Voltage- and Calcium-dependent Inactivation of Calcium Channels in Lymnaea Neurons

Shalini Gera and Lou Byerly

From the Department of Biological Sciences, University of Southern California, Los Angeles, California 90089-2520

abstract  

Ca$^{2+}$ channel inactivation in the neurons of the freshwater snail, Lymnaea stagnalis, was studied using patch-clamp techniques. In the presence of a high concentration of intracellular Ca$^{2+}$ buffer (5 mM EGTA), the inactivation of these Ca$^{2+}$ channels is entirely voltage dependent; it is not influenced by the identity of the permeant divalent ions or the amount of extracellular Ca$^{2+}$ influx, or reduced by higher levels of intracellular Ca$^{2+}$ buffering. Inactivation measured under these conditions, despite being independent of Ca$^{2+}$ influx, has a bell-shaped voltage dependence, which has often been considered a hallmark of Ca$^{2+}$-dependent inactivation. Ca$^{2+}$-dependent inactivation does occur in Lymnaea neurons, when the concentration of the intracellular Ca$^{2+}$ buffer is lowered to 0.1 mM EGTA. However, the magnitude of Ca$^{2+}$-dependent inactivation does not increase linearly with Ca$^{2+}$ influx, but saturates for relatively small amounts of Ca$^{2+}$ influx. Recovery from inactivation at negative potentials is biexponential and has the same time constants in the presence of different intracellular concentrations of EGTA. However, the amplitude of the slow component is selectively enhanced by a decrease in intracellular EGTA, thus slowing the overall rate of recovery. The ability of 5 mM EGTA to completely suppress Ca$^{2+}$-dependent inactivation suggests that the Ca$^{2+}$ binding site is at some distance from the channel protein itself. No evidence was found of a role for serine/threonine phosphorylation in Ca$^{2+}$ channel inactivation. Cytochalasin B, a microfilament disrupter, was found to greatly enhance the amount of Ca$^{2+}$ channel inactivation, but the involvement of actin filaments in this effect of cytochalasin B on Ca$^{2+}$ channel inactivation could not be verified using other pharmacological compounds. Thus, the mechanism of Ca$^{2+}$-dependent inactivation in these neurons remains unknown, but appears to differ from those proposed for mammalian L-type Ca$^{2+}$ channels.

key words: molluscs • cytochalasin B • intracellular Ca$^{2+}$ • Ca$^{2+}$ buffering

INTRODUCTION

Inactivation of Ca$^{2+}$ channels is thought to be of two types—voltage-dependent inactivation, like that of Na$^+$ channels (as originally proposed by Hodgkin and Huxley, 1952), and Ca$^{2+}$-dependent inactivation, which is determined by the intracellular Ca$^{2+}$ concentration (Brehm and Eckert, 1978; Tillotson, 1979). Ca$^{2+}$-dependent inactivation is an important example of negative feedback through which the Ca$^{2+}$ levels inside the cell can regulate the amount of Ca$^{2+}$ influx. It is, as yet, unknown whether Ca$^{2+}$ merely influences the rate constants of entry to and departure from the inactivated state, or if it actually causes the channels to enter an entirely different inactivation state.

In this paper, we have studied the inactivation of Ca$^{2+}$ channels in Lymnaea neurons. Ca$^{2+}$-dependent inactivation in molluscan neurons has received considerable attention; it was in these neurons that this phenomenon was first characterized (Tillotson, 1979; Eckert and Tillotson, 1981). However, it is necessary to reexamine the inactivation of molluscan Ca$^{2+}$ channels because the original studies did not take into account the outward proton current, which was discovered later in snail neurons (Thomas and Meech, 1982; Byerly et al., 1984a) and can easily be misinterpreted as Ca$^{2+}$ current inactivation. Also, the early studies were inconclusive about the amount of voltage-dependent inactivation present in molluscan neurons (Eckert and Chad, 1984). These studies, and others (Brehm and Eckert, 1978; Ashcroft and Stanfield, 1982; Lee et al., 1985) established that Ca$^{2+}$ channel inactivation under some conditions has a bell-shaped voltage dependence; i.e., depolarizations to potentials that elicit large Ca$^{2+}$ currents also cause maximal amounts of inactivation. This is consistent with the idea that inactivation is caused by Ca$^{2+}$ influx, and thus a bell-shaped inactivation curve is often interpreted to indicate the presence of Ca$^{2+}$-dependent inactivation.

In this study, we show that Ca$^{2+}$ channel inactivation in Lymnaea neurons has both Ca$^{2+}$- and voltage-dependent components, and that both of these components have a bell-shaped voltage dependence. From the kinetics of the development of and the recovery from inactivation, we infer that there are two distinct inactivation states, even in the absence of Ca$^{2+}$-dependent inactivation, and an increase in Ca$^{2+}$ causes a greater occu-
Ca\(^{2+}\)-channel Inactivation in Lymnaea Neurons

MATERIALS AND METHODS

Cell Preparation and Electrophysiology

Neurons were dissociated from the pedal, parietal, and visceral ganglia of adult Lymnaea stagnalis, and prepared for patch clamp experiments as previously described (Johnson and Byerly, 1993a). The cells used for this study were nearly spherical, and their diameters ranged from 50 to 75 \(\mu\)m. The Axopatch 200A patch clamp amplifier (Axon Instruments) was used in this study to measure currents. pClamp software (version 6.0) was used for data acquisition (Clampex) and analysis (Clampfit). The patch clamp electrodes typically had resistances of 1 M\(\Omega\) and tip diameters of 12-16 \(\mu\)m. Series resistance (usually \(\sim\)2-4 M\(\Omega\)) was electronically compensated to \(\geq\)90%. Inactivation measurements were taken at least 10 min after entering the whole-cell configuration, unless otherwise noted, to allow for the diffusion of the electrode solution into the cell. Junction potential errors (described in Hagiwara and Ohmori, 1982; Neher, 1995) have not been corrected for in these experiments and are expected to be approximately \(-10\) to \(-15\) mV (Byerly and Hagiwara, 1982). Linear leak currents and capacitive transients are subtracted using a P/4 protocol. Currents recorded in the standard solutions are comprised of voltage-gated Ca\(^{2+}\) and H\(^{+}\) currents (Byerly et al., 1984a). These currents are not contaminated with outward Cs\(^{+}\) or Cl\(^{-}\) currents, since replacing internal Cs\(^{+}\) with N-methyl-d-glucamine\(^{+}\), or external Cl\(^{-}\) with methanesulfonate\(^{-}\) (CH\(_3\)SO\(_3\)) does not produce any change in the shape of the recorded currents.

Internal Perfusion Experiments

Internal perfusion experiments were done following the method described by Neher and Eckert (1988). The intracellular solution of the cell could be completely exchanged in \(<6\) min, as estimated from measuring outward K\(^{+}\) currents while replacing intracellular K\(^{+}\) with Cs\(^{+}\).

Solutions

The Lymnaea saline used for dissociation and storage of cells contains 50 mM NaCl, 2.5 mM KCl, 4 mM MgCl\(_2\), 4 mM CaCl\(_2\), 10 mM HEPES (N-[2-hydroxyethyl] piperazine-N’-[2-ethane sulfonic acid]), adjusted to pH 7.4 with NaOH. The standard extracellular saline used for recording Ca\(^{2+}\) or Ba\(^{2+}\) currents is composed of 76 mM TrisCl and 10 mM CaCl\(_2\) or BaCl\(_2\), and is adjusted to pH 7.4. In some experiments where the concentration of Ca\(^{2+}\) in the external solution is reduced to 1 mM, 9 mM MgCl\(_2\) is added to keep the total concentration of divalent ions constant. All of the intracellular solutions contain 50 mM HEPES, 0.5 mM MgCl\(_2\), 3-12 mM CsCl, 15-20 mM aspartic acid, and 2 mM MgATP, with varying amounts of calcium buffers, adjusted to pH 7.3 with CsOH, thus making Cs\(^{+}\) the main intracellular cation. The different levels of intracellular Ca\(^{2+}\) buffers used in this study are 0.1 mM EGTA, 5 mM EGTA, 5 mM EGTA with 2.5 mM CaCl\(_2\), 5 mM EGTA with 4.5 mM CaCl\(_2\), and 11 mM 1,2-bis(2- amino phenoxyl) ethane-N, N’, N’-tetraacetic acid (BAPTA) with 1 mM CaCl\(_2\). These solutions adjusted to various free Ca\(^{2+}\) levels were used for calibrating Fura-2.

Fura-2 Measurements of Free Ca\(^{2+}\) in Cells

Free Ca\(^{2+}\) levels in cells were measured by loading the cells with an intracellular solution containing 10 \(\mu\)M Fura-2, a Ca\(^{2+}\)-sensitive ratiometric dye (Grynkiewicz et al., 1985). Fura-2 fluorescence was measured in a confocal arrangement using pinholes as previously described (Johnson and Byerly, 1993b). The calibration solutions described above were used for calibrating Fura-2 in microuettes (borosilicate microslides with an optical path length of 50 \(\mu\)m; VitroCom Inc.). The UV exciting Fura-2 was controlled by a Lambda 10-2 filter wheel (Sutter Instrument Co.) and limited to 200-ms pulses of 380 and 360 nm radiation for each measurement of Ca\(^{2+}\). Background fluorescence measurements were made before the patch membrane was ruptured, and later subtracted from all records.

Measurement of Inactivation

In the studies reported in this paper, inactivation has been measured using mainly a three-pulse protocol (see Fig. 1A). First a short test pulse (10 ms) to +40 mV is applied, then a conditioning pulse (150 ms) of variable amplitude, followed by a gap (20 ms) at the holding potential (−60 mV), and, finally, a second test pulse to +40 mV is applied. The inactivation caused by the conditioning pulse is calculated as the percent reduction in the test pulse current after the conditioning pulse. In an earlier study (Gera and Byerly, 1999), we have shown that this method of measuring Ca\(^{2+}\) channel inactivation avoids errors caused by H\(^{+}\) currents, which can be prominent in snail neurons under the conditions used to record Ca\(^{2+}\) currents (Byerly et al., 1984a). This is primarily because H\(^{+}\) current activates slowly at the test pulse po-
and Byerly, 1999). Byerly and Gera observing that as the permeant ion concentration approaches zero, the tail current measurement also approaches zero (Gera 100)

The conditioning-pulse voltage. The inward currents for each cell were normalized to the maximum inward current, and then averaged across all cells. Error bars in this and the following figures represent SEM.

**RESULTS**

**Ca**

Inactivation is measured using a three-pulse protocol (Fig. 1 A). In this protocol, a short test pulse is applied before and after a conditioning pulse to variable potentials, and inactivation is measured as the percentage reduction in the test pulse current due to the conditioning pulse. There is a 20-ms gap at the holding potential between the conditioning pulse and the second test pulse, which is necessary to allow background currents activated during the conditioning pulse to completely deactivate. However, it also results in some Ca2+ channels recovering from inactivation during the gap. Thus, our measurement of inactivation relates to the fraction of Ca2+ channels that are still inactivated at the end of the 20-ms gap. Inactivation measured this way in cells containing a high level of intracellular Ca2+ buffer (5 mM EGTA) is relatively small (the peak inactivation being ~0.25), and exhibits a bell-shaped voltage dependence (Fig. 1 B). We show below that the inactivation measured in these conditions is independent of Ca2+ influx.

A common test for the presence of Ca2+-dependent inactivation is to compare inactivation of Ca2+ currents with that of Ba2+ currents. The underlying assumption is that the intracellular site, which mediates Ca2+-dependent inactivation, is less sensitive to Ba2+ ions than to Ca2+ ions. Therefore, the current-dependent component of inactivation during Ba2+ influx would be reduced, as compared with that during Ca2+ influx. Exchanging external Ca2+ with Ba2+ does not cause a reduction in the levels of peak inactivation measured in Lymnae neurons containing 5 mM EGTA (Fig. 2 A), suggesting that there is very little Ca2+-dependent inac-

**Analysis**

Statistical analyses were performed using Systat software (Version 6; SPSS Inc.) and the corrected R2 parameter was used to determine the quality of fit of a model to the experimental data.

**Figure 1.** Ca2+-channel inactivation measurements in cells containing 5 mM EGTA. (A) Method for measuring Ca2+-channel inactivation. Command voltage traces for three different conditioning pulse voltages (top) and the corresponding current traces (bottom) are shown. A 10-ms test pulse (TP) to +40 mV is followed by a 150-ms conditioning pulse (CP), a 20-ms gap at the holding potential (-60 mV), and a second test pulse to +40 mV. (B) Average inactivation measurements ( ■, n = 31) and the normalized peak inward currents (□, n = 32) as a function of the conditioning-pulse voltage. The inward currents for each cell were normalized to the maximum inward current, and then averaged across all cells. Error bars in this and the following figures represent SEM.
tivation in this case. The leftward shift in the inactivation curve observed with external Ba\(^{2+}\) can be explained on the basis of the shift of Ba\(^{2+}\) current activation to potentials lower than those for Ca\(^{2+}\)-current activation.

Ca\(^{2+}\)-dependent inactivation also implies that inactivation should depend on the amount of Ca\(^{2+}\) influx. This is not true for Lymnaea neurons containing 5 mM EGTA. The standard extracellular solution in our experiments contains 10 mM Ca\(^{2+}\); reducing this concentration 10-fold to 1 mM does not cause a significant reduction in the amount of inactivation measured (Fig. 2 B), even though the current magnitude decreases fourfold.

We also compared inactivation in cells dialyzed with a 5 mM EGTA solution to that in cells dialyzed with an 11 mM BAPTA solution. In the second case, not only is there a higher concentration of a Ca\(^{2+}\) buffer, but the buffer used (BAPTA) is also substantially faster. Consequently, the Ca\(^{2+}\) transients in cells with 11 mM BAPTA should be substantially smaller than those in cells with 5 mM EGTA (Neher, 1986). However, the inactivation measured in the two cases is similar (Fig. 2 C), showing that the Ca\(^{2+}\) transients in 5 mM EGTA are already too small to affect inactivation.

Another way of assessing the contribution of Ca\(^{2+}\) influx in Ca\(^{2+}\)-channel inactivation is to eliminate all Ca\(^{2+}\) influx, and measure the inactivation of the Ca\(^{2+}\)-channel current carried by monovalents (Cox and Dunlap, 1994). However, replacing permeant divalent cations with impermeant ones (such as Mg\(^{2+}\)) in the extracellular solution causes Lymnaea neurons to develop a large leak current, possibly because of loss of K\(^{+}\) channel selectivity (Armstrong and Lopez-Barneo, 1987; Armstrong and Miller, 1990). As an alternative approach, we have used 100 \(\mu\)M external Cd\(^{2+}\), a Ca\(^{2+}\) channel blocker, to eliminate Ca\(^{2+}\) influx. Cd\(^{2+}\) was chosen because it is known to block Ca\(^{2+}\) channels in a voltage-dependent manner (Byerly et al., 1984b; Swandulla and Armstrong, 1989); it is a more effective blocker at positive potentials than at negative ones, and, consequently, currents during a positive pulse are blocked more than the tail currents (measured at negative potentials). Thus, using 100 \(\mu\)M Cd\(^{2+}\) in the external solution, we can block 90\% of the Ca\(^{2+}\) influx during a conditioning pulse and still measure inactivation using the tail currents, which are only blocked by 40\% (see materials and methods). We find that for cells containing 5 mM EGTA, the amplitude of peak inactivation does
not change when 100 μM Cd2+ is added to the external solution (Fig. 2D), which supports our conclusion that inactivation in 5 mM EGTA is independent of Ca2+ influx. (We consistently observe that Cd2+ also causes a change in the shape of the inactivation curve, which we do not understand.)

We conclude from the four types of experiments described above (Fig. 2, A–D) that Ca2+-channel inactivation in cells containing 5 mM EGTA is entirely voltage dependent and is independent of Ca2+ influx. Ca2+ channels in Lymnaea neurons are capable of exhibiting Ca2+-dependent inactivation when the intracellular Ca2+ buffering is lowered (shown below). Thus, we conclude that 5 mM EGTA reduces intracellular Ca2+ transients to a size where they are incapable of activating the site that mediates Ca2+-dependent inactivation.

Ca2+ Channel Inactivation in Cells Containing 0.1 mM EGTA Has a Current-dependent Component

To demonstrate that Ca2+ channels in Lymnaea neurons are capable of exhibiting Ca2+-dependent inactivation, we compared Ca2+-channel inactivation in two identical populations of neurons loaded with different amounts of intracellular Ca2+ buffer. Cells containing 0.1 mM EGTA show substantially more inactivation than those containing 5 mM EGTA (Fig. 3A), indicating that increased levels of intracellular Ca2+ lead to an increase in Ca2+ channel inactivation. This result was confirmed with internal perfusion experiments in which intracellular solutions were changed while recording from one cell. Inactivation was first measured when the cells were perfused with a solution containing 5 mM EGTA, and was found to increase if the intracellular solution was changed to one containing only 0.1 mM EGTA (Fig. 3B). In control experiments, in which the second intracellular solution perfused into the cells was the same as the first, inactivation was not affected by the exchange (data not shown).

To ensure that it is indeed the Ca2+-buffering properties of EGTA and not some other pharmacological property that contributes to Ca2+-channel inactivation, we measured inactivation in cells where the intracellular solution contained 5 mM EGTA, loaded with different amounts of CaCl2 (Fig. 3C). Cells containing 5 mM EGTA alone show smaller magnitudes of inactivation than those in which 5 mM EGTA has been loaded with 2.5 mM CaCl2 or 4.5 mM CaCl2, suggesting that it is the

![Figure 3. Inactivation of Ca2+ channels is dependent upon intracellular Ca2+ buffering. (A) Average inactivation curves measured for cells containing 0.1 (○, n = 41) and 5 (□, n = 31) mM EGTA. (B) Average inactivation measured in cells first when they contain an internal solution containing 5 mM EGTA (□), and after the internal solution has been changed to one that contains 0.1 mM EGTA (○), n = 6. In these experiments, the test pulse potential is +30 mV. (C) Inactivation was measured in cells containing different amounts of free EGTA. Each point (○) represents the peak inactivation of a cell containing the specified levels of EGTA and CaCl2 (in millimolar); n = 15 for 5 EGTA alone; n = 7 for 5 EGTA and 2.5 CaCl2; n = 3 for 5 EGTA and 4.5 CaCl2; n = 15 for 0.1 EGTA alone. All the measurements of inactivation included here were taken in the same time period from similar populations of cells. (D) Ca2+-dependent inactivation is independent of steady state levels of Ca2+. Cells were perfused with a solution containing either 0.1 mM EGTA (○) or 5 mM EGTA/2.5 mM CaCl2 (■). Inactivation is measured using the three-pulse protocol, while steady state levels of Ca2+ are monitored using Fura-2. Inactivation and Ca2+ levels within the cell are measured at different times after going whole-cell. No measurements are taken within the first 5 min to allow for equilibration of Fura-2 and the buffer solution concentrations within the cell. Each symbol represents the average inactivation and Ca2+ concentration in a single cell.]
concentration of free Ca²⁺ buffer that is important for determining the amount of inactivation. These solutions of a fixed amount of Ca²⁺ buffer and variable amounts of CaCl₂ also contain slightly different levels of free Ca²⁺. But the free Ca²⁺ levels in all these solutions are low (<10⁻⁶ M) and do not appear to determine the amount of Ca²⁺-dependent inactivation (Fig. 3 D, discussed later).

The increased Ca²⁺ channel inactivation in Lymnaea neurons containing 0.1 mM EGTA is dependent upon Ca²⁺ influx. In these cells, exchange of an external Ca²⁺-containing solution with a Ba²⁺-containing solution causes a significant decline in peak inactivation (Fig. 4 A), even though the magnitude of Ba²⁺ current is two to three times larger than that of Ca²⁺ current. Similarly, changing the external Ca²⁺ concentration from 10 to 1 mM also leads to a decrease in the total Ca²⁺-channel inactivation measured in cells containing 0.1 mM EGTA (Fig. 4 B). Furthermore, 100 μM Cd²⁺ in the extracellular solution causes a substantial decrease in inactivation measured in cells containing 0.1 mM EGTA (Fig. 4 C); the remaining inactivation is not significantly different from that in cells with 5 mM EGTA under similar conditions (Fig. 2 D).

From these experiments, we conclude that Lymnaea neurons containing 0.1 mM EGTA have both Ca²⁺- and voltage-dependent components, while those with 5 mM EGTA exhibit only the voltage-dependent component of Ca²⁺ channel inactivation. For the purposes of this study, we define Ca²⁺-dependent inactivation in these cells as the difference in inactivation observed with cells containing 0.1 and 5 mM EGTA.

Recovery from and Development of Inactivation

We measured the rates with which the Ca²⁺ channels recover from inactivation in cells containing 5 or 0.1 mM EGTA. This was done by varying the lengths of the gap after the conditioning pulse in a protocol that measures inactivation using tail currents (see materials and methods). Using this protocol, we find that the recovery of Ca²⁺ channels from inactivation has a biexponential time course at −60 mV (τfast = 15 ms, τslow = 600 ms; Fig. 5 A). After a conditioning pulse to +120 mV, the rate of recovery in 0.1 mM EGTA is not substantially different from that in 5 mM EGTA. However, after conditioning pulses to +40 and +60 mV (which cause maximal inactivation) the slow component of recovery is much larger for 0.1 mM EGTA compared with that for 5 mM EGTA. This is accompanied by a modest decrease in the magnitude of the fast component of recovery in 0.1 mM EGTA. The fast component of recovery decays rapidly in the first 20 ms; thus, most of the difference observed between inactivation measured in 0.1 and 5 mM EGTA measured using the 20-ms gap (as in Fig. 3 A) is due to the differences in the magnitude

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**Figure 4.** Inactivation in cells with 0.1 mM EGTA is dependent on Ca²⁺ influx. (A) Average inactivation curves measured in external Ca²⁺ (■), and after substitution with Ba²⁺ (□) in cells containing 0.1 mM EGTA (n = 9). (B) Average inactivation curves obtained in cells containing 0.1 mM EGTA in 10 (■) and 1 (□) mM external Ca²⁺, n = 6. (C) Average inactivation curves in control external solution (□), and upon the addition of 100 μM external Cd²⁺ (■) in cells containing 0.1 mM EGTA (n = 4).
of the slow component of recovery in the two conditions. The two separate time constants for the rate of recovery from inactivation indicate that there are two different inactivated states from which the channels are recovering—at negative potentials, recovery from one inactivated state takes place at a considerably faster rate than from the other inactivated state. The effect of Ca\(^{2+}\) is to increase the occupancy of the latter state.

We also measured the development of inactivation during conditioning pulses to +60 and +120 mV. The inactivation measurements were made after allowing the channels to recover for 20 ms after conditioning pulses of variable length, and thus are not independent of the recovery rates. During a pulse to +120 mV, inactivation develops with a single time constant of 250 ms in both 0.1 and 5 mM EGTA (Fig. 5 B). However, when the conditioning pulse is to +60 mV, there is an additional faster component (τ = 50 ms), and the amplitude of this component is three times as large in 0.1 as in 5 mM EGTA. These experiments also indicate that the Ca\(^{2+}\) influx during the tail currents at the end of the conditioning pulse does not contribute significantly towards inactivation, since inactivation approaches zero as the conditioning pulse becomes very short. In the discussion, we develop a model of Ca\(^{2+}\)-induced inactivation that can account for our observations regarding the kinetics of inactivation in 5 and 0.1 mM EGTA.

**Ca\(^{2+}\)-dependent Inactivation Does Not Depend Linearly on Ca\(^{2+}\) Influx**

We have shown that for cells containing 0.1 mM EGTA, the Ca\(^{2+}\)-dependent component of inactivation can be reduced by reducing Ca\(^{2+}\) influx (Fig. 4). The magnitude of Ca\(^{2+}\)-dependent inactivation, however, is not linearly related to the amount of Ca\(^{2+}\) influx. This conclusion is demonstrated by the experiment in which the Ca\(^{2+}\) channel blocker Cd\(^{2+}\) was used to reduce the influx of Ca\(^{2+}\) during conditioning pulses. This experiment is analogous to the one described above using Cd\(^{2+}\) to block the Ca\(^{2+}\) current, but Co\(^{2+}\) is a weaker Ca\(^{2+}\) channel blocker than Cd\(^{2+}\), and exerts a simpler, non-voltage-dependent block of current (Byerly et al., 1984b). In these experiments, 1 mM Co\(^{2+}\) was used to replace 1 mM of the 10-mM external Ca\(^{2+}\), resulting in a >50% block of Ca\(^{2+}\) currents; yet it was found to have very little effect on Ca\(^{2+}\)-channel inactivation (Fig. 6 A). Therefore, relatively small amounts of Ca\(^{2+}\) influx during the conditioning pulse may be sufficient to cause maximal Ca\(^{2+}\)-dependent inactivation. We also observe that Ca\(^{2+}\)-dependent inactivation reduces to a half when the external Ca\(^{2+}\) is reduced from 10 to 1 mM (Fig. 4 B), even though the peak Ca\(^{2+}\) current is reduced to a fourth. Ca\(^{2+}\)-dependent inactivation, in this case, is the difference between the inactivation measured in cells with 0.1 mM EGTA (Fig. 4 B) and that in cells containing 5 mM EGTA (Fig. 2 B).

Our conclusion that the amount of Ca\(^{2+}\)-dependent inactivation is not simply related to the magnitude of Ca\(^{2+}\) influx during the conditioning pulse is also supported by other observations. In cells perfused with 0.1 mM EGTA, the decrease in Ca\(^{2+}\) current over time (due to rundown) is not accompanied by any significant changes in inactivation. Also, there is no obvious correlation between the peak inactivation and the current density measured in the 53 cells containing 0.1 mM EGTA that we studied (Fig. 6 B). It should be noted, however, that peak inactivation is the sum of voltage- and Ca\(^{2+}\)-dependent inactivation. It is possible that in the absence of a strong dependence of Ca\(^{2+}\)-dependent inactivation on Ca\(^{2+}\) current density, the random variation in voltage-dependent inactivation obscures any weaker correlation between the two quantities in Fig. 6 B.

**Intracellular Sources of Calcium Do Not Contribute Towards Ca\(^{2+}\)-dependent Inactivation**

Effects of basal Ca\(^{2+}\) levels on inactivation. We investigated the possibility that the basal (i.e., steady state) levels of
Ca\textsuperscript{2+} within the cell may affect Ca\textsuperscript{2+}-channel inactivation. Fura-2 measurements indicated that steady state Ca\textsuperscript{2+} levels in cells containing 0.1 mM EGTA are much higher (70–300 nM) than those in cells containing 5 mM EGTA (5–20 nM). This difference occurs despite the fact that the two intracellular solutions, containing 5 and 0.1 mM EGTA, respectively, have similar low levels of free Ca\textsuperscript{2+} (2–10 nM, according to Fura-2 measurements in microcuvettes). The increase in free Ca\textsuperscript{2+} levels in cells perfused with a poorly buffered solution is probably due to a high rate of Ca\textsuperscript{2+} influx through the plasma membrane. Thus, it is possible that increased Ca\textsuperscript{2+}-channel inactivation observed in cells with 0.1 mM EGTA results from higher basal levels of free Ca\textsuperscript{2+} inside the cells.

To examine whether differences in basal Ca\textsuperscript{2+} levels can account for the variations in inactivation measured in different cells, we measured free Ca\textsuperscript{2+} levels (using Fura-2) and inactivation in cells perfused with an intracellular solution containing either 0.1 mM EGTA or 5 mM EGTA/2.5 mM Ca\textsuperscript{2+}. These two intracellular solutions were chosen since they result in comparable values of free intracellular Ca\textsuperscript{2+} levels, but have very different Ca\textsuperscript{2+}-buffering capacities. We find that while cells with 0.1 mM EGTA consistently show more inactivation than cells with 5 mM EGTA/2.5 mM Ca\textsuperscript{2+}, there is no correlation between peak inactivation and steady state Ca\textsuperscript{2+} levels for the 0.1 mM EGTA data (R\textsuperscript{2} = 0.021, Fig. 3 D). This leads us to believe that the site of inactivation is not sensitive to resting levels of Ca\textsuperscript{2+} (≤300 nM); instead, intracellular domains of high Ca\textsuperscript{2+} that are transiently set up when Ca\textsuperscript{2+} channels are activated must be mediating Ca\textsuperscript{2+}-channel inactivation.

Intracellular sources of calcium ions do not contribute to Ca\textsuperscript{2+}-dependent inactivation. Recent studies have shown that, in ventricular myocytes, an influx of Ca\textsuperscript{2+} through voltage-gated Ca\textsuperscript{2+} channels triggers a release of Ca\textsuperscript{2+} from the sarcoplasmic reticulum, and it is this release of Ca\textsuperscript{2+} from intracellular stores that is largely responsible for Ca\textsuperscript{2+} channel inactivation in these cells (Balke and Wier, 1991; Adachi-Akahane et al., 1996). Such a scenario could potentially explain the nonlinear dependence of Ca\textsuperscript{2+}-dependent inactivation on Ca\textsuperscript{2+} influx (described above). Therefore, we investigated if Ca\textsuperscript{2+}-induced Ca\textsuperscript{2+} release (CICR) may also play a role in the Ca\textsuperscript{2+}-dependent inactivation of Lymnaea neurons. Ryanodine is known to be an inhibitor of CICR (Friel and Tsien, 1992; Orkand and Thomas, 1995). Ryanodine (10 μM), applied extracellularly, does not produce any substantial change in inactivation in cells containing 0.1 mM EGTA (peak inactivation is 0.57 ± 0.07 in the control solution and 0.61 ± 0.08 in 10 μM ryanodine, n = 4). Low caffeine (1 mM) is thought to potentiate CICR, while 10 mM caffeine depletes the internal stores of calcium (Friel and Tsien, 1992; Orkand and Thomas, 1995). However, neither concentration of caffeine has any effect on Ca\textsuperscript{2+} channel inactivation: peak inactivation is 0.56 ± 0.03 (control), compared with 0.63 ± 0.05, n = 2 (1 mM caffeine), and 0.53 ± 0.03 (control), compared with 0.58 ± 0.05 (n = 3) in 10 mM caffeine. While intracellular Ca\textsuperscript{2+} levels in Lymnaea neurons were not monitored during these experiments, similar applications of ryanodine and caffeine are effective in changing intracellular Ca\textsuperscript{2+} levels in other mollusc neurons (Orkand and Thomas, 1995). Therefore, we conclude that CICR is not involved in Ca\textsuperscript{2+}-current inactivation in Lymnaea neurons, and that influx of extracellular Ca\textsuperscript{2+} through the voltage-gated Ca\textsuperscript{2+} channels is wholly responsible for the Ca\textsuperscript{2+}-dependent component of inactivation.

Intracellular Proteins Involved in Ca\textsuperscript{2+}-dependent Inactivation

Different mechanisms for Ca\textsuperscript{2+}-dependent inactivation have been proposed in the literature. Some researchers
have concluded that Ca\(^{2+}\) may bind directly to the Ca\(^{2+}\) channel (de Leon et al., 1995; Neely et al., 1994), or to a protein, such as calmodulin, that is closely associated with the Ca\(^{2+}\) channel (Peterson et al., 1999; Qin et al., 1999; Zühlke et al., 1999), causing Ca\(^{2+}\)-dependent inactivation. The result that 5 mM EGTA can completely suppress Ca\(^{2+}\)-dependent inactivation leads us to question this model in Lymnaea neurons (see discussion). We think it is likely that intracellular Ca\(^{2+}\) binds to some other cytoplasmic protein at some distance from the channel, which in turn influences the inactivation of Ca\(^{2+}\) channels. This intermediate protein could be involved in phosphorylation (as proposed by Chad and Eckert, 1986; Armstrong, 1989) or be a component of the cortical cytoskeleton (Johnson and Byerly, 1993a, 1994). We examine these hypotheses below.

Serine/threonine phosphorylation is not involved in Ca\(^{2+}\)-channel inactivation. It has been proposed that an increase in cytoplasmic Ca\(^{2+}\) levels may lead to an activation of a Ca\(^{2+}\)-dependent phosphatase (or a kinase) that may alter the phosphorylation state of the Ca\(^{2+}\) channel leading to an increase in inactivation (Chad and Eckert, 1986). In support of this theory, Schuhmann et al. (1997) have shown that cyclosporin A, an inhibitor of Ca\(^{2+}\)-activated protein phosphatase 2B (calcineurin), can substantially reduce the magnitude of Ca\(^{2+}\)-dependent inactivation of L-type Ca\(^{2+}\) channels in smooth muscle cells. However, Ca\(^{2+}\) channel inactivation in other preparations has been shown to be insensitive to phosphorylation (Fryer and Zucker, 1993; Imredy and Yue, 1994; Branchaw et al., 1997). We tested for an effect of phosphorylation in Lymnaea Ca\(^{2+}\)-channel inactivation by extracellularly applying the drug H-7, a broad-range serine/threonine kinase inhibitor. We find that H-7 has no significant effect upon inactivation in cells perfused with 0.1 mM EGTA solution (peak inactivation goes from 0.42 ± 0.04 in control external solution to 0.45 ± 0.04 in the presence of 250 μM H-7, n = 4). Okadaic acid, a phosphatase inhibitor, also has no effect upon inactivation in cells with 0.1 mM EGTA (peak inactivation is 0.60 ± 0.08 in the control solution and 0.61 ± 0.07 in the presence of 5 μM okadaic acid, n = 3). Since okadaic acid is not effective against calcineurin, we also tested for an effect of Cyclosporin A (CsA), a specific inhibitor of calcineurin, on inactivation of Ca\(^{2+}\) channels. Neither acute application of 10 μM CsA during a whole-cell experiment, nor pretreatment of cells with 10 μM CsA for at least 20 h, causes any change in the inactivation of Ca\(^{2+}\) channels (peak inactivation is 0.50 ± 0.02 for cells pretreated with CsA, n = 4, and is 0.52 ± 0.04, n = 7, for cells in the control saline under similar conditions). While we don’t have any direct evidence for an effect of these drugs on phosphorylation in Lymnaea neurons, they have been shown to inhibit phosphorylation, or de-phosphorylation in similar preparations of other molluscan neurons (Loechner et al., 1992; Yakel, 1992; Golowash et al., 1995). Therefore, we believe that serine/threonine phosphorylation does not play any role in the inactivation of Ca\(^{2+}\) channels in Lymnaea neurons; however, our experiments leave open the possibility that tyrosine phosphorylation may be involved.

Cytochalasin B greatly enhances inactivation of Ca\(^{2+}\) channels. Previous experiments done in our lab have shown that the rundown process of Ca\(^{2+}\) channels in giant inside-out patches is accelerated by the disruption of cytoskeleton (Johnson and Byerly, 1993a). To address the question of whether Ca\(^{2+}\)-channel inactivation in a whole-cell preparation is influenced by the cytoskeleton, we studied the effects of acute bath application of colchicine and cytochalasin B on Ca\(^{2+}\)-channel inactivation. We find that 100 μM colchicine, a microtubule disrupter, has no effect upon inactivation (peak inactivation, in cells containing 5 mM EGTA, is 0.26 ± 0.02 in the control solution and 0.27 ± 0.02 upon addition of 100 μM colchicine, n = 5). However, cytochalasin B, a disrupter of actin microfilaments, causes a large increase in inactivation of Ca\(^{2+}\) channels. This effect of cytochalasin B (which was applied extracellularly, being membrane permeant) is extremely robust and reproducible (Fig. 7 A). Addition of the vehicle by itself produces no change in inactivation. Cytochalasin B causes a rapid and stepwise increase in Ca\(^{2+}\) channel inactivation (Fig. 7 B, ■). The effect of cytochalasin B is selective for inactivation, since it has very little effect on the magnitude of Ca\(^{2+}\) current (Fig. 7 B, □), and its rate of rundown. Furthermore, the recovery of Ca\(^{2+}\) channels from inactivation is also much slower in the presence of cytochalasin B, primarily because of the increased amplitude of the slow component of recovery (Fig. 7 C), which is similar to the effect of reducing intracellular EGTA concentration. Surprisingly, the effect of cytochalasin B is readily reversible; the Ca\(^{2+}\)-channel inactivation returns to its pre–cytochalasin B levels after a few minutes of perfusion with the control external saline.

To quantify the effect of cytochalasin B on inactivation, we calculated an f parameter, the fractional increase in inactivation. The f parameter is defined as:

\[
f = \frac{\Delta I}{\Delta I_{\text{max}}},
\]

where \(\Delta I\) is the change in inactivation upon the addition of cytochalasin B, and \(\Delta I_{\text{max}}\) is the maximal possible change in inactivation. We chose f, calculated for a conditioning pulse to +40 mV, \(f_{40\text{mV}}\), to measure the effect of different concentrations of cytochalasin B upon inactivation in cells with 5 mM EGTA, and obtain the dose–response curve. The concentration of cytochalasin B that causes half the maximal effect on inactivation is ~100 μM. The effect of cytochalasin B in increasing
inactivation in cells with 5 mM EGTA was compared with that for cells with 0.1 mM EGTA. Surprisingly, $I_{\text{Ca}^{2+}}$ at $+40 \text{ mV}$, which is measured at $+40 \text{ mV}$, where Ca$^{2+}$ influx is maximal, is not significantly different in the two cases (Fig. 8), suggesting that the effect of cytochalasin B in increasing inactivation is independent of the Ca$^{2+}$-dependent inactivation.

While cytochalasin B is known to disrupt the actin cytoskeleton (Cooper, 1987), we could not confirm a role of actin filaments in Ca$^{2+}$-channel inactivation using other cytoskeletal agents. Cytochalasin D, another actin filament disrupter, has no significant effect on Ca$^{2+}$-channel inactivation in Lymnaea neurons (Fig. 8). Also, internally applied phalloidin, a stabilizer of microfilaments, does not decrease inactivation in cells with 0.1 mM EGTA (data not shown), and is unable to block the effect of extracellularly applied cytochalasin B in cells with 5 mM EGTA (Fig. 8). Moreover, cytochalasin B (50 μM), applied intracellularly, does not increase inactivation (data not shown). It is possible that phalloidin and cytochalasin B, added to the cell through the pipette solution, are unable to reach the cortical cytoskeleton, which alone may be involved in regulating inactivation. However, these results are also consistent with an extracellular effect of cytochalasin B on Ca$^{2+}$-channel inactivation, and suggest that the effect of cytochalasin B on Ca$^{2+}$-channel inactivation may be mediated by mechanisms independent of actin microfilaments.

**DISCUSSION**

In this study, we have characterized the Ca$^{2+}$ and voltage-dependent inactivation of Ca$^{2+}$ channels in Lymnaea neurons. Many of the earliest studies describing Ca$^{2+}$-dependent inactivation of Ca$^{2+}$ channels were done in molluscan neurons (Tillotson, 1979; Eckert and Tillotson, 1981). However, these early studies were done before outward H$^{+}$ currents had been identified in snail neurons (Thomas and Meech, 1982; Byerly et al., 1984a), and this may have resulted in erroneous measurements of Ca$^{2+}$-channel inactivation. Outward H$^{+}$ currents cannot be pharmacologically isolated from inward Ca$^{2+}$ currents (Byerly and Suen, 1989) and can give the impression of enhanced inactivation of inward currents. We have shown earlier that the three-pulse protocol used to measure inactivation in this study avoids errors due to H$^{+}$ currents (see materials and methods). For some experiments, we have used tail currents to measure inactivation. This method is also independent of H$^{+}$ currents because the H$^{+}$ tail currents deactivate faster than the Ca$^{2+}$ tail currents at negative potentials (Byerly et al., 1984a), and therefore do not contribute significantly to the measurement of peak Ca$^{2+}$ tail current (Gera and Byerly, 1999).

Much of the recent work in Ca$^{2+}$-channel inactivation has focussed on the study of recombinant channels expressed in heterologous systems (Neely et al., 1994; de Leon et al., 1995; Adams and Tanabe, 1997; Zhou et al., 1997; Peterson et al., 1999), and as such has
been successful in avoiding the problems of overlapping currents. However, Ca$^{2+}$-channel inactivation, especially the Ca$^{2+}$-dependent component, is greatly influenced by the intracellular environment of the channel (Isom et al., 1994; Johnson and Byerly, 1994; Schuhmann et al., 1997), which may be very different in a heterologous system compared with the native cell. Hence, study of Ca$^{2+}$ channels in their native environments provides a necessary complement to the molecularly defined study of these channels in heterologous systems.

**Bell-shaped Inactivation Curve in the Absence of Ca$^{2+}$-dependent Inactivation**

Ca$^{2+}$-channel inactivation in Lymnaea neurons perfused with 5 mM EGTA solution is not affected by replacing external Ca$^{2+}$ by Ba$^{2+}$ (Fig. 2 A), reducing the Ca$^{2+}$ influx (Fig. 2 B and D), or by increasing the level of intracellular Ca$^{2+}$ buffering (Fig. 2 C). We conclude from this that Ca$^{2+}$ channels in Lymnaea neurons containing 5 mM EGTA exhibit only voltage-dependent inactivation, even though the inactivation curve in 5 mM EGTA is bell shaped. A bell-shaped inactivation curve has often been taken as evidence for the presence of Ca$^{2+}$-dependent inactivation, though previous studies of native Ca$^{2+}$ channels in bullfrog sympathetic neurons (Jones and Marks, 1989) and of recombinant Ca$^{2+}$ channels in HEK 293 cells (Patil et al., 1998) have also shown that purely voltage-dependent mechanisms can yield bell-shaped inactivation curves. Our results extend those of the earlier studies, for we show that even in Ca$^{2+}$ channels capable of Ca$^{2+}$-dependent inactivation, a bell-shaped inactivation curve is not always correlated with the presence of Ca$^{2+}$-dependent inactivation.

Jones and Marks (1989) have proposed a model where the inactivated state of the channel is reached with highest probability from the open state, but the probability of this transition decreases with increasing voltage. Patil et al. (1998) have proposed a different model where the inactivated state is reached only from intermediate closed states. This model predicts that one long depolarizing step causes less inactivation than the cumulative inactivation caused by a number of shorter steps to the same voltage. However, experiments in Lymnaea neurons failed to conform to this prediction, but this model cannot be ruled out altogether. The model proposed by Correa and Bezanilla (1994) for Na$^{+}$ channel inactivation in squid giant axons predicts the existence of another open state that is preferentially reached from the inactivated state, and such a model also yields a bell-shaped inactivation curve. Another possibility is that the bell-shaped inactivation curve results from an inactivation process that increases monotonically with voltage combined with a “facilitation” process that increases at higher voltages. Such a facilitation of L-type current by very positive voltages has been reported, and is accompanied by a
slowing of deactivation kinetics after long, positive prepulses (Fleig and Penner, 1995; Slesinger and Lansman, 1996); this slowing of deactivation kinetics is also observed in Lymnaea channels (Gera and Byerly, 1999). However, we believe that the process that underlies the slowing of deactivation kinetics in Lymnaea neurons is of a different nature than the one that leads to the bell-shaped inactivation curve. This is because the change in deactivation kinetics lasts only a very short time after the conditioning pulse, and the 20-ms gap before the second test pulse eliminates it. However, the inactivation curve measured is consistently bell shaped. It is possible, though, that some other facilitatory process not involving a change in deactivation kinetics may be involved. While a number of models are adequate for explaining the bell-shaped inactivation curves that we obtain, we do not have any evidence that strongly favors one model over another.

Kinetics of Ca\(^{2+}\)-channel Inactivation

We show that increased intracellular Ca\(^{2+}\) concentration increases the time that Ca\(^{2+}\) channels require to recover from inactivation. The effect of Ca\(^{2+}\) in inhibiting the recovery of channels from inactivation at negative potentials has been observed before (Yatani et al., 1983; Gutnick et al., 1989; Branchaw et al., 1997); however, our results differ from those of the previous studies in that we do not observe a change in the time constants of inactivation due to Ca\(^{2+}\) influx, but only a change in the relative amplitudes of the fast and slow components.

It is more difficult to interpret our results pertaining to the development of inactivation (shown in Fig. 5 B) since the measurements of inactivation are dependent on the rate of recovery. To explain these observations, we have developed a simple model of the inactivation kinetics of Ca\(^{2+}\) channels at different voltages that fits the data shown in Fig. 5, A and B (the model is shown in Fig. 9, with rate constants at three different voltages given in Table I). In this model, there are two inactivated states, \(I_{FR}\) and \(I_{SR}\), that can be reached from the noninactivated states (which have been lumped together as N-I). At negative potentials, recovery from \(I_{FR}\) is considerably faster than that from \(I_{SR}\). Ca\(^{2+}\) influx during a conditioning pulse increases the occupancy of the inactivated state \(I_{SR}\), and thus increases the amplitude of the slow component of recovery at negative potentials. Our data can be fit by making only the forward rate constant \(\alpha_{SR}\) Ca\(^{2+}\) dependent, but we cannot rule out the possibility that the reverse rate constant \(\beta_{SR}\) may also be dependent on Ca\(^{2+}\) influx. After repolarization, \(I_{FR}\) is depleted rapidly, while \(I_{SR}\) is not greatly affected within the first 20 ms. Hence, the difference in inactivation that we measure between 0.1 and 5 mM EGTA with the three-pulse protocol is largely the difference in rate of recovery of channels from inactivation.

Table I

<table>
<thead>
<tr>
<th>Voltage</th>
<th>(-60) mV</th>
<th>(+60) mV</th>
<th>(+120) mV</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\alpha_{SR})</td>
<td>0</td>
<td>0.006</td>
<td>0.0035</td>
</tr>
<tr>
<td>(\beta_{SR})</td>
<td>0.06</td>
<td>0.001</td>
<td>0.001</td>
</tr>
<tr>
<td>(\alpha_{SR}) (in 0.1 mM EGTA)</td>
<td>0</td>
<td>0.0005</td>
<td>0.0009</td>
</tr>
<tr>
<td>(\beta_{SR})</td>
<td>0.0015</td>
<td>0.0001</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

All rate constants are expressed in units of ms\(^{-1}\). \(\alpha_{SR}\) is Ca\(^{2+}\) sensitive; its values in 0.1 mM EGTA are given in parenthesis.
ence in the $I_{\text{SR}}$ components under the two conditions. Also, the model predicts that the actual difference in inactivation between 0.1 and 5 mM EGTA (Fig. 9 B, continuous curves) during the course of a conditioning pulse is smaller than the difference we measure at the end of the 20-ms gap (Fig. 9 B, points and dashed curves). This is so because the proportion of channels in $I_{\text{FR}}$ is larger in 5 than in 0.1 mM EGTA; however, the difference measured after a 20-ms gap reflects primarily the difference in the $I_{\text{SR}}$ components in the two cases.

Since inactivation in 5 mM EGTA is completely voltage dependent, these results imply that $\alpha_{\text{SR}}$ is not zero at positive voltages even in 5 mM EGTA, and a conditioning pulse to +60 mV causes some occupancy of $I_{\text{SR}}$.

In this analysis, we have assumed that there is only one class of Ca$^{2+}$ channels in Lymnaea neurons. Pharmacological and kinetic studies done in our lab have failed to resolve more than one component of Ca$^{2+}$ current, though multiple types of Ca$^{2+}$ channels cannot be ruled out. In such a case, it is possible that Ca$^{2+}$- and voltage-dependent inactivation are due to different channel types that gate independently of each other? The results from our kinetic analyses (Fig. 5 A) show that the magnitude of voltage-dependent inactivation is reduced in the presence of Ca$^{2+}$-dependent inactivation (since the fast component of recovery is smaller in 0.1 than in 5 mM EGTA), which indicates that these phenomena are not independent of each other. It is, therefore, unlikely that different channel types underlie voltage- and Ca$^{2+}$-dependent inactivation.

Saturation of Ca$^{2+}$-dependent Inactivation

Our results with the Ca$^{2+}$ channel blocker Co$^{2+}$ indicate that just half of the Ca$^{2+}$ influx under standard conditions may be adequate to almost saturate the Ca$^{2+}$-dependent component of inactivation (Fig. 6 A). To illustrate the relation between Ca$^{2+}$-dependent inactivation and Ca$^{2+}$ influx, we plotted Ca$^{2+}$-dependent inactivation against Ca$^{2+}$ influx for a typical cell in Fig. 10. Fig. 10 (■, connected by a continuous line) shows the relation between these two quantities for each of the conditioning pulse potentials with 10 mM external Ca$^{2+}$. The general shape of these points shows that the Ca$^{2+}$-dependent inactivation is not linearly related to the Ca$^{2+}$ influx, and that Ca$^{2+}$ influx during pulses from +30 to +80 mV (with 10 mM external Ca$^{2+}$) causes considerable saturation of the Ca$^{2+}$-dependent component of inactivation. The shape of this curve is in good agreement with the 20% reduction in peak Ca$^{2+}$-dependent inactivation that is caused by a 50% reduction in Ca$^{2+}$ influx with 1 mM Co$^{2+}$ (Fig. 10, ◼). The result that a fourfold reduction in Ca$^{2+}$ influx, caused by reducing extracellular Ca$^{2+}$ from 10 to 1 mM, only blocks half of the peak Ca$^{2+}$-dependent inactivation also fits this same relationship (Fig. 10, △).

Mechanisms of Ca$^{2+}$-dependent Inactivation

Several researchers have proposed that Ca$^{2+}$ may cause inactivation by binding to a site on the channel itself (Sherman et al., 1990; Neely et al., 1994), or to an associated protein in the channel complex (Peterson et al., 1999; Qin et al., 1999; Zühlke et al., 1999). However, we do not think that is the case in Lymnaea Ca$^{2+}$ channels, mainly because of two observations. The first is that 5 mM EGTA is able to suppress Ca$^{2+}$-dependent inactivation completely. Since EGTA is a slow Ca$^{2+}$ buffer, 5 mM EGTA is fairly ineffective in reducing Ca$^{2+}$ concentration at the mouth of the channel and makes a significant contribution only at some distance from the channel. A mobile Ca$^{2+}$ buffer attenuates the intracellular Ca$^{2+}$ transient exponentially with distance from the channel, the space constant being ~60 nm for 5 mM EGTA (Neher, 1986). At distances smaller than the space constant, the buffer has little effect on the Ca$^{2+}$ transient. Hence, we think the Ca$^{2+}$-sensitive site mediating Ca$^{2+}$-dependent inactivation of Lymnaea Ca$^{2+}$
channels must be at least 100 nm away from the channel itself. Secondly, if there were a Ca\(^{2+}\)-binding site on the Ca\(^{2+}\)-channel complex that causes the inactivation of the channel, then this site would have a low affinity for Ca\(^{2+}\) (in the range of a few hundred micromolar, which is the concentration of Ca\(^{2+}\) at the inner mouth of an open channel). However, Ca\(^{2+}\) channels in Lymnaea neurons are sensitive to much lower levels of intracellular Ca\(^{2+}\). Perfusing these neurons with intracellular solutions buffered to 2 \(\mu M\) Ca\(^{2+}\) causes a rapid loss of Ca\(^{2+}\) current as soon as the intracellular saline diffuses within the cells (Byerly and Moody, 1984). Also, a sudden increase in the Ca\(^{2+}\) levels to a few hundred nanomolar by flash photolysis of Ca\(^{2+}\)-loaded DM-nitropheno causes a reversible decrease in Ca\(^{2+}\) current (Johnson and Byerly, 1993b). We interpret these results to mean that the site mediating Ca\(^{2+}\)-channel inactivation is sensitive to lower levels of intracellular Ca\(^{2+}\) and, therefore, cannot be on the channel. Here we are assuming that the inactivation of Ca\(^{2+}\) channels (as measured by 150-ms-long pulses), rundown (which develops with a much slower time course of minutes in whole-cell preparations and is largely irreversible) and block of Ca\(^{2+}\) current by intracellular Ca\(^{2+}\) (which has a time course of a few milliseconds) are all caused by similar mechanisms and, hence, have similar sensitivities to intracellular Ca\(^{2+}\). It is possible, however, that the mechanism of inactivation is independent of rundown and Ca\(^{2+}\) block; in which case, we cannot use the above results to estimate the Ca\(^{2+}\) sensitivity of the site mediating Ca\(^{2+}\)-dependent inactivation.

Studies on vertebrate L-type Ca\(^{2+}\) channels have led several researchers to conclude that Ca\(^{2+}\)-dependent inactivation in these channels is caused by Ca\(^{2+}\)-ion binding directly to the channel or to a site closely associated with it (Shirokov et al., 1993; Imredy and Yue, 1994; de Leon et al., 1995; Peterson et al., 1999; Qin et al., 1999; Zühlke et al., 1999). Giannattasio et al. (1991) showed that L-type Ca\(^{2+}\) channels in a smooth muscle-derived cell line have Ca\(^{2+}\)-dependent inactivation even in the presence of 10 mM intracellular BAPTA, suggesting that the Ca\(^{2+}\)-binding site is very close to the Ca\(^{2+}\) pore. This view has been supported by the identification of a putative Ca\(^{2+}\) binding domain (an E-F hand) on the COOH terminal of the mammalian \(\alpha_{1c}\) subunit of the Ca\(^{2+}\) channel that controls the Ca\(^{2+}\)-dependent inactivation of these channels (de Leon et al., 1995), although some studies suggest that the binding of Ca\(^{2+}\) ions to this site is unimportant for Ca\(^{2+}\)-dependent inactivation (Zhou et al., 1997; Bernatchez et al., 1998). Recent studies on Ca\(^{2+}\) channels containing the \(\alpha_{1c}\) subunit suggest that the channel is stably complexed with calmodulin, and it is the binding of Ca\(^{2+}\) ions to calmodulin that causes inactivation of the Ca\(^{2+}\) channel (Peterson et al., 1999; Qin et al., 1999; Zühlke et al., 1999). Considering that 5 mM intracellular EGTA completely eliminates Ca\(^{2+}\)-dependent inactivation in Lymnaea neurons, it is likely that vertebrate L-type Ca\(^{2+}\) channels have a different mechanism for Ca\(^{2+}\)-dependent inactivation than do the Ca\(^{2+}\) channels in Lymnaea neurons.

Effect of Cytochalasin B on Ca\(^{2+}\)-channel Inactivation

Several studies have shown that voltage-gated Ca\(^{2+}\) channels are sensitive to the state of the cytoskeleton (Fukuda et al., 1981; Johnson and Byerly, 1993a; Johnson and Byerly, 1994; Furukawa et al., 1997). However, the role of the cytoskeleton in modulating Ca\(^{2+}\) current is less clear. One study in chick ventricular myocytes shows that colchicine (a microtubule disrupter) and taxol (a microtubule stabilizer) strongly influence the inactivation kinetics of L-type Ca\(^{2+}\) channels (Galli and DeFelice, 1994). This is in contrast with the present study, where we find that the inactivation of Lymnaea channels is not affected by 100 \(\mu M\) colchicine.

In this study, we have shown that cytochalasin B, an actin microfilament disrupter, increases the inactivation of Ca\(^{2+}\) channels; however, there is some suggestion in our results that actin filaments may not be involved in this effect of cytochalasin B on inactivation. First, the concentration of cytochalasin B that results in half-maximal increase in inactivation is \(\sim 100 \mu M\), which is very high for an effect on actin filaments. Second, cytochalasin D does not yield the same effect as cytochalasin B (Fig. 8). Also, cytochalasin B applied intracellularly does not increase the amount of inactivation measured, and phalloidin applied intracellularly cannot block the effect of extracellularly applied cytochalasin B (Fig. 8). (Although, in the last two instances, it is questionable whether these drugs can effectively diffuse through the cytoplasm to reach the cytoskeleton close to the membrane.) Additionally, the time course of the onset of the cytochalasin B effect is very fast, limited only by the delay in application, and the effect is readily and completely reversible, inactivation returning to its original levels within minutes of perfusion with the control solution. This suggests that cytochalasin B may have only an extracellular effect.

The mechanism by which cytochalasin B increases inactivation currently remains unresolved. Cytochalasin B is known to inhibit the glucose transporter (Cooper, 1987); however, glucose is not present in the extracellular medium at the time of the experiments. Thus, the effect of cytochalasin B on the glucose transporter is unlikely to explain the increase in Ca\(^{2+}\) channel inactivation. Our results are consistent with the model where cytochalasin B acts as a weak open channel blocker, entering the pore from the external side. The slow time constant of recovery from inactivation that we observe.
in cytochalasin B may be due to the slow rate of dissociation of cytochalasin B from the pore. More experiments are required to distinguish whether actin filaments play a role in inactivation, or if cytochalasin B acts like an open channel blocker.

These studies leave unknown the mechanisms of Ca\(^{2+}\) channel inactivation in Lymnaea neurons. While we conclude that serine/threonine phosphorylation does not play any role in Ca\(^{2+}\) channel inactivation, it is possible that tyrosine phosphorylation may be involved. Indeed, tyrosine phosphorylation has been shown to modulate excision-activated Ca\(^{2+}\) channels in Lymnaea (Pafford et al., 1995). We cannot rule out a role of the actin cytoskeleton in Ca\(^{2+}\)-dependent inactivation, although the large effect of cytochalasin B on inactivation is unlikely to involve the cytoskeleton. Recent studies suggest that calmodulin may cause Ca\(^{2+}\)-dependent inactivation of mammalian L- and P/Q-type channels by interacting directly with the cytoplasmic domains of the α\(_{1C}\) subunits in a Ca\(^{2+}\)-dependent manner (Zühlke and Reuter, 1998; Lee et al., 1999; Peterson et al., 1999; Qin et al., 1999; Zühlke et al., 1999). This remains a possibility for Lymnaea Ca\(^{2+}\) channels also, although our results suggest that the Ca\(^{2+}\) receptor in Lymnaea neurons is not stably associated with the Ca\(^{2+}\) channel (as suggested by these studies for the L-type channels). Other intracellular proteins, such as G-proteins or phospholipid kinases, could also be involved in Ca\(^{2+}\)-dependent inactivation of Ca\(^{2+}\) channels.

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