Anomalous L-Type Calcium Channels of Rat Spinal Motoneurons

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Abstract

Single channel patch-clamp recordings show that embryonic rat spinal motoneurons express anomalous L-type calcium channels, which reopen upon repolarization to resting potentials, displaying both short and long reopenings. The probability of reopening increases with increasing voltage of the preceding depolarization without any apparent correlation with inactivation during the depolarization. The probability of long with respect to short reopenings increases with increasing length of the depolarization, with little change in the total number of reopenings and in their delay. With less negative repolarization voltages, the delay increases, while the mean duration of both short and long reopenings decreases, remaining longer than that of the openings during the preceding depolarization. Open times decrease with increasing voltage in the range −60 to +40 mV. Closed times tend to increase at V > 20 mV. The open probability is low at all voltages and has an anomalous bell-shaped voltage dependence. We provide evidence that short and long reopenings of anomalous L-type channels correspond to two gating modes, whose relative probability depends on voltage. Positive voltages favor both the transition from a short-opening to a long-opening mode and the occupancy of a closed state outside the activation pathway within each mode from which the channel reopens upon repolarization. The voltage dependence of the probability of reopenings reflects the voltage dependence of the occupancy of these closed states, while the relative probability of long with respect to short reopenings reflects the voltage dependence of the equilibrium between modes. The anomalous gating persists after patch excision, and therefore our data rule out voltage-dependent block by diffusible ions as the basis for the anomalous gating and imply that a diffusible cytosolic factor is not necessary for voltage-dependent potentiation of anomalous L-type channels.

Key words: Ca\(^{2+}\) channel • dihydropyridine • gating • voltage-dependent potentiation

Introduction

Among the different types of neuronal voltage-gated calcium channels, L-type channels play a specific role in regulating activity-dependent gene expression (Murphy et al., 1991; Bading et al., 1993; Deisseroth et al., 1998), neuronal survival and differentiation (Collins et al., 1991; Ghosh et al., 1994; Galli et al., 1995; Finkbeiner and Greenberg, 1996; Shitaka et al., 1996; Borsenitsch et al., 1998; Kirsch and Betz, 1998), and some forms of synaptic plasticity (Grover and Teyler, 1990; Anksztejn and Ben-Ari, 1991; Johnston et al., 1992; Kullmann et al., 1992; Bolshakov and Siegelbaum, 1994).

Multiple functionally and structurally different neuronal L-type channels have been described (Snutch et al., 1991; Williams et al., 1992; Forti and Pietrobon, 1993; Kavalali and Plummer, 1994, 1996; Ferroni et al., 1996; Cloues et al., 1997). The probability of reopening of these L-type channels is voltage dependent, increasing with increasing voltage of the previous depolarization.

Different authors have proposed and supported different mechanisms accounting for voltage-dependent reopenings of L-type channels. Forti and Pietrobon (1993) have proposed that long reopenings of anomalous L-type channels are a manifestation of a voltage-dependent equilibrium between gating modes, whereby increasing voltage drives the channel from a short- to a long-opening mode (Pietrobon and Hess, 1990), and also increases the occupancy of a closed state outside the activation pathway, which is connected to the open state through a voltage-dependent transition within each mode. This additional closed state accounts for both the delay with which long openings occur upon repolarization of the membrane and the anomalous voltage dependence of the mean open/closed times and of the open probability, found by the same authors.
(Forti and Pietrobon, 1993). Alternatively, reopenings have been interpreted as a manifestation of recovery from voltage- and/or current-dependent inactivation (Slesinger and Lansman, 1991; Thibault et al., 1993), resulting from voltage-dependent block of the channel pore by a positively charged cytoplasmic particle (Slesinger and Lansman, 1996). Kavalali and Plummer (1994, 1996) have proposed that reopenings reflect a particular form of voltage-dependent potentiation (LVP) in which the conditioning depolarization essentially reduces the voltage necessary to activate the channel. The different interpretations might reflect real differences in the L-type channels under study or simply derive from the emphasis of different aspects of the functional properties of essentially similar anomalous L-type channels. The data presented in this paper favor the second hypothesis.

As in most other neurons, the high-voltage activated whole-cell calcium current of embryonic and neonatal motoneurons can be dissected into four (L-, N-, P-, and R-type) pharmacological components (Mynlieff and Beam, 1992; Umemiya and Berger, 1994; Hivert et al., 1995; Magnelli et al., 1998). It is not known whether motoneurons express anomalous L-type channels. In the only single channel characterization of calcium channels in motoneurons (Umemiya and Berger, 1995), a classical L-type channel has been described. Here we show that embryonic rat spinal motoneurons express L-type channels that reopen at negative repolarization voltages and display anomalous gating properties similar to those of anomalous L-type channels of cerebellar granule cells. We have investigated the mechanism giving rise to the anomalous gating in motoneurons. Our data are consistent with the model proposed by Forti and Pietrobon (1993). They are also consistent with reversible block of the open pore by a positively charged cytoplasmic particle, but exclude block by diffusible ions, and imply that a diffusible cytosolic factor is not necessary for voltage-dependent potentiation of anomalous L-type channels.

Materials and Methods

Cell Culture

Spinal motoneurons from embryonic day 15 (E15) Wistar rat embryos were grown in primary culture after purification by a two-step metrizamide-panning method according to the procedure of Camu et al. (1993). In brief, ventral spinal cords were dissociated after trypsin digestion, and centrifuged over 6.5% metrizamide (Serva) cushions to eliminate the four-plate cells, which are dense enough to sediment through the cushion. The large cells were further enriched by immunopanning on Petri dishes coated with the IgG-192 antibody specific for the p75 neurotrophin receptor, which is specifically expressed by motoneurons at this stage. The hybridoma was generously provided by Dr. C. E. Henderson (University Mediterrane, Marseille, France). Routinely, 90% of purified neurons express p75 immunoreactivity (Camu et al., 1993). The cells were plated on polyornithine- (Sigma Chemical Co.) and laminin- (GIBCO BRL) coated glass coverslips, and cultured in Dulbecco’s modified Eagle medium (GIBCO BRL) supplemented with 17 mM glucose, 0.87 µM insulin, 0.99 mM putrescine, 0.93 µM sodium selenite, 1.32 µM transferrin, 0.19 µM progesterone, 0.51 µM triiodothyronine, 0.45 µM tiroxine, 1.32 µM BSA (all purchased from Sigma Chemical Co.), 2 mM glutamine, 100 IU/ml penicillin, and 100 µg/ml streptomycin (all purchased from GIBCO BRL). 12 h after plating, the culture medium was supplemented with 25% muscle-conditioned medium, obtained from myotubes cultures of newborn rat. Experiments were performed on motoneurons kept in culture from 1 to 5 d.

Patch-Clamp Recordings and Data Analysis

Single channel patch-clamp recordings followed standard techniques (Hamill et al., 1981). Currents were recorded with a DAGAN 3900 patch-clamp amplifier, low-pass filtered at 1 kHz (–3 dB), eight-pole Bessel filter), sampled at 5 kHz and stored for later analysis on a PDP-11/73 computer. Experiments were performed at room temperature (21–25°C).

Linear leak and capacitative currents were digitally subtracted from all records used for analysis. Current amplitude histograms were obtained from the data directly, with bin width equal to our maximal resolution (323.6 points/pA). For display, each histogram was normalized to the value of the zero current peak. Open probability, P_o, was computed by measuring the average current in a given single channel current. 12 h after plating, the unitary single channel current. Open-channel current amplitudes were measured by manually fitting cursors to well-resolved channel openings. A channel opening or closure was detected when more than one sampling point crossed a discriminator line at 50% of the elementary current. Histograms of open and closed times were fitted with sums of decaying exponentials. The best fit was determined by maximum likelihood maximization (Colquhoun and Sigworth, 1983) and the best minimum number of exponential components was determined by the maximum likelihood ratio test (Rao, 1973). Log binning and fitting of the binned distributions were done as described by Mclarnan et al. (1987) and Sigworth and Sine (1987). Openings occurring with a delay of more than one sampling point after repolarization of the membrane at –80 or –60 mV were considered as reopenings. In the measurement of the fraction of traces with long openings, reopenings were detected using a discriminator line at 33% of the elementary current, to decrease the number of missed short reopenings. To calculate the fraction of traces with long and short reopenings, we used a discriminating open time value, calculated from the double exponential open time histogram as the open time that equalized the number of openings of the fast exponential component falsely assigned as long openings and the number of openings of the slow exponential component falsely assigned as short openings (Demo and Yellen, 1991). Reopenings of duration longer than this value were considered as long reopenings and those of shorter duration as short reopenings. The pipette solution contained (mM) 90 BaCl_2, 10 TEACl, 15 CsCl, 10 HEPEs, pH 7.4 with TEAOH. The bath solution was (mM) 140 K-gluconate, 5 EGTA, 35 i-glucose, 10 HEPEs, pH 7.4 with KOH. The high-potassium bath solution was used to zero the membrane potential outside the patch. The dihydroyypiridine agonist (S)-202-791 (gift from Dr. Hof, Sandoz Co., Basel, Switzerland) was added (1 µM) to the bath solution in most recordings. Liquid junction potential at the pipette tip was +12 mV (pipette positive), and this value should be subtracted to all voltages to obtain the correct values of membrane potentials in cell attached recordings (Neher, 1992).
**Results**

Figs. 1 and 2 show that embryonic rat spinal motoneurons express L-type channels which reopen after repolarization of the membrane and display anomalous gating properties similar to those of anomalous L-type channels of cerebellar granule cells (Forti and Pietrobon, 1993). The single channel current recordings in Figures 1 and 2 were obtained from cell-attached membrane patches of rat spinal motoneurons in primary culture, which contained only one channel. The membrane was held at $-80$ mV and depolarized to four different voltages for 724 ms, every 4 seconds, in the presence of the dihydropyridine (DHP) agonist ($(+)-(S)$-202-791 in the bath in Figure 1 and in its absence in Figure 2. The representative current traces and the normalized current amplitude histograms from all traces with activity display the main unusual voltage-dependent properties of anomalous L-type channels.

The first unusual property is represented by the reopenings occurring with some delay after repolarization of the membrane at $-80$ mV, a voltage well below the threshold for channel activation. Some of these reopenings are quite long, much longer than the openings of the same channel during the preceding depolarization (compare Fig. 1, +30- and +40-mV traces). The second unusual property is represented by the voltage dependence of the open probability and of the open and closed times. The open probability ($P_o$) does not increase with voltage in the usual sigmoidal manner, but reaches a maximum at $+20$ mV, and then decreases with increasing voltage, remaining low in the entire voltage range (Fig. 3 C). In three single-channel patches, the maximal $P_o$ at $+20$ mV in the presence of DHP agonist was $0.12 \pm 0.02$. In each patch, average open probabilities were obtained from the traces with activity, without including nulls, which were a minority at each voltage (0–7% at

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1Abbreviation used in this paper: DHP, dihydropyridine.

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**Figure 1.** Single channel activity of the anomalous L-type Ca$^{2+}$ channel of rat spinal motoneurons in the presence of DHP agonist. Cell-attached recordings with 90 mM Ba$^{2+}$ as charge carrier from a patch containing a single L-type channel with anomalous gating, in the presence of 1 $M$ $(+)-(S)$-202-791 in the bath. Representative current traces at $+10$, $+20$, $+30$, and $+40$ mV are shown together with the normalized current amplitude histograms from all traces with activity at each voltage. Depolarizations were 724-ms long and were delivered every 4 s from holding potentials of $-80$ mV. Records were sampled and filtered at 5 and 1 kHz, respectively. Cell B58D.
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In the absence of DHP agonist, the activity of anomalous L-type channels is characterized by brief, mostly unresolved and infrequent openings and by an extremely low open probability at all voltages: the maximal value of \(P_o\) was 0.024 in the single channel patch of Fig. 2. Kinetic analysis of the open and closed time histograms reveals that the low open probability and its anomalous voltage dependence are due to the anomalous voltage dependence of both open and closed time constants. As shown in Fig. 3, the time constants of the two exponential components best fitting open time histograms both decrease with increasing voltage, and the two larger time constants of the three exponential components best fitting closed time histograms decrease with voltage up to +20 mV, and then start to increase (see also Fig. 10 B). Interestingly, the contribution of the slow exponential component in the open time histograms increases with increasing voltage, with a symmetrical decrease of the fast component. Thanks to these anomalous gating properties, anomalous L-type channels in the presence of DHP agonist could be easily distinguished from the other L- and non-L-type channels of rat spinal motoneurons. We have found that rat spinal motoneurons coexpress, together with the anomalous L-type channels characterized in this study, two additional DHP-sensitive channels, one similar to cardiac L-type channels and the other inactivating quite rapidly (Hivert and Pietrobon, 1995, 1997; and our unpublished observations). They can be distinguished from L-type channels with anomalous gating on the basis of their larger unitary current and conductance (24 vs. 20 pS), their larger mean open time and open probability at \(V > +20\) mV (not decreasing with increasing voltage), and the complete absence of reopenings. Moreover, rat spinal motoneurons express several different DHP-insensitive calcium channels, including two channels sharing the same conductance of 20 pS but differing in inactivation and pharmacological properties (Hivert and Pietrobon, 1995; and our unpublished observations).
In this study, we have investigated the mechanism giving rise to the unusual voltage-dependent properties of anomalous L-type channels of rat spinal motoneurons. To discriminate between different mechanisms, it was essential to be able to study the activity of a single anomalous channel during both the depolarization and repolarization periods. Since anomalous L-type channels represent only a small fraction of the different types of calcium channels with similar conductance expressed in motoneurons and patches with only one anomalous L-type channel were very rare, and since, in addition, the open probability of single anomalous L-type channels in the absence of DHP agonist is extremely low (Figs. 2 and 3 C), a property that they share with the more abundant inactivating L-type channel of 24 pS, it was necessary to prolong the openings of L-type channels with a DHP agonist to be sure that only one anomalous L-type channel was present in the patch. The comparison between Figs. 1 and 2 shows that the peculiar voltage-dependent properties of anomalous L-type channels above described are essentially similar with or without agonist, as previously shown in cerebellar granule cells (Forti and Pietrobon, 1993).

It has been proposed that reopenings of anomalous L-type channels reflect recovery from voltage- and/or current-dependent inactivation (Slesinger and Lansman, 1991, 1996; Thibault et al., 1993). If this interpretation is correct, then, in an experiment in which the membrane is depolarized at increasingly positive voltages, one should find a correlation between the extent of inactivation of single anomalous L-type channels during the depolarization and the fraction of traces with reopenings upon repolarization. Fig. 4 shows that such a correlation is absent. The ensemble average currents from a patch containing a single anomalous L-type channel in Fig. 4 A shows a lack of inactivation during long depolarizations at positive voltages (+10 to +40 mV) elicited from quite negative holding potentials (−80 mV). In the same voltage range, the fraction of traces with reopenings at −80 mV of the same channel increased as shown in Fig. 4 B (from 0 to 62%). Similar results were obtained for single anomalous L-type channels in cerebellar granule cells (Forti and Pietrobon, unpublished observations). Thus, the previously reported absence of inactivation of cerebellar anomalous L-type channels during depolarizations effective in inducing reopenings (Forti and Pietrobon, 1993) cannot be ascribed to the relatively depolarized holding potentials, as recently suggested (Slesinger and Lansman, 1996), but appears as a general distinctive property of anomalous L-type channels.

As already pointed out, after long depolarizations, both short and long reopenings at −80 mV could be observed. The existence of two clearly different open states, one short and the other long lasting, is even more evident if one analyzes the reopenings at −60 mV after a 400-ms long depolarization to +40 mV (Fig. 5, left). The recordings in Fig. 5 were obtained from a cell-attached patch containing only one anomalous L-type channel. The open time histogram of reopenings at −60 mV required two exponential components with time constants of 1.8 and 29 ms for best fit according to the maximum likelihood criterion. Strikingly, both open time constants were larger (1.6 ± 0.1 and 26 ± 3 ms, n = 5) than those measured for the anomalous L-type channel during the predepolarization at +40 mV (0.54 ± 0.19 and 1.4 ± 0.2 ms, n = 3; see Fig. 3).

A common feature of the different single L-type channels described so far is their voltage-dependent modal gating, whereby increasing voltage progressively drives the channels from a short-opening mode of activity (mode 1), prevailing at low voltages, to a long-
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opening mode (mode 2) prevailing at high positive voltages (Pietrobon and Hess, 1990; Forti and Pietrobon, 1993; Kavalali and Plummer, 1996). Forti and Pietrobon (1993) proposed that the anomalous gating arises from the presence of a nonadsorbing closed state outside the activation pathway connected to the open state through a voltage-dependent transition within each mode, and used the simplified kinetic scheme of Fig. 6 to explain, at least qualitatively, the peculiar properties of cerebellar anomalous L-type channels. In this kinetic model, the individual open and closed states underlying the two modes within the two boxes are lumped together and connected by single voltage-dependent forward ($k_f$) and backward ($k_b$) rate constants. To account for the anomalous voltage dependence of both open and closed times (see Fig. 3), the open states within each mode (O, O*) are connected to a closed state outside the activation pathway ($C_b$, $C_b^*$), and the rate constants $a$, $a^*$ for entry into the closed states $C_b$ in mode 1 and $C_b^*$ in mode 2 are assumed to increase with voltage, while the rate constants $\beta$, $\beta^*$ for exit from these closed states are assumed to decrease with voltage. Although the kinetic scheme in Fig. 6 is likely an oversimplification, we will use it here as a useful conceptual framework for the analysis and discussion of our data on anomalous L-type channels of motoneurons.

A possible interpretation of the presence of both short and long reopenings at negative voltages is that they represent reopenings of the channel in either mode 1 (from $C_b$) or mode 2 (from $C_b^*$), respectively. If this interpretation is correct, then any intervention that changes the probability of finding the channel in mode 2 at the end of the depolarization should change the relative proportion of long with respect to short reopenings, owing to the change in the relative probability of finding the channel in $C_b^*$ with respect to $C_b$ when the membrane is repolarized. The probability of finding the channel in the long-opening mode at the end of the depolarization can be changed by either changing the length or the amplitude of the depolarization. One expects that if the depolarization is shortened the relative proportion of long with respect to short reopenings should decrease. Indeed, Fig. 5 shows that when the depolarization was shortened from 400 to 50 ms, most of the reopenings of the single anomalous channel in the patch became short. The long reopenings were too few to define the second slower component in the open time histogram of the reopenings. The histogram was best fitted by a single exponential with a time constant of 2 ms, quite similar to the time constant of the fast component best fitting the histogram of reopenings of the same channel after 400 ms.

In three single channel patches, long reopenings were on average 40 ± 1% of the total number of reopenings after a 400-ms long depolarization, and de-
creased to 7 ± 4% of the total number of reopenings when the depolarization was shortened to 50 ms (Fig. 7 A). In one single channel patch, after shortening the depolarization at +40 mV from 400 to 50 ms, the voltage was increased to +150 mV keeping the duration constant at 50 ms. The long reopenings decreased from 38 to 12% of the total number of reopenings when the moderate depolarization was shortened, and increased again to 36% of the total number of reopenings when the amplitude of the short depolarization was increased. The relative increase of long with respect to short reopenings with increasing length and amplitude of the previous depolarization is consistent with the interpretation that short and long reopenings are associated with two different gating modes of the channel (mode 1 and mode 2) and with the existence of a voltage-dependent equilibrium between the gating modes whereby the probability of the long-opening mode (mode 2) increases with increasing voltage.

Consistent with this interpretation is also the finding that the probability of (long + short) reopenings, which depends on the probability of finding the channel in either C_b or C*_b at the end of the depolarization, was not much affected by changing the length of the depolarization. In three single-channel patches, the probability of reopenings changed from 63 ± 7% after 400 ms at +40 mV to 50 ± 5% after 50 ms at the same voltage (Fig. 7 B), but the change was not statistically significant (P < 0.2). The model in Fig. 6 predicts that changing the length of the depolarization should produce mirror changes in the probabilities of long and short reopenings, with a consequent unchanged probability of (long + short) reopenings, if the intrinsic probabilities of C_b and C*_b within the two individual
modes were identical and as long as the rate constants of the transitions to $C_c$ and $C^*_b$ were fast with respect to the duration of the depolarization.

As a consequence of the relative increase of long with respect to short reopenings with increasing length of the depolarization, the peak ensemble average current at $-60 \text{ mV}$ was larger and decayed more slowly after the longer depolarization (Fig. 5). In three single channel patches, the average time constant of decay of the ensemble current at $-60 \text{ mV}$ changed from $48 \pm 1 \text{ ms}$ after the long depolarization to $9.9 \pm 5 \text{ ms}$ after the short depolarization. On average, the ratio of the peak average currents at $-60 \text{ mV}$ after short and long depolarizations was $0.49 \pm 0.07$. The ensemble averages at $-60 \text{ mV}$ after both long and short depolarizations showed a clear rising phase, which originates from the delay with which most reopenings occurred after the repolarization. The time constant of the rising phase after short depolarizations ($1.5 \pm 0.3 \text{ ms}$) was slightly smaller than that after long depolarizations ($2.6 \pm 0.5 \text{ ms}$), but the difference did not reach statistical significance ($P < 0.1$).

Fig. 8 shows directly that long moderate depolarizations are able to drive anomalous L-type channels from a short- into a long-opening mode. The unitary current recordings in Fig. 8 A were obtained from a patch containing a single anomalous L-type channel, in an experiment in which after 400 ms at $+30 \text{ mV}$ the membrane was repolarized at $-10 \text{ mV}$, a voltage just above the threshold for channel activation. Control depolarizations to $-10 \text{ mV}$ for 800 ms were alternated with the prepulse protocol. In the large majority of control depolarizations at $-10 \text{ mV}$ (in 98 of 102 traces with openings), the unitary activity was characterized by relatively short openings and long closings and a low open probability, as shown by the first five representative traces in Fig. 8 A (right). The last trace represents a small minority of depolarizations (4 of 102 active traces) in which the channel shifted to a different mode of activity, characterized by long-lasting bursts with longer openings and shorter closings and by a much larger open probability.

The representative traces and the ensemble average current of Fig. 8 A (left) show that a preceding depolarization to $+30 \text{ mV}$ for 400 ms increased the probability of observing the long opening mode, with a consequent potentiation of the average current at $-10 \text{ mV}$. The fraction of active traces with the long opening mode increased from 4% in control depolarizations to $35\%$ after 400 ms at $+30 \text{ mV}$. These fractions were calculated using a discriminating open probability value of 0.1 to separate the traces with the long opening mode from those with activity similar to that in control. Fig. 8, B–D, shows that, while the open time histogram of all traces at $-10 \text{ mV}$ after the prepulse required two exponential components with time constants of 1.7 and 9.1 ms for best fit, the open time histogram of the traces with $P_o < 0.1$ could be best fitted by a single exponential with a time constant of 1.6 ms, similar to that of the fast component in the overall histogram and to

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\text{C} & \quad \text{C^*} \\ \rightarrow & \quad \text{O^*} \\ \text{p} & \quad \text{C^*} \\
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the time constant obtained from the best fit of the open time histogram of control traces (excluding the four traces with the mode shift having $P_o > 0.1$).

The average current at $-10$ mV after the predepolarization shows a clear rising phase, and after reaching a maximum value, slowly decays towards the control value. On average, the time constant of decay was $177 \pm 28$ ms ($n = 4$). The decay of the potentiated current should mainly reflect the kinetics of return of the channel from the long-opening mode to the short-opening mode prevailing in control. The slower decay of the average current at $-10$ mV with respect to that at $-60$ mV after the depolarization ($46 \pm 2$ ms, $n = 6$, see Fig. 5) is consistent with the voltage dependence of $k_o$, whereby $k_o$ decreases with increasing voltage (Pietrobon and Hess, 1990; Forti and Pietrobon, 1993). The rising phase of the current at $-10$ mV is clearly slower than the rising phase of the current at $-60$ mV after a similar prepulse (see Fig. 5), indicating that the rate of reopening from the closed states accessed during the depolarization ($C_o$ and $C^*_o$) increases with more negative repolarization voltages. On average, the time constant of the rising phase increased from $2.6 \pm 0.3$ ms ($n = 6$) at $-60$ mV to $11 \pm 2$ ms ($n = 4$) at $-10$ mV. This finding is consistent with and supports the voltage dependence of the rate constants of exit from the closed states $C_o$ and $C^*_o$ in the model in Fig. 6, whereby $\beta$ and $\beta^*$ decrease with increasing voltage. Consistent with this voltage dependence is also the peculiar lengthening of the closed times with increasing voltage of the depolarization (see Figs. 3 B and 10 B).

The two open time constants, obtained from the biexponential open time histogram at $-10$ mV were both smaller than those obtained for the reopenings at $-60$ mV after the prepulse (see Fig. 5), indicating that the rate of reopening from the closed states accessed during the depolarization ($C_o$ and $C^*_o$) increases with more negative repolarization voltages. On average, the time constant of the rising phase increased from $2.6 \pm 0.3$ ms ($n = 6$) at $-60$ mV to $11 \pm 2$ ms ($n = 4$) at $-10$ mV. This finding is consistent with and supports the voltage dependence of the rate constants of exit from the closed states $C_o$ and $C^*_o$ in the model in Fig. 6, whereby $\beta$ and $\beta^*$ decrease with increasing voltage. Consistent with this voltage dependence is also the peculiar lengthening of the closed times with increasing voltage of the depolarization (see Figs. 3 B and 10 B).

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-60 mV after a similar depolarization (compare Figs. 8 and 5). On the other hand, the open time constants at both -10 and -60 mV were larger than those obtained from the histogram of open times during the preceding depolarization (Fig. 3). In the voltage range -60 to +40 mV, the open time constants decreased as shown in Fig. 9 A. This anomalous voltage dependence of the open times supports the existence, within each mode, of a closed state outside the activation pathway to which the open state is connected through a transition whose rate constant increases with increasing voltage. Accordingly, in Fig. 6, the open states within each mode (O, O*) are connected to a closed state outside the activation pathway (C_o, C_o*), and the rate constants α, α* for entry into these closed states increase with increasing voltage. Our interpretation that the fast and slow components in the open-time histograms reflect sojourns in the open states of modes 1 and 2, respectively, is further supported by the finding that, as already pointed out, the contribution of the slow exponential component in the open time histograms increases with increasing depolarization voltage, with a symmetrical decrease of the fast component (Fig. 3 A, right).

At voltages higher than +20 mV, the two open time constants become similar (Figs. 3 A and 9 A). At these positive voltages, the anomalous L-type channel in mode 2 opens only for brief times, thus explaining the absence of bursts of activity with long openings during depolarizations effective in inducing the change to the long-opening mode as seen on repolarization (Fig. 8 A). One predicts that the potentiation of the anomalous L-type current by positive depolarizations should decrease with increasing repolarization voltage and there should be no potentiation of the current at repolarization voltages higher than +20 mV. Fig. 9 B shows that, in a single channel patch, the unitary activity of an anomalous L-type channel at +20 mV after a predepolarization to +40 mV for 400 ms was hardly distinguishable from that in control depolarizations at +20 mV. As a result, the same depolarization that produced a ro-

![Figure 9](image-url)
bust potentiation of the average current at $-10$ mV (Fig. 8), hardly potentiated the current at $+20$ mV.

Although Forti and Pietrobon (1993) did not speculate on the nature of $C_b$ and $C^*_b$, these states might correspond to either particular conformations of the channel or to open-pore blocked states, as proposed by Slesinger and Lansman (1996). In the latter case, $\alpha$, $\beta$ and $\alpha^*$, $\beta^*$ would be the rate constants of voltage-dependent block and unblock of the channel in the short- and long-opening modes, respectively, and their voltage dependence would be consistent with block by a positively charged cytoplasmic particle (Slesinger and Lansman, 1996). However, the result shown in Fig. 10 excludes any diffusible ion as the blocking particle. After excision in the K-gluconate/EGTA solution without divalents, the anomalous gating can persist unaltered for 40 min. Indeed, the data shown in Figs. 4 and 8 were derived after excision of the patch. The voltage-dependent induction of the long-opening mode in the inside-out patch in Fig. 8 shows that a diffusible cytosolic factor is not necessary for voltage-dependent potentiation of anomalous L-type channels.

discussion

In this study, we have shown that embryonic rat spinal motoneurons express anomalous L-type calcium channels, which reopen upon repolarization to resting potentials, displaying both short and long reopenings. The probability of reopening increases with increasing voltage of the preceding depolarization without any apparent correlation with inactivation during the depolarization. The probability of long with respect to short reopenings increases with increasing length of the depolarization, with little change in the total number of reopenings and in their delay. With less negative depolarization voltages, the delay increases, while the mean duration of both short and long reopenings decreases, remaining longer than that of the openings during the preceding depolarization. Open times decrease with increasing voltage in the range $-60$ to $+40$ mV, while closed times tend to increase with voltage at $V > 20$ mV. The open probability during depolarization pulses is low at all voltages and has an anomalous bell-shaped voltage dependence.

We have provided evidence that the two open states, leading to short and long reopenings, correspond to two gating modes of the channel, whose relative probability depends on voltage. Since the sojourn of the channel in both open states decreases with increasing voltage, the two open states must be connected to a closed state outside the activation pathway with a voltage-dependent transition whose rate constant increases with increasing voltage. The anomalous voltage dependence of the closed times suggests that the rate constant of reopening from this closed state decreases with increasing voltage. This voltage dependence leads to reopening of the channel upon repolarization and predicts a faster rate of reopening at more negative depolarization voltages, as found. According to our data,

![Figure 10](image_url)

**Figure 10. The anomalous gating persists after patch excision.** Single channel recordings from a patch containing one anomalous L-type channel, in the presence of 1 $\mu$M (+)-(S)-202-791 in the bath. After 3 min of recording in the cell-attached configuration, the patch was excised in the inside-out configuration; i.e., with the internal side facing the K-gluconate/EGTA bath solution. Cell B48L. (A) Representative current traces at $+20$ mV before and after excision are shown together with a diary plot displaying the open probability of the channel in successive depolarizations as a function of time during the recording. Depolarizations were 724- or 800-ms long and were separated by 4 s at $-80$ mV. During the recording, the test pulse voltage changed as indicated above the thin horizontal lines. The time after excision is indicated by the thick horizontal line. (B) Voltage dependence of the open probability and of the time constants of the exponential components best fitting open ($\tau_{o1}$ and $\tau_{o2}$) and closed ($\tau_{c1}$, $\tau_{c2}$ and $\tau_{c3}$) time distributions after patch excision are shown.
positive voltages favor both the transition from a short-
opening gating mode (mode 1) to a long-opening mode (mode 2), and the occupancy of a closed state within each mode from which the channel reopens on repolarization, displaying short reopenings when it re-
opens from the closed state of mode 1 and long re-
openings when it reopens from the closed state of mode 2 (Fig. 6, $C_b$ and $C^{*}_b$) (Forti and Pietrobon, 1993). The voltage dependence of the probability of reopenings reflects the voltage dependence of the occupancy of the closed states from which the channel reopens, while the relative probability of long with respect to short reopenings reflects the voltage dependence of the equilibrium between modes.

The properties of the first latency distribution of re-
openings of anomalous L-type channels of mouse cere-
bellar granule cells, measured by Slesinger and Lans-
man (1996) as a function of repolarization voltage, are consistent with our conclusions. To explain their data, Slesinger and Lansman (1996) assumed the existence of a positively charged cytoplasmic blocking particle that may reversibly block the pore during the depolar-
ization and be released upon repolarization at negative membrane potentials. According to this interpretation, $C_b$ and $C^{*}_b$ would correspond to open-pore blocked states. Our finding that the anomalous gating persists after excision of the patch in divalent-free solution rules out block by a diffusible ion. It does not rule out block by a membrane-bound particle. The abrupt switch from the anomalous gating to the cardiac-type gating, observed by Forti and Pietrobon (1993) in one single channel patch, tends to exclude part of the chan-
el as the blocking particle. An alternative interpreta-
tion, consistent with all the available data, is that $C_b$ and $C^{*}_b$ represent conformational states of the channel and that voltage-dependent pore block is not involved in anomalous gating.

Our data, both in motoneurons and cerebellar gran-
ule cells, do not show any apparent inactivation of sin-
gle anomalous L-type channels during depolarizations effective in inducing reopenings, even though the ki-
etic scheme in Fig. 6 is clearly compatible with inactiva-
tion and actually might seem to be inconsistent with lack of inactivation. Simulations performed using the model in Fig. 6 show that, depending on the rate con-
stants of the transitions between the states within each mode, the model can generate both noninactivating and inactivating currents during depolarizations effective in inducing reopenings (not shown). The extent of inactivation depends crucially on the ratio between for-
ward and backward rate constants, and increases with increasing ratios above a certain value. Thus, the model predicts that macroscopic inactivation should become apparent at sufficiently high voltages. The fact that it was not apparent from our single channel ensemble av-
erages in the range from +10 to +40 mV may signify that these voltages were not sufficiently high. Alterna-
tively, a small extent of inactivation at high voltages might have been missed, due to the stochastic behavior in the records.

Voltage-dependent potentiation is an interesting property shared by the different L-type channels de-
scribed so far. This name refers to the ability of earlier depolarization to transiently increase macroscopic L-type current. In different cells, different voltage dependence, different time course, and different duration of volt-
age-dependent potentiation of L-type channels have been reported, reflecting different L-type channels and/or different modulatory mechanisms (Fenwick et al., 1982; Hoshi et al., 1984; Lee, 1987; Pietrobon and Hess, 1990; Artalejo et al., 1991, 1992; Forti and Pietro-
bon, 1993; Nakayama and Brading, 1993; Sculptoreanu et al., 1993a,b, 1995; Bourinet et al., 1994; Johnson et al., 1994; Kleppisch et al., 1994; Fleig and Penner, 1996; Kavalali and Plummer, 1996; Parri and Lansman, 1996; Cloues et al., 1997). The progressive shift towards a long-opening mode induced by increasing voltage can explain the voltage-dependent potentiation of L-type channels of cardiac (Pietrobon and Hess, 1990) and smooth muscle cells (Kleppisch et al., 1994), as well as that of cardiac-type neuronal L-type channels (Bourinet et al., 1994; Kavalali and Plummer, 1996, and our unpublished observations). We have shown here and in our previous work (Forti and Pietrobon, 1993) that a voltage-dependent change in gating mode is also at the basis of voltage-dependent potentiation of anomalous L-type channels.

The closed states giving rise to the anomalous gating confer to the voltage-dependent potentiation of anom-
alous L-type channels two specific properties. The tran-
sient increase of anomalous L-type current following a depolarization is delayed, and falls with increasing re-
polarization voltage to become almost zero at +20 mV, where the open probability in the two modes becomes similar. In similar experimental conditions, potentia-
tion of cardiac L-type channels (Pietrobon and Hess, 1990) and brain $a_{1C}$ (Bourinet et al., 1994) is still ob-
served at +20 mV, where the open probability of the two modes is still quite different. Anomalous L-type channels differ from cardiac-type channels also in the voltage range controlling the mode change, which is shifted to lower voltages for anomalous L-type channels (Pietrobon and Hess, 1990; Kavalali and Plummer, 1996). Furthermore, the potentiation lasts longer for anomalous L-type channels with respect to cardiac channels (see Figure 4 of Forti and Pietrobon, 1993).

An important specific property of anomalous L-type channels is that even very short or small depolariza-
tions, insufficient to significantly shift the channels to-
wards the long-opening mode, can induce short re-
openings and thus transiently increase, for a brief time, the current upon repolarization. By delaying channel openings at resting potential, where the driving force is larger, the closed/ blocked state from which anomalous L-type channels reopen provides a mechanism to maximize calcium influx after a transient membrane depolarization such as an action potential. With increasing duration and amplitude of the depolarization, the relative contribution of long reopenings, due to the mode shift, increases, leading to a potentiated and longer lasting transient increase of the current upon repolarization. The anomalous gating warrants maximal potentiation at resting potentials. One can predict that a presynaptic train of action potentials at high frequency may lead to a progressive increase of calcium influx through presynaptic anomalous L-type channels during the train, and/or may generate a large surge of calcium influx through postsynaptic anomalous L-type channels at the end of the train. Reopenings of L-type channels after single action potentials, with increasing open times at more negative repolarization voltages, and a progressive shift of the same channels to a long-opening gating mode during trains of action potentials have been observed in sensory neurons (Ferroni et al., 1996). Reopenings of calcium channels induced by single back-propagating action potentials have been observed also in dendrites of hippocampal cells (Magee and Johnston, 1995). Most likely, anomalous L-type channels were involved in both cases. Kavalali and Plummer (1994, 1996) and Kavalali et al. (1997) have characterized L-type channels in hippocampal neurons with biophysical properties quite similar to those of anomalous L-type channels of cerebellar granule cells and motoneurons. The similarities include single channel current and conductance, low open probability, mean open times at +20 mV shorter than at −30 mV, potentiation after modest depolarizations (LVP) and lack of potentiation at repolarization voltages higher than +20 mV, slower decay of the potentiated current than that of cardiac L-type channels, and persistent activity in excised patches.

Given their presence in cerebellar, hippocampal, sensory, and motor neurons, L-type channels with anomalous gating are probably widely expressed in the nervous system, while they are absent from cardiac and pituitary endocrine cells (our unpublished observations). The predicted capability of anomalous L-type channels to produce a delayed influx of calcium at resting potentials after previous neuronal electrical activity, whose extent depends on the duration and strength of such activity, makes them particularly suited to play a critical role in coupling transient neuronal activity with long-term changes in nervous system development and function. Indeed, several of these changes have been found to be specifically inhibited by dihydropyridine drugs (see Introduction). Calcium entry through anomalous L-type channels might play an important role also in neurotoxic pathogenesis, since these channels are probably still active under conditions of metabolic stress such as hypoxia, which would favor the voltage-dependent change in gating mode leading to potentiation of calcium influx. In addition, anomalous L-type channels may play a role in neuronal deterioration during aging (Thibault and Landfield, 1996).

The molecular basis for the anomalous L-type channel is unknown. Its absence in pituitary endocrine cells, which express the α1D subunit in large amounts (Fomina et al., 1996), and, on the other hand, its abundance in rat cerebellar granule cells in primary culture, where the α1D subunit is either not expressed or expressed in small amounts (Schramm et al., 1999), suggest that, most likely, the pore-forming subunit of anomalous L-type channels is not α1D. Since the α1C subunit was detected as a major transcript in rat cerebellar granule cells in primary culture using degenerated oligonucleotide primer pairs under highly stringent conditions, and no new α1-related sequences were amplified under the same conditions (Schramm et al., 1999), it appears likely that the anomalous L-type channel is related to the α1C subunit. It remains to be established whether the anomalous behavior arises from an unknown splice variant of the α1C subunit, from a particular subunit composition, or from an unknown type of modulation.

We thank Dr. C.E. Henderson for providing the IgG-192 hybridoma and Dr. M. Cantini for providing the muscle-conditioned medium.

The financial support of Telethon-Italy to D. Pietrobon (grant 720) and to B. Hivert is gratefully acknowledged. This work was also partially supported by a grant from the Regione del Veneto (Giunta Regionale Ricerca Sanitaria Finalizzata-Venezia-Italia).

Original version received 19 January 1999 and accepted version received 25 March 1999.

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