

# 1 Functional and anatomical connectivity predict 2 brain stimulation's mnemonic effects

3 **Abbreviated Title:** Mediators of stimulation's effect on memory

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26

## 27 Abstract

28 Closed-loop direct brain stimulation is a promising tool for modulating neural activity and behav-  
29 ior. However, it remains unclear how to optimally target stimulation to modulate brain activity in  
30 particular brain networks that underlie particular cognitive functions. Here, we test the hypothesis  
31 that stimulation's behavioral and physiological effects depend on the stimulation target's anatomi-  
32 cal and functional network properties. We delivered closed-loop stimulation as 47 neurosurgical  
33 patients studied and recalled word lists. Multivariate classifiers, trained to predict momentary  
34 lapses in memory function, triggered stimulation of the lateral temporal cortex (LTC) during the  
35 study phase of the task. We found that LTC stimulation specifically improved memory when  
36 delivered to targets near white matter pathways. Memory improvement was largest for targets  
37 near white matter that also showed high functional connectivity to the brain's memory network.  
38 These targets also reduced low-frequency activity in this network, an established marker of suc-  
39 cessful memory encoding. These data reveal how anatomical and functional networks mediate  
40 stimulation's behavioral and physiological effects, provide further evidence that closed-loop LTC  
41 stimulation can improve episodic memory, and suggest a method for optimizing neuromodulation  
42 through improved stimulation targeting.

## 43 Introduction

44 Direct electrical stimulation of the human brain can manipulate circuits underlying perception,  
45 cognition, and action (Siddiqi et al., 2022; Scangos et al., 2021b). Such stimulation has been used  
46 to treat network syndromes of brain dysfunction, suggesting that stimulation influences a broader  
47 network of brain regions beyond the stimulated location (Mayberg et al., 2005; Scangos et al., 2021a;  
48 Limousin et al., 1998; Bouthour et al., 2019; Deuschl et al., 2006; Geller et al., 2017; Jobst et al., 2017;  
49 Lozano and Lipsman, 2013). Stimulation can also modulate behaviors, such as learning and  
50 memory, that depend on the coordinated activity of a network of brain regions (Mankin and Fried,  
51 2020; Das and Menon, 2021; Voytek and Knight, 2015; Keerativittayayut et al., 2018; Staresina and  
52 Wimber, 2019).

53 Although increasingly used as a therapeutic and experimental tool, variability in outcomes  
54 poses a critical challenge, in part because stimulation's mechanisms of action remain poorly  
55 understood. Theoretical accounts evolved from models of local disruption of pathological activity  
56 (Benabid et al., 2004) to modulation of the broader network of areas connected to the stimulated  
57 location (Ashkan et al., 2017; McIntyre and Hahn, 2010). If stimulation's effects are best understood  
58 at the network level, perhaps variability in individual network structure can explain the variability  
59 in physiological and behavioral outcomes.

60 In support of this idea, applying stimulation to gray matter, the gray-white matter boundary,  
61 or specific white matter fibers determines the spread of physiological effects through the network  
62 (Pault et al., 2022; Solomon et al., 2018). Compared to gray matter stimulation, white matter  
63 stimulation leads to more broadly distributed excitation in downstream areas (Crocker et al., 2021;  
64 Pault et al., 2022; Mohan et al., 2020). White matter pathways also constrain stimulation's down-  
65 stream functional effects (Lujan et al., 2013; Khambhati et al., 2019; Stiso et al., 2019). Behaviorally,  
66 stimulation of white matter has led to remission in depression (Mayberg et al., 2005), slowed  
67 cognitive decline in Alzheimer's (Lozano et al., 2016; Hamani et al., 2008), and enhanced memory  
68 in epilepsy (Suthana et al., 2012; Mankin et al., 2021; Titiz et al., 2017).

69 In addition to the brain's anatomical architecture, research shows that functional architecture  
70 also mediates the spread and persistence of stimulation's physiological effects (Keller et al., 2011;  
71 Fox et al., 2020; Fox et al., 2014; Keller et al., 2018). Previous work further suggests this relation

72 to be frequency-specific. For example, stimulating targets in the medial temporal lobe leads to  
73 greater downstream changes in low-frequency (5-13 Hz) activity in brain regions that are strongly  
74 connected, at low-frequencies, to the stimulated site (Solomon et al., 2018). There are a variety  
75 of cognitive functions, including episodic memory, that have been linked to modulation of low-  
76 frequency activity (Colgin, 2013; Burke et al., 2013; Solomon et al., 2017; Donoghue et al., 2020;  
77 Koster and Gruber, 2022; Griffiths et al., 2021). Therefore, these physiological findings suggest that  
78 stimulating targets with strong low-frequency network connectivity could reliably modulate such  
79 behaviors, although this idea has yet to be tested.

80 We hypothesized that anatomical and functional characteristics of the stimulation target rep-  
81 resent key variables that control the effect of stimulation on the brain's memory network. We  
82 applied stimulation in closed-loop in 47 patients while they participated in an episodic memory  
83 task (free recall). We stimulated 57 targets located in the lateral temporal cortex (LTC), with the  
84 timing of stimulation determined by multivariate classification of neural activity during the en-  
85 coding phase of the memory task. Using patient-specific data, we characterized each stimulation  
86 target based on its proximity to the nearest white matter pathway, as well as its low-frequency  
87 resting-state functional connectivity with the the brain's memory-encoding network. We found  
88 that closed-loop LTC stimulation improves memory performance relative to random stimulation,  
89 extending prior evidence that LTC stimulation modulates episodic memory (Ezyat et al., 2018;  
90 Kucewicz et al., 2018). Further, we reveal that stimulation target proximity to white matter and  
91 functional connectivity predict both stimulation's effects on memory performance and changes in  
92 rhythmic low-frequency activity involved in successful memory encoding.

## 93 **Experimental Procedures**

### 94 **Participants**

95 Forty-seven patients undergoing intracranial electroencephalographic monitoring as part of clin-  
96 ical treatment for drug-resistant epilepsy were recruited to participate in this study. In total,  $N = 57$   
97 brain locations were stimulated: 38 patients were stimulated in one location, 8 patients were stim-  
98 ulated in two separate locations, and 1 patient was stimulated in three separate locations. Only one  
99 location was stimulated per session. Of the current dataset, data from 14 patients were included

100 in an earlier publication (Ezzyat et al., 2018). All of the presently reported analyses and results are  
101 novel.

102 Data were collected as part of a multi-center project designed to assess the effects of electri-  
103 cal stimulation on memory-related brain function. Data were collected at the following centers:  
104 University of Texas Southwestern Medical Center (Dallas, TX), Dartmouth-Hitchcock Medical Cen-  
105 ter (Lebanon, NH), Thomas Jefferson University Hospital (Philadelphia, PA), Emory University  
106 Hospital (Atlanta, GA), Mayo Clinic (Rochester, MN), Hospital of the University of Pennsylva-  
107 nia (Philadelphia, PA), and Columbia University Medical Center (New York, NY). The research  
108 protocol was approved by the IRB at each hospital and informed consent was obtained from  
109 each participant. Electrophysiological data were collected from electrodes implanted subdurally  
110 (grid/strip configurations) on the cortical surface and/or electrodes within the brain parenchyma  
111 (depth electrodes). The clinical team determined the placement of the electrodes based on the  
112 epileptogenic monitoring needs of the patient.

113 **Anatomical localization**

114 Cortical surface regions were delineated on pre-implant whole brain volumetric T1-weighted MRI  
115 scans using Freesurfer (Fischl et al., 2004) according to the Desikan-Kiliany atlas (Desikan et al.,  
116 2006). Whole brain and high resolution medial temporal lobe volumetric segmentation was also  
117 performed using the T1-weighted scan and a dedicated hippocampal coronal T2-weighted scan  
118 with Advanced Normalization Tools (ANTS) (Avants et al., 2008) and Automatic Segmentation  
119 of Hippocampal Subfields (ASHS) multi-atlas segmentation methods (Yushkevich et al., 2015).  
120 Coordinates of the radiodense electrode contacts were derived from a post-implant CT and then  
121 registered with the MRI scans using ANTS. Subdural electrode coordinates were further mapped  
122 to the cortical surfaces using an energy minimization algorithm (Dykstra et al., 2012). Two neuro-  
123 radiologists reviewed cross-sectional images and surface renderings to confirm the output of the  
124 automated localization pipeline. Stimulation targets that were localized to the left inferior, middle,  
125 and superior temporal gyri were classified as LTC. For region of interest analyses, electrodes were  
126 assigned to regions using Freesurfer atlas labels (IFG: inferior frontal gyrus; MFG: middle frontal  
127 gyrus; SFG: superior frontal gyrus; MTLC: medial temporal lobe cortex; HIPP: hippocampus; ITG:

128 inferior temporal gyrus; MTG: middle temporal gyrus; STG: superior temporal gyrus; IPC: inferior  
129 parietal cortex; SPC: superior parietal cortex; OC: occipital lobe).

130 **Verbal memory task**

131 Across participants, data were collected from two behavioral tasks: standard delayed free recall  
132 and categorized delayed free recall. In both tasks, participants were instructed to study lists of  
133 words for a later memory test; no explicit encoding task was used. Lists were composed of 12  
134 words presented in either English or Spanish, depending on the participant's native language.  
135 In the standard free recall task, words were selected randomly from a pool of common nouns  
136 ([https://memory.psych.upenn.edu/Word\\_Pools](https://memory.psych.upenn.edu/Word_Pools)). In the categorized free recall task, the word  
137 pool was constructed from 25 semantic categories (e.g. fruit, furniture, office supplies). Each list  
138 of 12 items in the categorized version of the task consisted of four words drawn from each of  
139 three categories. Overall,  $N = 19$  participated in standard free recall only;  $N = 26$  participated in  
140 categorized free recall only; and  $N = 2$  participated in both free and categorized recall (in separate  
141 sessions).

142

143 Immediately following the final word in each list, participants performed a distractor task (to  
144 attenuate the recency effect in memory, length = 20 seconds) consisting of a series of arithmetic  
145 problems of the form  $A+B+C=??$ , where A, B and C were randomly chosen integers ranging from  
146 1-9. Following the distractor task participants were given 30 seconds to verbally recall as many  
147 words as possible from the list in any order; vocal responses were digitally recorded and later  
148 manually scored for analysis. Each session consisted of 25 lists of this encoding-distractor-recall  
149 procedure.

150 **EEG recording and analysis**

151 Electrophysiological recording and stimulation was conducted using a variety of systems across the  
152 sites over which the project was conducted. Recording and stimulation equipment included clinical  
153 EEG systems (Nihon Kohden EEG-1200, Natus XLTek EMU 128 or Grass Aura-LTM64), equipment  
154 from Blackrock Microsystems, as well as the External Neural Stimulator (ENS) (Medtronic, Inc.).

155 Data were sampled at 500, 1000, or 1600 Hz (depending on the clinical site). During the sessions,  
156 a laptop recorded behavioral responses (vocalizations, key presses), synchronized to the recorded  
157 EEG via transmitted network packets.

158 Intracranial electrophysiological data were filtered to attenuate line noise (5 Hz band-stop  
159 fourth order Butterworth, centered on 60 Hz). We referenced the data using a bipolar montage  
160 (Burke et al., 2013) by identifying all pairs of immediately adjacent contacts on every depth, strip  
161 and grid and taking the difference between the signals recorded in each pair. The resulting bipolar  
162 timeseries was treated as a virtual electrode and used in all subsequent analysis. For the purposes  
163 of anatomical localization, we used the midpoint of the bipolar pair as the location for this virtual  
164 electrode. We used the same midpoint approach to localize stimulation targets and to measure  
165 stimulation target distance to white matter (see below).

## 166 **Multivariate classification of memory**

167 We performed spectral decomposition (8 frequencies from 6-175 Hz, logarithmically-spaced; Mor-  
168 let wavelets; wave number = 5) for 1366 ms epochs from 0 to to 1366 ms relative to word onset.  
169 Mirrored buffers (length = 1365 ms) were included before and after the interval of interest to avoid  
170 convolution edge effects. The resulting time-frequency data were then log-transformed, averaged  
171 over time, and z-scored within session and frequency band across word presentation events. For a  
172 subset of participants, we also performed the same spectral decomposition procedure on record-  
173 only data from the memory recall phase of each list. These data were then used in addition to the  
174 encoding data to train the classifier (Kragel et al., 2017). To do so, we computed spectral power  
175 for the 500 ms interval preceding a response vocalization, as well as during unsuccessful periods  
176 of memory search (the first 500 ms of any 2500 ms interval in which no recall response was made).  
177 For both trial types (correct vocalizations and unsuccessful search periods), we further stipulated  
178 that no vocalization onsets occurred in the preceding 2000 ms.

179 Our closed-loop stimulation approach was based on using individualized memory classifiers  
180 to control the timing of stimulation in response to brain activity. Thus, after collecting at least  
181 three record-only sessions from an individual patient, we then used the data as input to a logistic  
182 regression classifier that would trigger closed-loop stimulation during the later stimulation ses-

183 sion(s). To build the classifier, we used patterns of brain activity collected during record-only  
184 sessions and trained the classifier to discriminate words that were recalled vs. not recalled. The  
185 input features were spectral power at the eight analyzed frequencies  $\times N$  electrodes (Fig 1A).  
186 We used L2-penalization to prevent overfitting (Hastie et al., 2001) and set the penalty parameter  
187 (C) to  $2.4 \times 10^{-4}$  based on the optimal penalty parameter calculated across our large pre-existing  
188 dataset of free-recall participants (Kragel et al., 2017; Ezzyat et al., 2018). We weighted the penalty  
189 parameter separately for each participant in inverse proportion to their number of recalled and  
190 not recalled words; this was done so that the model would learn equally from both classes (Hastie  
191 et al., 2001). Classification analyses were programmed using either the Matlab implementation of  
192 the LIBLINEAR library (Fan et al., 2008) or the Python library scikit-learn (Pedregosa et al., 2011).

193 For the Closed-loop group (34 participants,  $N = 40$  stimulation targets), classifiers were trained  
194 using the true mapping of features (spectral power  $\times$  electrodes) to recall outcomes. In contrast,  
195 for the Random group (13 participants,  $N = 17$  stimulation targets), a technical error in labeling  
196 features during classifier training led to classifiers that were trained on permuted data, eliminating  
197 the true mapping between neural activity on each trial and recall outcomes. This provided a natural  
198 experiment for testing whether the closed-loop nature of stimulation enhanced the efficacy of LTC  
199 stimulation.

200 To assess the importance of individual features to the classifier's performance, we calculated a  
201 forward model (Haufe et al., 2014):

$$A = \frac{\Sigma_x \mathbf{w}}{\sigma_{\hat{y}}^2}$$

202 where  $\Sigma_x$  is the data covariance matrix,  $\mathbf{w}$  is the vector of feature weights from the trained  
203 classifier, and  $\sigma_{\hat{y}}^2$  is the variance of the logit-transformed classifier outputs for all recalled/not  
204 recalled events  $\hat{y}$ . Positive values in  $A$  suggest a positive relation between power for a given feature  
205 and successful memory recall. We computed  $A$  separately for each participant (averaging features  
206 within anatomical regions of interest based on the Freesurfer labels derived from anatomical  
207 localization of electrodes) before conducting across-participant statistical tests (Fig 1D).

208 **Closed-loop stimulation**

209 At the start of each stimulation session, we determined the safe amplitude for stimulation using  
210 a mapping procedure in which stimulation was applied at 0.5 mA while a neurologist monitored  
211 for afterdischarges. This procedure was repeated, incrementing the amplitude in steps of 0.5 mA,  
212 up to a maximum of 1.5 mA for depth contacts and 3.5 mA for cortical surface contacts. These  
213 maximum amplitudes were chosen to be below the afterdischarge threshold and below accepted  
214 safety limits for charge density (Shannon, 1992). For each stimulation session, we passed electrical  
215 current through a single pair of adjacent electrode contacts. The locations of implanted electrodes  
216 were determined strictly by the monitoring needs of the clinicians (recording sites depicted in  
217 Figure 1B). We therefore used a combination of anatomical and functional information to select  
218 stimulation sites, prioritizing (if available) targets in the middle temporal gyrus (stimulation  
219 targets depicted in Figure 2A). This choice was guided by prior work identifying the middle  
220 temporal gyrus as an effective target for modulating memory with stimulation (Ezzyat et al., 2018;  
221 Kucewicz et al., 2018). Stimulation was delivered using charge-balanced biphasic rectangular  
222 pulses (pulse width = 300  $\mu$ s) at either 50, 100 or 200 Hz frequency (a single frequency was chosen  
223 for each subject), and was applied for 500 ms in response to classifier-detected poor memory states  
224 (see below). Participants performed one practice list followed by 25 task lists: lists 1-3 were used  
225 as a baseline for normalizing the classifier; lists 4-25 consisted of 11 lists each of Stim and NoStim  
226 conditions, randomly interleaved. NoStim lists were identically structured to Stim lists, except  
227 that stimulation was never delivered in response to classifier output.

228 To determine (in actuality) how well the classifier predicted recalled and forgotten words in a  
229 given participant's stimulation session, we again used AUC. We used the true classifier outputs  
230 and true recall outcomes from the NoStim lists to calculate the classifier generalization AUC for  
231 the stimulation sessions. To generate the corresponding receiver operating characteristic curves  
232 for visualization (Figure 1C), we modeled the classifier outputs for recalled and not recalled words  
233 using signal detection theory (Wixted, 2007). We did this by using the classifier outputs to estimate  
234 the mean and variance of hypothetical (normal) distributions of memory strength for recalled and  
235 not recalled words. We then generated a curve relating true and false positive rates by varying the  
236 assumed decision criterion (Wixted, 2007).

237 **Analysis of memory performance**

238 All participants completed at least three sessions of the record-only task (for purposes of classi-  
239 fier training) and at least one session of the stimulation task. For the stimulation session(s) we  
240 calculated stimulation's effect on recall performance as follows:

$$\Delta = \frac{R_S - R_{NS}}{R_{NS}} \times 100$$

241 where  $R_S$  is the average recall for stimulated lists and  $R_{NS}$  is the average recall for non-stimulated  
242 lists. Because the first three lists of every stimulation session were always non-stimulated (used  
243 for normalization of the classifier input features for that session), we excluded these lists from  
244 the calculation of  $R_{NS}$  to avoid introducing a temporal order confound (Ezzyat et al., 2018). All  
245 participants were required to demonstrate a minimum  $R_{NS} = 8.33\%$  (1 out of 12 words per list) for  
246 inclusion in the sample.

247 **Calculation of stimulation target distance to white matter**

248 Using Freesurfer to segment patients' T1 MRI scan, we identified white-matter vertex locations,  
249 then calculated the distance between the stimulation location (midpoint of the bipolar pair) and  
250 the nearest white matter vertex. These distances were then split into thirds in order to categorize  
251 stimulation sites as Near, Mid, or Far relative to the nearest white matter (Solomon et al., 2018;  
252 Mohan et al., 2020).

253 **Calculation of stimulation target node strength**

254 We adapted a previously reported method for calculating the resting-state functional connectivity  
255 between channels using the MNE-Python software package (Gramfort et al., 2014; Solomon et al.,  
256 2018). We extracted data from non-task periods of the record-only sessions of each patient and  
257 used the data to calculate the coherence between each pair of bipolar channels in the patient's  
258 montage. The coherence ( $C_{xy}$ ) between two signals is the normalized cross-spectral density. This  
259 measure reflects the consistency of phase differences between signals at two electrodes, weighted  
260 by the correlated change in spectral power at both sites:

$$C_{xy} = \left| \frac{S_{xy}}{S_{xx}S_{yy}} \right|$$

261 where  $S_{xy}$  is the cross-spectral density between signals at electrodes  $x$  and  $y$ ;  $S_{xx}$  and  $S_{yy}$  are the  
262 auto-spectral densities at each electrode. We used the multitaper method to estimate spectral  
263 density (Bronez, 1992). We used a time-bandwidth product of 4 and a maximum of 8 tapers  
264 (tapers with spectral energy < 0.9 were removed), computing coherence for frequencies between  
265 5 and 13 Hz. We computed inter-electrode coherence within non-overlapping 1-s windows of  
266 data collected during a 10-second baseline (countdown) period that occurred at the start of each  
267 word list. The resulting coherence values between each pair of electrodes were then regressed  
268 on the Euclidean distance between each pair of electrodes, to account for the correlation between  
269 inter-electrode coherence and distance (Solomon et al., 2018). This distance-residualized measure  
270 of coherence was then used in the node-strength calculation. We repeated this entire procedure  
271 for calculating high-frequency functional connectivity in the 45-90 Hz range.

## 272 **Analysis of physiological effects of stimulation**

273 To assess the effect of LTC stimulation on neural activity, we analyzed recording channels (i.e.  
274 those that were not stimulated) and we compared stimulation-evoked spectral power separately  
275 at low and high frequencies. We first excluded electrodes exhibiting non-physiological post-  
276 stimulation artifacts (such as amplifier saturation/relaxation) using three different measures of the  
277 EEG timeseries before and after stimulation. We compared intervals before and after stimulation  
278 for changes in variance using an  $F$ -test and for changes in signal amplitude using a  $t$ -test. We  
279 additionally fit a polynomial function to the timeseries before and after each stimulation event and  
280 used a  $t$ -test to compare the resulting parameter estimates for the quadratic term. We calculated  
281 these three measures using the signal from -400 ms to -100 ms relative to stimulation onset and  
282 100 ms to 400 ms relative to stimulation offset. In order to select statistical thresholds for each  
283 measure, we conducted the same analysis on each participant's record-only data. We then selected  
284  $P$ -value thresholds associated with a 5% detection rate in the record-only data (i.e. false positives).  
285 Any channel that was significant on any of the three measures was excluded from analysis.

286 To measure stimulation's effect on low-frequency power, we extracted spectral power from

287 -600 ms to -100 ms relative to stimulation onset and 100 ms to 600 ms relative to stimulation  
288 offset. We used Morlet wavelets (wave number = 5) to estimate spectral power for the same set  
289 of frequencies used to train the classifier with buffers to eliminate edge artifacts. The resulting  
290 spectral power estimates were then z-scored within each frequency, separately for each session. We  
291 then averaged power within each frequency across the time dimension for each pre-stimulation  
292 period and for each matched post-stimulation period. We then subtracted the pre-stimulation  
293 data from the post-stimulation data to yield a distribution of change in spectral power for each  
294 electrode.

295 We compared the distribution of power changes for stimulation events to the power changes  
296 from matched intervals on NoStim lists. To do so, we calculated spectral power using identical pa-  
297 rameters. However, because there were no actual stimulation events in NoStim lists, we generated  
298 a synthetic distribution of stimulation onset times by extracting the lag (in milliseconds) between  
299 each word onset and stimulation event in Stim lists, and sampling randomly from that distribution  
300 of onset times to determine when to extract data relative to word onset events in NoStim lists.

301 Finally, we used an independent samples *t*-test to compare the distribution of Stim list power  
302 differences to the distribution of NoStim list power differences within each electrode. The resulting  
303 distribution of *t*-statistics was then averaged across electrodes to estimate the stimulation-evoked  
304 change in power (Figure 5A). We then averaged these values separately within clusters of low  
305 and high frequencies that significantly predicted memory performance (based on classifier feature  
306 importance, Figure 1D).

## 307 Statistics

308 Data are presented as mean  $\pm$  standard error of the mean; scatterplots show the standard error of  
309 the estimate. All statistical comparisons were conducted as two-tailed tests. Non-parametric tests  
310 (Mann-Whitney; Spearman rank correlation) were used for non-normally distributed variables  
311 (e.g. white matter distance, Figure 3A); parametric tests (*t*-tests and Pearson correlation) were  
312 used for the remaining analyses. Data distributions were visually inspected or assumed to be  
313 normal for parametric tests.

314 **Data Availability**

315 Upon publication, all de-identified raw data and analysis code may be downloaded at [http://memory.psych.upenn.edu/Electrophysiological\\_Data](http://memory.psych.upenn.edu/Electrophysiological_Data).

317

318 **Results**

319 **Multivariate classifiers identify memory lapses**

320 Our stimulation strategy sought to intercept and rescue periods of poor memory encoding. To  
321 do so, we trained participant-specific multivariate classifiers to discriminate patterns of neural  
322 activity during record-only sessions of free recall (Figure 1A). For the Closed-loop group ( $N = 40$ ),  
323 classifiers were trained using the true mapping of features (spectral power  $\times$  electrodes) to recall  
324 performance; for the Random group ( $N = 17$ ), due to a technical error in labeling features (see  
325 *Methods*), classifiers were trained on permuted features. The recording electrode locations for the  
326 Closed-loop and Random groups appear as spheres in Figure 1B. After training the classifiers on  
327 record-only data, we used them in later (independent) sessions to identify poor memory states for  
328 targeting with stimulation.

329 Our first question was how well the classifiers predicted memory outcomes during the stim-  
330 ulation sessions (i.e. out-of-sample generalization). To answer this question, we used data from  
331 NoStim lists in which we obtained classifier predictions about the probability of recall for each  
332 word, but did not use these predictions to trigger stimulation (see *Methods*). Using area under  
333 the receiver operating characteristic curve as an index of classification accuracy, we found that  
334 classifiers for the Closed-loop reliably exceeded chance performance [Mean AUC = 0.62 (chance  
335 AUC = 0.50), Wilcoxon signed rank test  $P = 5.73 \times 10^{-7}$ ]. Closed-loop classifiers also outperformed  
336 classifiers for the Random group [Mann-Whitney  $U = 586.0, P = 1.85 \times 10^{-5}$ ]. As expected, Ran-  
337 dom classifiers did not exceed chance [Mean AUC = 0.49, Wilcoxon signed rank test  $P = 0.55$ ;  
338 Figure 1C].

339 To understand what features the classifier used to discriminate good vs. poor memory encoding  
340 states, we used a forward model for each participant to derive importance estimates for each

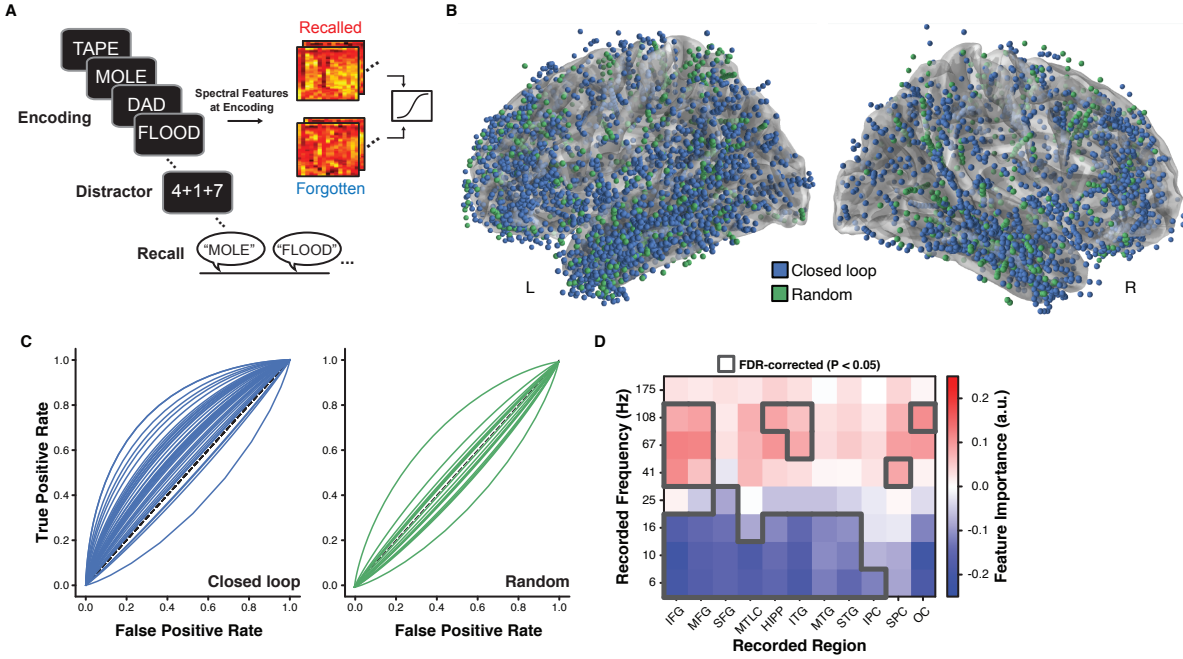


Figure 1: Stimulation strategy and classifier performance. **A.** Participants performed at least three sessions of the free recall task while being monitored with intracranial EEG. Multivariate classifiers trained on whole-brain patterns of spectral activity predicted subsequent recalled vs. not recalled words. **B.** Recording electrode locations for all participants in the Closed-loop (blue) and Random (green) groups, rendered on the Freesurfer average brain. **C.** Each participant's multivariate classifier then served as their personalized model to trigger stimulation. Classifiers trained on record-only data generalized to the stimulation session(s) for the Closed-loop group ( $P = 6.14 \times 10^{-7}$ ) and outperformed classifiers for the Random group ( $P = 2.73 \times 10^{-5}$ ). **D.** An analysis of feature importance for classifiers from the Closed-loop group showed that successful memory states were associated with decreases in low-frequency activity and increases in high-frequency activity.

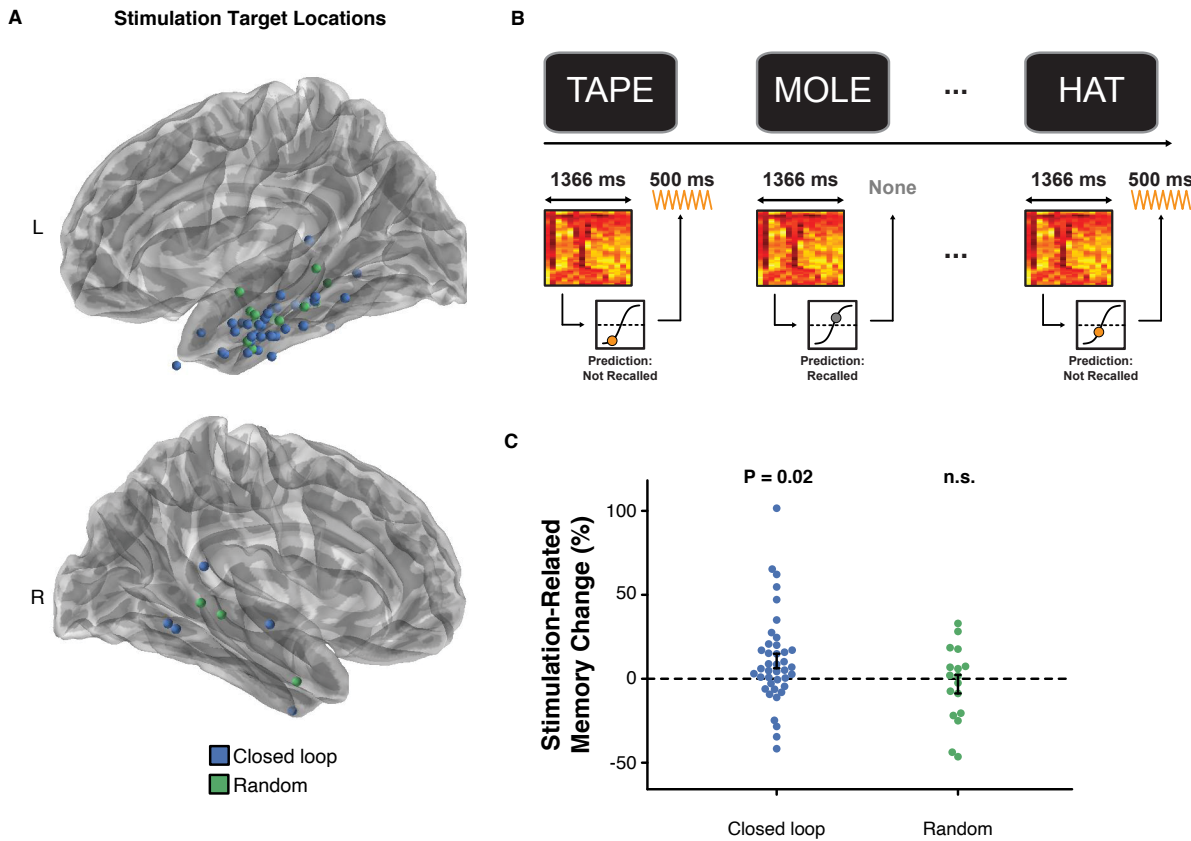
341 feature (Haufe et al., 2014). We averaged the feature importance values within a set of regions of  
 342 interest (ROIs) separately for each classifier frequency. Across participants, classifiers predicted  
 343 successful memory encoding based on increased high-frequency activity (especially in frontal,  
 344 lateral temporal, and medial temporal lobe areas) and decreased low-frequency activity across  
 345 much of the recorded cortex and subcortex (Figure 1D). This pattern, which we refer to as the  
 346 spectral tilt, has been observed in previous studies to be a biomarker of successful episodic memory  
 347 encoding and retrieval (Ezzyat et al., 2017; Burke et al., 2014; Long et al., 2014).

348 **Closed-loop LTC stimulation improves memory**

349 Having established that classifiers in the Closed-loop group reliably discriminate memory encod-  
350 ing states, we next asked if we could increase memory performance via stimulation of the LTC (Fig-  
351 ure 2A). Our stimulation strategy was based on detecting poor memory encoding states and inter-  
352 cepting them with stimulation (Figure 2B). For the Closed-loop and Random groups, we compared  
353 recall performance for lists in which we delivered stimulation (Stim lists) vs. identically structured  
354 lists in which we did not stimulate (NoStim lists, as described above). In the Closed-loop group,  
355 recall was higher on Stim lists compared to NoStim lists [ $\Delta = 10.6\% \pm 4.3$ ;  $t(39) = 2.45, P = 0.02$ ,  
356 Figure 2C], suggesting that intercepting poor memory encoding states with LTC stimulation en-  
357 hanced recall. In contrast, there was no difference in memory performance for the Random group  
358 [ $\Delta = -3.2\% \pm 5.5$ ;  $t(16) = -0.59, P = 0.57$ ]. There was a trend for greater memory enhancement  
359 for the Closed-loop compared to the Random group [ $t(55) = 1.83, P = 0.07$ ]. These findings  
360 are the first to directly compare closed-loop LTC stimulation with a random/open-loop stimula-  
361 tion control and, in a larger replication sample, demonstrate the robustness of previous studies  
362 showing memory enhancement via LTC stimulation (Ezzyat et al., 2018; Kucewicz et al., 2018;  
363 Kahana et al., 2023).

364 **White matter proximity mediates stimulation's effect on memory**

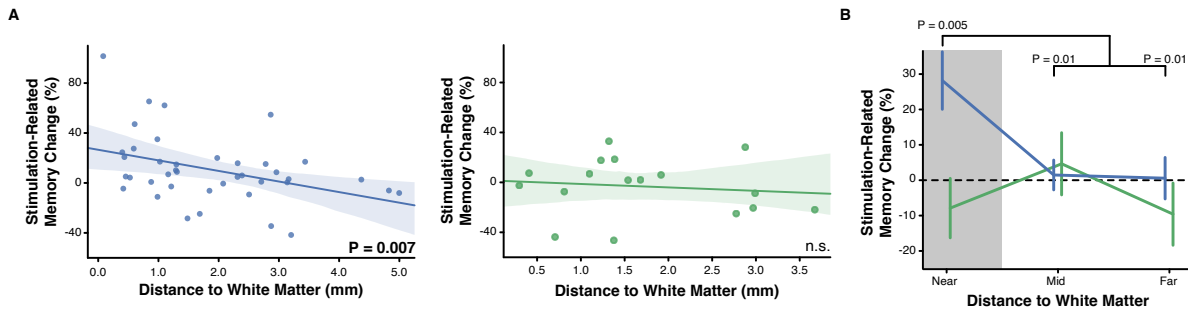
365 Motivated by physiological studies of electrical stimulation's effects on downstream targets (Mo-  
366 han et al., 2020; Solomon et al., 2018; Keller et al., 2018), we asked whether stimulating close to  
367 white matter tracts would produce greater positive or negative effects on memory. If so, this  
368 would suggest that the brain's anatomical network structure plays a key role in determining how  
369 effectively stimulation can modulate cognitive function (Stiso et al., 2019; Crocker et al., 2021). To  
370 answer this question, we examined how stimulation's effect on memory performance varied as a  
371 function of the stimulation target's proximity to white matter (see Figure 3A). For the Closed-loop  
372 group, lower distance to white matter predicted greater stimulation-related memory improvement  
373 [Spearman  $\rho(38) = -0.42, P = 0.007$ ; Figure 2B]. In the random stimulation group, we neither ex-  
374 pected nor observed a correlation between white matter distance and the memory effect [Spearman  
375  $\rho(15) = -0.17, P = 0.52$ ]. There was no difference between the distances to white matter for the



**Figure 2: Closed-loop stimulation improves memory performance.** A. Stimulation target locations for the Closed loop (blue) and Random (green) groups. B. Closed-loop stimulation strategy. C. Closed-loop LTC stimulation improved memory performance ( $P = 0.02$ ) while Random stimulation did not. Error bars in C reflect s.e.m.

376 Closed-loop and Random groups [Mann-Whitney  $U = 331.0, P = 0.44$ ] and the median distance  
 377 was in fact numerically greater for the Closed loop (1.58 mm) compared to the Random group  
 378 (1.39 mm). This suggests that distance to white matter alone does not explain the finding of im-  
 379 proved memory in the Closed-loop group. Instead, proximity to white matter appears to enhance  
 380 the effectiveness of closed-loop stimulation.

381 To further test this idea, we divided stimulation targets into terciles and asked whether stim-  
 382 ulation near white matter was particularly effective in modulating memory performance. Indeed,  
 383 Closed-loop stimulation targets near white matter enhanced memory performance on Stim lists  
 384 compared to NoStim lists [Near:  $M = 28.25\% \pm 8.14\%, t(13) = 3.23, P = 0.005$ ]. This memory  
 385 improvement was larger than for Closed-loop stimulation targets further away from white matter  
 386 [Mid:  $M = 1.55\% \pm 4.22\%, P = 0.01$ ; Far:  $M = 0.64\% \pm 5.87\%, P = 0.01$ ]. Closed-loop stimulation near

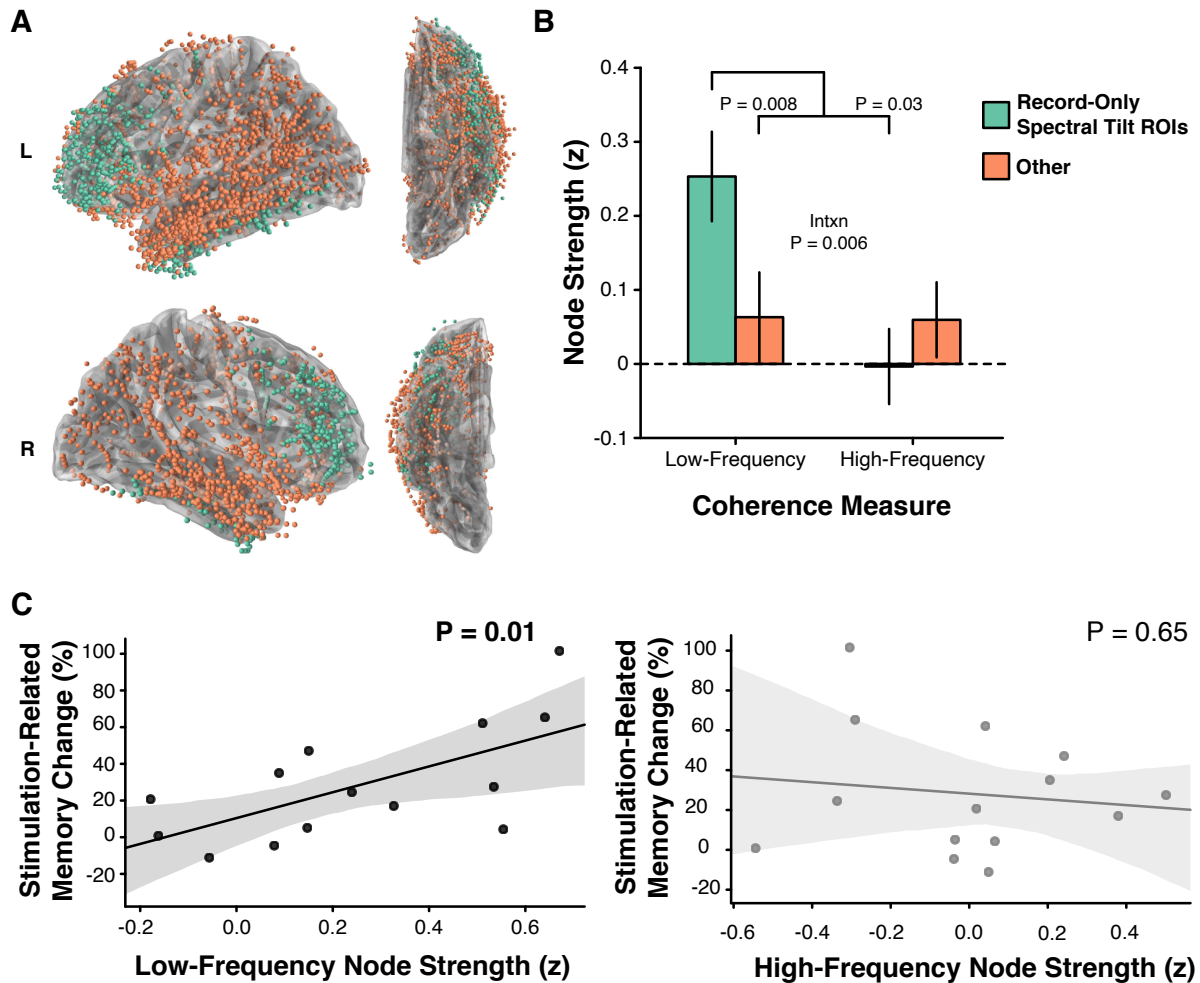


**Figure 3: Closed-loop stimulation near white matter enhances memory.** A. For the Closed-loop group, stimulation's effect on memory depended on the target distance from the nearest white matter [left,  $P = 0.007$ ]. The correlation was not significant for the Random group [right,  $P = 0.52$ ]. B. Closed-loop LTC stimulation improved memory performance for targets located nearest to white matter ( $P = 0.005$ ). There was no effect for the Random group ( $P = 0.62$ ). Error regions in A reflect the standard error of the estimate. Error bars in B reflect s.e.m.

387 white matter also significantly outperformed Random stimulation near white matter group [Rand-  
 388 dom  $M = -7.89\% \pm 8.40\%$ ,  $t(17) = 2.36$ ,  $P = 0.03$ , Figure 2C]. As expected, the Random group did  
 389 not show improved memory (Stim vs. NoStim within-participant) in any white matter distance bin  
 390 (all  $P > 0.38$ ). These data suggest that stimulating near white matter leads to greater modulation  
 391 of memory, and extend previous work that linked white matter proximity to stimulation's effect  
 392 on electrophysiology (Mohan et al., 2020; Solomon et al., 2018; Keller et al., 2018; Stiso et al., 2019;  
 393 Crocker et al., 2021).

### 394 Stimulation target functional connectivity predicts the change in memory

395 We next asked why closed-loop stimulation delivered near white matter reliably modulated mem-  
 396 ory function. One possibility is that stimulating near white matter allows more reliable and direct  
 397 access to the broader memory network connected to the stimulated location (Khambhati et al., 2019;  
 398 Stiso et al., 2019; Solomon et al., 2018; Mohan et al., 2020). We therefore measured functional con-  
 399 nectivity between the brain's memory encoding network and the stimulation targets located near  
 400 white matter. Critically, we constructed separate measurements of connectivity at low (5-13 Hz)  
 401 and high frequencies (45-90 Hz) by calculating coherence using participant-specific resting-state  
 402 data (see *Methods*). Then, to isolate the brain's memory encoding network, we identified all elec-  
 403 trodes that were in brain regions that showed a spectral tilt that predicted memory success during  
 404 the task, assessed using classifier feature importance Figure 3A. We then compared stimulation tar-



**Figure 4: Stimulation target functional connectivity.** A. We assigned each patient's record-only electrodes to two ROIs based on whether the electrode was located in a region that showed a memory-related spectral tilt or not (Other). B. Low-frequency connectivity was higher between the stimulation target and electrodes in classifier-defined memory regions, compared to electrodes in Other regions ( $P = 0.008$ ) and compared to high-frequency network connectivity ( $P = 0.03$ ). In contrast, there was no difference in stimulation target high-frequency network connectivity. C. For closed-loop targets nearest to white matter, there was a significant correlation between stimulation target low-frequency connectivity and stimulation's effect on memory [ $r(12) = 0.648, P = 0.01$ ]. There was no effect for high-frequency connectivity. Errorbars reflect s.e.m. Error regions reflect the standard error of the estimate.

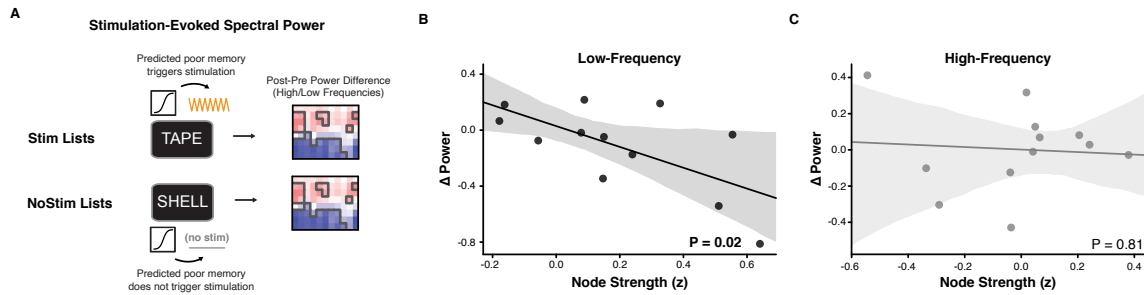
405 get connectivity to electrodes In vs. Out of the memory network, for both low and high-frequency  
 406 coherence (referred to as Node Strength). Stimulation targets showed stronger low-frequency con-  
 407 nectivity to electrodes in the memory network than to electrodes outside of the memory network  
 408 [ $t(13) = 3.14, P = 0.008$ , Figure 3B]. For memory network electrodes, low-frequency connectivity  
 409 was also higher than high-frequency connectivity [ $t(13) = 2.48, P = 0.03$ ]. In contrast, stimula-

410 tion targets showed equivalent high-frequency connectivity In vs. Out of the memory network  
411 [interaction:  $F(1, 13) = 10.54, P = 0.006$ , Figure 3B].

412 Although stimulation targets near white matter showed greater overall low-frequency con-  
413 nectivity with memory-predicting brain areas, this finding leaves open the question of whether  
414 variability in connectivity strength with the memory network predicts variability stimulation's  
415 effect on memory. To answer this question, we correlated low-frequency node strength with  
416 stimulation-related memory change. We found that low-frequency node strength predicted closed-  
417 loop stimulation's effect on memory [ $r(12) = 0.648, P = 0.01$ , Figure 3C] while high-frequency node  
418 strength did not ( $P = 0.65$ ). The difference in correlation for low vs. high-frequency node strength  
419 was also significant (two-tailed permutation test  $P = 0.03$ ). For all other targets that were further  
420 from white matter, there was no relation between node strength and stimulation-related memory  
421 change (all  $P > 0.21$ ).

#### 422 **Functional connectivity mediates stimulation's effect on downstream physiology**

423 The preceding results indicate that low-frequency functional connectivity to the memory network  
424 predicts stimulation effects on memory. Our final question was whether low-frequency connec-  
425 tivity also predicts stimulation's physiological effects across the memory network. To test this  
426 prediction we again examined Closed-loop stimulation targets near white matter and correlated  
427 each stimulation target's connectivity to the memory network with the stimulation-evoked spec-  
428 tral power in this network (Figure 5A). Two participants' data were excluded due to excessive  
429 stimulation artifact on the recording channels. In the remaining participants, we found that  
430 stimulation-target functional connectivity predicted stimulation-related changes in low-frequency  
431 power [ $r(10) = -0.65, P = 0.02$ , Figure 5B]. The correlation was not significant when using high-  
432 frequency connectivity and evoked power ( $P = 0.81$ , Figure 5C).



**Figure 5: Memory network connectivity predicts physiology.** **A.** Schematic of analysis of stimulation-evoked physiology. **B.** For stimulation targets near white matter, low-frequency functional connectivity predicted the stimulation evoked change in low-frequency power ( $P = 0.02$ ). **C.** High-frequency network connectivity did not predict stimulation's effect on high-frequency activity.

## 433 Discussion

434 Direct electrical stimulation has emerged as a powerful tool for manipulating neural activity. The  
435 present study evaluated the hypothesis that network properties of a stimulated brain location  
436 predict stimulation's effects on both memory and network physiology. Prior studies suggest that  
437 white matter pathways mediate stimulation's network-level physiological effects (Paulk et al., 2022;  
438 Solomon et al., 2018; Mohan et al., 2020; Khambhati et al., 2019; Stiso et al., 2019). Other studies  
439 demonstrate that measures of structural and functional connectivity predict stimulation's effects on  
440 downstream targets (Keller et al., 2011; Fox et al., 2020; Solomon et al., 2018). However, none have  
441 simultaneously linked structural/functional connectivity with both (1) a reliable improvement over  
442 baseline cognitive functioning and (2) concomitant changes in neurophysiology that explain the be-  
443 havioral effect. To directly address these questions, we asked whether white-matter proximity and  
444 functional connectivity underlie the degree to which stimulation of LTC produces improvements  
445 or impairments of memory, alongside changes in oscillatory signatures of mnemonic function.

446 We found that closed-loop stimulation of LTC reliably improved memory on stimulated vs.  
447 non-stimulated lists. Consistent with the hypothesis that white-matter pathways convey the  
448 effects of stimulation to the broader memory network, we found the benefits of closed-loop LTC  
449 stimulation to arise principally from stimulating in, or near, white matter pathways. For the  
450 electrodes nearest to white matter, stimulation yielded a 28% increase in recall performance,  
451 whereas we failed to observe any reliable increase when delivering stimulation far from these  
452 pathways (1%). In a subgroup of subjects who received randomly timed stimulation in LTC  
453 targets we failed to observe any improvement in memory performance.

454 To evaluate how stimulation–target functional connectivity mediates stimulation's behavioral  
455 and physiological effects, we analyzed participant-specific large-scale neural recordings obtained  
456 during prior record-only sessions. Prior studies have shown that brain networks become coherent  
457 at low-frequencies during successful memory encoding and retrieval (Solomon et al., 2017; Kragel  
458 et al., 2021a), so we used low-frequency coherence to measure the network node strength of  
459 each stimulation target. We then asked if greater node strength between LTC stimulation targets  
460 and downstream memory-predicting areas resulted in greater effects of stimulation on memory  
461 performance. Consistent with this hypothesis, we found a strong positive correlation ( $r = 0.648$ , see

462 Figure 4C) between low-frequency connectivity and stimulation-related memory improvement.  
463 Finally, LTC stimulation engaged low-frequency activity across a broader brain network in a way  
464 that matched the network position of the stimulated location (Figure 5).

465 Our data highlight how precise targeting improves stimulation efficacy by showing that de-  
466 livering stimulation near LTC white-matter leads to greater stimulation-related memory gains  
467 (Figure 3C). By linking low-frequency network connectivity with physiological and behavioral  
468 outcomes, our study also points to a neural mechanism for modulating memory with stimu-  
469 lation. This result extends earlier work that demonstrated the potential to modulate episodic  
470 memory by targeting LTC with stimulation (Ezzyat et al., 2018; Kucewicz et al., 2018). Di-  
471 rectly comparing closed-loop and open-loop stimulation strategies in the same study helps to  
472 establish a causal role for the closed-loop approach (Hampson et al., 2018; Ezzyat and Rizzuto,  
473 2018). Finally, our data from 57 stimulation targets (across 47 patients) also represents a sub-  
474 stantial increase compared to sample sizes described in related prior studies (Ezzyat et al., 2018;  
475 Hampson et al., 2018).

476 Prior work has linked successful memory function with theta power and coherence (Burke  
477 et al., 2013; Solomon et al., 2017; Herweg et al., 2020; Griffiths et al., 2019; Kragel et al., 2021b;  
478 Ter Wal et al., 2021; Osipova et al., 2006; Guderian and Düzel, 2005; Klimesch et al., 1997;  
479 Staudigl and Hanslmayr, 2013). Here, we investigated this physiological correlate of memory  
480 function by testing how memory-modulating LTC stimulation affects low-frequency physiology.  
481 We found that stimulation's effect on low-frequency activity depends on the low-frequency func-  
482 tional connectivity of the stimulation target. This suggests that identifying strong functional con-  
483 nections can produce stronger modulation of low-frequency activity within the memory network.  
484 Furthermore, we found that stimulation that modulated low-frequency activity also modulated  
485 memory performance.

486 Several prior studies found potential therapeutic benefits of closed-loop stimulation triggered  
487 by decoding of intracranial brain recordings (Ezzyat et al., 2018; Scangos et al., 2021a; Hampson  
488 et al., 2018; Kahana et al., 2023). However, with some important exceptions (Hampson et al.,  
489 2018), this work has lacked an open-loop or random stimulation control condition, leaving open  
490 the question of what *specific* role the closed-loop nature of stimulation played in its therapeutic  
491 effects. Here, we compared the effects of closed-loop stimulation with a random stimulation

492 condition. Closed-loop participants received stimulation only for those items predicted to be  
493 forgotten. Participants in the random group followed the same protocol, but using classifiers  
494 trained on permuted data, resulting in stimulation being applied without regard to predicted  
495 memory success. This led to reliable memory improvement for the closed-loop group and none  
496 for the random group, despite following an otherwise identical protocol (Figure 1C).

497 We found that closed-loop stimulation improved memory the most when it was delivered to  
498 LTC targets in or near white matter. This finding builds on a growing literature that indicates that  
499 stimulation is most effective when it is delivered in or near white matter pathways (Khambhati  
500 et al., 2019; Stiso et al., 2019; Mohan et al., 2020; Solomon et al., 2018; Pault et al., 2022). One  
501 explanation for this phenomenon is that only stimulation of white matter pathways successfully  
502 engages broader brain networks, perhaps via oscillatory synchronization. In contrast, gray matter  
503 stimulation tends to cause more local effects (Mohan et al., 2020; Pault et al., 2022). Though purely  
504 local effects may sometimes be desirable, the key cognitive and pathophysiological processes of  
505 greatest interest to neuroscientists tend to involve multiple interconnected brain regions.

506 Among its many applications for modulating cognition and behavior (Siddiqi et al., 2022;  
507 Fox et al., 2020; Sreekumar et al., 2017) a number of recent studies have evaluated stimulation's  
508 potential for enhancing episodic memory (Mankin and Fried, 2020; Suthana and Fried, 2014;  
509 Curot et al., 2017; Lee et al., 2013; Sankar et al., 2014). While our study investigated numerous  
510 stimulation targets within the LTC, future work should compare stimulation of this region to other  
511 brain areas within the broader episodic memory network. Recent work suggests that stimulating  
512 white matter pathways in the medial temporal lobe, for example, can also improve memory (Titiz  
513 et al., 2017; Mankin et al., 2021; Suthana et al., 2012). However, these previous studies used visual  
514 and/or spatial memoranda, while the present study focused on encoding and retrieval of verbal  
515 material. Thus, future research should compare stimulation to the lateral and medial temporal  
516 lobes, to determine whether stimulation target location interacts with the modality of the to-be-  
517 remembered information. This could contribute to other work that has used stimulation to study  
518 the component processes that contribute to successful episodic memory (El-Kalliny et al., 2019).

519 We delivered stimulation using macroelectrodes, consistent with its clinical applications (Krauss  
520 et al., 2021; Morrell, 2011; Sun et al., 2008). Macroelectrode stimulation alters local activity  
521 at the spatial scale of the distance between the anode and cathode (approximately 1 cm), but

522 can also alter more distant regions. Because memory relies on a broad network of cortical  
523 and subcortical regions, including the hippocampus (Kim, 2011; Keerativittayayut et al., 2018),  
524 stimulating a broader network may be necessary to impact cognitive function. On the other  
525 hand, memory also relies on the recapitulation of specific patterns of neuronal activity, espe-  
526 cially within the hippocampus (Foster, 2017; Staresina and Wimber, 2019). Thus, other work  
527 has stimulated through microelectrodes to mimic and reinstate memory-related hippocampal ac-  
528 tivity using a model-based closed loop approach (Hampson et al., 2018; Hampson et al., 2013;  
529 Deadwyler et al., 2017). An avenue for future work could use macroelectrode stimulation in a  
530 similar vein, by triggering stimulation at multiple macroelectrode contacts in order to synchronize  
531 a particular spatiotemporal pattern of activity across key memory-related regions (Kim et al., 2016;  
532 Kim et al., 2018).

533 In relating low-frequency network connectivity, physiology, and behavior, our study con-  
534 tributes to methodological development for invasive stimulation (Krauss et al., 2021; Cagnan et  
535 al., 2019) that illuminates the critical role of low-frequency networks in cognition (Voytek and  
536 Knight, 2015). In addition, the present study also suggests that other methods that manipulate  
537 low-frequency activity could be leveraged to modulate neural and cognitive function. Several  
538 recent studies using non-invasive methods have leveraged low-frequency theta-patterned stimu-  
539 lation to modulate episodic and working memory (Nilakantan et al., 2017; Hermiller et al., 2020;  
540 Tambini et al., 2018; Warren et al., 2019; Grover et al., 2022). Such low-frequency stimulation  
541 modulates electrophysiology perhaps by entraining low-frequency oscillations that are associated  
542 with cognitive function (Solomon et al., 2021; Reinhart and Nguyen, 2019; Reinhart et al., 2017;  
543 Hanslmayr et al., 2019).

544 In summary, our demonstration of improved memory with closed-loop stimulation supports  
545 the idea that memory function is dynamic, and that closed-loop algorithms that account for  
546 moment-to-moment variability in the brain's memory state can selectively deliver stimulation  
547 only when it is needed. The present study also links closed-loop stimulation efficacy to white  
548 matter targeting, brain-wide evoked physiology, and changes in episodic memory performance.  
549 The findings suggest future strategies for using the functional and anatomical network profile of  
550 putative stimulation targets to optimize downstream changes in oscillatory activity and cognition.

551 **Acknowledgments**

552 This work was supported by NIH grant NS106611 and MTEC project 20-06-MOM from the Army  
553 Medical Research and Development Command. We are indebted to the patients and their families  
554 for their participation and support. The views, opinions, and/or findings contained in this material  
555 are those of the authors and should not be interpreted as representing the official views or policies  
556 of the Department of Defense or the U.S. Government. B.C.J. receives research funding from  
557 NeuroPace and Medtronic not relating to this research. M.J.K. and D.S.R. each hold a greater than  
558 5% equity interest in Nia Therapeutics, LLC, a company intended to develop and commercialize  
559 brain stimulation therapies for memory restoration.

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