

Complex dynamics of compound bursting with burst episode composed of different bursts

Zhuoqin Yang · Qingyun Wang · Marius-F Danca · Jiaoying Zhang

Received: date / Accepted: date

Abstract Episodic or compound bursting arises from a transition between a burst episode composed of a long burst and several short bursts and a relatively long subthreshold oscillation in this work. The minimal and generic phantom bursting model proposed by Bertram et al is employed to produce compound bursting of a single pancreatic β -cell and compound bursting synchronization with anti-phase spikes of two electrical coupling pancreatic β -cells. Two different fast/slow analysis for the moderate and the slower slow variables in three-dimensional spaces are combined to highlight better how these two slow variables with different time scales commonly or separately result in complex dynamic of the compound bursting of both the single β -cell and the two electrical coupling β -cells. Especially for the compound bursting synchronization with anti-phase spikes, more is stressed on how a varying coupling strength leads to a change of the number of the short

bursts within the burst episode for different compound bursting.

Keywords Compound bursting · Slow variables with different time scales · Electrical coupling · Synchronization · Fast/slow analysis

1 Introduction

Electrical bursting in electrically excitable cells is characterized by alternation of fast spiking oscillations with silent periods. As a very complex bursting pattern, episodic or compound bursting is exhibited as a transition between a burst episode composed of a few bursts and a long silent phase or "desert". As reported in pancreatic islets the compound bursting is responsible for pulsatile insulin secretion, since its period is considerably longer than a single simple burst [1,2].

In a variety of bursting models the variables can typically be classified as either "fast" or "slow", and there is usually a single slow variable determining the period of bursting [3–8]. Instead the compound bursting can be represented in some bursting models with two or more disparate-time-scale slow variables, and the burst period is determined by some combination of the slow variable time constants [9,10].

It is usually thought that independent glycolytic oscillations play a key role in producing the compound bursting [11–13]. A minimal model for electrical activity and independent glycolysis oscillation demonstrates the occurrence of compound bursting [11]. More realistic models for both the electrical and the glycolytic components including the participation of mitochondria describe this mechanism for compound bursting in pancreatic β -cells [12,13]. As data have been gradually accumulated and some biological phenomena to

This work is supported by the National Natural Science Foundation of China (Nos.11072013 and 11172017).

Z. Yang
School of Mathematics and System Sciences, Beihang University, Beijing 100191, China

Q. Wang
Department of Dynamics and Control, Beihang University, Beijing 100191, China
E-mail: nmqingyun@163.com

Marius-F Danca
Department of Mathematics and Computer Science, Avram Iancu University, Cluj-Napoca 400380, Romania

Romanian Institute of Science and Technology, 400487 Cluj-Napoca, Romania

J. Zhang
School of Mathematics and System Sciences, Beihang University, Beijing 100191, China

be explained have expanded, the models become more and more complex, culminating in the so-called dual oscillator model [14, 15], where slow oscillations in glycolysis are combined with an already complex model of electrical activity. At this level of complexity, the classical fast-slow approach becomes intractable due to the large number of slow variables. For consideration of simplification, Bertram et al [16] uses a minimal and generic phantom bursting model [9] to produce episodic or compound bursting in some parameter settings. It is obtained from time courses of a moderate and a slower slow variable and membrane potential that dynamics of the moderate slow variable is responsible for fast bursts, while dynamics of the slower slow variable clusters bursts together with intervening deserts. So the compound bursting oscillation is examined by a fast/slow analysis treating the slower slow variable as the sole slow variable in [16]. Nevertheless, the compound bursting pattern is driven by a combination of these two slow variables with different time scales, which can commonly or separately play important but distinct role in different aspects of the compound bursting. It is obviously not enough to explain dynamic behavior of the complex compound bursting only considering the slower slow variable. Moreover, one need to investigate some unknown questions associated with the occurrence of the compound bursting listed as follows. What underlies the initiating and terminating of every burst, which is same or not to that of the beginning and ending of a burst episode? What leads to shape of the bursts in the burst episode? Why to a certain extent of amplitude the first burst in the burst episode can continue for a long time and appears to be long while the other bursts have to end very soon and be short? Also, why subthreshold oscillation exists in the quiescent state between the burst episodes but not between the bursts in the burst episode? In this work, different fast/slow analysis for the moderate and the slower slow variables, respectively, are exhibited in a three-dimensional space so as to investigate preferably how these two different-time-scale slow variables commonly or separately give rise to the compound bursting.

On the other hand, synchronization of bursting activity in pancreatic islets when pancreatic β -cells are coupled together by gap junctions plays an important role in the insulin secretion [17, 18]. In addition, an impact of the channel noise in the important ATP-sensitive K⁺ channels can be reduced when shared by all the cells in the islet [18–20]. The phenomenon of synchronization via gap junctions has been extensively investigated in the pancreatic β -cell models [21–24] and even in some other coupled biological oscillators [25–27]. However, compound bursting induced by electrical

coupling is little studied and is still a challenging problem because of complexity of two or more disparate-time-scale slow variables. Here, the minimal phantom bursting model of two electrical coupling pancreatic β -cells is applied to examine for the first time compound bursting synchronization phenomena with anti-phase spikes. In particular, it is more concerned what evokes a change of the number of the short bursts within a burst episode in different compound bursting.

The structure of this paper is settled as follows. Following this brief introduction in Section 1, the phantom bursting model is presented in Section 2. Numerical investigations and fast/slow analysis for both the single cell model and the two coupled cells model are described in Section 3 and 4, respectively. Finally, conclusions and discussions are given in Section 5.

2 The phantom bursting model

An important feature of the minimal compound bursting model [9] is two slow variables with different time scales interact with a fast subsystem. The fast subsystem is composed of two differential equations

$$\frac{dV}{dt} = -(I_{Ca} + I_{Kdr} + I_{Leak} + I_{K1} + I_{K2})/C_m \quad (1)$$

$$\frac{dn}{dt} = [n_{\infty}(V) - n]/\tau_n(V) \quad (2)$$

In equation (1), C_m is the membrane capacitance and V is membrane potential. A fast Ca²⁺ current I_{Ca} , a fast delayed rectifier current I_{Kdr} , a constant-conductance leakage current I_{Leak} , and two slow K⁺ currents I_{K1} and I_{K2} with slowly changing conductances are expressed as follows, respectively.

$$I_{Ca} = g_{Ca} m_{\infty}(V)(V - V_{Ca}) \quad (3)$$

$$I_{Kdr} = g_{Kdr} n(V - V_K) \quad (4)$$

$$I_{Leak} = g_{Leak}(V - V_{Leak}) \quad (5)$$

$$I_{K1} = g_{K1} s_1(V - V_K) \quad (6)$$

$$I_{K2} = g_{K2} s_2(V - V_K) \quad (7)$$

where V_{Ca} , V_K and V_{Leak} are the Nernst potentials of the Ca²⁺ current, the K⁺ current and the leakage current, respectively, and g_{Ca} , g_{Kdr} , g_{Leak} , g_{K1} and g_{K2} are the maximum conductances of the respective ionic currents. An equilibrium function $m_{\infty}(V)$ for the Ca²⁺ current activates very rapidly, and an activation variable n for the delayed rectifier current I_{Kdr} satisfies equation (2) with an equilibrium function $n_{\infty}(V)$ and a time scale function $\tau_n(V)$.

The other two activation variables s_1 and s_2 changing very slowly in virtue of large time constants τ_{s_1} and

τ_{s_2} , constitute the slow subsystem as described in the following differential equations.

$$\frac{ds_1}{dt} = [s_{1\infty}(V) - s_1]/\tau_{s_1} \quad (8)$$

$$\frac{ds_2}{dt} = [s_{2\infty}(V) - s_2]/\tau_{s_2} \quad (9)$$

Moreover, a big difference in values of τ_{s_1} and τ_{s_2} can induce disparate time scales of these two slow processes.

The equilibrium functions $m_\infty(V)$, $n_\infty(V)$, $s_{1\infty}(V)$ and $s_{2\infty}(V)$ have the form of $x_\infty(V)$ below,

$$x_\infty(V) = \frac{1}{1 + \exp((v_x - V)/s_x)} \quad (10)$$

Parameter values are fixed as shown in [16], except for the maximum conductance of the moderate slow K+ current g_{K1} as a key parameter. All numerical simulations and bifurcation diagrams were calculated with Matlab and XPPAUT [28] software packages.

3 Compound bursting in the single β -cell model

Compound bursting consisting of bursts episodes followed by long silent phases can appear in the minimal phantom bursting model (1)-(10) in Fig.1, due to the impact of these two slow variables with different time scales. Just as time course of the membrane potential V for $g_{K1}=21.5\text{pS}$ shown in Fig.1(a), a burst episode in the compounding bursting is composed of a long burst with many spikes, several short bursts with a few spikes and subthreshold oscillation with large amplitude on both sides but small on the middle during a long silent phase. At the meantime, the membrane potential of the compounding bursting is accompanied with a no-spike compound mode of the moderate slow variable s_1 in Fig.1(b) and a relatively large jagged oscillation of the slower slow variable s_2 in Fig.1(c).

The slow variables s_1 and s_2 with different time scales may commonly or separately contribute to complex dynamic of the compound bursting. So two different fast/slow analyses for the slow variables s_1 and s_2 given in three-dimensional V - s_1 - s_2 spaces in Fig.2 and 3, respectively, are combined to explore the dynamic behavior of the compound bursting.

In the fast/slow analysis in Fig.2, the moderate variable s_1 is treated as a bifurcation parameter of the fast subsystem (1) and (2) and the slower slow variable s_2 is fixed at different values between 0.40 and 0.47 presented in the oscillatory range in Fig.1(c). A slow manifold surface is composed of multiple Z-shaped bifurcation curves of the fast subsystem with respect to the bifurcation parameter s_1 for these different values of s_2 . Stable nodes on the lower branches and saddles on the

middle branches of the Z-shaped curves coalesce via fold bifurcation (LP) at left knees. As s_1 increases the minimum and the maximum of amplitude of stable limit cycles via Hopf bifurcations (H) on upper branches of the Z-shaped curves form a C-shaped surface, since the stable limit cycles hit saddles on middle branches of the Z-shaped curves and disappear via saddle homoclinic bifurcation. S_1 -nullcline surface for the different values of s_2 (blue) and trajectory of the compound bursting (red) are overlapped in Fig.2. Fig.2(b) is gotten when rotating Fig.2(a) 90° and making it bigger, and Fig.2(c) is a projection of Fig.2(a) in the V - s_1 plane.

At lower level of the s_1 the trajectory jumps up from the stable lower branch to the stable limit cycle branch, and the burst starts to appear in the active phase owing to coalescence of the lower branches and the middle branches via the fold bifurcation. A little tilted form at the start of the burst is resulted when it locating in the C-shaped stable limit cycles, and then the burst terminates and the trajectory returns to the lower branch as the stable limit cycles disappear via saddle homoclinic bifurcation. It follows that these two bifurcations with respect to the fast slow variable s_1 are primarily responsible for the beginning and the ending of every burst in burst episode. However, whether the causes of the beginning and ending of every burst are same to those of the burst episode? Then, why the burst episode can terminate and go into subthreshold oscillation in the quiescent state? In the burst episode, why the first burst can continue for a long time and appears to be long while the other bursts have to end very soon and be short?

It is worth noting that the slower slow variable s_2 gradually increases during the occurrence of the burst episode in the three-dimensional V - s_1 - s_2 space in Fig.2. Therefore, the fast/slow analysis for the moderate slow variable s_1 in Fig.2 should be combined with that for the slower slow variable s_2 in Fig.3, where the moderate slow variable s_1 is converted to a fast variable and a fast subsystem consists of (1), (2) and (8). The fast subsystem also forms a Z-shaped bifurcation curve as shown in Fig.3, where (b) and (c) are enlargements of two different parts in (a), respectively. But differently the stable limit cycle occurring via a supercritical Hopf bifurcation (H1) on the upper branch loses stability via a period-doubling bifurcation, and then the unstable limit cycle hits saddle on the middle branch and disappears via saddle homoclinic bifurcation. Apart from that, there is another subcritical Hopf bifurcation (H2) on the lower branch, before which only unstable limit cycle coexists with a stable focus and then hits the saddle on the middle branch.

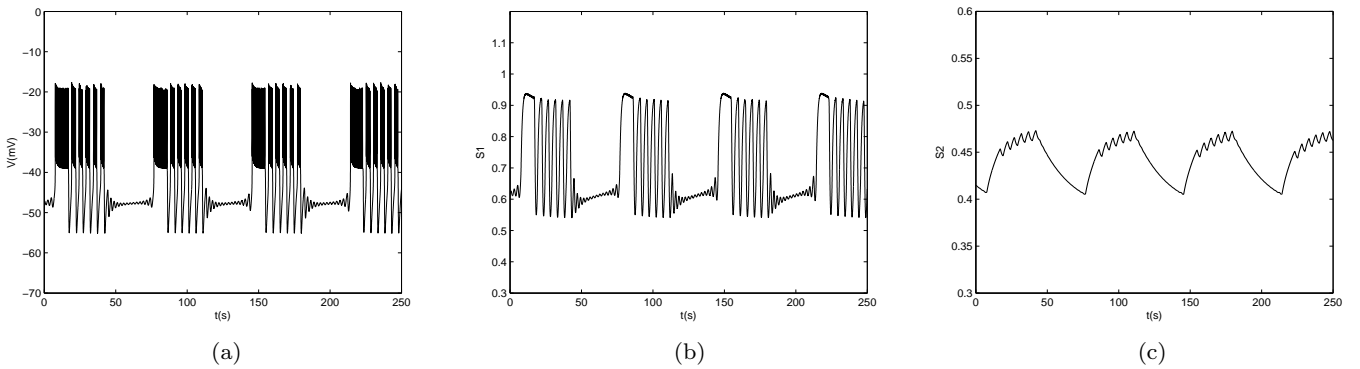


Fig. 1 Time courses of the membrane potential V (a), the moderate slow variable s_1 (b) and the slower slow variable s_2 (c) for $g_{K1}=21.5\text{pS}$ in the single cell model.

The first burst in burst episode above both s_1 -nullcline in Fig.2(b) and s_2 -nullcline in Fig.3(b) begins to move from forward to backwards along an increasing direction of s_2 . When the stable limit cycle hits saddle on the middle branch of the z-shaped bifurcation curve for the bifurcation parameter s_1 in Fig.2, the trajectory exactly reaches the stable limit cycle around the upper branch of the z-shaped bifurcation curve for the bifurcation parameter s_2 in Fig.3. Therefore, the fast/slow analysis for the slower slow variable s_2 in Fig.3 is employed to consider different dynamic behaviors of the long burst and the short bursts before their terminating via the saddle homoclinic bifurcation in Fig.2. The first burst at smaller values of s_2 can continue to possess so many spikes in a very long interval of s_2 until the stable limit cycle becomes unstable at the PD bifurcation point in Fig.3. However, the other bursts locating in the unstable limit cycles at larger values of s_2 are not able to continue and have to become the short bursts. As s_2 continues to rise, the whole burst episode composed of the long burst and the short bursts is totally terminated via the saddle homoclinic bifurcation for the slower slow variable s_2 in Fig.3. Thus, the initiation of the burst episode, that is, the first burst is induced by fold bifurcation for s_2 , while the termination of the burst episode, that is, the last short burst is resulted from collaboration of the different saddle homoclinic bifurcations for s_1 and s_2 . Furthermore, the trajectory falls down and stays in gradually decreasing oscillatory mode around the stable focus on the lower branch of the Z-shaped curve for the bifurcation parameter s_2 in Fig.3. As s_2 decreases, the stable focus becomes unstable through the subcritical Hopf bifurcation (H2) so that the trajectory goes away from the lower branch in gradually increasing oscillatory mode. That can account for the subthreshold oscillation with large amplitude on both sides and small one on the middle during the long silent phase.

4 Compound bursting synchronization with the different number of short bursts

Bursting synchronization induced by electrical coupling in pancreas islet plays an important role in the insulin secretion and it has been explored in many pancreatic β -cell models. However, synchronization of compound bursting induced by two or more disparate-time-scale slow variables is still a challenging problem. In the present work, the minimal phantom model is extended to two pancreatic β -cells connected by gap junction to explore synchronization behaviors of the compound bursting evoked by different electrical coupling strength. To model gap junctions of pancreatic β -cells, the two electrical coupling cells model is often handled by adding the following linear coupling term $g_c(V_i - V_j)$, where g_c is the coupling strength between two neighbor identical cells 1 and 2.

$$\frac{dV_i}{dt} = -[I_{Ca}(V_i) + I_{Kdr}(V_i, n_i) + I_{Leak}(V_i) + I_{K1}(V_i, s_{i1}) + I_{K2}(V_i, s_{i2}) + g_c(V_i - V_j)]/C_m \quad (11)$$

$$\frac{dn_i}{dt} = [n_\infty(V_i) - n_i]/\tau_n(V_i) \quad (12)$$

$$\frac{ds_{i1}}{dt} = [s_{1\infty}(V_i) - s_{i1}]/\tau_{s1} \quad (13)$$

$$\frac{ds_{i2}}{dt} = [s_{2\infty}(V_i) - s_{i2}]/\tau_{s2} \quad (14)$$

where $i, j=1, 2$ and $i \neq j$ represent the number of the cells. In this work, the coupling strength of gap junctions g_c is treated as a free parameter.

The compound bursting with the long burst, the short bursts and the subthreshold oscillation for $g_{K1}=21.5\text{pS}$ can be presented in the single cell model in Fig.1. Actually in a certain range of the parameter g_{K1} , the increasing electrically coupled strength g_c in the two coupled cells model (11)-(14) can make a transition from fast bursting to compound bursting even to slow bursting.

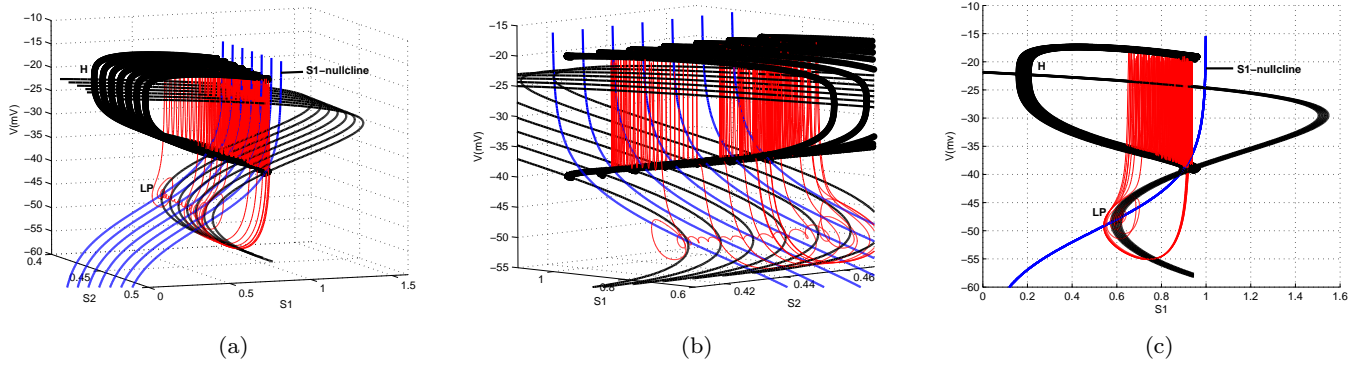


Fig. 2 (a) The fast/slow analysis of the compound bursting for $g_{K1}=21.5\text{pS}$ in the single β -cell model in the three-dimensional $V - s_1 - s_2$ space, when the moderate variable s_1 is treated as the bifurcation parameter of the fast subsystem (1) and (2) and the slower slow variable s_2 is considered as a constant. The s_1 -nullcline surface (blue) and trajectory of the compound bursting (red) are overlapped; (b) is gotten when rotating (a) 90° and making it bigger; (c) is a projection of (a) in the V - s_1 plane.

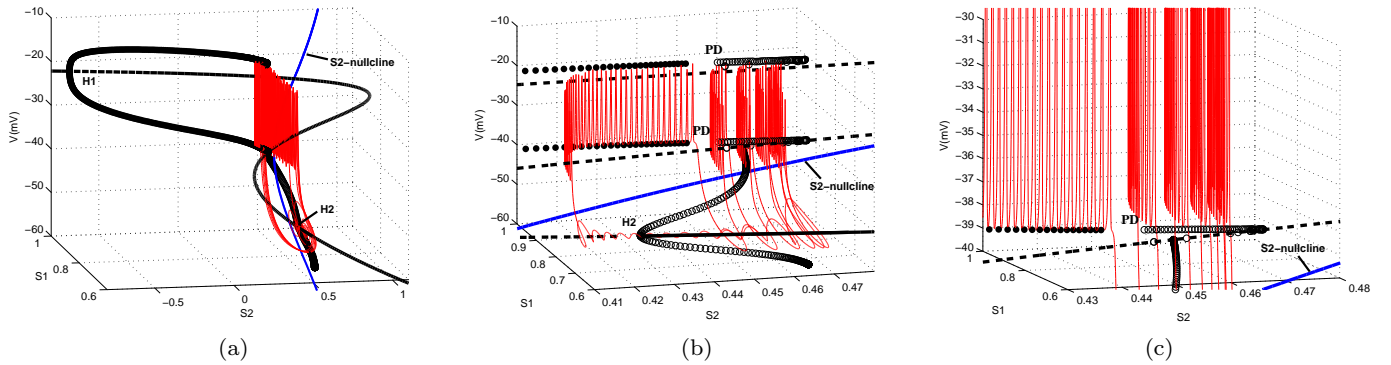


Fig. 3 The fast/slow analysis of the compound bursting for $g_{K1}=21.5\text{pS}$ in the single β -cell model in the three-dimensional $V - s_1 - s_2$ space, when the slower variable s_2 is treated as the bifurcation parameter of the fast subsystem (1), (2) and (8), where H1 and H2 appear in upper and lower branches, respectively; (b) and (c) are enlargements of two different parts in (a), respectively.

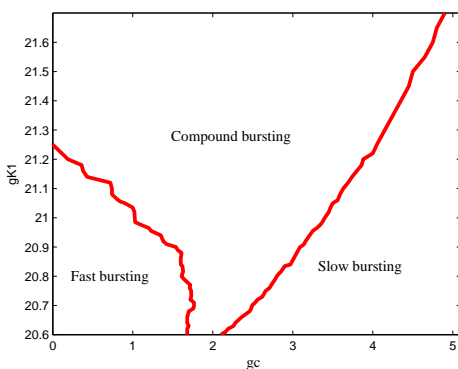


Fig. 4 The existing ranges of the different bursting patterns in the g_{K1} - g_c plane.

As shown in (g_{K1}, g_c) -plane in Fig.4, compound bursting can dominate a very wide region, especially it can always exist no matter coupled or uncoupled when g_{K1} varying between 21.3 and 21.7.

Here, we only consider different bursting behaviors induced by different electrically coupled strength g_c for $g_{K1}=21.2\text{pS}$. As shown in Fig.5, the fast bursting in the uncoupled case is a square-wave burster. From dynamic point of view, it is classified as "fold/homoclinic" bursting by the fast/slow analysis for the moderate slow variable s_1 and a fixed value of s_2 chosen from its very narrow oscillatory range from 0.47 to 0.48 in Fig.5(b). Furthermore, all these compound bursting patterns for the different electrically coupled strength g_c in the two electrical coupling cells model gradually synchronize in the membrane potential V_1 and V_2 , two moderate slow variables s_{11} and s_{21} and two slower slow variables s_{12} and s_{22} as shown in Fig.6. In every burst episode of these different synchronized compound bursting patterns, the number of the short bursts will decrease from 4 to 3, 2 and 1 when the coupled strength g_c is fixed as 1.6, 1.95, 2.5 and 3.3, respectively. Additionally, it can be seen in return maps of Inter-Burst Intervals (IBIs) in Fig.7 that time intervals between two adjacent short

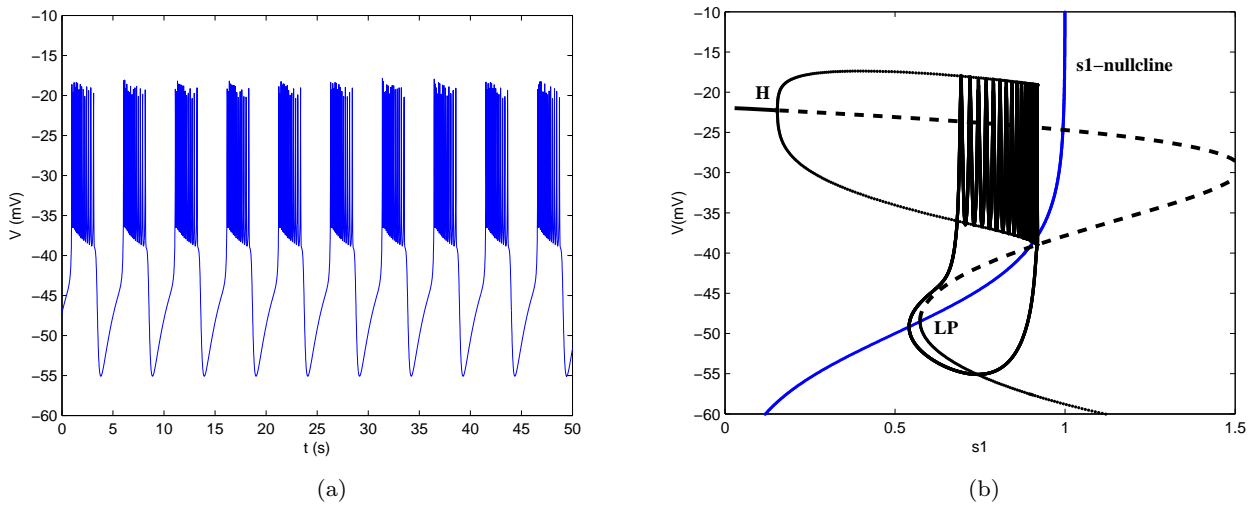
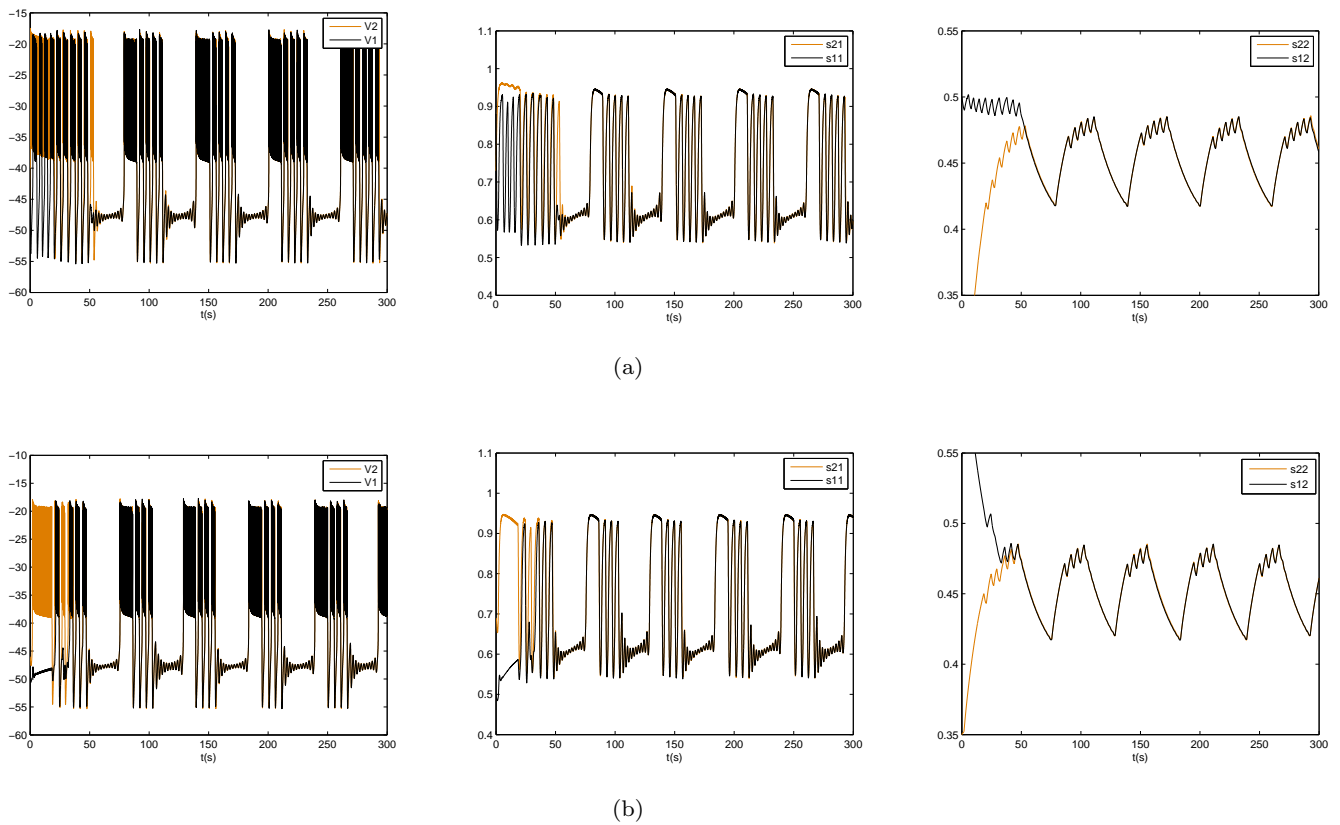


Fig. 5 Fast "fold/homoclinic" bursting for $g_{K1}=21.2\text{pS}$ in the single β -cell model, where (a) is time course of the membrane potential V and (b) is the fast/slow analysis for the bifurcation parameter s_1 as s_2 is fixed at 0.483.



bursts have little change but the number of the minimal IBIs decreases from 3 to 2, 1 and 0. That means the different number of the short bursts is decided by the difference in total time durations of all these short bursts. Therefore, the mechanism behind the different durations of all these short bursts in the burst episode for the coupled strength $g_c=1.6, 1.95, 2.5$ and 3.3 , re-

spectively, will be deep revealed by the fast/slow analysis of the two coupled β -cells model.

Without the loss of generality, we only give the fast/slow analysis of the compound bursting synchronization with a long burst and two short bursts for $g_c=2.5$. Just as the time courses of the moderate slower variables and the slower slow variables for cell 1 and 2 in Fig.6,

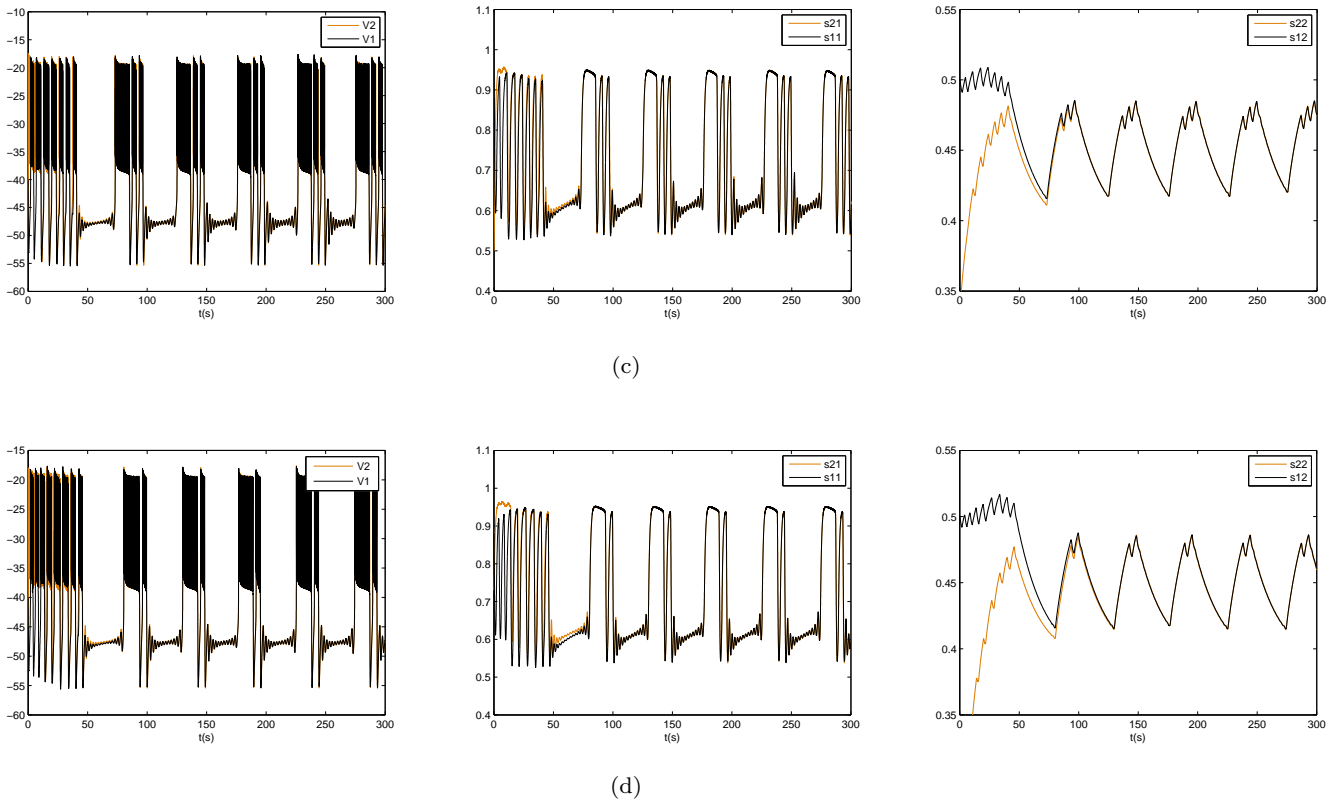


Fig. 6 Compound bursting synchronization shown in time courses of the membrane potentials V_1 and V_2 , two moderate slow variables s_{11} and s_{21} and two slower slow variables s_{12} and s_{22} in the two coupled β -cells model when the coupled strength $g_c=1.6$ (a), 1.95(b), 2.5(c) and 3.3(d), respectively.

the moderate slower variable $s_{11} \approx s_{21} = s_1$ for different values of $s_{12} \approx s_{22}$ and the slower slow variable $s_{12} \approx s_{22} = s_2$ are regarded as a bifurcation parameter in the fast/slow analyses in Fig.8 and Fig.9, respectively. Stable limit cycles via two Hopf bifurcation points H1 and H2 on upper branch of z-shaped bifurcation curves in the fast/ slow analyses in Fig.8 and Fig.9, correspond to in-phase (IP) and anti-phase (AP) branches, respectively. Since there is a function relationship between V_1 and V_2 , the dynamic behavior of V_1 is only considered here. The stable limit cycle along the AP branch decides anti-phase spike of burst synchronization in a burst, which can be seen in the time courses of V_1 and V_2 in Fig.10(a) and the V_1 - V_2 phase plane in Fig.10(b). In the fast/ slow analysis for the bifurcation parameter $s_{12} \approx s_{22} = s_2$ in Fig.9, the first anti-spike burst along the stable AP limit cycle branch exists in a long interval of s_2 before Torus bifurcation(TR). The other anti-spike bursts lying in the unstable limit cycle after the TR bifurcation can not continue but return to the stable lower branch via the saddle homoclinic bifurcation (HC). It follows that the total time duration of all the short bursts in every compound burst-

Table 1 The difference of the TR bifurcation points and the saddle homoclinic points for the increasing coupled strength g_c .

g_c	s_2 for TR	s_2 for HC	The difference between them
1.6	0.4596	0.4745	0.0149
1.95	0.4654	0.4787	0.0133
2.5	0.4742	0.4827	0.0085
3.3	0.4853	0.4916	0.0063

ing is determined by the existing range of the unstable limit cycle between the TR bifurcation point and the saddle homoclinic point for the bifurcation parameter $s_{12} \approx s_{22} = s_2$. Moreover, it can be seen in Table 1 that the difference of the TR bifurcation points and the saddle homoclinic points gradually decreases with the increasing coupled strength g_c , thereby it so that the total time duration of all these short bursts decrease with the increasing coupled strength g_c . Based on rough invariability in the time durations of every short burst and in the time intervals between the short bursts, the number of the short bursts gradually decreases from 4 to 3, 2 and 1.

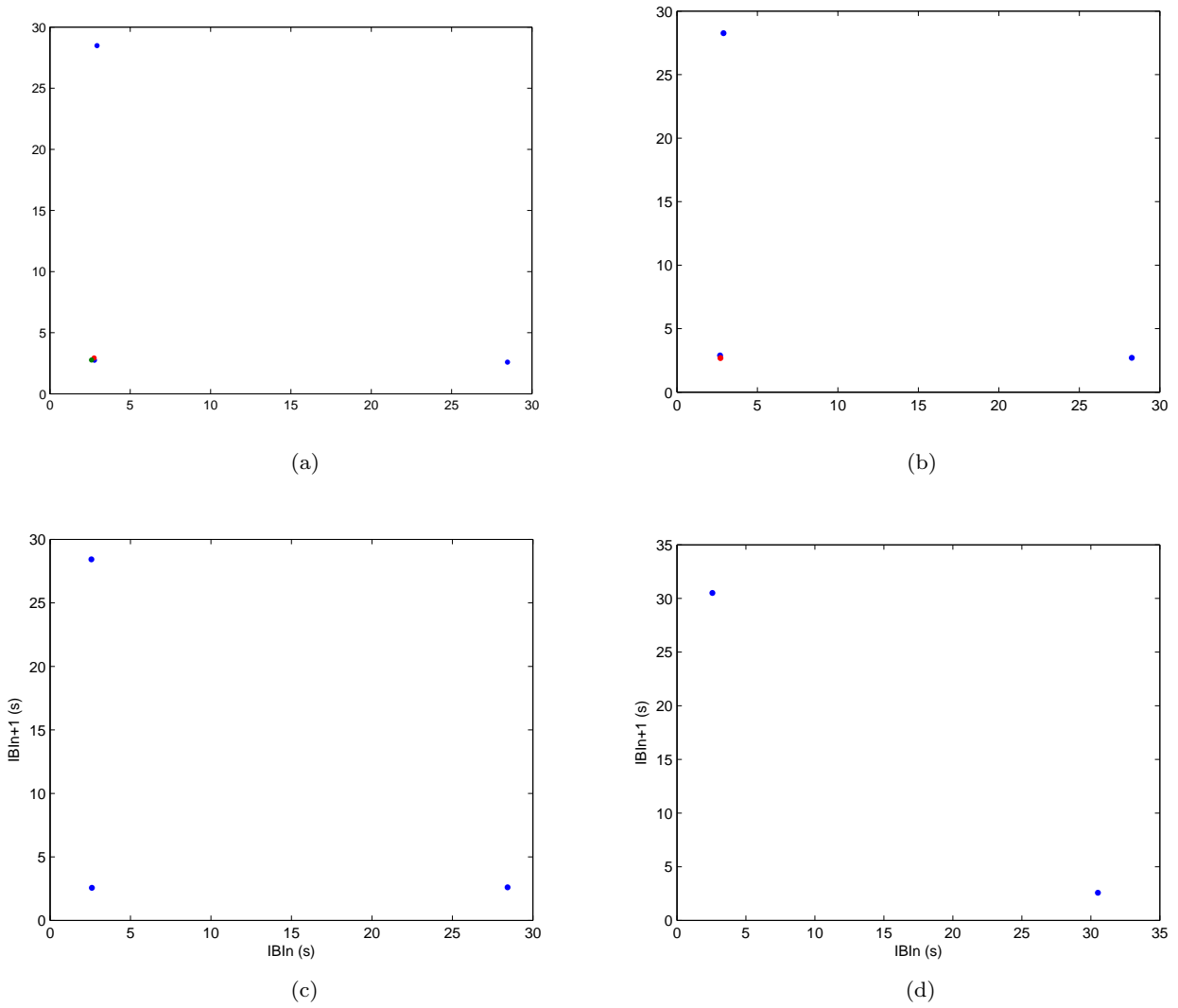


Fig. 7 Return maps of Interburst intervals (IBIs) of the compound bursting patterns for $g_c=1.6$ (a), 1.95(b), 2.5(c) and 3.3(d), respectively.

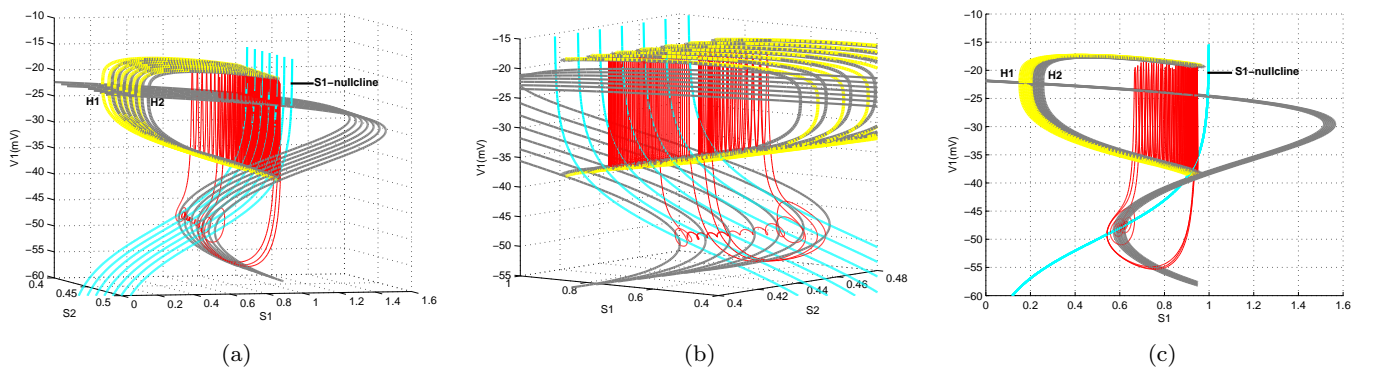


Fig. 8 (a) The fast/slow analysis for $s_{11} \approx s_{21} = s_1$ as the bifurcation parameter when the coupled strength $g_c=2.5$ in the three-dimensional $V-s_1-s_2$ space, where stable limit cycles via these two Hopf bifurcations H1 and H2 correspond to in-phase (IP, yellow) and anti-phase (AP, grey) branches, respectively. (b) is gotten when rotating (a) 90° and making it bigger. (c) is a projection of (a) in the V_1-s_1 plane.

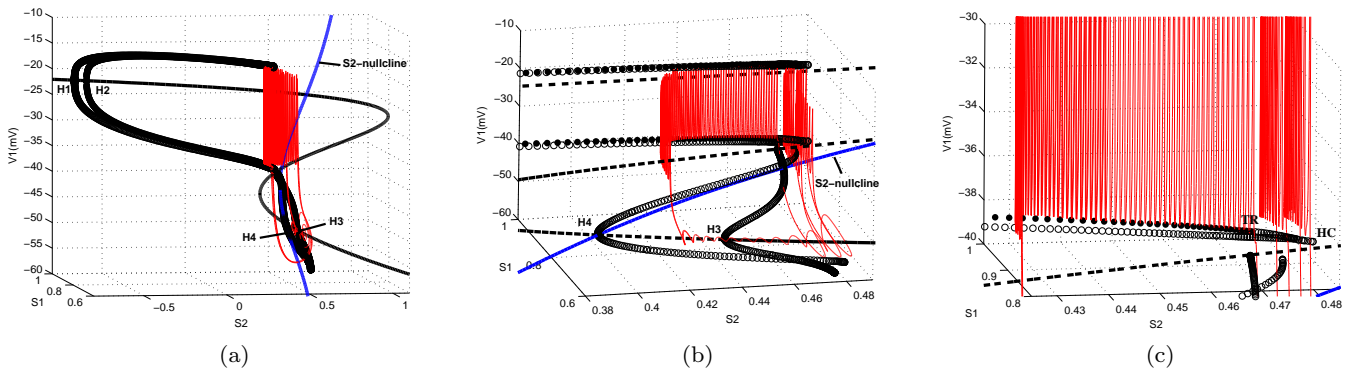


Fig. 9 (a) The fast/slow analysis for $s_{12} \approx s_{22} = s_2$ as the bifurcation parameter when the coupled strength $g_c=2.5$ in the three-dimensional $V-s_1-s_2$ space, where stable limit cycles via these two Hopf bifurcations H1 and H2 correspond to in-phase (IP) and anti-phase (AP) branches, respectively. (b) and (c) are enlargements of two different parts in (a), respectively.

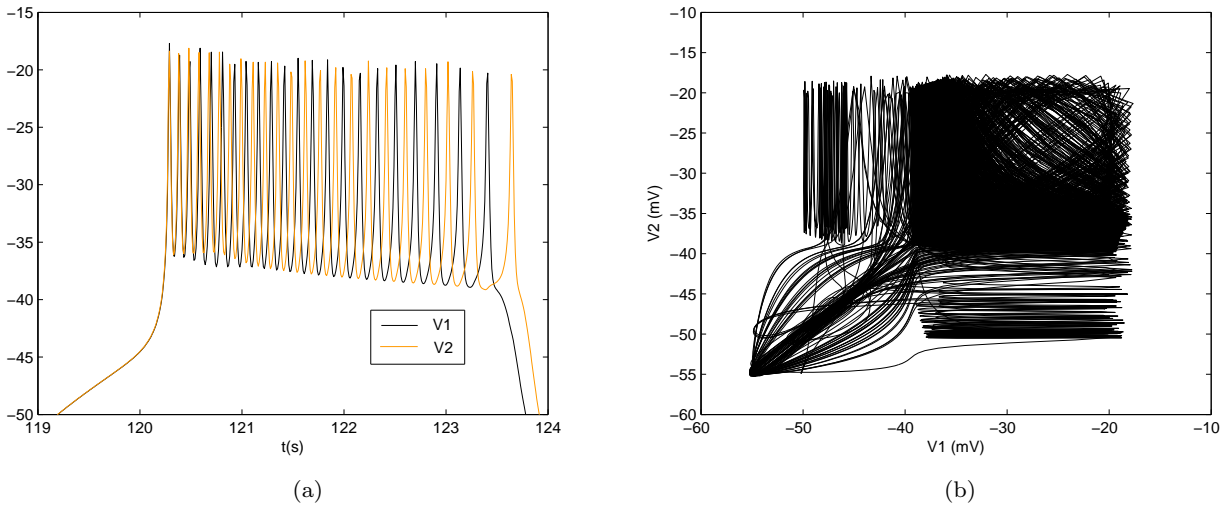


Fig. 10 An anti-spike burst synchronization in the compound bursting for $g_c=2.5$ is represented in (a) time courses of V_1 (black) and V_2 (orange) and (b) the trajectory in V_1 - V_2 phase plane.

5 Conclusions and discussions

Episodic or compound bursting is a very complex but important bursting pattern in bursting models with two or more disparate-time-scale slow variables. For consideration of simplification, the minimal and generic phantom bursting model [9] is used to produce episodic or compound bursting, when two slow variables with different time scales commonly or separately play different roles in different aspects of the compound bursting. It is obviously not enough to explain dynamic behavior of the complex compound bursting only considering the slower slow variable. Therefore, different fast/slow analysis for the moderate and the slower slow variables, respectively, are given in a three-dimensional space to reveal complex dynamic of the compound bursting. At first, the beginning and the ending of every burst in burst episode are decided by the fold bifurcation and

the saddle homoclinic bifurcation for the moderate slower variable s_1 , where the little tilted form at the start is resulted from locating in the C-shaped stable limit cycles. Nonetheless, the burst episode is terminated by common effects of the different saddle homoclinic bifurcations for s_1 and s_2 . Then, the first burst exactly reaches the stable limit cycle for the bifurcation parameter s_2 , when the stable limit cycle for the bifurcation parameter s_1 hits saddle on the middle branch. So the first burst existing in stable limit cycle for s_2 have many spikes but the others locating in the unstable limit cycles for s_2 are not able to continue and become the short bursts, which can explain different occurrence of the long burst and the short bursts. Finally, during the long silent phase the property of the stable focus on the lower branch for the bifurcation parameter s_2 accounts for the subthreshold oscillation with large amplitude on both sides and small amplitude on the middle.

Furthermore, synchronization of bursting activity plays an important role in the insulin secretion. Using the minimal phantom bursting model of two electrical coupling pancreatic β -cells, we examine for the first time compound bursting synchronization phenomena with anti-phase spikes. Moreover, we also discuss the fundamental mechanism of the change of the number of the short bursts within a burst episode for different compound bursting. Since the total time duration of all the short bursts is decided by the existing range of the unstable limit cycle between the TR bifurcation point and the saddle homoclinic point. So the decrease of the difference between these two points for the increasing coupled strength g_c leads to the decreasing durations of all these short bursts. All above can explain the number of the short bursts decreasing from 4 to 3, 2 and 1, in consideration of rough invariability of the time duration of every short burst and the time intervals between the short bursts.

In fact, heterogeneity in the coupled phantom-burster model with heterogeneous electrical coupling can reproduce many aspects of the measured islet electrical dynamics. Furthermore, it would be interesting to study the influence of the topology of the network and the coupling strength between β -cells inside the Langerhans islets. Using complex networks theory, one can investigate under what conditions synchronization emerges for β -cell network models. In addition, periodic insulin secretion appears to be controlled by metabolic oscillations within the cells. To explore the bases of the synchronization phenomenon, both electrical and metabolic features of a mathematical gap junction model will be taken into account to study biological functions of the β -cell cluster.

References

- Henquin, J.C., Meissner, H.P., Schmeer, W.: Cyclic Variations of Glucose-induced Electrical Activity in Pancreatic β -cells. *Pflugers Arch.* 393, 322-327 (1982)
- Zhang, M., Goforth, P., Sherman, A., Bertram, R., Satin, L.: The Ca^{2+} Dynamics of Isolated Mouse β -cells and islets: Implications for Mathematical Models. *Biophys. J.* 84, 2852-2870 (2003)
- Perc, M., Marhl, M.: Resonance effects determine the frequency of bursting Ca^{2+} oscillations. *Chem. Phys. Lett.* 376, 432-437 (2003)
- Perc, M., Marhl, M.: Different types of bursting calcium oscillations in non-excitable cells. *Chaos, Solitons & Fractals* 18, 759-773 (2003)
- Perc, M., Marhl, M.: Sensitivity and flexibility of regular and chaotic calcium oscillations. *Biophys. Chem.* 104, 509-522 (2003)
- Lu, Y., Ji, Q. B.: Control of intracellular calcium bursting oscillations using method of self-organization. *Nonlinear Dynamics.* 67(4), 2477-2482 (2012)
- Chay, T.R., Keizer, J.: Minimal Model for Membrane Oscillations in the Pancreatic β -cell. *Biophys. J.* 42, 181-190 (1983)
- Rinzel, J.: Bursting Oscillations in an Excitable Membrane Model. In: Sleeman, B.D., Jarvis, R.J.(Eds), *Ordinary and Partial Differential Equations, Lecture Notes in Mathematics.* New York: Springer. 304-316 (1985)
- Bertram, R., Previte, J., Sherman, A., Kinard, T.A., Satin, L.S.: The Phantom Burster Model for Pancreatic β -cells. *Biophys. J.* 79, 2880-2892 (2000)
- Rinzel, J.: A Formal Classification of Bursting Mechanisms in Excitable System. In: Teramoto, E., Yamaguti, M.(Eds.), *Mathematical Topics in Population Biology, Morphogenesis and Neurosciences.* Springer, Berlin (1987)
- Smolen, P., Terman, D., Rinzel, J.: Properties of a Bursting Model with Two Slow Inhibitory Variables. *SIAM J. Appl. Math.* 53, 861-892 (1993)
- Wierschem, K., Bertram, R.: Complex bursting in pancreatic islets: a potential glycolytic mechanism. *J. Theor. Biol.* 228, 513-521 (2004)
- Bertram, R., Satin, L., Zhang, M., Smolen, P., Sherman, A.: Calcium and Glycolysis Mediate Multiple Bursting Modes in Pancreatic Islets, *Biophys. J.* 87, 3074-3087 (2004)
- Bertram, R., Satin, L., Pedersen, M. G., Luciani, D. S., Sherman, A.: Interaction of Glycolysis and Mitochondrial Respiration in Metabolic Oscillations of Pancreatic Islets. *Biophys. J.* 92, 1544-1555 (2007)
- Goel, P., Sherman, A.: The Geometry of Bursting in the Dual Oscillator Model of Pancreatic β -cells. *SIAM J. Appl. Dyn. Sys.* 8(4), 1664-1693 (2009)
- Bertram, R., Rhoadsa, J., Cimborra, W.P.: A Phantom Bursting Mechanism for Episodic Bursting. *Bulletin of Mathematical Biology.* 70, 1979-1993 (2008)
- Sherman, A., Rinzel, J., Keizer, J.: Emergence of organized bursting in clusters of pancreatic β -cells by channel sharing. *Biophys. J.* 54, 411-425 (1988)
- Pedersen, M.G., Bertram, R., Sherman, A.: Intra- and inter-islet synchronization of metabolically driven insulin secretion. *Biophys. J.* 89, 107-119 (2005)
- Atwater, I., Rosario, L., Rojas, E.: Properties of the Ca-activated K^+ channel in pancreatic b-cells. *Cell Calcium* 4, 451-461 (1983)
- Chay, T.R., Kang, H.S.: Role of single-channel stochastic noise on bursting clusters of pancreatic β -cells. *Biophys. J.* 54, 427-435 (1988)
- Pedersen, M. G., Bertram, R. and Sherman, A.: Intra- and Inter-Islet Synchronization of Metabolically Driven Insulin Secretion. *Biophys. J.* 89, 107-119 (2005)
- Kitagawa, T., Murakami, N. and Nagano, S.: Modeling of the gap junction of pancreatic β -cells and the robustness of insulin secretion, *Biophys. J.* 6, 37-51 (2010)
- Barajas-Ramírez, J.G., Steur, E., Femat, R., Nijmeijer H.: Synchronization and activation in a model of a network of β -cells. *Automatica.* 47, 1243-1248 (2011)
- Gonze, D., Markadieu, N., and Goldbeter, A.: Selection of in-phase or out-of-phase synchronization in a model based on global coupling of cells undergoing metabolic oscillations. *Chaos.* 18, 037127 (2008)
- Wang, Z. L., Shi, X. R.: Chaos bursting synchronization of mismatched HindmarshRose systems via a single adaptive feedback controller. *Nonlinear Dynamics.* 67(3), 1817-1823 (2012)
- Shi, X. R.: Bursting synchronization of HindmarshRose system based on a single controller. *Nonlinear Dynamics.* 59(1-2), 95-99 (2010)

-
27. Perc, M., Marhl, M.: Synchronization of regular and chaotic oscillations: The role of local divergence and the slow passage effect. *Int. J. Bifurcat. Chaos.* 14, 2735-2751 (2004)
 28. Ermentrout, B.: *Simulating: Analyzing and Animating dynamical systems: a guide to XPPAUT for Researchers and students*, philadelphia. (2002)