A Large-Scale Circuit Mechanism for Hierarchical Dynamical Processing in the Primate Cortex

Highlights

- Large-scale model of the macaque cortex with a gradient of synaptic excitation
- Sensory areas show fast responses while cognitive areas show slow integrative activity
- Multiple temporal hierarchies in the same anatomical network
- Functional connectivity analysis needs to incorporate inter-areal heterogeneity

Authors

Rishidev Chaudhuri, Kenneth Knoblauch, Marie-Alice Gariel, Henry Kennedy, Xiao-Jing Wang

Correspondence

xjwang@nyu.edu

In Brief

Chaudhuri et al. report a large-scale model of the macaque cortex incorporating quantitative anatomical data and inter-areal heterogeneity. This model gives rise to a hierarchy of timescales and suggests a revision of functional connectivity analysis of global brain dynamics.
A Large-Scale Circuit Mechanism for Hierarchical Dynamical Processing in the Primate Cortex

Rishidev Chaudhuri,1,2 Kenneth Knoblauch,3,4 Marie-Alice Gariel,3,4 Henry Kennedy,3,4 and Xiao-Jing Wang1,5,*

1Center for Neural Science, New York University, New York, NY 10003, USA
2Center for Learning and Memory, University of Texas at Austin, Austin, TX 78712, USA
3INSERM U846, Stem Cell and Brain Research Institute, 69500 Bron, France
4Université de Lyon, Université Lyon I, 69003 Lyon, France
5NYU-ECNU Institute of Brain and Cognitive Science, NYU Shanghai, Shanghai 200122, China
*Correspondence: xjwang@nyu.edu
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SUMMARY

We developed a large-scale dynamical model of the macaque neocortex, which is based on recently acquired directed- and weighted-connectivity data from tract-tracing experiments, and which incorporates heterogeneity across areas. A hierarchy of timescales naturally emerges from this system: sensory areas show brief, transient responses to input (appropriate for sensory processing), whereas association areas integrate inputs over time and exhibit persistent activity (suitable for decision-making and working memory). The model displays multiple temporal hierarchies, as evidenced by contrasting responses to visual versus somatosensory stimulation. Moreover, slower prefrontal and temporal areas have a disproportionate impact on global brain dynamics. These findings establish a circuit mechanism for “temporal receptive windows” that are progressively enlarged along the cortical hierarchy, suggest an extension of time integration in decision making from local to large circuits, and should prompt a reevaluation of the analysis of functional connectivity (measured by fMRI or electroencephalography/magnetoencephalography) by taking into account interareal heterogeneity.

INTRODUCTION

The receptive field is a central concept in neuroscience, defined as the spatial region over which an adequate stimulus solicits rigorous response of a neuron (Sherrington, 1906). In the primate visual cortical system, the receptive field size of neurons progressively enlarges along a hierarchy (Hubel and Wiesel, 1962; Hubel, 1988; Wallisch and Movshon, 2008). As a result, higher areas can integrate stimuli over a greater spatial extent, which is essential for such functions as size-invariance of object recognition in the ventral (“what”) stream for visual perception (Kobatake and Tanaka, 1994).

Accumulating evidence suggests that the brain also displays a hierarchy in the temporal domain. This allows neurons in higher areas to respond to stimuli spread over a greater temporal extent and to integrate information over time, while neurons in early sensory areas rapidly track changing stimuli. In human studies, preserving the short timescale structure of stimuli while scrambling long timescale structure changes responses in association areas but not early sensory areas (Hasson et al., 2008; Lerner et al., 2011; Honey et al., 2012; Gauthier et al., 2012; Stephens et al., 2013). Notably, using electrocorticography (ECoG), Honey et al. (2012) found that cortical areas sensitive to long time structure in the stimulus also show slower decays in their temporal autocorrelation (and hence slower dynamics), and Stephens et al. (2013) made a similar observation with fMRI. In the macaque, Murray et al. (2014) found a hierarchical organization in the timescales of spontaneous fluctuations of single neurons across 7 cortical areas, and an area’s timescale was well predicted by its position in the anatomical hierarchy of Felleman and Van Essen (1991). Similarly, temporal correlations in neural activity reveal slower decay rates in the frontal eye fields than area V4 (Ogawa and Komatsu, 2010), the timescales of reward memory lengthen from parietal to dorsolateral prefrontal to anterior cingulate cortex (Bernacchia et al., 2011), and, more generally, persistent activity after a brief stimulus can last for seconds, even across inter-trial intervals, in association areas (Amit et al., 1997; Histed et al., 2009; Curtis and Lee, 2010). Finally, normative theories of predictive coding suggest that a hierarchy of timescales would allow animals to form a nested sequence of predictions about the world (Kiebel et al., 2008).

What underlying neurobiological mechanisms might give rise to such a range of temporal dynamics? For example, spatial patterns of convergence can produce increasing receptive field sizes in the visual hierarchy. Are there basic anatomical motifs that produce a hierarchy of timescales?

Here we report a large-scale circuit mechanism for the generation of a hierarchy of temporal receptive windows in the primate cortex. This hierarchy naturally emerges in a dynamical model based on a recent quantitative anatomical dataset containing directed and weighted connectivity for the macaque neocortex (Markov et al., 2011, 2013b, 2014a; Ercsey-Ravasz et al., 2013). The data were obtained using the same experimental conditions and measures, ensuring a consistent database (Kennedy et al., 2013), and include both the number of projections between areas and their laminar origins. Based on a separate anatomical study (Elston, 2000; Elston et al., 2011), we introduced heterogeneity across cortical areas in the form of a gradient of excitatory
connection strengths. Strong recurrent excitation has been proposed as a mechanism by which prefrontal cortex could implement “cognitive-type” computations, such as information integration and memory-related delay activity; we hypothesized that differences in recurrent excitation might allow the generation of a temporal hierarchy.

The model thus incorporates anatomically constrained variation in both within-area and inter-area connectivity and enables us to probe the interplay of local microcircuitry and long-range connectivity that underlies a hierarchy of timescales. Using different sensory inputs, we demonstrate the existence, in our model, of multiple dynamical hierarchies subserved by a single integrated global and local circuit. We then investigate the implications of local circuit heterogeneity for macroscopic dynamics measured by functional connectivity (i.e., correlations in activity across areas). Here we find a disproportionate role for slow dynamics in the prefrontal and other association cortices in shaping resting-state functional connectivity. This role is not predicted by long-range connections, suggesting that interpretations of brain imaging data will need to be revised to account for inter-areal heterogeneity.

While we have used the model to investigate the origin of a hierarchy of timescales, it can be a platform for future models relating connectivity to dynamics and the functions of cortical areas. Most statistical analyses of connectivity (Bullmore and Sporns, 2009; Sporns, 2014) and computational models (Ghosh et al., 2008; Deco and Corbetta, 2011; Honey et al., 2007, 2009; Deco et al., 2014) have lacked comprehensive high-resolution data, relying either on collating qualitative tract-tracing data across disparate experiments and conditions or on diffusion tensor imaging, which is noisy and cannot reveal the direction of a pathway. Moreover, such models typically treat cortical areas as identical nodes in a network, distinguished by connection patterns but not by local properties or computational capabilities. Although this approach is reasonable for certain purposes, it is doubtful that functional specialization of cortical areas can be elucidated without considering heterogeneity. Our model provides a framework to explore how dynamical and functional specialization can emerge from inter-areal pathways coupled with local circuit differences.

**RESULTS**

We developed the model in three steps. First, we used recent connectivity data for the macaque neocortex (Markov et al., 2014a), designed to overcome the limitations of collated anatomical datasets, and collected by the same group under similar conditions, with quantitative measures of connectivity. The connectivity weights are directionally specific and cover 29 widely distributed cortical areas, with 536 connections whose strengths span five orders of magnitude (Figure 1). The presence or absence of all projections in this network has been established; thus, there are no unknown pathways.

Second, each cortical area was described by a threshold-linear recurrent network with interacting excitatory and inhibitory populations and calibrated by the neurophysiology of the primary visual cortex (Binzegger et al., 2009), but rescaled as described below. This is a highly simplified description of the dynamics of an area and ignores most within-area variability. In particular, note that the model is large-scale in that it addresses macroscopic cortical dynamics but is not large-scale in the sense of having millions of neurons or very high-dimensional activity. However, this level of complexity allows us to parsimoniously capture essential requirements for a hierarchy of timescales. We extend our results in Figure 7 and suggest further extensions in the Discussion.

Third, we hypothesized that the local microcircuit is qualitatively canonical (Douglas and Martin, 1991), i.e., the same across
areas, but that quantitative inter-areal differences are crucial in generating the timescales of areas. Specifically, the number of basal dendritic spines on layer three pyramidal neurons increases sharply from primary sensory to prefrontal areas (Elston, 2000; Elston et al., 2011). Taking spine count as a proxy for excitatory synapses per pyramidal cell, we introduced a gradient of excitatory input strength across the cortex. We modeled this by scaling the strength of excitatory projections in an area according to the area’s position in the anatomical hierarchy described below.

Gradient of Excitation along the Cortical Hierarchy

The laminar pattern of inter-areal projections can be used to place cortical areas in a hierarchy: neurons mediating feedforward connections from one area to another tend to originate in supragranular layers of the source area, whereas feedback projections tend to originate in infragranular layers (Felleman and Van Essen, 1991; Barbas and Rempel-Clower, 1997). This was quantified by Barone et al. (2000), who observed that the fraction of projecting neurons located in the supragranular layers of the source area defines a hierarchical distance between two areas; this allowed them to reproduce the hierarchy of Felleman and Van Essen (1991) using data from connections to only two areas (V1 and V4).

The laminar data included with this paper (see Table S1) contain hierarchical distance measured this way for all pairs of cortical areas included in the model (Figure 2A). We follow the approach of Markov et al. (2014b), and use these to estimate each area’s position in an underlying hierarchy. We found that an area’s position in this anatomical hierarchy is strongly correlated with counts of spines on pyramidal neurons in that area (Elston, 2007). This allowed us to introduce a systematic gradient of excitatory connection strength per neuron along the cortical hierarchy, and to explore how such heterogeneity interacts with the pattern of long-range projections to produce large-scale dynamics.

As a visual and conceptual aid, in Figure 2C we use a two-dimensional embedding to plot hierarchy and connectivity for the 29 areas. The angle between two areas reflects connection strength (closer areas have stronger connections), and the distance of an area from the center reflects hierarchy (higher areas closer to the center). The low-dimensional embedding is approximate but captures broad features of cortical organization and provides intuitive understanding of the model’s behavior. It suggests two hierarchical streams of sensory input originating in area V1 (primary visual cortex) and area 2 (part of primary somatosensory cortex) respectively, and converging on densely connected association areas. We next explored the response of the network to these sensory inputs.

Response to Visual Inputs

We simulated the response of the network to a pulsed input to primary visual cortex (area V1). The response is propagated up
the visual hierarchy, progressively slowing as it proceeds (Figure 3A). Early visual areas, such as V1 and V4, exhibit fast, short-lived responses. Prefrontal areas, on the other hand, exhibit slower responses and longer integration times, with traces of the stimulus persisting several seconds after stimulation. As with the response to a pulse of input, white-noise input is integrated with a hierarchy of timescales: the activity of early sensory areas shows rapid decay of autocorrelation with time whereas cognitive areas are correlated across longer periods (Figures 3B and 3C). Thus, a hierarchy of widely disparate temporal windows or timescales emerges from this anatomically calibrated model system.

To quantitatively compare areas, we fit single or double exponentials to the decay of each area’s autocorrelation function (see Figure S2 for plots of the fits). These fits capture a dominant characteristic timescale for each area in our model in response to visual stimulation. The time constants from the fits are plotted in Figure 3D, with areas ordered by position in the anatomical hierarchy. As can be seen from the bar plot, the dominant time-scale of an area tends to increase along the hierarchy (i.e., left to right), suggesting an important role for a gradient of excitation in generating the temporal hierarchy.

Nevertheless, an area’s timescales are not entirely determined by its hierarchical position, and the plotted timescales do not increase monotonically with hierarchy. To gain some intuition for the role of long-range projections in the model, consider area 8m (part of the frontal eye fields), which is low in the hierarchy and would show a rapid decay of correlation in the absence of long-range projections (far-right image of Figure 5A) but instead demonstrates long timescales in the model (and in the empirical observations of Hasson et al., 2008). As can be seen from Figure 2C, area 8m participates in a strongly-connected core of prefrontal and association areas (Ercsey-Ravasz et al., 2013; Markov et al., 2013b), allowing it to show long timescales that emerge from inter-areal excitatory loops (these timescales are strongly attenuated in the absence of feedback projections). The shared slower timescales are particularly characteristic of prefrontal areas in our model (see Figure S2, especially areas best fit by two timescales). Conversely, whereas area TEPd is high in the hierarchy, it does not participate in this core and is instead strongly coupled to ventral stream visual areas. Thus, it reflects the faster timescales of visual input.

**Multiple Functional Hierarchies**

The response to visual input reveals an ascending hierarchy of timescales in the visual system. We next stimulated primary somatosensory cortex (area 2), which is weakly connected to the visual hierarchy and strongly connected to other somatosensory...
A hierarchy of timescales (Figure 4A). However, the somatosensory and motor areas (Figure 2C). As previously, input propagates up a hierarchy of timescales (Figure 4A). However, the somatosensory response uncovers a different dynamical hierarchy to visual stimulation. Primary somatosensory cortex shows the fastest timescale, followed by primary motor cortex (area F1) and somatosensory association cortex (area 5). Parietal and premotor areas show intermediate timescales and, as with visual stimulation, prefrontal areas show long timescales. Visual areas demonstrate much weaker responses than before and are mostly driven by top-down projections from association areas. Thus, in the absence of direct input, they reflect the slower timescales of a distributed network state. In Figure 4B, we contrast time constants for visual and somatosensory stimulation across areas.

An area’s timescales emerge from a combination of local circuit properties, the specificity of long-range projections, and the particular input to the network. Our model allows us to examine the contribution of each. These can be intuitively summarized by noting that each area in Figure 2C shows timescales approximately determined by its distance from the periphery (hierarchical position), proximity to the central clusters (long-range connectivity), and distance from the source of input.

**Role of Local and Long-Range Projections**

To further dissect the contributions of local and long-range projections, we examined time constants in response to visual input after removing either differences in local micrcircuitry or inter-areal projections. In the second image of Figure 5A, we show that the range of timescales is drastically reduced in the absence of differences in the microcircuit across areas. Moreover, there is no longer a relationship to an area’s position in the anatomical hierarchy. Thus, while differences in long-range inputs and outputs to each area are significant, they are insufficient to account for disparate timescales and local heterogeneity is needed.

In the third image of Figure 5A, we show the effect of removing long-range feedback projections, and for the far right image, we remove all long-range projections and stimulate individual areas separately. The range of time constants is lower, reflecting the propensity of slow areas to form long-range excitatory loops with each other. More significantly, once long-range projections are removed, an area’s time constant simply reflects its position in the hierarchy.

We extend our investigation of the role of long-range projections by contrasting the resting-state response (i.e., equal white-noise input to all areas) of the intact network to networks where long-range connections are scrambled while preserving the gradient of excitation. A number of these networks show responses that are poorly fit by exponentials, so we measure timescale non-parametrically as the time after pulse offset for activity to decay to within 5% of baseline. In Figure 5B, we show that scrambling almost entirely removes the hierarchy of timescales, further confirming that a gradient of excitation alone is insufficient to separate timescales.

The connectivity data show specificity in which projections exist and in their strengths, and both connection probability and strength decay exponentially with inter-areal distance (Markov et al., 2011, 2013b, 2014a; Ercsey-Ravasz et al., 2013). In Figure 5C, we preserve network topology (i.e., which areas are connected), but scramble the strengths of non-zero projections. Here the separation of timescales is strongly attenuated for most areas, suggesting that specificity in projection strengths and not just network topology is required for the timescales we see.

**Localized Eigenvectors and Separated Timescales**

The model for a single area is threshold-linear, meaning we ignore nonlinearities besides the constraint that firing rates be positive. This allowed us to explore the genesis of separated timescales with linear systems analysis. The activity of a linear network is the weighted sum of characteristic activity patterns, called eigenvectors (Rugh, 1995). Each eigenvector evolves on a timescale given by a corresponding eigenvalue and is differentially driven by different inputs.

The eigenvectors of the linearized network are localized: those with short timescales are broadly concentrated around sensory areas and those with long timescales are concentrated at frontal areas (Figure 6). In general, if an eigenvector is small at a node then its amplitude at that node in response to input will also be

![Figure 4. The Response to Somatosensory Input Reveals a Different Functional Hierarchy Subserved by the Same Anatomical Network](image-url)

(A) Autocorrelation of activity for areas that show strong responses to input to area 2 (part of primary somatosensory cortex). Area labels are arranged according to position in the underlying anatomical hierarchy. Inset: time constants fitted to the autocorrelation function for each area.

(B) Timescales in response to visual (left) and somatosensory input (right) shown with lateral (top) and medial (bottom) views of the cortex. See also Figure S4.
small, and the corresponding timescale will be weakly expressed. Thus, localization means that for most inputs network dynamics will be dominated by rapid timescales at sensory areas and slower timescales at cognitive areas. In previous theoretical work, we have shown how localized eigenvectors can arise in networks with gradients of local properties and produce a diversity of timescales (Chaudhuri et al., 2014).

Extension to Nonlinear Dynamics and Multistability

The threshold-linear local circuit let us highlight the requirements for a hierarchy of timescales and provide intuition from linear systems theory. Moreover, many systems can be linearly approximated, and neural responses are often near linear over a wide range of inputs (Wang, 1998; Chance et al., 2002), making linear and threshold-linear models useful for neural circuits (Dayan and Abbott, 2001).

Nevertheless, linear models show limited dynamics and cannot capture features such as persistent activity or multistability, which are thought to be important for cognitive capabilities in higher areas (Wang, 2013). We thus replaced our local circuit with a firing rate (“mean-field”) version of a spiking network with more realistic synaptic dynamics (Wang, 2002; Wong and Wang, 2006). When isolated, an area in this network can display qualitatively different regimes (Figure 7A). For relatively weak recurrent connections, an area shows a single stable state. As recurrent excitation is increased, there is a transition to a regime with two stable states, with low and high firing rates that correspond to a resting state and a self-sustained persistent activity state. In this regime, an area can integrate inputs over time and maintain activity in the absence of a stimulus. Such dynamical regimes have been proposed to underlie “cognitive-type” computations such as working memory and decision-making (Wang, 2002, 2013).

With this model for each area in the large-scale network, we introduced the previous gradient of excitation. Consequently, sensory areas show single stable states while areas further up...
the hierarchy can also show persistent activity when driven by strong inputs (Figure 7B). Small perturbations are insufficient to shift the state of a node but take longer to decay away in areas further up the hierarchy (Figure 7C).

For small inputs, the network response resembles the threshold-linear model: a brief input to V1 is propagated up the hierarchy, with rapid decays in sensory areas and slow decays in association areas (Figure 7D). Thus, the previous results extend to a nonlinear model with a larger dynamical repertoire. Exploring the complex dynamical behaviors that this network can show is beyond the scope of this paper, but one interesting consequence of the extended model is that the timescales of small fluctuations around baseline predict the ability of an area to show much longer timescales in response to larger inputs (Figure 7C and see Discussion), as observed in Honey et al. (2012) and Murray et al. (2014).

Functional Connectivity

We now investigate the implications of local heterogeneity for network organization as measured by correlations in resting-state activity (resting-state functional connectivity). In our model, frontal and association areas reflect a slowly varying network state, and we hypothesized that this state should strongly shape functional connectivity.

In Figure 8A, we show functional connectivity in our threshold-linear model with heterogeneity in local area properties, or without it (as typically assumed in models relating functional to anatomical connectivity). The inclusion of a gradient of local excitation reduced the correlation ($r^2$) between functional and anatomical connectivity from 0.83 to 0.53 (Figure S6 shows results using a BOLD kernel [Boynton et al., 1996]).

Multiple studies find that the strength of an anatomical connection between areas (“structural connectivity”) partially predicts correlations in neurophysiological signals from those areas (functional connectivity), but there are significant differences (Hagmann et al., 2008; Honey et al., 2009; Damoiseaux and Greicius, 2009; Honey et al., 2010; Deco and Corbetta, 2011; Deco et al., 2014). Our results also suggest that inter-areal connections are insufficient to predict functional connectivity. However, we find that heterogeneity in local connectivity could help account for the previously unexplained variance.

In our model, slower frontal and temporal areas in particular show enhanced functional connectivity. Consequently, areas with slow timescales play a predominant role in the network, as shown by “lesioning” individual areas (Figure 8B, left panel). For the simple case of identical input to each area, the effect of lesioning an area is well predicted by the time constant of intrinsic fluctuations (Figure 8B, right panel). Note that areas most important for functional connectivity are not simply those at the highest positions in the hierarchy (i.e., with the most recurrent connections), and hierarchy alone poorly predicts impact on functional connectivity ($r^2 = 0.18$). For instance, the caudal superior temporal polysensory region (STPc) and the rostral parabelt (PBr) are at intermediate hierarchical positions but have strong connections to other parts of STP (darker lines in Figure 8B) forming a cluster that shapes functional connectivity. In general, areas combining intermediate to high hierarchical position and strong connections to slow areas have the strongest influence on global activity patterns.

**DISCUSSION**

The main findings of this work are 3-fold. First, it establishes a circuit mechanism for a hierarchy of temporal receptive windows, which has received empirical support in recent human (Hasson et al., 2008; Lerner et al., 2011; Honey et al., 2012; Gauthier et al., 2012; Stephens et al., 2013) and single-unit monkey experiments (Murray et al., 2014). The model extends time integration in decision making from local circuits (Wang, 2008) to a large-scale system across multiple timescales (Hasson et al., 2015). Second, inter-areal heterogeneity implies that areas cannot be treated as identical nodes of a network and slow dynamics in association areas can play a disproportionate role in determining the pattern of functional connectivity. This suggests that functional connectivity analyses be revised. Third, this is the first large-scale dynamical model of the macaque cortex based on weighted and directed connectivity and incorporating heterogeneity across areas.

The ability to integrate and hold information across time is critical for cognition. On the other hand, the brain must rapidly and transiently respond to changing stimuli. Complex behavior thus requires a multitude of coexisting timescales. We demonstrate how such timescales (or temporal receptive windows) naturally emerge in a model of primate cortex, built with quantitative anatomical data. Our work reveals multiple functional hierarchies converging on a slow distributed network of densely connected frontal and other association areas.

A long-standing observation is that strong recurrent connections can produce slower dynamics (Wang, 2008), and we show how this basic anatomical motif can interact with the pattern of long-range connections to produce a hierarchy of timescales. The hierarchies we observe with different stimuli...
thus emerge from a combination of heterogeneity in excitatory connection strengths across areas and the profile of long-range connectivity (which is highly specific to each area (Markov et al., 2013a)), and neither alone can predict an area’s timescales. For example, while differences in local recurrence play a crucial role in generating timescales, the correlation between anatomical hierarchy and timescale is relatively weak ($r^2 = 0.25, 0.14, 0.22$ in the visual, somatosensory, and resting-state conditions, respectively). Moreover, areas can show quite different timescales in response to different inputs: as seen in Figure 4B, even early visual areas with relatively weak recurrence can have slower timescales. To characterize the dependence of timescales on local and long-range properties, we first removed the gradient of local properties and observed that the hierarchy of timescales vanishes. Separately, we preserved the local properties of areas and either removed (Figure 5A, right panels) or scrambled the long-range projections both globally and while preserving network topology (Figures 5B and 5C).

It will be important to further probe the interaction of local and long-range connectivity. This will require additional anatomical and physiological data, and our model can be a platform to explore the consequences of these data for large-scale dynamics. For example, following the finding of Markov et al. (2011) that the proportion of local to long-range synapses is roughly conserved across areas, we have chosen to scale both local and long-range projections by an area’s position in the hierarchy. Nevertheless, local and long-range synapses may have different strengths and properties and may differentially target cell types and dendritic locations. Relatedly, long-range inputs may be differentially gated depending on task demands and the local circuit regime. Conversely, in the nonlinear model, long-range input can shift the dynamical regime of the local circuit: an area that lacks persistent activity when isolated may show persistent activity in the presence of a weak long-range control signal. These interactions can provide the network with an enhanced computational repertoire.
To examine timescales in the clearest way possible, we modeled individual areas with a threshold-linear rate model, where time constants are mathematically well defined. However, the results hold for a nonlinear local circuit with multiple stable states. Note that this work did not focus on the latency of neural responses (Schmolesky et al., 1998; Bullier, 2001), for which a spiking model is needed. Nevertheless, single neurons in the monkey cortex display slow responses during stimulus presentation as shown in the model; for example, in decision tasks prefrontal and parietal neurons can show quasi-linear ramping with a time constant that may appear effectively infinite (Smith and Ratcliff, 2004; Gold and Shadlen, 2007; Wang, 2008; Brunton et al., 2013). Thus, the model is the simplest that is adequately designed to reveal a hierarchy of timescales in the cortex.

Our model has several testable predictions. Though there are multiple combinations of local time constants and network connection strengths that could produce a particular set of observed timescales, the model suggests that timescales of small fluctuations should reflect the intrinsic properties of areas (far right panel of Figure 5A), while larger responses should reflect time constants that emerge from the entire system (far left panel of Figure 5A). In the model, slow network timescales
are driven by strongly connected frontal and temporal areas, corresponding to a slowly varying global state. Inactivating these areas should decrease slow dynamics in connected areas lower in the hierarchy. The differential responses to visual and somatosensory input suggest that when a particular input is not involved in a task, the corresponding sensory areas better reflect slow changes in global cortical state. This may explain decreases in low-frequency EEG power (i.e., slow modes) when a subject engages in a task (He et al., 2010; Honey et al., 2012), as well as the observation of Stephens et al. (2013) that, despite fast timescales in response to visual input, early visual areas have slow timescales during auditory processing. Finally, we predict that areas with longer timescales, such as prefrontal and superior temporal areas, can shape functional connectivity to a greater degree. This highlights the importance of incorporating heterogeneous local dynamics in studying the determinants of functional connectivity and, intriguingly, suggests that functional connectivity might be used to probe local properties. Whereas there is some evidence that frontal and association areas show enhanced functional connectivity (Sepulcre et al., 2010) and of a correlation between enhanced functional connectivity and slow timescales (Baria et al., 2013), it would be interesting to use functional imaging to better understand the link between functional connectivity and response timescales (for example, as determined by the approach of Hasson et al. [2008], Lerner et al. [2011], Honey et al. [2012], and Gauthier et al. [2012]). The link between slow timescales and enhanced functional connectivity might also explain observations that functional connectivity is greater at low frequencies (Salvador et al., 2005). Moreover, because distant areas tend to lack strong direct connections, their functional connectivity will be primarily driven by slow distributed network modes and will be further biased toward low frequencies, as previously observed (Salvador et al., 2005).

We mostly used a threshold-linear model for local areas, but the hierarchy of timescales holds when areas are modeled by a nonlinear microcircuit, similar to one proposed as a model for general “cognitive-type” computations (Wang, 2002, 2013). Depending on connectivity and input parameters, such networks show a single stable state, multistability with persistent firing, or continuous slow fluctuations between metastable states. While we do not explore this broader range of behaviors, note that in the nonlinear model the timescales of small fluctuations around baseline predict an area's ability to show much longer timescales in response to larger inputs. This can be seen by comparing the timescales of Figure 7C with the steady states of Figure 7A, and by contrasting responses to large and small perturbations in Figures 7B and 7D (note that timescales in response to large perturbations tend to be slower than those from small perturbations even if the area is not bistable). This may explain why the timescales of spontaneous fluctuations in an area (on the order of hundreds of milliseconds) correlate with its sensitivity to temporal structure in stimuli across seconds (Honey et al., 2012) as well as with slow drifts in baseline neural activity and the timescales of reward memory (Murray et al., 2014).

Our model is parsimonious, designed to capture a basic mechanism underlying a hierarchy of timescales, and can be extended in several ways. First, the local area model could be made more complex, and an interesting direction is using the SLNs to incorporate a laminar structure. Second, in our model activity propagates along the hierarchy with significant attenuation. This attenuation can be substantially decreased by changing model parameters (M. Joglekar and X.-J.W., unpublished data) and may be removed by synchronous firing (Diesmann et al., 1999) or more sophisticated feedback projections (Moldakarimov et al., 2015). Third, we only consider cortico-cortical connections. Whereas these form the major input to a cortical area (Markov et al., 2011), subcortical projections will play an important role. For example, incorporating thalamocortical projections would allow us to more realistically model input and may help set network state and gate inter-areal interactions, whereas neuromodulators such as acetylcholine might modulate the excitability of local populations and enhance information transmission at other synapses. Fourth, as a first step, we used two global parameters to scale long-range connection strengths but emerging data relating long-range anatomy and physiology should be incorporated. Fifth, extensions should include other inter-areal heterogeneities, such as in interneuron types and densities (Medalla and Barbas, 2009) and in neuromodulatory signaling (Hawrylycz et al., 2012). For example, it would be interesting to model the higher numbers of dopaminergic projections to prefrontal areas. Finally, while we have focused on how areas are able to accumulate incoming information on different timescales, processing input requires synthesizing it with previous input. Future work should explore how different areas in our model integrate information from more realistic time-varying stimulation such as a movie or a song and to probe how these responses change when the correlation structure of the input is disrupted (for example, by scrambling).

In conclusion, we report a novel, quantitatively calibrated, dynamical model of the macaque cortex with directed and weighted connectivity. The identification of a specific circuit mechanism for a hierarchy of timescales (temporal receptive windows) represents a key advance toward understanding specialized processes and functions of different (from early sensory to cognitive-type) cortical areas. Our findings demonstrate the importance of heterogeneity in local areal properties, as well as the specific profile of long-range connectivity, in sculpting the large-scale dynamical organization of the brain.

**EXPERIMENTAL PROCEDURES**

**Anatomical Data**

Connectivity data are from an ongoing project to quantitatively measure all connections between cortical areas in the macaque (Markov et al., 2014). Inter-areal connection strengths are measured by counting projecting neurons labeled by retrograde tracer injections and normalizing by the total number of neurons labeled in the injection, yielding a fractional weight or FLN (fraction of labeled neurons) for each pathway:

\[
FLN_{B \rightarrow A} = \frac{\# \text{ neurons projecting to area A from area B}}{\text{total neurons projecting to area A from all areas}}
\]

So far, 29 areas have been injected and we use the subnetwork consisting of these areas. The presence or absence of all connections is known bidirectionally, and 66% of possible connections exist, with widely varying strengths.
We also use data on the fraction of neurons in each projection that originate in
the upper layers of the source area (SLN, for supragranular layer neurons [Markov et al., 2014b]) defined as:

\[
SLN_{A \rightarrow B} = \frac{\text{# supragranular neurons projecting to area A from area B}}{\text{# neurons projecting to area A from area B}}
\]

Data are in Table S1 and can also be accessed at http://core-nets.org/. Further
details of data collection can be found in Markov et al., 2014a, 2014b. All the
procedures used in the study followed the national and European regulations
concerning animal experiments (EC guidelines 86/609/EC) and were approved by
the authorized national and veterinary agencies.

Hierarchy and Connectivity Embedding
To extract the hierarchy, we follow observations from the visual system that the
fraction of projections originating in the supragranular layers of the source area
(the SLN) measures hierarchical distance between the source and target areas
(Felleman and Van Essen, 1991; Barone et al., 2000; Markov et al., 2014b).
We use a generalized linear model to assign hierarchical values to areas such as
that the differences in hierarchical values predict the SLNs (similar to the method
in Markov et al., 2014b).

For Figure 2C, we compute angles \( \theta \), so that the angular distances between
areas \( A_i \) and \( A_j \) correspond to dissimilarity measured as \(-\log(\text{SLN}_{A_iA_j})\). We
then plot the areas on a polar plot with \( \theta(A_i) = \theta_i \) and \( R(A_i) = \sqrt{1 - \theta_i} \).

See the Supplemental Experimental Procedures and Figure S1 for an
expanded discussion of the hierarchy and the circular embedding.

Model Architecture
Each area consists of an excitatory and an inhibitory population described by

\[
\tau_e \frac{d}{dt} v^E_k = -v^E_k + \beta^E \left[ 1 + \eta h_1 \right] \left( w_{EE} v^E_k + \mu_{EE} \sum_{j=1}^N \text{FLN}_{ij} v^E_j \right) - w_{EI} v^I_k + I_{\text{ext,E}}
\]

\[
\tau_i \frac{d}{dt} v^I_k = -v^I_k + \beta^I \left[ 1 + \eta h_1 \right] \left( w_{EI} v^E_k + \mu_{EI} \sum_{j=1}^N \text{FLN}_{ij} v^E_j \right) - w_{II} v^I_k + I_{\text{ext,I}}
\]

\( v^E_k \) is the firing rate of the \( k \)-th excitatory population, with intrinsic time
constant \( \tau_e \), couplings \( w_{EE} \) and \( w_{EI} \) from the local excitatory and inhibitory
population, and external input \( I_{\text{ext,E}} \) (both stimulus input and any noise we add
to the system). The inhibitory population has corresponding parameters \( \tau_i \), \( w_{EI} \), \( w_{II} \),
and \( I_{\text{ext,I}} \). The \( I \)-f curves are threshold linear, with slope \( I_{\text{th,E}} \) and \( I_{\text{th,I}} \) control the strengths of long-range input
to the excitatory and inhibitory populations, and do not vary between connect-
ions: all specifity comes from the FLNs. \( \eta \) scales both local and long-range excitatory
inputs to an area by its position in the hierarchy, \( h_i \). We set \( \tau_e = 20 \text{ ms}, \tau_i = 10 \text{ ms}, \text{IE} = 0.066 \text{ Hz/pA}, \text{IE} = 0.351 \text{ Hz/pA}, \text{EE} = 24.3 \text{ Hz/A}, \text{EI} = 12.2 \text{ pA/Hz}, \text{II} = 19.7 \text{ pA/Hz}, \text{EI} = 12.5 \text{ pA/Hz}, \text{IE} = 33.7 \text{ pA/Hz}, \text{II} = 25.3 \text{ pA/Hz} \) and \( \eta = 0.68 \). For more details, see the Supplemental
Experimental Procedures.

We mostly ignore inter-area conduction delays; however, see Figure S3 for
a network with conduction delays.

Pulse Input, Autocorrelation, and Fitted Time Constants
For Figures 3, 4, 5, and 6, we choose the background input for each area so that
the excitatory and inhibitory populations have rates of 10 and 35 Hz,
respectively.

In Figure 3A, V1 receives a 250 ms pulse of input that drives its rate to
100 Hz. For the remaining images of this figure and Figure 5A, the stimulus
to V1 is a white noise with a mean of 2 Hz and a SD of 0.5 Hz. The other areas
receive a small amount of background input (SD on the order of 10^{-3}), but are
primarily driven by long-range input propagating out from area V1. For
Figure 4, the currents are the same except that area 2 receives the stimulus
than V1.

For each area, we extract time constants by fitting both one and two expo-
nentials to the part of the autocorrelation function that decays from 1 to 0.05. If
the sum of squared errors of the single exponential fit is less than eight times
that of the double exponential, then we report that time-constant. Otherwise,
we use the sum of time constants from the double exponential fit, with each
weighted by its amplitude. Fits in response to V1 and area 2 input and for
resting state activity are shown in Figures S2, S4, and S5.

For Figure 4B, we map the time constants logarithmically to a heatmap and
plot them using Caret (Van Essen et al., 2001).

Functional Connectivity
To highlight the effect of intrinsic hierarchy, in Figure 6A we contrast a network
without hierarchy with a network that has a gradient of local excitatory connec-
tions but unlike in the remaining figures, no gradient in the long-range projec-
tion strengths (thus, these networks have the same long-range projection
strengths and differences emerge from local properties). We replace

\[
(1 + \eta h_1) \left( w_{EE} v^E_k + \mu_{EE} \sum_{j=1}^N \text{FLN}_{ij} v^E_j \right) + \mu_{EI} \sum_{j=1}^N \text{FLN}_{ij} v^I_k
\]

for the excitatory population, and similarly for the inhibitory population. For
Figure 6B, we use the same network as elsewhere; so, all ingoing excitatory
projections are scaled by an area’s hierarchical position.

We calculate functional connectivity as the correlation matrix of area activity
in response to equal white-noise input to all areas. For Figure 8B, we determine
this correlation matrix analytically (see Supplemental Experimental Proce-
dures). The effect of lesioning an area, \( A_i \), is measured as \(|C_{A_i \rightarrow \text{area}}(X) - |C_{\text{area} \rightarrow \text{area}}|X|\)
where \( C_{A_i \rightarrow \text{area}} \) is the correlation matrix after lesioning \( A_i \), \( C_{\text{area} \rightarrow \text{area}} \) is the intact correla-
tion matrix without the row and column corresponding to \( A_i \), and the double
lines indicate the norm. The values are then scaled to lie between 0 and 1.

Nonlinear Network
The nonlinear single area model is a variant of a model proposed in Wong and
Wang (2006) as an approximation to a spiking network with AMPA, GABA and
NMDA synapses (Wang, 2002). Each area is described by:

\[
v^E_k = \varphi \left( 1 + \eta h_1 \right) \left( w_{EE} s^E_k + \mu_{EE} \sum_{j=1}^N \text{FLN}_{ij} s^E_j \right) - w_{EI} s^I_k + I_{\text{ext,E}}
\]

\[
v^I_k = \gamma \left( 1 - s^E_k \right) v^E_k - \gamma \tau_k \left( 1 - s^I_k \right) v^I_k
\]

\( v^E_k \) and \( v^I_k \) are excitatory and inhibitory firing rates, \( s^E_k \) is a gating variable corre-
sponding to NMDA synapses (with decay timescale \( \tau_E \)) and \( s^I_k \) is a simplified
f-I curve from Abbott and Chance (2005). We set \( \tau_E = 60 \text{ ms}, \tau_I = 10 \text{ ms}, \gamma = 0.641, \text{WE} = 250.2 \text{ pA}, \text{WE} = 8.110 \text{ pA} \), \( \text{EE} = 330.9 \text{ pA}, \text{EI} = 12.5 \text{ pA} \).

For Figures 7A–7C, we remove long-range connections and characterize an
isolated area. The bifurcation diagram of Figure 7A shows network steady
states as we vary the hierarchy scaling (i.e., \( 1 + \eta h_1 \)), whereas Figure 7C shows the slowest timescale of the Jacobian around the low firing state.

For Figure 7B, we set \( \eta = 3.4 \) and give a 100 Hz pulse of input for 250 ms to the
two disconnected areas at opposite ends of the hierarchy (V1 and 24c). For
Figure 7D, we consider a connected network, with long-range projections only
targeting excitatory subpopulations, for simplicity, and set \( \text{EE} = 125.1 \text{ pA} \).
We give a 200 Hz pulse of input to area V1 for 250 ms.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures,
six figures, and one table and can be found with this article online at http://dx.
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REFERENCES


Supplemental Information

A Large-Scale Circuit Mechanism for Hierarchical Dynamical Processing in the Primate Cortex

Rishidev Chaudhuri, Kenneth Knoblauch, Marie-Alice Gariel, Henry Kennedy, and Xiao-Jing Wang
Figure S1. Related to Figure 2. Hierarchy fitted from pairwise SLN relationships. (A) Left panel: Hierarchy fitted from logistic regression (and used in main text). The hierarchical position of an area is normalized to lie between 0 and 1. Right panel: Hierarchy fitted from beta regression (Cribari-Neto and Zeileis, 2010). (B) SLN values predicted from logistic regression compared to observed SLNs.
Figure S2. Related to Figure 3. Timescales in response to white-noise input to V1. Data shown in grey, single exponential fits in blue and double exponential fits in dark red. For double exponential fits, $\tau_1$ and $\tau_2$ are the time-constants of individual exponentials, and $\tau$ is a weighted average of $\tau_1$ and $\tau_2$, with weights given by the amplitudes of the exponentials.
Figure S3. Related to Figure 3. Response of a network with inter-areal conduction delays. (A) Distances (in mm) between the nodes of the network (Ercsey-Ravasz et al., 2013). (B) Response of the network to a pulse of input to area V1. Conduction delays between nodes are imposed using the distances in panel A and a conduction velocity of 1.5 m/s (Deco et al., 2009).
Figure S4. Related to Figure 4. Timescales from exponential fits of activity in response to white-noise input to Area 2. Colors as in Figure S2.
Figure S5. Related to Figure 8. Timescales from exponential fits of resting-state activity (i.e., equal white-noise input to all areas). Colors as in Figure S2.
Figure S6. Related to Figure 8. Functional connectivity of simulated BOLD signal. (A) As in Figure 8A, the network on the left has the same local properties at each node, while the network on the right has a gradient of local recurrent strengths. Firing rate is convolved with a gamma function to generate a simulated BOLD signal (Boynton et al., 1996). Top panel: functional connectivity in response to background white noise input to each node. Bottom panel: functional connectivity (correlations in BOLD) vs. structural connectivity (FLN) for non-zero projections. (B) Effect of lesioning areas on functional connectivity measured via simulated BOLD signal. Plots are as in Figure 8B.
Supplemental Experimental Procedures

Several of these sections are expanded versions of the corresponding sections in Experimental Procedures in the main text. To make these descriptions self-contained, the relevant portions from the main text are repeated here.

Anatomical data

The connectivity data are from an on-going project to quantitatively measure all connections between cortical areas in the macaque cortex, with areas defined according to a 91 area parcellation scheme (Markov et al., 2014a). Descriptions of data collection can be found in Markov et al. (2011; 2014a). Briefly, connection strengths between areas are measured by counting the number of neurons labeled by retrograde tracer injections. The number of neurons labeled in a projection ranges from a few neurons to on the order of 100,000 neurons. To control for injection size, these counts are then normalized by the total number of neurons labeled in the injection, yielding a fractional weight or FLN (Fraction of Labeled Neurons) for each pathway, defined as

\[
FLN_{B \rightarrow A} = \frac{\# \text{ neurons projecting to area } A \text{ from area } B}{\text{total neurons projecting to area } A \text{ from all areas}}
\]

The corresponding weights span 5 orders of magnitude. So far, 29 areas have been injected and we use the subnetwork consisting of these areas. In this network the presence or absence of all connections is known bidirectionally, and 66% of possible connections exist in the network, though with widely varying strengths.

We also use data on the fraction of neurons in each projection that originate in the upper layers of the source area, which we call the SLN, for Supragranular Layer Neurons (Markov et al., 2014b). These are defined as

\[
SLN_{B \rightarrow A} = \frac{\# \text{ supragranular neurons projecting to area } A \text{ from area } B}{\# \text{ neurons projecting to area } A \text{ from area } B}.
\]

The data are included in Table S1, and all data can be downloaded from www.core-nets.org.

Hierarchy and low-dimensional connectivity embedding

In the visual system, projections directed from early visual areas to higher-order areas (i.e. increasing size of receptive field, position-invariance, and so on) tend to originate in the supragranular layers of the cortex and terminate in layer 4 (Felleman and Van Essen, 1991; Barone et al., 2000). Conversely, projections from higher-order areas to early visual areas
originate in the infragranular layers and terminate outside of layer 4. This observation was systematized by Felleman and Van Essen (1991), who used these anatomical constraints to place cortical areas in a hierarchical ordering.

Felleman and Van Essen used a discrete classification of projections: in their framework projections are either feedforward, feedback or lateral depending on where the majority of projections originate and terminate. However, such binary relations are typically insufficient to specify a unique hierarchy (Hilgetag et al., 1996). Subsequently, it was observed that rather than classifying a projection as feedforward, feedback or lateral, the fraction of neurons in a projection originating in the supragranular layers (the SLN) could be used as a continuous measure of hierarchical displacement: the difference of the SLN from 50% is positive for feedforward projections and negative for feedback projections, and its magnitude gets larger as a projection moves further away from lateral (Barone et al., 2000). For example, a projection with an SLN of 90% would be very strongly feedforward, while a projection with an SLN of 65% would be only moderately feedforward. Using these values, the Felleman and van Essen hierarchy could be reproduced using observations of connections to only two areas (V1 and V4) (Barone et al., 2000).

To construct the hierarchy we follow a similar framework to Markov et al. (2014b) and use a generalized linear model. We assign hierarchical values to each area such that the difference in values predicts the SLN of a projection. Specifically, we assign a value $H_i$ to each area $A_i$ such that

$$SLN_{A_j \rightarrow A_i} \approx g^{-1}(H_i - H_j).$$

We choose $g^{-1}$ to be a logistic function (logistic regression), which is standard for probabilities and fractional values, but we note that other functions yield similar values (Figure S1A). We have one such constraint for each projection (536 in total), and we find the set of hierarchical values that best fit these constraints. In the fit we weight the contribution of each projection by the log of its FLN to preferentially match stronger and less noisy projections. The resulting best fit hierarchy is shown in the left panel of Figure S1A. We then normalize by the maximum hierarchical value yielding $h_i = H_i/H_{max}$.

We extract the spine counts in Figure 2B from Elston (2007) and plot the areas in common with our data set. The parcellation in that paper is coarser than the parcellation we use, so we report the results in terms of that parcellation. For area 7 we average together the hierarchical positions of 7A, 7B and 7m; for 6 we average F2, F5 and F7; and for 46 we average together 46d, 9/46d and 9/46v.

For the two-dimensional circular embedding of Figure 2C, we convert the FLN to a measure of dissimilarity according to

$$d(A_i, A_j) = \begin{cases} -\log(\text{FLN}(A_i, A_j)) & \text{for } \text{FLN}(A_i, A_j) > 0, \\ -\log(\text{FLN}_{\text{min}}) & \text{for } \text{FLN}(A_i, A_j) = 0. \end{cases}$$

Here, $A_i$ is the $i$th area, and $\text{FLN}_{\text{min}}$ is some value less than the smallest FLN in the network.
We use $\text{FLN}_{\text{min}} = 10^{-7}$ but the results are robust to the precise choice of this value. We then assign angles $\theta_i$ to each area such that $d(A_i, A_j) \approx R \min(|\theta_i - \theta_j|, 2\pi - |\theta_i - \theta_j|)$, where $R$ is a single free parameter. We fix area $V_1$ to have $\theta = 0$, but choosing any other area to fix would simply rotate the plot. Finally, we plot the areas on a 2-dimensional polar plot with $\theta(A_i) = \theta_i$ and $R(A_i) = \sqrt{1-h_i}$.

**Model architecture**

Each of the 29 nodes consists of an excitatory and an inhibitory population, which summarize the effective dynamics of the area. Populations are described by

$$
\tau_E \frac{d}{dt} \nu_E = -\nu_E + \beta_E [I_E]^+ \\
\tau_I \frac{d}{dt} \nu_I = -\nu_I + \beta_I [I_I]^+. \tag{3}
$$

$\nu_E$ is the firing rate of the excitatory population, with intrinsic time constant $\tau_E$ and input current $I_E$, and for which the f-I curve has slope $\beta_E$. $[I_E]^+ = \max(I_E, 0)$. The inhibitory population has corresponding parameters $\tau_I$, $I_I$ and $\beta_I$. Values for $\tau_E$, $\tau_I$, $\beta_E$ and $\beta_I$ are given below and are taken from Binzegger et al. (2009).

At each node, the input currents have a component from within the area (i.e. local input) and another that comes from other areas:

$$
I_E^i = (1 + \eta h_i)(w_{EE} \nu_E^i + I_{lr,E}^i) - w_{EI} \nu_I^i + I_{ext,E}^i \\
I_I^i = (1 + \eta h_i)(w_{IE} \nu_E^i + I_{lr,I}^i) - w_{II} \nu_I^i + I_{ext,I}^i. \tag{4}
$$

$w_{EE}$ and $w_{EI}$ are couplings to the excitatory population from the local excitatory and inhibitory population respectively, $I_{lr,E}^i$ is the long-range input to the excitatory population, and $I_{ext,E}^i$ is external input (both stimulus input and any noise we add to the system). $w_{IE}$, $w_{II}$, $I_{lr,I}^i$ and $I_{ext,I}^i$ are corresponding parameters for the inhibitory population.

Following Binzegger et al. (2009), we write $w_{ij} = \alpha_j S_{ij}$, where $i$ and $j$ can be E or I. $\alpha_E$ ($\alpha_I$) measures charge introduced per excitatory (inhibitory) spike times transmitter release probability; both are slightly modified from Binzegger et al. (2009). $S_{ij}$ is the number of synapses from cells of type $j$ to cells of type $i$, taken from the counts for layer 2/3 cells in Binzegger et al. (2004). Inhibitory values are weighted averages of basket, double bouquet and chandelier cells, with weights chosen according to their projections to the excitatory population.

We scale the excitatory inputs to an area, both local and long-range, by its position in the hierarchy, $h_i$. $h_i$ is normalized between 0 and 1, and $\eta$ is a scaling parameter that controls the effect of hierarchy. By setting $\eta = 0$ we remove intrinsic differences between areas. Note
that we scale both local and long-range projections with hierarchy, rather than just local projections, in accordance with the observations of Markov et al. (2011), who find that the proportion of local to long-range connections is approximately conserved across areas.

Long-range input is modeled as excitatory current to both excitatory and inhibitory cells:

\[
\begin{align*}
I_{lr,E}^i &= \mu_{EE} \sum_{j=1}^{N} FLN_{ij} \nu_E^j \\
I_{lr,I}^i &= \mu_{IE} \sum_{j=1}^{N} FLN_{ij} \nu_I^j.
\end{align*}
\] (5)

Here \(j\) ranges over all areas. \(I_{lr,E}^i\) and \(I_{lr,I}^i\) are the inputs to the excitatory and inhibitory populations, \(\nu_E^j\) is the firing rate of the excitatory population in area \(j\) and \(FLN_{ij}\) is the FLN from area \(j\) to area \(i\). \(\mu_{EE}\) and \(\mu_{IE}\) are scaling parameters that control the strengths of long-range input to the excitatory and inhibitory populations, respectively, and do not vary between connections; all the specificity comes from the FLNs. Long-range connectivity is thus determined by three parameters: \(\mu_{EE}\) and \(\mu_{IE}\) control the connection strengths of long-range projections, and \(\eta\) maps the hierarchy into excitatory connection strengths.

We can choose the excitatory to inhibitory ratio of an input current, \(\gamma = I_{inp,E}/I_{inp,I}\), such that the steady-state firing rate of the excitatory population does not change when the current is present. Given input of \(I_{inp,E}\) to the excitatory population, an input of \(\gamma I_{inp,E}\) to the inhibitory population increases the inhibitory firing rate sufficiently to cancel out the additional input to the excitatory population. We call such inputs balanced. We choose \(\mu_{EE}\) and \(\mu_{IE}\) with a ratio slightly above this value so that projections are weakly excitatory.

Parameters are \(\tau_E=20\) ms, \(\tau_I=10\) ms, \(\beta_E=0.066\) Hz/pA, \(\beta_I=0.351\) Hz/pA, \(w_{EE} = 24.3\) pA/Hz, \(w_{IE} = 12.2\) pA/Hz, \(w_{EI} = 19.7\) pA/Hz, \(w_{II} = 12.5\) pA/Hz, \(\mu_{EE} = 33.7\) pA/Hz, \(\mu_{IE} = 25.3 \) pA/Hz and \(\eta = 0.68\).

**Network with conduction delays**

In our simulations we ignore conduction delays between areas. While these will be important for oscillations, synchronization and other fine temporal structure, the timescales we consider are typically slow enough that conduction delays do not play an important role.

In Figure S3 we demonstrate that our results hold in a network with realistic conduction delays. We use distances from the same data set as the connectivity strengths (Ercsey-Ravasz et al., 2013) and, to ensure a fair comparison, assume a relatively low conduction velocity of
1.5 m/s (Deco et al., 2009). As shown in Figure S3B, the response of this network to a pulse of input to area V1 is almost identical to that of a network without conduction delays.

Scrambled connectivity

For the simulations shown in Figure 5B, we scramble the connectivity matrix by permuting all entries of the matrix randomly. For Figure 5C, we preserve the absent entries and permute the non-zero entries. Note that the connectivity data show specificity both in terms of which projections exist and in their strengths, and both the probability of a connection and its strength decay exponentially with distance between areas (Markov et al., 2011; 2013; 2014a; Ercsey-Ravasz et al., 2013). In particular, nearby areas tend to be strongly connected and to have similar timescales (see Fig. 2C); thus scrambling projections should reduce the separation of timescales.

We examine the response of these scrambled networks to a pulse of input to all areas, similar to the “resting-state” condition. In the intact network, areas are dominated by a few timescales and are well fit by one or two summed exponentials. However, a number of the scrambled networks show responses that consist of many mixed timescales and are not well described by two exponentials. Thus we use a non-parametric measure of timescale: we compute the time taken after pulse offset for the area’s activity to decay to within 5% of its value at baseline. Scrambling the connection strengths makes about 20% of networks unstable, meaning that responses to input grow instead of decaying, and we exclude these networks. We then compute the median and the 5th, 10th, 90th and 95th percentile of the decay time distribution for each area, and contrast it with values for the intact network.

Functional connectivity for a linear network

If a linear network is driven by white noise input then, away from the threshold, it evolves according to the equation

\[ \dot{x}(t) = Ax(t) + I + B \xi(t), \]  

where \( I \) is the mean of the noise, \( B \) is its covariance matrix and \( A \) is the coupling matrix, which includes any intrinsic leak of activity.

In the steady-state the covariance, \( C \), of this matrix is the solution to the equation (Gardiner, 1985)

\[ AC + CA^\dagger + BB^\dagger = 0 \]  

This equation can be solved given the eigenvector basis (Deco et al., 2013). In the eigenvector decomposition, \( A = V \Lambda V^{-1} \), where \( \Lambda \) is the diagonal matrix of eigenvalues and the columns
of $V$ are the right eigenvectors of $A$. Define
\[
\tilde{Q} = V^{-1}BB^\dagger V^{-\dagger}
\]
(8)
\[
M_{ij} = -\frac{\tilde{Q}_{ij}}{(\lambda_i + \lambda_j^*)}
\]
Then $C = VMV^\dagger$.

As an aid to intuition, assume that $A$ is a normal matrix so that $V^{-1} = V^\dagger$. Then $\tilde{Q} = V^\dagger BB^\dagger V$, and the covariance matrix of the network is a rescaled version of the covariance structure of the input noise.

If, as in the simulations of Figure 8, the input noise is independent and identical at each node, then the covariance matrix of the noise is diagonal with constant entries (and all correlations come from the structure of the network). If this has the value $\sigma^2$ at each node then, for a normal matrix, $\tilde{Q}_{ij} = \sigma^2\delta_{ij}$, and $M$ is diagonal with $i$th entry $\tau^2_i\sigma^2 / 2$, where $\tau_i = -1 / \lambda_i$. Hence the covariance of the $i$th eigenmode is proportional to its corresponding timescale.

Now $C = VMV^\dagger$, meaning that the matrix is rotated out of the eigenvector basis giving a non-diagonal matrix. Thus eigenvectors that are more broadly shared contribute more to the functional connectivity. In this case $C \propto A^{-1}/2$.

We also note that Baria et al. (2013) conduct a similar analysis on a linear network with nodes having identical properties and binary connectivity, and find that nodes with more anatomical connections and, consequently, higher functional connectivity show greater activity at low frequencies (i.e., slower timescales).

**Functional connectivity with hemodynamic response function**

For Figure S6, we convolve the firing rates of the excitatory population at each node with a hemodynamic response function of the form
\[
H(t) = \frac{(t - d)e^{-(t-d)/\tau_h}}{\tau_h^2},
\]
with timescale $\tau_h = 1.25$ s and delay $d = 2.25$ s (Boynton et al., 1996). This yields a simulated BOLD signal, and we calculate the functional connectivity as the correlation matrix of this activity.
Nonlinear network

The single area model is a variant of the model developed in Wong and Wang (2006) as a simplified mean-field version of the spiking network of Wang (2002). There the dynamics were assumed to be dominated by the slow time-constant of NMDA synapses, and the activity of the inhibitory population was incorporated into the effective connection strengths between the excitatory populations. As in that study, we assume that the dynamics of the excitatory population are modeled by a dimensionless gating variable, $s_N$, reflecting the fractional activation of the NMDA conductance, with timescale set by the slow NMDA time-constant. However, we also consider an inhibitory population, modeled with a threshold-linear differential equation (as in the previous sections).

The equation for the excitatory population is

$$\nu^i_E = \phi \left( I^i_E \right) = \phi \left( \left( 1 + \eta h_i \right) \left( w_{EE} s^i_N + I^i_{lr,E} \right) - w_{EI} \nu^i_I + I^i_{ext,E} \right)$$

$$\tau_N \frac{d}{dt} s^i_N = -s^i_N + \gamma \tau_N (1 - s^i_N) \nu^i_E$$  \hspace{1cm} (9)

Here $\nu_E$ is the excitatory firing rate and $s_N$ is the NMDA gating variable, which is bounded between 0 and 1. $\phi$ models the firing rate-current dependence of a leaky integrate-and-fire neuron (Abbott and Chance, 2005) and is defined as

$$\phi(I_{syn}) = \frac{a I_{syn} - b}{1 - \exp(-d(a I_{syn} - b))}$$

with $a = 0.27$ Hz/pA, $b = 108$ Hz and $d = 0.154$ s.

The inhibitory population is described with a threshold-linear equation as before.

$$\tau_I \frac{d}{dt} \nu^i_I = -\nu^i_I + \beta_I \left[ I^i_I \right]_+ = -\nu^i_I + \beta_I \left[ \left( 1 + \eta h_i \right) \left( w_{IE} s^i_N + I^i_{lr,I} \right) - w_{II} \nu^i_I + I^i_{ext,I} \right]_+ .$$

Parameter values are: $\tau_N = 60$ ms, $\tau_I = 10$ ms, $\gamma = 0.641$, $w_{EE} = 250.2$ pA, $w_{EI} = 8.110$ pA/Hz, $w_{IE} = 303.9$ pA and $w_{II} = 12.5$ pA/Hz.

Supplemental References


