Scale-Change Symmetry in the Rules Governing Neural Systems

Vidit Agrawal, Srimoy Chakraborty, Thomas Knöpfel, Woodrow L. Shew

HIGHLIGHTS

We study the dynamical rules governing neuronal networks in models and mouse cortex

Dynamical rules can change when viewed at fine versus coarse spatiotemporal scales

Rules have scale-change symmetry when the network operates near a phase transition

Rules of cortical dynamics approach scale-change symmetry as a mouse awakens

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Scale-Change Symmetry in the Rules Governing Neural Systems

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SUMMARY

Similar universal phenomena can emerge in different complex systems when those systems share a common symmetry in their governing laws. In physical systems operating near a critical phase transition, the governing physical laws obey a fractal symmetry; they are the same whether considered at fine or coarse scales. This scale-change symmetry is responsible for universal critical phenomena found across diverse systems. Experiments suggest that the cerebral cortex can also operate near a critical phase transition. Thus we hypothesize that the laws governing cortical dynamics may obey scale-change symmetry. Here we develop a practical approach to test this hypothesis. We confirm, using two different computational models, that neural dynamical laws exhibit scale-change symmetry near a dynamical phase transition. Moreover, we show that as a mouse awakens from anesthesia, scale-change symmetry emerges. Scale-change symmetry of the rules governing cortical dynamics may explain observations of similar critical phenomena across diverse neural systems.

INTRODUCTION

The ongoing collective population activity of neurons in cerebral cortex exhibits complex spatiotemporal fluctuations. This ongoing activity is responsible for the majority of the brain’s energy consumption (Buzsáki et al., 2007), is closely related to past experiences (Han et al., 2008; Kenet et al., 2003; Luczak et al., 2009), contributes to memory consolidation (Gupta et al., 2010; Ji and Wilson, 2007), and modulates ongoing cortical processing (Arieli et al., 1996; Fox et al., 2007, 2006; Petersen et al., 2003). Thus understanding how ongoing cortical activity is organized is an important goal of systems neuroscience. Comparing ongoing cortical activity across diverse species, measured with different experimental methods over the past decade, a common phenomenon has been found with surprising consistency. The spatiotemporal sizes of ongoing fluctuations follow a specific statistical law; they are distributed according to a power law probability density function with the same exponent near $-1.5$. This phenomenon has been observed in functional magnetic resonance imaging (fMRI) in humans (Haimovici et al., 2013; Tagliazucchi et al., 2012); magnetoencephalography (MEG) in humans (Shriki et al., 2013); voltage imaging in mice (Fagerholm et al., 2016; Scott et al., 2014); multielectrode electrophysiology in monkeys (Petermann et al., 2009; Yu et al., 2017), cats (Hahn et al., 2017), rats (Gautam et al., 2015; Greesh and Plenz, 2008), and even turtles (Clawson et al., 2017; Shew et al., 2015); and in vitro brain-in-a-dish systems (Beggs and Plenz, 2003; Shew et al., 2011). What can explain such shared phenomena observed across such diverse cortical systems, measured with differing spatial and temporal resolution?

One hypothesis is that the cerebral cortex can operate near the critical point of a phase transition (Beggs and Timme, 2012; Hesse and Gross, 2014; Plenz et al., 2014; Shew and Plenz, 2013). This hypothesis builds upon well-established physics of critical phase transitions; at criticality, multiple material properties are expected to be power law distributed according to universal scaling laws that are insensitive to many details of the physical system (Stanley, 1987, 1971; Wilson, 1979, 1975). From theory and computational modeling work, it is clear that neural systems can be tuned into a variety of different dynamical regimes or phases (e.g., asynchronous, oscillatory, bursting), often with distinct boundaries separating different regimes (Brunel, 2000; Gautam et al., 2015; Haldeman and Beggs, 2005; Poil et al., 2012; Wang et al., 2011). Similar to physical systems, near certain regime boundaries these neural models exhibit power-law-distributed collective dynamics with the same power law exponents, despite many detailed differences among models (Gautam et al., 2015; Haldeman and Beggs, 2005; Wang et al., 2011). What does the physics of critical phenomena tell us about where this universality comes from?

Fundamental insight into the origins of universal critical phenomena in physical systems (and a Nobel Prize) came from the realization that the basic laws governing the system obey a peculiar symmetry. The laws are
the same across different scales; they have scale-change symmetry. Critical phenomena, including power-law-distributed observables, stem directly from this bizarre fractal symmetry of the governing physical laws (Wilson, 1979, 1975). Motivated by this basic fact about critical phenomena in physical systems, here we hypothesize that the governing laws for cortical dynamics also conform to scale-change symmetry. Despite more than a decade of intense research on criticality in neural systems, this basic question has not been addressed. How does one go about testing this hypothesis?

Here, we develop an approach inspired by renormalization group theory, but with a focus on practical applicability to real neural data. Renormalization group theory is the mathematical approach used to understand scale-invariance at criticality in equilibrium physical systems (Stanley, 1999; Stanley and Wong, 1972; Wilson, 1979, 1975), and in certain non-equilibrium dynamical systems (Loreto et al., 1995; Tauber, 2014; Vespignani et al., 1997, 1995). However, renormalization group ideas have not been developed in the context of neural systems. In brief, the idea begins with a coarse-graining procedure that transforms all the system variables and governing laws at one spatiotemporal scale to a new set of variables and laws at a coarser scale. If the system obeys the scale-change symmetry, then an appropriately chosen coarse-graining procedure will leave the governing laws unchanged and leave the system variables with identical statistics.

Here, we first applied our approach to two computational models, one simple and the other more biologically realistic. For both models, we found that dynamical rules were indeed maximally scale-invariant when operating near a phase transition. Next, we applied our approach to experimental data. We found that, in the awake state, the apparent rules governing dynamics of mouse cerebral cortex were more scale-invariant than in the anesthetized state. This suggests that the unconscious cortex deviates further from criticality than does the conscious cortex.

RESULTS
Simple Model
We developed our approach using a network of binary, probabilistic, excitable nodes similar to that used in previous studies of non-equilibrium critical phenomena in neural systems (Haldeman and Beggs, 2005; Kinouchi and Copelli, 2006; Larremore et al., 2011; Shew et al., 2009), but on a two-dimensional \( L \times L \) square lattice (\( L = 400 \)) with nearest-neighbor connections (including self). We interpret each node in the network as the aggregate state of a group of neurons, analogous to the signals measured using experimental techniques with somewhat coarse spatial resolution (e.g., local field potential, Petermann et al., 2009; voltage imaging, Scott et al., 2014; or fMRI, Haimovici et al., 2013, Tagliazucchi et al., 2012). The state \( X(t+1) \) of a node at time \( t+1 \) is either active (1) or inactive (0) depending on the number of neighbors \( n(t) \) that were active at time \( t \).

\[
X(t+1) = \begin{cases} 
1 \text{ with probability } \phi_n = 1 - (1 - C)^{n(t)} (1 - p) \\
0 \text{ otherwise } 
\end{cases} \tag{Equation 1}
\]

and \( C \) defines the strength of coupling between all connected pairs of nodes and \( p \) defines the probability of activation without any active neighbors (\( p = 0.001 \) unless stated otherwise). We interpret \( p \) as an external source of input to the neural system. The states of all the network nodes are updated synchronously. One way to characterize the collective network dynamics is by measuring the fraction of active nodes at a time, or network activity, given by \( S(t) = L^{-2} \sum_{i=1}^{L^2} X_i(t) \). By tuning the coupling strength, we tuned the model through a critical phase transition at \( C = C^* \), separating a low-activity (small \( S \)) from a high-activity (large \( S \)) regime (Figure 1). Most of our results consider the range of \( C \) from 0.15 to 0.35.

Our first goal was to examine how the dynamical rules governing the system change when considered at different scales of observation. For this, we consider six activation probabilities \( \phi_n = 1 - (1 - C)^{n(t)} (1 - p) \) with \( n = 0, 1, 2, 3, 4, \text{ or } 5 \) (\( n = 5 \) means all 4 neighbors plus self). Together, these six probabilities completely specify the rules of the system dynamics (actually they over-specify the rules, which are completely specified by \( C \) and \( p \), but over-specification can be helpful with noisy data). Importantly, we can estimate these probabilities directly from data, whether simulated or experimentally measured. For instance, \( \phi_0 \) is estimated as the fraction of instances with \( n(t) = 1 \) that lead to activation at time \( t+1 \). Based on \( 10^4 \) time steps of data from the model (excluding the transient of duration \( 10^2 \) time steps), we can estimate all six \( \phi_n \) probabilities with precision \( \leq 2\% \) for \( 0.15 < C < 0.35 \) (except \( \phi_0 \), which was less accurate for \( C > 0.3 \), see Figure S1).
The scale-change symmetry at the heart of critical phenomena in physical systems suggests that the Ornstein-Zernike integral could be invariant across observational scales when $C$ equals its critical value $C^*$, but not if $C$ deviates from $C^*$. The next step toward testing this possibility was to identify an appropriate coarse-graining renormalization transformation to map one observational scale onto a coarser scale.

Coarse-Graining Scheme
Motivated by the limited spatiotemporal resolution of typical experimental data and real-space block renormalization schemes originally used to gain insight into the critical phenomena in physical systems (Kadanoff, 1993; Stanley and Wong, 1972; Wilson, 1979), we devised a spatiotemporal transformation scheme described as follows. Each $r \times r \times r$ spatiotemporal block of nodes ($r \times r$ in space and of duration $t$) at scale $b$ transforms into a single node at coarser scale $b + 1$. Thus, an $L \times L \times T$ lattice at observational scale $b$ transforms into a coarse $L/r \times L/r \times T/r$ lattice at observational length scale $b + 1$, as illustrated for $r = 4$ and $t = 1$ in Figures 2A and 2B. The state $X^b_{b+1}$ of a coarse node depends on the average state $S^b = \sum_{i} X^b_{i}$ of the corresponding $r^3$ nodes at the finer scale according to

$$X^{b+1}_{b+1} = \begin{cases} 1 & \text{with probability } f(S^b) = [1 + \exp(-k(S^b - x_0))]^{-1}, \\ 0 & \text{otherwise.} \end{cases}$$

(Equation 2)

By tuning the transformation parameters $k$ and $x_0$, we can explore a family of logistic function coarse-graining procedures including majority rules and other previously studied functions. Similar coarse-graining schemes have been used in previous work on deep learning (Hinton and Salakhutdinov, 2006).

To quantify the change in the dynamical rules due to coarse graining, we define a parameter $\zeta$

$$\zeta = \sum_{n=0}^{S} |\phi^b_n - \phi^{b+1}_n|$$

(Equation 3)

where $\phi^b_n$ is the activation probability estimate at scale $b$ and $\phi^{b+1}_n$ is the activation probability estimate at the coarser scale $b + 1$. Note that root-mean-square differences or any other monotonic function of the differences of $\phi^b_n$ and $\phi^{b+1}_n$ would not change our following conclusions. We ran the model for a range of coupling strengths $0.15 < C < 0.35$, simulating $10^6$ time steps for each $C$. In line with expectations from critical phenomena in physical systems, we found that $\zeta$ was minimized for $C$ near $C^* = 0.23$ (Figure 2C). This result was obtained using transformation parameters $k = 76$, $x_0 = 0.22$, and block size $r = 8 \times 8 \times 1$ (Figure 2C, inset). This means that the dynamical rules governing the network activity are least changed at $C^*$.

However, some caution is appropriate; if a different transformation function $f$ was used, perhaps the minimum $\zeta$ would occur at a different $C$. Therefore, to draw a more definitive conclusion, we next systematically searched the two-dimensional space of all possible $f$ functions, seeking the function that minimizes $\zeta$ for
each C independently. We found that, when using such optimal f functions, the dynamical rules remained most scale-invariant (i.e., lowest $z$) for C near C* (Figures 2D and 2E). Thus our first conclusion is that scale-invariance of the governing laws at criticality, one of the most fundamental concepts of critical phenomena in physical systems, can also manifest in neural systems.

We also verified our results for coarse-graining blocks with different spatial sizes $r = 4, 8, 16$ (Figures 2D and 2E), different durations $\tau = 1, 2, 4, 8$ (Figures 2D and 2E), and different levels of noise $p = 10^{-2}, 10^{-3}, 10^{-4}$ (Figure 2F). Optimal $x_0$ increased with C and r; optimal k increased with r and was lowest near C = C* (Figure 2D). Changes in r, $\tau$, and p did not qualitatively change our conclusion that coarse graining causes the least change in the dynamical rules near criticality. However, increasing p, decreasing r, and increasing $\tau$ all had the effect of broadening the minimum in $\zeta_{min}$ and shifting the minimum toward slightly smaller values of C. For increasing p, this shift in the minimal C may be due to the phase transition becoming “smeared out,” less sharp with increased noise (Williams-Garcia et al., 2014), but further investigation is needed to test this possibility. We chose the range 0.15 < C < 0.35 because at lower values (C < 0.15) the rates of activity became so small that our estimates of the $\phi_0$ became poor due to subsampling, which made $\zeta_{min}$ an unreliable measure (see Figure S2). A skeptical reader may note that $\zeta$ is not zero at criticality for our model; the scale-invariance we observe is imperfect. Why might this be? In the calculation of $\zeta$, we found that the largest contribution to $\zeta$ came from the $\phi_0$ term in the sum, which represents spontaneous activation of a node. This is because the coarse-graining procedure tends to cause periods with very low S at scale b to become S = 0 at scale b+1, thus creating excessive apparent spontaneous activation at scale b+1. When $\zeta$ is calculated by excluding $\phi_0$, it still minimizes for C near C* but with a much lower value of $\zeta$, indicating higher degree of scale-invariance for the dynamical rules governing interactions, i.e., those with n > 0 (see Figure S3).

**Realistic Model**

The abstract binary model we have discussed so far is simple to interpret and directly comparable to previous work on critical phenomena in neural systems using similar models. Moreover, it is relatively similar to models of spreading dynamics in previous renormalization group studies (e.g., sand piles, Vespignani et al., 1995, and forest fires, Loreto et al., 1995). In this sense, it is not terribly surprising that our simple
model conform to scale-change symmetry of governing rules, which is expected from theory of critical phenomena. However, our simple binary model is not very biologically realistic. For example, the dynamics of real cerebral cortex never exhibits a sustained high-firing regime as we see in the simple model with $C > C^*$. In reality, if the excitability of real cortex is enhanced (or other parameters analogous to $C$ are enhanced) the dynamics tends to exhibit large, repetitive bursts of activity that are suppressed by depressor adaptive mechanisms before reaching a sustained high-firing regime (Gautam et al., 2015; Shew et al., 2009). Indeed, activity-dependent adaptive effects can act to make the critical regime more robust (Levina et al., 2007; Shew et al., 2015), which could make the possibility of scale-invariant dynamical rules even more plausible in real brains. Next, we set out to test our ideas in a more realistic model of a neural network that generates more realistic network activity. Similar to other recent model studies (di Santo et al., 2018; Gautam et al., 2015), the phase transition we examine in our more realistic model separates an asynchronous phase from an oscillatory ordered phase, which better matches experimental observations.

Building on our simple model, we kept binary, probabilistic, integrate-and-ﬁre neurons on a two-dimensional square lattice ($L \times L = 160 \times 160$), but now we introduced inhibitory neurons (20% of all neurons), spike-frequency adaptation, refractoriness, and different distance-dependent connectivity for excitatory and inhibitory neurons. At each time $t$, the binary state $X(t) = 1$ with probability $p_i(t)$, where

$$
p_i(t) = \begin{cases} 
1 & \text{for } \chi_i(t)h_i(t)^{-1} \geq 1 \\
\chi_i(t)h_i(t)^{-1} & \text{for } 0 \leq \chi_i(t)h_i(t)^{-1} < 1 \\
0 & \text{for } \chi_i(t)h_i(t)^{-1} < 0
\end{cases}
$$

(Equation 4)

Input from other neurons is $\chi_i(t) = \sum_{j \neq i} W_{ij}X(t-1)$ and the activity-dependent adaptation is modeled by $h_i(t) = \sum_{\tau=1,\tau \neq 0} X(t')$ where $\tau = 80$. If this sum is zero, we set $h$ to 1. At each time step a neuron can also be activated due to external sources with probability $p_{ext} = 0.001$ (treated independently of $p$). After a neuron fires, its state is set to 0 for a refractory period of 1 time step. The default synaptic weight matrix $W$ is constructed with long-range inhibition relative to shorter-range excitation. Weights of inhibitory synapses were $W_i(d) = -I + \exp \left( -\frac{d^2}{2} \right)^2$ and those of excitatory synapses were $W_{ii}(d) = -E + \exp \left( -\frac{d^2}{2} \right)^2$,

where $d$ is the distance from the presynaptic neuron to the postsynaptic neuron. We consider $C_1 = 3$ and $C_2 = 2$ (in units of lattice spacing) and fixed excitatory input strength $E = 1$. By tuning the inhibitory input strength $I$, we tuned the model through a phase transition near $I = 0.65$ separating a synchronous oscillatory firing state (high mean pairwise correlation) from asynchronous low-firing (low mean pairwise correlation) regime (Figures 3A–3C).

Like the simple model, we found that $\zeta_{min}$ is minimized near the phase transition (Figure 3D). We emphasize that this more realistic model is outside the bounds of well-understood critical phenomena in physics. Although it shares some features with simple models of spreading dynamics (e.g., excitable nodes with refractory periods), the presence of inhibition and nonlocal connectivity makes it quite different from these previously studied models. Thus there is no guarantee that the phase transition we consider here has anything to do with criticality. In this sense, it is substantially more surprising that we observe minimal scale change of dynamical rules near this phase transition. For the results in Figure 3D, we only considered $\phi_1$ with $n = 1, 2, 3, 4$ for $\zeta$ calculation, because $\phi_1$ tended to be most prone to error as discussed above (Figure S3). We did not include $\phi_5$ because the refractory period precluded the occurrence of $\phi_5$ at scale $b$.

So far, we have applied our approach to spike data from our two models. However, many experimentally measured signals are continuous and more closely related to membrane potential, particularly at the larger spatial scales (e.g., local field potential (LFP), fMRI, electroencephalography, voltage imaging, wide-field calcium imaging). How might we apply our approach to such continuous signals? To address this question, here we first used our more realistic model. We generated a continuous signal from the model, using the “membrane potential” from the model neurons. We defined the membrane potential of the $i^{th}$ neuron to be $\chi_i(t)h_i(t)^{-1}$, as defined above. Next, we binarized the continuous membrane potential signal. Following a previously established approach (Scott et al., 2014; Tagliazucchi et al., 2012), we defined time points of activation when the voltage imaging time course crossed above a threshold of 0.5 SD beyond the mean from below (results consistent for different thresholds 0.25 SD and 1 SD) (Figure 4). We found that, like the spike data presented above, this binarized continuous signal also supported our main claim: minimal $\zeta_{min}$ near the phase transition.
One advantage of our approach is that it can readily be applied to experimental data, provided the data have sufficient spatial resolution and coverage. Here, we demonstrate this for measurements of cerebral cortex in a mouse awakening from anesthesia. Previous work (Scott et al., 2014) with these data suggested that as the mouse awakens the cortical dynamics transitioned from a supercritical regime, similar to the oscillatory regime in our realistic model, toward criticality. The brain activity was measured over the dorsal surface of nearly an entire hemisphere of mouse cortex with high spatial resolution using genetically encoded voltage-sensitive fluorescence imaging. Each dataset was acquired during 1 min with 50 Hz sample rate and 33 \( \times \) 33\( \mu \text{m}^2 \) per pixel spatial resolution. Multiple such 1-min recordings were performed over a period of 200 min as the mouse awoke. Each pixel represents the aggregate activity of many neurons within cortical layers 2 and 3. This voltage imaging signal is a continuous signal. To apply our approach, we followed the same binarization approach described above for the membrane potential analysis of the realistic model (Figure 4). We converted each pixel voltage time series into binary form (Figure 5A).

For each pixel and each 1-min recording, time points of activation were defined at times when the voltage imaging time course crossed above a threshold of 0.5 SD beyond the mean from below (results consistent for different thresholds 0.25 SD and 1 SD).

Under the effect of anesthesia, we found that the mouse exhibited synchronous burst firing, similar to the oscillatory regime of our realistic model. As the mouse woke up, more asynchronous firing was observed (Figure 5B), similar to the activity near the onset of the oscillatory regime of our realistic model. This change manifested as a decrease in pairwise correlations as the mouse awoke from anesthesia (Figure 5C). Next, we assumed nearest-neighbor interactions among pixels and proceeded to estimate the activation probabilities \( \phi^b_n \). Then, we applied our course-graining procedure and estimated \( \phi^{b+1}_n \). As with our analysis of the realistic model, only
four activation probabilities, $n = 1, 2, 3, 4$ were used to estimate $z_{\text{min}}$. We found that $z_{\text{min}}$ decreased as the mouse awoke from anesthesia. This finding demonstrates that the rules governing cerebral cortex dynamics approach scale-change symmetry during the transition from anesthetized to awake.

We note that, for higher values of $t$, there was increased noise in the trend relating $z_{\text{min}}$ to time since drug delivery. As $t$ increases, we are left with fewer time points with which we estimate the rules $\phi_n$ and calculate $z_{\text{min}}$. This decrease in samples may be responsible for poorer estimates of the rules $\phi_n$ and a noisier $z_{\text{min}}$ versus time trend. Future studies with longer-duration experimental recordings could better test this possibility.

**Relating Rules to Dynamics**

The approach we have developed here offers a new way to learn about whether a system is operating near criticality by examining how the rules governing dynamics change with scale. In contrast, the traditional way to assess whether a system is near criticality is by examining the dynamics, rather than the rules. For instance, the experimental data we analyze here (Figure 5) has previously been shown to exhibit dynamics that are consistent with an approach to criticality as the mouse wakes up (Fagerholm et al., 2016; Scott et al., 2014). This claim was largely based on examining cascades of propagating neural activity, often called neuronal avalanches. At criticality, theory predicts that different sizes of avalanches occur with probability that is related to size according to a power law with exponent near $-1.5$. In contrast, in the ordered phase (like the oscillatory phase in our realistic model) avalanche sizes are expected to deviate from a power law distribution, with very large cascades becoming more prominent. When Scott et al. analyzed our experimental data, they found that the anesthetized state was consistent with an ordered phase, whereas the awake state was more consistent with criticality. This finding is consistent with our conclusion that scale-change symmetry increases as the mouse wakes up. However, we emphasize that a power law distribution of avalanche sizes is certainly not equivalent to our finding of scale-change symmetry of rules. Indeed, power law distributions of observed dynamics can arise due to mechanisms that are totally unrelated to criticality (Beggs and Timme, 2012; Mitzenmacher, 2003; Reed and Hughes, 2002; Sornette, 1998; Stumpf and Porter, 2012). If criticality is responsible for a power law distribution of an observable, then we would expect to also find scale-change symmetry of rules. To clarify how scale-change symmetry of rules is related to power law avalanche size distributions, we studied our models further. We examined avalanche size distributions for the simple model as $C$ was tuned through $C^*$, and for the more realistic model as inhibition ($I$) was tuned from strong ($I = 2$) to weak ($I = 0.01$). We used the previously developed measure $\kappa$ to quantify deviation from a power law (Scott et al., 2014; Shew et al., 2009). In brief, $\kappa = 1$ for a perfect match to the reference power law with exponent $-1.5$, $\kappa > 1$ when large avalanches become prominent (as expected in the ordered phase), and $\kappa < 1$ when large avalanches become rare (as expected in the asynchronous phase). Further description of $\kappa$ is given in the Methods section.

As expected for the simple model, we found that the distribution of avalanche sizes was close to a power law ($\kappa$ near 1) when $C$ was tuned near $C^*$ (Figures 6A and 6B). Thus, in this case, $z_{\text{min}}$ was also smallest near $\kappa = 1$ (Figure 6C). Comparing $\kappa$ and $z_{\text{min}}$, we found that $\kappa$ is more sensitive to changes in $C$ for $C > C^*$,
whereas $z_{\text{min}}$ is more sensitive for $C < C^*$. For the more complex model, the oscillatory regime (low inhibition) showed a clear bump in the tail of the avalanche distributions similar to previous studies of dynamics in the ordered (supercritical) phase (Figure 6D). As inhibition was tuned from weak to strong, the avalanche size distributions did approach approximate, but rather imperfect power laws. The changes in avalanche distributions we observed for increasing inhibition is similar to what has been shown previously for our experimental data as the mouse wakes up (Scott et al., 2014). Despite the imperfect power laws found in the realistic model, we computed $k$ and found a clear tendency for minimal $z_{\text{min}}$ near $\kappa = 1$ (Figures 6E and 6F). These results demonstrate a close relationship between avalanche statistics and the scale-change symmetry of rules governing a neural system.

**DISCUSSION**

Here we have shown that in neural network models, the rules that govern the system behavior obtain a degree of scale-change symmetry that is most pronounced when the system operates near criticality. In the spirit of renormalization group theory in physical systems, our approach offers a potential explanation of why diverse neural systems can exhibit very similar critical phenomena.

Another useful outcome of our work is that the tools we develop provide a way to assess whether a change in system dynamics takes the system closer to or further from criticality. Our approach complements traditional approaches for seeking evidence for criticality based on power laws and scaling relations. Such power law distributions constitute necessary, but insufficient, evidence for the criticality hypothesis (Beggs and Timme, 2012; Klaus et al., 2011; Stumpf and Porter, 2012). Our results suggest that a change toward criticality should be accompanied by an increase in scale-invariance of effective dynamical rules. To be
more specific, considering our model work (both models, Figures 2, 3, and 4), it is clear that, for a wide range of parameters (C or I) around criticality, a decrease in $\zeta$ does not happen unless the parameters have been tuned toward criticality—tuning C toward $C^*$ (Figure 2) or tuning I toward the value that corresponds to the onset of oscillations (Figures 3 and 4). However, for a given observed decrease in $\zeta_{\text{min}}$, we cannot make precise conclusions about how much closer to criticality the system has shifted. This is not possible considering the quantitative differences in the shape of the $\zeta_{\text{min}}$ versus C and $\zeta_{\text{min}}$ versus I curves and how they depend on the details of analysis parameters (e.g., block size). The stronger conclusions that can be made are about (1) the direction of change and (2) about the relative size of changes within one system. For example, our methods could be used to ascertain which kinds of pharmacological manipulations result in a bigger shift away or toward criticality in a single system with consistent measurement tools. This is an important value of our approach—measuring relative changes in proximity to criticality.

**Limitations of Study**

One limitation of our approach is that it currently requires binary data (Figures 1, 2, and 3) or binarization of continuous data (Figures 4 and 5). Although spike data are well suited to a binary approach, it would be useful for future work to generalize the approach to continuous data, which is more common in experiments. However, we emphasize that our results based on binarization of continuous data from our realistic model (Figure 4) were qualitatively consistent with our results based on spikes. This suggests that the binarization of continuous data provides a useful strategy for studying scale-change symmetry of dynamical rules.

Another limitation of our approach is that we have based our assessment of scale-change symmetry $\zeta_{\text{min}}$ on one step of renormalization. Renormalization group theory, in contrast, emphasizes that many steps of renormalization may be required before true scale-invariance is found. Each step of renormalization is thought to remove irrelevant details that could corrupt scale-invariance. This fact may contribute to why our scale change measure $\zeta_{\text{min}}$ is not very close to zero even when our simple model is at criticality. However, the fact that we do find a prominent minimum in $\zeta_{\text{min}}$ near criticality indicates that we can detect the scale-change symmetry in spite of only performing one step of renormalization. This is important, because, when working with real data from finite systems, it is not plausible to do many steps of coarse graining especially with a large block size.
We also note that there are some trivial types of scale-change symmetry that we are not interested in and have not highlighted in our results above. For instance, for a completely saturated system—all neurons firing at every time step—the dynamics and rules would be identical upon coarse graining. The coarse level would still be completely saturated. Likewise, a completely silent network would also exhibit such trivial scale-invariance. These trivial types of invariance are not the subject of our work, but should be noted for completeness.

Our initial application of our approach to experimental data provided interesting results. We showed that scale-invariance of the rules governing cortical network dynamics increased during the transition from unconsciousness to consciousness. This finding is in line with other recent work that suggests that the anesthetized cortex deviates from critical dynamics and approaches criticality as it wakes up (Bellay et al., 2015; Fagerholm et al., 2016; Scott et al., 2014). A recent coarse-graining study of experimental data obtained in hippocampus also revealed interesting scaling, in line with the scale-change symmetry we report here (Meshulam et al., 2018). Our finding raises interesting questions about the functional consequences of scale-invariant rules. What does it mean that local interactions, say among cortical columns, are governed by the same rules as larger-scale interactions, say among cortical regions? We expect that our approach will be useful for future studies of these questions and how the dynamical rules governing brain dynamics differ across scales and brain states.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information includes Transparent Methods and three figures and can be found with this article online at https://doi.org/10.1016/j.isci.2019.01.009.

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AUTHOR CONTRIBUTIONS
W.L.S. and V.A. conceived the study. V.A., W.L.S., and S.C. performed the simulations. T.K. performed the experiments. V.A. analyzed the experimental data. W.L.S. and V.A. wrote the paper.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental Information

Scale-Change Symmetry in the Rules

Governing Neural Systems

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Figure S1. Accuracy of estimating rules $\phi$. (Related to Figure 2 in main text.) To test the accuracy of estimation of $\phi_n$ we took network activity data for $10^4$ timesteps after $10^2$ transient timesteps at coupling strength $0.15 < C < 0.35$. We compared the activation probability estimated from the simulation $\phi_{n}^{\text{sim}}$ to analytically calculated $\phi_{n}^{\text{th}} = 1 - (1 - C)^n (1 - p)$ at each $C$ value. Our accuracy is better than 2% for all $\phi_n$ in the range $0.15 < C < 0.35$ except $\phi_0$ where the accuracy is around 2% for $C > 0.3$. All calculations done for network size, $L \times L = 400 \times 400$ and $p = 0.001$. 
Figure S2. Dependency of rule-change $\zeta_{min}$ on simulation timesteps. (Related to Figure 2 in the main text.) Shown are $\zeta_{min}$ values calculated using data from simulations with varying numbers of timesteps at coupling strengths, $C = 0.1$ (yellow), $C = 0.15$ (green), $C = 0.17$ (cyan). When the network activity is too low, the estimation of a consistent $\zeta_{min}$ require data from increased simulation timesteps. All calculations done for network size, $L \times L = 400 \times 400$, $r = 8$ and $p = 0.001$. 
Figure S3. Alternative estimation of $\zeta_{min}$ by excluding $\phi_0$. (Related to Figures 2-5 in the main text.) Shown is the change in dynamical rules $\zeta_{min}$ based on excluding $\phi_0$. $r = 4$ (blue); $r = 8$ (green); $r = 16$ (red). All calculations done for network size, $L \times L = 400 \times 400$, $r = 8$ and $p = 0.001$. A major contribution to $\zeta_{min}$ comes from the difference in $\phi_0$ when the network activity is transformed from observational length scale $b$ to $b + 1$. The main reason behind this are the cases when the transformation scheme probabilistically transforms a partially active block of nodes to an inactive node at one timestep and at next timestep partially active block of nodes do changes only slightly but is transformed to an active node. Even with the 5 activation probabilities $\phi_n$ for $n = 1, 2, 3, 4, 5$, $\zeta_{min}$ still minimize at $C$ near critical point $C^*$.
Transparent methods

Experiments

Animal experiments were performed in accordance with the National Institutes of Health guidelines for animal research and were approved by the Institutional Animal Care and Use Committees of the RIKEN Wako Research Center (Japan). The following methods for obtaining the experimental data in Fig. 5 have been reported in previous publications as described previously (Akemann et al., 2012, 2010; Scott et al., 2014). We describe them again here. In utero electroporation was performed on the mouse (day E15) with the pCAG-VSFP Butterfly 1.2 plasmid. This resulted in expression of the voltage indicator Butterfly 1.2 in pyramidal cells in cortical layers 2/3 of one hemisphere. For imaging, the skull was thinned and a head-post implanted 2-6 months following electroporation. Three days following head-post implantation the imaging measurements reported here were performed. The mouse was first anesthetized (pentobarbital 0.9 g/kg i.p.), then head-fixed, and imaged with a dual emission wide-field epifluorescence microscope (halogen excitation). The voltage imaging signal was the ratio of mKate2 to mCitrine fluorescence, taken after offset subtraction and equalization of heartbeat-related modulation of fluorescence. Between consecutive 1 min imaging periods were 1 min pauses. Preprocessing of voltage signals included baseline normalization by the average over the duration of each recording, spatial and temporal smoothing, and high-pass filtering at 0.5 Hz.

Avalanche size distributions and κ calculation

The following methods for defining avalanches and computing κ (Figure 6) have been reported in previous publications (Fagerholm et al., 2016; Scott et al., 2014; Tagliazucchi et al., 2012). First, avalanches were defined as spatiotemporally contiguous clusters of active pixels. Clusters of active pixels were identified in each frame, based on the detection of connected pixels in a coactive first neighbors graph. Avalanches were then defined as starting with the activation of a
previously inactive cluster, continuing while 1 contiguous cluster was active in the next time point. Avalanche size probability density distributions (Figure 6a, d) were calculated by counting the number of avalanches in each size bin and normalizing by the total number of avalanches and bin size. Avalanche size probability distributions were compared with a power law with exponent -1.5 using a measure called $\kappa$, first developed in our previous work (Shew et al., 2009). First, the measured distribution is recast as a cumulative density function (CDF), called $F(\beta)$, which specifies the fraction of measured cluster sizes $s < \beta$. Then a reference CDF is created corresponding to a perfect -1.5 power law, called $F^{NA}(\beta)$

$$F^{NA}(\beta) = \left(1 - \sqrt{l/L}\right)^{-1} \left(1 - \sqrt{l/\beta}\right)$$

for $l < s < L$, where $l$ is the smallest avalanche size considered and $L$ is the largest. A nonparametric measure, $\kappa$, is defined to quantify the difference between the measured avalanche size CDF, $F(\beta)$, and the theoretical reference CDF, $F^{NA}(\beta)$, as

$$\kappa = 1 + \frac{1}{m} \sum_{k=1}^{m} (F^{NA}(\beta_k) - F(\beta_k))$$

where $\beta_k$ are $m = 10$ avalanche sizes logarithmically spaced between $l$ and $L$. When computing $\kappa$, avalanches below a minimum size, $x_{min}$, were excluded. The rationale for this is that some measurement noise is inevitable and likely to be uncorrelated across pixels, resulting in some small “noise cascades.”

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