Functional Triads Consisting of Ryanodine Receptors, Ca\textsuperscript{2+} Channels, and Ca\textsuperscript{2+}-activated K\textsuperscript{+} Channels in Bullfrog Sympathetic Neurons

Plastic Modulation of Action Potential

TENPEI AKITA and KENJI KUBA

From the Department of Physiology, Nagoya University School of Medicine, Nagoya 466-8550, Japan

ABSTRACT Fluorescent ryanodine revealed the distribution of ryanodine receptors in the submembrane cytoplasm (less than a few micrometers) of cultured bullfrog sympathetic ganglion cells. Rises in cytosolic Ca\textsuperscript{2+} ([Ca\textsuperscript{2+}],) elicited by single or repetitive action potentials (APs) propagated at a high speed (150 \mu m/s) in constant amplitude and rate of rise in the cytoplasm bearing ryanodine receptors, and then in the slower, waning manner in the deeper region. Ryanodine (10 \mu M), a ryanodine receptor blocker (and/or a half opener), or thapsigargin (1–2 \mu M), a Ca\textsuperscript{2+}-pump blocker, or \omega-conotoxin GVIA (\omega-CgTx, 1 \mu M), a N-type Ca\textsuperscript{2+} channel blocker, blocked the fast propagation, but did not affect the slower spread. Ca\textsuperscript{2+} entry thus triggered the regenerative activation of Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release (CICR) in the submembrane region, followed by buffered Ca\textsuperscript{2+} diffusion in the deeper cytoplasm. Computer simulation assuming Ca\textsuperscript{2+} release in the submembrane region reproduced the Ca\textsuperscript{2+} dynamics. Ryanodine or thapsigargin decreased the rate of spike repolarization of an AP to 80%, but not in the presence of iberiotoxin (IbTx, 100 nM), a BK-type Ca\textsuperscript{2+}-activated K\textsuperscript{+} channel blocker, or \omega-CgTx, both of which decreased the rate to 50%. The spike repolarization rate and the amplitude of a single AP-induced rise in [Ca\textsuperscript{2+}], gradually decreased to a plateau during repetition of APs at 50 Hz, but reduced less in the presence of ryanodine or thapsigargin. The amplitude of each of the [Ca\textsuperscript{2+}], rise correlated well with the reduction in the IbTx-sensitive component of spike repolarization. The apamin-sensitive SK-type Ca\textsuperscript{2+}-activated K\textsuperscript{+} current, underlying the afterhyperpolarization of APs, increased during repetitive APs, decayed faster than the accompanying rise in [Ca\textsuperscript{2+}], and was suppressed by CICR blockers. Thus, ryanodine receptors form a functional triad with N-type Ca\textsuperscript{2+} channels and BK channels, and a loose coupling with SK channels in bullfrog sympathetic neurons, plastically modulating AP.

KEY WORDS: Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release • intracellular Ca\textsuperscript{2+} dynamics • spike broadening • afterhyperpolarization • plasticity of excitability

INTRODUCTION

Intracellular Ca\textsuperscript{2+} ([Ca\textsuperscript{2+}],)\textsuperscript{1} in neurons plays important roles in exocytosis, regulation of cell membrane excitability, cell growth, and gene expressions (see review by Berridge, 1998). To achieve these actions effectively, Ca\textsuperscript{2+}-sensing molecules responsible for these functions should exist close to sources of the Ca\textsuperscript{2+}. At presynaptic terminals, voltage-gated Ca\textsuperscript{2+} channels are coupled with proteins involved in exocytosis of neurotransmitters (Llinàs et al., 1992; Heidelberger et al., 1994; Stanley, 1997). In some sorts of neurons, specific types of voltage-gated Ca\textsuperscript{2+} channels are closely located to specific kinds of Ca\textsuperscript{2+}-activated K\textsuperscript{+} channels, presumably for the effective regulation of cell membrane excitability (Marrion and Tavalin, 1998).

These regulatory actions of Ca\textsuperscript{2+} can be amplified, if Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release (CICR; Endo et al., 1970) via Ca\textsuperscript{2+} release channels augments the rise in [Ca\textsuperscript{2+}], created near the primary source of Ca\textsuperscript{2+}, as known in muscles (reviewed by Endo, 1977; Kuba, 1994; Berridge, 1998; Verhratsky and Petersen, 1998). In several types of neurons, CICR is known to generate rhythmic membrane hyperpolarizations (Kuba and Nishi, 1976) and the afterhyperpolarization (AHP) of an action potential (AP; Kuba et al., 1983; Kawai and Watanabe, 1989; Sah and McLachlan, 1991; Yoshizaki et al., 1995; Cohen et al., 1997; Tanabe et al., 1998) and to modulate transmitter exocytosis (Smith and Cunnane, 1996; Narita et al., 1998). There is little evidence, however, for the close functional coupling of ryanodine receptors to both the Ca\textsuperscript{2+}-sensing target molecules and Ca\textsuperscript{2+} channels providing Ca\textsuperscript{2+} to trigger CICR, although the functional coupling of ryanodine receptors, voltage-gated Ca\textsuperscript{2+} channels and exocytotic machinery was recently reported (Narita et al., 2000).

\textsuperscript{1}Abbreviations used in this paper: \omega-CgTx, \omega-conotoxin GVIA: AHP, afterhyperpolarization; AP, action potential; [Ca\textsuperscript{2+}]\textsubscript{i}, intracellular Ca\textsuperscript{2+}; CICR, Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release; ER, endoplasmic reticulum; IbTx, iberiotoxin; PIS, pseudo-intracellular solution.
In bullfrog sympathetic ganglion cells, CICR occurs in response to the action of caffeine (Kuba and Nishi, 1976; Hernández-Cruz et al., 1990) or a long (>200 ms) depolarization (Kuba et al., 1992; Friel and Tsien, 1992; Hua et al., 1993). This activation of CICR appears to take place predominantly in the cytoplasm within 5 μm from the cell membrane and around the nucleus (Hua et al., 2000). In the present study, we have investigated where, how and in what timing CICR occurs in response to Ca²⁺ entry evoked by APs and what types of Ca²⁺-dependent K⁺ channels are activated by CICR in bullfrog sympathetic neurons, using whole-cell patch-clamp and confocal laser-scanning fluorescence imaging techniques. To minimize the exogenous buffering effect of a Ca²⁺ indicator, we used Oregon green BAPTA-1 at a concentration (10 μM) much lower than the concentrations used in the previous studies (50 ~ 100 μM, Fura-2 or Indo-1; Nohmi et al., 1992; Hua et al., 1993).

The results demonstrated that CICR indeed occurs in the submembrane region in response to Ca²⁺ influx via N-type Ca²⁺ channels activated during the spike of an AP. This CICR directly shapes the spike repolarization of the AP by opening BK-type Ca²⁺-dependent K⁺ channels (BK channels) and the AHP of the AP by activating SK-type Ca²⁺-dependent K⁺ channels (SK channels). The activation of BK channels was gradually decreased during repetition of APs by the graded reduction of CICR, broadening the spike duration. Whereas the CICR-dependent SK channel activity was also decreased during repetitive APs, the total activation of SK channels was increased by the increase in Ca²⁺ entry due to the prolonged spike duration and the accumulation of Ca²⁺. Ryanodine receptors thus form a functional triad with N-type Ca²⁺ channels and BK channels and a loose coupling with SK channels in bullfrog sympathetic ganglion cells, which plastically modulate the cell membrane excitability. Part of these findings was published in abstract form (Akita and Kuba, 1999a,b).

**Materials and Methods**

**Cell Preparations**

Adult bullfrogs (*Rana catesbeiana*) were killed by pithing. Neurons of paravertebral sympathetic ganglia were dissociated and cultured as previously described (Hua et al., 1995). In brief, after dissociation in the presence of collagenase (3 mg/ml; Sigma-Aldrich) and trypsin (0.5 mg/ml; Sigma-Aldrich), neurons were placed on coverglasses and incubated at room temperature over 1 wk. The culture medium was based on L-15 (diluted to 70%; GibCO BRL), supplemented with 6% fetal calf serum (Nakashibetsu), 20 ng/ml nerve growth factor (Wako Chemicals, Inc.) and 50 μg/ml gentamicin (GibCO BRL). Large, spherical cells, 40–60 μm in diameter along their long axis and presumably B cell origin, were selected for the experiments.

**Electrophysiology**

A whole-cell patch-clamp technique (Hamill et al., 1981) was applied to cultured neurons at room temperature by using single electrode amplifiers (8100-I from Dagan Corp., Minneapolis, MN; or CEZ-3100 from Nihon-Kohden, Tokyo, Japan). Patch electrodes (6–8 MΩ) were filled with a solution containing (mM): 120 KCl; 2 MgCl₂; 4 Na₃ATP; 0.3 NaGTP; 10 HEPES, and 0.01 Oregon green BAPTA-1 (OGB-1), pH 7.2. The composition of Ringer’s solution was (mM): 115 NaCl; 2.5 KCl; 1 MgCl₂; 2 CaCl₂; 5 HEPES, and 10 D-glucose, pH 7.2. Voltage-gated Ca²⁺ currents were recorded in a solution containing (mM): 119 TEA-Cl; 2 CaCl₂; 5 HEPES; 10 D-glucose; supplemented with 100 nM tetrodotoxin, pH 7.2, using patch pipettes filled with a solution, in which K⁺ was replaced by Cs⁺. The liquid junction potential of Ringer’s solution to the K⁺-based electrode solution and that of the TEA-based solution to the Cs⁺-based electrode solution were 11 and 20 mV, respectively.

APs were evoked by brief current pulses of 1–2 ms duration to avoid overlapping the first derivatives of the spike repolarization with the current pulses. The derivative of the spike of AP was taken to estimate the membrane current underlying spike repolarization. The rate of spike repolarization showed either single or double negative peaks. When BK channels were blocked by ibotenic acid (IbTX), the spike repolarization occasionally became diphasic in the cells, which normally showed a monophasic spike repolarization (see Fig. 10 B, b). Furthermore, IbTX predominantly decreased the initial negative peak of the diphasic derivative of spike repolarization in a considerable number of cells (see Fig. 10 A, b). The initial peak was therefore analyzed to estimate the relative change in the BK channel component of spike repolarization. The data were stored on DAT tapes through TEAC RD-135T recorder, and then analyzed with software (pCLAMP6 from Axon Instruments, Inc.; or Excel from Microsoft Co.). The current and voltage records were filtered at 3–5 kHz with a 4-pole Bessel filter (NF Corporation) before the analysis. Drugs were applied by extracellular perfusion. To avoid photobleaching, nifedipine was applied under the nearly dark condition except for laser light for line scanning.

To record the current underlying AHP (I_AHP), a bridge current clamp mode was electronically switched to a voltage clamp mode at the end of the spike of AP (hybrid clamp; Lancaster and Adams, 1986). For analysis, the current obtained was fitted to an exponential decaying function with a nonlinear least square fitter (Levenberg-Marquadt routine) of software (Mathcad2000; Mathsoft Inc.). The switching was made at the time when the spike repolarization of a single AP or the last of repetitive APs reached the holding potential level (~50 mV) to avoid a capacitative surge. The timing of the switching was determined under the control condition but fixed throughout a series of the experiment. This yielded comparison of I_AHP (i.e., membrane conductance) under different conditions at the same timing after the beginning of Ca²⁺ entry and the resultant CICR. When the duration of the spike of an AP became longer in some conditions, however, there was a positive shift of membrane voltage relative to the holding potential at the time of the switching. In this case, the starting point of the fitting was set at 1–2 ms after the switching and the amplitude of I_AHP was estimated at the time of the switching by extrapolation. Another drawback was that the comparison of voltage-sensitive components of the current at the same voltage level became not possible due to the voltage shift. The switching at the higher voltage and the faster timing (1–2 ms at most) might overestimate the initial amplitude of the current relaxation of voltage-gated K⁺ currents relative to that expected to be obtained at ~50 mV. On the other hand, it was possible that a component of the voltage-sensitive K⁺ conductance that would have been activated during the rest of repolarization phase (that was abolished by voltage-clamping) was lost in the record of the current relaxation produced at the earlier switching. Thus, these two opposite effects could be compromised. The
extent of this overestimation would be less than the difference between the amplitude of I_{AP} at the time of switching and the amplitude at the time, at which the prolonged spike repolarization was expected to reach −50 mV. The difference was 28%, for instance, when IbTx prolonged the fast decay time constant of the I_{AP} evoked by 1 AP to 8 ms (see Fig. 14 B, b).

$[Ca^{2+}]$, Measurement

Neurons were loaded with a Ca^{2+} indicator, OGB-1 (10 μM), through a patch pipette. Equilibrium of OGB-1 loading was attained 15–20 min after the opening of a membrane patch. Neurons were scanned with blue laser (488 nm, 0.45 mW) using a confocal scan unit (MRC-600; Nippon BIO-RAD Laboratories) attached to an inverted microscope (TMD-300; water immersion 40× objective, NA 1.15; Nikon). The diameter of the confocal aperture was set to 3.1 μm, yielding the lateral and axial resolutions of 0.15 and 1.25 μm, respectively (measured at the half maximum decay length of the fluorescence of a 0.21-μm fluorescent bead). The whole image of a neuron was initially obtained by a XY scan mode, and then the neuron was line-scanned at the time resolution of 2 or 4 ms across the cell soma.

The in vivo $K_t$ of OGB-1 was determined by the modified version of a calibration method (Tokimasa et al., 1997). In brief, cells were immersed in a pseudo-intracellular solution (PIS) containing each $[Ca^{2+}]_i$, (0, 0.1, 0.3, 1, and 100 μM) and 20 μM digitonin for 25 min. The cells were then incubated in digitonin-free PIS containing 10 μM OGB-1 for 15 min. After removal of the PIS, the cells were briskly rinsed once with dye-free PIS and then covered with liquid paraffin. The fluorescence intensity in the cells (see Fig. 2, A and B). $[Ca^{2+}]_i$ values were checked by confocal laser microscopy with two-wavelength Ca^{2+} indicators in the previous study (Nohmi et al., 1992). In most cases, the maximum amplitude at the time, at which the prolonged spike repolarization (1 ms) every 20 ms was involved in the outermost shell for the plasma membrane. (Although this formulation does not represent the real mechanism of CICR, it is appropriate at least to test whether the additional source of Ca^{2+} plays a role in the spatial and temporal characteristics of AP-induced Ca^{2+} transients recorded experimentally.) The timing of the flux in each shell was delayed by 0.5 ms from the outer adjacent shell. The amplitude of the flux was assumed to decrease exponentially (time constant; 50 ms) until the end of the fifth pulse. The distribution of the sources of Ca^{2+} was divided into two parts of 2 μm in depth; the outer high-density region and the inner low-density re-
The initial amplitude of the flux in each shell was set to be 8.8 \times 10^{-5} \text{ mol/s} (equivalent to 1.7 nA of Ca^{2+} current) in the high-density region and 4.4 \times 10^{-5} \text{ mol/s} (equivalent to 0.85 nA of Ca^{2+} current) in the low-density region. Noncooperative, Michaelis-Menten type Ca^{2+} pumps and leakage at the cell membrane were included in the outermost shell (Sala and Hernández-Cruz, 1990). The maximum speed of the pump at the cell membrane was set to be 1.4 \times 10^{-4} \text{ mol/cm}^2 \text{s}. The Ca^{2+} concentration for its half-maximal activation was 0.83 \text{ mM}. Endogenous fixed rapid buffers were assumed only in the submembrane region of 4 \mu m in depth. \beta_i was set to be 75 (Neher and Augustine, 1992). D_m of 120 \mu m^2/s yielded the best result, a slightly higher value than that of fura-2 in axons of Aplysia neurons (102.5 \mu m^2/s; Gabso et al., 1997).

The second type of simulation (see Fig. 4, A and B, b) incorporated only Ca^{2+} entry at the plasma membrane and the mobile buffer (for the Ca^{2+} indicator) in the cytoplasm. Ca^{2+} release, endogenous fixed buffers (except for the outermost shell), Ca^{2+} extrusion and uptake were not included. A greater flux was assumed for Ca^{2+} entry (3.2 nA). Other parameters were the same as in the first type.

The third type of simulation (see Fig. 4, A and B, c) incorporated Ca^{2+} entry, Ca^{2+} release, fixed and mobile buffers, Ca^{2+} extrusion and uptake. The amount of Ca^{2+} entry was the same as in the first type of simulation. Ca^{2+} release was similarly defined as those of the first type, but the magnitude was fixed constant during its repetition. The maximum rate of Ca^{2+} pumping at Ca^{2+} storing organelles was assumed 64\% \ (0.9 \times 10^{-4} \text{ mol/cm}^2 \text{s}) of that at the cell membrane with the same affinity. The area occupied by the pumps in each compartment was assumed equal to the surface area of the cell for the outer 2-\mu m region and its half for the deeper 2-\mu m region. Other parameters were the same as those in the first type.

**Chemicals**

Thapsigargin, ryanodine, and nifedipine were purchased from Wako Chemicals, Inc. Iberiotoxin, apamin, \omega-conotoxin GVIA, digitonin, TEA, and HEPES were from Sigma-Aldrich. OGB-1 and BODIPY FL-X ryanodine were from Molecular Probes. TTX was from Alomone Labs.

**Statistics**

Each data is shown by mean \pm SEM. Student’s paired t test was performed for statistical comparison with Microsoft Excel software.

**RESULTS**

**CICR Occurs in the Submembrane Regions**

Localization of Ryanodine Receptors in Bullfrog Sympathetic Neurons. Ryanodine receptors in bullfrog sympathetic neurons were stained with fluorescent ryanodine, BODIPY FL-X ryanodine (0.5 \mu M). To subtract nonspecific binding of fluorescent ryanodine, the following protocol was adopted. Ganglion cells in culture (Fig. 1...
Akita and Kuba

A) were first incubated in a solution containing both fluorescent ryanodine (0.5 μM) and ryanodine (10 μM) for 10 min. Under this condition, there was weak fluorescence in narrow regions close to the cell membrane and perinuclear regions (Fig. 1 B). After washing with Ringer’s solution for 30 min (Fig. 1 C), fluorescent ryanodine (0.5 μM) alone was applied for 8 min. This procedure gave strong fluorescence in almost the same cytoplasmic regions (Fig. 1 D) as those showing weak fluorescence in Fig. 1 B. The subtraction of the image in Fig. 1 B from that in Fig. 1 D showed a similar distribution of the fluorescence (Fig. 1 E) to those in Fig. 1, B and D, indicating the negligible nonspecific binding of fluorescent ryanodine in the cytoplasm of ganglion cells. Thickness of the submembrane distribution ranged from 1 to 3 μm in most cases (n = 14). With some cells, as in Fig. 1, the distribution of ryanodine receptors was also recognized around the inner boundary of the nucleus. In such cases, the perinuclear fluorescence was likely to reach the submembrane region, comprising a wider (5–6 μm) distribution of the fluorescence. This indicates the intracellular network of Ca^{2+} stores endowed with ryanodine receptors, which may transmit Ca^{2+} signals from the cell membrane toward the nucleus. In this study, we focus on the mode of CICR activation in the submembrane region related to the modulation of cell membrane excitability, whereas the mechanism of CICR activation in the perinuclear region will be reported elsewhere.

Fast and Slow Propagation of a Ca^{2+} Transient in the Cytoplasm Evoked by an AP or APs. A region of the cytoplasm within 12 μm from the cell membrane (Fig. 2 A) was line-scanned with a confocal laser microscope before, during, and after current pulse stimulation. This provided the time courses of single-dimensional changes in fluorescence evoked by 1–10 APs (50 Hz), representing those of [Ca^{2+}], (Ca^{2+} transients; Fig. 2, B–D). Ca^{2+} transients initially occurred in the cytoplasm just beneath the cell membrane and spread toward the deeper cytoplasm. The propagation front of the increased [Ca^{2+}] showed two phases; fast and slow. The fast propagation front was almost flat and clearly demarcated from the resting fluorescence, spanning 1.5–2 μm beneath the plasma membrane (Fig. 2 B). The width of the fast phase was analogous to that of the distribution of ryanodine receptors identified by fluorescent ryanodine (Fig. 1 E). On the other hand, the slow phase following the fast phase showed the dull front...
plane of the propagation and became wider as the number of APs was increased (Figs. 2 B and 3 B).

Regenerative Activation of CICR in the Fast Propagation Front of a Ca$^{2+}$ Transient. Similarity between the width of the distribution of ryanodine receptors and that of the fast propagation phase of a Ca$^{2+}$ transient suggests the regenerative activation of CICR during the fast propagation. The time courses of a single AP-induced Ca$^{2+}$ transient at discrete points in the fast propagating region were quite similar. The amplitude and rate of rise of the single AP-induced Ca$^{2+}$ transient only slightly decreased with an increase in distance from the cell membrane and slightly delayed in peak time (traces 1 and 2 in Fig. 3, D and E, a). The peak amplitude of the Ca$^{2+}$ transient induced by a single AP or the first of APs in a train was on average 58.5 ± 13.3 nM ($n = 31$). Likewise, the time courses of a five AP-induced Ca$^{2+}$ transient in the fast propagation phase followed a similar trend (traces 1 and 2 in Fig. 3, D and E, b; see filled squares in Fig. 6 B). The peak amplitude of a five APs-induced Ca$^{2+}$ transient within the fast phase was 167.2 ± 28.8 nM.

The characteristics of the propagation of Ca$^{2+}$ transients are also shown in their spatial profiles (Fig. 4 A). As APs were successively delivered on the cell, the spatial profiles of [Ca$^{2+}$]$_i$, within 2 μm beneath the plasma membrane, corresponding to the fast propagation region, became gradually flat. Computer simulation assuming only the diffusion process without fixed buffers and extrusion processes (both of which should slow Ca$^{2+}$ diffusion) for intracellular Ca$^{2+}$ dynamics failed to demonstrate the formation of the flat propagation region (Fig. 4 A, b). Instead, the simulation including the additional sources of Ca$^{2+}$ in the submembrane region nicely reproduced the experimental data (Fig. 4 A, a). With this assumption of the additional sources of Ca$^{2+}$, the time courses of 5APs-induced Ca$^{2+}$ transients were also reconstructed well (Fig. 4 B, a), whereas the assumption of no additional Ca$^{2+}$ sources failed to reproduce them (Fig. 4 B, b). Thus, the flat gradient of
the rise in \([\text{Ca}^{2+}]\), in the submembrane region must be caused by the activation of additional sources of \(\text{Ca}^{2+}\) during the fast propagation, namely the regenerative activation of CICR, producing a “\(\text{Ca}^{2+}\) wave” in the fast phase. The propagation velocities of the \(\text{Ca}^{2+}\) wave (see MATERIALS AND METHODS) evoked by a single and five APs were \(183.7 \pm 17.8 \text{ m/s}\) and \(142.5 \pm 15.9 \text{ m/s}\) (\(n = 31\)), respectively.

**Blockade of the Fast \(\text{Ca}^{2+}\) Wave Propagation by Ryanodine and Thapsigargin.** The final test of the activation of CICR in the submembrane region is to observe the actions of blockers of \(\text{Ca}^{2+}\) release or uptake on AP-induced \(\text{Ca}^{2+}\) transients. Ryanodine (10 \(\mu\text{M}\)) or thapsigargin (1–2 \(\mu\text{M}\)) raised the basal level of \([\text{Ca}^{2+}]\), (Fig. 2, C and D). The magnitudes of the increases were \(65.8 \pm 26.6 \text{ nM} (n = 11)\) or \(58.8 \pm 22.3 \text{ nM} (n = 12)\), respec-
tively, at 10 min after the application. Although the increased basal \([Ca^{2+}]\), remained over 30 min in most cells, some cells showed recovery and/or reduction of the basal level within 30 min by 17.9 \(\pm\) 4.3 nM (4/15) or 9.2 \(\pm\) 2.4 nM (3/15) from the base level of the control even in the presence of ryanodine or thapsigargin, respectively. Sometimes a transitory, spatially inhomogeneous increase in the basal \([Ca^{2+}]\), occurred during the course of the application of CICR blockers (Fig. 2 C; 1AP and 2APs). Its extent differed from cell by cell, but it finally disappeared in most cases (Fig. 2 D).

Ryanodine (10 \(\mu\)M) decreased \(Ca^{2+}\) transients in the submembrane region evoked by a single AP or the first AP in a train to 49.1 \(\pm\) 9.1\% of the control, and the total amplitude of five APs-induced \(Ca^{2+}\) transients to 50.1 \(\pm\) 7.2\% at 10 min after its application (see Fig. 7 C, a for a single AP or the first AP in a train). Likewise, thapsigargin (1–2 \(\mu\)M) depressed \(Ca^{2+}\) transients evoked by a single AP or the first AP in a train to 46.6 \(\pm\) 10.7\%, and the total amplitude of five APs-\(Ca^{2+}\) transients to 50.4 \(\pm\) 8.9\% at 10 min after the application (Figs. 5, 6 B, and 7, A and B, a). The rate of rise in \([Ca^{2+}]\), by a single AP or the first AP in a train was also reduced similarly to 47.0 \(\pm\) 7.6\% by ryanodine and to 42.1 \(\pm\) 8.9\% by thapsigargin. Under this condition, the \(Ca^{2+}\) wave in the fast phase disappeared leaving only the slow, waning mode of propagation (Figs. 5 and 6). The full blocking action of ryanodine or thapsigargin on AP-induced \(Ca^{2+}\) transients was seen within 1–3 min in some cells, but mostly it took over 30 min. The actions of CICR blockers were not due to the saturation of \(Ca^{2+}\) indicators (see MATERIALS AND METHODS, and also Fig. 2 legend) and the decrease in \(Ca^{2+}\) entry caused by the action of the blockers on voltage-gated \(Ca^{2+}\) channels (see the next section and Fig. 8). Computer simulation with the assumption of the reduction in \(Ca^{2+}\) release to 50\% of the control and the increased basal level of \([Ca^{2+}]\), with unchanged \(Ca^{2+}\) entry also reproduced the effects of CICR blockade on the time course and the spatial spread of five APs-induced \(Ca^{2+}\) transients (Fig. 6 A).

The spatial gradient of \(Ca^{2+}\) transients induced by one or five APs disappeared at 400–500 ms after the beginning of stimuli (Fig. 3 E). The decay time course of an increased \([Ca^{2+}]\), over that period would reflect the rate of \(Ca^{2+}\) clearance in the cytoplasm, i.e., \(Ca^{2+}\) extrusion and uptake. The blockade of \(Ca^{2+}\) pump or the opening of ryanodine receptors on \(Ca^{2+}\) stores is then expected to retard the rate of this \(Ca^{2+}\) clearance. Unexpectedly, however, both ryanodine and thapsigargin shortened the decay time constant of the AP-induced \(Ca^{2+}\) transient over 400 ms after the beginning of stimuli in most cases (Fig. 7 B, a and inset). The decay time constant of the five APs-transient (822.9 \(\pm\) 98.4 ms) decreased to 72.8 \(\pm\) 14.2\% \((P < 0.01)\) by the application

![Figure 5. Effects of thapsigargin on AP-evoked \(Ca^{2+}\) transients.](https://example.com/image5.png)
Akita and Kuba

of ryanodine for 10 min and to 70.2 ± 12.3% (P < 0.01) by thapsigargin. One possible explanation might be that CICR remains to be activated to some extent after the end of Ca²⁺ entry and is eliminated by the blockers. Alternatively, Ca²⁺ release from mitochondria, which may normally occur during the late decay phase of Ca²⁺ transient following Ca²⁺ uptake during the rising phase (Colegrove et al., 2000), could be reduced for the suppression of Ca²⁺ transients under the blockade of CICR. This issue needs to be studied.

Reduction of Ca²⁺ Entry Does Not Explain the Depressant Actions of CICR Blockers. Run-down of Ca²⁺ current (I_{Ca}) during the course of whole-cell recording or the blockade of voltage-gated Ca²⁺ channels by CICR blockers might explain the depressant actions of the drugs on the Ca²⁺ transients. We first examined the run-down of I_{Ca} induced by a voltage pulse (10–50 ms). The rate of run-down of I_{Ca} varied among cells and patch-clamp conditions. The amplitude of I_{Ca} decreased on average by 15–20% (n = 24) at 8 min after the opening of the membrane patch (Fig. 8 A) and so was the accompanying Ca²⁺ transient (by 23.2 ± 6.5%, n = 8; data not shown). These reductions were much smaller than the decrease (50%) in AP-induced Ca²⁺ transients produced by ryanodine or thapsigargin (Fig. 7, A–C, a). Next, we examined effects of ryanodine (10 μM) and thapsigargin (1–2 μM) on I_{Ca} and its run-down. The blockers affected neither the amplitude nor the rate of run-down of I_{Ca} (Fig. 8 A). In some cells, the depressant effects of ryanodine or thapsigargin on the amplitude of Ca²⁺ transients appeared long after its application (Fig. 8 B, c). Even in these cells, where the I_{Ca} rundown proceeded to a fair extent, the amplitude of Ca²⁺ transient was not recovered after raising the extracellular Ca²⁺ and restoring the amplitude of the I_{Ca} (Fig. 8 B, d). Accordingly, the depressant actions of ry-
Functional Triads of Ion Channels

anodine and thapsigargin did not result from the reduction of ICa due to its blockade or run-down. The results also indicate that the principal mechanism of the increase in [Ca2+]i evoked by an AP or APs is CICR in bullfrog sympathetic ganglion cells, similar to the mechanism in cardiac muscles. This is also supported by the successful simulation of AP-induced Ca2+ transients with the assumption of a large Ca2+ release 17 times greater than Ca2+ entry (see MATERIALS AND METHODS).

Graded Reduction of the Submembrane CICR during Repetitive APs. Another remarkable feature of APs-induced Ca2+ transients was the progressive decrease in the rises in [Ca2+]i produced by individual APs during repetitive stimulation. The amplitude and rate of rise in [Ca2+]i evoked by the first AP were the largest and those by the subsequent APs became progressively smaller (Fig. 3 D, b, and Fig. 7 B, a). The amplitude of the rise evoked by the fifth AP (25.2 ± 3.9 nM) was 43.4 ± 3.6% of the first. Similarly, the rate of rise in [Ca2+]i induced by the
first AP was 7.1 ± 1.5 μM/s, whereas that induced by the fifth AP was 46.7 ± 5.5% of the first.

There are several possible mechanisms for the progressive decrease in the [Ca\(^{2+}\)]\(_i\) rise. The acceleration of Ca\(^{2+}\) uptake into Ca\(^{2+}\) stores, say via thapsigargin-sensitive pumps, might have reduced the net Ca\(^{2+}\) flux during the later part of stimuli. This is unlikely, however, according to the simulation assuming the Ca\(^{2+}\) entry and release of constant magnitude and the Michaelis-Menten type Ca\(^{2+}\) pumping in the submembrane region, the strength of which was adjusted so that the peak [Ca\(^{2+}\)]\(_i\) values reached by the individual simulated Ca\(^{2+}\) rises were matched to those of the experimental data. Although the simulation apparently reproduced the spatial profiles of [Ca\(^{2+}\)]\(_i\), averaged over the period of 10 ms after the first and third AP (Fig. 4 A, c), it was unable to reconstruct the progressive reduction in amplitude and the decay phase of the individual [Ca\(^{2+}\)]\(_i\) rise (Fig. 4 B, c). Varying the pump speed, cooperativity or the affinity for Ca\(^{2+}\) also failed to improve the large discrepancy between the results of simulation and the observations (not shown). Mitochondrial Ca\(^{2+}\) uptake that responds to the microdomains of high [Ca\(^{2+}\)]\(_i\) (Rizzuto et al., 1998) is also unlikely for the mechanism of the progressive reduction of Ca\(^{2+}\) transients, since its operation could be regarded as the rapid buffering system (i.e., β in our simulation) for its fast kinetics (see reviews by Carafoli, 1987 and Rizzuto et al., 1999).

Another possible mechanism would be the progressive decline in Ca\(^{2+}\) release during repetition of APs. This was again tested by computer simulation assuming the exponential decay in Ca\(^{2+}\) release flux until the end of repetitive pulses. This simulation not only yielded the progressive reduction in the amplitude of individual Ca\(^{2+}\) rises during repetitive APs, but also reproduced the spatial profiles of the Ca\(^{2+}\) transient and the decay phases of the individual Ca\(^{2+}\) rises during repetitive APs, which may be the inactivation of Ca\(^{2+}\) release (Fabiato, 1985) or Ca\(^{2+}\) depletion. Ca\(^{2+}\) depletion, however, may have not occurred during repetition of 5–10 APs, since CICR blockers largely in-

![Figure 8](image_url)
increased the basal \([\text{Ca}^{2+}]_{\text{i}}\) (Fig. 2, C and D) and took longer time to cease CICR. We will report the details of the property of the inactivation of CICR elsewhere.

**Ca2**\(^{+}\) Influx through N-type Ca2**\(^{+}\) Channels Triggers CICR.** Bullfrog sympathetic ganglion cells are endowed predominantly with N-type Ca2**\(^{+}\) channels, and less with L- and other types of Ca2**\(^{+}\) channels (Jones and Marks, 1989a; Elmslie et al., 1994). \(\omega\)-CgTx at 1 \(\mu\)M drastically blocked Ca2**\(^{+}\) transients induced by a single AP to 18.3 ± 6.5\% (\(n = 8\); Fig. 9 A, a) and those by 10 APs to 32.2 ± 8.2\% (not shown) at 10 min. Nifedipine (20 \(\mu\)M), however, did not affect single AP-induced Ca2**\(^{+}\) transients (96.6 ± 12.9\%, \(n = 6\); Fig. 9 B, a) and slightly reduced 10 APs-induced Ca2**\(^{+}\) transients to 79.6 ± 8.2\% (\(n = 6\)), which was the same as that caused by the run-down of ICa (see above). Thus, Ca2**\(^{+}\) entry via N-type Ca2**\(^{+}\) channels activates CICR in bullfrog sympathetic neurons.

**Passive Propagation of Ca2**\(^{+}\) Transients in the Deeper Cytoplasm.** As single or five APs-induced Ca2**\(^{+}\) transients spread into the region deeper than a few micrometers from the plasma membrane, they decreased progressively in amplitude, rate of rise, and peak time (traces 3–5 in Fig. 3 B, D and E; open squares in Fig. 6 B). The passive property of Ca2**\(^{+}\) propagation in the slow phase was clearly shown in the spatial and temporal profiles of the rise in \([\text{Ca}^{2+}]_{\text{i}}\), (Figs. 3 and 4 A). During the train of stimuli, the spatial profiles in the slow phase showed gradual decreases in \([\text{Ca}^{2+}]_{\text{i}}\), toward the center of the cell (Fig. 4 A). As the amplitude of the rise in \([\text{Ca}^{2+}]_{\text{i}}\) in the fast phase decreased after the end of the stimuli, the \([\text{Ca}^{2+}]_{\text{i}}\) in the deeper region of the slow phase slowly increased until the spatial gradient of \([\text{Ca}^{2+}]_{\text{i}}\) disappeared (traces 3–5 in Fig. 3 D), and then decayed over hundreds of milliseconds to several seconds (Fig. 3 E). Thus, only the buffered Ca2**\(^{+}\) diffusion process would contribute to the propagation in the deeper cytoplasm. Computer simulation with no assumption of Ca2**\(^{+}\) release, fixed Ca2**\(^{+}\) buffering and pumping in the deeper cytoplasm also reconstructed well the spatial profiles of the rise in \([\text{Ca}^{2+}]_{\text{i}}\), both in the absence and presence of CICR blockers (Figs. 4 A and 6 A, a).

The passive property of the propagation can also be demonstrated by the spatial decay in the amplitude and rate of rise of a five APs-induced Ca2**\(^{+}\) transient (Fig. 6 B, a and b). The spatial decay of the amplitude and rate of rise were fitted by single exponentials, yielding their “length constants” (\(\lambda_{\text{amp}}\) and \(\lambda_{r}\)) in the slow propagation phase. \(\lambda_{\text{amp}}\) and \(\lambda_{r}\) were found to be 6.4 ± 0.6 \(\mu\)m and 3.4 ± 0.3 \(\mu\)m, respectively. These length constants were not significantly affected by ryanodine and thapsigargin (99.1 ± 2.3\% or 99.7 ± 2.0\% of the control for \(\lambda_{\text{amp}}\) and 100.5 ± 3.8\% or 102.5 ± 3.8\% for \(\lambda_{r}\), respec-
The results also support the predominance of the diffusion process in the deeper cytoplasm.

### CICR Shapes the Spike Repolarization and AHP of APs

Possible target molecules of CICR in the submembrane region would be two types of Ca²⁺-activated K⁺ channels: BK channels (Adams et al., 1982) facilitating the spike repolarization and SK channels generating AHP of APs (Pennefather et al., 1985). Thus, it is possible that CICR shapes directly the spike and AHP of an AP that triggers CICR.

Blocking CICR gave three modes of actions in shaping APs. First, the maximum rate of spike repolarization of an AP or the first AP in a train was reduced to 79.1 ± 8.3% (P < 0.01, n = 11) of the control (−76.7 ± 6.9 V/s, n = 37) by ryanodine (10 μM; Fig. 7 C, d) and to 81.9 ± 4.4% (P < 0.01, n = 12) by thapsigargin (1–2 μM; Fig. 7 A, d) after their application for 10 min. Those of the fifth spike in a train at 50 Hz were 70.1 ± 5.8% (not shown) and 70.3 ± 4.1% (Fig. 7 B, d), respectively. These reductions were not due to the rundown of any types of ion channels because of intracellular perfusion, since there was no change in the slope of spike repolarization during the same period in the absence of the blockers (102.7 ± 1.4%, P > 0.1; not shown). Second, the amplitude of the AHP of an AP was decreased to 85.5 ± 6.8% (n = 11, P < 0.01) of the control (20.4 ± 0.9 mV) by ryanodine (Fig. 7 C, c and d) or to 88.9 ± 2.8% (n = 12; P < 0.01) by thapsigargin (Fig. 7 A, c and d). Those of the fifth AP in a train at 50 Hz were 85.9 ± 5.2% (not shown) and 86.4 ± 3.7% (Fig. 7 B, c) of the control (17.2 ± 1.1 mV), respectively, the degree of which were not significantly different from those of a single AP (P > 0.1). Third, the half decay time of the AHP of an AP was shortened to 85.7 ± 9.3% (control, 34.0 ± 4.7 ms) by ryanodine (Fig. 7 C, c) or to 80.1 ± 11.7% by thapsigargin (Fig. 7 A, c). Those of the fifth AP in a train were 86.1 ± 7.8% (control, 144.6 ± 28.7 ms) by ryanodine (not shown) or to 78.7 ± 7.2% by thapsigargin (Fig. 7 B, c). Again, the extent of the decrease was not different from that of a single AP (P > 0.1).

**Spike-triggered CICR Activates BK-type Ca²⁺-dependent K⁺ Channels for Spike Repolarization.** It is likely that the broadening of the spike of AP and the reduction in the amplitude of AHP under the blockade of CICR are caused by the reduction in BK channel activity, for the known role of BK channels in spike repolarization. To examine how many fractions of spike repolarization is attributed to BK channel activity, we first examined the effect of IbTx on spike repolarization. IbTx (100 nM) markedly prolonged the spike of an AP, decreased its maximum rate of fall to 49.0 ± 3.7% (n = 13) at 10 min after the application (black traces in Fig. 10 A, a and b) and enhanced the single AP-induced Ca²⁺ transient in the submembrane region (Fig. 10 A, c, black trace). In many cells, the derivative of spike repolarization was diphasic (Fig. 10 A, see also Figs. 7 B and 9 A). IbTx decreased its initial phase, but not the late phase (Fig. 10 A, b). In contrast, a blocker of SK channel, apamin (100 nM), had no effect on spike repolarization (100.0 ± 2.7%, P > 0.1, n = 5; Fig. 10 C, a and b) and Ca²⁺ transient.

---

**Figure 10.** CICR activates BK channels during the spike repolarization. (A) Effects of sequential applications of IbTx and ryanodine on the spike (a), its first derivatives (b), and a single AP-induced Ca²⁺ transient (c). Ca²⁺ transients were measured within the region of 2 μm below the plasma membrane and shown in net changes in [Ca²⁺]. IbTx (100 nM) was first applied for 10 min, and then ryanodine (10 μM) was applied for 5 min. Gray traces indicate the records before the application of blockers, whereas black and red traces represent those after the application of IbTx and ryanodine, respectively. The derivatives of the spike during current stimulation were omitted. Vertical arrows indicate the points at which the rates of spike repolarization were measured. Dotted lines were drawn to illustrate no apparent changes in the second component of the rate of spike repolarization by the drugs. (B) Effects of sequential applications of ryanodine and IbTx on the spike (a), its first derivatives (b), and a single AP-induced Ca²⁺ transient (c). Ryanodine was first applied for 10 min, and then IbTx was applied for 10 min. Gray traces indicate the records before the application of the blockers, whereas red and black traces represent those after the application of ryanodine and IbTx, respectively. (C) Effects of apamin (100 nM) on the spike repolarization of AP (a and b) and the Ca²⁺ transient (c). Gray and black traces are the records before and 10 min after the application of apamin, respectively.
sents (Fig. 10 C, c). Thus, the activation of BK channel contributes to 50% of the rate of spike repolarization, predominantly to its initial phase. In the presence of IbTx, ryanodine had little effects on the spike of AP (n = 5, \( P > 0.1 \); red traces in Fig. 10 A, a and b), although it considerably suppressed the accompanying \( \Delta [Ca^{2+}] \) transient (Fig. 10 A, c, red trace). Furthermore, blocking \( Ca^{2+} \) entry by \( \omega \)-CgTx (1 \( \mu \)M) reduced the rate of spike repolarization, preferentially its initial phase, to 52.2 ± 2.6% (n = 7, Fig. 9 A, c and d), whereas nifedipine had no effect (100.1 ± 1.0%, \( P > 0.1 \), n = 8, Fig. 9 B, c and d). Thus, only the BK channel participates in the \( Ca^{2+} \)-dependent \( K^+ \) current for spike repolarization in response to \( Ca^{2+} \) entry via N-type \( Ca^{2+} \) channels and the resultant CICR. (Enhancement of the AP-induced \( Ca^{2+} \) transient by blocking BK channels with IbTx is obviously caused by an increase in \( Ca^{2+} \) entry.)

Next, we asked how many fractions of BK channels are activated by CICR for spike repolarization. We first blocked the CICR-activated component of spike repolarization and the accompanying \( Ca^{2+} \) transients during repetitive APs. (A) Time courses of increases in \([Ca^{2+}]\), induced by 20 APs in the submembrane region and the accompanying AHPs. Upper records are the net increases in \([Ca^{2+}]\), \( \Delta [Ca^{2+}] \) within 2 \( \mu \)m from the plasma membrane, whereas lower records are the AHPs. Gray and black traces are those before and after the application of ryanodine (10 \( \mu \)M), respectively. Each trace of the records of \([Ca^{2+}]\), changes was the average of two records and smoothed by a moving average over 6 ms. Vertical dashed lines indicate the timing of the first to fifth, tenth, and twentieth spike, corresponding to each of those of AP-evoked \( \Delta [Ca^{2+}] \). Horizontal short arrows and bars were added to note the decrease in the individual \( \Delta [Ca^{2+}] \) during the repetitive APs. The first, second, fifth, tenth, and twentieth APs and their first derivatives in a 50-Hz train in the absence and presence of ryanodine, IbTx and \( \omega \)-CgTx. (a) Control APs; (b) APs recorded at 10 min after the application of ryanodine (10 \( \mu \)M). (c) APs recorded at 10 min after the application of IbTx (100 nM) following ryanodine. (d) APs in the presence of \( \omega \)-CgTx (1 \( \mu \)M) applied for 10 min in another cell. The derivatives of the spike during the rising phase were omitted. Vertical arrows indicate the points at which the rates of spike repolarization were measured. (C) Reduction in the rate of spike repolarization during repetitive APs. The maximum rate of fall of each spike was normalized to that of the first in the absence of blockers. The mean ± SEM of pooled data obtained before (circles, n = 31) and after the application of ryanodine (10 \( \mu \)M, squares, n = 11), thapsigargin (1–2 \( \mu \)M, triangles, n = 12), \( \omega \)-CgTx (1 \( \mu \)M, diamonds, n = 8) and IbTx (100 nM, crosses, n = 15) were plotted and bound with straight lines.

**Figure 11.** Reduction in the rate of spike repolarization and the accompanying \( Ca^{2+} \) transients during repetitive APs. (A) Time courses of increases in \([Ca^{2+}]\), induced by 20 APs in the submembrane region and the accompanying AHPs. Upper records are the net increases in \([Ca^{2+}]\), \( \Delta [Ca^{2+}] \) within 2 \( \mu \)m from the plasma membrane, whereas lower records are the AHPs. Gray and black traces are those before and after the application of ryanodine (10 \( \mu \)M), respectively. Each trace of the records of \([Ca^{2+}]\), changes was the average of two records and smoothed by a moving average over 6 ms. Vertical dashed lines indicate the timing of the first to fifth, tenth, and twentieth spike, corresponding to each of those of AP-evoked \( \Delta [Ca^{2+}] \). Horizontal short arrows and bars were added to note the decrease in the individual \( \Delta [Ca^{2+}] \) during the repetitive APs. (B) The first, second, fifth, tenth, and twentieth APs and their first derivatives in a 50-Hz train in the absence and presence of ryanodine, IbTx and \( \omega \)-CgTx. (a) Control APs; (b) APs recorded at 10 min after the application of ryanodine (10 \( \mu \)M). (c) APs recorded at 10 min after the application of IbTx (100 nM) following ryanodine. (d) APs in the presence of \( \omega \)-CgTx (1 \( \mu \)M) applied for 10 min in another cell. The derivatives of the spike during the rising phase were omitted. Vertical arrows indicate the points at which the rates of spike repolarization were measured. (C) Reduction in the rate of spike repolarization during repetitive APs. The maximum rate of fall of each spike was normalized to that of the first in the absence of blockers. The mean ± SEM of pooled data obtained before (circles, n = 31) and after the application of ryanodine (10 \( \mu \)M, squares, n = 11), thapsigargin (1–2 \( \mu \)M, triangles, n = 12), \( \omega \)-CgTx (1 \( \mu \)M, diamonds, n = 8) and IbTx (100 nM, crosses, n = 15) were plotted and bound with straight lines.
polarization with ryanodine or thapsigargin. Then, we eradicated all the remaining BK channel activity with IbTx. As already shown (Fig. 7 C, d), ryanodine reduced the rate of spike repolarization to 80% (red traces in Fig. 10 B, a and b), and the subsequent application of IbTx further decreased it to 50% (black traces). The sequential depressant actions of ryanodine and IbTx occurred preferentially on the initial phase of spike repolarization so that its monophasic shape changed to diphasic one (Fig. 10 B, b; see also Fig. 7 A), indicating the involvement of the CICR-dependent activation of BK channels in the initial phase of spike repolarization. Combination of the applications of thapsigargin and IbTx showed similar results (not shown). As the rate of voltage change (i.e., the first derivatives) reflected the actual membrane current, these values yielded the fraction of BK channels activated by CICR, which was at least 40% ([100 – 80] × 100/[100 – 50]). It may be noted that the Ca$^{2+}$ current component of spike repolarization, which is in the opposite direction, would have been negligible, as estimated from the difference between the effects of ω-CgTx and IbTx (<3% [52–49%]; no statistical significance).

**Decrease in CICR-dependent BK Channel Activity during a Train of APs.** The progressive decrease in each AP-induced [Ca$^{2+}$], rise in the submembrane region during a 5APs-train (Fig. 3 D, b, and Fig. 7 B, a and b) suggests the waning of BK channel activation by CICR during repetitive APs. We explored this possibility in detail by evoking a train of 20 APs at 50 Hz. During repetition of APs, the spike of AP broadened progressively (Fig. 11 B, a). The rate of spike repolarization decreased and attained a plateau value of 79.0 ± 2.2% at the tenth to twentieth AP (Fig. 11 B, B and C, circles). The peak of the corresponding rise in [Ca$^{2+}$], evoked by each AP in a train also decreased with the progress of stimulation and reached a plateau value of 35.2 ± 9.0% of the first (Figs. 11 A and 12 A). On the other hand, repetition of APs in the presence of ω-CgTx (1 μM) or IbTx (100 nM) decreased the rate of spike repolarization only by 5% (from 50% to 44.8 ± 3.8% or 44.9 ± 6.8%, respectively, at the twentieth spike; Fig. 11 B, c and d, and diamonds and crosses in Fig. 11 C, respectively). These results obviously indicate that the spike broadening during repetitive generation of APs results predominantly from the progressive reduction in BK channel activity and less from that in other K$^{+}$ currents (compare Aldrich et al., 1979; Ma and Koester, 1996). The subtraction of the rates of repolarization in the presence of IbTx from those in the absence of IbTx yielded the time course of changes in the BK channel component of spike repolarization during repetitive APs (open circles in Fig. 12 B). The magnitude of the BK channel component was significantly correlated with that of the corresponding rise in [Ca$^{2+}$], evoked by each AP (Fig. 12 C).

The maximum reduction of BK channel activation ascribed to AP repetition was 32% ([100 – 79 – 5] × 100/50; Fig. 12 B).

Ryanodine (10 μM) or thapsigargin (1 ~ 2 μM) decreased the extent of the reduction in the rate of spike repolarization induced by repetitive APs, which was only 12% (from 79.1% to 68.5 ± 7.4% or from 81.9% to 69.0 ± 4.7%, respectively, at the twentieth spike). Accordingly, the maximum net decreases in the BK channel component under these circumstances were 14% ([12 – 5] × 100/50; open triangles and squares, respectively, in Fig. 12 B). These results strongly indicate that the spike broadening during repetitive APs results from the progressive reduction of CICR in the submembrane region evoked by each AP. The lesser extent of the broadening in the presence of the blockers

**Figure 12.** Correlation of BK channel activity to individual AP-induced rise in [Ca$^{2+}$] during repetition of APs. (A) Changes in each AP-induced increase in [Ca$^{2+}$], in the submembrane region (∆[Ca$^{2+}$]) during a train of 20 APs at 50 Hz. Each change in ∆[Ca$^{2+}$] was normalized to that evoked by the first AP in a train in the absence of blockers. (B) Changes in IbTx-sensitive fraction of the maximum rate of spike repolarization (peak dV/dt BK) in the absence (open circles; n = 14) and the presence of ryanodine (squares; 10 μM, 10 min, n = 7) or thapsigargin (triangles; 1-2 μM, n = 7). The maximum rate of spike repolarization in the presence of IbTx was subtracted from each of those in the absence of IbTx. This procedure yielded the IbTx-sensitive component of spike repolarization in the presence and absence of CICR blockers. All the data were normalized to the maximum fractional decrease of the first AP in the presence of IbTx (50% of the control; see Fig. 11 C). (C) Correlation between ∆[Ca$^{2+}$], shown in A and peak dV/dt (BK) shown in B. Error bars indicate the means ± SEM. A dashed line in C expresses the slope of linear regression (0.57).
Functional Triads of Ion Channels

could be explained by the same type of decrease in residual CICR. These progressive decreases in each of AP-induced Ca\(^{2+}\) transients as well as BK channel activation during repetitive APs could not be due to the inactivation of Ca\(^{2+}\) channels because of their slow inactivation property (Jones and Marks, 1989b). Ryanodine or thapsigargin reduced the amplitude of each AP-induced rise in [Ca\(^{2+}\)] and the plateau level during the course of stimulation [Fig. 11 A]. However, the faithful quantitative measurement of the individual AP-induced rises in [Ca\(^{2+}\)] was not possible because of the high noise level due to the high background fluorescence.) Consequently, the sum of the reduction in the ryanodine- or thapsigargin-sensitive component of the spike repolarization of the first AP (40%) and that brought about by repetition of APs in the presence of CICR blockers (14%) provides an estimate of the total fraction of BK channels activated by CICR for spike repolarization, which was 54%. The remaining fraction of Ca\(^{2+}\)-dependent spike repolarization must therefore depend on another source of Ca\(^{2+}\), i.e., Ca\(^{2+}\) entry.

**The Shift in Predominance from BK to SK Channels for AHP Generation during Repetitive APs.** Shortening of AHP by suppression of CICR (Fig. 7, A–C, c) suggests that CICR also regulates SK channels. Apamin (100 nM), a SK channel blocker (Pennefather et al., 1985; Tanaka et al., 1986), reduced the half-decay time of AHP of an AP to 72.2 ± 9.1%, but not the amplitude (99.8 ± 2.8% of the control, n = 5; Fig. 13 A, a). In contrast, IbTX (100 nM) markedly decreased the amplitude of AHP following a single spike to 76.5 ± 3.5% (n = 7) and the half decay time to 81.8 ± 5.7% (Fig. 13 B, a). This indicates that the initial component of AHP of an AP is produced by the activation of BK channels, whereas the later component is caused by the activation of SK channels. In addition, ω-CgTx suppressed both
the amplitude (50.7 ± 4.8%) and half-decay time of AHP of an AP (48.1 ± 6.1%, n = 8, Fig. 9 A, b), whereas nifedipine did not affect the AHP (peak amplitude; 100.2 ± 1.1%, half-decay time; 91.5 ± 9.3%, n = 8, Fig. 9 B, b). Ca\(^{2+}\) entry via N-type Ca\(^{2+}\) channels and the subsequent CICR thus activate Ca\(^{2+}\)-dependent K\(^+\) channels that contribute to 50% of AHP formation of an AP.

This mode of AHP generation, however, changed with repetition of APs due to the progressive decrease in BK channel activity. IbTx (100 nM), which significantly reduced the amplitude of the first AHP (105.0 ± 3.8%) and reduced the half-decay time only slightly (74.6 ± 9.3%; Fig. 13 B, b). On the other hand, apamin (100 nM), which did not affect the amplitude of the first AHP (Fig. 13 A, a), decreased that of the tenth AP to 89.0 ± 5.1% and markedly the half-decay time to 49.3 ± 4.8% (Fig. 13 A, b). Thus, SK channel activity increased during repetitive APs, compensating for the reduction in BK channel activity in AHP formation. This increase in SK channel activity as well as the decrease in BK channel activity was also recognized in the membrane current underlying the AHP (\(I_{\text{AHP}}\)).

The time course of \(I_{\text{AHP}}\) was best fitted with a double exponential function in most cases. The amplitude of the fast decay component of \(I_{\text{AHP}}\) (fast \(I_{\text{AHP}}\)) following a single AP was decreased after repeating 10 APs at 50 Hz (Fig. 14 A, a), whereas the decay time constant was increased (Fig. 14 B, a). On the other hand, the amplitude of the slow component of \(I_{\text{AHP}}\) (slow \(I_{\text{AHP}}\)) was increased after repeating 10 APs (Fig. 14 C, a). Under this condition, the time constant of the slow \(I_{\text{AHP}}\) was also increased (Fig. 14 D, a).

IbTx (100 nM) reduced the amplitude of the fast \(I_{\text{AHP}}\) following 1 AP and 10 APs (Fig. 13 B and Fig. 14 A, b) and increased their time constants (Fig. 14 B, b). In contrast, apamin (100 nM) had no significant effects on the fast \(I_{\text{AHP}}\) (Fig. 14 A and B, c). Thus, BK channel activity is largely involved in the fast \(I_{\text{AHP}}\). The lesser contribution of BK channel activity to the fast \(I_{\text{AHP}}\) (38% for 1 AP and 33% for 10 APs; see Fig. 14 A, b) than that estimated from the rate of repolarization (51% for 1 AP and 34% for 10 AP; see above) may indicate the faster decay in BK channel activity during spike repolarization than other voltage-gated K\(^+\) currents. (Prolongation of the time constant of fast \(I_{\text{AHP}}\) by repeating repetitive APs or the action of IbTx would reflect the longer time constants of other K\(^+\) channel currents, which became apparent after the reduction of BK channel component.) Ryanodine and thapsigargin decreased the amplitude of the fast \(I_{\text{AHP}}\) (Fig. 14 A, d and e), indicating again the activation of BK channels by CICR.

Apamin applied for 10 min decreased the amplitude of the slow \(I_{\text{AHP}}\) following 1 and 10 APs (Fig. 13 A) to 69 and 50%, respectively (Fig. 14 C, c). In contrast, IbTx did not affect the amplitude of the slow \(I_{\text{AHP}}\) after a single AP (Fig. 14 C, b). Thus, SK channel is involved in the slow \(I_{\text{AHP}}\) and its fraction was increased from 31 to 50% by repetition of APs. (IbTx increased the amplitude of the slow \(I_{\text{AHP}}\) following 1 AP and 10 APs (Fig. 13 B and Fig. 14 A, b) and increased their time constants (Fig. 14 B, b).)

\(I_{\text{AHP}}\) was fitted by the equation, \(I_{\text{AHP}} = I_{\text{fast}} \times \exp(-t/\tau_{\text{fast}}) + I_{\text{slow}} \times \exp(-t/\tau_{\text{slow}})\) [nA]. (A and B) The amplitude and time constants of fast \(I_{\text{AHP}}\) (C and D) The amplitude and time constants of slow \(I_{\text{AHP}}\). Effects of the blockers are shown in percentages of control, reflected in the relative length of bars for the control. Pale and dark gray dashed lines indicate 100% level of control for each component of \(I_{\text{AHP}}\) after 1 AP and 10 APs, respectively. Asterisks (*) indicate that changes are significantly different from the corresponding control values (\(P < 0.01\)). Numerical values are shown only for the mean in or outside each column, whereas the SEM is shown by a horizontal bar.
tude of the slow $I_{\text{AHP}}$ after 10 APs (Fig. 14 C, b) and the time constants of the slow $I_{\text{AHP}}$ after single and 10 APs (Fig. 14 D, b). These increases can be accounted for by an increase in $\text{Ca}^{2+}$ entry due to the broadened spike duration; see the following section.) These results together with those in preceding paragraphs demonstrate that the shift in predominance from BK to SK channels for AHP formation occurs during repetition of APs.

**Changes in the Mode of SK Channel Activation during Repetitive APs.** The foregoing results demonstrated that the component of SK channel activity in AHP formation increased during a train of APs, whereas BK channel activity decreased progressively due to the reduction of CICR. Since the suppression of AHP by ryanodine or thapsigargin (Fig. 7, A–C, c) suggests that SK channels are also activated by CICR, the CICR-dependent component of SK channel activity is expected to decrease during repetitive stimulation. This was the case.

Ryanodine (10 µM) and thapsigargin (1 µM) reduced the amplitude of the slow $I_{\text{AHP}}$ (reflecting SK channel activity) after a single AP by 19% and 18%, respectively, whereas they decreased the amplitude after 10 APs by 11% and 9% (Fig. 14 C, d and e, and Fig. 15, A and B, b and c, respectively). The difference between those of single and 10 APs in each condition is statistically significant ($P < 0.01$). Since the apamin-sensitive SK channel component of the slow $I_{\text{AHP}}$ amplitude were 31% for a single AP and 50% for 10 APs (see above), the CICR-dependent fraction of SK channel activity were calculated to be 61% (19/31) and 22% (11/50), respectively (Fig. 16). Thus, the CICR-dependent SK channel activity decreases with repetition of APs as in the case of BK channels. The increase in the total activity of SK channels during repetitive APs must then be explained by other mechanisms.

Two mechanisms can be deduced for the increase in SK channel activity as follows: First, this increase in SK channel activity in contrast to the decrease in BK channel activity indicates that SK channel is not activated by the $[\text{Ca}^{2+}]$, in the microdomain sensed by BK channel.
Furthermore, the duration of the activation of SK channels, reflected in the time courses of the slow \( I_{\text{AHP}} \) and AHP, were comparable (albeit shorter; see below) to that of the accompanying submembrane rise in \( [\text{Ca}^{2+}]_i \), and both were concomitantly shortened by CICR blockers (Fig. 7 B, c, and Fig. 15, A and B, b). The \( \text{Ca}^{2+} \)-sensing molecule, calmodulin, which couples with SK channels (Xia et al., 1998), thus senses the mixed, accumulated rise of \( [\text{Ca}^{2+}]_i \) in the submembrane region during a train of APs (Fig. 16; see discussion). Since the accumulated rise of \( [\text{Ca}^{2+}]_i \) is less affected by changes in individual \( \text{Ca}^{2+} \) entry or release during their repetitive activation than the rise in the \( \text{Ca}^{2+} \) microdomains, the total activity of SK channels should be less sensitive to changes in CICR and be increased by the cumulative effect of total \( \text{Ca}^{2+} \) entry and release. Second, total \( \text{Ca}^{2+} \) entry induced by repetitive APs must be increased by the spike broadening, as supported by the enhancement of AP-induced \( \text{Ca}^{2+} \) transients by \( \text{IbTx} \) (Fig. 10 A, c) and the slowly inactivating property of N-type \( \text{Ca}^{2+} \) channel (Jones and Marks, 1989b). This increase in \( \text{Ca}^{2+} \) entry should maintain the averaged rise in \( [\text{Ca}^{2+}]_i \), and enhance SK channel activation, thus compensating for the decrease in CICR during the repetitive APs.

Finally, another evidence suggests that SK channels are likely to respond to the higher \( [\text{Ca}^{2+}]_i \), accumulated in the submembrane region. The time constant of the slow \( I_{\text{AHP}} \) of 10 APs (0.46 s; Figs. 14 D, a, 15) was shorter than that of the slow decay component of 10 APs-induced \( \text{Ca}^{2+} \) transients in the submembrane cytoplasm (1.66 \( \pm \) 0.13 s, \( n = 35 \)). It was still smaller than those of the \( \text{Ca}^{2+} \) transients in the presence of ryanodine (0.76 \( \pm \) 0.09 s, \( n = 14 \); Fig. 15 A, b) or thapsigargin (1.10 \( \pm \) 0.13 s, \( n = 21 \); Fig. 15 B, b). Thus, the activation of SK channels disappears before the decay of an increased \( [\text{Ca}^{2+}]_i \), in the submembrane region. This indicates that the \( \text{Ca}^{2+} \) sensor of SK channels have relatively low \( \text{Ca}^{2+} \) affinity and sense the higher \( [\text{Ca}^{2+}]_i \), in the submembrane space (see discussion).

**DISCUSSION**

**Activation of CICR by an Action Potential in the Submembrane Cytoplasm**

The present study demonstrates that \( \text{Ca}^{2+} \) influx accompanying an AP activates CICR in the submembrane region of cultured bullfrog sympathetic ganglion cells. AP-induced \( \text{Ca}^{2+} \) transients were regeneratively propagated at constant amplitude and rate of rise in the submembrane region (a few micrometers), whose width was almost identical to that of the distribution of ryanodine receptors revealed by fluorescent ryanodine. The propagation was blocked by ryanodine, thapsigargin, or \( \text{o-CgTx} \). The \( \text{Ca}^{2+} \) transient initiated in the submembrane region spread toward the deeper cytoplasm with decreasing in amplitude and the rate of rise, reflecting the passive buffered diffusion of \( \text{Ca}^{2+} \). Furthermore, computer simulation assuming \( \text{Ca}^{2+} \) release in the submembrane region with \( \text{Ca}^{2+} \) entry at the cell membrane nicely reconstructed the spatial and temporal profiles of AP-induced \( \text{Ca}^{2+} \) dynamics.
Previous attempts (Nohmi et al., 1992) to prove the involvement of CICR in the AP-induced Ca$^{2+}$ transients in intact bullfrog sympathetic ganglion cells were not successful. In that study, Ca$^{2+}$ transients induced by repetitive APs were little affected by ryanodine (10 μM), although caffeine-induced [Ca$^{2+}$]i oscillation was completely blocked under the same condition. There are three technical reasons for the failure to reveal the activation of CICR by APs in the previous study. First, in contrast to the high time resolution (500 Hz) in the present study, fluorescence was measured at 0.5 Hz for the experiments to observe the effects of ryanodine in the previous study (see Methods in Nohmi et al., 1992). Second, averaged fluorescence changes were measured through an objective of low NA (0.5 or 0.75) predominantly from the core cytoplasm (more precisely from the square inscribed to the center of a cell image; Nohmi et al., 1992). Under those conditions, the fast regenerative phase of AP-induced Ca$^{2+}$ transient in the submembrane region would have not been fully resolved and partly blunted, although the extent of an increase in the measured [Ca$^{2+}$]i would have been enhanced by CICR. The situation is quite a contrast to that in the present study, in which confocal imaging with a high NA (1.15) objective was used. Third, in contrast to a low concentration (10 μM) of OGB-1 in the present study, a high concentration (~50 μM) of Fura-2 was used in the previous study (Nohmi et al., 1992). This large amount of exogenous Ca$^{2+}$ buffer would have retarded the regenerative activation of CICR. Studies using 50 μM of Indo-1 or Fura-2 (and also 1 mM EGTA in patch pipettes) showed that a long depolarizing pulse (>200 ms), high K+-induced depolarization or the action of caffeine activated the submembrane CICR (Hua et al., 1993, 2000). Rather long lasting Ca$^{2+}$ entry seemed to be needed to overcome the exogenous buffering effects of Indo-1 and EGTA. Ca$^{2+}$ entry during the later part of long depolarization (during which exogenous buffers would have been saturated) therefore activated CICR. In support of this, CICR that was activated by a voltage pulse increased with an increase in the total amount of Ca$^{2+}$ entry in the previous studies. In addition, depolarizing pulse (100 ms)-induced Ca$^{2+}$ transients were facilitated by repeating pulses (Hua et al., 1993). The activation of CICR by caffeine appears to be less affected by exogenous Ca$^{2+}$ buffers, since the activation occurs as a result of an increase in the Ca$^{2+}$ sensitivity of ryanodine receptors (compare to Endo, 1975).

The activation of CICR by Ca$^{2+}$ entry accompanying a single AP was also found in other neurons. Ca$^{2+}$ transients through the activation of CICR by a single AP were reported in rabbit vagal afferent neurons (Cohen et al., 1997) and rat CA1 pyramidal neurons (Sandler and Barbara, 1999). The ryanodine- or thapsigargin-sensitive component of AHP following a single AP was seen in rat sympathetic ganglion cells (Kawai and Watanabe, 1989), guine pig vagal motoneurons (Sah and McLachlan, 1991), rabbit otic ganglion cells (Yoshizaki et al., 1995), rabbit vagal afferent neurons (Moore et al., 1998), and rat CA3 pyramidal neurons (Tanabe et al., 1998). Thus, the functional coupling of voltage-gated Ca$^{2+}$ channels to ryanodine receptors appears to be the common mechanism in a variety of neurons. The strength of such a coupling, however, may vary among these neurons. The coupling of Ca$^{2+}$ channels and ryanodine receptors in bullfrog sympathetic neurons seems to be as strong as the coupling in cardiac muscles (see below). However, N-type Ca$^{2+}$ channels, instead of L-type, are involved in the coupling.

The Functional Triad Consisting of Ryanodine Receptors, N-type Ca$^{2+}$ Channels and BK Channels

The present study showed that blocking CICR by ryanodine or thapsigargin reduced the rate of spike repolarization of an AP. The extent of the blockade by either of the CICR blockers was 40% of that brought about by ω-CgTx or IbTx. CICR also decreased during repetition of APs, further decreasing the rate of the spike repolarization by 12–14% under the effect of CICR blockers. If we assume that the remaining CICR in the presence of CICR blockers disappeared during the repetitive APs, at least 54% of the population of BK channels is activated by CICR in response to Ca$^{2+}$ entry through N-type Ca$^{2+}$ channels. The rest of population (46%) was directly activated by the Ca$^{2+}$ entry (Fig. 16).

There are two limiting conditions required for the effective couplings of ryanodine receptors to Ca$^{2+}$ channels and BK channels. First, the time delay for the sequential activation of ryanodine receptors and BK channels after the entry of Ca$^{2+}$ through N-type Ca$^{2+}$ channels must be less than the time period from the peak of spike to the maximum rate of the spike repolarization. This latency, which was 0.64 (0.53 ~ 0.82) ms in this study, would provide the upper bound for the sum of the distance for Ca$^{2+}$ diffusion between N-type Ca$^{2+}$ channels and ryanodine receptors and that between ryanodine receptors and BK channels. The diffusion time of 0.64 ms for Ca$^{2+}$ would give the diffusion path of 196 nm [= (6Dt)$^{1/2}$ = (6 × 10$^{-7}$ cm$^2$/s × 0.64 × 10$^{-3}$ s)$^{1/2}$], yielding 98 nm of the distance from ryanodine receptor to Ca$^{2+}$ channels or BK channels (assuming the equivalent distances for the two diffusion paths). The time delay for the activation of BK channels by CICR would be even shorter, because the effect of CICR blockers on APs had already appeared around the peak of the spike (Fig. 7). The distances between ryanodine receptor to Ca$^{2+}$ channels and BK channels could therefore be much shorter than this value.
The second limiting condition for the effective coupling may be the Ca\textsuperscript{2+} affinity of BK channels. The Ca\textsuperscript{2+} sensitivity of BK channel was reported to be \(>10\ \mu\text{M} \) at +20 \(\sim\) –40 mV (Barrett et al., 1982; Smart, 1987; McManus and Magleby, 1991). This indicates that BK channels are activated by the high [Ca\textsuperscript{2+}], in Ca\textsuperscript{2+} microdomains ([Ca\textsuperscript{2+}]_{\text{M}} \approx [Ca\textsuperscript{2+}]_{\text{D}} in Fig. 16) produced by both Ca\textsuperscript{2+} entry through Ca\textsuperscript{2+} channels and Ca\textsuperscript{2+} release via ryanodine receptors, rather than the [Ca\textsuperscript{2+}], in the bulk phase of the cytoplasm measured. Such a low Ca\textsuperscript{2+} sensitivity of BK channels obviously requires the close proximity of BK channels to ryanodine receptors. Assuming rapid Ca\textsuperscript{2+} buffering, numerical analysis of the [Ca\textsuperscript{2+}], in the Ca\textsuperscript{2+} domain predicted that the range of an increased [Ca\textsuperscript{2+}], \(>10\ \mu\text{M} \) within 1 ms after instantaneous Ca\textsuperscript{2+} entry at a point was \(\sim100\ \text{nm} \) from the Ca\textsuperscript{2+} source (see Fig. 1 of Smith, 1996; Smith et al., 1996). This value is similar to that estimated from the temporal limitation (see above). Thus, ryanodine receptors in bullfrog sympathetic neurons must be closely coupled to both N-type Ca\textsuperscript{2+} channels and BK channels within this distance (Fig. 16).

The findings in bullfrog sympathetic neurons are consistent with the earlier observations of spontaneous miniature transient outward currents (SMOCs) due to the activation of BK channels by presumptive spontaneous Ca\textsuperscript{2+} release (Brown et al., 1983; Satin and Adams, 1987). Similar transient outward currents in smooth muscle cells were nicely correlated with the occurrence of localized spontaneous Ca\textsuperscript{2+} transients, Ca\textsuperscript{2+} sparks (Pérez et al., 1999; Zhu Ge et al., 1999). Moreover, SMOCs in mudpuppy parasympathetic neurons were blocked by Ca\textsuperscript{2+} channel antagonists, nifedipine and ω-CgTx, as well as ryanodine (Merriam et al., 1999). This indicates the existence of a similar coupling of ryanodine receptors to Ca\textsuperscript{2+} channels and BK channels in the parasympathetic neurons, although the mode of CICR activation by Ca\textsuperscript{2+} influx was not selective as in bullfrog sympathetic neurons.

Functional Coupling of Ryanodine Receptors to N-type Ca\textsuperscript{2+} Channels at the Plasma Membrane

CICR triggered by Ca\textsuperscript{2+} entry accompanying the spike of AP activates another type of Ca\textsuperscript{2+}-dependent K\textsuperscript{+} channels, SK channels, and participates in the generation of AHP. This was indicated in the reduction of the apamin-sensitive components of AHP and the slow \(I_{\text{AHP}}\) by thapsigargin or ryanodine. This is consistent with the previous findings in the same (Kuba et al., 1983) or other (Kawai and Watanabe, 1989; Sah and McLachlan, 1991; Yoshizaki et al., 1995; Moore et al., 1997; Tanabe et al., 1998) types of neurons.

As already described in the results, the increase in the total SK channel activity during repetitive APs and its slow decay (reflected in AHP and slow \(I_{\text{AHP}}\)), which are comparable with the time course of repetitive APs-induced Ca\textsuperscript{2+} transient, suggest that SK channels would sense the mixed, accumulated rise of [Ca\textsuperscript{2+}], during the train of APs. This mode of activation of SK channel can be achieved only by the location of SK channels somehow remote from both Ca\textsuperscript{2+} channels and ryanodine receptors. This is consistent with the previous observations. The intracellular injection of EGTA shortened the duration of AHP with no change in the peak amplitude (Kuba et al., 1983; Tanaka and Kuba, 1987), indicating the distance of more than 100 nm (Naraghi and Neher, 1997) between the site of CICR and SK channels (Fig. 16). However, the activity of SK channels should reflect a relatively high level of [Ca\textsuperscript{2+}], in the submembrane region, since the decay time course of \(I_{\text{AHP}}\) was shorter than that of the submembrane [Ca\textsuperscript{2+}], rise (Fig. 15). The affinity of the Ca\textsuperscript{2+} sensing molecule forming complex with SK channels was reported to be 0.3 \(\mu\text{M} \) (Xia et al., 1998), which is greater than the peak of AP-induced Ca\textsuperscript{2+} transients measured in this study. In this context, SK channels could be regarded to couple functionally with N-type Ca\textsuperscript{2+} channels and ryanodine receptors, but less tightly than BK channels.

Plastic Modulation of Cell Membrane Excitability by the Functional Coupling

The activation of BK and SK channels by CICR in response to Ca\textsuperscript{2+} entry through N-type Ca\textsuperscript{2+} channels and its dependence on the frequency of APs may flexibly regulate the cell membrane excitability under the physiological condition. For sparsely coming presynaptic inputs, the fast spike repolarization of an AP through the activation of BK channels by CICR would prepare for the brisk subsequent firing of the neuron. This effect of CICR, activating 52 \(\sim\) 54\% of the population of BK channel, however, is not maintained for repeated high-frequency synaptic inputs. The magnitude of CICR decreases during the course of repetitive APs presumably due to inactivation of CICR. This reduces the activation of BK channel by CICR to a level, for instances, less than 20 \(\sim\) 23\% at 50 Hz, broadening the spike of AP (Fig. 11; see also Fig. 16). Under this condition, the broadened spike would prolong the refractory period of Na\textsuperscript{+} channels at the axon hillock, limiting the rate of impulse transmission to target cells.

On the other hand, the broadened spike increases the duration of N-type Ca\textsuperscript{2+} channel activation. This increases the amount of Ca\textsuperscript{2+} entry, thereby compensates for the decrease in CICR and maintains the averaged total rise in [Ca\textsuperscript{2+}], around SK channels for their activation. Thus, the AHP following repetitive APs is prolonged in spite of the reduced CICR. The long-lasting AHP produced mainly by the SK channels activation increases the threshold for the subsequent firing. These modes of activation of BK and SK channels by CICR...
during repetitive nerve activity may favor intermittent bursting discharges. Consequently, this bullfrog sympathetic neuron regulates its membrane excitability by shifting the dependence of spike and AHP formation over two different kinds of Ca\(^{2+}\)-activated K\(^+\) channels, through changing the extent of CICR. The efficacy of this mechanism would be variable among the cells, depending on the density of ryanodine receptors or the amount of Ca\(^{2+}\) stores.

Functional coupling between two types of ion channels has been known for those between ryanodine receptors and L-type Ca\(^{2+}\) channels or Ca\(^{2+}\)-dependent K\(^+\) channels (Pérez et al., 1999; Zhu Ge et al., 1999) in muscles. There are also several lines of evidence for more or less similar functional couplings in neurons. Selective activation of SK channels by L-type Ca\(^{2+}\) channels and that of BK channels by N-type Ca\(^{2+}\) channels were reported in hippocampal neurons (Marion and Tavalin, 1998) and their opposite roles in rat superior cervical ganglion (Davies et al., 1996). The regulation of L-type Ca\(^{2+}\) channels at the cell membrane by ryanodine receptors was reported in mouse cerebellar granule cells (Chavis et al., 1996). There is no evidence, however, for the functional coupling of three types of ion channels. The coupling of ryanodine receptor to N-type Ca\(^{2+}\) channel and Ca\(^{2+}\)-activated K\(^+\) channel in bullfrog sympathetic neurons would be the first evidence for the functional coupling of ion channels, which operates in a cascade of Ca\(^{2+}\) entry, Ca\(^{2+}\)-induced Ca\(^{2+}\) release and Ca\(^{2+}\)-activated opening of K\(^+\) channel within a short period of time (<1 ms). Moreover, the graded reduction of the submembrane CICR during repetitive APs provides plasticity in the strength of this coupling, which seems to be also the first evidence for the plastic modulation of cell functions by CICR. Furthermore, the modulation of Ca\(^{2+}\) channels by adrenaline, acetylcholine, and other neurotransmitters are known in bullfrog sympathetic neurons (Kuba and Koketsu, 1976; Minota and Koketsu, 1977; Bley and Tsien, 1990; Elmslie et al., 1990). Such modulation should affect the extent of the submembrane CICR due to the close coupling. Thus, the functional triad formed by ryanodine receptors, Ca\(^{2+}\) channels, and Ca\(^{2+}\)-activated K\(^+\) channels operates as one of the important mechanisms of postsynaptic plasticity in bullfrog sympathetic neurons and may provide a novel mechanism in general for the plastic regulation of cell membrane excitability in various kinds of cells.

This work was supported by Grants in Aid for Scientific Research from the Japanese Ministry of Education, Science and Culture to K. Kuba.

Submitted: 21 June 2000
Revised: 26 September 2000
Accepted: 28 September 2000

References


Akita, T., and K. Kuba. 1999a. Ca\(^{2+}\) release from submembrane and perinuclear Ca\(^{2+}\) stores are involved in action potential-induced Ca\(^{2+}\) transients and modulation of membrane excitability in cultured bullfrog sympathetic neurons. Jpn. J. Physiol. 49(Suppl.): S120. (Abstr.).


