Tricritical behavior in a neural model with excitatory and inhibitory units

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While the support for the relevance of critical dynamics to brain function is increasing, there is much less agreement on the exact nature of the advocated critical point. Thus, a considerable number of theoretical efforts are currently concentrated on which mechanisms and what type(s) of transition can be exhibited by neuronal network models. In that direction, the present work describes the effect of incorporating a fraction of inhibitory neurons on the collective dynamics. As we show, this results in the appearance of a tricritical point for highly connected networks and a nonzero fraction of inhibitory neurons.

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I. INTRODUCTION

A large repertoire of diverse spatiotemporal activity patterns in the brain is the basis for adaptive behavior. Understanding the manner in which the brain is able to form and reconfigure a large range of cortical configurations, in a flexible manner, remains an unsolved challenge. A leading proposal interprets this large repertoire as the expected, generic, large diversity of states near instabilities, which is composed, by its own nature, of a mixture of ordered and disordered patterns. In more technical terms, near the critical point of a second-order phase transition it is known that the system exhibits the largest number of metastable states which is only limited by the system size. In the brain, these metastable states would correspond to cortical configurations, which is only limited by the system size. The paper is organized as follows: In Sec. II we describe the model as well as the simulation and the finite-size scaling methods used. The results are presented in Sec. III, and the relevance of the main findings is discussed in Sec. IV.

II. MODEL AND METHODS

A. The model

In this work we use a generalization of the neural model presented in Ref. [9] in which a fraction $f$ of neurons are inhibitory. To this end, we associate a variable $\epsilon_i = \pm 1$ to each neuron $i$, where $\epsilon = -1$ represents an inhibitory neuron and $\epsilon = +1$ an excitatory one. The value of each of the variables $\{\epsilon_i\}$ is chosen independently with probability $f$ to be $\epsilon_i = -1$ and $1 - f$ to be $\epsilon_i = +1$. Those values are kept fixed during the network evolution. The model runs over a small-world network with a weighted adjacency matrix $w_{ij}$. The network topology is obtained following the usual Watts-Strogatz recipe [10]. That is, we start from a ring of $N$ nodes in which each node is connected symmetrically to its $2m$ nearest neighbors. Then, for each node each vertex connected to a clockwise neighbor is rewired to a random node with a probability $\pi$ and preserved with probability $1 - \pi$, so the average degree $\langle k \rangle = 2m$ is preserved [11]. This algorithm provides a nonweighted symmetric adjacency matrix $A_{ij} = A_{ji} = 0, 1$. Then, the weighted adjacency matrix $w_{ij} = w_{ij}$ is obtained by

with excitatory interactions running on a Watts-Strogatz topology. By investigating the effects of adding inhibitory interactions we uncover the presence of a tricritical point for a nonzero fraction of inhibitory neurons, in the regime of high connectivity. The paper is organized as follows: In Sec. II we describe the model as well as the simulation and the finite-size scaling methods used. The results are presented in Sec. III, and the relevance of the main findings is discussed in Sec. IV.
assigning to every non-null link $A_{ij} \neq 0$ a random real value chosen from an exponential distribution $p(w) = \lambda e^{-\lambda w}$ with $\lambda = 12.5$. This procedure mimics the weights distribution of the human connectome [5,12].

The node dynamics of the neural model responds to the Greenberg-Hastings cellular automaton [13], in which each node $i$ of the network has associated a three state dynamical variable $x_i = 0, 1, 2$, corresponding to the following dynamical states: quiescent ($x_i = 0$), excited ($x_i = 1$), and refractory ($x_i = 2$). The transition rules are the following: if a node at the discrete time $t$ is in the quiescent state $x_i(t) = 0$ it can make a transition to the excited state $x_i(t + 1) = 1$ with a small probability $r_1$ or if $\sum_j w_{ij} x_j(t) > T$, where $T$ is a threshold and $\delta(x,y)$ is a Kronecker delta function; otherwise, $x_i(t + 1) = 0$. If it is excited $x_i(t) = 1$, then it becomes refractory $x_i(t + 1) = 2$ always. If it is refractory $x_i(t) = 2$, then it becomes quiescent $x_i(t + 1) = 0$ with probability $r_2$ and remains refractory $x_i(t + 1) = 2$ with probability $1 - r_2$. Following Refs. [5,9] we set $r_1 = 10^{-3}$ and $r_2 = 0.3$.

**B. Analysis of the dynamical transition**

We focus on dynamical clusters of coherent activity, namely, groups of simultaneously activated nodes ($x_i = 1$) which are linked through nonzero weights $w_{ij}$. It is known that for $f = 0$ the system presents a dynamical phase transition separating a regime where the active clusters are isolated from one where such clusters span across the whole system. The transition can be continuous or discontinuous depending on the values of the topological parameters $\langle k \rangle$ and $\pi$ [9,14]. As we will show in the next section, varying $f$ can also change the transition from continuous to discontinuous at fixed topological parameters. Here we explain the metrics used to characterize the transition.

We simulate the model at several values of $\pi$, $\langle k \rangle$, and $f$, and different network sizes $N$. Each simulation is started from a random distribution of activated sites, and the system is let to run 500 time steps before starting data collection. We found this time interval to be enough for the system to reach a stationary state for any system size and for any value of the network parameters. We compute several observables to describe a percolation-like transition as a function of $T$ and $f$. Specifically, we calculate the average size of the largest (i.e., giant) cluster, $\langle S_1 \rangle$. For very large systems, this quantity provides the standard percolation order parameter $P_\infty = \lim_{N \rightarrow \infty} \langle S_1 \rangle / N$, namely, the probability of an arbitrary node to belong to the infinite percolating cluster. We also compute the average size of the second largest cluster $\langle S_2 \rangle$, together with the average cluster size (or susceptibility),

$$\langle s \rangle = \frac{\sum_{\text{size}} s^2 N_s}{\sum_{\text{size}} s N_s},$$

where the primed sum runs over all cluster sizes except the giant one and $N_s$ is the number of clusters of size $s$ [11,15].

We find that on varying the control parameter ($T$ or $f$), $\langle S_1 \rangle$ can change from zero to finite both continuously or discontinuously.

When the transition is continuous, we analyze it as in standard percolation. In this case both $\langle s \rangle$ and $\langle S_2 \rangle$ are expected to exhibit (size-dependent) maxima for a certain pseudocritical value of the control parameter (the threshold $T$ or the fraction $f$), that scales with system size as $[16] \langle s \rangle \sim N^{\nu / d}$, $S_2 \sim N^{\eta / d}$. Here $\nu$ and $\eta$ are the standard susceptibility and correlation length critical exponents, $d$ is the effective dimension of the system, and $d_f$ the fractal dimension of the percolating cluster.

To characterize transition in the discontinuous case we used two different methods.

(a) **Order parameter hysteresis analysis** [14]. For fixed values of $N$, $\langle k \rangle$, $\pi$, and $f$, we keep track of $S_1$ as $T$ is slowly increased at a fixed rate from some initial value $T_0$ up to some maximum value $T_F$, and then decreased again down to $T_0$ at the same rate, without resetting the neuron states when changing $T$. We set the rate of change of the control parameter by changing $T \rightarrow T + \Delta T$ every $t_1$ steps. The values of $T_0$ and $T_F$ were chosen such that the location of the maxima of $\langle s \rangle$ and $S_2$ fall inside the interval $[T_0, T_F]$. As in the $f = 0$ case [14], we verified in many cases the presence of well-defined hysteresis loops for values of $T < T < T_F$, where the border values $T_b$ depend on $\langle k \rangle$, $\pi$, and $f$. In all the simulations we used $\Delta T = 5 \times 10^{-4}$ and for every set of parameters we performed several checks using values of $t_1$ between $10^2$ and $10^4$. If the values $T_b$ turned out to be independent of $t_1$ in that range (within errors) the transition threshold was estimated as the average of the hysteresis loop, $T_c = (T_b + T_b)/2$. When the hysteresis loop showed a strong dependency on $t_1$, we switched to the next method to estimate the transition threshold.

(b) **Order parameter histograms analysis**. For fixed values of $N$, $\langle k \rangle$, $\pi$, and $f$, we computed a histogram of the values of the order parameter $S_1$ along a single, long simulation run, for different values of $T$. Close to a discontinuous transition, one expects such distribution to show a two-peak structure for long enough simulation times (i.e., for periods of time such that the system evolution provides a good sampling of both phases). The transition threshold can then be estimated as the value of $T$ for which both peaks are the same height. This method is useful when the probability of jumping from one phase to the other is relatively high (moderate system sizes and/or close enough to a critical point), so that the characteristic flip time between phases is small compared with the simulation time. The histogram method is very well established for studying first-order phase transitions in systems under thermodynamic equilibrium [17]. The consistency of our results shows that the method can also work in nonequilibrium discontinuous transitions.

**III. RESULTS**

For $f = 0$, i.e., in the absence of inhibitory neurons, the model corresponds to the case studied in Ref. [9]. It can exhibit different dynamical regimes, including a percolation-like phase transition between high and low activity regimes, depending on the topological parameters of the underlying network. Such transition can be of second order (i.e., critical) for intermediate values of $\langle k \rangle$ and high enough values of $\pi$, or first-order-like (discontinuous) for large enough values of $\langle k \rangle$ [9].
We started our analysis by considering the effect of including inhibitory neurons in a network whose topological parameters correspond to the second-order region for \( f = 0 \). We found that the presence of inhibitory neurons does not eliminate the continuous transition. On the contrary, \( f \) acts as a new control parameter for the transition, as shown in Fig. 1 for \( \langle k \rangle = 16 \) and \( \pi = 0.6 \). In other words, the transition can be observed (i.e., a size-dependent maximum of \( \langle s \rangle \) at the point where the order parameter almost falls to zero) either by changing \( T \) for fixed \( f \) (see Fig. 1) or by changing \( f \) for fixed \( T \) (not shown). Hence, we have a line of critical points in the \((f, T)\) space, whose universality class will be analyzed later. Very similar results were obtained for other values of \((\langle k \rangle, \pi)\) in the critical region for \( f = 0 \) (see Fig. 4 of Ref. [9]).

Next, we analyzed the influence of inhibitory neurons on the dynamics when the topological parameters for \( f = 0 \) give rise to a discontinuous transition. The typical behavior of the order parameter and \( \langle s \rangle \) as a function of \( T \) for fixed \( f \) is shown in Fig. 2 for \( \langle k \rangle = 30 \) and \( \pi = 0.6 \). We see that for small fractions of inhibitory neurons the transition remains discontinuous, giving rise to a first-order transition line. However, as \( f \) increases, the nature of the transition changes smoothly to second order, where the maximum of \( \langle s \rangle \) starts to exhibit a strong size dependency. This suggests the presence of a tricritical point where the first- and second-order transition lines meet. In order to better characterize this phenomenon, we first performed a detailed calculation of the first-order transition line, using the two methods described in Sec. II B. Hereafter we will focus on the \( \langle k \rangle = 30 \) and \( \pi = 0.6 \) case. For small enough values of \( f \) (i.e., up to \( f \approx 0.25 \)) we observe well-defined hysteresis loops (namely, independent of the rate of change of \( T \)) as shown in Fig. 3. We also observe that the area of the hysteresis loops shrinks as \( f \) increases and tends to disappear for \( f \approx 0.3 \), giving a first estimation of the tricritical point location. However, a strong dependency on the rate of change of \( T \) emerges for \( f > 0.25 \) and the method loses accuracy in that region. As we depart from the
The threshold $T$ is changed $T \to T + \Delta T$ with $\Delta T = 5 \times 10^{-4}$, every $t_1$ simulation steps. Empty black symbols correspond to $t_1 = 100$ and filled red symbols to $t_1 = 10^3$. The network parameters are $\pi = 0.6$, $(k) = 30$, and $N = 2 \times 10^3$.

An example for $f = 0.8$ is shown in Fig. 4. We summarize all the previous results in the phase diagram in $(f, T)$ space shown in Fig. 5. We see that both transition lines (first and second order) meet at the tricritical point (in the thermodynamic limit). An example is shown in the insets of Figs. 6(a) and 6(c). Such pseudo-first-order behavior has already been observed in the two-dimensional Potts model with $q = 4$ [19]. We estimated the tricritical point location as that where the above-described change in finite-size behavior occurs.

The fact that the minimum goes to zero for $N \to \infty$ corresponds to the existence of two well-defined and distinct phases in the thermodynamic limit. An example of such behavior (although weak due to the closeness of the tricritical point) is illustrated in Fig. 6(a). On the other hand, close to the tricritical point (but on the continuous side), both maxima and the minimum tend to collapse into a single maximum when $N \to \infty$, as shown in Figs. 6(b) and 6(c). Such pseudo-first-order behavior has already been observed in the two-dimensional Potts model with $q = 4$ [19]. We estimated the tricritical point location as that where the above-described change in finite-size behavior occurs.

Finally, we consider the universality class of the second-order transitions, by estimating the critical exponents $\gamma/\nu d$ and $d_{f}/d$ from the finite-size scaling behavior of the maxima of $\langle s \rangle$ and $\langle S_2 \rangle$. An example is shown in the insets of Fig. 4 for $(k) = 30$, $\pi = 0.6$, and $f = 0.8$. We found that all along the continuous transition line of Fig. 5 the exponents...
are compatible with the mean-field percolation universality class \( \gamma / \nu d = 1/3 \) and \( d_f / d = 2/3 \), as observed for \( f = 0 \) and smaller values of \( k \) [9]. We observed the same behavior even for values of \( f \) relatively close to the tricritical point, i.e., down to \( f = 0.359 \) (although fluctuations become larger as we approach the tricritical point, thus increasing the error bars), so we were not able to clearly detect a crossover to a different set of exponents. To further check the consistency with the mean-field percolation universality class we also analyzed the associated behavior of the cluster size distribution (CCDF) \( P(s) \equiv N_s / N \) at different critical values \( T = T_c(f) \). Figure 7 shows that the associated behavior of the cumulative cluster size distribution exhibits the expected behavior \( P_s(s) \equiv \sum_{s' \geq s} P(s') \sim s^{-(\tau - 1)} \exp(-s/S^*) \), with an exponent \( \tau \approx 5/2 \) and \( S^* \propto (S_2) \) (thus satisfying the scaling law \( \tau = d/d_f + 1 \)). A similar analysis with similar results was performed for points along the second-order line for \( k = \{16\} \) and \( \pi = 0.6 \).

**IV. CONCLUSIONS**

At first sight, the effects of changing the interaction sign on a fraction of neurons could be interpreted as nothing more than a trivial rescaling of the excitability control parameter (i.e., the threshold \( T \)). In fact, as shown in Fig. 5, this holds only for relatively small fractions of inhibitory neurons: the discontinuous transition as a function of increasing numbers of inhibitory neurons occurs now for relatively smaller values of \( T \). However, for \( f \sim 0.35 \) a novel dynamics appears; as the parameter \( f \) is increased the line meets the tricritical point and then continues as a second-order phase transition. To interpret its biological relevance, it may be important to recall that the condition for the tricritical point to appear is (besides a large enough fraction of inhibition) that the network connectivity is very large (i.e., high \( k \)). For such highly connected networks, in the absence of inhibition there is typically an explosion of highly synchronous bursts in which a very large number of neurons is active, even in response to very small perturbations. This dynamics, corresponding to a first-order phase transition, has no behavioral or cognitive value since high synchrony impedes any information processing or storage. Since the connectivity of cortical neurons is typically in the thousands, a given fraction of inhibitory neurons can prevent such synchrony. It is intriguing that the
percentage of inhibitions is usually set around 20%, but probably such quantity cannot be predicted without accounting for the more complex topology of the real brains compared with the simple Watts-Strogatz network topology studied here.

In summary, these results demonstrate that the addition of inhibitory neurons enriches the dynamical phase diagram observed in Greenberg-Hasting neural models defined on small-world networks [9]. The present work adds to the plethora of dynamics exhibited by extended excitable media, including the tricritical behavior reported recently in related models (see Refs. [20,21]). The fraction of inhibitory neurons acts then as an alternative control parameter (in addition to the usual activation threshold) for the dynamical phase transitions between a low activity phase and a percolated, highly active one. Moreover, we observed that the presence of inhibitory neurons allows the emergence of a tricritical point in highly connected networks, i.e., a critical region in parameter space where a second-order (i.e., critical) transition hypersurface and a first-order (i.e., discontinuous) transition one join smoothly. We found evidence that, both for large and low values of the connectivity ($k$) the second-order surface belongs to the mean-field percolation universality class. On the other hand, we were not able to observe a crossover to a different set of exponents on approaching the tricritical point for large values of ($k$), due to a large increase in fluctuations, which make an accurate estimation difficult. This scenario suggests the existence of a tricritical fixed point associated to the tricritical surface (in the sense of renormalization group) located far away from the region here analyzed in the parameters space.

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