Neural resources shift under Methylphenidate: A computational approach to examine anxiety-cognition interplay

Manish Saggar\textsuperscript{a,1,*}, Jennifer Bruno\textsuperscript{a,1}, Claudie Gaillard\textsuperscript{b}, Leonardo Claudino\textsuperscript{b}, Monique Ernst\textsuperscript{b,**}

\textsuperscript{a}Department of Psychiatry and Behavioral Sciences, Stanford University, Stanford, CA, USA

\textsuperscript{b}Section on Neurobiology of Fear and Anxiety, National Institute of Mental Health, Bethesda, MD, USA

Abstract

The reciprocal interplay between anxiety and cognition is well documented. Anxiety negatively impacts cognition, while cognitive engagement can down-regulate anxiety. The brain mechanisms and dynamics underlying such interplay are not fully understood. To study this question, we experimentally and orthogonally manipulated anxiety (using a threat of shock paradigm) and cognition (using methylphenidate; MPH). The effects of these manipulations on the brain and behavior were evaluated in 50 healthy participants (25 MPH, 25 placebo), using an n-back working memory fMRI task (with low and high load conditions). Behaviorally, improved response accuracy was observed as a main effect of the drug across all conditions. We employed two approaches to understand the neural mechanisms underlying MPH-based cognitive enhancement in safe and threat conditions. First, we performed a hypothesis-driven computational analysis using a mathematical framework to examine how MPH putatively affects cognitive enhancement in the face of induced anxiety across two levels of cognitive load. Second, we performed an exploratory data analysis using Topological Data Analysis (TDA)-based Mapper to examine changes in spatiotemporal brain activity across the entire cortex. Both approaches provided converging evidence that MPH facilitated greater differential engagement of neural resources (brain activity) across low and high working memory load conditions. Furthermore, load-based differential management of neural resources reflects enhanced efficiency that is most powerful during higher load and induced anxiety conditions. Overall, our results provide novel insights regarding brain mechanisms that facilitate cognitive enhancement under MPH and, in future research, may be used to help mitigate anxiety-related cognitive underperformance.
1. Introduction

Anxiety disorders are highly prevalent in the United States: 19.1% of adults reported having an anxiety disorder in the past year (Website, 2007), and the lifetime prevalence estimate is 31% (Website, 2007). In addition to the severe emotional burden, anxiety interferes with cognition (Moran, 2016) and is associated with cognitive deficits (Airaksinen et al., 2005; Robinson et al., 2013; Eysenck and Derakshan, 2011), further reducing life quality. Paradoxically, performing a task with a high cognitive load can reduce anxiety (Balderston et al., 2016; Clarke and Johnstone, 2013; Patel et al., 2016; Vytal et al., 2012; Vytal et al., 2013; Vytal et al., 2016). This latter effect is of particular interest as a potential strategy to optimize anxiety disorder treatment. Studies have sought to understand the impact of pharmacological cognitive enhancement (via Methylphenidate, MPH) on the anxiety-cognition interplay, but the results have been mixed (Kritchman et al., 2019; Sánchez-Pérez et al., 2012). Here, we use a computational framework that includes both hypothesis-driven and exploratory approaches to advance our understanding of the mechanisms that underlie cognitive enhancement and its impact on state anxiety. It is important to point out that our results in state anxiety may not generalize to persons with pathological anxiety. Nonetheless, from a clinical neuroscience perspective, this work represents an essential “proof of concept” step toward clarifying neural mechanisms and laying the initial groundwork for harnessing this phenomenon for clinical interventions.

Previous theoretical and empirical work has laid a foundation for understanding the interplay between anxiety and cognitive enhancement. First, anxiety (both state and trait) can reduce the i) efficiency of the central executive network; ii) ability to inhibit responses; iii) ability in cognitive switching; and iv) processing efficiency (Eysenck and Derakshan, 2011; Eysenck and Calvo, 1992; Eysenck et al., 2007; Bishop, 2007; Ishai et al., 2004). Here, processing efficiency in the context of working memory is defined by the relationship between accuracy and the extent of neural resources mobilized (Eysenck and Derakshan, 2011). Thus, high efficiency would indicate higher accuracy while using fewer resources. Second, work in rats has demonstrated that MPH enhances neuronal activity and reduces latency to correct response, showing greater efficiency when performing a visual signal detection task (Navarra et al., 2017). Finally, neuroimaging during a working memory task has demonstrated that MPH increased activation within the frontoparietal network (FPN) while reducing deactivation within the default mode network (DMN) (Gaillard et al., 2021). This shift in resources was found only during induced anxiety and thus may reflect optimization of the balance between core networks: FPN for regulating cognition (Zanto and Gazzaley, 2013) and DMN for regulating emotion (Raichle, 2015; Broyd et al., 2009).

While the findings mentioned above present important information regarding how complementary brain networks regulate cognition and anxiety, two critical gaps remain. First, we need to better understand how cognitive load impacts the interplay between induced anxiety and enhanced cognition. Second, there is a lack of understanding about how the whole-brain activity patterns adapt across cognitive load conditions in the presence of induced anxiety or enhanced cognition. The present study gathered data from a randomized trial designed to examine neural mechanisms in response to MPH-related cognitive enhancement and threat-of-shock induced anxiety (Gaillard et al., 2021). Anxiety
and cognitive load were experimentally and orthogonally manipulated using threat-of-shock vs. safety and a verbal n-back working memory task at low- and high-load, providing a three-factor design, with the factors being drug (MPH vs. placebo), cognitive load (1-back vs. 3-back), and anxiety (safe vs. induced anxiety) (Fig. 1).

We employed complementary hypothesis-driven and exploratory analytical approaches to further understand how boosting cognition via MPH impacts the anxiety/cognitive interplay. These approaches differ fundamentally from a previous study that used a generalized linear model (GLM) to examine changes in brain activity associated with the three-factor design, using the same dataset as here (Gaillard et al., 2021). The hypothesis-driven and exploratory approaches we employed here have unique advantages to the previous GLM-based work. Here, in the first hypothesis-driven computational analysis, we explicitly constructed a mathematical framework to examine how MPH putatively affects cognitive enhancement in the face of induced anxiety across two levels of cognitive load. Hypothesis-driven models typically encapsulate a theoretical (usually mechanistic) understanding of the underlying phenomena and use a comparatively small number of parameters to represent theoretically meaningful constructs (Huys et al., 2016). Once fitted to individual data (first-level individual analysis), these parameters can be compared across groups (Huys et al., 2013). The hypothesis-driven models are beneficial for measuring hidden variables and their interactions (Huys et al., 2013), which are otherwise difficult to measure directly.

In our hypothesis-driven modeling approach, we defined explicit parameters for cognitive load and anxiety (Fig. 2) to further understand the group differences (MPH vs. placebo) in core network engagement. Additionally, when compared to a more traditional and equally effective approach, such as a mixed-effects model, our computational approach has putatively greater interpretability and visualization. For example, the model parameters (load and anxiety) can be used to embed the data into lower, more interpretable dimensions.

Our prior whole-brain voxel-wise examination (Gaillard et al., 2021) indicated that MPH increased activation within the (FPN) while reducing deactivation within the DMN. These results were interpreted as an expansion of overall cognitive resources and were only found during the induced anxiety condition. To probe the anxiety/cognitive interaction further, our present, hypothesis-driven analysis was limited to the examination of the DMN and FPN. However, for completeness, we also report results from other networks.

Our second approach used exploratory data analysis to understand how MPH modulates cortical activity patterns at the whole-brain level across varying cognitive load and anxiety. Hypothesis-free (or exploratory) methods designed to detect patterns in whole-brain dynamics without an explicit hypothesis or search restriction have been shown to be essential for understanding the overall dynamic response of the brain to a given task (Gonzalez-Castillo et al., 2012). How the brain dynamically responds to changing cognitive demands is a critical question in clinical neuroscience because aberrant dynamics have been associated with several clinical conditions, including anxiety (Chen et al., 2020) and ADHD. To examine changes in brain activity patterns without collapsing data in space, time, or across participants at the outset, we used Topological Data Analysis (TDA) based Mapper approach (Saggar et al., 2018). Mapper attempts to reveal the underlying shape (or manifold) of high-dimensional data by embedding it into a low-dimensional space as a graph. At the same time, the information loss associated with dimensionality reduction is partly salvaged...
by performing a partial clustering step in the original high-dimensional space (Saggar et al., 2018; Geniesse et al., 2019, Singh et al., 2007). We have previously used Mapper to capture task-evoked transitions in whole-brain activity patterns at the acquired spatiotemporal resolution, as well as to reveal the rules that govern transitions in spontaneous brain activity at rest (Saggar et al., 2018; Saggar et al., 2022). We used whole-brain activity patterns to construct “shape” graphs that portray the manifold governing changes in activity patterns. The nodes in these graphs represent whole-brain configurations, while the edges represent similarity across configurations (Fig. 3). We subsequently annotated (or colored) the graphs based on task conditions and studied differences in topological properties across groups. Together, the hypothesis-driven and exploratory analyses hold the potential for revealing novel, converging evidence to further inform clinical neuroscience.

2. Methods

The present study uses data collected previously as part of the original GLM study (Gaillard et al., 2021). Here we only present brief details about data acquisition.

2.1. Participants

Seventy healthy volunteers were recruited from the Washington D.C. metropolitan area in our parallel-group design, a randomized, double-blind, placebo-controlled study. Participants were randomly assigned to receive either a single oral dose of 20 mg methylphenidate (MPH) or placebo (PLA), according to a randomization schedule established by the National Institutes of Health (NIH) pharmacy. The placebo capsule contained an inert substance. MPH and placebo capsules were identical in color (white) and shape (oblong). They were prepared by the NIH pharmacy. All subjects provided written informed consent approved by the National Institute of Mental Health (NIMH) Combined Neuroscience Institutional Review Board (CNS IRB). Briefly, participants were aged between 18 and 50 years, with no current psychiatric disorders or past significant psychiatric conditions (assessed by psychiatric interview using the Structured Clinical Interview for DSM-IV (SCID) (First, 2002), no medical conditions (assessed by clinical interview and physical exam), and no contraindications to magnetic resonance imaging (MRI). Additionally, participants were excluded if they had prior treatment with stimulants, intelligence quotient (IQ) lower than 80 as assessed via the Wechsler Abbreviated Scale of Intelligence (WASI (Wechsler, 2012)), pregnancy or a positive pregnancy test, current or past alcohol/drug dependence, alcohol/drug abuse in the past year, or positive toxicology urine screen. Twenty participants were excluded from analyses for the following reasons: missing task-based fMRI sequences (n = 4), incomplete behavioral data (n = 6), and excessive head motion (n = 10). Therefore, the final sample consisted of 50 healthy, right-handed adults (26 F; age mean = 28.2 years, SD = 6.9 years).

2.2. Drug administration

Participants received a single oral dose of PLA or immediate-release MPH 20 mg (Ritalin, Novartis, Basel, Switzerland), both presented in identical-appearing capsules. The MPH dose was based on the lowest dose reported to be effective on cognitive function (Mehta et al., 2000; Moeller et al., 2014; Pauls et al., 2012). To maximize MPH plasma levels during
cognitive testing, the drug was administered approximately 90 minutes before the beginning of the experimental working memory task in the scanner (Fig. 1) (Mehta et al., 2000; Pauls et al., 2012; Nandam et al., 2014). Potential side effects and adverse reactions were monitored by a clinician using a 34-item inventory to assess physical and mental symptoms (e.g., stomachache, nausea, lightheadedness, rash, drowsiness, headache). This assessment was performed after the study visit.

2.3. Experimental design

As detailed previously (Gaillard et al., 2021), the cognitive task consisted of the widely-used verbal N-back working memory (WM) paradigm (Balderston et al., 2017; Braver et al., 1997; Ernst et al., 2016). Participants viewed letters displayed sequentially on the screen. For each presented letter, participants were instructed to indicate (via button press) if each presented letter was identical to (matched) or different from (did not match) the letter presented N letters before. Thirty-three percent of trials were ‘match’ trials in which the presented letter was identical to the letter presented N letters before. We included two levels of difficulty: 1-back and 3-back, using a block design. At the beginning of each block, participants viewed instructions (8s) regarding the upcoming difficulty level (1-back or 3-back). The complete task included two runs, each with eight blocks of 18 letters. Each letter was presented for 0.5 s at 2 s intervals. The order of the runs was counter-balanced across participants. Within each run, we also manipulated the level of anxiety by including safe blocks and blocks that included the threat of unpredictable electrical shocks (threat blocks). Each safe and threat block was paired with a level of difficulty (1-back or 3-back). Thus, each run contained two blocks per condition (i.e., 1-back/safe, 3-back/safe, 1-back/threat, 3-back/threat). A color surrounded each letter; either blue to indicate a safe block or orange to indicate a threat block. Participants were explicitly told that they would never receive an electrical shock during the safe blocks (blue) but that they could receive unpredictable electrical shocks at any time during the threat blocks (orange). Three shocks were delivered per run for a total of six shocks throughout the task.

After each run of the n-back task, we collected subjective anxiety ratings on a 10-point Likert scale ranging from 1 (“not at all”) to 10 (“extremely”). Before the working memory task, the shock level intensity was titrated so that each participant experienced it as “highly uncomfortable and aversive, but not painful” according to a 10-point Likert scale ranging from 1 (“not at all aversive, maybe you barely felt it”) to 10 (“being highly uncomfortable, but tolerable”). Shock discomfort was retrospectively assessed following each run of the N-back task via a rating of “how unpleasant were the electric shocks” on an 11-point Likert scale ranging from 0 (not at all) to 10 (extremely).

2.4. Data acquisition

Two runs of 225 multi-echo EPI images were collected using a 3T Siemens MAGNETOM Skyra (Erlangen, Germany) fMRI system and a 32-channel head coil. Thirty-two interleaved 3mm slices (matrix = 64 mm × 64 mm) were collected parallel to the AC-PC line with an anterior-to-posterior phase encoding direction (TR = 2000 ms; TEs = 12 ms, 24.48 ms, 36.96 ms; flip angle = 70°). Prior to the first functional task-based run, we acquired two additional sets of 10 multi-echo EPI images using the same parameters, with
one “forward” series using the same phase-encoding gradient (anterior-to-posterior phase encoding direction) and the second “reverse” series using a reverse phase-encoding gradient with opposite polarity (posterior-to-anterior phase encoding direction). These additional series were used to correct for EPI spatial distortion related to phase-encoding direction. Additionally, a multi-echo T1-weighted Magnetization-Prepared Rapid Gradient-Echo (MPRAGE) image (TR = 2530 ms; TEs = 1.69 ms, 3.55 ms, 5.41 ms, 7.27 ms; flip angle = 7°) was acquired. T1-weighted MPRAGE images consisted of interleaved 1 mm axial slices (matrix = 256 mm × 256 mm), which were later co-registered to the combined EPI images.

2.5. Data preprocessing

Results included in this manuscript come from a preprocessing step performed using fMRIPrep 1.5.9 (Esteban et al., 2019; Esteban et al., 2018) RRID:SCR_016216), which is based on Nipype 1.4.2 (Gorgolewski et al., 2018; Gorgolewski et al., 2011); RRID:SCR_002502).

Anatomical data preprocessing—The T1-weighted (T1w) image was corrected for intensity non-uniformity (INU) with N4BiasFieldCorrection (Tustison et al., 2010), distributed with ANTs 2.2.0RRID:SCR_004757) (Avants et al., 2008), and used as T1w-reference throughout the workflow. The T1w-reference was then skull-stripped with a Nipype implementation of the ants BrainExtraction.sh workflow, using OASIS30ANTs as the target template. Brain tissue segmentation of cerebrospinal fluid (CSF), white matter (WM), and gray matter (GM) were performed on the brain-extracted T1w using fast (FSL 5.0.9, RRID:SCR_002823) (Zhang et al., 2001). Volume-based spatial normalization to two standard spaces (MNI152NLin6Asym, MNI152NLin2009cAsym) was performed through nonlinear registration with antsRegistration (ANTs 2.2.0), using brain-extracted versions of both T1w reference and the T1w template.

Functional data preprocessing—The following preprocessing was performed for each of the 2 BOLD runs per subject. First, a reference volume and its skull-stripped version were generated using a custom methodology of fMRIPrep. The BOLD reference was then co-registered to the T1w reference using flirt (FSL 5.0.9) (Jenkinson and Smith, 2001) with the boundary-based registration (Greve and Fischl, 2009) cost function. Co-registration was configured with nine degrees of freedom to account for distortions remaining in the BOLD reference. Head-motion parameters with respect to the BOLD reference (transformation matrices and six corresponding rotation and translation parameters) are estimated before any spatiotemporal filtering using mcflirt (FSL 5.0.9) (Jenkinson et al., 2002). The BOLD time series were resampled onto their original native space by applying the transforms to correct for head motion. First, a reference volume and its skull-stripped version were generated using a custom methodology of fMRIPrep. Confounding time series were calculated based on the preprocessed BOLD: framewise displacement (FD) and DVARS. FD and DVARS are calculated for each functional run, both using their implementations in Nipype (Power et al., 2014). To further reduce the effect of head movement, framewise displacement (FD) was used to create a temporal mask to remove motion-contaminated frames. We used a threshold of FD = 0.2mm to flag frames as motion contaminated. For each such motion-contaminated frame, we also flagged a back and two forward frames as motion
contaminated. Following the construction of the temporal mask for censuring, the data were processed with the following steps: (i) demeaning and detrending, (ii) multiple regression using six motion parameters, while temporally masked data were ignored during beta estimation, (iii) interpolation across temporally masked frames using linear estimation of the values at censored frames so that continuous data can be passed through (iv) a band-pass filter (0.009 Hz < f < 0.08 Hz). The temporally masked (or censored) frames were removed for further analysis.

The cortical data was then parcellated into 400 regions using the Schaefer parcellation, which implements a gradient-weighted Markov Random Field (gwMRF) model that integrates both local gradient and global similarity to produce higher functional homogeneity (Schaefer et al., 2018).

2.6. Hypothesis-based approach: construction of the mathematical framework

We created an explicit mathematical framework to perform a hypothesis-driven examination of the interplay between drug-induced cognitive enhancement and threat-induced anxiety. We operationalized the framework using two parameters: alpha ($\alpha$) and beta ($\beta$), where the $\alpha$ parameter accounted for the load-related changes in activation (i.e., 3-back > 1-back) and the $\beta$ parameter accounted for the anxiety-related changes (i.e., threat > safe).

Both parameters were estimated using the BOLD signal strength, $SS_{3-back, 1-back}$, for respective networks $Net_j$ and participant $S_i$, as follows,

$$\alpha_{S_i}^{Net_j} = \frac{SS_{Safe}^{3-back} - SS_{1-back}^{Safe}}{2} + \frac{SS_{Threat}^{3-back} - SS_{1-back}^{Threat}}{2}$$

$$\beta_{S_i}^{Net_j} = \frac{SS_{Threat}^{3-back} - SS_{3-back}^{Safe}}{2} + \frac{SS_{Threat}^{1-back} - SS_{1-back}^{Safe}}{2}$$

Using these two parameters, for each brain network, we modeled three scenarios: (A) network is only affected by changes in load and not anxiety (i.e., $\alpha > 0$, $\beta = 0$); (B) network is only affected by changes in anxiety and not load (i.e., $\alpha = 0$, $\beta > 0$); and (C) network is affected by both load and anxiety (i.e., $\alpha > 0$, $\beta > 0$). We ignored the null condition, where the network is not affected by either load or anxiety (i.e., $\alpha = 0$, $\beta = 0$). See Fig. 2 for a cartoon depicting the parameters and modeled scenarios. Based on previous work, this analysis was limited to the default mode network (DMN) and frontoparietal control network (FPN) (Gaillard et al., 2021). The network definitions were based on the 7-network Yeo parcellation (Yeo et al., 2011), which was applied to the data that had been previously parcellated according to the Schaefer parcellation (Schaefer et al., 2018).

2.7. Hypothesis-free approach: TDA-based Mapper pipeline

The TDA (Tenenbaum et al., 2000) based Mapper pipeline was run on each participant. Complete details of the Mapper analysis pipeline are presented elsewhere (Saggar et al., 2018, Geniesse et al., 2019; Saggar et al., 2022). The Mapper pipeline consists of four
main steps. First, Mapper embeds the high-dimensional input data into a lower dimension \( d \), using a filter function \( f \). For ease of visualization, we chose \( d = 2 \). The choice of filter function dictates which data properties are to be preserved in the lower dimensional space. Several studies using animal models and computational research suggest that inter-regional interactions in the brain are multivariate and nonlinear. Thus, we used a non-linear filter function based on neighborhood embedding (Saggar et al., 2018). We implemented the filter function \( f \) as a nonlinear dimensionality reduction step, where the geodesic distances were computed in the original high-dimensional space and were later embedded into a low \( d \)-dimensional space using classical multidimensional scaling (MDS). Using geodesic distances, instead of Euclidean (or other similar metrics), allow for better preservation of the local structure evident in the original high-dimensional space after projection into a lower dimensional space. Similar functions have been used previously in manifold learning (Tenenbaum et al., 2000). Recently, we showed the efficacy of neighborhood embedding (with \( k \) neighbors) in capturing the landscape of whole-brain configurations extracted from a continuous multitask paradigm and task-evoked data from the Human Connectome Project (HCP). A similar filter function is used in our more recent Mapper application to resting state fMRI data as well (Saggar et al., 2022).

The second step of Mapper creates overlapping bins in \( n \)-dimensional space to allow for compression, thereby reducing the effect of noisy data points. Based on previous work using fMRI data (Saggar et al., 2018), we divided the lower-dimensional space into overlapping bins using a resolution parameter \( (r; \# \text{bins}) \) of 18. The percent overlap between bins \( (g) \) was kept at 70%. Mapper-generated graphs have been previously shown to be stable for a large variation across parameters for resolution and percent overlap (Saggar et al., 2018; Saggar et al., 2022). Here, we varied \( r \) and \( g \) across a range of values to examine the robustness of the results (see Section 2.8).

The third step of Mapper includes partial clustering within each bin, where the original high-dimensional information is used for coalescing (or separating) data points into nodes in the low-dimensional space. Partial clustering allows recovering the loss of information due to dimensional reduction in step one (Geniesse et al., 2019; Lum et al., 2013). This partial clustering step can (and often does) create multiple nodes within a bin.

Lastly, to generate a graphical representation of the “shape” of input data, nodes from different bins are connected if any data points lie in the overlap between bins. See Fig. 3 below for a pictorial representation of the Mapper pipeline.

### 2.8. Network measures

The Mapper-generated graphs can be annotated (or colored) using meta-information not used to construct the graphs. Some examples of meta-measures are condition labels and task performance. Here, we annotated these graphs using task condition labels to assess whether whole-brain activation patterns are similar or different across task conditions. To quantify the extent of divergence (high condition specificity) or overlap (low condition specificity) across task conditions, we used a graph theoretical measurement of the participation coefficient (Guimerà and Nunes Amaral, 2005). The participation coefficient of a node is defined as:

\[
\text{Participation Coefficient}_{\text{node}} = \frac{\sum_{\text{bins}} \left| \text{Node in bin} \right|}{\text{Total Number of Nodes} \times \text{Total Number of Bins}}
\]
\[ P_i = 1 - \sum_{s=1}^{N_M} \left( \frac{K_{is}}{k_i} \right)^2 \]

Where \( K_{is} \) is the number of links of node \( i \) to nodes in community \( s \), \( k_i \) is the total degree of node \( i \) and \( N_M \) is the total number of communities. The \( P_i \) of a node \( i \) is close to 1 if its links are uniformly distributed among all communities of the graph (and hence overlapping), and it is close to 0 if its links are primarily within its own community (and therefore diverging).

### 2.9. Effect of Mapper parameters

A parameter perturbation analysis was run to examine whether the results were stable across different parameter choices. Two Mapper parameters, number bins \( r \) and %overlap (or gain \( g \)), were varied across a range to show that the observed results using \( r = 18 \) and \( g = 70 \) were robust across other parameter values (Fig. S1).

### 3. Results

#### 3.1. Behavioral results

Using a linear mixed-effects model, we observed a significant effect of MPH on working memory performance while controlling for age and sex, such that participants in the MPH group outperformed those in the placebo group in terms of accuracy \((\beta = 0.05, t(323) = 1.87, p_{\text{one-tailed}} = 0.03; \) we hypothesized that participants in the MPH group would show enhanced performance relative to the PLA thus 1-tailed significance was tested). We used a similar mixed effects model to examine anxiety ratings and observed a significant main effect of the condition (higher anxiety for threat vs. safety \((b = 3.64, t(350) = 8.85, p < 0.001), \) thereby validating our induced anxiety condition. Mean shock ratings for each run and group (MPH vs. placebo) were above 7.88 (out of 10) and did not differ significantly between groups for either run one or run two \((p's > 0.10). \) A more detailed analysis of behavioral results was previously presented \((Gaillard et al., 2021).\)

#### 3.2. Hypothesis-driven examination using explicit parameters for load- and anxiety-related changes in activation

Based on our previous work \((Gaillard et al., 2021), \) we limited the present hypothesis-driven examination of load and anxiety parameters to the default mode (DMN) and frontoparietal (FPN) networks. After estimating load (alpha) and anxiety (beta) parameters for each network, a repeated-measures ANOVA with estimated parameters and networks as within-subject factors and the group as a between-subject factor was run while covarying for age and sex. A significant group x network x parameter interaction was found \((F(1,46) = 5.335, p = 0.025). \) Post-hoc pairwise comparisons (adjusted for multiple comparisons using Bonferroni correction) revealed significantly higher values of the load parameter in the MPH group (as compared to PLA) for both networks \((p = 0.043 \text{ for DMN and } p = 0.009 \text{ for FPN).} \) No significant group differences were evident for the anxiety parameter for either network. These results fit with model A (Fig. 2). Fig. 4 shows these results and visualizes each network’s average BOLD signal strength across the two groups and four task blocks.
For completeness and to inform future studies, group differences in the load and anxiety parameters of the other five networks (from the Yeo network parcellation (Yeo et al., 2011)) were also examined. We used univariate analysis of variance to compare group differences while controlling for age and sex. All five networks showed significant group differences for the load (alpha) parameter (Visual network: $F(1,46) = 4.42, p = 0.041$; Somatomotor network: $F(1,46) = 6.79, p = 0.012$; Dorsal Attention network: $F(1,46) = 5.07, p = 0.029$; Ventral attention network: $F(1,46) = 7.68, p = 0.008$; Limbic network: $F(1,46) = 6.19, p = 0.017$), while no group differences were observed for the anxiety (beta) parameter. Thus, suggesting a global shift in activation for higher load conditions was facilitated by MPH.

3.3. Hypothesis-free examination of induced spatiotemporal changes across the cortex

We used a TDA-based Mapper approach to assess spatiotemporal changes in the entire cortex under different task conditions for the hypothesis-free examination. Mapper graphs were separately generated for each individual using their total task scan (i.e., data combined across both runs). Mapper-generated graphs were later colored (annotated) by load (3-back vs 1-back) and anxiety (safety vs threat) information. See Fig. 5 for Mapper-generated graphs of representative individuals from both groups. Supplementary Figures S2–S5 provide Mapper-generated graphs for all individuals, annotated by load and anxiety separately.

To estimate the amount of similarity (or divergence) between different degrees of load and anxiety, Mapper-generated graphs colored by load and anxiety were analyzed using the graph theoretical metric of Participation Coefficient (PC) (Guimerà and Nunes Amaral, 2005). PC values were then estimated for each graph node. Nodes with higher values of PC indicate higher similarity between different degrees of load (or anxiety), and lower values indicate higher divergence between different degrees of load (or anxiety). A repeated-measures ANOVA with a within-subject factor of annotation (load vs. anxiety) and between-subject factor of group was run, with age and sex as covariates. A significant group × annotation interaction was found $F(1,46) = 10.154, p = 0.003)$. Post-hoc pairwise comparisons showed the effect of load to be significantly different across the two groups ($p = 0.018$; Bonferroni corrected), such that significantly lower values of PC were observed for the load-based annotation in the MPH group (as compared with the PLA group). These results suggest higher divergence (or lower similarity) between spatial activity profiles based on the load in the MPH group. No such group differences were found for the anxiety-based annotation of the Mapper graphs. The parameter perturbation analysis showed similar results across different Mapper parameter choices (Fig. S1).

Because the participation coefficient implicitly assumes that communities are equally sized, we examined differences in community size and intramodular connectivity across communities and confirmed that there are no differences between conditions for load and anxiety ($ps > 0.05$; Fig. S6). Hence, our use of non-normalized PC estimation is justified.

Next, we examined whether individual differences in load-related divergence (i.e., lower PC value) were associated with behavioral performance on the working memory task using Spearman’s rank correlation while controlling for age and sex. Data were combined across groups to optimize statistical power. Correlation results indicated that the load-related
divergence of Mapper graphs was not only associated with better behavioral performance during higher load (rho (46) = −.31, p = .033) but also during induced anxiety (threat; rho (46) = −.32, p = .025) conditions (Fig. 5D).

4. Discussion

Using two complementary analytical approaches, we advance our understanding of how cognitive enhancement (under MPH) alters brain activity patterns in the face of induced anxiety and increased cognitive load. In the first approach, we develop an explicit mathematical framework to parametrically investigate the role of two core networks (DMN and FPN) implicated in the anxiety-cognition interplay. In the second approach, we use a hypothesis-free TDA-based analysis to examine the whole-brain dynamical response to our paradigm. Both approaches yield converging evidence that cognitive enhancement under MPH facilitates greater differential engagement of neural resources (activation) across conditions of low and high working memory load. This load-based differential management of neural resources facilitated better task performance during both higher load and higher anxiety conditions. Overall, our results provide novel insight into how cognitive enhancement under MPH putatively diminishes anxiety. Such information represents a critical preliminary step toward informing neurobiologically targeted treatment approaches employing cognitive enhancement to reduce anxiety and, in turn, reduce anxiety-related cognitive deficits.

Our findings include critical neurobiological and behavioral processes underlying the complex, reciprocal relationship between anxiety and cognition. Previous neuroimaging work has indicated that both DMN, which subserves self-referential and emotional processes (Raichle, 2015; Broyd et al., 2009), and FPN, which subserves executive processes (Zanto and Gazzaley, 2013), demonstrate aberrant connectivity in association with anxiety disorders (Xiong et al., 2020; Coutinho et al., 2016; Xu et al., 2019) and with high trait anxiety (Sylvester et al., 2012). Furthermore, emotion regulation is supported by the greater efficiency of both FPN and DMN (Pan et al., 2018). Recent neuroimaging work, using the same dataset as here, has shown that cognitive enhancement (via MPH) was associated with increased engagement of the FPN as well as reduced deactivation of the DMN during high anxiety and high WM load conditions (Gaillard et al., 2021). This recent finding suggests that expansion of cognitive resources under MPH provides an optimal balance between recruitment of cognitive processing and emotion regulation resources. Using different analytical approaches, we provide complementary evidence suggesting the recruitment of both core networks (DMN and FPN) under MPH. For completeness, we also examined other five networks from the Yeo parcellation and observed a similar response of MPH across networks. Thus, our present findings indicate that a global shift in activation during difficult (higher cognitive load) conditions was facilitated by the MPH.

Additionally, we further specify the nature of the network modulation under MPH according to task difficulty. Specifically, we found that the parameter for the load (model A; Fig 2) was significantly higher in the MPH group relative to the PLA group for all networks. Thus, our findings indicate that cognitive enhancement (via MPH) results in a differential engagement in response to a higher working memory load globally. Interestingly, no similar
group differences were observed for the anxiety parameter, i.e., cognitive enhancement (via MPH) did not result in differential engagement in response to higher anxiety, suggesting a lack of direct interaction between MPH and anxiety processing in our cohort.

Our hypothesis-free TDA results provide converging evidence for differential neural engagement in response to higher working memory load under MPH and extend these findings in two important ways. First, the TDA-based Mapper approach is a whole-brain approach and thus provides evidence of whole-brain dynamical response under MPH in addition to the aforementioned network-specific results. Second, our Mapper approach reveals how brain activity patterns differ (i.e., segregate) or collapse (i.e., integrate) at the level of individual timeframes (TRs). We used the participation coefficient (Guimerà and Nunes Amaral, 2005) (PC), an established graph-theoretical metric, to quantify the degree of segregation (vs. integration) in brain activity patterns across task factors. Using WM load as the task factor revealed lower PC (i.e., higher segregation across WM load) for the MPH group relative to the PLA group. This suggests that cognitive enhancement under MPH was facilitated by differential engagement of neural resources under low and high-load WM conditions. Thus, in line with our network-based approach, the TDA results also suggest load-based differential engagement at the whole-brain level. No such group differences were found for the PC when the Mapper-generated graphs were annotated (colored) by anxiety. Furthermore, WM load-based differential engagement was associated with better behavioral performance during higher load and anxiety conditions. This suggests that by modulating the level of neural engagement, MPH putatively facilitates higher cognitive efficiency during challenging conditions based on required cognitive demand.

Overall, competition for cognitive resources can explain the interactions between anxiety and cognition (King and Schaefer, 2011). When demands for one process increase, the resources available for other processes decrease. The present study was built upon foundational work demonstrating that enhancing cognition with exercise (Lago et al., 2019) or with MPH (Ernst et al., 2016) results in increased cognitive capacity facilitating enhanced cognitive and threat processing. Furthermore, previous work suggested that improving working memory performance through training (practice on high and low load working memory conditions) led to increased anxiety which, as the authors argued, may be due to the availability of increased resources to process threats (Grillon et al., 2020). Here, we extend previous work by providing converging evidence that cognitive enhancement associated with MPH is likely facilitated by load-appropriate (and hence efficient) engagement of neural resources. According to the neural efficiency hypothesis, individuals with higher intelligence demonstrate more efficient (i.e. lower) brain activation while performing cognitive tasks (Haier et al., 1988). Further, fMRI brain activation levels also adapt to the particular task demands (Dunst et al., 2014). Thus, one plausible interpretation of our present results is that MPH enhances the neural efficiency phenomenon by facilitating increased activation during increased task difficulty (high WM load). A potential mechanism by which MPH may enhance neural efficiency is via MPH’s action to increase dopaminergic neurotransmission which increases the signal-to-noise ratio in neural networks thereby reducing neural noise (Pertermann et al., 2019). However, it is also important to point out threat evidence for the neural efficiency hypothesis is not found in all studies and important factors moderate the relationship between brain activation and intelligence (e.g. sex, task type, task complexity,
brain area investigated and level of training, see (Neubauer and Fink, 2009) for a review of these issues). Therefore, our aforementioned interpretation should be considered with caution as assessment of these important moderators and their interactions with our variables of interest was outside the scope of the present study.

In the future, similar TDA-based analytical approaches can be measured and tracked over time and, as such, may represent a useful metric for informing and tracking response to interventions. Effective treatment of cognitive interference associated with anxiety disorders is an area of great clinical need. Our study was focused on state anxiety in healthy individuals and thus cannot be generalized to patients with anxiety disorders. The 20mg dose of MPH chosen for this study, while based on previous research (Ernst et al., 2016), was relatively low, yet it resulted in significant neural divergence. This speaks to the sensitivity of our hypothesis-driven and hypothesis-free approaches. However, examining the effects of higher dose MPH will be informative for understanding the whole dynamics of the anxiety/cognition interplay.

In summary, we provide novel mechanistic evidence of load-appropriate engagement of neural resources under MPH. Such efficient load-based engagement was associated with improved behavioral performance in the WM task during high load and anxiety conditions. We hope these results can provide a novel avenue for using computational approaches in improving mechanistic understandings of pharmacological interventions.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

M.S. was supported by the NIH Director’s New Innovator Award (DP2; MH119735) and Stanford’s MCHRI Faculty Scholar Award. Financial support for this study was provided by the Intramural Research Program of the National Institutes of Mental Health, project no. ZIAMH002798 to M.E.

**Data and code availability statement**

The data used in this paper can be made available upon reasonable request to Dr. Monique Ernst and may require a formal data sharing agreement with the NIMH.

The code used in this study will be made available upon publication at this address: https://github.com/braindynamicslab/tda-metpro.

**References**

Website, 2007. Available: Harvard Medical School. National Comorbidity Survey (NCS).

Moran TP., 2016. Anxiety and working memory capacity: a meta-analysis and narrative review. Psychol. Bull. 142, 831–864. [PubMed: 26963369]

Airaksinen E, Larsson M, Forsell Y, 2005. Neuropsychological functions in anxiety disorders in population-based samples: evidence of episodic memory dysfunction. J. Psychiatr. Res. >207–214. doi:10.1016/j.jpsychires.2004.06.001.
Robinson OJ, Vytal K, Cornwell BR, Grillon C, 2013. The impact of anxiety upon cognition: perspectives from human threat of shock studies. Front. Hum. Neurosci. 7, 203. [PubMed: 23730279]

Eysenck MW, Derakshan N, 2011. New perspectives in attentional control theory. Personal. Individ. Differ 955–960. doi:10.1016/j.paid.2010.08.019.

Balderston NL, Quispe-Escudero D, Hale E, Davis A, O’Connell K, Ernst M, et al., 2016. Working memory maintenance is sufficient to reduce state anxiety. Psychophysiology 1660–1668. doi:10.1111/psyp.12726. [PubMed: 27434207]

Clarke R, Johnstone T, 2013. Prefrontal inhibition of threat processing reduces working memory interference. Front. Hum. Neurosci. 7, 228. [PubMed: 23750133]

Patel N, Vytal K, Pavletic N, Stoodley C, Pine DS, Grillon C, et al., 2016. Interaction of threat and verbal working memory in adolescents. Psychophysiology 518–526. doi:10.1111/psyp.12582. [PubMed: 26589772]

Vytal K, Cornwell B, Arkin N, Grillon C, 2012. Describing the interplay between anxiety and cognition: from impaired performance under low cognitive load to reduced anxiety under high load. Psychophysiology 49, 842–852. [PubMed: 22332819]

Vytal KE, Cornwell BR, Letkiewicz AM, Arkin NE, Grillon C, 2013. The complex interaction between anxiety and cognition: insight from spatial and verbal working memory. Front. Hum. Neurosci. 7, 93. [PubMed: 23542914]

Vytal KE, Arkin NE, Overstreet C, Lieberman L, Grillon C, 2016. Induced-anxiety differentially disrupts working memory in generalized anxiety disorder. BMC Psychiatry 16, 62. [PubMed: 26976146]

Kritchman M, Koubi M, Mimouni Bloch A, Bloch Y, 2019. Effect of methylphenidate on state anxiety in children with ADHD—a single dose, placebo controlled, crossover study. Front. Behav. Neurosci. 13, 106. [PubMed: 31156406]

Sánchez-Pérez AM, García-Avilés Á, Gascó HA, Sanjuán J, Olucha-Bordonau FE., 2012. Effects of methylphenidate on anxiety. Rev. Neurol. 55, 499–506. [PubMed: 23055432]

Eysenck MW, Calvo MG., 1992. Anxiety and performance: the processing efficiency theory. Cogn. Emot. 6, 409–434.

Eysenck MW, Derakshan N, Santos R, Calvo MG. Anxiety and cognitive performance: attentional control theory. Emotion. 2007. pp. 336–353. doi:10.1037/1528-3542.7.2.336. [PubMed: 17516812]

Bishop SJ., 2007. Neurocognitive mechanisms of anxiety: an integrative account. Trends Cogn. Sci. 11, 307–316. [PubMed: 17553730]

Ishai A, Pessoa L, Bikle PC, Ungerleider LG., 2004. Repetition suppression of faces is modulated by emotion. Proc. Natl. Acad. Sci. USA 101, 9827–9832. [PubMed: 15210952]

Navarra RL, Clark BD, Gargiulo AT, Waterhouse BD., 2017. Methylphenidate enhances early-stage sensory processing and rodent performance of a visual signal detection task. Neuropsychopharmacology 42, 1326–1337. [PubMed: 27910862]

Gaillard C, Lago TR, Gorka AX, Balderston NL, Fuchs BA, Reynolds RC, et al., 2021. Methylphenidate modulates interactions of anxiety with cognition. Transl. Psychiatry 11, 544. [PubMed: 34675189]

Zanto TP, Gazzaley A, 2013. Fronto-parietal network: flexible hub of cognitive control. Trends Cogn. Sci. 602–603. [PubMed: 24129332]

Raichle ME., 2015. The brain’s default mode network. Annu. Rev. Neurosci. 38, 433–447. [PubMed: 25938726]

Broyd SJ, Demanuele C, Debener S, Helps SK, James CJ, Sonuga-Barke EJS., 2009. Default-mode brain dysfunction in mental disorders: a systematic review. Neurosci. Biobehav. Rev. 33, 279–296. [PubMed: 18824195]

Huys QJM, Maia TV, Frank MJ, 2016. Computational psychiatry as a bridge from neuroscience to clinical applications. Nat. Neurosci. 19, 404–413. [PubMed: 26906507]

Huys QJ, Pizzagalli DA, Bogdan R, Dayan P, 2013. Mapping anhedonia onto reinforcement learning: a behavioural meta-analysis. Biol. Mood Anxiety Disord. 3, 12. [PubMed: 23782813]
Gonzalez-Castillo J, Saad ZS, Handwerker DA, Inati SJ, Brenowitz N, Bandettini PA., 2012. Whole-brain, time-locked activation with simple tasks revealed using massive averaging and model-free analysis. Proc. Natl. Acad. Sci. USA 109, 5487–5492. [PubMed: 22431587]

Chen Y, Cui Q, Xie A, Pang Y, Sheng W, Tang Q, et al., 2020. Abnormal dynamic functional connectivity density in patients with generalized anxiety disorder. J. Affect. Disord. 261, 49–57. [PubMed: 31600587]

Saggar M, Sporns O, Gonzalez-Castillo J, Bandettini PA, Carlsson G, Glover G, et al., 2018. Towards a new approach to reveal dynamical organization of the brain using topological data analysis. Nat. Commun. 9, 1–14. [PubMed: 29317637]

Geniesse C, Sporns O, Petri G, Saggar M, 2019. Generating dynamical neuroimaging spatiotemporal representations (DyNeuSR) using topological data analysis. Netw. Neurosci. 3, 763–778. [PubMed: 31410378]

Singh G, Mémoli F, Carlsson G, 2007. Topological methods for the analysis of high dimensional data sets and 3D object recognition. In: Botsch M, Pajarola R (Eds.), Eurographics Symposium on Point-Based Graphics.

Saggar M, Shine JM, Liégeois R, Dosenbach NUF, 2022. Fair D. Precision dynamical mapping using topological data analysis reveals a hub-like transition state at rest. Nat. Commun. 13, 4791. [PubMed: 35970984]

First MB., 2002. The DSM series and experience with DSM-IV. Psychopathology 67–71. doi:10.1159/000065121. [PubMed: 12145486]

Wechsler D, 2012. Wechsler abbreviated scale of intelligence. PsycTESTS Dataset doi:10.1037/t15170-000.

Mehta MA, Owen AM, Sahakian BJ, Mavaddat N, Pickard JD, Robbins TW., 2000. Methylphenidate enhances working memory by modulating discrete frontal and parietal lobe regions in the human brain. J. Neurosci. RC65. doi:10.1523/jneurosci.20-06-j0004.2000. –RC65.

Moeller SJ, Honorio J, Tomasi D, Parvaz MA, Woicik PA, Volkow ND, et al., 2014. Methylphenidate enhances executive function and optimizes prefrontal function in both health and cocaine addiction. Cereb. Cortex 643–653. doi:10.1093/cercor/bhs345. [PubMed: 23162047]

Pauls AM, O’Daly OG, Rubia K, Riedel WJ, Williams SCR, Mehta MA, 2012. Methylphenidate effects on prefrontal functioning during attentional-capture and response inhibition. Biol. Psychiatry 142–149. doi:10.1016/j.biopsych.2012.03.028. [PubMed: 22552046]

Nandam LS, Sanjay Nandam L, Hester R, Bellgrove MA, 2014. Dissociable and common effects of methylphenidate, atomoxetine and citalopram on response inhibition neural networks. Neuropsychologia 263–270. doi:10.1016/j.neuropsychologia.2014.01.023.

Balderston NL, Vytal KE, O’Connell K, Torrisi S, Letkiewicz A, Ernst M, et al., 2017. Anxiety patients show reduced working memory related dlPFC activation during safety and threat. Depress. Anxiety 25–36. doi:10.1002/da.22518. [PubMed: 27110997]

Braver TS, Cohen JD, Nystrom LE, Jonides J, Smith EE, Noll DC., 1997. A parametric study of prefrontal cortex involvement in human working memory. Neuroimage 49–62. doi:10.1006/nimg.1996.0247. [PubMed: 90375392]

Esteban O, Markiewicz CJ, Blair RW, Moodie CA, Ilkay Isik A, Erramuzpe A, et al., 2019. fMRIPrep: a robust preprocessing pipeline for functional MRI. Nat. Methods 111–116. doi:10.1038/s41592-018-0235-y. [PubMed: 30532080]

Esteban O, Blair R, Markiewicz CJ, Berleant SL, Moodie C, Ma F, et al., 2018. FMRIPrep. Software Zenodo doi:10:5281/zenodo.

Gorgolewski KJ, Esteban O, Markiewicz CJ, Ziegler E, Ellis DG, Notter MP, et al., 2018. Nipype. Software. Zenodo.

Gorgolewski K, Burns CD, Madison C, Clark D, Halchenko YQ, Waskom ML, et al., 2011. Nipype: a flexible, lightweight and extensible neuroimaging data processing framework in python. Front. Neuroinform doi:10.3389/fninf.2011.00013.
Tustison NJ, Avants BB, Cook PA, Zheng Y, Egan A, Yushkevich PA, et al., 2010. N4ITK: improved N3 bias correction. IEEE Trans. Med. Imaging 1310–1320. doi:10.1109/tmi.2010.2046908. [PubMed: 20378467]

Avants B, Epstein C, Grossman M, Gee J, 2008. Symmetric diffeomorphic image registration with cross-correlation: evaluating automated labeling of elderly and neurodegenerative brain. Med. Image Anal. 26–41. doi:10.1016/j.media.2007.06.004. [PubMed: 17659998]

Zhang Y, Brady M, Smith S, 2001. Segmentation of brain MR images through a hidden Markov random field model and the expectation-maximization algorithm. IEEE Trans. Med. Imaging 45–57. doi:10.1109/42.906424. [PubMed: 11516708]

Jenkinson M, Smith S, 2001. A global optimisation method for robust affine registration of brain images. Med. Image Anal. 143–156. doi:10.1016/s1361-8415(01)00036-6. [PubMed: 11293691]

Zhang Y, Brady M, Smith S, 2001. Segmentation of brain MR images through a hidden Markov random field model and the expectation-maximization algorithm. IEEE Trans. Med. Imaging 45–57. doi:10.1109/42.906424. [PubMed: 11293691]

Power JD, Mitra A, Laumann TO, Snyder AZ, Schlaggar BL, Petersen SE., 2014. Methods to detect, characterize, and remove motion artifact in resting state fMRI. Neuroimage 320–341. doi:10.1016/j.neuroimage.2013.08.048.

Schaefer A, Kong R, Gordon EM, Laumann TO, Zuo X-N, Holmes AJ, et al., 2018. Local-global parcellation of the human cerebral cortex from intrinsic functional connectivity MRI. Cerebral Cortex 3095–3114. doi:10.1093/cercor/bhx179. [PubMed: 28981612]

Yeo BT, Krienen FM, Sepulcre J, Sabuncu MR, Lashkari D, Hollinshead M, et al., 2011. The organization of the human cerebral cortex estimated by intrinsic functional connectivity. J. Neurophysiol. 106, 1125–1165. [PubMed: 21653723]

Tenenbaum JB, de Silva V, Langford JC., 2000. A global geometric framework for nonlinear dimensionality reduction. Science 2319–2323. doi:10.1126/science.290.5500.2319.

Lum PY, Singh G, Lehman A, Ishkanov T, Vejdemo-Johansson M, Alagappan M, et al., 2013. Extracting insights from the shape of complex data using topology. Sci. Rep. doi:10.1038/srep01236.

Guimerà R, Nunes Amaral LA, 2005. Functional cartography of complex metabolic networks. Nature 895–900. doi:10.1038/nature03288.

Xiong H, Guo R-J, Shi H-W, 2020. Altered default mode network and salience network functional connectivity in patients with generalized anxiety disorders: an ICA-based resting-state fMRI study. Evid. Based Complement. Alternat. Med. 2020, 4048916. [PubMed: 32855650]

Coutinho JF, Fernandes SV, Soares JM, Maia L, Gonçalves OF, Sampaio A, 2016. Default mode network dissociation in depressive and anxiety states. Brain Imaging Behav. 147–157. doi:10.1007/s11682-016-9377-7. [PubMed: 25804311]

Xu J, Van Dam NT, Feng C, Luo Y, Ai H, Gu R, et al., 2019. Anxious brain networks: a coordinate-based activation likelihood estimation meta-analysis of resting-state functional connectivity studies in anxiety. Neurosci. Biobehav. Rev. 21–30. doi:10.1016/j.neubiorev.2018.11.005.

Sylvester CM, Corbetta M, Raichle ME, Rodebaugh TL, Schlaggar BL, Sheline YI, et al., 2012. Functional network dysfunction in anxiety and anxiety disorders. Trends Neurosci. 35, 527–535. [PubMed: 22658924]

Pan J, Zhan L, Hu CL, Yang J, Wang C, Gu L, et al., 2018. Emotion regulation and complex brain networks: association between expressive suppression and efficiency in the fronto-parietal network and default-mode network. Front. Hum. Neurosci. 12, 1–12. [PubMed: 29387003]

King R, Schaefer A, 2011. The emotional startle effect is disrupted by a concurrent working memory task. Psychophysiology 48, 269–272. [PubMed: 20636296]

Lago TR, Hsiung A, Leitner BP, Duckworth CJ, Balderston NL, Chen KY, et al., 2019. Exercise modulates the interaction between cognition and anxiety in humans. Cogn. Emotion 863–870. doi:10.1080/02699931.2018.1500445.

Grillon C, Lago T, Stahl S, Beale A, Balderston N, Ernst M, 2020. Better cognitive efficiency is associated with increased experimental anxiety. Psychophysiology doi:10.1111/psyp.13559.
Haier RJ, Siegel BV, Nuechterlein KH, Hazlett E, Wu JC, Paek J, et al., 1988. Cortical glucose metabolic rate correlates of abstract reasoning and attention studied with positron emission tomography. Intelligence 12, 199–217.

Dunst B, Benedek M, Jauk E, Neubauer AC., 2014. Neural efficiency depending on subjective and objective level of task difficulty – an fMRI study. Personal. Individ. Differ S36–S37. doi:10.1016/j.paid.2013.07.087.

Pertermann M, Bluschke A, Roessner V, Beste C, 2019. The modulation of neural noise underlies the effectiveness of methylphenidate treatment in attention-deficit/hyperactivity disorder. Biol. Psychiatry Cogn. Neurosci. Neuroimaging 4, 743–750. [PubMed: 31103546]

Neubauer AC, Fink A, 2009. Intelligence and neural efficiency. Neurosci. Biobehav. Rev. 33, 1004–1023. [PubMed: 19580915]
Fig. 1.
Study timeline for each participant. Drug (Methylphenidate) was administered 90 minutes prior to the beginning of the working memory task in the MR scanner.
Mathematical formulation and scenarios modeled. We operationalized the framework using two parameters: alpha ($\alpha$) and beta ($\beta$), where the $\alpha$ parameter accounted for the load-related changes in activation (i.e., 3-back > 1-back) and the $\beta$ parameter accounted for the anxiety-related changes (i.e., threat > safe). Using these two parameters, we modeled three different scenarios for each brain network: load-driven, anxiety-driven, and both load and anxiety-driven.
Fig. 3.
Mapper pipeline is shown pictorially within the dashed border. In the first step, each participant’s high-dimensional neuroimaging data matrix for the entire scan session (time frames × cortical parcels) is embedded into a lower dimension set $d_2$ using a non-linear filter function $f$. In the second step, overlapping $d$-dimensional binning is performed to allow for compression and reduce the effects of noisy data points. In the third step, within each bin, partial clustering is performed such that data points that are closer to each other in the original high-dimensional space coalesce together in the low dimensional space. This partial clustering step allows for recovering information loss incurred due to initial dimensionality reduction. As a final step, to generate a graphical representation of the landscape, nodes from different bins are connected if any data points are shared. Once constructed, Mapper graphs can be annotated (colored) using meta-information, e.g., task condition, and their graph properties can be examined.
Fig. 4.
Results of the explicit parameters using default mode network (DMN) and frontoparietal network (FPN). [A] The parameter for the load (alpha) was observed to be significantly higher for the MPH group (than PLA group) for both frontoparietal and default mode networks. No significant group difference was observed for the anxiety parameter (beta). [B-C] Shows FPN and DMN networks activation for each of the four conditions across the two groups.
Fig. 5. Hypothesis-free examination of induced spatiotemporal changes across the cortex using the topological data analysis (TDA)-based Mapper approach. [A-B] Showing Mapper graphs for two representative participants (methylphenidate [MPH] on the left and placebo [PLA] on the right), annotated by load condition. In the case of the MPH group participant, higher separation between load conditions was observed. To better illustrate the separation between 3-back and 1-back conditions, we added background clusters (yellow and green) to the graph from the MPH participant. [C] The participation coefficient (PC) for load-based annotation was significantly different between the two groups. [D] Relating individual differences in participation coefficient of Mapper graphs (extracted from load-based annotation) with behavioral performance on the task, suggesting lower PC values were associated with better performance under both high load (rho (46) = −.31, p = .033) and induced anxiety (rho (46) = −.32, p = −.025) conditions.