Effect of Postnatal Development on Calcium Currents and Slow Charge Movement in Mammalian Skeletal Muscle

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ABSTRACT Single- (whole-cell patch) and two-electrode voltage-clamp techniques were used to measure transient ($I_{\text{trans}}$) and sustained ($I_{\text{slow}}$) calcium currents, linear capacitance, and slow, voltage-dependent charge movements in freshly dissociated fibers of the flexor digitorum brevis (FDB) muscle of rats of various postnatal ages. Peak $I_{\text{trans}}$ was largest in FDB fibers of neonatal (1–5 d) rats, having a magnitude in 10 mM external Ca of $1.4 \pm 0.9 \ pA/pF$ (mean ± SD; current normalized by linear fiber capacitance). Peak $I_{\text{trans}}$ was smaller in FDB fibers of older animals, and by ~3 wk postnatal, it was so small as to be unmeasurable. By contrast, the magnitudes of $I_{\text{slow}}$ and charge movement increased substantially during postnatal development. Peak $I_{\text{slow}}$ was $3.6 \pm 2.5 \ pA/pF$ in FDB fibers of 1–5-d rats and increased to $16.4 \pm 6.5 \ pA/pF$ in 45–50-d-old rats; for these same two age groups, $Q_{\text{max}}$, the total mobile charge measurable as charge movement, was $6.0 \pm 1.7$ and $23.8 \pm 4.0 \ nC/\mu F$, respectively. As both $I_{\text{slow}}$ and charge movement are thought to arise in the transverse-tubular system, linear capacitance normalized by the area of fiber surface was determined as an indirect measure of the membrane area of the t-system relative to that of the fiber surface. This parameter increased from $1.5 \pm 0.2 \ \mu F/cm^2$ in 2-d fibers to $2.9 \pm 0.4 \ \mu F/cm^2$ in 44-d fibers. The increases in peak $I_{\text{trans}}$, $Q_{\text{max}}$, and normalized linear capacitance all had similar time courses. Although the function of $I_{\text{slow}}$ is unknown, the substantial postnatal increase in its magnitude suggests that it plays an important role in the physiology of skeletal muscle.

INTRODUCTION

An action potential causes a skeletal muscle fiber to contract, a process known as excitation-contraction (E-C) coupling. Two important structural elements for E-C coupling are infoldings of the surface membrane, termed transverse-tubules (t-tubules or t-system), and a wholly internal membranous compartment termed the sarcoplasmic reticulum (SR). During E-C coupling, depolarization of the t-system causes the SR to release calcium, which in turn causes contraction (Costantin, 1975; Endo, 1977). The events that link t-system depolarization to SR calcium release...
remain to be established. One event likely to be involved is measurable as an electrical signal termed charge movement. Charge movement, which has been measured in both amphibian (Schneider and Chandler, 1973; Chandler et al., 1976; Adrian and Almers, 1976) and mammalian (Hollingworth and Marshall, 1981; Dulhunty and Gage, 1983; Simon and Beam, 1983, 1985a; Lamb, 1986a) skeletal muscle, has been hypothesized to represent the voltage-induced change in the configuration of a “sensor” molecule embedded in the t-system membrane; this reconfiguration in turn is hypothesized to cause SR calcium release (Schneider and Chandler, 1973).

An alternative hypothesis for E-C coupling is that a small amount of calcium entering the fiber from the extracellular space triggers a large release of calcium ions from the SR (Endo et al., 1970; Ford and Podolsky, 1970). Consistent with this hypothesis, a slowly activated calcium current (Islow) has been measured in both amphibian (Beaty and Stefani, 1976; Stanfield, 1977; Sanchez and Stefani, 1978; Almers and Palade, 1981) and mammalian (Potreau and Raymond, 1980; Donaldson and Beam, 1983; Walsh et al., 1986) skeletal muscle fibers. However, the slow time course of this current compared with the action potential makes it seem unlikely that it is involved in E-C coupling (Sanchez and Stefani, 1978). Moreover, blocking the current pharmacologically does not abolish E-C coupling in adult frog muscle (Gonzalez-Serratos et al., 1982). Thus, the function of Islow is not known.

A possibility is that Islow is vestigial in adult muscle and important only to the function of developing muscle. Specifically, one might imagine that Islow is important for E-C coupling in developing muscle. E-C coupling might well be different since the diameter of neonatal mammalian skeletal muscle fibers is only about one-fifth that of adult fibers (Luff and Atwood, 1971), and neonatal fibers have very little t-system (Schiaffino and Margreth, 1969; Luff and Atwood, 1971). Calcium entering into small-diameter fibers with a sparse t-system might well have a function that is no longer important in the much larger adult fibers with their extensive t-system. With this hypothesis in mind, we examined calcium currents and charge movement in acutely dissociated fibers of the flexor digitorum brevis muscle of rats of various ages.

We have found two calcium currents in neonatal muscle, Ifast and Islow. Ifast disappears after the first few weeks postnatal. By contrast, both Islow and charge movement are present both neonatally and later in development. In fact, the magnitude of both increases more than fivefold during postnatal development. The postnatal increase in the magnitude of Islow is the opposite of what one might expect for a vestigial current, and suggests instead that Islow reflects a functionally important component of adult muscle.

A preliminary account of some of these results has appeared (Knudson et al., 1986).

METHODS

All experiments were carried out with fibers enzymatically dissociated (Beam and Knudson, 1988) from the flexor digitorum brevis (FDB), a fast-twitch (Carlsen et al., 1985) muscle of the rat. Currents were measured using the whole-cell configuration of the patch-clamp technique or, for the larger fibers from older animals, a two-electrode voltage clamp. For both the whole-cell patch clamp and the two-electrode clamp, pipettes were pulled from borosilicate
glass and filled with a solution that contained (millimolar): 140 Cs-aspartate, 5 Mg-aspartate, 5 MgCl₂, 10 Cs₄EGTA, 10 HEPES, pH 7.4 with CsOH. Pipette resistances ranged from 1.5 to 2.5 MΩ. The whole-cell patch clamp was implemented as described by Hamill et al. (1981; additional details are given in Beam and Knudson, 1988). For the two-electrode clamp, one pipette was used to measure potential and the other was used to pass current. The output of the current-passing pipette was controlled via a conventional, high-voltage-output, two-electrode voltage-clamp circuit. Each of the two pipettes was brought up to the surface of the fiber and gentle suction was applied until a high-resistance (>100 MΩ) seal was established; strong suction was then used to rupture the patch of membrane beneath the pipette. To monitor seal formation and patch rupture, the potential at the top of the pipette was measured in response to a square wave applied to the pipette via a 1-GΩ resistor. To decrease their capacitance, both pipettes were coated with Q-dope (G.C. Electronics, Rockford, IL). Separate agar-bridge electrodes were used for sensing the bath potential and for providing the bath ground for current flow. The bath solutions used were (millimolar): "50 Ca": 50 Ca²⁺, 85 TEA⁺, 185 Cl⁻; "10 Ca": 10 Ca²⁺, 145 TEA⁺, 165 Cl⁻ (or 20 Cl⁻ and 145 Br⁻). Both solutions were buffered with 10 HEPES, titrated to pH 7.4 with CsOH.

Nonlinear capacitative currents were measured with fibers bathed in 10 Ca solution; usually 10 μM TTX was also present, and in some cases 1 mM Cd was present as well. For measurements made with the whole-cell patch clamp, the feedback resistor in the headstage was 10 MΩ. For measurements made with the two-electrode clamp, the clamp current was determined as the voltage drop across a 1-MΩ resistor in series with the current-passing pipette. For both voltage-clamp methods, linear components of capacitative and leak currents were canceled as completely as possible with a three-time-constant analog leak-subtractor before additional gain was taken on the signal from the current-measuring amplifier. Remaining linear components of current were removed from test currents by digital scaling and subtraction of a control current elicited by a 20–30-mV depolarization or hyperpolarization from the holding potential (the same control current was used to calculate the linear fiber capacitance). Because the measured charge movements were similar for the depolarizing and hyperpolarizing control pulses, we concluded that little charge moves at membrane potentials less than or equal to −50 mV. To help prevent saturation or slew-rate limiting of amplifiers, voltage-clamp command pulses were exponentially rounded with a time constant of 50–300 μs. After subtraction of a sloping baseline as necessary (Horowicz and Schneider, 1981), Q₁₉ and Q₁₅ were calculated by integrating the ON and OFF charge movement transients.

The dependence of Q₁₉ and Q₁₅ on test potential (V) was quantified by fitting the data according to a two-state Boltzmann model for the distribution of charge:

$$Q = Q_{\text{max}}/[1 + \exp (-V/\tilde{V})/k],$$

where $Q_{\text{max}}$ represents the maximum charge that can be moved, $\tilde{V}$ is the potential at which half the charge has moved, and $k$ is a slope factor. The parameters $Q_{\text{max}}$, $\tilde{V}$, and $k$ were initially adjusted manually to give a visual best fit of the data. The computer was then used to calculate the sum of the squared deviations between the calculated curve and the data points as each of these three parameters was stepped through a range of values: ±8 mV in 1-mV steps for $\tilde{V}$ and ±40% in 5% steps for $Q_{\text{max}}$ and $k$. The combination of parameters that minimized the sum of the squared deviations was taken as the best fit.

The two-state Boltzmann expression was also used to quantify activation of the slow calcium current. Specifically, plots of conductance vs. voltage were fitted according to the expression

$$g(V) = g_{\text{max}}/[1 + \exp (-V/\tilde{V})/k],$$

where $g(V)$ is the slow calcium conductance at test potential $V$, $g_{\text{max}}$ represents the maximum conductance activated by strong test potentials, $\tilde{V}$ is the midpoint of the activation curve (i.e.,
the potential for activation of half-maximal conductance), and $k$ has the same meaning as in Eq. 1. The parameters $g_{\text{max}}, V$, and $k$ were adjusted to minimize the sum of the squared deviations between the data points and the curve calculated with Eq. 2. The conductance $g(V)$ was calculated from the measured slow calcium current with the expression

$$g(V) = \frac{I_{\text{slow}}}{V - V_{\text{rev}}}.$$

where $I_{\text{slow}}$ represents the peak inward slow calcium current measured at test potential $V$, and $V_{\text{rev}}$ was determined by linear extrapolation of the falling limb of the peak $I-V$ curve. (In cells with a large $I_{\text{fast}}$, care was taken to calculate peak $I_{\text{slow}}$ at a time when the decay of $I_{\text{fast}}$ was essentially complete.)

RESULTS

Postnatal Changes in Calcium Currents

Fig. 1 compares calcium currents recorded from FDB muscle fibers of a 2-d-old (A) and a 19-d-old (B) rat. In the fiber from the 2-d rat, two distinct calcium currents are present, a transient current ($I_{\text{fast}}$) selectively activated by test potentials from $-40$ to $-10$ mV, and an additional, maintained current ($I_{\text{slow}}$) activated by test depolarizations $\geq 0$ mV. The presence of these two calcium currents in primary cultures of embryonic rat and mouse muscle, and in FDB muscle fibers of neonatal rats, has been previously documented (Beam et al., 1986; Cognard et al., 1986a; Beam and Knudson, 1988). For the fiber in Fig. 1 A, the amplitudes of $I_{\text{fast}}$ and $I_{\text{slow}}$ were comparable. This was typical of muscle fibers from animals 1 or 2 d old. The calcium currents recorded from the fiber of the 19-d-old rat (Fig. 1 B) are different in two major respects from those of the 2-d rat fiber. First, the test potentials from $-40$ to $-10$ mV, which elicited $I_{\text{fast}}$ in the 2-d fiber, failed to elicit any appreciable time-dependent current in the 19-d fiber. Second, the $I_{\text{slow}}$ current that is evoked by test pulses to $\geq 0$ mV is $>15$-fold larger than $I_{\text{slow}}$ in the 2-d fiber. Although fiber size increases between 2 and 19 d ($\sim 1.4$-fold in diameter and 2.5-fold in length), the increase in the fiber surface area is much smaller than the increase in the magnitude of $I_{\text{slow}}$. The growth of a membrane system (i.e., the t-system) where slow calcium channels reside may be involved in the increase of $I_{\text{slow}}$ (see below).

To get a more complete picture of the effect of postnatal development on $I_{\text{fast}}$ and $I_{\text{slow}}$, calcium currents were measured in fibers acutely dissociated from FDB muscles of various-aged rats. For each fiber, the value at the peak of the largest $I_{\text{fast}}$ and of the largest $I_{\text{slow}}$ were recorded. These values were normalized by the linear capacitance of the fiber, since the capacitance provides an indirect measure of the fiber’s membrane area (see below). Fig. 2 illustrates data collected from rats aged 0–50 d. Peak $I_{\text{fast}}$ is large at birth and then undergoes a gradual decline and disappears entirely by 3–4 wk postnatal. By contrast, peak $I_{\text{slow}}$ is relatively small at birth but increases about fivefold during the first 4 wk postnatal, after which it remains at relatively constant level. The postnatal developmental increase in the magnitude of peak $I_{\text{slow}}$ suggests that the slow calcium channel protein plays an important role in skeletal muscle function.

Postnatal Changes in Charge Movement

As described in the Introduction, a nonlinear capacitative current, or “charge movement,” which is thought to be involved in E-C coupling, has been found to be present in adult skeletal muscle of several different species. Fig. 3, A and C, illustrates similar
Developmental change in muscle calcium currents. Calcium currents are illustrated for FDB muscle fibers from a 2-d (A) and a 19-d (B) rat. (A) Fiber A76; four points blanked; 10 Ca bath solution; holding potential (HP) = -90; C = 175 pF. (B) Fiber D48, three points blanked; 10 Ca solution + 10 μM TTX; HP = -80; C = 430 pF.

nonlinear capacitative currents measured in neonatal rat muscle fibers. These were measured in response to 10-ms depolarizations from the holding potential to the test potential indicated to the right of each trace. By integrating the ON transient that follows the onset of the test pulse, one obtains $Q_{\text{ON}}$, the amount of charge that moves upon depolarization. Similarly, $Q_{\text{OFF}}$, the charge that moves upon repolarization, is the integral of the OFF transient that follows the cessation of the test pulse. Plots of
$Q_{ON}$ and $Q_{OFF}$ vs. test potential are illustrated in Fig. 3, B and D. As in adult skeletal muscle, the absolute magnitudes of $Q_{ON}$ and $Q_{OFF}$ are nearly equal. Also, both $Q_{ON}$ and $Q_{OFF}$ initially increase in size as the test pulse is made larger and then saturate for sufficiently large depolarizations.

The charge movements in Fig. 3 A were measured in an FDB fiber of a 5-d rat, while the charge movements in Fig. 3 C were measured in an FDB fiber of a 21-d rat. The charge movements from the two different-aged rats have similar properties, with the exception that $Q_{max}$, the maximum amount of charge that moves for a strong depolarization, is ~20 times larger in the older rat. After normalization by linear fiber capacitance, $Q_{max}$ in the older animal is still more than three times larger. However, as discussed later, this may not represent an increase in $Q_{max}$ with respect to the membrane area of the t-system. The postnatal developmental increase in the normalized $Q_{max}$ is illustrated in more detail in Fig. 4. As for peak $I_{slow}$, $Q_{max}$ increased about fivefold during the first 4 wk postnatal, after which it remained at a fairly constant level. To permit a more direct comparison of the postnatal development of $Q_{max}$ and peak $I_{slow}$, the same smooth curve that was shown superimposed on peak $I_{slow}$ (Fig. 2 B) is also shown superimposed on the $Q_{max}$ data.
Although the increase in $Q_{\text{max}}$ represented the major postnatal change in muscle charge movements, there was also a small change in the charge movements' steady state voltage dependence. To characterize this voltage dependence, a least-squares procedure was used to fit Eq. 1 to plots of $Q_{\text{ON}}$ and $Q_{\text{OFF}}$ vs. test potential. This procedure yields values for $V$, the midpoint of the charge-voltage relationship, and $k$, a parameter that determines the steepness of this relationship. The values for $V$ were shifted positively in older rats. Specifically, the values (mean ± SD) of $V$ were $-9.3 ± 7.9$ (N = 6), $-0.6 ± 7.9$ (9), and $2.6 ± 4.2$ mV (5) for <15-, 15-25-, and >25-d FDB fibers, respectively. For the same three age groups, the values for $k$ showed little difference, being $11.2 ± 4.9$, $11.1 ± 3.5$, and $11.5 ± 2.6$ mV.

The midpoint of the activation curve for $I_{\text{slow}}$ also appeared to shift positively with postnatal development. Thus, fits of the Boltzmann expression to conductance vs. voltage plots (see Methods) yielded values for $V$ of $10.3 ± 7.5$ (24), $12.4 ± 6.3$ (30), and $16.1 ± 4.8$ mV (15), respectively, for <15-, 15-25-, and >25-d FDB fibers; the values of $k$ for these three age groups were $5.8 ± 1.7$, $4.9 ± 1.4$, and $7.1 ± 2.5$ mV, respectively.

**Postnatal Development of the T-System**

As detailed in the Discussion, both the slow calcium current and charge movement are thought to arise within the t-system. Morphometric measurements of mouse skeletal muscle have revealed that the percentage of fiber volume occupied by the t-system increases about fivefold during the first few weeks postnatal (Luff and Atwood, 1971). This increase implies a concomitant increase in the membrane area of the t-system relative to that of the fiber surface. Thus, the developmental increases in $Q_{\text{max}}$ and peak $I_{\text{slow}}$ may be related to the developmental increase in relative t-tubular area. As an indirect measure of the latter, we determined the linear capacitance ($C$) and the fiber surface area ($A_f$) in individual fibers from rats of various ages. $A_f$ was calculated, under the assumption of right-cylindrical geometry, from visual measurements of fiber length and diameter. Fig. 5 illustrates that $C/A_f$ increases as a function of age. This increase implies that with postnatal development, membrane infolding is increasing the total membrane area compared with that calculated from fiber dimensions. In skeletal muscle, the t-tubular system accounts for a large component of membrane infolding. Thus, the data of Fig. 5 suggest that there is a postnatal increase in relative t-tubular area. The time course of this increase is strikingly similar to that of the postnatal increases in $Q_{\text{max}}$ and $I_{\text{slow}}$, as can be seen by comparison of the data points with the smooth curve (the same one as in Figs. 2 B and 4). That all three increase postnatally suggests that each is important to the function of adult muscle. The similarity of time course may indicate that there is coordinate developmental regulation of slow calcium channels, charge movement, and the t-system.

**DISCUSSION**

During the first few weeks of postnatal development of the rat FDB muscle, the magnitude of the calcium current $I_{\text{slow}}$, the quantity of mobile charge measurable as voltage-dependent charge movement ($Q_{\text{max}}$), and the relative contribution of the t-system to linear fiber capacitance increase severalfold. A postnatal developmental increase is consistent with an important role for all three in the function of adult
FIGURE 3. Slow, voltage-dependent charge movement is present in developing muscle fibers. The upper panels show charge movements from a 5-d (A) and a 21-d (C) FDB fiber; the charge movements were elicited by a 10-ms depolarization from the holding potential to the indicated test potentials. The lower panels illustrate, for the same two fibers, the dependence of $Q_{ON}$ (filled triangles) and $Q_{OFF}$ (open triangles) on the test potential. The smooth curves in B and D are plots of text Eq. 1 with values for $Q_{max}$, $k$, and $V$ that were determined by a least-squares fitting procedure. (A and B) Fiber D29; 5-d FDB; whole-cell (single-electrode) patch clamp; 10 Ca solution with 1 mM CdCl$_2$ and 10 µM TTX; HP = -80; $C$ = 135 pF. The control current was a 20-mV depolarization from the holding potential. At each test potential, the illustrated trace is the average of four separate test currents. For this fiber, $Q_{max} = 3.5$ nC/µF, $k = 5.2$ mV, and $V = -24$ mV. (C and D) (opposite) Fiber D81; 21-d FDB; two-electrode voltage clamp; 10 Ca solution + 10 µM TTX; HP = -80; $C$ = 875 pF. The control pulse was a 30-mV depolarization; the traces illustrated are individual test currents without signal averaging. For this fiber, $Q_{max} = 11.3$ nC/µF, $k = 12.2$ mV, and $V = -6$ mV.
muscle cells. By contrast, the disappearance of the calcium current $I_{\text{fast}}$ during postnatal development is obviously inconsistent with a role for this current in adult muscle.

**Function of $I_{\text{fast}}$**

One possible function for $I_{\text{fast}}$, to couple membrane excitation to contraction in developing muscle, is not supported by the available data. Thus, blocking $I_{\text{fast}}$ (and also $I_{\text{slow}}$) does not prevent membrane depolarization from causing contraction of rat or mouse
Figure 4. Postnatal developmental increase in \( Q_{\text{max}} \). Each data point represents the average value obtained for fibers spanning a 5-d range of age. Charge movements were measured as described in the Methods. \( Q_{\text{max}} \) was determined as the area under the ON transient for a test pulse to +40 mV. For each fiber, the value of \( Q_{\text{max}} \) was normalized by the linear fiber capacitance. The filled square is the value determined by Simon and Beam (1985a) from three-microelectrode measurements on adult rat omohyoid muscle fibers. Error bars indicate \( \pm \) SEM. The smooth curve is the same one as in Fig. 2 B.

myotubes or neonatal rat FDB fibers (Knudson et al., 1986). Similarly, blocking calcium currents does not abolish E-C coupling in embryonic amphibian skeletal muscle (Huang, 1986). Because \( I_{\text{fam}} \) is seen in acutely dissociated fibers only during the first 2–3 wk postnatal, it is tempting to speculate that it plays a role in developmental events occurring during this period. One such event is myogenesis; in the rat lumbrical muscle, the number of fibers doubles during the first 2 wk postnatal (Betz et al., 1979). Perhaps \( I_{\text{fam}} \) plays a role in the fusion of myoblasts to form myotubes. Another significant developmental event that occurs during the first 2–3 wk postnatal is the shift from polyneuronal motor innervation to single-neuronal innervation of muscle fibers (Betz, 1987). Thus, one might imagine that \( I_{\text{fam}} \) somehow guides the elimination of supernumerary synapses. Alternatively, the presence or absence of \( I_{\text{fam}} \) may be under neural control such that maturation of the motor innervation causes the elimination of \( I_{\text{fam}} \) as has been demonstrated for the apamin-sensitive \( Ca^{2+} \)-activated K\(^+\) channel in mammalian skeletal muscle (Schmid-Antomarchi et al., 1985). Establishing the developmental factors that cause the disappearance of \( I_{\text{fam}} \) should provide insights into the function of this current.

Figure 5. Postnatal developmental increase in capacitance normalized by fiber surface area. For each fiber, the linear capacitance was measured as described in the Methods. Fiber surface area was calculated, under the assumption of right-cylindrical geometry, from visual measurements of fiber diameter and length. Error bars indicate \( \pm \) SEM. The smooth curve is the same one as in Figs. 2 B and 4.
Developmental Shift in the Voltage Dependence of Charge Movement and $I_{\text{slow}}$

The steady state charge-voltage distribution for developing fibers was characterized in terms of the parameters $k$ and $V$, which appear in Eq. 1. The values of $k$ determined in developing FDB fibers are similar to those determined for adult mammalian fast-twitch muscles (Hollingworth and Marshall, 1981; Dulhunty and Gage, 1983; Simon and Beam, 1985a; Lamb, 1986a). The value of $V$ determined for FDB fibers from the oldest animals (>25 d) is ~15–20 mV more positive than the values for $V$ determined for adult mammalian muscle (comparison of values for $V$ requires taking into account differences in measurement conditions; see Discussion in Simon and Beam, 1985a).

Independent of the absolute values for $V$, our data suggest that there may be a developmental change in the steady state charge-voltage relationship. Thus, in FDB fibers from newborn animals, $V$ was ~12 mV hyperpolarized compared with $V$ for older animals. Denervation of adult mammalian fast-twitch muscle has been reported to produce a hyperpolarizing shift in $V$ of about the same magnitude (Hollingworth et al., 1984; Dulhunty and Gage, 1985). During postnatal development, the voltage dependence for activation of $I_{\text{slow}}$ shifted in the same direction as steady state charge distribution, although the shift for $I_{\text{slow}}$ (~6 mV) was smaller.

Comparison of the Voltage Dependence of Charge Movement and $I_{\text{slow}}$

At least a portion of charge movement in skeletal muscle is necessarily involved in the gating of the slow calcium current, making it of interest to compare the voltage dependence of these two processes. At any given age, the average parameters determined by Boltzmann fits indicate that the dependence of the slow calcium conductance on potential is steeper and is centered at a more positive potential than steady state charge distribution. Thus, for the slow calcium conductance, the value of $V$ is 12–20 mV depolarized, and the value of $k$ (which is inversely related to steepness) is about half, relative to these parameters for steady state charge distribution. Interestingly, in single frog skeletal muscle fibers, the voltage dependence of the peak rate of SR calcium release is steeper and shifted positively compared to the voltage dependence of steady state charge distribution (Melzer et al., 1986), and these differences are similar in magnitude to those found by us for slow calcium conductance relative to charge movement. Thus, it appears that the voltage dependence of SR calcium release is similar to that for activation of slow calcium conductance. Although a direct comparison of these two processes has not been reported, Brum et al. (1987) have compared calcium current amplitudes with changes in myoplasmic calcium concentrations in single frog muscle fibers. Their results are consistent with a similar voltage dependence for activation of slow calcium conductance and peak SR calcium release.

Given a similar voltage dependence for activation of slow calcium conductance and SR calcium release, it seems a reasonable hypothesis that the same process (e.g., charge movement) controls both (Rios et al., 1986; Beam et al., 1986; Lamb and Walsh, 1987). Melzer et al. (1986) have described simple physical models in which charge movement accounts for the observed voltage dependence of SR calcium release. The same models would also be able to account for the voltage dependence of $I_{\text{slow}}$ activation, although they would not explain why $I_{\text{slow}}$ reaches its peak at a time...
long after the movement of charge is essentially complete. For this, it seems necessary to postulate that after the movement of charge, at least one slow and relatively voltage-independent transition must occur before the slow calcium channel opens.

**Tubular Origin of $I_{\text{slow}}$ and Charge Movement**

Studies of adult skeletal muscle imply that both $I_{\text{slow}}$ and charge movement arise mainly, if not entirely, within the t-system. In frog skeletal muscle, the slow calcium channel current is eliminated by glycerol shock (Nicola Siri et al., 1980; Potreau and Raymond, 1980), a treatment that uncouples the t-system from the surface of the fiber. Also in frog skeletal muscle, studies of cut fibers demonstrate that the time-dependent decay of the slow calcium current during a prolonged depolarization is due to depletion of calcium from a restricted extracellular space having properties consistent with the t-system (Almers et al., 1981). In this regard, it is interesting that $I_{\text{slow}}$ recorded from FDB fibers of older rats showed a more prominent decay during prolonged depolarization than $I_{\text{slow}}$ from FDB fibers of younger rats (cf. Fig. 1). Because the fibers of older animals have a larger diameter and more extensive t-system, one would expect more significant calcium depletion than in fibers of younger animals (to be replenished, calcium must diffuse into the t-system from the fiber surface). In rat skeletal muscle, the effects of temperature on the charge movement time course (Simon and Beam, 1985a) and mathematical modeling of charge movement kinetics (Simon and Beam, 1985b) both indicate that the mobile charge is in the t-system. Experimental manipulations that alter the electrical coupling of the t-system to the fiber surface also imply a tubular location for the mobile charge in both frog (Campbell, 1986) and rabbit (Lamb, 1986a) skeletal muscle.

**Does T-Tubular Growth Account for the Increases in $I_{\text{slow}}$ and $Q_{\text{max}}$?**

When normalized by total linear capacitance, peak $I_{\text{slow}}$ and $Q_{\text{max}}$ both increase postnatally. However, early in development, the t-system contributes a smaller proportion of the total linear capacitance than later in development (Fig. 5). Thus, the question arises: do $Q_{\text{max}}$ and peak $I_{\text{slow}}$ simply scale with the membrane area of the t-system? Our data do not permit us to answer this question. If the surface membrane is assumed to have a capacitance of 1 $\mu$F/cm$^2$, then the fraction of total capacitance contributed by the t-system approximately doubled during postnatal development (i.e., from 0.5/1.5 to 2.0/3.0). This doubling is much smaller than the approximately fivefold increase in normalized $Q_{\text{max}}$ and $I_{\text{slow}}$. However, if the surface membrane is assumed to have a capacitance of 1.35 $\mu$F/cm$^2$, then the fractional contribution of the t-system to the total capacitance increased slightly more than fivefold postnatally (i.e., from 0.15/1.5 to 1.65/3.0), an increase sufficient to account for the increases in $I_{\text{slow}}$ and $Q_{\text{max}}$. In frog skeletal muscle fibers, where it has been most extensively investigated, the capacitance referred to the right-cylindrical area of smooth fiber surface exceeds 1 $\mu$F/cm$^2$, with different workers having determined values ranging from 1.1 to $>2.0$ $\mu$F/cm$^2$ (Chandler and Schneider, 1976; Mathias et al., 1977; Takashima, 1985). The capacitance values exceeding 1.0 $\mu$F/cm$^2$ appear to be accounted for by folding and indentation of the fiber surface (Dulhunty and Franzini-Armstrong, 1975). Unfortunately, there are no data as to whether the folding and indentation of
the surface membrane of neonatal FDB fibers are sufficient to increase the surface capacitance to 1.35 μF/cm² and whether these geometric factors remain constant during development. Moreover, the subcellular distribution (tubular vs. surface) of charge movement and slow calcium channels may change postnatally. Thus, although \( Q_{\text{max}} \) and \( I_{\text{slow}} \) both increase faster than the total linear capacitance, it is uncertain whether \( Q_{\text{max}} \) and \( I_{\text{slow}} \) normalized only by t-system area remain constant or increase during postnatal development. However, even if \( Q_{\text{max}} \) and \( I_{\text{slow}} \) increase only as fast as the t-system, it still implies that both are important to adult muscle.

\( I_{\text{slow}}, \) Charge Movement, and Dihydropyridine Receptors

Kazazoglou et al. (1983) have determined that the number of dihydropyridine-binding sites (per milligram of protein) in rat leg skeletal muscle increases ~4.5-fold during the first 21 d postnatal. Both the time course and the magnitude of this increase are very similar to the time course and magnitude of the postnatal increases in \( I_{\text{slow}}, \) \( Q_{\text{max}}, \) and the t-system. Furthermore, dihydropyridine-binding sites appear to be localized preferentially in the t-system (Fosset et al., 1983; Glossman et al., 1983), just as are the channels responsible for \( I_{\text{slow}} \) and the mobile charge giving rise to charge movement (see above). These parallels are consistent with the growing body of evidence that, although calcium entry mediated by \( I_{\text{slow}} \) is not involved in E-C coupling, the slow calcium channel protein, or a closely related protein, is essential for E-C coupling. Thus, under appropriate conditions, the organic calcium channel antagonist D-600 causes paralysis of frog skeletal muscle (Eisenberg et al., 1983). Under these same conditions, D-600 abolishes charge movements (Hui et al., 1984). Similarly, nifedipine reduces both the magnitude of charge movement (Lamb, 1986b; Rios and Brum, 1987) and the transient of calcium release from the SR (Rios and Brum, 1987); both these effects are potentiated at depolarized holding potentials (Rios and Brum, 1987), as is the dihydropyridine block of calcium current (Bean, 1984; Cognard et al., 1986b; Beam and Knudson, 1988). Further evidence linking \( I_{\text{slow}}, \) charge movement, and dihydropyridine receptors is provided by studies of mice with muscular dysgenesis, a recessive mutation that causes E-C coupling in skeletal muscle to fail (Powell and Fambrough, 1973; Klaus et al., 1983). Dysgenic skeletal muscle cells lack \( I_{\text{slow}} \) (Beam et al., 1986) and have only about one-fifth as many dihydropyridine receptors as normal muscle (Pincon-Raymond et al., 1985).

During E-C coupling, calcium is released from the SR in response to a change in potential across the t-system membrane. Thus, there must be “sensor” molecules in the t-tubular membrane whose state alters when potential across the t-system changes. Schneider and Chandler (1973) suggested that these hypothetical sensor molecules are responsible for the signal measured as charge movement. Recently, Rios and Brum (1987) have proposed that the molecule that generates charge movement is the skeletal muscle dihydropyridine receptor. Since dihydropyridines block \( I_{\text{slow}}, \) one would expect that the slow calcium channel is also a dihydropyridine receptor. Perhaps the same molecule that generates charge movement also gives rise to \( I_{\text{slow}} \) (Rios et al., 1986; Beam et al., 1986; Lamb and Walsh, 1987). Specifically, one might imagine that the charge movement molecule represents a modified calcium channel protein whose main function is to sense t-system potential and control SR calcium...
release, but which also maintains the function of gating calcium entry across the t-tubular membrane, even though this calcium movement is not directly involved in E-C coupling. Alternatively, the molecule that generates charge movement and the one that gives rise to $I_{\text{slow}}$ may represent closely related proteins, both of which are affected by dihydropyridines in a voltage-dependent manner. In either case, the function, if any, of $I_{\text{slow}}$ itself remains unidentified.

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