Role of inhibitory feedback for information processing in thalamocortical circuits

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The information transfer in the thalamus is blocked dynamically during sleep, in conjunction with the occurrence of spindle waves. In order to describe the dynamic mechanisms which control the sensory transfer of information, it is necessary to have a qualitative model for the response properties of thalamic neurons. As the theoretical understanding of the mechanism remains incomplete, we analyze two modeling approaches for a recent experiment by Le Masson *et al.* [Nature (London) **417**, 854 (2002)] on the thalamocortical loop. We use a conductance based model in order to motivate an extension of the Hindmarsh-Rose model, which mimics experimental observations of Le Masson *et al.* Typically, thalamic neurons posses two different firing modes, depending on their membrane potential. At depolarized potentials, the cells fire in a single spike mode and relay synaptic inputs in a one-to-one manner to the cortex. If the cell gets hyperpolarized, *T*-type calcium currents generate burst-mode firing which leads to a decrease in the spike transfer. In thalamocortical circuits, the cell membrane gets hyperpolarized by recurrent inhibitory feedback loops. In the case of reciprocally coupled excitatory and inhibitory neurons, inhibitory feedback leads to metastable self-sustained oscillations, which mask the incoming input, and thereby reduce the information transfer significantly.

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I. INTRODUCTION

Spindle oscillations are waxing and waning waves that originate in the thalamus at the transition to slow wave sleep in mammals [1]. In a pioneering experiment [2], spindle oscillations have been investigated in a hybrid thalamic circuit consisting of a biological thalamocortical (TC) and an artificial reticular (RE) thalamic cell [2]. An artificial cell was necessary to manipulate selectively the synaptic connection of the inhibitory feedback from the RE to the TC cell (Fig. 1). Spindle oscillations depended critically on the presence of the synaptic connections between TC and RE cells [3,4]. Further, the occurrence of spindle oscillations comes along with a significant decorrelation between input and the output of the TC cells. Clarification of the relationship between the occurrence of spindle oscillations and the decrease in information transmission could help to gain more insight into the mechanisms which deprive the sensory information from the consciousness while mammals are sleeping.

Spindle, or bursting, oscillations in excitable media are widespread observed in a variety of physical [5], chemical [6], and biological [7] oscillators. Typically, the potential of a bursting neural cell undergoes subsequent shifting between active and silent phases. In the active phase, the membrane potential oscillates quickly, and in the silent phase, it evolves slowly without or only with subthreshold oscillations [8]. In this work, we focus on bursting or spindle oscillations in neural systems, and investigate their role for information transfer through the thalamus. The thalamus, our major gateway for all sensory information, relays the incoming information depending on the state of arousal [9,10]. To process and relay information [11] is a generally important, but not equally well understood, feature of neural systems; here we contribute to the understanding of the underlying dynamics of a key mechanism in the thalamocortical loop.

The paper is organized as follows. In Sec. II, we use biophysical models of TC and RE neurons interconnected with realistic model synapses as introduced in [3] and also used in the hybrid network investigated by Le Masson in [2]. In Sec. III, we establish an extension of the Hindmarsh-Rose model [12]. For this purpose, we use dynamically coupled Hindmarsh-Rose neurons to reproduce the experimental results of [2]. We show that the Hindmarsh-Rose system has to be extended by a fourth variable working on a very slow time scale, in order to describe the experimental results.

For both models, we investigate the influence of the inhibitory synaptic connection on the information transfer of the TC cell and compare the results with the experiment of Le Masson *et al.* [2]. An additional goal is to answer the question whether the complicated biophysical conductancebased model can be replaced by the much simpler Hindmarsh-Rose model without loss of the most important dynamical features. We show how to achieve this by introducing an extended Hindmarsh-Rose model with an additional degree of freedom. Some important features are better reproduced by the extended Hindmarsh-Rose neuron than by the biophysical model. Our model is by far less computationally costly, yet simple enough to understand dynamical mechanisms.



FIG. 1. Structure of synaptic connections in a pair of reciprocally coupled RE-TC cells.

II. INFORMATION TRANSFER IN A BIOPHYSICAL MODEL OF A THALAMOCORTICAL OSCILLATOR

A common approach to model the thalamocortical system are conductance-based models [13-16]. Descriptions of the properties of single neural cells can be found for example in [3,4]. The case of two interconnected cells without external forcing has been investigated in [4]. To our knowledge the case of two interconnected cells as shown in Fig. 1 which get excited by an unregularly distributed realistic synaptic bombardment as used in the experiment of Le Masson *et al.* [2] has not been studied in detail upto now. In this section we will investigate the influence of the inhibitory synaptic connection on the information transfer of the TC cell for this case. Destexhe *et al.* [4] model the TC cell and the RE thalamic cell by using a conductance-based single compartment model. The time evolution of the membrane potentials is governed by the following cable equations:

$$C_m V_T = -I_{T_L} - I_T - I_h - I_{T_{Na}} - I_{T_K} - I_{GABA_a} - I_{GABA_b}, \quad (1)$$

$$C_m V_R = -I_{R_L} - I_{Ts} - I_{K[Ca]} - I_{CAN} - I_{R_{Na}} - I_{R_K} - I_{GLU}, \quad (2)$$

where V_T is the membrane potential of the TC cell; C_m is the capacity of the membrane. According to [4], we included the following ion currents: The leakage current I_{T_L} and I_{R_L} , the low-threshold Ca²⁺ currents I_T and I_{Ts} , the hyperpolarization activated current I_h , the Ca²⁺-activated currents $I_{K[Ca]}$ and I_{CAN} , and, like in the Traub and Miles [17] model, the fast Na⁺ and K⁺ currents $I_{T/R_{Na}}$ and I_{T/R_K} , which are responsible for the generation of action potentials. The synaptic currents I_{GABA_a} and I_{GABA_b} represent the $GABA_a$ and $GABA_b$ receptors in the synapses from RE to TC cells, while I_{GLU} describes the excitatory synapse from the TC to RE cells.

Destexhe *et al.* [4] refer to several sources for the description of the ion currents. The kinetic equations of the TC cell have been described in detail in [15]. The I_T kinetics was taken from Wang *et al.* [18]. I_h was described by the model of Destexhe *et al.* [15], which incorporates a regulation by intracellular calcium. For the RE cell I_{Ts} was taken from Huguenard and Prince [19]. The kinetics of the Ca²⁺-dependent currents $I_{K[Ca]}$ and I_{CAN} are adjusted to the clamp data of RE neurons [20]. For both cells, I_{Na} and I_K were taken from [17]. The calcium dynamics in both cells are described by a simple model which was introduced in [15]. All details are described in Appendix A.

In biology, the typical input signal for the TC cells is a spike train with unregularly distributed interspike intervals, which can be modeled by a Poisson process with a refractory period. For example, the retinal cells fire such unregularly distributed spikes in darkness or under constant illumination. The refractory period is necessary because every biological cell needs some time τ_r to recover after it has fired a spike, so the cell cannot fire spikes with arbitrary low interspike intervals (ISIs). In our model, the ISI are distributed according to the following distribution:

$$\widetilde{f}(\tau) = \begin{cases} 0 & \text{for } \tau < \tau_r \\ \overline{r}e^{\overline{r}\tau_r}e^{-\overline{r}\tau} & \text{for } \tau \ge \tau_r, \end{cases}$$
(3)

where $e^{\overline{r}\tau_r}$ is a scaling factor, $\overline{r}=1/100$ and $\tau_r=30$ ms here. If we stimulate our model by such a modified Poisson process, we have a computational model for the experiment of Le Masson *et al.* [2], which allows us to compare our computational model with the experiment.

A. Simulation results of the biophysical model: Spindle oscillations, waxing and waning, and influence on information transfer

With the TC and RE cells interconnected, the computational thalamic circuit receiving this realistic synaptic bombardment [2] showed waxing and waning oscillations. These spindle sequences consisted of bursts of 8-10 Hz oscillations lasting a few seconds, and were separated by silent phases of around 8-10 s. The periodical occurrence of this oscillations is very similar to biological spindle waves which occur in sleep-related states. As in the experiment, each spindling state is terminated by an after-depolarization of the membrane potential. The reason for this effect is the hyperpolarization activated current I_h (see Appendix A); when this after-depolarization adds to the synaptic bombardment, it forms a depolarization which deactivates the low threshold Ca²⁺ current I_T and subsequently blocks the input signal transfer.

B. Measuring the information transfer

During the spindling state, the firing pattern of the TC cells (excitor), which is very different from the input [see Figs. 2(a) and 2(b), shows that obviously the information transfer of the input is low. Le Masson *et al.* [2] answer the question whether this low transfer is still reliable, by calculating two different indices. First, the contribution index (T_{Cl}) examines the TC cell discharge and quantifies the percentage of output spikes Nout which are precisely correlated with retinal input spikes. It estimates the reliability with which a TC spike can be considered as being triggered by an input spike rather than being spontaneous. It is computed as the peak of the cross correlation, normalized by the number of output spikes. Second, the correlation index (T_{CC}) measures the global efficiency of the input-output spike transfer and indicates the ratio of input spikes being actually transmitted as output spikes in the TC neuron. It is computed as the peak of the cross correlation between retinal and TC neuron spikes, normalized by the number of retinal cell spikes. To be able to compare our numerical results with the experimental results of Le Masson *et al.* [2] quantitatively, we will use exactly these measures here.

To be able to compare the numerical results obtained from the biophysical model with the simplified model introduced in this paper, we will define two new measures for the information transfer which are equivalent to the measures used by Le Masson *et al.* in [2], but do not depend on the detailed form of the signals. First, the signal-to-noise ratio (T_{SN}) examines the amplitude of the TC-cell signals and quantifies



FIG. 2. (Color online) (a) Spontaneous spindle activity in the computational thalamic circuit. The TC cell receives artificial synaptic retinal bombardment, modeled by a Poisson-distributed spike train. The RE-TC oscillator shows spindle activity. (b) Detail of (a). (c) Total synaptic current injected into the biological TC cell and simultaneous voltage of the TC and RE cell. Bottom: The hyperpolarization of the TC cell activates the I_T current, what initiates a post-inhibitory rebound burst (see also Fig. 5). Calibration bars: 10 s, 20 mV (a); 1 s, 20 mV (b); 10 ms, 6 nA, 20 mV, 1 nA (c).

the percentage N_{tr} of output spikes N_{out} which are exactly triggered by an input spike; such it is a measure equivalent to the T_{CI} used. An output spike is considered to be triggered by an input spike if it occurs within a delay of <50 ms after an input spike. As the T_{CC} , it is a measure for the reliability with which an output spike can be considered as being triggered by an input spike rather than being due to the intrinsic dynamic of the circuit

$$T_{SN} = \frac{N_{tr}}{N_{out}}.$$
 (4)

Second, the transfer efficiency T_{TE} measures the global efficiency in the input-output spike transfer, and indicates the ratio of input spikes being actually transmitted as output spikes, obviously it is equivalent to the measure T_{CC} . It is computed by the number of triggered spikes N_{tr} divided by the total number of input spikes

$$T_{TE} = \frac{N_{tr}}{N_{in}}.$$
 (5)



FIG. 3. (Color online) Percentage of output spikes triggered by an input spike (T_{SN}) (solid) and transfer efficiency T_{TE} (dashed) for different values of the inhibitory feedback. Both indices show no significant change when the inhibitory feedback is varied within the stability borders.

The difference between the measures (T_{SN}) and (T_{CI}) is that our measure checks if an output spike occurs within a certain delay after an input spike, while the contribution index counts the spikes at a fixed time delay defined by the maximum of the cross correlation. The T_{CC} and T_{TE} differ in the same way as mentioned above, further T_{TE} only takes values between zero and one. As our measures only count the spikes, and such treat spikes like binary events, they can be used for any spiking system and allow a comparison of the transmission properties of different models. The disadvantage is that they do not shed light on the amplified details of a signal system such as T_{CI} and T_{CC} do.

These two measures, the transfer efficiency and the transfer reliability, are necessary to characterize the information transfer appropriately. The usage of two different transfer measures allows us to answer the question whether the transfer gets blocked or if incoming signals get masqued by autonomous oscillations of the system. The results are given in Fig. 3.

In the presence of strong inhibitory feedback (g_{GABA_a} =0.1 mS and g_{GABA_b} =0.01 mS) the (T_{CI}) and (T_{SN}) were low (\approx 0.4), indicating that less than half of the output spikes were triggered by an input spike; thus the thalamus is not transferring spikes in a one-to-one manner. To answer the question whether the degree of inhibition produced by the RE neurons could directly control the precision of spike transfer, we varied the strength of inhibition. Our simulations show that the influence of the inhibitory feedback is by far not so strong as in the experiment [2]. In our computational model, the T_{SN} varied only between \approx 0.6 and \approx 0.4. In agreement with the experimental data [2], the global efficiency of spike transfer T_{TE} was not significantly different in the absence or presence of inhibitory feedback.

C. Discussion of the biophysical model

Altogether, our simulation cannot reproduce the transmission behavior of the thalamic circuit observed [2] experimentally. When we compare the numerical achieved values for



FIG. 4. Averaged contribution (T_{Cl}) (a) and correlation (T_{CC}) (b) indices versus synaptic strength.

the T_{CI} and T_{CC} (Fig. 4) quantitatively with the values from the experiment, we find that the effect of input-output spike decorrelation is present, but lacks some features. First, even without or very low inhibitory feedback, the T_{CC} is not more than ≈ 25 ; in the experiment the T_{CC} varied in a range from about 85–45 (see Fig. 3(e) in [2]). For a biological system this would mean that the animal would always be drowsy. Second, the system does not allow precise up and down regulation of the signal reliability in a wide range, the T_{CI} only varies in range from around 0.7–0.35, that is about a factor 2, while in the experiment the T_{CI} varies from about 0.8–0.2 (see Fig. 3(d) in [2]) which corresponds to a factor 4. Further, spindle oscillations occurred even with very little or no inhibitory feedback, which is also in contrast to the experimental results.

At this point it appears questionable whether the detailed biophysical model described above—with its high dimensionality and parameter space quite too detailed to gain systematic understanding—provides the appropriate level of description to analyze coupled systems, e.g., of the thalamocortical loop investigated here. We will however use the biophysical model to establish our simplified model. As the biophysical model incorporates many features of realistic biological neurons, it allows us to identify the key elements which may be responsible for sleep spindle oscillations. The slow repetition rate of spindling (0.3-0.1 Hz) is due to intrinsic mechanisms of the TC cell. The slow variables F_2 and S_2 (Fig. 11) of the I_h current (see Sec. III D and Appendix A) play the key role for these long silent phases. The spindling starts in the TC cell due to the hyperpolarization low threshold Ca²⁺ current I_T [see Figs. 2(c) and 5]. The spindling terminates due to the activation of the Ca2+ activated current I_H (see Fig. 5). The frequency of the spindling is determined by intrinsic properties of the RE cell, as shown by Destexhe in [4], but also the intrinsic properties of the inhibitory synapses play a role. To provide an overview, Fig. 5 shows a schematic diagram of the mechanisms leading to spindle oscillations.

Unfortunately, due to the complexity of the highdimensional biophysical system, the connection between biophysics and dynamics is hard to unravel. Further, the computational effort is very high, making studies of networks almost impossible. To solve these problems, we propose a reduced model to reproduce the experimental data of Le Masson *et al.* by using the simpler Hindmarsh-Rose [12] type neurons, and modify them in a way that the features described above are reproduced.

III. THE REDUCED SYSTEM: AN EXTENDED HINDMARSH-ROSE MODEL

A. The Hindmarsh-Rose model

The Hindmarsh-Rose equations are a simple polynomial model of bursting in thalamic cells, which reproduce several features, like for example rebound bursts, of more complicated biophysical models [3,21,22]. The original Hindmarsh-Rose equations are given by



Input Spike

FIG. 5. Schematic diagram illustrating the generation mechanism of spindle oscillations. We distinguish two cycles: First, the spindle cycle is responsible for the spiking; the system runs several time through it before the waxing and waning cycle is activated, leading to bursts of oscillations. The slowly decaying current I_h leads to the long silent epochs during which spike transfer is blocked. Altogether, the interplay of the currents I_T and I_h plays an important role for the genesis of spindle oscillations. (See also Figs. 2 and 3.)



FIG. 6. Maximum and or minimum bifurcation diagram of v in the fast subsystem (4) as a function of z. Heavy lines indicate stable fixed points (limit cycles) and thin lines indicate unstable fixed points (limit cycles). (See Appendix B 3.)

$$\dot{v} = w - v^3 + 3v^2 - z + I(t), \tag{6}$$

$$\dot{w} = 1.8 - 5v^2 - w, \tag{7}$$

$$\dot{z} = \epsilon [3.3(v+1.56) - z],$$
 (8)

where v is the voltage, w is a recovery variable, I(t) is an external forcing, and z is a slow variable. In order to understand the behavior of Eqs. (6)–(8), we first consider the limit $\epsilon \rightarrow 0$, so z(t)=z, which results in a planar reaction-diffusion system. This fast subsystem is responsible for the spiking dynamics of the Hindmarsh-Rose system. The dynamics of this subsystem is also crucial for $\epsilon \neq 0$. This procedure is called singular approximation, or slaving principle, and is widely used to separate fast and slow subsystems [23–25]. Slow variables (here z) can be treated as slowly varying parameters and the rest of the system can be studied as a function of these new parameters. We apply this method to the Hindmarsh-Rose model.

To understand the influence of the variable *z* on the (v, w) subsystem, Eqs. (6) and (7) with I(t)=0 are transformed to a Lienard system [7,26], yielding (Appendix B 2)

$$0 = \ddot{v} + f(v)\dot{v} + g(v),$$
(9)

where $g(v)=v^3+2v^2-1$ can be considered as the gradient of a potential $\Phi(v,z)$, and $f(v)=1-6v+3v^2$ as a damping term. The roots of g(v,z) give the fixed points of the equation. Depending on the parameter *z*, there exist either one or three real roots.

The bifurcation diagram of Eqs. (6) and (7) as a function of z is shown in Fig. 6 (the bifurcation diagram was computed with the auto interface [27] of xpp [28]). If we start with z=2 from the fixed point (v=-2.05, w=-19.67) of the fast subsystem and go to the left in the bifurcation diagram, the stable fixed point loses its stability by a saddle node bifurcation at $z \approx 0.6148$. For values of z between [0.6148,0.8891], a stable limit cycle and a stable fixed point coexist. At z=-10.4, the upper branch of the steady state



FIG. 7. (Color online) v(t) (black) and z(t) (red, upper curve) with I=1.0. When z(t) is periodic, v(t) oscillates with the same period; thus the slow variable z(t) slaves v(t).

loses stability by a Hopf bifurcation (for further details see [7,29]). Here we will only focus on values of z between [-5,+5].

B. Periodic forcing of the Hindmarsh-Rose neuron

Another important case is that of z being a periodic function p(t); this is the case for certain values of a constant external forcing I [29], or when the neuron is coupled to another periodic spiking or bursting neuron. So in a next step we consider the fast (v, w) subsystem with periodic forcing

$$\dot{v}(t) = w - v^3 + 3v^2 + p(t),$$

$$\dot{w}(t) = 1.8 - 5v^2 - w,$$
 (10)

or, in Lienard form

$$p(t) = \ddot{v} + f(v)\dot{v} + g(v),$$
 (11)

where f and g are defined as above. According to [30], theorem 4.3.1 and 4.3.3, Eq. (11) has a nonconstant periodic solution with the same period as the forcing term p(t) if several conditions (i–vi) (given in Appendix B 4) are fulfilled. Hence, we make the following proposition: When p(t)is a *T*-periodic function, then (11) has a periodic solution with the same period *T*. The detailed proof is in Appendix B 4.

For illustration, we show the behavior of the Hindmarsh-Rose Eqs. (6)–(8) excited with a constant forcing $I(t)=I_0$ (see Fig. 7), thus again z(t) is a periodic function. This is an example for a typical intrinsic burster, where the slow variable slaves the fast subsystem [27]. The theorem presented above proves that the fast (v,w) subsystem of the Hindmarsh-Rose model shows mode locking when it is driven by a periodic forcing, no matter what frequency or amplitude this forcing has.

If we assume ϵ to be small (≈ 0.006 as used by [31]), z can be treated like a slowly varying parameter in the (v, w) system (10). To understand the influence of v on Eq. (8), we write it in the form of a relaxation process



FIG. 8. (Color online) v(t), z(t), and I(t). The system shows a post-inhibitory rebound burst after a hyperpolarizing step of I(t)=-0.5 and a duration of 70 ms.

$$\dot{z} = -\epsilon [z - z_{\infty}(v)], \qquad (12)$$

where $z_{\infty}(v)=3.3(v+1.56)$. As ϵ is very small, the fast dynamics of v has only little influence on Eq. (12), as its own dynamics is too slow to follow the fast $z_{\infty}(v)$. Because of this, z will arrive in a steady state if v is spiking quickly, and then just oscillate with a very little amplitude around it. This effect leads to the so-called spike frequency adaption (for details see [12]), which is also observed in biological neurons [22]. For small enough values of I(t) this effect can also lead to an effect similar to intrinsic bursting [29].

C. Post-inhibitory rebound bursts in the Hindmarsh-Rose neuron

Post-inhibitory rebound bursts are a dynamical feature being also present in the biophysical neuron model [3]. As it is crucial for spindle oscillations and for the information transfer, we will describe it in detail here, see also Fig. 8. If the Hindmarsh-Rose model [12] is hyperpolarized for a period similar to the burst duration, the adaption current z will de-

FIG. 9. (Color online) Right: Reciprocally coupled RE and TC cell. Left: If the excitor cell gets activated by an external pulse, it excites the inhibitor cell, which leads to an inhibitory current. This current hyperpolarizes the excitor cell and evokes a rebound burst in it, and the mechanism repeats, which leads to self-sustained oscillations.

FIG. 10. (Color online) Reciprocally coupled inhibitor and excitor; without the h current, the oscillation does not terminate.

crease below its steady state. If the hyperpolarizing current is released, the z current stays below its steady state for some time due to its slow timescale. As a consequence, the model behaves as if some extra current had been applied, so it will cross the saddle node bifurcation in Fig. 6 and stay on the limit cycle. Due to the spikes, z will increase above its steady state, stopping the burst (see Figs. 8 and 9). Due to the slow time scale, a post-inhibitory rebound burst will only appear if slowly decaying dynamic synapses are used, as otherwise z will not decrease below its steady state. In the biophysical model, the same effect occurs due to the low threshold calcium current I_T [see Fig. 2(c)] [3], so the variable z can be interpreted as a calcium current.

In reciprocally coupled neurons, the effect described above leads to self-sustained oscillations if the excitor gets activated by a single input spike; this effect is an example for so-called hard-excited self-oscillations [25]. During this oscillation, the two neurons exhibit antiphase synchronization. We show this for the case of two Hindmarsh-Rose neurons, which are coupled like in a typical thalamocortical circuit (see Fig. 9).

D. Motivation of a simplified calcium current: Extending the Hindmarsh-Rose model

The self-sustained oscillations in Fig. 10 are stable and do not terminate; this is contrary to the experimental observations of Le Masson *et al.* [2]. A spindle oscillation or a burst consists of a series of metastable self-oscillations enhanced by rebound bursts which terminate after a few seconds. Above we argumented that the slow variable z can be inter-

preted as a calcium current, and such corresponds to the low threshold calcium current I_T in the biophysical model. The schematic diagram in Fig. 5 shows that the reason for the termination of the oscillation is a further ion current I_h [2–4,32]. As the original Hindmarsh-Rose model does not incorporate a variable which corresponds to the I_h current in the biophysical model, we will extend the Hindmarsh-Rose model by an equation motivated directly by the dynamics of I_h in the biophysical model. Destexhe *et al.* [4] model I_h by a double activation kinetic, consisting of slow and fast activation variables, regulated by intracellular Ca²⁺

$$S_0 \rightleftharpoons S_1,$$
 (13)

$$F_0 \rightleftharpoons F_1,$$
 (14)

were $S_{0/1}$ and $F_{0/1}$ represent the closed and open states of the slow and fast activation gates, respectively. The open state gates are assumed to have *n* bindings for Ca²⁺ which lead to the open bounded gates S_2 and F_2 ,

$$S_1 + n\mathrm{Ca}^{2+} \rightleftharpoons S_2, \tag{15}$$

$$F_1 + n\mathrm{Ca}^{2+} \rightleftharpoons F_2. \tag{16}$$

In this model the activation function of I_h is shifted during the oscillatory phase by the entry of Ca²⁺, and thus terminates the oscillation. Destexhe et al. [4] find that the length of the silent phase and of the oscillatory phase were directly proportional to the time constant of intracellular Ca²⁺ binding to I_h channels, k_2^{-1} . Further it is assumed that the binding of Ca²⁺ is critical for the onset and termination of the oscillatory phase [4]. So the length of the oscillatory phase depends on the rate of rise of the variables S_2 and F_2 , while the length of the silent phase depends on the rate of relaxation of S_2 and F_2 back to their resting values (see Fig. 11 and [4]). According to (A20), both the length of rising and relaxation of S_2 (red) and F_2 are proportional to k_2^{-1} . As Fig.11 shows, F_2 only displays small variations of amplitude compared to S_2 and therefore plays a less important rule. In our further simplification we will neglect the influence of F_2 . The slow variable S_2 slaves the system and switches between the oscillatory and the resting state leading to waxing and wanning oscillations [4]. Our simplified model will be motivated from the kinetic equation of S_2 . As the transition from S_0 to S_1 is much faster than the one from S_1 to S_2 , we will assume it to be instantaneous, what leads to $S_2 \approx S_0$, so we get

$$S_0 + n\mathrm{Ca}^{2+} \rightleftharpoons S_2. \tag{17}$$

In Eq. (5) in [4], the number of binding sites *n* for Ca²⁺ is assumed to be 2, in our model we use n=1 with the backward rate $k_2=4\times10^{-4}$, the forward rate $k_1=k_2/(5\times10^{-4})$ and $C=[\text{Ca}_{in}]/(5\times10^{-4})$, where $[\text{Ca}_{in}]$ is the intracellular calcium concentration, thus altogether we get

$$\dot{S}_2 = -k_2[S_2 - CS_0]. \tag{18}$$

The essential features of Eq. (18) can be summarized as follows: S_2 opens and closes proportional to the same rate constant k_2 ; due to C, the activation depends critically on the

FIG. 11. (Color online) Top: Time course of the gating variables S_2 (red, upper curve) and F_2 (green, lower curve) during a spindle cycle. Bottom: The membrane potential V_{Tc} of the biophysical model.

concentration of intracellular Ca²⁺. This features should also be present in our simplified model. In the Hindmarsh-Rose model the calcium current is mimicked by the variable z[12], in our model for the *h* current will get activated by the variable *z*. The rate of activation and deactivation will be equal to the constant used in the biophysical model. For convenience and to be consistent with the notation used in biophysics, we call this current *h*

$$\dot{h}_{TC} = -k_2 [h_{TC} - 0.88(0.9 - z_{TC})].$$
 (19)

As z decreases when the cell gets hyperpolarized, we use the difference between the maximum magnitude of z and z itself as an activation term in (19). In order to reproduce the behavior of the biological TC cell, we extend the Hindmarsh-Rose equations as follows:

$$\dot{v}_{TC} = w_{Tc} - v_{TC}^3 + 3v_{TC}^2 - z_{TC} - h_{TC}, \qquad (20)$$

$$\dot{w}_{TC} = 1.8 - 5v_{TC}^2 - w_{TC},\tag{21}$$

$$\dot{z}_{TC} = \epsilon [3.3(v_{TC} + 1.56) - z_{TC}],$$
 (22)

$$\dot{h}_{TC} = -0.0004[h_{TC} + 0.88(0.9 - z)].$$
 (23)

FIG. 12. (Color online) Top: Time course of *h* during a spindle cycle. Bottom: The membrane potential v_{Tc} of the extended Hindmarsh-Rose model.

Equations (20)–(23) define our central model, and will be used to model the reciprocally coupled neurons in the remainder. As Fig. 12 shows, the time course of the variable his very similar to the time course of s_2 in the biophysical model. To understand the influence of the variable h we proceed in the same way as before. The time scale of the dynamics of h is less than a tenth than the time scale of z. As declared above, the reason for the self-sustained oscillations is the after-hyperpolarization activation by the variable z. In the presence of h, the slowly varying parameter in the (v, w)system is the sum [z(t)+h(t)]. As h(t) gets activated by z(t), after some rebound bursts the sum [z(t)+h(t)] is below the threshold of the (v, w) system, as a consequence, the oscillation terminates. When the excitor cell is inactive, there are no more inhibitory currents, that means that t is inactive too. so h decays slowly until it is small enough that the system can get activated again by an input spike.

E. Reciprocally coupled TC-RE neurons using the extended Hindmarsh-Rose model

As the *h* current is absent in the biological RE neuron, we have to extend the Hindmarsh-Rose system by the *h* Eq. (23) for the TC neuron only

FIG. 13. (Color online) Spontaneous spindle activity in our computational model (the numerical values of the voltages are scaled by a factor of 30 to match with the biophysical values). Similar rescalings have been also necessary in the underlying Hindmarsh-Rose model (see, e.g. [22]). (a) The computational circuit: A TC cell (excitor) modeled by the extended Hindmarsh-Rose model is reciprocally coupled to a model RE cell (inhibitor) represented by the original Hindmarsh-Rose model. The TC cell receives artificial synaptic retinal bombardment modeled by a Poisson distributed spike train. Like in the experiment the system shows spindle activity. (b) Detail of (a). (c) Like in Fig. 8.

$$\begin{aligned} T_{TC} &= w_{TC} - v_{TC}^{3} + 3v_{TC}^{2} - z_{TC} - h_{TC} + I_{GABA} + I(t), \\ &\dot{w}_{TC} = 1.8 - 5v_{TC}^{2} - w_{TC}, \\ &\dot{z}_{TC} = 0.006[4(v_{TC} + 1.56) - z_{TC}], \\ &\dot{h}_{TC} = -0.0004[h_{TC} + 0.88(0.9 - z)], \\ &\dot{v}_{RE} = w_{RE} - v_{RE}^{3} + 3v_{RE}^{2} - z_{RE} + I_{GLU}, \\ &\dot{w}_{RE} = 1.8 - 5v_{RE}^{2} - w_{RE}, \\ &\dot{z}_{RE} = 0.006[4(v_{RE} + 1.56) - z_{RE}]. \end{aligned}$$

$$(24)$$

v

FIG. 14. (Color online) Percentage of output spikes triggered by an input spike. An increase of the strength of the inhibitory coupling g_{GABA} comes along with a significant decrease in the reliability of the information transfer. As in the experiment of Ref. [2] and in the biophysical model, the transfer efficiency does not vary significantly.

 I_{GABA} and I_{GLU} are governed by Eqs. (B4) and (B5) in Appendix B 1, respectively. The external input was modeled as described above for the biophysical model, with modified parameters adapted to the Hindmarsh-Rose model, \bar{r} was 1/100 and τ_0 was 30 ms. If we stimulate our model by such a modified Poisson process, we have a computational model for the experiment of Le Masson *et al.* [2], which allows us to compare our computational model with the experiment and the biophysical model [3], so that we can validate our extension of the Hindmarsh-Rose model.

A comparison of Fig. 13 and ([2], Fig. 2) shows that our system reproduces the experimental results of Le Masson *et al.* [2] quite well. During the spindling state, the firing pattern of the TC cells (excitor), which is very different from

the input, shows that the information transfer of the input is low. The question whether this low transfer is still reliable is calculated in the same way as for the biophysical system. With a strong inhibitory feedback, the T_{SN} was low, showing that most of the TC spikes where not triggered by an input spike and thus that the system is not transferring spikes in a one-to-one manner. This result encouraged us to screen the strength of inhibition to test if recurrent feedback inhibition could be a way to control the precision of input-output transfer in a wide range, or if there is just a switching between a relay state and masquerading state. Our numerical investigation leads to the following results. An increase of the inhibitory coupling strength leads to a smooth decrease in T_{SN} from a maximum value of around 1 to a minimum value of less than 0.3 (Fig. 14). Despite this significant decrease in the reliability of spike transfer, the efficiency of spike transfer T_{TE} was not significantly diminished by the strength of the inhibitory coupling (Fig. 14). Thus inhibitory feedback has a direct decorrelating effect, which is able to reduce the reliability of the spike transfer.

F. Dynamical behavior of coupled TC-RE neurons using the extended Hindmarsh-Rose model

As we argued in Sec. III A for the Hindmarsh-Rose model, Destexhe *et al.* [4] use singular approximation to characterize the spindle oscillations as a transition between a hyperpolarizing stationary phase and an oscillatory phase. Here we will apply this method to our simplified model of a thalamocortical oscillator. In Fig. 13, the *h* current evolves according to a much slower time scale as the dynamics of the coupled TC-RE neurons.

We assume *h* to be a slow varying parameter in (24) with I(t)=0, so we consider the limit $k_2 \rightarrow 0$ in (19). We will consider *h* as bifurcation parameter of (24) in an interval be-

FIG. 15. Bifurcation diagram of (24) with h_{TC} treated as a slowly varying parameter; solid lines indicate stable fixed points, dashed lines unstable fixed points, filled circles stable limit cycles, open circles unstable limit cycles. For extreme values of *h*, the system is monostable, i.e., either a stable fixed point or a stable limit cycle coexist.

tween -1 and 1.5 as a parameter. In this range of h the dynamical state of the system can be divided in three areas: For high values of h the system is in a stable resting state, for intermediate values of h both a stable resting state and a stable oscillatory state exist, separated by an unstable limit cycle. For negative values of h the stable resting state gets unstable, while the stable oscillatory state persists. The transition between the stable fixed point and the oscillatory state occurs by a subcritical Hopf bifurcation [33]. The existence of a region of parameters where stable solutions overlap is typical for a subcritical Hopf bifurcations; our system shows this typical behavior in a wide range of the bifurcation parameter. In this states the system is bistable, i.e., a stable fixed point and a stable limit-cycle coexist (see Fig. 15). In this area, the state of the system depends on its history: If the initial conditions or the past state of the system lie within the basin of attraction of the limit cycle, the system exhibits stable oscillations; if the starting point lies within the basin of attraction of the fixed point, the system rests on the fixed point. These two attractive areas are separated by an unstable limit cycle. Further, the system can be switched between these two states by an input pulse.

G. Discussion of the transfer properties of the extended Hindmarsh-Rose model

The *h* current switches the system between two existing dynamical states, namely a fixed point and a bistable state where a limit cycle and a fixed point coexist. Without this further current (i.e., h=0), two reciprocally coupled Hindmarsh-Rose neurons exhibit so-called hard selfexcitation [25] as response to an input spike, resulting in stable self-sustained oscillations. The h current makes this oscillatory state metastable, resulting in waxing and waning oscillations. An increase of the inhibition changes the responsivity of the system: Without, or with only little inhibition, the system responds to an input pulse with an output spike, so that spikes get transmitted in a one-to-one manner. With high gain in inhibitory feedback, the system responds to an input spike by an output burst (range B in Fig. 15), followed by a silent period where transmission is totally blocked (range A in Fig. 15). During the burst, the system is in an autonomous self-oscillatory state which masquerades the input, this leads to an increase of the T_{SN} . As the system is still excitable, but with a different responsivity the T_{TE} does not change as strong as the T_{SN} . So the transmission behavior depends critically on the rate of rise and fall of the h equation. This insight might also help to improve the biophysical model in terms of information transfer.

IV. CONCLUSIONS

We have shown that an extended Hindmarsh-Rose model is able to reproduce the behavior of a biological thalamocortical relay neuron in the recent experiment of Le Masson *et al.* To gain more insight into the dynamical mechanisms, a simplification or reduction of the detailed biophysical model, which did not convincingly reproduce the experimentally observed decrease of the signal-to-noise ratio, was necessary. The widely used Hindmarsh-Rose model however does not show the characteristic waxing and waning oscillations. Further it does not exhibit the quiescence periods necessary for transfer of information. Especially, we have analytically proven that the fast subsystem of the Hindmarsh-Rose model gets slaved by periodic forcing, which can lead to intrinsic bursting if the forcing is slow. To account for the waxing and waning mechanism, we propose an extended Hindmarsh-Rose neuron model for the TC cell, directly motivated from the biophysical model, taking the low threshold calcium current explicitly into account.

In this paper, we have demonstrated that our extended Hindmarsh-Rose model serves as a computational model for the setup in the Le Masson *et al.* experiment, showing that the information transfer can be adjusted within a wide range by the gain of the recurrent feedback inhibition. Our numerical investigations strongly suggest that the low threshold calcium current plays an important role for the information transfer in thalamocortical circuits. From the technical simplicity of our approach and its agreement with the experiment, this approach may give rise to similar models for neural systems where a third time scale is apparent, and a second slow degree of freedom is necessary to describe the dynamics.

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APPENDIX A: THALAMOCORTICAL CIRCUIT

The thalamocortical circuit consists of a pair of TC and thalamic RE neurons connected as shown in Fig. 1. For each TC and RE cell several ion currents were included which will be described in detail here. All intrinsic currents are described by the same general equation [16]

$$I_i = \overline{g}_i m^M h^N (V - E_i). \tag{A1}$$

Here the current I_j is the product of the maximal conductance, g_j , activation m and inactivation h variable, and the difference between the membrane potential V and the Nernst reversal potential E_j . The powers N and M are the respective numbers of ion channels to be open synchroneously. The gating of a membrane channel is described by a first order kinetic scheme

$$\begin{array}{l} \overset{\alpha(V)}{C \to O}, \end{array} \tag{A2}$$

$$\begin{array}{c} \beta(V)\\ C \leftarrow O, \end{array} \tag{A3}$$

where *O* and *C* stand for the open and the closed state of the gate, with $\alpha(V)$ and $\beta(V)$ as the transition rates. According to the Hodgkin-Huxley model [20], the variables *m* and *h* represent the fraction of independent gates in the open state and are described by simple first order differential equations

$$\dot{m} = -\frac{1}{\tau_m(V)} [m - m_\infty(V)], \qquad (A4)$$

$$\dot{h} = -\frac{1}{\tau_h(V)} [h - h_\infty(V)].$$
 (A5)

The steady states $m_{\infty}(V)$ and $h_{\infty}(V)$ and the time constants $\tau_{m/h}(V)$, respectively, can be written as functions of the transition rates α and β , using $x \in \{m, h\}$, as

$$\tau_x(V) = \frac{1}{\alpha_x(V) + \beta_x(V)},$$
$$x_\infty(V) = \alpha_x(V)\tau_x(V).$$
(A6)

1. The TC cell

Here we summarize the results of the previously mentioned investigations of TC cell membrane properties. The membrane potential of the TC neuron is given by

$$C_m V_T = -I_{T_L} - I_T - I_h - I_{T_{Na}} - I_{T_K} - I_{GABA_a} - I_{GABA_b}.$$
(A7)

The area of a TC cell membrane is about 2.9×10^{-4} cm² which is according to a cell capacity of C_m =0.29 nF. All constants used in the simulations correspond to a cell of this size.

The leakage current I_{T_L} is the only passive current, and is governed by the Ohm law

$$I_{T_{I}} = g_{T_{I}}(V_{T} - E_{L}), \tag{A8}$$

where $g_{T_{t}} = 0.05 \text{ mS}, E_{L} = -86 \text{ mV}.$

All the other currents are active currents, and more complicated, as detailed below.

The sodium current I_{Na} has the form

$$I_{T_{Na}} = g_{Na}m(t)^{3}h(t)(V_{T} - E_{Na}),$$
(A9)

with $g_{T_{\text{Na}}}$ =30 mS and E_{Na} =50 mV. In addition we have for the gating variables m(t) and h(t)

$$m_{\infty}(V_T) = \frac{\alpha_m(V_T)}{\alpha_m(V_T) + \beta_m(V_T)},$$

$$h_{\infty}(V_T) = \frac{\alpha_h(V_T)}{\alpha_h(V_T) + \beta_h(V_T)},$$

$$\tau_m(V_T) = \frac{1}{\alpha_m(V_T) + \beta_m(V_T)},$$

$$\tau_h(V_T) = \frac{1}{\alpha_h(V_T) + \beta_h(V_T)},$$
(A10)

where

$$\alpha_m(V_T) = 0.32 \frac{(V_T + 37)}{1 - \exp\left(-\frac{V_T + 37}{4}\right)},$$

$$\beta_{m}(V_{T}) = 0.28 \frac{(V_{T} + 10)}{\exp\left(\frac{V_{T} + 10}{5}\right) - 1},$$

$$\alpha_{h}(V_{T}) = 0.128 \exp\left(-\frac{V_{T} + 33}{5}\right),$$

$$\beta_{h}(V_{T}) = \frac{4}{\exp\left(-\frac{V_{T} + 10}{5}\right) + 1}.$$
 (A11)

With Eqs. (A4) and (A5) the sodium current is completely described.

The potassium current $I_{\rm K}$ has the form

$$I_{T_{\rm K}} = g_{T_{\rm K}} m(t)^4 (V_T - E_k), \qquad (A12)$$

with $g_{T_{\rm K}}$ =2 mS and E_k =-95 mV. The gating and transition variables are given by

$$m_{\infty}(V_T) = \frac{\alpha_m(V_T)}{\alpha_m(V_T) + \beta_m(V_T)},$$

$$\tau_m(V_T) = \frac{1}{\alpha_m(V_T) + \beta_m(V_T)},$$

$$\alpha_m(V_T) = 0.032 \frac{(V_T + 35)}{1 - \exp\left(-\frac{V_T + 35}{5}\right)},$$

$$\beta_m(V_T) = 0.5 \exp\left(-\frac{V_T + 40}{5}\right).$$
 (A13)

With Eq. (A4) the potassium current is completely described. *The low threshold* Ca²⁺ *current* I_T is taken to be

$$I_T = g_{Ca}m(t)^3h(t)(V - E_T),$$
 (A14)

where $g_{Ca}=1.75$ mS and the reversal potential E_T depends on the Ca²⁺ concentration inside ([Ca]_{in}) and outside ([Ca]_{out}) the cell. It is defined by the Nernst equation

$$E_T = 1000 \frac{RT}{2F} \ln \frac{[\text{Ca}]_{out}}{[\text{Ca}]_{in}},$$
 (A15)

where $R=8.314 \, 41 \, J/K \, \text{mol}$, $T=309.15 \, \text{K}$, $F=96489 \, C/\text{mol}$, and $[\text{Ca}]_{out}=2 \, \text{mM}$ is considered to be constant. In the remainder, all concentrations will be denoted by square brackets, following an usual convention (see e.g., [13].

The calcium dynamics in the cell are described by a simple model which was introduced in [13],

$$\frac{d[\operatorname{Ca}]_{in}}{dt} = -AI_T - K_T[\operatorname{Ca}]_{in}/([\operatorname{Ca}]_{in} + K_d), \quad (A16)$$

where $A=0.179 \ m \ mol/ms \ \mu A$, $K_T=10^{-4} \ m \ mol/ms$ and $K_d=K_T$.

The gating variables are governed by the following system which was proposed by Wang [18]

$$\dot{m} = [\tau_m(V_T)]^{-1}[m - m_{\infty}(V_T)],$$

$$\dot{h} = \alpha_1(V_T)[1 - h - d - K(V_T)h],$$

$$\dot{d} = \alpha_2(V_T)[K(V_T)(1 - h - d) - d].$$
 (A17)

In addition, for this current the constants are

$$m_{\infty}(V_T) = \left[1 + \exp\left(\frac{V_T + 65}{7.8}\right)\right]^{-1},$$

$$\tau_m(V_T) = 0.15m_{\infty}(V_T) \left[1.7 + \exp\left(-\frac{V_T + 30.8}{13.5}\right)\right],$$

$$\alpha_1(V_T) = \exp\left(-\frac{(V_T + 162.3)}{17.8}\right) / 0.26,$$

$$K(V_T) = \sqrt{0.25 + \exp\left(\frac{(V_T + 85.5)}{6.3}\right)} - 0.5,$$

$$\alpha_2(V_T) = \frac{1}{\tau_2(V_T)[K(V_T) + 1]},$$

$$\tau_2(V_T) = \frac{62.4}{1 + \exp\left(\frac{V_T + 39.4}{30}\right)}.$$
 (A18)

The numerical values are fits to experimental data [19].

The hyperpolarization-activated cation current I_h finally is described by

$$I_h = g_h (S_1 + S_2) (F_1 + F_2) (V_T - E_h), \qquad (A19)$$

where $g_h = 0.15$ mS and $E_h = -43$ mV. The gating variables for this current are governed by

$$S_{1} = \alpha_{s}(V_{t})S_{0} - \beta_{s}(V_{t})S_{1} + k_{2}(S_{2} - CS_{1}),$$

$$\dot{F}_{1} = \alpha_{f}(V_{t})F_{0} - \beta_{f}(V_{t})F_{1} + k_{2}(F_{2} - CF_{1}),$$

$$\dot{S}_{2} = -k_{2}(S_{2} - CS_{1}),$$

$$\dot{F}_{2} = -k_{2}(F_{2} - CF_{1}),$$
 (A20)

where $C = [Ca_{in}]/(5 \times 10^{-4})$, $k_2 = 4 \times 10^{-4}$, the rate constants $\alpha_{s/f}(V_t)$ and $\beta_{s/f}(V_t)$ are related to the activation function H_{∞} and the time constants $\tau_{s/f}$

$$\alpha_s(V_t) = H_{\infty}/\tau_s,$$

$$\alpha_f(V_t) = H_{\infty}/\tau_f,$$

$$\beta_s(V_t) = (1 - H_{\infty})/\tau_s$$

$$\beta_f(V_t) = (1 - H_{\infty})/\tau_f,$$
(A21)

$$H_{\infty} = \left[1 + \exp\left(\frac{V_t + 68.9}{6.5}\right) \right]^{-1},$$

$$\tau_s = \exp[(V_t + 183.6)/15.24],$$

$$\tau_f = \exp\left(\frac{V_t + 158.6}{11.2}\right) / \left[1 + \exp\left(\frac{V_t + 75}{5.5}\right) \right].$$
(A22)

This makes the biophysical description of TC cell complete. While it contains a lot of simplifications, for example the single compartment assumption, it reproduces almost all important features of TC cells quite good. We will use this model in our computer experiment to verify theories obtained by using simplified models. For more details see [15].

2. The RE cell

The membrane potential of the thalamic RE neuron is governed by the cable equation

$$C_m \dot{V}_R = -I_{R_L} - I_{TS} - I_{K[Ca]} - I_{CAN} - I_{R_{Na}} - I_{R_K} - I_{GLU}.$$
(A23)

For the RE cell the membrane capacity is $C_m = 0.143$ nF, the leakage conductance is $g_{R_l} = 0.05$ mS and the reversal potential of the leakage current is $E_L = -80$ mV. This model of thalamic RE cell was introduced by Destexhe *et al.* in [34].

The equations for the sodium $I_{R_{Na}}$ and potassium $I_{R_{K}}$ current in the RE cell are the same as for the TC cell, except that the conductance $g_{Na}=100$ mS is different. The description of the RE cell potassium current is reached by putting $g_{K} = 10$ mS.

The kinetics of the low threshold Ca^{2+} current I_{Ts} was established as a model for voltage clamp data on rat RE cells using a so-called m^2h formalism [19]

$$I_{Ts} = g_{Ts}m(t)^2h(t)(V_R - E_{Ts}),$$
(A24)

where $g_{Ts} = 1.75 \ \mu\text{S}$ and E_{Ts} depends on the calcium concentration in the same way as described for the TC cell. The calcium dynamics are described by Eq. (A9), when I_T is replaced by I_{Ts} . In addition we have for this current

$$m_{\infty}(V_R) = \left[1 + \exp\left(-\frac{V_R + 52}{7.4}\right)\right]^{-1},$$

$$\tau_m(V_R) = 1 + \frac{1}{3} \left[\exp\left(\frac{V_R + 27}{10}\right) + \exp\left(-\frac{V_R + 102}{15}\right)\right]^{-1} h_{\infty}(V_R),$$

$$= \left[1 + \exp\left(\frac{V_R + 80}{5}\right)\right]^{-1},$$

where

ROLE OF INHIBITORY FEEDBACK FOR INFORMATION ...

$$\tau_h(V_R) = \frac{85}{3} + \frac{1}{3} \left[\exp\left(\frac{V_R + 48}{4}\right) + \exp\left(-\frac{V_R + 407}{50}\right) \right]^{-1}.$$
(A25)

The full kinetics of the I_T current is reached if m_{∞} , τ_m , h_{∞} , and τ_h are inserted in Eqs. (A4) and (A5).

The calcium dependent currents: The RE cell possesses further two calcium dependent currents: the slow K⁺ current $I_{K[Ca]}$, and the slow nonspecific cation current I_{CAN} . According to [15] they are modeled as voltage-independent currents described by Eq. (A1) with M=2 and N=0. The activation variable *m* obeys

$$\dot{m} = \alpha [\operatorname{Ca}_i]^2 (1 - m) - \beta m, \qquad (A26)$$

where α and β are rate constants and [Ca_i] is the intracellular calcium concentration. For $I_{K[Ca]} \alpha = 48 \text{ ms}^{-1}$, mM^{-2} , and $\beta = 0.03 \text{ ms}^{-1}$, for I_{CAN} , $\alpha = 20 \text{ ms}^{-1}$, mM^{-2} , and $\beta = 0.002 \text{ ms}^{-1}$. With this the description of the RE cell is complete.

3. Synaptic currents

Kinetic models of synaptic currents have been directly fitted to experimental data [3], based on whole cell recorded synaptic currents obtained in hippocampal neurons [35]. The $GABA_A$ and GLU (=glutamate) receptor mediated currents are both represented by a first order kinetic scheme [36]

$$C + T \leftrightarrows O, \tag{A27}$$

where the transition between closed (C) and open (O) states depend on the binding of the transmitter (T). The current is given by

$$I_{syn} = g_{syn}[O](t)[V(t) - E_{syn}],$$
 (A28)

where g_{syn} is the maximal conductivity and E_{syn} is the reversal potential. For AMPA receptors $E_{syn}=0$ mV and for $GABA_A$ receptors $E_{syn}=-80$ mV. $g_{syn}=0.05$ for $GABA_A$ and $g_{syn}=0.1$ for glutamate synapses. [O](t) is the fraction of open channels

$$\frac{d[O](t)}{dt} = \alpha \{1 - [O](t)\}[T](t) - \beta[O](t), \quad (A29)$$

and [T](t) is the concentration of transmitter released from time t to time t_{max}

$$[T](t) = A\Theta(t_{max} - t)\Theta(t), \qquad (A30)$$

where $\Theta(x)$ is the Heaviside function. The synaptic parameter values used in the simulation are A=0.5 and t_{max} =3 ms for $GABA_A$ and 9 ms for AMPA synapses, the rate constants were chosen as $\alpha=5$ ms and $\beta=0.16$ ms for $GABA_A$ synapses and $\alpha=0.94$ ms and $\beta=0.18$ ms for GLU synapses. The activation scheme of $GABA_B$ receptors is more complex, as it involves the activation of K⁺ channels by *G* proteins (for details see [37]). The model used by Le Masson *et al.* [2] is a modified version of the $GABA_B$ kinetic model introduced in [38]

$$R_0 + T \rightleftharpoons R \leftrightarrows D, \tag{A31}$$

$$R + G_0 \leftrightarrows RG \to R + G, \tag{A32}$$

$$G \to G_0,$$
 (A33)

$$C_1 + 4G \leftrightarrows O, \tag{A34}$$

where the transmitter, T, binds to the receptor, R_0 , leading to its activated form, R, and desensitized form, D. The G protein is transformed from an inactive (GDP bound) form, G_0 to an activated form, G, catalyzed by R. Finally, G binds to open the K⁺ channel, with four independent binding sites. With some assumptions (see [3]) the kinetic model for this system reduces to

$$I_{GABA_B} = g_{GABA_B} \frac{[G]^4}{[G]^4 + K_d} [V(t) - E_k],$$

$$\frac{d[R]}{dt} = K_1 (1 - [R])[T] - K_2[R],$$

$$\frac{d[G]}{dt} = K_3[R] - K_4[G],$$
 (A35)

where [*R*] is the fraction of activated receptors and [*G*] is the concentration of *G* proteins. In these equations g_{GABA_B} , $K_1 = 0.52$, $K_2 = 0.0013$, K_3 , $K_4 = 0.033$, and $K_d = 100$.

The strengths of the synaptic couplings $g_{GABA_{A/B}}$ and g_{AMPA} are varied in different simulations.

APPENDIX B: THE REDUCED MODEL

1. Synaptic currents

The simplest way to model synaptic currents is a two state scheme of the binding of a neurotransmitter T to postsynaptic receptors [3]. So the receptor mediated currents are both represented by a first order kinetic scheme

$$C + T \leftrightarrows O$$
,

where the transition between closed (*C*) and open (*O*) states depend on the binding of the transmitter (*T*) with forward and backward rates α and β respectively. As in [39] we only consider voltage-independent rate constants. With this assumptions we get the following kinetic equation

$$\frac{d[O]}{dt} = \alpha(1 - [O])[T] - \beta[O] = \alpha[T] - [O]\alpha[T] - \beta[O].$$
(B1)

The synaptic current is given by

$$I_{syn} = g_{syn}[O](t)[V(t) - E_{syn}], \qquad (B2)$$

where g_{syn} is the maximal conductivity and E_{syn} is the reversal potential. For glutamate receptors $E_{syn}=0$ mV and for $GABA_A$ receptors $E_{syn}=-50$ mV. The release and clearance of transmitter are extremely fast processes compared to the open/close kinetics, resulting in a very brief presence of transmitter in the synaptic cleft [40], which allows us to con-

sider *O* to be constant during the transmitter release. For simplicity we assume the transmitter release time course to be a square pulse $A\Theta(V_{pre})$ which occurs when the presynaptic neuron fires a spike, i.e., when the presynaptic potential V_{pre} gets depolarized towards positive potentials. If we assume that initially all synaptic channels are in the closed state, we get

$$\frac{d[O]}{dt} = \gamma \alpha A \Theta(V_{pre}) - \beta[O], \tag{B3}$$

were $\gamma = \alpha A$. As $GABA_B$ synapses contribute less than a tenth of the total inhibitory synaptic current, they are neglected here. The rate constants and amplitude were taken from the biophysical model A3. The constants for the *GABA* synapse were modified because of the absence of the *GABA*_B synapse: $\alpha = 5 \text{ ms}^{-1}$ and $\beta = 0.05 \text{ ms}^{-1}$. For the GLU synapse we used the same constants as in the biophysical model: α =0.94 ms⁻¹ and $\beta = 0.18 \text{ ms}^{-1}$. We chose A = 0.5, and respectively, $\gamma = 2.5$ for *GABA* synapses and $\gamma = 0.47$ for glutamate synapses. The synaptic currents are governed by the following equations:

$$I_{GABA} = g_{GABA}[O]_{GABA}(v_{TC} - 2.5),$$
 (B4)

$$I_{GLU} = g_{GLU}[O]_{GLU} v_{RE}, \tag{B5}$$

the reversal potential of the $GABA_A$ synapse was rescaled by a factor of 30 to adapt it to the scale of the Hindmarsh-Rose model.

2. Transformation to a Lienard system

We begin with the autonomous (v, w) subsystem (6) and (7)

$$\dot{v}(t) = w - v^3 + 3v^2, \tag{B6}$$

$$\dot{w}(t) = 1.8 - 5v^2 - w. \tag{B7}$$

Then $\ddot{v} = \dot{w} + (6v - 3v^2)\dot{v}$, $\dot{w} = \ddot{v} - (6v - 3v^2)\dot{v}$, and

$$w = -\ddot{v} + (6v - 3v^2)\dot{v} + 1.8 - 5v.$$
 (B8)

Inserting (B8) in (B6) yields

$$0 = \ddot{v} + (1.8 - 6v + 3v^2)\dot{v} + (v^3 + 2v^2 - 1.8),$$

$$\Leftrightarrow 0 = \ddot{v} + f(v)\dot{v} + g(v). \tag{B9}$$

(B9)is the so-called Lienard form of Eqs. (B6) and (B7), where $f(v)=1-6v+3v^2$ and $g(v)=v^3+2v^2-1.8$.

3. Stability of the equilibrium points

The stability of equilibrium points in the fast (v, w) subsystem is investigated by a linear approximation to the system (6) and (7). Suppose the equilibrium point has the *v* coordinate v_0 , then the linear approximation is

TABLE I. Equilibrium points depending on v_0 (see text). Only stable nodes and unstable spirals and saddle nodes occur.

Region	Values of v_0	Sign of $\operatorname{Tr} A(v_0)$	Sign of Det $A(v_0)$	Type of node
I	$v_0 < -4/3$	_	+	stable
II	$-4/3 < v_0 < Tr$	-	-	saddle
III	$Tr_{-} < v_0 < 0$	+	-	saddle
IV	$0 < v_0 < Tr_+$	+	+	unstable spiral
V	$\mathrm{Tr}_{+} < v_{0}$	+	-	saddle

$$\begin{pmatrix} \dot{x} \\ \dot{y} \end{pmatrix} = A(v_0) \begin{pmatrix} x \\ y \end{pmatrix}, \tag{B10}$$

where (x, y) are new coordinates whose origin is in the equilibrium point, and $A(v_0)$ is the Jacobian in v_0

$$A(v_0) = \begin{pmatrix} -3v_0^2 + 6v_0 & 1\\ -10v_0 & -1 \end{pmatrix}.$$
 (B11)

The kind of equilibrium point may be determined by the signs of the trace $Tr[A(v_0)] = -3v_0^2 + 6v_0 - 1$ and determinant $Det[A(v_0)] = 3v_0^2 + 4v_0$ [12,41]. The eigenvalues $\lambda_{1/2}$ of the Jacobian $A(v_0)$ are expressed by

$$\lambda_{1/2} = \frac{1}{2} \{ \operatorname{Tr}[A(v_0)] \pm \sqrt{\{\operatorname{Tr}[A(v_0)]\}^2 - 4\operatorname{Det}[A(v_0)]\}} \}.$$
(B12)

Table I gives the type of equilibrium point according to the region v_0 belongs to; here $\text{Tr}_{-}=(3-\sqrt{6})/3$ and $\text{Tr}_{+}=(3+\sqrt{6})/3$ are the negative and positive zeroes of $\text{Tr}[A(v_0)]$, respectively.

4. Periodic solutions for periodic forcing

The conditions for constants b, m, M > 0 are (i) for $|v| \ge b, f(v) > m$, (ii) $\forall v \in \mathbb{R}, f(v) > -M$, (iii) for $|v| \ge b, vg(v) > 0$, (iv) g(v) is monotone increasing in $(-\infty, -b)$, (v) $|g(v)| \to \infty$ for $|v| \to \infty$, (vi) $g(v)/G(v) \to 0$ for $|v| \to \infty$, para $G(v) = \int_{-\infty}^{v} g(v) dv$

where $G(v) = \int_0^v g(u) du$.

The proof can be sketched as follows:

(a) The existence of M is easy to establish as f(v) has a minimum at v=1, further as f(v) is a convex function this minimum is the global one. If a M > |f(1)=-2| is chosen (ii) holds.

(b) Set M=m, then b_1 may be chosen arbitrarily.

(c) Next vg(v) is a fourth degree polynomial where the coefficient of the v^4 term is positive. So there exists a b_2 with vg(v) > 0 if $v > b_2 > 0$.

(d) As g(v) is a third degree polynomial, there is a b_3 , such that g(v) is monotonously increasing in $(-\infty, -b_3)$ and (b_3, ∞) .

Choose $b = \max(b_1, b_2, b_3)$ then (i), (iii), and (iv) hold. Finally (v) and (vi) are obvious; so the proof is complete.

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