FEATURE ARTICLE
Linking Microcircuit Dysfunction to Cognitive Impairment: Effects of Disinhibition Associated with Schizophrenia in a Cortical Working Memory Model

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Excitation–inhibition balance (E/I balance) is a fundamental property of cortical microcircuitry. Disruption of E/I balance in prefrontal cortex is hypothesized to underlie cognitive deficits observed in neuropsychiatric illnesses such as schizophrenia. To elucidate the link between these phenomena, we incorporated synaptic disinhibition, via N-methyl-D-aspartate receptor perturbation on interneurons, into a network model of spatial working memory (WM). At the neural level, disinhibition broadens the tuning of WM-related, stimulus-selective persistent activity patterns. The model predicts that this change at the neural level leads to 2 primary behavioral deficits: 1) increased behavioral variability that degrades the precision of stored information and 2) decreased ability to filter out distractors during WM maintenance. We specifically tested the main model prediction, broadened WM representation under disinhibition, using behavioral data from human subjects performing a spatial WM task combined with ketamine infusion, a pharmacological model of schizophrenia hypothesized to induce disinhibition. Ketamine increased errors in a pattern predicted by the model. Finally, as proof-of-principle, we demonstrate that WM deteriorations in the model can be ameliorated by compensations that restore E/I balance. Our findings identify specific ways by which cortical disinhibition affects WM, suggesting new experimental designs for probing the brain mechanisms of WM deficits in schizophrenia.

Keywords: disinhibition, NMDAR hypofunction, prefrontal cortex, schizophrenia, working memory

Introduction

A basic principle of cortical computation is the dynamically balanced interaction of excitatory pyramidal cells and inhibitory interneurons (Shadlen and Newsome 1994; van Vreeswijk and Sompolinsky 1996; Shu et al. 2003). Disruption of the ratio of excitation and inhibition (E/I balance) can give rise to profound behavioral deficits and may play a key role in serious mental illnesses such as schizophrenia (Yizhar et al. 2011). Specifically, there is growing support for a hypothesis that cortical disinhibition occurs in schizophrenia, due to the disrupted functioning of inhibitory interneurons, which results in elevated E/I ratio (Lewis et al. 2005; Marin 2012). However, it remains unclear how disinhibition may produce deficits in higher cognition, which are a prominent feature of neuropsychiatric diseases.

Cognitive deficits are postulated to be at the “core” of schizophrenia neuropathology (Elvevåg and Goldberg 2000; Barch and Ceaser 2012), with a severe deficit in working memory (WM), the ability to transiently maintain and manipulate information internally (Goldman-Rakic 1994; Lee and Park 2005). In addition to maintenance over time, robust WM requires shielding internal representations from interference by both internal noise and external distraction. This “filtering” function, in particular, may be severely compromised in schizophrenia (Anticevic et al. 2011).

The prefrontal cortex is a key node in the distributed cortical network recruited during WM (Owen et al. 2005; Fuster 2008) and exhibits altered inhibitory microcircuitry in schizophrenia (Lewis et al. 2005; Bitanihirwe et al. 2009). Primate electrophysiological studies show that persistent firing of prefrontal pyramidal cells supports robust and stable WM representations (Funahashi et al. 1989) that depend critically on E/I balance (Rao et al. 2000). Biophysically realistic computational models have delineated a neural circuit basis of WM activity through 2 key network properties: Strong recurrent excitation to sustain persistent activity and recruitment of lateral inhibition to shape selectivity of representations (Compte et al. 2000; Brunel and Wang 2001).

An open question is whether cortical disinhibition, specifically within a WM microcircuit, can give rise to the types of deficits that may be observed in schizophrenia. To test this hypothesis, we examined the neural and behavioral effects of disinhibition in a spiking neural network model of spatial WM (Compte et al. 2000). Specifically, we induce disinhibition in the model via perturbation of N-methyl-D-aspartate receptors (NMDARs) on inhibitory interneurons, which weakens the recruitment of feedback inhibition. This mechanism may be linked to the pathophysiology of schizophrenia (Lisman et al. 2008; Nakazawa et al. 2012) and possibly accounts for some of the key effects of NMDAR antagonists, such as ketamine, which are a leading pharmacological model of schizophrenia (Krystal et al. 1994).

The model predicts that, at the neural level, disinhibition elevates baseline firing rates and broadens WM activity patterns. At the behavioral level, disinhibition increases deterioration over time of the precision of stored information during WM and increases the range of distractors that can interfere with WM. To
experimentally investigate synaptic mechanisms and test the main model prediction of broadened WM representations under disinhibition, we used behavioral data from healthy human subjects performing a spatial WM match/nonmatch test following ketamine administration. It was found that performance errors increased with ketamine specifically for those nontarget test probes that are similar to the remembered targets, as predicted by the model. Finally, as proof-of-principle, we demonstrate that synaptic compensations that restore E/I balance in the model can ameliorate the effects of disinhibition.

**Materials and Methods**

We used an extensively validated spiking network model of spatial WM, consisting of pyramidal cells and interneurons, structured in a ring architecture (Fig. 1A; Compte et al. 2000; Carter and Wang 2007). Parameters were modified starting from the original “modulated parameter set” of Compte et al. (2000) to produce 1) a narrower persistent activity pattern (adjusting the connectivity profile), 2) drift more comparable with experimental human behavior (Badcock et al. 2008; adjusting background input), and 3) robustness of multistability to small (~1%) decreases in recurrent excitatory conductances (adjusting recurrent conductance parameters). Notably, all reported effects were present in the original parameter set of Compte et al. (2000) and were quite robust to parameter modulations.

**Single-Neuron Models**

Pyramidal cells and interneurons are modeled as leaky integrate-and-fire neurons (Tuckwell 1988), characterized by total capacitance $C_m$, leak conductance $g_L$, resting potential $V_r$, spike threshold potential $V_{th}$, reset potential $V_{res}$, and refractory time $t_{ref}$. For pyramidal cells, $C_m=0.5 \text{ nF}$, $g_L=25 \text{ nS}$, $V_r=-70 \text{ mV}$, $V_{th}=-50 \text{ mV}$, $V_{res}=-60 \text{ mV}$, and $t_{ref}=2 \text{ ms}$; for interneurons, $C_m=0.2 \text{ nF}$, $g_L=20 \text{ nS}$, $V_r=-70 \text{ mV}$, $V_{th}=-50 \text{ mV}$, $V_{res}=-60 \text{ mV}$, and $t_{ref}=1 \text{ ms}$. The sub-threshold membrane potential, $V(t)$, follows:

$$C_m \frac{dV(t)}{dt} = -g_L(V(t) - V_r) - I(t),$$

where $I(t)$ is the total input current to the cell.

### Synaptic Interactions

The network consists of $N_E=2048$ pyramidal cells and $N_I=512$ inhibitory interneurons. Neurons receive recurrent, background, and external inputs. Excitative synaptic currents are mediated by 2-amino-3-(3-hydroxy-5-methyl-isoxazol-4-yl)propanoic acid (AMPA) receptors (AMPARs) and NMDARs, and inhibitory synaptic currents are mediated by $\gamma$-aminobutyric acid type A (GABA$_A$) receptors (GABARs). The total input current to each neuron is:

$$I = I_{\text{NMDA}} + I_{\text{AMPA}} + I_{\text{GABA}} + I_{\text{ext}},$$

where $I_{\text{ext}}$ delivers stimulus input to pyramidal cells. The dynamics of synaptic currents for neuron $i$ follow:

$$I_{i,\text{AMPA}} = \langle V_i - V_E \rangle \sum_j g_{j,i,\text{AMPA}} s_{j,i,\text{AMPA}};$$

$$I_{i,\text{NMDA}} = \langle V_i - V_E \rangle \frac{1}{1 + [\text{Mg}^{2+}] \exp(-0.062 V_i/5.57 \text{ mV})};$$

$$I_{i,\text{GABA}} = \langle V_i - V_I \rangle \sum_j g_{j,i,\text{GABA}} s_{j,i,\text{GABA}};$$

where $V_E=0 \text{ mV}$ and $V_I=-70 \text{ mV}$, and $g_{j,i,\text{syn}}$ denotes the synaptic conductance strength on neuron $i$ from neuron $j$. NMDAR-mediated currents exhibit voltage dependence controlled by the extracellular magnesium concentration ($[\text{Mg}^{2+}]=1 \text{ mM}$ (Jahr and Stevens 1990)).

Given a spike train $[\delta(t)]$ in the presynaptic neuron $j$, the gating variables $s_{j,i,\text{AMPA}}$ and $s_{j,i,\text{GABA}}$ for AMPAR- and GABAR-mediated currents, respectively, are modeled as:

$$\frac{ds}{dt} = \frac{-s}{\tau_x} + \sum_k s_k \delta(t - t_k);$$

$$\frac{dx}{dt} = \frac{-x}{\tau_x} + \sum_k s_k \delta(t - t_k);$$

$$\frac{ds}{dt} = \frac{-s}{\tau_x} + \alpha_x (1 - s);$$

with $\alpha_x=0.5 \text{ kHz}$ and $\tau_x=2 \text{ ms}$. The decay time constant $\tau_x$ is 2 ms for AMPA, 10 ms for GABA, and 100 ms for NMDA. For simplicity, background inputs are mediated entirely by AMPARs, and recurrent excitatory inputs are mediated entirely by NMDARs, as they are critical for the stability of persistent activity (Wang 1999; Compte et al. 2000). Background input to each neuron is provided by an independent Poisson spike train with the rate of 0.6 kHz and AMPAR conductances of 9.3 nS on pyramidal cells and 7.14 nS on interneurons.

### Network Model

As noted, pyramidal cells are organized in a ring architecture and are tuned to the angular location on a circle ($0-360^\circ$), with uniform distribution of their preferred angles. The network structure follows a columnar architecture, such that pyramidal cells with similar stimulus selectivity are preferentially connected to each other. The synaptic conductance on neuron $i$ from neuron $j$, $g_{j,i,\text{syn}} = W(\theta_j - \theta_i) G_{\text{syn}}$, where $\theta_i$ is the preferred angle of neuron $i$, and $W(\theta_j - \theta_i)$ is the connectivity profile normalized such that:

$$\int_{0}^{360} d\theta W(\theta)d\theta = 1.$$

For pyramidal-to-pyramidal connection, $W(\theta_j - \theta_i) = J^+ + J^- \exp [-(|\theta_j - \theta_i|)^2/2\sigma^2]$. We take $J^+=3$ and $\sigma=9^\circ$. All other synaptic connection profiles are unstructured. Synaptic conductance strengths are given by $G_{\text{ER}}=1003.9/N_E \text{nS}$, $G_{\text{EI}}=717.6/N_E \text{nS}$, $G_{\text{II}}=807.2/N_I \text{nS}$, and $G_{\text{II}}=566.2/N_I \text{nS}$. The recurrent excitatory conductances, $G_{\text{ER}}$ and $G_{\text{II}}$.##
are parametrically reduced to study the effects of altered E/I ratio on neural activity and behavior.

**Stimulus**
As previously validated (Compte et al. 2000), we followed the stimulus protocol of Funahashi et al. (1989). Inputs are modeled as injected current with a Gaussian profile, I(θ) = J0 exp[−[(θ − θ0)/r]2/2r2], where the maximum current J0 = 375 pA, θ0 is the stimulus location, and the width parameter r0 = 6°. WM robustness to external disruption was tested with distractors. That is, we examined how stable the WM trace is, for control versus disinhibition conditions, when explicitly perturbed by an additional distracting input. Distractors were modeled identically to the initial cues, with same intensity and duration, but with a different stimulus location, such that the distractor appeared at a given angle relative to the original cue. Such a distractor manipulation allowed for the testing of specific hypotheses under disinhibition (discussed below).

**Synaptic Manipulations**
Disinhibition is implemented through a reduction of NMDAR conductance only on interneurons, G_{EI} (Fig. 1B). For all simulations except those in Fig. 3, we make the approximation of reducing recurrent excitatory conductance only on interneurons, since that site is hypothesized to be the primary site of the action of NMDAR antagonists such as (Krystal et al. 2003; Kotermanski et al. 2009). To test the ability of disinhibitions to restore E/I balance, we modeled 2 other synaptic manipulations: 1) the reduction of presynaptic glutamate release, by decreasing α0 in the glutamatergic synaptic update (Equation 8); 2) the enhancement of GABAR conductance on pyramidal cells, by increasing G_{IE}. If disinhibition results in elevated E/I ratio, the glutamatergic or GABAergic compensation can restore E/I balance by reducing excitation or increasing inhibition, respectively.

**Simulation**
Simulations were implemented with the Brian neural simulator (Goodman and Brette 2008). Simulation code is available from the authors upon request.

**Analysis of Simulation Data**
To decode the behavioral report from neural activity, we use the population vector approach (Georgopoulos et al. 1986): The angular location encoded by the network is given by θ_{report} = arg[∑i r_i exp(i θ_i)]. To visualize the scatter of WM reports in Figure 4D, we used the method described in Renart et al. (2003) to convert 1-dimensional (1D) angular position in the model into a 2-dimensional position. Population vectors were computed with a 50-ms temporal window. As done previously, spatiotemporal plots of network activity were smoothed over time (50 ms) and the pyramidal-cell population (40 neurons; Compte et al. 2000).

To measure the tuning width of the WM population activity pattern (bump attractor), we fit the firing rate profile of the population using a von Mises profile with sigmoidal saturation: r(θ) = r0 + r1[1 + exp(−B4·exp(a·cos(2π(θ − θ0)))))]−1. The bump width calculated as the full width at half maximum of the fit curve. Firing rate profile fits were computed with a 500-ms temporal window.

**WM Task with Ketamine Manipulation**
Complete experimental details are reported in Anticevic et al. (2012). Briefly, healthy human subjects performed a previously validated delayed spatial WM task (Leung et al. 2002; Driesen et al. 2008). Task details are described in Fig. 6A. Two or 4 targets are sequentially presented, followed by a 16-s delay period during which the target locations are to be held in WM. At the end of the delay, a probe is presented, either at a target or nontarget location, and subjects provided a button-press response of match or nonmatch, respectively. Subjects performed a series of such WM trials first during a session of saline infusion to act as a control, followed by a low-dose ketamine administration (achieved plasma concentration of 183 ng/mL (0.77 μM) using the pharmacokinetic parameters of a 3-compartment model (Domino et al. 1982). Subjects completed a total 32 WM trials and 16 control task trials with no WM encoding and maintenance requirement but requiring a probe response, to control for motor effects per infusion (64 WM and 32 control trials in total). The experiment also included 2 additional randomized sessions in which subjects were pretreated with 2 dosages of a positive allosteric modulator of Group II metabotropic glutamate receptors (Patil et al. 2007). Behavior from these pretreatment sessions was not included in present analyses, as it does not directly relate to model predictions (due to multiple compound interactions), and will be reported elsewhere. As is standard, statistical practice (Howell 2012), here, we report a simple effect for the factor involving no pretreatment, explicitly relevant to the model predictions.

**Match/Nonmatch Simulation**
To allow for a comparison between simulated and experimental ketamine data, we implemented a simple model of the comparison and decision computations involved in a WM match/nonmatch task (Engel and Wang 2011). For the firing rate profile of the WM bump, we use the fit function described above for calculating bump width. Next, we center that profile at the target location, removing the effect of drift. The probe representation is given by a Gaussian profile centered at the probe angle θ_p = exp[−((θ − θ_0)^2/2σ_p^2)], with σ_p = 6°. Following recent theoretical models (Tagamets and Horwitz, 1998; Engel and Wang 2011), the match/nonmatch decision is based on the overlap between target and probe representations. We calculate such overlap by taking the dot product of the 2 profiles. This overlap is normalized to be 1 when the probe is at the target location with control network parameters, and 0 when the probe is 180° away with control network parameters. To model the downstream decision circuit, the probability of match response, P_M, is then given by a sigmoidal function of the normalized overlap, x: P_M(x) = p_0 + (p_1 − p_0)/(1 + exp(−(x − x_c)/σ))^−1. To replicate the experimental error rates, we set the parameters of the decision circuit to P_0 = 0.18, P_1 = 0.78, x_c = 0.44, and σ = 0.075.

**Results**
The network model, consisting of interconnected excitatory pyramidal cells and inhibitory interneurons, represents a local microcircuit in prefrontal cortex. The pyramidal cells exhibit selectivity to angular location on a circle (Fig. 1A; Funahashi et al. 1989). To observe WM, the circuit exhibits multistability (Amit and Brunel 1997; Wang 1999; Compte et al. 2000). In addition to supporting a uniform, low-firing baseline state, the circuit allows for a continuous family of WM states that are supported by the strong recurrent excitation and capable of encoding stimulus location across a delay. A brief localized stimulus can activate a bell-shaped persistent activity pattern (bump attractor), consistent with location-selective sustained activity observed in prefrontal neurons during the delay period of WM tasks (Funahashi et al. 1989). Excitation in the network is dynamically balanced with inhibition, since increased pyramidal cell activity enhances the activity of interneurons that deliver feedback inhibition.

**Disinhibition of Neural Activity**
We implemented disinhibition in the model by reducing the strength of NMDAR conductance on interneurons, G_{EI} (Fig. 1B). Here, we set our disinhibition condition as G_{EI} reduced by 3.25% (for a complete parameter space characterization of our disinhibition manipulation, Fig. 3). Figure 2A shows spatiotemporal plots of the network activity during a...
WM trial for control and disinhibition conditions. Before the stimulus is applied, the network is in the uniform baseline state. A brief stimulus excites a subset of pyramidal cells selective to the stimulus angle. After removal of the stimulus, the stimulus angle is encoded throughout the delay by a persistent WM bump. Disinhibition substantially broadens the firing-rate profile of the bump (Fig. 2B, C). Because inhibitory interneurons are less strongly recruited by pyramidal-cell activity, more pyramidal cells can be activated by recurrent collaterals. Notably, network multistability is preserved at this level of disinhibition (i.e. the baseline state and WM bump states remain stable).

We also examined the effects of disinhibition on the spontaneous firing rate (Fig. 2D). As expected, disinhibition increases firing rates for pyramidal cells, since feedback inhibition is weakened. For strongly recurrent networks, excitation is dynamically balanced by inhibition via the recruitment of interneurons by pyramidal cells. Disinhibition also increases the firing rates of interneurons, though to a lesser degree than in pyramidal cells. In a strongly recurrent network such as ours, even though the strength of excitatory synaptic conductance to interneurons is reduced, their firing rates can still increase. This is because the disinhibited pyramidal cells increase their firing rates by a large enough proportion that consequently the net excitatory input to interneurons is also increased (Tsodyks et al. 1997).

Persistent WM activity patterns depend on both targets of recurrent excitatory synapses, interneurons, and pyramidal cells. Pyramidal-to-pyramidal synapses, with strength $G_{EE}$, are critical for the recurrent excitation necessary for persistent activity (Wang 1999). NMDAR antagonists likely act on both interneurons and pyramidal cells, although they may affect postsynaptic cell types differentially (Kotermanski and Johnson 2009). For the sake of parsimony in the model, all recurrent excitatory synapses are mediated by NMDARs. This approximation allows the direct comparison of the network's structure and function.

Figure 2. Disinhibition broadens WM representation. (A) The spatiotemporal plot of persistent activity patterns for control (upper panel) and disinhibition (3.25% reduction of $G_{EI}$ condition (lower panel). A stimulus is presented at $90^\circ$ for 250 ms and, during the subsequent delay, the stimulus location is encoded by persistent activity of a WM bump. Disinhibition broadens the network activity pattern. (B) The firing rate profile of the WM bump for control and disinhibition conditions averaged over 500 ms in a single trial. (C) Increase of bump width as a function of the degree of disinhibition. (D) Increase of baseline firing rates as a function of the degree of disinhibition for both pyramidal cells and interneurons, from their control values of 1 and 6.4 Hz, respectively. Error bars show standard error of the mean.
sensitivity to changes in the net E/I ratio, but not the dissection of the relative roles of recurrent AMPARs and NMDARs. The tolerance of the model to NMDAR perturbations would be greater if a large fraction of the recurrent excitatory conductance was mediated by AMPARs rather than NMDARs. This is important in light of recent studies showing that the AMPAR/NMDAR ratio differs between pyramidal cells and fast-spiking interneurons (Wang and Gao 2009; Rotaru et al. 2011). To explicitly examine the model's sensitivity to E/I ratio, we parametrically decreased both \( G_{\text{EI}} \) and \( G_{\text{EE}} \), while testing multistability and measuring bump width (Fig. 3).

Here, we found that if \( G_{\text{EI}} \) and \( G_{\text{EE}} \) are reduced together in a particular proportion to each other, the bump width is unaltered, that is, E/I balance is maintained. This suggests that E/I ratio is the critical parameter for network function, rather than the absolute strength of either \( G_{\text{EI}} \) or \( G_{\text{EE}} \) alone. If \( G_{\text{EE}} \) is reduced in greater proportion, then the network is in a state of disinhibition, and the WM bump width is increased (lower right corner in Fig. 3). In contrast, if \( G_{\text{EE}} \) is reduced in greater proportion, the bump width is decreased (upper left corner in Fig. 3). These bump-width effects are present within the parameter region of network multistability around a diagonal line in the parameter space of \( G_{\text{EI}} \) and \( G_{\text{EE}} \). Taken together, these results show that the net effect of mild reductions in excitatory drive on neural WM activity can be described by the net change to E/I balance.

If E/I imbalance is substantial, either too elevated or too reduced, we observe a loss of network multistability. If disinhibition is too strong (via elevated E/I ratio), then the spontaneous state is no longer stable, and multistability is lost (lower right corner of Fig. 3). In that regime, a WM bump spontaneously appears at random locations in the network in the absence of an external stimulus. Conversely, if recurrent excitation is too weak (via reduced E/I ratio), then the circuit cannot support persistent activity. In that regime, the persistent activity pattern collapses during the delay period and the memory is lost (upper left corner in Fig. 3). Of note, the loss of multistability that we observe, after only a moderate reduction of \( G_{\text{EI}} \) or \( G_{\text{EE}} \), may result from the model's requirement that a single microcircuit be capable of both multistability and continuous representation of a memorandum. An extended model, which separates bistability and representation between different populations of cells, may exhibit greater robustness of multistability against disruption of E/I balance. However, the sensitivity of bump width to E/I ratio would still be present in such a configuration.

**Disinhibition Increases Random Drift and Decreases Memory Precision**

For robust WM, the precision of the encoded memorandum is critical and should be maintained throughout the delay. Stochastic activity internal to the neural system may interfere with maintenance, causing WM precision to decline continuously with increasing delay duration. In memory-guided saccade tasks, variability of saccade endpoints increases with delay duration (Funahashi et al., 1989; White et al. 1994; Ploner et al. 1998). Individuals with schizophrenia exhibit greater variability in memory-guided saccades, due in part to impaired maintenance (Badcock et al. 2008). Motivated by the observation that individuals with schizophrenia exhibit degraded precision during visuospatial WM maintenance, we characterized the effect of disinhibition on behavioral variability in the model. Specifically, we tested the time course of the variance of the WM report encoded in the network activity pattern. That is, we examined how the variability in WM report changes over time as a function of disinhibition, which would imply increased vulnerability to noise internal to the neural system.

Similar to experimental findings, the model exhibits a random drift of the angular location encoded by the WM bump. We extract the encoded angle, or report, from the population vector, which is the center of mass of the network activity pattern (Georgopoulos et al. 1986; see Materials and Methods). During the delay period, the encoded angle undergoes random drift, due to internal noise introduced by the Poisson background input. In essence, the network possesses a continuous family of possible bump states, so noise can shift the bump along this continuum. As the network integrates noise, the variability of report increases with delay duration (Compte et al. 2000; Chow and Coombes 2006; Carter and Wang 2007).

Figure 4A shows example traces of the encoded angle for control and disinhibition conditions. As expected, the variance of the encoded angle grows throughout the delay (Fig. 4B). We found that disinhibition increases the rate at which the variance grows (Fig. 4BE). The increased variance is due to a deficit in maintenance rather than in initial encoding of the stimulus, since the variance at the start of the delay is similarly low for the 2 conditions. This effect of disinhibition can be understood through the effect of E/I ratio on bump width. Random drift occurs because the active neurons in the WM bump receive noisy background input. Pyramidal cells that are far from the bump are hyperpolarized by lateral inhibition and, therefore, farther from spike threshold than are cells that constitute the active flanks of the bump. As a result, background noise is more likely to produce output spikes in the depolarized cells on the flanks, consistent with an expansive nonlinearity in the firing rate versus mean current curve (e.g. Roxin et al. 2011). These noise-induced spikes then recurrently propagate to drive nearby cells and
potentially to shift the bump’s center of mass. A broadened bump under disinhibition is therefore subjected to more background noise input.

For visualization, we converted the variance of encoded angle into a 2D scatter of the memory report (Renart et al. 2003; see Materials and Methods), illustrating the degraded precision under disinhibition (Fig. 4C). If we introduce an accuracy threshold for correct responses (indicated with gray circles in Fig. 4C), the error rate is increased under disinhibition (Fig. 4D). This greater loss of WM precision over time captures prior experimental observations of behavior in schizophrenia (Badcock et al. 2008).

**Disinhibition Increases the Range of Vulnerability to External Distractors**

In addition to robust stimulus maintenance across a delay, a neural circuit that subserves WM should be resistant to intervening external distractors during the delay (Jonides et al. 2008). There is a rich literature suggesting that schizophrenia is associated with deficits in gating of external stimuli (Geyer et al. 2001; Turetsky et al. 2007) and abnormalities in resistance to distractors during WM (Hahn et al. 2010; Anticevic et al. 2011). In experimental settings, the effect of a distractor on WM appears to be largely dependent on the similarity between a distractor and the target held in memory. In visuospatial WM, similarity is determined by the spatial separation between the target and the distractor. Prior experimental studies in healthy human subjects found that a distractor can attract the memory report toward its location, but only if the distractor appears within a “distractibility window” around the target location (Van der Stigchel et al. 2007; Herwig et al. 2010), as predicted by the similarity hypothesis. In other words, WM report can be perturbed toward the distractor if target–distractor similarity is high enough (i.e. the distractor is near the target and overlaps the representation), but is unperturbed if target–distractor similarity is low enough (i.e. the distractor is far from the target and does not overlap the representation).

**Figure 4.** Disinhibition degrades memory precision due to increased noise-induced drift. (A) Decoding by the population vector of the WM activity pattern drifts across a 6-s delay. Left: Shown are 5 example traces each for control (blue) and disinhibition (orange). Right: Histogram of reports at the end of a 3-s delay. (B) The variance of the encoded angle fluctuations grows with time during the delay. Variance starts from a similar value for the 2 conditions, but the variance grows at a higher rate under disinhibition. (C) Visualization of reports after a 3-s delay for a memory-guided saccade task, with 96 trials at each of 8 stimulus angles. Gray circles mark the accuracy equivalent to within ±10° in the 1-dimensional model. (D) Error rates. Errors are counted if the reports shown in C lie outside of their corresponding gray circles. (E) Increase of the report variance, after a 3-s delay, as a function of the degree of disinhibition.
similarity is low (i.e. the distractor is far away from the target and can be effectively filtered).

Previous work has shown that the present computational model possesses an inherent resistance to distractors (Compte et al. 2000; Gruber et al. 2006). Interestingly, model behavior also exhibits a dependence on target–distractor similarity. In that sense, there are 2 qualitative types of distractors, “near” and “far”. When the distractor is spatially far from target, there is no overlap between the neural representations of the distractor and of the target in WM. In this regime, the interaction between the memory and distractor is “winner-take-all” competition mediated by lateral inhibition: If the distractor strength is low, the distractor will have negligible effect on the report, but if the distractor strength is too high, the memory bump will switch completely to the distractor location. In contrast, when the distractor is near the target, there is overlap between the distractor and target representations. If the distractor input excites neurons on the flanks of the bump, the target–distractor interaction is better described as “vector-averaging”, and the resulting bump will lie at an intermediate location (Compte et al. 2000; Liu and Wang 2008). We define the model’s distractibility window as the angular separation that results in the largest deviation in the report from the original cue location.

Since both experimental results in humans and model results exhibit a distractibility window within which distractors are able to attract the WM report, the model readily allows us to study the neural circuit basis of increased distractibility by disinhibition. In the model, the distractibility window is dependent on the width of the WM bump. As noted, attraction by the distractor occurs when there is overlap between the neural representations of the target in WM and of the distractor. Since disinhibition broadens WM representations, we specifically hypothesized that the distractibility window will be similarly expanded. This would increase the range of locations at which distractors can disrupt WM following disinhibition.

![Figure 5](http://cercor.oxfordjournals.org/)
As predicted, disinhibition increases the width of the bump such that a far distractor is transformed to a near distractor (Fig. 5A). Put differently, the distractibility window increases under disinhibition (Fig. 5B,C), and disinhibition therefore widens the range of distractors that can disrupt WM. This model finding also suggests that manipulating target–distractor similarity (to measure the distractibility window) may be a highly sensitive experimental probe of changes in the neural WM representation (see Discussion).

In addition to widening the distractibility window, disinhibition changes the shape of the distractibility curve shown in Figure 5B. Interestingly, for an intermediate range within the distractibility window, disinhibition decreases the magnitude of deflection toward the distractor, which may be attributed to greater “inertia” of the broadened bump. The shapes of the distractibility curves in Figure 5B can be roughly decomposed into 4 regions: 1) very close (<20–30°) where neither network loses fidelity; 2) near (approximately 30–75°) where the control network is more distractible than the disinhibited network; 3) medium-range (approximately 75–120°) where the disinhibited network is affected by the distractor but the control network is not; and 4) distal (>120°) where neither network is affected by the distractor. We know of no relevant experimental studies that have combined 1) parametric control of target–distractor similarity; 2) direct, continuous report; and 3) direct comparison between control condition and a condition of putative disinhibition. The characterization of how disinhibition affects the similarity dependence of distractibility should be addressed by future psychophysical studies in healthy adults, clinical populations, and animal models.

**Experimental Evidence for Broadened WM Representation**

As described above, the core model prediction is that WM representations are broadened under disinhibition, due to disrupted E/I balance. We tested this prediction experimentally in humans using a pharmacological manipulation that is hypothesized to induce disinhibition (Anticevic et al. 2012). Specifically, we employed NMDAR antagonism via acute, low-dose administration of ketamine. As a pharmacological model of schizophrenia, ketamine allows for controlled and transient NMDAR manipulation in healthy volunteers (Krystal et al. 2003). Ketamine-induced neural disinhibition effects can be observed in both in vitro and in vivo (Greene 2001). While the precise pathways by which ketamine induces disinhibition are not fully resolved, at the low concentrations employed in human experimental work, there is evidence that its effects may involve a preferential blockade of NMDAR subunits (Kotermanski and Johnson 2009) that are expressed more in interneurons than in pyramidal cells (Xi et al. 2009).

In the experiment, healthy subjects performed a delayed spatial WM match/nonmatch task, following either saline (control) or ketamine infusion. In this task, subjects encoded multiple target locations, and after a delay period, indicated whether a probe was presented at a target location (match) or nontarget location (nonmatch; Fig. 6A). There are 2 types of errors that can occur in this task: Misses (nonmatch response to a probe at a target location), and false alarms (match response to a probe at a nontarget location).

In neural circuit models of match/nonmatch decision making, the probability of match response is a monotonically increasing function of the overlap between the neural representations of the probe and of the target in WM (Tagamets and Horwitz 1998; Engel and Wang 2011). For nontarget probes, the rate of false alarms increases with the overlap between the target memory and probe representations. Put simply, a match response to a nontarget probe is more likely when there is a larger overlap between the probe and the target memoranda.

Therefore, changes to the WM representation by disinhibition can potentially alter the target–probe overlap and thereby affect the pattern of responses. The model predicts that the increase in errors is selective, with an increased rate of false alarms when the probe is presented at near, but not far, nontarget locations. This is because the overlap between the target WM representation and the probe representation is increased substantially when the probe is presented at near nontarget locations (due to increased bump width), but not when presented at far nontarget locations. Similarly, the model predicts that misses should not increase since the overlap does not substantially decrease when the probe is presented at the target location. In other words, because the peak firing rate of the WM bump does not change substantially following disinhibition, there is similar overlap for a probe presented at the target location.

Figure 6B shows the effects of ketamine on error rates for false alarms to near nontarget probes, false alarms to far nontarget probes, and misses to target probes. To quantitatively test the model prediction using experimental data, we computed a repeated-measures ANOVA with the number of false alarms per subject as the dependent measure, and infusion (saline vs. ketamine), and distance (near vs. far nontarget probes) as within-subject factors. Experimental results revealed a significant infusion × distance interaction ($F_{1,19} = 7.55, P < 0.015$). As predicted, the interaction effect was driven by a significantly greater number of false alarms following ketamine infusion for near but not far distraction ($t_{19} = 4.09, P < 0.0007$, Cohen’s $d = 1.1$), whereas this effect was substantially attenuated and did not reach significance for placebo trials ($t_{19} = 0.99, P = 0.33$, Cohen’s $d = 0.25$). Moreover, when examining miss trials there was no significant difference across infusions ($t_{19} = 0.72, P = 0.35$). This selective increase in the number of errors for false alarms to near nontarget probes, but neither for false alarms to far nontarget probes nor for misses, is in line with the model prediction.

Moreover, as an explicit demonstration of the model prediction, we used disinhibition in the model to generate the selective pattern of errors we observed experimentally (Fig. 6C,D). The match/nonmatch decision is based on the overlap between the neural representations of the target in WM and of the probe (see Materials and Methods). The probability of match response is taken as a sigmoidal function of this target–probe overlap. As shown in Figure 6D, broadening of the WM bump by disinhibition is sufficient to capture the pattern of errors observed experimentally, with a selective increase in false alarms to near nontarget probes, but little change in false alarms to far nontarget probes or in misses. Of note, Figure 6B shows error rates for trials of each probe type for which subjects responded, to allow comparison to model performance (Figure 6D).
Interestingly, a recent psychophysical study using a similar spatial WM match/nonmatch task found that, relative to healthy controls, individuals with schizophrenia also exhibit a selective increase in false alarms for near nontarget probes but not for far nontarget probes (Mayer and Park 2012). In addition to its effects on the maintenance of WM representations, NMDAR antagonism may also induce deficits downstream of WM, in the neural circuitry implementing the target–probe comparison and match/nonmatch decision. Future experimental and modeling studies should differentiate deficits due to WM from deficits due to decision making. Furthermore, a circuit model that incorporates the diversity of functional cell types in prefrontal cortex may be necessary to explain the diverse effects of NMDAR antagonists at both the neural and behavioral levels.

Restoring E/I Balance Ameliorates Deficits

The model suggests that E/I balance is the crucial parameter that determines WM performance. As illustrated in the parameter space characterization in Figure 3, if E/I balance is disrupted by a change at one site in the network, it can be restored through a compensatory change at another site (e.g. reduction in $G_{EE}$ compensating for reduction in $G_{EI}$). Therefore, we explicitly tested the hypothesis that compensations that restore E/I balance in the model can ameliorate the observed WM deficits induced by disinhibition.

To this end, we implemented 2 compensations, modifying glutamatergic and GABAergic neurotransmission, respectively. Our first compensation is to decrease the strength of presynaptic glutamate release at recurrent synapses. Reducing presynaptic glutamate release has been proposed as compensation for the elevated E/I ratio and behavioral symptoms arising from schizophrenia (Large et al. 2005) and NMDAR antagonists (Moghaddam and Adams 1998; Anand et al. 2000; Brody et al. 2003; Krystal et al. 2005). Our second compensation is to increase the inhibitory conductance onto pyramidal cells (i.e. $G_{IE}$). This in effect strengthens the recruitment of inhibition, which is weakened by the disinhibition manipulation in the model. Lewis et al. (2004) proposed that this could be realized using activators of the α2 subunit of the GABAR (which mediates perisomatic inhibition to the axon initial segment of pyramidal cells), and an α2-subunit agonist was found to ameliorate WM deficits induced by ketamine in monkeys (Castner et al. 2010).

We examined whether these proof-of-principle compensations affected 4 key model properties outlined above: 1) baseline firing rate, 2) bump width, 3) drift due to internal noise, and 4) distractibility window in response to external
distractors. Implemented in conjunction with our disinhibition manipulation, both compensations reduced the width of the bump attractor (Fig. 7A, B). We set the strength of the compensation such that bump width was restored to near its control value, and then characterized neural and behavioral effects.

In addition to restoring bump width, both compensations reduce the baseline firing rates from their values under disinhibition (Fig. 7C). Both compensations therefore ameliorate the neural changes induced by disinhibition, both at baseline and during active WM maintenance. We observed that the decrease in baseline firing was smaller for the GABAergic compensation, indicating that strengthening GABAR conductance on pyramidal cells is more effective at reducing bump width than at reducing baseline firing rates in the model.

At the behavioral level, both compensations ameliorated WM deficits observed under disinhibition. Report variance and distractibility windows reduced toward their control values (Fig. 7D,E). These manipulations are a simple proof-of-principle, rather than detailed models of any given experimental or actual treatment manipulation. However, that these 2 manipulations restored behavior, despite different sites of action, highlights that the critical parameter of the network is E/I ratio, which controls bump width.

It is also important to note the magnitudes of the 2 compensations. We implemented a 25% decrease in presynaptic glutamate release for the first compensation, but only a 2% increase in GABAR conductance on pyramidal cells for the second compensation. This order-of-magnitude difference reflects that the glutamatergic compensation in the model acted both on pyramidal cells (decreasing E/I ratio) and on interneurons (increasing E/I ratio). Because these 2 changes partially counteract one another, the level of the manipulation must be relatively large to restore E/I balance. In contrast, because the GABAergic compensation in the model acted only on pyramidal cells, a relatively lower level of manipulation is required to restore E/I balance. Together, this suggests that compensations with greater specificity, with strong changes in either pyramidal cells or interneurons but not both, may be more efficacious than those that act equally on pyramidal cells and interneurons.

**Discussion**

In this study, we examined the neural and behavioral effects of disinhibition in a cortical microcircuit model of WM. We found that recurrent circuit function is highly sensitive to perturbations of E/I balance; a slight weakening of inhibition
results in marked neural and behavioral changes. The salient neural changes are elevated E/I ratio and broadened WM representation. They are linked to behavioral deficits, namely increased susceptibility to both internal noise and external distraction. Our proof-of-principle compensations further highlight that E/I balance is a critical property of the model that relates to observed deficits induced by disinhibition.

This work focuses on the consequences of a reduced overall strength of connections from excitatory cells to interneurons, which could be related, at least in part, to interneuron dysfunction hypothesized to operate in schizophrenia (Lewis et al. 2005; Lisman et al. 2008; Marín 2012). Previous modeling studies have suggested that NMDA receptors on pyramidal cells may be important for providing sufficiently strong recurrent synaptic excitation that underlies persistent activity during WM (Wang 1999; Durstewitz et al. 2000). The present work provides a different, complementary perspective that emphasizes the importance of the width of the population firing pattern, rather than the peak firing rate. The width of the WM bump is controlled by E/I balance. Increasing E/I ratio, and thereby bump width, degrades WM robustness in a manner that yields specific types of errors. Such errors are dissociable from those predicted by a complete loss of WM maintenance, which may result from a collapse of persistent activity due to insufficient recurrent excitation.

Implications for Task Design

Present findings offer important implications for the paradigm design of WM tasks used to study cognitive dysfunction in neuropsychiatric disease. First, the present model suggests that a continuous, veridical measure of the report is sensitive to detecting degraded precision due to drift. This increase in scatter may be subtle and difficult to detect with a categorical (e.g. match/nonmatch) design. The veridical report of memoranda can also help to distinguish deficits in maintenance from potential deficits in the downstream target–probe comparison, and match/nonmatch decision computations inherent in the match/nonmatch paradigm (Lencz et al. 2003; Zhang and Luck 2008). Building on this continuous measure of report, future studies can vary delay duration to better distinguish increased delay-dependent drift during maintenance from impaired encoding (Lee and Park 2005; Wei et al. 2012).

Our findings also underscore that target–distractor similarity is a critical parameter in determining the effect of distractors on WM (Van der Stigchel et al. 2007; Herwig et al. 2010). To characterize how this changes under disinhibition, future studies should employ stimulus sets with well-defined target–distractor similarity and manipulate target–distractor similarity parametrically. The veridical report is critical in this setting, in order to measure how distractors alter the memory of the target. We find that the distractibility window varies with bump width as E/I balance is perturbed. This effect suggests an interesting interpretation of behavior in distractor tasks, which extends beyond their use to characterize how dysfunction changes distractibility. That is, distractors can serve as a sensitive probe of the underlying neural WM representation, enabling one to measure behaviorally how neural dysfunction perturbs WM representation.

Moreover, the simple tasks that we modeled (i.e. single-item spatial WM with continuous report and parametric control of target–distractor similarity) explicitly build upon electrophysiological studies in awake behaving monkeys (Funahashi et al. 1989). These tasks can therefore be carried out in nonhuman primates as well as clinical populations and healthy volunteers undergoing pharmacological manipulations. This framework in turn allows for the complete translation from the model to electrophysiology, pharmacology, and ultimately clinical phenomena. Our proof-of-principle demonstration of compensations for disinhibition points toward future computational studies of the neural and behavioral effects of pharmacological treatments.

Relation to Other Modeling Studies

Present findings complement recent studies using attractor-based WM models to link the synaptic changes to behavioral deficits in schizophrenia (Loh et al. 2007; Cano-Colino and Compte 2012). Both of these previous computational studies focused on very strong microcircuit perturbations, to such an extent that they compromise network multistability. In contrast, our study focused on more subtle effects that occur within the multistability regime (Fig. 3). Both prior studies describe a more extreme disinhibition regime in which the baseline state is destabilized, and a WM pattern at a random location (“false memory”) can spontaneously emerge anywhere in the network without a sensory cue. In addition, the discrete-pool model used in Loh et al. (2007) cannot capture features that emerge naturally from a continuous model described here: Broadening of WM representations, smooth random drift of memory, and sensitivity to target–distractor similarity. Detecting all of these effects is dependent on the continuous nature of the present model.

As noted, because these previous studies put their networks in regimes of compromised multistability, they predict the different patterns of degraded behavior and errors from those predicted by the present subtle perturbation—namely, they predict the random formation of erroneous WM patterns (false memory). In the context of a match/nonmatch task, spontaneous bump creation at random locations would increase false alarms to far nontarget probes. Increased miss rate would be expected if the persistent activity pattern was completely lost in the target cells (Loh et al. 2007), or a bump spontaneously appeared in nontarget cells, interfering with target encoding (Cano-Colino and Compte 2012). Neither of these effects, the increased false alarm rate to far nontarget probes or the increased miss rate, were observed in the reported ketamine experiment (Fig. 6C). It is worth noting that the same model (Compte et al. 2000) was used by Cano-Colino and Compte (2012) and in the present work. In that sense, even relatively modest changes of parameters, such as the degree of disinhibition, can give rise to qualitatively different behavioral effects with vastly different clinical implications (namely loss of internal precision vs. false memory formation).

Indeed, in a single-item spatial WM experiment, behavioral results for schizophrenia patients were consistent with degraded precision, manifested as increased scatter of reports around the target location (Badcock et al. 2008). This behavioral pattern and those reported by other studies (Mayer and Park 2012) appear inconsistent either with the complete loss of the memory or with false memories, since both of these regimes would correspond with reports made to random, far locations. However, given schizophrenia population heterogeneity, it is possible that some patients indeed show patterns
consistent with more extreme forms of impairment. Therefore, it will be vital for future studies to examine whether any experimental conditions match the more extreme forms of disinhibition regimes proposed in previous modeling studies (Loh et al. 2007; Cano-Colino and Compte 2012). Together, these differing model predictions underscore the utility of distinguishing different error types. The detailed characterization of behavior reports and error types can be harnessed to constrain which network regimes in models may correspond to experimentally studied disease states.

**Future Directions**

In this work, we focused on single-item spatial WM, since it has a better understood neural circuit basis compared with other forms of WM (Funahashi et al. 1989). There is evidence that WM capacity is impaired in schizophrenia (Gold et al. 2010). Computational models suggest that the maintenance of multiple items within a WM circuit depends on the strength of lateral inhibition (Edin et al. 2009) and on the ability to keep activity patterns separated from each other (Wei et al. 2012). Future work with those models could implement disinhibition and test whether elevated E/I ratio, with broadened representations, reduces WM capacity.

Elevated E/I ratio has been implicated in the neuropathology of other psychiatric disorders, including autism spectrum disorders (Rubenstein and Merzenich 2003; Yizhar et al. 2011) and attention-deficit/hyperactivity disorder (Won et al. 2011), both of which exhibit WM deficits (Martinussen et al. 2005; Williams et al. 2005). Present findings on cortical disinhibition may generally describe a core neuropathology underlying multiple neuropsychiatric disorders. Sensitive psychophysical characterization of WM deficits across clinical populations could clarify similarities and differences in WM behavior and potentially elucidate underlying neural substrates of dysfunctions.

Elevated prefrontal E/I ratio may result from alterations in multiple pathways that are implicated in schizophrenia. These include GABAergic deficits, such as reduced GAD67 production (Lewis et al. 2005); increased background input from other areas such as the thalamus or hippocampus (Lisman et al. 2008); or abnormal levels of neumodulators (Laruelle et al. 2003). We predict that multiple mechanisms that elevate E/I ratio may converge in their neural and behavioral consequences. Similarly, there may be multiple mechanisms to restore E/I balance, through compensations in the glutamatergic, GABAergic, or neuromodulatory systems. Computational models linking synaptic perturbations to behavior provide a powerful platform to explore network sensitivity to compensation by various interventions.

As a future extension, the study of disinhibition in recurrent networks should incorporate the diversity of inhibition observed in the cortex. In particular, 2 important classes of interneurons are those that target the perisomatic region of pyramidal cells, and those that target distal dendrites of pyramidal cells (Markram et al. 2004). This segregation of targets may allow these interneuron classes to serve distinct roles in cortical computation (Lovett-Barron et al. 2012; Royer et al. 2012). Both interneuron cell classes are altered in schizophrenia (Lewis et al. 2005; Morris et al. 2008; Bitanihwre et al. 2009). Interestingly, these interneuron classes and pyramidal cells may be differentially sensitive to NMDAR antagonists (Neymotin et al. 2011) and express different proportions of AMPARs and subunits of NMDARs across developmental stages (Wang and Gao 2009; Xi et al. 2009; Rotaru et al. 2011), potentially contributing to the complex developmental trajectory of schizophrenia. Circuit models that incorporate a division of labor among different classes of interneurons (Wang et al. 2004) may be able to link the disruption of each class to specific behavioral deficits. Since soma- and dendrite-targeting interneurons are differentially engaged in gamma- and theta-band oscillations (Gonzalez-Burgos and Lewis 2008; Wang 2010), more detailed microcircuit models have the potential to link the specific aspects of cognitive dysfunction to the diversity of abnormal cortical oscillations observed in schizophrenia (Ulhaas and Singer 2010).

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