Intrinsic noise and deviations from criticality in Boolean gene-regulatory networks

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Gene regulatory networks can be successfully modeled as Boolean networks. A much discussed hypothesis says that such model networks reproduce empirical findings the best if they are tuned to operate at criticality, i.e. at the borderline between their ordered and disordered phases. Critical networks have been argued to lead to a number of functional advantages such as maximal dynamical range, maximal sensitivity to environmental changes, as well as to an excellent tradeoff between stability and flexibility. Here, we study the effect of noise within the context of Boolean networks trained to learn complex tasks under supervision. We verify that quasi-critical networks are the ones learning in the fastest possible way—even for asynchronous updating rules—and that the larger the task complexity the smaller the distance to criticality. On the other hand, when additional sources of intrinsic noise in the network states and/or in its wiring pattern are introduced, the optimally performing networks become clearly subcritical. These results suggest that in order to compensate for inherent stochasticity, regulatory and other type of biological networks might become subcritical rather than being critical, all the most if the task to be performed has limited complexity.

The central dogma of molecular biology is that each single gene is transcribed into RNA, which in turn is translated into a protein, which—usually in cooperation with different proteins—can regulate the expression of other genes, giving rise to a complex network of regulatory interactions and different possible patterns of gene expression. Genetic regulation, protein-protein interactions, as well as cell metabolic and signaling pathways are essential biological processes that can all be represented as networks. The network picture encapsulates the complexity of cellular processes and provides us a natural framework for a systems-perspective approach to extremely complicated biological problems. As a matter of fact, the study of information processing in living systems has shifted from the analysis of single pathways to increasingly complex regulatory networks, allowing for a visualization of the collective effects of a host of units acting at unison. Since the pioneering work of Kauffman3–7, genetic regulatory systems have been modeled as Boolean networks, in which the expression level of each gene is represented by a binary (on/off) variable and where mutual regulatory interactions are described as arbitrary random Boolean functions operating synchronously at discrete time steps. Even if admittedly simplistic and limited in a number of ways (e.g. continuous levels of gene expression might be essential to understand some cellular processes), such a binary description is particularly useful when dealing with large networks because it simplifies the overwhelming complexity of the real problem reducing it to a logical one. In particular, the Boolean approach has shed light on important conceptual problems such as the possibility of diverse (phenotypic) states emerging from a unique given genetic network, as well as the possibility of transitions among them (as happens in cell differentiation and reprogramming), and the emergence of cycles in cell states. The trajectory of the segment polarity network in the fly Drosophila melanogaster8 and the yeast cell cycle9 are two specific examples in which the most relevant features of gene expression have been fully elucidated on the basis of Boolean models10 (for more details we refer to the literature4–6,11,12).

Random Boolean networks (RBNs) can operate in different regimes including ordered and chaotic phases as well as a critical point (or line or surface) separating them in parameter space. Ordered or frozen phases (typically obtained for small network connectivities) are characterized by a small set of stable attractors which are largely

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robust to perturbations, while in the disordered or chaotic phase (typically obtained for densely connected networks) perturbations rapidly propagate all through the network hindering the existence of truly stable states. As formalized mathematically by Derrida and Pomeau, separating these two phases there is a critical line (that used to be called the “edge of chaos”) at which perturbations propagate marginally. It was conjectured some time ago that critical RBNs might be optimal to represent actual biological networks: the underlying idea is that operating at criticality might provide such systems with an optimal tradeoff between being exceedingly ordered/stable (thus, barely responsive to environmental changes, signals, and clues) and being too disordered/noisy (thus enormously sensitive to the effects of noise, lacking the required robustness and accuracy that biological machinery demands). The criticality hypothesis states that the marginal situation between these two impractical tendencies constitutes an excellent compromise. This conjecture (which was developed in the machine-learning and neural-network community), proposes that—by operating nearby criticality—networks exhibit an optimal tradeoff between stability to perturbations and sensitivity/responsiveness to signals. Similarly, at larger timescales, it also provides an excellent compromise between robustness and evolvability. Moreover, it entails optimization of information storage and transmission, response and sensitivity, computational capabilities, and a number of other functional advantages.

In parallel, the development of powerful experimental high-throughput technologies in molecular biology has paved the way to experimental analyses of gene-expression patterns in large regulatory networks. Recent empirical results, analyzing hundreds of microarray experiments to infer regulatory interactions among genes and implementing these data into Boolean models, seem to support the hypothesis that regulatory networks of Saccharomyces cerevisiae, Escherichia coli, Bacillus subtilis, the murine macrophage, as well as some subnetworks of Drosophila melanogaster and Arabidopsis thaliana are indeed very close to criticality (in the sense of marginal propagation of perturbations) while some other empirical analyses leave the door open to regulatory networks being ordered/subcritical.

Recent work, aimed at rationalizing why and how criticality might come about in living systems, relies on adaptive/evolutionary models, in which communities of agents—each of them modeled as a Boolean network—are selected for if they succeed at performing some complex tasks which may change in time. For instance, Hidalgo et al. showed by employing an information-theoretic approach that critical networks may emerge as optimal solutions in such a setting (however, the networks employed as a specific example in ref. 30 are fully connected and thus lack the structural richness of usual RBNs). Similarly, Goudarzi et al. considered an ensemble of RBN’s able to experience “mutations” in their topological structure and employed a genetic algorithm to select for those able to perform a given computational task (see Fig. 1); i.e. networks which have learnt have a larger fitness than those that have not. Under these conditions the ensemble converges to a state in which all networks operate close to criticality. In other words, critical networks emerge as the optimal solution out of the combined selective pressures of having to learn different tasks (i.e. having to produce different outcomes/attractors) and being able to readily shift among them following changes in the inputs in real time.

Given that living cells typically possess very low copy numbers of important regulatory molecules (e.g. for the 80% of genes in Escherichia coli genome the copy number of their associated proteins is less than 100) stochastic effects are unavoidable and ubiquitous in gene regulatory networks. Even if noise is usually assumed to be detrimental to reliable information transfer and, more in general, to cell functioning, stochastic effects can lead to beneficial outcomes; for instance, noise accounts for the observed (phenotypic) variability in identical (isogenic) populations and can help cells to adapt to fluctuating environments. Within the framework of RBN the role of stochasticity and noise has been addressed in a number of works.

In this paper, we further delve in the problem of investigating the mechanisms and the conditions under which networks may become critical (or not), focusing on the role played by noise, and ask the question whether—in the presence of strongly noisy conditions—regulatory networks, modeled as RBNs having to perform some complex computational task, operate in ordered, critical or supercritical regimes (see Fig. 1). In other words: what is the role of noise in the emergence of criticality? Does it foster or hinder critical behavior? In order to gauge the effect of noise on the dynamics of RBNs having to perform a complex task we consider a setting very similar to that of Goudarzi et al., including different additional sources of stochasticity. In particular, our approach differs from the previous one in three main aspects: (i) we consider asynchronous updating rather than the usual deterministic one, thus introducing the effect of stochasticity in the updating timings, (ii) both the structure and the dynamics of the networks are subjected to noise (be it intrinsic or external), and (iii) we do not consider an evolutionary algorithm to search for the best possible network connectivity, but rather we work in a constant-connectivity ensemble and explore how the network performance depends on the network connectivity, i.e. on the network dynamical state.

As we shall illustrate, criticality emerges as the solution providing the fastest route to learning complex tasks but, on the other hand, once additional sources of stochasticity are explicitly taken into account, ordered dynamical states perform better than critical ones. That is, networks need to compensate the excess of noise by becoming progressively more subcritical.

Model and Training Protocol
As said above, we consider a setting (similar to that of previous approaches) to train Random Boolean networks to perform a computational task, but we introduce a number of changes—to be detailed in this section—to devoted mostly to implement stochasticity in various ways.

Network architecture and dynamics. Random Boolean networks (RBNs) consist of random Erdős–Rényi (ER) structures with N nodes, L directed links, and averaged connectivity or degree \( K = L/N \). Self-loops and multiple edges with the same directionality linking two nodes are explicitly excluded, and otherwise the wiring pattern is randomly set. Each node \( i \) represents a gene, whose dynamical state is characterized by a Boolean variable...
\[ \sigma_i = \{0, 1\}; 1 \text{ for the ”on” state and } 0 \text{ for the ”off” one.} \]

The node is updated according to a random Boolean function, \( f_i \), which depends on the state of the \( K_{in}(i) \) neighbor nodes regulating it (restricted to a maximum value of 8 for computational convenience), and it contributes to regulating the state of \( K_{out}(i) \) out-neighbors (see Table I in Methods for an example of random Boolean functions). The averaged fraction of 1’s in the outputs of the random Boolean function, \( p \), can be fixed a priori and taken as a control parameter, determining the bias toward ”on” or ”off” states (here, we consider the unbiased case \( p = 1/2 \) in all analyses). In contrast with most studies of RBNs and in order to implement a first source of stochasticity, nodes are updated in an asynchronous way\(^{42–44} \), i.e. a given node is randomly selected with homogeneous probability, its state is updated according to:

\[
\sigma_{i+\Delta t}(t) = f_i(\sigma_{n_1^i}(t), \sigma_{n_2^i}(t), ..., \sigma_{n_{K_{in}}^i}(t)),
\]

where \( n_j^i \) identifies the \( j \)-th neighbor of node \( i \), time is incremented in \( \Delta t = 1/N \) units, and the process is iterated. A time step of the dynamics corresponds to one update per node on average. In order to implement computational tasks or learning rules in RBNs we consider a slight variation of the just-described general architecture, in which some pre-defined input and output nodes are included (see Fig. 1B). By construction, input nodes are imposed to have \( K_{in} = 0 \), so that they are not influenced by others and \( K_{out} \geq 0 \), so that they are not isolated, while –on the contrary– output nodes have \( K_{out} = 0 \) and \( K_{in} \geq 1 \) (in particular, we take \( n_{input} = 3 \) input nodes and one single output or readout node (\( n_{output} = 1 \) as in Fig. 1B). The set of \( N - n_{input} \) non-input nodes is called the network core.

**Assessing the network dynamical state.** In the infinite size limit, synchronous RBNs are known to exhibit a critical point –in the sense of marginal propagation of perturbations\(^{5,13} \)– at a value of the connectivity

\[
K_{c}(p) = \frac{1}{2p(1-p)}
\]

being ordered/subcritical for \( K < K_{c}(p) \) and disordered/supercritical otherwise. In particular, in the unbiased case, \( p = 1/2, K_{c} = 2 \) (see Fig. 1A) which is often quoted as ”the” critical connectivity for RBNs. However, these results hold only for infinite networks; for finite ones, critical values are shifted toward slightly larger connectivity values by corrections of order \( \mathcal{O}(N^{-1}) \). Here, instead of calculating such critical values analytically, and thus to quantify possible deviations from criticality, we explicitly compute in numerical simulations the...
dynamical state of any given finite-size network. For this, we determine whether individual site perturbations do grow or shrink on average; i.e. we measure the branching parameter, $B$, defined as the averaged Hamming distance—after one timestep—between the original and all possible network-states differing from the original one at just a single (flipped) site (see Methods). Branching parameters $B > 1$ (resp. $B < 1$) reflect supercritical (resp. subcritical) networks while the marginal case $B = 1$ is the trademark of criticality.

### Computational tasks.

The task to be learned can be codified in a "truth table", i.e. for each specific input configuration (out of a total of $I = 2^{\text{num}}$) there is an output value to be reproduced. A given truth table defines a specific computational task. An example is the odd-even classifier (rule R150 in the Wolfram's classification of cellular automata), which assigns a Boolean variable to each input accounting for its parity. Other examples that we consider are rules number R51 and R60 in Wolfram's classification. These rules can be categorized accordingly to their "complexity", understanding as such, the number of nodes in the input that do change the output state when altered (and how often they do so for different values of the remaining nodes). In particular, out of the three rules that we study here, the most complex one is the odd-even classifier (R150) whose output obviously depends on all input nodes, R60 is an intermediate case, while the less complex one is R51 whose output is the opposite of one particular input unit, being insensitive to the other two. A more precise definition on how to quantify task complexity—unnecessary for our purposes here—has been discussed by Goudarzi et al.

### Network fitness.

The goal of the trained networks is to produce—as fast as possible—the correct output when exposed to each of the $I = 2^{\text{num}}$ specific input states; i.e. the network learns the computational task as defined by a given truth table. To implement this, we sequentially expose the network to $I$ randomly chosen inputs. The resulting random order of inputs can be viewed as a form of stochasticity, mimicking environmental variability. Moreover, the environment is assumed to change rapidly so that, in order to cope with that, networks are trained to reach the correct output within just $t_{\text{max}}$ (usually fixed to 10) timesteps, after which the input is changed (while the network state is left unaltered). The first half of this time interval allows for the network to adapt to the new input configuration, while in the second half we measure the average state of the output node $\langle \sigma_{\text{output}} \rangle$ and compute the value of the network fitness, $F$.

### Network mutations.

Having established the fitness of a given network, $M$, we now allow it to "mutate" by rewiring some existing links—thus preserving its overall connectivity $K$—and generate a slightly modified network $M'$. The technicalities of how the mutation process is implemented are deferred to the Methods section.

### Network evolution and convergence.

The network with the largest fitness value, between $M$ and its mutated counterpart $M'$, is selected (while the original one is kept if the two fitnesses coincide). This mutation and selection process defines an evolutionary time step (to be distinguished from a time step of the dynamics; there is a factor $t_{\text{mut}}$: $t$ between both). The evolutionary process is iterated until $F$ reaches its maximal possible value $F = 1$. Observe, however, that as the $I$ inputs are randomly chosen at each evolutionary step, observation of $F = 1$ at a given step does not necessarily imply $F = 1$ at successive time. Therefore, in order to impose that the network robustly "learns" the computational task, we continue to measure its fitness, when exposing it to a much large number of randomly chosen inputs ($100I$, instead of just $I$ as in the fitness-computation Eq. (2)); if $F = 1$ all across this long checking time window, the network is classified as having learned. Otherwise, the mutation/selection process is restarted until an optimally performing network is found. The final number of evolutionary steps required to reach an optimal network is called convergence time, $T$.

### Ensemble averages.

Keeping fixed specific values of the network size $N$ and connectivity $K$, the previous evolutionary process is iterated a large number of times (typically from $10^3$ to $5 \cdot 10^5$) giving rise to an ensemble of trained networks. The ensemble averaged convergence time, $\overline{T} = \overline{T}(N, K)$, is a proxy for the network performance: the best network ensemble is the one with the smallest $\overline{T}$. In this set of networks—once they have been trained—we also measured the ensemble-average of the branching parameter, $B$. In the approach of Goudarzi et al., $K$ is allowed to change during the evolutionary process; thus the fastest learning networks are selected for; instead, we explore different fixed-$K$ ensembles and determine a posteriori which is the optimal one. Both approaches are obviously equivalent to determine the optimal connectivity $K$.

### Dynamics under noisy conditions.

To investigate the effect of fluctuations in the system dynamics, we allow the dynamics to be exposed to noise. In particular, we consider that either (i) with a small probability, $\eta$, nodes can invert their state every time they are updated (accounting for errors/fluctuations in gene expression levels) or (ii) with some small probability, $\xi$, (which is proportional to the network connectivity) the network topology experiences a mutation process at each evolutionary step, and the mutated network is kept/selected regardless of its fitness value (this describes physical damage in the network produced, for example, by the lack or...
excess of some regulatory factors). For the sake of simplicity, we refer to the first possibility as “dynamical” noise and to the second one as “structural” noise.

Results

Convergence times and dynamical phases of learning networks. Even in the absence of explicit noise sources, the dynamics based on asynchronous updating—which is the one we adopt here—has a stochastic component (i.e. nodes are updated in a random order), which could be more adequate to represent real genetic networks than synchronously updated RBNs as it avoids spurious effects associated with perfectly synchronous updating

We consider a complex computational task—the odd-even classifier—and analyze networks of variable N and K. We let them evolve to learn this task and measure the average convergence time, \( T \), to do so. Results are shown in Fig. 2 for sizes from \( N = 6 \) to \( N = 64 \) as a function of the network connectivity \( K \) (from \( K = 0.5 \) to \( K = 3.5 \)). First of all (upper Fig. 2B), observe that for all values of \( N \), \( T \) exhibits a characteristic (pseudo)parabolic shape with a minimum at some optimal connectivity value, \( K_T \), at which networks learn the computational task in the fastest possible way. It is important to stress that networks with connectivities other than \( K_T \) also learn, even if after longer evolutionary times. In Fig. 2A the same data are represented, but rescaling \( T \) for each \( N \) with its minimum, \( T_{\min}(N) \) (this is done to help the eye to compare the location of the different minima). In both cases, there is a convergence toward a value close to 2 in the large N limit (blue squares) the red line is a guide to the eye and corresponds to a decay \( O(N^{-1}) \) toward 2. (D) \( \Delta = K_C - K_T \) plotted as a function of \( N \) showing explicitly that the distance to criticality diminishes with network size; i.e. the larger the network the closer to criticality the fastest learning networks.

Figure 2. (A) Averaged convergence time, \( T \), divided by its minimum value \( T_{\min} \) plotted as a function of the network connectivity \( K \) for different network sizes \( N \) (solid lines); ensemble averages performed over up to \( 5 \times 10^5 \) network realizations. Similarly, the non-normalized convergence times are plotted in (B). Discontinuous lines in (A) represent the value of the branching parameter, \( B \) as measured in the network after the learning process is completed; dashed-dotted lines stand for \( B \) averaged after perturbing only input nodes, while dotted lines have been obtained after perturbing nodes in the network core. Note that as \( T/T_{\min} \) and \( B \) are both dimensionless quantities, they have been plotted in the same scale; the same color code has been used for all curves. (C) Scaling of the connectivity at which the minimum \( T \) is obtained, \( K_T \), as a function of \( N \) (blue squares), plotted together with the position of the critical point \( K_C \) as estimated from the condition \( B \approx 1 \) (orange diamonds). In both cases, there is a convergence toward a value close to 2 in the large \( N \) limit (blue squares) the red line is a guide to the eye and corresponds to a decay \( O(N^{-1}) \) toward 2. (D) \( \Delta = K_C - K_T \) plotted as a function of \( N \) showing explicitly that the distance to criticality diminishes with network size; i.e. the larger the network the closer to criticality the fastest learning networks.
more responsive (larger $B$) to input perturbations than to changes in the core. For example, networks with connectivity $K = 2$ are supercritical to input perturbations (fostering network sensitivity to external changes) and subcritical for core perturbations (as required for a robust convergence to the attractor/output).

To obtain the overall branching parameter $B$ (given $N$ and $K$) – for all nodes in the network – we need to average these two contributions (weighted with $n_{\text{input}} = 3$ and $N - 3$ nodes, respectively). For these averaged curves (which, for the sake of clarity, are not explicitly shown in Fig. 2A) the crossing $B = B_1$ indicates overall critical dynamics, and corresponds to a critical connectivity $K_C$.

$K_C$ turns out to be larger than $K = 2$ and shifts toward lower connectivity values as $N$ grows; indeed, its distance to $K = 2$ decreases with $N$ (see Fig. 2C; orange diamonds), suggesting that learning networks have critical connectivity $K \approx 2$ (within our resolution) in the infinite size limit, as happens with random networks.

Moreover, we have measured the difference $\Delta = K_C - K_T$ to gauge how far optimal connectivities (in the sense of achieving the fastest possible learning) are from critical dynamics (in the sense of the branching parameter as close as possible to 1). As shown in Fig. 2D (magenta circles), $\Delta$ decreases monotonically upon increasing $N$, indicating that – for sufficiently large networks – the optimal connectivity is as close to criticality as desired, but for any finite size they are slightly subcritical ($\Delta > 0$). Thus optimal learning occurs for slightly subcritical networks, arbitrarily close to criticality for sufficiently large system sizes.

Figure 3A illustrates results for other, less complex (see above) computational tasks. As before, there is a well-defined minimum for $T$ in all cases, but these times are significantly shorter for lesser complex tasks (about two orders of magnitude less for a fixed size). Observe also that for the simplest, R51 rule, $T$ hardly depends on $K$ (Fig. 3B), indicating that, as the task complexity decreases $K$ progressively becomes a lesser relevant parameter. Observe also (Fig. 3D) that the distance of optimal networks to criticality, $\Delta$, decreases with increasing network complexity. Therefore, it is reasonable to conjecture that for more complex tasks than the ones we considered (e.g., involving larger values of $n_{\text{input}}$), the benefits derived from operating at optimality/criticality are progressively more crucial.

Finally, we also scrutinized the network topology (in-degree distribution) after learning and, interestingly, we did not detect significant structural changes, as the overall network skeleton was in all cases very close to a random network.

Summing up, in order to achieve the fastest possible learning of complex tasks, RBNs with a connectivity such that their dynamics turns out to be critical (or slightly subcritical for finite sizes) are the best possible option. The larger the network size and the more complex the task, the more evolutionarily favourable to be close to criticality.

Learning under noisy conditions. Dynamical noise. Figure 4 is analogous to Fig. 2 but has been obtained in the presence of dynamical noise, $\eta \neq 0$ (results for $\eta = 0$ are also plotted for the sake of comparison); observe that we present results for a fixed size $N = 16$ and variable noise strengths (from $\eta = 10^{-5}$ to $\eta = 10^{-3}$). It is
increased (see also Fig. 5C where $K_T$ Fig. 5B). In these cases, the optimal connectivity is observed to shift toward lower values of $K$ as the noise level is increased (see also Fig. 5C where $K_T$). For intermediate noise amplitudes, the larger $\xi$ the longer the learning process takes (see Fig. 4C, are always close to $K = 2$ (for the considered sizes). Comparing these values with the optimal connectivities for learning, i.e. measuring, $\Delta = K_c - K$, one observes (see Fig. 4D) that $\Delta$ increases monotonically with $\eta$. This occurs for the different system sizes we studied allowing us to conclude that under noise conditions, it takes longer to learn, and the larger the dynamical-noise strength the more subcritical the optimal networks.

**Structural noise.** Figure 5 shows results analogous to those in Fig. 4. Also in this case we present results for a fixed size $N = 16$ and variable noise strengths (from $\xi = 10^{-3}$ to $\xi = 10^{-2}$). In parallel with the site-noise case, there is a noise intensity threshold above which the mutation probability is exceedingly high for the networks to learn, while for too small strengths, the same results as for $\xi = 0$ are observed within the operational checking time windows we have. For intermediate noise amplitudes, the larger $\xi$ the longer the learning process takes (see Fig. 5B). In these cases, the optimal connectivity is observed to shift toward lower values of $K$ as the noise level is increased (see also Fig. 5C where $K_c$ is plot as a function of $\xi$). Also, as above, the branching parameter, $B$ (measured keeping a fixed network structure) does not have a strong dependence on $\xi$ (Fig. 5A). The associated critical point $K_c$ is slightly above $K = 2$ for small noises, and moves progressively to smaller connectivity values as $\xi$ grows. Also, as in the previous case, $\Delta$ increases monotonically with $\eta$, so that, as above, we can safely conclude that, in general, the larger the structural noise strength the more subcritical the optimal networks.

Summing up, we conclude that while in the case of noiseless dynamics the optimal solution—to achieve the fastest possible learning—is obtained at connectivities for which the network is about critical (actually slightly
subcritical, but closer and closer to criticality as the network size and/or the complexity of the task are increased), the situation is different in the presence of additional stochasticity, be it dynamical or structural noise. Under noisy conditions, the optimal solutions lie clearly well within the ordered/subcritical phase. A straightforward interpretation of this result is that the network dynamics needs to compensate for the excess of noise, and does so by reducing its internal level of uncertainty, i.e. by shifting deep into the ordered/subcritical phase.

Empirical networks. We have collected a set of empirical data from the literature and compiled a set of real directed networks. This includes public empirical datasets with biological genetic regulatory networks and networks of metabolic interactions. Specific examples of networks collected from the literature are the metabolic networks of *Chlamydomonas reinhardtii* (*K* = 2.05) and *Bacillus subtilis* (*K* = 1.03), and the gene regulatory networks of *Escherichia coli* (*K* = 1.24, *K* = 2.32), *Arabidopsis thaliana* (*K* = 2.755), *Mycobacterium tuberculosis* (*K* = 2.05, *K* = 1.98), *Pseudomonas aeruginosa* (*K* = 1.48) and *Saccharomyces cerevisiae* (*K* = 1.85).

Figure 6 presents a scatter plot of all networks in our dataset, representing the averaged connectivity *K* and network size *N* of each one. As it can be seen, the averaged connectivity of this dataset is well below the value *K* = 2, the critical connectivity for large random networks, suggesting that they could operate in subcritical regimes. It is noteworthy that it has been suggested that some empirical networks with high connectivity values (such as some of the outliers in Fig. 6) might result from systematic errors in correlation analyses (giving rise to false positives). Being more precise –given the absence of knowledge on dynamical aspects of the specific dynamics of each empirical network– it is not possible to properly ascertain the dynamical state (critical or not) of each of them.

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Conclusions and Discussion

The hypothesis that living systems may operate in the vicinity of critical points of their internal dynamics has inspired and tantalized scientists for some time. In particular, it has been claimed that genetic regulatory networks might operate close to criticality, achieving in this way an optimal balance between sensitivity to signals and stability to noise, and/or between adaptability and robustness on large evolutionary scales. A few works have recently explored different mechanisms allowing for networks to self-organize or evolve to critical or quasi-critical dynamics.

Here –inspired by the set up proposed by Goudarzi et al.31—we have shown that random Boolean network models that are trained to perform a given computational task, can learn it much faster if they have a connectivity $K$ such that their dynamics turns out to be close to criticality, as defined by a marginal averaged propagation of perturbations. This does not mean that networks far from criticality cannot learn; indeed they do, but it takes much longer to do so. Two important differences between the present work and previous ones are as follows. First, we work with networks with constant connectivity, i.e. the allowed mutations keep $K$ constant, while in previous work there was no such constraint31. This difference implies that our evolutionary process does not converge to the optimal connectivity for fast learning, $K_c$; by studying the constant-connectivity ensemble, we are able to put forward that learning (not necessarily in the fastest possible way) is compatible with rather diverse connectivity patterns and, thus, with the network being critical, subcritical or supercritical. The second important difference is that we implement a stochastic updating scheme, which introduces stochasticity in the dynamics; we find, however, that results are mostly insensitive to this change. Moreover, we have seen that in all cases, the distance to criticality of the optimal-connectivity networks diminishes monotonically upon enlarging system size and upon enlarging the task complexity. Indeed, very simple tasks, establishing simple relationships between (a few) inputs...
new link corresponding to input \( i \); the outputs in rows 0, 1, 4, and 5 can be flipped with probability \( p = 0.25 \); (b) Addition of a new link corresponding to input \( i \) (\( K_i \) of the node increases from 2 to 3): outputs for rows 4, 5, 6, and 7 are randomly chosen (represented as □).

| \( \sigma_{i_0} \) | \( \sigma_{i_1} \) | \( \sigma_{i_2} \) | \( \sigma_{\text{out}} \) | \( \sigma_{i_0} \) | \( \sigma_{i_1} \) | \( \sigma_{i_2} \) | \( \sigma_{\text{out}} \) |
|---|---|---|---|---|---|---|---|
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 1 | 1 | 1 | 0 | 1 |
| \( \times 2 \) | 0 | 1 | 0 | 2 | 0 | 1 | 0 |
| \( \times 3 \) | 0 | 1 | 1 | 0 | 3 | 0 | 1 | 0 |
| 4 | 1 | 0 | 0 | 4 | 1 | 0 | 0 |
| 5 | 1 | 0 | 1 | 5 | 1 | 0 | 1 |
| \( \times 6 \) | 1 | 1 | 0 | 6 | 1 | 1 | 0 |
| \( \times 7 \) | 1 | 1 | 1 | 7 | 1 | 1 | □ |

(a) (b)

Table 1. Examples of the modification of Boolean functions—initially with 3 inputs and hence 2\(^3\) possible input configurations—after the addition or removal of an input node. (a) Link \( i \), is removed (the connectivity \( K_i \) of the node decreases from 3 to 2) the rows 2, 3, 6, and 7 (corresponding to \( \sigma_{i_2} = 1 \)) are canceled out (marked with \( \times \)); the outputs in rows 0, 1, 4, and 5 can be flipped with probability \( p = 0.25 \); (b) Addition of a new link corresponding to input \( i \) (\( K_i \) of the node increases from 2 to 3): outputs for rows 4, 5, 6, and 7 are randomly chosen (represented as □).

and the output, can be readily learned by networks in the ordered/subcritical regime, where such a direct correspondence can be robustly realized. On the other hand, complex tasks, in which the output is sensitive to many different possible changes in the input nodes, require of much larger responsiveness/susceptibility, and thus, shift the network optimal connectivity toward larger values, closer and closer to criticality. In any case, we do not find under any circumstances the optimal connectivity to lie within the disordered/supercritical regime; it seems as if the requirement to learn a task was incompatible with the network being disordered.

Biological systems must have homeostasis, i.e. the capacity to maintain their internal conditions even in the presence of fluctuations and noise, be it internal or external. In the second part of our study we posed ourselves the question of how do these results depend upon the explicit introduction of noise. To this end, we have introduced more extreme forms of noise, be it dynamical or structural, within the same RBN model. Dynamical noise allows network nodes to invert their dynamical state with a small probability each time they are updated, introducing perturbations that can potentially propagate through the system, compromising the network performance. Similarly, structural noise, implying that the network topology itself is exposed to random changes with some small probability, also producing potential damage in the learned patterns. Both of these noise sources have clear correspondence with stochastic effects in real biological networks. In both cases, there is a threshold in noise strength above which networks do not learn the computational task in a reliable and robust way; i.e. they end up being plagued with errors, hindering network learning. Such thresholds clearly depend on the criterion imposed to declare that networks have learned; put differently, if the time in which one checks for network robustness are increased, i.e. if the criterion becomes more stringent, the noise-strength thresholds diminish. Remarkably, in both of the cases, dynamical and structural noise, we find that the optimal connectivity to achieve the fastest possible learning lies deep-inside the subcritical region, far away from criticality, and the distance to criticality increases upon enlarging the noise strength and does not diminish upon increasing the system size (as it happens in the absence of explicit noise).

Our results suggest that real biological networks, in order to perform the complex tasks required for information processing and survival in a noisy world, should operate in sub-critical regimes rather than in critical ones as it has been argued. As a matter of fact, the collection of empirical (genetic and metabolic) networks that we have compiled from the recent literature shows a rather sparse averaged connectivity in most cases, with only a few outlier networks. If the dynamics underlying these networks could be modeled by random Boolean functions, one could safely conclude that they are typically subcritical. However, in most cases, the dynamics remains mostly unknown, and a clear cut conclusion about the dynamical state of each specific network instance cannot be derived. To fill this gap, recent analyses have employed high throughput data from hundreds of microarray experiments to infer regulatory interactions among genes. This type of approach leads to more detailed information on dynamical aspects (e.g. switching on a given gene it is possible to follow the cascade of modifications it generates through the whole network). The resulting data, implemented into Boolean models, seem to support the hypothesis that regulatory networks for a number of species (Saccharomyces cerevisiae, Escherichia coli, etc) are close to criticality\(^{25,26}\), but some other analyses leave the door open for the networks to operate in an ordered/subcritical phase\(^{25,26}\). Therefore, given the present state of affairs, one can only conclude that more accurate and extensive experimental approaches (including, in particular, more accurate direct measurements of the bias \( p \)) would be extremely valuable to shed further light on this fascinating problem.

An important observation to be made is that the tasks we have employed to be learned are relatively simple (as they only involve a maximum of 3 input nodes and a single readout). Thus, one can wonders what would happen if a more extensive use of the network potentiality was necessary (by employing for instance, two or more tasks simultaneously, and/or involving a much larger number of inputs in each single task). Under the light of our results for the noiseless case –where we found that upon considering far more complex tasks, involving many more input and output nodes, the dynamics becomes progressively more critical– it would not be surprising that if one could analyze much more complex tasks –as the ones probably controlling real biological networks– the dynamics could become closer to criticality even in the presence of noise. Furthermore, in such more complex
cases, one should also relax the criterion to declare that networks have learned, and look for “fuzzy” types of learning (i.e. accept networks with fitnesses slightly smaller than one). The combination of much more complex rules together with less rigid criteria for learning, could very likely shift the optimal solutions toward more critical states. A detailed analysis of these issues is left as an open challenge for future work.

It is also noteworthy that, even if network topology is known to play a very important role in the outcome of RBNs, here we have focused mostly on random Erdős-Rényi networks and left the analysis of important topological features of empirical networks –such as scale-free connectivity distributions, and hierarchical and modular organization– for future work. These aspects might also play an important role in determining the network dynamical state. Finally, we also plan to extend the studies beyond the limit of the Boolean approach and to implement more complex and biologically realistic tasks. Hence, our summary is that the criticality hypothesis remains as a valid and fascinating possibility, but that it needs to be critically evaluated under each set of specific circumstances, avoiding making exceedingly general claims.

Methods

Network mutations.

1. Given a original network, $M$, we perform a rewiring, which consists in choosing a link (say from node $i$ to node $j$), removing it, and introducing a new one (from $i$ to $j'$) assuming this one did not exist before (and keeping the topological constraints described above).
2. This change of the network topology requires some modifications in the random Boolean functions $f_i$ and $f_j$ (see Table 1). For $f_i$ one needs to eliminate the input $\sigma_j$ thus $f_i$ changes from being a function of $K_m(j)$ arguments to a function of $K_m(j) - 1$. The new function coincides with the original one fixing $\sigma_j = 0$, i.e. for the case when the driving node $j$ was off. After this, each output in its table is changed with probability $1/4$, defining the “mutated” Boolean function. Similarly, for node $j'$ a new argument, $\sigma_j$, is introduced to the Boolean function $f_j$: all values for $\sigma_j = 1$ (‘on’ $i$ node) are assigned randomly, while for $\sigma_j = 0$ (the new input is off) we keep the pre-existing Boolean-function values.
3. This whole rewiring process is performed the first time with prob. one; after that a second rewiring is attempted with prob. 1/2; if it occurs, then a third one happens with prob. 1/3 and so on, giving rise to a mutated network, $M'$. This sequential process allows for the possibility of large mutations, involving many rewirings.
4. Observe that these mutations keep the out degree sequence, as well as the overall connectivity $K$ fixed, so it can be understood as a sort of “micro-canonical ensemble” of Boolean networks.

Assessing network criticality. We employ the standard method of plotting the Derrida curve in order to determine the dynamical phase of any specific RBN –specified by its topology and the set of its Boolean functions– and assess how far it operates from criticality. The method is based in damage spreading dynamics and involves the next steps: (1) take a network $M$ in one specific state, and a copy of it $M'$ in which a single randomly chosen node has changed its state, (2) compute the Hamming distance, $H_f$, between these two networks after one time step ($t = 1$; in the asynchronous case nodes are updated following the same random order in both networks), (3) average such a Hamming distance by considering all the possible nodes in the network that can host the initial one-node perturbation, (4) average the previous result over network states. We define the branching parameter $B$, as the averaged $H_f$ after perturbing the different nodes in the network (in some cases, we present results for perturbations only at input/core nodes). If $B < 1$ perturbations shrink on average and the network is said to be subcritical (or in the ordered phase), while if $B > 1$ perturbations proliferate and grow on average and the network is supercritical (chaotic or disordered phase). Finally, in the intermediate case, $B = 1$, in which perturbations propagate marginally, the network is critical.

Observe that in networks with some fixed input and output nodes, we can measure $B$ in different ways, depending on whether we flip input nodes or not and on whether we compute the Hamming distance in the whole network or just in the core (excluding input nodes); therefore the concept of criticality might refer to just the core or to the full network. Finally, in order to determine the critical regime of an ensemble of networks –and not just an individual one– it is necessary to measure the ensemble average, $\overline{B}$, of $B$.

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Acknowledgements
We acknowledge the Spanish-MINECO grant FIS2013-43201-P (FEDER funds) for financial support. We thank P. Moretti for very useful comments.

Author Contributions
J.H. and M.A.M. conceived the project, P.V. and J.R.-F. performed the numerical simulations and prepared the figures. P.V., J.H. and M.A.M. wrote the main manuscript text. All authors reviewed the manuscript.

Additional Information
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Villegas, P. et al. Intrinsic noise and deviations from criticality in Boolean gene-regulatory networks. Sci. Rep. 6, 34743; doi: 10.1038/srep34743 (2016).

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