Dynamics of networks of excitatory and inhibitory neurons in response to time-dependent inputs

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1 INTRODUCTION

Networks of neurons in the central nervous system are driven by stimuli that vary on a wide range of time scales, and need to encode these stimuli by the pattern of firing of their constituent neurons. To understand how this encoding is performed, one needs to understand the relationship between the input to the network (the set of spike trains of all neurons that are presynaptic to a given network) and its output (the set of spike trains of all neurons in the network). This is a question that is extremely hard to address experimentally, since one has typically access only to a very small fraction of neurons in a network (with the retina being a notable exception Pillow et al., 2008), or access to variables that only reflect population activity, such as the multi-unit activity (MUA) or local field potential (LFP). It is also very hard to address in general in theoretical models, since inputs and outputs live in a high-dimensional space.

A simpler question, and a useful first step, is to understand the relationship between the mean inputs to a network and its population firing rate (i.e., average instantaneous probability that a neuron in the network emits an action potential). However, even this question can be difficult to answer in the space of all possible time-dependent inputs. A natural strategy to study this question is to start with time-independent inputs. One then computes how the mean population rate depends on the mean input, and therefore oscillate with the same frequency as the input. One can then characterize the dynamics of the network with only two quantities: the amplitude and the phase of the population firing rate modulation as a function of frequency. These two quantities together characterize the transfer function of the network, and can be used to reconstruct the dynamics of the population in response to arbitrary time-dependent inputs, provided the amplitude of the time-dependent variations in the input is sufficiently small.

In the present paper, we investigate systematically the properties of this transfer function as a function of the connectivity of the network, for networks composed of two-populations, one excitatory (E), one inhibitory (I). Such networks have been extremely popular as simplified models of local networks in neocortex (Wilson and Cowan, 1972; van Vreeswijk and Sompolinsky, 1996; Amit and Brunel, 1997), hippocampus (Tsodyks et al., 1997), as well as other structures. Surprisingly however, extensive characterization of this transfer function is lacking, even for simple models. In fact, the goal of the present paper is to study this question systematically in two classes of models (1) “firing rate” models in which individual neurons are not described explicitly, rather all dynamical variables describe population quantities like average firing rates or average synaptic inputs (Wilson and Cowan, 1972; Ermentrout, 1998; Dayan and Abbott, 2001) (2) fully connected networks of leaky
integrate-and-fire (LIF) neurons, in the presence of white noise (Brunel and Hansel, 2006). These two types of models have the advantage that the transfer function can be computed analytically, as a function of network parameters, provided the network is in a so-called “asynchronous state,” i.e., a state in which the population firing rate goes to a stable fixed point for time-independent inputs.

For each network model, we first characterize the region of stability of asynchronous states in the space of the parameters characterizing the coupling. Instabilities can arise through four qualitatively different mechanisms, which have been characterized extensively: (i) the “rate” instability (when the firing rate of the network can no longer be held at a given value, because of the excitatory feedback), present in all models; (ii) an oscillatory instability due to the E–I loop (Brunel and Wang, 2003), also present in both models; (iii) an oscillatory instability due to the delays in the I–I connections (Brunel and Hakim, 1999, 2008; Brunel and Wang, 2003; Brunel and Hansel, 2006), present in both models, provided delays are present in I–I connections, and finally; (iv) instability due to single neuron resonances at integer multiples of their firing rates, when the noise level is sufficiently small (Abbott and van Vreeswijk, 1993; Hansel and Mato, 2003; Brunel and Hansel, 2006). The last type of instability is only present in networks of spiking neurons.

We then characterize regions with qualitatively different types of transfer functions, in the space of coupling parameters, in both models. Transfer functions are characterized both in terms of the behavior of the gain as a function of frequency (i.e., monotonically decaying as a function of frequency, or with a single or multiple resonant peaks), and in terms of the phase relationship between E and I cells, again as a function of frequency. We relate qualitatively different types of transfer function with nearby instabilities. Finally, we compare the behaviors of rate model and network of integrate-and-fire neurons, and point out similarities and differences between both models.

2 MODELS

We investigate the dynamics of a network composed of two-populations of neurons, one excitatory (E), the other inhibitory (I; see Figure 1). The network received average, time-dependent inputs \( \mu_{E,\text{ext}} \) and \( \mu_{I,\text{ext}} \) to E and I populations, respectively. All types of connections are potentially present in the model, with strengths \( J_{EE} \) (E → E), \( J_{E,I} \) (E → I), \( J_{I,E} \) (I → E), \( J_{I,I} \) (I → I). We consider in this paper two types of models: a rate model and a network of LIF neurons.

2.1 RATE MODELS

In the simplest rate model, the instantaneous firing rates at time \( t \), \( r_{E,I}(t) \) of E and I populations, are given by (see, e.g., Wilson and Cowan, 1972; Tsodyks et al., 1997; Ermentrout, 1998; Dayan and Abbott, 2001; Murphy and Miller, 2009):

\[
\tau_e \frac{dr_e(t)}{dt} = -r_e(t) + F_{E,E}(\mu_{E,\text{ext}}(t) + J_{EE} r_e(t) - J_{E,I}(t))
\]

\[
\tau_i \frac{dr_i(t)}{dt} = -r_i(t) + F_{I,I}(\mu_{I,\text{ext}}(t) + J_{I,E} r_e(t) - J_{I,I}(t))
\]

where \( \tau_{E,I} \) characterize the time constant of firing rate dynamics, while \( F_{E,I}(x) \) are the static transfer functions that describe the relationship between a mean synaptic input and an average firing rate in stationary conditions. Various types of response functions \( F_{E,I}(x) \) have been used in the literature. Here, in order to simplify the analysis, we take this function to be threshold-linear (see, e.g., Ben-Yishai et al., 1995; Hansel and Sompolinsky, 1998; Roxin et al., 2005)

\[
F_{E,I}(x) = \begin{cases} 
    x & \text{if } x \geq 0 \\
    0 & \text{else}
\end{cases}
\]

In the simplest rate model [Eqs (1,2)], synaptic time constants are ignored. Another class of rate models assumes instantaneous firing rate dynamics, and dynamical variables are assumed to represent average synaptic currents (Ermentrout, 1998). Since synaptic time constants are known to play an important role in shaping the dynamics of the activity of neurons, we investigate a rate model in which both neuronal and synaptic time constants are present. Real synaptic currents are characterized by three qualitatively distinct time constants: a latency (interval between pre-synaptic spike and the beginning of the post-synaptic response), a rise time, and a decay time (see, e.g., Destexhe et al., 1998). The simplest way to implement such synaptic currents in a rate model is through two synaptic variables \( s_{ab}(t) \) and \( x_{ab}(t) \) (where \( a, b = E, I \) denote the post/pre-synaptic population, respectively) whose dynamics are described by the following equations

\[
\tau_{ab} \frac{ds_{ab}(t)}{dt} = -s_{ab}(t) + r_a(t - D_{ab})
\]

\[
\tau_{ab} \frac{dx_{ab}(t)}{dt} = -x_{ab}(t) + s_{ab}(t)
\]

where \( D_{ab}, \tau_{ab}, \tau_{ab} \) are the latencies, rise times, and decay times of synaptic currents from population \( b \) to population \( a \) (see, e.g., Brunel and Wang, 2003). In this model, Eqs (1,2) are replaced by

\[
\tau_a \frac{dr_a(t)}{dt} = -r_a(t) + [\mu_{a,\text{ext}}(t) + J_{a,b} s_{ab}(t) - J_{a,a} x_{ab}(t)]
\]
where \( a = E, I \). Note that Eqs (1,2) are recovered when \( D_{ab} = 0 \), \( \tau_{\text{ras}} = 0, \tau_{\text{sab}} = 0 \) for all \( a, b \).

We also consider a version of the model where only the latency is described (Roxin et al., 2005). In this case, Eqs (4,5,6) are replaced by

\[
\tau_a \frac{dr_a(t)}{dt} = -r_a(t) + \mu_{a,\text{ext}}(t) + \sum_{i} \mu_{a,i}(t) - j_{a,A}(t) - j_{a,B}(t) \tag{7}
\]

This model corresponds to a special case of Eqs (4–6) with \( \tau_{\text{ras}} = \tau_{\text{sab}} = 0 \) for all \( a, b \).

### 2.2 NETWORK OF LEAKY INTEGRATE-AND-FIRE NEURONS

In this model E and I populations are composed of \( N_E, N_I \) LIF neurons (see, e.g., Tuckwell, 1988; Dayan and Abbott, 2001), with \( N_E + N_I = N \). Each neuron \( i \) of population \( a \) is described by its membrane potential \( V(t) \), which obeys

\[
\tau_a \frac{dV(t)}{dt} = V_{\text{rest}} - V_i(t) + \mu_{a,\text{ext}}(t) + \frac{j_{a}}{N_{E,i = E}} \sum_{j} s_{ij}(t) - j_{a}(t) - \sigma \eta(t) \tag{8}
\]

where \( \tau_{\text{mem}} \) is the membrane time constant of neurons in population \( a = E, I \), \( V_{\text{rest}} = -70 \text{ mV} \) is the resting membrane potential, \( \sigma \eta(t) \) is a noise term modeling the variability in external inputs received by neurons in addition to the “signal” \( \mu_{a,\text{ext}}(t) \), where \( \sigma \) is the standard deviation of the noise and \( \eta(t) \) is a white noise of unit variance, uncorrelated from neuron to neuron. Finally, \( s_{ij}(t) \) are individual synaptic currents from neuron \( j \in b = E, I \) to neuron \( i \in a = E, I \), modeled as in the rate model with the help of an additional synaptic variable \( x_{ij} \):

\[
\tau_{\text{ras}} \frac{dx_{ij}(t)}{dt} = -x_{ij}(t) + \tau_{\text{sab}} \sum_{k} \delta(t - t_k) - D_{ab} \tag{9}
\]

\[
\tau_{\text{sab}} \frac{ds_{ij}(t)}{dt} = -s_{ij}(t) + x_{ij}(t) \tag{10}
\]

where again \( D_{ab}, \tau_{\text{ras}}, \tau_{\text{sab}} \) are the latencies, rise times, and decay times of synaptic currents from population \( b \) to population \( a \). The sum \( \delta(t) \) represents a sum over all spikes of pre-synaptic neuron \( j \), that occur at times \( t_k \). Note the factor \( \tau_{\text{sab}} \) in front of this sum, which is chosen such that (i) both \( x \) and \( s \) variables are dimensionless; (ii) the total charge of individual PSCs is independent of synaptic time constants. In this model, action potentials occur when the voltage crosses \( V_{\text{reset}} = -50 \text{ mV} \). At the time of the action potential, the voltage is reset at \( V_{\text{reset}} = -60 \text{ mV} \). For simplicity, no refractory period is included.

The instantaneous firing rate of each population \( r_{E,I}(t) \) is given by the sum of the number of spikes of the corresponding population in a time window \( dt \)

\[
r_{E,I}(t) = \frac{1}{N_{E,I}} \sum_{k} \delta(t - t_k) \tag{11}
\]

In numerical simulations, \( r_{E,I}(t) \) is computed from simulating a network of \( N_E = 2000 \) and \( N_I = 500 \) neurons, except in Figure 9 where \( N_E = 8000 \) and \( N_I = 2000 \), using a Runge–Kutta second order method with a time step \( dt = 0.01 \text{ ms} \). In the mean-field analysis, one takes the limit \( N \to \infty, dt \to 0 \) (see below).

### 2.3 EXTERNAL INPUTS AND RESPONSE OF THE NETWORK

In this paper, we consider external inputs of the form

\[
\mu_{a,\text{ext}}(t) = \mu_{a,\text{ext}}(t) + \mu_{a,\text{int}}(t) \tag{12}
\]

where \( \mu_{a,\text{ext}} \) is a constant external drive, and \( \mu_{a,\text{int}} \) is the amplitude of a sinusoidal modulation in the average input at a frequency \( \omega = 2\pi f \).

We assume \( \mu_{a,\text{int}} \) is sufficiently small so that the response of the network firing rate is linear in \( \mu_{a,\text{int}} \). If this assumption is satisfied, then the firing rate response is given by

\[
r_{a}(t) = r_{a,0} + \text{Re}(r_{a,\omega} e^{i \omega t}) \tag{13}
\]

where \( r_{a,0} \) is a constant firing rate in absence of time-dependent modulation in the input (in other words, the background activity of the network), while \( r_{a,\omega} \) describes how the instantaneous firing rate is modulated by a sinusoidal input at frequency \( \omega \). It is a complex function which describes the gain and the phase of the modulation at frequency \( \omega \). In the following, we call \( r_{a,\omega} \) the static transfer function of the network, while \( r_{a,\omega} \) is the dynamical transfer function of the network. The motivation for computing the response to sinusoidal inputs is twofold: (1) one can then reconstruct the response of the network to arbitrary-time dependent inputs; (2) oscillations are observed in many regions of the central nervous system, and are therefore a common type of input for post-synaptic structures.

### 3 ANALYTICAL METHODS

The analysis of the network response is performed in three steps. (1) we compute the background activity as a function of the mean drive (static transfer function); (2) we check whether background activity with a constant firing rate is stable; (3) we compute the dynamical response of the network. The outcome of the analysis is therefore both the region of stability of states with time-independent firing rate (i.e., asynchronous states in networks of spiking neurons) in the space of parameters characterizing the network, and the dynamical transfer function as a function of such parameters. We now briefly outline how these three steps are performed in practice.

#### 3.1 STATIC TRANSFER FUNCTION

##### 3.1.1 Rate models

In the rate model, with constant external inputs (\( \mu_{a,\text{int}} = 0 \) for \( a = E, I \)), the stationary firing rates are given by

\[
r_{a,0} = [I_{a,0}]_1 \tag{14}
\]

\[
I_{a,0} = \mu_{a,\text{ext}} + I_{a,\text{int}} - J_{a,E} f_0 - J_{a,I} f_0 \tag{15}
\]

where \( I_{a,0} \) is the synaptic input received by population \( a \). In this paper, we only consider the situation where both input currents are suprathreshold \( (I_{a,0} > 0 \) for both \( a = E, I \)).

When \( I_{a,0} > 0 \) for both \( a = E, I \), we have

\[
r_{E,0} = \frac{\mu_{E,0}(1 + J_{E,E}) - \mu_{E,0} J_{E,I}}{(1 - J_{E,E})(1 + J_{E,E}) + J_{E,I}} \tag{16}
\]

\[
r_{I,0} = \frac{\mu_{I,0}(1 + J_{I,E}) - \mu_{I,0} J_{I,I}}{(1 - J_{I,E})(1 + J_{I,I}) + J_{I,E}} \tag{17}
\]
It is sometimes useful to rewrite both equations as follows
\[
\begin{align*}
\lambda^2 \tau_L \tau_x + \lambda \left[ \tau_x (1 + J_E) + \tau_t (1 - J_E) \right] \\
+ (1 + J_E)(1 - J_E) + J_{EE} I = 0
\end{align*}
\]  

As soon as delays are included, the eigenvalues no longer have simple analytical expressions. In order to compute the eigenvalues, we use the following strategy. The first step is, for a given perturbation in the rate \( \delta r_I(t)e^{\omega t} \), to compute the resulting perturbation in the average synaptic variables
\[
\delta s_{ab}(t) = S_{ab}(\lambda) \delta r_I(t)e^{\omega t}
\]
where, for synaptic currents defined by Eqs (4,5),
\[
S_{ab}(\lambda) = \frac{e^{-\lambda t_0}}{(1 + \lambda \tau_{ab})(1 + \lambda \tau_{ab})}
\]

For \( \lambda = i\omega \), \( S \) behaves as a low-pass filter with an additional delay (see Figures 2C,D).

From the perturbation in the individual synaptic variables, we can easily compute the perturbation in total average currents to neurons in a population
\[
\delta I_a(t) = J_{aa} \delta s_{aa}(t) - J_{ab} \delta s_{ab}(t),
\]
The second step is, from a perturbation in total average currents, to compute the perturbation in the instantaneous firing rate,
\[
\delta r_I(t)e^{\omega t} = R_\omega(\lambda) \delta I_a(t)
\]
where, for the rate model,
\[
R_\omega^{RM}(\lambda) = \frac{1}{1 + \lambda \tau_a}
\]
while for the network of integrate-and-fire neurons (Brunel and Hakim, 1999; Brunel et al., 2001)
\[
R^{\omega}_{\omega}^{IF}(\lambda) = \frac{\tau_{ma} e^\tau}{ \sigma (1 + \lambda \tau_{ma})} \left( \frac{\partial U}{\partial y}(y, \lambda \tau_{ma}) - \frac{\partial U}{\partial y}(y, \lambda \tau_{ma}) \right)
\]
where \( y = V - (V_{rest} + I_{I/I})/\sigma, \)
\[
U(y, \lambda) = \frac{e^{y^2}}{\Gamma(1 + \lambda/2)} \left[ \frac{1 - \lambda}{2} \right] [M \left( 1 + \frac{3}{2}, -y^2 \right) + 2ye^y] M \left( 1 - \frac{3}{2}, -y^2 \right)
\]
where \( M \) is a confluent hypergeometric function (Abramowitz and Stegun, 1970). The function \( R^{IF} \) is shown in Figure 2 for \( \lambda = i\omega \). This figure shows some of the main qualitative features of \( R^{IF} \): (i) when noise is large, it behaves as a low-pass filter, with a decay at high frequency in \( 1/\sqrt{\omega} \), while resonances are present when noise is small; (ii) the gain depends markedly on both noise level, and mean firing rate.

The last step is to insert Eqs (27,29) in Eq. (30), and to divide both sides by \( e^{\omega t} \). Eliminating \( \delta r_I \), we find that the eigenvalues \( \lambda \) should satisfy
Figure 2 | Neuronal ($R^\alpha$) and synaptic (S) transfer functions. (A) Modulus (amplitude) of $R^\alpha$ as a function of frequency, for several values of the mean firing rate $r_0$ and the amplitude of the noise $\sigma$. $|R^\alpha|$ is proportional to the slope of the static transfer function at low frequencies, and decays as $r_0^{1/2}$ at high frequency. It shows resonances at integer multiples of $r_0$ at high firing rates and low noise. (B) Argument (phase shift) of the $R^\alpha$ as a function of frequency, for several values of the mean firing rate $r_0$ and the amplitude of the noise $\sigma$. (C) Modulus of $S$, for $\tau_r = 2$ ms, $\tau_i = 0$ ms (full line) and $\tau_r = 2$ ms, $\tau_i = 1$ ms (dashed line). Note that the modulus is independent of the delay. It decreases at high frequency as $1/(\pi f)$ when the rise time is zero, and $1/(\pi f \tau_r)$ when both rise and decay times are non-zero. (D) Argument of $S$, for three parameter sets: $D = 0$ ms, $\tau_r = 2$ ms, $\tau_i = 0$ ms (full line); $D = 0$ ms, $\tau_r = 2$ ms, $\tau_i = 1$ ms (dashed line); and $D = 1$ ms, $\tau_r = 2$ ms, $\tau_i = 1$ ms (dotted line).

\begin{equation}
1 = A_{\alpha}(\lambda)(1 + A_{\alpha}(\lambda)) - A_{\alpha}(\lambda)A_{\alpha}(\lambda) \quad (34)
\end{equation}

where

\begin{equation}
A_{\alpha}(\lambda) = J_{\alpha}R_0(\lambda)S_{\alpha}(\lambda) \quad (35)
\end{equation}

describes the net effect of the firing rate of population $b$ on population $a$. Note that the terms in the r.h.s. of Eq. (34) potentially lead to three distinct instabilities, mediated by $E \rightarrow E, I \rightarrow I$, or the $E \rightarrow I \rightarrow E$ feedback loop, respectively.

Note that in the simplest rate model, $S_{\alpha}(\lambda) = 1$, $R_0(\lambda)$ is given by Eq. (31), and Eq. (34) reduces to Eq. (26).

Equation (34) is solved numerically, using a globally convergent Newton routine which tracks the solutions $\lambda = \alpha + i\beta$ to this equation, starting from many different initial conditions. Models are considered stable only if the routine converges toward negative $\alpha$ for any initial condition.

3.3 Dynamical Transfer Function

Once linear stability of the background state has been checked, we can calculate the response of the system when the input has an additional oscillatory component. The oscillatory response of population firing rates is given by

\begin{equation}
r_{\alpha}^{(\omega)}(\omega) = \frac{(1 + A_{\alpha})R_0(\mu_{z,1}) - A_{\alpha}R_0(\mu_{z,1})}{(1 - A_{\alpha})(1 + A_{\alpha}) + A_{\alpha}A_{\alpha}} \quad (36)
\end{equation}

\begin{equation}
r_{\alpha}^{(\omega)}(\omega) = \frac{A_{\alpha}R_0(\mu_{z,1}) + (1 - A_{\alpha})R_0(\mu_{z,1})}{(1 - A_{\alpha})(1 + A_{\alpha}) + A_{\alpha}A_{\alpha}} \quad (37)
\end{equation}

where in the above equations $A_{\alpha}$ is given by Eq. (35) in which $\lambda = i\omega$, and $R_0$ is given by Eq. (31) for the rate models, Eq. (32) for the network of integrate-and-fire neurons, in which $\lambda = i\omega$.

4 RESULTS

4.1 One-population Model

It is useful to start by summarizing the dynamics of a network of a single population of neurons, before moving to the two-population network. In a rate model, the stationary firing rate $r_0$ is given by

\begin{equation}
r_0 = \frac{\mu_0}{1 - J}
\end{equation}

where $\mu_0$ is the external input, and $J$ is the recurrent coupling (excitatory if $J > 0$, inhibitory if $J < 0$). In a network of LIF neurons, the stationary firing rate $r_0$ is given by

\begin{equation}
r_0 = \Phi(\mu + Jr).
\end{equation}

Linear stability analysis of the system shows that the eigenvalues $\lambda$ satisfy

\begin{equation}
1 = A(\lambda) = JR(\lambda)S(\lambda) \quad (38)
\end{equation}

where $R(\lambda)$ is given by Eq. (31/32) for rate model/network of LIF neurons, respectively, and $S(\lambda) = 1$ for the simplest rate model, and is otherwise given by Eq. (28).

When the asynchronous state is stable, the dynamical transfer function is

\begin{equation}
r_0(\omega) = \frac{R(\omega)}{1 - A(\omega)} \quad (39)
\end{equation}
In the absence of coupling, the network firing rate behaves as a low-pass filter (see the gray curves in Figure 3), except in the LIF network in the small noise regime where resonances appear at the single neuron firing frequency (see, e.g., Knight, 1972; Brunel et al., 2001; Fourcaud and Brunel, 2002; Fourcaud-Tromé et al., 2003 and Figure 2 – regime not shown in Figure 3).

In an excitatory network \(J > 0\), the asynchronous state is stable provided \(J < J^*\) where \(J^* = 1/\Phi'(\Phi^{-1}(r_c))\) (LIF network). When \(J = J^*\), one of the eigenvalues becomes equal to zero, signaling the onset of a saddle-node bifurcation. This is the much-studied rate instability: because of the excitatory feedback, the network cannot maintain its rate at a value \(r_c\). Rather, the rate either decreases to a smaller value (typically close to zero), or increases to a larger value (typically close to maximal firing rate). When \(J < J^*\) the effect of excitatory coupling is to amplify the response at low frequencies (the much-studied amplification effect, see, e.g., Seung, 2003), while at the same time slowing down the dynamics (through a reduction of the cut-off frequency, see, e.g., Seung, 2003). This effect is present in all models, independently of synaptic characteristics.

In the simplest inhibitory network (no synaptic time constants), the asynchronous state is always stable. Introducing a delay in synaptic transmission leads to an oscillatory instability (through a Hopf bifurcation) at a frequency \(\omega\). In the rate model in which no rise and decay times are present, the frequency \(\omega\) satisfies the equation \(\tan(\omega t) = -\tau_0\omega\) whose solution is \(\omega = \pi/2\tau_0\). This instability leads to an oscillation due to delayed inhibitory feedback when \(|J| > |J^*|\), which has been characterized in many different inhibitory network models (see Brunel and Hakim, 2008 for a review). When \(|J| < |J^*|\), the effect of inhibitory coupling is to suppress the response at low frequencies, which leads to an increase in the cut-off frequency (see, e.g., Seung, 2003). In the absence of delays, the response of the network is basically the response of a low-pass filter. In the presence of a delay, the response shows a resonance at a frequency close to the frequency of the oscillation at the Hopf bifurcation (see Figures 3B,D, blue curves).

### 4.2 TWO-POPULATION RATE MODELS

#### 4.2.1 Stability of the asynchronous state in the simplest rate model

In the simplest rate model, the eigenvalues of the linear stability matrix are given by Eq. (26). From this equation it is easy to compute the conditions the coupling parameters must satisfy on the two possible types of instability lines.

A saddle-node bifurcation occurs when \(\lambda = 0\); this happens whenever

\[
(1 + J_1)(1 - J_{EE}) + J_{IJ}J_{II} = 0
\]

or equivalently \(1 + Y = 0\). This instability corresponds to the rate instability induced by excitatory feedback onto the excitatory population, which is already present in the one-population model. The presence of feedback inhibition (as measured by the parameter \(Y\)) stabilizes the network with respect to this instability.

A Hopf bifurcation occurs when \(\lambda = i\omega\); this happens whenever

\[
\tau_0 (1 + J_1) + \tau_0 (1 - J_{EE}) = 0 \text{ and } (1 + J_1)(1 - J_{EE}) + J_{IJ}J_{II} > 0
\]

This bifurcation is a new feature of the two-population model; it depends both on E–E feedback and the E–I feedback loop. The two bifurcation lines are shown in Figure 4.
where $\tau_a$ is the effective time constant of the population $a$, 

$$r_{EI} = \frac{\mu_{EI}}{1 - J_{EE}} \left( \frac{1 - X + i\omega \tau_\xi}{(1 + i\omega \tau_\xi)} + Y \right)$$  \hspace{1cm} (40)$$

$$r_{IE} = \frac{\mu_{IE}}{1 - J_{EE}} \left( \frac{Y + (1 + i\omega \tau_\xi) X}{(1 + i\omega \tau_\xi)(1 + i\omega \tau_\xi) + Y} \right)$$  \hspace{1cm} (41)$$

In writing Eqs (40,41), we have assumed that $a_1 \equiv m_{IE}/m_{EI}$ is equal to $a_0 \equiv m_{IE}/m_{EI}$. This would be the case if both E and I networks receive inputs from the same structures. If E and I networks receive different types of inputs, then we could in principle have $a_1 \neq a_0$. In this case, we would need to define two effective feed-forward parameters $X_0$ (strength of "static" feed-forward inhibition) and $X_1$ (strength of "dynamic" feed-forward inhibition), proportional to $a_0$ and $a_1$, respectively.

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4.2.2 Dynamical transfer function in the simplest rate model

For a two-population rate model, the dynamical transfer function is given by Eqs (36,37), which can be rewritten for the simplest rate model as

$$r_{EI} = \frac{\mu_{EI}}{1 - J_{EE}^E} \left( \frac{1 - X + i\omega \tau_\xi}{(1 + i\omega \tau_\xi)} + Y \right)$$  \hspace{1cm} (40)$$

$$r_{IE} = \frac{\mu_{IE}}{1 - J_{EE}^I} \left( \frac{Y + (1 + i\omega \tau_\xi) X}{(1 + i\omega \tau_\xi)(1 + i\omega \tau_\xi) + Y} \right)$$  \hspace{1cm} (41)$$

where $\tau_\xi$ is the effective time constant of the population $a$, $\tau_\xi = \tau_\xi^E/(1 - J_{EE}^E) = \tau_\xi^I/(1 + J_{EE}^I)$.

In writing Eqs (40,41), we have assumed that $\alpha_1 = \mu_{IE}/\mu_{EI}$ is equal to $\alpha_0 = \mu_{IE}/\mu_{EI}$. This would be the case if both E and I networks receive inputs from the same structures. If E and I networks receive different types of inputs, then we could in principle have $\alpha_1 \neq \alpha_0$. In this case, we would need to define two effective feed-forward parameters $X_0$ (strength of "static" feed-forward inhibition) and $X_1$ (strength of "dynamic" feed-forward inhibition), proportional to $\alpha_0$ and $\alpha_1$, respectively.
We focus here on two quantities: the amplitude of the excitatory transfer function \( |r_e(\omega)| \) and the phase shift between the two-populations, \( \Phi_{j} \). In this model, it is easy to show (see Appendix) that the amplitude of the transfer function can have only two qualitatively different behaviors: (i) a monotonically decaying function of the frequency \( \omega \) (“low-pass” behavior – see, e.g., Figure 4B, panel with the cross); or (ii) a function that first increases with frequency, peaks at a finite frequency, and then decays toward zero at high frequencies (see Figure 4B, panel with the star). Note that in the rate model the amplitude of the response does not depend on \( r_{\omega} \). Therefore the examples in the Figures 4, 5, and 6 are shown with amplitudes normalized by \( r_{\omega}(f = 0) \) and not \( r_{\omega} \). The regions where both types of behaviors occur are shown in the plane of couplings involving the inhibitory network (self-inhibitory coupling \( I_{j} \) vs strength of EI loop \( I_{e}f_{\omega} \)), for different values of \( I_{e}f_{\omega} \). We see that strong values of \( I_{j} \) tend to favor low-pass behavior, while strong values of \( I_{e}f_{\omega} \) tend to favor resonant behavior. Close to the saddle-node bifurcation, the transfer function is low-pass, with a zero-frequency response that diverges on the bifurcation line. Close to the Hopf bifurcation, the transfer function is resonant, with a peak that diverges on the bifurcation line, at the frequency of the oscillation that appears on the line.

The boundary separating the two regions depends on the strength of feed-forward inhibition \( X \); the larger \( X \), the more pronounced the resonant behavior. This is because feed-forward inhibition tends to suppress inputs at low frequencies. In fact, feed-forward inhibition can create a resonance even in the absence of feedback E–I coupling (see Figure 4A, case where \( I_{e}f_{\omega} = 0 \)). In the limit \( X \to 1 \), the resonant behavior becomes present in the whole stable region. Note also that the presence of a resonance is not equivalent to the presence of complex eigenvalues of the linear stability analysis, as happens in a two-variable single neuron model (Richardson et al., 2003): the network can be resonant in the absence of complex eigenvalues, and conversely, the network can be low-pass in the presence of such complex eigenvalues.

Another quantity of interest is the phase shift between excitatory and inhibitory populations. In this model, one can show (see Appendix for details) that again only two qualitatively different behaviors can occur: (i) the excitatory population leads the inhibitory population at all frequencies \( f \) (see, e.g., Figure 4D, panel with the cross); or (ii) the opposite occurs and the inhibitory population leads the excitatory population at all \( f \) (see, e.g., Figure 4D, panel with the star). The regions where both types of behavior occur are shown in the plane of couplings involving the inhibitory network (self-inhibitory coupling \( I_{j} \) vs strength of EI loop \( I_{e}f_{\omega} \)), for different values of \( I_{e}f_{\omega} \) in Figure 4C. Close to the rate instability, inhibition tends to lead excitation, while the opposite occurs close to the Hopf bifurcation. Note that for all parameters, the phase shift between the two-populations tends to zero in both \( f \to 0 \) and \( f \to \infty \) limits, and that it reaches a maximum at some finite value. In the resonant region, this value is close to the location of the resonant peak (compare Figures 4B,D, panels with the star).

### 43 Rate Model with Non-Zero Synaptic Time Constants

How are the dynamics modified when the synaptic time scales are taken into account in the rate model? We show here that new behaviors emerge, due to the presence of delays in synaptic interactions.

#### 4.3.1 Stability of the asynchronous state

The stability of the asynchronous state can be evaluated by computing the eigenvalues, i.e., the solutions of Eqs (34, 35, 28, 31). We first discuss the stability analysis when synaptic interactions are simply described by a delay (i.e., rise and decay times are equal to zero), and both E and I populations have the same rate time constant. In fact, all the qualitative features of the general case (i.e., with strictly positive rise times and decay times, and/or with \( \tau_{e} \neq \tau_{i} \)) are already present in this simplified model.

The main qualitative changes that occur when delays are introduced are as follows: (1) there is a new oscillatory instability induced by I–I connectivity, which is reached when \( I_{j}f_{\omega} \) becomes strong enough (Brunel and Hakim, 1999; Roxin et al., 2005); (2) the oscillatory instability induced by the E–I feedback loop is now present at all values of \( I_{e}f_{\omega} \). As a result of the new oscillatory instability, the space of parameters for which the asynchronous state is stable is now bounded for any value of \( I_{j}f_{\omega} \) (see Figure 5). The number of distinct bifurcation lines that form the boundary of the region of stability can vary between one and three, depending on parameters. First, the rate instability line is only present when \( I_{j}f_{\omega} > 1 \), as in the simplest rate model without synaptic time constant. Second, there can be either one or two distinct oscillatory instability lines, depending on the relative values of the excitatory and inhibitory delays, \( D_{e} \) and \( D_{i} \).

When \( D_{e} > D_{i} \) there exist two distinct instability lines, characterizing the I–I and E–I scenarios; whereas the two lines cross at some point in the \( I_{j}f_{\omega} \) plane (see Figures 5A,B). The frequencies of the corresponding instabilities remain well separated. The frequency of the I–I instability is generically higher than the frequency of the E–I instability, since the I–I induced-oscillations involve only two synapses from peak to peak (increase in I rate leads to a decrease in I rate due to feedback, which in turn leads to the next increase in I rate), while E–I induced-oscillations involve four synapses (E rate increase, leading to an increase in I rate, leading to a decrease in E rate, then decrease in I rate, followed by the next increase in E rate). For example, for the parameters of Figure 5A, I–I instability leads to frequencies in the 150 to 200-Hz range, while E–I instability leads to frequencies which are in the 10 to 80-Hz frequency range. When \( D_{e} = D_{i} \) the two instability lines meet at a cusp; at this point the frequencies of the two instabilities become equal (Figure 5C).

Finally, when \( D_{e} < D_{i} \) there is a single oscillatory instability line, smoothly connecting the “pure” I–I and “pure” E–I scenarios in the \( I_{j}f_{\omega} \) plane (Figure 5F). In this case the frequency decreases smoothly along the line as the \( \frac{I_{j}f_{\omega}}{D_{e}} \) ratio is decreased. As a result, the region of stability of the asynchronous state exhibits either a triangular shape, when \( D_{e} \geq D_{i} \) (see Figures 5B,F) or a D shape, when \( D_{e} < D_{i} \) (see Figure 5F).

#### 4.3.2 Dynamical transfer function

The next step is to compute the dynamical transfer function, Eqs (36, 37). We show in Figure 5 how the qualitative shape of this transfer function depends on synaptic parameters and on the connectivity strengths. In the vicinity of the rate instability, the transfer function has either low-pass characteristics (black regions in Figure 5, see a representative transfer function in Figure 5G), or shows a trough followed by a resonant peak, when the system is
Figure 5 | Qualitative behaviors of the rate model with delays, for $r_f = r_i = 10$ ms, $D_{j_1} = 2$ ms, $D_{i} = 1$ ms; (E,F) $D_f = 1$ ms, $D_i = 2$ ms. (A,B) Frequency of the oscillatory instability on the bifurcation lines, as a function of $J_{E_1} - J_{I_1}$; (B,D,F) behaviors of the gain of the E network in the $J_{E_1} - J_{I_1}$ plane for three different values of $X$ (indicated at the top of the corresponding panels). White areas show regions in which the asynchronous state is unstable due to the rate or Hopf instabilities, shown as black lines. The stable region is subdivided in regions of different colors, indicating qualitatively different behaviors of the E transfer function. Colors indicate the number of extrema at non-zero frequencies of the amplitude vs frequency curve (0: black, 1: green, 2: red, 3: orange, 4: yellow). (G) Representative transfer functions (top: amplitude; bottom: phase) for each of the regions. From left to right, $J_{E_1} = 5, 18, 23, 80, 19$; $J_{I_1} = 5, 4, 15.5, 14, 7$. Symbols on the top left of the amplitude plots refer to the location of the network parameters in (B,F). Black dots on top of the red curves indicate the location of the extrema.
close to the I–I instability (red region in Figure 5, see a representative transfer function in Figure 5G). In the vicinity of the Hopf bifurcation lines, the transfer function always exhibits resonant peaks, but in qualitatively different ways. There can be a single resonant peak — the typical situation when the stability region is D-shaped, i.e., there is a single bifurcation line (see green, red, and yellow regions in Figure 5, and the corresponding transfer functions in Figure 5G). Note that in the yellow region, there is a second peak in the transfer function at a frequency which is larger than the one of the resonant peak, but its magnitude is always small compared to the amplitude of the other peak. Alternatively, there can be two separate resonant peaks, typically when the stability region is triangular, i.e., with two separate bifurcation lines, with two separate frequencies (see orange regions in Figure 5B, and the transfer function in Figure 5G), and the network is close to the intersection between the two lines. The frequency of the resonant peaks is close to the frequency of the corresponding instabilities. For parameters of Figure 5, the resonant frequency is typically in the gamma range when the system is close to the E–I bifurcation line, while it is typically very high (more than 100 Hz) when the system is close to the I–I line. As in the simplest rate model, the boundary between the different regions also depends on the strength of feed-forward inhibition: in particular, the pure low-pass region tends to disappear as $X$ increases, since a high value of $X$ decreases the response at low frequencies.

The transfer functions can also be classified according to the behavior of the phase shift between excitatory and inhibitory populations. The main new behavior introduced by the presence of synaptic delays is that it is now possible for the phase shift to change sign when the frequency is increased. This leads to a larger number of qualitatively different behaviors for the phase shift. Some of the typical behaviors of the phase shift are represented in Figure 5G. Note in particular that resonances are associated with sharp changes in the phase shift. Close to the E–I instability line, the E population leads the I population at the corresponding resonant frequency.

The behavior of a model with $\tau_y \neq \tau_x$ is qualitatively very similar to the one of the model with $\tau_y = \tau_x$. The main difference is that the transition between the 1 Hopf and 2 Hopf bifurcation behaviors no longer occurs at $D_\theta = D_\delta$, but for slightly different values of the delays. For example, for $\tau_y = 20 \text{ ms}$, $\tau_x = 10 \text{ ms}$, and $D_\theta = 1.5 \text{ ms}$, the transition between both behaviors occurs at $D_\delta \sim 1.55 \text{ ms}$.

### 4.3.3 Behavior of the model with finite rise and decay times

A similar analysis can be performed for the rate model with finite delay, rise, and decay times. The qualitative behaviors of the transfer function in the region of stability of the asynchronous state are shown in Figure 6. This figure shows that the same types of behaviors are observed as in Figure 5. At the quantitative level, the frequencies of the instabilities are reduced by the presence of finite rise/decay times (compare Figures 6A, C and 6B, D with 6C, E).

**FIGURE 6** | Qualitative behaviors of the rate model with non-zero delays, rise, and decay times ($\tau_{\text{rise}} = \tau_{\text{delay}} = 2 \text{ ms}, \tau_{\text{rise}} = \tau_{\text{delay}} = 5 \text{ ms}, \text{ and } \tau_{\text{rise}} = \tau_{\text{delay}} = 5 \text{ ms}$). Other parameters: $\tau_y = 20 \text{ ms}, \tau_x = 10 \text{ ms}, J_{\text{EE}} = 1.5$. (A) Frequency of the oscillatory instabilities as a function of $J_{\text{EE}}/J_{\text{EI}}$, for $D_{\delta} = 2 \text{ ms}, D_{\theta} = 1 \text{ ms}$. (B) Stability region of the asynchronous state in the $J_{\text{EI}/J_{\text{IE}}}$ plane, and qualitative behaviors of the E transfer function for $D_{\delta} = 2 \text{ ms}, D_{\theta} = 1 \text{ ms}$, and three values of $X$(same color codes as in Figure 5). (C) Frequency of the oscillatory instability for $D_{\theta} = 1 \text{ ms}, D_{\delta} = 2 \text{ ms}$. (D) Stability region of the asynchronous state, and qualitative behaviors of the E transfer function for $D_{\theta} = 1 \text{ ms}, D_{\delta} = 2 \text{ ms}$, and three values of $X$(same color codes as in Figure 5).
Figures 5A,E). This decrease in frequency is stronger for the 1–I instability. Consequently, the two possible resonant frequencies are now much closer than in the case in which only delays are present, which decreases the size of the region in which both resonances are observed (compare orange regions in Figures 5 and 6). Another quantitative difference is that phase differences between E and I populations tend to be smaller.

4.4 NETWORK OF LIF NEURONS

The same analysis can be performed for a network of LIF neurons, as outlined in Section 3.2. Interestingly, the stability analysis of the asynchronous state revealed the same types of instabilities as in the rate model with synaptic time constants, i.e., a rate instability (mostly governed by E–E interactions), and two types of oscillatory instabilities (governed by E–I or I–I interactions). As in the rate model, the two types of oscillatory instabilities merge smoothly in the space of synaptic connection strengths if $D_e < D_p$, while they remain disjoint if $D_e = D_p$. There are however a few qualitative differences between the two models:

- The high frequency behavior of the LIF model is different from the rate model: the amplitude of the response decays as $1/\sqrt{\lambda}$, and the phase shifts go to $\pi/4$ in the LIF network, as compared to a $1/f$ decay with a phase shift of $\pi/2$ in the rate model (see Section 5)
- In the LIF network, the single neuron transfer function depends both on its average inputs (and therefore its average firing rate) and on the level of noise. As a consequence, the network transfer function changes as a function of both mean and variance of the external inputs.

In this section we first show how the qualitative behaviors of the LIF network compare with those of the rate model (at fixed mean firing rate and level of noise), and then show how external inputs qualitatively change the network transfer function in the LIF network. This property is not present in a rate model with threshold-linear static transfer function, but can be reproduced with a suitable non-linearity in the static transfer function.

4.4.1 Comparison between LIF network and rate model

Figure 7 shows the behaviors of the transfer function of the LIF network in strong noise regime ($\sigma = 5$ mV). To make an explicit comparison with the rate model, we choose to fix the firing rates of E and I populations to specific values (10 Hz and 30 Hz, respectively), independently of the matrix of coupling. This means that on any point of the phase diagrams showed in Figure 7 the external inputs must be computed such that the network has the required mean firing rates. We also need to generalize the definition of the feed-forward and feedback inhibition parameters to the LIF network. A natural generalization is

$$X = \frac{\Phi_e'(I_{io})}{\Phi_e'(I_{io})} \frac{I_E I_e \tau_m \Phi_e'(I_{io})}{\mu_{e1} (1 + \tau_m \Phi_e'(I_{io}))} \tag{42}$$

$$Y = \frac{I_E I_e \tau_m \Phi_e'(I_{io})}{(1 - I_E I_e \tau_m \Phi_e'(I_{io}))(1 + \tau_m \Phi_e'(I_{io}))} \tag{43}$$

In Eqs (42,43), we see that these parameters are now governed by “effective coupling strengths,” where the coupling strength $J_m$ is multiplied by the gain of the transfer function of the pre-synaptic population $\Phi_e'$ at the corresponding firing rate.

To compare the LIF network with the rate model, we then modulated independently the sinusoidal inputs to the E and I populations, but with a ratio between amplitudes taken in order to keep the feed-forward inhibition parameter $X$ constant in each plot. Synaptic parameters was taken to be identical to the rate model. Figure 7 shows that, qualitatively, the LIF network behaves essentially as the rate model. There are of course quantitative differences due to the fact the transfer function of the LIF neuron is different from that of the rate model. Importantly, the high frequency behavior is different, with a $1/\sqrt{\lambda}$ decay of the gain, and a phase shift of $45^\circ$ at high frequencies in the LIF model, as already mentioned above.

4.4.2 Dependence of the network transfer function on external inputs

In the LIF model, the single-cell transfer function $R^{ef}$ depends on the inputs to the network. In particular, changing the overall firing rate alters the static gain of the transfer function $\Phi$, which changes the effective coupling strengths and in turn the effective feed-forward and feedback inhibition parameters.

Figure 8 shows an example of a network where an increased input to both populations led to qualitative changes of the network transfer function. The coupling strengths were chosen such that when the external inputs to both populations increase, the firing rates of both populations first increase. Then, inhibition becomes strong enough to reduce excitatory rates even with the increased external inputs. Interestingly, this leads to a feed-forward parameter $X$ larger than one. The network transfer function goes through a number of changes as the external inputs are varied: at low input strength/low rates, the transfer function is low-pass, due to the fact that the gain of single neuron F–I curves is small, and hence effective coupling strengths are small. When the inputs increase, the slopes of the transfer functions increase, and therefore the effective couplings increase. This leads to a resonant transfer function, due to feedback between E and I populations. Finally, when the inputs increase further, the E rate reaches a peak and then decreases with increased inputs. This leads to a major change of the response to external inputs: the E and I populations are now in antiphase for slowly varying inputs (see Figure 8D). This is due to the fact that $X > 1$. When $X = 1$, then the phase shift between the two-populations is $90^\circ$, as shown by Figure 8D. In both cases, the phase shift between the two-populations decreases with frequency, and goes to zero in the high frequency limit.

A qualitative change in network transfer function with increasing external input is a feature that is not present in a rate model with threshold-linear static transfer function, but it can be obtained using a suitable non-linear transfer function. In particular, Figure 8D shows when one uses a rate model with the static transfer function of the LIF neuron, Eq. (24), one obtains network transfer functions which are very similar to the LIF network.

4.4.3 Dependence of the network transfer function on external noise

The single-cell transfer function also depends on the magnitude of external noise, as shown by Figure 2. In particular, as was characterized in a number of previous studies (Knight, 1972; Brunel et al.,
Figure 7 | Qualitative behaviors of the network of LIF neurons.
Synaptic time constants are the same as in Figure 6. Color codes as in Figure 5. In all cases, \( r_E = 10\) Hz, \( r_I = 30\) Hz and \( \mu_E = 1\) mV, \( \tau_{E} = 20\) ms, \( \tau_{I} = 10\) ms, \( J_{EIE} = 25\) mV. (A) Frequency of the oscillatory instabilities for \( D_E = 2\) ms, \( D_I = 1\) ms. (B) Stability region of the asynchronous state, and qualitative behaviors of the E transfer function for \( D_E = 2\) ms, \( D_I = 1\) ms, and three values of \( X \). (C) Frequency of the oscillatory instability for \( D_I = 1\) ms, \( D_E = 2\) ms. (D) Stability region of the asynchronous state, and qualitative behaviors of the E transfer function for \( D_I = 1\) ms, \( D_E = 2\) ms, and three values of \( X \). (E) Representative transfer functions of each behavior. Lines: analytical expressions, Eqs (36,37). Squares: numerical simulations of a network of 2000 excitatory neurons and 500 inhibitory neurons. From left to right: \( J_{EI} = 600, 1300, 2000, 8000, 1100\) mV and \( J_{II} = 20, 30, 100, 150, 40\) mV.

Inhibitory and excitatory connection strengths (i.e., we choose \( I_{EI} = I_{II} = g_{EI} = g_{II} \)), and the strength of external inputs. Parameters are chosen such that the network can potentially have a double resonance, due to E–I feedback coupling and I–I coupling, respectively (the orange region on the plots). The top graphs show the “phase diagram” in the high noise regime (\( \sigma = 5\) mV) that has been explored so far. We see that the asynchronous state is stable in a finite range of external inputs. As the external inputs increase, the network transfer function develops a single or several resonances,
Figure 8 | External inputs can change qualitatively the network transfer function. Parameters of the network are $\mu_E = 1\, \text{mV}$, $\tau_w = 20\, \text{ms}$, $\tau_m = 10\, \text{ms}$, synaptic time constants as in Figure 6B. $J_{EE} = 25\, \text{mV}$, $J_{EI} = 6400\, \text{mV}$, $J_{II} = 70\, \text{mV}$. (A) Firing rate of E (red) and I (blue) populations as a function of external input. (B) Feed-forward inhibition parameter $X$ as a function of $\mu_E$. (C) Feedback inhibition parameter $Y$ as a function of $\mu_E$. (D) Network transfer functions for three different values of $\mu_E$, indicated by the corresponding symbols in (A–C). From left to right panel, $\mu_E = 10, 17.4, 30\, \text{mV}$, corresponding to $X = 0.2, 1, 1.2$, $r_E = 0.9, 2.8, 1.7\, \text{Hz}$ and $r_I = 0.3, 4.3, 12.3\, \text{Hz}$. Solid lines: analytical expressions for the LIF network, Eqs (36,37). Squares: numerical simulations of the LIF network. Dashed lines: analytical expressions for a rate model with a LIF static transfer function.

Depending on the value of $g$, above a critical value of the external inputs the asynchronous state destabilizes due to a Hopf bifurcation, as in Brunel (2000).

The other two graphs show how the picture is modified when the level of external noise decreases. There are two main effects: first, the region of stability of the asynchronous state becomes smaller, as expected from previous studies in purely inhibitory networks (Brunel and Hakim, 1999; Brunel and Hansel, 2006); second, qualitatively new behaviors of the network transfer function emerge. In particular, the network can develop a new resonance which is associated with the single-cell resonance (see Figures 9D,F).

5 DISCUSSION

In this paper, we have investigated how the transfer function of an E–I network is controlled by its connectivity and the temporal parameters of its synapses. We have shown that a two-population rate model, as well as a fully connected network of excitatory
and inhibitory integrate-and-fire neurons, exhibit many qualitatively different behaviors, from low-pass to multiple resonances. Resonance phenomena appear generically in both models provided inhibition is sufficiently strong. With realistic synaptic currents, oscillatory instabilities can arise due both to the E–I feedback loop and to the I–I connectivity. Close to these instabilities, a single or double resonance can appear, depending on synaptic parameters. We have also shown that for a given level of external input, and in a high noise regime, the behaviors of the network transfer function of LIF networks is qualitatively similar to the rate models with identical synaptic dynamics. However, an important difference is that in the network of LIF neurons the transfer function depends on the level of external inputs, and external noise. In a rate model with a threshold-linear transfer function, the “single-cell” dynamical transfer function is independent of the input. If a sigmoidal static transfer function had been used instead of the threshold-linear

**FIGURE 9** Network transfer functions of LIF model depend on external noise. (A) Qualitative behaviors of the E transfer function in the $g - \mu_E/V$ plane for $\sigma = 5$ mV. Same color codes as in the Figure 7, with a new region (with five extrema – three maxima and two minima, in brown). Dotted lines are iso-excitatory firing rate lines. Dashed line indicates where the input current is equal to the firing threshold. (B) Representative transfer function in the yellow region [see + sign in (A)]. Full lines: analytical expressions, Eqs (36,37). Squares: numerical simulations of a network of 8000 excitatory neurons and 2000 inhibitory neurons. (C) Same as A, but with $\sigma = 3$ mV. (D) Representative transfer function in the brown region in (C). (E) Same as A, but with $\sigma = 1$ mV. (F) Representative transfer function in the brown region in E. In all cases, $\alpha = 1$, $\mu_E = 1$ mV (except case F where $\mu_E = 0.5$ mV), $J_E = 75$ mV. Same time constants as in Figure 6. For (B,D,F) respectively: $\sigma = 5, 3, 1$ mV, $r_E = 11.4, 8.8, 5.9$ Hz and $r_I = 22.8, 17.7, 11.8$ Hz.
one, then the gain at zero-frequency (i.e., the slope of the f–I curve) would depend on the input, which could also lead to qualitative changes of the network transfer function as a function of the input. However, the LIF neuron has the additional feature that the single-cell dynamical transfer function can exhibit qualitative changes as a function of its inputs, such as the appearance of a resonance at single-cell firing frequency when the noise is sufficiently weak.

There is an extensive literature describing the bifurcations of fixed points in firing rate/neural field/neural mass models (see, e.g., Wilson and Cowan, 1972; Borisyuk and Kirillov, 1992; Jansen and Rit, 1995; Roxin et al., 2005; Deco et al., 2008; Spiegler et al., 2010) and of asynchronous states in networks of LIF neurons (see, e.g., Abbott and van Vreeswijk, 1993; Brunel and Hakim, 1999; Brunel, 2000; Hansel and Mato, 2003; Brunel and Hansel, 2006), which has already characterized at length the bifurcations that have been described here. On the other hand, only a few studies have investigated systematically how networks respond to dynamical inputs. Tsodyks et al. (1997) and Murphy and Miller (2009) in particular investigated the dynamics of a rate model in the “inhibition-stabilized” regime, i.e., \( I_{\text{inh}} > 1 \) in the rate model. Our analysis is also closely related to finite-size dynamics of networks of integrate-and-fire neurons in the presence of constant external inputs (Brunel and Hakim, 1999; Brunel, 2000; Mattia and Del Giudice, 2002, 2004; Kriener et al., 2008). Dynamics of stochastic versions of rate models have also been recently studied in detail (Bressloff, 2009; Buice et al., 2010). Transfer functions of networks of IF neurons with a specific architecture have also been derived previously in Doiron et al. (2004) and Lindner et al. (2005), where it was shown that global oscillations depend on the correlations in inputs. Transfer functions of gap-junction coupled networks have been studied by Ostojic et al. (2009) and Dugue et al. (2009), where it was shown that gap-junction coupling enhances resonances at the firing frequency of single-cells. The functional consequences of feed-forward inhibition for information transmission have also been studied in a number of recent studies (e.g., Akam and Kullmann, 2010; Kremkov et al., 2010a,b). The present paper is to our knowledge the first detailed comparative study of network transfer functions of both rate models and synaptically coupled networks of LIF neurons.

Input-dependent changes in the network transfer function could underlie several prominent phenomena observed \textit{in vivo} in different areas. In visual cortex, the power spectrum of the LFP typically decays monotonically at low contrast, while a plateau or even a peak in the gamma frequency emerges at high contrast (Henrie and Shapley, 2005). Input-dependent changes of the transfer function at gamma frequency (due to the emergence of a resonance) has been modeled in a framework similar to ours by Kang et al. (2010). Such input-dependent changes could underlie the high information content of the gamma range in the LFP (Belitski et al., 2008), as shown by Mazzoni et al. (2008). It could also underlie the phenomenon of cross-frequency coupling, through which the power of a fast oscillation is strongly modulated by the phase of a slower oscillation, as is the case in the hippocampus (Bragin et al., 1995) as well as in primary sensory cortices (Whittingstall and Logothetis, 2009; Mazzoni et al., 2010).

This study could be generalized in a number of ways. First, it could be generalized to more realistic single-cell models such as the exponential integrate-and-fire (EIF) neuron (Fourcaud-Trocmé et al., 2003), since an efficient method has been introduced to compute both the static and dynamic single-cell transfer function for this model (Richardson, 2007). Interestingly, the transfer function of the EIF model has a high frequency behavior which is qualitatively identical to the rate model (a 1/f decay at high frequency, associated with a phase shift of 90°). This led Ostojic and Brunel (2011) to propose a reduction of the EIF model to an (adaptive) rate model, whose parameters (time constant and static transfer function) can be computed analytically from the parameters of the underlying EIF model. From these results, we expect the dynamics of networks of EIF neurons to be even closer to rate models, compared to LIF networks, provided of course the noise is sufficiently strong.

A second generalization would be to study networks of more than two-populations. It would be of interest, for example, to understand how the presence of functionally different classes of interneurons affect network dynamics. Finally, it would also be of interest to study the transfer function of randomly connected networks (Brunel and Hakim, 1999; Brunel, 2000), or of networks with more structured connectivity (Ben-Yishai et al., 1995; Roxin et al., 2005).

In this paper, we have computed how the population firing rate reacts to time-dependent inputs. Experimentally, the firing rate of small populations of neurons can be observed through MUA. Another experimental observable is the LFP. In cortex and hippocampus, the LFP is assumed to be related to the average synaptic inputs received by pyramidal cells of the main network. Given a model of how the LFP depends on synaptic currents (e.g., Mazzoni et al., 2008), one could compute the dynamics of the LFP signal using the methods used in this paper.

Finally, this work offers insight into the relationship between network connectivity and experimental observables such as phase shifts of different populations in the network. It could be used to predict network connectivity from observed patterns of resonances and phase shifts, in areas where this information is available.

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APPENDIX
ANALYTICAL RESULTS FOR THE SIMPLEST RATE MODEL

Qualitative behaviors of the amplitude of the excitatory transfer function

Using Eq. (40), the amplitude of the modulation of the excitatory population can be written as

$$|r_{EI}(\omega)| = \frac{\mu_{EI}}{(1 - J_{EE})} \sqrt{\frac{(1 - X)^2 + (\omega \tau_E)^2}{(1 + Y - \omega^2 \tau_E \tau_I)^2 + \omega^2 (\tau_E + \tau_I)^2}}$$

(44)

The derivative of Eq. (44) with respect to \(\omega\) is therefore

$$\frac{dr_{EI}(\omega)}{d\omega} = \frac{-\mu_{EI} \omega \tau_I Q(\omega)}{(1 - X)^2 + (\omega \tau_E)^2 \left[(1 + Y - \omega^2 \tau_E \tau_I)^2 + \omega^2 (\tau_E + \tau_I)^2\right]}$$

(45)

where \(Q\) is a second order polynomial in \(\omega^2\),

$$Q(\omega) = \tau_E^2 \omega^4 + 2 \tau_E^2 (1 - X) \omega^2 + (1 - X)^2 \left[2 (1 + Y) \frac{\tau_E}{\tau_I} - (1 + Y)^2\right] - (1 + Y)^2 \tau_I$$(46)

From Eq. (45) it is clear that for \(\omega > 0\), \(dr_{EI}(\omega) = 0\) implies \(Q(\omega) = 0\). \(Q\) is an increasing function of \(\omega^2\) and therefore can have either 0 or 1 zeros. It has a zero at a strictly positive frequency if and only if

$$\Delta = (1 - X)^2 \left[2 (1 + Y) \frac{\tau_E}{\tau_I} - (1 + Y)^2\right] - \tau_I < 0.$$ (47)

If this condition is satisfied, \(dr_{EI}\) is an increasing function of \(\omega\) for small \(\omega\), reaches a maximum at the frequency \(\omega_0\) given by

$$\omega_0 = \sqrt{\frac{(1 - X)^4}{\tau_I^2} - \Delta \tau_I^2 (\tau_E + \tau_I)^2}$$

(48)

and then decays monotonically with frequency. If this condition is not satisfied \(dr_{EI}\) is a monotonically decaying function of frequency.

Qualitative behaviors of the phase difference between E and I

The phase shift between E and I populations can be obtained from Eq. (40):

$$\Delta \phi_{EI}(\omega) = \tan^{-1} \left[\frac{\omega X \tau_E}{1 - X} - \frac{\tau_E}{\tau_I} \right]$$

(49)

The difference of these two arctangent functions is either positive at all frequencies, or only negative. The phase shift is positive (excitatory population leads the inhibitory one) when

$$\tau_E (X + Y) - \tau_I X (1 - X) > 0$$

(50)

and negative (inhibitory population leads the excitatory one) otherwise.

ANALYTICAL RESULTS FOR THE RATE MODEL WITH DELAY

In the rate model with finite delays but zero rise and decay times, Eq. (34) becomes

$$\left(1 + i \omega \tau_E - J_{EE} e^{-i \omega D_E}\right) \left(1 + i \omega \tau_I + J_{II} e^{-i \omega D_I}\right) + J_{II} e^{-i \omega (D_E + D_I)} = 0,$$

(51)

or equivalently

$$1 - \omega^2 \tau_E \tau_I - J_{EE} \left(\cos(\omega D_E) + \omega \tau_I \sin(\omega D_I)\right) + J_{II} \left(\cos(\omega D_I) + \omega \tau_E \sin(\omega D_E)\right)$$

$$- J_{II} e^{i \omega (D_E + D_I)} = 0$$

(52)

We first consider the case \(\tau_E \neq \tau_I\), and/or \(D_E \neq D_I\), Eqs (52,53) give

$$J_{II} = \frac{J_{EE} \left(\sin(\omega D_I) + \omega \tau_I \cos(\omega D_I)\right) - \left(1 - \omega^2 \tau_E \tau_I\right) \sin(\omega (D_E + D_I))}{\sin(\omega D_E) + \omega \tau_E \cos(\omega D_E) - \frac{-\omega \tau_E \tau_I \cos(\omega (D_E + D_I))}{\sin(\omega D_E) + \omega \tau_I \cos(\omega D_I)}}$$

(54)

$$J_{II} = \frac{-\left(1 + \omega^2 \tau_E \tau_I\right) + J_{EE} \left(\cos(\omega D_E) + \omega \tau_I \sin(\omega D_I)\right)}{\cos(\omega (D_E + D_I))}$$

(55)

Thus, bifurcation lines can be obtained parametrically by varying \(\omega\) and plotting the resulting values of \(J_{II}\) and \(J_{II}'\).

In the case \(D_E = D_I = D\) and \(\tau_E = \tau_I = \tau\), there is a solution to Eqs (52,53) which is simply \(\sin(\omega D) + \omega \tau \cos(\omega D) = 0\), i.e., \(\tan(\omega D) = -\omega \tau\). Solving this equation gives the frequency of the 1–I instability, which occurs when

$$J_{II} = -\frac{1}{\cos(\omega D) + \frac{J_{EE} \omega \tau \sin(\omega D)}{1 - J_{II} e^{-i \omega D}}}$$

(56)

NUMERICAL PROCEDURE FOR THE PHASE DIAGRAMS IN FIGURES 5, 6, 7, AND 9

In the rate models with delay and the LIF model, the whole region where the asynchronous state is stable is discretized. On each point of the resulting grid, the number of extrema in the amplitude of the excitatory transfer function is counted in the interval 0–500 Hz. The color of the corresponding location then encodes the number of extrema in this interval, as described in the figure legends.