Effect of small-world connectivity on fast sparsely synchronized cortical rhythms

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ABSTRACT

Fast cortical rhythms with stochastic and intermittent neural discharges have been observed in electric recordings of brain activity. For these fast sparsely synchronized oscillations, individual neurons fire spikings irregularly and sparsely as Geiger counters, in contrast to fully synchronized oscillations where individual neurons exhibit regular firings like clocks. We study the effect of network architecture on these fast sparsely synchronized rhythms in an inhibitory population of suprathreshold fast spiking (FS) Izhikevich interneurons (which fire spontaneously without noise). We first employ the conventional Erdős–Rényi random graph of suprathreshold FS Izhikevich interneurons for modeling the complex connectivity in neural systems, and study emergence of the population synchronized states by varying both the synaptic inhibition strength J and the noise intensity D. Fast sparsely synchronized states of relatively high degree are found to appear for large values of J and D. However, in a real cortical circuit, synaptic connections are known to have complex topology which is neither regular nor random. Hence, for fixed values of J and D we consider the Watts–Strogatz small-world network of suprathreshold FS Izhikevich interneurons which interpolates between regular lattice and random graph via rewiring, and investigate the effect of small-world synaptic connectivity on emergence of fast sparsely synchronized rhythms by varying the rewiring probability p from short-range to long-range connection. When passing a small critical value $p^*_c$, fast sparsely synchronized population rhythms are found to emerge in small-world networks with predominantly local connections and rare long-range connections. This transition to fast sparse synchronization is well characterized in terms of a realistic “thermodynamic” order parameter. For further understanding of this transition, we also investigate the effect of long-range connections on dynamical correlations between neuronal pairs, and find that for $p > p^*_c$, global synchronization appears in the whole population because the spatial correlation length covers the whole system thanks to sufficient number of long-range connections. The degree of fast sparse synchronization for $p > p^*_c$ is also measured in terms of a realistic “statistical–mechanical” spiking measure. As p is increased from $p^*_c$, the degree of population synchrony becomes higher, while the axon “wire length” of the network also increases. At a dynamical-efficiency optimal value $p^*_E$, there is a trade-off between the population synchronization and the wiring economy, and hence an optimal fast sparsely-synchronized rhythm is found to occur at a minimal wiring cost in an economic small-world network.

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1. Introduction

Recently, brain rhythms have attracted much attention [1]. Particularly, we are interested in fast sparsely synchronized cortical rhythms, associated with diverse cognitive functions [2,3]. In some experimental data [4–10], synchronous small-amplitude fast oscillations (e.g., gamma rhythm (30–100 Hz) during awake behaviors and rapid eye movement sleep and sharp-wave ripple (100–200 Hz) during quiet sleep and awake immobility) have been observed in local field potential recordings, while individual neuron recordings have been found to show stochastic and intermittent spike discharges. Thus, single-cell firing activity differs markedly from the population oscillatory behavior. We note that these sparsely synchronized rhythms are in contrast to fully synchronized rhythms. For the fully synchronized rhythms, individual neurons fire regularly at the population frequency like the clock oscillators [11]. Hence, fully synchronized oscillations may be well described by using the conventional coupled-oscillator model composed of suprathreshold spiking neurons above a threshold in the absence of noise or for weak noise [12]. However, such coupled-oscillator models seem to be inappropriate for describing sparse synchronization because of stochastic and intermittent individual neural discharges like the Geiger counters. By taking an opposite view from that of coupled oscillators, the authors in Refs. [13–20] developed a framework appropriate for description of sparse synchronization. When the external noise is strong, suprathreshold spiking neurons discharge irregular firings as Geiger counters, and then the population state becomes unsynchronized. However, as the inhibitory recurrent feedback becomes sufficiently strong, a synchronized population state with stochastic and sparse neural discharges emerges. In this way, under the balance between strong external excitation and strong recurrent inhibition, fast sparse synchronization was found to occur in networks of suprathreshold neurons for both cases of random coupling [13–16] and global coupling [17–20]. Similar sparsely synchronized rhythms were also found to appear through cooperation of noise-induced spikings of subthreshold Morris–Lecar neurons (which cannot fire spontaneously without noise) [21–23].

In this paper, we study the effect of network architecture on fast sparsely synchronized cortical rhythms in an inhibitory population of suprathreshold fast spiking (FS) Izhikevich interneurons [24–27]. The conventional Erdös–Rényi random graph has been usually used for modeling complex connectivity occurring in diverse fields such as social, biological, and technological networks [28]. So, we first consider a random network of suprathreshold FS Izhikevich interneurons, and investigate occurrence of the population synchronized states by varying the inhibition strength and the noise intensity. Fast sparsely synchronized oscillations are found to appear when both the inhibition and the noise are sufficiently strong. Global efficiency of information transfer becomes high for random connection because its average path length (i.e., typical separation between two neurons represented by average number of synapses between two neurons along the minimal path) is short due to long-range connections [29,30]. However, random networks have poor clustering (i.e., low cliquishness of a typical neighborhood) and they are non-economic ones because the (axon) wiring cost becomes expensive due to appearance of short-range and long-range connections with equal probability [31,32]. In a real cortical circuit, synaptic connections are known to have complex topology which is neither regular nor completely random [31–39]. Hence, we consider the Watts–Strogatz model for small-world networks which interpolates between regular lattice with high clustering and random graph with short path length via rewiring [40–42]. The Watts–Strogatz model may be regarded as a cluster-friendly extension of the random network by reconciling the six degrees of separation (small-worldness) [43,44] with the circle of friends (clustering). Many recent works on various subjects of neurodynamics have been done in small-world networks with predominantly local connections and rare long-distance connections [38,45–56]. Here, we investigate the effect of small-world connectivity on emergence of fast sparsely synchronized rhythms by varying the rewiring probability \( p \) from local to long-range connections. As \( p \) is increased, long-range short-cuts that connect distant neurons begin to appear, and the average path length can be dramatically decreased only by a few short-cuts. Thus, global effective communication between distant neurons may be available via shorter synaptic paths. Eventually, when \( p \) passes a critical value \( p^*_c \), fast sparsely synchronized rhythm emerges in the whole population because dynamical correlation length covers the whole system thanks to sufficient number of long-range connections. However, with increasing \( p \), the (axon) wiring length also becomes longer due to appearance of long-range connections. Longer axonal projections are expensive due to their material and energy costs. Hence, we must take into account the (axon) wiring economy for the dynamical efficiency because wiring cost is an important constraint of the brain evolution [1,2,31,32,47,50,57–62]. At a dynamical-efficiency optimal value \( p^*_c \) an optimal fast sparse synchronization is found to occur via trade-off between synchrony and wiring cost at a minimal wiring cost in an economic small-world network [32].

This paper is organized as follows. In Section 2, we describe an inhibitory population of suprathreshold FS Izhikevich interneurons. The Izhikevich neurons are not only biologically plausible, but also computationally efficient [24–27], and they interact through inhibitory GABAergic synapses (involving the GABA\(_A\) receptors). In Section 3, we first consider the conventional Erdös–Rényi random graph [28], and study appearance of the population synchronized states by varying the noise intensity \( D \) and the inhibition strength \( J \). We fix \( J \) and \( D \) at appropriately strong values where sparsely synchronized rhythms of relatively high degree emerge. Then, we consider the Watts–Strogatz model for the small-world network which interpolates between the regular lattice and the random graph [40], and investigate the effect of the small-world connectivity on fast sparsely synchronized rhythms by increasing the rewiring probability \( p \). For the regular connection of \( p = 0 \), the average path length is very long because there exist only short-range connections, and hence an unsynchronized population state appears. However, with increasing \( p \), long-range connections begin to appear, and hence the average path length becomes shorter. Consequently, when passing a critical value \( p^*_c (>0.12) \), the unsynchronized state is destabilized and then fast sparsely synchronized population rhythm emerges. This transition to fast sparse synchronization is well described by using a realistic “thermodynamic” order parameter, based on the instantaneous population spike rate (IPSR) [63]. In order
to further investigate the effect of geometrical long-range connections on dynamical spatial correlation length for occurrence of population synchronization, we also make a dynamical-correlation analysis between neuronal pairs. It is thus found that the spatial correlation length for $p < p^*_c$ is so small that global synchronization cannot occur. On the other hand, for $p > p^*_c$, the correlation length is found to cover the whole system thanks to sufficient number of long-range connections, and consequently global synchronization appears in the whole population. Furthermore, the degree of fast sparse synchronization is well measured by employing a realistic “statistical–mechanical” spiking measure, based on IPSR [63]. At an optimal value $p^*_c$ ($\approx 0.26$), a dynamical-efficiency factor given by the ratio of the synchronization degree to the (axon) wiring cost is found to become maximal. Thus, an optimal fast sparsely-synchronized rhythm is found to appear at a minimal wiring cost in an economic small-world network. Finally, a summary is given in Section 4.

2. Inhibitory network of suprathreshold FS Izhikevich interneurons

A neural circuit in the major parts of the brain consists of a few types of excitatory principal cells and diverse types of inhibitory interneurons. By providing a synchronous oscillatory output to the principal cells, interneuronal networks play the role of the backbones of many brain rhythms [1,2,12,32]. We consider an inhibitory population of $N$ sparsely-coupled neurons equidistantly placed on a one-dimensional ring of radius $N/2\pi$. As an element in our neural system, we choose the FS Izhikevich interneuron model [24–27]. The population dynamics in this neural network is governed by the following set of ordinary differential equations:

$$\frac{d v_i}{dt} = k(v_i - v_r)(v_i - v_t) - u_i + I_{DC} + D\xi_i - I_{syn,i},$$

$$\frac{du_i}{dt} = a[U(v_i) - u_i], \quad i = 1, \ldots, N,$$

with the auxiliary after-spike resetting:

if $v_i \geq v_p$, then $v_i \leftarrow c$ and $u_i \leftarrow u_i + d$,

where

$$U(v) = \begin{cases} 0 & \text{for } v < v_b, \\ b(v - v_b)^3 & \text{for } v \geq v_b, \end{cases}$$

$$I_{syn,i} = \frac{f}{d^{N}} \sum_{j(i) \neq i} w_{ij} s_{j}(t)(v_i - V_{syn}),$$

$$s_{j}(t) = \sum_{f=1}^{f_i} E(t - t_{f}^{(j)} - \tau_{i}); \quad E(t) = \frac{1}{\tau_{d} - \tau_{r}} \left( e^{-t/\tau_{d}} - e^{-t/\tau_{r}} \right) \Theta(t).$$

Here, the state of the $i$th neuron at a time $t$ is characterized by two state variables: the membrane potential $v_i$ and the recovery current $u_i$. In Eq. (1), $C$ is the membrane capacitance, $v_r$ is the resting membrane potential, and $v_t$ is the instantaneous threshold potential. After the potential reaches its apex (i.e., spike cutoff value) $v_p$, the membrane potential and the recovery variable are reset according to Eq. (3). The units of the capacitance $C$, the potential $v$, the current $u$ and the time $t$ are pF, mV, pA, and ms, respectively.

Unlike Hodgkin–Huxley-type conductance-based models, the Izhikevich model matches neuronal dynamics by tuning the parameters instead of matching neuronal electrophysiology. The parameters $k$ and $b$ are associated with the neuron's rheobase and input resistance, $a$ is the recovery time constant, $c$ is the after-spike reset value of $v$, and $d$ is the total amount of outward minus inward currents during the spike and affecting the after-spike behavior (i.e., after-spike jump value of $u$). Tuning these parameters, the Izhikevich neuron model may produce 20 of the most prominent neuro-computational features of cortical neurons [24–27]. Here, we use the parameter values for the FS interneurons (which do not fire post-inhibitory rebound spikes) in the layer 5 Rat visual cortex [26]: $C = 20$, $v_r = -55$, $v_t = -40$, $v_p = 25$, $v_b = -55$, $k = 1$, $a = 0.2$, $b = 0.025$, $c = -45$, $d = 0$.

Each Izhikevich interneuron is stimulated by using the common DC current $I_{DC}$ (measured in units of pA) and an independent Gaussian white noise $\xi_i$ (see the 3rd and the 4th terms in Eq. (1)) satisfying $\langle \xi_i(t) \rangle = 0$ and $\langle \xi_i(t) \xi_j(t') \rangle = \delta_{ij} \delta(t - t')$, where $\langle \cdots \rangle$ denotes the ensemble average. The noise $\xi_i$ is a parametric one that randomly perturbs the strength of the applied current $I_{DC}$, and its intensity is controlled by using the parameter $D$ (measured in units of pA ms$^{1/2}$). In the absence of noise (i.e., $D = 0$), the Izhikevich interneuron exhibits a jump from a resting state to a spiking state via subcritical Hopf bifurcation for $I_{DC, h} = 73.7$ by absorbing an unstable limit cycle born via a fold limit cycle bifurcation for $I_{DC, l} = 72.8$. Hence, the Izhikevich interneuron shows type-II excitability because it begins to fire with a non-zero frequency, as shown in Fig. 1(a1) [64,65]. As $I_{DC}$ is increased from $I_{DC, h}$, the mean firing rate $f$ increases monotonically (see Fig. 1(a2)). Throughout this paper, we consider a suprathreshold case of $I_{DC} = 1500$. For this case, the membrane potential $v$ oscillates very fast with $f = 633$ Hz, as shown in Fig. 1(b).

The last term in Eq. (1) represents the synaptic coupling of the network. $I_{syn,i}$ of Eq. (5) represents a synaptic current injected into the $i$th neuron. The synaptic connectivity is given by the connection weight matrix $W (\equiv \{w_{ij}\})$ where $w_{ij} = 1$.
if the neuron $j$ is presynaptic to the neuron $i$; otherwise, $w_{ij} = 0$. Here, the synaptic connection is modeled by using both the conventional Erdős–Rényi random graph and the Watts–Strogatz small-world network. Then, the in-degree of the $i$th neuron, $d^m_i$ (i.e., the number of synaptic inputs to the neuron $i$) is given by $d^m_i = \sum_{j \neq i} w_{ij}$. The fraction of open synaptic ion channels at time $t$ is denoted by $s(t)$. The time course of $s_i(t)$ of the $j$th neuron is given by a sum of delayed double-exponential functions $E(t - t_f^{(j)} - \tau_i)$ (see Eq. (6)), where $\tau_i$ is the synaptic delay, and $t_f^{(j)}$ and $F_j$ are the $j$th spike and the total number of spikes of the $j$th neuron at time $t$, respectively. Here, $E(t)$ (which corresponds to contribution of a presynaptic spike occurring at time 0 to $s(t)$ in the absence of synaptic delay) is controlled by the two synaptic time constants: synaptic rise time $\tau_r$ and decay time $\tau_d$, and $\Theta(t)$ is the Heaviside step function: $\Theta(t) = 1$ for $t \geq 0$ and $0$ for $t < 0$. For the inhibitory GABAergic synapse (involving the GABA_A receptors), $\tau_i = 1$ ms, $\tau_r = 0.5$ ms, and $\tau_d = 5$ ms [18]. The coupling strength is controlled by the parameter $J$ (measured in units of $\mu$S), and $V_{syn}$ is the synaptic reversal potential. Here, we use $V_{syn} = -80$ mV for the inhibitory synapse.

Numerical integration of Eqs. (1)–(2) is done using the Heun method [66] (with the time step $\Delta t = 0.01$ ms). For each realization of the stochastic process, we choose a random initial point $[u_i(0), u_i(0)]$ for the $i$th ($i = 1, \ldots, N$) neuron with uniform probability in the range of $u_i(0) \in (-50, -45)$ and $u_i(0) \in (10, 15)$.

3. Effect of small-world connectivity on fast sparsely synchronized rhythms

In this section, we study the effect of network architecture on fast sparsely synchronized rhythms with stochastic and intermittent neural discharges. For modeling the complex connectivity in neural systems, we first use the conventional Erdős–Rényi random network of suprathreshold FS Izhikevich interneurons [28], and study occurrence of population oscillatory states by varying the inhibition intensity $J$ and the noise intensity $D$. But, in a real cortical circuit, synaptic connections are known to be neither regular nor completely random. Hence, we consider the Watts–Strogatz model for the small-world network which interpolates between the regular lattice and the random graph [40], and investigate the effect of small-world connectivity on fast sparsely synchronized rhythms by varying the rewiring probability $p$ for fixed values of $J$ and $D$. Particularly, we search for an optimal fast sparse synchronization occurring at a minimal wiring cost in an economic small-world network.

We first consider the conventional Erdős–Rényi random graph of $N$ sparsely-connected suprathreshold FS Izhikevich interneurons equidistantly placed on a one-dimensional ring of radius $N/2\pi$. A postsynaptic neuron $i$ receives a synaptic input from another presynaptic neuron $j$ with a connection probability $P_{syn} (= M_{syn}/N)$, where $M_{syn}$ is the average number of synaptic inputs per neuron (i.e., $M_{syn} = \langle d_i \rangle$; $d_i$ is the number of synaptic inputs to the neuron $i$ and $\langle \cdots \rangle$ denotes an ensemble-average over all neurons). Here, we consider a sparse case of $M_{syn} = 50$. By varying the inhibition intensity $J$ and the noise intensity $D$, we investigate occurrence of population synchronized states. In computational neuroscience, an ensemble-averaged global potential,

$$V_G(t) = \frac{1}{N} \sum_{i=1}^{N} v_i(t),$$

\(\text{(7)}\)
is often used for describing emergence of population neural synchronization. However, to directly obtain \( V_c \) in real experiments is very difficult. To overcome this difficulty, instead of \( V_c \), we use an experimentally-obtainable IPSR which is often used as a collective quantity showing population behaviors [2,13–18]. The IPSR is obtained from the raster plot of neural spikes which is a collection of spike trains of individual neurons. Such raster plots of spikes, where population spike synchronization may be well visualized, are fundamental data in experimental neuroscience. For the synchronous case, “stripes” (composed of spikes and indicating population synchronization) are found to be formed in the raster plot. Hence, for a synchronous case, an oscillating IPSR appears, while for an unsynchronized case the IPSR is nearly stationary. To obtain a smooth IPSR, we employ the kernel density estimation (kernel smoother) [67]. Each spike in the raster plot is convoluted (or blurred) with a kernel function \( K_h(t) \) to obtain a smooth estimate of IPSR, \( R(t) \):

\[
R(t) = \frac{1}{N} \sum_{i=1}^{N} \sum_{j=1}^{n_i} K_h(t - t_i^{(0)}),
\]

where \( t_i^{(0)} \) is the \( i \)th spiking time of the \( j \)th neuron, \( n_i \) is the total number of spikes for the \( i \)th neuron, and we use a Gaussian kernel function of bandwidth \( h \):

\[
K_h(t) = \frac{1}{\sqrt{2\pi}h} e^{-t^2/2h^2}, \quad -\infty < t < \infty.
\]

We first study the case of \( D = 0 \). For small \( J \), individual interneurons fire too fast to be synchronized. Fig. 2(a1) shows the raster plot of spikes and the IPSR kernel estimate \( R(t) \) for an unsynchronized case of \( J = 10 \). Spikes in the raster plot are completely scattered and hence \( R(t) \) is nearly stationary. However, as \( J \) is increased mean firing rates, \( f_i \), of individual interneurons decrease, and full synchronization occurs when \( J \) passes a critical value \( J^* \) (\( \geq 12 \)). For a synchronous case of \( J = 100 \), clear stripes (composed of spikes and indicating population spike synchronization) are formed in the raster plot, and hence \( R(t) \) shows regular oscillation with population frequency \( f_p = 197 \) Hz, as shown in Fig. 2(a2). The interspike interval (ISI) histogram with a single peak appearing at 5.1 ms is shown in Fig. 2(a3), and hence individual neurons fire regularly with mean firing rate \( f_i \) which is the same as \( f_p \). Thus, complete full synchronization with \( f_p = f_i \) occurs for \( J = 100 \). We next consider the effect of noise on the full synchronization for a fixed \( J \). As \( D \) is increased, the full synchronization for \( D = 0 \) evolves, depending on the values of \( J \), and eventually desynchronization occurs when passing a critical value \( D^* \). Fig. 2(b) shows the state diagram in the \( J-D \) plane. For the full synchronization, mean firing rates, \( f_i \), of individual neurons are the same as the population frequency \( f_p \), while for the partial and sparse synchronization, \( f_i \) is less than \( f_p \) (i.e., individual neurons fire at lower rates than the population frequency). For the sparsely synchronized cortical rhythms, \( R_i = f_i \sim 4 : 1 \) [13–16]. Hence, when the population frequency is much higher than the mean firing rate of individual interneurons \( f_p > 4 \) \( f_i \), the synchronization will be referred to as sparse synchronization. Plots of \( f_p \) and \( f_i \) versus \( D \) are shown in Fig. 2(c1)–(c4) for \( J = 100, 400, 1400, \) and 2000. For small \( J (J^* < J < 162) \), only the full synchronization occurs because \( f_p = f_i \) (e.g., see the case of \( J = 100 \)). However, for \( J > 162 \), the full synchronization is developed into partial synchronization at some threshold value \( D_{th} \) via pitchfork-like bifurcations (e.g., see the cases of \( J = 400, 1400, \) and 2000). With increasing \( J \), the difference between \( f_p \) and \( f_i \) increases abruptly when passing \( D_{th} \). For \( J > 1275 \), the partial synchronization evolves into sparse synchronization with \( f_p > 4f_i \) (e.g., see the cases of \( J = 1400 \) and 2000).

For further understanding of the full and the partial synchronization, we present two explicit examples showing how the full synchronization is evolved into unsynchronized states as \( D \) is increased. For \( J^* < J < 162 \), the full synchronization occurs for \( D = 0 \) develops directly into an unsynchronized state without any other type of intermediate synchronization stage. As an example consider the case of \( J = 100 \). The raster plots and the IPSR kernel estimate \( R(t) \) for various values of \( D \) are given in Fig. 3(a1)–(a4). As \( D \) is increased, stripes of spikes in the raster plot become more and more smeared, and hence the amplitudes of \( R(t) \) become smaller (i.e., the pacing degree of spikes decreases). When passing a critical value \( D^* \approx 173 \), stripes become overlapped and \( R(t) \) becomes nearly stationary. Thus, a transition to an unsynchronized state occurs (e.g., see Fig. 3(a4)). The ISI distributions for various values of \( D \) are also shown in Fig. 3(b1)–(b4). As \( D \) is increased from 0, the height of the ISI histogram becomes smaller and its width becomes wider. During this process, \( f_p = f_i \approx 197 \) Hz, as shown in Fig. 2(c1). Thus, as \( D \) passes a critical value \( D^* \), a direct transition from full synchronization to an unsynchronized state occurs for \( J = 100 \). For \( J > 162 \), with increasing \( D \) the full synchronization for \( D = 0 \) evolves into the partial synchronization with \( f_p > f_i \). As an example, we consider the case of \( J = 1400 \). Fig. 3(c1)–(c4) show the raster plots and the IPSR kernel estimate \( R(t) \) for various values of \( D \). For \( 0 < D < D_{th} \) (\( \approx 144 \)), full synchronization with \( f_p = f_i \) occurs (see Fig. 2(c3)). As \( D \) is increased from 0 to \( D_{th} \), the degree of full synchronization decreases because the stripes of the raster plot become smeared, and hence the amplitude of \( R(t) \) also becomes smaller (see Fig. 3(c1)). As the case of \( J = 100 \), the width of the ISI histogram becomes wider due to decrease in the pacing degree (see Fig. 3(d1)). However, for \( D > 144 \), partial synchronization with \( f_p > f_i \) appears via a pitchfork-like bifurcation, as shown in Fig. 2(c3). As \( D \) is increased from \( D = 144 \), the interval between stripes of the raster plot becomes smaller, and hence the population frequency \( f_p \) increases (see Fig. 3(c2)). For this case, the stripes of the raster plot become smeared, and hence the pacing degree of spikes decreases. Furthermore, the density of stripes becomes smaller because smaller fraction of total neurons fire in each stripes. Thus, with increasing \( D \) from \( D = 144 \) both the pacing and the occupation degrees of spikes decrease, and consequently a large decrease in the amplitude of \( R(t) \) occurs. In contrast to the case of full synchronization, the ISI histogram has multiple peaks appearing at multiples of the period \( T_e \) of \( R(t) \) (see Fig. 3(d2)).
Fig. 2. Erdös–Rényi random graph of $N = 10^3$ suprathreshold FSI Izhikevich interneurons for $I_{\text{DC}} = 1500$ and $M_{\text{syn}} = 50$. Raster plots of spikes and plots of the IPSR kernel estimate $R(t)$ versus $t$ for (a1) $J = 10$ and (a2) $J = 100$; the bandwidth of the Gaussian kernel estimate is 1 ms. (a3) ISI histogram for $J = 100$ (ISI histogram is composed of $5 \times 10^4$ ISIs and the bin size for the histogram is 0.5 ms). (b) State diagram in the $J$–$D$ plane. For the full synchronization, the individual frequency $f_i$ is the same as the population frequency $f_p$, while for the partial and sparse synchronization, $f_i$ is less than $f_p$. Particularly, the cases of $f_p > 4f_i$ are referred to as the sparse synchronization. Plots of $f_p$ and $f_i$ versus $D$ for (c1) $J = 100$, (c2) $J = 400$, (c3) $J = 1400$, and (c4) $J = 2000$. Here, the circles and crosses denote $f_p$ and $f_i$, respectively.

have also been found in networks of coupled inhibitory neurons in the presence of noise where noise-induced hopping from one cluster to another one occurs [68], in single noisy neuron models exhibiting stochastic resonance due to a weak periodic external force [69,70], and in inhibitory networks of coupled subthreshold neurons showing stochastic spiking coherence [21–23]. Stochastic spike skipping in coupled systems is a collective effect because it occurs due to a driving by a coherent ensemble-averaged synaptic current, in contrast to the single case driven by a weak periodic force where stochastic resonance occurs. Due to this stochastic spike skipping, partial occupation occurs in the stripes of the raster plot. Thus, the mean firing rates $f_i$ of individual interneurons become less than the population frequency $f_p$, and hence partial synchronization occurs. Particularly, for $D > 448$, sparse synchronization with $f_p > 4f_i$ appears. As $D$ is further increased from $D = 448$, both the pacing and the occupation degrees of spikes (seen in the raster plot) decrease and multiple peaks in the ISI histogram overlap (see Fig. 3(c3) and (d3)). Eventually, when passing a critical value $D^* \simeq 741$, an unsynchronized state appears (e.g. see the case of $D = 800$).

As shown in the state diagram of Fig. 2(b), fast sparsely synchronized rhythms appear when both the inhibition strength and the noise intensity are strong in the Erdös–Rényi random graph of suprathreshold FSI Izhikevich interneurons. For random connectivity, the average path length is short due to long-range connections, and hence global efficiency of information transfer becomes high [29,30]. However, unlike the regular lattice, the random network has poor clustering, and it becomes non-economic due to appearance of short-range and long-range connections with equal probability [31,32]. Real synaptic connectivity is known to have complex topology which is neither regular nor completely random [31–38]. To study the effect of network structure on fast sparsely synchronized oscillations, we consider the Watts–Strogatz model
for small-world networks which interpolates between regular lattice and random graph via rewiring [40]. By varying the rewiring probability $p$ from local to long-range connection, we investigate the effect of small-world connectivity on fast sparse synchronization for fixed values of $J = 1400$ and $D = 500$. We start with a directed regular ring lattice with $N$ suprathreshold FS Izhikevich interneurons where each Izhikevich interneuron is coupled to its first $M_{\text{syn}}$ neighbors ($M_{\text{syn}}/2$ on either side) via outward synapses, and rewire each outward connection at random with probability $p$ such that self-connections and duplicate connections are excluded. As in the above random case, we consider a sparse but connected network with a fixed value of $M_{\text{syn}} = 50$. Then, we can tune the network between regularity ($p = 0$) and randomness ($p = 1$); the case of $p = 1$ corresponds to the above Erdős–Rényi random graph. In this way, we investigate emergence of fast sparsely synchronized rhythm in the directed Watts–Strogatz small-world network of $N$ suprathreshold FS Izhikevich interneurons by varying the rewiring probability $p$ for $J = 1400$ and $D = 500$.

The topological properties of the small-world connectivity have been well characterized in terms of the clustering coefficient and the average path length [40]. The clustering coefficient, denoting the cliquishness of a typical neighborhood in the network, characterizes the local efficiency of information transfer, while the average path length, representing the typical separation between two vertices in the network, characterizes the global efficiency of information transfer. The regular lattice for $p = 0$ is highly clustered but large world where the average path length grows linearly with $N$, while the random graph for $p = 1$ is poorly clustered but small world where the average path length grows logarithmically with $N$ [40]. As soon as $p$ increases from zero, the average path length decreases dramatically, which leads to occurrence of a small-world phenomenon which is popularized by the phrase of the “six degrees of separation” [43,44]. However, during
Fig. 4. Watts–Strogatz small-world network of suprathreshold FSIzhikevich interneurons for \(I_{\text{DC}} = 1500, M_{\text{syn}} = 50, J = 1400, \) and \(D = 500\). (a) Plots of \(\log_{10} \Theta\) versus \(p\). Unsynchronized state for \(p = 0\); raster plots of spikes and plots of the IPSR kernel estimate \(R(t)\) versus \(t\) for \(p = 0\). (b) \(N = 10^3\). Synchronized state for \(p = 0.25\); raster plots of spikes and plots of the IPSR kernel estimate \(R(t)\) versus \(t\) for \(p = 0.25\) for \(N = 10^3\). The band width of the Gaussian kernel estimate is 1 ms.

this dramatic drop in the average path length, the clustering coefficient remains almost constant at its value for the regular lattice. Consequently, for small \(p\) small–world networks with short path length and high clustering emerge [40].

As is well known, a conventional order parameter, based on the ensemble-averaged global potential \(V_c\), is often used for describing transition from asynchrony to synchrony in computational neuroscience [71–73]. Recently, instead of \(V_c\), we used an experimentally-obtainable IPSR kernel estimate \(R(t)\), and developed a realistic order parameter, which may be applicable in both the computational and the experimental neuroscience [63]. The mean square deviation of \(R(t)\),

\[
\Theta \equiv \langle R(t) - \langle R(t) \rangle \rangle^2,
\]

plays the role of an order parameter \(\Theta\). (Here the overbar represents the time average.) The order parameter may be regarded as a thermodynamic measure because it concerns just the macroscopic IPSR kernel estimate \(R(t)\) without any consideration between \(R(t)\) and microscopic individual spikes. In the thermodynamic limit of \(N \to \infty\), the order parameter \(\Theta\) approaches a non-zero (zero) limit value for the synchronized (unsynchronized) state. Fig. 4(a) shows a plot of the order parameter versus the rewiring parameter \(p\). For \(p < p_c^*\) (>0.12), unsynchronized states exist because the order parameter \(\Theta\) tends to zero as \(N \to \infty\). As \(p\) passes the critical value \(p_c^*\), a transition to synchronization occurs because the values of \(\Theta\) become saturated to non-zero limit values for \(N \geq 3 \cdot 10^7\). These synchronized states seem to appear because global efficiency of information transfer between distant neurons for \(p > p_c^*\) becomes enough for occurrence of population synchronization. Here we present two explicit examples for the synchronized and unsynchronized states. First, we consider the population state in the regular lattice for \(p = 0\). As shown in Fig. 4(b1) for \(N = 10^3\), the raster plot shows a zigzag pattern intermingled with inclined partial stripes of spikes with diverse inclinations and widths, and \(R(t)\) is composed of coherent parts with regular large-amplitude oscillations and incoherent parts with regular small-amplitude fluctuations. For \(p = 0\), the clustering coefficient is high, and hence partial stripes (indicating local clustering of spikes) seem to appear in the raster plot of spikes. As \(N\) is increased to \(10^4\), partial stripes become more inclined from the vertical, and hence spikes become more difficult to keep pace with each other. As a result, \(R(t)\) shows noisy fluctuations with smaller amplitudes, as shown in Fig. 4(b2). Hence the population state for \(p = 0\) seems to be unsynchronized because \(R(t)\) tends to be nearly stationary as \(N\) increases to the infinity. As \(p\) is increased from 0, long-range short-cuts begin to appear, and hence average path length becomes shorter. Eventually, when passing the critical value \(p_c^*\), synchronized population state emerges because of sufficient global efficiency of information transfer between distant neurons, which will be discussed below in more detail. As a second example, we consider a synchronized case of \(p = 0.25\). For \(N = 10^3\), the degree of zigzaggness for partial stripes in the raster plot is much reduced when compared with the \(p = 0\) case, and hence \(R(t)\) shows a regular oscillation, as shown in Fig. 4(c1). Its amplitudes are much larger than that for the case of \(p = 0\), although there is a little variation in the amplitude. As \(N\) is increased to \(N = 10^4\), \(R(t)\) shows regular oscillations, and the amplitudes in each oscillating cycle are nearly the same, in contrast to the case of \(N = 10^3\) (see Fig. 4(c2)). Hence, \(R(t)\) displays more regular oscillations with nearly the same amplitudes for \(N = 10^4\). Consequently, the population state for \(p = 0.25\) seems to be synchronized because \(R(t)\) tends to show regular oscillations as \(N\) goes to the infinity.

In order to further understand the above transition to global spike synchronization, we investigate the effect of geometrical long-range connections on the dynamical cross-correlations between neuronal pairs. As examples, we reconsider the same cases of \(p = 0\) and 0.25 as in Fig. 4. Fig. 5(a1) shows a raster plot of spikes in the longer time interval (where the horizontal structure may be more clearly seen than the vertical zigzag structure when compared with the raster
Fig. 5. Watts-Strogatz small-world network of suprathreshold FSI Izhikevich interneurons for $I_{\text{DC}} = 1500$, $M_{\text{syn}} = 50$, and $D = 500$. Unsynchronized state for $p = 0$ and $J = 1400$: (a1) raster plots of spikes and MFR distribution of individual neurons for $N = 10^3$ and the spatial correlation function $C_L$ for $N = (a2) 10^3$ and (a3) $10^4$. For this case of $J = 1400$, the MFR distribution exhibits spatially modulated oscillation. Unsynchronized state for $p = 0$ and $J = 200$: (b1) raster plots of spikes and MFR distribution of individual neurons for $N = 10^3$ and the spatial correlation function $C_L$ for $N = (b2) 10^3$ and (b3) $10^4$. The MFR distribution for $J = 200$ shows noisy fluctuation around the average MFR ($\leq 140$ Hz) without any regular spatial oscillation. Hereafter, the values of $J$ in (c1)–(b8) are fixed at $J = 1400$. Synchronized state for $p = 0.25$: (c1) raster plots of spikes and MFR distribution of individual neurons for $N = 10^3$ and the spatial correlation function $C_L$ for $N = (c2) 10^3$ and (c3) $10^4$. Spatial correlation functions $C_L$ for unsynchronized states for various values of $p$ when (d1)–(d3) $N = 10^3$ and (d4)–(d6) $10^4$. Spatial correlation functions $C_L$ for synchronized states for various values of $p$ when (e1)–(e3) $N = 10^3$ and (e4)–(e6) $10^4$. Flat spatial correlation functions $C_L$ for synchronized states for various values of $p$ when (f1)–(f4) $N = 10^3$ and (f5)–(f8) $10^4$. The bandwidth of the Gaussian kernel estimate is 1 ms. The temporal cross-correlation $C_{ij}(\tau)$ is obtained through average over 20 realizations, and the number of data used for the calculation of each temporal cross-correlation function $C_{ij}(\tau)$ is $2 \times 10^4$. 
plot (in the shorter time interval) in Fig. 4(b1)) and the distribution of the mean firing rates (MFRs) of individual neurons for the case of \( p = 0 \) (without long-range connections). Dense and sparse horizontal stripes of spikes appear alternatively in the raster plot with the average spatial period \( \tau_{L} \approx 36 \). In accordance to this structure of the horizontal stripes, the MFR distribution also shows a regular oscillation with large amplitude around the average MFR \( \bar{\nu} \approx 33 \text{ Hz} \): the peak (valley) regions of the MFR correspond to the centers of the dense (sparse) stripes. This spatial oscillation of the MFR distribution affects the behavior of the spatial cross-correlations between neuronal pairs. For obtaining dynamical pair cross-correlations, each spike train of the \( i \)-th neuron is convoluted with a Gaussian kernel function \( K_{h}(t) \) of band width \( h \) to get a smooth estimate of instantaneous individual spike rate (IISR), \( r_{i}(t) \):

\[
r_{i}(t) = \sum_{s=1}^{n_{i}} K_{h}(t - t_{i}^{(s)}),
\]

where \( t_{i}^{(s)} \) is the \( s \)-th spiking time of the \( i \)-th neuron, \( n_{i} \) is the total number of spikes for the \( i \)-th neuron, and \( K_{h}(t) \) is given in Eq. (9). Then, the normalized temporal cross-correlation function \( C_{ij}(\tau) \) between the IISRs \( r_{i}(t) \) and \( r_{j}(t) \) of the \((i,j)\) neuronal pair is given by:

\[
C_{ij}(\tau) = \frac{\Delta r_{i}(t + \tau) \Delta r_{j}(t)}{\sqrt{\Delta r_{i}^{2}(t) \Delta r_{j}^{2}(t)}},
\]

where \( \Delta r_{i}(t) = r_{i}(t) - \bar{r}_{i} \) and the overline denotes the time average. We now introduce the spatial cross-correlation \( C_{L}(L = 1, \ldots, N/2) \) between neuronal pairs separated by a spatial distance \( L \) through average of all the temporal cross-correlations between \( r_{i}(t) \) and \( r_{i+L}(t) \) \((i = 1, \ldots, N)\) at the zero-time lag:

\[
C_{L} = \frac{1}{N} \sum_{i=1}^{N} C_{i,i+L}(0) \quad \text{for } L = 1, \ldots, N/2.
\]

Fig. 5(a2) shows the plot of the spatial cross-correlation \( C_{L} \) versus \( L \) for \( N = 10^{3} \) in the case of \( p = 0 \). We note that the spatial cross-correlation function \( C_{L} \) exhibits a "damped" oscillation with respect to \( L \). Local maxima (minima) of \( C_{L} \) come from cross-correlations when the distances \( L \) between neuronal pairs are multiples (odd multiples) of the spatial period \( \tau_{L} \) (half-period \( \tau_{L}/2 \)) of the MFR. In this way, \( C_{L} \) makes an oscillatory decay to zero. To obtain the exponential decay rate of \( C_{L} \), the maximal envelope (denoted by a dotted line) may be well fitted with an exponential function with a characteristic correlation \( \eta \):

\[
C_{L} = A \cdot e^{-L/\eta}; \quad A = 0.31, \quad \eta = 143.
\]

Then, one can think that the whole system is composed of independent partially synchronized blocks of size \( \eta \). To examine occurrence of population synchronization, we increase the number of neurons as \( N = 10^{4} \). For this case, the spatial cross-correlation function \( C_{L} \) in Fig. 5(a3) also shows a damped oscillation, and it decays to zero. Then, the maximal envelope also is well fitted with the same exponential function \( (C_{L} = A \cdot e^{-L/\eta}; A = 0.31, \eta = 143) \). Hence, the correlation length \( \eta \) remains unchanged, although the system size is increased 10 times. As a criterion for occurrence of synchronization, we introduce a normalized correlation length \( \tilde{\eta} \):

\[
\tilde{\eta} = \frac{\eta}{N},
\]

which represents the ratio of the correlation length \( \eta \) to the system size \( N \). As \( N \) is increased, \( \tilde{\eta} \) tends to zero (i.e., the relative size of partially synchronized blocks (when compared to the whole system size) tends to zero). Consequently, no global synchronization occurs for the case of \( p = 0 \). Although the population dynamical state for \( p = 0 \) is unsynchronized, it seems to be worth noting the spatially modulated MFR distribution which exhibits regular spatial oscillation for \( J = 1400 \).

To examine appearance of spatially modulated phase, we decrease the inhibitory coupling strength \( J \), and find spatially non-modulated phase for small \( J \) less than a critical value \( J_{c} \approx 540 \). Fig. 5(b1) shows an example of spatially non-modulated phase for \( J = 200 \) where the MFR distribution exhibits a noisy fluctuation around the average MFR \( \bar{\nu} \approx 140 \text{ Hz} \) without any regular oscillation. The average MFR for \( J = 200 \) is much larger than that for \( J = 1400 \) because the inhibition strength \( J \) is decreased. In this case of non-modulated phase, the spatial cross-correlation function \( C_{L} \) exhibits a direct exponential decay to zero without any oscillation, as shown in Fig. 5(b2) and (b3), unlike the case of modulated phase for \( J = 1400 \) where \( C_{L} \) shows an oscillatory decay to zero. Furthermore, the correlation length for \( J = 200 \) is \( \eta = 32 \), independently of the system size \( N \), which is smaller than that for the case of \( J = 1400 \) because the decay rate to zero for \( J = 200 \) is faster than that for \( J = 1400 \). Although the topological clustering coefficient is high for \( p = 0 \), no distinct dynamical clustering seems to occur when the inhibitory coupling strength \( J \) is small. The clustering effect seems to be distinctly evident only when \( J \) becomes sufficiently strong. Hence, when \( J \) passes the critical value \( J_{c} \), this type of spatially modulated phase seems to emerge spontaneously. Further in-depth study of this transition to spatially modulated phase is beyond our present subject, and it will be done in a future work. Then, we return to the original subject on the effect of small-world connectivity for a fixed value of \( J = 1400 \) and consider another case of \( p = 0.25 \) where long-range connections appear. The raster plot of spikes
and the MFR distribution are shown in Fig. 5(c1). Some of dense horizontal stripes become smeared and overlap vertically with neighboring dense stripes, and then some parts of sparse horizontal stripes between merging dense stripes disappear. The MFR distribution shows irregular oscillation with relatively small dispersion about the average MFR ($\approx 33$ Hz) (when compared with the MFR distribution for $p = 0$). Unlike the case of $p = 0$, the spatial correlation function $C_t$ for $N = 10^3$ becomes nearly non-zero constant ($\approx 0.045$) in the whole range of $L$, as shown in Fig. 5(c2), and hence the correlation length $\eta$ becomes $N/2 (=500)$ covering the whole system (note that the maximal distance between neurons is $N/2$ because of the ring architecture on which neurons exist). Consequently, the whole system is composed of just one single synchronized block (i.e., global synchronization occurs in the whole system), in contrast to the case of $p = 0$. For $N = 10^4$, the flatness of $C_t$ in Fig. 5(c3) also extends to the whole range ($L = N/2 = 5000$), and the correlation length becomes $\eta = 5000$, which also covers the whole system. Hence, the number of long-range connections for $p = 0.25$ seems to become enough to make the correlation length $\eta$ cover the whole system, independently of $N$. Consequently, as $N$ is increased, the normalized correlation length $\hat{\eta}$ has a non-zero limit value, $1/2$, and global synchronization emerges in the whole population, in contrast to the case of $p = 0$. In addition to the above cases of $p = 0$ and 0.25, we also make an extensive dynamical-correlation analysis for several values of $p$ in the subcritical and the supercritical subregions of $p$. Fig. 5(d1)–(d6) shows the spatial correlation functions $C_t$ in the subcritical cases of $p = 0.05, 0.08, and 0.11$ (less than the critical values $p_c^*$ ($\approx 0.12$)). For $N = 10^3$, $C_t$ makes an oscillatory decay to zero. As $p$ is increased, the decay rate becomes slower, and hence the correlation length $\eta$ becomes longer, as shown in Fig. 5(d1)–(d3). When the system size is increased to $N = 10^4$, $C_t$ also exhibits an oscillatory decay to zero in the same way as the case of $N = 10^3$ (see Fig. 5(d4)–(d6)). Hence, the correlation length $\eta$ becomes the same, independently of $N$, and the normalized correlation length $\hat{\eta}$ ($=\eta/N$) tends to zero in the thermodynamic limit of $N \to \infty$ due to the insufficient number of long-range connections. As a result, no global synchronization occurs for the subcritical case. On the other hand, in the supercritical case of $p = 0.13, 0.14, and 0.15$ (larger than $p_c^*$), $C_t$ seems to make an oscillatory decay to a non-zero limit, as shown in Fig. 5(e1)–(e3), and the convergence rate to non-zero limit becomes faster with increasing $p$. As $N$ is increased, this kind of convergence to non-zero limit in $C_t$ may be more clearly seen (e.g., see Fig. 5(e4)–(e6) for $N = 10^4$). As $p$ is further increased, the oscillatory part in $C_t$ disappears gradually, and then $C_t$ becomes nearly positive constant, as shown in Fig. 5(f1)–(f8) for both $N = 10^3$ and $10^4$. Hence, unlike the subcritical case, with increasing $N$ the correlation length $\eta$ increases as $N/2$, covering the whole system, and the normalized correlation length $\hat{\eta}$ has a non-zero limit value, $1/2$, in the thermodynamic limit of $N \to \infty$. Consequently, global population synchronization occurs for the supercritical case because $\eta$ covers the whole system thanks to sufficient number of long-range connections. We also note that, with increasing $p$ the constant values of $C_t$ increase, as shown in Fig. 5(f1)–(f8). However, their values become saturated for $p = p_{max} (=0.4)$ (i.e., $C_t = 0.075$ for $p \geq p_{max}$), because long-range connections which appear up to $p_{max}$ seem to play a sufficient role for obtaining the maximal degree of pair cross-correlations.

We now study the population and individual behaviors of synchronized states for various values of $p > p_c^*$. Through comparison of the population behaviors with individual behaviors, one can understand fast sparsely synchronized states well. With increasing $p$ the zigzagness degree of partial spikes in the raster plots of spikes becomes reduced (see Fig. 6(a1)–(a5)), and eventually for $p = p_{max} (=0.4)$, the raster plot becomes composed of vertical stripes without zigzag, and then the pacing degree between spikes becomes nearly the same. Hence, the amplitude of the IPSR kernel estimate $R(t)$ increases up to $p_{max}$, and then its value becomes saturated. For these values of $p$, $R(t)$ shows regular oscillation with the population frequency $f_p = 147$ Hz, corresponding to the ultrafast rhythm (100–200 Hz). In contrast to population rhythm, individual neurons make stochastic and sparse discharges as Geiger counters. We collect $5 \times 10^5$ ISIs from all individual neurons and get the ISI histograms which are shown in Fig. 6(b1)–(b5). Multiple peaks appear at multiples of the period $T_c (=1/f_p \approx 6.8$ ms)
of $R(t)$. Hence, individual neurons exhibit stochastic phase locking leading to stochastic spike skipping (i.e., intermittent spikings phase-locked to $R(t)$ at random multiples of the period of $R(t)$). For these values of $p$, mean firing rates $f_i$ of individual neurons, corresponding to the inverse of the average ISI, is 33 Hz, and hence each neuron makes an average firing very sparsely once during 4.5 population cycles. Consequently, for $p > p^* \approx 22$ fast sparsely synchronized rhythms emerge.

By varying $p$ in the whole range of fast sparse synchronization, we also measure the degree of fast sparsely synchronized rhythms in terms of a realistic statistical–mechanical spiking measure $M_i$, based on the IPSR kernel estimate $R(t)$, which was developed in our recent work [63]. As shown in Fig. 6(a1)–(a5), population spike synchronization may be well visualized in a raster plot of spikes. For a synchronized case, the raster plot is composed of partially-occupied stripes (indicating sparse development in our recent work [63]). As shown in Fig. 6(a1)–(a5), populationspike synchronization may be well visualized by $M_i$ of spikes in the stripes [63]. The spiking measure $M_i$ of the $i$th stripe is defined by the product of the occupation degree $O_i$ of spikes (representing the density of the $i$th stripe) and the pacing degree $P_i$ of spikes (denoting the smearing of the $i$th stripe):

$$M_i = O_i \cdot P_i.$$  \hspace{1cm} (16)

The occupation degree $O_i$ in the $i$th stripe is given by the fraction of spiking neurons:

$$O_i = \frac{N^{(s)}_i}{N},$$  \hspace{1cm} (17)

where $N^{(s)}_i$ is the number of spiking neurons in the $i$th stripe. For sparse synchronization, $O_i \ll 1$, while $O_i = 1$ for full synchronization. The pacing degree $P_i$ of each microscopic spike in the $i$th stripe can be determined in a statistical–mechanical way by taking into account its contribution to the macroscopic IPSR kernel estimate $R(t)$. Each global cycle of $R(t)$ begins from its left minimum, passes the central maximum, and ends at the right minimum; the central maxima coincide with centers of stripes in the raster plot (see Fig. 6(a1)–(a5)). An instantaneous global phase $\Phi(t)$ of $R(t)$ is introduced via linear interpolation in the two successive subregions forming a global cycle [63,74]. The global phase $\Phi(t)$ between the left minimum (corresponding to the beginning point of the $i$th global cycle) and the central maximum is given by

$$\Phi(t) = 2\pi(i - 3/2) + \pi \left( \frac{t - t_i^{(\text{min})}}{t_i^{(\text{max})} - t_i^{(\text{min})}} \right) \quad \text{for} \quad t_i^{(\text{min})} \leq t < t_i^{(\text{max})} \quad (i = 1, 2, 3, \ldots),$$  \hspace{1cm} (18)

and $\Phi(t)$ between the central maximum and the right minimum (corresponding to the beginning point of the $(i + 1)$th global cycle) is given by

$$\Phi(t) = 2\pi(i - 1) + \pi \left( \frac{t - t_i^{(\text{max})}}{t_{i+1}^{(\text{min})} - t_i^{(\text{max})}} \right) \quad \text{for} \quad t_i^{(\text{max})} \leq t < t_{i+1}^{(\text{min})} \quad (i = 1, 2, 3, \ldots),$$  \hspace{1cm} (19)

where $t_i^{(\text{min})}$ is the beginning time of the $i$th global cycle (i.e., the time at which the left minimum of $R(t)$ appears in the $i$th global cycle) and $t_i^{(\text{max})}$ is the time at which the maximum of $R(t)$ appears in the $i$th global cycle. Then, the contribution of the $k$th microscopic spike in the $i$th stripe occurring at the time $t_k^{(i)}$ to $R(t)$ is given by $\cos \Phi_k$, where $\Phi_k$ is the local phase at the $k$th spiking time (i.e., $\Phi_k \equiv \Phi(t_k^{(i)})$). A microscopic spike makes the most constructive (in-phase) contribution to $R(t)$ when the corresponding global phase $\Phi_k$ is $2\pi n$ ($n = 0, 1, 2, \ldots$) while it makes the most destructive (anti-phase) contribution to $R(t)$ when $\Phi_k = 2\pi(n - 1/2)$. By averaging the contributions of all microscopic spikes in the $i$th stripe to $R(t)$, we obtain the pacing degree of spikes in the $i$th stripe:

$$P_i = \frac{1}{S_i} \sum_{k=1}^{S_i} \cos \Phi_k,$$  \hspace{1cm} (20)

where $S_i$ is the total number of microscopic spikes in the $i$th stripe. By averaging $M_i$ of Eq. (16) over a sufficiently large number $N_i$ of stripes, we obtain the statistical–mechanical spiking measure $M_i$:

$$M_i = \frac{1}{N_i} \sum_{i=1}^{N_i} M_i.$$  \hspace{1cm} (21)

By varying $p$, we follow $3 \times 10^3$ stripes and characterize sparse synchronization in terms of $O_i$ (average occupation degree), $P_i$ (average pacing degree), and the statistical–mechanical spiking measure $M_i$ for 12 values of $p$ in the sparsely synchronized region, and the results are shown in Fig. 6(c1)–(c3). We note that the average occupation degree $O_i$ (denoting the average density of stripes in the raster plot) is nearly the same ($O_i \simeq 0.22$), independently of $p$; only a fraction (about 1/4.5) of the total neurons fires in each stripe (see Fig. 6(a1)–(a5)). This partial occupation in the stripes results from stochastic spike skipping of individual neurons and is seen well in the multi-peaked ISI histograms (see Fig. 6(b1)–(b5)). The average...
neuron by varying $M$ degree is given by the statistical–mechanical spiking measure synchronized rhythmmay emerge via tradeoff between the synchronization degree and the wiring cost. The synchronization dynamicalefficiency we search for optimal population rhythm emerging at a minimal wiring cost. An optimal fast sparselysynchronized rhythm is shown in Fig. 7(c). Sincethe economical small-world network has a moderate clustering coefficient $C(p_c)$, the raster plot of spikes shows a zigzag pattern due to local clustering of spikes, and the IPSR kernel estimate $K(t)$ exhibits a regular ultrafast oscillation at a population frequency $f_b$ ($=147$ Hz). In contrast to population rhythm, individualneurons fire irregularly and sparsely with $f_i = 33$ Hz as Geiger counters, as shown well in the multi-peaked ISI histogram of Fig. 7(d).

4. Summary

We have investigated the effect of network architecure on fast sparsely synchronized cortical rhythms with stochastic and intermittent neural discharges. These fast sparsely synchronized neural oscillations are in contrast to fully synchronized oscillations with regular neural discharges. For modeling of complex connections in neural systems, we first used the conventionel Erdős–Rényi random graph of suprathreshold FS Izhikevich interneurons, and studied occurrence of the population synchronized states by varying the inhibition strength $J$ and the noise intensity $D$. Fast sparsely synchronized states have been found to appear for large values of $J$ and $D$. However, real synaptic connections are known to be neither regular nor random. Hence, we considered the Watts–Strogatz model for small-world networks which interpolates between the regular lattice and the random graph, and for fixed values of $J$ and $D$ ($J = 1400$ and $D = 500$), we investigated the effect of small-world connectivity on emergence of fast sparsely synchronized rhythms by varying the rewiring probability $p$ from local to
Fig. 7. Watts–Strogatz small-world network of suprathreshold FSIzhikevich interneurons for $N = 10^3$, $I_{DC} = 1500$, $M_{syn} = 50$, $J = 1400$, and $D = 500$. (a) Statistical–mechanical spiking measure $M_s$ and normalized wiring length $L$ versus $p$. (b) Dynamical efficiency $\varepsilon$ versus $p$. The values of $M_s$, $L$, and $\varepsilon$ at an optimal value $p^*_s$ ($\simeq 0.26$) are denoted by the symbol “+.” Optimally fast sparsely synchronized rhythm for $p = p^*_s$; (c) raster plot of neural spikes and plot of the IPSR kernel estimate $R(t)$ versus $t$ and (d) ISI histogram (ISI histogram is composed of $5 \times 10^4$ ISIs, the bin size for the histogram is 0.5 ms, and vertical dotted lines in (d) denote integer multiples of the global period $T_G$ ($\simeq 6.8$ ms) of $R(t)$).

Through calculation of a realistic thermodynamic order parameter $\Theta$, fast sparsely synchronized rhythms have been found to emerge as $p$ passes a small critical value $p^*_c$ ($\simeq 0.12$). For $p > p^*_c$, the IPSR kernel estimate $R(t)$ has been found to oscillate with population frequency of 147 Hz. However, individual neurons discharge spikes stochastically at low rates ($\sim 33$ Hz) which is much lower than the population frequency. We have also investigated the effect of geometrical long-range connections on dynamical correlations between neuronal pairs for occurrence of global synchronization. It has thus been found that for $p > p^*_c$, the dynamical correlation length covers the whole system, thanks to sufficient number of long-range connections, and consequently global synchronization appears in the whole population. The degree of fast sparse synchronization has been well measured in terms of the realistic statistical–mechanical spiking measure $M_s$ introduced by considering both the occupation and the pacing degrees of spikes in the raster plot of neural spikes. As $p$ is increased, the synchrony degree increases, while the network axon wiring length also becomes longer because more long-range connections appear. Hence, wiring economy must be taken into account for dynamical efficiency. A ratio of the synchrony degree to the geometrical wiring cost is found to be maximal at a dynamical-efficiency optimal value $p^*_s$ ($\simeq 0.26$). For this case, an optimal fast sparsely synchronized rhythm is found to emerge at a minimal wiring cost in an economic small-world network.

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