Energy cost reduction in the synchronization of a pair of nonidentical coupled Hindmarsh-Rose neurons

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Abstract Many biological processes involve synchronization between nonequivalent systems, i.e., systems where the difference is limited to a rather small parameter mismatch. The maintenance of the synchronized regime in these cases is energetically costly [1]. This work studies the energy implications of synchronization phenomena in a pair of structurally flexible coupled neurons that interact through electrical coupling. We show that the forced synchronization between two nonidentical neurons creates appropriate conditions for an efficient actuation of adaptive laws able to make the neurons structurally approach their behaviors in order to decrease the flow of energy required to maintain the synchronization regime.

1 Introduction

When a given oscillator moves freely on its natural attractor its oscillatory regime consists of a balanced exchange of energy between the system and its environment that occurs spontaneously through the divergent components of the systems structure without concurrence of any additional device. If, on the other hand, the system is forced to synchronize to a different guiding system its oscillatory regime occurs...
on an unnatural region of the state space where there is a nonzero net average exchange of energy with its environment. This net flow of energy per unit time requires the concurrence of a coupling device that includes an external source of energy. This flow of energy is necessary to maintain the synchronized regime and constitutes a cost for the synchronization process [1]. This consumption of energy can be reduced if the guided system itself adapts its structure to become closer to the one of the guiding system [3]. Ideally, if the systems become identical their joint dynamics is attracted toward a regime of zero error in the variables.

Many biological processes involve synchronization between different members of the same family of systems that have similar, although not identical, values of some distinctive parameters. This work studies the energy implications of synchronization phenomena in a pair of structurally flexible coupled neurons that interact through electrical coupling. We show that the forced synchronization between two nonidentical neurons creates appropriate conditions for an efficient actuation of adaptive laws able to make the neurons structurally approach each other in order to decrease the flow of energy required to maintain the synchronization regime. The neuron has been modelled by a four-dimensional Hindmarsh-Rose model [6, 7, 8, 9, 10]. This model is described by the following equations of movement:

\[
\begin{align*}
\dot{x} &= ay + bx^2 - cx^3 - dz + \xi I, \\
\dot{y} &= e - fx^2 - y - gw, \\
\dot{z} &= m(-z + s(x + h)), \\
\dot{w} &= n(-kw + r(y + 1)),
\end{align*}
\]

where \(a, b, c, d, \xi, I, e, f, g, m, s, h, n, k, r, \) and \(l\) are the parameters that govern the

![Fig. 1 Time series and 3D projections of the dynamical variables \(x(t), y(t), z(t), w(t)\) of the four-dimensional Hindmarsh-Rose neuron model](image-url)
dynamics of the neural system. The variable $x$ is a voltage associated to the membrane potential, variable $y$ although in principle associated to a recovery current of fast ions has been transformed into a voltage, variable $z$ is a slow adaptation current associated to slow ions, and variable $w$ represents an even slower process than variable $z$. $I$ is an external current input, and is the main parameter we used to control the modes of spiking and bursting activity of the model. For the numerical results of this work we fix the parameters to the values $a = 1, b = 3.09 (mV)^{-1}, c = 1(mV)^{-2}, d = 0.99 M\Omega, \xi = 1 M\Omega, e = 1.0128 (mV)^{-1}, g = 0.0278 M\Omega, m = 0.00215, s = 3.966 \mu S, h = 1.605 mV, n = 0.0009, k = 0.9573, r = 3.0 \mu S, l = 1.619 mV.

Figure 1 shows a chaotic time series of the four variables. The complexity achieved by the incorporation of a slow variable $w$ that increases the realism of the description of slow Calcium currents can be observed in the projections of the attractor on the $(x, y, z), (x, y, w)$ and $(x, z, w)$ axes.

In Sec. 2 we report the energy-like function associated to a four-dimensional Hindmarsh-Rose model. Sec. 3 briefly summarizes the adaptation mechanism we used to adapt the structure of the postsynaptic neuron, and presents computational results of the synchronization process of two electrically nonidentical coupled neurons. We consider that the presynaptic (sending) neuron always signal in a chaotic regime, while the postsynaptic (receiving) neuron is set to its quiescent state at a low value of its external current. In a first stage, the postsynaptic neuron has been forced to synchronize with the presynaptic one, then we initiate an adaptive process that adapts some parameters of the postsynaptic neuron to ones of the presynaptic neuron. We have analysed the energy dissipation of the receiving neuron during the synchronization process without and with structural adaptation.

2 Four-dimensional Hindmarsh-Rose model energy

In the Hindmarsh-model given by Eq. (1) the energy function $H(x)$ is given by [2]

$$H = \frac{p}{a} \left( \frac{2}{3} f x^3 + \frac{msd - gn r}{a} x^2 + ay^2 \right) + \frac{p}{a} \left( \frac{d}{ams} (msd - gn r) z^2 - 2dyz + 2gxw \right) \tag{2}$$

where $p$ is a parameter. As in the model time is dimensionless and every adding term in Eq. (2) has dimensions of square voltage, function $H$ is dimensionally consistent with a physical energy as long as parameter $p$ has dimensions of conductance. In this paper we fix parameter $p$ to the arbitrary value $p = -1 S$. The minus sign has been chosen to make consistent the outcome of the model with the usual assumption of a demand of energy associated with the repolarization period of the membrane potential and also with its refractory period (see Fig. 3).

And the corresponding energy derivative $\dot{H}$ is given by [2]
\[ \dot{H} = \frac{2p}{a} \left( f x^2 + \frac{msd-gnr}{ay-dz} z + gw \right) \left( bx^2 - cx^3 + \xi I \right) \left( \frac{dx}{ay} (msd-gnr) z - dy \right) \left( msh - mz \right) \left( nrl - nkw \right) \]  

(3)

is also dimensionally consistent with a dissipation of energy. As the states of an isolated Hindmarsh-Rose neuron are confined to an attractive manifold the range of possible values of its energy is recurrent and the long term average of its energy derivative is zero.

This energy and energy derivative functions are used to evaluate the energy consumption of the neuron in isolation and also when it is connected to other neurons through electrical synapses, and provide the basis for all the computational results presented in this work. The procedure followed to find this energy function has been reported in detail in [1].

![Graphs](image)

**Fig. 2** (a) Action potentials, (b) energy and (c) energy derivative for the Hindmarsh-Rose model neuron. (d), (e) and (f) Details of the action potential, energy and energy derivative associated to two spikes

Figure 2(a) shows a series of action potentials (variable \( x \) in the model neuron). Fig. 2(b) and Fig. 2(c) show both energy and energy derivative corresponding to that action potentials. Fig. 2(e) and Fig. 2(f) show detail of energy and energy derivative associated to a train of two action potentials. For each action potential it can be appreciated (see Fig. 2(e,f)) that the energy derivative is first negative, dissipation of energy while the membrane potential depolarizes during the rising period of the spike, and then positive, contribution of energy to repolarize the membrane potential during its descending period. During the refractory period between the two spikes the energy derivative remains slightly positive, still demanding energy, until the onset of the following action potential.
3 Synchronization energy of two electrically coupled neurons

In this section we analyze the energy aspects of the synchronization of two non-identical neurons coupled by an electrical synapse. The presynaptic neuron is set in the chaotic spiking-bursting regime corresponding to an external current $I_1 = 3.024$. While the postsynaptic neuron is set to its quiescent state at a low value $I_2 = 0.85$ of its external current. The two neurons are coupled unidirectionally according to the following equations:

$$
\begin{align*}
\dot{x}_i &= ay_i + bx_i^2 - cx_i^3 - dz_i + \xi I_i + K_1(x_j - x_i), \\
\dot{y}_i &= e - f x_i^2 - y_i - g w_i, \\
\dot{z}_i &= m(-z_i + s(x_i + h)), \\
\dot{w}_i &= n(-kw_i + r(y_i + 1)),
\end{align*}
$$

(4)

where $K_1 = 0$ and $K_2 \geq 0$ is the coupling strength. $i, j = 1, 2; i \neq j$ are the indices for the neurons. Note that the coupling affects only the first variables $x_2$ of the postsynaptic neuron.

The coupling scheme given in Eq. 4 can be written in general terms as follows:

$$
\begin{align*}
\dot{y} &= f(y, p) \\
\dot{x}_k &= f(x_k, q) + K(y - x_k),
\end{align*}
$$

(5)

where $p$ and $q$ stand for the parameters of the presynaptic and postsynaptic neurons, and $x_k(t)$ indicates the states of the postsynaptic neuron when the coupling strength is set to $k$. Notice that $K(y - x_k)$ is the coupling interface required in order to be physically able to implement the coupling of both neurons $\dot{x} = f(x)$ and $\dot{y} = f(y)$.

If the coupling strength $K$ is large enough as to make the errors in the variables $e = x_k - y$ small, an operational law that adapts the parameters of the postsynaptic neuron to the ones of the presynaptic neuron is given by

$$
\dot{e}_l^p = - \left[ \sum_{l=1}^{q} \left( \frac{\partial f_l(x_k, q)}{\partial q_l} \right) (y, p) e_l \right]
$$

(6)

where $e^p = q - p$ denotes the vector of parameter errors, and the summation is over every component of the vector field $f$. The above law is general and can be used to find specific adaptive laws to any kind of homochaotic systems provided they are coupled through a feedback scheme of large enough coupling strength.

In the following we analyse the change in the balance of energy of the postsynaptic neuron when its external current parameter is governed by an adaptive law in order to reach the nominal value of the external current in the presynaptic neuron. The adaptation law has been implemented following Eq. 6. For this experiment, we have used a coupling strength with value $k = 5$. We started the adaptation procedure at $t = 100$ and registered data between $t = 50$ and 200 for proper observation of the evolution of both energy and energy derivative during the process. The registrated values has been averaged over a convenient length of time in order to avoid large fluctuations. The dissipated energy (energy derivative) has been averaged over five units of time, while the proper energy has been averaged over ten units of time.
Figure 3 shows the average values of both energy and energy derivative per unit time. In Fig. 3(a) we can see that in a first stage ($t < 100$), the receiving neuron is forced to synchronize with the sending neuron, and oscillates in an unnatural region of the state space characterized by an average energy of about 66 (arb. units), and a nonzero energy derivative average of about 14 (arb. units), i.e., a nonzero net average exchange of energy with its environment (see Fig. 3(b)). After adaptation take place, the two neurons become structurally close each other, and enter in a completely synchronized regime of balanced exchange of energy between the system and its environment corresponding to zero value of energy derivative ($\dot{H} = 0$).
To illustrate the ability of this adaptive laws to decrease the energy dissipation, we have computed, for the receiving neuron, the average energy and average energy dissipation over a convenient length of time at different values of the coupling strength ranging from $K = 0$ to $K = 2$. The adaptation run at time $t = 100$.

Figure 4 shows average over ten units of time of the energy per unit time at different values of the coupling strength ranging from $k = 0$ to $k = 2$. It can be seen that before the adaptation occurs ($t < 100$), and for low value of the coupling strength not sufficient to induce a certain degree of synchrony, the receiving neuron shows a waving average energy pattern. This oscillating regime of energy average disappears once the adaptation process start, and decreases to low values. For high values of the coupling strength, the receiving neuron is forced to synchronize with the sending neuron and moves in a region of state space where the average of its energy is greater than that of the sending neuron. When the coupling strength is large enough as to make the errors in the variables small, and as soon as the adaptation process starts the average energy quickly decreases to values corresponding to the average energy of the sending neuron.

Figure 5 shows average over five units of time of the energy derivative per unit time at different values of the coupling strength ranging from $k = 0$ to $k = 2$. When $K = 0$, ie, no guidance at all, the receiving neuron moves on its natural region of state space and its averaged dissipated energy is zero. As soon as the coupling device is
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connected the average energy derivative per unit time becomes negative, that is, it
start to dissipate on average an energy that the coupling device will have to provide
in order to maintain the forced regime. The required energy increases with coupling
strength as it can appreciated in Fig.5 (c). Once the adaptation process starts the
average energy derivative quickly decreases to zero, reflecting the fact the receiving
neuron has become structurally so close to the sending neuron that they can reach a
regime of identical synchronization.

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