code table. RT was measured as the time from the onset of the stimulus (i.e., code table and probe-pair presentation) to the time that the subject made a response. Participants were required to respond within the 3.5 s that the stimuli appeared on the screen. To discourage WM-based strategies, the digit–symbol pairings in the code table changed randomly from trial to trial. We used an event-related design that allowed us to examine blood-oxygenation-level-dependent (BOLD) signal changes separately during each trial event.

MRI Technique

Imaging was performed on a 3-T head-only Allegra scanner (Siemens Medical Systems, Ehrangen, Germany) equipped with a fast gradient system for echoplanar imaging. A standard radiofrequency head coil was used with foam padding to comfortably restrict head motion. High-resolution T1-weighted sagittal images were collected. A gradient echo, echoplanar sequence (repetition time [TR] = 2000 ms, echo time [TE] = 30 ms, DSST = 150 vol, Vigilance = 180 vol) was used to acquire data sensitive to the BOLD signal. Resolution was 3.5 × 3.5 mm in-plane and 4 mm between planes (thus 32 axial slices were acquired). Eighteen seconds of gradient and radiofrequency pulses preceded the actual data acquisition to allow tissue to reach steady state magnetization.
Image Analysis

fMRI data were analyzed using AFNI software (Cox 1996). Participant-level task-related effects were identified using conventional linear deconvolution. A regressor was constructed by convolving a hemodynamic response model (a gamma-variate function; Cohen 1997, parameters $b = 8.6, c = 0.547$) with each trial onset in a task-reference function. The $t$-value matrix for each subject was resampled to a 2-mm isovoxel resolution and then spatially normalized to Talairach space (Talairach and Tournoux 1988). Each participant’s 3D structural image (coregistered to the functional data) was transformed, via a 12-parameter affine transformation, to fit it to a Talairach template (i.e., the Colin-brain template), and then the t-value matrix was transformed to Talairach space based on structural image transformation parameters. Regressors for motion correction estimates and linear, quadratic, and cubic trends for each run were included in the baseline regression model. Any subjects with greater than 3 mm of motion were not considered for further analysis. No subjects met this criterion for exclusion (see Supplementary Table 1). For each participant, the preprocessed BOLD data per voxel were then regressed on the resulting model to obtain model scaling parameter estimates (i.e., task-related percent signal-change estimates) and corresponding $t$-values.

To plot mapwise activation at the group level, the data for individual participants were corrected for slice-timing offset, motion corrected, and then spatially filtered with a Gaussian kernel (full-width at half-maximum = 8 mm). For each run, data were then scaled by the mean for that run (i.e., $100 \times y/M$) in each voxel so that the deconvolution parameter estimates would be expressed in terms of percent signal change.

GCA was performed using code written in MATLAB. Data analyses were performed on the time series from 12 regions of interest (ROIs). These ROIs were drawn on individual subjects’ anatomical images to include Brodmann’s areas (BAs) where DSST activation was observed. ROIs were drawn on each subject’s T1 axial slices by one of the authors (D.A.E.) using software from the VoxBo statistical package. These regions included middle and superior frontal gyri, corresponding to BAs 9 and 46, ventral PFC ROIs including inferior frontal gyrus corresponding to BAs 44, 45, and 47, and superior parietal gyrus corresponding to BAs 39 and 40, in each hemisphere, according to the Talairach and Tournoux (1988) and Duvernoy (1999) atlases. For individual participants, time series from all voxels within an ROI were averaged to create a single time series for each ROI.

Each time series was detrended to remove any systematic variation in the data sets that could result from machine system noise leading to linear, cubic, or quadratic drift. To minimize the effect of physiological noise sources such as respiration rate and heart rate (so-called nuisance covariates), a low-pass filter (with a cutoff frequency of 0.10 Hz) was used. Granger causality between 2 regions can be defined as the extent to which the data from 1 region at 1 point in time significantly improves the prediction of another region’s data at a later point in time (Goebel et al. 2003).

Bivariate Granger analyses were performed using F-statistics to test whether lagged data from a time series (variable $y$) improved the prediction of a later value in a time series (variable $x$) to a degree that was statistically significant over that provided by lagged $x$ alone. If not, then $y$ did not influence $x$. The model assumed a model order (i.e., lag length) $p = 5$ TRs, and estimated a residual for the following unrestricted equation by ordinary least squares (OLS):

$$x(t) = c1 + \sum_{i=1}^{p} a(i) x(t-i) + \sum_{i=1}^{p} b(i) y(t-i) + u(t),$$

(1)

where $x(t)$ and $y(t)$ are the 2 time series being evaluated for influence, $t$ is the current time point, $a(i)$ and $b(i)$ are the linear prediction coefficients for $x$ and $y$, $u$ is the residual error of the fit, and $p$ is the lag length.

The residual variance from this full model (eq. 1) was compared with the residual variance from the following reduced autoregressive model:

$$x(t) = c1 + \sum_{i=1}^{p} g(i) x(t-i) + e(t),$$

(2)

where $x(t)$ is the time series being evaluated for influence, $t$ is the current time point, $g(i)$ is the linear prediction coefficient for $x$, $e$ is the residual error in fit, and $p$ is the lag length.

If the F-test comparing the residual variance from the full model to the residual variance from the reduced model

$$F_y \rightarrow x = \left( \frac{\sum_{i=1}^{p} e(i)^2 - \sum_{i=1}^{p} u(i)^2}{p} \right) / \left( \frac{\sum_{i=1}^{p} u(i)^2}{(T-2p-1)} \right)$$

(3)

(where $T$ is the total number of time points, $p$ is the model order) was greater than tabular significance values, then the null hypothesis ($y$ does not influence $x$) was rejected (Greene 2008).

Thus, a time series from each ROI was fit using a full autoregressive model. Briefly, in an autoregressive process the time-series data sets assume that the current time point is functionally related to its $N$ previous time points. For this study a 5th order autoregressive process was used for each of the 12 ROIs.

Five time points (10 s) in the $y$ time series were sequentially assessed for their effect in the prediction of a point immediately preceding them in the $x$ time series as the difference in error terms between the full and reduced models. Thus, for each such assessment, the time point in the $x$ time series served as the dependent variable whereas the time points in the $y$ time series served as independent variables.

Ten-second model orders were adopted to account for delay that may arise due to differences between time series in the hemodynamic response. Although 10 s is orders of magnitude larger than neuronal delays, the fMRI signal represents a convolution of the neuronal signal with the vascular response function. A number of methods have been proposed to estimate the appropriate model order, including Bayesian and Akaike information criterion methods that permit selection between models on the basis of the extent to which variance can be explained with the fewest parameters. Application of these methods to determine a model-order parameter for GCA of fMRI data depends on the assumption that a single model fits the data across all voxels involved in the analysis. FMRI signal, however, is composed of several noise components that arise from respiration, cardiac pulsatility, and machine noise, differentially affect different voxels, and impose influences upon the signal arising from neural activity. Therefore different model orders may be obtained for different voxels, as calculated by information criterion methods. Based on these considerations, we sought to determine an optimal model order based on known properties of the hemodynamic response function.

We determined an optimal autoregressive order based on prior estimates of onset-delay and phase-delay variances of the vascular response function, observed to be around $10-12$ s by a number of different investigators (Lee et al. 1995; Boynton et al. 1996; Saad et al. 2003). Thus, our 5-TR model order was relatively conservative in the context of the hemodynamic filter.

Submodel fits were then carried out for each time series data set compared with the other time series data sets. The significance level for each of them was tabulated for group analysis. For each bivariate model, the $F$ value for each individual subject was calculated, resulting in an interregional matrix consisting of $M(M-1)/2$ values (where $M$ is the number of matrix elements) that were then transformed. An average z-score map was obtained for each task and converted to significance values. We used a false discovery rate procedure to correct for multiple comparisons at a $q$ value of 0.05 (Benjamini and Yekutieli 2001).

This bivariate procedure was performed on all possible ROI pairs such that Granger influences were computed in both directions for all ROI pairs (Brovelli et al. 2004). Unidirectional influences between each ROI pair were calculated for 2 different significance levels. Influences were considered significant for $P < 0.05$, and they were considered trends when $0.05 < P < 0.10$. Influences were considered "unidirectional" if the influence of 1 region of the pair was significant. They were considered "bidirectional" when the influences of both pairs were significant.

Results

Behavioral Performance

One participant’s data were lost due to equipment failure. Behavioral analyses (using the SAS statistical package, SAS Institute, Cary, NC) of the vigilance data indicated uniformly high accuracy with minimal interindividual variability ($M = 98.8\%$,
standard deviation [SD] = 0.001). RTs were fast, also with minimal interindividual variability (M = 507.5 ms, SD = 41.0). Analysis of DSST data also indicated uniformly high with minimal interindividual variability (M = 95.8%, SD = 0.01) that was not significantly different from vigilance task accuracy. DSST RTs were significantly slower than button-press RT (M = 164.5 ms, SD = 238.8), t(10) = 16.45, P < 0.005, and showed greater variability.

**DSST Activation**

Figure 2 shows the average t-values (thresholded at t ≥ 2.00, P < 0.05 uncorrected). The results of this analysis indicated task-related signal-change in the previously identified target regions, including dorsolateral PFC (BA 9 and BA 46), ventrolateral (BA 44, BA 45, and BA 47) PFC, and inferior parietal cortex (BA 39 and BA 40).

**FMRI Connectivity: between-Task Differences**

Analysis of DSST fMRI data (filtered for temporal drift and high-frequency noise to minimize nuisance-covariate effects) using a modified (to account for serially correlated error terms; Worsley and Friston 1995) general linear model indicated activity in several cortical regions including the 12 ROIs across the 2 hemispheres, dorsal PFC (DPPC; Brodmann Area 9; BA 9), a relatively more posterior and inferior regions of PFC (BA 46), ventral PFC (BA 44, BA 45, and BA 47), and parietal cortex (BA 40). Anatomical ROIs were drawn on these regions and GCA methods were used to assess influences between them. GC between 2 regions can be defined as the extent to which the data from one region at one point in time improves the prediction of another region's data at a later point in time (Goebel et al. 2003). GCA was used to evaluate causal influences between ROIs by measuring the extent to which activation changes in one region affected (i.e., reliably preceded) those in other regions at later points in time. Thus, it permitted characterization of the strength and direction of influence between discrete brain regions (Goebel et al. 2003).

Figure 3 shows the GCA results for the vigilance task (A) and for the DSST (B). For the results of both tasks, the results were arranged in a circular fashion with rostral information represented on the left side of each circle, relatively ventral and posterior regions are illustrated in the middle portions so that the caudal-most ROIs are on the right side of the circle. Arrows indicate significant influences; black dashed-line arrows represent influences with P < 0.05; thinner gray arrows represent influences with P < 0.10.

The results illustrated in Figure 3 show 2 differences between the vigilance task and the DSST. First it can be observed that, overall, there were more significant influences between ROIs for the vigilance task than for the DSST. This difference was significant t(10) = 14.96, P < 0.0005. Second, it can be observed that there were more bidirectional influences between ROIs for the vigilance task than for the DSST. This difference is illustrated in Figure 4 which shows mean numbers of unidirectional and bidirectional influences for both task types. It can be seen in Figure 4 that unidirectional influences are equivalent between the 2 tasks but that there are more bidirectional influences for the vigilance task than for the DSST. The interaction of Task-type (vigilance vs. DSST) and Influence type (unidirectional vs. bidirectional) was significant, F(1, 10) = 11.4, P < 0.007. Posthoc t-tests indicated significant differences between tasks for bidirectional but not for unidirectional influences.

**FMRI Connectivity: between-Subject Differences**

To test relationships between the nature of directional influences in each ROI and performance, we made 2 calculations for each subject and each task within each ROI. First, we calculated the number of regions that exerted influences upon each ROI (i.e., the number of “input influences”) and second, we calculated the number of regions that each ROI had influences upon (i.e., the number of “output influences”; Brovelli et al. 2004). To examine how brain-behavior relationships during DSST activity differ from vigilance activity, we first calculated, for each ROI in each subject, differences in the numbers of influences for the vigilance task and the DSST. These calculations were performed separately for input and output influences. It is worth noting that, in correlation-based analyses, these types of connections are considered to be identical. GCA however permits separate assessment of both input influences from other regions and output influences to other regions for each ROI. Miniscule intersubject variability in vigilance RT performance obviated meaningful analysis with these behavioral data. Thus, we performed a series of linear regressions, with Bonferroni correction (Holm 1979), on individual participants’ DSST RT, and differences between tasks in the numbers of input and output influences. This approach is similar to that performed by Brovelli et al. (2004).
There were no input influences to any ROIs that were significantly different between the 2 tasks after Bonferroni correction. The linear regression of DSST RT and task differences in numbers of output influences from BA 9 to inferior PFC and parietal regions in the right hemisphere was significant (slope = -0.78; \( r^2 = 0.49; \) \( t = -3.3; P < 0.01 \)) in the vigilance task, but positively correlated with their DSST RT (slope = 0.62; \( r^2 = 0.31; t= 2.4; P < 0.04 \); Fig. 5) in the DSST.

**Discussion**

In this study, we tested the hypothesis that the nature of activation–performance relations varies with the extent of cognitive involvement required by the task. We compared effective connectivity differences between brain regions where neural activity was elicited in 2 simple tasks that varied in the extent of cognitive demand. We used an analysis method that permitted unambiguous assignment of the direction of connectivity, GCA. Participants were faster when they were cued periodically to press a button than when they were required to determine the presence or absence of a probe digit-symbol pair among a string of such pairs. There was greater effective connectivity in the vigilance task compared with the DSST. Analyses of directional connectivity indicated that there were significantly more bidirectional influences in the vigilance task than in the DSST. Unidirectional influences were, however, equivalent between the 2 tasks. Finally, analysis of individual differences in Granger causal influences indicated that individuals with faster DSST RTs showed reduced dorsal PFC influence extending to ventral PFC and posterior parietal regions than slower individuals during DSST performance. These same faster individuals, however, showed increased dorsal PFC influence compared with slower individuals during vigilance task performance.

**Vigilance-Related Activity**

The present results clearly suggest an association between activation–performance relations and task demand. When task demand was low, requiring participants only to maintain vigilance for a signal to press a button, processing-speed ability
Astrocytes could mediate arteriole dilation. Ca²⁺ was relatively high, requiring participants to not only maintain low-frequency fluctuations (Biswal and Hudetz 1996). A shift in the balance between cell signaling mechanisms could mediate a modal shift from vigilant rest to rest-related activity. Ca²⁺ opposing cellular signaling mechanisms may govern vigilant vigilance has important implications for speculation regarding the mechanisms that could give rise to interregional connectivity when cognitive mechanisms are relatively minimally engaged. To support such activity, brief neural pulses may fluctuate in synchrony between functionally related regions. Such spontaneous fluctuation has been observed in low-frequency ranges during other tasks that minimally involve cognition (e.g., Fransson 2006). Increased bidirectional connectivity might suggest an increased equilibrium in signal transmission between brain regions during vigilance compared with DSST performance. Increased bidirectional fluctuation during sustained vigilance has important implications for speculation regarding the mechanisms that could give rise to interregional connectivity. This increased vigilance-related connectivity mainly resulted from increases in bidirectional connectivity between brain regions. Such activity may reflect in part vigilance operations of participants awaiting the response signal. Passive vigilance has been known to elicit increased activity in previous studies, compared with that elicited during task performance (e.g., Fransson 2006). Increased bidirectional connectivity might suggest an increased equilibrium in signal transmission between brain regions during vigilance compared with DSST performance. Increased bidirectional fluctuation during sustained vigilance has important implications for speculation regarding the mechanisms that could give rise to interregional connectivity when cognitive mechanisms are relatively minimally engaged. To support such activity, brief neural pulses may fluctuate in synchrony between functionally related regions. Such spontaneous fluctuation has been observed in low-frequency ranges during other tasks that minimally involve cognition (e.g., Cordes et al. 2001; Goldman et al. 2002).

A number of mechanisms have been hypothesized that could give rise to such phenomena. For instance, Zonta et al. (2003) have suggested that glutamate-mediated Ca²⁺ fluctuation in astrocytes could mediate arteriole dilation. Ca²⁺ transient pulses in astrocyte end feet (part of the astrocyte that makes contact with the arteriole) also cause cerebrovascular contraction (Mulligan and McVicar 2004). Thus, the balance between these opposing cellular signaling mechanisms may govern vigilant rest-related activity. Ca²⁺ channel inhibition led to disruption of this signaling. A shift in the balance between cell signaling mechanisms could mediate a modal shift from vigilant rest to active task performance. Indeed, other similar mechanisms, including nitric oxide synthase have also been shown to disrupt low-frequency fluctuations (Biswal and Hudetz 1996).

**DSST-Related Activity**

In contrast to vigilance-related connectivity, when task demand was relatively high, requiring participants to not only maintain vigilance but also to search an array for the presence of a digit-symbol target, connectivity was 1) reduced relative to the lower demand vigilance task and 2) dominated by more unidirectional than bidirectional activity, as measured by GCA. The relative reduction in bidirectional connections during DSST suggests that task-related activity reflects disengagement of equilibrium mechanisms that dominate low-demand task activity and engagement of mechanisms that involve more directed activity between brain regions. This directed activity may reflect the goal-oriented executive control that dorsal PFC regions exert upon more ventral and more posterior brain regions that are involved in the execution of the visual search, target-detection, and response-selection processes required by the DSST. The relationship we observed between connections that extended from dorsal PFC to other brain regions replicates earlier results from our laboratory (Rypma et al. 2006; Motes MA, Rypma B, unpublished data) and suggests support for the hypothesis that individuals vary in the extent to which cognitive processes can be implemented automatically. It may be that some individuals implement the cognitive operations necessary for successful task performance (mediated by ventral PFC and parietal brain regions) automatically, with minimal reliance on PFC regions involved in executive operations (e.g., dorsolateral PFC; Jung and Haier 2007; Prabhakaran and Rypma 2007). Other individuals may implement these operations in a more controlled fashion. In these individuals, PFC mediation may serve to guide more ventral and posterior brain regions in the service of successful task performance.

**Task-Dependent Variation in Brain-Behavior Relationships**

Relationships between neural activity and behavioral state have been observed in studies comparing waking, sleeping, and anesthesia in animals and humans. When humans are minimally engaged in cognitive activity (i.e., during “resting state”), neuro-imaging signal exhibits higher amplitude and interregional correlation depending upon whether individuals are anesthetized or not (Kiviniemi et al. 2005), whether they are asleep or awake (Fukunaga et al. 2006; Horovitz et al. 2008), or whether they have their eyes open or closed (e.g., Yang et al. 2007; Bianciardi et al. 2009). Although the origin of these state-related signal changes are not yet completely understood (e.g., Birn et al. 2006), these findings have important implications for a complete theory of functional neural circuitry at rest, during task-related activity, and the interaction of resting and active functional circuitry. For instance, some studies indicate that distinct networks of activity during spontaneous and evoked activity interact such that increases in spontaneous network activity result in reduced activity evoked by sensory stimulation (Sachdev et al. 2004; Hasenstaub et al. 2007). Such results suggest a dynamic balancing mechanism in which resting neural activity levels mediate the responsiveness of networks to stimulation. The differences observed in DSST-related Granger connectivity, compared with vigilance-related connectivity, suggest that such balancing mechanisms play an important role in determining levels of activation observed during task performance.

**Participant-Dependent Variation in Functional Connectivity**

In the present results, task-dependent connectivity changes differed between individuals. Specifically, within dorsal PFC, faster participants (as measured by DSST) showed more...
vigilance-related connectivity than slower participants, but faster participants showed less DSST-related connectivity than slower participants. The results suggest that the balancing mechanisms that mediate differences in low- versus high-demand PFC connectivity are intimately related to individual differences in cognitive efficiency (as measured by DSST).

The finding of participant-dependent connectivity differences has implications for both cognitive and neural explanations of individual differences. At a cognitive level, one possibility is that dorsal PFC regions subserve the simultaneous monitoring of task-demand information and performance accuracy in the service of task learning. This region has been implicated in performance monitoring (e.g., Sharp et al. 2004). Such task- and performance-monitoring may aid in the development of more efficient, or “automatic” task-performance. Those individuals without such extra-task processing capability may rely on less efficient “controlled” processing (cf. Schneider and Shiffrin 1977), mediated by right-hemisphere regions of dorsal PFC. Consistent with our findings, frontal activity declines have been observed with skill improvements that reflect development of task automaticity (Petersen et al. 1998).

At a neural level, participant-dependent connectivity changes may reflect individual differences in the integrity of large-scale networks composed of computational nodes comprised of physically distant but functionally related brain regions that must coordinate and integrate functions (e.g., Bressler and Kelso 2001). Findings of activation synchrony across relatively distal brain regions support this notion. It may be that white-matter integrity affects neural transmission efficiency between these brain regions which in turn, affects performance (e.g., Jensen 1982; Vernon 1983; Rypma and D'Esposito 1999; Grabner et al. 2003; Rypma and Prabhakaran 2009). Precise explication of the mechanisms that govern differences in white-matter integrity has only begun to emerge as measures that distinguish these mechanisms in pathological and aging populations have been developed (e.g., Song et al. 2003; Nair et al. 2005; Bennett et al. 2009). Studies utilizing these measurement methods indicate that such mechanisms could take several forms including changes in axon number, size, and myelination extent. More research is certainly required before any precise inferences could be made about the nature of the white-matter differences that distinguish between relatively good and poor performers on cognitive tasks.

**Granger Causality and Neural Connectivity**

Other multivariate statistical methods including ICA and PCA have been used to decompose fMRI data into independent components on the basis of distinct sets of linear parameters. The “images,” or time-series data sets, derived from these analyses represent functional connectivity maps that have been important for understanding connectivity relations between regions (e.g., McKeown et al. 1998; Biswal and Ulmer 1999).

GCA improves upon these methods because Granger regression explicitly accounts for interregional temporal variability that has been demonstrated in previous reports (Lee et al. 1995; Buckner et al. 1996; Miezin et al. 2000), whereas ICA and PCA assume that the exact sequence of information flow cannot be obtained from the data. Thus, Granger correlation adds to information obtained from ICA and PCA about functionally connected brain regions by explicitly accounting for interregional temporal variability.
Joy S, Fein D, Kaplan E, Freedman M. 2000. Speed and memory in WAIS-R-NL digit symbol performance: among healthy older adults. J Int Neuropsychol Soc. 6:770-780.

Jung RE, Haier RJ. 2007. The parieto-frontal integration theory (P-FIT) of intelligence: converging neuroimaging evidence. Behav Brain Sci. 30:135-189.

Just MA, Carpenter PA. 1992. A capacity theory of comprehension: individual differences in working memory. Psychol Rev. 99:122.

Kalchman D. 1973. Attention and effort. Englewood Cliffs (NJ): Prentice-Hall.

Kayser AS, Sun FT, D’Esposito M. 2009. A comparison of Granger causality and coherency in fMRI-based analysis of the motor system. Hum Brain Mapp. 30:3475-3494.

Klimesch W. 1997. EEG alpha rhythms and memory processes. Int J Psychophysiol. 26:319-340.

Klimesch W. 1999. EEG alpha and theta oscillations reflect cognitive and memory performance: a review and analysis. Brain Res Brain Res Rev. 29:169-195.

Landau SM, Garavan H, Schumacher EH, D’Esposito M. 2007. Regional specificity and practice: dynamic changes in object and spatial working memory. Brain Res. 1180:78-89.

Larson GE, Haier RJ, LaCasse L, Hazen K. 1995. Evaluation of a ‘mental effort’ hypothesis for correlations between cortical metabolism and intelligence. Intelligence. 21:267-278.

Lee AT, Glover GH, Meyer CH. 1995. Discrimination of large venous vessels in time-course spiral blood-oxygen-level-dependent magnetic resonance imaging. Mag Res Med. 33:745-754.

Maccotta L, Buckner RL. 2004. Evidence for neural effects of repetition that directly correlate with behavioral priming. J Cog Neurosci. 16:1625-1632.

Mckown MJ, Jung TP, Makeig S, Brown G, Kindermann SS, Lee TW, Sejnowski TJ. 1998. Spatially independent activity patterns in functional MRI data during the stroop color-naming task. Proc Natl Acad Sci USA. 95:803-810.

Miczin FM, Maccotta L, Ollinger JM, Petersen SE, Buckner RL. 2000. Characterizing the hemodynamic response: effects of presentation rate, sampling procedure, and the possibility of ordering brain activity based on relative timing. Neuroimage. 11:735-759.

Mulligan SJ, McVicar BA. 2004. Calcium transients in astrocyte endfeet cause cerebrovascular constrictions. Nature. 431:195-199.

Nair G, Tanahashi Y, Low HP, Billings-Gagliardi S, Schwartz WJ, Duong TQ. 2005. Myelination and long diffusion times alter diffusion-tensor-imaging contrast in myelin-deficient shiverer mice. Neuroimage. 28:165-174.

Neubauer AC, Preuss MT, Bressler SM, Seifert V, Gratton G. 2004. Decreased EEG coherence between prefrontal electrodes: a correlate of high language proficiency? Exp Brain Res. 163:109-113.

Newman SD, Carpenter PA, Varma S, Just M. 2003. Frontal and parietal participation in problem solving in the Tower of London. Acta Psychol. 116:55-74.

Poldrack RA, Desmond JE, Glover GH, Gabrieli JDE. 1998. The neural basis of visual skill learning: an fMRI study of mirror reading. Cereb Cortex. 8:1-10.

Prabhakaran V, Rypma B. 2007. P-fit and the neuroscience of intelligence: how well does P fit? Behav Brain Sci. 30:166.

Reiterer S, Berger ML, Hemmelmann C, Rappelsberger P. 2005. Decreased EEG coherence between prefrontal electrodes: a correlate of high language proficiency? Exp Brain Res. 163:109-113.

Rypma B. 2006. Factors controlling neural activity during delayed-response task performance: testing a memory organization hypothesis of prefrontal function. Neuroscience. 139:223-235.

Rypma B, Berger JS, D’Esposito M. 2002. The influence of working memory demand and subject performance on prefrontal cortical activity. J Cog Neurosci. 14:721-731.

Rypma B, Berger JS, Genova HM, Rebbechi D, D’Esposito M. 2005. Dissociating age-related changes in cognitive strategy and neural efficiency using event-related fMRI. Cortex. 41:582-594.

Rypma B, Berger JS, Prabhakaran V, Bly BM, Kimberg DY, Biswal BH, D’Esposito M. 2006. Neural correlates of cognitive efficiency. Neuroimage. 33:145-156.

Rypma B, D’Esposito M. 1999. The roles of prefrontal brain regions in components of working memory: effects of memory load and individual differences. Proc Nat Acad Sci. 96:6558-6563.

Rypma B, D’Esposito M. 2000. Isolating the neural mechanisms of age-related changes in human working memory. Nat Neurosci. 3:509-515.

Rypma B, Prabhakaran V. 2009. When less is more and when more is more: the mediating roles of capacity and speed in brain-behavior efficiency. Intelligence. 37:207-222.

Rypma B, Prabhakaran V, Desmond JE, Glover GH, Gabrieli JDE. 1999. Load-dependent roles of frontal brain regions in the maintenance of working memory. Neuroimage. 37:216-226.

Saa'd Z, DeYoe EA, Ropella KM. 2003. Estimation of fMRI response delays. Neuroimage. 18:949-954.

Sachdev RN, Ebner FF, Wilson CJ. 2004. Effect of subthreshold up and down states on the whisker-evoked response in somatosensory cortex. J Neurophys. 92:3511-3521.

Schneider W, Shiffrin R. 1977. Controlled and automatic human information processing I: Detection, search, and attention. Psychol Rev. 84:1-66.

Sharp DJ, Scott SK, Wise RJ. 2004. Monitoring and the controlled processing of meaning: distinct prefrontal systems. Cereb Cortex. 14:1-10.

Smith EE, Jonides J. 1999. Storage and executive processes in the frontal lobes. Science. 283:1657.

Song SK, Sun SW, Ju WK, Lin SJ, Cross AH, Neufeld AH. 2003. Diffusion tensor imaging detects and differentiates axon and myelin degeneration in mouse optic nerve after retinal ischemia. Neuroimage. 20:1714-1722.

Spearman C. 1904. General intelligence, objectively determined and measured. Am J Psychol. 15:201.

Talairach J, Tournoux P. 1988. Co-planar stereotactic atlas of the human brain. New York: Thieme Medical Publishers.

Tolftan J, Johnson A, de Jong R, Martens S. 2007. Rethinking neural causality and coherency in fMRI-based analysis of the motor system. Neuroimage. 36:144-152.

Toffanin P, Johnson A, de Jong R, Martens S. 2007. Rethinking neural causality and coherency in fMRI-based analysis of the motor system. Neuroimage. 36:144-152.

Wechsler D. 1981. Manual for the Wechsler Adult Intelligence Scale—revised. New York: The Psychological Corporation.

Worsley KJ, Friston KJ. 1995. Analysis of fMRI time-series revisited—again. Neuroimage. 36:144-152.

Young H, Long XY, Yang Y, Yan H, Zhu CZ, Zhou XP, Zang YF, Gong QY. 2007. Amplitude of low frequency fluctuation within visual areas revealed by resting-state functional MRI. Neuroimage. 36:144-152.

Zonta M, Angulo MC, Gobbo S, Rosengarten B, Hossmann KA, Pozzan T. 2003. Neuron-to-astrocyte signaling is central to the dynamic control of brain microcirculation. Nat Neurosci. 6:43-50.