Review

Anesthetics and Cell–Cell Communication: Potential Ca\(^{2+}\)-Calmodulin Role in Gap Junction Channel Gating by Heptanol, Halothane and Isoflurane

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Abstract: Cell–cell communication via gap junction channels is known to be inhibited by the anesthetics heptanol, halothane and isoflurane; however, despite numerous studies, the mechanism of gap junction channel gating by anesthetics is still poorly understood. In the early nineties, we reported that gating by anesthetics is strongly potentiated by caffeine and theophylline and inhibited by 4-Aminopyridine. Neither Ca\(^{2+}\) channel blockers nor 3-isobutyl-1-methylxanthine (IBMX), forskolin, CPT-cAMP, 8Br-cGMP, adenosine, phorbol ester or H7 had significant effects on gating by anesthetics. In our publication, we concluded that neither cytosolic Ca\(^{2+}\) nor pH\(_i\) were involved, and suggested a direct effect of anesthetics on gap junction channel proteins. However, while a direct effect cannot be excluded, based on the potentiating effect of caffeine and theophylline added to anesthetics and data published over the past three decades, we are now reconsidering our earlier interpretation and propose an alternative hypothesis that uncoupling by heptanol, halothane and isoflurane may actually result from a rise in cytosolic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)\(_i\)]\(_i\)) and consequential activation of calmodulin linked to gap junction proteins.

Keywords: anesthetics; heptanol; halothane; isoflurane; gap junctions; connexins; channel gating; calcium; calmodulin; cell communication; cell-to-cell channels; cell coupling; cell uncoupling; membrane channels

1. Introduction

Direct cell–cell communication is mediated by gap junction channels that enable free exchange of small cytosolic molecules among neighboring cells. Each channel is formed by two hemichannels that create a hydrophilic passageway, spanning the plasma membranes of two neighboring cells and a narrow extracellular space (gap). A gap junction hemichannel is made of six radially arranged proteins named connexins in vertebrates and innexins in invertebrates. Connexins/innexins contain two extracellular lops (EL\(_1\) and EL\(_2\)) and three cytoplasmic domains: an NH\(_2\)-terminus (NT), a cytoplasmic loop (CL) and a COOH-terminus domain (CT; rev, in [1]).

Gap junction channels are physiologically regulated by a chemical gating mechanism that is activated by changes in cytosolic ionic homeostasis, resulting from cell damage, inhibition of the metabolism, acidification and hypoxia, among others. Over the years, numerous studies have suggested that channel gating results from a rise in cytosolic calcium concentration ([Ca\(^{2+}\)\(_i\)]\(_i\)) [2–4] in the high nanomolar–low micromolar range [1,5–7]. Since the early 1980s, we have proposed that Ca\(^{2+}\)\(_i\) causes gating by activating calmodulin (CaM) [1,7–10] via a cork-like pore-plugging mechanism [11–13] probably involving conformational changes in connexins as well.

In the last four decades, many studies have reported that gap junction channels are sensitive to anesthetics. Channel gating induced by long-chain alcohols (heptanol and octanol) was first reported by Johnston and coworkers [14] in crayfish septate axons. Soon
after, gating by volatile anesthetics such as halothane and isoflurane was also reported in a variety of vertebrate and invertebrate systems [15,16].

In spite of over four decades of research, the mechanism by which anesthetics cause gap junction channel gating is still poorly understood. Johnston and coworkers [14] proposed an extracellular site of action for alkanols. Similarly, Eskandari and coworkers [17] reported that, in inside-out patches of lens connexin hemichannels, the addition of 1 mM octanol did not affect the channel’s open probability or the unitary conductance, while in outside-out patches, addition of 1 mM octanol to the bath (extracellular surface of hemichannels) significantly reduced single-channel open probability without altering the unitary current. Therefore, they concluded that octanol inhibits lens connexin hemichannels by acting on a site accessible only from the extracellular space. However, the possibility that a soluble intermediate was washed away by the internal perfusion of crayfish axons [14] and by the bathing solution of inside-out patches [17] was not considered.

In 1991, our data seemed to also exclude the role of cytosolic Ca$^{2+}$ and pH in crayfish axons uncoupled by heptanol, halothane and isoflurane [16]. However, based on the potentiating effect of caffeine and theophylline added to anesthetics, and the numerous studies published in the past three decades on the effect of anesthetics on both CaM's Ca$^{2+}$ sensitivity and Ca$^{2+}$ release from the sarcoplasmic reticulum (SR) via inositol trisphosphate (IP3) and/or ryanodine receptor (RyR) channels, as well as evidence that ion-selective electrodes may be unreliable in the presence of anesthetics [18], we are now reconsidering our earlier interpretation [16]. As a result, we propose an alternative hypothesis, that cell–cell uncoupling by anesthetics may result from CaM's activation caused by anesthetic-induced [Ca$^{2+}$]i rise, and increased Ca$^{2+}$ sensitivity in CaM.

2. Measurement of Junctional Resistance (Rj), [Ca$^{2+}$]i and [H$^{+}$]i in Crayfish Septate Axons

In our 1990s study [16], crayfish axons were superfused with a standard saline solution for crayfish (SES) [19], containing (in mM): NaCl, 205; KCl, 5.4; CaCl$_2$, 13.5 and 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), 5 (pH 7.5). Either 2.8–5.6 mM 1-heptanol, 9.5–28.5 mM halothane or 23.6 mM isoflurane was added to SES in the presence and absence of either 10–20 mM caffeine or 10-20 mM theophylline. For testing the effect of low pH$_i$ [20,21], the axons were superfused with a sodium acetate saline solution (Ac) containing (in mM): Na acetate, 205, KCl, 5.4 and CaCl$_2$, 13.5 (pH 6.3).

Four microelectrodes were inserted into a lateral giant axon, two on each side of the septum (Figure 1A), and hyperpolarizing square current pulses (150 nA, 300 ms) were passed every 10 s alternatively into the posterior (C$_1$) and anterior (C$_2$) axon segments. The resulting electrotonic potentials $V_1$ and $V_2$ (from current injection in C$_1$), $V_1^*$ and $V_2^*$ (from injection in C$_2$) and the membrane potentials ($E_1$ and $E_2$) were recorded with two voltage microelectrodes through a voltage follower. The voltage signals were displayed on an oscilloscope and a chart recorder, and were digitized. Both membrane ($R_{m1}$, $R_{m2}$) and junctional ($R_{j1}$, $R_{j2}$) resistances were calculated from current ($I_1$, $I_2$) and voltage ($V_1$, $V_2$, $V_1^*$, $V_2^*$) records (Figure 1B) [16].

[Ca$^{2+}$]$i$ and [H$^{+}$]$i$ were measured with ion-sensitive microelectrodes based on neutral-carrier sensors. Ca$^{2+}$ microelectrodes used the calcium cocktail (ETH 129) and H$^{+}$ microelectrodes used the proton cocktail tri-n-dodecylamine. At the time, we thought that these ion-selective microelectrodes were insensitive to anesthetics. Later on, however, ion-selective microelectrodes were proven unreliable in the presence of anesthetics [18]. In contrast, these ion-selective microelectrodes are reliable for measuring changes in [Ca$^{2+}$]$i$, or [H$^{+}$]$i$, caused by acetate-induced cytosolic acidification [20,21].
Figure 1. Diagram of electrical recording setup (A). Current pulses are injected via current microelectrodes (I1, I2) alternatively in the posterior (C1) and anterior (C2) axon segments. The resulting potentials (V1, V2, V1*, V2*) are displayed in the chart recorder and the oscilloscope. The voltage signal (Vr) recorded by the ion-selective microelectrode, is displayed in the chart recorder after subtraction of membrane potential (Vm). (B) Equivalent circuit and equations used to calculate junctional (Rj1, Rj2) and non-junctional (Rm1, Rm2) resistances. Adapted from [21].

3. Gating by Heptanol in the Presence and Absence of Caffeine or Theophylline

Lateral giant axons have a membrane potential that ranges from −80 to −95 mV (Figures 2A and 3A) and are electrically coupled at the septum with an initial Rj of 150 ± 53.7 kΩ (mean ± SD; n = 28). Superfusion of the axons with 2.8–5.6 mM heptanol results in a small depolarization (Figures 2A and 3A) and increases Rj by 191.3 ± 83% (mean ± SD; n = 28) of control values (Figures 2B and 3B). Note that the peaks of depolarization and Rj correspond well (Figure 2A,B). Recovery of both Rj and membrane potential (MP) are quicker than their onset (Figures 2A,B and 3B).

Addition of 10–20 mM theophylline to heptanol solutions increases Rj maxima by 309.3 ± 265% (mean ± SD; n = 24) of controls treated with heptanol alone (Figures 2B and 3B) and causes greater depolarization (Figures 2A and 3A). Significantly, when caffeine is added to heptanol several minutes after the beginning of heptanol treatment, the rates of depolarization (Figure 3A) and Rj rise (Figure 3B) greatly increase. Indeed, a 7 min superfusion of heptanol–caffeine, following a 22 min superfusion of heptanol alone, virtually doubles the Rj rise induced by heptanol alone (Figure 3B). Following the first heptanol–caffeine application the Rj maxima with heptanol alone or heptanol–caffeine significantly decreased (Figure 4, blue and red arrows, respectively). This may suggest that the first heptanol–caffeine treatment reduced the calcium content of the endoplasmic reticulum (ER). Addition of 10 µM ryanodine partially reduces the Rj maxima reached with heptanol–caffeine (Figure 5), suggesting a partial inhibition of calcium release from the ER stores.

Addition of 10–20 mM theophylline to heptanol solutions also dramatically enhanced the heptanol effects on Rj (Figure 6). Indeed, Rj maxima with heptanol–theophylline are 676 ± 386% (mean ± SD; n = 4) greater than those with heptanol alone. Neither different external [Ca²⁺]o, ranging from 7 to 27 mM, nor blockers of Ca²⁺ entry, such as Cd²⁺ (500 µM) or nisoldipine (10 µM), significantly change the heptanol-uncoupling efficiency [16].
Addition of caffeine to heptanol causes a rapid change in the amplitude of the electrotonic potentials (A), indicative of rapid increase in $R_j$ (B). $R_j$ increases with heptanol-caffeine 2–3 times as much as with heptanol alone (B, red arrow). The second heptanol treatment shows that the maximal rate of $R_j$ rise with heptanol alone is $14 \, \text{k}\Omega/\text{min}$ (B, green arrow), while with heptanol-caffeine (first treatment) the rate more than doubles ($35 \, \text{k}\Omega/\text{min}$) (B, red and green arrows). Note that the membrane depolarization caused by heptanol (A, left green arrow) is increased by caffeine addition (A, red and green arrows). The maximum depolarization (A) corresponds to the maximum $R_j$ increase (B). Adapted from [16].
heptanol-induced uncoupling [16]. The possible involvement of protein kinase C (PKC) was tested by superfusing the axons with heptanol solutions containing either 162 nM TPA (4β-phorbol-12β-myristate-13α-acetate) or 100 μM H7 (1-(5-isoquinoliny sulfonyl)-2-methylpiperazine); neither TPA (an activator of PKC) nor H7 (an inhibitor of PKC) significantly affected the magnitude of heptanol-induced uncoupling [16].

The possibility that the effect of caffeine or theophylline is due to an increase in cyclic nucleotides was tested by exposing the axons to 3-isobutyl-1-methylxanthine (IBMX, a phosphodiesterase inhibitor), forskolin (an activator of adenylate cyclase) or diffusible cAMP and cGMP (CPT-cAMP and 8Br-cGMP) [16]. Additions to heptanol of 1 mM IBMX (Figure 7), a phosphodiesterase inhibitor 200 times more potent than caffeine [22], 5 μM forskolin, 500 μM CPT-cAMP or 200 μM 8Br-cGMP do not significantly affect Rj maxima [16]. The possible involvement of protein kinase C (PKC) was tested by superfusing the axons with heptanol solutions containing either 162 nM TPA (4β-phorbol-12β-myristate-13α-acetate) or 100 μM H7 (1-(5-isoquinoliny sulfonyl)-2-methylpiperazine); neither TPA (an activator of PKC) nor H7 (an inhibitor of PKC) significantly affected the magnitude of heptanol-induced uncoupling [16].
Figure 6. Time course of Rj in crayfish axons uncoupled by heptanol in the presence and absence of 20 mM theophylline. As with caffeine (Figures 2–4), theophylline dramatically increases Rj maxima with heptanol. Adapted from [16].

Figure 7. Time course of Rj in crayfish axons uncoupled by heptanol in the presence and absence of 1 mM IBMX. IBMX (an inhibitor of phosphodiesterases and adenosine receptors) does not affect heptanol-induced Rj rise. Adapted from [16].

In view of the fact that caffeine is also a powerful inhibitor of adenosine receptors, the effect of adenosine on Rj was tested both in the presence and absence of heptanol. Superfusion of 1.3–5 mM adenosine, added to either SES or heptanol solutions, does not significantly change either control Rj values or Rj maxima with heptanol, indicating that adenosine receptors do not participate in the effect of heptanol and caffeine on gap junction channel conductance [16]. Lack of an involvement of adenosine receptor inhibition is also provided by the absence of an effect of IBMX on heptanol-induced Rj rise (Figure 7) [16]; indeed, IBMX, as caffeine and theophylline, is also an inhibitor of adenosine receptors [23,24].

Curiously, the K+ channel blocker 4-aminopyridine (4-AP) strongly inhibits the heptanol-induced uncoupling. With heptanol solutions containing 5 mM 4-AP, the Rj maxima are 26.2 ± 20% (mean ± SD; n = 12) lower than those with heptanol alone (Figure 8A). In contrast, addition of 4-AP (5 mM) to acetate solutions does not alter their uncoupling effects (Figure 8B). No effect on Rj was seen with 4-AP alone, with the only change being a 3–4 mV depolarization [16], probably caused by the inhibition of K+ channels.
with heptanol (\(\text{mean} \pm \text{SD}; n = 8\)) greater than those with halothane alone [16].

\[ R_j = 329 \pm 147\% \text{ of control values} \]

\[ V_1^* \text{ and } V_2^* \text{ are greater than with halothane alone (Figure 8).} \]

**Figure 8.** Time course of Rj in crayfish axons uncoupled by heptanol (A) or acetate (B) in the presence or absence of 5 mM 4-aminopyridine (4-AP). 4-AP dramatically reduce the Rj maximum with heptanol (A). In contrast, addition of 4-AP to acetate solutions (Ac) does not affect the acetate-uncoupling efficiency (B). Adapted from [16].

**4. Gating by Halothane or Isoflurane in the Presence and Absence of Caffeine**

Superfusion of crayfish axons with halothane (28.5 mM) causes a small depolarization (Figure 9A) and increases Rj by 155.6 \(\pm\) 56\% \(\text{mean} \pm \text{SD}; n = 9\) of control values of 226 \(\pm\) 73 k\(\Omega\) \(\text{mean} \pm \text{SD}; n = 9\) \(\text{Figures 9B and 10}\) [16]. Addition of 20 mM caffeine to halothane solutions causes a more marked depolarization (Figure 9A) and a greater increase in Rj (Figures 9B and 10). The Rj maxima with halothane–caffeine are 329 \(\pm\) 147% \(\text{mean} \pm \text{SD}; n = 8\) greater than those with halothane alone [16].

**Figure 9.** Time course of electrotonic potentials (A) and Rj (B) in crayfish axons uncoupled by halothane in the presence and absence of 20 mM caffeine. With halothane-caffeine, the increase in \(V_1^*\) and \(V_2^*\), and the decrease in \(V_1^*\) and \(V_2\) are greater than with halothane alone ((A), red and green arrows), due to a grater increase in Rj (B). Note that with caffeine, there is a greater depolarization ((A), green and red arrows), whose peak corresponds to the peak or Rj maximum. Following the last halothane-caffeine treatment, coupling recovered only partially (A,B). Adapted from [16].
Similar results were obtained with isoflurane (23.6 mM), which causes a 4–5 mV depolarization (data not shown) and increases Rj by ~125% of control values (Figure 10) [16]. With addition of 20 mM caffeine to isoflurane solutions, Rj maxima are ~170% greater than those with isoflurane alone (Figure 10).

![Figure 10. Time course of Rj in crayfish axons treated with either isoflurane or halothane in the presence or absence of 20 mM caffeine. As for heptanol and halothane, caffeine has a pronounced effect on the uncoupling efficiency of isoflurane. However, isoflurane has a significantly weaker effect on Rj than halothane. Adapted from [16].](image)

5. Potential Mechanism of Channel Gating by Anesthetics

In our 1991 study, we concluded that the uncoupling mechanism of the anesthetics heptanol, halothane and isoflurane involves a direct interaction between anesthetics and amphiphilic chains of gap junction proteins. Our interpretation was based primarily on the apparent drop in [Ca$^{2+}$], and [H$^+$]]. However, as mentioned in the previous section, after our publication, ion-selective electrodes have been reported not to be very reliable in the presence of anesthetics [18], such that a role of Ca$^{2+}$ could not be excluded. Indeed, we are now reconsidering our earlier interpretation and propose a hypothesis suggesting the potential role of Ca$^{2+}$-activated CaM.

5.1. Does Uncoupling by Anesthetics Result from Increased [Ca$^{2+}$], Potentiated by Caffeine and Theophylline?

Meda and coworkers [25] reported no significant change in [Ca$^{2+}$], measured with Quin-2, in exocrine pancreas treated with heptanol or octanol at concentrations that greatly increased amylase release. However, this is curious because amylase release is known to be caused by a sustained [Ca$^{2+}$], rise [26–28]; therefore, one may wonder whether the Ca$^{2+}$ sensitivity of Quin-2 might have been affected by alkanols. In agreement with this report [25], some studies have reported the normal uncoupling efficiency of alkanols and halothane with patch pipette solutions buffered for Ca$^{2+}$ [29–32]. However, these data conflict with several other studies which reported an alkanol-induced [Ca$^{2+}$], rise. In a careful double whole-cell patch-clamp study on embryonic chick cardiac cells, Veenstra and DeHaan [33] reported that octanol uncoupled the cells by reducing junctional conductance (Gj) to 4.3 ± 3.4% of the control level with patch pipette solutions weakly buffered for Ca$^{2+}$ (0.1 mM EGTA), but octanol reduced Gj by only 29.0 ± 19.1% with more strongly Ca$^{2+}$-buffered solutions (5 mM EGTA), in their words [33]: “A possible mechanism for the octanol effect comes from studies by Vassort et al. ([34]) indicating that long-chain alcohols inhibit cytoplasmic Ca-buffering. These workers reported that octanol caused a rise in [Ca]c (cytosolic
Ca$^{2+}$) in squid axons, associated with an alkalinization of the axoplasm, and they presented evidence that Ca$^{2+}$ is released from an intracellular binding site in exchange for H$^+$ ions ... these observations are consistent with our finding that octanol uncoupled cell pairs when calcium buffering was minimal, and that the effect was reduced when the cells were infused with Lo-Ca intracellular pipette solutions containing 5 mM EGTA.

Furthermore, halothane and n-alkanols increased resting [Ca$^{2+}$], by 20–70% in mouse whole-brain synaptosomes [35], and several studies reported a [Ca$^{2+}$] rise, monitored by the Ca$^{2+}$-indicator arsenazo III, in octanol-treated squid axons [34,36,37]. In addition, a study on smooth muscle cells of porcine airway reported that halothane increases [Ca$^{2+}$], by a Ca$^{2+}$ leak through both inositol (1,4,5)-trisphosphate-(IP3)- and ryanodine-receptor channels (RyR) [38]. In pancreatic acinar cells, halothane and octanol induced a sustained [Ca$^{2+}$] increase [39]. In a study on malignant hyperthermia (MH), halothane, used in contracture testing for MH susceptibility, caused large elevations of myoplasmic [Ca$^{2+}$] [40].

In vascular smooth muscles, halothane causes both Ca$^{2+}$ release from stores and stimulates Ca$^{2+}$ uptake [41]; the halothane-induced Ca$^{2+}$ release from the stores is sensitive to both caffeine and IP3, suggesting that both RyR and IP3 channels of the ER play a role [41]. In a more recent study, volatile anesthetics were found to cause cell damage by abnormal calcium release from the ER via excessive activation of IP3-receptor channels [42], and the anesthetics’ neuroprotective and neurotoxic mechanisms have been shown to involve Ca$^{2+}$ release from the ER’s IP3-receptor channels [43]. In frog skeletal muscle, halothane was found to increase [Ca$^{2+}$], by releasing it from the sarcoplasmic reticulum (SR) via the RyR’s Ca$^{2+}$-release channel [44].

Morphological studies also reported changes in gap junction particle size and spacing with heptanol and other uncouplers known to increase [Ca$^{2+}$], in heart [45,46] and pancreas [47]. While the structural changes in gap junctions most likely are due to Ca$^{2+}$-activated CaM, it should be kept in mind that halothane has been reported to change the domain structure of a model membrane [48]. X-ray and neutron-diffraction studies of a binary lipid membrane demonstrate that halothane at physiological concentrations produces a pronounced redistribution of lipids between domains of different lipid types identified by different lamellar d-spacings and isotope composition. The redistribution of lipids between domains induced by anesthetics could in principle contribute to changes in gap junction particle size and spacing as well. Furthermore, gap junctions are rich in cholesterol [49–52]. Indeed, Bastiaanse and coworkers [53] have suggested that heptanol decreases gap junction channel conductance by decreasing the fluidity of cholesterol-rich domains in cardiac cells.

Plasma membranes of cells show asymmetric lipid distribution between the bilayer leaflets with a negative charge of the inner bilayer leaflet [54]. Phospholipid unsaturation is dramatically asymmetric, with the cytoplasmic leaflet being approximately twofold more unsaturated than the exoplasmic leaflet [55]. Atomic simulations and spectroscopy of leaflet-selective fluorescent probes reveal that the outer PM leaflet is more packed and less diffusive than the inner leaflet. The tightly packed outer leaflet may serve as an effective permeability barrier, while the more fluid inner leaflet may allow for rapid signal transmission. Thus, it is conceivable that the solubility of anesthetics is different in the two halves of the membrane bilayer.

The effect of caffeine and theophylline on uncoupling by anesthetics also indicates a participation of Ca$^{2+}$ release from stores, as previously shown with acidification [20,21]. Indeed, both caffeine and theophylline are known to increase [Ca$^{2+}$], by releasing it from Ca$^{2+}$ stores [56,57]. At mM concentrations, caffeine exerts a powerful effect on the SR by activating Ca$^{2+}$ release via RyR channels and perhaps also by inhibiting calcium reuptake [57]. Remarkably, the uncoupling mechanism of halothane seems to parallel in some way the mechanism of MH (reviewed in [58]), as in both cases the halothane-induced Ca$^{2+}$ release from the stores is potentiated by caffeine (see the caffeine–halothane contracture test). Perhaps, heptanol and halothane release Ca$^{2+}$ from stores by activating either the
IP3 or the RyR receptor (Figures 2A, 3A, 9A and 11, inset b, green arrows), while the addition of caffeine or theophylline releases Ca\(^{2+}\) by activating both IP3 and RyR receptors (Figures 2A, 3A and 9A, green and red arrows).

Although, in crayfish axons, Ca\(^{2+}\) release from IP3- and/or RyR-receptor channels seems most likely, one should keep in mind that the effect of xanthines, such as caffeine and theophylline, and anesthetics on Ca\(^{2+}\) release from stores is complex and depends on cell type. Indeed, Bezprozvanny and coworkers [59] demonstrated caffeine-induced inhibition of IP3-gated calcium channels from cerebellum incorporated into planar lipid bilayers. Furthermore, Parker and Ivorra [60] found that caffeine inhibits IP3-mediated liberation of intracellular Ca\(^{2+}\) in *Xenopus* oocytes, and Saleem and coworkers [61] showed that caffeine is a low-affinity antagonist of type 1 IP3 receptors (IP3R1), while it had no significant effect on IP3-evoked Ca\(^{2+}\) release via IP3R2 or IP3R3.

Joseph and coworkers [62] found that isoflurane modulates IP3R channel sensitivity to IP3 only at low, sub-saturating concentrations of IP3 (<0.1 µM), and showed that isoflurane causes Ca\(^{2+}\) release from the ER via this activation of IP3R which can regulate intracellular Ca\(^{2+}\) homeostasis and apoptosis. In frog skeletal muscle, halothane was found to increase [Ca\(^{2+}\)]\(_i\) by releasing it from the SR via the RyR’s Ca\(^{2+}\)-release channel [44]. Later, Laver and coworkers [63] found that halothane activation of RyR2 is different from that seen in the skeletal isofrom RyR1. Unlike RyR1, RyR2 was reported to be responsive to halothane and enflurane when recorded in bilayers [64] as well as in myocardial cells [65]. For gap junction channel gating in the heart, the cardiac RyR2 isofrom is more relevant than halothane has been reported to activate the cardiac-ryanodine-receptor channel, while isoflurane proved ineffective in activating RyR2 [63,64].

The magnitude of the effect of anesthetics of gap junction channel gating is also related to the type of gap junction protein expressed. Indeed, He and Burt [66] have reported that halothane has only a small uncoupling effect in cells expressing human Cx40, but has a great effect in cell expressing heteromeric (human) Cx40/Cx43 channels. This is interesting and may support the Ca\(^{2+}\) role, because Xu and coworkers [67] reported that ionomycin application increases [Ca\(^{2+}\)]\(_i\) and causes Gj to drop by 95% in N2a cells expressing human Cx43, but not in cells expressing human Cx40. Significantly, the human Cx40 does not have the CaM-binding site at the second half of the cytoplasmic loop (CL2), while Cx43 does [1,67].

The inhibitory action of the K\(^+\) channel blocker 4-aminopyridine (4-AP) on heptanol-induced gating (Figure 8A) is puzzling. It is interesting to note, however, that the potentiating effect of 4-AP on excitatory postsynaptic potentials (EPSP) in frog motoneurons is opposite to that of heptanol, which is an EPSP blocker [68,69]. Evidence that 4-AP fails to induce porcine MH [70], in spite of the fact that it mobilizes Ca\(^{2+}\) from intracellular stores [71,72], is also puzzling. Interestingly, 4-AP has been shown to modify phospholipid metabolism [73] and to stimulate protein phosphorylation in a Ca\(^{2+}\)-dependent manner [74].

Indirect evidence of [Ca\(^{2+}\)]\(_i\) rise with anesthetics is also provided by changes in membrane potential (MP). Indeed, the peak of membrane depolarization with heptanol (Figures 2A, 3A and 11, inset b) or halothane (Figure 9A), both alone (green arrows) or in the presence of caffeine (green and red arrows), matches the peak of Rj rise well (Figures 2B, 3B and 9B). Sauviat and coworkers [44] reported that the effects of halothane on membrane depolarization are likely to result from increased [Ca\(^{2+}\)]\(_i\), due to Ca\(^{2+}\) release via the RyR channel, perhaps as well as by the activation of Ca\(^{2+}\)-dependent Cl\(^-\) channels.

It is significant to compare the MP and Rj changes caused by anesthetics with those caused by cytosolic acidification, which is known to increase [Ca\(^{2+}\)]\(_i\) [21,75] (Figure 11). While with anesthetics the MP change is monophasic (Figures 2A, 3A, 9A and 11, inset b), with acidification it is biphasic (Figure 11A and inset a). The initial mild hyperpolarization with acidification is likely to result from Ca\(^{2+}\)-activation of K\(^+\) channels [76]—it is absent with anesthetics probably because they inhibit K\(^+\) channels [7]. The depolarization with both anesthetics and acidification is most likely caused by Ca\(^{2+}\) release from stores.
Figure 11. Time course of electrotonic potentials and $R_j$ in crayfish septate axons uncoupled by acetate (Ac) in the presence and absence of 10 mM caffeine (A, B). (A). Low-speed chart recording of membrane and electrotonic potentials. With Ac, $V_1$ and $V_2^*$ increase and $V_1^*$ and $V_2$ decrease (A), reflecting an increase in $R_j$ (B). Ac-caffeine causes a larger change in electrotonic potentials (A), indicating a larger increase in $R_j$ (B). Note that $R_j$ increases with Ac-caffeine 3–4 times as much as with Ac alone (B). With Ac or Ac-caffeine, there is a biphasic change in the membrane potential: a moderate hyperpolarization followed by depolarization ((A), and inset a, green and red arrows, respectively). In contrast, with heptanol, there is only depolarization (inset b, blue arrow, and Figures 2A and 3A). (A, B) and inset a were adapted from [20]. Inset b was adapted from [16].

5.2. Role of a Potential Soluble Intermediate—Calmodulin

Johnson and coworkers [14] proposed that alkanols act on an extracellular binding site, but did not consider the possible loss of a soluble intermediate, which might have been washed out by the internal perfusion of crayfish axons. Indeed, the standard internal solution (SIS) used contained (in mM): NaCl 15, K-fluoride 109, K-citrate 37, mannitol 96, HEPES 1, pH 7.5; the presence of K-fluoride and K-citrate might have lowered the $[Ca^{2+}]_i$, sufficiently to release CaM from binding sites and wash it away. Similarly, Eskandari and coworkers [17], who reported that octanol did not close lens connexin hemichannels in inside-out patches, did not consider the possibility that CaM was lost by exposing the cytosolic side of the membrane to solutions containing 5 mM EGTA and 1 mM CaCl$_2$. Furthermore, the idea that molecules such as heptanol, octanol and halothane and other general anesthetics only bind to an extracellular membrane site is hard to conceive because these molecules easily cross the plasma membrane, such that their concentration on both sides of the membrane is rapidly equilibrated. Consistent with the idea that CaM may play a role in the gating by anesthetics both provides evidence that halothane binds to CaM [78] and provides evidence that halothane, isoflurane and alcohols increase CaM’s $Ca^{2+}$ sensitivity [79,80].

On the other hand, Zhou and coworkers [81] showed that volatile anesthetics (VA) inhibit the activity of calmodulin by interacting with its hydrophobic core. Accordingly, Streiff and coworkers [82] predicted that volatile anesthetics bind to $Ca^{2+}$-bound CaM.
(holo-CaM), but not to apo-CaM. The VA-binding sites predicted for the structures of holo-CaM are located in hydrophobic pockets that form when the Ca\(^{2+}\)-binding sites in CaM are saturated. Volpi and coworkers [83] reported the antagonism of CaM by local anesthetics and the inhibition of the CaM-stimulated calcium transport of erythrocyte in an inside-out membrane vesicle. Levin and Blanck [79] observed a biphasic effect of halothane and isoflurane on calmodulin: at low concentrations of the anesthetics, the affinity of calmodulin for Ca\(^{2+}\) was decreased, while at higher concentrations, the affinity for Ca\(^{2+}\) was increased. Moreover, Rudnick et al. suggested that halothane mimics calmodulin-blocking agents and may alter CaM interaction with Ca\(^{2+}\)-dependent kinases.

6. Summary and Conclusions

When we studied the effect of anesthetics on channel gating in crayfish lateral giant axons, we were puzzled by the apparent contradiction between the striking potentiating effect of caffeine and theophylline and the apparent drop in [Ca\(^{2+}\)]\(_i\) [16]. Indeed, the effect of caffeine and theophylline on Rj and MP, and the fact that both the peaks of Rj rise and depolarization coincided, pointed to an increase in [Ca\(^{2+}\)]\(_i\). The reason that now, three decades later, we are reevaluating the earlier interpretation of the lack of Ca\(^{2+}\)\(_i\) participation in the effect of heptanol, halothane and isoflurane on gating is based on a number of facts, as follows:

1. The ion-selective electrodes [16] may not be reliable in the presence of anesthetics [18].
2. Anesthetics increase the CaM’s Ca\(^{2+}\) sensitivity [79,80]. This might have activated many CaM molecules, resulting in significant drop in [Ca\(^{2+}\)]\(_i\).
3. Halothane binds to CaM [78].
4. Anesthetics cause Ca\(^{2+}\) release via both IP3- and RyR-receptor channels [38].
5. Both caffeine and theophylline increase [Ca\(^{2+}\)], by releasing Ca\(^{2+}\) from stores [56,57].
6. Heptanol and other uncouplers known to raise [Ca\(^{2+}\)]\(_i\) cause gap junction particle crystallization in heart [45,46] and pancreas [47].
7. The interpretation of an extracellular gap junction binding site for anesthetics’ effect on coupling in internally perfused axons and inside-out membrane patches [14,17] is questionable, and may in fact suggest the loss of a cytosolic soluble intermediate (possibly CaM).
8. Lack of cyclic nucleotide role, excluded by experiments with IBMX (a phosphodiesterase inhibitor), forskolin (an activator of adenylate cyclase) or diffusible cAMP and cGMP.
9. Absence of kinase C role, tested with TPA (activator) or H7 (inhibitor).
10. Lack of adenosine-receptor role, tested with adenosine and IBMX treatments.
11. Evidence that the peak of depolarization caused by anesthetics or anesthetics–caffeine matches the peak of Rj rise.
12. Ryanodine-induced inhibition of Rj rise with heptanol–caffeine.
13. There is evidence that the second exposure to heptanol or heptanol–caffeine has a much smaller effect on Rj, suggesting partial depletion of Ca\(^{2+}\)\(_i\) stores by the previous exposure.

In view of these findings, while we cannot exclude a direct effect of anesthetics on gap junction proteins and/or lipids, we propose the hypothesis that heptanol, halothane and isoflurane induce gap junction channel gating by releasing Ca\(^{2+}\) from internal stores and activating CaM. We realize that this hypothesis needs to be further tested experimentally and it should be stressed that our data were obtained entirely on crayfish axons, which express innexins rather than connexins. Furthermore, there are mysterious data; for example, there is evidence for a strong inhibitory action of 4-aminopyridine on the heptanol effect (Figure 8A). Since 4-aminopyridine is a K\(^+\) channel blocker, the effect of other K channel blockers should be tested as well. Moreover, the CaM hypothesis needs to be tested with CaM inhibitors, inhibition of CaM expression and overexpression of CaM mutants.
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