Small-world structure induced by spike-timing-dependent plasticity in networks with critical dynamics

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The small-world property in the context of complex networks implies structural benefits to the processes taking place within a network, such as optimal information transmission and robustness. In this paper, we study a model network of integrate-and-fire neurons that are subject to activity-dependent synaptic plasticity. We find the learning rule that gives rise to a small-world structure when the collective dynamics of the system reaches a critical state which is characterised by power-law distributions of activity clusters. Moreover, by analysing the motif profile of the networks, we observe that bidirectional connectivity is impaired by the effects of this type of plasticity.

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I. INTRODUCTION

In their seminal paper, Watts and Strogatz [1] described a class of networks that lie halfway between completely random and regular networks. This class of networks is characterised by a small average path length (which is typical for dense random networks) and an average clustering coefficient significantly larger than expected by chance (as in many regular lattices). However, the notion of a special class of small-world networks is somewhat misleading and it seems more appropriate to speak of a small-world property, i.e. this property is not a binary but a gradual one and is stronger in some networks as compared to others as can be expressed by quantitative measures [2].

The small-world property offers a structural benefit to the processes taking place within the network, e.g. with respect to information transmission by speeding-up the communication among distant nodes in a regular network that can be reached via short-cuts in the small-world case. Moreover, this class of networks is a suitable model in social, technological and biological contexts etc. [2]. For example, it has been observed that the electrical power grid network of western United States, the co-stardom network comprising movie actors and their collaborations and the neural network of the nematode worm C. elegans have the structural properties of low average path length along with a high average clustering coefficient [1].

In another seminal paper, Barabási and Albert [3] proposed a model, known as the BA model, to explain the emergence of scale invariance in the degree distribution e.g. in the World Wide Web (WWW) [4]. In these scale-free networks the probability $P(k)$ that a node connects to $k$ other nodes follows a power-law $P(k) \sim k^{-\gamma}$ [3]. It implies the existence of many poorly connected nodes co-existing with very few but not negligible massively connected nodes hubs. Scale-invariant degree distribution and the small-world property are by no means exclusive and in fact many scale-invariant networks are also small-world.

A look at the degree distribution of a network and the estimation of some network statistics like local clustering coefficients and shortest path lengths provide an insightful initial description of network structure. A more detailed picture of the network structure can be obtained by considering the relative frequency of particular configurations of small blocks in comparison to random networks [5]. In particular, 3-node subgraphs, the motifs, describe the relationship among nodes taken in threes. Song et al. [6] observed that the motif distribution of acute slices taken from the visual cortex of rats exhibits an overrepresentation of some 3-node patterns when compared to random networks. Moreover, networks from a similar context share similarities in their motif profiles, so that these 3-node subgraphs might even define broad classes, or super-families, of networks [5,7]. Similar motif distributions could point to similar dynamical processes in the network, e.g. food networks represent the flow of energy from bottom to top of the food web resulting in a particular motif configuration, whereas brain networks represent a flow of information without necessarily implying a particular direction of the flow, which results in a different motif profile [5].

When it comes to the dynamics occurring in a system that comprises a large number of simple elements interacting in a network, a large number of studies have been dedicated to the occurrence of power-law behaviour and its relationship with the concept of self-organised criticality (SOC) [8]. The concept of SOC has been suggested to explain the dynamics of phenomena as diverse as plate tectonics [9], piles of granular matter [10], forest fires [11], neuronal avalanches [12] (see below), and mass extinctions [13], among several others. SOC implies the existence of a critical point that becomes an attractor in the collective dynamics of a system which resides thus...
at the boundary between two different phases of the system (e.g. ‘order’ and ‘chaos’). SOC can be characterised by power-law event-size distributions, divergence of the correlation length, critical slowing-down of the decay of perturbation as well as the existence of avalanche-like causation of events in the system [8].

In neural systems, an activated neuron can trigger a chain reaction which takes the form of a cascade of activity. These neuronal avalanches have been observed in biological neurons [14] as well as in models (see Sect. 1A). It has been reported that the distribution of avalanche sizes and avalanche durations can be approximated by power laws with resp. exponents $\gamma = -3/2$ and $\delta = -2$ in the thermodynamic limit in globally-coupled networks [12, 13]. Interestingly, the theoretical predictions are in good agreement with experimental results in real brain tissue in local field potentials of cultured slices of rat cortex [14], and in the superficial cortical layers of awake, resting primates [16].

Critical dynamics of brain networks have been studied thoroughly in artificial models, and it has been found that the critical regime implies several computational benefits for the system, namely: optimal information transmission and maximum dynamic range [17], maximum information storage [18, 19], stability of information transmission [20], among others.

An important area of research in network science is focused on studying the mechanisms by which nodes connect and disconnect during the development of a network. Complex networks often possess feedback mechanisms by which the node dynamics and their interactions affect the structure of the network, which in turn alters the behaviour of the nodes etc. It is thus useful to consider both the dynamics on networks as well as the dynamics of networks [21, 22].

In brain networks, synaptic plasticity refers to the structural changes that neuronal networks undergo through the strengthening or weakening of synaptic connections in response to the in-going activity. In particular, spike-timing dependent plasticity (STDP) captures the existing temporal correlations between the spikes of pre- and post-synaptic units resulting in a temporally asymmetric learning rule that has been proposed as a mechanism for learning and memory in the brain [23]. STDP is a local rule that emphasises the precise timing of each individual spike, it also incorporates and extends the essential mechanism of Hebbian learning by including the notion of long-term potentiation (LTP) and long-term depression (LTD) of synapses based on the differences in activation times for pre- and post-synaptic neurons [24].

In this paper we study the effects of STDP mechanisms over systems comprising integrate-and-fire neurons poised at criticality and whose structures follow a complex topology. In the following sections we report how these two concepts combined, that of STDP and that of criticality, have an effect on structural properties of the network which are summarised in the concept of the small-world property.

II. MODEL

A. The Eurich model

The model consists of $N$ non-leaky integrate-and-fire nodes and was formulated for all-to-all connectivity [12], but here we will also consider heterogeneous directed networks.

In the model, every node $j$ is characterised by a membrane potential $h_j$ which is a continuous variable updated in discrete time according to the following equation:

$$h_j(t+1) = h_j(t) + \sum_{i=1}^{N} A_{ij} w_{ij} s_i(t) + I_{ext}$$  \hspace{1cm} (1)

where $A$ represents the adjacency matrix with entries $A_{ij} = 1$ if node $i$ sends an edge to node $j$, and $A_{ij} = 0$ otherwise; $w_{ij}$ denotes the synaptic strength from node $i$ to node $j$; $s_i(t) \in \{1, 0\}$ represents the state of node $i$ (active or quiescent, respectively) at time $t$; and $I_{ext}$ denotes external input which is supplied to a node depending on the state of the system at time $t$. If there is no activity at time $t$, then a node is chosen uniformly at random and its membrane potential is increased by a fixed amount through the variable $I_{ext}$ that represents the external driving of the system. When the potential exceeds a neural threshold, i.e. $h_i(t) \geq \theta$, the node becomes active, $s_i(t) = 1$. Afterwards, this node is reset, i.e. $h_i = 0$.

In the simulations we study networks of different sizes and with the following network topologies:

i. fully connected,

ii. random,

iii. scale-free with low mean clustering coefficient (CC) and power-law out-degree distribution,

iv. scale-free with high CC and power-law out-degree distribution,

v. scale-free with low CC and power-law in-degree distribution,

vi. scale-free with high CC and power-law in-degree distribution.

It is worth mentioning that for the case of random and scale-free structures, the networks considered possess the same number of edges per system size, which results in the same average connectivity per network. In other words, as they have the same number of edges, their structure remains per a permutation of the edges. For a random network, edges are inserted independently as in the Erdös-Renyi model [25], for scale-free networks we follow Ref. [26]. The latter algorithm achieves tunable clustering by performing a triad-formation step in addition to growth and preferential attachment (which are the same as in the BA model [3]).
Thus, we consider two levels of mean clustering for scale-free networks (low and high) by tuning a simple parameter \(20\). The small-world property is not a binary one, and as such, there exists a degree of what we would call small-world-ness. The process of tuning the mean clustering coefficient in our scale-free networks has an immediate effect in the degree of small-world-ness of such structures.

B. Broadcasting hubs and absorbing hubs

In directed networks, the in-degree and the out-degree distribution are generally not the same. Some directed networks possess a power-law distribution in both their in-degree and out-degree distributions, but with different exponents (e.g. the WWW \(25\)), whereas others have a power law only in one of the two directions (e.g. citation networks \(25\)). In either case, the presence of a long-tail in the out-degree distribution of a network implies the existence of broadcasting hubs, that is, nodes that have massive outgoing connections compared with other nodes in the system. On the contrary, the presence of a long-tail in the in-degree distribution implies the existence of absorbing hubs, that is nodes that have a big amount of incoming connections. Here, we are interested in analysing how collective dynamics develop for the case of networks with broadcasting hubs and for networks with absorbing hubs.

C. Spike-timing dependent plasticity (STDP)

As mentioned in Sect. 1, STDP is a temporally asymmetric form of Hebbian learning induced by temporal correlations between pre- and postsynaptic neurons. Synaptic weight between pre- and postsynaptic nodes is potentiated (increased), if the postsynaptic neuron fires shortly after the presynaptic neuron. It is depressed (decreased) if the opposite happens, namely the post-synaptic neuron fires shortly before the pre-synaptic neuron. We implemented STDP mechanisms in our model through the following set of equations:

\[
 w_{ij}(t + 1) = w_{ij}(t) + \Delta w_{ij}(\Delta t)
\]

where \(\Delta t = t_{\text{post}} - t_{\text{pre}}\) denotes the difference between spikes of pre- and post-synaptic neurons, and:

\[
 \Delta w_{ij}(\Delta t) = \begin{cases} 
 a_p \exp\left(-\frac{\Delta t}{T_p}\right) & \text{if } \Delta t \geq 0 \\
 -a_d \exp\left(-\frac{\Delta t}{T_d}\right) & \text{if } \Delta t < 0 
\end{cases}
\]

where parameters \(a_p\) and \(T_p\) set the amount and duration of LTP, whereas \(a_d\) and \(T_d\) set the amount and duration of LTD. In our experiments we set \(a_p = a_d = 0.1\). Observations of STDP in brain tissue suggest that the time window for potentiation is typically shorter than the depression time window \(23\), for that reason we let \(T_p = 10\) time-steps and \(T_d = 20\) time-steps. However, it also has been observed that time-windows and amount of potentiation/depression vary across species and brain structures \(24\).

We impose hard bounds on synaptic weights, that is, \(0 < w_{\text{min}} < w_{ij} < w_{\text{max}} \forall i, j\), which prevents unbounded weight growth, gives rise to strong competition between inputs to a neuron and results in a bimodal distribution of the synaptic weights at the end of simulation time \(27\). In order to allow for activity-dependent pruning of synapses, we set \(w_{ij} = 0\), if \(w_{ij} \leq w_{\text{min}}\), following application of Eq. 2. If the connection is to be terminally deleted, we set also \(A_{ij} = 0\) once \(w_{ij} = 0\).

D. Estimation of small-world-ness.

Following the ideas in Ref. 2, we estimate the degree of small-world-ness in a network through the following process. Let \(G\) be a network consisting of \(n\) nodes and \(e\) edges. To test whether \(G\) exhibits the small-world property we construct a random network \(R\) with same number of nodes and same number of edges. Then, we estimate the mean clustering coefficient of both networks \(CC_G\) and \(CC_R\) along with their mean path length \(L_G\) and \(L_R\). Finally, we compute \(S\) as the ratio of those values:

\[
 S = \frac{CC_G/CC_R}{L_G/L_R}
\]

If \(S > 1\), then \(G\) possesses the small-world property \(2\), which implies a more abundant presence of cliques and long-range connections among nodes than expected by chance.

E. Numerical implementation

When starting simulations, all membrane potentials are initialised at random, whereas states are set to inactive. The neural threshold \(\theta\) will always be set to unity. We let the system reach the critical state, which is identified by a power-law approximation of the distribution of avalanches (see below), which implies that large events coexist with small events during running time. Afterwards we apply STDP mechanisms to update the synaptic strength among nodes based on their activity. At the end of simulations, we analyse the resulting network and the dynamics of the system.

Both the relaxation time towards the critical state as well as the sampling time needed to assess criticality depend on the system size. We consider network sizes \(N = 128, 256\) and \(512\) and choose the initial ‘settling’ time of the networks to, resp., \(10^6, 2 \times 10^6, 3 \times 10^6\) before introducing STDP mechanisms in the system. This selection of times is appropriate for large events to take place during simulation time.
Results are fitted by power-law distributions and the quality of the fit is evaluated by the mean-squared deviation $\Delta \gamma$. This is obtained based on the best-matching power-law exponent $\gamma$ from linear regression in log-log scales. Our choice of using this method is due to its simplicity and justified by the asymptotic unbiasedness of the estimation. When this error function is at its minimum, that is, when the data is best approximated by a power-law distribution with exponent $\gamma$, is then when the system is at criticality.

Moreover, following Ref. \[28\] we inspect the value of the largest eigenvalue of the matrices $W$ associated to each network and whose entries $w_{ij}$ denote the synaptic weight between node $i$ and $j$. The authors in Ref. \[28\] observe that the largest eigenvalue of the weight matrix governs the dynamics of the system. Through an analytical derivation, it is reported that when the largest eigenvalue equals unity the system is at the critical state.

In Table I we report the value of the largest eigenvalue $\Lambda$ of the weight matrices associated to our networks. In our experiments the critical state is not only identified by the power-law distribution of avalanche sizes (see Fig. 3) before the onset of STDP mechanisms, but also by the value close to unity of $\Lambda$.

After the system has reached such a state, we let STDP mechanisms to set in at the synaptic level, that is, at the level of individual coupling strengths. For all system sizes and topologies we allow four million time steps of STDP regime, after which the simulation is over and we proceed to analyse the structural changes of the system. Unlike the model in Ref. \[29\] we do not distinguish between STDP-neurons and non-STDP-neurons, rather in our model every synapse in every unit is susceptible of plasticity.

For our experiments we consider 50 different networks per class (b to d in Sect. II A) and system size for the sake of statistical robustness. In the case of fully-connected networks, as there exists only one fully-connected network of size $N$, randomness is introduced in our code for each realisation of the experiment rather than in the structure as for the other network classes considered. After each realisation of the experiment we generate 100 random networks to compute the metric $S$ (see Sect. II D) for each of the networks that result from the simulation. Experiments were carried out in the EDDIE computer cluster of the University of Edinburgh.

III. RESULTS

A. Small-world structure emerges in fully-connected networks

We observe that after simulation time a small-world structure emerges from fully-connected networks as measured by the metric $S$. In Fig. 1 we show the evolution of $S$ per time step for the fully-connected and random topologies considered. (We present mean values and standard deviations estimated from the different realisations of our experiment.)

Unlike the fully-connected case, the random structure does not show any particular trend regarding the evolution of $S$. Therefore, STDP does not imply any improvement in the structure of random networks. Moreover, as we can see in Fig. 1 STDP acts faster in smaller systems than in larger ones, thus its effect is clearer in the networks comprising 128 nodes than in those of 512 nodes. In any case, we observe a positive trend in the evolution of $S$.

What is the effect of STDP in the scale-free networks considered? As mentioned previously, these structures already possess the small-world topology, that is $S > 1$ for all of them, however the degree of small-world-ness in them vary. We considered high and low values of
the mean clustering coefficient for this type of networks, this results respectively in high and low degrees of small-world-ness in the networks considered. The reason we considered such classes is because we are interested in assessing how STDP mechanisms affect scale-free structures with varying clustering coefficients. Additionally, as stated previously, we consider scale-free networks with broadcasting hubs and networks with absorbing hubs.

For these topologies, STDP has a negative effect regarding their small-world-ness, identified by a decrease in the value of $S$ once STDP sets in. For example, in Fig. 2a we show a comparison of the evolution of $S$ for two scale-free networks with 512 nodes and low mean clustering coefficient (which implies low degree of small-world-ness). The network identified by the continuous line has a power-law out-degree distribution (which yields broadcasting hubs), whereas the dashed line represents its transpose, that is, a network with a power-law in-degree distribution (which yields absorbing hubs). As it can be seen, STDP affects negatively the small-world property in these networks.

Similarly, Fig. 2b traces the changes of $S$ for two scale-free networks of same size as above but with high degree of small-world-ness. The continuous line represents a scale-free network with power-law out-degree distribution, whereas the dashed line represents its transpose, a network with a power-law in-degree distribution. In the best of cases, $S$ does not exhibit an noticeable decrease. These situations are verified in all other systems sizes considered for scale-free networks.

It is clear that STDP does not have the same effects when the power-law of the degree distribution is present in the out-degree distribution or in the in-degree distribution; nor when the clustering coefficient of network is low or high. Nevertheless, we find that STDP affects negatively the degree of small-world-ness in scale-free networks.

**B. STDP impairs criticality**

As mentioned in Sect. II E, we assess the quality of the power-law approximation to the distribution of avalanches by estimating the deviation from the best power-law fit. When such an error function reaches a minimum value of less than 0.05, we consider the event-size distribution as well approximated by a power-law and conclude that the system is in a critical state. After the system has approached a power-law behaviour, we allow STDP mechanisms to set in. We observe that criticality is generally lessened while STDP is modifying the synaptic strengths. With STDP the difference between the distribution of avalanche sizes and the best-fit power-law increases again, such that eventually a power-law ceases to appropriately model the system, see Fig. 2a. A similar behaviour as shown there is observed in all the networks that we considered.

The deterioration of the power-law approximation is explained by the fact that once STDP sets in large avalanches cease to occur and only small avalanches take place. In this sense, it can be said that STDP favours local, clustered events. Fig. 2a shows an example of this behaviour. Here, the continuous line shows the system during criticality before STDP mechanisms take place, and can also be identified in the minimum value shown in Fig. 2b around the time step $2 \times 10^6$. However, after STDP there are no more large avalanches to be added up to the distribution, only small ones, which results in the particular shape of the dashed line in Fig. 2a. As well, in Fig. 2b after STDP has set in right after the $2 \times 10^6$ time step the deviation from a power-law distribution increases and we observe a larger error as time passes.

As mentioned before, this same phenomenon is ob-
served across all networks considered in our model. This behaviour is captured by the largest eigenvalue of the weight matrix \( W \) denoted by \( \Lambda \). Before STDP the value of \( \Lambda \) is close to unity. However, once STDP mechanisms set in, the synaptic modulation results in a weight matrix whose largest eigenvalue is less than unity, thus reflecting the deviation from criticality (see Table I). From this we might conclude that in our model the critical state vanishes as STDP sets in.

Does this imply that self-organised criticality and spike-timing-dependent plasticity are two biological phenomena that cannot coexist? The model proposed by Levina et al. [15], based on dynamical synapses, exhibits critical behaviour that coexists with STDP. The modulation induced by such dynamical synapses results in a compensatory mechanism that recovers the critical regime.

C. STDP prunes direct-feedback connections

We analyse the motif profile of the networks after the STDP regime. There are 13 different motifs representing the possible relations among nodes taken in threes from a directed network. The different configurations are shown in Fig. 4.

A motif profile shows the distribution of these 13 different 3-node configurations for a single network. For example, before STDP, a fully-connected network possesses a single motif distribution: all 3-node relationships are of the XIII type (see Fig. 4). However, after STDP the motif XIII breaks apart and instead motifs of the types I, II, IV, V and IX abound. Interestingly, none of these contain direct feedback connections. Fig. 5 shows the motif profiles of the fully-connected networks considered (sizes 256 and 512 inside the inset) after STDP. For the smallest system size it is more evident how motifs I, II, IV, V and IX grow where previously there was only motif XIII. As the system grows, STDP requires more time to prune the network and profiles differ from one another.

In the case of heterogeneous topologies, STDP attacks motifs with bidirectional connections affecting in this way the local clustering of the global structure, which in turn affects the degree of small-world-ness of the network. Motifs with bidirectional connections are: III, VI, VII, VIII, X, XI, XII and XIII; all of them impaired by the action of STDP. Fig. 6 shows this behaviour for our networks of 128 nodes. However, we observe this particular behaviour in all the different system sizes considered.

In particular, motif II is severely impaired in all cases considered. This motif represents the most basic feed-forward flow of information comprising 3 nodes, and as such the most elementary pre- and post-synaptic relationship among three nodes, and as such it is expected to be a target for STDP. Interestingly, some motifs vanish completely from the profile. These are motifs III, VI, VII, VIII, X, XI, XII and XIII; all of them involving direct feed-back connectivity.

Therefore, in a sense it can be said that STDP prevents direct feedback connectivity, preferring indirect feedback flow in which a third node serves as intermediary. For example, motif X includes a direct feedback connection between two nodes; however, after STDP this motif might transform into motif IX, in which the feedback flow is now mediated by a third node.

STDP favours one-way connections rather than two-way connections, thus establishing a direction for the flow of stimuli given by the current activity in the system. Simply put, if node \( i \) sends an edge to node \( j \), but this latter node fires shortly before the former most...
FIG. 4: All configurations for 3-node connected subgraphs.

FIG. 5: Post-STDP motif profile for fully connected networks of size $N = 128$. (Inset: sizes $N = 256$ and $N = 512$.) After STDP, motif $XIII$ breaks apart and motifs $I$, $II$, $IV$, $V$ and $IX$ emerge. This is more evident for smaller system sizes, where STDP requires less time for pruning.

of the time, then STDP acts by severing this somewhat erroneous connection.

D. Other effects on topology

Previously, we mentioned how STDP affects the degree of small-world-ness in the networks considered. The process of activity-dependent pruning has other consequences in topology, namely, the decrease in edge density and eventually, the disconnection of the network and the emergence of different connected components across the system.

The density of a network is the ratio of the number of edges to the number of possible edges. As such, a fully-connected network possess all possible connections in the network resulting in the maximum possible value of the density, which is 1. As mentioned before, for any of the system sizes considered, the heterogeneous structures have the same number of edges, which yields the same density for all of them. However, how STDP affects this ratio varies depending on the particular type of network considered (fully-connected, random or scale-free).

Network density is an indicator of sparseness in a structure. Our heterogeneous networks are sparse, a feature that has an effect on the way STDP acts upon the structure. Fig. 4 shows the effects of STDP mechanisms on network density for all the fully-connected cases considered and for the heterogeneous structures of size 128. We observe the same behaviour for the other system sizes considered.

The effects of STDP are soon visible in a dense network topology. In a fully-connected network, STDP acts fast removing a considerable amount of edges as soon as it sets in. In contrast, for sparse-networks, either scale-free
or random, STDP acts more slowly not even removing 1% of the initial configuration under otherwise similar conditions.

However, STDP does not act the same in every heterogeneous topology. Scale-free networks in which the power-law is present in the in-degree distribution (that is, networks with absorbing hubs) lose density faster than any other, specially when the mean clustering coefficient is low. This can be explained by the fact that in in-degree scale-free networks absorbing hubs are more susceptible to fire, which in turn has a strong influence on STDP mechanisms. However, clustering seems to serve as a protective mechanism to prevent edge pruning. This can be observed in the fact that our scale-free networks with low mean clustering coefficients lose edges faster than their high clustered counterparts.

Nevertheless, out-degree scale-free networks (that is, networks with broadcasting hubs) decrease the size of their largest connected component faster than any other topology, specially when combined with high mean clustering coefficient.

During simulation time, the effects of STDP pruning result on a disconnected network where components of different sizes emerge. All our observations regarding topology refer to the largest connected component (LCC) of the network. Because STDP behaves differently across the different topologies considered, we expect the LCC to be different as well. As we have just mentioned, the LCC of scale-free networks with broadcasting hubs suffer a shrinkage of size faster than random networks and scale-free networks with absorbing hubs, even if it is this latter type of networks the one that loses density faster than any other topology. This behaviour is shown in Fig. 8b and it is observed across all system sizes considered.

In summary, what is happening is that scale-free networks with absorbing hubs lose edges faster than any other structure. However, these networks show more resilience to become disconnected (Fig. 8a). On the contrary, scale-free networks with broadcasting hubs show more resilience to lose edges, but they are more prone to become disconnected when losing edges (Fig. 8b).

E. Effects of spike triplets

In this section we briefly discuss the effects of extending the implementation of STDP mechanisms from pairs of spikes to triplets of spikes. The computational implementation of STDP described in Sect. II C takes into account
pairs of spikes to estimate the synaptic modulation at each connection, namely, one from the pre-synaptic node and another from the post-synaptic node, whose order determine the nature of the modulation. In the following, we refer to paired-wise STDP as pSTDP.

It has been reported that pSTDP fails to replicate observations in experimental data. Intuitively, a pre-post pairing followed by a post-pre pairing of the same magnitude would cancel out any synaptic modulation triggered, however this is not what it has been observed in biological experiments. In experiments, models of pSTDP are not able to explain the synaptic modulation triggered by higher-order plasticity rules such as triplets and quadruplets of spikes \[30, 31\]. For this reason, pSTDP has been extended in order to consider triplets of spikes rather than just pairs of them. In the following, we refer to triplet-wise STDP as tSTDP to differentiate it from pSTDP.

A triplet rule for tSTDP involves sets of three spikes: two pre- and one postsynaptic, or one pre- and two postsynaptic spikes. With a triplet rule of this form it is possible to fit experimental data from visual cortical slices as well as from hippocampal cultures. Interestingly, when this rule is based on Poisson spike trains, the learning rule can be mapped to a Bienenstock-Cooper-Munro (BCM) learning rule \[30, 32\]. Moreover, it has been proposed that the triplet rule for STDP is a mechanism used by neuronal networks to perform computations that resemble those of independent component analysis (ICA) \[32\].

When we consider triplets of spikes for STDP, we might feel inclined to extend the model further in order to consider higher-order STDP rules. However, experiments show that spike triplets are able to reproduce data generated by higher-order terms \[30\].

A model for tSTDP is implemented in the following way. The mechanism is similar to that of pSTDP in the sense that the difference between spike times in pre- and post-synaptic neurons determines the amount of synaptic modulation, however one extra term is added to the weight update function. This extra term considers the temporal difference between the two most recent spikes of one of the spiking nodes. Hence, the spike triplet. Potentiation occurs in a similar way as in pSTDP, however the amount of synaptic modulation is also in function of the difference between the two most recent spikes of the post-synaptic unit. Therefore, this rule is identified as post-pre-post. Similarly, depression occurs analogously to pSTDP, but the synaptic update is in function of the temporal difference between the two most recent spikes of the pre-synaptic neuron. So, the rule is summarized as pre-post-pre. Figure 9 shows a schematic representation of this two rules.

Thus, in order to extend pSTDP into tSTDP we add some extra terms to the equation for \( \Delta w_{ij}(\Delta t) \) described in Sect. 11C which yields:

\[
\Delta w_{ij}(\Delta t) = \left\{ \begin{array}{ll}
  a_p \exp\left\{ -\frac{\Delta T_1}{T_p} \right\} \exp\left\{ -\frac{\Delta T_2}{T_p} \right\} & \text{if } \Delta T_1 \geq 0 \\
  -a_d \exp\left\{ -\frac{\Delta T_1}{T_d} \right\} \exp\left\{ -\frac{\Delta T_2}{T_d} \right\} & \text{if } \Delta T_1 < 0 
\end{array} \right.
\]

where parameters \( a_p, P \) and \( T_p \) set the amount and duration of LTP, whereas \( a_d \) and \( T_d \) set the amount and duration of LTD, as with pSTDP (see above); \( T_1 \) represents the difference between pre- and post-synaptic spikes, \( T_2 \) denotes the temporal difference between the two most immediate post-synaptic spikes (if \( \Delta T_1 \geq 0 \)) or the temporal difference between the most immediate pre-synaptic spikes (if \( \Delta T_1 < 0 \)); and \( T_d \) and \( T_p \) are two parameters that in a similar fashion as with parameters \( T_d \) and \( T_p \) set the amount of influence of immediate spikes for depression and potentiation, respectively.

With these ideas in mind we report the following observations when considering tSTDP mechanisms in our model rather than simple pSTDP. Similar to pSTDP the synaptic modulation triggered by tSTDP mechanisms alter the quality of the power-law approximation to the distribution of avalanche sizes. As expected, this in turn has effects over the error function and the exponent of the power-law fit. This behaviour is also captured by the largest eigenvalue \( \Lambda \) of the weight matrix \( W \). Similar to pSTDP, when tSTDP sets in the synaptic modulation takes \( \Lambda \) away from unity reflecting the deviation from criticality induced by tSTDP (see Table II).

However, although tSTDP has an effect on the modulation of the synaptic strength between two connected nodes, this type of plasticity is less effective than pSTDP on severing connections; to the point that almost all edges survive after simulation time. Because of this, we do not observe any small-world structure emerging from fully-connected networks as we did with pSTDP. At criticality, tSTDP acts as a mild modulator of synaptic weights based on the firing activity of pre- and post-synaptic nodes.

Why tSTDP does not achieve the same amount of pruning as pSTDP? Let node \( i \) send an edge to node \( j \); and let \( j \) spike before \( i \). For potentiation to take place, a second spike from \( j \) to take place. Let us refer to the first spike of \( j \) as \( j_1 \), and to its second spike as \( j_2 \), whereas
the only spike of \( i \) will be denoted as \( i_1 \). If node \( j \)'s first spike and node \( i \)'s only spike belong to the same avalanche we write \( j_1, i_1 \in Av_1 \). As said above, for potentiation to occur, a second spike from \( j \) must take place. These leaves the following possibilities:

1. \( j_1, i_1 \in Av_1 \) and \( j_2 \in Av_2 \). Here, for the maximum potentiation to occur the inter-avalanche interval must be very small. Such a regime, in which firing rate is high, is associated with super-criticality and not with criticality.

2. \( j_1 \in Av_1 \) and \( i_1, j_2 \in Av_2 \). Same as above. Here spikes are separated in two different avalanches that require to be as close as possible in order to potentiation be at its fullest.

3. \( j_1, i_1, j_2 \in Av_1 \). Here what is required is that the three spikes occur in the same avalanche. For this to happen node \( j \) must be active twice in a single avalanche giving place to a non-Hamiltonian avalanche, that is, an avalanche in which there is a node becomes active more than once. Although not shown here, this occurs naturally in scale-free and random networks at criticality due to the sparseness of the network. Therefore we expect this to happen in this type of networks and to be non-existent in fully-connected structures.

We observe that the third case occurs in scale-free networks and this will result in weights reaching the largest allowed value of synaptic weight \( (w_{\text{max}}) \), whereas fully-connected networks fall in cases (1) and (2) described above, in which the weights do not settle in the maximum synaptic weight and rather give rise to a bell-shaped distribution for the weights (not shown here).

Let us consider the case where \( i \) fires before \( j \). For depression to take place we require a second spike from node \( i \). Similar to the case for potentiation, we have the following possibilities:

1. \( i_1, j_1 \in Av_1 \) and \( i_2 \in Av_2 \). As with the cases presented above, this situation requires a small inter-avalanche interval in order to get the maximum possible amount of synaptic depression.

2. \( i_1 \in Av_1 \) and \( j_1, i_2 \in Av_2 \). Same as above.

3. \( i_1, j_1, i_2 \in Av_1 \). As with potentiation, this case requires non-Hamiltonian avalanches to be present in the system. As mentioned earlier, only scale-free and random structures exhibit this kind of behaviour at criticality.

Unlike potentiation, we do not observe a large amount of synaptic depression occurring in our networks, and consequently no synaptic pruning either. This is due to the asymmetry of the STDP learning rule, in which potentiation is benefited over depression plus the fact that the rule requires either a small inter-avalanche interval in the system or non-Hamiltonian avalanches.

IV. DISCUSSION

One important observation regarding small-world-ness in networks is that of the case of fully-connected topologies. It can be argued that this particular type of structure possesses the small-world property \textit{ab initio} as it has both the maximum value of mean clustering coefficient and the lowest value of mean path length possibles, therefore any alteration to its structure can only impair its degree of small-world-ness. This is a very valid observation. However, we should point out that a fully-connected network is a blank slate, in which no dynamic process has yet taken place. STDP carves a structure out of this topology (in particular, pSTDP), that changes the ratio of the mean path length and the mean clustering coefficient when compared to a random network with the same number of nodes and edges. The resulting structure is better than the one that would have emerged from a process pruning edges randomly. Moreover, the value of \( S \) for a fully-connected structure is 1 always, as the comparison with a random network with same number of nodes and edges yields exactly the same fully-connected structure. However, we observe that after STDP the fully-connected network becomes a network in which clearly \( S > 1 \) (see Fig. 1a), which implies the presence of the small-world property.

The model of Basalyga et al. \cite{29} is, unlike ours, not based on critical dynamics. They consider Erdős-Renyi networks of slightly more complex unit but only at a network size of \( N = 100 \) neurons, while we tried to consider scaling relations for system sizes larger than that. Since in there model only excitatory synapses are subject to STDP, the topological effects are biased. While the main observation that STDP can impair criticality is reproduced here, we find a clear improvement of the small-world-ness of the emerging networks.

Our analysis includes an inspection of the motif profile that resulted from STDP mechanisms. Song et al. \cite{6} estimated the motif distribution of acute slices from the visual cortex of rats and observed that the motif profile of such networks differs from their random counterparts, in particular bidirectional connections were found to be more frequent than expected by chance. In contrast, we have obtained a slight decrease of bidirectional motifs which is related to the particular form of the STDP rule used here \cite{33} when compared against a random network, which is not our case. As Song et al. claim, their counts are relative to random, whereas ours are absolute counts, that is, real counts which are not compared against a random network \cite{6}. In their work, they present a ratio of actual counts to that predicted by their null hypothesis. In this context, bidirectional connections show an overrepresentation, a situation which is not in clash with our observations. When the ratio of motifs containing bidirectional connections is compared to that of non-bidirectional connections in the same network profile, we observe that the latter is much larger than the former, an observation that is in agreement with ours. If
we compare the resulting state of our model to the state before STDP was applied, we find that many bidirectional connections have been deleted. Moreover, it can be said that from a certain point of view, their observations are just snapshots of a dynamic process happening in the networks that they considered. Such a process would have another effect in the long run, in which bidirectional connections are found more rarely than before. Also, these authors do not mention anything related to the type of dynamic process that gave rise to the structures that they observed. Thus, the question regarding STDP as a mechanism that prunes two-way connections is not settled by their work.

Ex nihilo nihil fit. The brain is shaped not only by genetics but also by activity-dependent processes during development. It is known that several brain structures possess the features commonly associated to small-world networks [34]. In particular, the only neural network that has been mapped in its entirety, namely the nervous system of the nematode *C. elegans*, is known to possess the small-world property [1]. In higher organisms, genetic processes cannot fully account for the existence of this particular network structure, as during the lifetime of an individual, modification of the network are present due to activity-dependent processes, e.g., in connection to learning and memory. Here, we claim that massively connected structures combined with critical dynamics can give rise to a small-world structure already if a standard STDP rule is in place for adapting the network towards a better-than-random structure which is beneficial for information transmission across the system.

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| Type                          | Subtype       | Size | $\Lambda_{\text{static}}$ | $\Lambda_{\text{pSTDP}}$ | $\Lambda_{\text{tSTDP}}$ |
|------------------------------|---------------|------|---------------------------|---------------------------|---------------------------|
|                              | Low Mean CC   | 128  | 0.906 ± 0.029             | 0.27 ± 0.009              | 0.35 ± 0.01               |
|                              |               | 256  | 0.9 ± 0.02                | 0.45 ± 0.007              | 0.52 ± 0.016              |
|                              |               | 512  | 0.95 ± 0.01               | 0.7 ± 0.005               | 0.85 ± 0.002              |
| Out-degree scale-free        | High Mean CC  | 128  | 0.89 ± 0.04               | 0.26 ± 0.01               | 0.33 ± 0.02               |
|                              |               | 256  | 0.91 ± 0.03               | 0.43 ± 0.01               | 0.51 ± 0.01               |
|                              |               | 512  | 0.91 ± 0.01               | 0.63 ± 0.01               | 0.73 ± 0.009              |
| In-degree scale-free         | Low Mean CC   | 128  | 0.98 ± 0.02               | 0.21 ± 0.005              | 0.32 ± 0.009              |
|                              |               | 256  | 0.99 ± 0.01               | 0.4 ± 0.006               | 0.49 ± 0.01               |
|                              |               | 512  | 1.0006 ± 0.006            | 0.62 ± 0.006              | 0.68 ± 0.002              |
|                              | High Mean CC  | 128  | 0.96 ± 0.02               | 0.23 ± 0.013              | 0.307 ± 0.02              |
|                              |               | 256  | 0.99 ± 0.02               | 0.408 ± 0.016             | 0.47 ± 0.01               |
|                              |               | 512  | 1.002 ± 0.014             | 0.61 ± 0.01               | 0.62 ± 0.007              |
| Random                       | Low Mean CC   | 128  | 0.92 ± 0.012              | 0.28 ± 0.004              | 0.34 ± 0.003              |
|                              |               | 256  | 0.99 ± 0.022              | 0.48 ± 0.003              | 0.53 ± 0.004              |
|                              |               | 512  | 0.97 ± 0.001              | 0.77 ± 0.001              | 0.87 ± 0.001              |
|                              | High Mean CC  | 128  | 0.91 ± 0.034              | 0.098 ± 0.007             | 0.63 ± 0.0001             |
|                              |               | 256  | 0.93 ± 0.0068             | 0.23 ± 0.0062             | 0.78 ± 0.0002             |
|                              |               | 512  | 0.95 ± 0.001              | 0.33 ± 0.004              | 0.88 ± 0.0005             |

TABLE I: Largest eigenvalue $\Lambda$ of matrix $W$ of synaptic weights. It has been found analytically that $\Lambda = 1$ is associated with a system at criticality [28]. The synaptic weight matrices of our networks have $\Lambda \approx 1$ before starting STDP mechanisms. Once STDP (either paired- or triplet-wise) mechanisms set in the synaptic weights are modulated based on neuronal activity. This affects the spectral nature of the weight matrix such that $\Lambda \ll 1$ post-STDP. Columns $\Lambda_{\text{static}}$, $\Lambda_{\text{pSTDP}}$ and $\Lambda_{\text{tSTDP}}$ refer to the largest eigenvalue $\Lambda$ for pre-STDP, post-paired-wise STDP and post-triplet-wise STDP, respectively. (We present mean values and standard deviations.)