Integrative Frontal-Parietal Dynamics Supporting Cognitive Control

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Competing Interests
None declared.
Abstract
Coordinating among the demands of the external environment and internal plans requires cognitive control supported by a fronto-parietal control network (FPCN). Evidence suggests that multiple control systems span the FPCN whose operations are poorly understood. Previously (Nee and D’Esposito, 2016; 2017), we detailed frontal dynamics that support control processing, but left open their role in broader cortical function. Here, I show that the FPCN consists of an external/present-oriented to internal/future-oriented cortical gradient extending outwardly from sensory-motor cortices. Areas at the ends of this gradient act in a segregative manner, exciting areas at the same level, but suppressing areas at different levels. By contrast, areas in the middle of the gradient excite areas at all levels, promoting integration of control processing. Individual differences in integrative dynamics predict higher-level cognitive ability and amenability to neuromodulation. These data suggest that an intermediary zone within the FPCN underlies integrative processing that supports cognitive control.

Impact Statement
This study shows that intermediary areas within the FPCN are critical for integrating control processing, cognitive ability, and amenability to neuromodulation.
Introduction

While habits rigidly link stimuli to actions, cognitive control enables flexible behavior that can adapt to present conditions, prevailing contexts, and future plans1–4. The human capacity for flexible behavior is thought to arise from the expansion of transmodal cortices that are synaptically distant from primary cortices5–7 such that synaptic distance may untether transmodal areas from the processing hierarchies that link stimulus to action in canonical circuits8. Such transmodal areas are prominent in the prefrontal (PFC) and posterior parietal cortices (PPC).

The relationship between cognitive control and areas of the PFC and PPC is so ubiquitous that a co-active set of PFC-PPC areas is frequently termed the “frontoparietal control network” (FPCN)9–12. Recognizing their involvement across a diverse array of tasks, a similar constellation of areas is also referred to as the “multiple demand network”13–15. This network is thought to implement cognitive control by flexibly coordinating activity among diverse brain systems to integrate brain-wide processing in a goal-directed manner 16,17. The integrative capacity of these areas enables them to reconfigure the brain into difficult-to-reach states18, thereby conferring the flexibility needed to act in an adaptive, rather than habitual manner. In this way, the integrative capacity of cognitive control is central to higher-level cognition.

Mounting evidence suggests that there is not a single FPCN, but multiple networks10–12. These networks are situated upon global brain gradients such that increasingly sensory-motor distal areas of the PFC and PPC are increasingly distant from areas of the brain involved in external processing6,7. This has led to the proposal that the more sensory-motor distal aspects of the FPCN are involved in more internally-oriented control processes12. The proposal that gradients in the PFC and PPC can be classed along an external-internal axis19,20 offers a unifying perspective of the different forms of cognitive control which has remained elusive4.

Ultimately, cognitive control is grounded in behavior. This means that internally-oriented aspects of control, such as those that plan for the future, must be integrated with externally-oriented aspects of control, such as those that select appropriate sensory features for processing and action. How and where such integration takes place is an open question. One possibility is that the control gradient that expands outwardly from sensory-motor cortices doubles as an integration gradient such that the most sensory-motor distal areas are the most integrative. This possibility is consistent with theories of PFC function that posit that the rostral-most (i.e. most sensory-motor distal) areas act as apex controllers that exert the widespread influences that can coordinate brain-wide activity under a single goal21. Another possibility is that a cascade of control signals progresses from sensory-motor distal to sensory-motor proximal areas with integration of those signals progressing along the way22,23. A third possibility is that those areas situated between externally-oriented and internally-oriented control are responsible for their integration. This last possibility is consistent with recent data indicating the importance of mid-lateral PFC in integrative control4,24–26. This latter possibility would suggest a nested structure to gradients in the brain: just as the FPCN is situated in intermediary zones of the brain to flexibly guide sensation to action, so too are intermediary zones of the FPCN essential for flexibly guiding control itself.

Here, two datasets that have demonstrated gradients of cognitive control in the PFC25,26 are examined to investigate dynamics in the broader FPCN. The same gradients of activation observed in the PFC are mirrored in the PPC and tied to specific control functions and behavior. Interactions among areas of the PFC and PPC are measured by examining both static and dynamic indices of effective connectivity, revealing how control integrates processing to support adaptive behavior. Finally, static and dynamic
Figure 1. Comprehensive Control Task. On each trial, participants observed a letter at a spatial location and made a keypress in response. Keypresses mapped onto “yes”/”no” responses. Participants either responded “no” to the stimulus without regard to the stimulus features, or made a choice response based on a pre-learned sequence to a color cued feature (T-A-B-L-E-T for letters; star trace for locations) thereby engaging sensory-motor control. The correct response was either based upon a reference stimulus for which participants responded regarding whether the present stimulus followed a reference stimulus (sequence-back), or whether the stimulus was the start of the sequence (sequence-start). Switching among these tasks engages contextual control. Finally, the reference stimulus could either be the last item presented, or a distant item. In the latter case, the reference item had to be sustained over several trials requiring temporal control to prepare for the future. Colored frames indicated relevant stimulus features (letter or location) while frame shapes indicated cognitive control demands which were manipulated in the middle of each block (sub-task). Stimulus domain and cognitive control demands were independently manipulated in a factorial design wherein orthogonal contrasts separately isolated sensory-motor control, contextual control, and temporal control.
measures of integration are used to predict trait-level cognitive ability, and susceptibility of cognitive control to neuromodulation showing the importance of integration for higher-level cognition and interventions more broadly. Collectively, these analyses indicate the integrative organization of the FPCN whose insights may be useful to understand other transmodal networks.

**Results**

Two independent samples (n=24, n=25) completed a Comprehensive Control Task that independently manipulated demands on stimulus domain (verbal vs spatial processing), sensory-motor control (associating stimuli to actions), contextual control (context-dependence of appropriate stimulus-response associations), and temporal control (planning for the future). These processes can be classed along an external-internal continuum such that sensory-motor control acts upon the sensory environment, contextual control informs those actions based upon an internalized rule, and temporal control informs the rule based upon an additional internalized representation (Figure 1). These processes can be classed by timescale such that sensory-motor control specifies an action for the present stimulus, contextual control specifies a prevailing task context, and temporal control sustains a prospective memory for the future. More generally, the processes are classed by abstraction with sensory-motor control being the most concrete and temporal control being the most abstract. Behavioral data are reported in Supplemental Material.

**Mirrored Control Gradients in the PFC and PPC**

Figure 2A depicts the voxel-wise, whole-brain activations for each cognitive control contrast. In the lateral PFC, activations progressed in a caudal to rostral fashion as a function of abstraction of cognitive control from sensory-motor control to contextual control, to temporal control\(^{25,26}\). In the PPC, activations progressed in a rostral to caudo-lateral fashion as a function of abstraction of cognitive control\(^{27}\). That is, in both the lateral PFC and PPC, increasingly abstract control was associated with activations increasingly distant from sensory-motor cortices of the pre/post-central gyri.

Rather than activating discrete patches of cortex, different cognitive control demands produced overlapping activations eliciting gradients along the lateral surface (Figure 2B). Although the extent of overlap depends upon preprocessing (e.g. smoothing, volumetric vs surface processing), similar gradients of activation were observed using minimal, surface-based smoothing (Supplemental Figure 1). The mirrored gradients observed in the lateral PFC and PPC are suggestive of a mirrored functional organization extending outwardly from sensory-motor areas consistent with macroscale gradients of cortical function\(^5\)–\(^7\).

To better characterize the functions of these areas, spherical regions-of-interest (ROIs) were centered on activation peaks of the contrasts (Figure 2B; Table 1). Activations across the eight conditions of the factorial design are depicted in Figure 3A providing a graphical profile of the activation patterns in each area. PFC-PPC areas closer to sensory-motor cortex (blue, green) tended to be sensitive to multiple demands activating for both sensory-motor and contextual control. These areas also had a tendency to activate preferentially to one stimulus domain or the other. By contrast, areas distal from sensory-motor cortices (red) tended to be more specialized towards temporal control and did not show a preference for stimulus domain.
Figure 2. A) Activations for temporal control (red), contextual control (green), and sensory-motor control (blue). Activations are depicted separately for each sample, as well as the samples combined. B) Activations overlayed to demonstrate the gradient of cognitive control. Spheres indicate the location of regions-of-interest based upon activation peaks. FPl – lateral frontal pole; MFG – middle frontal gyrus; VLPFC – ventrolateral prefrontal cortex; cMFG – caudal middle frontal gyrus; IFJ – inferior frontal junction; SFS – superior frontal sulcus; aIPS – anterior intra-parietal sulcus; IPL – inferior parietal lobule; mIPS – mid intra-parietal sulcus; SPL – superior parietal lobule.
Figure 3. Activation profiles. Activations across the eight conditions of the task design are depicted as radar plots (SB – spatial baseline; VB – verbal baseline; SS – spatial switching; VS – verbal switching; SP – spatial planning; VP – verbal planning; SD – spatial dual; VD – verbal dual). The top panel depicts idealized profiles for areas sensitive solely to temporal control (red), contextual control (green), sensory-motor control (blue), verbal stimulus domain (purple), and spatial stimulus domain (orange). Inset: results of multi-dimensional scaling of the activation profiles across regions. Colored by abstraction refers to coloring as a function of position along the control gradient (blue – sensory-motor proximal; green – intermediary; red – sensory-motor distal). Colored by stim domain refers to coloring as a function of sensitivity to stimulus domains (orange – spatial; purple – verbal; gray – neither).
To provide a more compact, data-driven description of these profiles, the data were subjected to multi-dimensional scaling (MDS). Two dimensions accounted for 89% of the variance of the data (Figure 3B). The first dimension recapitulated the abstraction gradient, placing PFC-PPC areas proximal to sensory-motor cortices on one end of the dimension (e.g. superior frontal sulcus - SFS; anterior intra-parietal sulcus - aIPS), and areas distal to sensory-motor cortices on the other (e.g. middle frontal gyrus - MFG; inferior parietal lobule - IPL). Hence, across the different conditions of the task, this dimension collapsed abstraction of cognitive control into a single axis. The second dimension reflected sensitivity to stimulus domain with areas preferentially engaged by spatial processing at one end and areas preferentially engaged by verbal processing at the other. These data are consistent with the idea that the cortex is organized along two principle gradients reflecting abstraction and modality5–7. Collectively, these dimensions provided a data driven way to operationalize the factors of the task design.

| Area      | X   | Y   | Z   | Sample 2 X | Y   | Z   |
|-----------|-----|-----|-----|-----------|-----|-----|
| FPl       | -44 | 48  | 4   | -36       | 52  | 0   |
| MFG       | -38 | 28  | 44  | -36       | 34  | 38  |
| VLPFC     | -52 | 20  | 28  | -42       | 30  | 20  |
| cMFG      | -34 | 10  | 60  | -30       | 6   | 56  |
| IFJ       | -38 | 6   | 26  | -42       | 10  | 24  |
| SFS       | -22 | 0   | 54  | -20       | 0   | 56  |
| IPL       | -54 | 50  | 44  | -56       | -52 | 42  |
| mIPS      | -28 | 60  | 42  | -26       | -56 | 44  |
| aIPS      | -34 | 40  | 46  | -30       | -42 | 42  |
| SPL       | -14 | 52  | 64  | -12       | -60 | 58  |

Table 1. Coordinates of regions-of-interest reported in MNI space. FPl – lateral frontal pole; MFG – middle frontal gyrus; VLPFC – ventrolateral prefrontal cortex; cMFG – caudal middle frontal gyrus; IFJ – inferior frontal junction; SFS – superior frontal sulcus; IPL – inferior parietal lobule; mIPS – mid intra-parietal sulcus; aIPS – anterior intra-parietal sulcus; SPL – superior parietal lobule

**Control Gradient is related to Present-Future Behavior**

Next, areas were characterized as a function of their relationship to behavior in the task. Activations were assessed during the sub-task phases that manipulated cognitive control demands (Figure 1). Cognitive control behaviors were expressed both during the sub-task phases, as well as during return trials that immediately followed the sub-task phase. For example, temporal control requires sustaining an internal representation (i.e. a reference stimulus) during the sub-task phase to be utilized on the return trial (i.e. does the stimulus on the return trial follow the reference stimulus in the sequence?). Therefore, activations during temporal control that sustain the internal representation would be expected to relate to behavior on the return trial, but not necessarily the sub-task trials. Hence, activations can be separately correlated with behavior measured at the same time as the activations (sub-task trials: present behavior) or the behavior for which the activations are preparing (return trials: future behavior).

In the PFC, increasing rostral areas were increasingly associated with future behavior and decreasingly associated with present behavior25,26. This pattern was mirrored in the PPC (Figure 4A) such that sensory-motor proximal areas (aIPS, SPL) were positively associated with present, but not future behavior, while
Figure 4. Brain-behavior relationships. A) Top: Repeated measures correlations between activation and present behavior (i.e. behavior during sub-task trials; see Figure 1). Areas related to sensory-motor control (blue) and contextual control (green) showed positive associations, while areas related to temporal control (red) showed no or negative associations. Bottom: Repeated measures correlations between activation and future behavior (i.e. behavior during return trials; see Figure 1). Areas related to temporal control (red) and contextual control (green) tended to show positive associations, while areas related to sensory-motor control (blue) tended to show no associations. ** indicates Bonferroni-corrected p < 0.05. * indicates uncorrected p < 0.05. B) Voxel-wise partial correlations between activation and present behavior (blue), future behavior (red), and both (green). Results are visualized at p < 0.001 with 124 voxel cluster extent. C) Linear mixed effects modeling of the activation-present behavior slope (top) and activation-future behavior (slope) bottom using the first dimension of multi-dimensional scaling (MDS) depicted in Figure 3. * indicates p < 0.05; ** indicates p < 0.005.
sensory-motor distal areas (IPL) were positively associated with future, but not present behavior. Areas in-between (mIPS) were associated with both present and future behavior. These patterns were confirmed with voxel-wise analyses (Figure 4B). These data are consistent with an integrative role of contextual control areas positioned between internal, future-oriented temporal control areas and external, present-oriented sensory-motor control areas.

To better quantify these relationships, the abstraction dimension uncovered by MDS was used to account for areal relationships with behavior. Separate linear mixed effects models were fit for the activation-present behavior and activation-future behavior relationships across areas. Consistent with the impressions above, the more abstract the PFC-PPC area, the less it related to current behavior (sample 1: t(238) = -9.18, p = 2.19e-17; sample 2: t(248) = -8.73, p = 3.81e-16) and more it related to future behavior (sample 1: t(238) = 2.72, p = 0.007; sample 2: t(248) = 4.39, p = 0.00002). Hence, these analyses indicate that progressively abstract areas in both the PFC and PPC are increasingly future-oriented.

Establishing Source-Target Relationships

Next, interactions among PFC-PPC areas were examined. Control is embodied by source-target relationships such that controllers effect processing in controlled targets. Effective connectivity, which estimates directed influences, offers the most straightforward means to assess such source-target relationships. There are both static (stationary) and dynamic (non-stationary) interactions among the PFC and PPC\(^{16,28-30}\). To examine static interactions and their directed nature, a biophysically plausible generative model of how neuronal interactions produce cross spectra in the fMRI signal was employed\(^{31-33}\).

The method was validated in two ways. First, the method was applied to the task data of both samples and estimates of directed interactions were compared across the samples. These estimates had excellent correspondence across the samples (r = 0.96; Figure 5, top). Second, estimates of directed interactions were used to examine the putative pre-eminent role of the PFC in cognitive control\(^1\). Cognitive control is exemplified by source-target relationships such that drivers of control asymmetrically influence targets. To characterize such asymmetries on the lobular level, the magnitude of effective connectivity (i.e. deviations from zero) were separately combined for each of the 2 x 2 combinations of source-target lobe (i.e. PFC->PFC, PFC->PPC, PPC->PFC, PPC->PPC). In both samples, the magnitude of effective connectivity arising from the PFC was significantly stronger than that arising from the PPC (sample 1: F(1,23) = 7.1, p = 0.0138; sample 2: F(1,24) = 6.26, p = 0.0195; Figure 5, bottom). Such data are consistent with the idea that the PFC is the primary driver of PFC-PPC dynamics. Collectively, these data indicate that the method produces replicable and theoretically sensible estimates.

An Intermediary Integration Zone of Control

To examine integration of control signals, source-target relationships were assessed at the network level. Areas were assigned into networks based upon their position along the control gradient (i.e. as colored-coded in the analyses above). In both samples, a significant source x target network interaction was observed in effective connectivity (sample 1: F(4,46) = 25.1, p = 4.56e-14; sample 2: F(4,48) = 17.81, p =5.74e-11; Figure 6). In all cases, within-network directed interactions were significant and numerically
Figure 5. Estimates of static effective connectivity. Bars are color coded by source>target lobe pairs. Top: estimates for each connection. Inset: correlation among sample averaged parameter estimates. Bottom: estimates averaged over source>target lobe pairs. Influences arising from the PFC were significantly stronger in magnitude than those arising from the PPC. * indicates $p < 0.05$. 
**Figure 6.** Static Network Interactions. Top: Effective connectivity was averaged as a function of source>target network. Bars are colored as a function of the source network: TC – temporal control (red); CC – contextual control (green); SC – sensory-motor control (blue). ** denotes Bonferroni-corrected p < 0.05. * denotes p < 0.05 uncorrected. Bottom: Effective connectivity organized by abstraction (first dimension of multi-dimensional scaling depicted in Figure 3). Circles denote positive interactions and inverted triangles denote negative interactions. Markers are scaled by the magnitude of effective connectivity. Markers are colored by the network assignments.
the most positive connections for each network. This is to be expected given that areas within a cortical network are assumed to excite one another, and also provides validation of the network assignment. For both nodes in the abstract, temporal control network and the concrete, sensory-motor control network, between-network directed interactions tended to be negative. Such patterns suggest that these networks dampen activity in other networks, thereby segregating processing. By contrast, between-network directed interactions arising from contextual control nodes were significantly positive. Control analyses revealed that these dynamics were not driven by task-related signals since regressing out task-related activity demonstrated the same pattern of results (Supplemental Figure 2). Such patterns suggest that the contextual control network elevates activity in other networks, thereby promoting integrative processing.

To examine these patterns in a more continuous form that does not depend upon assigning nodes to networks, linear mixed effects models were fit using the abstraction dimension uncovered by MDS. The model sought to explain effective connectivity as a function of abstraction of the source area, abstraction of the target area, and the interaction among source and target abstraction. Additionally, quadratic forms of each of these effects was included to capture U shaped and inverted-U shaped relationships. In both samples, a significant source x target abstraction interaction was observed (sample 1: t(1097) = 5.08, p = 4.43e-07; sample 2: t(1143) = 4.94, p = 9.03e-07) reflecting that network interactions depend upon whether sources and targets are drawn from similar/different levels of abstraction. Critically, there was also a quadratic effect of source abstraction (sample 1: t(1097) = -4.51, p = 7.05e-06; sample 2: t(1143) = -4.52, p = 6.99e-06), which was driven by areas at mid-levels of abstraction having consistently positive outwards influences, while areas at high and low levels of abstraction demonstrated both positive and negative outward influences which yielded lower net influences when averaged. No other effects were significant across samples. Collectively, these patterns are consistent with the segregation/integration patterns described above and show that these patterns do not strictly depend on network assignment.

**Dynamic Integration by Contextual Control**

The static directed interactions described above indicate the potential for integration/segregation of the PFC-PPC during cognitive control. Actualization of this potential would require the activation of particular networks and corresponding activity flow. That is, activation of intermediary, contextual control nodes should produce more integrated network interactions. To examine such actualization, psychophysiological interaction (PPI) analysis was performed to estimate changes in effective connectivity as a function of different cognitive control demands.

Validation of the method by comparing estimates across samples (Supplemental Figure 3) indicated that the context-independent effective connectivity estimates were replicable (r = 0.86), PPI’s of temporal control showed modest replicability (r = 0.32), and PPI’s of contextual control showed strong replicability (r = 0.78). PPI’s of sensory-motor control were not replicable (r = -0.06). However, those areas engaged by sensory-motor control could largely be recapitulated by contrasts of stimulus domain (Supplemental Figure 4). PPI’s of stimulus domain showed strong replicability (r = 0.88). Therefore, PPI’s of stimulus domain were used as a proxy of interactions generated by activation of sensory-motor control areas.
Figure 7. Dynamic Interactions. A) Psychophysiological interactions (PPIs) for temporal control, contextual control, and stimulus domain contrasts. Solid lines indicate positive modulations of effective connectivity while dashed lines indicate negative modulations of effective connectivity. Arrows are colored by the network of the source node. Thickness of the arrows indicates the strength of modulations. Modulations are visualized at p < 0.05 uncorrected. B) Averaged within- and between-network modulations for the temporal control (red), contextual control (green), and stimulus domain (blue) contrasts. C) Integration indices computed by contrasting between- minus within-network modulations for temporal control (TC), contextual control (CC), and stimulus domain (SD) contrasts.
For each cognitive control demand, source-target relationships among PFC-PPC areas were estimated. In this case, the interactions reflect the changes in effective connectivity induced by the cognitive control demands. In contrast to the stationary dynamics, within-network PPI’s were generally weak across demands (Fig 7A,B). This indicates that within-network interactions are weakly modulated by task demands, which has been previously been observed in functional connectivity28–30. Stronger modulations were observed between networks which was especially prominent during contextual control. To quantify these effects, within- and between-network interactions were separately averaged for each PPI contrast. These were then submitted to a 2 x 3 ANOVA with factors of network interaction (within, between) and contrast (temporal, contextual, stimulus domain; Fig 7B). This analysis revealed a main effect of network interaction (sample 1: F(1,23) = 15.01, p = 0.0008; sample 2: F(1,24) = 10.34, p = 0.0037), driven by overall stronger between- than within-network interactions. There was no main effect of contrast (both samples p > 0.15). However, there was a network x contrast interaction (sample 1: F(2,46) = 11.84, p = 0.0001; sample 2: F(2,48) = 9.86, p = 0.0003). This interaction was driven by stronger between-network interactions for the contextual control contrast relative to the other contrasts. These results demonstrate that contextual control is associated with integration across PFC-PPC networks (Fig 7C).

Static and Dynamic Integration Relates to Higher-Level Cognitive Ability

The analyses above indicate that a PFC-PPC network involved in contextual control statically integrates processing among PFC-PPC networks, and that PFC-PPC networks are also more dynamically integrated when contextual control is required. Next, the relationship between individual differences in these network interactions (Fig 8A,B) and higher-level cognitive ability was examined. Each participant completed several tasks measuring short-term memory, working memory, and fluid intelligence. These measures were combined using principle components analysis (PCA) to form a composite measure of higher-level cognitive ability (Fig 8C). Measures of static and dynamic integration were then used to predict higher-level cognitive ability using 2-fold cross-validated ridge regression. The static integration measure was intended to capture individual differences in the degree to which contextual control areas integrate PFC-PPC processing in a stationary manner. In other words, this measure captured the general integration of the FPCN without regard to specific demands. By contrast, the dynamic integration measure was intended to capture individual differences in the degree to which the FPCN became more integrated when contextual control was required. That is, this measure reflects integration “on demand.” The data were separated by sample with one sample used to estimate regression weights, and the estimated regression weights used to predict higher-level cognitive ability in the other sample.

A significant prediction effect was observed (r = 0.32, p = 0.029; Fig 8D) indicating that individual differences in integration dynamics are related to higher-level cognitive ability. While static integration was associated with lower cognitive ability, dynamic integration was associated with higher cognitive ability (Fig 8E). These data indicate the importance of control network integration for higher-level cognition.

Static and Dynamic Integration Relates to Transcranial Magnetic Stimulation Susceptibility

The fMRI data for sample 2 was used to localize targets for transcranial magnetic stimulation (TMS). TMS was performed on nodes in each sub-network: FPl – temporal control; VLPFC – contextual control; SFS – sensory-motor control, as well as a control site (S1). As previously reported26, we anticipated behavioral
Figure 8. Integration predicts higher-level cognitive ability. A) Static integration was computed by contrasting between minus within-network effective connectivity of the contextual control network while controlling for (regressing out) the same contrast of the temporal control network and sensory-motor control network and individual differences in head motion. B) Dynamic integration was based upon the integration index of the contextual control PPI’s while controlling for (regressing out) the integration indices of the temporal control and stimulus domain PPI’s and individual differences in head motion. C) Higher-level cognitive ability was computed as the first principle component of a battery of cognitive tests measuring capacity. D) Cross-validated ridge regression was used to predict cognitive capacity based upon static and dynamic integration. The scatterplot depicts correlations between the predicted cognitive ability and actual cognitive ability. E) Correlations among static integration, dynamic integration, and cognitive ability.
effects on temporal control following FPl stimulation, stimulus domain processing following SFS stimulation, and a contextual control x stimulus domain processing interaction following VLPFC stimulation. These expectations were observed\(^{26}\). However, individuals varied in their susceptibility to these effects. To examine whether such susceptibilities were related to the organization of cognitive control networks, the static and dynamic integration measures described above were used to predict TMS effects.

A PCA on the TMS effects was performed to derive a general susceptibility of cognitive control to PFC TMS (Fig 9A). Static and dynamic integration measures were used to predict the cognitive control TMS effect using leave-1-out cross-validated ridge regression. This analysis revealed that TMS effects on cognitive control could be predicted by static and dynamic integration \(r = 0.56, p = 0.01\); Fig 9B). In particular, those individuals with stronger static integration tended to show increased TMS-induced cognitive control impairments, while those individuals with stronger dynamic integration tended to show decreased TMS-induced cognitive control impairments. Such data integration that integration of control networks is related to susceptibility to neuromodulation.

Discussion

The data revealed the existence of an integrative set of control areas whose dynamics relate to higher-level cognitive ability and amenability to neuromodulation. These integrative areas are situated in an intermediary zone of a mirrored PFC-PPC gradient extending outwardly from sensory-motor areas wherein sensory-motor proximal areas respond to external, present-oriented demands while sensory-motor distal areas respond to internal, future-oriented demands. Networks on either end of this control gradient acted in a segregative manner, exciting within-network nodes, while suppressing between-network nodes. Such dynamics may support selective processing of one time horizon (present/future) or medium (external/internal) at the exclusion of another. By contrast, the integration of both external/present-oriented and internal/future-oriented control networks may support hierarchical control wherein appropriate behaviors are jointly contingent on external and internal representations. The broader importance of such integrative processing was underscored by the relationship of individual differences in integrative dynamics and higher-level cognitive ability on the one hand, and amenability to neuromodulation on the other. Hence, the dynamics may be useful to specify optimal cognitive function, and predict responsivity to interventions.

There has been a recent surge of interest in macroscale gradients across the cortex\(^{6,7,36}\). This interest stems from the thought that gradients provide a scaffold for functional processing thereby offering a window into how large-scale networks support cognition. Following Mesulam\(^{5}\), Margulies et al\(^{6}\) proposed the existence of two macroscale axes upon which gradients are built with the first reflecting connectivity distance from primary cortices and the second reflecting modality. Huntenburg et al\(^{7}\) further proposed that the first gradient matches a temporal gradient reflecting the fundamental timescale over which a cortical area operates\(^{37–39}\), and the abstractness of the mental representations processed by the cortical area. These ideas are similar to those proposed by Fuster\(^{40}\) who proposed that mirrored frontal and posterior abstraction gradients supported the temporal organization of controlled behaviors. However, within these frameworks, it is unclear exactly whether or how control should be distinguished from...
Figure 9. Integration predicts transcranial magnetic stimulation (TMS) effects. A) Previously reported cognitive control impairments induced by continuous theta-burst TMS were combined using principle components analysis to derive a general susceptibility to PFC TMS. B) Correlations between cognitive control TMS effects and static and dynamic integration. Leave-one-out cross-validated ridge regression was used to predict TMS effects using static and dynamic integration. Scatterplot depicts the correlation between predicted and actual TMS impairments.
processing. That is, if the cortex can be organized along a principle axis of abstraction, does control emerge at the far end of that axis, all throughout, or in-between?

The position of the FPCN with respect to macroscale gradients of cortex is intermediary between canonical circuits involved in modality-specific processing, and the default-mode network involved in internal mentation\textsuperscript{10,41}. The intermediary positioning of the FPCN may be optimal for the orchestration of diverse brain regions in the service of adaptive behavior\textsuperscript{16}. Consistent with prior data\textsuperscript{10–12}, the data here indicate that the FPCN is not unitary but itself has a gradient organization. In particular, the data show that an intermediary zone of the FPCN, which itself is an intermediary zone of macroscale cortical gradients, is critical for the integration of control networks for adaptive behavior. This FPCN intermediary zone likely corresponds to what has been referred to as FPCN B\textsuperscript{10,12} situated between the more sensory-motor proximal dorsal attention network and the more sensory-motor distal FPCN A. Such data suggest the importance of examining gradients at multiple scales to understand brain-wide function. For example, the default-mode network has also been recently revealed to be comprised of multiple networks\textsuperscript{10,20}. Extrapolating from the findings here, default-mode network B may act as an intermediary between the memory-related default-mode network A, and the FPCN, positioned to integrate memory with control to guide internally-oriented cognition. Testing this hypothesis would be an interesting future endeavor.

Previous work has indicated the presence of an integrative core of regions important for multiple tasks\textsuperscript{13–16,42–44}. For example, Shine et al\textsuperscript{43} recently performed a spatiotemporal principle components analysis across multiple tasks of the Human Connectome Project. The first principle component of this analysis rose and fell in cadence with task demands across multiple domains with greater rising associated with greater fluid intelligence. Similarly, Cole et al\textsuperscript{42} found that the global brain connectivity of a region in mid-lateral PFC predicted individual differences in fluid intelligence. Cocchi et al\textsuperscript{24} found that a similar region showed enhanced dynamic coupling with a diverse set of brain areas as the complexity of reasoning demands increased. Collectively, these data indicate the need for integration across tasks and demands for higher-level cognition. Our data situate the areas associated with such integration in an intermediary zone of the FPCN.

In contrast to the integrative dynamics of intermediary areas, sensory-motor proximal and distal areas acted in a segregative fashion. Segregation is likely to be important to select relevant information while suppressing irrelevant information. Consistent with the data here, Shine\textsuperscript{45} posited that rostral areas of the PFC are involved in segregation while mid-lateral areas are involved in integration. He theorized that segregation is mediated by cholinergic modulations while integration is mediated by noradrenergic modulations. Further research into the influences of distinct PFC-PPC networks on neuromodulatory mechanisms would be fruitful to elucidate such effects.

The data demonstrated both static and dynamic forms of integration. Areas involved in contextual control tended to excite other control networks, providing a potential substrate for integration through binding. Moreover, demands on contextual control increased inter-network communication producing a dynamic form of integration. Both static and dynamic integration were related to individual differences in higher-level cognitive ability. Interestingly, the different forms of integration tended to be associated with opposite effects. On the one hand, increased static integration was associated with decreased higher-level cognitive ability. It has been posited that an appropriate balance of segregation and integration into distinct networks or modules is important to optimize brain efficiency\textsuperscript{46}. In particular, it has been observed that more segregation between networks, and integration within networks (i.e. modularity) at rest is
associated with memory capacity. Therefore, integration across networks in a stationary manner may be sub-optimal for cognition. On the other hand, increased dynamic integration was associated with increased higher-level cognitive ability. These data are consistent with previous studies that have shown that integrative reconfigurations of brain networks from rest to task are related to improved performance on complex tasks. These data suggest that the brain’s ability to integrate “on-demand” is beneficial to cognitive processing.

Areas and networks that bind together processing of multiple brain systems are integral to multimodal processing. A potential cost of such an organization is that integral processing hubs convey a vulnerability to cognitive processing such that damage to key areas/networks will have widespread impacts. The data here suggest that in the domain of control processing, static integrative dynamics provide increased susceptibility to TMS-induced control deficits. The form of TMS explored here was aimed to be inhibitory to cause deficits in control processing. Many therapeutic approaches to TMS utilize excitatory protocols in order to enhance irregular hypo-function (e.g. ). It would be interesting to see whether the widespread impacts of stimulation to integrative nodes can enhance therapeutic effects when excitatory protocols are employed.

The FPCN is not the only network important for integration and cognitive control. Extensive work has indicated the importance of the cingulo-opercular network. Moreover, a cognitive control organization that parallels the PFC has recently been observed in the cerebellum. In the cerebellum, as in here, intermediary areas involved in contextual control showed an integrative role, mediating relationships among control areas at either end of the external/internal, present/future axis. Detailing interactions among the FPCN and these other networks would provide valuable insights into the brain-wise basis of cognitive control.

Materials and Methods

Participants

Twenty-four participants (13 female; age 18-28, mean 19.9 years) formed sample 1, previously described. Twenty-five participants (16 female; age 18-27; mean 20.6 years) formed sample 2, previously described. Collectively, the total sample size was forty-nine.

All participants were screened to be right-handed, native English speakers or fluent by the age of 6, and had no reported history of neurological or psychiatric disorders. Informed consent was obtained in accordance with the Committee for Protection of Human Subjects at the University of California, Berkeley.

The targeted number of participants was based upon previous work with related paradigms. The empirical replicability of univariate activations from these data has been previously established indicating that power is sufficient for the estimation of cognitive control networks at the group level at the collected sample sizes. To ensure replicability of more complex analyses, bi-variate, and effective connectivity analyses in the present study are reported separately for each sample when possible, providing indications of replicability and power sufficiency.
Experimental Design and Statistical Analyses

Comprehensive Control Task

The Comprehensive Control Task was adapted from prior work, and designed to manipulate multiple forms of cognitive control within a single, well-controlled paradigm. These different forms of control can be classed by different levels of abstraction with concrete processing acting on external stimuli in the present moment, and abstract processing supporting the maintenance of internal representations to guide future behavior. At the lowest level, sensory-motor control selects relevant stimulus features and associated actions. At the mid-level, contextual control selects the rules that guide the appropriate stimulus-response associations. Finally, at the highest level, temporal control supports temporally-extended representations that prepare for future control demands.

On each trial of the task (Figure 1), participants were presented with a letter at one of five spatial locations surrounded by a colored shape. Each stimulus required a choice decision indicated by a left or right keypress. Choice-to-keypress mappings were counter-balanced between participants. The correct decision depended upon a combination of 1) a stimulus feature, 2) a contextual rule, and 3) a temporal epoch.

Stimulus feature: choice decisions were based either on the letter (verbal task) or spatial location (spatial task). Participants pre-learned a sequence of letters (t-a-b-l-e-t) and locations (trace of a star, starting at the top position). Choice decisions were based upon these sequences.

Contextual rule: for a given stimulus feature, participants determined either whether the present stimulus was the first item of the relevant sequence (‘t’ for the verbal task; top position for the spatial task; sequence start task) or whether the present stimulus followed a reference stimulus in the sequence (sequence back task).

Temporal epoch: for the sequence back task, the reference stimulus was either the immediately preceding stimulus, or a stimulus that appeared multiple trials ago.

Loads on each of these factors were orthogonally varied to form a 2 x 2 x 2 factorial design. Trials were grouped into blocks wherein each block sampled one cell of the 2 x 2 x 2 design. In each block, the relevant stimulus feature, either letter or location, was cued by the color of the frame. The relevant stimulus feature remained constant throughout a block. Participants performed the sequence start task on the relevant stimulus feature on the first trial of a given block. Thereafter, each block was divided into three phases: a first baseline phase, a sub-task phase, and a second baseline phase. Transitions among these phases were indicated by the shape of the colored frames. In all blocks, the baseline phases were cued by square frames. Sub-tasks were cued by triangle, diamond, or cross frames with shape-to-sub-task mappings counter-balanced across participants. For ease of exposition in what follows, we will assume the following associations: circle-switching, triangle-planning, diamond-dual. Only one sub-task was cued in a given block.

In Baseline blocks, participants continued to perform the sequence back task throughout the block thereby keeping the contextual rule and temporal epoch constant. On Switching blocks, shape-switches (i.e. from square to circle or from circle to square) indicated the need to switch tasks (i.e. from sequence back to sequence start). Thus, these blocks placed demands on responding based upon the appropriate contextual rule engaging contextual control. On Planning blocks, triangle shapes indicated that the
presented stimulus could be ignored. Participants acknowledged the presence of each triangle-framed stimulus with a “no” keypress. All the while, the last square-shaped stimulus had to be retained as a reference for the next square-shaped stimulus. Hence, Planning blocks minimized processing of present stimuli, but placed demands on planning for the future in order to respond according to the correct temporal epoch thereby requiring temporal control. Finally, on Dual blocks, diamond frames indicated the need to both switch contextual rules, and plan for the future. That is, participants started and continued a new sequence on all diamond framed stimuli, but backwards-matched to the last square-framed stimulus when the frame reverted to a square. Hence, Dual blocks required both contextual and temporal control.

Different cognitive control process can be detailed through orthogonal contrasts of the factorial design. At the highest level, temporal control is isolated by contrasting blocks that require planning with those that do not (Dual + Planning > Switching + Baseline). At the middle level, contextual control is isolated by contrasting blocks that require switching with those that do not (Dual + Switching > Planning + Baseline). At the lowest level is sensory-motor control which can be examined through the contrast of (Dual + Baseline > Planning + Switching). To understand this contrast, consider that the Dual condition is effectively Switching + Planning. Then, using subtraction logic, what remains after subtraction is the Baseline condition which consists of the demands of selecting the appropriate feature and responding to it. The utility of using the contrast over-and-above simply examining the Baseline condition alone is that it better controls for ancillary demands, and it keeps all of the main contrasts of interest statistically orthogonal to one another. Finally, those areas involved in sensory-motor control can be further fractionated by contrasting stimulus features with one another (i.e. verbal > spatial or spatial > verbal), thereby emphasizing verbal articulatory processes (verbal > spatial) or spatial attention processes (spatial > verbal).

Participants in sample 1 completed 24 blocks of each cell of the 2 x 2 x 2 design over the course of 2 fMRI sessions. These blocks consisted of 1920 total trials with each block containing 7 to 13 trials. Participants in sample 2 completed 12 blocks of each cell of the 2 x 2 x 2 design during a single fMRI session. These blocks consisted of 864 total trials with each block containing 7 to 11 trials. In both samples, the task was split across 6 runs of 16 blocks each in each session.

Each stimulus was presented for 500 ms followed by a variable inter-trial interval of 2600-3400 ms. At the end of each block, feedback indicating the number of correct trials in the block out of the total number of trials in the block was presented for 500 ms. A variable 2600-3400 ms interval separated each block. Self-paced breaks were administered in-between runs.

Within a week prior to scanning, participants performed a practice session to learn the task. During the practice session, participants received extensive written instruction and clarification from an experimenter. Given the numerous rules and complexities of the task, instruction was broken up such that participants first learned the verbal sequence, then the spatial sequence, and then the sub-tasks. Participants continued to repeat instruction and practice under experimental supervision until they were comfortable with the rules. Thereafter, participants completed 3 runs of the task on their own. During each scanning session, participants completed one additional practice run in the scanner prior to fMRI data collection.
Cognitive Battery

To obtain trait-level measures of cognitive ability, participants performed computerized versions of letter span, spatial span, operation span, and symmetry span. Operation span and symmetry span provide measures of working memory capacity. Letter span and spatial span provide simple measures of short-term memory capacity. Participants also performed a paper version of Raven’s Advanced Progressive Matrices (www.perasonassessments.com), wherein Set I was used as practice, and Set II was used to measure fluid intelligence. As previously reported, performance across these measures were correlated. Therefore, the measures were combined using principle components analysis with the first principle component providing an index of general higher-level cognitive ability.

Image Acquisition

As previously reported, brain imaging data were acquired at the Henry H. Wheeler, Jr. Brain Imaging Center at the University of California, Berkeley using a Siemens TIM/Trio 3T MRI equipped with a 32-channel head coil. Stimuli were projected to a coil-attached mirror using an MRI-compatible Avotec projector (Avotec, Inc.; https://avotecinc.com). Experimental tasks were created using E-Prime software version 2.0 (Psychology Software Tools, Inc; https://pstnet.com). Eye position was monitored using an Avotec system (RE-5700) and Viewpoint software (http://www.arringtonresearch.com). Response data were collected on an MR-compatible button box (Current Designs, Inc.; https://curdes.com).

T2* weighted fMRI was performed using gradient echo planar imaging (EPI) with 3.4375 mm² in-plane resolution and 35 descending slices of 3.75 mm thickness. TR=2000ms, echo time = 25 ms, flip angle = 70, field of view = 220 mm². The first three images of each run were automatically discarded to allow for image stabilization. Field maps were collected to correct for magnetic distortion. High-resolution T1-weighted MPRAGE images were collected for anatomical localization and spatial normalization (240 x 256 x 160 matrix of 1 mm³ isotropic voxels; TR = 2300 ms; echo time = 2.98 ms; flip angle = 9). Each session included a 6-minute eyes open resting state run in addition to six task runs. In sample 1, the resting-state run was collected prior to the task in the first session, and after the task in the second session. In sample 2, the resting-state run was collected prior to the task.

Behavioral Data Preprocessing

Behavioral data was preprocessed using custom MATLAB code (MathWorks; https://www.mathworks.com). Data were sorted by phase (first baseline, sub-task, second baseline). Transitions between phases were considered separately. Of primary importance in this report are the sub-task trials (i.e. all trials in the sub-task phase excluding the first trial of the phase) and what are referred to as “return” trials which mark the transition from the sub-task phase to the second baseline phase. For example, in Planning blocks, the sub-task trials themselves require minimal processing since all stimuli during this phase require a “no” keypress. On the return trial, participants perform sequence back with reference to the stimulus feature that preceded the sub-task phase. Thus, the sub-task trials reflect a combination of cognitive control in the moment and preparing for the future, while the return trials reflect actualization of sub-task preparation. Sub-task and return trials were separately sorted for each cell of the 2 x 2 x 2 design. Reaction times were computed on trials with a correct response only. Reaction times less...
than 200 ms were discarded as anticipatory and reaction times greater than 2000 ms were discarded as inattentive. Furthermore, reaction times greater than 2.5 standard deviations of the mean of a given condition were removed as outliers. These procedures resulted in the removal of 0.76% of the trials in sample 1, and 0.65% of the trials in sample 2.

**Image Preprocessing**

Unless otherwise specified, preprocessing was performed using SPM8 ([http://www.fil.ion.ucl.ac.uk/spm/](http://www.fil.ion.ucl.ac.uk/spm/)). Raw data was converted from DICOM into nifti format. Origins for all images were manually set to the anterior commissure. AFNI’s 3dDespike function ([http://afni.nimh.nih.gov/afni](http://afni.nimh.nih.gov/afni)) was applied to the functional data to remove outlier time-points. Correction for differences in slice-timing and spatial realignment were performed next. Images were then corrected for magnetic distortion using the FieldMap toolbox implemented in SPM8. Functional data were co-registered to the T1-weighted image. Segmented normalization was performed on the T1-weighted image. Spatial normalization parameters were subsequently applied to the functional task data. This step also resampled the functional images to 2 mm³ isotropic. Task data were subsequently smoothed using an 8mm³ full-width/half-maximum isotropic Gaussian kernel. Prior to spatial normalization, resting data underwent additional preprocessing using custom-written MATLAB scripts. Resting data were high-pass filtered at 0.009 Hz using SPM8’s `spm_filter` function. Next, nuisance signals from white matter and cerebrospinal fluid were extracted and regressed out of the data along with linear, squared, differential, and squared differential motion parameters (motion parameters were estimated during spatial realignment described above). The residualized data were low pass filtered at 0.08 Hz using a second-order Butterworth filter. The filtered, artifact-corrected resting data were then spatially normalized and smoothed as described above.

The bulk of the report focuses on data that were preprocessed in the manner described above. However, large volumetric smoothing kernels can blur and obscure activation gradients. To ensure that the observed activation gradients were not artfactually caused by these processing choices, the data were re-processed using Freesurfer version 6.0 ([https://surfer.nmr.mgh.harvard.edu/](https://surfer.nmr.mgh.harvard.edu/)). The T1-weighted image was registered to the fsaverage surface using recon-all. Despiked, slice-timing corrected, realigned, unwarped functional data described above (i.e. before spatial normalization) were sampled onto the fsaverage surface using preproc-sess (note, motion correction was omitted using the –nomc flag since motion correction was previously performed). 4mm smoothing was performed on the surface and then the functional data was returned to the volume using surfsmooth-sess. In this case, returning the data to the volume enabled the same model estimation procedures to be performed on both the volume smoothed and surface smoothed data, providing a well-matched comparison.

**Univariate Image Analysis**

Univariate modeling of task activation has been previously described[25] and was implemented using SPM8. Of main interest are the sub-task phases which were separately modeled for each cell of the 2 x 2 x 2 design as epochs starting after the first sub-task trial through the end of the final sub-task trial. The first and second baseline trials were also modeled as epochs, separately for each phase and stimulus domain,
also starting from the second trial of the epoch through the last. Transient regressors were included to model the first trial of each block, separately for each stimulus domain, the first trial of each sub-task, separately for each cell of the 2 x 2 x 2 design, and the first trial of the second baseline (also known as the “return” trial), separately for each cell of the 2 x 2 x 2 design. Additional transient regressors were included to model left and right keypresses, as well as error trials. All regressors were convolved with the canonical hemodynamic response function implemented in SPM. Each run was modeled separately containing a separate intercept term, high-pass filter at 128s, AR(1) modeling to account for temporal autocorrelation, and scaling such that run-wide global signal averaged 100. For participants demonstrating greater than 3mm/degrees of motion over the course of the session or a single framewise movement greater than 0.5mm/degrees, 24 motion regressors were included reflecting linear, squared, differential, and squared differential to remove motion-related artifacts.

At the first level, contrasts were created to isolate the sub-task phase of each cell of the 2 x 2 x 2 design by averaging parameter estimates across runs. These first level parameter estimates were then carried forward to a second level 2 x 2 x 2 ANOVA. Second-level contrasts were calculated as t-tests within the ANOVA framework. Five orthogonal contrasts of interest were considered: Temporal Control (Dual + Planning > Switching + Baseline), Contextual Control (Dual + Switching > Planning + Baseline), Sensory-Motor Control (Dual + Baseline > Planning + Switching), Verbal Stimulus Domain (Verbal > Spatial), and Spatial Stimulus Domain (Spatial > Verbal). Note that the contrasts with the “Control” modifier collapsed across Stimulus Domain (e.g. Dual is a combination of Verbal Dual and Spatial Dual), while contrasts with the Stimulus Domain modifier collapsed across the Control conditions (e.g. Verbal is a combination of Verbal Dual, Verbal Planning, Verbal Switching, and Verbal Baseline). In keeping with previous reports on these datasets, whole-brain voxel-wise activation maps were thresholded at p < 0.001 at the voxel level with 124 voxel extent providing correction for family-wise error (FWE) at p < 0.05 according to AlphaSim.

Regions-Of-Interest

The five contrasts of interest produced wide swaths of activation throughout the PFC and PPC. To examine the activations in a more granular manner, regions-of-interest (ROIs) were created as 6mm³ spheres centered on peak activation coordinates. PFC ROIs have been previously described. Given that the contrasts produced partially overlapping activations, distinct peaks were defined as local maxima that were distanced at least 1.5 cm from another peak. The location of PFC peaks was determined by sample 1 and peaks similar in location revealed by the same contrast were identified in sample 2. This ordering was due to the temporal ordering of data collection and analysis (i.e. sample 1 was collected and analyzed first and sample 2 was analyzed in an effort to replicate findings in sample 1 as detailed previously). This resulted in six PFC ROIs detailed in Table 1: lateral frontal pole (FPl), middle frontal gyrus (MFG), ventrolateral prefrontal cortex (VLPFC), caudal middle frontal gyrus (cMFG), inferior frontal junction (IFJ), and superior frontal sulcus (SFS).

ROIs in the PPC were defined separately for sample 1 and sample 2, again using the five contrasts-of-interest. Again, distinct peaks were identified as local maxima distanced from other peaks by at least 1.5 cm. This procedure results in four PPC ROIs detailed in Table 1: inferior parietal lobule (IPL), mid intraparietal sulcus (midIPS), superior parietal lobule (SPL), and anterior intraparietal sulcus (antiIPS).
In both the PFC and PPC, some peaks could be identified by two or three contrasts (i.e. local maximum within 1.5 cm from one another were observed across multiple contrasts). The contrast chosen to define a given peak was done with consideration of the contrast that most unambiguously identified the peak in individual subjects. This choice was made since the effective connectivity analyses detailed below were tailored to each individual’s activation peaks, providing individual-specific parcellation. The end result of this analysis choice was that the least expansive contrast (where expansive refers to the number of activated voxels of a contrast) tended to be chosen to define an ROI for a given peak.

Functional Profiling and Multi-Dimensional Scaling

Since ROIs could respond to multiple demands, we sought to quantify each ROI’s activation profile. Because the ROIs were formed from peaks of a particular contrast in a particular sample, simply plotting the activation profile of the ROIs defined above constitutes circularity that biases the profile towards the ROI-defining contrast. To provide unbiased estimates, the ROIs from sample 1 were used to functionally profile sample 2, and vice versa. Hence, data used to define ROIs and test ROIs were independent. Within these unbiased ROIs, statistics for the five contrasts-of-interest were calculated using repeated-measures 2 x 2 x 2 ANOVAs. For each sample, statistical significance was Bonferroni corrected for the number of ROIs.

To further characterize each ROI, the ROI data were submitted to multi-dimensional scaling. The data for each individual and each ROI were z-scored across the eight conditions-of-interest. Then, for each ROI, the individual-level data were stacked, resulting in a two-dimensional matrix wherein one-dimension was composed of individuals x conditions, and the other dimension was composed of ROIs. Dissimilarity among ROIs was computed using 1 – r, where r is the Pearson’s correlation. This dissimilarity matrix was submitted to MATLAB’s cmdscale function. Each ROI was then plotted as a point in the space defined by the first two dimensions wherein the first two dimensions captured 89.2% of the variance in the data (71.0% for the first, 18.3% for the second, 0.04% for the third). As explained in the Results, the first dimension appeared to map well onto an abstraction axis, and the second dimension appeared to map well onto a stimulus domain axis.

Brain-Behavior Relationships

To better understand how activation in each ROI relates to cognitive control, relationships between behavioral performance in the Comprehensive Control Task and activation were calculated similar to previous reports. Three differences between the analyses reported here and those reported previously is that here, 1) each ROI is considered separately (pairs of ROIs were combined previously), 2) ROI data were extracted to ensure independence as explained above (i.e. sample 1 defined ROIs for sample 2, and vice versa), and 3) repeated measures correlations were computed using the rmcorr package in R (previously correlations were computed by regressing out subject-specific terms). Mean RT for the eight conditions of interest were calculated for each individual separately for the sub-task trials and return trials, as described above. The eight element vector was then z-scored separately for each phase. The sub-task trial RTs were regarded as “Present Behavior” since the RTs reflect behavior at the time that the fMRI signal was estimated (i.e. activations of interest were drawn from the sub-task phase).
The return trial RTs were regarded as “Future Behavior” since the RTs reflect the time directly after the fMRI signal was estimated. Future Behavior reflects the behavior that is being prepared for during the sub-task phase, which might not be reflected in the sub-task trial RT per se (e.g. Planning blocks have fast Present RTs, but slow Future RTs). For each ROI, a repeated measures correlation was used to assess the relationship between activation and Present Behavior (controlling for Future Behavior), and activation and Future Behavior (controlling for Present Behavior). These procedures were performed separately for each sample. Statistical significance was assessed after Bonferroni correction for number of ROIs. Correlations that did not survive multiple comparisons correction, but were present at uncorrected levels are also noted.

Linear mixed effects models were fit in order to quantify how ROIs differ in terms of their relationships to Present and Future Behavior. For each individual and ROI, Present and Future Behavior were simultaneously regressed onto the activations. Parameter estimates reflecting the slope relating activation to Present and Future Behavior, respectively were submitted to the linear mixed effects models. For each ROI, the loading of the ROI on the first dimension of the multi-dimensional scaling described above was used as a continuous, objective measure of abstraction. Separate linear mixed effects models tested whether the relationship between Present+ RT and activation varied as a function abstraction, and whether the relationship between Future RT and activation varied as a function of abstraction. That is, these analyses tested whether the slope estimates varied by abstraction. Models of the form: Present Behavior Slope ~ Abstraction + (Abstraction|Subject), and Future Behavior Slope ~ Abstraction + (Abstraction|Subject) were estimated. These analyses were performed separately for each sample.

ROI analyses offer an incomplete picture of brain-behavior relationships. To better visualize whole-brain patterns, voxel-wise partial correlations were performed. The data were stacked across individuals. For each voxel, the partial correlation between activation and Present Behavior, controlling for Future Behavior and individual, as well as the partial correlation between activation and Future Behavior, controlling for Present Behavior and individual were calculated. The resultant maps were thresholded at p < 0.001 at the voxel-level, and clusters of 124 voxels or more were retained for visualization.

**Resting-State Functional Connectivity**

Of main interest in this report is effective connectivity which estimates directed relationships among areas. To constrain these analyses, seed-based functional connectivity was performed on the resting-state data in order to identify ROI’s that may be connected to one another (and prune connections that are likely to not be present). For each ROI, voxel-wise correlations were computed between the ROI time-series and each voxel time-series. This was done separately for each resting-state run in sample 1 (note, only sample 1 was used here to maintain compatibility with the connectivity matrix defined previously). For a given seed ROI, the correlation values of each voxel were ranked after excluding voxels within 1.5 cm of the seed ROI. A correlation value r at cost c was computed wherein r reflects the correlation value of voxel at the cth percentile. Here, c was set to 18 as detailed previously. Then, four counts were made: whether ROI 1 was connected to ROI 2 when ROI 1 was the seed in run 1, whether ROI 1 was connected to ROI 2 when ROI 2 was the seed in run 1, whether ROI 1 was connected to ROI 2 when ROI 2 was the seed in run 2, and whether ROI 1 was connected to ROI 2 when ROI 2 was the seed in run 2. A pair of ROIs were considered “connected” if the average correlation value in the target ROI exceeded r. If at least 3 of the 4 counts were tallied, the connection between the areas was considered a candidate for future
The resultant candidate connectivity matrix was utilized for effective connectivity analyses. The intent of this procedure was to be somewhat liberal, and allows the effective connectivity modeling to further prune connections. Hence, this procedure should be considered a scaffold for analyses to follow rather than an attempt to determine the true connectivity structure of the data.

**Spectral Dynamic Causal Modeling**

Time-invariant effective connectivity was estimated using spectral dynamic causal modeling (spDCM) implemented in SPM12 and DCM12.5. spDCM inverts a biophysically plausible generative model of the fMRI cross-spectral density\textsuperscript{31–33}. The model estimates the effective connectivity among hidden neural states and includes a hemodynamic forward model that accounts for how synaptic activity translates into regional hemodynamics and observed connectivity in the fMRI signal. spDCM estimated the effective connectivity among the six PFC and four PPC areas that are the focus of this report. Given that each individual has a specific topography in the PFC\textsuperscript{10}, volumes of interest (VOIs) were defined based on each individual’s activation peaks. A 6mm\textsuperscript{3} sphere was created around each individual’s activation peak that was close to the peak in the group contrast. To reduce noise in the VOIs, tiered thresholding was performed to eliminate unresponsive voxels. This threshold was started at p < 0.001 (80.0% of VOIs), was lowered to p < 0.05 if voxels did not survive (14.4% VOIs). If voxels still did not survive, the VOI was centered at the group peak and the threshold was lowered further to p < 0.5 (5.1% of VOIs), and the threshold was eliminated if this was still insufficient (0.004% of VOIs). The model architecture utilized the candidate connectivity matrix estimated from the resting-state data described above.

spDCM’s required the estimation of 82 parameters using the model architecture based upon the candidate connectivity matrix. To provide sufficient data to estimate these parameters, pairs of runs were collapsed together using a modified version of spm_fmri_concatenate. The modifications allowed a flexible number of runs to be concatenated (default behavior is to concatenate all runs), and added a regressor to account for edge effects caused by run-wise (i.e. not concatenated) high-pass filtering. Other changes relative to the univariate modeling was that models of all participants included linear, squared, differential, and squared differential motion parameters, as well as a regressor to capture framewise displacement calculated across the six motion parameters\textsuperscript{65}. The time-series’ for every VOI was extracted after regressing out all regressors related to confounds of non-interest (i.e. run means, motion, filter edge effects).

For each participant, two spDCM’s were estimated. The first modeled effective connectivity on time-series’ that excluded only confounds signals. The second modeled effective connectivity on time-series’ that additionally regressed out all task-related regressors included in the univariate modeling. Although spDCM is designed to estimate time-invariant effective connectivity, it is possible that task-related activity may nevertheless influence effective connectivity estimates\textsuperscript{66}. However, each of these spDCM’s produced similar results with no evidence of bias produced by including/excluding task-related regressors (Supplemental Figure 2) consistent with the idea that spDCM estimated time-invariant effective connectivity as designed.

The following defaults were changed: the order of the AR model was set to 4 (this was the default for DCM12, but the default is 8 in DCM12.5). This was done after observing systematically poorer fits at the default levels in preliminary analysis. The maximum number of nodes to model was increased to 10 to
accommodate the number of areas of interest. Finally, the maximum number of iterations for model fitting was increased to 1000 to ensure that all models converged.

spDCM was performed separately for each sample. This enabled the examination of the replicability of effective connectivity estimates and derivations thereof.

**Psychophysiological Interaction Analysis**

To examine dynamic (time-varying) effective connectivity, psychophysiological interaction (PPI) analysis was performed. Here, a variant of the method detailed in Cole et al\(^ {16} \) was employed. Time-series’ from the same VOIs that were submitted to spDCM were modeled (in this case, the VOIs without the task regressed out since the task regressors are regressed out as part of PPI). For each connection in the probable connectivity matrix, the data were modeled as:

\[
Y_{\text{target}} = B_{\text{conn}} Y_{\text{source}} + B_{\text{uni}} X + B_{\text{dep}} (\text{bin}(X) * C * Y_{\text{source}})
\]

Here, \( Y_{\text{target}} \) is the time-series of the target VOI, \( Y_{\text{source}} \) is the time-series of the source VOI, \( X \) is the univariate design matrix, \( \text{bin}(X) \) is a binarized version of the univariate design matrix wherein time-points that are above 0 are set to 1 and other time-points are set to 0, and \( C \) is the contrast matrix. This results in estimations of several contributions to the signal in the target VOI: \( B_{\text{conn}} \) which represents the time-invariant functional connectivity among the target and source, \( B_{\text{uni}} \) which represents the univariate time-invariant task signals, and of main interest, \( B_{\text{dep}} \) which is the context-dependent changes in effective connectivity from the source to the target. This formulation is identical to Cole et al\(^ {16} \) except that contrasts are included within the model itself rather than being computed after parameter estimation. This change was performed because it reduces multi-collinearity, which is particularly problematic for PPI among highly correlated nodes. Preliminary analyses revealed that the more connected two regions were, the more negatively biased \( B_{\text{dep}} \) became using the method of Cole et al\(^ {16} \). This bias was eliminated by utilizing contrasts within the model.

Context-dependent connectivity (i.e. the PPI) is dictated by the contrasts defined in \( C \). Here, five contrasts were defined which correspond to the four univariate contrasts of interest (Temporal Control, Contextual Control, Sensory-motor Control, Stimulus Domain) plus an additional interaction contrast that modeled Contextual Control x Stimulus Domain.

PPI analyses were performed separately for each sample. This enabled the examination of the replicability of the effective connectivity estimates and derivations thereof. PPI’s of Sensory-motor Control and Contextual Control x Stimulus Domain were unreliable across samples (Supplemental Figure 3). Therefore, analyses were restricted to PPI’s of Temporal Control, Contextual Control, and Stimulus Domain.

**Predicting higher-level cognitive ability**

spDCM and PPI provided measures of static and dynamic effective connectivity, respectively. In particular, spDCM analyses revealed a role of contextual control-related areas in integrating cognitive control networks, and PPI analyses revealed greater integration across cognitive control networks during contextual control conditions. To examine the relevance of these measures to higher-level cognition,
static and dynamic effective connectivity was used to predict individual-differences in general higher-level cognitive ability.

General higher-level cognitive ability was assessed using the first principle component of the scores on the cognitive battery described above. To provide an index of static integration, for each network, between-network effective connectivity was contrasted against within-network effective connectivity. To isolate static integration of the contextual control network, the residualized static integration of the contextual control network was utilized after regressing out the static integration indices of the temporal control and sensory-motor control networks, as well as the root-mean square of framewise displacement (RMS FD) as a confound. Dynamic integration was calculated in a parallel manner to static integration. In this case, between-network effective connectivity was contrasted with within-network effective connectivity across all networks, but isolated to particular contrasts. For example, dynamic integration of contextual control indexed the degree to which there was greater between- relative to within-network communication for the contextual control contrast. Intuitively, whereas static integration of the contextual control network indicates the potential for contextual control areas to integrate networks, dynamic integration during contextual control indicates the actualization of this integration during the conditions that maximally activate contextual control areas. Similar to the static integration index, the residualized dynamic integration during contextual control was utilized after regressing out the dynamic integration indices during temporal control and stimulus processing, along with RMS FD as a confound.

To ensure independence among our samples, each of these indices (general higher-level cognitive ability, static integration, dynamic integration) were calculated separately on sample 1 and sample 2.

Ridge regression was performed to predict general higher-level cognitive ability from static and dynamic integration. This was done in a two-fold manner, wherein each fold divided the dataset into training and test sets as a function of the samples (i.e. sample 1 and sample 2). For example, sample 1 was used to estimate weights using ridge regression, and then those weights were used to predict general higher-level cognitive ability in sample 2. Ridge regression was performed with an arbitrarily chosen penalty parameter $= 1$, although significant prediction was achieved at penalties as high as 100 (which was the highest tested). The correlation between the observed and predicted general higher-level cognitive abilities was used to assess prediction performance. The observed correlation was compared to a permutation distribution formed by randomly permuting the rows of the prediction matrix 1000 times.

**Predicting Transcranial Magnetic Stimulation Susceptibility**

As previously reported$^{26}$, sample 2 was used to test the causal role of frontal areas to different forms of cognitive control. Each participant in sample 2 underwent an additional four sessions of the Comprehensive Control Task following continuous theta-burst stimulation (cTBS) to each of FPl, VLPFC, SFS, and S1 (control site). cTBS is patterned form of transcranial magnetic stimulation (TMS) thought to inhibit cortical tissue$^{33}$. Each TMS target was a node in a distinct PFC-PPC network and was predicted to produce distinct impairments in cognitive control. To examine whether static and dynamic integration of cognitive control networks is related to TMS susceptibility, the three observed TMS effects of interest were combined: the effect of FPl TMS on temporal control on error-rate, the effect of VLPFC TMS on the contextual control x stimulus-domain interaction on reaction time, and the effect of SFS TMS on stimulus-domain on error-rate (see$^{26}$ for additional details). For each effect, the frontal TMS effect was isolated by
regressing out the corresponding effect following S1 TMS. Then, all TMS effects were combined using the first principle component.

Leave-1-out ridge regression was performed to predict the general cognitive control TMS effect from static and dynamic integration, wherein static and dynamic integration were computed identically to the above. The correlation between the observed and predicted general higher-level cognitive abilities was used to assess prediction performance. The observed correlation was compared to a permutation distribution formed by randomly permuting the rows of the prediction matrix 1000 times.

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Data Availability

All data and code needed to reproduce the figures in this report can be found at https://osf.io/938dx/.

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Supplemental Material

Supplemental Behavioral Results

Behavioral data have been previously reported\textsuperscript{1,2}. Here, the focus is on two phases of the data: the sub-task phase wherein cognitive control demands are manipulated, and the return trials, wherein preparatory control processes readied during the sub-task phase are expressed.

Due to the complex design, it is useful to highlight particular data patterns that provide assurance that the task manipulations operate as desired. The interested reader is directed to Supplemental Tables 1 and 2 for complete factorial statistics.

First, effects of stimulus domain were examined. In sample 1, RTs on verbal and spatial tasks were well-matched during the sub-task (verbal: 644.32 ms; spatial: 642.4 ms; $F(1,23) = 0.03, p = 0.87$) and return trials (verbal: 829.5 ms; spatial: 823.7 ms; $F(1,23) = 0.12, p = 0.74$). Accuracies were also similar during sub-task (verbal: 92.8%; spatial: 93.9%; $F(1,23) = 3.12, p = 0.09$), and return trials (verbal: 97.9%; spatial: 98.1%; $F(1,23) = 3.01, p = 0.10$), though trending towards more errors on the verbal task. In sample 2, RT data provided some evidence that the verbal task was more difficult than the spatial task on sub-task trials (verbal: 792.9 ms; spatial: 757.8 ms; $F(1,24) = 5.01, p = 0.03$), with a similar trend on return trials (verbal: 932.7 ms; spatial: 908.0 ms; $F(1,24) = 3.80, p = 0.06$). Accuracies were not significantly different on sub-task (verbal: 94.5%; spatial: 95.4%; $F(1,24) = 2.90, p = 0.10$) nor return trials (verbal: 84.5%; spatial: 84.6%; $F(1,24) = 0.01, p = 0.91$). Combining the samples, there was no significant difference in verbal and spatial RTs during the sub-task ($F(1,48) = 3.55, p = 0.07$) and return trials ($F(1,48) = 2.32, p = 0.13$). There was a significant difference in verbal and spatial accuracies during the sub-task ($F(1,48) = 6.09, p = 0.02$), but not return trials ($F(1,48) = 1.62, p = 0.21$). Collectively, these data provide some scattered evidence that the verbal task was modestly more challenging than the spatial task, but overall, the stimulus domains were reasonably well-matched.

Second, it is desirable that the cognitive control manipulations produce clear and consistent effects on behavior. To illuminate these effects, I consider simple comparisons to the Baseline condition, although clear and consistent effects are observed using the full factorial framework and detailed in Supplemental Tables 1 and 2. Sensory-motor control involves selecting an appropriate action based upon the stimulus. Demands on these processes are minimized during the Planning sub-task phase, during which participants can reflexively respond without processing the verbal or spatial stimulus features. Correspondingly, RTs are expected to be speeded during the Planning sub-task phase. These expectations were met in both samples (sample 1: mean difference $= -184.1$ ms; Cohen’s $d = 2.09$; $t(23) = -10.24, p = 4.85e-10$; sample 2: mean difference $= -211.7$ ms; Cohen’s $d = 1.61$; $t(24) = -8.06, p = 2.75e-8$). Accuracies were also elevated in the Planning sub-task phase relative to Baseline indicating that RT speeding did not tradeoff from accuracy (sample 1: mean difference $= 5.3$%; Cohen’s $d = 0.87$; $t(23) = 4.25, p = 0.0003$; sample 2: mean difference $= 3.6$%; Cohen’s $d = 0.71$; $t(24) = 3.56, p = 0.002$)

Next, contextual control involves selecting the appropriate task set over-and-above sensory-motor control. This should incur an additional cost in RT reflected when comparing the Switching sub-task phase to the Baseline condition. These expectations were met in both samples (sample 1: mean difference $= 60.8$ ms; Cohen’s $d = 1.08$; $t(23) = 5.31, p = 2.17e-5$; sample 2: 189.0 ms; Cohen’s $d = 1.53$; $t(24) = 7.65, p = 6.90e-8$). Accuracy was either the same or slightly worse in the Switching condition, indicating that
slowing was not a result of speed-accuracy tradeoff; sample 1: mean difference = 0.4%; Cohen’s d = 0.10; t(23) = 0.51, p = 0.62; sample 2: mean difference = -1.8%; Cohen’s d = -0.43; t(24) = -2.16, p = 0.04).

Finally, temporal control involves utilizing temporally extended representations to guide action. In the Planning condition, these representations are sustained during the sub-task phase, and utilized during the return trial. Hence, demands on temporal control are most clearly manifest in behavior on the return trial. In contrast to the above analyses that showed speeded RTs during the sub-task phase of the Planning condition relative to the Baseline condition, this same comparison on return trials showed slowed RTs for the Planning condition in both samples (sample 1: mean difference = 88.8 ms; Cohen’s d = 0.65; t(23) = 3.20, p = 0.004; sample 2: mean difference = 71.9 ms; Cohen’s d = 0.59; t(24) = 2.96, p = 0.007). Accuracies were also diminished (sample 1: mean difference = -13.1%; Cohen’s d = 0.93; t(23) = -4.54, p = 0.0001; sample 2: mean difference = -6.5%; Cohen’s d = 0.63; t(24) = -3.16, p = 0.004).

Collectively, the behavioral data produced the expected cognitive control-related patterns.
Supplemental Figure 1. Contrasts of temporal control (red), contextual control (green), and sensory-motor control (blue) and their overlap using surface-based processing and reduced smoothing to minimize blurring.
Supplemental Figure 2. A) Comparison of effective connectivity parameters with and without regressing out task signals. Bias (e.g. task-induced increases) would be apparent as a shift off of the diagonal (red line). No bias was observed. B) Source x target network interactions after regressing out task signals. The same interaction was observed after regressing out task signals as was observed in the main text indicating that the interactions were not driven by task signals.
Supplemental Figure 3. Sample averaged psychophysiological interaction (PPI) parameter estimates. TC – temporal control; CC – contextual control; SC – sensory-motor control; SD – stimulus domain.
Supplemental Figure 4. Comparison of the sensory-motor control contrast and stimulus domain contrasts. Stimulus domain is jointly determined by verbal > spatial and spatial > verbal.
| Effect       | Samp 1 |          | Samp 2 |          | Combine |          |
|--------------|--------|----------|--------|----------|---------|----------|
|              | RT     | ACC      | RT     | ACC      | RT      | ACC      |
| ME Stim Domain | 0.03   | 0.8691   | 3.12   | 0.0905   | 5.01    | 0.0348   |
| ME Context Control | 150.96 | <0.0001  | 16.22  | 0.0005   | 144.39  | <0.0001  |
| ME Temp Control | 24.23  | 0.0001   | 5.71   | 0.0255   | 11.22   | 0.0027   |
| Stim Domain X Context Control | 0.04   | 0.8395   | 0.04   | 0.8392   | 0.14    | 0.7091   |
| Stim Domain X Temp Control | 7.09   | 0.0139   | 0.13   | 0.7172   | 1.82    | 0.1900   |
| Context Control X Temp Control | 96.13  | <0.0001  | 23.79  | 0.0001   | 105.04  | <0.0001  |
| Stim Domain X Context Control X Temp Control | 0.03   | 0.8559   | 0.24   | 0.6321   | <0.01   | 0.9729   |
## Supplemental Table 2 – Return Behavioral Data

|                      | Samp 1 |         | Samp 2 |         | Combine |         |
|----------------------|--------|---------|--------|---------|---------|---------|
|                      | RT     | ACC     | RT     | ACC     | RT      | ACC     |
| **Effect**           | p-value| p-value | p-value| p-value | p-value | p-value |
| ME Stim Domain       | 0.12   | 0.7369  | 3.01   | 0.0961  | 3.80    | 0.063   | 0.01    | 0.9057  | 2.32    | 0.1345  | 1.62    | 0.2091  |
| ME Context Control   | 85.74  | <0.0001 | 20.00  | 0.0002  | 145.46  | <0.0001 | 38.88  | <0.0001 | 216.36  | <0.0001 | 57.08  | <0.0001 |
| ME Temp Control      | 3.79   | 0.064   | 11.44  | 0.0026  | 3.09    | 0.0913  | 3.79   | 0.0634  | 6.9     | 0.0116  | 14.43  | 0.0004  |
| Stim Domain X Context Control | 0.74 | 0.3986 | 3.95   | 0.0588  | 0.35    | 0.5576  | 0.49   | 0.4907  | 0.96    | 0.3329  | 2.79   | 0.1012  |
| Stim Domain X Temp Control | 1.02 | 0.3223 | 0.12   | 0.7326  | 0.39    | 0.5357  | 1.18   | 0.2352  | 0.01    | 0.937   | 0.64   | 0.4269  |
| Context Control x Temp Control | 8.66 | 0.0073 | 19.19  | 0.0002  | 6.77    | 0.0156  | 1.66   | 0.2103  | 15.45   | 0.0003  | 10.22  | 0.0025  |
| Stim Domain X Context Control x Temp Control | 1.48 | 0.2364 | 1.99   | 0.172   | 0.39    | 0.5384  | 0.01   | 0.9306  | 1.68    | 0.2015  | 0.32   | 0.5761  |
Supplemental References

1. Nee, D. E. & D'Esposito, M. The hierarchical organization of the lateral prefrontal cortex. *eLife* 5, (2016).

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