The effects of degree distributions in random networks of Type-I neurons

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We consider large networks of theta neurons and use the Ott/Antonsen ansatz to derive degree-based mean field equations governing the expected dynamics of the networks. Assuming random connectivity we investigate the effects of varying the widths of the in- and out-degree distributions on the dynamics of excitatory or inhibitory synaptically coupled networks, and gap junction coupled networks. For synaptically coupled networks, the dynamics are independent of the out-degree distribution. Broadening the in-degree distribution destroys oscillations in inhibitory networks and decreases the range of bistability in excitatory networks. For gap junction coupled neurons, broadening the degree distribution varies the values of parameters at which there is an onset of collective oscillations. Many of the results are shown to also occur in networks of more realistic neurons.

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I. INTRODUCTION

It is well-known that the structure of a network can have a significant effect on its dynamics. One type of network of great interest is that of networks of neurons, and much effort has gone into investigating this issue [1-5]. In networks of neurons, the nodes are normally thought of as individual neurons, while the edges describe connections between neurons. These connections can be directed (in the case of synaptic connections [6]) or undirected (in the case of gap junction connections [7, 8]). One of the important properties of a node in a network is its degree: a node’s in-degree is the number of connections to it, and a node’s out-degree is the number of connections from it. In the case of undirected connections a node simply has a degree, as there is no distinction between incoming or outgoing connections. Previous work on understanding the dynamics of networks of neurons has considered the effects of correlations between the in- and out-degrees of individual neurons [3, 4, 9, 12] and of degree assortativity, in which the probability that two neurons are connected is influenced by the degrees of the two neurons [2, 13, 15].

In this paper we consider the effects of varying the widths of the distributions of in-degrees and out-degrees on large, randomly connected networks of Type-I neurons, i.e., neurons whose onset of firing is through a saddle-node-on-invariant-circle (SNIC) bifurcation. The analysis is done of networks of theta neurons, as these are the normal form of the SNIC bifurcation. They also have the property of being amenable to the use of the Ott/Antonsen ansatz [16, 17], a now well-used method for deriving equations governing the evolution of order parameter-like quantities, valid for large networks with particular forms of heterogeneity [18-22].

A common assumption in creating random networks of neurons is that there is a fixed probability of connecting any two neurons [23, 24]. Such networks are often referred to as Erdős-Rényi and have a binomial distribution of degrees, with the ratio of mean degree to standard deviation of degrees going to zero as network size goes to infinity. However, real networks of neurons are observed to have properties incompatible with this assumption [25].

Previous work on investigating the effects of degree distributions includes that of Roxin [1]. He found in an inhibitory network of leaky integrate-and-fire neurons that broadening the distribution of in-degrees suppressed macroscopic oscillations. This result was reproduced in a simplified rate model which included the heterogeneity in neuronal input due to the in-degree of cells. He also considered a network with both excitatory and inhibitory neurons and investigated the effects of varying the in- and out-degree distributions of the recurrent excitatory connections. In other work, [26] studied the effects of degree distribution in feedforward networks.

We consider synaptic coupling in Sec. [11] and gap junction coupling in Sec. [11] Both sections start with a derivation of the relevant equations using the Ott/Antonsen ansatz and then give some numerical results. Most results use
Lorentzian distributions of heterogeneous parameters, and in Sec. [IV] we briefly discuss results using normal and uniform distributions. We conclude in Sec. [V] Appendix [A] contains the description of the network of Morris-Lecar neurons used to verify some of the results derived for networks of theta neurons.

II. SYNAPTIC COUPLING

We first consider networks of theta neurons coupled by synaptic input currents with a timescale $\tau$. Alternative formulations could model synaptic input by using dynamic synapses [27] or input current pulses with both a rise time and a decay time [6].

A. Model and Theory

We consider a network of $N$ theta neurons, each of which has dynamics described by

$$\frac{d\theta_i}{dt} = 1 - \cos \theta_i + (1 + \cos \theta_i)(\eta_i + I_i(t))$$

(1)

for $i = 1, \ldots, N$, where the input current to neuron $i$ is

$$I_i(t) = \frac{K}{\langle k \rangle} \sum_{j=1}^{N} A_{ij} u_j(t)$$

(2)

and the synaptic variables have dynamics given by

$$\tau \frac{du_j}{dt} = \sum_{m \in \mathbb{Z}} \delta(t - T_j^m) - u_j,$$

(3)

where $T_j^m$ is the $m$th firing time of neuron $j$, defined to happen every time $\theta_j$ increases through $\pi$, and $\delta(\cdot)$ is the Dirac delta. Thus every time $\theta_j$ increases through $\pi$, $u_j$ is instantaneously incremented by an amount $1/\tau$, and between firing times it decays as $e^{-t/\tau}$. $K$ is the strength of connections between neurons (which may be positive or negative), $\langle k \rangle$ is the average in-degree in the network, and $A$ describes the connectivity of the network, i.e., $A_{ij} = 1$ if there is a connection from neuron $j$ to neuron $i$ and $A_{ij} = 0$ otherwise. We have $\langle k \rangle = \sum_{i,j} A_{ij}/N$. The $\eta_i$ are randomly chosen from a Lorentzian

$$g(\eta) = \frac{\Delta/\pi}{(\eta - \eta_0)^2 + \Delta^2}$$

(4)

with centre $\eta_0$ and half-width-at-half-maximum (HWHM) $\Delta$, which introduces heterogeneity to the network. The Lorentzian is chosen so that analytical progress can be made. We discuss results for other distributions below.

If the input current $I_i$ is constant, the theta neuron shows one of two types of behaviour. For $\eta_i + I_i < 0$, (1) has two fixed points, one stable and one unstable. For $\eta_i + I_i > 0$, (1) has no fixed points and $\theta_i$ increases with time, showing periodic oscillations with frequency $\sqrt{\eta_i + I_i}/\pi$ [28]. The bifurcation at $\eta_i + I_i = 0$ is a SNIC bifurcation. Note that under the transformation $V = \tan(\theta/2)$ a network of theta neurons is exactly equivalent to a network of quadratic integrate-and-fire neurons with infinite threshold and reset values [20]. We now proceed to analyse the network dynamics, using ideas similar to those in [8, 13, 21, 22, 30].

We assume that the network is characterised by two functions: firstly the degree distribution $P(k)$, normalised to sum to 1, where $k = (k_{in}, k_{out})$ and $k_{in}$ and $k_{out}$ are the in- and out-degrees of a neuron with degree $k$, respectively, and secondly the assortativity function $\alpha(k' \rightarrow k)$ giving the probability of a connection from a neuron with degree $k'$ to one with degree $k$, given that such neurons exist. We also make the mean field assumption that the dynamics of a neuron depend only on its degree $k$, thus effectively averaging the dynamics of all neurons with the same degree.

In the limit of large $N$ and large in- and out-degrees, the network is described by the distribution \( f(\theta, \eta|k, t) \) where \( f(\theta, \eta|k, t) d\theta d\eta \) is the probability that a neuron with degree $k$ has phase in $[\theta, \theta + d\theta]$ and value of $\eta$ in $[\eta, \eta + d\eta]$ at time $t$. This distribution satisfies the continuity equation

$$\frac{\partial f}{\partial t} + \frac{\partial}{\partial \eta}(vf) = 0$$

(5)

where (from (1)-(3))

$$v(\theta, k, \eta, t) = 1 - \cos \theta + (1 + \cos \theta)|\eta + I(k, t)|,$$

(6)

$$I(k, t) = \frac{K N}{\langle k \rangle} \sum_{k'} P(k') \alpha(k' \rightarrow k) u(k', t),$$

(7)

and

$$\tau \frac{du(k, t)}{dt} = \tilde{F}(k, t) - u(k, t),$$

(8)

where $\tilde{F}(k, t)$ is the firing rate of neurons with degree $k$ at time $t$.

The Ott/Antonsen ansatz gives the dynamics for the order parameter for neurons with degree
where $b(k, t)$ is the expected value of $e^{i\theta}$ for neurons with degree $k$, i.e.

$$b(k, t) = \int_{-\infty}^{\infty} \int_{0}^{2\pi} f(\pi, \eta; k, t) e^{i\theta} d\theta d\eta.$$  \hfill (9)

The firing rate of neurons with degree $k$ at time $t$ is the expected value of the flux through $\theta = \pi$ \cite{31, 32}, i.e.

$$\hat{F}(k, t) = \int_{-\infty}^{\infty} f(\pi, \eta; k, t) d\eta = \frac{1}{\pi} \text{Re} \left( \frac{1 - b(k, t)}{1 + b(k, t)} \right)$$  \hfill (10)

where the overline indicates complex conjugate.

We define $F(b(k, t)) \equiv \hat{F}(k, t)$.

With neutral assortativity \cite{21},

$$a(k' \rightarrow k) = \frac{k_{\text{out}}' k_{\text{in}}}{N(k)}$$  \hfill (11)

and with independent in- and out-degrees the degree distribution $P(k')$ factorises as $P(k') = p_{\text{in}}(k_{\text{in}}') p_{\text{out}}(k_{\text{out}}')$, where $p_{\text{in}}$ and $p_{\text{out}}$ are the marginal distributions of the relevant degrees, so

$$\frac{KN}{\langle k \rangle} \sum_{k'} P(k') a(k' \rightarrow k) u(k', t)$$  \hfill (12)

which is independent of $k_{\text{out}}$ and contributes to the “input” to neurons with degree $k$. Thus $b(k, t)$ must also be independent of $k_{\text{out}}$ and so must $F(b(k, t))$ and $u(k, t)$. Thus (12) simplifies to

$$\frac{KN}{\langle k \rangle} \sum_{k'} P(k') a(k' \rightarrow k) u(k', t) = \frac{K k_{\text{in}}}{\langle k \rangle} \sum_{k_{\text{in}}'} p_{\text{in}}(k_{\text{in}}') u(k_{\text{in}}', t)$$  \hfill (13)

and we see that the distribution of out-degrees does not affect the expected dynamics. This was observed by Roxin in \cite{1}, although he observed that broadening the out-degree distribution increases the amplitude of the cross-correlation of synaptic currents, something we do not consider here.

We have

$$\frac{\tau}{dt} u(k_{\text{in}}, t) = F(b(k_{\text{in}}, t)) - u(k_{\text{in}}, t)$$  \hfill (14)

and defining

$$s(t) \equiv \sum_{k_{\text{in}}} p_{\text{in}}(k_{\text{in}}) u(k_{\text{in}}, t)$$  \hfill (15)

we see that $s$ satisfies

$$\frac{\tau}{dt} s = \sum_{k_{\text{in}}} p_{\text{in}}(k_{\text{in}}) F(b(k_{\text{in}}, t)) - s$$  \hfill (16)

and the dynamics of $b$ are given by

$$\frac{\partial b(k_{\text{in}}, t)}{\partial t} = \frac{-i[b(k_{\text{in}}, t) - 1]^2}{2} + \frac{[b(k_{\text{in}}, t) + 1]^2}{2} \left[ -\Delta + i\eta + iK k_{\text{in}} s \right]$$  \hfill (17)

Equations (16)-(17) form a set of $N_{k_{\text{in}}} + 1$ ordinary differential equations (ODEs) governing the network’s dynamics, where $N_{k_{\text{in}}}$ is the number of distinct in-degrees in the network. In the next section we give some numerical results showing the possible dynamics of this set of equations.

B. Results

We first consider inhibitory coupling, i.e., $K < 0$.

1. Inhibitory coupling

Consider the parameter values $\eta_0 = 1, \Delta = 0.05, \tau = 1, K = -2, \langle k \rangle = 100$. Having $\eta_0 > 0$ indicates that when uncoupled, most neurons would be firing rather than quiescent. We choose $p_{\text{in}}$ to be uniform with mean $\langle k \rangle = 100$ and write its support as $[100 - \sigma, 100 + \sigma]$. For $\sigma = 5$ (a narrow distribution) we obtain global oscillations, see Fig. 1(a). However, when the in-degree distribution is made broader ($\sigma = 50$), the oscillations die out: see Fig. 1(b). Note the independence of the dynamics on the distribution of out-degrees, $p_{\text{out}}(k_{\text{out}})$, as expected.

To numerically solve (16)-(17) we treat $k_{\text{in}}$ as a continuous variable and discretise the support of $p_{\text{in}}$ using 100 evenly spaced points, and use $p_{\text{in}} = 1/100$ at each of those points, effectively using the midpoint rule. To create the network used in (1)-(3) we randomly sample $N$ in-degrees from $p_{\text{in}}(k_{\text{in}})$ and $N$ out-degrees from $p_{\text{out}}(k_{\text{out}})$.
choosing until the sum of the in-degrees equals the sum of the out-degrees, then use the configuration model to connect the network. Any self or multiple connections are then removed by random rewiring, keeping the degrees fixed. To solve (1)-(3) we used Euler’s method with a step-size of 0.001.

The destruction of oscillations seen in Fig. 1 seems due to a Hopf bifurcation. Using pseudo-arclength continuation to follow the stable fixed point of (16)-(17) as \( \sigma \) is decreased we find a Hopf bifurcation at \( \sigma \approx 31.4 \), and continuing that bifurcation as both \( \sigma \) and \( \tau \) are varied we obtain the curve in Fig. 2. For any \( \tau \) for which an oscillation occurs, increasing \( \sigma \) will destroy the oscillations, and for small \( \sigma \), a value of \( \tau \) which is either too large or too small will also destroy oscillations. (We varied \( \tau \) here just as an example; we could equally well vary other parameters such as \( \eta_0 \) or \( \Delta \).) The destruction of oscillations in an inhibitory network by broadening the in-degree distribution was also observed by Roxin [1]. He analysed a heuristic rate model containing a fixed delay (since he used delayed synapses) and found a Hopf bifurcation in that model, in agreement with the results shown here.

To investigate the generality of our result we now consider a beta distribution of in-degrees, with equal parameters greater than one, shifted to have mean 100 and support on \([50, 150]\) (\( \sigma = 50 \)). For (1)-(3) \( N = 500 \) neurons were used and the initial conditions were \( \theta_i = u_i = 0 \), and for (16)-(17) we used \( b = 1 \) and \( s = 0 \).

\[ p_{in}(k_{in}) = \begin{cases} Cx^{\alpha-1}(1-x)^{\alpha-1}, & 0 \leq x \leq 1 \\ 0, & \text{otherwise} \end{cases} \]

(18)

where \( x = (k_{in} - 50)/100 \) and \( C \) is a normalisation factor. Increasing \( \alpha \) narrows the distribution, as shown in the inset of Fig. 3. Varying \( \alpha \) and \( \tau \) we find a curve of Hopf bifurcations, shown in Fig. 3, which shows the same qualitative behaviour as for the uniform distribution.

To further demonstrate the generality of our results we now consider a network of 500 Morris-Lecar neurons [34], known to undergo a SNIC bifurcation as the input current is increased [9]. The network equations are given in Appendix A. The in-degree distribution is uniform on \([100 - \sigma, 100 + \sigma]\) and the out-degree is uniform on \([50, 150] \). For each different value of \( \sigma \) we generate a network as explained above and for each network we vary \( \tau \), the synaptic timescale, integrating for 50 seconds at each value of \( \tau \). Defining \( \hat{s} = N^{-1} \sum_{i=1}^{N} s_i \) we discard data from the first 45 seconds and calculate the standard deviation of \( \hat{s} \) over the last 5 seconds, plotting that in Fig. 4. Large values indicate oscillations while small values indicate an approximate steady state. We see results consistent with those in Figs. 2 and 3. Thus we con-
clude that broadening the in-degree distribution of an inhibitory network of Type-I neurons acts to destroy global oscillatory behaviour. This is presumably due to having a wider range of dynamics for neurons with different in-degrees, making them harder to synchronise. We now consider excitatory coupling, i.e., $K > 0$.

2. Excitatory coupling

Consider the parameter values $\Delta = 0.05, \tau = 1, K = 5, \langle k \rangle = 100$. Varying $\eta_0$ we expect a region of bistability between a high activity steady state and a low activity steady state, as is often found in excitatory networks [35]. This is found, as shown in Fig. 5 where saddle-node bifurcations mark the boundaries of the bistable region. Varying the width of the in-degree distribution ($\sigma$) varies the width of the bistable region. In particular, widening the in-degree distribution narrows the width of the bistable region. Following the saddle-node bifurcations seen in Fig. 5 we obtain Fig. 6. Similar behaviour was seen for a beta distribution of in-degrees (not shown).

We reproduced this behaviour in a network of $N = 500$ Morris-Lecar neurons with parameters as given in Appendix A. For networks with $\sigma = 10$ and $90$ we quasistatically varied $I_0$, integrating for 10 seconds at each value. We define $\bar{s}$ to be the mean of $\dot{s}$ over the last 2 seconds of simulation and plot this in Fig. 7. The results are qualitatively the same as in Fig. 5: $\bar{s}$ is lower when $\sigma = 90$ than when $\sigma = 10$, and the left-most saddle-node bifurcation is moved more than the right-most when $\sigma$ is varied. The threshold for firing for single neuron is $I_0 \approx 39.69$ so the jumps occur at values of $I_0$...
less than this, consistent with the results in Fig. 6. Thus we conclude that for an excitatory network of Type-I neurons, broadening the in-degree distribution narrows the range of values of the mean input for which the network is bistable. The effects of varying other parameters could equally well be investigated using the techniques shown here.

III. GAP JUNCTIONS

We now consider theta neurons coupled by gap junctions [7, 36]. Gap junctional coupling is well-known to induce synchrony in networks of neurons [8, 37, 38]. The quadratic integrate-and-fire (QIF) neuron [39] with infinite firing threshold and reset to \( V = -\infty \) is equivalent under the transformation \( V = \tan(\theta/2) \) to a theta neuron [40], and since gap junction coupling is through voltage differences it is easier to start with a network of QIF neurons. Our analysis is similar to that in [31]; also see [41, 43]. A theta neuron is an excitable system, so our results add to those on coupled excitable systems [44, 47].

A. Model and Theory

Consider a network of \( N \) gap-junction coupled QIF neurons governed by

\[
\frac{dV_j}{dt} = \eta_j + V_j^2 + g \frac{k_j}{\langle k \rangle} \sum_{l=1}^{N} A_{jl}(V_l - V_j) \tag{19}
\]

for \( j = 1, \ldots, N \) together with the rule that if \( V_j(t^-) = -\infty \) then \( V_j(t^+) = -\infty \), and neuron \( j \) is said to fire at this time \( t \). \( A \) describes the connectivity of the network, where \( A_{ji} = 1 \) if neurons \( j \) and \( l \) are connected and zero otherwise. Since gap junctional coupling is not directional we have \( A_{ji} = A_{ij} \). \( k_j \) is the degree of the \( j \)th neuron, i.e. \( k_j = \sum_l A_{jl} \), \( \langle k \rangle \) is the mean degree, as above, and \( g \) is the strength of coupling (non-negative). The \( \eta_j \) are randomly chosen from a distribution \( h(\eta) \).

We rewrite (19) as

\[
\frac{dV_j}{dt} = \eta_j + V_j^2 - \frac{g k_j V_j}{\langle k \rangle} + g \frac{\langle k \rangle}{\sum_{l=1}^{N} A_{jl} V_l} \tag{20}
\]
Now let \( V_j = \tan(\theta_j/2) \). Then
\[
\frac{dV_j}{dt} = \frac{d\theta_j/dt}{2 \cos^2(\theta_j/2)} = \eta_j + \tan^2(\theta_j/2) - \frac{g k_j}{\langle k \rangle} \tan(\theta_j/2) + \frac{g}{\langle k \rangle} \sum_{l=1}^{N} A_{jl} \tan(\theta_l/2)
\]
so
\[
\frac{d\theta_j}{dt} = 1 - \cos \theta_j + (1 + \cos \theta_j)\eta_j - \frac{g k_j}{\langle k \rangle} \tan(\theta_j/2) + \frac{g}{\langle k \rangle} \sum_{l=1}^{N} A_{jl} \tan(\theta_l/2).
\]
Noting that
\[
\tan(\theta/2) = \frac{\sin \theta}{1 + \cos \theta},
\]
we have
\[
\frac{d\theta_j}{dt} = 1 - \cos \theta_j - \frac{g k_j}{\langle k \rangle} \sin \theta_j + (1 + \cos \theta_j)\eta_j + \frac{g}{\langle k \rangle} \sum_{l=1}^{N} A_{jl} \tan(\theta_l/2).
\]
When a neuron fires, at \( \theta = \pi \), the term involving tan becomes infinite. To avoid this problem we follow [8] and replace \( \tan(\theta/2) \) in [24] by
\[
q(\theta) = \frac{\sin \theta}{1 + \cos \theta + \epsilon},
\]
where \( 0 < \epsilon \ll 1 \), thereby removing the singularity. We take the limit \( \epsilon \to 0 \) below.

We analyse the system in a similar way as in Sec. II. The system is described by the probability density function \( f(\eta, \theta|k, t) \) which satisfies [46, 50]
\[
\frac{\partial f}{\partial t} + \frac{\partial}{\partial \theta}(fv) = 0,
\]
where
\[
v(\eta, \theta, k, t) \equiv 1 - \cos \theta - \frac{g k_{in}}{\langle k \rangle} \sin \theta + (1 + \cos \theta) [\eta + g T(k, t)],
\]
and
\[
T(k, t) = \frac{N}{\langle k \rangle} \sum_{k'} P(k') a(k' \rightarrow k) Q(k', t),
\]
so
\[
Q(k', t) = \int_{-\infty}^{\infty} \int_{0}^{2\pi} f(\eta, \theta|k', t) q(\theta) \, d\theta \, d\eta,
\]
and \( k = (k_{in}, k_{out}) \), \( P(k') \) is the distribution of degrees of a neuron, and \( a(k' \rightarrow k) \) is the probability of a connection from a neuron with degree \( k' \) to one with degree \( k \). But connections are undirected, so a neuron just has a degree. Thus we write
\[
T(k, t) = \frac{N}{\langle k \rangle} \sum_{k'} P(k') a(k' \rightarrow k) Q(k', t)
\]
where \( P(k) \) is the degree distribution, and
\[
Q(k', t) = \int_{-\infty}^{\infty} \int_{0}^{2\pi} f(\eta, \theta|k', t) q(\theta) \, d\theta \, d\eta
\]
is the expected value of \( q(\theta) \) for neurons with degree \( k' \) at time \( t \). From [31], assuming that \( h(\eta) \) is a Lorentzian with median \( \eta_0 \) and HWHM \( \Delta \), we have
\[
\frac{\partial b(k, t)}{\partial t} = (i \eta_0 - \Delta)(1 + b)^2 - i (1 - b^2) + i (1 + b)^2 g T + g (1 - b^2)
\]
where \( T(k, t) \) and \( Q(k', t) \) are as above and
\[
b(k, t) = \int_{-\infty}^{\infty} \int_{0}^{2\pi} f(\eta, \theta|k, t) e^{i \theta} \, d\theta \, d\eta
\]
is the complex-valued order parameter for neurons with degree \( k \) at time \( t \).

By expanding \( q(\theta) \) in a Fourier series it was shown in [31] that
\[
Q(k', t) = \sum_{m=1}^{\infty} [c_m b(k', t)^m + \text{c.c.}]
\]
where “c.c.” is the complex conjugate of the previous term and
\[
c_m = i [(\sqrt{2 \epsilon + \epsilon^2} - 1 - \epsilon)^{m+1} - (\sqrt{2 \epsilon + \epsilon^2} - 1 - \epsilon)^{m-1}] / 2 \sqrt{2 \epsilon + \epsilon^2}
\]
Recently, [32] showed that
\[
\lim_{\epsilon \to 0} c_m = i (-1)^m
\]
and thus
\[
Q(k', t) = i \sum_{m=1}^{\infty} (-1)^m [b(k', t)^m - \bar{b}(k', t)^m] / 2 \text{Im}[b(k', t)] / [1 + b(k', t)][1 + \bar{b}(k', t)]
\]
where we have summed the geometric series.
Defining
\[ w \equiv \frac{1 - \bar{b}}{1 + b} \]  
(38)

(so \( b = (1 - \bar{w})/(1 + \bar{w}) \)) we find that \( w \) satisfies
\[ \frac{\partial w(k,t)}{\partial t} = i\eta_0 + \Delta - i[|w(k,t)|^2 + gT(k,t) - gw(k,t)], \]
(39)

and writing \( w = \pi \phi + iV \) where \( \phi \) and \( V \) are real we find that \( Q(k',t) = V(k',t) \) and the real and imaginary parts of (39) give

\[ \frac{\partial \phi(k,t)}{\partial t} = \frac{\Delta}{\pi} + 2\phi(k,t)V(k,t) - g\phi(k,t) \]  
(40)

\[ \frac{\partial V(k,t)}{\partial t} = \eta_0 - \pi^2[\phi(k,t)]^2 + [V(k,t)]^2 \]
\[ + g[T(k,t) - V(k,t)] \]  
(41)

where
\[ T(k,t) = \frac{N}{\langle k \rangle} \sum_{k'} P(k')a(k' \rightarrow k)V(k',t). \]  
(42)

The interpretation of \( \phi \) and \( V \) is that \( \phi(k,t) \) is the expected firing frequency of neurons with degree \( k \) at time \( t \), and \( V(k,t) \) is the mean voltage of QIF neurons with degree \( k \) at time \( t \) where voltage \( V \) and \( \theta \) are related through \( V = \tan(\theta/2) \) [32]. Note that (40)-(41) are completely equivalent to (39).

Assuming neutral assortativity we have
\[ a(k' \rightarrow k) = \frac{k'k}{N\langle k \rangle} \]  
(43)

so that
\[ T(k,t) = \frac{k}{\langle k \rangle^2} \sum_{k'} k'P(k')V(k',t). \]  
(44)

Note that if all neurons have the same degree then \( T = V \), so the last term in (41) vanishes, and (40)-(41) reduce to a pair of ODEs. A special case of this is all-to-all coupling, which was studied in [22]. Also, (44) is invariant under the scaling \( k \rightarrow \alpha k, P(k) \rightarrow P(k/\alpha) \), i.e. only degree relative to mean degree is of relevance.

B. Results

First consider a network with \( \Delta = 0.01 \), and \( g = 0.4 \), with \( \langle k \rangle = 100 \). As above we consider a uniform distribution of degrees on \([100 - \sigma, 100 + \sigma]\). Consistent with [11][32] we find that when increasing \( \eta_0 \) the transition to periodic firing is through a SNIC bifurcation, as shown in Fig. 8. Increasing the width of the degree distribution increases the value of \( \eta_0 \) at which collective oscillations start. Note that even for \( \sigma = 0 \) (i.e. identical degrees) \( \eta_0 \) can be small and positive yet the network is quiescent, as also found by [11] [42]. Note also the small range of \( \eta_0 \) values as \( \sigma \) is varied.

Now consider a more heterogeneous network with \( \Delta = 0.05 \) and \( \eta_0 = 0.2 \), i.e. well above threshold so that most neurons would fire if uncoupled, again with \( \langle k \rangle = 100 \) and a uniform degree distribution on \([100 - \sigma, 100 + \sigma]\). Consistent with the results in [11][32] we find that upon increasing \( g \) (the strength of coupling) the transition to firing is through a Hopf bifurcation, as shown in Fig. 9. Increasing the width of the degree distribution decreases the value of \( g \) at which collective oscillations start. This bifurcation is reminiscent of that which occurs in all-to-all connected networks of Winfree oscillators [31]: increasing the coupling strength causes the onset of oscillations through a Hopf bifurcation [30][32][33]. As is also seen in networks of Winfree oscillators, decreasing \( \Delta \) (the level of heterogeneity) has the same effect as increasing \( g \), producing oscillations via a Hopf bifurcation (not shown).

We tried to reproduce the trend in Fig. 8 using a network of gap junction coupled Morris-Lecar neurons, the equations of which are given in Appendix A. We chose the \( I_0 \) from a Lorentzian with HWHM \( \Delta = 0.01 \) and coupling strength \( \epsilon = 0.3 \). We were unable to reproduce the trend. This is likely due to the sensitivity of the network to the value of \( I_0 \): notice the very small range of \( \eta_0 \) values in Fig. 8. The variation in values of \( I_0 \) at

\[ \text{FIG. 8: SNIC bifurcation curve of a fixed point of (40)-(41). A stable fixed point exists to the left of the curve and stable periodic oscillations to the right. Parameters: } \Delta = 0.01, \ g = 0.4, \langle k \rangle = 100, \ P(k) \text{ is a uniform distribution on } [100 - \sigma, 100 + \sigma]. \]
IV. GAUSSIAN OR UNIFORM DISTRIBUTION OF $\eta_i$

All of the results so far have involved a Lorentzian distribution of a heterogeneous parameter, either the input currents to theta neurons or to Morris-Lecar neurons. In this section we investigate whether we obtain qualitatively similar results for other distributions.

Using either a Gaussian or uniform distribution of $I_i$ in a Morris-Lecar network we obtained qualitatively the same results as in Figs. 4 and 7 for synaptic coupling (results not shown). We investigated the effects shown in Fig. 8 in a network of gap-junction coupled Morris-Lecar neurons for both uniform and normally distributed $I_i$. The results are shown in Fig. 11. For each value of $\sigma$ we created a network and a realisation of the $I_i$, and then used bisection in $I_0$ to approximately determine the transition from quiescence to periodic firing (with large period). For broader distributions (panel (a)) we obtained the same trend as in Fig. 8 while for narrower distributions we seem to obtain the opposite trend (panel (b)). Note that the threshold for firing for a single neuron is $I_0 \approx 39.693455$, so all bifurcations occur for $I_0$ less than this, in contrast with the results in Fig. 8. Such an effect has been seen before in excitable systems [44] indicating that the Lorentzian distribution of heterogeneity, while providing analytical insight, may not give generic results.

We reproduced the results in Fig. 9 with the $I_i$ taken from a unit Gaussian (normal distribution), as shown in Fig. 12(a). Choosing the $I_i$ from a uniform distribution on $[-1, 1]$ we obtain the results in Fig. 12(b). Quasistatically sweeping $\epsilon$ up and down for $\sigma = 10$ we found a region of bistability between an approximate steady state and a macroscopic oscillation, suggesting that the Hopf bifurcation seen is subcritical. For clarity, we only show the results of increasing $\epsilon$ for both networks. The effect of varying the width of the in-degree distribution is the same: broadening the distribution moves the Hopf bifurcation to a lower value of $\epsilon$.

V. SUMMARY

We have derived approximate equations describing the expected dynamics of large networks of theta neurons, under the assumption that the heterogeneous parameter has a Lorentzian distribution. We have chosen the case of neutral asessorativity within the networks and independent in- and out-degrees, concentrating on the effects...
of varying the widths of the in- and out-degree distributions. We have investigated both synaptic and gap junctional coupling. Numerical bifurcation analysis has enabled us to determine the effects of varying the degree distributions on the networks’ dynamics.

For synaptically coupled inhibitory neurons, broadening the in-degree distribution destroys macroscopic oscillations while for excitatory networks, it narrows the range of values of mean input for which the network is bistable. The dynamics are independent of the out-degree distribution. For gap junctional coupling, broadening the degree distribution causes SNIC or Hopf bifurcations associated with the onset of collective firing to move in parameter space. Most of the results are confirmed to occur in networks of more realistic neurons with different distributions of heterogeneity, but the use of Lorentzian distributions may not always give generic results.

This work could be generalised in several ways. One is to extend it to cover coupled populations of excitatory and inhibitory neurons [1, 27, 54]. This would naturally result in more parameters to explore, as distributions of degrees relating to up to four types of connection would need to be specified. Another area of interest is the inclusion of noise in the neurons’ dynamics. Noise is known to play a significant role in neural dynamics [6, 55] yet its presence would invalidate the use of the Ott/Antonsen ansatz which lies behind the derivations in this paper. Goldobin et al. have made progress in this area, developing perturbation theory away from noise-free case [56–60], and Ratas and Pyragas have applied these ideas to networks of theta neurons [61].

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FIG. 11: SNIC bifurcations for gap-junction coupled Morris-Lecar neurons. (a): $I_i$ chosen from a uniform distribution on $[-1/2,1/2]$ (blue circles, solid curve) or a normal distribution with centre zero and standard deviation $1/3$ (red triangle, dashed curve). (b): $I_i$ chosen from a uniform distribution on $[-1/8,1/8]$ (blue circles, solid curve) or a normal distribution with centre zero and standard deviation $1/10$ (red triangle, dashed curve). The curves result from fitting the values of $I_0$ as a cubic function of $\sigma$ and are to guide the eye. Parameters: $N = 2500, \epsilon = 0.3$.

FIG. 12: Evidence of Hopf bifurcations in networks of gap-junction coupled Morris-Lecar neurons. (a): $I_i$ taken from a unit Gaussian distribution. The Hopf bifurcations appear to be supercritical. (b): $I_i$ taken from a uniform distribution on $[-1,1]$. The Hopf bifurcation for $\sigma = 10$ appears to be subcritical, as explained in the text. For each panel, on the vertical axis we plot the standard deviation of $\hat{s}$ (the mean of the $s_i$) over a period of 10 seconds, having discarded the first 10 seconds as transients. $N = 2500$. 

Appendix A: Morris-Lecar equations

1. Synaptic coupling

For synaptic coupling the equations for $N$ neurons are

\[ C \frac{dV_i}{dt} = g_L(V_L - V_i) + g_{Ca}m_\infty(V_i)(V_{Ca} - V_i) \]

\[ + g_K n_i(V_K - V_i) + I_0 + \frac{\epsilon}{\langle k \rangle} \sum_{j=1}^{N} A_{ij}s_j \]

\[ \frac{dn_i}{dt} = \frac{\lambda_0(w_\infty(V_i) - n_i)}{\tau_n(V_i)} \]  

\[ \tau \frac{ds_i}{dt} = s_\infty(V_i) - s_i \]  

where

\[ m_\infty(V) = 0.5(1 + \tanh [(V - V_1)/V_2]) \]  

\[ w_\infty(V) = 0.5(1 + \tanh [(V - V_3)/V_4]) \]  

\[ \tau_n(V) = \frac{1}{\cosh ((V - V_3)/(2V_4))} \]  

\[ s_\infty(V) = 1 + \tanh (V/10) \]

Parameters are $V_1 = -1.2, V_2 = 18, V_3 = 12, V_4 = 17.4, \lambda_0 = 1/15msec^{-1}, g_L = 2, g_K = 8, g_{Ca} = 4, V_L = -60, V_{Ca} = 120, V_K = -80, C = 20\mu F/cm^2$ and these are unchanged throughout the paper. Voltages are in mV and conductances are in mS/cm$^2$. These are taken from [34] but we have added synaptic dynamics. The threshold for firing for single neuron is $I_0 \approx 39.693455$.

For Fig. 4 we set $\epsilon = -1mS/cm^2, I_0 = 41$. The $I_i$ were taking from a Lorentzian with mean zero and HWHM 0.01. For Fig. 7 we set $\epsilon = 25mS/cm^2, \tau = 20, N = 500$. The $I_i$ were taking from a Lorentzian with mean zero and HWHM 0.01.

2. Gap junction coupling

For gap-junctional coupling we use

\[ C \frac{dV_i}{dt} = g_L(V_L - V_i) + g_{Ca}m_\infty(V_i)(V_{Ca} - V_i) \]

\[ + g_K n_i(V_K - V_i) + I_0 + \frac{\epsilon}{\langle k \rangle} \sum_{j=1}^{N} A_{ij}s_j \]

\[ \frac{dn_i}{dt} = \frac{\lambda_0(w_\infty(V_i) - n_i)}{\tau_n(V_i)} \]  

where $m_\infty(V), w_\infty(V)$ and $\tau_n(V)$ and all other parameters are as above. For Fig. 10 we set $I_0 = 40$ and choose the $I_i$ from a Lorentzian with mean zero and HWHM 0.5.

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