1	Predicting distributed working memory activity			
2	in a large-scale mouse brain:			
3	the importance of the cell type-specific connectome			
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17 Abstract

Recent advances in connectomic and neurophysiological tools make it possible to probe 18 whole-brain mechanisms in the mouse that underlie cognition and behavior. Based on 19 experimental data, we developed a large-scale model of the mouse brain for a cardinal 20 cognitive function called working memory, the brain's ability to internally hold and process 21 information without sensory input. In the model, interregional connectivity is constrained 22 by mesoscopic connectome data. The density of parvalbumin-expressing interneurons in the 23 model varies systematically across the cortex. We found that the long-range cell type-specific 24 targeting and density of cell classes define working memory representations. A core cortical 25 subnetwork and the thalamus produce distributed persistent activity, and the network exhibits 26 numerous attractor states. Novel cell type-specific graph theory measures predicted the 27 activity patterns and core subnetwork. This work highlights the need for cell type-specific 28 connectomics, and provides a theory and tools to interpret large-scale recordings of brain 29 activity during cognition. 30

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31 Introduction

In contrast to our substantial knowledge of local neural computation, such as orientation 32 selectivity in the primary visual cortex or spatial map of grid cells in the medial entorhinal 33 cortex, much less is understood about distributed processes in multiple interacting brain 34 regions underlying cognition and behavior. This has recently begun to change, as advances in 35 new technologies enable neuroscientists to probe neural activity at single-cell resolution and 36 on a large-scale by electrical recording or calcium imaging of behaving animals (Jun et al. 37 2017; Steinmetz et al. 2019; Stringer et al. 2019; Musall et al. 2019; Steinmetz et al. 2021), ushering in a new era of neuroscience investigating distributed neural dynamics and brain 39 functions (Wang 2022). 40

To be specific, consider a core cognitive function called working memory, the ability 41 to temporally maintain information in mind without external stimulation (Baddeley 2012). 42 Working memory has long been studied in neurophysiology using delay-dependent tasks, where 43 stimulus-specific information must be stored in working memory across a short time period 44 between a sensory input and a memory-guided behavioral response (Fuster and Alexander 45 1971; Funahashi et al. 1989; Goldman-Rakic 1995; Wang 2001). Delay-period mnemonic 46 persistent neural activity has been observed in multiple brain regions, suggesting distributed 47 working memory representation (Suzuki and Gottlieb 2013; Leavitt et al. 2017; Christophel 48 et al. 2017; Xu 2017; Dotson et al. 2018). Connectome-based computational models of the 49 macaque cortex found that working memory activity depends on interareal connectivity 50 (Murray et al. 2017; Jaramillo et al. 2019), macroscopic gradients of synaptic excitation 51 (Wang 2020; Mejias and Wang 2022) and dopamine modulation (Froudist-Walsh et al. 2021a). 52 Mnemonic neural activity during a delay period is also distributed in the mouse brain 53 (Liu et al. 2014; Schmitt et al. 2017; Guo et al. 2017; Bolkan et al. 2017; Gilad et al. 2018). 54 The new recording and imaging techniques as well as optogenetic methods for causal analysis 55 (Yizhar et al. 2011), that are widely applicable to behaving mice, hold promise for elucidating 56 the circuit mechanism of distributed brain functions in rodents. Recurrent synaptic excitation 57 represents a neural basis for the maintenance of persistent neural firing (Goldman-Rakic 58 1995; D. J. Amit 1995; Wang 2021). In the monkey cortex, the number of spines (sites 59 of excitatory synapses) per pyramidal cell increases along the cortical hierarchy, consistent 60 with the idea that mnemonic persistent activity in association cortical areas including the 61 prefrontal cortex is sustained by recurrent excitation stronger than in early sensory areas. 62 Such a macroscopic gradient is lacking in the mouse cortex (Gilman et al. 2017), raising the 63 possibility that the brain mechanism for distributed working memory representations may be 64 fundamentally different in mice and monkeys. 65

In this paper we report a cortical mechanism of distributed working memory that does not depend on a gradient of synaptic excitation. We developed an anatomically-based model of the mouse brain for working memory, built on the recently available mesoscopic connectivity data of the mouse thalamocortical system (Oh et al. 2014; Gămănuţ et al. 2018; Harris

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et al. 2019; Kim et al. 2017). Our model is validated by capturing large-scale neural activity
observed in recent mouse experiments (Guo et al. 2017; Gilad et al. 2018). Using this model,
we found that a decreasing gradient of synaptic inhibition mediated by parvalbumin (PV)
positive GABAergic cells (Kim et al. 2017; Fulcher et al. 2019; Wang 2020) shapes the
distributed pattern of working memory representation.

A focus of this work is to examine whether anatomical connectivity can predict the 75 emergent large-scale neural activity pattern underlying working memory. Interestingly, 76 traditional graph-theory measures of inter-areal connections, which ignore cell types of 77 projection targets, are uncorrelated with activity patterns. We propose new cell typespecific graph theory measures to overcome this problem, and differentiate contributions of 79 cortical areas in terms of their distinct role in loading, maintaining, and reading out the 80 content of working memory. Through computer-simulated perturbations akin to optogenetic 81 inactivations, a core subnetwork was uncovered for the generation of persistent activity. This 82 core subnetwork can be predicted based on the cell type-specific interareal connectivity, 83 highlighting the necessity of knowing the cell type targets of interareal connections in order 84 to relate anatomy with physiology and behavior. This work provides a computational and 85 theoretical platform for cross-scale understanding of cognitive processes across the mouse 86 cortex.

$\mathbf{Results}$

⁸⁹ A decreasing gradient of PV interneuron density from sensory to ⁹⁰ association cortex

Our large-scale circuit model of the mouse cortex uses inter-areal connectivity provided by 91 anatomical data within the 43-area parcellation in the common coordinate framework v3 atlas (Oh et al. 2014) (Fig. 1A, Fig 1 - supplement 1A). The model is endowed with area-to-area 93 variation of parvalbumin-expressing interneurons (PV) in the form of a gradient measured ٥z from the qBrain mapping platform (Fig 1 - supplement 1B) (Kim et al. 2017). The PV cell 95 density (the number of PV cells per unit volume) is divided by the total neuron density, to 96 give the PV cell fraction, which better reflects the expected amount of synaptic inhibition 97 mediated by PV neurons (Fig. 1B-C, neuron density is shown in Fig 1 - supplement 1C). 98 Cortical areas display a hierarchy defined by mesoscopic connectome data acquired using 99 anterograde fluorescent tracers (Oh et al. 2014) (Fig. 1D-E). In Fig. 1F, the PV density is 100 plotted as a function of the cortical hierarchy, which shows a moderate negative correlation 101 between the two. Therefore, primary sensory areas have a higher density of PV interneurons 102 than association areas, although the gradient of PV densities does not align perfectly with 103 the cortical hierarchy. 104



Figure 1: Anatomical basis of the multi-regional mouse cortical model. (A). Flattened view of mouse cortical areas. Figure adapted from (Harris et al. 2019). (B). Normalized PV cell fraction for each brain area, visualized on a 3d surface of the mouse brain. Five areas are highlighted : VISp, Primary somatosensory area, barrel field (SSp-bfd), primary motor (MOp), MOs and PL. (C). The PV cell fraction for each cortical area, ordered. Each area belongs to one of five modules, shown in color. (Harris et al. 2019). (D). Hierarchical position for each area on a 3d brain surface. Five areas are highlighted as in (B), and color represents the hierarchy position. (E). Hierarchical positions for each cortical area. The hierarchical position is normalized and the hierarchical position of VISp is set to be 0. Same as (C), colors represent which module an area belongs to. (F). Correlation between PV cell fraction and hierarchy (Pearson correlation coefficient r = -0.35, p < 0.05).

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¹⁰⁵ A whole-mouse cortex model with a gradient of interneurons

In our model, each cortical area is described by a local circuit (Fig. 2A), using a mean-field 106 reduction (Wong and Wang 2006) of a spiking neural network (Wang 2002). A minimal 107 version of it has two neural pools selective for different stimuli and a shared inhibitory 108 neural pool. The model makes the following assumptions. First, local inhibitory strength is 109 proportional to PV interneuron density across the cortex. Second, the inter-areal long-range 110 connection matrix is given by the anterograde tracing data (Oh et al. 2014; Knox et al. 2018; 111 Wang et al. 2020). Third, targeting is biased onto inhibitory cells for top-down compared 112 with bottom-up projections, therefore feedforward connections have a greater net excitatory 113 effect than feedback connections (counterstream inhibitory bias, CIB) (Mejias and Wang 114 2022; Javadzadeh and Hofer 2022; Wang 2022). 115

¹¹⁶ Distributed working memory activity depends on the gradient of¹¹⁷ inhibitory neurons and the cortical hierarchy

We simulated the large-scale network to perform a simple visual delayed response task that requires one of two stimuli to be held in working memory. We shall first consider the case in which the strength of local recurrent excitation is insufficient to generate persistent activity when parcellated areas are disconnected from each other. Consequently, the observed distributed mnemonic representation must depend on long-range interareal excitatory connection loops. Later in the paper we will discuss the network model behavior when some local areas are capable of sustained persistent firing in isolation.

The main question is: when distributed persistent activity emerges after a transient visual 125 input (a 500 ms current pulse to a selective excitatory population) is presented to the primary 126 visual cortex (VISp), what determines the spatial pattern of working memory representation? 127 After we remove the external stimulus, the firing rate in area VISp decreases rapidly to 128 baseline. Neural activity propagates throughout the cortex after stimulus offset (Fig. 2B). 129 Neural activities in the higher visual cortical areas (e.g. VISrl and VISpl) show similar 130 dynamics to VISp. In stark contrast, many frontal and lateral areas (including prelimbic 131 (PL), infralimbic (ILA), secondary motor (MOs) and ventral agranular insula (AIv) areas) 132 sustained a high firing rate during the delay period (Fig. 2B). Areas that are higher in the 133 cortical hierarchy show elevated activity during the delay period (Fig. 2C). This persistent 134 firing rate could last for more than 10 seconds and is a stable attractor state of the network 135 (Inagaki et al. 2019). 136

The cortical hierarchy and PV fraction predict the delay period firing rate of each cortical area (Fig. 2C-E). Thus the activity pattern of distributed working memory depends on both local and large-scale anatomy. The delay activity pattern has a stronger correlation with hierarchy (r = 0.91) than with the PV fraction (r = -0.43). The long-range connections thus play a predominant important role in defining the persistent activity pattern.

Activity in early sensory areas such as VISp displays a rigorous response to the transient



Figure 2: Distributed working memory activity depends on the gradient of PV density and the cortical hierarchy. (A). Model design of the large-scale model for distributed working memory. Top, connectivity map of the cortical network. Each node corresponds to a cortical area and an edge is a connection, where the thickness of the edge represents the strength of the connection. Only strong connections are shown (without directionality for the sake of clarity). Bottom, local and long-range circuit design. Each local circuit contains two excitatory populations (red), each selective to a particular stimulus and one inhibitory population (blue). Long-range connections are scaled by mesoscopic connectivity strength (Oh et al. 2014) and follows counterstream inhibitory bias (CIB) (Mejias and Wang 2022). (B). The activity of 6 selected areas during a working memory task is shown. A visual input of 500ms is applied to area VISp, which propagates to the rest of the large-scale network. (C). Delay period firing rate for each area on a 3d brain surface. Similar to Fig. 1B, the positions of 5 areas are labeled. (D). Delay-period firing rate is positively correlated with cortical hierarchy (r = 0.91, p < 0.05). (E). Delay-period firing rate is negatively correlated with PV cell fraction (r = -0.43, p < 0.05).

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input but returns to a low firing state after stimulus withdrawal. In contrast, many frontal
areas show strong persistent activity. When the delay period firing rates are plotted versus
hierarchy, we observe a gap in the distribution of persistent activity (Fig. 2D) that marks an
abrupt transition in the cortical space. This leads to the emergence of a subnetwork of areas
capable of working memory representations.

We also used our circuit model to simulate delayed response tasks with different sensory 148 modalities (Fig 2 - supplement 1), by stimulating primary somatosensory area SSp-bfd and 149 primary auditory area AUDp. The pattern of delay period firing rates for these sensory 150 modalities is similar to the results obtained for the visual task: sensory areas show transient 151 activity, while frontal and lateral areas show persistent activity after stimulus withdrawal. 152 Moreover, the cortical hierarchy could predict the delay period firing rate of each cortical 153 area well (r = 0.89, p < 0.05). Our model thus predicts that working memory may share 154 common activation patterns across sensory modalities, which is partially supported by cortical 155 recordings during a memory-guided response task (Inagaki et al. 2018). 156

¹⁵⁷ Thalamocortical interactions maintain distributed persistent activity

To investigate how thalamocortical interactions affect the large-scale network dynamics, 158 we designed a thalamocortical network similar to the cortical network (Fig. 3A). Several 159 studies have shown that thalamic areas are also involved in the maintenance of working 160 memory (Bolkan et al. 2017; Guo et al. 2017; Schmitt et al. 2017). However, the large-scale 161 thalamocortical mechanisms underlying memory maintenance are unknown. We set the 162 strength of connections between the thalamus and cortex using data from the Allen Institute 163 (Oh et al. 2014) (Fig 3 - supplement 1). All thalamocortical connections in the model are 164 mediated by AMPA synapses. There are no recurrent connections in the thalamus within 165 or across thalamic nuclei (Jones 2007). The effect of thalamic reticular nucleus neurons 166 was included indirectly as a constant inhibitory current to all thalamic areas. Similarly to 167 cortical areas, the thalamus is organized along a measured hierarchy (Harris et al. 2019). For 168 example, the dorsal part of the lateral geniculate nucleus (LGd) is lower than the cortical 169 area VISp in the hierarchy, consistent with the fact that LGd sends feedforward inputs to 170 VISp. Thalamocortical projections in the model are slightly more biased toward excitatory 171 neurons in the target area if they are feedforward projections and towards inhibitory neurons 172 if they are feedback. 173

Here, we weakened the strength of cortical interareal connections as compared to the cortex model of Fig. 2. Now, persistent activity can still be generated (Fig. 3B, blue) but is maintained with the help of the thalamocortical loop, as observed experimentally (Guo et al. 2017). Indeed, in simulations where the thalamus was inactivated, the cortical network no longer showed sustained activity (Fig. 3B, red).

In the thalamocortical model, the delay activity pattern of the cortical areas correlates with the hierarchy, again with a gap in the firing rate separating the areas engaged in



Figure 3: Thalamocortical interactions help maintain distributed persistent activity. (A). Model schematic of the thalamocortical network. The structure of the cortical component is the same as our default model in Fig. 2A, but with modified parameters. Each thalamic area includes two excitatory populations (red square) selective to different stimuli. Long range projections between thalamus and cortex also follow the counterstream inhibitory bias rule as in the cortex. Feedforward projections target excitatory neurons with stronger connections and inhibitory neurons with weaker connections; the opposite holds for feedback projections. (B). The activity of 6 sample cortical areas in a working memory task is shown during control (blue) and when thalamic areas are inhibited in the delay period (red). Black dashes represent the external stimulus applied to VISp. Red dashes represent external inhibitory input given to all thalamic areas. (C). Delay period firing rate of cortical areas in the thalamocortical network. The activity pattern has a positive correlation with cortical hierarchy (r = 0.78, p < 0.05). (D). Same as (C) but plotted against PV cell fraction. The activity pattern has a negative correlation with PV cell fraction, but it is not significant (r = -0.26, p = 0.09). (E). Delay firing rate of thalamic areas in thalamocortical network. The firing rate has a positive correlation with thalamic hierarchy (r = 0.94, p < 0.05). (F). Delay period firing rate of cortical areas in thalamocortical network has a positive correlation with delay firing rate of the same areas in a cortex-only model (r = 0.77, p < 0.05). Note that only the areas showing persistent activity in both models are considered for correlation analyses.

persistent activity from those that do not (Fig. 3B, Fig. 3C). Sensory areas show a low 181 delay firing rate, and frontal areas show strong persistent firing. Unlike the cortex, the firing 182 rate of thalamic areas continuously increases along the hierarchy (Fig. 3E). On the other 183 hand, cortical dynamics in the thalamocortical and cortical models show many similarities. 184 Early sensory areas do not show persistent activity in either model. Many frontal and lateral 185 areas show persistent activity and there is an abrupt transition in cortical space in the 186 thalamocortical model, like in the cortex only model. Quantitatively, the delay firing pattern 187 of the cortical areas is correlated with the hierarchy and the PV fraction (Fig. 3C, Fig. 188 3D). Furthermore, the delay period firing rate of cortical areas in the thalamocortical model 189 correlates well with the firing rate of the same areas in the cortical model (Fig. 3F). This 190 comparison suggests that the cortical model captures most of the dynamical properties in 191 the thalamocortical model; therefore in the following analyses, we will mainly focus on the 192 cortex-only model for simplicity. 193

¹⁹⁴ Cell type-specific connectivity measures predict distributed persistent

¹⁹⁵ firing patterns

Structural connectivity constrains large-scale dynamics (Mejias and Wang 2022; Froudist-196 Walsh et al. 2021a; Cabral et al. 2011). However, we found that standard graph theory 197 measures could not predict the pattern of delay period firing across areas. There is no 198 significant correlation between input strength and delay period firing rate (r = 0.25, p = 0.25, 199 Fig. 4A(i), A(ii)) and input strength cannot predict which areas show persistent activity 200 (prediction accuracy = 0.51, Fig. 4A(iii)). We hypothesized that this is because currently 201 available connectomic data used in this model do not specify the type of neurons targeted by 202 the long-range connections. For instance, when two areas are strongly connected with each 203 other, such a loop would contribute to the maintenance of persistent activity if projections are 204 mutually excitatory, but not if one of the two projections predominantly targets inhibitory 205 PV cells. Therefore, cell type-specificity of interareal connections must be taken into account 206 in order to relate the connectome with the whole-brain dynamics and function. To examine 207 this possibility, we introduced a cell type projection coefficient (see Calculation of network 208 structure measures in the Methods), which is smaller with a higher PV cell fraction in 209 the target area (Fig 4 - supplement 1). The cell type projection coefficient also takes cell 210 type targets of long range connections into account, which, in our model, is quantified by 211 counterstream inhibitory bias (CIB). As a result, the modified cell type-specific connectivity 212 measures increase if the target area has a low density of PV interneurons and/or if long-range 213 connections predominantly target excitatory neurons in the target area. 214

We found that cell type-specific graph measures accurately predict delay-period firing rates. The cell type-specific input strength of the early sensory areas is weaker than the raw input strength (Fig. 4B(i)). The firing rate across areas is positively correlated with cell type-specific input strength (Fig. 4B(ii)). Cell type-specific input strength also accurately



Figure 4: Cell type-specific connectivity measures are better at predicting firing rate pattern than nonspecific ones. (A(i)). Delay period firing rate (orange) and input strength for each cortical area. Input strength of each area is the sum of connectivity weights of incoming projections. Areas are plotted as a function of their hierarchical positions. Delay period firing rate and input strength are normalized for better comparison. (A(ii)). Input strength does not show significant correlation with delay period firing rate for areas showing persistent activity in the model (r = 0.25, p = 0.25). (A(iii)). Input strength cannot be used to predict whether an area shows persistent activity or not (prediction accuracy = 0.51). (B(i)). Delay period firing rate (orange) and cell type-specific input strength for each cortical area. Cell type-specific input strength considers how the long-rang connections target different cell types and is the sum of modulated connectivity weights of incoming projections. Same as (A(i)), areas are sorted according to their hierarchy and delay period firing rate and input strength are normalized for better comparison. (B(ii)). Cell type-specific input strength has a strong correlation with delay period firing rate of cortical areas showing persistent activity (r = 0.89, p < 0.05). Inset: Comparison of the correlation coefficient for raw input strength and cell type-specific input strength. (B(iii)). Cell type-specific input strength predicts whether an area shows persistent activity or not (prediction accuracy = 0.95). Inset: comparison of the prediction accuracy for raw input strength and cell type-specific input strength.

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predicts which areas show persistent activity (Fig. 4B(iii)). Similarly, we found that the cell
type-specific eigenvector centrality, but not standard eigenvector centrality (Newman 2018),
was a good predictor of delay period firing rates (Fig. 4 - supplement 2).

²²² A core subnetwork for persistent activity across the cortex

Many areas show persistent activity in our model. However, are all active areas equally important in maintaining persistent activity? When interpreting large-scale brain activity, we must distinguish different types of contribution to working memory. For instance, inactivation of an area like VISp impairs performance of a delay-dependent task because it is essential for a (visual) "input" to access working memory; on the other hand a "readout" area may display persistent activity only as a result of sustained inputs from other areas that form a "core", which are causally important for maintaining a memory representation.

We propose four types of areas related to distributed working memory: input, core, 230 readout, and nonessential (Fig. 5A). External stimuli first reach input areas, which then 231 propagate activity to the core and non-essential areas. Core areas form recurrent loops and 232 support distributed persistent activity across the network. By definition, disrupting any of 233 the core areas would affect persistent activity globally. The readout areas also show persistent 234 activity. Yet, inhibiting readout areas has little effect on persistent activity elsewhere in 235 the network. We can assign the areas to the four classes based on three properties: a) the 236 effect of inhibiting the area during stimulus presentation on delay activity in the rest of the 237 network; b) the effect of inhibiting the area during the delay period on delay activity in the 238 rest of the network; c) the delay activity of the area itself on trials without inhibition. 239

In search of a core working memory subnetwork in the mouse cortex, in model simulations 240 we inactivated each area either during stimulus presentation or during the delay period, akin 241 to optogenetic inactivation in mice experiments. The effect of inactivation was quantified 242 by calculating the decrement in the firing rate compared to control trials for the areas that 243 were not inhibited (Fig. 5B). The VISp showed a strong inhibition effect during the stimulus 244 period, as expected for an Input area. We identified eight areas with a substantial inhibition 245 effect during the delay period (Fig. 5C), which we identify as a core for working memory. 246 Core areas are distributed across the cortex. They include frontal areas PL, ILA, medial part 247 of the orbital area (ORBm), which are known to contribute to working memory (Liu et al. 248 2014; Bolkan et al. 2017). Other associative and sensory areas (AId, VISpm, ectorhinal area 249 (ECT), perihinal area (PERI), gustatory area (GU)) are also in the core. Similarly, we used 250 the above criteria to classify areas as Readout or Non-essential (Fig. 5D). 251

The core subnetwork can be identified by the presence of strong excitatory loops

Inhibition protocols across many areas are computationally costly. We sought a structural indicator that is easy to compute and is predictive of whether an area is engaged in working

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Figure 5: A core subnetwork generates persistent activity across the cortex. (A). We propose four different types of areas. Input areas (red) are responsible for coding and propagating external signals, which are then propagated through synaptic connections. Core areas (blue) form strong recurrent loops and generate persistent activity. Readout areas (green) inherit persistent activity from core areas. Nonessential areas (purple) may receive inputs and send outputs but they do not affect the generation of persistent activity. (B). Delay period firing rate for cortical areas engaged in working memory (Y axis) after inhibiting different cortical areas (X axis). Areas in the X axis and Y axis are both sorted according to hierarchy. Firing rates of areas with small firing rate (<1Hz) are partially shown (only RSPv and RSPd are shown because their hierarchical positions are close to areas showing persistent activity). (C). The average firing rate for areas engaged in persistent activity under each inhibition simulation. The X axis shows which area is inhibited, and the Y axis shows the average delay period activity for all areas showing persistent activity. Note that when calculating the average firing rate, the inactivated area was excluded in order to focus on the inhibition effect of one area on other areas. Average firing rates on the Y axis are normalized using the average firing in control (no inhibition) simulation. (D). Classification of 4 types of areas based on their delay period activity after stimulus- and delay-period inhibition. The inhibition effect, due to either stimulus or delay period inhibition, is the change of average firing rate normalized by the average firing rate in the control condition. Areas with strong inhibition effect during stimulus period are classified as Input areas; areas with strong inhibition effect during delay period and strong delay period firing rate are classified as Core areas; areas with weak inhibition effect during delay period but strong delay period firing rate during control are classified as Readout areas; areas with weak inhibition effect during delay period and weak delay period firing rate during control are classified as Nonessential areas.

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memory function. Such an indicator could also guide the interpretation of large-scale neural recordings in experimental studies. In the dynamical regime where individual cortical areas do not show persistent activity independently, distributed working memory patterns must be a result of long-range recurrent loops across areas. We thus introduced a quantitative measurement of the degree to which each area is involved in long-range recurrent loops (Fig. 6A).

The core subnetwork can be identified by the presence of strong loops between excitatory 262 cells. Here we focus on length-2 loops (Fig. 6A); the strength of a loop is the product of 263 two connection weights for a reciprocally connected pair of areas; and the loop strength 264 measure of an area is the sum of the loop strengths of all length-2 loops that the area is part 265 of. Results were similar for longer loops (Fig. 6B, also see Fig. 6 - supplement 1 for results 266 of longer loops). The raw loop strength had no statistical relationship to the core working 267 memory subnetwork (Fig. 6C(i), Fig. 6C(ii)). We then defined cell type-specific loop strength 268 (see Methods). The cell type-specific loop strength is the raw loop strength multiplied by 269 the cell type projection coefficient. The cell type-specific loop strength, but not the raw 270 loop strength, predicts which area is a core area with high accuracy (Fig. 6D(i), Fig. 6D(ii), 271 prediction accuracy = 0.93). This demonstrates that traditional connectivity measures are 272 informative but not sufficient to explain dynamics during cognition in the mouse brain. Cell 273 type-specific connectivity, and new metrics that account for such connectivity, are necessary 274 to infer the role of brain areas in supporting large-scale brain dynamics during cognition. 275

²⁷⁶ Multiple attractor states emerge from the mouse mesoscopic connec-²⁷⁷ tome and local recurrent interactions

Different tasks lead to dissociable patterns of internally sustained activity across the brain, 278 described as separate attractor states. We developed a protocol to identify other attractor 279 states, then analyzed the relationship between network properties and the attractor states 280 (Fig. 7A-C). For different parameters, the number of attractors and the attractor patterns 281 change. Two parameters are especially relevant here. These are the long-range connection 282 strength (μ_{EE}) and local excitatory connection strength $(g_{E,self})$. These parameters affect the 283 number of attractors in a model of the macaque cortex (Mejias and Wang 2022). Increasing 284 the long range connection strength decreases the number of attractors (Fig. 7D). Stronger 285 long-range connections implies that the coupling between areas is stronger. If areas are 286 coupled with each other, the activity state of an area will be highly correlated to that of its 287 neighbors. This leads to less variability and fewer attractors. 288

To quantify how the patterns of attractors change for different parameters, two quantities are introduced. The *attractor fraction* is the fraction of all detected attractor states to which an area belongs. An area "belongs" to an attractor state if it is in a high activity state in that attractor. The *attractor size* is defined by the number of areas belonging to that attractor. As we increased the long-range connection strength, the attractor size distribution became



Figure 6: The core subnetwork can be identified structurally by the presence of strong excitatory loops. (A). Distribution of length-2 loops. X axis is the single loop strength of each loop (product of connectivity strengths within loop) and Y axis is their relative frequency. (B). Loop strengths of each area calculated using different length of loops (e.g., length 3 vs length 2) are highly correlated (r = 0.96, p < 0.05). (C(i)). Loop strength (blue) is plotted alongside Core Areas (orange), a binary variable that takes the value 1 if the area is a Core Area, 0 otherwise. Areas are sorted according to their hierarchy. (C(ii)). A high loop strength value does not imply that an area is a core area. Blue curve shows the logistic regression curve fits to differentiate the core areas versus non core areas. (D(i)). Same as (C), but for cell type-specific loop strength. (D(ii)). A high cell type-specific loop strength predicts that an area is a core area (prediction accuracy = 0.93). Same as (C), but for cell type-specific loop strength.

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²⁹⁴ bimodal. The first mode corresponded to large attractors, with many areas. The second ²⁹⁵ mode corresponded to small attractors, with few areas (Fig. 7D).

When the local excitatory strength is increased, the number of attractors increased as 296 well (Fig. 7E). In this regime some areas are endowed with sufficient local reverberation to 297 sustain persistent activity even when decoupled from the rest of the system, therefore the 298 importance of long-range coupling is diminished and a greater variety of attractor states is 299 enabled. This can be understood by a simple example of two areas 1 and 2, each capable of 300 two stimulus-selective persistent activity states; even without coupling there are $2 \times 2 = 4$ 301 attractor states with elevated firing. Thus, local and long-range connection strength have 302 opposite effects on the number of attractors. 303

The cell type-specific input strength predicted firing rates across many attractors. In 304 an example parameter regime ($\mu_{EE} = 0.04$ nA and $g_{E,self} = 0.44$ nA), we identified 143 305 attractors. We correlated the input strength and cell type-specific input strength with the 306 many attractor firing rates (Fig. 7F). The raw input strength is weakly correlated with 307 activity patterns. The cell type-specific input strength is strongly correlated with activity 308 across attractors. This shows that the cell type-specific connectivity measures are better at 309 predicting the firing rates in many scenarios. These results further prove the importance of 310 having cell type-specific connectivity for modeling brain dynamics. 311

Different attractor states rely on distinct subsets of core areas. In one example attractor, 312 we found 5 areas that show persistent activity: VISa, VISam, FRP, MOs and ACAd (Fig. 313 7G). We repeated the previous inhibition analysis to identify core areas for this attractor 314 state. Inhibiting one area, MOs, during the delay had the strongest effect on delay activity 315 in the other parts of the attractor (Fig. 7H). MOs also showed strong persistent activity 316 during delay period. This is consistent with its role in short-term memory and planning (Li 317 et al. 2015; Inagaki et al. 2019). According to our definition, MOs is a core area for this 318 attractor. To calculate a loop strength that was specific to this attractor, we only examined 319 connections between these five areas. The cell type-specific loop strength was strongest in 320 area MOs (Fig. 7I). Thus, we can identify likely core areas for individual attractor states 321 from cell type-specific structural measures. This also demonstrates that different attractor 322 states can be supported by distinct core areas. 323

324 Discussion

We developed a connectome-based dynamical model of the mouse brain. The model was capable of internally maintaining sensory information across many brain areas in distributed activity in the absence of any input. To our knowledge this is the first biologically-based model of the entire mouse cortex and the thalamocortical system for a cognitive function. Together with our recent work (Mejias and Wang 2022; Froudist-Walsh et al. 2021a; Froudist-Walsh et al. 2021b), it provides a study of contrast between mice and monkeys.

Our main findings are threefold. First, mnemonic activity pattern is shaped by the differing





Figure 7: Multiple attractors coexist in the mouse working memory network. (A-C) Example attractor patterns with a fixed parameter set. Each attractor pattern can be reached via different external input patterns applied to the brain network. Delay activity is shown on a 3D brain surface. Color represents the firing rate of each area. (D-E) The distribution of attractor fractions (left) and number of attractors as a function of size (right) for different parameter combinations are shown. Attractor fraction of an area is the ratio between the number of attractors that include the area and the total number of identified attractors. In (D), local excitatory strengths are fixed ($g_{E,self} = 0.44$ nA) while long-range connection strengths vary in the range $\mu_{EE} = 0.01-0.05$ nA. Left and right panels of (D) show one specific parameter $\mu_{EE} = 0.03$ nA. Inset panel of (D) shows the number of attractors under different long-range connection strengths while $g_{E,self}$ is fixed at 0.44 nA. In (E), long range connection strengths are fixed ($\mu_{EE} = 0.02 \text{ nA}$) while local excitatory strengths varies in the range $g_{E,self} = 0.4-0.44$ nA. Left and right panels of (E) show one specific parameter $g_{E,self} = 0.43$ nA. Inset panel of (E) shows the number of attractors under different local excitatory strengths, while μ_{EE} is fixed at 0.02 nA. (F). Prediction of the delay period firing rate using input strength and cell type-specific input strength for each attractor state identified under $\mu_{EE} = 0.04$ nA and $g_{E,self}$ = 0.44 nA. 143 distinct attractors were identified and the average correlation coefficient using cell type-specific input strength is better than that using input strength. (G). A example attractor state identified under the parameter regime $\mu_{EE} = 0.03$ nA and $g_{E,self} = 0.44$ nA. The 5 areas with persistent activity are shown in red. (H). Effect of single area inhibition analysis for the attractor state in (G). For a regime where 5 areas exhibit persistent activity during the delay period, inactivation of the premotor area MOs yields a strong inhibition effect (<0.95 orange dashed line) and is therefore a Core area for the attractor state in (G). (I). Cell type-specific loop strength (blue) is plotted alongside Core areas (orange). Only 5 areas with persistent activity are used to calculate the loop strength. Loop strength is normalized to be within the range of 0 and 1. High cell type-specific loop measures predict that an area is a Core area (prediction accuracy is 100% correct). The number of areas is limited, so prediction accuracy is very high.

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densities of PV interneurons across cortical areas. Areas with high PV density encoded 332 information only transiently. Those with low PV density sustained activity for longer periods. 333 Thus, the gradient of PV cells (Kim et al. 2017) has a definitive role in separating rapid 334 information processing in sensory areas from sustained mnemonic information representation 335 in associative areas of the mouse cortex. This is consistent with the view that each local area 336 operates in the "inhibition-stabilizing regime" where recurrent excitation alone would lead 337 to instability but the local network is stabilized by feedback inhibition even in the primary 338 visual cortex (R. J. Douglas et al. 1995; Murphy and Miller 2009). Second, we deliberately 339 considered two different dynamical regimes: when local recurrent excitation is not sufficient to sustain persistent activity and when it does. In the former case, distributed working 341 memory must emerge from long-range interactions between parcellated areas, thereby the 342 concept of synaptic reverberation (Lorente de Nó 1933; P. S. Goldman-Rakic 1995; Wang 343 2001; Wang 2021) is extended to the large-scale global brain. Note that currently it is unclear 344 whether persistent neural firing observed in a delay dependent task is generated locally or 345 depends on long-distance reverberation among multiple brain regions. Our work made the 346 distinction explicit and offers specific predictions to be tested experimentally. Third, presently 347 available connectomic data are not sufficient to account for neural dynamics and distributed 348 cognition, and we propose cell type-specific connectomic measures that are shown to predict 349 the observed distributed working memory representations. 350

We found that the cortical structures form recurrent loops with the thalamus, and the 351 thalamocortical loops aided in sustaining activity throughout the delay period. The specific 352 pattern of cortico-cortical connections was also critical to working memory. However, standard 353 graph theory measures based on the connectome were unable to predict the pattern of working 354 memory activity. By focusing on cell type-specific interactions between areas, we were able to 355 reveal a core of cortical areas. The core is connected by excitatory loops, and is responsible for 356 generating a widely distributed pattern of sustained activity. Outside the core, we identified 357 "readout" areas that inherited activity from the core. Readout areas could use this information 358 for further computations. This clarifies the synergistic roles of the connectome and gradients 359 of local circuit properties in producing a distributed cognitive function. This additionally highlights the need for a cell type-specific connectome. 361

Previous large-scale models of the human and macaque cortex have replicated the func-362 tional connectivity (Deco et al. 2014; Demirtaş et al. 2019; Honey et al. 2007; Schmidt et al. 363 2018; Shine et al. 2018; Cabral et al. 2011; Wang et al. 2019) and propagation of information 364 along the cortical hierarchy (Chaudhuri et al. 2015; Joglekar et al. 2018; Diesmann et al. 1999). 365 Recently, large-scale models of brain activity during cognitive tasks have been developed 366 (Mejias and Wang 2022; Froudist-Walsh et al. 2021a; Klatzmann et al. 2022). In these models, 367 the number of dendritic spines per pyramidal cell increases along the hierarchy. This enables 368 associative regions of the cortex to maintain information in working memory. The increase 369 of dendritic spines along the hierarchy is a robust feature of primate cortical organisation 370 (Elston 2007), which does not exist in the mouse cortex (Gilman et al. 2017). Yet, in the 371

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mouse cortex, other properties do vary along the cortical hierarchy (Kim et al. 2017; Fulcher 372 et al. 2019). We took advantage of the recent discovery of a gradient of PV interneurons in 373 the mouse cortex (Kim et al. 2017), and implemented it directly in our large-scale model. We 374 demonstrated how the increasing gradient of excitation along the primate cortical hierarchy 375 and the decreasing gradient of PV inhibition in the mouse cortex could serve a similar role. 376 Both gradients enable sustained activity to emerge in associative areas. Thus, while the neural 377 activity underlying working memory may be widely distributed in both rodents and primates, 378 the circuit-level mechanisms may differ. This should be considered when interpreting studies 379 of working memory in rodent models of cognition and disease.

In the macaque, long-range connectivity is a strong predictor of the working memory 381 activity (Mejias and Wang 2022; Froudist-Walsh et al. 2021a). Thus, at least some of the 382 functional specialization of brain areas is due to differences in interareal connections. In 383 contrast, we found that traditional graph theory metrics of connectivity were unable to 384 predict the working memory activity in the mouse brain. This may be due to the almost 385 fully connected pattern of interareal connectivity in the mouse cortex (Gămănut et al. 2018). 386 This implies that, qualitatively, all areas have a similar set of cortical connections. In our 387 model, we allowed the cell type target of interareal connections to change according to the 388 relative position of the areas along the cortical hierarchy. Feedforward connections had a 389 greater net excitatory effect than feedback connections. This hypothesis (Mejias and Wang 390 2022) has received some recent experimental support (Yoo et al. 2021; Huang et al. 2019; 391 Javadzadeh and Hofer 2022). 392

By introducing cell type-specific graph theory metrics, we were able to predict the pattern and strength of delay period activity with high accuracy. Connectome databases are an invaluable resource for basic neuroscience. However, they may be insufficient for constraining models of brain activity. In the future, connectome databases should be supplemented by cell type-specific information.

We demonstrated how cell type-specific graph-theory measures can accurately identify 398 the core subnetwork, which can also be identified independently using a simulated large-399 scale optogenetic experiment. We found a core subnetwork of areas that, when inhibited, 400 caused a substantial drop in activity in the remaining cortical areas. This core working 401 memory subnetwork included frontal cortical areas with well documented patterns of sustained 402 activity during working memory tasks, such as prelimbic (PL), infralimbic (ILA) and medial 403 orbitofrontal cortex (ORBm) (Schmitt et al. 2017; Liu et al. 2014; Wu et al. 2020). However, 404 the core subnetwork for the visual working memory task we assessed was distributed across 405 the cortex. It also included temporal and higher visual areas, suggesting that long-range 406 recurrent connections between the frontal cortex and temporal and visual areas are responsible 407 for generating persistent activity and maintaining visual information in working memory in 408 the mouse. 409

The core visual working memory subnetwork generates activity that is then inherited by many readout areas. Readout areas also exhibit persistent activity. However, inhibiting

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readout areas does not significantly affect the activity of other areas (Figure 5). Readout 412 areas can use the stored information for further computations or to affect behavior. The 413 readout areas in our model were a mixture of higher visual areas, associative areas and 414 premotor areas of cortex. Notably, we classified the secondary motor cortex (MOs), which 415 contains the anterior lateral motor (ALM) area, as a readout area despite its high level 416 of persistent activity. ALM has received a lot of attention in mouse studies of working 417 memory and motor preparation (Guo et al. 2017; Guo et al. 2014; Inagaki et al. 2019; Li 418 et al. 2015; Wu et al. 2020; Voitov and Mrsic-Flogel 2022). If ALM is indeed a readout area 419 for sensory working memory tasks, (e.g., Schmitt et al. 2017), then the following prediction arises. Inhibiting ALM should have a relatively small effect on sustained activity in core 421 areas (such as PL) during the delay period. In contrast, inhibiting PL and other core areas 422 may disrupt sustained activity in ALM. Even if ALM is not part of the core for sensory 423 working memory, it could form part of the core for motor preparation tasks (Figure 7G). We 424 found a high cell type-specific loop strength for area ALM, like that in core areas, which 425 supports this possibility (Figure 7I). Furthermore, we found some attractor states for which 426 the MOs was classified as a core area, while PL was not even active during the delay period. 427 Our result is supported by a recent study that found no behavioral effect after PL inhibition 428 in a motor planning task (Wang et al. 2021). Therefore, the core subnetwork required for 429 generating persistent activity is likely task-dependent. Outside of this core subnetwork, there 430 is a large array of readout brain areas that can use the stored information to serve behavior. 431 Future modeling work may help elucidate the biological mechanisms responsible for switching 432 between attractor landscapes for different tasks. 433

Neuroscientists are now observing task-related neural activity at single-cell resolution 434 across much of the brain (Stringer et al. 2019; Steinmetz et al. 2019). This makes it important 435 to identify ways to distinguish the core areas for a function from those that display activity 436 that serves other purposes. We show that a large-scale inhibition protocol can identify the 437 core subnetwork for a particular task. We further show how this core can be predicted based 438 on the interareal loops that target excitatory neurons. Were such a cell type-specific interareal 439 connectivity dataset available, it may help interpretation of large-scale recording experiments. 440 This could also focus circuit manipulation on regions most likely to cause an effect on the 441 larger network activity and behavior. Our approach identifies the brain areas that are working 442 together to support working memory. It also identifies those that are benefiting from such 443 activity to serve other purposes. Our simulation and theoretical approach is therefore ideally 444 suited to understand the large-scale anatomy, recording and manipulation experiments which 445 are at the forefront of modern systems neuroscience. 446

Neuroscience has rapidly moved into a new era of investigating large-scale brain circuits.
Technological advances have enabled the measurement of connections, cell types and neural activity across the mouse brain. We developed a model of the mouse brain and theory of working memory that is suitable for the large-scale era. Previous reports have emphasized the importance of gradients of dendritic spine expression and interareal connections in sculpting

task activity in the primate brain (Mejias and Wang 2022; Froudist-Walsh et al. 2021a). 452 Although these anatomical properties from the primate cortex are missing in the mouse 453 brain (Gămănuț et al. 2018; Gilman et al. 2017), other properties such as interneuron density 454 (Kim et al. 2017) may contribute to areal specialization. Indeed, our model clarifies how 455 gradients of interneurons and cell type-specific interactions define large-scale activity patterns 456 in the mouse brain during working memory, which enables sensory and associative areas 457 to have complementary contributions. Future large-scale modeling studies can leverage cell 458 type-specific connectivity to study other important cognitive computations beyond working 459 memory, including learning and decision making (Abbott et al. 2017; Abbott et al. 2020).

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467 Inclusion and Diversity

468 N/A.

469 Author contributions

XJW: designed the research, worked with the other authors throughout the project and
co-wrote the paper; XYD: carried out all the computer simulations and analysis of simulation
data, and co-wrote the paper; SFW, JJ and JJJ: contributed to all aspects of this project in
interactions with XYD and co-wrote the paper.

474 Declaration of Interests

⁴⁷⁵ No competing interests declared.

$_{476}$ Methods

477 Anterograde tracing, connectivity data

We used the mouse connectivity map from Allen institute (Oh et al. 2014) to constrain our large-scale circuit model of the mouse brain. The Allen Institute measured the connectivity among cortical and subcortical areas using an anterograde tracing method. In short, they injected virus and expressed fluorescent protein in source areas and performed fluorescent

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imaging in target areas to measure the strength of projections from source areas. Unlike 482 retrograde tracing methods used in other studies (Markov et al. 2014b), the connectivity 483 strength measured using this method does not need to be normalized by the total input or 484 output strength. This means that connectivity strength between any two areas is comparable. 485 The entries of the connectivity matrix from the Allen Institute can be interpreted as 486 proportional to the total number of axonal fibers projecting from unit volume in one area 487 to unit volume in another area. Before incorporating the connectivity into our model, we 488 normalized the data as follows. In each area, we model the dynamics of an "average" neuron, 489 assuming that the neuron receive inputs from all connected areas. Thus, we multiplied the connectivity matrix by the volume Vol_j of source area j and divided by the average neuron 491 density d_i in target area i: 492

$$W_{norm,ij} = W_{raw,ij} \frac{Vol_j}{d_i},\tag{1}$$

where $W_{raw,ij}$ is the raw, i.e., original, connection strength from unit volume in source area jto unit volume in target area i, Vol_j is the volume of source area j (Wang et al. 2020), and d_i is the neuron density in source area i (Erö et al. 2018). $W_{norm,ij}$ is the matrix that we use to set the long rang connectivity in our circuit model. We can define the connectivity between thalamus and cortex, $W_{ct,norm,ij}$ and $W_{tc,norm,ij}$ in a similar manner.

⁴⁹⁸ Interneuron density along the cortex

Kim and colleagues measured the density of typical interneuron types in the brain (Kim et al. 2017). They expressed fluorescent proteins in genetically labeled interneurons and counted the number of interneurons using fluorescent imaging. We took advantage of these interneuron density data and specifically used the PV density to set local and long-range inhibitory weights. We first normalized the PV density in each area:

$$PV_i = \frac{PV_{raw,i} - min(PV_{raw,i})}{max(PV_{raw,i}) - min(PV_{raw,i})}$$
(2)

⁵⁰⁴ $PV_{raw,i}$ is the PV interneuron density of all layers in area *i*, and PV_i is a normalized value of ⁵⁰⁵ PV_{raw} , which will be used in subsequent modeling.

⁵⁰⁶ Hierarchy in the cortex

The concept of hierarchy is important for understanding the cortex. Hierarchy can be 507 defined based on mapping corticocortical long range connections onto feedforward or feedback 508 connections (Felleman and Essen 1991; Markov et al. 2014a; Harris et al. 2019). Harris and 509 colleagues measured the corticocortical projections and target areas in a series of systematic 510 experiments in mice (Harris et al. 2019). Projection patterns were clustered into multiple 511 groups and the label "feedforward" or "feedback" was assigned to each group. Feedforward 512 and feedback projections were then used to determine relative hierarchy between areas. For 513 example, if the projections from area A to area B are mostly feedforward, then area B has 514

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⁵¹⁵ a higher hierarchy than area A. This optimization process leads to a quantification of the ⁵¹⁶ relative hierarchy of cortical areas $h_{raw,i}$. We defined the normalized hierarchy value h_i as

$$h_i = \frac{h_{raw,i} - \min(h_{raw,i})}{\max(h_{raw,i}) - \min(h_{raw,i})}$$
(3)

where $h_{raw,i}$ is the raw, i.e., original hierarchical ordering from (Harris et al. 2019). Due to data acquisition issues, 6 areas did not have a hierarchy value assigned to them (SSp-un, AUDv, GU, VISC, ECT, PERI) (Harris et al. 2019). We estimated hierarchy through a weighted sum of the hierarchy value of 37 known areas, while the weight is determined through the connectivity strength. The parameters α_h and β_h are selected so that $h_{i,estimate}$ are close to h_i for areas with known hierarchy.

$$h_{i,estimate} = \alpha_h \frac{\sum_{j=1}^{37} w_{raw,ij} h_j}{\sum_{j=1}^{37} w_{raw,ij}} + \beta_h.$$
(4)

For the thalamocortical model, we also used the hierarchy value for thalamic areas (Harris et al. 2019). The hierarchy of thalamic areas are comparable to cortical areas, so in order to use it in the model, we also normalized them.

$$h_{th,i} = \frac{h_{th,raw,i} - min(h_{raw,i})}{max(h_{raw,i}) - min(h_{raw,i})}$$
(5)

To estimate the hierarchy value of thalamic areas with missing values, we used the known hierarchy value of the thalamic area next to the missing one as a replacement.

528 Description of the local circuit

Our large-scale circuit model includes 43 cortical areas. Each area includes two excitatory populations, labeled A and B, and one inhibitory population, C. The two excitatory populations are selective to different stimuli. The synaptic dynamics between populations are based on previous firing rate models of working memory (Wang 1999; Wong and Wang 2006). The equations that define the dynamics of the synaptic variables are

$$\frac{dS_A}{dt} = -\frac{S_A}{\tau_N} + \gamma (1 - S_A) r_A \tag{6}$$

$$\frac{dS_B}{dt} = -\frac{S_B}{\tau_N} + \gamma (1 - S_B) r_B \tag{7}$$

$$\frac{dS_C}{dt} = -\frac{S_C}{\tau_G} + \gamma_I r_C \tag{8}$$

where S_A and S_B are the NMDA synaptic variables of excitatory populations A and B, while S_C is the GABA synaptic variable of the inhibitory population C. r_A , r_B and r_C are the firing rates of populations A, B and C, respectively. τ_N and τ_G are the time constants of NMDA and GABA synaptic conductances. γ and γ_I are the parameters used to scale the contribution of presynaptic firing rates. The total currents received I_i (i = A, B, C) are given by

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$$I_A = g_{E,self} S_A + g_{E,cross} S_B + g_{EI} S_C + I_{0A} + I_{LR,A} + x_A(t)$$
(9)

$$I_B = g_{E,self} S_B + g_{E,cross} S_A + g_{EI} S_C + I_{0B} + I_{LR,B} + x_B(t)$$
(10)

$$I_C = g_{IE}S_A + g_{IE}S_B + g_{II}S_C + I_{0C} + I_{LR,C} + x_C(t).$$
(11)

In these equations, $g_{E,self}$, $g_{E,cross}$ denote the connection strength between excitatory neurons 535 with same or different selectivity, respectively. These connection strengths are the same for 536 different areas, since there is no significant gradient for excitatory strength in mice. g_{IE} 537 are the connection strengths from excitatory to inhibitory neurons, while g_{EI} , and g_{II} are 538 connection strengths from inhibitory to excitatory neurons and from inhibitory to inhibitory 539 neurons, respectively. These connections will be scaled by PV density in the corresponding 540 area. We will discuss the details in the next section. I_{0i} (i = A, B, C) are constant background 541 currents to each population. $I_{LR,i}$ (i = A, B, C) are the long range (LR) currents received 542 by each population. The term $x_i(t)$ where i = A, B, C represents noisy contributions from 543 neurons external to the network. It is modeled as an Ornstein-Uhlenbeck process: 544

$$\tau_{noise} \frac{dx_i}{dt} = -x_i + \sqrt{\tau_{noise}} \sigma_i \zeta_i(t), \qquad (12)$$

where $\zeta_i(t)$ is Gaussian white noise, τ_{noise} describes the time constant of external AMPA synapses and σ_i sets the strength of the noise for each population. $\sigma_A = \sigma_B = 5pA$ while $\sigma_C = 0pA$.

The steady state firing rate of each population is calculated based on a transfer function $\phi_i(I)$ of input current received by each population I_i (i = A, B, C) given by

$$\phi_{A,B}(I_{A,B}) = \frac{aI_{A,B} - b}{1 - exp[-d(aI_{A,B} - b)]}$$
(13)

$$\phi_C(I_C) = \left[\frac{1}{g_I}(c_1 I - c_0) + r_0\right]^+ \tag{14}$$

Note that the transfer functions $\phi_i(t)$ are the same for two excitatory populations. x^+ denotes the positive part of the function x. The firing rate of each population follows equations:

$$\tau_r \frac{dr_{A,B}}{dt} = -r_{A,B} + \phi_{A,B}(I_{A,B})$$
(15)

$$\tau_r \frac{dr_C}{dt} = -r_C + \phi_C(I_C). \tag{16}$$

548 Interneuron gradient and local connections

We scaled local interneuron connectivity with the interneuron density that was obtained using fluorescent labeling (Kim et al. 2017). Specifically, local I-I connections and local I-E connections are scaled by the interneuron density by setting the connection strength $g_{k,i}(k = EI, II)$ as a linear function of PV density PV_i in area *i*.

$$g_{EI,i} = J_{EI,min} + J_{EI,scaling} PV_i \tag{17}$$

$$g_{II,i} = J_{II,min} + J_{II,scaling} PV_i \tag{18}$$

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where $J_{k,min}$ (k = EI, II) is the intercept and $J_{k,scaling}$ (k = EI, II) is the slope.

⁵⁵⁰ Hierarchy and long range connections

Long range (LR) connections between areas are scaled by connectivity data from the Allen Institute (Oh et al. 2014). We consider long-range connections that arise from excitatory neurons because most long-range connections in the cortex correspond to excitatory connections (Petreanu et al. 2009). Long-range connections will target excitatory populations in other brain areas with the same selectivity (Zandvakili and Kohn 2015) and will also target inhibitory neurons. These long-range connections are given by the following equations:

$$I_{A,B,LR,i} = \mu_{EE} W_{E,ij} S_{A,B,j} \tag{19}$$

$$I_{C,LR,i} = \mu_{IE} W_{I,ij} (S_{A,j} + S_{B,j}),$$
(20)

where W_E is the normalized long-range connectivity to excitatory neurons, and W_I is the normalized long-range connectivity to inhibitory neurons. μ_{EE} and μ_{IE} are coefficients scaling the long-range E to E and E to I connection strengths, respectively.

Here, we assume that the long-range connections will be scaled by a coefficient that is based 554 on the hierarchy of source and target area. To quantify the difference between long-range 555 feedforward and feedback projections, we introduce m_{ij} to measure the "feedforwardness" of 556 projections between two areas. According to our assumption of counterstream inhibitory bias 557 (CIB), long-range connections to inhibitory neurons are stronger for feedback connections 558 and weaker for feedforward connections, while the opposite holds for long range connections 559 to excitatory neurons. Following this hypothesis, we define m_{ij} as a sigmoid function of 560 the difference between the hierarchy value of source and target areas. For feedforward 561 projections, $m_{ij} > 0.5$; for feedback projections, $m_{ij} < 0.5$. Excitatory and inhibitory 562 long-range connection strengths are implemented by multiplying the long-range connectivity 563 strength W_{ij} by m_{ij} and $(1 - m_{ij})$, respectively: 564

$$m_{ij} = \frac{1}{1 + \beta e^{-(h_i - h_j)}} \tag{21}$$

$$W_{E,ij} = m_{ij}W_{ij} \tag{22}$$

$$W_{I,ij} = (1 - m_{ij})W_{ij} \tag{23}$$

565 with

$$W_{ij} = (W_{norm,ij})^{k_{scale}} \tag{24}$$

The normalized connectivity $W_{norm,ij}$ is then rescaled to translate the broad range of connectivity values (over five orders of magnitude) to a range more suitable for our firing rate models. k_{scale} is the coefficient used for this scaling. $k_{scale} < 1$ effectively makes the range much smaller than the original normalized connectivity $W_{norm,ij}$.

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570 Thalamocortical network model

⁵⁷¹ Corticothalamic connectivity. We introduced thalamic areas in the network to examine their
⁵⁷² effect on cortical dynamics. Each thalamic area includes 2 excitatory populations, A and B,
⁵⁷³ with no inhibitory population. These two populations share the same selectivity with the
⁵⁷⁴ corresponding cortical areas. Unlike cortical areas, there are no recurrent connections between
⁵⁷⁵ thalamic neurons (Sherman 2007). Thalamic currents have the following contributions (tc
⁵⁷⁶ stands for thalamocortical connections and ct for corticothalamic connections):

$$I_{th,A,B} = I_{ct,A,B} + I_{th,0,A,B} + I_{th,noise,A,B}$$

$$\tag{25}$$

where $I_{th,i}$ (i = A, B) is the total current received by each thalamic population, $I_{ct,i}$ (i = A, B)is the long range current from cortical areas to target thalamic area, $I_{th,0,i}$ (i = A, B) is the background current for each population, and $I_{th,noise,i}$ (i = A, B) is the noise input to thalamic population A and B, which we set to 0 in our simulations. $I_{ct,i}$ (i = A, B) has the following form:

$$I_{ct,A,B,i} = g_{ct} W_{ct,E,ij} S_{k,j} \tag{26}$$

where $W_{ct,E,ij}$ is the LR connectivity to thalamic neurons, and $S_{k,j}$ is the synaptic variable of population k (k = A, B) in cortical area j. Since all thalamic neurons are excitatory, we model corticothalamic projections as in the previous section:

$$m_{ct,ij} = \frac{1}{1 + \beta e^{-(h_{th,i} - h_j)}}$$
(27)

$$W_{ct,E,ij} = m_{ct,ij} W_{ct,ij} \tag{28}$$

(29)

585 where

$$W_{ct,ij} = (W_{ct,norm,ij})^{k_{scale}}$$
(30)

 $W_{ct,norm,ij}$ is the normalized connection strength from cortical area j to thalamic area i. $m_{ct,ij}$ is the coefficient quantifying how the long range connections target excitatory neurons based on cortical hierarchy h_j and thalamic hierarchy $h_{th,i}$.

589 The thalamic firing rates are described by:

$$\tau_r \frac{dr_{th,A,B}}{dt} = -r_{th,A,B} + \phi_{th,A,B}(I_{th,A,B})$$
(31)

⁵⁹⁰ with the activation function for thalamic neurons given by:

$$\phi_{th,A,B}(I_{th,A,B}) = \frac{aI_{th,A,B} - b}{1 - exp[-d(aI_{th,A,B} - b)]}$$
(32)

⁵⁹¹ Thalamic neurons are described by AMPA synaptic variables (Jaramillo et al. 2019):

$$\frac{dS_{th,A,B}}{dt} = -\frac{S_{th,A,B}}{\tau_A} + \gamma_A r_{th,A,B}$$
(33)

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Thalamocortical connectivity. The connections from thalamic neurons to cortical neurons follow these equations

$$I_{tc,A,B,i} = g_{E,tc} W_{E,tc,ij} S_{th,A,B,j} \tag{34}$$

$$I_{tc,C,i} = g_{I,tc} W_{I,tc,ij} (S_{th,A,j} + S_{th,B,j})$$
(35)

and connectivity

$$m_{tc,ij} = \frac{1}{1 + \beta e^{-(h_i - h_{th,j})}}$$
(36)

$$W_{E,tc,ij} = m_{tc,ij} W_{tc,ij} \tag{37}$$

$$W_{I,tc,ij} = (1 - m_{tc,ij})W_{tc,ij}$$
 (38)

⁵⁹² and connectivity matrix

$$W_{tc,ij} = (W_{tc,norm,ij})^{k_{scale}}$$
(39)

⁵⁹³ The thalamocortical input is added to the total input current of each cortical population.

$$I_A = g_{E,self} S_A + g_{E,cross} S_B + g_{EI} S_C + I_{0A} + I_{LR,A} + I_{tc,A} + x_A(t)$$
(40)

$$I_B = g_{E,self}S_B + g_{E,cross}S_A + g_{EI}S_C + I_{0B} + I_{LR,B} + I_{tc,B} + x_B(t)$$
(41)

$$I_C = g_{IE}S_A + g_{IE}S_B + g_{II}S_C + I_{0C} + I_{LR,C} + I_{tc,C} + x_C(t)$$
(42)

⁵⁹⁴ Calculation of network structural measures

⁵⁹⁵ We considered three types of structural measures. The first one is input strength. Input ⁵⁹⁶ strength of area i is the summation of the connection strengths onto node i. It quantifies the ⁵⁹⁷ total external input onto area i.

$$W_{input,i} = \sum_{j=1}^{n} W_{ij} \tag{43}$$

The second one is eigenvector centrality (Newman 2018). Eigenvector centrality of area i is the ith element of the leading eigenvector of the connectivity matrix. It quantifies how many areas are connected with the target area i and how important these neighbors are.

$$W = Q\Lambda Q^{-1} \tag{44}$$

$$C_{eig,i} = q_{i1} \tag{45}$$

The third structural measure is loop strength, which quantifies how each area is involved in strong recurrent loops. We first define the strength of a single loop k

$$L_k = \prod_{A_i, A_j \in loop_k} w_{ij},\tag{46}$$

and then the loop strength S_{A_i} of a single area A_i

$$S_{A_i} = \sum_{A_i \in loop_k} L_k \tag{47}$$

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We now focus on cell type-specific structural measures. Cell type specificity is introduced via a coefficient k_{cell} that scales all long range connection strengths (cell type projection coefficient):

$$k_{cell} = m_{ij} - PV_i(1 - m_{ij})$$
(48)

⁶⁰⁴ Thus, we can define cell type-specific input strength as:

$$W_{input,i,cellspec} = \sum_{j=1}^{n} (m_{ij} - PV_i(1 - m_{ij}))W_{ij}$$
(49)

Similarly, cell type-specific eigenvector centrality is defined as

$$\tilde{W}_{ij} = (m_{ij} - PV_i(1 - m_{ij}))w_{ij}$$
(50)

$$\tilde{W} = \tilde{Q}\tilde{\Lambda}\tilde{Q}^{-1} \tag{51}$$

$$C_{eig,i,cellspec} = \tilde{q}_{i1} \tag{52}$$

and cell type-specific loop strength:

$$L_{k,cellspec} = \prod_{A_i, A_j \in loop_k} (m_{ij} - PV_i(1 - m_{ij}))w_{ij}$$
(53)

$$S_{A_i,cellspec} = \sum_{A_i \in loop_k} L_{k,cellspec}$$
(54)

⁶⁰⁵ Stimulation protocol and inhibition analysis

We simulate a working memory task by applying an external current I_{stim} to one of the excitatory populations. The external current is a pulsed input with start time T_{on} and offset time T_{off} . Without losing generality, we assume that the external input is provided to population A. In most of the simulations in this study, we simulate a visual working memory task, with the external applied to VISp. The simulation duration is T_{trial} and we used a time step of dt.

We apply inhibition analysis to understand the robustness of attractors and, more 612 importantly, to investigate which areas play an important role in maintaining the attractor 613 state. Excitatory input was applied to the inhibitory population I to simulate opto-genetic 614 inhibition. The external input I_{inh} is strong as compared to I_{stim} and results in an elevated 615 firing rate of the inhibitory population, which in turn decreases the firing rate of the excitatory 616 populations. Usually the inhibition is applied to a single area. When inhibition is applied 617 during the stimulus period, its start and end times are equal to T_{on} and T_{off} , respectively. 618 When inhibition is applied during delay period, its start time is later than T_{off} to allow the 619 system settle to a stable state. Thus, the onset of inhibition starts 2 seconds after T_{off} and 620 lasts until the end of trial. In the case of thalamocortical network simulations, we inhibit 621 thalamic areas by introducing a hyperpolarizing current to both excitatory populations, since 622 we do not have inhibitory populations in thalamic areas in the model. 623

To quantify the effect of single area or multiple areas inhibition, we calculate the average firing rate of areas that satisfy two conditions: i) the area shows persistent activity before

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inhibition and ii) the area does not receive inhibitory input. The ratio between such average 626 firing rate after inhibition and before inhibition is used to quantify the overall effect of 627 inhibition. If the ratio is lower than 100%, this suggests that inhibiting certain area(s) 628 disrupts the maintenance of the attractor state. Note that the inhibition effect is typically 629 not very strong, and only in rare cases, inhibition of a single area leads to loss of activity of 630 other areas (Fig. 5B, Fig. 5C). To quantify such differences, we use a threshold of 5% to 631 differentiate them. We will use (relatively) "weak inhibition effect" and "strong inhibition 632 effect" to refer to them afterwards. 633

We used the three measures to classify areas into 4 types (Fig. 5D): i) inhibition effect during delay period, ii) inhibition effect during stimulus period, and iii) delay period firing rate. Areas with strong inhibition effect during stimulus period are classified as input areas; areas with strong inhibition effect during delay period and strong delay period firing rate are classified as core areas; areas with weak inhibition effect during delay period but strong firing rate are classified as readout areas; areas with weak inhibition effect during delay period and weak firing rate during delay period are classified as nonessential areas.

⁶⁴¹ Simulation of multiple attractors

Multiple attractors coexist in the network and its properties and number depends on the 642 connectivity and dynamics of each node. In this study we did not try to capture all the 643 possible attractors in the network, but rather compare the number of attractors for different 644 networks. Here we briefly describe the protocol used to identify multiple attractors in the 645 network. We first choose k areas and then generate a subset of areas as the stimulation areas. 646 We cover all possible subsets, which means we run 2^k simulations in total. The external 647 stimulus is given to all areas in the subset simultaneously with same strength and duration. 648 The delay period activity is then quantified using a similar protocol as the standard simulation 649 protocol. The selection of k areas corresponds to a qualitative criterion. First we choose 650 the areas with small PV fraction or high hierarchy, since these areas are more likely to show 651 persistent activity. Second, the number of possible combination grows exponentially as we 652 increase k, and if we use k = 43, the number of combinations is around 8.8e+12, which is 653 beyond our simulation power. As a trade-off between the simulation power and coverage of 654 areas, we choose k = 18, which correspond to 2.6e+5 different combinations of stimulation. 655 For each parameter setting, we run 2.6e+5 simulations to capture possible attractor patterns. 656 For each attractor pattern, a binary vector is generated by thresholding delay firing rate using 657 a firing rate threshold of 5Hz. An attractor pattern is considered distinct if and only if the 658 binary vector is different from all identified attractors. In these way we can identify different 659 attractors in the simulation. We also apply same simulation pipeline to identify attractors 660 for different parameters. Specifically we change the long range connectivity strength μ_{EE} 661 and local excitatory connections $g_{E,self}$. 662

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Figure 1 - Supplement 1. Anatomical details of the mouse cortex. (A). Connectivity matrix depicting cortico-cortical connections between 43 cortical areas. Areas are sorted according to their hierarchy. (B). The raw PV cell density for each cortical area (Y axis), with areas sorted (X axis). Each area belongs to one of five modules, shown in color (see also Figure 1). (Harris et al. 2019). (C). Neuron density for each cortical area. The data is from Erö et al. 2018.



Figure 2 - Supplement 1. Example simulation for different sensory modalities. The simulation protocol is the same as the default one in Figure 2, except that the external input is applied to primary sensory areas related to two other sensory modalities: somatosensory and auditory. (A). The activity of 6 selected areas during the working memory task is shown. A somatosensory input of 500ms is applied to primary somatosensory area SSp-bfd, which propagates to the rest of the large-scale network. (B). Similar to the simulation where a primary visual area is stimulated (Fig 2D), delay period firing is positively correlated with cortical hierarchy. (r = 0.89, p < 0.05) (C) and (D) are similar to (A) and (B) except that the input is given to primary auditory area AUDp. (D). Delay period firing is also positively correlated with cortical hierarchy. (r = 0.89, p < 0.05)



Figure 3 - Supplement 1. Anatomical data of thalamus and cortical connectivity. (A). Connectivity matrix of corticothalamic connections: 43 cortical areas to 40 thalamic areas. (B). Connectivity matrix of thalamocortical connections: 40 thalamic areas to 43 cortical areas.



Figure 4 - Supplement 1. Details of cell type-specific connectivity measures. (A). The matrix of cell type projection coefficients between cortical areas. The cell type projection coefficient is given by the formula $k_{cell} = m_{ij} - PV_i(1 - m_{ij})$. (B). The matrix of connectivity strengths, modified by cell type projection coefficient between cortical areas. The modified connectivity strength is given by $\tilde{W}_{ij} = (m_{ij} - PV_i(1 - m_{ij}))w_{ij}$.



Figure 4 - Supplement 2. Cell type-specific eigenvector centrality measures are better at predicting firing rate patterns than raw eigenvector centrality measures. The analysis is the same as in Figure 4, where we compared cell type-specific input strength and raw input strength. Eigenvector centrality (EC, eigencentrality) of area i is the ith element of the leading eigenvector of the connectivity matrix. It quantifies how many areas are connected with the target area i and how important are these neighbors. Details are in the Methods section. (A(i)). Delay period firing rate (orange) and eigenvector centrality for each cortical area (blue). (A(ii)). Eigenvector centrality does not show a significant correlation with delay period firing rate for areas showing persistent activity in the model (r = 0.24, p = 0.29). (A(iii)). Eigenvector centrality cannot be used to predict whether an area shows persistent activity or not (prediction accuracy = 0.46). (B(i)). Delay period firing rate (orange) and cell type-specific eigenvector centrality for each cortical area (blue). (B(ii)). Cell type-specific eigenvector centrality has a strong correlation with the firing rate of cortical areas showing persistent activity (r = 0.94, p < 0.05). (B(iii)). Cell type-specific eigenvector centrality predicts whether an area shows persistent activity or not (prediction accuracy = 0.79). (C). Comparison of the correlation coefficient r for raw eigenvector centrality and cell type-specific eigenvector centrality in predicting delay firing rate. Raw input strength and cell type-specific input strength are also included for comparison. (D). Comparison of the prediction accuracy for raw eigenvector centrality and cell type-specific eigenvector centrality. Raw input strength and cell type-specific input strength are also included for comparison.



Figure 6 - Supplement 1. Cell type-specific loop strengths (Length 3 loops) are also better at predicting firing rate patterns than raw loop measures. Loop strengths (length 3 loops or L3) is calculated using similar method as loop strengths (length 2 loops). The only difference is we considered loops with length 3 (eg. A1->A2->A3->A1). The analysis is the same as in Figure 6, where we compared cell type-specific loop strengths (length 2 loops) and raw loop strengths. (A(i)). Loop strength (blue) is plotted alongside Core Areas (orange), a binary variable that takes the value 1 if the area is indeed a Core Area, 0 otherwise. (A(ii)). A high loop strength value does not imply that an area is a Core Area. (B(i)). Same as (A), but for cell type-specific loop strength. (B(ii)). High cell type-specific loop measures predicts that an area is a Core Area (prediction accuracy = 0.95). Same as (A), but for cell type-specific loop strength.

Main text: Distributed working memory in the mouse brain

Area	Supporting literature		
ALM (MOs)	(Kopec et al. 2015; Guo et al. 2014; Li et al. 2016),		
	(Inagaki et al. 2019; Erlich et al. 2011; Guo et al. 2017),		
	(Gilad et al. 2018; Gao et al. 2018; Wu et al. 2020)		
mPFC (PL/ILA)	(Liu et al. 2014; Schmitt et al. 2017),		
	(Bolkan et al. 2017)		
OFC	(Wu et al. 2020)		
PPC (VISa)	(Harvey et al. 2012)		
AIa (AId,AIv)	(Zhu et al. 2020)		
Area p (VISpl)	(Gilad et al. 2018)		
dorsal cortex	(Pinto et al. 2019)		
entorhinal (in vitro persistent activity)	(Egorov et al. 2002)		
piriform	(Zhang et al. 2019; Wu et al. 2020)		
VM/VAL	(Guo et al. 2017)		
MD	(Schmitt et al. 2017 ; Bolkan et al. 2017)		
superior colliculus	(Kopec et al. 2015)		
cerebellar nucleus	(Gao et al. 2018)		

Table 1: Supplementary experimental evidence. The listed literature include experiments that provide supporting evidence for working memory activity in cortical and subcortical brain areas in the mouse or rat. These studies show either that a given area is involved in working memory tasks and/or exhibit delay period activity. Area name corresponds to what has been reported in the literature. Some areas do not correspond exactly to the names from the Allen common coordinate framework.

MAIN TEXT: DISTRIBUTED WORKING MEMORY IN THE MOUSE BRAIN

Parameter	Description	Task/Figure	Value	
	Cortical circuit parameters			
$ au_{NMDA}$	NMDA synapse time constant	All figures	60 ms	
$ au_{GABA}$	GABA synapse time constant	All figures	5 ms	
$ au_{AMPA}$	AMPA synapse time constant	All figures	2 ms	
τ_{rates}	neuron time constant	All figures	20 ms	
τ_{noise}	noise time constant	All figures	2 ms	
a, b, d	parameters in excitatory F-I curve.	All figures	140 Hz/nA, 54 Hz, 308 ms	
g_I, c_1, c_0, r_{0I}	parameters in inhibitory F-I curve.	All figures	4, 615 Hz/nA, 177 Hz, 5.5 Hz	
γ	parameters in NMDA excitatory synaptic equations.	All figures	1.282	
γ_I	parameters in GABA synaptic equations.	All figures	2	
γ_A	parameters in AMPA excitatory synaptic equations.	All figures	2	
$g_{E,self}$	local self excitatory connections	Figures 1-6	0.4 nA	
$g_{E,cross}$	local cross population excitatory connections	All figures	10.7 pA	
g_{IE}	local E to I connections	All figures	0.4087 nA	
$J_{EI,min}, J_{EI,scaling}$	local I to E connection strength	All figures	0.192 nA, 0.16 nA	
$J_{II.min}, J_{II.scaling}$	local I to I connection strength	All figures	0.105 nA, 0.075 nA	
I_{0A}, I_{0B}	background current for excitatory neurons	All figures	0.305 nA	
I_{0C}	background current for inhibitory neurons	All figures	0.26 nA	
σ_A, σ_B	standard deviation of excitatory noise current	All figures	5 pA	
σ_C	standard deviation of inhibitory noise current	All figures	0 pA	
r_{0E}	background current for excitatory neurons	All figures	5 Hz	
r_{0I}	background current for excitatory neurons	All figures	$5.5~\mathrm{Hz}$	
μ_{EE}	long range E to E connection strength	Figures 1, 2, 4, 5, 6	0.1 nA	
μ_{IE}	long range E to I connection strength	Figures 1, 2, 4, 5, 6	0.167 nA	
β	parameters in m_{ij}	All figures	2.42	
kscale	parameters for scaling the connectivity matrix	All figures	0.3	
α_h, β_h	parameters for estimation of hierarchy	All figures	1.33, -0.22	
Istim	external stimulus strength	All figures	0.5 nA	
I _{inh}	external input to inhibitory neurons	All figures	5 nA	
T_{on}	stimulus start time	All figures	2 s	
T_{off}	stimulus end time	All figures	2.5 s	
T_{trial}	simulation time for each trial	All figures	10 s	
dt	simulation time step	All figures	$0.5 \mathrm{\ ms}$	
Thalamocortical network				
μ_{EE}	long range E to E connection strength	Figure 3	0.01 nA	
μ_{IE}	long range E to I connection strength	Figure 3	0.0167 nA	
g_{ct}	cortico thalamic connections strength	Figure 3	0.32 nA	
$g_{E,tc}$	thalamo-cortical connections to excitatory neurons	Figure 3	0.6 nA	
$g_{I,tc}$	thalamo-cortical connections to inhibitory neurons	Figure 3	1.38 nA	
	Simulation of multiple attractors	D: -		
μ_{EE}	long range E to E connection strength	Figure 7	0.01, 0.02, 0.03, 0.04, 0.05 nA	
μ_{IE}	long range E to I connection strength	Figure 7	0.0167, 0.033, 0.05, 0.066, 0.083 nA	
$g_{E,self}$	local self excitatory connections	Figure 7	0.4, 0.41, 0.2, 0.43, 0.44 nA	

Table 2: Parameters for numerical simulations

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