



# SORN: a self-organizing recurrent neural network

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Understanding the dynamics of recurrent neural networks is crucial for explaining how the brain processes information. In the neocortex, a range of different plasticity mechanisms are shaping recurrent networks into effective information processing circuits that learn appropriate representations for time-varying sensory stimuli. However, it has been difficult to mimic these abilities in artificial neural network models. Here we introduce SORN, a self-organizing recurrent network. It combines three distinct forms of local plasticity to learn spatio-temporal patterns in its input while maintaining its dynamics in a healthy regime suitable for learning. The SORN learns to encode information in the form of trajectories through its high-dimensional state space reminiscent of recent biological findings on cortical coding. All three forms of plasticity are shown to be essential for the network's success.

**Keywords:** synaptic plasticity, intrinsic plasticity, recurrent neural networks, reservoir computing, time series prediction

## INTRODUCTION

The mammalian neocortex is the seat of our highest cognitive functions. Despite much effort, a detailed characterization of its complex neural dynamics and an understanding of the relationship between these dynamics and cognitive processes remain elusive. Cortical networks present an astonishing ability to learn and adapt via a number of plasticity mechanisms which affect both their synaptic and neuronal properties. These mechanisms allow the recurrent networks in the cortex to learn representations of complex spatio-temporal stimuli. Interestingly, neuronal responses are highly dynamic in time (even when the stimulus is static) (Broome et al., 2006) and contain a rich amount of information about past events (Brosch and Schreiner, 2000; Bartlett and Wang, 2005; Broome et al., 2006; Nikolic et al., 2006).

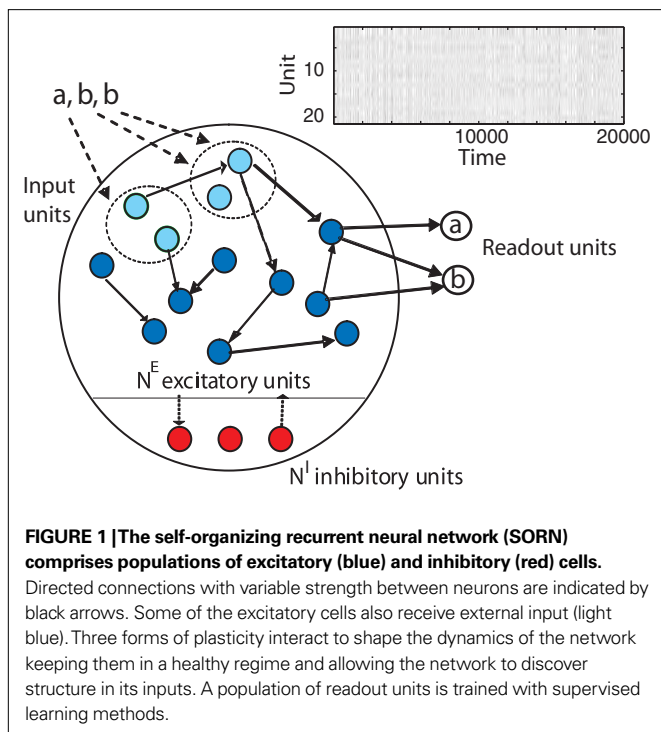
But mimicking these features in artificial neural networks has proven to be very difficult. The first models that could address temporal tasks have incorporated in their structure an explicit representation of time (Elman and Zipser, 1988). Recurrent neural networks (RNNs) were the first models to represent time implicitly, through the effect that is has on processing (Hopfield, 1982; Elman, 1990). In the recently developed framework of 'reservoir' computing (Jaeger, 2001; Maass et al., 2002), a randomly structured RNN non-linearly transforms a time varying input signal into a spatial representation. At each time step, the network combines the incoming stimuli with a volley of recurrent signals containing a memory trace of recent inputs. For a network with  $N$  neurons, the resulting activation vector at a discrete time  $t$ , can be regarded as a point in a  $N$ -dimensional space. Over time, these points form a pathway through the state space also referred to as a *neural trajectory*. A separate read-out layer is trained, with supervised learning techniques, to map different parts of the state space to desired outputs. In real cortical networks, experimental evidence has shown that different stimuli elicit different trajectories while for a given stimuli the activity patterns evolve in time in a reproducible manner (Broome et al., 2006; Churchland et al., 2007). Furthermore, identical trials can present a high response variability, but the resulting trajectories are not dominated by noise

(Mazor and Laurent, 2005; Broome et al., 2006; Churchland et al., 2007). Reservoir networks do not require classical attractor states and are compatible with the view that cortical computation is based on transient dynamics (Mazor and Laurent, 2005; Durstewitz and Deco, 2008; Rabinovich et al., 2008). It has been shown that neural systems may exhibit transients of long durations which carry more information about the stimulus than the steady states towards which the activity evolves (Mazor and Laurent, 2005).

Attempts to endow RNNs with unsupervised learning abilities by incorporating biologically plausible local plasticity mechanisms such as spike-timing-dependent plasticity (STDP) (Markram et al., 1997; Bi and Poo, 1998) have remained largely unsuccessful (and often unpublished). The problem is most difficult, because structural changes induced by plasticity will impact the network's dynamics giving rise to altered firing patterns between neurons. These altered firing patterns can further induce changes in connectivity through the plasticity mechanisms and so forth. Understanding and controlling the ensuing self-organization of network structure and dynamics as a function of the network's inputs is a formidable challenge.

The key to the brain's solution to this problem may be the synergistic combination of multiple forms of neuronal plasticity. There has been extensive evidence that synaptic learning is accompanied by homeostatic mechanisms. Synaptic scaling regulates the total synaptic drive received by a neuron but maintains the relative strength of synapses established during learning (Turrigiano et al., 1998). At the same time, intrinsic plasticity (IP) was shown to directly regulate neuronal excitability (Desai et al., 1999; Zhang and Linden, 2003). In a RNN, IP induced robust homeostatic effects on the network dynamics (Steil, 2007; Schrauwen et al., 2008). But there is only little work combining several forms of plasticity in RNNs (Lazar et al., 2007).

In the following, we present a RNN of threshold units combining three different forms of plasticity that learns to efficiently represent and "understand" the spatio-temporal patterns in its input. The SORN model (self-organizing recurrent network) consists



of a population of excitatory cells and a smaller population of inhibitory cells (Figure 1). The connectivity among excitatory units is sparse and subject to a simple STDP rule. Additionally, synaptic normalization (SN) keeps the sum of an excitatory neuron's afferent weights constant, while IP regulates a neuron's firing threshold to maintain a low average activity level. The network receives input sequences composed of different symbols and learns the structure embedded in these sequences in an unsupervised manner. The three types of plasticity mechanisms induce changes in network dynamics which we assess via hierarchical clustering and principal component analysis (PCA). In addition, we train a separate readout layer with supervised learning techniques and compare the performance of our network with that of fixed random networks constructed in the spirit of reservoir computing.

We show that only the combination of all three types of plasticity allows the network to (a) learn to effectively represent the spatio-temporal structure of its inputs, (b) maintain 'healthy' dynamics<sup>1</sup> that make efficient use of all the network's resources, and (c) perform much better on prediction tasks compared to random networks without plasticity. Furthermore, the network dynamics are consistent with a range of neurophysiological findings.

## MATERIALS AND METHODS

### THE SORN MODEL

#### Network definition

We consider a network with  $N^E$  excitatory ( $E$ ) and  $N^I = 0.2 \times N^E$  inhibitory ( $I$ ) threshold units. Neurons are coupled through weighted synaptic connections, where  $W_{ij}$  is the connection strength from unit  $j$  to unit  $i$ , with  $i \neq j$ . All possible connections between the excitatory and inhibitory neuron populations are present

<sup>1</sup>Dynamics suitable for computation.

( $W^{IE}$  and  $W^{EI}$ ), while the excitatory–excitatory connections ( $W^{EE}$ ) are sparse and random with a mean number  $\lambda^W$  of incoming and outgoing connections per neuron. Direct connections between inhibitory units are not present. The weight strengths are drawn from the interval  $[0, 1]$  and subsequently normalized such that the incoming connections to a neuron sum up to a constant value:  $\sum_j W_{ij}^{IE} = 1$ ,  $\sum_j W_{ij}^{EI} = 1$  and  $\sum_j W_{ij}^{EE} = 1$ . Inputs are time series  $U(t)$  of different symbols (letters or digits). Each symbol is associated with a specific group of  $N^U$  input units which all receive a positive input drive ( $v^U = 1$ ) when that particular symbol is active.

The network state, at a discrete time  $t$ , is given by the binary vectors  $x(t) \in \{0, 1\}^{N^E}$  and  $y(t) \in \{0, 1\}^{N^I}$  corresponding to the activity of the excitatory and inhibitory units, respectively. The evolution of the network state is described by:

$$x_i(t+1) = \Theta \left( \sum_{j=1}^{N^E} W_{ij}^{EE}(t)x_j(t) - \sum_{k=1}^{N^I} W_{ik}^{EI}y_k(t) + v_i^U(t) - T_i^E(t) \right) \quad (1)$$

$$y_i(t+1) = \Theta \left( \sum_{j=1}^{N^E} W_{ij}^{IE}x_j(t) - T_i^I \right). \quad (2)$$

The  $T^E$  and  $T^I$  are threshold values for the excitatory and inhibitory units. They are initially drawn from a uniform distribution in the interval  $[0, T_{\max}^E]$  and  $[0, T_{\max}^I]$ , respectively. The heaviside step function  $\Theta(\cdot)$  constrains the activation of the network at time  $t$  to a binary representation: a neuron fires if the total drive it receives is greater than its threshold, otherwise it stays silent.

At each time step the activity of the network is determined both by the inputs  $v_i^U(t)$  and the propagation of the previously emitted spikes through the network. This recurrent drive received by unit  $i$  is given by:

$$R_i(t+1) = \sum_{j=1}^{N^E} W_{ij}^{EE}(t)x_j(t) - \sum_{k=1}^{N^I} W_{ik}^{EI}y_k(t) - T_i^E(t). \quad (3)$$

Based on this, we define a "pseudo state"  $x'(t)$  that only depends on the recurrent drive:

$$x'_i(t) = \Theta(R_i(t)); \quad (4)$$

This equation is identical to Eq. 1, but lacking the input drive  $v_i^U(t)$ . Most of our analysis focuses on the pseudo states  $x'(t)$  as the network's internal representation of previous inputs, although it may contain less information than  $R(t)$  due to the thresholding operation.

#### Plasticity mechanisms

The network relies on three forms of plasticity: STDP, synaptic scaling of the excitatory–excitatory connections, and IP regulating the thresholds of excitatory units.

Learning with STDP is constrained to the set of  $W^{EE}$  synapses. We use a simple model of STDP that strengthens the synaptic weight  $W_{ij}^{EE}$  by a fixed amount  $\eta_{\text{STDP}} = 0.001$  whenever unit  $i$  is active in the time step following activation of unit  $j$ . When unit  $i$  is active in the time step preceding activation of unit  $j$ ,  $W_{ij}^{EE}$  is weakened by the same amount:

$$\Delta W_{ij}^{EE}(t) = \eta_{\text{STDP}} (x_i(t)x_j(t-1) - x_i(t-1)x_j(t)); \quad (5)$$

STDP changes the synaptic strength in a temporally asymmetric “causal” fashion. The changes introduced by STDP can push the activity of the network to grow or shrink in an uncontrolled manner. To keep the activity balanced during learning we make use of additional homeostatic mechanisms that are sensitive to the total level of synaptic efficacy and the post-synaptic firing rate.

SN proportionally adjusts the values of incoming connections to a neuron so that they sum up to a constant value. Specifically, the  $W^{EE}$  connections are rescaled at every time step according to:

$$W_{ij}^{EE}(t) \leftarrow W_{ij}^{EE}(t) / \sum_j W_{ij}^{EE}(t). \quad (6)$$

This rule does not change the relative strengths of synapses established by STDP but regulates the total incoming drive a neuron receives.

An IP rule spreads the activity evenly across units, such that on average each excitatory neuron will fire with the same target rate  $H_{\text{IP}}$ . To this end, a unit that has just been active increases its threshold while an inactive unit lowers its threshold by a small amount:

$$T_i^E(t+1) = T_i^E(t) + \eta_{\text{IP}} (x_i(t) - H_{\text{IP}}), \quad (7)$$

where  $\eta_{\text{IP}} = 0.001$  is a small learning rate. We set the target rate to  $H_{\text{IP}} = 2 \times N^U / N^E$  in which the input spikes are approximately half of the total number of spikes. Other settings of  $H_{\text{IP}}$  do not necessarily lead to the desired improvements in prediction performance (see Appendix).

The implementation of the model described above and the simulations presented in Section “Results” were performed in Matlab.

## RESULTS

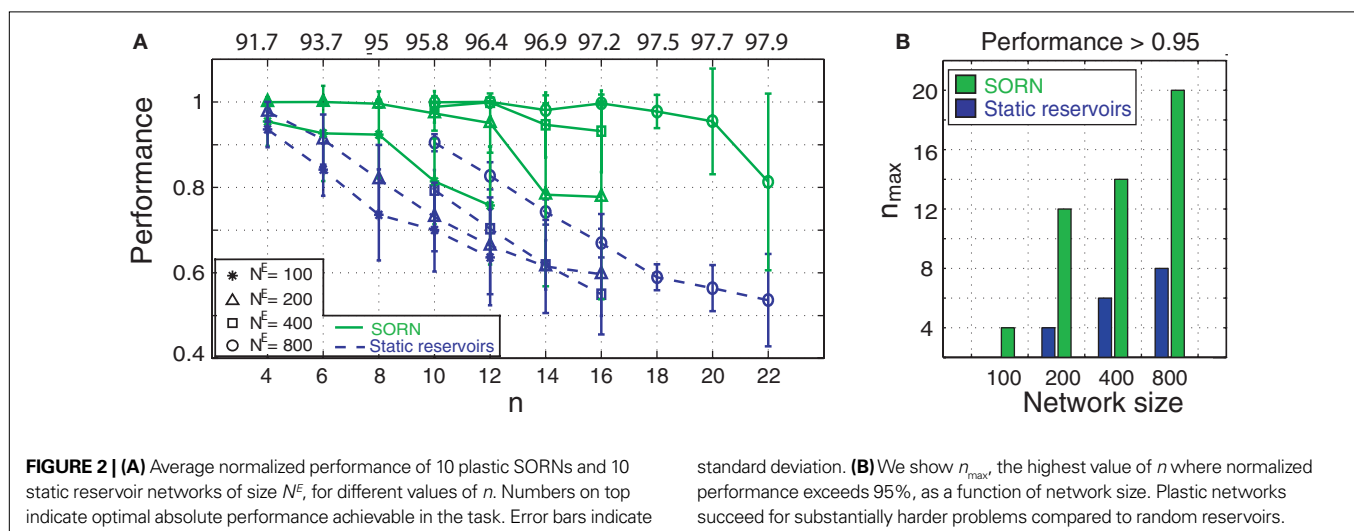
### SORNs OUTPERFORM STATIC RESERVOIRS

We demonstrate the SORN’s ability to learn spatio-temporal structure in its inputs with a “counting” task, especially designed to test the memory property of the reservoir. To this end, we construct input sequences  $U(t)$  as random alternations of two “words”

‘abbb...bc’ and ‘eddd...df’, composed of  $n + 2$  “letters”, with letters ‘b’ and ‘d’ repeating  $n$  times. In order to predict the next input letter correctly, the network has to learn to “count” how many repetitions of letters ‘b’ and ‘d’ it has already seen. Increasing  $n$  raises the difficulty of the task. We compare SORNs with all three forms of plasticity to static networks without plasticity. Networks of different sizes  $N^E$  have their initial parameters set to  $N^U = 5\% \times N^E$ ,  $T_{\text{max}}^E = 0.5$ ,  $T_{\text{max}}^I = 1$  and  $\lambda^W = 10$ . For small static reservoirs, the parameters are tuned such that their dynamics is critical and the networks’ firing rate is similar to the rate exhibited by SORNs structured by plasticity (see Supplementary Material and Section “Occluder Task”). It has been argued that a tuning of network dynamics to criticality should bring the performance of static reservoir networks close to the optimal performance (Bertschinger and Natschläger, 2004). To compute prediction performance, 5000 steps of network activity are simulated and a readout is trained in a supervised fashion to predict the next input  $[U(t)]$ , e.g., ‘a’, or ‘c’, or 5th repetition of ‘b’, etc., based on the network’s internal state  $[x'(t)]$  after presentation of the preceding letter  $[U(t-1)]$ . We use the Moore–Penrose pseudoinverse method that minimizes the squared difference between the output of the readout neurons and the target output value. The quality of the readout (the network performance) is assessed on a second sample of 5000 steps of activity using an independent input sequence.

The SORNs are exposed to the input sequences for 50,000 time steps. Then, all their weights and thresholds are frozen and a readout is trained in the same manner.

Since the input sequences are partly random – the order of letters within a word is fixed but the order of words is random – prediction performance is inherently limited. We define a normalized performance measure that obtains a score of 1 when the network always correctly predicts the next letter and its position within a word but is at chance level for guessing the first letter of the next word (either ‘a’ or ‘e’). **Figure 2** compares the performance of SORNs and static reservoir networks. For any given network size ( $N^E$ ) and any given task difficulty ( $n$ ), the plastic SORNs perform considerably better than their randomly structured counterparts (**Figure 2A**). For the same task difficulty  $n$ , larger networks perform better than smaller



networks. For a given network size the SORNs achieve a performance greater than 0.95 for much higher values of  $n$  compared to the static reservoirs (**Figure 2B**). A more detailed analysis of the network performance as a function of various initial parameter settings is given in the Appendix.

### SORNs LEARN EFFECTIVE INTERNAL REPRESENTATIONS

To better understand the reason underlying the performance advantage of SORNs over static reservoirs, we performed hierarchical clustering and PCA on the networks' internal representations.

We performed agglomerative hierarchical clustering of the networks' internal state representations ( $x'$ ). Each pattern of activity  $x'(t)$  is a point in a  $N^E$ -dimensional space. Agglomerative clustering starts by considering each of these points as centers of their own cluster. The distance between two clusters is computed as the Euclidean distance between their centers. Repeatedly, the two closest clusters are merged into a single cluster, until the entire data are collapsed.

In **Figures 3A,E** we present a snapshot of the last 20 clusters of agglomerative clustering, for an example network with  $N^E = 200$ ,  $N^U = 10$ ,  $T_{\max}^E = 0.5$ ,  $T_{\max}^I = 0.8$ ,  $\lambda = 10$  during a counting task with  $n = 8$ . In the case of randomly structured reservoir networks, the cluster structure of internal representations only weakly reflects the underlying input conditions (**Figure 3A**). Many of the emerging clusters combine network states resulting from distinct input conditions, i.e., the networks internal representation easily confuses, say, the 5th repetition of letter 'b' with its 6th repetition. In fact most clusters lump together as many as seven input conditions (**Figure 3B**). In contrast after 50,000 steps of plasticity, the SORN learns an internal representation that tends to map different input conditions on to distinct network states falling into separate clusters (**Figure 3E**). Here, each cluster will combine at most two different input conditions (**Figure 3F**). For a parallel with the performance tests from the previous section, the analysis was performed on 5000 steps of activity with frozen weights and thresholds but the network presents similar clustering properties in the presence of plasticity.

We also performed PCA on the internal network states. In the case of random networks a single input condition produces a cloud of network states that is substantially overlapping with those from other input states within the projection space of the first three PCs (**Figure 3C**). In contrast, the SORN develops an internal representation where an input conditions produces a tight cluster of network states that is well separated from those of other input conditions (**Figure 3G**). In particular, it learns to internally distinguish different states that have a very similar history of inputs, say, five vs. six repetitions of letter 'b'. This leads to more orderly and stereotyped trajectories through the network state space in the case of SORNs. This is in line with the greater amount of variance explained by the first few PCs in the SORNs compared to random networks (compare **Figures 3D,H**).

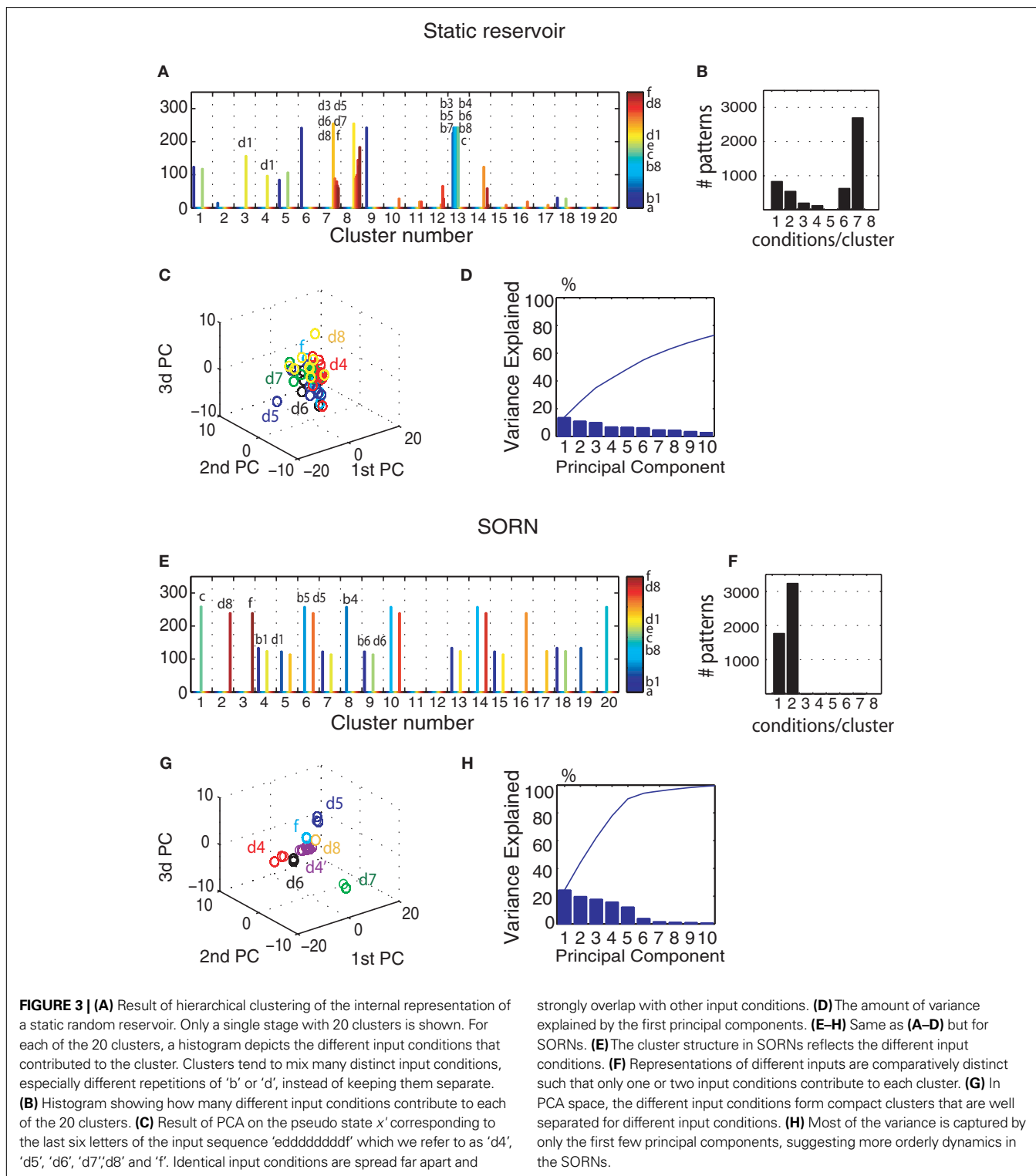
Interestingly, as long as plasticity is switched on, the internal representation will keep changing, i.e., the network does not converge. The internal representations of different input conditions tend to change gradually with time. For example, in **Figure 3G** the input condition d4 is shown after an additional 5000 time steps of plasticity, as d4'. To function properly, the network requires re-training of the readout as soon as the network's internal representations change significantly.

### OCCLUDE TASK

We demonstrate the ability of the SORN to learn effective representations on a second difficult task. Specifically, we consider an input sequence containing random alternations of the following four "words": '12345678', '87654321', '19999998', '89999991'. If we associate different spatial positions with the numbers 1–8, we can interpret these stimuli as left to right and right to left motion of an object along an axis. The symbol '9' can be interpreted as an occluder that obstructs the sight of the object at locations 2–7. This task is more difficult than the counting task in that several words share start and end letters and the repetitive symbol '9' is common in the last two sequences. The bidirectional quality of this stimuli might impose difficulties for the causal STDP rule. The interference of enforced synaptic pathways could decrease the prediction performance of SORNs. On the other hand, due to synaptic competition STDP might encourage one direction of motion and prune away the other. Our results suggest that both of these effects are avoided and SORNs present prediction advantages over random reservoirs.

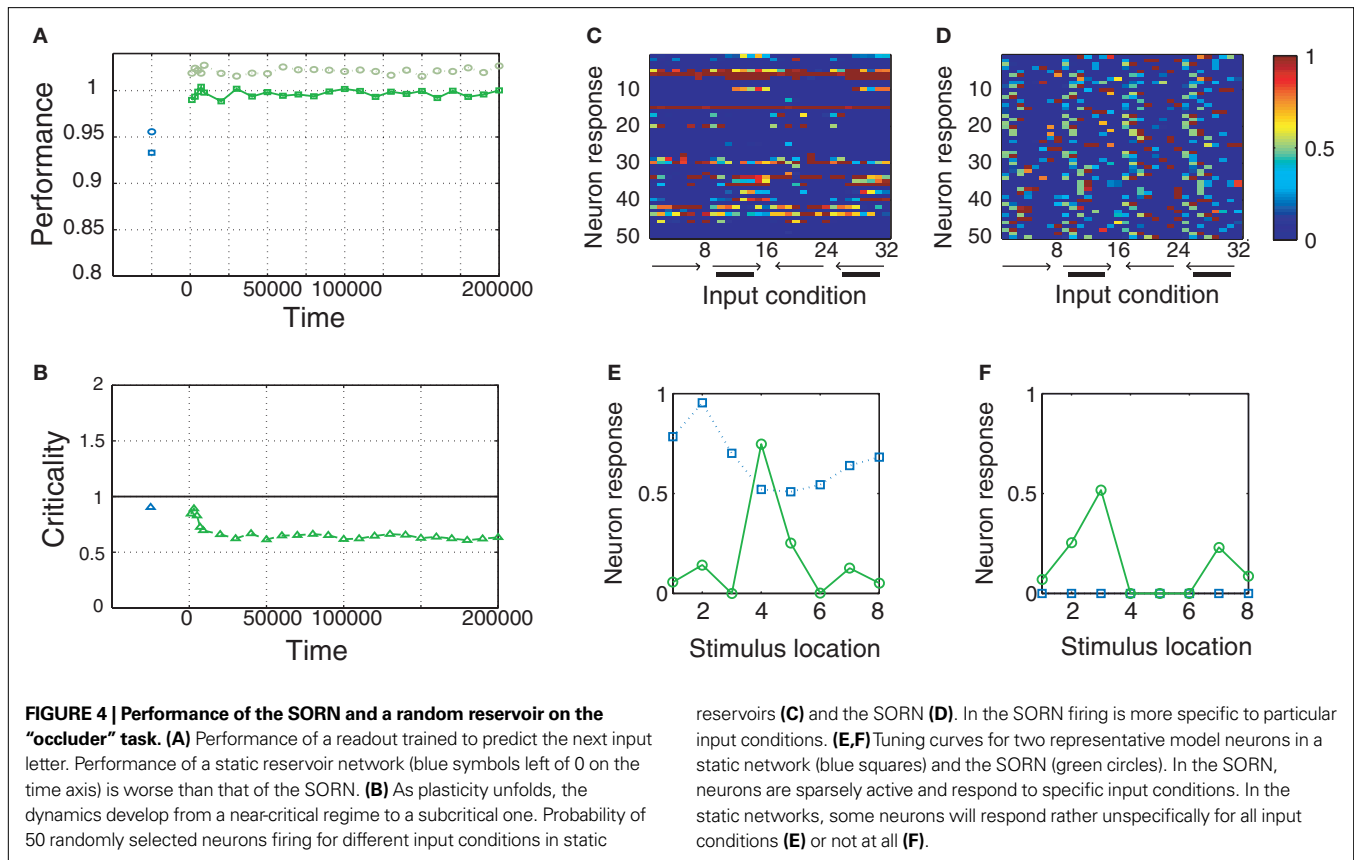
We choose a network with  $N^E = 200$ ,  $N^U = 15$ ,  $T_{\max}^E = 0.75$ ,  $T_{\max}^I = 1.4$ , and  $\lambda^W = 10$ . We run the SORN for 200,000 time steps and take snapshots of weights  $W$  and thresholds  $T$  at every 1000 steps of self-organization through plasticity. We evaluate each of these networks in terms of prediction performance for the one step prediction task. Similarly to the previous experiment, the performance drastically improves (**Figure 4A**) and is close to the theoretical optimum for all the different time intervals of self-organization with plasticity. We also assess the criticality of the network dynamics by performing a perturbation analysis. For every state  $x(t)$ , we perturb the activation of a randomly chosen excitatory neuron (from active to inactive or from inactive to active) creating an altered state  $\tilde{x}(t)$ . The Hamming distance between  $x(t)$  and its perturbed version  $\tilde{x}(t)$  is 1 [ $d(t) = 1$ ]. We calculate the successor states of  $x(t)$  and  $\tilde{x}(t)$  by applying Eq. 1 and obtain  $x(t+1)$  and  $\tilde{x}(t+1)$  with the Hamming distance  $d(t+1)$ . If the average distance  $\bar{d}(t+1) > 1$  the network amplifies perturbations and is in a supercritical regime. If  $\bar{d}(t+1) < 1$  the network has self-correcting properties and is in a subcritical dynamical regime. When  $\bar{d}(t+1) \approx 1$  the dynamics is said to be critical. Performing perturbation analysis, we find that the network dynamics changes from a critical regime, in the case of static reservoirs, to a subcritical regime for SORNs (**Figure 4B**). Interestingly, in the case of SORNs this corresponds to a higher network performance for prediction.

We also compare the tuning of the random reservoir network with the SORN after 50,000 steps of plasticity. For each of these two networks we consider 5000 time steps of network activity (in both cases without plasticity) and count the number of neuron responses corresponding to each of the 32 input conditions: left–right motion ('12345678'), left–right motion with occluder ('19999998'), right–left motion ('87654321') and right–left motion with occluder ('89999991'). For the random network we find that a number of neurons are silent and do not fire for any of the input conditions (**Figure 4C**). Also the neurons responding to the occluder sequences are not very selective in terms of either location or direction. In contrast, for the SORN all neurons take part in the activity and their responses are input specific (**Figure 4D**). We calculated "tuning curves" of two example neurons to illustrate this



point in more detail. To this end, we summed the neurons' responses for each of the eight locations of the visual space irrespective of motion direction or occluder presence. The neuron in **(Figure 4E)** responded unselectively to all eight locations before any plasticity (static reservoir case, blue squares) and after learning it has developed a clear preference for location 4 (SORN case, green circles).

The neuron in **(Figure 4F)** was silent in the initial network setup (static reservoir case). Through plasticity, it developed selectivity for locations 3 and 7 (SORN case). Interestingly, this selectivity is also specific with regard to motion direction. The neuron fires when a stimulus is at location 3 moving to the right, or when the stimulus is at location 7 moving to the left (not shown).



### HOMEOSTATIC PLASTICITY MECHANISMS ARE CRITICAL FOR MAINTAINING HEALTHY DYNAMICS

To better understand the role of the homeostatic plasticity mechanisms accompanying STDP-learning in SORNs, we compare SORNs with plastic reservoirs in which either the synaptic scaling or the IP is switched off. We consider networks receiving unstructured inputs, here in the form of random alternations of six symbols. Thus, there is no specific spatio-temporal structure in the inputs that could be learned during these experiments. The networks ( $N^E = 200$ ,  $N^U = 10$ ,  $T_{\max}^E = 0.5$ ,  $T_{\max}^I = 1$  and  $\lambda^W = 10$ ) are shaped in the presence of all three forms of plasticity for 50,000 steps.

The results are summarized in **Figure 5**. When SN is missing, the network dynamics develop into a regime with seizure-like synchronous bursts of activity (**Figure 5A**), even though the network is driven by random inputs. We compared the distribution of the total number of spikes per time step for 10 networks with and without SN (**Figure 5B**). In networks with SN the distribution is unimodal and centered at a low activity level. In contrast, networks without SN will show a bimodal distribution such that most units are either active or inactive at the same time. This is also expressed in the average correlation coefficient between neurons. In networks with SN the average correlation coefficient remains close to 0 with an average value of 0.025. For networks that lack SN the average correlation coefficient increases as a function of time to values beyond 0.8 within 50,000 steps of simulation, indicating a high degree of synchronization (**Figure 5C**).

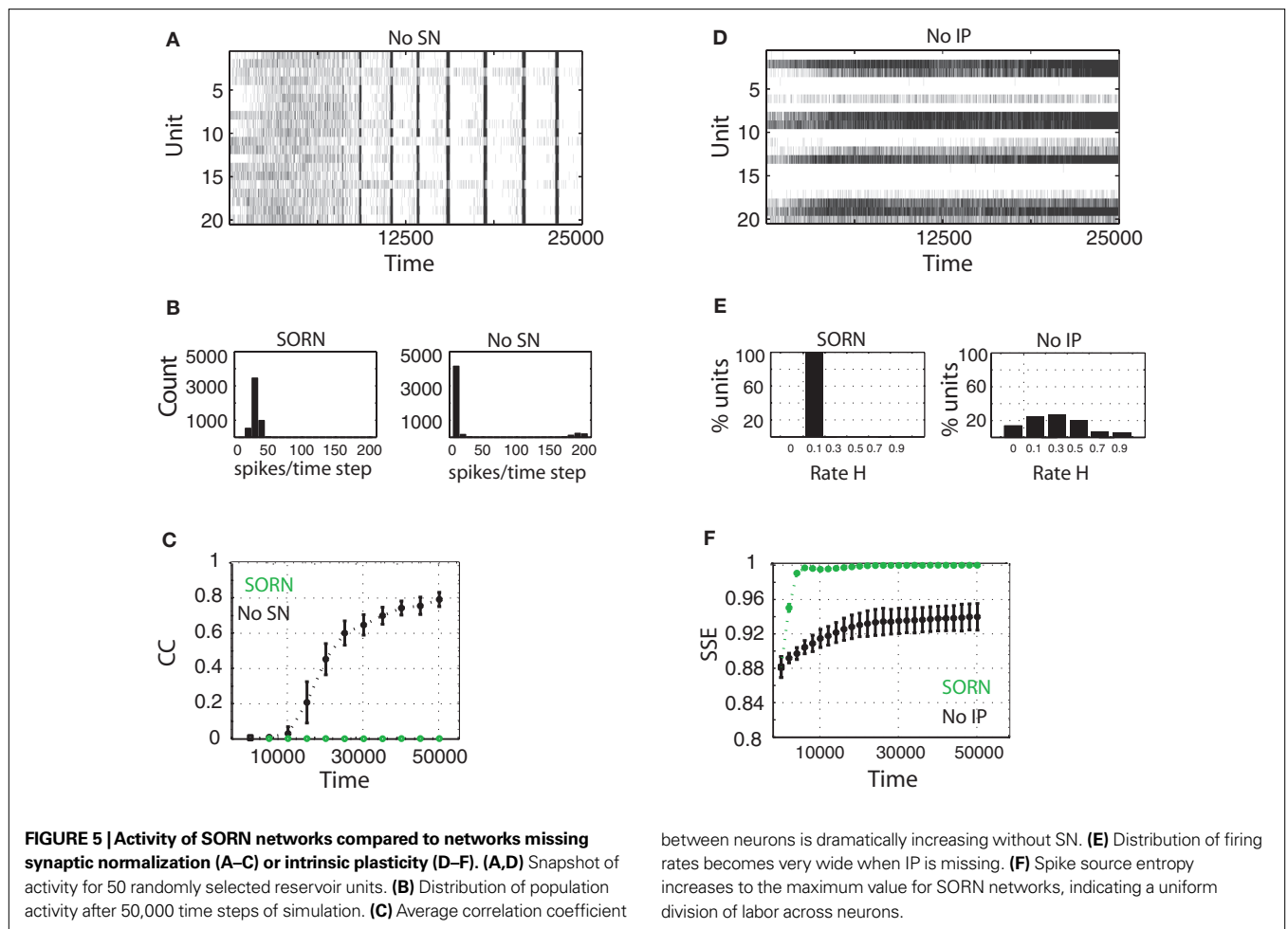
When IP is missing a number of neurons remain permanently silent, while others develop an unnaturally high activity (**Figure 5D**). We calculated the distribution of average firing rates for 10 such networks. In networks with IP, all excitatory units develop average firing rates close to the desired target rate, which was 0.1 in these experiments. Without IP, the distribution is more spread out with some units staying completely silent and others being active in almost every time step (**Figure 5E**). We quantified this effect by following the time evolution of the spike source entropy, which measures how much uncertainty there is about the origin of a spike in the network. It is defined as:

$$\text{SSE} = \left( \frac{\sum_{i=1..N^E} p_i \times \log_2 p_i}{\log_2(1/N^E)} \right), \quad (8)$$

where  $p_i$  is the probability that a spike is generated by the unit  $i$ . SSE achieves its maximum value of 1 if all units fire at the same rate ( $p_i \propto H_i$ , where  $H_i$  is the firing rate of neuron  $i$ ). SORNs show an abrupt increase in SSE to a value close to 1, which indicates identical rates across the neuronal population, compared to a smaller value of 0.94 for networks missing IP (**Figure 5F**). Due to IP, SORNs make efficient use of all the network's resources.

### DISCUSSION

Self-organizing recurrent networks are the substrate for neural information processing in the brain. Such networks are shaped by a wealth of plasticity mechanisms which affect synaptic as well as neuronal properties and operate over various time scales (from



seconds to days and beyond). Somehow these mechanisms must work together to allow the brain to learn efficient representations for the various tasks it is facing. They shape the neural code and form the foundation on which our higher cognitive abilities are built.

While great progress has been made in characterizing these mechanisms individually, we only have a poor understanding of how they work together at the network level. In a non-linear system like the brain, any local change to, say, a synaptic efficacy potentially alters the pattern of activity at the level of the entire network and may induce further plastic changes to it. To investigate these processes, recent methods for observing the activities of large populations of neurons simultaneously need to be combined with careful measurements of the evolution of their synaptic and intrinsic properties – a formidable task for experimental neuroscience.

Computational modeling and theoretical analysis can contribute to this quest by providing simplified model systems that hopefully capture the essence of some of the brain's mechanisms and that can reveal underlying principles. In this article, we have introduced the SORN (self-organizing RNN). It combines three different kinds of plasticity and learns to represent and in a way “understand” the structure in its inputs. Maybe its most striking feature is the ability to map identical inputs onto different internal representations based on temporal context. For example,

it learns to distinguish the 5th repetition of an input from the 6th repetition by finding distinct encodings (internal representations) for the two situations (compare **Figure 3**). All this is happening in a completely unsupervised way without any guidance from the outside. The “causal” nature of the STDP rule is at the heart of this mechanism. It allows the network to incorporate predictable input structure into its own dynamics. At the same time, we have shown that STDP needs to be complemented by two homeostatic plasticity mechanisms. Without them the network will lose its favorable learning properties and may even develop seizure-like activity bursts (compare **Figure 5**).

Our network can be contrasted to recurrent networks without plasticity. Such static networks have received significant attention in the recent past, giving rise to the field of reservoir computing (Jaeger, 2001; Maass et al., 2002). The performance of a reservoir network relies on two requirements: (a) that different inputs to the network result in separable outputs based on the reservoir's response (the separation property) and that (b) the network activity states maintain information about recent inputs (the fading memory property). Given the high dimensionality of the reservoir, the separation property is easy to meet. Dockendorf et al. (2009) have confirmed this property for in vitro networks of cortical neurons. The memory property has been addressed in a series of experimental studies, across different brain areas, that compare the

neuronal response to a stimulus B vs. the response to B when it was preceded by stimulus A (Brosch and Schreiner, 2000; Bartlett and Wang, 2005; Broome et al., 2006; Nikolic et al., 2006). For example in (Nikolic et al., 2006), the authors analyzed neuronal responses in cat primary visual cortex, area 17, to a sequence of two letter images and were able to recover the identity of the first and second letter reliably using a simple linear classifier.

The most important force shaping the representations in the SORN is STDP. Although the STDP model we used is much simplified, it captures what is arguably the essence of STDP: a “causal” modification of synaptic strengths. In recent years much evidence has accumulated suggesting that the brain’s encoding of stimuli is subject to modifications due to STDP-like mechanisms. Several studies showed that repetitive stimulation with temporally patterned inputs causes a rapid STDP-based synaptic reorganization (Yao and Dan, 2001; Fu et al., 2002; Yao et al., 2007). Specifically, in Yao et al. (2007) a short repeated exposure to natural movies induced a rapid improvement in response reliability in cat visual cortex. Interestingly, the movie stimulation also left a “memory trace” which could be picked up in subsequent spontaneous activity.

It is interesting to note that in all the example tasks we considered the SORNs outperformed optimized versions of recurrent networks without plasticity. We find it unsurprising but rather reassuring that networks that try to discover and incorporate the temporal structure of their inputs into their dynamics outperform static reservoirs. Under repetitive stimulation with temporally structured inputs, SORNs selforganize in efficient ways that boost the network memory and separation properties. In our results, the SORNs could incorporate much longer input sequences as compared to the static reservoirs of similar size (Figure 2). SORNs developed internal representations where each input condition, reflecting both spatial and temporal aspects of the input, produced a tight cluster of network states that was well separated from those of other input conditions. This results in orderly and stereotyped trajectories through the network state space, that can be easily separated by a linear readout.

Reservoir computing architectures are thought to function best when their dynamics are critical (which we also found true for random reservoirs). It has been proposed that self-organization based on neuronal plasticity is able to achieve critical dynamics (Lazar et al., 2007; Gómez et al., 2009). Interestingly, the SORNs develop dynamics that are subcritical (compare Figure 4). This raises two questions. First, what is the exact mechanism that gives rise to the subcritical dynamics? Second, why are the subcritical dynamics of SORNs superior to the critical dynamics of static networks? Regarding the latter, we speculate that SORNs’ ability to incorporate the *predictable* sequence of inputs into their internal dynamics makes it unnecessary to maintain criticality, which should give the best fading memory for *arbitrary* input sequences. But if there is predictable structure in the input, the recurrent network should try to exploit and use its resources to model this specific structure rather than striving to have a general purpose fading memory.

The current model is particularly suited for efficient hardware implementation due to the simplicity of the chosen model neurons. In the current design individual neurons do not have any

intrinsic memory properties, which makes a strong point that all memory information is maintained by the recurrent dynamics. An open problem is to investigate the generality of these ideas in the context of more elaborate network models based on integrate and fire neurons or conductance based neurons, which also include direct connections between inhibitory units. Future work needs to address if the performance advantage of SORNs over static networks transfers from the simple problems studied here to more difficult engineering problems in time series prediction, speech recognition, etc.

We have shown how the synergistic combination of different local plasticity mechanisms can shape the global structure and dynamics of RNNs in meaningful and adaptive ways. This emergent property could not have been easily predicted on the basis of the individual mechanisms – the whole is more than the sum of its parts. This implies that as we try to understand neural plasticity and how it shapes the brain’s representation and processing, it is insufficient to study individual mechanisms in isolation. Only by studying their interaction at the network level, we have a chance to unravel this mystery.

## APPENDIX

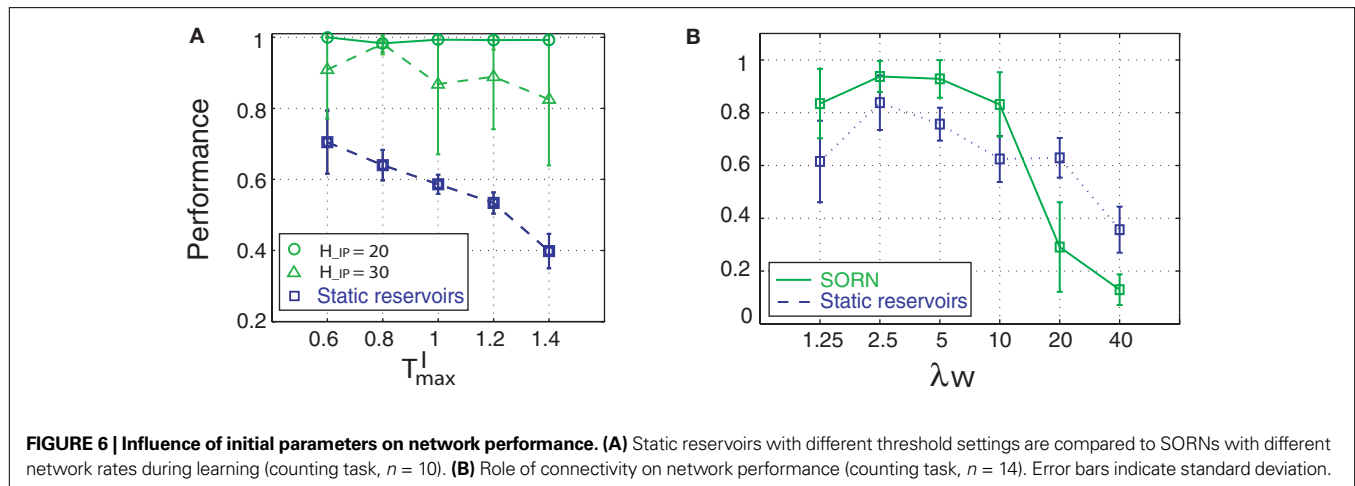
### PERFORMANCE AND NETWORK SETTINGS

For static reservoirs the choice of threshold values for excitatory ( $T_{\max}^E$ ) and inhibitory units ( $T_{\max}^I$ ) plays an important role in determining the network rate  $H_0$ , defined as the mean fraction of firing neurons per unit of time. Furthermore, the setting of  $T_{\max}^E$  and  $T_{\max}^I$  has an impact on the reservoir’s dynamics in terms of criticality and performance for prediction. A detailed analysis of the dependence between initial threshold settings and network dynamics for static reservoirs and SORNs is given in the supplementary online material.

In Figure 6A example networks with  $N^E = 200$ ,  $N^U = 10$ ,  $\lambda^W = 10$ ,  $T_{\max}^E = 0.25$  and various values of  $T_{\max}^I$  present significant improvements in prediction scores for SORNs (green) over static reservoirs (blue). The fraction of input spikes at the beginning of training is approximately  $N^U/N^E$ . A higher  $H_{\text{ip}}$  ( $H_{\text{ip}} = 3 \times N^U/N^E$ ) leads to a higher fraction of reservoir spikes compared to input spikes and results in a smaller increase in performance for SORNs (Figure 6A green triangles). These results suggest that a purposeful self-organization with significant improvements in performance relies on a balanced representation of input drive and internally generated drive (Figure 6A green circles).

In addition, we varied the number of synaptic connections per neuron ( $\lambda^W = 1.25, 2.5, 5, 10, 20, 40$ ). Figure 6B compares the prediction performance of networks with  $N^E = 200$ ,  $N^U = 10$ ,  $T_{\max}^E = 0.5$ ,  $T_{\max}^I = 0.8$  performing a counting task with  $n = 14$ . We find that a sparse connectivity is preferable both for static networks (blue) as well as SORNs (green). A high network connectivity induced seizure-like bursts of activity at the expense of computation (not shown). For a sparse connectivity SORNs perform better than the corresponding static reservoirs.

To summarize, SORN’s prediction performance: (a) is independent of the rate, criticality and performance of the initial static reservoir, (b) requires sparse network connectivity and (c) relies on a balanced representation of input spikes vs. reservoir spikes during learning.



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