NMDA receptor function in large-scale anticorrelated neural systems with implications for cognition and schizophrenia

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Glutamatergic neurotransmission mediated by N-methyl-D-aspartate (NMDA) receptors is vital for the cortical computations underlying cognition and might be disrupted in severe neuropsychiatric illnesses such as schizophrenia. Studies on this topic have been limited to processes in local circuits; however, cognition involves large-scale brain systems with multiple interacting regions. A prominent feature of the human brain's global architecture is the anticorrelation of default-mode vs. task-positive systems. Here, we show that administration of an NMDA glutamate receptor antagonist, ketamine, disrupted the reciprocal relationship between these systems in terms of task-dependent activation and connectivity during performance of delayed working memory. Furthermore, the degree of this disruption predicted task performance and transiently evoked symptoms characteristic of schizophrenia. We offer a parsimonious hypothesis for this disruption via biophysically realistic computational modeling, namely cortical disinhibition. Together, the present findings establish links between glutamate’s role in the organization of large-scale anticorrelated neural systems, cognition, and symptoms associated with schizophrenia in humans.

Drug treatments for serious mental illnesses and investigations of the neurochemical bases of healthy cognition have, for the most part, targeted the slow neuromodulatory neurotransmitters, dopamine and serotonin (1). However, rapid excitatory glutamatergic and inhibitory γ-aminobutyric acid (GABA) signals mediate local and long-range cortical computations (2) and play a critical role in cognition and severe psychiatric illnesses such as schizophrenia (3–5). We investigated how disrupting the N-methyl-D-aspartate (NMDA) receptor component of fast glutamatergic neurotransmission via the administration of the NMDA receptor antagonist ketamine altered cognitive performance and systems-level neural activity and connectivity in healthy volunteers. Furthermore, we related these system-level neural changes to behavior and transiently evoked psychotic symptoms associated with schizophrenia.

Studies investigating glutamate’s role in cognition have largely focused on local circuits (5–7); however, cognition involves large-scale brain systems with multiple interacting regions. Recent neuroimaging work highlights the competitive relationships between two large-scale neural systems: a set of brain regions preferentially engaged during tasks that require goal-directed cognition and attention (task-positive) and the regions associated with resting-state neural activity and connectivity in healthy volunteers. Further, we related these system-level neural changes to behavior and transiently evoked psychotic symptoms associated with schizophrenia.

Here, we administered ketamine to healthy volunteers (Methods), which safely, transiently, and reversibly perturbs regional brain responses in a number of nodes integral to large-scale brain systems (15). Ketamine also alters synaptic function in the long-range connections that provide the connective basis of distributed brain systems (16). Infusing ketamine induces a transient state resembling schizophrenia in healthy volunteers (14), effects that have been related to individual differences in regional functional brain responses to cognitive tasks (13, 15, 17, 18). Here we investigated ketamine’s effects while subjects performed a demanding working
memory (WM) task, a cognitive process involving short-term encoding and maintenance of information (19) that is profoundly impaired in schizophrenia (20) and that robustly modulates task-positive and DMN networks (10). Using blood oxygen-level-dependent (BOLD) imaging, we examined how modulating glutamatergic neurotransmission alters system-level activation and deactivation and the interplay between large-scale neural systems during WM, as well as the relevance of these changes for the psychosis-like phenomenology induced by ketamine. We hypothesized that reduction in signaling via NMDA receptors would attenuate task-related activation but also reduce the degree of DMN suppression: a phenomenon critical for cognitive performance (21, 22) but disrupted in neuropsychiatric illness (11, 23).

Lastly, we related observed experimental effects to a well-validated biophysically realistic computational model of WM (24) to study a leading hypothesized synaptic mechanism of NMDA blockade on neural activity, namely disrupted balance between cortical excitation/inhibition (5). With our computational modeling approach, we attempt to provide a framework for relating synaptic-level hypotheses to observed neural activity and cognition to better understand the complex and puzzling symptoms of serious mental illness (25). Together, the present findings highlight the functional utility of large-scale brain systems and the role of intact NMDA receptor operation in their interrelationship.

Results

Behavioral Effects. We first assessed the impact of ketamine on performance of the spatial WM task (Fig. 1A). Confirming hypothesized effects, results revealed a significant accuracy reduction for WM vs. control trials following ketamine administration $[t(18) = 5.65, P < 0.0001]$ (Fig. 1C; for details, see SI Text and Fig. S1), which was in line with well-established deleterious effects of ketamine on WM across species (26, 27).

Task-Based Activation and Deactivation. Performing WM tasks robustly modulates task-positive regions (28) but also involves deactivation of the DMN (10). We first verified that the cognitive task engaged regions previously shown to be active during WM (28) by computing a main effect of task condition (WM vs. control trials) during encoding, delay, and probe phases of the trial (Fig. 2 A–C and SI Text, fMRI Analyses). Analyses confirmed robust WM effects across the three WM phases, in terms of both task-based activation (Fig. 2, Left) and deactivation (Fig. 2, Right), in correspondence with the DMN (8) (for all WM-modulated regions, see Tables S1–S3).

Ketamine Modulation of Task-Based Activation and Deactivation. Next, using a stringent conjunction masking technique (29) (Fig. 1B), we independently tested for effects of ketamine only in areas that were explicitly engaged during WM (Fig. 2), ensuring a principled method for isolating ketamine modulation of task-related regions. We computed a whole-brain–corrected task condition $\times$ infusion interaction map and then computed a conjunction (logical AND) between surviving results and the map showing main effects of task condition. We repeated this procedure across task phases. This approach ensures that observed BOLD modulations by ketamine are present in regions explicitly modulated by the WM task (SI Text, fMRI Analyses). Results revealed a number of WM-related regions modulated by ketamine across encoding and delay (Fig. S2 and Table S4). We highlight two exemplar regions exhibiting identified effects: the dorsolateral prefrontal cortex [Brodman area (BA) 46; Fig. 3A, Upper] and the precuneus (BA 31; Fig. 3A, Lower) (see Fig. S3 for all regions). These effects illustrate that ketamine attenuated signal in task-activated regions, as well as deactivation of regions overlapping the DMN (8). No regions exhibited a ketamine modulation of WM signals at the probe phase. Consistent with hypotheses, these results indicate that ketamine not only attenuates encoding and early delay signals during WM but also disrupts suppression of regions typically deactivated during demanding cognitive operations (10). To further verify preferential effects of ketamine in regions involved in WM (and to rule out potential vascular confounds), we examined a motor cortex region that showed a strong response to the button press during the probe phase. There was no BOLD signal attenuation by ketamine in this motor region; in fact, the response for ketamine was numerically higher (Fig. S3).

![Image](https://example.com/image.png)

Fig. 1. (A) Subjects encoded four or two (not shown; see Methods) circle locations and, after a delay, indicated whether the circle was presented at that location or not (probe). Subjects also completed a control task where four gray circles appeared but were explicitly asked not to encode the circles. During the probe phase, another gray circle was shown, requiring a motor response but no recall. (B) Second-level conjunction fMRI analysis strategy: (i) we computed a main effect of task (i.e., WM vs. control condition) type I error corrected at the whole-brain level (Fig. 2); and (ii) we computed a task $\times$ infusion interaction, revealing regions differentially modulated by ketamine across task conditions. Regions identified this way are not guaranteed to be involved in WM. That is, regions showing a task $\times$ infusion interaction may not show engagement during WM (i.e., main effect of task). Thus, we computed a conjunction (logical AND) between these effects. The surviving regions were ensured to show both a task main effect and modulation of this effect by ketamine (Fig. 3A and Fig. S2). (C) Percentage drop in accuracy (% correct) is shown for the control (white bar) and WM (black bar) tasks following ketamine vs. placebo infusion (difference plotted). **P < 0.0001 (see SI Text, Behavioral Results for complete behavioral analysis). Error bars reflect ±1 SEM.
Examing Hypothesized Synaptic Effects of Ketamine via Computational Modeling. The above analysis suggests a breakdown in WM-related BOLD signal in both task-activated and task-deactivated regions. As noted, a leading hypothesis for ketamine’s effects at the synaptic level, derived from both animal and human work, postulates disinhibition via preferential antagonism of NMDA receptors on inhibitory interneurons (3, 30). How can we reconcile the observed reduction in BOLD signal and WM performance with disinhibition? To investigate hypothesized effects of ketamine on the cortical microcircuit level, which may sheds light on observed BOLD effects, we adapted a well-validated biophysically plausible computational model of WM (24). Our model is comprised of two modules: a task-activated module, which is a recurrent microcircuit capable of WM computations at the time scale relevant to a single WM trial; and a task-deactivated module representing the DMN characterized by high baseline firing rate and deactivation at task onset (31). The microcircuit modules interact through long-range, net inhibitory projections (SI Text, Computational Modeling). The biophysical realism in the model allows for direct implementation of pharmacological manipulations at the synaptic level (32) (Methods and SI Text, Computational Modeling for model operation and pharmacological implementation). Here, we related hypothesized ketamine effects, highlighted by preclinical findings (3, 30) to observed BOLD effects: we preferentially disrupted NMDA conductance onto GABAergic interneurons (i.e., E-I conductance). We contrasted E-I reduction with a preferential reduction in recurrent excitation on pyramidal cells (i.e., E-E reduction; see Figs. S5 and S6). To ensure model simulations describe processes qualitatively similar to empirical observations, we computed BOLD signal based on model-generated activity, which reproduced described effects (Fig. 3B; see SI Text, Computational Modeling for details). Although at achieved

Fig. 2. WM effects for encoding (A), delay (B), and probe (C) phases are shown. Maps illustrate task-based WM activations (orange-yellow) and deactivations (blue) (Left and Right, respectively). All displayed foci met a whole-brain correction (SI Text, fMRI Analyses). Time courses are shown for exemplar regions identified using an assumed hemodynamic response function (SI Text, fMRI Analyses), exhibiting significant WM (black squares) vs. control task (dashed lines) effects. Approximate trial epochs (encoding, delay, probe) are marked with gray vertical bars. Canonical WM responses are evident across all epochs. Region coordinates are marked in boxes. For complete list of task-modulated foci, see Tables S1–S3.

Fig. 3. (A) Regions exhibiting WM effects and modulation of this effect by ketamine for task-based activation (Upper) and task-based deactivation (Lower). WM time courses are shown for dorso-lateral prefrontal cortex (Upper) and precuneus (Lower) for ketamine (red) and placebo (blue) (coordinates are shown in boxes). Complete list of foci exhibiting significant effects across encoding and delay phases is presented in Table S4 and Fig. S2. The probe phase analysis did not reveal significant modulation by ketamine. Note: less negative value for DMN under ketamine reflects less deactivation relative to baseline compared with the placebo condition. (B) Computational model scheme, comprised of task-activated (Upper) and task-deactivated (Lower) modules followed by modeling results. We modeled the effects of ketamine as a reduction of NMDA conductance onto inhibitory interneurons (gE-I). We examined whether “disinhibition” via reduced NMDA conductance onto GABA cells (E-I) would result in effects similar to BOLD findings under ketamine. Here, we present predicted BOLD signal derived from the simulated local field potential (LFP) on the time scale comparable to a single WM trial in the experiment to appropriately juxtapose model simulations to BOLD empirical observations. For complete modeling implementation, BOLD simulation details, and comparison with firing-rate results, see SI Text, Computational Modeling and Figs. S4–S6.
concentrations ketamine likely also disrupts recurrent excitation (i.e., E-E conductance), E-I reduction alone was sufficient to account for both the attenuation in task-related activation and the lack of DMN suppression (see Figs. S5 and S6). Furthermore, there are two mechanisms by which reduced E-I strength could potentially disrupt the proper pattern of activation and deactivation: (i) long-range (net inhibitory) connections between modules are weakened, impairing the ability of the task-activated module to shut down the task-negative module; and (ii) local disinhibition renders a hyperactive microcircuit less sensitive to the long-range input, so that the already high-firing task-deactivated module cannot be shut down even with an equal-strength, long-range suppressive input. We found that local microcircuit E-I reduction, as opposed to long-range reduction, plays the dominant role in disrupting model function (Fig. S4).

Ketamine Modulation of Task-Based Connectivity. Given that glutamate plays a role in both local and long-range cortical computations, we also examined the possibility that NMDA receptor blockade may induce a reduction in task-based functional connectivity (tb-fcMRI) between large-scale anticorrelated systems. This second-level hypothesis was compelling because ketamine modulated task-related activation and deactivation in these systems. To this end, we selected a set of independent seeds previously identified as part of the fronto-parietal (FP) and DMN (8, 33). We focused on the delay phase of the trial using a previously validated technique developed to circumvent the possibility that overall task structure drives observed relationships (21) (for details see SI Text, tb-fcMRI). As hypothesized, there was a significant modulation of tb-IMRI between the FP-DMN networks during the delay, confirmed statistically by a task × infusion interaction ($F(1,18) = 11.09; P < 0.004$) (Fig. 4). This effect was preferential for the FP-DMN tb-fcMRI (when examining the cingulo-opercular system there was no task × infusion interaction, although there was a main effect of infusion, Fig. S7). Given emerging concerns that spurious head movement can confound connectivity results (34, 35), we implemented an additional volume censoring (“scrubbing”) movement correction (36, 37). The tb-fcMRI results remained significant after movement scrubbing ($F(1,18) = 6.1; P < 0.025$). For completeness, we present both sets of results in Fig. S7 and additional movement analysis details (SI Text, tb-fcMRI).

Ketamine Disrupts Performance-Related Deactivation. Next, we examined whether ketamine effects on WM in the task-based analysis relate to behavioral performance. We computed an additional model with WM accuracy as a covariate to enable examination of the within-subject, trial-by-trial relationship between behavioral performance and brain activity (38) (SI Text, fMRI Analyses). We tested whether foci showing a ketamine modulation of WM-related signals also exhibit a differential pattern of responses related to performance accuracy. To test this hypothesis, we computed an accuracy (correct vs. incorrect WM trials) × infusion (ketamine vs. placebo) interaction specifically for WM trials. We did so across task phases, but only delay-related results revealed significant findings. Among the foci identified in Fig. S4, three DMN regions showed a significant accuracy × infusion interaction: middle temporal gyrus [$F(1,18) = 7.93; P < 0.015$]; precuneus [$F(1,18) = 5.29; P < 0.035$], and superior frontal gyrus [$F(1,18) = 5.05; P < 0.04$]. We highlight effects for the precuneus (Fig. S4) given the consistent pattern of results across all three regions: during placebo infusion there was a greater reduction of task-based deactivation for correct vs. incorrect trials. However, following ketamine administration there was less task-based suppression on accurate trials. These results replicate prior findings showing that DMN suppression is vital for WM performance (21, 22) and suggest that NMDA blockade attenuates the ability for such performance-relevant suppression to take place. Notably, these patterns were evident in task-deactivated regions, but we did not observe an accuracy × infusion modulation of regions exhibiting WM-related activation.

DMN Signal and Ketamine-Induced Psychiatric Symptoms. We also examined the degree to which ketamine-induced modulations of brain responses related to symptom severity: establishing links between cognition, brain function, and psychiatric symptoms. To that end, we computed a correlation between positive, negative, and dissociative symptoms measures (for clinical measure details, see SI Text, Clinical Measures) and the degree of DMN deactivation during WM (we averaged across all DMN regions modulated by ketamine; see SI Text, fMRI Analyses). Only the relationship with negative symptoms reached significance [r = 0.61; P < 0.006; two-tailed, Bonferroni-corrected] (Fig. 5B), indicating that the most symptomatic subjects had the least ability to suppress DMN during WM.

Discussion

Present results suggest that intact NMDA receptor function is critical for optimal cognitive performance, at least in the context of WM, as well as seamless engagement and deactivation of distributed anticorrelated systems in the human brain. Here, we focused on pharmacologically disrupting fast glutamatergic neurotransmission. Prior work attempting to improve cognition found that enhancing slow neuromodulation via modafinil resulted in more potent deactivation of the DMN (39). Minzenberg et al. (39) hypothesized that disrupted reciprocal inhibition, mediated via GABA interneurons, might be responsible for the breakdown of the “mirror” anticorrelated structure of these large-scale systems. Indeed, one leading hypothesis of ketamine’s effects is preferential antagonism of NMDA conductance on GABA cells, which would lead to disinhibition (3, 30, 40). Our experimental results imply NMDA receptor activity in optimal local and long-range circuit function. When induced via ketamine, NMDA receptor hypofunction may result in cognitive deficits and negative symptoms associated with schizophrenia. This may be driven by disrupting the balance of excitation/inhibition in the cortical microcircuit, as recently suggested by preclinical work (5). Our modeling simulations offer a framework for relating this hypothesis, at the

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**Fig. 4.** tb-fcMRI during the delay phase between independently selected FP and DMN regions following ketamine (red) and placebo (blue) infusion, shown for the control (CT) and WM task conditions (all seed coordinates are listed in Table S5; for tb-fcMRI details, see SI Text, tb-fcMRI). Error bars reflect ±1 SEM.
synaptic level, to experimental results, suggesting that disrupted NMDA conductance on GABA cells could contribute to the observed effects.

The model is a proof-of-principle instantiation of one leading hypothesis of what may be ketamine’s effects on task-based BOLD response in certain regions and its modulation of cellular-level phenomena (30). The reciprocal inhibition between microcircuit modules is an important assumption based on leading hypotheses of the intrinsic configuration of these large-scale systems (8). We explicitly used this assumed antagonistic configuration to relate synaptic-level hypotheses to systems-level observations. Using this framework, we found that local E/I balance is the crucial property of the model, suggesting a perspective on the importance of local circuit properties in controlling the nature of large-scale interactions between brain areas during cognitive tasks. That is, the observed relationship between modules was very sensitive to NMDA receptor manipulation within each module. The modeling results also highlight that a subtle E-I perturbation resulted in disinhibition and was robust across a range of E-I reductions (Fig. S6). In contrast, E-E perturbations result in a set of regimes that were not observed experimentally, namely reduced firing across both task-activated and task-deactivated modules. This pattern might be observed under a higher level of ketamine that may result in elevated level of NMDA blockade that also strongly affects E-E conductance (e.g., anesthesia).

The model provides a biologically plausible framework for relating experimental findings following neuropharmacological manipulations to cellular-level hypotheses. As recently proposed by Montague et al. (25), this approach can help hone our understanding of synaptic disruptions, relate these hypotheses to system-level observations and ultimately to behavior observed in severe mental illness, for instance, the significant relationship between DMN signal and behavior in the present experiment.

Indeed, at the systems level, a growing literature associates DMN abnormalities with neuropsychiatric illness (23). A recent investigation examined the effect of transcranial magnetic stimulation (TMS) on modulating DMN function but did not report psychiatric symptomatology in healthy volunteers (41). Conversely, a recent study of schizophrenia patients linked DMN deactivation, cognition, and symptom severity (11). It may be possible that these two investigations capture the phenomenon on different time-scales: perturbing the DMN transiently via TMS may not be sufficient to potent or persistent to induce profound psychiatric phenomena, but more extended “tonic” manipulation via NMDA receptor blockade may begin to resemble what is observed in patients. Our experimental findings further highlight the functional relevance of suppressing the DMN during WM. As argued previously, DMN activity may reflect “passive” mental phenomena, such as self-referential, future-oriented thought (42). Whitfield-Gabrieli et al. suggest that “over-engagement” of the default network could lead to an exaggerated focus on one’s own thoughts and feelings, as well as an ambiguous integration between one’s internal and external environment (11). In that sense, tonic activation of the DMN in schizophrenia, if unregulated, may result in a state marked by exaggerated self-relevant processing, blurring the boundary between neural computations relevant for internal percepts and external reality. Furthermore, inadequately suppressed tonic activity in the DMN may obstruct goal-directed cognition and interfere with motivated pursuits, both characteristic of the negative syndrome (43). Indeed, failure to deactivate the DMN has been linked with poor cognitive performance in healthy populations (22) and lack of task-based DMN deactivation has been observed in schizophrenia, even before DMN literature evolved (44). Therefore, overactivity in the DMN, resulting from cortical disinhibition, may exacerbate cognitive dysfunction observed in neuropsychiatric conditions by reducing cortical efficiency and increasing noise during goal-directed cognition. The present findings also implicate the role of fast glutamatergic neurotransmission in the emergence of negative symptoms, which are suboptimally treated by medication targeting dopamine and serotonin (45).

Taken together, these results offer evidence that modulating glutamatergic neurotransmission in humans alters the relationship between large-scale, task-positive and task-negative neural systems in a performance and symptom-related manner. Our experimental findings and simulations are in line with one hypothesized pathophysiological mechanism for cognitive impairment and negative symptoms: that of reduced NMDA conductance on GABA interneurons, consistent with cortical disinhibition proposed previously (4, 5). These observations provide a possible framework for treatment development aimed at ameliorating these debilitating and inadequately treated symptoms associated with schizophrenia.

Methods

Subjects. Nineteen healthy, neurologically and psychiatrically intact right-handed volunteers (10 male) with a mean ± SD age of 27.5 ± 6.3 y completed the study (see SI Text, Subjects for complete recruitment details).

Experimental Task Design and Infusion. While in the scanner, subjects completed a well-validated delayed spatial WM task, described in our prior work (46), developed to mimic primate physiology experiments (47) (Fig. 1A). During scanning two phases of testing occurred. First, subjects were administered saline while they completed a series of scans. Second, subjects underwent the same series of scans during administration of ketamine. Subjects completed 32 WM trials and 16 control task trials (i.e., no WM encoding and maintenance requirement but requiring a probe response to control for motor effects) per infusion, resulting in 64 WM and 32 control trials per visit. Subjects completed three such visits (see SI Text, Overall Experimental Design and SI Text, Infusion Protocol for experimental design and infusion details).

The WM task included trials with two and four encoding locations, the former with extended inter-trial interval (ITI) (2500 ms), to make both the same length. The control task always contained four locations. All reported analyses collapse across two and four encoding locations given no load effects on reported behavioral (SI Text, Behavioral Results) and neuroimaging analyses (SI Text, fMRI Analyses). Following scanning, subjects underwent a battery of clinical measures to quantify the presence, nature, and severity of psychotic symptoms (SI Text, Clinical Measures).
fMRI Acquisition and Analysis. All data were acquired using a 3T Tim Trio (Siemens) scanner at the Yale University School of Medicine (see Si Text, fMRI Acquisition for acquisition details). For the fMRI task-based analyses, we used FIDL software to preprocess and construct the general linear models (developed at Neuroimaging Laboratories, Washington University in St. Louis, version 2.64) (48). For functional connectivity, we implemented validated in-house Matlab software developed at Washington University in St. Louis in our prior work (21, 49-51). For preprocessing and analysis details, see Si Text, fMRI Preprocessing, Si Text, fMRI Analyses, and Si Text, tb-fcMRI.

Computational Modeling. Complete model implementation, interaction between modules, pharmacological implementation, and dependence on parameters are presented in Si Text, Computational Modeling and Figs. 54-56.

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Supporting Information

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SI Text

Subjects. All subjects provided informed consent approved by Yale University's Institutional Review Board. We recruited 58 subjects from the local community by advertisement to participate in the present study. Of those, 40 were eligible and entered the study. Nineteen healthy, neurologically and psychiatrically intact right-handed volunteers (10 male) with a mean ± SD age of 27.5 ± 6.3 y completed the study. Subjects withdrew for the following reasons: (i) necessary time commitment for this study (n = 6); (ii) strong phenomenological response to ketamine (n = 4); (iii) noncompliance with the protocol procedures (n = 1); (iv) minor adverse events (n = 3); (v) concerns over taking ketamine (n = 3); and (vi) rescreened (n = 3). One subject’s fMRI data were unusable. Subjects were first screened using an initial telephone interview and underwent a subsequent diagnostic interview using the Structured Clinical Interview for the DSM-IV (SCID) (1). Subjects also underwent a physical examination by a physician and a drug screen. Subjects were excluded for any psychiatric or major physical illness [e.g., severe endocrine disorder (Cushing syndrome, lupus), heart disorder (past history of heart attacks, angina), or other major systemic medical conditions (kidney, multiple sclerosis, cerebral palsy, blindness, serious physical disability] or chronic/acute condition including any managed by medication, head injury, history of neurological symptoms, loss of consciousness, drug or alcohol dependence, and smoking, as well as family history of psychiatric history and alcohol problems.

Overall Experimental Design. Present data were collected as one part of a larger study examining effects of a positive allosteric modulator (PAM) of the metabotropic glutamate receptor (mGluR) on effects of ketamine. Effects of mGluR were not relevant for the reported effects of the current study but will be reported in subsequent manuscripts. We used a double-blind, placebo-controlled, randomized, within-subjects design. Subjects completed three sessions before which they were randomly assigned to pretreatment of different dose of mGluR PAM [0 mg (placebo), 50 mg, or 180 mg], which they received a day before the scan. The aim of this aspect of the design was to ascertain whether pretreatment with an mGluR PAM exerted an ameliorative effect on the glutamatergic deficit induced via NMDA receptor blockade produced by ketamine administration, motivated by prior results in patients (2). mGluR PAM pretreatment did not alter any reported effects (i.e., behavior, fMRI activation, functional connectivity, or relationship with symptoms) in the context of the present WM task. Therefore, for all reported analyses, we collapsed across the pretreatment condition to increase statistical power and we focused on ketamine-related WM effects explicitly. The order of pretreatment visits was counterbalanced across subjects and spaced by at least 2 wk.

Infusion Protocol. As done in our prior work (3), subjects were administered racemic ketamine (1 mg/mL) i.v. via initial bolus (0.23 mg/kg over 1 min), followed by subsequent continuous target-controlled infusion (0.58 mg/kg over 1 h; plasma target, 200 ng/mL) using a computerized pump (Sigma Spectrum pump; P/N-35162). This resulted in achieved plasma concentrations of 183 ng/mL (±0.77 μM) using the pharmacokinetic parameters of a three-compartment model (4).

Clinical Measures. Subjects underwent clinical ratings the morning before the scan and immediately following ketamine infusion: (i) the Clinician-Administered Dissociative States Scale (CADSS) (5); and (ii) the Positive and Negative Syndrome Scale (PANSS), which is designed to assess positive, negative, and general aspects of schizophrenia psychopathology (6). These scales are extensively validated, standardized, and frequently used to assess schizophrenia symptoms.

Behavioral Results. As noted, the overall design involved two load levels and three levels of pretreatment with an mGluR PAM. No term involving either pretreatment or load interacted with effects of ketamine. Therefore, we report overall effect of ketamine on WM relative to the control task (Fig. 1C and Fig. S1). We computed a repeated-measures ANOVA with task condition (WM vs. control) × infusion (ketamine vs. placebo) as factors. Results revealed a highly significant main effect of task condition [F(1,18) = 48.56; P < 0.0001], as well as a task condition × infusion interaction [F(1,18) = 29.14; P < 0.0001]. Reaction time (RT) results revealed a highly significant main effect of task condition [F(1,18) = 43.79; P < 0.0001]. No other terms were significant for RT. For simplicity, we computed a percent drop in accuracy following ketamine vs. placebo for each subject, across both control and WM tasks separately. These effects are presented in the main text to facilitate visual inspection of ketamine effects on WM (Fig. 1C).

fMRI Acquisition. Functional images were acquired using an asymmetric spin-echo, echo-planar sequence maximally sensitive to blood oxygenation level-dependent (BOLD) contrast (T2*) [repetition time (TR), 1500 ms; echo time (TE), 30 ms; field of view (FOV), 200 mm; flip angle, 15°; voxel size, 3.125 × 3.125 × 5 mm], with 24 axial slices parallel to the AC-PC line. All functional acquisitions lasted 4.15 min and produced 166 volumetric images per BOLD run. Structural images were acquired using a T1-weighted, 3D magnetization-prepared rapid gradient-echo (MPRAGE) sequence [TR/TE/TI, 1500/2.83/800 ms; flip angle, 15°; voxel size (isotropic), 1 mm; FOV, 200 mm] with axial slices parallel to the Anterior Commissure (AC)-Posterior Commissure (PC) line. During each visit, subjects completed the following sequence of scans: (i) MPRAGE scan; (ii) one resting-state BOLD scan (not reported here), during which initial i.v. infusion occurred; and (iii) WM task acquisition across nine BOLD runs always following the initial resting-state scan. This sequence was performed during both saline and ketamine infusions.

fMRI Preprocessing. Preprocessing included: (i) slice-time correction; (ii) removal of the first 6 images from each run to reach steady state; (iii) elimination of odd/even slice intensity differences given interpolated acquisition; (iv) rigid body motion correction; (v) intensity normalization to a whole-brain mode value of 1,000 without bias or gain field correction; (vi) registration using a 12-parameter affine transform of the structural image to a template image in the Talairach coordinate system; (vii) coregistration of fMRI volumes to the structural image with 3-mm resampling; and (viii) smoothing using a 6-mm full-width at half-maximum (FWHM) Gaussian kernel. All preprocessing results were inspected for movement and signal-to-noise (SNR) characteristics. Movement across BOLD runs never exceeded one functional voxel along any axis and no BOLD run had SNR of <100. Furthermore, there was no evidence of lower SNR for BOLD runs during ketamine vs. saline infusions (mean SNR ketamine, 279.03; mean SNR saline, 230.84). As done before (7), we calculated SNR after preprocessing but before atlas transformation (i.e., in each subject’s native space) by obtaining the
mean signal and SD for a given slice across the BOLD run, while excluding all nonbrain voxels across all frames. To further rule out possible effects of head motion on functional connectivity results (8), we have implemented an additional volume censoring (scrubbing) movement correction as reported by Power and colleagues (9, 10) (described comprehensively in SI Text, tbf-cfMRI, Preprocessing and analysis).

fmri Analyses. We used a general linear model (GLM) approach to estimate magnitudes of task-related activity for each voxel. We concatenated all of the BOLD runs across all three pretreatment sessions (i.e., 0, 50, and 180 mg) to estimate a “global” baseline across all three visits. Once concatenated, we estimated effects for 24 regressors: task condition (control vs. WM), infusion (ketamine vs. placebo), treatment (0, 50, 180 mg), and load (2 vs. 4 items). Treatment and load effects were modeled to ensure complete model specification, although reported effects did not interact with these factors. Given the focus on encoding, maintenance and probe phases of the task, we specifically isolated phase-specific activation using an assumed hemodynamic response function (HRF) GLM approach (11), as done in our prior work (12). To facilitate visual inspection, we also computed all activation and deactivation time courses using an unassumed HRF GLM approach across the first 24 frames of each trial (13) (see sections below for GLM details, second level analyses, symptom analyses, and task-based connectivity approach).

Second-level GLM analysis: WM effects. At the second level, we computed appropriate t tests and ANOVAs using the assumed GLM magnitudes for each trial component treating subjects as a random factor. All analyses were computed at the whole-brain level (i.e., voxel-wise) with the appropriate whole-brain type I error correction ascertained via AFNI’s AlphaSim (14). AlphaSim considers voxel-wise and cluster-volume thresholds to establish a false-positive rate of 5%. Only regions corrected at $P < 0.05$ with a combined height and cluster level were considered to be significantly activated or deactivated in the whole-brain analysis. For coordinate reporting purposes, given the large number of active regions meeting a $P < 0.05$ correction in the whole-brain analyses, all identified maps were partitioned using a peak-splitting algorithm such that peaks were considered as separate if they were more than 10 mm apart (15, 16). Confirming the validity of our GLM approach, all of the results using the unassumed GLM analysis did not differ from the results obtained using an assumption-based model (13). All whole-brain analyses were visualized using Caret 5.5 software (http://brainvis.wustl.edu/wiki/index.php/Caret:Download) and projected onto the Population Average Landmark and Surface-based (PALS) atlas (17).

Second-level GLM analysis: ketamine effects. All reported second-level ketamine analyses followed a stringent and principled conjunction approach ensuring maximal specificity of task-relevant effects (see Fig. 1B for visual illustration). First, we identified areas, at each WM phase, showing a significant task condition (WM vs. control task contrast) effect. Second, we identified all regions exhibiting a significant infusion (ketamine vs. placebo) by task condition (WM vs. control) interaction. Next, we corrected these two maps at the appropriate whole-brain type-I error level and formed a conjunction (logical AND) of the two maps. This yielded a set of regions that were independently identified to be modulated by WM (first effect), as well as a modulation of WM by ketamine (second effect) (see Fig. S2 for resulting foci). This approach ensured that all surviving regions are modulated by task condition and that the task signal is modulated by ketamine. Within-subject trial-to-trial relationship with accuracy. We also computed a second GLM for each subject that included an accuracy variable (correct vs. incorrect) as a covariate for each WM trial to enable examination of the within-subjects relationship between behavioral performance and brain activity, as done in our prior work (18). In other words, this allowed us to capture trial-to-trial variability in task response that was associated with correct vs. incorrect WM performance. As with all other analyses, the reported effects were computed using the assumed GLM estimates, but all visualized time courses were computed using an unassumed approach (13) (Fig. 5).

Across-subject symptom analyses. We investigated the relationship between the lack of DMN suppression and clinical symptom measures. Specifically, we averaged the signal across all DMN regions showing both a modulation by WM and by ketamine (i.e., surviving our conjunction approach) for each subject (given no a priori motivation to focus on any one region). Next, we computed the association between the obtained DMN values during WM specifically and standard clinical symptom measures obtained during an interview that took place immediately after the scan. We computed four correlations: (i) negative symptoms using the PANSS Negative Scale; (ii) positive symptoms using the PANSS Positive Scale; (iii) overall psychopathology as derived using the PANSS General Psychopathology Scale (6); and (iv) severity of dissociative symptoms using the sum of all items on the CADSS scale (5). To avoid false positives, we applied Bonferroni correction across these four analyses.

tb-fcMRI. Preprocessing and analysis. We further preprocessed the BOLD time series to remove sources of spurious variance that can drive between-region coupling: (i) high-pass filtering (>0.009 Hz) to remove low frequencies and scanner drift; and (ii) removal of motion correction parameters, ventricle, deep white matter, and global mean (GMS) signals, as well as their first derivatives using the GLM framework. We conducted all subsequent tb-fcMRI analyses on the residual values as done previously (18). We acknowledge that prior studies have shown that GMS removal can possibly induce some negative relationships (19). However, there is competing evidence showing that this preprocessing step is critical to optimize specificity of findings (20) and is widely used in the literature (21). Furthermore, this step was performed in an identical fashion across conditions; therefore, ketamine vs. placebo comparisons cannot be confounded by GMS removal. (iii) We implemented additional volume censoring (scrubbing) movement correction, as reported by Power and colleagues (9, 10), to ensure that head motion artifacts are not driving observed tb-fcMRI effects (8, 22). Briefly, image frames with possible artifactual fluctuations in intensity were identified using two criteria with a procedure suggested by Power et al. (10). First, frames in which the sum of the displacement across all six rigid body movement correction parameters exceeded 0.5 mm (assuming a 50-mm cortical sphere radius) were identified. Second, root mean square (rms) of differences in intensity between the current and preceding frame was computed across all voxels and divided by mean intensity. Frames in which normalized rms exceeded the value of 3 were identified. The frames flagged by either criterion were marked for exclusion, as well as the one preceding and two frames following the flagged frame. Given that the average of two frames was used to compute per trial activity estimates for functional connectivity analysis (see below), trials that had either of the two frames marked for exclusion were omitted from the analyses. For completeness, we present effects both prior and after movement scrubbing (Fig. S7). In addition, we verified that the number of trials remaining after movement scrubbing did not differ across conditions, because one possibility is that movement scrubbing eliminated substantially more trials for the ketamine infusion. To this end, we computed an ANOVA with task condition (WM vs. control) × infusion (ketamine vs. placebo) as factors with the mean number of trials per subject as the dependent measure. There was no significant interaction $F(1,18) = 0.65; P = 0.4$ (not significant). Moreover, the main effect of infusion $F(1,18) = 4.94; P < 0.04$ was actually driven by there being a slightly higher fraction of frames removed from the placebo infusion. This effect is in the opposite direction to that expected by there being more movement-flagged trials for the
ketamine infusion, indicating that the number of removed frames across infusions was not preferentially higher for ketamine.

Next, we computed the average BOLD signal value for the approximate delay period (time points 8 and 9) at each trial for every voxel in the image, as validated in our prior studies (18, 23). As noted, we averaged two time-points to reduce variability attributable to possible outlier frames. Next, we concatenated the values into 4D (brain volume x trial) time series that represented trial-to-trial variability. Extracting only specific time-locked components of the time series, as demonstrated in our prior work (7, 18, 23), ensured that the correlations are driven primarily by trial-to-trial variability and not overall task response. Furthermore, the issue of overall task response driving trial-to-trial variability is minimized given the slow event-related nature of the design.

**Network definition and analysis.** Our hypotheses focused on the relationship between the FP, cingulo-opercular (CO) control systems as defined by Dosenbach et al. (24), and the DMN as defined by Fox and colleagues (25) during the delay phase of WM. We included the CO system to examine specificity of the hypothesized FP-DMN relationship (all regions coordinates listed in Table S5). To control for individual anatomical variability, regions of interest (ROIs) were defined for each individual in two steps: (i) we created spherical ROIs (15 mm in diameter) in standard Talairach space centered on the reported coordinates for each region, as done previously (26); and (ii) we masked the resulting group ROIs with the individual subject-derived FreeSurfer (http://surfer.nmr.mgh.harvard.edu; version 4.1) segmentation of the high-resolution structural image that was previously registered to the standard Talairach space (27). This way, we excluded any voxels within the group-defined ROIs that did not represent the relevant gray matter for a given individual subject. We extracted the time series for each of these ROIs and computed the ROI-ROI correlation matrix across all ROIs for each participant for FP-DMN and CO-DMN pairs at the delay phase of the trial. All obtained correlations for each subject were converted to Fisher Z (Fz) values. Given no prior motivation to focus on any one specific ROI-to-ROI connection, we averaged Fz values across all connections between the nodes of two networks of interest to produce a single “mean Fz” index of network connectivity [as done previously (18)]. Using this mean Fz index as the dependent measure, we computed a two-way repeated measures ANOVA with task condition (WM vs. control) x infusion (ketamine vs. placebo) as factors. All analyses are shown in Fig. S7. The FP-DMN results are also shown in Fig. 4.

**Computational Modeling.** To further relate observed BOLD effects to cellular-level hypothesized effects of ketamine, we constructed a parsimoniouscomputational model of reciprocal interactions between task-activated and deactivated networks during WM. The current simulations are based on prior well-validated biophysically realistic models (28, 29), which are spiking local circuit models capable of WM and decision-making computations. The present model is comprised of two modules, one task-activated and one task-deactivated, each a local circuit of spiking excitatory (E) and inhibitory (I) cells. E cells interact with one another through horizontal connections mediating recurrent excitation via NMDA receptors (model scheme shown in Fig. 3B) and a pool of I cells that mediate feedback synaptic inhibition. We modeled the acute effects of ketamine as a reduction of NMDA conductance for both local and long-range E-I projections.

**Model parameter details.** Each module contains NE = 2,048 pyramidal cells and NI = 512 interneurons. The task-activated module is based on a well-validated model of spatial WM (28). We used the “modulated parameter set” of Compte et al. (28) with the modified E-E connectivity for increased WM robustness: JEE = 1.64 (height of the Gaussian connectivity profile) and σEE = 14.4° (width of the Gaussian connectivity profile). The task-deactivated module contains a homogenous population of E cells and a population of I cells, with the following parameters changes from the modulated parameter set of Compte et al.: JII = 1 and JEE = 1.64 (i.e., recurrent excitatory conductance) to attain a uniform high firing-rate state. Excitatory projections between modules, mediated by NMDA receptors as in recurrent excitatory connections, are unstructured all-to-all, and target both E cells and I cells, to avoid assumptions about preferential anatomical connectivity patterns. Projection strengths from the task-activated module to the task-deactivated module are: gE-E = gE-I = 200/N0 nS. Projection strengths from the task-deactivated module to the task-activated module are: gE-E = gE-I = 60/N0 nS. These strengths were chosen to instantiate appropriate model behavior and patterns of both task-based activation and deactivation. Stimulus input is a current pulse to the E cells in the task-activated module with maximum of 200 nA and Gaussian profile width of 14.4°. Stimulus duration was 4.75 s for simulated BOLD traces as in the experiment (Fig. S3B and Fig. S5C), and 1 s for firing-rate traces (Figs. S4, S5B, and S6). Disinhibition by ketamine was implemented by a 1.25% reduction in the strength of all NMDA conductances onto I cells. Simulations were implemented with the Brian neural simulator (30); code is available from the authors upon request. Firing rate traces are calculated using a 50-ms exponential filter and averaged over 64 neurons (centered at the stimulus location for the task-activated module).

**Interaction between modules.** The interaction between task-activated and task-deactivated modules was modeled as follows: the task-activated module receives task-related sensory input and enables spatial WM through selective persistent firing. The task-deactivated module is characterized by high baseline firing rate at rest and deactivation at task onset, an effect shown across species (6, 31). The task-deactivated module does not have stimulus-selective cells and is only characterized as tonically active or deactivated. Long-range reciprocal projections between modules are from E cells and target both E cells and I cells, with the strengths biased onto I cells so that the net interaction between the modules is inhibitory, inducing anticorrelation in their activities (Fig. 3B). Strong stimulus input selectively excites a subset of E cells in the task-activated module. Activation of the task-activated module sends signals to the task-deactivated module that induce a deactivation attributable to biased projections onto I cells. In turn, deactivation of the task-deactivated module relieves the task-activated module of the inhibition by the task-deactivated module, which was previously shown to occur. These dynamics facilitate storage of the memorandum through persistent firing via reduction of excessive signals in the task-deactivated module. In this way, proper deactivation of the task-deactivated module supports successful WM performance (18).

**Effects of ketamine.** We modeled the acute effects of ketamine as a reduction of NMDA conductance for both local and long-range projections. There are the two sites for this reduction: onto I cells and onto E cells. By selectively reducing NMDA conductance onto one cell type, we explored preferential NMDA receptor antagonism on interneurons by ketamine (32, 33). Reduction of NMDA conductance onto I cells induces disinhibition of the local circuit by impairing the recruitment of feedback inhibition. As a result of disinhibition, the task-deactivated module’s E cells have stronger reverberatory excitation and a higher baseline firing rate. Disinhibition, therefore, impedes suppression of the task-deactivated module by task onset. The task-deactivated module does not deactivate adequately and continues to exert inhibition on the task-activated module, impeding WM delay activity. As described in the main text, it is important to note that ketamine administration likely reduced NMDA conductance onto E cells as well (i.e., E-E conductance). However, exclusively modeling reduction in E-I conductance was sufficient to produce model behavior similar to our observed BOLD effects (Fig. S5). Furthermore, at achieved ketamine concentration (likely less than 50% occupancy), it is possible that ketamine more preferentially reduced conductance of NMDA receptors on inhibitory cells (32), which would be in
to further relate our modeling – in visual working memory: A and Neuroimage of 13 distribution function of the form: We used this hemodynamic response function because it o = 1.25 s, delay Neuroimage f(t) = \left(\frac{1-o}{d}\right)^{p-1} \left[\exp\left(-\frac{1-o}{d}\right) \right]^{p-1} (d(p-1))! The BOLD signal was then calculated by convolving the LFP signal with a sinusoidal signal was then calculated by convolving the LFP signal with a sinusoidal model of LFP has been used successfully to link spiking circuit models to experimental LFP recordings (37). The BOLD signal was then calculated by convolving the LFP signal with a single γ distribution function of the form:

where timescale \( d = 1.25 \) s, delay \( o = 2.25 \) s, and shape parameter \( p = 2 \). We used this hemodynamic response function because it was also used to compute the assumed HRF for distinct task phases in the WM trail from the experimental data (38).

**Local vs. long-range E-I conductance manipulation.** Lastly, it is important to clarify a key insight the microcircuit model provides regarding the breakdown in task-based coordination between brain areas induced by ketamine. In the antagonistic architecture between modules, there are two mechanisms by which reduced E-I strength could potentially disrupt the proper pattern of activation and deactivation: (i) long-range (net inhibitory) connections between modules are weakened, impairing the ability of the task-activated module to shut down the task-deactivated module; and (ii) local disinhibition renders a hyperactive microcircuit less sensitive to the long-range input, so that the already high firing-task-deactivated module cannot be shut down even with an equal-strength, long-range suppressive input. A model implementation that does not instantiate biophysically realistic detail in each module would not suggest which mechanism plays the dominant role. We have carried out further simulations to isolate and test these two mechanisms. We found that local E/I balance is the crucial aspect of the model, suggesting a perspective on the importance of local circuit properties in controlling the nature of large-scale interactions between brain areas during cognitive tasks (Fig. S4).
Fig. S1. Behavioral results. Accuracy (% correct) is shown across both control and WM tasks following placebo (white bars) and ketamine infusions (black bars). Error bars reflect ±1 SEM.

A Regions Modulated by WM and by Ketamine - Encoding

B Regions Modulated by WM and by Ketamine - Delay

Fig. S2. Regions showing effects of WM and modulation by ketamine. All of the shown regions survived the stringent conjunction approach [i.e., both main effect of task condition (WM vs. control) and a task condition x infusion (ketamine vs. placebo) interaction]. Reconstructed time courses are displayed for regions showing effects at encoding (A) and delay (B) phases of the trial. All effects were isolated using an assumed HRF but visualized using an unassumed HRF analysis to allow inspection of time courses across the entire trial. As noted in the main text, no regions at the probe phase survived the conjunction. x, y, and z coordinates above each figure are represented in Talairach stereotaxic space. Coordinates for all regions are also listed in Table S4.
Fig. S3. Illustrating preferential ketamine effects across regions and tasks. (A) Motor cortex region (proximal to BA 4) response is shown for the control task (Left) and WM task (Right). As highlighted by green arrows, the BOLD response across both task conditions at the probe phase, where a motor response is required, is not attenuated by ketamine (in fact it is numerically higher). Note: we plotted the left motor cortex response given that all subjects were right-handed. (B) We also highlight preferential ketamine effects on WM encoding/delay signal in a region activated by the WM task. (Left) No difference is found between ketamine and placebo response during the control task early in the trial and during the probe phase, highlighted with green arrows. (Right) Attenuation of BOLD signal is shown for ketamine relative to placebo during WM encoding/delay phases of the trial (black arrow) but, again, no difference at the probe phase. Together, these results are inconsistent with the possibility that ketamine administration compromised BOLD responses in general but suggest a more preferential disruption of computations critical for WM-related processing.

Fig. S4. Effects of reducing local vs. long-range E-I synapses. (A) Model scheme illustrating effects of a small reduction in NMDA conductance at local microcircuit E-I synapses (red box) vs. long-range between-module E-I synapses (green box), as well as global reduction in E-I synapses (black box surrounding the smaller boxes). (B) Modeling results illustrating reduction in NMDA conductance for global (black), long-range (green), and local microcircuit (red) E-I synapses. In the global case, all NMDA conductances onto interneurons were reduced by 1.25%, as in the main text. For both local and long-range cases, this strength of reduction was the same value (i.e., 1.25%). The local reduction regime produces similar model behavior dynamics as observed in the global reduction regime. In contrast, the long-range reduction regime produces similar model behavior dynamics as observed in the control regime. Therefore, the observed disruption to model performance is not primarily driven by reduced inhibitory coupling between modules but instead driven primarily by local microcircuit disinhibition. That is, under local disinhibition, the already high-firing, task-deactivated module cannot be shut down even though long-range connections are unaffected.
Effects of reducing E-I vs. E-E synapses and simulated BOLD signal. (A) Model scheme illustrating effects of a small reduction in NMDA conductance at E-E synapses (green box) vs. E-I synapses (red box). (B) Modeling results illustrating reduction in NMDA conductance for E-E (green) vs. E-I synapses (red). The E-E manipulation facilitates deactivation of the task-deactivated module, contrary to the experimental results presented in the main text. (C) We juxtapose the predicted BOLD signal (also shown in the main text; Fig. 3B) with the model-generated firing rate traces. As noted, the BOLD signal is derived from the simulated LFP on the time scale comparable to a single WM trial in the experiment. Differences between the simulated and experimental BOLD traces highlighted in the main text likely reflect contributions from multiple cell types that are not instantiated in the model. That is, the model contains only two subtypes of cells that are involved in generating persistent WM-related activity. Nevertheless, the simulated BOLD signal still qualitatively captured the observed task-dependent activation/deactivation that was observed experimentally. For comparison with experimental results, the stimulus duration in C was extended to 4.75 s, as in the experiment. The WM delay was simulated over 16 s as done experimentally. Notes: for reduced E-E conductance case, all NMDA conductances onto pyramidal cells were reduced by 0.5%; for reduced E-I conductance case, all NMDA conductances onto interneurons were reduced by 1.25%. See Fig. 56 for systematic characterization of model dependence on these two parameters. The different magnitudes of BOLD signal deflection across modules are attributable to the different proportions of neurons with a significant activity change (i.e., for the task-activated module only a fraction of preferentially-stimulated cells increase their firing rate and contribute to the LFP).

Parameter space illustrating dependence of model regimes on parameter selection. (A) Dependence of the modules’ delay-period firing rates on reductions to NMDA conductance onto interneurons (G_{EI}) and onto pyramidal cells (G_{EE}). There are generally three observed regimes in this parameter space: (i) along the middle diagonal (approximately equal reduction in E-I and E-E conductances), E/I balance is roughly maintained and the model functions properly during the WM delay, with the task-activated module preserved at high firing rate and the task-deactivated module at low firing rate. (ii) In the lower right portion of the parameter space (preferential reduction of E-E conductance), disinhibition disrupts model function, so that during the delay the task-deactivated module is still at high firing rate and the task-activated module exhibits low firing rate and fails to sustain WM representation. (iii) In the upper left portion of the parameter space (preferential reduction of E-I conductance), both modules exhibit low firing rates during the delay. This is because with the low E/I ratio neither module has sufficient recurrent excitation to sustain a high-firing state. The dashed gray line marks the minimum E/I ratio necessary for the task-deactivated module to sustain a uniform high-firing state before stimulus onset (i.e., before deactivation). Below the line, the module supports a high-firing state before stimulus onset. Above the line, there is insufficient recurrent excitation to support the high-firing state. Therefore, before stimulus onset, both modules are in low-firing states. (B) Model traces corresponding to four selected points from the parameter space: control condition (blue); disinhibition via subtle E-I reduction (cyan) as shown in the main text; subtle E-E reduction (green); and hypothesized higher level of ketamine, which may result in higher E-E and E-I reduction (purple). These selected parameters illustrate that with a subtle E-E reduction (green) model results did not match observed experimental effect in that the task-deactivated module is still successfully suppressed in the model. In contrast, at higher levels of both E-E and E-I reduction (purple), there was a collapse of high-rate states across both modules (as may be expected in anesthesia). Both of these sets of regimes were less consistent with our experimental effects. The visualization method for complex multi-parameter space in A was provided by Dr. Eve Marder and Gabrielle Gutierrez (1).

Fig. S7. Effects of ketamine on tb-fcMRI findings and effects of movement. We illustrate the effects of ketamine (red) vs. placebo (blue) on tb-fcMRI for the FP and default mode network (DMN) (upper graphs), as well as for the cingulo-opercular (CO): DMN network relationships (lower graphs) during the delay phase of WM. We show the pattern of results across the two network analyses without additional volume censoring (scrubbing) movement correction implemented (A and B), after movement scrubbing (1, 2) (C and D), and after additionally removing three subjects for whom movement scrubbing removed a somewhat larger number of trials, resulting in <10 useable trials for any one condition (E and F). As evidenced across all analyses, there was a significant task condition (control vs. WM) x infusion (ketamine vs. placebo) interaction for the DMN-FP networks during the delay phase [F(1,18) = 11.09; P < 0.004], which remained significant even after movement scrubbing [F(1,18) = 6.1; P < 0.025] and after removal of three additional subjects that had <10 trials left for any given condition after movement scrubbing was implemented [F(1,15) = 5.13; P = 0.038]. The effect of ketamine on WM trials was preferential to the FP-DMN network, because there was no task condition (control vs. WM) x infusion (ketamine vs. placebo) interaction across the CO-DMN comparisons (but there was a main effect of infusion for the CO-DMN tb-fcMRI). Error bars reflect ±1 SEM.


### Table S1. Regions showing significant WM effects during the encoding phase

<table>
<thead>
<tr>
<th>x</th>
<th>y</th>
<th>z</th>
<th>Hemisphere</th>
<th>Anatomical landmark</th>
<th>Peak Z statistic</th>
<th>Size (mm³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>−2</td>
<td>61</td>
<td>Right</td>
<td>Middle frontal gyrus (BA 6)</td>
<td>6.02</td>
<td>10,746</td>
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<td>−15</td>
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<td>45</td>
<td>Left</td>
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<td>5.74</td>
<td>10,692</td>
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<tr>
<td>19</td>
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<td>53</td>
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<td>5.60</td>
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<tr>
<td>−30</td>
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<td>51</td>
<td>Left</td>
<td>Precentral gyrus (BA 6)</td>
<td>5.53</td>
<td>7,722</td>
</tr>
<tr>
<td>41</td>
<td>−61</td>
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<td>5.44</td>
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<tr>
<td>29</td>
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<td>5,184</td>
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<td>−32</td>
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<td>48</td>
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<td>5.17</td>
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</tr>
<tr>
<td>35</td>
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<td>50</td>
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<td>5.17</td>
<td>8,775</td>
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<td>−81</td>
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<td>5</td>
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<td>−30</td>
<td>18</td>
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</tr>
<tr>
<td>−42</td>
<td>−37</td>
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<td>4.60</td>
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<td>8</td>
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<td>Cerebellum/ulmen</td>
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<td>3,159</td>
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<tr>
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**Task-negative**

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Table S3. Regions showing significant WM effects during the probe phase

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Table S4. Regions showing both a significant effect of WM, as well as a modulation by ketamine

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Table S5. Independently selected regions used for task-based functional connectivity analyses across the three networks

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