The Calcium and Voltage Clocks in Sinoatrial Node Automaticity

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ABSTRACT

Recent evidence indicates that the voltage (cyclic activation and deactivation of membrane ion channels) and Ca2+ clocks (rhythmic spontaneous sarcoplasmic reticulum Ca2+ release) jointly regulate sinoatrial node (SAN) automaticity. Since the intact SAN is a heterogeneous structure that includes multiple different cell types interacting with each other, the relative importance of the voltage and Ca2+ clocks for pacemaking may be variable in different regions of the SAN. Recently, we performed optical mapping in isolated and Langendorff-perfused canine right atria. We mapped the intracellular calcium (Ca2+) and membrane potentials of the intact SAN simultaneously. Using previously described criteria of the timing of the late diastolic Ca2+ elevation (LDCAE) relative to the action potential upstroke to detect Ca2+ clock activity, we demonstrated that the sinus rate increased and the leading pacemaker shifted to the superior SAN with the robust LDCAE during β-adrenergic stimulation. We also showed that the LDCAE was caused by spontaneous diastolic SR Ca2+ release and was closely related with heart rate changes. We conclude that the Ca2+ and voltage clocks work synergistically to generate SAN automaticity. (Korean Circ J 2009; 39:217-222)

KEY WORDS: Calcium; Sinoatrial node; Sarcoplasmic reticulum; Sympathetic nervous system.

Sinoatrial node (SAN) automaticity is responsible for controlling heart rhythm. SAN function is therefore essential for normal cardiac physiology. Although it was shown that spontaneous diastolic depolarization of SAN cells periodically initiates action potentials to set the rhythm of the heart, the mechanism by which heart rhythm is generated is unclear. Sick sinus syndrome is an abnormality involving the generation of the action potential by the SAN and is characterized by an atrial rate inappropriate for physiologic requirements. The sick sinus syndrome occurs in 1 of every 600 cardiac patients with >65 years of age and accounts for approximately one-half of implantations of pacemakers in the United States. A better understanding of the mechanisms of SAN automaticity is therefore clinically important.

The Voltage Clock and Sinoatrial Node Automaticity

The mechanism of spontaneous diastolic depolarization has traditionally been attributed to a “voltage clock” mechanism, mediated by voltage-sensitive membrane currents, such as the hyperpolarization-activated pacemaker current (Ii) regulated by cyclic adenosine monophosphate (cAMP). This current is also referred to as a “funny” current because, unlike the majority of voltage-sensitive currents, it is activated by hyperpolarization rather than depolarization. The funny channel becomes activated at the end of the action potential (at voltages from -40/-50 mV to -100/-110 mV), which corresponds to the range in which diastolic depolarization occurs. The funny channel then depolarizes the membrane to a level at which Na+ channels open to initiate the action potential. Ii is a mixed Na+-K+ inward current activated by hyperpolarization and modulated by the autonomic nervous system. Because the membrane ionic channels open and close according to the membrane potential, this process is referred to as a membrane voltage clock. The f-channels are encoded by the hyperpolarization-activated, cyclic nucleotide-gated (HCN) channel gene family. Of the four known HCN subunits, HCN4 is the most highly expressed in the mammalian SAN. The major role of Ii has been reinforced by the fact that mutations in the Ii channel are associated with a reduced baseline heart rate, and drugs which block Ii (such as ivabradine) do the same. However, while point mutations of HCN4 are associated with baseline sinus bradycardia, the maximum heart rate achieved during exercise is normal. The latter finding implies that Ii is not
the only mechanism of SAN automaticity, especially during sympathetic activation.

The Calcium Clock, a New Mechanism for Sinoatrial Node Automaticity

The traditional mechanism for heart rhythm generation has been restricted to sarcolemmal ion channels. Recently, spontaneous Ca$^{2+}$ release from the sarcoplasmic reticulum (SR) has been suggested to be the mechanism for sinus rhythm generation. When the SR is full, the probability of spontaneous Ca$^{2+}$ release increases. On the other hand, when the SR is empty, the chances for spontaneous Ca$^{2+}$ release decreases. The rhythmic alternations of SR Ca$^{2+}$ release is referred to as the Ca$^{2+}$ clock. Because the SR Ca$^{2+}$ content is controlled in part by the membrane voltage, it is important to recognize that the activation of the Ca$^{2+}$ clock and the membrane ionic clock are interdependent. The involvement of Cai in heart rhythm generation was first suggested by the observation that application of ryanodine slowed subsidiary pacemakers in cat right atria. Subsequent work showed the involvement of the electrogenic Na$^{+}$-Ca$^{2+}$ exchange current ($I_{\text{NCX}}$) in pacemaker activity and also raised the possibility that increased Ca$^{2+}$ release, followed by activation of $I_{\text{NCX}}$, could play a role in the positive chronotropic effect of β-adrenergic stimulation in latent pacemaker cells. Lakatta et al. suggested that spontaneous rhythmic Ca$^{2+}$ release from the SR in SAN cells, manifested as Ca$^{2+}$ sparks, work as the “Ca$^{2+}$ clock.”

The idea of a Ca$^{2+}$ clock suggests that the mechanism of automaticity is the same as that of delayed afterdepolarization (DAD), which occurs when there is a SR Ca$^{2+}$ overload. This suggests that the SAN must exist normally in a state of calcium overload. Vinogradova et al. demonstrated that a high basal level of protein kinase A (PKA) activity in the SAN might contribute to a state of Ca$^{2+}$ overload. A disease associated with Ca$^{2+}$ clock malfunction has also been reported in patients with a genomic deletion of the RyR2 exon-3. Patients with that mutation develop catecholaminergic polymorphic ventricular tachycardia, along with SAN and atrioventricular node dysfunction, atrial fibrillation, and atrial standstill. It is possible that Ca$^{2+}$ clock malfunction might contribute to the bradycardia and atrial arrhythmias in the latter patients.

Pacemaker Hierarchy and the Importance of the Ca$^{2+}$ Clock in an Intact Sinoatrial Node

Cardiac automaticity at the organ level is a very complex phenomenon and, in addition to cellular mechanisms, integrative factors are involved in cardiac pacemaking. The intact SAN is a heterogeneous structure that includes multiple different cell types interacting with each other. The relative importance of the voltage and Ca$^{2+}$ clocks for pacemaking in different regions of the SAN, and in response to neurohumeral stimuli, such as β-agonists, may be different. Indeed, activation maps in intact canine right atria (RA) have shown that the SAN impulse origin is multicentric, and sympathetic stimulation predictably results in a cranial (superior) shift of the pacemaking site in humans and dogs. Based on evidence from isolated SAN myocytes, late diastolic Ca$^{2+}$ elevation (LDCAE) relative to the action potential upstroke is a key signature of pacemaking by the Ca$^{2+}$ clock. It is possible that the same phenomenon could provide insight into the relative importance of the Ca$^{2+}$ and voltage clock mechanisms in pacemaking in intact SAN tissue. We therefore designed a study to determine the role of Ca$^{2+}$ and voltage clocks on heart rhythm generation and on the mechanisms of pacemaker hierarchy in the intact SAN.

Heterogeneous Ca$^{2+}$ Dynamics in an Intact Sinoatrial Node

At baseline, the spontaneous diastolic SR Ca$^{2+}$ release, which is manifested by the LDCAE, was observed in only a small percentage of the preparations. However, the LDCAE occurred in all preparations during isoproterenol infusion, and was associated with a superior shift of the leading pacemaker site coincident with the appearance of robust LDCAEs (Fig. 1) in this region. Most importantly, the site of maximum LDCAE slope always co-localized with the leading pacemaking site, suggesting a shift in which the voltage clock now lagged the Ca$^{2+}$ clock (Fig. 2). This observation indicates a strong association between LDCAE and pacemaking during β-adrenergic stimulation, and provides new insight into pacemaker hierarchy in the canine RA.

The Ca$^{2+}$ dynamics of SAN were characterized not only by the earliest onset of LDCAE, but also by the fastest Ca$^{2+}$ reuptake as compared with other RA sites. The baseline 90% Ca$^{2+}$ relaxation time was shorter at the superior SAN than at other RA sites. This resulted in the formation of the Ca$^{2+}$ sinkhole, which was facilitated by a rapid decline (short relaxation time) of the Ca$^{2+}$ fluorescence at the superior SAN during isoproterenol infusion and suggests that Ca$^{2+}$ reuptake by the SR is fastest in the superior SAN (Fig. 1D). The key protein regulator of SR Ca$^{2+}$ uptake is phospholamban, which inhibits SERCA2a in the dephosphorylated state. There was a significantly lower SERCA2a/phospholamban ratio at SAN sites than at RA sites, suggesting more phospholamban molecules are available to regulate SERCA2a molecules in SAN.
than in RA. Isoproterenol infusion phosphorylates phospholamban and relieves phospholamban inhibition of SERCA2a, which may account for more robust Ca\(^{2+}\) uptake in SAN than in RA during isoproterenol infusion.\(^{30}\)

**The Importance of Sarcoplasmic Reticulum Function on Pacemaking**

The LDCAE was closely related to SR Ca\(^{2+}\) release. Caffeine sensitizes the ryanodine receptor 2 to activation, resulting in increased SR Ca\(^{2+}\) release.\(^{39}\) The superior shift of the LDCAE and the pacemaking site was also consistently observed with caffeine infusion. Ryanodine, which blocks ryanodine receptors, caused a dose-dependent suppression of sinus node activity, and impaired isoproterenol-induced LDCAE.

The combination of ryanodine and thapsigargin also suppressed sinus node activity, and impaired isoproterenol-induced LDCAE. In contrast, the \(I_f\) blocker, ZD 7288 (3 \(\mu\)mol/L), did not prevent the LDCAE in the superior SAN.

**Mechanisms of Diastolic Depolarization**

Multiple time- and voltage-dependent ionic currents have been identified in cardiac pacemaker cells, which contribute to diastolic depolarization, including \(I_Ca,L\), \(I_Ca,T\), \(I_{Kf}\), and various types of delayed rectifier K currents.\(^{33}\) Many of these membrane currents are known to respond to \(\beta\)-adrenergic stimulation. Some of these currents, such as \(I_Ca,L\), also promote the LDCAE and the acceleration of the sinus rate by the Ca\(^{2+}\) clock, as well as the voltage clock. In an intact SAN, SR inhibitors and \(I_f\) blockade slowed the sinus rate under basal conditions and blunted the isoproterenol-induced increase in the sinus rate. Therefore, the interdependence and synergy between the two clocks was evident.

**Sympathetic Stimulation and Tachybrady Syndrome**

It is known that heart failure is frequently associated
Fig. 2. Co-localization of the LDCAE and leading pacemaker site. A: upward shift of the leading pacemaker site with the LDCAE during an isoproterenol infusion. (a) Ca ratio maps of SAN at each sinus rate. (b) corresponding Ca tracings from the superior (1, 2), middle (3, 4), and inferior (5, 6) SAN. At 95 bpm, sites 4 and 5 had the most prominent LDCAEs (*). As the sinus rate gradually increased, the sites of Ca elevation progressively moved upward. At the maximum sinus rate of 173 bpm, site 2 had the most apparent LDCAE. (a) the Ca and Vm tracings from the inferior, middle, and superior SAN sites at different sinus rates. (b) the LDCAE and DD slopes of the superior SAN at different sinus rates. This figure was reproduced with permission from Joung et al.\textsuperscript{31} LDCAE: late diastolic Ca elevation, SAN: sinoatrial node, DD: diastolic dysfunction.

Fig. 3. Satellite stellate ganglion nerve activity (SGNA) and prolonged sinus pause. A: intermittent low amplitude burst discharge activity (LABDA) (arrows) associated with intermittently increased heart rate, and a large LABDA (+)-induced tachycardia. Sudden SGNA withdrawal resulted in a 5.5-s prolonged sinus pause. B: the data within the dotted line in panel A: vagal nerve activity (VNA). This figure was reproduced with permission from Ogawa et al.\textsuperscript{31}
with SAN remodeling, resulting in decreased SAN reserve.\textsuperscript{32} We performed nerve recording in a canine model of pacing-induced heart failure and found intermittent tachybradycardia episodes.\textsuperscript{33} Interestingly, the prolonged (>3 s) sinus pauses were triggered not by vagal activation, but by short bursts of sympathetic activity (Fig. 3). Typically, a burst of sympathetic activity is associated with tachycardia. When there is sympathetic withdrawal, the tachycardia terminates, followed by prolonged pauses during which no activation was observed. Preliminary studies\textsuperscript{34} showed that cryoablation of the stellate ganglia during which no activation was observed. Preliminary findings suggest that the left stellate ganglion nerve activity is causally related to the tachy-arrhythmias. The authors suggested that the down-regulation of the stellate sympathetic activity might reduce the prolonged sinus pulse episodes in the same model. These findings suggest that the left stellate ganglion nerve activity is causally related to the tachybrady syndrome. The molecular mechanism of this association, however, remains unclear.

**Future Directions**

Our understanding of cardiac automaticity has progressed considerably during the last 10 years. Several questions regarding how the heart rhythm is generated and controlled in physiologic and pathologic conditions remain unanswered. A recent study by Yeh et al.\textsuperscript{35} showed funny current downregulation is associated with atrial tachyarrhythmias. The authors suggested that the down-regulation of funny current may underlie the mechanisms tachybrady syndrome. Whether or not Ca\textsuperscript{2+} clock malfunction also contributes to the mechanisms of tachybrady syndrome remains unclear. Our laboratory is currently evaluating whether the Ca\textsuperscript{2+} clock is impaired in a canine atrial fibrillation model with sinus dysfunction. Secondarily, we are trying to reproduce the human sick sinus syndrome model by manipulating the Ca\textsuperscript{2+} and voltage clocks in a canine isolated RA preparation. We anticipate that these studies will help us better understand the mechanisms of SAN dysfunction and facilitate design mechanism-based therapy of this disease.

**Acknowledgments**

Supported by NIH Grants P01 HL78931, R01 HL78932, 71140, a Korean Ministry of Information and Communication through research and develop support project (BJ), a Medtronic-Zipes Endowments (PSC) and an American Heart Association (AHA) Established Investigator Award (SFL).

Medtronic, Inc. and Cryocath, Inc. donated equipment to support our research laboratory.

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