

**Department of Neurology Faculty Papers** 

Department of Neurology

9-9-2021

# Language Tasks and the Network Control Role of the Left Inferior Frontal Gyrus

John D Medaglia Drexel University

Denise Y Harvey Drexel University

Apoorva S Kelkar Drexel University

Jared P Zimmerman University of Pennsylvania

Joely A Mass Thomas Jefferson University Follow this and additional works at: https://jdc.jefferson.edu/neurologyfp

Part of the Neurology Commons, and the Psychiatry Commons

# **Recommended Citation**

Medaglia, John D; Harvey, Denise Y; Kelkar, Apoorva S; Zimmerman, Jared P; Mass, Joely A; Bassett, Danielle S; and Hamilton, Roy H, "Language Tasks and the Network Control Role of the Left Inferior Frontal Gyrus" (2021). *Department of Neurology Faculty Papers*. Paper 259. https://jdc.jefferson.edu/neurologyfp/259

This Article is brought to you for free and open access by the Jefferson Digital Commons. The Jefferson Digital Commons is a service of Thomas Jefferson University's Center for Teaching and Learning (CTL). The Commons is a showcase for Jefferson books and journals, peer-reviewed scholarly publications, unique historical collections from the University archives, and teaching tools. The Jefferson Digital Commons allows researchers and interested readers anywhere in the world to learn about and keep up to date with Jefferson scholarship. This article has been accepted for inclusion in Department of Neurology Faculty Papers by an authorized administrator of the Jefferson Digital Commons. For more information, please contact: JeffersonDigitalCommons@jefferson.edu.

# Authors

John D Medaglia, Denise Y Harvey, Apoorva S Kelkar, Jared P Zimmerman, Joely A Mass, Danielle S Bassett, and Roy H Hamilton



Research Article: Confirmation | Cognition and Behavior

# Language Tasks and the Network Control Role of the Left Inferior Frontal Gyrus

https://doi.org/10.1523/ENEURO.0382-20.2021

Cite as: eNeuro 2021; 10.1523/ENEURO.0382-20.2021

Received: 2 September 2020 Revised: 30 April 2021 Accepted: 3 May 2021

This Early Release article has been peer-reviewed and accepted, but has not been through the composition and copyediting processes. The final version may differ slightly in style or formatting and will contain links to any extended data.

Alerts: Sign up at www.eneuro.org/alerts to receive customized email alerts when the fully formatted version of this article is published.

Copyright © 2021 Medaglia et al.

This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license, which permits unrestricted use, distribution and reproduction in any medium provided that the original work is properly attributed.

# Title Page

1 2

3

4

5

10

11 12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27 28

29

34

1. Manuscript Title (50 word maximum): Language Tasks and the Network Control Role of the Left Inferior Frontal Gyrus

2. Abbreviated Title (50 character maximum): Language and Brain Network Controllability

3. List all Author Names and Affiliations in order as they would appear in the published article:

John D. Medaglia<sup>1,2,3</sup>; Denise Y. Harvey<sup>2</sup>, Apoorva S. Kelkar<sup>1</sup>, Jared P. Zimmerman<sup>3</sup>, Joely A. Mass<sup>4</sup>, Danielle S. Bassett<sup>3,5,6,7,8,9</sup>, Roy H. Hamilton<sup>3</sup>

<sup>1</sup>Department of Psychology, Drexel University Philadelphia, Pennsylvania, 19104, USA <sup>2</sup>Department of Neurology, Drexel University Philadelphia, Pennsylvania, 19104, USA <sup>3</sup>Department of Neurology, University of Pennsylvania Philadelphia, Pennsylvania, 19104, USA <sup>4</sup>Sidney Kimmel Medical College, Thomas Jefferson University Philadelphia, Pennsylvania, 19107, USA <sup>5</sup>Department of Psychiatry, University of Pennsylvania Philadelphia, Pennsylvania, 19104, USA <sup>6</sup>Department of Bioengineering, University of Pennsylvania Philadelphia, Pennsylvania, 19104, USA <sup>7</sup>Department of Physics and Astronomy, University of Pennsylvania Philadelphia, Pennsylvania, 19104, USA <sup>8</sup>Department of Electrical and Systems Engineering, University of Pennsylvania Philadelphia, Pennsylvania, 19104, USA 9Santa Fe Institute Santa Fe, New Mexico, 87501, USA

4. Author Contributions: JDM, DYH, DSB & RHH designed research; JDM, ASK, DYH, JAM, JPZ performed research; JDM, DYH, ASK, JPZ Analyzed data; JDM wrote the paper.

30 5. Correspondence should be addressed to (include email address): John D. Medaglia, 31 johnmedaglia@gmail.com 32 6. Number of Figures: 6

33 7. Number of Tables: 13

35 8. Number of Multimedia: 0

36 9. Number of words for Abstract: 256

- 37 10. Number of words for Significance Statement: 88
- 38 11. Number of words for Introduction: 814

39 12. Number of words for Discussion: 2345

40 13. Acknowledgements: 0

41 14. Conflict of Interest: A. No ( 'Authors report no conflict of interest')

- 15. Funding sources: NIH Office of the Director under award DP5-OD021352-02, Perelman School of 42
- 43 Medicine under a Translational Neuroscience Initiative Award.

<sup>44</sup> 

# 46 Abstract

47 Recent work has combined cognitive neuroscience and control theory to make predictions about cognitive 48 control functions. Here, we test a link between whole-brain theories of semantics and the role of the left inferior 49 frontal gyrus (LIFG) in controlled language performance using network control theory, a branch of systems 50 engineering. Specifically, we examined whether two properties of node controllability - boundary and modal 51 controllability - were linked to semantic selection and retrieval on sentence completion and verb generation tasks. 52 We tested whether the controllability of the left IFG moderated language selection and retrieval costs and the 53 effects of continuous theta burst stimulation (cTBS), an inhibitory form of transcranial magnetic stimulation 54 (TMS) on behavior in 41 human subjects (25 active, 16 sham). We predicted that boundary controllability - a 55 measure of the theoretical ability of a node to integrate and segregate brain networks – would be linked to word 56 selection in the contextually-rich sentence completion task. In contrast, we expected that modal controllability – a 57 measure of the theoretical ability of a node to drive the brain into specifically hard-to-reach states - would be 58 linked to retrieval on the low-context verb generation task. Boundary controllability was linked to selection and 59 to the ability of TMS to reduce response latencies on the sentence completion task. In contrast, modal 60 controllability was not linked to performance on the tasks or TMS effects. Overall, our results suggest a link 61 between the network integrating role of the LIFG and selection and the overall semantic demands of sentence 62 completion.

# 63 Significance Statement

Our understanding of language systems and responses to neural stimulation is incomplete. Here, we demonstrate that the effects of neuromodulation (transcranial magnetic stimulation, TMS) on verbal language production are linked to the role of the left inferior frontal gyrus in mediating communication across white matter anatomical networks. We replicate prior findings in weighted anatomical networks, and further identify a link between the role of the LIFG in word selection demands. These findings provide a critical basis to reconcile local and whole brain models of language in the brain.

# 71 Introduction

Effective language production requires cognitive control: the mental processes that support flexible, contextually driven thought and action (Snyder et al., 2011). In contrast to cognitive control tasks that require inhibition of single prepotent exemplars, language tasks are frequently underdetermined - multiple responses might be appropriate (Snyder et al., 2014). Fluent language requires the ability to meet word retrieval (recalling task-appropriate words) and selection (selecting a subset of retrieved words to speak) demands when speaking. However, selection and retrieval demands vary based on the nature of specific tasks, sentence structures, and word combinations. In some cases, retrieving and selecting words is difficult and accompanied by a sense of subjective effort, such as when the appropriate words do not readily come to mind or when many appropriate, alternative words compete for selection.

Cognitive control facilitates language production by activating the relevant representations and resolving competition among the activated representations (Badre and Wagner, 2007). Broca's area, part of the left inferior frontal gyrus (LIFG), has been linked to retrieval and selection via interactions with temporal lobe regions that mediate semantic knowledge (Anwander et al., 2007; Harvey et al., 2013). However, debates about the neuroanatomical basis of cognitive control in language remain. It is unclear whether retrieval and selection localize to the same region or different subdivisions within the LIFG, reflecting the same or different mechanistic roles (Fedorenko et al., 2012; Souza et al., 2009). Conflicting accounts have asserted that the LIFG is implicated only in selecting a single response from among competing alternatives (Botvinick et al., 2001; Thompson-Schill et al., 2997), only in effortful retrieval of responses from semantic memory (Martin and Cheng, 2006; Wagner et al., 2001), or in both retrieval and selection through different neural substrates within the LIFG (Badre and Wagner, 2007) or through shared neural substrates with different, albeit not unrelated, mechanisms (Snyder et al., 2011).

Whereas localizationist accounts focus on the role of LIFG and left temporal regions in language production, the role of domain general and specific cognitive control and their representation in brain networks remains a persistent issue (Crinion et al., 2006; Fedorenko and Thompson-Schill, 2014; Diachek et al., 2019; Ryskin et al., 2020). Moreover, the role of distributed brain networks in semantic processing is an open question, with some accounts contending that the entire brain contributes to semantic representation (Patterson et al., 2007; Huth et al., 2012; Cukur et al., 2013; Shahdloo et al., 2019; Bruffaerts et al., 2019). The focus of the current study is on multiple network roles the IFG may play based on its anatomical position in brain networks. However, the extent to which these roles relate to selection and retrieval demands in language production has not been established.

To investigate the network roles of the LIFG relevant to language demands, we applied an emerging area of engineering called network control theory (NCT) (Liu et al., 2011) to brain networks. Network control theory evaluates the nature and costs of control strategies in networks used to achieve target states. Network controllability is the ability of parts of a network (e.g., specific regions in the brain) to guide the network to target states. In a broad sense, cognitive control in the language domain is a special case of a network control problem for the brain (Medaglia, 2019): how does the brain achieve the neural states necessary to produce context-appropriate responses? Since the first theoretical network controllability analyses in large scale diffusion MRI networks (Gu et al., 2015), NCT has been used to characterize the energy required to integrate or segregate network activity (Betzel et al., 2016; Gu et al., 2017; Tang et al., 2017; Wu-Yan et al., 2018), identify correlates of cognitive function in and out of the executive domain (Kenett et al., 2018a, 2019); Lee et al., 2019), and predict or correlate the effects of brain stimulation on the brain and behavior (Khambhati et al., 2019; Stiso et al., 2019).

Building on our previous study (Medaglia et al., 2018a), the current study specifically investigated (1) retrieval and selection demands in verbal language production (2) task-level differences in sentence completion and verb generation using weighted anatomical networks. We used NCT to compute the controllability of the LIFG within distributed brain networks. In NCT, a brain network can be represented as graphs that comprise nodes (e.g., brain regions) and edges (e.g., anatomical connections between regions) (Gu et al., 2015; Medaglia et al., 2018a; Patankar et al., 2020). We asked whether LIFG network controllability influenced language performance variability related to task- and item-level differences in demands. We expected that LIFG controllability would predict performance variability during sentence completion and verb generation tasks. We hypothesized that boundary controllability - the theoretical ability of a region to drive networks into integrated or segregated states - would be positively related to sentence completion performance, facilitating semantic

123 processes that rely on multiple networks. For both tasks, we predicted that higher boundary controllability would 124 be associated with reduced selection costs prior to administering TMS. In contrast, we expected that modal 125 controllability - the ability of a region to easily drive the brain into difficult-to-reach states - would be more 126 related to the decontextualized, single-exemplar retrieval demands required in a verb generation task, since that 127 task requires subjects to generate a single word in response to a cue, where there is no contextual information/meaning (unlike a sentence). Regarding neuromodulation effects, we expected that boundary 128 129 controllability would moderate transcranial magnetic stimulation (TMS) effects on overall sentence completion 130 performance and selection demands. In contrast, we expected that TMS effects would interact with retrieval 131 demands in verb generation and would be moderated by modal controllability. These relationships would provide 132 further evidence of demand-controllability associations within the LIFG.

# 133 Methods

# 134 Subjects

Forty-one healthy individuals (mean age = 25.3, St.D. = 5.9, 23 female) were scanned on a 3T Prisma scanner at 135 136 the University of Pennsylvania in the present study. There were 16 subjects (Age: 25.67, St.D. = 7.03) in the 137 sham group and 25 subjects (Age: 25.20, St.D. = 4.9) in the active group. Our previous study included n=32 (12 Sham, 20 Active) subjects (Medaglia et al., 2018a). From the previous n=32 sample, two left-handed subjects 138 139 (from the active group) and 2 subjects with English as a Second Language (from the sham group) were excluded 140 for the current study, leaving 28 subjects from the previous study included in the current study. The 13 new 141 subjects were right handed native English speakers with 7 subjects in the sham group and 6 subjects in the active 142 group. All procedures were approved in a convened review by the University of Pennsylvania's Institutional 143 Review Board and were carried out in accordance with the guidelines of the Institutional Review Board/Human 144 Subjects Committee, University of Pennsylvania. All participants volunteered with informed consent in writing 145 prior to data collection.

# 146 **Overview of Methods**

147 Network controllability characterizes the theoretical ability of a node in a network (e.g., a region in the brain) to 148 drive the state of network activity Liu et al. (2011). Here, we built on our previous work linking boundary 149 controllability to performance on open-ended language tasks and modal controllability to closed-ended language 150 tasks Medaglia et al. (2018a). Specifically, the current study focused on task-level differences between two 151 open-ended tasks - sentence completion and verb generation - and two dimensions of language demands -152 selection and retrieval Snyder and Munakata (2008); Snyder et al. (2014). Sentence completion task stimuli 153 contain additional grammatical structure and contextual semantics than verb generation task stimuli. Intuitively, 154 we expected that these processing demands would rely on multiple brain networks, and the theoretical role of the 155 LIFG in mediating among networks could be measured with boundary controllability. In contrast, verb generation task stimuli might place greater demands on the LIFG when subjects must obtain associations in the 156 157 absence of additional task structure or cues. We expected that if these demands are reflected in the LIFG's role in 158 achieving difficult-to-reach states (i.e., specifically states of activation that are otherwise difficult to activate in 159 the network), we would find a relationship between performance on verb generation and modal controllability. In 160 addition, both tasks stratified selection and retrieval demands at the item level, and we expected that the effects 161 of these demands on performance would be moderated by boundary and modal controllability, respectively. We 162 anticipated that boundary controllability would facilitate the ability to activate and select among multiple 163 competing options according to the associative, multi-network demands of semantic cognition. In contrast, we 164 anticipated that modal controllability would facilitate the ability to retrieve specific exemplars from memory, 165 prehaps facilitating cognitive associations when cues are weaker.

To test our hypotheses, subjects participated in two experimental sessions (henceforth "pre-TMS" and "post-TMS") in which subjects performed two language tasks with open-ended selection demands (verb generation and sentence completion) and one number naming task with a single appropriate response for comparison (not discussed here; see (Medaglia et al., 2018a)). Between the two task sessions, we administered either active or sham TMS. In the active TMS group, we administered continuous theta burst stimulation (cTBS), 171 a form of TMS thought to induce neural inhibition for 60 minutes or more (Huang et al., 2005), to the pars 172 triangularis within the left inferior frontal gyrus. We chose this target given its role in generalized selection in 173 semantic processing (Badre et al., 2005; Badre and Wagner, 2007), mediating cross-modal representation of 174 spoken and written words (Liuzzi et al., 2017), and patient improvements in naming after inhibitory TMS to the 175 right hemispheric homotope (Naeser et al., 2011; Harvey et al., 2017, 2019). In the sham TMS group, we 176 administered TMS to the vertex in each subject. After the experiment was complete, we constructed anatomical 177 brain networks from diffusion spectrum imaging (DSI) data acquired from each subject (Methods, Fig. 1A). Each 178 network contained 111 brain regions defined by the Lausanne anatomical parcellation (Cammoun et al., 2012) 179 and cerebellum (Diedrichsen et al., 2009) (Fig. 1B), and each pair of regions was connected by an edge weighted 180 by the number of streamlines linking those regions (Fig. 1C). We defined a simplified model of brain dynamics 181 and simulated network control to quantify modal and boundary controllability (Fig. 1D).

182

183

184 Figure 1: Overview of Methods (A) Continuous theta burst stimulation was administered to each subject's pars 185 triangularis (pictured with the bullseve) or the cranial vertex. (B) Diffusion tractography was computed for each subject. 186 A cortical parcellation was registered to each individual's anatomical T1 image to identify anatomical divisions. (C) A 187 region x region anatomical adjacency matrix was constructed representing the streamline counts between pairs of regions 188 corrected for region volume. (D) We applied a community detection algorithm to identify an initial consensus partition 189 based on partitions identified within subjects. (E) Modal and boundary controllability were computed for each node (brain 190 region) in the network for each individual. Each node received a rank representing its strength of control within the 191 individual. (F) Maps representing the variability in modal controllability (top) and boundary controllability (bottom).  $P_{1,N}$  represent different participants. The relationship between controllability values at the LIFG stimulation site and 192

193 task response times before and after stimulation were examined using mixed effects models.

# 194 Neuroimaging: Diffusion Tractography

Diffusion spectrum images (DSI) were acquired for all 41 subjects along with a T1-weighted anatomical scan at each scanning session. We followed a parallel strategy for data acquisition and construction of streamline adjacency matrices as in previous work applying network controllability statistics in human diffusion imaging networks (Gu et al., 2015; Medaglia et al., 2018a; Betzel et al., 2016). DSI scans sampled 257 directions using a Q5 half-shell acquisition scheme with a maximum *b*-value of 5,000 and an isotropic voxel size of 2.4 mm. We utilized an axial acquisition with the following parameters: repetition time (TR) = 5 s, echo time (TE) = 138 ms, 52 slices, field of view (FoV) (231, 231, 125 mm).

202 DSI data were eddy distortion corrected and reconstructed in DSI Studio (dsi-studio.labsolver.org) using 203 q-space diffeomorphic reconstruction (QSDR) (Yeh et al., 2011). QSDR first reconstructs diffusion-weighted 204 images in native space and computes the quantitative anisotropy (QA) in each voxel. These QA values are used 205 to warp the brain to a template QA volume in Montreal Neurological Institute (MNI) space using a nonlinear 206 registration algorithm. Once in MNI space, spin density functions were again reconstructed with a mean diffusion 207 distance of 1.25 mm using three fiber orientations per voxel. Fiber tracking was performed in DSI Studio with an 208 angular cutoff of 35°, step size of 1.0 mm, minimum length of 10 mm, spin density function smoothing of 0.0, 209 maximum length of 400 mm and a QA threshold determined by DWI signal in the cerebrospinal fluid. 210 Deterministic fiber tracking using a modified FACT algorithm was performed until 1,000,000 streamlines were 211 reconstructed for each individual. DSI Studio placed starting points within seeding "voxels" at subvoxel 212 resolution to account for potential partial volume influences on the fiber estimates (Campbell et al., 2005). The 213 actual seeding points were determined randomly and uniformly within the voxels. DSI Studio used a 214 deterministic random generator to place the seeds, and thus the seeding sequence was both deterministic and 215 random. These features ensured that the tracking result is reproducible using the same tracking parameters. DSI 216 Studio drew a point within the voxel range using a uniform distribution. The point was then used as the starting 217 point within the selected voxel.

218 Anatomical (T1) scans were segmented using FreeSurfer (Fischl, 2012) and parcellated using the connectome 219 mapping toolkit (Cammoun et al., 2012) plus the Diedrichsen spatially unbiased cerebellum atlas (Diedrichsen et 220 al., 2009). Compared to other functional parcellation schemes, our anatomical parcellation scheme ensures that 221 we obtained networks from a consistent anatomical location within each subject, which is essential to supporting 222 anatomical inferences and maintaining a consistent anatomical network location in each subject. The final 223 parcellation scheme including n=111 regions was registered to the B0 volume from each subject's DSI data. The 224 B0 to MNI voxel mapping produced via QSDR was used to map region labels from native space to MNI 225 coordinates. To extend region labels through the grey-white matter interface, the atlas was dilated by 4 mm 226 (Cieslak and Grafton, 2014). Dilation was accomplished by filling non-labeled voxels with the statistical mode of 227 their neighbors' labels. In the event of a tie, one of the modes was arbitrarily selected. Each streamline was 228 labeled according to its terminal region pair. From these data, we constructed a anatomical connectivity matrix, A 229 whose element  $A_{ii}$  represented the number of streamlines connecting different regions, divided by the sum of

volumes for regions *i* and *j* (Hagmann et al., 2008). Notably, there are numerous free parameters in diffusion tractography, image parcellation, and graph representations of anatomical connectivity (e.g., weighted *versus* binarized –or unweighted– graphs).

# **Cognitive Testing**

Participants performed a verb generation and sentence completion task administered with ePrime 3.0 software on a desktop computer before and after receiving TMS (Snyder and Munakata, 2008; Snyder et al., 2014; Medaglia et al., 2018a) (see Fig. 2). All stimuli were written words presented on the screen in English. Subjects were asked to provide spoken responses to the tasks.

Figure 2: Selection and Retrieval Demands within the Tasks. Items with high selection and low retrieval demands are those with many highly associated responses, and items with low selection and high retrieval demands are those with one weakly associated response. The stimuli were either verb cues in the verb generation task, or sentence cues in the sentence completion task. Even if selection and retrieval demands are similar in latent semantic analyses, each task places different predictive and syntactic demands on the semantic system that could influence performance. Selection and retrieval demands were measured continuously in a relative semantic space using LSA entropy and association strength, respectively, computed at the item level separately for each task.

248

239 240

eNeuro Accepted Manuscript

267 268 269

270

271

272 273

274

275

276

277

278

279

280

281

282

283

284

249 The order of tasks and order of task items (sentences/words) were counterbalanced across subjects, but within 250 subject's session, the order of tasks remained the same pre-TMS vs. post-TMS. Each task required а 251 approximately 5 minutes. In addition, about 5 minutes were required to set up and administer the cTBS sequence. 252 Thus, the pre-TMS session (two language tasks), TMS administration, and post-TMS session (two language 253 tasks) lasted a total of approximately 25 minutes. Items (sentences/words) were not repeated within or between 254 the sessions; half of the items per task were presented in the pre-TMS session and the other half were presented 255 in the post-TMS session for a given subject. For the verb generation task, a single written word was presented on 256 the screen, which remained on the screen for 10 seconds or until the participant made a response. For the 257 sentence completion task, segments of 1-2 words were presented serially (1,000 ms per segment) from left to 258 right, starting with the beginning of the sentence. The sentences were presented accumulatively (the prior words 259 remained on the screen until the response was given). Then, the whole sentence remained on the screen for 10 260 seconds starting from the onset of the final segment or until the participant made a response. The proportion of 261 acceptable verb responses during the sentence completion task was low (12/100) and stratified across selection 262 demands. For both tasks, trials were separated by the presentation of a fixation cross "+" for 500 ms. Subjects were given an example and five practice trials in the first administration of each language task (i.e., pre-TMS), 263 264 and were reminded of the instructions before performing the task a second time (i.e., post-TMS). In each of the 265 pre- and post- TMS sessions, subjects completed 50 trials for a total of 100 trials per task. 266

For the verb generation task, subjects were instructed to generate the first verb that came to mind when presented with a noun stimulus (e.g., "cat"). The verb could be either something the noun does (e.g., "meow") or something that is done with it (e.g., "feed"). Response times (RTs) were collected from the onset of the noun cue to the onset of the verb response. For the sentence completion task, participants were presented with a sentence, such as "They left the dirty dishes in the -----.", and were instructed to generate a single word that appropriately completes the sentence, such as "sink". RTs were computed as the latency between the onset of the subject's response. For both tasks, all items in the high vs. low selection demand conditions were matched on retrieval demands (association strength) (Snyder and Munakata, 2008).

The items for the verb generation task were identical to those used in (Snyder et al., 2011) and the items for the sentence completion task were those from (Snyder et al., 2014). The difficulty of items was sampled to cover a distribution of values computed via latent semantic analysis (LSA) applied to corpus data. In particular, items were sampled to represent a range of LSA entropy and LSA association strength (Snyder and Munakata, 2008), which represent the selection and retrieval demands of each item, respectively (Snyder and Munakata, 2008). An LSA association value of 0 means that the cue word or sentence is not strongly associated with any word in particular, whereas a value of 1 means that the cue word or sentence is strongly associated with at least one word, implying that it is easy to retrieve. An LSA entropy value of 0 indicates that the word is not related to any words, whereas higher values indicate higher relatedness to many words, which theoretically increases competition among appropriate words (Snyder and Munakata, 2008).

Verbal responses for all tasks were collected from a computer headset microphone. The microphone was calibrated to reduce sensitivity to environment background noise prior to the collection of data for each session such that the recording software was not triggered without clear verbalizations. List order was counterbalanced across participants and session (before or after active or sham stimulation). Item presentation order within each task was fully randomized across participants.

# 290 Transcranial Magnetic Stimulation

291 The Brainsight system (Rogue Research, Montreal) was used to co-register MRI data with the location of the 292 subject and the TMS coil. The stimulation site was defined as the posterior extent of the pars triangularis in each individual subject's registered T1 image. A Magstim Super Rapid<sup>2</sup> Plus<sup>1</sup> stimulator (Magstim; Whitland, UK) 293 294 was used to deliver cTBS via a 70 mm diameter figure-eight coil. cTBS consisted of 50 Hz triplets administered 295 every 200 ms (i.e., 5 Hz) (Huang et al., 2005) for 600 total pulses. To calibrate the intensity of stimulation, cTBS 296 was delivered at 80% of each participant's active motor threshold (Huang et al., 2005). Each subject's threshold 297 was determined prior to the start of the experimental session using a standard up-down staircase procedure with 298 stimulation to the motor cortex (M1). In the sham condition, the coil was held against the head at a 90-degree 299 angle at the subject's vertex to introduce a degree of induced electrical stimulation of the scalp. We administered

sham at vertex to reduce the possibility that subjects could see the orientation of the coil in the sham condition, as subjects were not naïve to TMS.

# **302** Network Controllability

To study the ability of a certain brain region to influence other regions in arbitrary ways we adopt the control theoretic notion of *controllability*. Controllability of a dynamical system refers to the possibility of driving the state of a dynamical system to a specific target state by means of an external control input (Liu et al., 2011; Ruths and Ruths, 2014; Pasqualetti et al., 2014). In the current paper, we follow the procedures applied in (Gu et al., 2015; Medaglia et al., 2018a) and focus on two network controllability statistics: *boundary* and *modal* controllability. Consistent with prior studies, we note that these statistics use linear discrete time dynamics that approximate nonlinear effects in simulations (Muldoon et al., 2016; Tiberi et al., 2017).

# **Mathematical Models**

# 311 Network Control Theory

312 All network controllability measures were computed in MATLAB. We follow previous applications of network 313 control theory in diffusion weighted imaging data as the basis for our examination of controllability and 314 cognitive control. We briefly describe the mathematical basis for the approach taken here. For a full discussion of 315 anatomical network controllability in the context of diffusion weighted imaging networks, see (Gu et al., 2015). 316 For a full discussion of the mathematical basis for anatomical network controllability see (Liu et al., 2011; Ruths 317 and Ruths, 2014; Pasqualetti et al., 2014). In contrast to traditional graph theory, network control theory offers 318 mechanistic predictors of network dynamics. Mechanistic models can provide rich tests of causal dynamics in the 319 human connectome by explicitly including a dynamic model (Medaglia et al., 2015).

The controllability of a networked system can be examined by defining a network represented by the graph G=(V,E), where V and E are the vertex (node, or here, brain region) and edge (connection, here anatomical streamline density) sets, respectively. Let  $a_{ij}$  be the weight associated with the edge  $(i,j) \in E$ , and define the weighted adjacency matrix of G as  $A=[a_{ij}]$ , where  $a_{ij}=0$  whenever  $(i,j) \notin E$ . We associate a real numeric value

324 (state) with each node, collect the node states into a vector (network state), and define the map  $x:N_{>0} \rightarrow R^n$  to

describe the evolution (*network dynamics*) of the network state over time. Using the observed network and node dynamics, network control theory can theoretically examine how the anatomical network structure relates to the types of control that nodes can exert.

# 328 **Dynamic Model of Neural Processes**

329 Following prior work, we define anatomical brain networks by subdividing the entire brain into anatomically 330 distinct brain areas (network nodes) in a commonly used anatomical atlas (Hagmann et al., 2008). Consistent 331 with prior work (Bassett et al., 2011; Hermundstad et al., 2013, 2014; Gu et al., 2015), we connect nodes by the 332 number of white matter streamlines identified by a commonly used deterministic tractography algorithm (Bassett 333 et al., 2011; Hermundstad et al., 2013, 2014; Gu et al., 2015; Betzel et al., 2016; Tang et al., 2017; Cornblath et 334 al., 2018; Stiso et al., 2019; Medaglia et al., 2018b) (for details on the tractography implementation, see 335 (Medaglia et al., 2018a)). This procedure results in sparse, weighted, undirected anatomical brain networks for 336 each subject. Properties of this network include high clustering, short path length, and strong modularity, 337 consistent with prior studies of similar network data (Bassett et al., 2011; Hagmann et al., 2008). The definition 338 of anatomical brain networks based on tractography data in humans follows from our primary hypothesis that 339 control features of neural dynamics are in part determined by the anatomical organization of the white matter in 340 the brain.

As a simplified estimate of controllability at the region of interest, we drew from intuitions applied in other work linking network anatomy and function. (Honey et al., 2009, 2010; Abdelnour et al., 2014). Although neural activity evolves through neural circuits as a collection of *nonlinear* dynamic processes, these prior studies have 345 predicted from simplified linear models. Based on this literature, we employ a simplified noise-free linear 346 discrete-time and time-invariant network model:  $\mathbf{x}(t+1) = \mathbf{A}\mathbf{x}(t) + \mathbf{B}\mathbf{u}(t),$ 347 348

where  $\mathbf{x}:R_{>0} \rightarrow R^N$  describes the state (e.g., a measure of the electrical charge, oxygen level, or firing rate) of

demonstrated that a significant amount of variance in neural dynamics as measured by resting state fMRI can be

(1)

brain regions over time, and  $\mathbf{A} \in \mathbb{R}^{N \times N}$  is a symmetric and weighted adjacency matrix. In this case, we construct a 349 weighted adjacency matrix whose elements indicate the number of white matter streamlines connecting two 350 351 different brain regions – denoted here as i and j – and we stabilize this matrix by dividing by the mean edge 352 weight. While the model used above is a discrete-time system, the controllability Gramian is statistically similar 353 to that obtained in a continuous-time system (Gu et al., 2015).

354 The diagonal elements of the matrix **A** satisfy  $A_{ii}=0$ . The input matrix **B**<sub>K</sub> identifies the control points K in and

the brain, where  $K = \{k_1, \dots, k_m\}$ 355

344

356 357

$$\mathbf{B}_K = [e_{k_1} \dots e_{k_m}], \quad (2)$$

and  $e_i$  denotes the *i*-th canonical vector of dimension N. The input  $\mathbf{u}: R_{>0} \rightarrow R^m$  denotes the control energy. 358

### **Boundary Controllability.** 359

360 Boundary controllability, a metric developed in network control theory, quantifies the role of a network node in 361 controlling dynamics between modules in hierarchical modular networks (Pasqualetti et al., 2014). Boundary 362 controllability identifies brain areas that can theoretically steer the system into states where different cognitive 363 systems are either coupled or decoupled. A region's boundary controllability describes its theoretical ability to 364 regulate the extent to which it can drive major networks to increase or decrease communication with one another. High boundary controllers are conceptually akin to the "gatekeepers" of communication between major brain 365 366 networks. Here, we applied a similar approach to that taken in (Gu et al., 2015; Medaglia et al., 2018a) to 367 quantify boundary controllability in our diffusion tractography networks and associate controllability variability 368 with cognitive performance. Specifically, we partition the brain into modules by maximizing the modularity 369 quality function (Newman, 2006) using a Louvain-like (Blondel et al., 2008) locally greedy algorithm (Jutla et 370 al., 2011). Because the modularity quality function has many near-degeneracies, we optimized the algorithm 371 multiple (100) times (Good et al., 2010).

372 Our approach differed from (Medaglia et al., 2018a) to include (1) full, weighted streamline networks and (2) 373 partitions estimated within individuals. Given that anatomical network topology can vary across subjects and is 374 explicitly of interest in examining the relationship between brain network organization, TMS, and behavior, we 375 applied a tiered strategy to obtain a consistent partition threshold. First, we obtained partitions in each of 100 376 optimizations per subject at each value of gamma from 1.0 to 4.0 in increments of 0.1. Next, we obtained the 377 mean z-Rand coefficient for each subject and obtained the mean across subjects. We observed that the peak 378 z-Rand across the sample was observed at  $\gamma$  at 2.0 (mean z-Rand score = 74.06, standard deviation = 3.8). We 379 therefore used the consensus partition at  $\gamma=2.0$  obtained from optimizations within each subject for the remainder 380 of the analysis in this study. High ranking boundary controllers were identified as the highest ranking set of 381 boundary regions between modules, and the remaining boundary regions were found within modules in the 382 network.

### 383 Modal Controllability.

384 Modal controllability refers to the ability of a node to control each evolutionary mode of a dynamical network 385 (Hamdan and Nayfeh, 1989), and can be used to identify the least controllable theoretical state from a set of 386 control nodes. Modal controllability is computed from the eigenvector matrix  $V=[v_{ij}]$  of the network adjacency

matrix A. By extension from the PBH test (Kailath, 1980), if the entry  $v_{ij}$  is small, then the *j*-th mode is poorly 387

controllable from node *i*. Following (Pasqualetti et al., 2014), we define  $\varphi_i = \sum_{j=1}^{N} (1 - \lambda_j^2(A)) v_{ij}^2$  as a scaled measure of the controllability of all *N* modes  $\lambda_1(A), \dots, \lambda_N(A)$  from the brain region *i*. Regions with high modal

390 controllability are able to control all the dynamic configurations of the network, and hence to drive the dynamics 391 towards hard-to-reach configurations. A hard-to-reach state is one that requires a high amount of energy to reach. 392 In the case of human brain networks, many competing and cooperating dynamics occur over time. As a result, the 393 high-energy states typically involve the activation of a few, specific regions in the network that would otherwise 394 express many coactivation patterns. High modal controllers are conceptually akin to dynamic "specialists" 395 driving specific, otherwise unachievable states. Intuitively, a modal controller could correspond to one that is 396 specialized to activate a single or small set of regions in the network, potentially supporting a few specific 397 computational processes at a single location in the brain.

# Statistical Analysis: Examining the Relationship Between Controllability, Cognition, and TMS effects

400 This was a mixed study design with between-subjects effects of stimulation condition (active or sham TMS) and 401 LIFG controllability, and within-subjects effects of item & selection and retrieval demands. To account for the 402 study design, analyses were conducted using multilevel modeling with maximum-likelihood estimation (Baayen 403 et al., 2008) implemented in the lme4 v.1.1-9 (Bates et al., 2014) package of R version 3.2.1 (R Core Team, 404 2016). This technique allows classical regression analyses to be performed on repeated measures data by 405 accounting for the non-independence of observations collected from each participant (i.e., multiple behavioral 406 observations obtained during the language tasks), without resorting to computing separate regression equations 407 for each subject (Lorch and Myers, 1990; Baayen et al., 2008; Baayen, 2008). Critically, multilevel modeling 408 accounts for the variances of the conditions of interest across subjects when estimating fixed effects, which is 409 appropriate due to the potentially different effects of TMS across subjects (Hamada et al., 2013; Lüders et al., 410 1985). Multilevel modeling also accounts for violations of the sphericity assumption by modeling 411 heteroskedasticity in the data when necessary, improving statistical power over other methods commonly 412 employed for analyzing repeated-measures data.

413 We excluded from analyses trials on which participants responded incorrectly (i.e., semantic and paraphasic 414 errors, hesitations, false starts) and experimenter error/equipment failures (such as false triggers for voice 415 recording), constituting a mean of 4.25% and 4.67% of all trials, respectively. In addition, responses of less than 416 200ms or greater than 10,000ms were excluded. We excluded responses below 200ms because they are likely 417 impulsive errors rather than those that reflect fast cognitive selection and retrieval and oral motor onsets 418 (Indefrey and Levelt, 2004). In addition, compared to closed-ended language tasks with a single appropriate 419 response, longer windows ensure that we measure task-relevant responses. Higher selection and retrieval 420 demands tend to increase the central tendency and tail of response times (Snyder and Munakata, 2008; Snyder et 421 al., 2014). In early piloting we found that subjects occasionally provided semantically relevant responses after 422 8-9 second delay, and the 10s cutoff allowed us to be inclusive of some of these slower responses. See Table 1 423 for total trial rejection percentages for each task, TMS session, and group.

Response times (RTs) were log-transformed due to non-normal distribution of raw RTs. For interactions with task variables, we discretized association and entropy values with a median split prior to computing interactions. Association and entropy values were centered and left continuous for interactions with the continuous controllability values.

428 Our modeling strategy was designed to test whether we replicated a prior finding that boundary 429 controllability moderated performance on the tasks when considered together (Medaglia et al., 2018a). Then, we 430 tested whether LIFG controllability was linked to TMS effects (1) between-task differences that suggest overall 431 influences of semantic processing demands or (2) the within task selection and retrieval demands. First, we tested 432 whether LIFG boundary controllability moderated TMS effects when both tasks were examined together as 433 observed in our prior study (Medaglia et al., 2018a) in this larger sample with a modified data processing stream 434 (i.e., full, weighted adjacency matrices and partitions for boundary controllability computed within subjects).

Then, we tested whether selection and retrieval demands (i.e., those measured by entropy and association strength in latent semantic analyses (Snyder et al., 2011, 2014) induced the same effect across the sentence

424

425

426

437 completion and verb generation tasks. This would determine if task-level distinctions due to differences in 438 overall semantic integration demands exist before neuromodulation. In our models, a selection cost was 439 represented by the main effect of entropy on response times: slowed response times in items with higher selection 440 demands (i.e., greater entropy). Likewise, a *retrieval cost* was represented in our models by the main effect of 441 association strength on response times: slowed response times for items with higher retrieval demands (i.e., lower 442 association strengths). To test whether these costs were moderated by controllability, we examined whether 443 baseline selection and retrieval costs were moderated by LIFG boundary and modal controllability in each task. Next, we tested whether session effects in the sham group differed across the tasks to examine if interference 444 445 observed in (Medaglia et al., 2018a) increased in both. This established an important test for whether TMS 446 alleviates interference observed in successive runs of language production as we speculated previously (Medaglia 447 et al., 2018a). After testing for session effects (i.e., pre-TMS versus post-TMS outcome) in the sham group that 448 could imply influences of increasing semantic interference (as indicated by slowed response times (Medaglia et 449 al., 2018a), we tested whether cTBS affected response times on each task. Then, we examined whether LIFG 450 controllability moderated observed TMS effects for each task. This analysis allowed us to determine if the TMS 451 effect was to mitigate this accumulated interference. The random effects structure for all models included a 452 random slope for trial order nested within subjects (Barr et al., 2013).

# 453 **Code and data availability**

454 Code for controllability measures can be found at: https://github.com/johnmedaglia/eneuro\_controllability/. Data 455 are available upon request.

# 456 **Results**

457 Across all sentence completion and verb generation data combined, we replicated the finding that LIFG boundary 458 controllability was related to performance when both tasks were examined together (main effect of boundary controllability:  $\beta = -0.002$ , p = 0.004, Table 2). In addition, boundary controllability moderated the TMS effect 459 460 (stimulation \* session \* boundary controllability:  $\beta$ =0.003, p=0.009, Table 2) In comparing the tasks, behavioral 461 evidence revealed that the costs of these demands differed across the tasks overall before TMS. Selection costs 462 (the effects of higher selection demands on performance) can be measured along a dimension as the parameter 463 weight associated with item entropy values. Accordingly, retrieval costs (the effects of higher retrieval demands 464 on performance) can be modeled as the parameter weight associated with item association strengths. Behavioral 465 data revealed a task dissociation in pre-TMS selection and retrieval costs. Specifically, selection costs were 466 greater in sentence completion (task by selection demand interaction:  $\beta = -0.180$ , p < 0.001, Table 3), whereas 467 retrieval costs were greater in verb generation ( $\beta$ =0.122, p<0.001, Table 4). These differences suggest that 468 differences in semantic demands exist at the task-level in addition to within-task variation in demands across 469 items. See Fig. 3 for estimated effects of selection and retrieval costs in the verb generation and sentence 470 completion tasks pre-TMS.

474 Figure 3: Selection and retrieval costs differ across language tasks. Selection costs were higher during the sentence
 475 completion task, whereas retrieval costs were higher in the verb generation task.

476 After detecting task differences in selection and retrieval demands, we investigated whether LIFG network 477 controllability moderated performance in response to cognitive demands at baseline. Following our behavioral 478 data, we tested the link between LIFG boundary and modal controllability on (1) sentence completion and 479 selection costs and (2) verb generation and retrieval costs. We found that the baseline selection costs were 480 moderated by LIFG boundary controllability in sentence completion (LIFG boundary controllability by entropy 481 interaction:  $\beta$ =0.001, p=0.002, see Table 5). The moderating influence of LIFG boundary controllability on the 482 effects of entropy is illustrated in Fig. 4. Modal controllability did not moderate selection demands during 483 sentence completion ( $\beta$ =-0.006, p=0.063, Table 6). Neither boundary nor modal controllability significantly 484 moderated baseline retrieval costs on verb generation ( $\beta$ =-0.001, p=0.587, Table 7;  $\beta$ =0.003, p=0.702, Table 8). 485

Figure 4: **Boundary controllability moderates selection costs during sentence completion.** Increased entropy values are associated with higher selection demands. A steeper positive slope of the relationship between entropy and response times represents higher *selection costs*. Selection costs were higher at baseline in individuals with higher boundary controllability. To visualize the effects of the continuous boundary controllability values as a third dimension, we used a split of estimated regression lines from the models at -1 and 1 standard deviations of boundary controllability across the sample at baseline. Please see Table 5 for the exact model estimates for the main effects of entropy and LIFG boundary controllability and their interaction.

In addition to differences in selection and retrieval costs across the tasks, we were interested in whether semantic interference in the sham group increased equally from the first to second session in each task. Differences across tasks could suggest that spreading activation causes increased competition in one task relative to the other with sustained task performance (Saunders and MacLeod, 2006; Nozari and Pinet, 2020). Session did not influence performance in both tasks: sentence completion response times increased overall ( $\beta$ =0.072, p=0.002) whereas verb generation did not ( $\beta$ =-0.022, p=0.319; see Tables 9 & 10). Thus, the increased context-driven nature of this task might induce more persistent, widespread activation of the semantic system that slows performance. See Fig. 5, blue dots.

502 503

504 505

Figure 5: TMS Effects. In the sham group, responses on sentence completion slowed, whereas responses on verb
 generation slightly quickened. Inhibitory TMS improved sentence completion performance relative to sham.

508 As illustrated in Fig. 5, TMS influenced response times only on sentence completion (stimulation by session 509 interaction:  $\beta = -0.092$ , p = 0.001, see Table 11; stimulation by session interaction in verb generation:  $\beta = 0.009$ . 510 p=0.750, see Table 12), improving performance by removing the slowing effect observed in the sham group. 511 Further dissociating the tasks, LIFG boundary controllability moderated the effect of inhibitory TMS only in 512 sentence completion (LIFG boundary controllability by TMS by session interaction:  $\beta$ =-0.002, p=0.046, see 513 Table 13; verb generation:  $\beta$ =-0.002, p=0.146, see Table 14). Thus, TMS effects were moderated by LIFG 514 boundary controllability in the more semantically context-rich task. See Fig. 6 for the estimated influence of 515 boundary controllability on the TMS effect. Given the complex interaction, we conducted post hoc analyses of 516 the boundary controllability values across individuals, finding that subjects in the active group had higher 517 average boundary controllability values than those in the sham group (Wilcoxon unpaired two-samples ranked 518 sum test: W=1776167, p<<0.001, see Extended Data Figure 6-1).

519 For further evaluation of whether accumulating interference or other temporal effects occurred during the 520 tasks before and after TMS, we additionally explored trial-wise effects in the pre-TMS and post-TMS sentence 521 completion data. Pre-TMS, subjects did not exhibit slowing overall (main effect of trial:  $\beta$ =0.001, p=0.113), but 522 greater slowing was observed among the items with higher selection demands (trial by selection interaction: 523  $\beta$ =0.002, p=0.005). Post-TMS, subjects exhibited slowing overall (main effect of trial:  $\beta$ =0.003, p=0.002), which 524 was also greater among items with higher selection demands (trial by selection interaction:  $\beta = 0.002$ , p = 0.001). 525 See Extended Data Figure 6-2 for response time distributions for all conditions of the data. See also Extended 526 Data Tables 6-3 and 6-4 for the complete modeling results for the trialwise pre-TMS and post-TMS effects. 527

eNeuro Accepted Manuscript

530 Figure 6: LIFG Boundary controllability moderates TMS effects. TMS effects were moderated by LIFG boundary 531 controllability specifically in sentence completion, where a crossover interaction was observed. Inhibitory TMS in individuals with higher boundary controllability attenuated the slowed performance observed pre-TMS among the 532 533 active subjects. However, in verb generation, changes in response times were consistently related to baseline 534 performance in both the active and sham condition. Boundary controllability is plotted as the zero-centered rank 535 controllability values at the LIFG across the sample. Please see Extended Data 6-1 illustrating baseline differences in 536 boundary controllability values between the Active and Sham groups. See Extended Data 6-2 for a plot of all raw RT 537 distributions by group, session, task, and selection and retrieval demands. See also Extended Data 6-3 and 6-4 for 538 trialwise modeling effects.

539

 540
 541 Extended Data Figure 6-1: Boundary controllability differed between the active and sham groups. Subjects in the active stimulation condition had higher average values of boundary controllability. The upper and lower extents of the boxes represent the mean upper 75 percentile and lower 25th percentile of values, respectively. The whiskers represent the maximum and minimum range of values.

547 Extended Data Figure 6-2: Raw RT distributions separated by task, TMS session, group, and selection and
 548 retrieval demands. Histograms represent the counts of RTs in each condition. Panels A-D in each subplot subdivide t

retrieval demands. Histograms represent the counts of RTs in each condition. Panels A-D in each subplot subdivide the data by median split along the selection (entropy) and retrieval (association strength) dimensions from the LSA analyses.

# 550 **Discussion**

546

549

We revealed novel associations between network controllability at the LIFG and controlled language functions. We found evidence linking boundary controllability to word selection and TMS effects during sentence completion. In partial agreement with our hypotheses, we revealed a link in the IFG between boundary controllability – the capacity for integrating and segregating activity across brain networks – and word selection in the context of the semantic demands of sentence processing. We did not find links between modal controllability and performance on either task or on selection and retrieval demands.

557 Consistent with theories that take a broad, whole-brain perspective on semantic processing (Patterson et al., 558 2007; Huth et al., 2012; Cukur et al., 2013; Shahdloo et al., 2019; Bruffaerts et al., 2019), part of the LIFG's role 559 in controlled language function could be to mediate the complex task of selecting context-dependent responses. 560 In individuals whose LIFG is positioned to mediate between major brain networks (i.e. those with high LIFG 561 boundary controllability), selection costs are increased. This suggests that as the LIFG increasingly mediates 562 between brain networks, it is less able to either mitigate coactivation across semantic representations (Collins and 563 Loftus, 1975; Anderson and Pirolli, 1984; Masson, 1995; De Deyne et al., 2016; Griffis et al., 2017; Mattheiss et 564 al., 2018) or select among them (Abdel Rahman and Melinger, 2019; Beaty et al., 2017; Canini et al., 2016; 565 Musz and Thompson-Schill, 2017). Moreover, task performance tends to slow on the second task administration 566 in the sham group among individuals, especially on the sentence completion task. This effect could represent 567 overall competition among representations increases over time on this task due to semantic priming. In addition, 568 because higher boundary controllability indicates a stronger role in mediating inter-network communication, 569 higher boundary controllability in the LIFG could imply that it is involved in managing additional demands in or 570 outside the language domain (de Bruin et al., 2014). Though we cannot fully distinguish between the potential 571 influences of fatigue or cognitive control in the absence of feedback and reward (Dreisbach and Fischer, 2012; 572 Hockey, 2011; Shenhav et al., 2017), these possibilities could also explain part of the TMS effect that we 573 observed.

574 Our results did not suggest a clear link between LIFG modal controllability and performance on either task or 575 a relationship with either selection or retrieval demands. In anatomical brain networks, high modal controllability 576 is strongly inversely related to node weighted degree (i.e., overall connectivity with nearest neighbors in the 577 network) (Gu et al., 2015). Thus, in persons with high LIFG modal controllability, the LIFG is more weakly 578 connected with anatomical sites one step away in the network. These weaker connections may facilitate more 579 limited, specific interactions with a few regions. This anatomical property might be especially relevant to 580 retrieval demands when subjects attempt to recall single noun-verb pairs without the additional context provided 581 by a complete sentence. For instance, when a noun is presented without context, it is potentially advantageous to 582 interact with a smaller set of brain regions to increase the speed with which a simple association with an 583 appropriate word can occur. This stands in contrast to the much richer semantic context required for sentence 584 processing, which requires sequenced, persistent engagement of large set of brain networks to guide responses 585 (Cooke et al., 2006; Friederici, 2002; Fedorenko and Thompson-Schill, 2014; Ni et al., 2000; Vigneau et al., 586 2006; Binder et al., 2009; Rogalsky and Hickok, 2009). In a prior study, modal controllability was only linked to 587 performance on the closed-ended number reading task (Medaglia et al., 2018a). Thus, it is possible that modal 588 controllability at the LIFG is restricted to cases without underdetermined competition, such as when only a 589 single, well-associated exemplar (e.g., a number associated with a lexical form) is appropriate. If modal 590 controllability is more generally linked to specific, well-learned representations, it is possible that it is more 591 relevant to retrieving specific episodes and items with no competition.

592 Our TMS effects further provide evidence that LIFG boundary controllability moderates processing demands 593 in language tasks with multiple processing demands. Pre-TMS, selection costs were more pronounced on

594 sentence completion than verb generation and higher in those with stronger LIFG boundary controllability. Over 595 sessions, slowed response times occurred in the sham group only on sentence completion. Higher LIFG boundary 596 controllability was associated with improved sentence completion performance after TMS. Thus, it is possible 597 that the LIFG manages multi-network processing demands. Stronger multi-network anatomical connectivity 598 could increase subjects' proneness to semantic satiation (a transient loss of meaning) via repeated performance of 599 the semantically rich sentence completion task. Further, inhibitory stimulation to the LIFG in individuals with 600 higher boundary controllability might reduce more general demands on this region that are incurred by mediating 601 among networks across the brain. For example, competition between the goal to stay on task versus attend to 602 other tasks might further tax the LIFG in these individuals over time. Alternatively, domain-general cognitive 603 control mechanisms could mediate slowed performance in the absence of reward, which is one basis of widely 604 observed potential effort-reward tradeoffs in behavior (Shenhav et al., 2017), and a potential explanation of 605 cognitive fatigue (Dobryakova et al., 2013; Fukuda et al., 2010; Milyavskaya et al., 2019). To test these 606 possibilities, future studies could manipulate demands within and out of the language domain over several 607 interleaved blocks of task performance. The role of reward on performance could be strong when high effort is 608 predicted or required (Kool and Botvinick, 2014; Kool et al., 2017; Kool and Botvinick, 2018). Manipulating task demands and rewards in neuromodulation studies could further distinguish how variability in the network 609 610 role of the LIFG mediates domain general and specific demands.

611 While our analyses focused on the anatomical connectivity of the LIFG, the mechanism of inhibitory TMS's 612 beneficial effect presumably involves local effects at the site of stimulation. Specifically, cTBS is thought to 613 induce inhibition involving complex effects on GABA-ergic neurons (Cárdenas-Morales et al., 2010; Trippe et 614 al., 2009; Gong et al., 2009; Stagg et al., 2009; Li et al., 2019). Previously, behavioral and computational work 615 suggested that word selection can be facilitated using GABA agonists (Snyder et al., 2011). Our current findings 616 point to the intriguing possibility that GABA-mediated mechanisms might parse the multi-network demands on 617 the LIFG. For instance, the LIFG's ability to efficiently select task-relevant words might be especially challenged 618 with sustained task effort when overall network demands on the LIFG are high. If the LIFG is inhibited (e.g., 619 with TMS), the neural gains on task-relevant information in the network may be enhanced when the overall 620 activity in this node is decreased (e.g., (Houghton and Tipper, 1996; Katzner et al., 2011; Ingham and McAlpine, 621 2005)), facilitating task-relevant responses (Houghton and Tipper, 1996; Herd et al., 2006). This benefit in 622 healthy individuals could be linked to evidence in individuals with aphasia after stroke. Some individuals with 623 aphasia benefit from inhibitory TMS to "noisy" node in the right inferior frontal gyrus, which sometimes 624 inherits the role of the damaged LIFG post-stroke (Torres et al., 2013). This notion could be examined by 625 applying inhibitory stimulation to the right IFG post stroke in individuals with aphasia and observing if language 626 task performance improves.

627 More broadly, we note that the task demands and cognitive control in sentence completion and verb 628 generation remain incompletely understood. Selection and retrieval demands might recruit anatomically different 629 brain networks, which could explain the relative lack of findings linking retrieval to LIFG controllability. In 630 addition, while we focused on the role of the LIFG with respect to the entire brain in order to be consistent with 631 broad, whole-brain semantic theories, it is reasonable to suspect that classic theories of more specialized, 632 left-lateralized language functions implicate a smaller set of networks to mediate these demands (Fedorenko, 633 2014). For example, circuits involving LIFG-anterior temporal lobe might be most relevant to selection (Musz 634 and Thompson-Schill, 2017; Piai and Knight, 2018), while those involving the hippocampus might be more 635 relevant to retrieval (Eldridge et al., 2000; Greenberg et al., 2005; Whitney et al., 2009). However, invasive 636 neural recordings also suggest that these processes transiently recruit a wide swath of the cortex across the entire 637 brain (Riès et al., 2017), challenging the assumption that a single-circuit model will be sufficient to account for 638 these functions. Future studies could examine the role of single circuits and networks (Chai et al., 2016) with 639 EEG and especially electrocorticography paired with anatomical diffusion tractography to obtain a more 640 comprehensive, multi-network model with good spatial and temporal resolution. Moreover, finer distinctions 641 between domain-general and language domain-specific processes and regions could improve how we 642 conceptualize task-level, selection, and retrieval demands (Fedorenko and Thompson-Schill, 2014; Fedorenko, 643 2014; Diachek et al., 2019; Blank and Fedorenko, 2017; Ridderinkhof et al., 2004). For instance, prior work 644 applying TMS has dissociated semantic processing and phonological processing in the anterior & posterior LIFG, 645 respectively (Hartwigsen et al., 2010; Ishkhanyan et al., 2020), with both contributing to grammatical sentence 646 production (Hartwigsen et al., 2016). In addition, an important difference between the sentence completion and 647 verb generation tasks is that sentences could be more likely to recruit predictive processes mediated through the

LIFG Arai and Keller (2013); Altmann and Mirković (2009); Grisoni et al. (2017); Vasishth et al. (2019); Yoshida et al. (2013), which we are not able to fully distinguish in the current study. Thus, investigating specific anatomical and functional pathways with tasks that dissociate these processes would further inform the relationship between LIFG anatomical connectivity and selection, retrieval, and other language production processes. Last and significantly, reward could be manipulated to dissociate task-related semantic satiation in the sentence completion task from reward-related processes (Kool and Botvinick, 2014; Kool et al., 2017; Kool and Botvinick, 2018; Savine and Braver, 2010; Shenhav et al., 2013).

655 Several limitations could be addressed with future studies. While our use of mixed effects modeling 656 statistically accounts for unequal sample sizes and variances, the between-subject design and unequal samples are 657 limitations. Future studies could use within-subjects crossover research designs with equal simple sizes. We used 658 an anatomically-based approach to investigate the link between LIFG controllability and demands in controlled 659 language performance. Here, our findings suggest that investigators should consider matching network measures 660 of interest (controllability or others) across active and sham groups at the site of stimulation when feasible. As 661 mentioned above, additional tasks that manipulate demand within and outside the language domain might further 662 elucidate the relationship between the network control role of the LIFG and cognitive control. In addition, while 663 we chose our anatomical network and tractography approach to be consistent with prior work using an 664 anatomically-based atlas, diffusion tractography is fundamentally limited (Thomas et al., 2014; Maier-Hein et al., 665 2017) and other tractography and parcellation schemes are available. In particular, integrating well-established 666 functional parcellations to focus on specific networks and their interactions could refine system-level predictions 667 about the relationships between network controllability, language performance, and TMS-induced network 668 effects (Bevnel et al., 2019).

669 In our behavioral data, we also observed some pre-TMS differences across individuals with high and low 670 boundary controllability in the active and sham groups. Most notably, boundary controllability was higher on 671 average in the active group that was accompanied by an inversion in the model-estimated brain-behavior relationship in sentence completion Pre-TMS. The TMS effect on this task appears to mitigate the slowing effect 672 673 of boundary controllability on RTs in the active group subjects. In the current data, our results are unlikely to be 674 accounted for by these pre-TMS differences. Our mixed effects modeling accounted for deviations in the active 675 relative to the sham group. In the Pre-TMS session, the relationship between boundary controllability and time 676 was positive, meaning that subjects with higher boundary controllability were slower. Post-TMS, the relationship 677 between boundary controllability and RTs was flattened. Thus, among individuals with relatively stronger 678 boundary controllability in the LIFG, TMS could mitigate the influence of inter-network processing demands on 679 average response times during sentence completion. Nevertheless, it is clear that additional studies would be 680 beneficial. Specifically, if sampling effects introduced pre-TMS differences at random, larger or prospectively 681 assigned studies could obtain better matched pre-TMS for controllability or other network measures of interest. 682 In addition, it is possible that other psychological differences that moderate controlled language functions such as 683 anxiety could influence results (Snyder et al., 2014). Further, subjects responded to the verb generation task with 684 verbs, whereas most responses to sentence completion were nouns. While we are unaware of specific prior data 685 suggesting that the cognitive processes mediating spoken noun and verb production differ specifically with 686 respect to the selection and retrieval demands studied here, this could be a topic for future studies. Moreover, our 687 choice to stimulate *pars triangularis* might be more relevant to word selection than retrieval, and future studies 688 could investigate whether controllability in the pars opercularis moderates performance in retrieval (Badre et al., 689 2005; Badre and Wagner, 2007). Lastly, the use of network controllability in diffusion tractography has several 690 challenges. Questions remain about the appropriateness of linear approximations (Gu et al., 2015; Friston, 2008; 691 Schiff, 2012), single-node control schemes (Tu et al., 2018; Suweis et al., 2019; Pasqualetti et al., 2019), and the 692 relevance of network-wide estimations to processes involving local (cognitive) computations (Medaglia, 2019).

# 693 Conclusion

The emerging synergy between cognitive neuroscience and neural engineering provides many opportunities. Here, drawing from whole-brain theories of semantics, a potential link between the role of the left inferior frontal gyrus in inter-network communication was examined with network control theory. Overall, we found evidence that an increased role for the LIFG at the boundaries of major networks is potentially associated with resolving competetion when processing sentences. This effect can be mitigated with inhibitory TMS in individuals whose LIFG serves a stronger role in inter-network connectivity. The mapping between general measures of node controllability and specific regional cognitive functions will require us to refine our models of cognitive control
 in language alongside our network imaging. Combining static anatomical measures with dynamic data (fMRI,
 EEG, electrocorticography) and neuromodulation could allow us to more specifically parse the distributed neural
 signals that mediate controlled language performance. In the long term, refined models could allow us to enhance
 this critical human function in health and disease.

# 706 Acknowledgments

JDM, DSB, and RHH acknowledge support pertinent to this work from the Office of the Director at the National
Institutes of Health and the National Institute of Mental Health (NIMH) through grant number
1-DP5-OD-021352-01 and the Perelman School of Medicine via a Translational Neuroscience Initiative. DSB
also acknowledges support for TMS and network control theory via NIMH grant RF1-MH116920. The content is
solely the responsibility of the authors and does not necessarily represent the official views of any of the funding
agencies.

# 714 **References and Notes**

715	Abdal Pahman P. Malingar A (2010) Samantic processing during language production: an undate of the
715	Auder Kalman K, Menniger A (2019) Semantic processing during language production, an update of the minging laviage hotwork. Language Cognition and Nauroscience 24:1176–1102
710	Abdelingurg E. Voos IIII. Dai A. (2014). Network diffusion accurately models the relationship between
719	Addeniour F, voss fro, Kaj A (2014) Network unfusion accurately modes une relationship between
710	structural and functional brain connectivity networks. <i>Neuroimage</i> 90:353–347.
719	Authania 11, Mirkovic J (2009) incrementanty and prediction in numan sentence processing. <i>Cognitive</i>
720	
721	Anderson JK, Pirolli PL (1984) Spread of activation. Journal of Experimental Psychology: Learning,
722	Memory, and Cognition 10:791.
723	Anwander A, Hitgemeyer M, von Cramon DY, Friederici AD, Knosche TR (2007) Connectivity-based
724	parcellation of broca's area. Cerebral cortex 17:810–825.
725	Aral M, Keller F (2013) The use of verb-specific information for prediction in sentence processing.
726	Language and Cognitive Processes 28:525–560.
727	Baayen RH (2008) Analyzing linguistic data: A practical introduction to statistics using R Cambridge
728	University Press.
729	Baayen RH, Davidson DJ, Bates DM (2008) Mixed-effects modeling with crossed random effects for
730	subjects and items. <i>Journal of memory and language</i> 59:590–412.
/31	Badre D, Poldrack KA, Pare-Blagoev EJ, Insier KZ, Wagner AD (2005) Dissociable controlled retrieval
732	and generalized selection mechanisms in ventrolateral pretrontal cortex. Neuron 4/:90/–918.
/33	Badre D, Wagner AD (2007) Left ventrolateral prefrontal cortex and the cognitive control of memory.
/34	Neuropsychologia 45:2883–2901.
735	Barr DJ, Levy R, Scheepers C, Tily HJ (2013). Random effects structure for confirmatory hypothesis
/36	testing: Keep it maximal. Journal of memory and language 68(3):255-18.
/3/	Bassett DS, Wymbs NF, Porter MA, Mucha PJ, Carlson JM, Gratton S1 (2011) Dynamic reconfiguration
/38	of human brain networks during learning. Proceedings of the National Academy of
/39	Sciences 108: /641–/646.
740	Bates D, Maechler M, Bolker B, Walker S et al. (2014) Ime4: Linear mixed-effects models using eigen and
741	s4. R package version 1:1–23.
742	Beaty RE, Christensen AP, Benedek M, Silvia PJ, Schacter DL (2017) Creative constraints: Brain activity
743	and network dynamics underlying semantic interference during loca production. <i>Neuroimage</i> 148:189–196.
744	Betzel RF, Gu S, Medaglia JD, Pasqualetti F, Bassett DS (2016) Optimally controlling the human
745	connectome: the role of network topology. Scientific reports 6:307/0.
746	Beynel L, Deng L, Crowell C, Dannhauer M, Paimer H, Hilbig S, Petercnev AV, Luber B, Lisanby SH,
747	Cabeza R et al. (2019) Structural controllability predicts functional patterns and neuromodulatory benefits $-704208$
748	associated with working memory. <i>bioKxiv</i> p. 794388.
749	Binder JR, Desal RH, Graves W W, Conant LL (2009) where is the semantic system? a critical review and
/50	meta-analysis of 120 tunctional neuroimaging studies. <i>Cerebral cortex</i> 19:2767–2796.
/51	Blank IA, Fedorenko E (2017) Domain-general brain regions do not track linguistic input as closely as
752	anguage-selective regions. Journal of Neuroscience 31:3999–10011.
153	Biondel VD, Guillaume JL, Lambiotte R, Lefebvre E (2008) Fast unfolding of communities in large
/54	networks. Journal of statistical mechanics: theory and experiment 2008;P10008.
155	Bolymick, Braver 13, Barch DM, Carler CS, Cohen JD (2001) Coninct monitoring and cognitive control.
/50	Psychological review 108:024.
151	Brunaerts R, De Deyne S, Meersmans R, Eluzzi AG, Storns G, Vandenbergne R (2019) Redelining the
/58	resolution of semantic knowledge in the brain: advances made by the introduction of models of semantics
139	ni neuronnagnig. Neuroscience & Biobenaviorai Keviews.
/00	Cammoun L, Gigandet A, Meskaldji D, Thiran JP, Sporns O, Do KQ, Maeder P, Meuli K, Hagmann P (2012) Magning the human connectance of multiple geoleg with diffusion constructions in the second
761	(2012) mapping the number connectome at multiple scales with diffusion spectrum mri. <i>Journal of</i>
/02	neuroscience methods 205:580–597.
103	Campoen JS, Studiqi K, Kymar VV, Sadikot AF, Pike GB (2005) Flow-based fiber tracking with diffusion
/04	tensor and q-ball data: validation and comparison to principal diffusion direction techniques.

NeuroImage 27:725–736.

- Canini M, Della Rosa PA, Catricalà E, Strijkers K, Branzi FM, Costa A, Abutalebi J (2016) Semantic interference and its control: A functional neuroimaging and connectivity study. Human brain mapping 37:4179-4196.
- Cárdenas-Morales L, Nowak DA, Kammer T, Wolf RC, Schönfeldt-Lecuona C (2010) Mechanisms and applications of theta-burst rtms on the human motor cortex. Brain topography 22:294-306.
- Chai LR, Mattar MG, Blank IA, Fedorenko E, Bassett DS (2016) Functional network dynamics of the language system. Cerebral Cortex 26:4148-4159.
- Cieslak M, Grafton S (2014) Local termination pattern analysis: a tool for comparing white matter morphology. Brain imaging and behavior 8:292-299.
- Collins AM, Loftus EF (1975) A spreading-activation theory of semantic processing. Psychological
- Cooke A, Grossman M, DeVita C, Gonzalez-Atavales J, Moore P, Chen W, Gee J, Detre J (2006) Large-scale neural network for sentence processing. Brain and language 96:14-36.
- Cornblath EJ, Tang E, Baum GL, Moore TM, Adebimpe A, Roalf DR, Gur RC, Gur RE, Pasqualetti F,
- Satterthwaite TD et al. (2019) Sex differences in network controllability as a predictor of executive function in youth. NeuroImage 188:122-134.
  - Cornblath EJ, Tang E, Baum GL, Moore TM, Roalf DR, Gur RC, Gur RE, Pasqualetti F, Satterthwaite TD, Bassett DS (2018) Sex differences in network controllability as a predictor of executive function in youth. arXiv preprint arXiv:1801.04623.
- Crinion J, Turner R, Grogan A, Hanakawa T, Noppeney U, Devlin JT, Aso T, Urayama S, Fukuyama H, Stockton K et al. (2006) Language control in the bilingual brain. Science 312:1537-1540.
- Cukur T, Nishimoto S, Huth AG, Gallant JL (2013) Attention during natural vision warps semantic representation across the human brain. Nature neuroscience 16:763.
- de Bruin A, Roelofs A, Dijkstra T, FitzPatrick I (2014) Domain-general inhibition areas of the brain are involved in language switching: Fmri evidence from trilingual speakers. NeuroImage 90:348-359
- De Deyne S, Navarro DJ, Perfors A, Storms G (2016) Structure at every scale: A semantic network account of the similarities between unrelated concepts. Journal of Experimental Psychology: General 145:1228. Diachek E, Blank I, Siegelman M, Fedorenko E (2019) The domain-general multiple demand (md) network
- does not support core aspects of language comprehension: a large-scale fmri investigation. BioRxiv p.
  - Diedrichsen J, Balsters JH, Flavell J, Cussans E, Ramnani N (2009) A probabilistic mr atlas of the human cerebellum. Neuroimage 46:39-46.
- Dobryakova E, DeLuca J, Genova HM, Wylie GR (2013) Neural correlates of cognitive fatigue: cortico-striatal circuitry and effort-reward imbalance. Journal of the International Neuropsychological Society 19:849-853.
- Dreisbach G, Fischer R (2012) The role of affect and reward in the conflict-triggered adjustment of cognitive control. Frontiers in Human Neuroscience 6:342.
- Eldridge LL, Knowlton BJ, Furmanski CS, Bookheimer SY, Engel SA (2000) Remembering episodes: a selective role for the hippocampus during retrieval. Nature neuroscience 3:1149-1152.
- Fedorenko E (2014) The role of domain-general cognitive control in language comprehension. Frontiers in
- Fedorenko E, Duncan J, Kanwisher N (2012) Language-selective and domain-general regions lie side by side within Broca's area. Current Biology 22:2059-2062.
- Fedorenko E, Thompson-Schill SL (2014) Reworking the language network. Trends in cognitive sciences 18:120-126.
- Fischl B (2012) Freesurfer. Neuroimage 62:774-781.
- Friederici AD (2002) Towards a neural basis of auditory sentence processing. Trends in cognitive
- Friston K (2008) Hierarchical models in the brain. PLoS computational biology 4.
- Fukuda S, Yamano E, Joudoi T, Mizuno K, Tanaka M, Kawatani J, Takano M, Tomoda A,
- Imai-Matsumura K, Miike T et al. (2010) Effort-reward imbalance for learning is associated with fatigue in 817 school children. Behavioral Medicine 36:53-62.

818 Gong N, Li Y, Cai GQ, Niu RF, Fang Q, Wu K, Chen Z, Lin LN, Xu L, Fei J et al. (2009) Gaba 819 transporter-1 activity modulates hippocampal theta oscillation and theta burst stimulation-induced 820 long-term potentiation. Journal of Neuroscience 29:15836-15845. 821 Good BH, De Montjoye YA, Clauset A (2010) Performance of modularity maximization in practical 822 contexts. Physical Review E 81:046106. 823 Greenberg DL, Rice HJ, Cooper JJ, Cabeza R, Rubin DC, LaBar KS (2005) Co-activation of the amygdala, 824 hippocampus and inferior frontal gyrus during autobiographical memory retrieval. 825 Neuropsychologia 43:659–674. 826 Griffis JC, Nenert R, Allendorfer JB, Szaflarski JP (2017) Linking left hemispheric tissue preservation to 827 fmri language task activation in chronic stroke patients. *Cortex* 96:1–18. 828 Grisoni L, Miller TM, Pulvermüller F (2017) Neural correlates of semantic prediction and resolution in 829 sentence processing. Journal of Neuroscience 37:4848-4858. Gu S, Betzel RF, Mattar MG, Cieslak M, Delio PR, Grafton ST, Pasqualetti F, Bassett DS (2017) Optimal 830 831 trajectories of brain state transitions. Neuroimage 148:305-317. 832 Gu S, Pasqualetti F, Cieslak M, Telesford QK, Alfred BY, Kahn AE, Medaglia JD, Vettel JM, Miller MB, 833 Grafton ST et al. (2015) Controllability of structural brain networks. Nature communications 6:1-10. 834 Hagmann P, Cammoun L, Gigandet X, Meuli R, Honey CJ, Wedeen VJ, Sporns O (2008) Mapping the 835 structural core of human cerebral cortex. PLoS Biol 6:e159. 836 Hamada M, Murase N, Hasan A, Balaratnam M, Rothwell JC (2013) The role of interneuron networks in 837 driving human motor cortical plasticity. Cerebral cortex 23:1593-1605. 838 Hamdan A, Nayfeh A (1989) Measures of modal controllability and observability for first-and second-order 839 linear systems. Journal of guidance, control, and dynamics 12:421-428. Hartwigsen G, Price CJ, Baumgaertner A, Geiss G, Koehnke M, Ulmer S, Siebner HR (2010) The right 840 841 posterior inferior frontal gyrus contributes to phonological word decisions in the healthy brain: evidence 842 from dual-site tms. Neuropsychologia 48:3155-3163. 843 Hartwigsen G, Weigel A, Schuschan P, Siebner HR, Weise D, Classen J, Saur D (2016) Dissociating 844 parieto-frontal networks for phonological and semantic word decisions: a condition-and-perturb tms study. 845 Cerebral cortex 26:2590-2601. 846 Harvey DY, Mass JA, Shah-Basak PP, Wurzman R, Faseyitan O, Sacchetti DL, DeLoretta L, Hamilton RH 847 (2019) Continuous theta burst stimulation over right pars triangularis facilitates naming abilities in chronic 848 post-stroke aphasia by enhancing phonological access. Brain and language 192:25-34. 849 Harvey DY, Podell J, Turkeltaub PE, Faseyitan O, Coslett HB, Hamilton RH (2017) Functional 850 reorganization of right prefrontal cortex underlies sustained naming improvements in chronic aphasia via 851 repetitive transcranial magnetic stimulation. Cognitive and behavioral neurology: official journal of the 852 Society for Behavioral and Cognitive Neurology 30:133. 853 Harvey DY, Wei T, Ellmore TM, Hamilton AC, Schnur TT (2013) Neuropsychological evidence for the 854 functional role of the uncinate fasciculus in semantic control. Neuropsychologia 51:789-801. 855 Herd SA, Banich MT, O'reilly RC (2006) Neural mechanisms of cognitive control: An integrative model of 856 stroop task performance and fmri data. Journal of cognitive neuroscience 18:22-32. 857 Hermundstad AM, Bassett DS, Brown KS, Aminoff EM, Clewett D, Freeman S, Frithsen A, Johnson A, 858 Tipper CM, Miller MB et al. (2013) Structural foundations of resting-state and task-based functional 859 connectivity in the human brain. Proceedings of the National Academy of Sciences 110:6169-6174. 860 Hermundstad AM, Brown KS, Bassett DS, Aminoff EM, Frithsen A, Johnson A, Tipper CM, Miller MB, 861 Grafton ST, Carlson JM (2014) Structurally-constrained relationships between cognitive states in the 862 human brain. PLoS Comput Biol 10:e1003591. 863 Hockey GRJ (2011) A motivational control theory of cognitive fatigue. . 864 Honey CJ, Thivierge JP, Sporns O (2010) Can structure predict function in the human brain? 865 Neuroimage 52:766-776. Honey C, Sporns O, Cammoun L, Gigandet X, Thiran JP, Meuli R, Hagmann P (2009) Predicting human 866 867 resting-state functional connectivity from structural connectivity. Proceedings of the National Academy of 868 Sciences 106:2035-2040. 869 Houghton G, Tipper SP (1996) Inhibitory mechanisms of neural and cognitive control: Applications to 870 selective attention and sequential action. Brain and Cognition

- 871 Huang YZ, Edwards MJ, Rounis E, Bhatia KP, Rothwell JC (2005) Theta burst stimulation of the human 872 motor cortex. Neuron 45:201-206. 873 Huth AG, Nishimoto S, Vu AT, Gallant JL (2012) A continuous semantic space describes the 874 representation of thousands of object and action categories across the human brain. Neuron 76:1210–1224. 875 Indefrey P, Levelt WJ (2004) The spatial and temporal signatures of word production components. 876 Cognition 92:101-144. 877 Ingham NJ, McAlpine D (2005) Gabaergic inhibition controls neural gain in inferior colliculus neurons 878 sensitive to interaural time differences. Journal of Neuroscience 25:6187-6198. 879 Ishkhanyan B, Michel Lange V, Boye K, Mogensen J, Karabanov A, Hartwigsen G, Siebner HR (2020) 880 Anterior and posterior left inferior frontal gyrus contribute to the implementation of grammatical 881 determiners during language production. Frontiers in Psychology 11:685. 882 Jutla IS, Jeub LG, Mucha PJ et al. (2011) A generalized louvain method for community detection 883 implemented in matlab. URL http://netwiki. amath. unc. edu/GenLouvain . 884 Kailath T (1980) Linear systems, Vol. 156 Prentice-Hall Englewood Cliffs, NJ. 885 Katzner S, Busse L, Carandini M (2011) Gabaa inhibition controls response gain in visual cortex. The 886 Journal of Neuroscience 31:5931-5941. 887 Kenett YN, Beaty RE, Medaglia JD (2018b) A computational network control theory analysis of depression 888 symptoms. Personality neuroscience 1. 889 Kenett YN, Medaglia JD, Beaty RE, Chen O, Betzel RF, Thompson-Schill SL, Oiu J (2018a) Driving the 890 brain towards creativity and intelligence: A network control theory analysis. Neuropsychologia 118:79-90. 891 Khambhati AN, Kahn AE, Costantini J, Ezzyat Y, Solomon EA, Gross RE, Jobst BC, Sheth SA, Zaghloul 892 KA, Worrell G et al. (2019) Functional control of electrophysiological network architecture using direct 893 neurostimulation in humans. Network Neuroscience 3:848-877. 894 Kool W, Botvinick M (2014) A labor/leisure tradeoff in cognitive control. Journal of Experimental 895 Psychology: General 143:131. 896 Kool W, Botvinick M (2018) Mental labour. Nature human behaviour 2:899-908. 897 Kool W. Shenhav A. Botvinick MM (2017) Cognitive control as cost-benefit decision making. 898 Lee WH, Rodrigue A, Glahn DC, Bassett DS, Frangou S (2019) Heritability and cognitive relevance of 899 structural brain controllability. Cerebral Cortex . 900 Li CT, Huang YZ, Bai YM, Tsai SJ, Su TP, Cheng CM (2019) Critical role of glutamatergic and gabaergic 901 neurotransmission in the central mechanisms of theta-burst stimulation. Human brain 902 mapping 40:2001-2009. 903 Liu YY, Slotine JJ, Barabási AL (2011) Controllability of complex networks. nature 473:167-173. 904 Liuzzi AG, Bruffaerts R, Peeters R, Adamczuk K, Keuleers E, De Deyne S, Storms G, Dupont P, 905 Vandenberghe R (2017) Cross-modal representation of spoken and written word meaning in left pars 906 triangularis. NeuroImage 150:292-307. 907 Lorch RF, Myers JL (1990) Regression analyses of repeated measures data in cognitive research. Journal of 908 Experimental Psychology: Learning, Memory, and Cognition 16:149. 909 Lüders H, Lesser R, Dinner D, Hahn J, Salanga V, Morris H (1985) The second sensory area in humans: 910 evoked potential and electrical stimulation studies. Annals of neurology 17:177-184. 911 Maier-Hein KH, Neher PF, Houde JC, Côté MA, Garyfallidis E, Zhong J, Chamberland M, Yeh FC, Lin 912 YC, Ji Q et al. (2017) The challenge of mapping the human connectome based on diffusion tractography. 913 Nature communications 8:1–13. 914 Martin RC, Cheng Y (2006) Selection demands versus association strength in the verb generation task. 915 Psychonomic bulletin & review 13:396-401. 916 Masson ME (1995) A distributed memory model of semantic priming. Journal of Experimental 917 Psychology: Learning, Memory, and Cognition 21:3. 918 Mattheiss SR, Levinson H, Graves WW (2018) Duality of function: activation for meaningless nonwords 919 and semantic codes in the same brain areas. Cerebral Cortex 28:2516-2524. 920 Medaglia JD (2019) Clarifying cognitive control and the controllable connectome. Wiley Interdisciplinary 921 Reviews: Cognitive Science 10:e1471. 922 Medaglia JD, Harvey DY, White N, Kelkar A, Zimmerman J, Bassett DS, Hamilton RH (2018a) Network 923 controllability in the inferior frontal gyrus relates to controlled language variability and susceptibility to
- tms. *Journal of Neuroscience* pp. 0092–17.

925 Medaglia JD, Huang W, Karuza EA, Kelkar A, Thompson-Schill SL, Ribeiro A, Bassett DS (2018b) 926 Functional alignment with anatomical networks is associated with cognitive flexibility. Nature human 927 behaviour 2:156-164. 928 Medaglia JD, Lynall ME, Bassett DS (2015) Cognitive network neuroscience. Journal of cognitive 929 neuroscience. 930 Milyavskaya M, Inzlicht M, Johnson T, Larson MJ (2019) Reward sensitivity following boredom and 931 cognitive effort: A high-powered neurophysiological investigation. Neuropsychologia 123:159-168. 932 Muldoon SF, Pasqualetti F, Gu S, Cieslak M, Grafton ST, Vettel JM, Bassett DS (2016) Stimulation-based 933 control of dynamic brain networks. PLoS computational biology 12:e1005076. 934 Musz E, Thompson-Schill SL (2017) Tracking competition and cognitive control during language 935 comprehension with multi-voxel pattern analysis. Brain and language 165:21-32 936 Naeser MA, Martin PI, Theoret H, Kobayashi M, Fregni F, Nicholas M, Tormos JM, Steven MS, Baker 937 EH, Pascual-Leone A (2011) Tms suppression of right pars triangularis, but not pars opercularis, improves 938 naming in aphasia. Brain and language 119:206-213. 939 Newman ME (2006) Modularity and community structure in networks. Proceedings of the national 940 academy of sciences 103:8577-8582. 941 Ni W, Constable R, Mencl W, Pugh K, Fulbright R, Shaywitz S, Shaywitz B, Gore J, Shankweiler D (2000) 942 An event-related neuroimaging study distinguishing form and content in sentence processing. Journal of 943 Cognitive Neuroscience 12:120-133. 944 Nozari N, Pinet S (2020) A critical review of the behavioral, neuroimaging, and electrophysiological 945 studies of co-activation of representations during word production. Journal of Neurolinguistics 53:100875. 946 Pasqualetti F, Gu S, Bassett DS (2019) Re: Warnings and caveats in brain controllability. 947 NeuroImage 197:586-588. 948 Pasqualetti F, Zampieri S, Bullo F (2014) Controllability metrics, limitations and algorithms for complex 949 networks. IEEE Transactions on Control of Network Systems 1:40-52. 950 Patankar SP, Kim JZ, Pasqualetti F, Bassett DS (2020) Path-dependent connectivity, not modularity, 951 consistently predicts controllability of structural brain networks. Network Neuroscience 4:1091–1121. 952 Patterson K, Nestor PJ, Rogers TT (2007) Where do you know what you know? the representation of 953 semantic knowledge in the human brain. Nature Reviews Neuroscience 8:976-987. 954 Piai V, Knight RT (2018) Lexical selection with competing distractors: Evidence from left temporal lobe 955 lesions. Psychonomic bulletin & review 25:710-717. 956 R Core Team (2016) R: A Language and Environment for Statistical Computing R Foundation for 957 Statistical Computing, Vienna, Austria. 958 Ridderinkhof KR, Van Den Wildenberg WP, Segalowitz SJ, Carter CS (2004) Neurocognitive mechanisms 959 of cognitive control: the role of prefrontal cortex in action selection, response inhibition, performance 960 monitoring, and reward-based learning. Brain and cognition 56:129-140. 961 Riès SK, Dhillon RK, Clarke A, King-Stephens D, Laxer KD, Weber PB, Kuperman RA, Auguste KI, 962 Brunner P, Schalk G et al. (2017) Spatiotemporal dynamics of word retrieval in speech production revealed 963 by cortical high-frequency band activity. Proceedings of the National Academy of 964 Sciences 114:E4530-E4538. 965 Rogalsky C, Hickok G (2009) Selective attention to semantic and syntactic features modulates sentence 966 processing networks in anterior temporal cortex. Cerebral Cortex 19:786-796. 967 Ruths J, Ruths D (2014) Control profiles of complex networks. Science 343:1373–1376. 968 Ryskin R, Levy RP, Fedorenko E (2020) Do domain-general executive resources play a role in linguistic 969 prediction? re-evaluation of the evidence and a path forward. Neuropsychologia 136:107258. 970 Saunders J, MacLeod MD (2006) Can inhibition resolve retrieval competition through the control of 971 spreading activation? Memory & Cognition 34:307-322. 972 Savine AC, Braver TS (2010) Motivated cognitive control: reward incentives modulate preparatory neural 973 activity during task-switching. 974 Schiff SJ (2012) Neural control engineering: the emerging intersection between control theory and 975 neuroscience MIT Press. 976 Shahdloo M, Çelik E, Çukur T (2019) Biased competition in semantic representation during natural visual 977 search. NeuroImage p. 116383.

- 978 Shenhav A, Botvinick MM, Cohen JD (2013) The expected value of control: an integrative theory of 979 anterior cingulate cortex function. Neuron 79:217-240. 980 Shenhav A, Musslick S, Lieder F, Kool W, Griffiths TL, Cohen JD, Botvinick MM (2017) Toward a 981 rational and mechanistic account of mental effort. Annual review of neuroscience 40:99-124. 982 Snyder HR, Banich MT, Munakata Y (2011) Choosing our words: retrieval and selection processes recruit 983 shared neural substrates in left ventrolateral prefrontal cortex. Journal of cognitive 984 neuroscience 23:3470-3482. 985 Snyder HR, Banich MT, Munakata Y (2014) All competition is not alike: Neural mechanisms for resolving 986 underdetermined and prepotent competition. Journal of Cognitive Neuroscience 26:2608-2623. 987 Snyder HR, Munakata Y (2008) So many options, so little time: The roles of association and competition in 988 underdetermined responding. Psychonomic Bulletin & Review 15:1083-1088. 989 Souza MJ, Donohue SE, Bunge SA (2009) Controlled retrieval and selection of action-relevant knowledge 990 mediated by partially overlapping regions in left ventrolateral prefrontal cortex. Neuroimage 46:299-307. 991 Stagg CJ, Wylezinska M, Matthews PM, Johansen-Berg H, Jezzard P, Rothwell JC, Bestmann S (2009) 992 Neurochemical effects of theta burst stimulation as assessed by magnetic resonance spectroscopy. Journal 993 of neurophysiology 101:2872–2877. 994 Stiso J, Ankit K, Menara T, Kahn A, Stein J, Das S, Gorniak R, Tracy J, Litt B, Davis K, Pasqualetti F, 995 Lucas T, Bassett D (2019) White matter network architecture guides direct electrical stimulation through 996 optimal state transitions. Cell. 997 Suweis S, Tu C, Rocha RP, Zampieri S, Zorzi M, Corbetta M (2019) Brain controllability: Not a slam dunk 998 vet. NeuroImage 200:552-555. 999 Tang E, Giusti C, Baum GL, Gu S, Pollock E, Kahn AE, Roalf DR, Moore TM, Ruparel K, Gur RC et al. 1000 (2017) Developmental increases in white matter network controllability support a growing diversity of 1001 brain dynamics. Nature communications 8:1-16. 1002 Thomas C, Frank QY, Irfanoglu MO, Modi P, Saleem KS, Leopold DA, Pierpaoli C (2014) Anatomical 1003 accuracy of brain connections derived from diffusion mri tractography is inherently limited. Proceedings of 1004 the National Academy of Sciences 111:16574–16579. 1005 Thompson-Schill SL, D'Esposito M, Aguirre GK, Farah MJ (1997) Role of left inferior prefrontal cortex in 1006 retrieval of semantic knowledge: a reevaluation. Proceedings of the National Academy of 1007 Sciences 94:14792-14797. 1008 Tiberi L, Favaretto C, Innocenti M, Bassett DS, Pasqualetti F (2017) Synchronization patterns in networks 1009 of kuramoto oscillators: A geometric approach for analysis and control. arXiv preprint arXiv:1709.06193. 1010 Torres J, Drebing D, Hamilton R (2013) Tms and tdcs in post-stroke aphasia: Integrating novel treatment 1011 approaches with mechanisms of plasticity. Restorative neurology and neuroscience 31:501-515. 1012 Trippe J, Mix A, Aydin-Abidin S, Funke K, Benali A (2009) Theta burst and conventional low-frequency 1013 rtms differentially affect gabaergic neurotransmission in the rat cortex. Experimental brain 1014 research 199:411 1015 Tu C, Rocha RP, Corbetta M, Zampieri S, Zorzi M, Suweis S (2018) Warnings and caveats in brain 1016 controllability. NeuroImage 176:83-91. 1017 Vasishth S, Nicenboim B, Engelmann F, Burchert F (2019) Computational models of retrieval processes in 1018 sentence processing. Trends in cognitive sciences 23:968-982. 1019 Vigneau M, Beaucousin V, Herve PY, Duffau H, Crivello F, Houde O, Mazoyer B, Tzourio-Mazoyer N 1020 (2006) Meta-analyzing left hemisphere language areas: phonology, semantics, and sentence processing. 1021 Neuroimage 30:1414-1432 1022 Wagner AD, Paré-Blagoev EJ, Clark J, Poldrack RA (2001) Recovering meaning: left prefrontal cortex 1023 guides controlled semantic retrieval. Neuron 31:329-338. 1024 Whitney C, Weis S, Krings T, Huber W, Grossman M, Kircher T (2009) Task-dependent modulations of 1025 prefrontal and hippocampal activity during intrinsic word production. Journal of Cognitive 1026 Neuroscience 21:697-712. 1027 Wu-Yan E, Betzel RF, Tang E, Gu S, Pasqualetti F, Bassett DS (2018) Benchmarking measures of network 1028 controllability on canonical graph models. Journal of Nonlinear Science pp. 1–39.
- 1029Yeh FC, Wedeen VJ, Tseng WYI (2011) Estimation of fiber orientation and spin density distribution by1030diffusion deconvolution. Neuroimage 55:1054–1062.

1031Yoshida M, Dickey MW, Sturt P (2013) Predictive processing of syntactic structure: Sluicing and ellipsis in1032real-time sentence processing. Language and Cognitive Processes 28:272–302.

# **Tables for**

# Language Tasks and the Network Control Role of the Left Inferior Frontal Gyrus

1040 Key

1041

1034

1035

1036

1037 1038 1039

1042	TMS =	Transcranial	Magnetic	Stimulation
------	-------	--------------	----------	-------------

1043 LIFG = Left Inferior Frontal Gyrus

1044 Boundary = LIFG ranked boundary controllability

1045 Modal = LIFG ranked modal controllability

1046 Session = effect of session (Pre-TMS vs. Post-TMS). Baseline is the reference condition.

1047 Stimulation = stimulation condition (active vs. sham). Sham is the reference condition.

Task = effect of task (sentence completion vs. verb generation). Sentence completion is the reference condition.

1050 Entropy =continuous effect of entropy. Higher entropy is associated with greater selection 1051 demands.

Association = continuous effect of association. Higher association is associated with lower
 selection demands.

All tables report the model estimates and parameter significance tests using Satterthwaite's approximation. All mixed effects models included a random intercept for trials nested within subjects. Significant p-values are denoted by **bold** text. The dependent variable in all models is the log of response times during the tasks. In all models, CI = 95% confidence interval for the fixed effects estimates.

1054

1070

1071

**Table 1:** Total trial rejection percentages for each session, task, and group.

_	Session	Task	Group	Trial rejection percentage
	Pre-TMS	Sentence Completion	Active	7.20
		Sentence Completion	Sham	10.125
	Pre-TMS	Verb Generation	Active	13.44
		Verb Generation	Sham	10.500
	Post-TMS	Sentence Completion	Active	3.36
		Sentence Completion	Sham	4.500
	Post-TMS	Verb Generation	Active	8.48
		Verb Generation	Sham	8.125

# Table 2: Transcranial magnetic stimulation (TMS) effects depend on left inferior frontal gyrus (LIFG) boundary controllability across both tasks.

Predictors	Estimates	CI	df	Statistic	р
(Intercept)	7.233	7.206 - 7.260	5609.031	522.049	<0.001
Stimulation	-0.030	-0.065 - 0.005	5608.513	-1.706	0.088
Session	0.024	-0.013 - 0.061	5770.483	1.256	0.209
Boundary	-0.002	-0.0030.001	5675.063	-2.913	0.004
Stimulation * Session	-0.035	-0.083 - 0.012	5775.291	-1.464	0.143
Stimulation * Boundary	0.004	0.002 - 0.005	5650.508	5.440	<0.001
Session * Boundary	0.001	-0.001 - 0.002	5824.119	0.718	0.473
Stimulation * Session * Boundary	-0.003	-0.0040.001	5809.284	-2.629	0.009

# **Table 3:** Selection costs differ across the tasks at baseline.

Predictors	Estimates	CI	df	Statistic	р
(Intercept)	6.846	6.818 - 6.874	3676.921	481.311	< 0.001
Task	0.580	0.541 - 0.619	2899.285	29.214	<0.001
Selection	0.266	0.226 - 0.305	3638.267	13.177	<0.001
Task * Selection	-0.180	-0.2360.124	3615.092	-6.253	<0.001

# **Table 4:** Retrieval costs differ across the tasks at baseline.

Predictors	Estimates	CI	df	Statistic	р
(Intercept)	6.925	6.897 - 6.952	3676.981	489.575	<0.001
Task	0.432	0.393 - 0.471	2893.463	21.924	<0.001
Retrieval	0.109	0.070 - 0.149	3644.480	5.394	<0.001
Task * Retrieval	0.122	0.066 - 0.179	3628.856	4.228	<0.001

1088	Table 5: LIFG boundary controllability moderates baseline selection costs in sentence
1089	completion.

Predictors	Estimates	CI	df	Statistic	р
(Intercept)	6.998	6.938 - 7.059	44.004	226.817	<0.001
Boundary	0.001	-0.001 - 0.004	40.167	1.284	0.199
Entropy	0.155	0.135 - 0.175	1833.751	15.293	<0.001
Boundary * Entropy	0.001	0.000 - 0.002	1826.762	3.150	0.002

**Table 6:** LIFG modal controllability does <u>not</u> moderate baseline selection costs in sentence
 completion.

Predictors	Estimates	CI	df	Statistic	р
(Intercept)	7.422	7.353 - 7.490	53.769	211.851	<0.001
Modal	0.005	-0.004 - 0.013	42.251	1.106	0.269
Entropy	0.221	0.173 - 0.269	1757.626	8.990	<0.001
Modal * Entropy	-0.006	-0.012 - 0.000	1736.954	-1.862	0.063

# **Table 7:** LIFG boundary controllability does <u>not</u> moderate baseline retrieval costs in verb generation.

Predictors	Estimates	CI	df	Statistic	р
(Intercept)	7.521	7.453 - 7.588	52.950	217.976	<0.001
Boundary	0.001	-0.001 - 0.003	43.007	0.772	0.440
Association	-0.666	-0.7740.557	1757.621	-12.029	<0.001
Boundary * Association	-0.001	-0.005 - 0.003	1747.836	-0.543	0.587

# **Table 8:** LIFG modal controllability does <u>not</u> moderate retrieval costs in verb generation.

Predictors	Estimates	CI	df	Statistic	р
(Intercept)	7.521	7.453 - 7.588	52.957	218.841	<0.001
Modal	0.003	-0.005 - 0.012	43.028	0.826	0.409

Association		-0.665 -	0.773 –	-0.556	1757.60	6 -12	2.018	<0.001
Modal * Assoc	ciation	0.003	-0.011 –	0.017	1741.31	5 0.	383	0.702
Table 9: Per         Predictors       Est	formance	e on sent CI	ence co	mpletic	on slows Statistic	in the	e sham	n group ac
(Intercept) 6	5.993 6	.959 – 7.0	27 14	74.687	398.217	<0.0	01	
Session (	).072 0	.026 – 0.1	17 76	53.964	3.057	0.00	02	
Predictors Est (Intercept) 7	timates 7.482 7	<i>CI</i> 7.448 – 7.5	516 14	<i>df</i> 409.983	<u>Statistia</u> 432.892	cnang <u>c p</u> 2 < <b>0.</b>	2 001	ie snam gi
Session (	0.022	0.064 0	021 7	11 236	0 007	0.3	10	
sham group.	VIS signi Estimates	s C	speeds	df	stance on	tistic	p	
(Intercept)	6.993	6.960 -	7.027	3802.5	23 406	.959	<0.001	1
Stimulation	-0.025	-0.068 -	- 0.018	3800.2	11 -1.	139	0.255	
Session	0.071	0.028 -	0.115	2000.4	52 3.2	204	0.001	
Stimulation * Session	-0.092	-0.148 -	-0.036	1986.2	31 -3.	245	0.001	
Table 12: TM Predictors	S does no Estimates	ot signific	antly af	fect perf df	ormance Stati	on ver	rb gene	eration.
(Intercept)	7.477	7.385 -	7.570	46.143	3 158.	547 <	<0.001	
Stimulation	-0.022	-0.141 -	0.096	46.311	-0.3	67	0.713	
Session	-0.018	-0.062 -	0.025	3637.31	0 -0.8	329	0.407	
Stimulation *	0.009	-0.047	0.065	3637 7/	14 03	18	0 750	

to the

1114

Session

# **Table 13:** LIFG boundary controllability moderates the TMS effect in sentence completion.

Predictors	Estimates	CI	df	Statistic	р
(Intercept)	6.992	6.958 - 7.025	3799.047	407.782	<0.001
Stimulation	-0.025	-0.068-0.018	3797.234	-1.157	0.247
Session	0.070	0.027 - 0.114	1993.170	3.156	0.002
Boundary	-0.001	-0.003 - 0.000	3799.107	-1.837	0.066
Stimulation * Session	-0.088	-0.1440.033	1982.086	-3.123	0.002
Stimulation * Boundary	0.003	0.002 - 0.005	3796.519	3.775	<0.001
Session * Boundary	-0.000	-0.002 - 0.002	2019.562	-0.196	0.845
Stimulation * Session * Boundary	-0.002	-0.0050.000	1996.384	-1.998	0.046

Predictors	Estimates	CI	$d\!f$	Statistic	р
(Intercept)	7.478	7.445 - 7.512	3615.889	434.784	<0.001
Stimulation	-0.022	-0.065 - 0.021	3618.283	-0.993	0.321
Session	-0.019	-0.063 - 0.024	1894.733	-0.871	0.384
Boundary	-0.001	-0.003 - 0.000	3623.787	-1.571	0.116
Stimulation * Session	0.010	-0.045 - 0.066	1902.848	0.368	0.713
Stimulation * Boundary	0.003	0.001 - 0.005	3624.616	3.540	<0.001
Session * Boundary	0.001	-0.001 - 0.002	1942.348	0.558	0.577
Stimulation * Session * Boundary	-0.002	-0.004 - 0.001	1940.439	-1.455	0.146

# **Table 14:** LIFG boundary controllability does not interact with TMS in verb generation.

1	1	1	9







# eNeuro Accepted Manuscript

**Response Time** 







