Dynamics and computation in mixed networks containing neurons that accelerate towards spiking

Paul Manz, Sven Goedeke, and Raoul-Martin Memmesheimer

Neural Network Dynamics and Computation, Institute for Genetics, University of Bonn, 53115 Bonn, Germany

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Networks in the brain consist of different types of neurons. Here we investigate the influence of neuron diversity on the dynamics, phase space structure, and computational capabilities of spiking neural networks. We find that already a single neuron of a different type can qualitatively change the network dynamics and that mixed networks may combine the computational capabilities of ones with a single-neuron type. We study inhibitory networks of concave leaky (LIF) and convex “antileaky” (XIF) integrate-and-fire neurons that generalize irregularly spiking nonchaotic LIF neuron networks. Endowed with simple conductance-based synapses for XIF neurons, our networks can generate a balanced state of irregular asynchronous spiking as well. We determine the voltage probability distributions and self-consistent firing rates assuming Poisson input with finite-size spike impacts. Further, we compute the full spectrum of Lyapunov exponents (LEs) and the covariant Lyapunov vectors (CLVs) specifying the corresponding perturbation directions. We find that there is approximately one positive LE for each XIF neuron. This indicates in particular that a single XIF neuron renders the network dynamics chaotic. A simple mean-field approach, which can be justified by properties of the CLVs, explains the finding. As an application, we propose a spike-based computing scheme where our networks serve as computational reservoirs and their different stability properties yield different computational capabilities.

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I. INTRODUCTION

Biological neural networks consist of a large variety of interconnected neurons, which communicate via short stereotypical electrical pulses called action potentials or spikes. After a neuron has generated a spike, this travels along the axon and is transmitted to other neurons at synaptic contacts. The electrical membrane potential of the receiving neuron is then changed by an excitatory or inhibitory current pulse. Sufficiently many excitatory inputs in turn lead to spike generation in a receiving neuron. Many biological neural networks generate irregular and asynchronous spiking. This is likely caused by a dynamically balanced network state, in which the average inhibitory and excitatory input current to each neuron sum to a value that is insufficient for frequent spike generation [1–4]. Spikes are caused by fluctuations in the inputs, and the resulting spiking dynamics appear random and irregular.

Irregular dynamics are often chaotic, implying that the dynamics are sensitive to perturbations: initially small ones can strongly grow with time, which results in ultimately large quantitative differences between perturbed and unperturbed trajectories. A powerful tool to quantify this sensitivity and therewith the local phase space structure are the Lyapunov exponents (LEs) and associated with them the covariant Lyapunov vectors (CLVs) [5,6]. The sign of the largest LE indicates whether the system is chaotic, and its magnitude equals the long-term average growth or decay rate of generic infinitesimal perturbations. The spectrum of LEs describes the long-term average evolution of volumes spanned by tangent vectors and the change of infinitesimal perturbations in nongeneric directions, which are specified by the CLVs.

To each LE, there is a CLV. The size of a perturbation in the CLV’s direction changes with an average rate of plus or minus the corresponding LE for long-term forward or backward time evolution, respectively. The CLVs thereby indicate the directions of the unstable and stable manifolds along a trajectory. Furthermore, the spectrum of LEs can be used to derive dynamical quantities such as the Kaplan-Yorke fractal dimension of a chaotic attractor [7].

In our study, we consider purely inhibitory networks of current-based, oscillating integrate-and-fire-type neurons with postsynaptic currents of infinitesimally short duration and instantaneous reset. It has been shown numerically [8,9] and analytically [10,11] that if such networks contain only leaky integrate-and-fire (LIF) neurons, the networks’ irregular balanced state dynamics are stable against infinitesimal and small finite-size perturbations and are thus not chaotic but a realization of stable chaos [12,13]. The dynamics ultimately converge to a periodic orbit; the durations of the preceding irregular transients, however, grow exponentially with system size. The stability of the network dynamics is robust against introducing excitatory connections and considering synaptic currents of finite temporal extent [9,11], and there is a smooth transition to chaos upon increasing the number of excitatory connections and the duration of synaptic currents. The computational abilities of the stable precise spiking dynamics have not yet been explored, even though the specific structure of the phase space, which is composed of “flux tubes,” may be beneficial and exploitable [14].

LIF neurons incorporate a leak current as found in biological neurons [15]. This increases linearly with increasing membrane potential and leads to dissipation (contraction of
phase space volume) in the subthreshold dynamics. When driven by a constant depolarizing input current, the membrane potential therefore has negative second derivative; the neuron has a purely concave so-called rise function. In the considered class of networks, this implies the stability of the microscopic dynamics if only LIF neurons are present [10,11]. In biological neurons as well as in neuron models that explicitly model spike generation, such as the quadratic and the exponential integrate-and-fire neuron [16], the membrane potential accelerates towards a spike for larger membrane potentials. The rise function thus has a convex part. Reference [17] showed that networks of quadratic integrate-and-fire neurons that are otherwise similar to those considered in Refs. [9–11] exhibit chaos. Furthermore, Ref. [17] computed the spectrum of LEs and quantities that are derivable from them, as well as the statistics of the first CLV, which points to the directions to which a generic perturbation vector aligns in the long term.

Motivated by the above results and by the fact that there are many different types of cortical inhibitory interneurons [18], in the present study we investigate the impact of inserting a different type of neuron, with nonconcave rise function, into inhibitory networks of LIF neurons. To be specific, we insert “antileaky” integrate-and-fire (XIF) neurons with purely convex rise function. We choose the letter ‘X’ in the abbreviation to highlight this convexity and the expansion of phase space volume by the flow of the subthreshold dynamics. XIF neurons may be interpreted as a model for a class of biological neurons whose membrane potential lingers in a region where it accelerates towards spiking. Simultaneously, these neurons maintain similar analytical tractability as their leaky counterparts because of their mostly linear subthreshold dynamics. We describe our neuron and network models in detail in the next section. Thereafter, self-consistent firing rates and membrane potential probability distributions for both types of neurons are analytically derived, assuming Poisson input with finite size spike impacts. We then consider the dynamical stability properties and local phase space structures of the network dynamics, computing the entire spectra of LEs both numerically and analytically in a mean-field approximation. We also compute their CLVs to investigate how the stable and unstable directions are related to the different neuron types within the network.

Finally, we consider computations in pure and mixed networks of the considered types and show how the richer phase space structure in mixed ones can be exploited. For this, we propose a reservoir computer based entirely on precisely timed spikes. Reservoir computing has been introduced several times at different levels of elaborateness and in different flavors, in machine learning, and in neuroscience [19–22]. A reservoir computer consists of a high-dimensional, nonlinear dynamical system, the reservoir or liquid, and a comparatively simple readout. The reservoir “echoes” the input in a complicated, nonlinear way; it acts like a random filter bank with finite memory as each of its units generates a nonlinearly filtered version of the current input and its recent past while forgetting more remote inputs [19,21–23]. The simple, often linear readout can then be trained to extract the desired results, while the reservoir is static. In our scheme, the output neuron is spiking and thus nonlinear, and the desired outputs are trains of precisely timed spikes. The learning thus requires different approaches than learning of conventional continuous targets; gradient-descent-based methods [24] fail due to the discontinuity at the threshold as well as methods that require errors to be small but finite [25]. A number of algorithms have been suggested to learn precisely timed spikes [26–33], mostly using heuristic approaches. For our readout neuron, we can use the Finite Precision Learning scheme [30]. It has been shown to generically converge if the input-output relation is realizable at all, which explains its numerically found superior learning abilities [31].

II. MIXED NETWORKS OF NEURONS WITH CONCAVE AND CONVEX RISE FUNCTION

We consider a recurrent network with $N$ neurons. The $k$th spike of neuron $j$, which is sent at time $t_{jk}$, generates a postsynaptic current pulse $h_i(V_i^-)C_{ij}\delta(t-t_{jk})$ in neuron $i$. Here $C_{ij} \leq 0$ is the weight of the inhibitory connection and $h_i(V_i^-)$ is a possible voltage-dependent modulation, which depends on the membrane potential of neuron $i$ just before input arrival, given by the left-hand side limit $V_i^- = V(t^-) = \lim_{\varepsilon \to 0} V_i(t-\varepsilon)$. We assume that all excitatory inputs to neuron $i$ can be gathered into a constant excitatory external input current $I_{i\text{ext}} > 0$ and that the remaining explicitly modeled recurrent inhibition is fast [10,14,34]. We further assume that there is a leak term with prefactor $\gamma_i \neq 0$. Taken together, we model the subthreshold membrane potential dynamics of neuron $i$ by

$$V_i = -\gamma_i V_i + I_{i\text{ext}} + h_i(V_i^-) \sum_{j=1}^{N} C_{ij} \sum_{k} \delta(t-t_{jk}). \quad (1)$$

When $V_i$ reaches the spike threshold at time $t$, $V_i^- = V_{\text{th}} > 0$, it is reset, $V_i(t) = V_{\text{re}} = 0$, and a spike is emitted. This, in turn, generates in a postsynaptic neuron $l$ a current pulse as introduced above, which causes $V_l$ to decrease in jumplike manner from $V_l^- = h_l(V_l^-)C_{il}$. The rise function, i.e., the membrane potential dynamics with $V_l(0) = 0$ in absence of recurrent inhibitory input [35,36], reads

$$V_l(t) = \frac{I_{l\text{ext}}}{\gamma_l}[1 - \exp(-\gamma_l t)]. \quad (2)$$

It is concave for $\gamma_l > 0$ and convex for $\gamma_l < 0$.

There are two types of neurons in our networks: LIF neurons with dissipation and concave rise function, which obey Eq. (1) with $\gamma_l > 0$, and antileaky XIF neurons with convex rise function, which obey Eq. (1) with $\gamma_l < 0$; see Fig. 1. The membrane potential dynamics of an LIF neuron has a globally attracting fixed point at $V_{\infty,i} = \frac{I_{i\text{inh}}}{\gamma_l}$, if there is no threshold for spike generation and no inhibitory input. We assume $V_{\infty,i} > V_{\text{th}}$, so neurons without inhibitory input periodically spike and reset. For our study it is sufficient to endow the LIF neurons with a simple, current-based synapse model, setting $h_i(V_i^-) = 1$. A coarse approximation of the membrane potential dynamics without threshold and neglecting input fluctuations yields \( \dot{V}_i = -\gamma_i V_i + I_{i\text{ext}} + I_{i\text{inh}} \), where $I_{i\text{inh}}$ is the average inhibitory input current. In the balanced state, its attractor at $\bar{V}_{\infty,i} = (I_{i\text{ext}} + I_{i\text{inh}})/\gamma_l$ is below or close to the spike threshold, such that spikes are always or typically
generated by input fluctuations, more specifically by periods of less than average inhibition.

In the absence of inhibitory input XIF neurons have an unstable, repelling fixed point at \( V_{-\infty,i} = \frac{I_i^{\text{ext}}}{\gamma_i} < 0 \). If the membrane potential starts above this separatrix, it increases exponentially towards the threshold. When it reaches there, the neuron spikes, its membrane potential resets to zero, increases towards the threshold again, and so forth: XIF neurons oscillate and spike periodically for any \( I_i^{\text{ext}} > 0 \), if there is no inhibitory input. If the membrane potential starts below the separatrix, it decreases exponentially to \(-\infty\). Also in the presence of recurrent inhibitory inputs an XIF neuron is uncovery switched off once its membrane potential falls below \( I_i^{\text{ext}}/\gamma_i \), since the inputs only decrease the membrane potential further. Averaging over the inhibitory inputs as before yields an effective separatrix at \( \bar{V}_{-\infty,i} = \left( I_i^{\text{ext}} + I_i^{\text{inh}} \right)/\gamma_i \). Membrane potentials falling below it have a tendency to further decrease, causing the neuron to effectively switch off. This can be also seen from the phase response curve of XIF neurons, which gets steeper for negative phases, in contrast to that of LIF neurons, which becomes flatter; see Fig. 1(c). In other words, in XIF neurons an incoming inhibitory input at a low potential still above the separatrix (and thus at a low phase) has a larger effect in the sense that it delays the next spiking more than the same input arriving at a higher potential. As a consequence, we observe in networks containing XIF neurons with purely current-based input \( [h(V_i) = 1] \) that many of these neurons are first effectively and then uncovery switched off, if the network dynamics are irregular, and the inhibitory inputs are therefore strongly fluctuating. In order to prevent this biologically implausible phenomenon, we introduce a voltage dependence

\[
h(V_i^-) = \Theta(V_i^- - V_{\text{cutoff}})
\]

de the input coupling strength, where \( \Theta \) is the Heaviside theta function. Inhibitory inputs arriving at a membrane potential lower than \( V_{\text{cutoff}} \) then do not induce a further decrease. This provides a simple conductance-based model for the synapses, where the driving force of the current vanishes below \( V_i = V_{\text{cutoff}} \) and is constant above. We assume \( V_{-\infty,i} < V_{\text{cutoff}} + C_{ij} \) for all \( j \) to exclude unrecoverable switching off and \( V_{\text{cutoff}} \leq V_{\text{rec}} \). We exemplarily checked that the overall network dynamics and their stability properties remain qualitatively unchanged, if we also endow the LIF neurons with these synapses.

For simplicity, we choose the parameters of all LIF and of all XIF neurons identical, i.e., \( \gamma_i = \gamma_{\text{LIF}} \), \( I_i = I_{\text{LIF}} \), etc., if neuron \( i \) is an LIF neuron, and \( \gamma_i = \gamma_{\text{XIF}} \), \( I_i = I_{\text{XIF}} \), etc., if neuron \( i \) is an XIF neuron. The spike threshold and reset potentials are \( V_{th} = 1 \) and \( V_{re} = 0 \), independent of the neuron type. We set \( V_{\text{cutoff}} = V_{\text{rec}} \) to avoid any effective switching off of XIF neurons. Coupling strengths are homogeneous, \( C_{ij} = C \), if the coupling is present. To keep the number of relevant parameters small, we further choose \( I_i^{\text{LIF}}/\gamma_i = V_{-\infty,\text{LIF}} = -V_{-\infty,\text{XIF}} = -I_i^{\text{XIF}}/\gamma_{\text{XIF}} \). The additional choice \( \gamma_{\text{XIF}} = -\gamma_{\text{LIF}} \) leads already in the absence of recurrent inhibition to a higher spike rate \( \rho_{\text{free,XIF}} \) in XIF neurons, since

\[
\rho_{\text{free,XIF}} = \frac{-\gamma_{\text{XIF}}}{\ln \left( \frac{V_{-\infty,\text{XIF}} + V_{\text{rec}}}{V_{-\infty,\text{XIF}}} \right)},
\]

\((4)\)
neurons. (a), (b) Spiking activity for a subset of the LIF (a) and the balanced state with asynchronous irregular activity in both types of neuron (b) neurons with convex rise function can exhibit a separate. (c)–(e) Distribution of coefficients of variation of inter-spike intervals for all neurons (c) and for LIF (d) and XIF (e) neurons separately. (f)–(h) Distribution of the average spike rates of all neurons 100). (c)–(e) Distribution of coefficients of variation of inter-spike intervals for all neurons (c) and for LIF (d) and XIF (e) neurons separately. (f)–(h) Distribution of the average spike rates of all neurons (f) and of LIF (g) and XIF (h) neurons separately. The analytically derived rate \( \rho \approx 26.1 \, \text{s}^{-1} \) [Eqs. (18) and (19)] is indicated by a black dashed vertical line. We use \( \gamma_{\text{XIF}} = -0.1 \, \text{ms}^{-1} \), \( \gamma_{\text{LIF}} = 0.169 \, \text{ms}^{-1} \), \( V_{\infty, \text{LIF}} = -V_{\infty, \text{XIF}} = 2 \) and a randomly connected network with fixed indegrees \( K = 50 \) and nonzero synaptic strength \( C_{ij} = C = -0.2 \).

whereas in LIF neurons,

\[
\rho_{\text{prox,LIF}} = \frac{\gamma_{\text{LIF}}}{\ln \left( \frac{V_{\infty, \text{LIF}}}{V_{\infty,LIF} - V_{\text{th}}} \right)}. \tag{5}
\]

As a consequence, we observe that in a mixed network the XIF suppress the LIF neurons, which become quiescent. Using the analytical results of the next section, we therefore rescale \( \gamma_{\text{LIF}} \) such that the spike rates in both populations are identical. Further, we fix the neurons’ indegree to the same number \( K \), implying that \( \sum_j C_{ij} \) is identical for each neuron \( i \). This reduces quenched noise [40] and avoids strong differences in average spike rates and switched off neurons.

With the described network model setup, we observe balanced states of asynchronous irregular spiking activity for any ratio of neurons with concave and convex rise function; see Fig. 2 for an illustration.

III. NETWORK FIRING RATE AND MEMBRANE POTENTIAL DISTRIBUTIONS

Mean-field theories have been developed in statistical physics [41] and are frequently used in computational neuroscience; see, for example, Refs. [40,42–44]. The basic idea is to average the interactions in a high-dimensional system to obtain for each element an effective action, which is not influenced by this element anymore. One can thereby reduce a high-dimensional problem to low-dimensional ones. In this section we analytically determine the steady-state firing rate and the voltage probability densities for LIF and XIF neurons in mixed networks using a mean-field approximation. We use the results to obtain neuron parameters that lead to the same average firing rates for both neuron types and thus to homogeneous firing rates in the entire network. In addition we employ the firing rates to analytically approximate the Lyapunov spectrum of the network dynamics using a mean-field approach in Sec. IV A.

We approximate the superposed input spike trains to a neuron by a Poisson spike train with a given rate; i.e., we assume that all input spikes are sent independently of each other. A common approach is to additionally consider the limit of a large number of small inputs. The neuron dynamics can then be approximated by a diffusion process, which allows us to compute firing rates and membrane potential distributions [45,46]. This diffusion approximation assumes that the inputs have (infinitesimally) small amplitude and arrive at (infinitely) high rate. Here we use a shot-noise approach, which accounts for the finite input rate and size of individual inputs [45,46], in the recent formulation of Refs. [34,47]. This allows us to more accurately obtain the firing rates and membrane potential distributions. In particular, the fact that in our networks the voltage probability density does not go to zero at threshold is reflected. We shortly review the approach for LIF neurons [34,47,48] and then extend it to XIF neurons with the voltage-dependent coupling (3).

The shot-noise approach (like the diffusion approximation) is based on the continuity equation for the voltage probability density \( \rho(V, t) \). For our neuron models it reads

\[
\frac{\partial \rho}{\partial t} + \frac{\partial j}{\partial V} = \rho_{\text{inh}} + \sigma_{\text{reset}}, \tag{6}
\]

where \( j(V, t) = V(V)\rho(V, t) \) is the drift probability current with velocity \( V(V) = -\gamma V + f_{\text{ext}} \); \( \sigma_{\text{inh}}(V, t) \) and \( \sigma_{\text{reset}}(V, t) \) are source terms incorporating the effects of inputs and resets of the neuron’s membrane potential \( V \).

For the LIF neuron without the voltage-dependent input, inhibitory input spikes arriving when the considered neuron is at a voltage \( V \) give rise to a sink at \( V \), whereas spikes arriving when the neuron is at a voltage \( V - C > V \) give rise to a source at \( V \). We therefore have a first source term

\[
\sigma_{\text{inh}}(V, t) = r(t)[\rho(V - C, t) - \rho(V, t)] \tag{7}
\]

with the rate \( r(t) \) of input spikes. We note that Refs. [34,47,48] include this term in the probability current. The second source term is due to the spike and reset mechanism of the neuron model. Its threshold and reset act as Dirac delta sink and source at the corresponding discrete voltages,

\[
\sigma_{\text{reset}}(V, t) = \rho(t)\delta(V - V_{\text{th}}) - \delta(V - V_{\text{th}}). \tag{8}
\]

This term is proportional to the instantaneous firing rate \( \rho(t) \) of the stochastic neuron dynamics or, in other words, to the probability current through the threshold \([\rho(t) = j(V_{\text{th}}, t) \geq 0] \).
We investigate stationary network dynamics, which are described by constant $r$ and $\rho$ and time-independent $p(V)$. For these Eq. (6) reduces to the linear delay differential equation (or differential-difference equation)
\[
\frac{d}{dV} p(V)(-\gamma V + f^{\text{ext}}) = r[p(V - C) - p(V)] + \rho[\delta(V - V_{\text{re}}) - \delta(V - V_{\text{inh}})].
\]

Dividing Eq. (9) by $\rho > 0$ yields an equation for the rescaled density $q(V) = p(V)/\rho$, which is independent of the unknown steady-state firing rate $\rho$. This equation can be integrated, for example, with the method of steps [49]. The integration starts with the “initial conditions” $q(V) = 0$ for $V > V_{\text{th}}$ and thus $q(V_{\text{th}}) = 1/(\gamma V_{\text{th}} + f^{\text{ext}})$ slightly below $V_{\text{th}}$. The normalization of $p(V)$ allows us to compute $\rho$ via
\[
\frac{1}{\rho} = \int_{-\infty}^{\infty} q(V) dV.
\]

To obtain an analytic expression for $\rho$, one applies a bilateral Laplace transform $f(s) = \int_{-\infty}^{\infty} f(V) e^{V s} dV$. We can focus on $s \geq 0$; $\tilde{q}(0)$ yields $\rho^{-1}$. The Laplace transform of the rescaled Eq. (9) results in a linear first-order ordinary differential equation for $\tilde{q}(s)$,
\[
\frac{d}{ds}\tilde{q}(s) = \left[\frac{f^{\text{ext}}}{\gamma} + \frac{r(e^{Cs} - 1)}{\gamma s}\right] \tilde{q}(s) + \frac{e^{V_{\text{th}} s} - e^{V_{\text{ext}} s}}{\gamma s}.
\]

It can be solved by variation of constants. The solution of the homogeneous equation is
\[
Z_0(s) = Ae^{\Psi(s)}
\]
with an arbitrary constant $A$ and
\[
\Psi(s) = \frac{f^{\text{ext}}}{\gamma} s + \frac{r}{\gamma} \int_{-\infty}^{s} e^{Cu} - 1 \ du = \frac{f^{\text{ext}}}{\gamma} s + \frac{r}{\gamma} [\text{Ei}(Cs) - \log(-Cs) - \Gamma].
\]

Here $\text{Ei}(s)$ is the exponential integral $\text{Ei}(x) = -\int_{-\infty}^{x} e^{-t} \ dt$ and $\Gamma$ is the Euler-Mascheroni constant. The solution of the full equation then reads
\[
\tilde{q}(s) = e^{\Psi(s)} \left[\tilde{q}(0) - \int_{0}^{s} e^{-\Psi(u)} \frac{e^{Vu} - e^{V_{\text{th}} u}}{\gamma u} \ du\right].
\]

Since the support of $q(V)$ is bounded from above by $V_{\text{th}}$, $\tilde{q}(s) = \int_{-\infty}^{\infty} q(V) e^{V s} dV \leq e^{V_{\text{th}} s}/\rho_0$. To balance the faster exponential growth $-\exp(f^{\text{ext}} s/\gamma)$ of its prefactor $\exp[\Psi(s)]$, the bracket on the right-hand side of Eq. (14) needs to vanish for large $s$. We thus have
\[
\tilde{q}(0) = \int_{0}^{\infty} e^{-\Psi(u)} \frac{e^{Vu} - e^{V_{\text{th}} u}}{\gamma u} \ du = \frac{1}{\rho}.
\]

For an XIF neuron without voltage-dependent synapses there is no stationary membrane potential probability density $p(V)$. This is because for any time $t > 0$ there is a finite probability that the membrane potential of a neuron jumps below $I^{\text{ext}}/\gamma$ and thereafter tends to minus infinity. In contrast, for an XIF neuron with the voltage dependence (3), $p(V)$ exists, and we may use the same approach as for the LIF neuron to determine it together with the firing rate. Since membrane potentials do not drop below $V_{\text{cutoff}} + C$, we focus on the interval $[V_{\text{cutoff}} + C, V_{\text{inh}}]$, where $p(V)$ can be nonzero. The couplings’ voltage dependence enters the source term $\sigma_{\text{inh}}$ in Eq. (6): If $V$ is below $V_{\text{cutoff}}$, incoming spikes have no effect and the sink term due to them vanishes. Equation (7) therefore changes to
\[
\sigma_{\text{inh}}(V, t) = r(t)[p(V - C, t) - h(V)p(V, t)],
\]
where we used that $V - C \geq V_{\text{cutoff}}$ in the relevant voltage range such that a modification of the source term is unnecessary. The stationary continuity equation becomes
\[
\frac{d}{dV} p(V)(-\gamma V + f^{\text{ext}}) = r[p(V - C) - h(V)p(V)] + \rho[\delta(V - V_{\text{re}}) - \delta(V - V_{\text{inh}})],
\]
which can be rescaled and integrated using the method of steps to obtain $q(V)$, $\rho$ and $p(V)$ as before. The nonlinear prefactor $h(V)$, however, impedes the derivation of $\rho$ via the Laplace transform.

We apply the above results to find mixed networks in which LIF and XIF neurons have similar firing rates. Equation (15) provides a map $G_{\text{LIF}}$ from the input to the output rate, $G_{\text{LIF}}(r) = \rho$. Equation (17) implicitly defines such a map $G_{\text{XIF}}$ for XIF neurons. The firing rate $\rho$ of the LIF and XIF neurons in the desired mixed network needs to solve both self-consistency equations
\[
G_{\text{LIF}}(K\rho) = \rho,
\]
\[
G_{\text{XIF}}(K\rho) = \rho,
\]
with the neurons’ indegree $K$. We employ Eq. (19) to compute $\rho$ for XIF neurons. Thereafter, we adapt the parameters of Eq. (18) such that the same $\rho$ becomes a solution. Specifically, we solve for $\gamma_{\text{LIF}}$, keeping the other parameters fixed.

Figure 3 compares the voltage densities $p(V)$ and rates $\rho$ obtained from the shot-noise approach with those of an LIF and an XIF neuron that receive input spike trains as they are generated in the recurrent network of Fig. 2. There is a pronounced discrepancy between the densities and rates for an LIF neuron for $K = 50$ and small $N$, because both the individual [see Figs. 2(a)–2(e)] and the superposed input spike trains in these dense networks are more regular than Poisson spike trains. Removing spatial correlations, for example, by increasing $N$ reduces the discrepancy; see Figs. 3(b)–3(d) and Appendix A for further analysis. Such input spike trains reduce the variance of the voltage and generate a $p(V)$ that is more concentrated around the value $(I^{\text{ext}} + I^{\text{inh}})/\gamma$, where $I^{\text{inh}}$ is the average inhibitory input current as discussed in Sec. II. For the XIF neuron, the input spike train statistics has less impact on $p(V)$. Presumably, this is because voltage
FIG. 3. Analytically and numerically estimated voltage probability densities and spike rates of LIF and XIF neurons. (a), (b) Voltage probability densities for networks of (a) $N = 100$ and (b) $N = 10,000$ neurons. The dark blue and dark red curves show the analytical results (9) and (17) of $\rho(V)$ for LIF and XIF neurons, where $\rho$ is obtained self-consistently from Eqs. (18) and (19). The light blue and light red curves show representative numerically sampled voltage densities of an LIF and an XIF neuron, where the input spike trains are superpositions of simultaneous output spike trains of $K$ neurons in the recurrent network. (c), (d) Spike rates of neurons in networks of (c) different sized $N$ and indegree $K = 50$ and (d) size $N = 10000$ and different indegree $K$. In panels (d) the presynaptic weights are scaled with $1/K$ such that their sum is independent of $K$. Numerically measured average spike rates of LIF and XIF neurons in the different networks are shown by blue and red dots. Error bars display the standard deviations of the rate distributions. Analytical results obtained from Eqs. (18) and (19) are displayed by dashed black lines. Remaining parameters are as in Fig. 2.

excursions due to input fluctuations are suppressed by the voltage dependence of the input strength (for potentials near $V_{\text{cutoff}}$) and by the drive towards threshold (for larger potentials). We note that the assumption of Poisson input spike trains is the only approximation in the chosen approach, such that sampled membrane potential distributions of neurons with Poisson input match the analytical ones up to the sampling noise as shown in Appendix A.

IV. GROWTH OF DYNAMICAL PERTURBATIONS

A. Mean-field approach

After obtaining the spike rates and membrane potential distributions using a statistical mean-field theory, we investigate the mixed network dynamics from a dynamical systems perspective. We first analytically determine the Lyapunov spectrum using again a mean-field approach. It focuses on the evolution of perturbations to a single neuron and treats the input from other neurons as external. Specifically, we disregard perturbations of the rest of the network including those generated by the considered neuron’s changed spiking.

Inputs thus arrive at the same times in the perturbed and in the unperturbed system and do not change the neuron’s perturbation. Figure 4 illustrates this and compares the resulting evolution of a perturbation of an XIF neuron and of an LIF neuron: The perturbation of the XIF neuron gradually increases as long as it is not spiking, while that of the LIF neuron decreases. Conversely, in the XIF neuron spiking and resetting reduces perturbations, while it increases them in the LIF neuron; compare the (finite-size) distance $|\delta V(t)| = |\tilde{V}(t) - V(t)|$ between two neighboring trajectories $\tilde{V}(t)$ and $V(t)$ in Figs. 4(a) and 4(b) before and after a spike event has taken place in both the perturbed and the unperturbed dynamics. To assess the influence of

FIG. 4. Evolution of perturbations during subthreshold evolution and spiking in (a) an LIF and (b) an XIF neuron. During subthreshold evolution the distance (perturbation) $|\delta V(t)|$ between two neighboring trajectories shrinks for LIF and grows for XIF neurons, while spike generation partially resets it. Due to the receiving of inhibitory spikes, the intervals between spike generations are generally longer than for freely oscillating neurons. The impact of the subthreshold dynamics therefore dominates and overall the perturbation in the LIF neuron decays while that in the XIF neuron grows. Here we use $V_{\infty,\text{LIF}} = 1.33$, $V_{\infty,\text{XIF}} = -1$, $\gamma_{\text{LIF}} = 0.5\,\text{ms}^{-1}$, and $\gamma_{\text{XIF}} = -0.3\,\text{ms}^{-1}$ for better illustration of the mechanism.
these two processes, we first note that in a freely oscillating neuron they need to cancel each other such that perturbations persist on average and the LE is zero. We then note that the inhibitory inputs do not affect perturbations but prolong the subthreshold evolution between spikes. Its impact therefore dominates, and perturbations in XIF neurons grow over time, while they shrink in LIF neurons. This does not depend on the specifics of the LIF and XIF dynamics but is a consequence of the curvature of the rise function and the inhibitory inputs.

In Appendix B we make the gained intuitive understanding precise by quantifying the growth of perturbations and the resulting LE. For this, we describe the dynamics by a sequence of discrete maps from the state at a time (infinitesimally) shortly after generation of a spike to the state at a time shortly after generation of the next spike. The discrete time dynamics of small perturbations are then given by the “single-spike Jacobians” [14,17] $J(k)$. For the effective single-neuron dynamics here, they reduce to scalar factors

$$J(k) = \frac{\partial V(t_{k+1}^+)}{\partial V(t_{k}^+)} = \exp \left[ \frac{\nu}{\rho_{\text{free}}} - \gamma (t_{k+1} - t_k) \right]$$

with the free firing rate $\rho_{\text{free}}$ [Eq. (4) or (5)] of the neuron.

The growth rate of perturbations and thus the mean-field LE are given by the long-term average of Eq. (20),

$$\lambda_{\text{mf}} = \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} \ln |J_{\text{mf}}(k)| = -\gamma \left( 1 - \frac{\rho}{\rho_{\text{free}}} \right).$$

(21)

This expression confirms the intuitive understanding that without perturbed inputs the growth rate depends (1) on the growth rate during subthreshold evolution and (2) on the prevalence of subthreshold evolution ($\rho < \rho_{\text{free}}$) or spike sending ($\rho > \rho_{\text{free}}$) relative to the free neuron case. In particular, without input we have $\lambda_{\text{mf}} = 0$ and if the neuron is silenced $\lambda_{\text{mf}} = -\gamma$. In our inhibitory networks we have $\rho < \rho_{\text{free}}$ such that $\lambda_{\text{mf}} > 0$ for XIF and $\lambda_{\text{mf}} < 0$ for LIF neurons. In networks in the balanced state, the actual spike rate is much smaller than the spike rate of a neuron if only excitation is present. Since in our networks the latter equals the spike rate of the freely oscillating neuron, we have $\rho_{\text{free}} \ll 1$. Thus the mean-field approach indicates that the growth rate of perturbations is mainly given by the subthreshold growth. The mean-field approach further indicates that a single XIF neuron renders the entire network dynamics unstable and that the number of unstable directions equals the number of XIF neurons in the network, while the number of stable directions equals the number of LIF neurons. This, however, does not give rise to a zero LE, which occurs in the full autonomous network due to time-translation symmetry. The mean-field spectrum and the rule for the number of stable and unstable directions can thus be only an approximation to the exact results.

Equation (21) together with the analytical results Eqs. (18) and (19) for $\rho$ give a fully analytical estimate of the Lyapunov spectrum. Since all LIF or XIF neurons have the same analytical rate estimates and leak strengths, the spectrum consists of $N_{\text{LIF}}$ identical negative and $N_{\text{XIF}}$ identical positive exponents; see Fig. 5. Due to quenched noise from random coupling, the rates in the actual network are distributed. We can account for this by inserting the numerically measured rates into Eq. (21); see Fig. 5.

### B. Network single-spike Jacobian

To derive exact Lyapunov spectra we need to take into account the spreading of perturbations in the network. For this, we compute the full single-spike Jacobian $J(k)$, which is a map from tangent vectors at the point $V(t_{k}^+)$ in phase space to tangent vectors at $V(t^+_{k+1})$, where $V(t) = [V_1(t), \ldots, V_N(t)]^T$ is the state of the system at time $t$. The resulting components of $J(k)$ read

$$J_{ij}(k) = \frac{\partial V_i(t_{k+1}^+)}{\partial V_j(t_{k}^+)} = \delta_{ij} e^{-\gamma(t_{k+1} - t_k)} + \delta_{ij} \frac{\gamma}{\rho_{\text{free}}} C_{ii} h_i [V_i(t_{k+1}^+)] - \frac{\rho_{\text{free}}}{\gamma} - V_i(t_{k}^+).$$

(22)

for an LIF or an XIF neuron $i$, where $i$ is the index of the neuron sending the $(k + 1)$th spike; see Appendix C for details. We note that the mean-field theory accounts for the diagonal terms of this Jacobian.

### C. Volume contraction

Owing to the simple form of the single-spike Jacobians we can find an analytical expression for the full network dynamics’ expansion rate of infinitesimal phase space volumes or, equivalently, for the sum of the LEs. The result in terms of the neuronal spike rates in the network is exact. It allows us to analytically compute the Lyapunov spectra for two-neuron systems and offers a test for the accuracy of their numerical estimates in larger networks.

The volume expansion and the sum of LEs are given by the time-averaged logarithms of the determinants of the Jacobians [5]. We thus have

$$\sum_{i=1}^{N} \lambda_i = \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} \ln |\det J(k)|$$

(23)

in terms of single-spike Jacobians [14]. In Appendix D we exploit the specific form of $J(k)$ to compute $\det J(k)$ with the matrix determinant lemma. The subsequent time averaging yields

$$\sum_{i=1}^{N} \lambda_i = - \sum_{j=1}^{N} \gamma_j \left( 1 - \frac{\rho_j}{\rho_{\text{free},j}} \right).$$

(24)

Notably, this shows that our mean-field theory yields an exact expression for the volume contraction rate and the sum of LEs: the estimate $\sum_{i=1}^{N} \lambda_{\text{mf},i}$ with Eq. (21) agrees with the exact expression (24).

### D. Numerical computation of the Lyapunov spectrum

The single-spike Jacobians (22) allow us to iteratively compute the largest LE and the full Lyapunov spectrum [5,17,50]; see also Appendix F. In short, for the largest LE, one iterates an initial random perturbation vector by the single-spike
FIG. 5. Lyapunov spectra of mixed networks. The numbers of LIF and XIF neurons in the networks are (a) $N_{\text{LIF}} = 100$ and $N_{\text{XIF}} = 0$, (b) $N_{\text{LIF}} = 99$ and $N_{\text{XIF}} = 1$, (c) $N_{\text{LIF}} = 75$ and $N_{\text{XIF}} = 25$, (d) $N_{\text{LIF}} = 50$ and $N_{\text{XIF}} = 50$, (e) $N_{\text{LIF}} = 75$ and $N_{\text{XIF}} = 25$, (f) $N_{\text{LIF}} = 0$ and $N_{\text{XIF}} = 100$. Blue circles display the numerically computed Lyapunov spectra using Eq. (22). Red diamonds display the mean-field result Eq. (21) with numerically measured neuron rates, and black lines display the mean-field result Eq. (21) using the analytically obtained rates (18) and (19). Insets show closeups of the positive and/or negative parts of the spectra.

Jacobians, stores its growth every few steps, and thereafter renormalizes it to its initial magnitude. The long-term average of the growth rate equals $\lambda_1$. For the full spectrum, one iterates a system of $N$ orthogonal perturbation vectors with the single-spike Jacobians. Every few steps, one records the growth of the different vectors. Thereafter one reorthogonalizes, always in the same order, and finally renormalizes the vectors. The long-term average growth rate of the first vector then equals $\lambda_1$, that of the second equals $\lambda_2$, etc. Reference [50] suggested an efficient method to compute the Lyapunov spectrum and applied it to large networks; we use some of the ideas in our implementation.

For networks consisting only of LIF neurons we find in agreement with previous work [8–11] and our mean-field theory that the largest nontrivial LE is negative; see Fig. 5(a). However, we also find that already the presence of a single XIF neuron renders the largest LE positive [see Fig. 5(b)], indicating chaos, in agreement with the mean-field theory. The computations also confirm that the destabilization of a network by a single XIF neuron is a special case of a general rule, namely, that each XIF neuron introduces about one positive LE. This holds independently of $N$ and $K$; see Fig. 5 and Appendix E. The trivial (zero) exponent is an exception to the rule. Our numerical results indicate that it replaces a negative exponent if there are more LIF than XIF neurons in the network and a positive exponent otherwise. There is also good quantitative agreement with the mean-field spectrum, in particular the exponents are close to $\gamma_{\text{LIF}}$ and $\gamma_{\text{XIF}}$. However, also when inserting the measured spike rates into Eq. (21) some discrepancy remains, showing that the spread of perturbations in the network and their transfer between neurons have a pronounced effect on their growth.

V. STABLE AND UNSTABLE DIRECTIONS

A. Lyapunov vectors and perturbation growth

To further elucidate the local phase space structure, we numerically investigate the characteristics of the perturbations that grow according to the individual LEs, i.e., how they are distributed across neurons and how they change during evolution. This will, in particular, allow us to understand why the mean-field theory works well. The directions of the perturbations are given by the CLVs or, in other words, by the stable and unstable manifolds along the trajectory [5,6].

The $i$th CLV $v_i(V_0)$ at a point $V_0$ in phase space is a normalized tangent vector that grows with long-term average rates $\lambda_i$ and $-\lambda_i$ when evolved forward and backward in time. We call it a stable CLV if $\lambda_i < 0$ and an unstable one if $\lambda_i > 0$. We assume for simplicity that all LEs are different; the vector is then unique up to its orientation. Consider a trajectory $V(t)$ that reaches shortly after the spike time $t_0$ the state $V(t_0^+)$ = $V_0$. Using the single-spike Jacobians $J(k)$, $v_i(V_0)$ may be defined as the tangent vector...
satisfying

\[
\prod_{k=0}^{L} J(k) v_i(V(0)) \sim e^{\lambda_i t},
\]

\[
\prod_{k=-M}^{-1} J^{-1}(k) v_i(V(0)) \sim e^{\lambda_i - t M},
\]

where \(L\) and \(M\) are chosen sufficiently large. The definition can be straightforwardly extended to states between spiking events. Both of its parts are important: The first part alone does not uniquely define the direction of \(v_i(V(0))\), since adding any vector with growth rate less than \(\lambda_i\) yields the same asymptotics. The second part excludes such an addition, since its shrinkage rate is slower than \(-\lambda_i\) and thus yields a different dominant asymptotics of backward evolution. As anticipated by the notation, the vector depends only on the state but not on the time when \(V(t)\) reaches it. Furthermore, the definition ensures covariance, that is, the evolution of infinitesimal perturbations (the tangent flow) maps CLVs to CLVs. At subsequent spike times we thus have

\[
J(k) v_i(V(t_k^+)) \propto v_i(V(t_{k+1}^-)).
\]

This extension to states between spike times is again straightforward: the covariance implies that we can obtain CLVs \(v_i(V(t))\) at a state between spike times \(t_k\) and \(t_{k+1}\) by propagating \(v_i(V(t_k^+))\) forward with the Jacobian \(J_j\) \((t - t_k) = \delta_{ij} \exp(-\gamma_i (t - t_k))\) of subthreshold evolution.

We compute the CLVs in a dynamical manner by forward and restricted backward propagating sets of vectors, following Refs. [5,51]. Appendix F provides a short description of the method. We note that the dynamics of our system are not invertible: given a state there is no unique way of propagating it back in time. This is because an ambiguity can arise at states where one neuron is at the reset potential; we generally cannot tell whether it was reset or crossed the reset potential from below (unless some postsynaptic neuron is too near to threshold to be able to have just received a spike). It is, however, still possible to compute the Lyapunov vectors by backward propagating along the trajectory that was previously taken for the forward propagation [51].

B. Stable and unstable directions in mixed networks

The CLVs yield the directions in which small but finite perturbations evolve according to the different LEs as shown in Fig. 6. We find that they generally contain perturbations to a variety of neurons and that they strongly change their direction during evolution. More specifically, we observe that the stable and unstable CLVs stay approximately confined to the subspaces of (strictly speaking: perturbations to) LIF and XIF neurons, respectively. Figures 7(a)–7(d) illustrate this by displaying the lengths \(\sqrt{\sum_{j=N_{LIF}+1}^{N} v_{i,j}^2(V(t))}\) and \(\sqrt{\sum_{j=N_{XIF}+1}^{N} v_{i,j}^2(V(t))}\) of the projections of different CLVs \(v_i\) onto the subspaces of LIF and XIF neurons. Here and in the following we assume that the LIF and XIF neurons have the indices \(1, \ldots, N_{LIF}\) and \(N_{LIF}+1, \ldots, N\), respectively. Figures 7(e) and 7(f) further illustrate the confinement and show the large temporal variability of single CLV components \(v_{i,j}\) that are not close to zero. The confinement does not hold exactly since perturbations of LIF neurons usually also give rise to perturbations of XIF neurons and vice versa.

In networks with inhomogeneous spike rates, we observe that single neurons that are strongly suppressed by inhibition have CLVs more aligned to them, because their perturbation spreads less in the network due to their lack of spiking.

We further quantify the localization of the CLVs using an inverse participation ratio (number) [51,52], which we define for the \(i\)th CLV as

\[
P_i^{-1} = \left( \sum_{j=1}^{N} v_{i,j}^2[V(t_k)] \right)^{-1}.
\]

Here \(\langle \cdot \rangle_k\) is an average over sufficiently many events and we use that the CLVs are normalized, \(\sum_{j=1}^{N} v_{i,j}^2[V(t_k)] = 1\). The participation ratio \(P_i\) measures how many components contribute to a vector. If, for example, the vector \(v_i(V(t_k))\)
FIG. 7. Stable and unstable CLVs are mostly confined to the subspaces of (perturbations to) LIF or XIF neurons, respectively. (a)–(d) The time evolution of the lengths of the projections of different CLVs onto the subspaces of LIF (blue) and XIF (red) neurons: (a) unstable CLV $v_6$, (b) stable CLV $v_{88}$, (c) unstable CLV $v_{25}$ (corresponding to the LE closest to the trivial one), and (d) trivial CLV $v_{26}$. (e), (f) The time evolutions of the projections of different CLVs onto the subspace of a single LIF neuron (neuron index $n = 5$, blue) and a single XIF neuron ($n = 95$, red): (e) unstable vector ($v_{15}$), (f) stable vector ($v_{85}$). Values are plotted at event times $t = t_k$. (g) The participation ratios $P_i$ of the unstable (purple), trivial (black), and stable (yellow) CLVs averaged over 10 000 events. The network is the same as in Figs. 2, 5(c), and 6. (h) The median $\tilde{P}$ of the participation ratios of the stable (yellow) and unstable (purple) CLVs, in networks as in panels (a)–(g) with $N = 100$ neurons but with different fractions of LIF and XIF neurons. Bars indicate the first and third quartiles of the distribution.

always has only one nonzero component, $P_i = 1$. If there are always $m$ nonzero components of equal size, $P_i = m$. We observe that the participation ratio of unstable CLVs increases approximately linearly with the number of XIF neurons starting with $P_i \approx 1$ at $N_{\text{XIF}} = 1$, consistent with a delocalization of these CLVs between the present XIF neurons; see Figs. 7(g) and 7(h). $P_i$ for stable CLVs increases likewise with the number of LIF neurons. The trivial CLV has a participation ratio close to $N$, because the components of the tangential vector $dV_j(t)/dt$ and thus the components of the CLV have roughly similar size.

Our mean-field approach uses the assumption that each LE is independently generated by the growth or shrinkage of a single-neuron perturbation, with negligible influence of the perturbation’s spread and backreaction in the network. Its suitability can now be understood as follows: The approximately $N_{\text{LIF}}$ stable CLVs are confined to the $N_{\text{LIF}}$-dimensional subspace of perturbations to LIF neurons. The stable CLVs thus form a basis of the subspace of perturbations to LIF neurons. Likewise the unstable CLVs form a basis of the subspace of perturbations to XIF neurons. At each time point, a perturbation to a single LIF neuron can therefore be expressed as a linear combination of stable CLVs, while a perturbation to an XIF neuron can be expressed as a linear combination of unstable CLVs. The stable CLVs have similar decay rates (negative LEs), and the unstable CLVs have similar growth rates (positive LEs); see Fig. 5. Any linear combination of only stable or only unstable CLVs inherits this decay or growth rate. This holds in particular for the perturbation of a single neuron. At each time point the perturbation to a single LIF or XIF neuron thus grows according to the negative or according to the positive LEs, respectively. The mean-field approach therefore yields good results.

VI. COMPUTATIONS WITH PRECISELY TIMED SPIKES

A. Network architecture and task design

In the following, we employ our networks for computations. In particular, we investigate how their different phase space structures and CLVs may be exploited in specific tasks. This requires a computational scheme based on precise spiking, which is affected by the phase space structure. We design a setup where one of our recurrent neural networks acts as a kind of computational reservoir [14,21,22], in the sense that it randomly nonlinearly filters its inputs. An output neuron receives the generated spikes and learns to generate desired outputs; see Fig. 8(a).

Inspired by experimental and computational neuroscience paradigms [53,54], we assume that the networks receive inputs from context neurons, whose spiking defines the computation to be executed in the specific trial and from input neurons. Their synaptic weights as well as the recurrent ones are static; only the output weights are learned. At the beginning of each trial, all membrane potentials are reset to
FIG. 8. Network for precise spike-based computations and solution of the XOR/AND task. (a) Network architecture. The recurrent network (middle, blue and red for LIF and XIF neurons) receives input spikes from context (top, black) and from input neurons (gray). The output neuron (bottom) changes its plastic weights (red) according to Finite Precision Learning to learn the task. (b) Spiking of the context and input neurons (top), voltage traces of recurrent neurons (middle, LIF neurons 1–7, XIF neurons 76–78, spikes highlighted by vertical lines) and voltage traces of the output neuron (bottom) after learning the XOR/AND task. Dashed lines indicate the times of possible context ($t_{\text{ctxt}}$) and input neuron ($t_{\text{in},+}, t_{\text{in},-}$) spiking as well as the possible desired output spike times ($t_{\text{out},+}, t_{\text{out},-}$). The output neuron sends its spike in the tolerance window around the desired time of the specific trial (gray rectangle). (c) Output weights $w_j$ from LIF (blue) and XIF neurons (red), $V_{\text{th}}$ (black line) and $V_{\infty, \text{LIF}}$ (gray dashed line), before (left) and after (right) learning. The weights have overall decreased during learning, while $V_{\text{th}}$ (black line) and $V_{\infty, \text{LIF}}$ have increased. The specific weight pattern after learning is crucial for executing the task; random weight shuffling leads to erroneous output spiking. (d) Overview of the eight spike patterns of the task after learning. In context 1, the system generates a temporal XOR computation, in context 2 a temporal AND computation. The output spikes are in the desired tolerance windows (gray rectangles) for all patterns.

The shapes of the postsynaptic potentials in our single-neuron dynamics are different from those in Ref. [30], and there is an additional constant driving term. The learning rule can be readily adapted to this: We consider $V_{\text{out}}(t) - V_{\text{th, out}}$ and cast it into the form $V_{\text{out}}(t) = V_{\text{th, out}} - \sum_{k=1}^{N+2} w_k x_k(t)$. Spikes are generated when $V_{\text{out}}(t) - V_{\text{th, out}}$ reaches zero. At each time $t$, we thus have a kind of perceptron classification task, where $w_j$, $V_{\text{th, out}}$ and $V_{\infty, \text{LIF}}$ are the “weights” to be learned. The “inputs” belonging to these weights are

\begin{align}
    x_j(t) &= \sum_{t_j < t} e^{-\gamma_{\text{LIF}}(t-t_j)} \\
    x_{N+1}(t) &= -\sum_{t_{\text{sp}} < t} e^{-\gamma_{\text{LIF}}(t-t_{\text{sp}})} - 1 \\
    x_{N+2}(t) &= 1 - e^{-\gamma_{\text{LIF}} t}.
\end{align}

Following Ref. [30], we assume a tolerance window of size $\varepsilon$ around each desired spike (we use $\varepsilon = 1$ ms throughout).
There are now two kinds of errors: (1) undesired spikes, i.e., spikes out of a tolerance window or second spikes within a tolerance window ($\text{Err} = 1$, the error time $t_{\text{err}}$ is the spike time) and (2) missing spikes within a tolerance window ($\text{Err} = -1$, $t_{\text{err}}$ is the end of the tolerance window). The dynamics are stopped at the first error, and $w_j$, $V_{\text{th,out}}$, and $V_{\text{sc,out}}$ are corrected according to the perceptron rule,

$$
\Delta w_j = -\eta \text{Err} \sum_{k:j<k<\text{Err}} e^{-\gamma \|w\|_1 (t_k - t_j)},
$$

$$
\Delta V_{\text{th,out}} = \eta \text{Err} \left( \sum_{j:p<j<\text{Err}} e^{-\gamma \|w\|_1 (t_p - t_j)} + 1 \right),
$$

$$
\Delta V_{\text{sc,out}} = -\eta \text{Err} (1 - e^{-\gamma \|w\|_1 t_j}),
$$

with learning rate $\eta$ (we use $\eta = 0.01$). To focus on networks with inhibitory neurons throughout the article, we restrict the output weights to be inhibitory by clamping them at zero when they would become excitatory during learning. We note that a missed spike generates increases in $w_j$ and $V_{\text{sc,out}}$ as well as a decrease in $V_{\text{th,out}}$ to foster spiking. If an undesired spike occurs, the signs are reversed.

### B. Switchable temporal XOR/AND

We exemplarily consider two tasks. In the first, the network of Fig. 8(a) learns to execute in context 1 a temporal XOR and in context 2 a temporal AND computation; see Figs. 8(b)–8(d). The weights from context and input neurons to the recurrent network are drawn randomly from the uniform distribution over $[2C, 0]$. At the beginning of a trial, at $t = 0$ ms, context neuron 1 or 2 sends a spike, specifying the context. Thereafter each input neuron sends a spike, either at time $t_{\text{in,}} = 5$ ms ("+"-input) or at $t_{\text{in,}} = 10$ ms ("-"-input). The desired output spike is at $t_{\text{out,}} = 15$ ms ("+"-output) or at $t_{\text{out,}} = 20$ ms ("-"-output), depending on the context and the input spike times.

The considered networks learn the task easily, whether the reservoir consists of LIF or XIF neurons or of a mixture of both. The example with a mixed network displayed in Fig. 8 [the same network as in Figs. 2, 5(c), 6, and 7] required 53 learning cycles, where in each cycle the four input-desired output patterns of both contexts were presented. The networks cannot learn the task, if the recurrent network dynamics at the desired output times are too similar for different contexts and input conditions. This happens for recurrent LIF networks, if the context or input neurons have coupling strengths that are so weak that the perturbations due to different input timing are small. The states are then within the same flux tube and the perturbation decays up to a time shift. In XIF and mixed networks, the recurrent dynamics are too similar if there is insufficient time for the perturbation to grow and spread before the first desired output.

### C. Detect or ignore input time differences

In the second task, the system has to ignore a difference in input timing in context 1 and to detect it in context 2. The network setup is as in Fig. 8(a), except that there is only one input neuron. This sends a spike at $t_{\text{in,}} = t_1 - \Delta t$ or at $t_{\text{in,}} = t_1 + \Delta t$ ($t_1 = 1$ ms, $\Delta t = 0.1$ ms). The output neuron shall generate in context 1 for both input conditions the same output, a burst of four spikes at $t = 105$ ms, 110 ms, 115 ms, and 120 ms. In context 2 it shall detect the difference and highlight it by sending one spike at $t = 100$ ms (input at $t_{\text{in,}}$) or two spikes at $t = 130$ ms and 135 ms (input at $t_{\text{in,}}$). For this task, for simplicity we assume that the impacts of input neurons do not depend on the membrane potential, i.e., for them $h_i(V_j) = 1$. Further, we allow context and input weights to be excitatory and inhibitory.

We find that networks with the previously chosen random parameters of external weights drawn from $[2C, 0]$ usually cannot solve the task (criterion: no convergence within 50,000 cycles). The reason is different for pure LIF reservoirs and for reservoirs containing XIF neurons: In a pure LIF reservoir, the small difference in input times leads to state perturbations that are usually in the same flux tube. These decay to a temporal shift until the time of the desired outputs. The readout neuron thus cannot learn to generate two different output patterns as required in context 2. In presence of XIF neurons, the dynamics are locally unstable. The small input difference is amplified in both contexts, and the reservoir spiking is different for all four patterns at the times of the desired output. The network therefore has to learn four input-output relations with 11 output spikes and silence periods in between, without being able to take advantage of the fact that two of the four output patterns are identical. This typically exceeds its learning capacity. We also observe for our parameters that the dynamics of the pure LIF reservoir can leave its flux tube due to the perturbation. If this happens only for context 2, the system can often learn the task.

To solve the problem, we design the network such that, reliably, in context 1 but not in context 2 the input differences leave the reservoir spiking at the time of the desired outputs largely unaffected. This can be achieved by choosing the context and input couplings such that the input difference generates a state perturbation along a stable CLV of the reservoir dynamics in context 1. In contrast, for context 2 the state perturbation should have a component in the direction of an unstable CLV such that it is quickly amplified. The setup requires mixed networks with both types of CLVs. We note that an alternative approach might exploit the dichotomy of large and small perturbations, which do and do not leave the flux tubes of pure LIF networks.

To derive appropriate weights, we compute the state perturbations in the reservoir assuming that in the “unperturbed” system the input arrives at $t_1$. We there have

$$
V_j(t_1^+) = V_j(t_1^-) + C_j^m,
$$

where $C_j^m$ is the coupling strength from the input neuron to neuron $j$. In the “perturbed” system, the input arrives shifted by $\delta t$ (here $\delta t = \pm \Delta t$), such that we have in linear approximation

$$
\tilde{V}_j(t_1 + \delta t^+) = V_j(t_1^-) + \tilde{V}_j(t_1^-) \delta t + C_j^m.
$$

To compute a perturbation in the $V_j(t_1^+)$ that corresponds to the perturbation due to the temporal shift of input, we propagate the perturbed potential in linear approximation
from $t_1 + \delta t^+$ to $t_1^+$,
\[ \delta V_j(t^+) \approx \hat{V}_j(t_1 + \delta t^+) - \hat{V}_j(t_1 + \delta t^+) \delta t - V_j(t_1^+) \]
\[ \approx \hat{V}_j(t_1^+) \delta t - \left\{ - (\gamma_j) [V_j(t_1^+) + C_{in}^j] + I_{ext}^j \right\} \delta t \]
\[ = \gamma_j C_{in}^j \delta t. \] (38)

A temporal input difference that should be ignored should be proportional to a stable CLV $v_i$ at the state $V(t_1^+)$,
\[ \gamma_j C_{in}^j \propto v_i J[V(t_1^+)]. \] (39)

We choose the same recurrent network as in Figs. 2, 5(c), 6, and 7 and the same CLV as in Fig. 6(d) at $t = 0$ ms, i.e., $i = 90$ and
\[ V(t_1^+) = V_0, \] (40)
where $V_0$ is the state at which the vector was recorded. Context input 1 determines the state at $t_1^+$ by fixing the initial conditions of the dynamics. We choose as context input weights
\[ C_{ext.1}^j = V_j(0^+) = e^{\gamma_j t}(V_0 C_{in}^j + V_\infty(1 - e^{\gamma_j t})), \] (41)
which lead to Eq. (40) after free propagation until $t_1$ and receiving of the input $C_{in}^j$. To ensure that the perturbation in context 2 has a component in the direction of an unstable CLV, it suffices to choose a random context weight vector, such that $V(t_1^+) \neq V_0$ and $v_i(V_0)$ is typically not a stable CLV or a linear combination of stable CLVs at the state $V(t_1^+)$. We randomly permute the entries of $C_{ext.1}$ to obtain $C_{ext.2}$.

We find that the network constructed in this way can reliably learn the task. The example displayed in Fig. 9 uses a proportionality factor of 0.01 in Eq. (39); the output weights converged after 146 cycles.

VII. DISCUSSION

In the present article we investigate the spiking and membrane potential statistics, the stability properties, and the phase space structure of mixed networks containing conventional LIF neurons and XIF neurons with a convex rise function. The recurrent connections are inhibitory, and the synaptic currents have an infinitesimal temporal extent. We employ two analytical mean-field approaches, one for the statistics and one for the dynamical stability properties; numerical simulations yield additional features of the dynamics and a better understanding of the analytical approximations. Finally, we apply the networks for computation with spikes, exploiting our insights into the dynamics.

We investigate networks in the balanced state. To establish it in our networks, we introduce a voltage dependence in the XIF neuron inputs: below a certain potential, further input has no impact. This simple model of a conductance-based synapse prevents XIF neurons from switching off and provides a good-natured nonlinearity, which leaves the dynamics analytically tractable.

The balanced state is typically investigated using spiking network models with an excitatory and an inhibitory neuron population or with a single population of hybrid excitatory-inhibitory or inhibitory neurons [3,4,17,42,55]. While detailed models of small circuits with specific abilities such as central pattern generators commonly consider multiple neuron types [56], studies on the impact of mixed populations of multiple neuron types on the collective dynamics of larger networks are rare. Reference [57] simulated networks with excitatory and inhibitory populations containing resonator-and integrator-type neurons. These mixed networks both persistently generated activity and quickly changed their overall rate in response to inputs, thereby combining abilities of their pure counterparts. References [58,59] considered models for working memory and visual processing with different types of interneurons that were grouped into distinct populations with different connectivities.

We characterize the balanced dynamics of inhibitory mixed LIF and XIF networks first from a statistical perspective, adopting a shot-noise approach, which accounts for the finite input rate and finite size of individual inputs [34,45–47]. We extend this approach to XIF neurons and derive their steady-state firing rate and voltage probability density. In contrast to the case of LIF neurons, the final continuity equation needs to be integrated numerically, due to the nonlinearity in the XIF input. We apply the results to obtain neuron parameters that lead to homogeneous firing rates for our further considered networks. We insert these rates into the mean-field expressions of the Lyapunov exponents (LEs) and thus analytically determine the dynamical stability properties of the network.

![FIG. 9. Network trained to detect or ignore input time differences, after learning. (a) Overview of the four spike patterns of the task. In context 1, the network ignores the small time difference in the input (right subpanels: closeups around input times); in context 2 it detects and highlights it by generating different numbers and timings of output spikes. After learning, the output spikes are in the desired tolerance windows (gray rectangles, appearing as lines at the displayed timescale) for all patterns. (b) Top] The spiking dynamics in the recurrent reservoir (green and yellow dots) are in context 1 similar for both input times, due to the chosen context and input weights. This generally fosters and here enables learning of the same output. (Middle) The temporal differences $\delta t$ between reservoir spikes display the typical pattern of first shrinkage then growth of perturbations along stable CLVs; cf. Fig. 6(d). (Bottom) The different spiking dynamics in the recurrent reservoir for different input times in context 2 allow the generation of different output.](output image)
While networks of LIF neurons have stable dynamics [8–11,14], we find that already one XIF neuron gives rise to a positive largest LE indicating chaos, in contrast to the robustness against introducing excitatory connections [9,11]. We give an analytical argument for this and expand it to a mean-field estimate of the entire Lyapunov spectrum. Simply put, the destabilizing effect of excitatory inputs will be compensated by receiving inhibitory ones, if the latter dominate and the period of spiking is overall increased compared to the free neurons. If one introduces an XIF neuron, there is nothing which could counteract the increase of its perturbation through inhibitory input other than an unlikely network back-reaction triggered by its perturbed output spikes. We note that in the phase representation of LIF neurons used in Ref. [11], in contrast to our voltage representation an excitatory input explicitly increases a perturbation, while an inhibitory input decreases it, unless the excitatory input is suprathreshold [60,61].

While computing the largest LE is a standard procedure, few studies have so far obtained a large part or the entire spectrum of balanced spiking dynamics. They considered a single homogeneous or an excitatory and an inhibitory neuron population [14,17,50,62–64]. We analytically and numerically obtain the full spectrum for mixed networks of inhibitory LIF and XIF neurons. Interestingly, we find that it separates into two parts, in contrast to the ones reported previously, including those of networks with separate excitatory and inhibitory populations. Furthermore, we compute the covariant Lyapunov vectors (CLVs) of the dynamics [5,6]. They provide us with further insight into the phase space structure and the approximations underlying the mean-field analysis of LEs. The stable (unstable) CLVs are approximately aligned to the subspace of perturbations to LIF (XIF) neurons.

Our mean-field analysis predicts that the number of negative (positive) LEs is equal to the number of LIF (XIF) neurons. Since the underlying arguments do not depend on the neurons’ specifics, we expect this to hold for any types of neurons with purely concave and convex rise functions. The mean-field analysis further indicates that the size of the LEs is approximately given by the strength of the leak and the quotient of free and actual spike frequency. The LEs are thus largely independent of the collective dynamics but rather reflect properties of individual neurons. This implies in particular that the typical perturbation growth rate does not change with network size. It further implies that in the balanced state, where the ratio of actual and free spike rate is low, the LEs are mainly determined by the single-neuron leak strengths; see Ref. [14] for a similar finding in large networks of LIF neurons with high indegree. The result is a consequence of the linear subthreshold dynamics of the neurons, which imply that the increase or decrease of a perturbation is independent of the state of the neuron when receiving a spike. We note that Ref. [65] defined the Lyapunov spectrum as consisting of mean-field LEs in a numerical study on LIF neuron networks.

Our numerical computations of the Lyapunov spectrum show that the mean-field result is a good approximation. We explain this by analyzing the CLVs. Furthermore, we derive an exact expression for the change of phase space volume, which agrees with the mean-field result.
computational capabilities that can be selected with different input vectors. While we have chosen the input weights by hand, plasticity rules for spiking networks in the brain as well as future artificial ones may allow finding them by learning.

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APPENDIX A: VOLTAGE PROBABILITY DISTRIBUTION OF LIF NEURONS

In the following, we further discuss the discrepancy between the voltage probability density of an LIF neuron obtained by the shot-noise approach and the one observed if the inputs are spike trains recorded in the network of Fig. 2; cf. Figs. 3(a) and 10(a). The analytical density obtained by the shot-noise approach, Eq. (9), with rate given by Eq. (18) matches that of an LIF neuron receiving a Poisson spike train with the same rate and spike impact strength; see Fig. 10(b). Hence, we can attribute the observed discrepancy for LIF neurons with network spike train input to deviations of the spike trains’ rate and the assumed Poisson statistics.

We expect that the discrepancy is mainly caused by spatial correlations that arise in a rather dense network of $N = 100$ neurons with an indegree of $K = 50$. To substantiate this we reduce the correlations in two ways: First, we use spike trains from a sparse network with $N = 10,000$ and $K = 50$ to generate the neuron input. Second, we randomly shift the individual spike trains of the original $N = 100$ network in time before superposing them to generate the input; this eliminates spatial correlations while keeping the temporal correlations of the individual spike trains intact. Figures 10(c) and 10(d) show that both manipulations strongly reduce the discrepancy to the analytical density. Some of the remaining discrepancy is due to the difference between the network spike rate and the result of Eq. (18); see Fig. 10(d).

Finally we explore the impact of the reduced variability of the interspike intervals. For this, we use Poisson and Gamma process input spike trains. The latter are completely characterized by their rate and the coefficient of variation of the interspike-interval distribution, which we match to those of the superposed spike trains of the network of Fig. 2. The quality of approximation increases when taking into account the reduced variability; see Fig. 10(e). Also if the input is a superposition of shifted spike trains, accounting for it still slightly improves the similarity between the resulting $p(V)$; compare Figs. 10(f) and 10(d) (which matches the result for Poisson input).

APPENDIX B: MEAN-FIELD LYAPUNOV EXPONENTS

In the following we compute the mean-field LEs. For this, we describe the dynamics by a sequence of discrete maps from the state at a time (strictly speaking: infinitesimally) shortly after generation of a spike to the state at a time shortly after generation of the next spike. We take a stroboscopic map approach; i.e., the times remain unchanged if a small perturbation is applied to the dynamics. The dynamics of small perturbations are encoded in the Jacobian matrices at each time point. Specifically, for our discrete description we need the single-spike Jacobians [14,17] $J(k)$. They generally describe the linear evolution of infinitesimal perturbations.
The growth rate of perturbations and thus the mean-field LE are given by the long-term average of Eq. (B8),
\[
\lambda_{\text{mf}} = \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} |J_{\text{mf}}(k)| = \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} \gamma \rho_{\text{tree}} - \gamma = -\gamma \left(1 - \frac{\rho}{\rho_{\text{tree}}} \right).
\]

**APPENDIX C: NETWORK SINGLE-SPIKE JACOBIAN**

The components of the single-spike Jacobian \( J(k) \) are given by
\[
J_{ij}(k) = \frac{\partial V_i(t_{k+1}^+)}{\partial V_j(t_k^+)}.
\]

To compute them, as in our mean-field approach we need to take into account the decay of perturbations between spikes as well as the reset of the neuron sending the \((k + 1)\)th spike, say, neuron \( l \). \( V_l(t_{k+1}^-) = V_{th} \) implies an implicit dependence of the spike time \( t_{k+1} \) on \( V_l(t_k^+). \) Additionally, we now have to include the jumplike potential change by \( C_l h_l[V_l(t_{k+1}^-)] \) that the spike induces in neuron \( i \), such that
\[
V_i(t_{k+1}^+) = V_i(t_k^+)e^{-\gamma(t_{k+1}^- - t_k)} + \frac{\text{ext}}{\gamma} [1 - e^{-\gamma(t_{k+1}^- - t_k)}] + C_l h_l[V_l(t_{k+1}^-)] - \delta_i V_{th}.
\]

The stroboscopic description yields a dependence
\[
\delta V_i(t_{k+1}^+) = e^{-\gamma(t_{k+1}^- - t_k)} \delta V_i(t_k^+) + \frac{\text{ext}}{\gamma} \left[1 - e^{-\gamma(t_{k+1}^- - t_k)}\right] + C_l h_l[V_l(t_{k+1}^-)] - \delta_i V_{th}.
\]

The stroboscopic description yields a dependence
\[
\delta V_i(t_{k+1}^+) = e^{-\gamma(t_{k+1}^- - t_k)} \delta V_i(t_k^+) + \frac{\gamma}{\rho_{\text{tree}}} \delta_i V_{th} - C_l h_l[V_l(t_{k+1}^-)]
\]

of the perturbation \( \delta V_i(t_{k+1}^+) \) on the perturbations at the state at \( t_k^+ \). This is analogous to Eq. (B5), with the difference that the neuron that sends the spike and determines the shift in \( t_{k+1} \) (neuron \( l \)) may be different from neuron \( i \). The Jacobian thus reads
\[
J_{ij}(k) = \delta_{ij} e^{-\gamma(t_{k+1}^- - t_k)} + \delta_j \left[ V_l(t_{k+1}^-) - V_l(t_{k+1}^+) \right] \frac{\partial h_{i|l}}{\partial V_l(t_k^+)}.
\]

Application of the implicit function theorem and inserting Eqs. (1), (2), (3), (4), and (5) results in
\[
J_{mf}(k) = \frac{V_i(t_{k+1}^+)}{V_l(t_k^+)} e^{-\gamma(t_{k+1}^- - t_k)} = \exp \left[ \frac{\gamma}{\rho_{\text{tree}}} - \gamma (t_{k+1}^- - t_k) \right].
\]

**APPENDIX D: VOLUME CONTRACTION**

The volume expansion and the sum of LEs are given by the time-averaged logarithms of the determinants of the Jacobians
\[
\lambda_{\text{mf}} = \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} \ln |J_{\text{mf}}(k)| = \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} \gamma \rho_{\text{tree}} - \gamma = -\gamma \left(1 - \frac{\rho}{\rho_{\text{tree}}} \right).
\]

for an LIF or an XIF neuron \( i \).

We note that another, equivalent derivation of \( J(k) \) first computes the voltages at a fixed time \( t' \) between \( t_{k+1} \) and \( t_{k+2} \) in terms of the voltages at another fixed time \( t \) between \( t_k \) and \( t_{k+1} \). Taking derivatives leads to the Jacobian for the dynamical evolution from \( t \) to \( t' \). The limits \( t \searrow t_k \) and \( t' \searrow t_{k+1} \) then yield \( J(k) \).
We thus have
\[
\sum_{i=1}^{N} \lambda_i = \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} \ln |\det J(k)| \quad (D1)
\]
in terms of single-spike Jacobians [14]. The specific form of our \( J(k) \) allows us to split it into a diagonal matrix \( \hat{J}(k) \) covering the perturbation change during subthreshold evolution and a rank one correction,
\[
J(k) = \hat{J}(k) + u(k)v(k)^T. \quad (D2)
\]
where
\[
\hat{J}_{ij}(k) = \delta_{ij} e^{-\gamma(t_{i+1} - t_k)}, \quad (D3)
\]
\[
v_j(k) = \delta_{j}, \quad (D4)
\]
\[
u_i(k) = \frac{\delta_{i} V_{th} - C_{il} h_i [V_j(\hat{t}_{k-1}^l)]}{\gamma_i - V_j(\hat{t}_{k}^i)} \quad (D5)
\]
The matrix determinant lemma now allows us to compute \( \det J(k) \) via
\[
\det J(k) = (1 + v(k)^T \hat{J}(k)^{-1} u(k)) \det \hat{J}(k). \quad (D6)
\]
Equation (B4) and the relation \( 1 + V_{th}/(Q_j^{ext}/\gamma_i - V_{th}) = \exp(\gamma_i/\tilde{\rho}_{free,i}) \) for the free spike frequency \( \tilde{\rho}_{free,i} \) of neuron \( j \) [see Eqs. (4) and (5)] lead to
\[
\det J(k) = \exp\left[ \frac{\gamma_i}{\tilde{\rho}_{free,i}} - (\sum_{l=1}^{N} \gamma_l) (t_{k+1} - t_k) \right] \quad (D7)
\]
Time averaging yields
\[
\sum_{i=1}^{N} \lambda_i = \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} \ln \det J_k
\]
\[
= \lim_{L \to \infty} \frac{1}{L} \sum_{k=0}^{L-1} \frac{\gamma_i(k)}{\tilde{\rho}_{free,i}(k)} - \sum_{j=1}^{N} \frac{\gamma_j}{\tilde{\rho}_{free,j}}
\]
\[
= -\sum_{j=1}^{N} \gamma_j \left( 1 - \frac{\rho_j}{\tilde{\rho}_{free,j}} \right), \quad (D8)
\]
where the index \( i(k) \) denotes the neuron that spikes at time \( t_k \) and \( \rho_j \) is the spike rate of neuron \( j \) in the network.

**APPENDIX E: DEPENDENCE OF THE LYAPUNOV SPECTRUM ON INDEGREE AND NETWORK SIZE**

The rule that the number of negative (positive) LEs approximately equals the number of LIF (XIF) neurons holds independent of \( N \) and \( K \); see Fig. 11. Figures 11(a)–11(c) indicate that for large \( N \) and sufficiently large fixed indegree the Lyapunov spectrum assumes a fixed shape, which differs from the result of our mean-field approach. This is because the mean-field approach neglects the nonzero off-diagonal entries of the single-spike Jacobians, whose strength and average number \( K \) do not depend on \( N \); see Appendix C. The shape of the Lyapunov spectrum varies with the indegree of the network. For larger ratios \( K/N \) the positive and negative parts of the Lyapunov spectrum become flatter; cf. Fig. 11(d).

**FIG. 11.** Lyapunov spectra of networks with different sizes and indegrees. (a) Lyapunov spectra for different network sizes \( N \) and constant indegree \( K = 50 \) (numerical results: color coded dots; result of the mean-field theory: horizontal black lines). (b), (c) Close-ups of the positive and negative parts of the spectra in panel (a). (d) Lyapunov spectra for different indegrees \( K \) and size \( N = 500 \) (numerical results: color coded dots; results of the mean-field theory: color coded dashed lines). Remaining network parameters are as in Fig. 2.

Note, however, that also the spiking becomes more regular. We observe for very sparse but still strongly connected networks that the Lyapunov spectrum is no longer well approximated by our mean-field theory; cf. Fig. 11(d).

**APPENDIX F: COMPUTING COVARIANT LYAPUNOV VECTORS**

We compute the CLVs in a dynamical manner [5,51]. In short, if we want to compute them at \( t = 0 \), we start sufficiently long before with an arbitrary set of \( N \) orthonormal vectors \( q_i(0) \), which forms a basis of the tangent space. We evolve this basis forward until zero and further to a sufficiently long time \( \tau_f \), using the single-spike Jacobians. Every few steps, we reorthonormalize the basis. The orthogonalizations leave the first vector \( q_1(t) \) unchanged. It thus evolves freely (up to normalization) until it has aligned with the first covariant Lyapunov vector at \( t = 0 \) and thus also at \( t = \tau_f \). The second vector, \( q_2(t) \), is kept orthogonal to \( q_1(t) \). Since it otherwise evolves freely, \( q_2(0) \) will lie in the subspace of the first and the second CLV at \( t = 0 \), which are in general not orthogonal; the same holds for \( q_3(t_f) \) at \( t = \tau_f \). Analogously \( q_3(0) \) will be in the subspace of \( v_1(0), v_2(0), \) and \( v_3(0) \), and so on. As noted in Sec. IV D, the growth rates of the vectors...
already yield the LEs. In order to find the CLVs one uses the time reversal property: we evolve the vectors \( q_i(t_f) \) along the previously taken forward trajectory back in time until \( t = 0 \). During this, we keep them restricted to their respective subspaces, which are known from the forward propagation. The vectors will then align with the least expanding directions of their subspaces, so the backpropagated \( v_2 \) will align with \( q_2(t_f) \) along \( t_f \) with \( v_2 \), and so on. We concretely implemented the simple and efficient algorithm derived in Ref. [51], which performs the backpropagation by representing and mapping the vectors in terms of their components in the bases \( q_i(t) \). After obtaining the CLVs at \( t = 0 \), those in the not too distant future can be obtained using Eq. (27).

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