# A scalar Poincaré map for anti-phase bursting in coupled inhibitory neurons with synaptic depression.

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# 2 ABSTRACT

Short-term synaptic plasticity is found in many areas of the central nervous system. In the inhibitory half-centre central pattern generators involved in locomotion, synaptic depression is believed to act as a burst termination mechanism, allowing networks to generate anti-phase bursting patterns of varying periods. To better understand burst generation in these central patter generators, we study a minimal network of two neurons coupled through depressing synapses. 8 Depending on the strength of the synaptic conductance between the two neurons, this network 9 can produce symmetric n-n anti-phase bursts, where neurons fire n spikes in alternation, with the period of such solutions increasing with the strength of the synaptic conductance. Relying on the timescale disparity in the model, we reduce the eight-dimensional network equations to a fully-explicit scalar Poincaré burst map. This map tracks the state of synaptic depression from one burst to the next and captures the complex bursting dynamics of the network. Fixed points of this map are associated with stable burst solutions of the full network model, and are created through fold bifurcations of maps. We derive conditions that describe period-increment bifurcations between stable n-n and (n+1)-(n+1) bursts, producing a full bifurcation diagram of the burst cycle period. Predictions of the Poincaré map fit excellently with numerical simulations of the full network model and allow the study of parameter sensitivity for rhythm generation.

19 Keywords: Synaptic depression, Poincaré map, Dynamical system, Neuronal bursting, Central pattern generator

# 1 INTRODUCTION

# 2 MATERIALS AND METHODS

We consider a pair of identical Morris-Lecar neurons [1], with parameters adapted from [2]. The Morris-Lecar model is a set of two first-order differential equations that describe the membrane dynamics of a spiking neuron. The depolarisation is modelled by an instantaneous calcium current, and the hyperpolarisation by a slow potassium current and a leak current. The membrane potential  $v_i$  and potassium activation  $w_i$  of neuron i (i, j = 1, 2) is described by:

$$\dot{v}_i = f(v_i, w_i) - \bar{g}s_i(v_i - v_s),\tag{1}$$

$$\dot{w}_i = h(v_i, w_i). \tag{2}$$

Here  $v_s$  is the inhibitory reversal potential, and  $\bar{g}$  and  $s_j$  are the maximal synaptic conductance and the synaptic gating, respectively, constituting the total inhibitory conductance  $\bar{g}s_j$  from neuron j to neuron i.

22 Function  $f(v_i, w_i)$  describes the membrane currents of a single cell:

$$f(v_i, w_i) = -g_{Ca} m_{\infty}(v_i)(v_i - v_{Ca}) - g_K w_i(v_i - v_K) - g_L(v_i - v_L) + I.$$
(3)

- 23 The currents include a constant current I, and three ionic currents: an instantaneous calcium current, a
- potassium current, and a leak current, with respective reversal potentials  $v_{\rm Ca}$ ,  $v_{\rm K}$ , and  $v_{\rm L}$ , as well as
- 25 maximum conductances  $g_{Ca}$ ,  $g_{K}$ , and  $g_{L}$ . The function  $h(v_i, w_i)$  models the kinetics of the potassium
- 26 gating variable  $w_i$ , and is given by

$$h(v_i, w_i) = \frac{w_{\infty}(v_i) - w_i}{\tau_w} . \tag{4}$$

- 27 The steady-state activation functions  $m_{\infty}$  and  $w_{\infty}$  as well as the default model parameters are described in the **Supplementary Material S1**.
- The dynamics of the synaptic interactions between the neurons are governed by a synaptic gating variable  $s_i$  and a depression variable  $d_i$ :

$$\dot{d}_i = \begin{cases} (1 - d_i)/\tau_a & \text{if } v_i < v_\theta, \\ -d_i/\tau_b & \text{if } v_i > v_\theta, \end{cases}$$

$$(5)$$

$$\dot{s}_i = \begin{cases} -s_i/\tau_\kappa \text{ if } v_i < v_\theta, \\ (d_i - s_i)/\tau_\gamma \text{ if } v_i > v_\theta. \end{cases}$$
 (6)

Variable  $d_i$  describes a firing rate dependent depletion mechanism that governs the amount of depression acting on the synapse. The model is agnostic with respect to the exact mechanism of this depletion, be it pre- or post-synaptic. When the voltage is below firing threshold, depression variable  $d_i$  recovers with time constant  $\tau_a$ , while synaptic variable  $s_i$  decays with time constant  $\tau_k$ . Because synaptic depression occurs on a much slower timescale than synaptic inhibition, we assume  $\tau_d \gg \tau_k$ . When the voltage is above firing threshold, variable  $d_i$  decays to zero with  $\tau_b$ , while  $s_i$  quickly approaches the respective value of  $d_i$ . Since  $\tau_\gamma$  is the shortest time constant of the system with  $\tau_\gamma \ll 1$ , we can assume that whenever  $v > v_\theta$  we can we have  $s_i = d_i$ . The equations for the depression model were adapted from the Bose et al. [3] model. These equations are a mathematically tractable simplification of the established phenomenological

depression model previously described by Tsodyks and Markram [4].

When the cells are uncoupled  $(\bar{g}=0)$ , the membrane dynamics are determined by the cubic v-nullcline  $v_{\infty}(v_i)$  and the sigmoid w-nullcline  $w_{\infty}(v_i)$ , satisfying  $\dot{v}_i=0$  and  $\dot{w}_i=0$ , respectively. The two curves intersect along the middle branch of  $v_{\infty}$ , creating an unstable fixed point  $p_f=(v_f,w_f)$  with a surrounding stable limit cycle of period  $T=T_{act}+T_{inact}$  (fig. 1A). Here  $T_{act}$  is the amount of time the cell spends in the active state when  $v>v_{\theta}$ , while  $T_{inact}$  is the time it spends in the silent state when  $v< v_{\theta}$ . Trajectories along that limit cycle have the familiar shape of the action potential (fig. 1B). The trajectory of an action potential can be dissected into four phases: (1) a silent phase, (2) a jump up, (3) an active phase, and (4) a jump down [see e.g. 5]. During the silent phase the trajectory evolves along the left branch  $(v_i < v_{\theta})$  of the cubic v-nullcline. Once the trajectory reaches the local minimum of  $v_{\infty}$ , it "jumps up" to the right branch  $(v_i > v_{\theta})$ , crossing the firing threshold  $v_{\theta}$ . During the active phase the trajectory then evolves along the right branch of the cubic until it arrives at the local maximum, where it "jumps down" to the left branch commencing a new cycle.

The two-cell network model is numerically integrated using an adaptive step-size integrator for stiff differential equations implemented with XPPAUT [6] and controlled through the Python packages SciPy [7] and PyXPP [8]. The following mathematical analysis is performed on the equations of a single cell. Unless required for clarity, we will therefore omit the subscripts i, j from here on.

# 3 RESULTS

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# 3.1 Anti-phase burst solutions

Short-term synaptic depression of inhibition in a half-centre oscillator acts as a *burst termination* mechanism [9] and is known to produce n-n anti-phase burst solutions of varying period. Such n-n

solutions consist of cells firing bursts of n spikes in alternation. Figure 2A shows the timecourse of a typical 4-4 burst. While one cell is firing a burst it provides an inhibitory conductance to the other cell, preventing it from firing. Therefore, at any given moment one cell is spiking while the other is inhibited. Consistent with Bose and Booth [2] we will refer to the currently firing cell as "free" and we will call the inhibited cell "quiet". Additionally, we will distinguish between two phases of a n-n solution: We will refer to the burst duration of a cell as the "free phase", which is the time between the first spike and the last spike in a burst. And we will call the remaining duration of a cycle, when a cell is not spiking, the "quiet phase".

With each action potential of the free cell, short-term depression leads to a step-wise decrease of d, and consequently of s (fig. 2B). If d depresses faster at spike time than it can recover in the interspike-intervals (ISIs), the total synaptic conductance  $\bar{g}s$  will eventually become sufficiently small to allow for the quiet cell to be released and start firing, thus inhibiting the previously free cell. While a cell is quiet its depression variable can recover. Once the quiet cell becomes free again its synaptic inhibition will be sufficient to terminate the burst of the previously free cell and commence a new cycle. As previously demonstrated by Bose and Booth [2], in a two-cell reciprocally inhibitory network with synaptic depression the coupling strength  $\bar{g}$  determines the type of n-n solution. Increasing  $\bar{g}$  produces higher n-n burst solutions with more spikes per burst and a longer cycle period. Figure 3 shows numerically stable n-n solutions for varying values of  $\bar{g}$ . For small values of  $\bar{g}$  the network produces anti-phase spiking 1-1 solutions. As  $\bar{g}$  is increased the network generates solutions of increasing n, that is 2-2, 3-3, and 4-4. When  $\bar{g}$  is sufficiently large (bottom of fig. 3), one of the cells continuously spikes at its uncoupled period T while the other cell remains fully suppressed. Depending on the initial conditions either of the two cells can become the suppressed cell, which is why the suppressed solution is numerically bistable.

Branches of numerically stable n-n solutions and their associated limit cycle period for varying values of  $\bar{g}$  are depicted in fig. 4A (see **Supplementary Material S2** for algorithm description). Not only do higher n-n solutions branches require stronger coupling  $\bar{g}$ , but also within n-n branches the period increases with  $\bar{g}$ . In line with Bose and Booth [2] we find small overlaps between solution branches indicating numerical bistability, for example such as between the 2-2 and 3-3 solution branches. Branches of higher n-n burst solutions occur on increasingly smaller intervals of  $\bar{g}$ , for instance is the  $\bar{g}$  interval of the 5-5 branch shorter than that of the 4-4 branch and so on. The interval between the 5-5 branch and the suppressed solution (region between dotted lines in fig. 4A) not only contains even higher numerically stable n-n solutions, such as 11-11 bursts, but also other non-symmetric n-m solutions as well irregular, non-periodic solutions. However, the analysis in the following sections will only be concerned with the numerically stable and symmetric n-n solutions.

# 3.2 Mathematical analysis of two-cell network

The goal of the following mathematical analysis is to reduce the complexity of the eight-dimensional system to some easily tractable quantity. As we will see later this quantity is the value of the depression variable d of either of the two cells. We will construct the solution of d in a piecewise manner from one spike to the next, first during the free phase, and then during the quiet phase. This construction will require two assumptions about the membrane and synaptic dynamics. The first assumption states that during a burst the free cell fires at its uncoupled period  $\hat{T}$ , which simplifies the construction of the solution of d. The second assumption states that once the inhibitory conductance acting on the quiet cell drops below a critical threshold, the cell is immediately released and fires. The second assumption is necessary to predict the release time of the quiet cell, which allows us to model the recovery of d during the quiet phase. In other words, the second assumption requires that the release of the quiet cell from inhibition depends only on the timecourse of the inhibition, and not on the membrane dynamics of the quiet cell. Both assumptions can be observed in coupled relaxation-oscillator types of neurons such as the Morris-Lecar model we use, and will be numerically verified below. Both assumptions were first explored in [2] to derive algebraic conditions that guarantee the periodicity of the depression variable for different n-n solutions. However here we will use these assumptions to construct a Poincaré map of d, which will provide a geometric intuition for the dynamics of the full two-cell network and its dependence on model parameters.

Our first assumption about the model states that the free cell fires at its uncoupled period T, that is, during the free phase of a burst we have ISI = T. Solution profiles in fig. 3 suggest that the ISIs are indeed approximately constant. We can further numerically confirm this observation by capturing

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the ISIs of the stable solutions from the bifurcation diagram in fig. 4A. In addition to ISIs, Figure 4B 115 also shows inter-burst intervals (IBIs), which correspond to the time interval between the last spike of the burst and the full cycle period, and which lie in the quiet phase of the burst. IBIs lie on multiple 116 branches, each branch associated with a stable n-n solution, and are monotonically increasing with  $\bar{g}$ . 117 In contrast, ISIs are calculated from the spikes within the free phase and do not vary significantly with  $\bar{q}$ , 118 but are approximately  $ISI \approx T$ , affirming our first assumption. Assuming ISI = T allows us to ignore 119 120 the non-linear membrane dynamics during the free phase, and to construct the evolution of the synaptic variables iteratively from spike to spike. Assuming ISI = T seems reasonable given that inhibition acting 121 on the quiet cell decays exponentially to zero on a much shorter timescale than the duration of the ISI, 122 and therefore, once the quiet cell is released its trajectory quickly approaches the spiking limit cycle. 123

Our second assumption states that the quiet cell is released and spikes as soon as inhibition from the free cell drops below a constant threshold. We will now define such a "release condition" by exploiting the discrepancy in timescales between the fast membrane dynamics, and the slower synaptic dynamics. Let us first consider the dynamics of a single Morris-Lecar neuron. We fix the synaptic variable that acts on the cell by setting s = 1, which also makes the applied synaptic conductance  $\bar{g}s$  constant. Recall from fig. 1A that in case of a single uncoupled cell ( $\bar{q}s=0$ ), the v- and w-nullclines intersect at some unstable fixed point  $p_f = (v_f, w_f)$ , while trajectories revolve around a stable spiking limit cycle. Increasing  $\bar{g}s$  moves the cubic  $v_{\infty}$  with the ensuing unstable fixed point  $p_f$  down the sigmoid  $w_{\infty}$  in the (v-w)-plane (fig. 5). When  $\bar{g}s$  is large enough, the fixed point  $p_f$  becomes stable, attracting all previously periodic trajectories. There exists a unique value  $\bar{g}s = g^*$  when  $p_f$  changes stability and the stable limit cycle vanishes. Thus, when a constant inhibitory conductance is applied,  $\bar{q}s < q^*$  acts as a necessary condition for a cell to spike. In contrast, when  $\bar{g}s > g^*$  inhibition is strong enough to prevent a cell from spiking [2].

Now let us analyse the nullclines of the quiet cell when the two cells are coupled via synaptic inhibition 137 with depression. Let  $\bar{q}s$  here denote the total synaptic conductance which acts on the quiet cell and is produced by the free cell, and let  $p_f$  be the fixed point associated with the quiet cell. At the start of 138 139 the burst of the free cell we have  $\bar{g}s > g^*$  and  $p_f$  is stable. When the free cell spikes,  $\bar{g}s$  peaks (??), and the v-nullcline with the ensuing stable  $p_f$  move down the w-nullcline. Then in between spikes  $\bar{g}s$ 140 decays exponentially (eq. (6)) causing the v-nullcline and  $p_f$  to move up the w-nullcline while attracting trajectories of the quiet cell. Once depression causes the synaptic conductance to become small enough to satisfy  $\bar{g}s < g^*$  and the quiet cell is released, fixed point  $p_f$  becomes unstable allowing the quiet cell to fire. If the trajectory of the quiet cell remains sufficiently close to the stable  $p_f$  at the time when it changes 144 stability, then  $\bar{g}s < g^*$  acts as a release condition that is not only necessary, but also sufficient for firing of the quiet cell. In this case the release of the quiet cell occurs precisely when

$$\bar{g}s = g^* \tag{7}$$

147 is satisfied.

Whether the (v, w)-trajectory of the quiet cell can remain close enough to  $p_f$  to make eq. (7) sufficient for firing depends largely on the coupling strength  $\bar{g}$  and the timescale disparity between membrane 149 dynamics and synaptic dynamics [2]. It is straightforward to test our assumption of a release condition by numerically integrating the full system of ODEs and calculating the time interval between the first spike of the quiet cell and the time when  $\bar{g}s$  first crosses  $g^*$ . We will call this time interval the "release delay". If our assumption holds, we would expect an approximately zero release delay. Figure 6 shows the numerically computed graph of the release delay for varying  $\bar{g}$ . The graph shows three distinct branches, next to each branch we also plot the timecourse of a corresponding sample solution of the total synaptic conductance  $\bar{g}s$  of both cells. For the rightmost branch where  $\bar{g} > 0.592 \text{ mS/cm}^2$  the release delay is approximately zero. Here the first spike of the quiet cell can be accurately predicted by the release condition in eq. (7). The leftmost and middle branches for  $\bar{q} < 0.592 \,\mathrm{mS/cm^2}$  show a release delay greater than zero, the quiet cell does not immediately fire when the release condition is first satisfied, and eq. (7) does not accurately predict the release of the quiet cell. The leftmost and middle branches contain 1-1 and 2-2 solutions respectively. In both cases the coupling  $\bar{g}$  is not sufficiently strong to allow trajectories of the quiet cell to be close enough to  $p_f$  to guarantee spiking once  $\bar{g}$  crosses  $g^*$ . Note that in the middle branch  $\bar{g}s$  crosses  $g^{\star}$  twice, and only after the second crossing does the quiet cell fire. The following map construction relies on the assumption that the release condition in eq. (7) can accurately predict the release time of the

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quiet cell. Given our model parameters this is only possible for sufficiently large  $\bar{g} > 0.592 \text{ mS/cm}^2$ .

166 For completeness, however, we will also consider values  $\bar{g} < 0.592 \text{ mS/cm}^2$  in the following analysis,

bearing in mind that in this parameter range our map will not be accurate.

In summary: For  $\bar{g} > 0.592 \text{ mS/cm}^2$  the release condition is sufficient to predict when the quiet cell 168 is released. Due to the symmetry of n-n solutions the release occurs at exactly half the period of 169 the full cycle, that is at P/2. The release time therefore uniquely determines the type of n-n solution. 170 Furthermore, computation of the release time does not depend on the membrane nor the synaptic dynamics 171 of the quiet cell. Instead, the solution of the synaptic variable s of the free cell is sufficient to predict when 172  $\bar{g}s = g^*$  is satisfied. Finally, s is solely determined by the evolution of the depression variable d of the free 173 cell. Constructing a solution of d during the free phase of either cell will therefore uniquely determine the 174 solution of the full eight-dimensional network. However, finding the solution d requires us to know the 175 176 initial value d(0) at the start of a cycle at t=0. In the next section we will construct a scalar return map that tracks these initial values d(0) from cycle to cycle of stable n-n solutions.

# 178 3.3 Construction of the scalar Poincaré map

In this section we construct the scalar Poincaré map  $\Pi_n: d^\star \mapsto d^\star$ . Here the discrete variable  $d^\star$  tracks the values of the continuous depression variable d at the beginning of each n-n burst. The map  $\Pi_n$  therefore describes the evolution of d, of either of the two cells, from the beginning of one cycle to the beginning of the next cycle. To simplify the map construction we will assume that a free cell fires exactly n times before it becomes quiet. Later we will relax this assumption. We will construct  $\Pi_n$  by evolving d first during the free phase and then during the quiet phase of the n-n limit cycle. First, let us give explicit definitions of the free and quiet phases. A schematic illustration of both phases is given in fig. 7.

Suppose that at t=0 cell 1 becomes free with some initial d(0). Cell 1 then fires n spikes at the uncoupled period  $T=T_{act}+T_{inact}$ . Let s(t) and d(t) be the corresponding solutions of the synaptic and depression variables of cell 1. After n spikes the total conductance  $\bar{g}s(t)$  acting on the quiet cell 2 has decayed sufficiently to satisfy the release condition (7), that is at some time  $t=(n-1)T+\Delta t$ , where  $\Delta t < T_{inact}$ , we have  $\bar{g}s(t)=g^*$ . Cell 2 is then released and prevents cell 1 from further spiking. Here  $\Delta t$  is the time between the last spike of cell 1 and the first spike of cell 2 [2]. Once released, cell 2 also fires n spikes until cell 1 becomes free once again at the cycle period. Let  $P_n$  denote the full cycle period of a n-n solution:

$$P_n = 2(n-1)T + 2\Delta t. (8)$$

We can now define the free and quiet phases of cell 1 explicitly. The free phase is the time interval between the first and last spikes of the burst, that is for time 0 < t < (n-1)T. During the free phase of cell 1, the quiet cell 2 is inhibited sufficiently strong to prevent it from firing, hence  $\bar{g}s > g^*$ . The quiet phase of cell 1 is the remaining duration of the cycle when the cell is not firing, that is for  $(n-1)T < t < 2(n-1)T + 2\Delta t$ .

Note that only the quiet phase depends on  $\Delta t$  which will play a central role in the construction of  $\Pi_n$ . 199 From eq. (8)  $\Delta t$  can be be computed as

$$\Delta t = \frac{1}{2}P_n - (n-1)T. \tag{9}$$

We can use eq. (9) and the numerically computed bifurcation diagram of the period for stable n-n solutions in fig. 4A to obtain the graph of  $\Delta t$  as a function of  $\bar{g}$  (fig. 8). Each continuous branch of  $\Delta t$  is monotonically increasing and corresponds to a n-n burst: Stronger coupling  $\bar{g}$  increases the total synaptic conductance  $\bar{g}s$  that acts on the quiet cell, thus delaying its release. It is easy to see that for any n-branch we have  $\Delta t < T$ : Once  $\Delta t$  crosses T, the free cell can "squeeze in" an additional spike and the solutions bifurcate into a (n+1)-(n+1) burst.

Distinguishing between the active and silent phases of a cycle allows us to describe the dynamics of the depression variable d explicitly for each phase. As can be seen from fig. 7C, during the active phase d depresses during the active phase of spikes and recovers during the inactive phases of spikes. In contrast, during the silent phase d only recovers and does not depress. Given the initial  $d^* = d(0)$  at the beginning of the cycle and the number of spikes in the free phase n, we can now construct the burst map  $\Pi_n$ . The

211 map

$$\Pi_n(d^*) = Q_n(F_n(d^*)) \tag{10}$$

is a composition of two maps. Map

$$F_n: d^* \mapsto \Delta t \tag{11}$$

models the evolution of d in the free phase.  $F_n$  takes an initial value  $d^*$  and calculates the inter-burst-213

interval  $\Delta t$ . Map

$$Q_n: \Delta t \mapsto d^* \tag{12}$$

- models the recovery of d in the quiet phase. Given some  $\Delta t$  map  $Q_n$  computes  $d^*$  at the start of the next 215 216
- Our aim in the following analysis is to elucidate the properties of  $\Pi_n$  and to understand the structure of 217
- its parameter space by exploring how the stable and unstable fixed points of  $\Pi_n$  are created. To that effect 218
- it will be useful to include not only positive, but also negative values of  $d^*$  to the domain of  $\Pi_n$ . But it is
- important to add that values  $d^* < 0$  are biologically impossible as the depression variable models a finite 220
- 221 pool of neurotransmitters, and therefore must be positive. Because  $\Pi_n$  maps first from  $d^*$  to  $\Delta t$ , and then back to  $d^*$ , we will also consider negative values of  $\Delta t$ , interpreting them as n-n solutions with partially 222
- overlapping bursts. As will become evident,  $\Delta t < 0$  is only a formal violation of the biological realism of 223
- the map  $\Pi_n$ , as numerically stable n-n solutions of the full system of ODEs only exist for  $\Delta t > 0$ . 224
- 225 We start the construction of  $\Pi_n$  by first considering the free phase and building the map  $F_n$ . At each
- spike time  $t_k$  where  $d(t_k) = d_k$ , variable d decays first for the duration of the active phase of the spike for  $T_{act}$ , as described by the solution to eq. (5). At  $t = t_k + T_{act}$  we have 226

$$d(t_k + T_{act}) = d_k e^{-T_{act}/\tau_b}. (13)$$

The depression variable then recovers during the inactive phase of the spike until  $t_{k+1}$ , where for  $0 < t < t_{k+1}$ 228

 $T_{inact}$  we get

$$d(t_{k+1}) = 1 - (1 - d_k e^{-T_{act}/\tau_b})e^{-t/\tau_a}.$$
(14)

- 230 By substituting  $t = T_{inact}$  we can build a linear map that models the depression of d from spike time  $t_k$
- to the subsequent spike time  $t_{k+1}$  during the free phase:

$$d_{k+1} = \lambda \rho d_k + (1 - \rho), \tag{15}$$

where to keep the notation simple we let

$$\lambda = \exp(-T_{act}/\tau_b),\tag{16}$$

$$\rho = \exp(-T_{inact}/\tau_a). \tag{17}$$

- Given constant  $T_{act}$  and  $T_{inact}$ , parameter  $\lambda$  determines how much the synapses depresses during the 232
- active phase of the spike, while  $\rho$  determines how much it recovers during the inactive phase. Since 233
- $0 < \lambda, \rho < 1$ , map eq. (15) is increasing and contracting, with a fixed point at

$$d_s = \frac{1 - \rho}{1 - \lambda \rho},\tag{18}$$

- where  $0 < d_s < 1$ . The value  $d_s$  is the maximum depression value that can be observed in the suppressed
- 236 solution where the active cell fires at its uncoupled period T (see fig. 3E). Using the release condition
- in eq. (7) allows us to derive the value of the the minimum coupling strength that will produce the full suppressed solution, denoted as  $\bar{g}_s$ . Solving eq. (6) for s(t) with t=T and setting the initial value 237
- 238
- $s(0) = d_s$  gives

$$\bar{g}_s d_s e^{-T/\tau_\kappa} = g^*. \tag{19}$$

By further substituting the definition of  $d_s$  in (18) and rearranging, we can write  $\bar{g}_s$  as a function of  $\lambda$  and 240 241

$$\bar{g}_s(\lambda, \rho) = g^* e^{T/\tau_\kappa} \frac{1 - \lambda \rho}{1 - \rho}.$$
 (20)

- Note that the above dependence of  $\bar{g}_s$  on  $\lambda$  is linear and monotonically decreasing. Increasing  $\lambda$  reduces
- the strength of the depression of the free cell. This in turn allows the free cell to fully suppress the quiet
- cell at smaller values of  $\bar{q}$ . 244

Solving (15) gives us the linear map  $\delta_n: d^* \mapsto d_n$ , that for some initial  $d^*$  computes the depression at 245 the *n*th spike time,  $d_n = d(t_n^-)$ : 246

$$\delta_n(d^*) = (\lambda \rho)^{n-1} d^* + (1 - \rho) \sum_{i=0}^{n-2} (\lambda \rho)^i.$$
 (21)

- Since  $\lambda < 1$ , function  $\delta_n$  is a linearly increasing function of  $d^\star$  with a fixed point at  $d_s$  for all n. Having identified d after n spikes, we can now use the release condition  $\bar{g}s = g^\star$  (eq. (7)) to find  $\Delta t$ . After the 247
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- last spike of the free phase at time  $t_n = (n-1)T$  the synapse variable s has the value of d for the duration 249
- of  $T_{act}$ , which is given by  $\delta_n(d^*)\lambda$ . s then decays exponentially for  $\Delta t < T_{inact}$ . Solving eq. (6) (case  $v < v_{\theta}$ ) with initial condition  $s(0) = \delta_n(d^*)\lambda$  yields: 250

$$s(\Delta t) = \delta_n(d^*)e^{-\Delta t/\tau_\kappa}.$$
 (22)

Substituting  $s(\Delta t)$  into s of the release condition (eq. (7)) gives then 252

$$\bar{q}\delta_n(d^*)e^{-\Delta t/\tau_\kappa} = q^*. \tag{23}$$

- Our assumption of the release condition guarantees that the quiet cell 2 spikes and becomes free when 253
- $\bar{q}s g^*$  crosses zero. Solving eq. (23) for  $\Delta t$  allows us to compute the inter-spike-interval as a function 254
- of  $d^*$ , which defines our map  $F_n$ : 255

$$F_n(d^*) := \tau_\kappa \ln \left( \frac{\bar{g}}{g^*} \delta_n(d^*) \right) = \Delta t.$$
 (24)

- Figure 9A shows  $F_n$  for various n, which is a strict monotonically increasing function of  $d^*$  as well as  $\bar{g}$ . Larger values of  $d^*$  and  $\bar{g}$ , respectively, cause stronger inhibition of the quiet cell, and therefore prolong its release time and the associated  $\Delta t$ . Map  $F_n$  is defined on  $d^* > d_a$ , where  $d_a$  is a vertical asymptote 256
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- found by solving  $\delta_n(d^*) = 0$  in eq. (21) for  $d^*$ , which yields

$$d_a(n) = -\frac{(1-\rho)\sum_{i=0}^{n-2}(\lambda\rho)^i}{(\lambda\rho)^{n-1}} \le 0.$$
 (25)

- We now turn to the construction of map  $Q_n$ , which describes the recovery of the depression variable 260
- during the quiet phase. As we have identified earlier, the recovery in the quiet phase of a n-n solution 261
- is of duration  $2\Delta t + (n-1)T$ . Substituting that into the solution for d(t) (??) with the initial condition 262
- $d(0) = \delta_n(d^*)$  yields the map  $Q_n$ : 263

$$Q_n(\Delta t) := 1 - (1 - \lambda \delta_n(d^*)) e^{-(2\Delta t + (n-1)T)/\tau_d}.$$
(26)

Given  $\Delta t$ , we can find  $\delta_n(d^*)$  by rearranging the release condition in eq. (23):

$$\delta_n(d^*) = \frac{g^*}{\bar{g}} e^{\Delta t/\tau_s}.$$
 (27)

- Map  $Q_n$  is shown in fig. 9B for various values n. Note that  $Q_n$  is monotonically increasing as larger values  $\Delta t$  imply a longer recovery time, and hence  $Q_n$  grows without bound. All curves  $Q_n$  intersect at some
- 266
- $\Delta t = \tau_s \ln \left[ \bar{g}/(g^* \lambda) \right]$  where

$$Q_n \left[ \tau_s \ln \left( \frac{\bar{g}}{g^* \lambda} \right) \right] = 1. \tag{28}$$

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- 269
- As we will show in the next section, all fixed points of the full map  $\Pi_n$  occur for  $d^\star < 1$ . We will therefore restrict the domain of  $Q_n$  to  $(-\infty, \ln{[\bar{g}/(g^\star\lambda)]}\tau_s)$  and the codomain to  $(-\infty, 1)$ . Additionally, while values  $\Delta t > T$  will be helpful in exploring the geometry of  $\Pi_n$ , recall from fig. 8 that in the flow

- system all n-n solutions bifurcate into (n+1)-(n+1) solutions exactly when  $\Delta t=T$ , and we will address this concern in the last part of our map analysis. 272
- Having found  $F_n$  and  $Q_n$ , we can now construct the full map  $\Pi_n(d^*) = Q_n(F_n(d^*))$ : 273

$$\Pi_n(d^*) = 1 - \frac{\rho^{n-1} g^{\star^{\tau}}}{\bar{q}^{\tau}} \delta_n^{-\tau}(d^*) \left(1 - \lambda \delta_n(d^*)\right), \tag{29}$$

- where we substituted  $\tau=2\tau_s/\tau_d$ . Since d is the slowest variable of the system and  $\tau_d\gg\tau_s$ , we will also assume  $\tau<1$ . Figure 10A depicts  $\Pi_n$  for various n. Intersections of  $\Pi_n$  with the diagonal are fixed points of the map. Figure 10B shows  $\Pi_2$  with n=2. Varying the synaptic strength  $\bar{g}$  moves the curves 274
- 275
- 276
- $\Pi_n$  up and down the  $(d^*, \Pi_n)$ -plane. For  $\bar{g} < 0.03 \text{ mS/cm}^2$  map  $\Pi_2$  has no fixed points. As  $\bar{g}$  is increased 277
- 278
- to  $\bar{g} \approx 0.03~\mathrm{mS/cm^2}$ , curve  $\Pi_2$  coalesces with the diagonal tangentially. When  $\bar{g} > 0.03~\mathrm{mS/cm^2}$ , a pair of fixed points emerge, one stable and one unstable fixed point, indicating the occurrence of a fold 279
- bifurcation of maps. 280

From eq. (29) it is evident that  $\Pi_n$  is monotonically increasing with respect to  $\bar{g}$  and also  $d^*$ :

$$\frac{\mathrm{d}\Pi_n}{\mathrm{d}\bar{q}} > 0,\tag{30}$$

$$\frac{\mathrm{d}\Pi_n}{\mathrm{d}d^*} > 0,\tag{31}$$

and in the following sections we will heavily rely on this monotonicity property of  $\Pi_n$ . Just as  $F_n$ , curves 281  $\Pi_n$  spawn at the asymptote  $d_a$  (eq. (25)), and because

$$\lim_{\bar{q} \to \infty} \Pi_n = 1 \text{ for all } n, \tag{32}$$

fixed points of  $\Pi_n$  lie in  $(d_a, 1)$ . 283

## Existence and stability of fixed points 284

We introduce the fixed point notation  $d_f^{\star}$  with  $\Pi_n(d_f^{\star}) = d_f^{\star}$ . The existence of fixed points  $d_f^{\star}$  for  $\bar{g}$ 285 sufficiently large can be shown from the strict monotonicity of  $\Pi_n$  with respect to  $\bar{q}$  and  $d^*$  (eqs. (30) 286 and (31)), as well as the fact that the slope of  $\Pi_n$  is monotonically decreasing, 287

$$\left(\frac{\mathrm{d}}{\mathrm{d}d^{\star}}\right)^{2}\Pi_{n}<0. \tag{33}$$

In the limit  $d^* \to d_a$  the value of  $\Pi_n$  decreases without bound for any  $\bar{g} > 0$ . In the limit  $\bar{g} \to 0$ ,  $\Pi_n$  also decreases without bound, but as  $\bar{g} \to \infty$  values of  $\Pi_n$  approach 1. It follows from eq. (30) and the intermediate value theorem that for some  $\bar{g}$  large enough  $\Pi_n$  intersects the diagonal. Moreover, because  $\Pi_n$  and its slope are monotonic with respect to  $d^{\star}$ , there exists some critical fixed point  $(d_b^{\star}, \bar{g}_b)$  where  $\Pi_n$ aligns with the diagonal tangentially with

$$\Pi_n(d_b^{\star}; \bar{g}_b) = d_b^{\star},\tag{34}$$

$$\frac{\mathrm{d}}{\mathrm{d}d^{\star}}\Pi_{n}(d_{b}^{\star};\bar{g}_{b}) = 1. \tag{35}$$

- Equations (30) and (33) constitute the non-degeneracy conditions for a codimension-1 fold bifurcation of maps, indicating that in a neighbourhood of  $(d_b^{\star}, \bar{q}_b)$  map  $\Pi_n$  has the topological normal form described 289
- by the graph of 290

$$x \mapsto \beta + x - x^2,\tag{36}$$

with a stable and unstable fixed point  $x = \pm \sqrt{\beta}$ , and slopes  $dx/d\beta = \mp (2\sqrt{\beta})^{-1}$ , respectively.

# 292 3.5 Fold bifurcations

293 Fixed points of  $\Pi_n$  satisfy the fixed point equation

$$\Phi_n(d^*; \bar{g}) = 0, \tag{37}$$

294 where

$$\Phi_n(d^*, \bar{q}) := \Pi_n(d^*, \bar{q}) - d^*. \tag{38}$$

As we have already shown, for  $\bar{g} > \bar{g}_b(n)$  solutions to eq. (37) exist in pairs of stable and unstable fixed points. Solving eq. (37) explicitly for  $d^*$  it not trivial, but solving for  $\bar{g}$  is straightforward and given by  $\bar{q} = G_n(d^*)$ , where

$$G_n(d^*) := g^* \left( \frac{\rho^{n-1} \delta_n^{-\tau} (d^*) (1 - \lambda \delta_n(d^*))}{1 - d^*} \right)^{1/\tau}.$$
 (39)

Plotting  $d^*$  against  $\bar{g}$  gives the fixed point curves, which are shown in fig. 11A. Note the typical quadratic shape of a fold bifurcation of maps. It is also evident that the fold bifurcations occur for increasingly smaller  $\bar{g}$  as n is increased. Moreover, we can observe that unstable fixed points have negative values of  $d^*$  for n > 1.

Equation (39) also allows us to find the critical fixed point connected with the fold bifurcation, namely  $(d_b^*(n), \bar{g}_b(n))$ , which is the global minimum of  $G_n(d_f^*)$ :

$$d_b^{\star}(n) = \operatorname{argmin} G_n(d_f^{\star}), \tag{40}$$

$$\bar{g}_b(n) = \min G_n(d_f^*). \tag{41}$$

Function  $G_n$  is strictly monotonic on the respective intervals of  $d_f^{\star}$  that correspond to the stable and unstable fixed points, that is

$$\frac{\mathrm{d}G_n}{\mathrm{d}d_f^{\star}} < 0, \text{ for } d_f^{\star} > d_b^{\star}(n) \text{ stable}, \tag{42}$$

$$\frac{\mathrm{d}G_n}{\mathrm{d}d_f^\star} > 0, \text{ for } d_f^\star < d_b^\star(n) \text{ unstable}, \tag{43}$$

which allows us to express the stable and unstable fixed points as the inverse of  $G_n$  on their respective intervals of  $d_f^*$ . Because we are primarily interested in the stable fixed points, we define the stable fixed

304 point function  $d_f^{\star} = \phi_n(\bar{g})$  as

$$\phi_n(\bar{g}) := G_n^{-1}(d_f^*) \text{ for } d_f^* > d_b^*(n).$$
 (44)

Function  $\phi_n(\bar{g})$  is also monotonic, and is therefore straightforward to compute numerically via rootfinding. Here we use the Python package Pynverse [10] for that purpose.

Having found the stable fixed points  $d_f^*$  as a function of the coupling strength  $\bar{g}$ , we can now compute the associated cycle period. Recall that the period is given by eq. (8), which we can be written as a function of  $\bar{g}$ :

$$P_n(\bar{g}) = 2(n-1)T + 2F_n(\underbrace{\phi_n(\bar{g})}_{d_f^*}, \bar{g}), \tag{45}$$

- 310 where map  $F_n$  (eq. (24)) calculates the inter-burst-interval  $\Delta t$  given a stable fixed point  $d_f^{\star} = \phi_n(\bar{g})$ . We
- 311 plot the predicted period  $P_n(\bar{g})$  versus the cycle period that was computed from numerically integrating
- the full system of ODEs in fig. 11B. For n > 1 our map  $\Pi_n$  accurately predicts the period. When laying
- out our assumptions in section 3.2, we have already predicted an inaccuracy for n=1 (see fig. 6), since
- here  $\bar{g}$  is not sufficiently strong to guarantee the validity of our release condition (eq. (7)).

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It is evident from fig. 11A that  $\phi_n$  is strictly increasing with  $\bar{g}$ . This property follows directly from the 315 normal form of the fold bifurcation (eq. (36)), but can also be shown using implicit differentiation and the 316 fixed point equation  $\Phi_n(\phi_n(\bar{g}), \bar{g}) = 0$  in eq. (37). For  $d_f^{\star} = \phi_n(\bar{g}) > d_b(\bar{n})$  we get: 317

$$\frac{\mathrm{d}\phi_n}{\mathrm{d}\bar{g}} = -\frac{\partial \Phi_n/\partial \bar{g}}{\partial \Phi_n/\partial d^*} = \frac{\partial \Pi_n/\partial \bar{g}}{1 - \partial \Pi_n/\partial d^*} > 0. \tag{46}$$

The inequality follows from  $\partial \Pi_n/\partial \bar{g}>0$  and the fact that  $\partial \Pi_n/\partial d^\star<1$  for  $d^\star>d_b(n)$ . Equation (46) allows us to explain why the period  $P_n$  increases with  $\bar{g}$ , as seen in fig. 11B. Differentiating  $P_n$  gives:

$$\frac{\mathrm{d}P_n}{\mathrm{d}\bar{g}} = 2\nabla F_n(d_f^{\star}, \bar{g}) \cdot \begin{bmatrix} \partial \phi_n / \partial \bar{g} \\ 1 \end{bmatrix} > 0, \tag{47}$$

where the partial derivatives of  $F_n(d_f^{\star}, \bar{g})$  are:

$$\frac{\partial F_n}{\partial d_f^{\star}} = \tau_s \frac{(\lambda \rho)^{n-1}}{\delta_n(d_f^{\star})} > 0, \tag{48}$$

$$\frac{\partial F_n}{\partial \bar{g}} = \frac{\tau_s}{\bar{g}} > 0. \tag{49}$$

Equations (46) and (47) have an intuitive biological interpretation: Increasing the coupling strength between the neurons leads to overall stronger inhibition of the quiet cell, which delays its release and leads to a longer cycle period. The latter allows more time for the synapse to depress in the free phase and 322 recover in the quiet phase, resulting in overall larger values of  $d_f^{\star}$ , that is weaker depression at the burst 323 onset. 324

While fixed points of our Poincaré map predict the cycle period of the flow system excellently, its construction relies on the strong assumption that the free phase contains exactly n spikes. As is evident from fig. 11B this assumption is clearly violated in the flow system, as stable n-n bursts exists only on certain parameter intervals of  $\bar{g}$ . In the last sub-section we will analyse the mechanisms that guide how the stable n-n are created and destroyed, and use our previous analysis to derive the corresponding parameter intervals of  $\bar{q}$  where such solutions exist.

### 3.6 Period increment bifurcations with co-existent attractors 331

Let  $\bar{g}_{+}(n)$  and  $\bar{g}_{-}(n)$  denote the left and right parameter borders on  $\bar{g}$  where stable n-n solutions exist. That is, as  $\bar{g}$  is increased stable n-n solutions are created at  $\bar{g}_+(n)$  and destroyed at  $\bar{g}_-(n)$ . When  $\bar{g}$ is reduced beyond  $\bar{g}_+(n)$ , n-n solutions bifurcate into (n-1)-(n-1) solutions, while when  $\bar{g}$  is increased beyond  $\bar{g}_-(n)$ , n-n solutions bifurcate into (n+1)-(n+1) solutions. Let us briefly recap our observations regarding  $\bar{q}_{+}(n)$  and  $\bar{q}_{-}(n)$  from the numerical bifurcation diagram in fig. 11B. For n>1there are the following relations:

$$\bar{q}_{+}(n) < \bar{q}_{-}(n), \tag{50}$$

$$\bar{g}_{+}(n) < \bar{g}_{+}(n+1) \text{ and } \bar{g}_{-}(n) < \bar{g}_{-}(n+1),$$
 (51)

$$\bar{g}_{+}(n+1) < \bar{g}_{-}(n),$$
 (52)

$$\bar{g}_{-}(n+1) - \bar{g}_{+}(n+1) < \bar{g}_{-}(n) - \bar{g}_{-}(n).$$
 (53)

Equations (50) and (51) are self-explanatory. Equation (52) formally describes occurrence of co-existence 332 between stable n-n and (n+1)-(n+1) solutions. Equation (53) implies that the parameter interval on  $\bar{g}$  of n-n solutions decreases with n, in other words, bursts with more spikes occur on increasingly smaller intervals of the coupling strength. All of the above relations are reminiscent of the period increment bifurcations with co-existent attractors, first described for piecewise-linear scalar maps with a single discontinuity by Avrutin and colleagues [e.g. see 11, 12, 13, 11]. While our maps  $\Pi_n$  are fully continuous, the above observation suggests that a different piecewise-linear scalar map that captures the

period increment bifurcations of the full system might exist. We will explore what such a map might look like in the discussion.

Let us now find algebraic equations that will allow us to calculate the critical parameters  $\bar{g}_+(n)$  and  $\bar{g}_-(n)$  associated with the period increment bifurcations. Recall that the period  $P_n$  derived from the fixed points of  $\Pi_n$  is an increasing function of  $\bar{g}$ :

$$\frac{\mathrm{d}P_n}{\mathrm{d}\bar{g}} = 2\frac{\mathrm{d}F_n(\phi_n(\bar{g}), \bar{g})}{\mathrm{d}\bar{g}} > 0,\tag{54}$$

the value of the release conductance, which delays the release of the quiet cell, and  $\Delta t$  becomes larger. Once  $\Delta t = T$ , the free cell can produce another spike and the solution bifurcates into a (n+1)-(n+1) solution. Note, however, that at  $\bar{g}_+(n)$  the bifurcation into a (n-1)-(n-1) does not occur when  $\Delta t = 0$ . Here the mechanism is different: A sufficient reduction of  $\bar{g}$  causes the total synaptic conductance to drop below the release conductance in the *previous* ISI, which allows the quiet cell to be released one spike earlier.

that is, as the coupling strength increases, it takes longer for the total synaptic conductance to fall below

Using the above reasoning we can now formulate the conditions for both bifurcations at  $\bar{g}_+(n)$  and  $\bar{g}_-(n)$ . As in the previous sections, we will only restrict ourselves to the analysis of the stable fixed points given implicitly by  $d_f^\star = \phi_n(\bar{g})$  (eq. (44)). At the right bifurcation border  $\bar{g}_-(n)$  we have  $\Delta t = T$ , and after substituting our  $F_n$ -map (eq. (24)) this translates into

$$F_n(\phi_n(\bar{g}), \bar{g}) = T, \tag{55}$$

355 which lets us define a function

$$R_n(\bar{q}) := F_n(\phi_n(\bar{q}), \bar{q}) - T, \tag{56}$$

356 whose root is the desired right bifurcation border  $\bar{g}_{-}(n)$ . In case of the left bifurcation border at  $\bar{g}_{+}(n)$  the release condition is satisfied just before the free cell has produced its nth spike, and after the depression variable has been reset n-1 times, which gives the condition

$$\bar{g}\delta_{n-1}(\phi_n(\bar{g}))e^{-T/\tau_s} = g^*, \tag{57}$$

359 and can be rewritten as a function

$$L_n(\bar{g}) := \bar{g}\delta_{n-1}(\phi_n(\bar{g}))e^{-T/\tau_s} - g^*, \tag{58}$$

360 whose root is  $\bar{g}_+(n)$ . Both  $R_n$  and  $L_n$  are increasing with respect to  $\bar{g}$ , which makes finding their roots numerically straightforward.

Figure 12 shows the period  $P_n(\bar{g})$  as predicted by the fixed points of  $\Pi_n$  (eq. (45)) plotted on their respective intervals  $\bar{g} \in [\bar{g}_+(n), \bar{g}_-(n)]$  (blue), as well as the cycle period acquired from numerical integration of the full system of ODEs (orange). Note that the width of n-n branches decreases with n, which confirms the inequality in eq. (53). That is, bursts with more spikes occur on increasingly smaller intervals of  $\bar{g}$ , which can be interpreted as a lost of robustness with respect to the coupling strength of long-cyclic solutions. We also note the occurrence of bi-stability between pairs of n-n and (n+1)-(n+1) branches, which also confirms our initial observation in eq. (52).

As previously observed in fig. 11B our maps prediction of the cycle period is accurate for n > 1. Recall 369 that our reduction assumptions required a sufficiently large coupling strength, which we numerically 370 estimated to be  $\bar{g} \approx 0.592 \text{mS/cm}^2$  in fig. 6. The mismatch in period for 1-1 solutions, but also the 371 mismatch in the left bifurcation border  $\bar{g}_{-}(n=2)$  of the 2-2 solution can be attributed to the violation 372 of that assumption. However, even for branches of large n-n solutions there is a mismatch between the 373 bifurcation borders. Presumably our assumptions on the time scales of w and s dynamics do not hold here, 374 375 and can only be captured by more complex approximations. Nevertheless, our map allows approximate extrapolation of the cycle period and the respective bifurcation borders where numerical integration of the 376 ODEs would require a very small time step. 377

# 4 DISCUSSION

Synaptic depression of inhibition is believed to play an important role in the generation of rhythmic activity involved in many motor rhythms such as in leech swimming [14] and leech heart beat [15], and in the lobster pyloric system [16, 17]. In inhibitory half-centre CPGs, such as believed to be found in the struggling network of *Xenopus* tapdoles, synaptic depression can act as a burst termination mechanism, enabling the alternation of bursting between the two sides of the CPG [18]. Mathematical modelling can shed light on the underlying mechanisms that enable the generation of such anti-phase bursts, and help identify the components that control this rhythm allowing it to switch between different patterns.

To study the mechanisms of burst generation in half-centre CPGs we have analysed a neuronal model network that consists of a pair of inhibitory neurons that undergo a frequency dependent synaptic depression. When the strength of synaptic inhibition between the neurons is varied, such a simple network can display a range of different n-n burst patterns. Using the timescale disparity between neuronal and synaptic dynamics, we have reduced the network model of eight ODEs to a scalar first return map  $\Pi_n$  of the slow depression variable d. This map  $\Pi_n$  is a composition of two maps,  $F_n$  and  $Q_n$ , that model the evolution of the depression during the free and quiet phases of n-n solutions respectively. Both  $F_n$  and  $Q_n$  maps are constructed by using the dynamics of single uncoupled neurons. Fixed points of  $\Pi_n$  are created in pairs through a fold bifurcation of maps, where the stable fixed point correspond to stable n-n burst solutions of the full two-cell system of ODEs. The results from our one-dimensional map match excellently with numerical simulation of the full network. Our results are in line with Brown's 1911 rhythmogenesis hypothesis, namely that synaptic depression of inhibition is a mechanism by which anti-phase bursting may arise.

We have studied n-n solutions assuming that the synaptic coupling  $\bar{g}$  between the two cells is symmetrical. However, Bose and Booth [2] have shown that asymmetrical coupling  $(\bar{q}_1, \bar{q}_2)$  can result in network solutions of type m-n, where one cell fires m spikes, while the other n spikes. It is conceivable that our map construction can be extended to also capture such m-n solutions. Remember, in the case of symmetrical coupling with n-n solutions, the timecourse of the depression variables  $d_1$  and  $d_2$  were in anti-phase, and it was therefore sufficient to track only one of the two variables. To capture the full network dynamics in case of asymmetrical coupling one would also have to account for burst patterns of type m-n, where the solutions of the depression variables  $d_1$  and  $d_2$  are not simply time-shifted versions of each other. To do that, one could track the state of both variables by constructing a two-dimensional Poincaré map  $\Pi(d_1, d_2)$ . While geometrical interpretation of two-dimensional maps remains challenging, there exist a number of recent studies which have employed novel geometrical analysis methods to understand the dynamics of two-dimensional maps of small neuronal networks [19, 20, 21]. Generally speaking, our map construction approach is applicable to any small network, even with more than two neurons. As long as the network dynamics occur on separable timescales the main challenges to the map construction lie in identifying the slowest variables, and finding an appropriate, simplified description of their respective timecourses. In theory, the reduction approach can be also applied to neuronal systems with more than two timescales [e.g. see 22].

In tadpoles, struggling is believed to be initiated by an increase in the firing frequency of reciprocally inhibitory commisural interneurons, which has been hypothesised to lead to stronger synaptic depression of inhibition and result in the iconic anti-phase bursting [18]. It would therefore be interesting to study how varying the cell intrinsic firing period T could affect the network rhythm. While we have laid out the framework to perform such an investigation, due to the choice of neural model we have avoided varying T. Recall that T is a derived parameter in the Morris and Lecar [1] model, and can therefore not be varied in isolation of other model parameters. This makes verifying any analytical results from our map analysis via numerical integration of the ODEs difficult. A more abstract model such as the quadratic integrate-and-fire model [23] allows varying T independently of other model parameters, and could be more fitting in this scenario.

Our simulations of the network showed that n-n solutions lose robustness as their period is increased. That is, solutions with a larger cycle period occur on increasingly smaller intervals of the coupling strength. We were able to replicate this finding by numerically finding the respective left and right borders of stable n-n branches of fixed points of  $\Pi_n$ , and showing that the distance between these borders shrinks with n. We have also noted the resemblance of our bifurcation diagram to one where such n-n branches are created via a period-increment bifurcation with co-existent attractors of scalar linear maps

- with a discontinuity [24, 13]. It is worthwhile noting that the bifurcations of piecewise linear maps studied 431
- 432 by Avrutin et al. and colleagues result from a "reinjection" mechanism, first described by Perez [25]. Here
- the orbit of a map performs multiple iterations on one side of the discontinuity, before jumping to the other 433
- side and being *reinjected* back into the initial side of the discontinuity. The stark difference of such a map 434
- to our map is that reinjection allows a *single* scalar map to produce periodic solutions of varying periods. 435 In contrast, we rely on n different maps  $\Pi_n$  to describe the burst dynamics without explicitly capturing the 436
- period increment bifurcations. It is therefore conceivable that despite the complexity and non-linearity of 437
- the dynamics of our two-cell network, a single piecewise-linear map might be already sufficient to capture 438
- the mechanisms that shape the parameter space of the full system. In their discussion, Bose and Booth [2] 439
- briefly outline ideas about how such a linear map could be constructed. 440
- 441 In addition to stable n-n solutions, the numerical continuation by Bose and Booth [2] also revealed
- 442 branches of unstable n-n solutions. While we have identified fold bifurcations of our burst map, we have
- not found corresponding bifurcations of the flow ODE system, and have generally ignored the significance 443
- 444 of unstable map fixed points. However, the quadratic nature of the period bifurcation curve is reminiscent
- of a saddle-node on an invariance circle (SNIC) bifurcation, where the oscillation period lengthens and 445
- finally becomes infinite as a limit cycle coalesces with a saddle point. SNIC bifurcations have been studied 446
- 447 in great detail [e.g. 26], and a next step would be to provide a rigorous explanation of not only the map
- 448 dynamics, but also of the flow dynamics of the ODE system.
- 449 We have shown that when the strength of the maximum synaptic conductance is varied, synaptic
- depression of inhibition can enable our two-cell network to produce burst solutions of different periods. 450
- This result is in line with the idea that one role of synaptic depression in the nervous system may be 451
- 452 to allow a finite size neuronal network to participate in different tasks by producing a large number
- of rhythms [2, 27, 18]. To change from one rhythm to another would only require a reconfiguration 453
- of the network through changes in synaptic coupling strength, which can occur through the process of 454
- 455 learning. Thus short-term synaptic depression of inhibition may provide means for a network to adapt to
- environmental challenges without changing its topology, that is without the introduction or removal of 456
- 457 neurons.

# CONFLICT OF INTEREST STATEMENT

- 458 The authors declare that the research was conducted in the absence of any commercial or financial
- 459 relationships that could be construed as a potential conflict of interest.

# **AUTHOR CONTRIBUTIONS**

- MO and CH contributed to conception and design of the study. MO performed numerical computation 460
- 461 and analysis. All authors contributed to manuscript writing and revision.

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# DATA AVAILABILITY STATEMENT

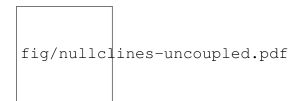
The datasets generated for this study are available on request to the corresponding author.

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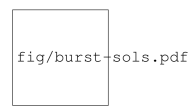
# FIGURE CAPTIONS



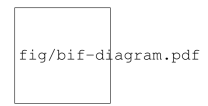
**Figure 1.** Periodic solution of relaxation-oscillator model neuron. (A) Projection of limit cycle onto (v, w)-phase plane with v-nullcline (blue,  $v_{\infty}$ ) and w-nullcline (orange,  $w_{\infty}$ ). Unstable fixed point  $p_f$  is indicated by an orange dot. (B) Corresponding voltage trace v(t) of an action potential.



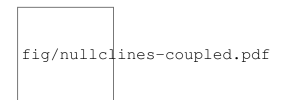
**Figure 2.** Solution profiles of a 4-4 burst. (A) Membrane potentials of cell 1  $(v_1, \text{ blue})$ , and cell 2  $(v_2, \text{ orange})$ . (B) Synaptic variables  $d_1$  (blue) and  $s_1$  (grey) of cell 1.



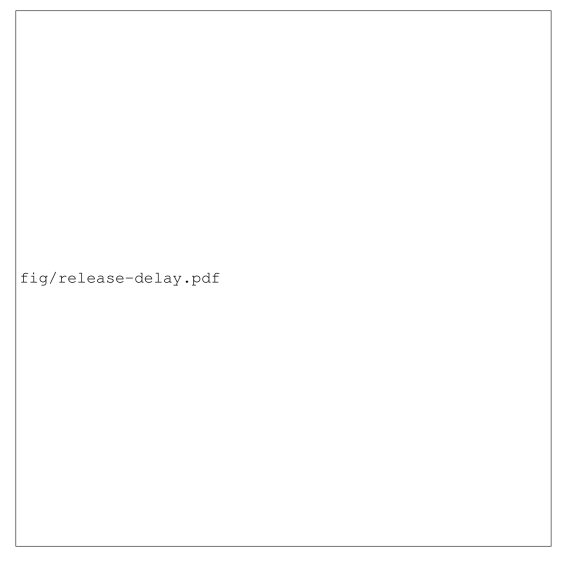
**Figure 3.** Voltage traces of both cells of numerically stable solutions for increasing values of the coupling strength  $\bar{q}$  (increasing top to bottom).



**Figure 4.** Numerically computed bifurcation diagrams of stable n-n solutions for increasing coupling strength  $\bar{g}$ . (A) Period of stable solutions. Dashed lines show the interval between the 5-5 and the suppressed solutions, where higher period n-n solutions occur on increasingly smaller intervals of  $\bar{g}$ . (B) ISIs corresponding to n-n solutions in (A). Long ISIs are associated with the quiet phase of a burst, short ISIs with the free phase. During the free phase, ISIs are of approximately constant duration T.



**Figure 5.** Nullclines  $v_{\infty}$  (blue) and  $w_{\infty}$  (orange) in the (v,w)-phase plane for different values of the total synaptic conductance  $\bar{g}s$ . For small  $\bar{g}s < g^{\star}$ , fixed point  $p_f$  is unstable (orange point). Larger values  $\bar{g}s$  move  $v_{\infty}$  down in the (v,w)-plane until  $p_f$  changes stability (half orange, half blue point) at some critical total conductance value  $\bar{g}s = g^{\star}$ , and becomes stable (blue point) for  $\bar{g} > g^{\star}$ .



**Figure 6.** Numerically computed values of the release delay for varying  $\bar{g}$ . Each of the three branches (A, B, C) also shows the timecourse of the total synaptic conductance  $\bar{g}s$  of a sample stable solution of both cells (blue and orange), as well as the release conductance  $g^*$  (dashed green line). (A) Branch corresponding to the 1-1 solution. Here the quiet cell only spikes after a significant release delay. (B) Branch with a long release delay associated with a subset of 2-2 solutions. Here the release condition is briefly satisfied after the first spike of cell 1. This does not cause firing of cell 2, which only occurs after the second spike of cell 1. (C) Branch with n-n solutions where release delay is approximately zero and the release condition is sufficient for firing of cell 2.



**Figure 7.** Schematic diagram of the free and quiet phases for a 3-3 solution. (A) Membrane potentials of cell 1 ( $v_1$ ) and cell 2 ( $v_2$ ). The grey patches depict inter-burst-intervals  $\Delta t$ . (B) Total synaptic conductance of cell 1 ( $\bar{g}s_1$ ) as it crosses the release conductance  $g^*$ . (C) Solution  $d_1(t)$  of depression variable of cell 1, during free (blue) and quiet phases (orange).



**Figure 8.** Numerically computed bifurcation diagram of  $\Delta t$  for varying  $\bar{g}$ . Each continuous branch is associated with a stable n-n burst solution. Increasing  $\bar{g}$  increases  $\Delta t$  until the solutions bifurcate at  $\Delta t \approx T$ .



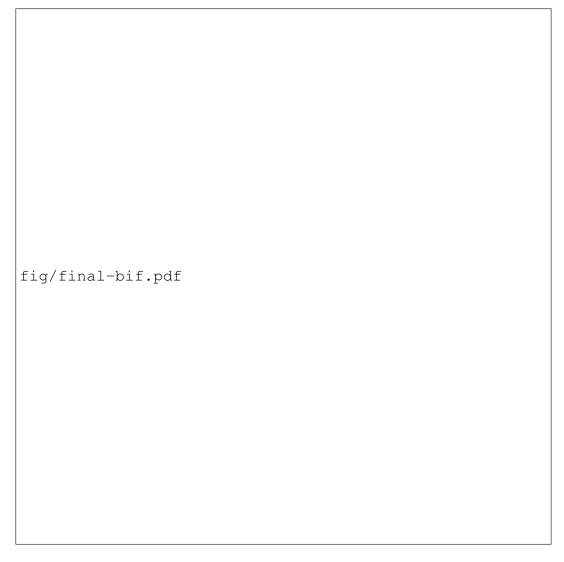
**Figure 9.** Maps  $F_n$  (A) and  $Q_n$  (B) for  $\bar{g} = 0.6 \text{ mS/cm}^2$  and n = 1, 2, 3, 4. Curves  $F_n$  intersect at  $d_s$  which is indicated by a dashed vertical line.



Figure 10. Map  $\Pi_n: d^*$ . (A)  $\Pi_n$  for n=1,2,3,4 at  $\bar{g}=0.6 \text{ mS/cm}^2$ . (B)  $\Pi_2$  with n=2 for varying  $\bar{g}\approx 0.01,0.034,0.3 \text{ mS/cm}^2$ . The identity function is illustrated by a diagonal line.



**Figure 11.** (A) Fold bifurcation diagrams of stable (continuous curves) and unstable (dotted curves) fixed points of  $\Pi_n$  for varying n. (B) Cycle periods computed from stable fixed points (blue), and the corresponding solution period from numerical integration of the system of ODEs (orange).



**Figure 12.** Bifurcation diagrams of stable n-n solutions computed analytically from fixed points of  $\Pi_n$  and plotted on the respective intervals of  $\bar{g} \in \left[\bar{g}_+(n), \bar{g}_-(n)\right]$  (blue), and computed from numerical integrations of the ODEs (orange).