Calcium flux in turtle ventricular myocytes

Gina L. J. Galli,1,2 Edwin W. Taylor,2 and Holly A. Shiels1

1Faculty of Life Sciences, The University of Manchester, Core Technology Facility, Manchester, United Kingdom; and 2School of Biosciences, The University of Birmingham, Birmingham, United Kingdom

Submitted 15 June 2006; accepted in final form 2 August 2006

Galli, Gina L. J., Edwin W. Taylor, and Holly A. Shiels. Calcium flux in turtle ventricular myocytes. Am J Physiol Regul Integr Comp Physiol 291: R1781–R1789, 2006.—The relative contribution of the sarcoplasmic reticulum (SR), the L-type Ca2⁺ channel and the Na⁺/Ca2⁺ exchanger (NCX) were assessed in turtle ventricular myocytes using confocal microscopy and electrophysiology. Confocal microscopy images of turtle myocytes revealed spindle-shaped cells, which lacked T-tubules and had a large surface-to-volume ratio. Myocytes loaded with the fluorescent Ca2⁺ indicator fura-2 elicited Ca2⁺ transients, which were insensitive to ryanodine and thapsigargin, indicating the SR plays a small role in the regulation of contraction and relaxation in the turtle ventricle. Sarcolemmal Ca2⁺ currents were measured using the perforated-patch voltage-clamp technique. Depolarizing voltage steps to 0 mV elicited an inward current that could be blocked by nifedipine, indicating the presence of Ca2⁺ currents originating from L-type Ca2⁺ channels (ICa,L). The density of ICa,L was 3.2 ± 0.5 pA/pF, which led to an overall total Ca2⁺ influx of 64.1 ± 9.3 μM/L. NCX activity was measured as the Ni²⁺-sensitive current at two concentrations of intracellular Na⁺ (7 and 14 mM). Total Ca2⁺ influx through the NCX during depolarizing voltage steps to 0 mV was 58.5 ± 7.7 μM/L and 26.7 ± 3.2 μM/L at 14 and 7 mM intracellular Na⁺, respectively. In the absence of the SR and L-type Ca2⁺ channels, the NCX is able to support myocyte contraction independently. Our results indicate turtle ventricular myocytes are primed for sarcolemmal Ca2⁺ transport, and most of the Ca2⁺ used for contraction originates from the L-type Ca2⁺ channel.

receptor; excitation-contraction coupling; sarcoplasmic reticulum; Na⁺/Ca2⁺ exchanger; L-type Ca2⁺ channel

FROM THE SIMPLE TWO-CHAMBERED heart of fish to the completely divided four-chambered heart of mammals, the structure and function of the vertebrate heart are remarkably varied. Nevertheless, the basic cellular process that underpins the cardiac contraction and relaxation cycle is common to all vertebrate hearts, there are important interspecific differences in the way Ca2⁺ is cycled to and from the myofilaments.

SR and sarcolemmal Ca2⁺ cycling can vary between species (3, 6, 16, 21, 32, 33, 36), different stages of development (14, 27, 28), and regionally within the heart (6, 25). In most fish and amphibians, transsarcolemmal Ca2⁺ influx is the primary source of activator Ca2⁺ responsible for initiating contraction (1, 26, 31, 39, 44). The majority of this extracellular Ca2⁺ enters the cell through L-type Ca2⁺ channels, although in some species, the NCX may also contribute significant amounts of activator Ca2⁺ (19, 34, 42). The contribution of intracellular Ca2⁺ cycling through the sarcoplasmic reticulum (SR) is minimal in most ectothermic vertebrates and varies subject to experimental conditions (6, 12, 13, 21, 31). In contrast, activation of the myofilaments in most adult mammalian hearts occurs mainly through the mobilization of the intracellular Ca2⁺ stores of the SR, with sarcolemmal Ca2⁺ influx primarily acting as a trigger for SR Ca2⁺ release (6).

The regulation of contraction with respect to Ca2⁺ cycling has never been studied on a cellular level in any reptilian species. Recently, we have shown isolated turtle ventricular muscle to be relatively insensitive to ryanodine, a compound that inhibits the SR Ca2⁺ release channel. These results suggest that, similar to fish, the SR contributes little Ca2⁺ to turtle heart contraction and relaxation (12). In the absence of a functional SR, we would hypothesize that sarcolemmal Ca2⁺ transport is the primary source of Ca2⁺ and that the NCX is the primary Ca2⁺ efflux pathway. The aim of the present study was to confirm this hypothesis experimentally. Here, we have examined the contribution of the cardiac L-type Ca2⁺ channel, the NCX, and the SR to contraction and relaxation in single isolated ventricular myocytes from the turtle heart. Our results confirm our hypothesis that transsarcolemmal flux is the primary route for Ca2⁺ transport in turtle ventricular myocytes. Moreover, our data provide the groundwork for future studies on ion regulation of contractility in reptilian cardiomyocytes.

MATERIALS AND METHODS

Animal Origin and Care

Yellow-bellied turtles, Trachemys scripta scripta (body mass = 218.5 ± 23.9 g, heart mass = 0.61 ± 0.1 g, n = 12) were obtained from Monkfield Nutrition Ltd (Hertfordshire, UK). Turtles were held in 1.5 × 0.5 × 0.5 m plastic tanks containing water maintained at 20–21°C (40 cm depth) and dry basking platforms, allowing access to heating lamps for behavioral thermoregulation.

The costs of publication of this article were defrayed in part by the payment of page charges. The article must therefore be hereby marked “advertisement” in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.
Solutions

The isolation solution contained (in mM) 100 NaCl, 10 KCl, 1.2 K2HPO4, 4 MgSO4, 50 taurine, 20 glucose, and 10 HEPES, with pH adjusted to 6.9 via KOH. For enzymatic digestion, 1.5 mg/ml collagenase (type IA), 1 mg/ml trypsin (type IX), and 1.5 mg/ml fatty acid-free BSA were added to the solution. The extracellular solution perfusing the myocytes contained (in mM) 150 NaCl, 5.4 CsCl, 1.5 MgSO4, 0.4 NaH2PO4, 2 CaCl2, 10 glucose, 10 HEPES, with pH adjusted to 7.7 via CsOH. For whole cell voltage-clamp electrophysiological measurements, the pipette solution contained (in mM) 130 CsCl, 1 MgCl2, 5 MgATP, 5 Na3-phosphocreatine (unless stated otherwise), 10 HEPES, 15 tetraethylammonium chloride (TEA), and 0.03 Na2GTP. The EGTA concentration was 0.025 mM unless stated otherwise. The pH was adjusted to 7.2 with CsOH. Including TEA and CsOH abolished all K+ currents. In perforated-patch, voltage-clamp experiments, 240 mg/ml amphotericin was added to the pipette solution. To be certain that cells were indeed perforated and not in the whole cell configuration, a high concentration of CaCl2 (10 mM) was also included in the pipette solution. All drugs were purchased from Sigma Aldrich.

Isolated Myocyte Preparation

Myocytes were obtained by adaptation of isolation protocols previously described for fish (32, 34, 42). All procedures were made in accordance with UK Home Office regulations. Briefly, turtles were euthanized by decapitation, and the head was immediately submerged into liquid nitrogen to fully destroy all neural connections. A 2 cm × 2 cm piece of the ventral plastron was removed above the heart using a bone saw. The heart was excised and a cannula was inserted into the right aortic arch and advanced into the ventricle for perfusion. The solution was removed from the shaker, left to settle, and shaken at 21°C. The heart was then perfused for 15–20 min with extracellular solution for deoxygenation. All fluorescent measurements were conducted at room temperature. To examine the contribution of the SR to Ca2+ flux in turtle myocytes, cells were field stimulated to contract at 0.2 and 0.5 Hz in the presence and absence of SR blockade [10 μM ryanodine and 2 μM thapsigargin (Sigma Aldrich)]. To ensure complete inhibition of the SR, cells were perfused with ryanodine and thapsigargin for a period of 5 min before measurements were commenced. The ratio of the fluorescence emitted at 510 nm in response to alternate illumination with light at 340 and 380 nm (Cairn Research) was used as our index of [Ca2+]i.

Electrophysiological Measurements of Sarcolemmal Currents

Samples of myocytes were recorded to the recording chamber and left to settle and attach to the bottom. Cells were perfused with extracellular solution at a rate of 1–2 ml/min at room temperature (~21°C). Both whole cell and perforated-patch, voltage-clamp experiments were performed using an Axopatch 200B amplifier (Axon Instruments) with a CV203BU headstage (Axon Instruments). Patch pipettes were pulled with borosilicate glass (Harvard Apparatus) and had a resistance of 2.4 ± 0.02 MΩ when filled with pipette solution. In perforated-patch, voltage-clamp experiments, once a gigaohm seal had formed, patch pipette resistance (5–6 MΩ) was compensated for, and Ra was monitored using the membrane test function to assess the extent of perforation. Once electrical access to the cell was gained, the cell capacitive currents were compensated for by manually adjusting series resistance (Ra) and the cell capacitance compensation circuits. Series resistance [whole cell Ra = 8.2 ± 0.9 MΩ (n = 38); perforated Ra = 17.2 ± 1.5 MΩ (n = 40)] and capacitance [Cm = 42.4 ± 1.9 pF (n = 78)] were measured using the membrane test function of pClamp 9.0 software (Axon Instruments). In the perforated-patch configuration, Ra was monitored throughout the experiments. Signals were analyzed offline using Clampfit 9.0 software (Axon Instruments).

The voltage-clamp waveform protocols for each experiment are provided in the figures. The amplitude of ICa was calculated as the difference between peak inward current and the current at the end of the depolarizing pulse. ICa was normalized to the cell area by dividing the amplitude of ICa by the cell capacitance to give pA/pF. To assess the rate of inactivation of ICa, tau fast (τf) and tau slow (τs) inactivation components were derived by fitting a second-order exponential function to the decaying portion of ICa using the Chebyshev procedure (Clampfit software, Axon Instruments). Steady-state kinetic parameters were determined by fitting steady-state activation and inactivation data to Boltzmann equations to calculate the half-activating and half-inactivating potential (V1/2) and the slope of activation and inactivation (κ), as previously described (44). Recovery from inactivation of ICa was assessed by normalizing current amplitude at a constant test pulse (500 ms, –70 to 0 mV) to the constant prepulse value (500 ms, –70 to 0 mV) after various interpulse durations (50–350 ms, –70 mV, see Fig. 6). The contribution of ICa to total

Table 1. Morphometric measurements of turtle ventricular myocytes

<table>
<thead>
<tr>
<th>Cell length, μm</th>
<th>Cell width, μm</th>
<th>Cell depth, μm</th>
<th>Cell capacitance, pF</th>
<th>Cell volume, pl</th>
<th>Ratio, pF/pl</th>
</tr>
</thead>
<tbody>
<tr>
<td>189.1 ± 10.3</td>
<td>7.2 ± 0.4</td>
<td>5.4 ± 0.3</td>
<td>42.4 ± 1.9</td>
<td>2.3 ± 0.1</td>
<td>18.3</td>
</tr>
</tbody>
</table>

Data are expressed as means ± SE, n = 18. Cell length, width, and depth were measured by analysis of confocal microscopy images of myocytes. Cell volume was derived from cell capacitance.

Fluorescent Measurements of [Ca2+]i, Using Fura-2 AM

To record changes in [Ca2+]i, ventricular myocytes were loaded with the AM form of the Ca2+-sensitive fluorescent dye Fura-2 (Molecular Probes) to a final concentration of 4 μmol/l. Myocytes were shaken gently for 10 min to allow loading; then a sample of the cells were placed on the stage and left to settle for a further 5 min. Cells were then perfused for 15–20 min with extracellular solution for deesterification. All fluorescent measurements were conducted at room temperature. To examine the contribution of the SR to Ca2+ flux in turtle myocytes, cells were field stimulated to contract at 0.2 and 0.5 Hz in the presence and absence of SR blockade [10 μM ryanodine and 2 μM thapsigargin (Sigma Aldrich)]. To ensure complete inhibition of the SR, cells were perfused with ryanodine and thapsigargin for a period of 5 min before measurements were commenced. The ratio of the fluorescence emitted at 510 nm in response to alternate illumination with light at 340 and 380 nm (Cairn Research) was used as our index of [Ca2+]i.
cellular \([\text{Ca}^{2+}]\) was calculated from the transferred charges and cell volume. Charge transfer was determined by integrating the inactivating portion of the \([\text{Ca}^{2+}]\) current for 500-ms square-wave voltage pulses from \(-70\) mV to 0 mV. Cell volume was calculated from the measured cell capacitance (42.4 ± 1.9) and the surface-to-volume ratio of the cells. The myocytes were considered to be flat elliptical cylinders with an axis ratio of 1.2 for the elliptical cross section (34, 43, 44). The change in total cellular \([\text{Ca}^{2+}]\) due to \([\text{Ca}^{2+}]\) influx through L-type \([\text{Ca}^{2+}]\) channels was expressed as a function of nonmitochondrial volume (34, 44).

**Statistical Methods**

With the exception of original traces and voltage protocols, data are given as mean values ± SE. N values are for number of cells in which the minimum number of animals is \(n = 4\). Statistical tests are supplied in the appropriate figure legends.

**RESULTS**

**Cell Morphology**

Dissociation of the turtle heart required a higher concentration of digestive enzymes, longer perfusion times, and higher temperatures than those used previously for various fish species (32, 34, 43, 44). These differences may be due to variation in the structure of the extracellular matrix of the turtle heart compared with the fish heart or may relate to the complication of perfusing a three-chambered ventricle. Light and confocal microscopy images of isolated turtle ventricular myocytes are displayed in Fig. 1. Ventricular myocytes were typically spindle-shaped, being \(~190\) μm in length and \(5-7\) μm in width and depth (Table 1). When comparing the light and confocal images, it is apparent that the sarcomeres of the myocytes are not associated with T-tubules. Myocytes had a small cell volume \((\sim 2\) pl), leading to a large surface area-to-volume ratio, typical of ectothermic vertebrates.

**Turtle \([\text{Ca}^{2+}]\) Transients and the Functional Significance of the SR**

The functional significance of the SR was assessed via pharmacological blockade with ryanodine and thapsigargin \((10^{-6}\) M and \(2^{-6}\) M, respectively). Turtle myocytes were loaded with Fura 2 and field stimulated at 0.2 Hz in the absence and presence of SR blockade (Fig. 2). The amplitude of \([\text{Ca}^{2+}]\) was not significantly altered following inhibition of the SR (Fig. 2B), indicating the SR plays a small role in \([\text{Ca}^{2+}]\) cycling on a beat-to-beat basis.

**L-Type \([\text{Ca}^{2+}]\) Channel Properties**

Characterization of inward currents. A depolarizing voltage step from \(-80\) mV to \(-40\) mV elicited a fast-inactivating \(\text{Na}^{+}\) current \((\text{INa})\) (Fig. 3A). As both \(\text{INa}\) and the L-type \([\text{Ca}^{2+}]\) current \((\text{ICa})\) are activated within the same voltage range, we used TTX, a specific \(\text{Na}^{+}\) channel blocker, to inhibit \(\text{INa}\) (Fig. 3A). Thus a voltage step from \(-80\) mV to 0 mV in the presence of TTX blocks \(\text{INa}\) and the \(\text{ICa}\), allowing the \(\text{ICa}\) to be studied in isolation.

**Fig. 1.** Morphology of live ventricular myocytes from the turtle heart. A: light microscopy image. B: confocal microscopy image. Scale bar applies to both images. Mean morphometric data are provided in Table 1.

**Fig. 2.** The effect of inhibiting the sarcoplasmic reticulum (SR) on the turtle intracellular calcium transient \([\text{Ca}^{2+}]\). A: relative reduction in \([\text{Ca}^{2+}]\) due to SR inhibition with ryanodine \((10\) μM) and thapsigargin \((2\) μM). As both \(\text{INa}\) and the L-type \([\text{Ca}^{2+}]\) current \((\text{ICa})\) are activated within the same voltage range, we used TTX, a specific \(\text{Na}^{+}\) channel blocker, to inhibit \(\text{INa}\) (Fig. 3A). Thus a voltage step from \(-80\) mV to 0 mV in the presence of TTX blocks \(\text{INa}\) and the \(\text{ICa}\), allowing the \(\text{ICa}\) to be studied in isolation.
In many species, $I_{Ca}$ is known to deteriorate or “rundown” over time, particularly when measured in the whole-cell configuration. We assessed $I_{Ca}$ rundown in turtle myocytes using the whole cell voltage-clamp technique (with various intracellular $Ca^{2+}$ buffering) compared with the perforated-patch, voltage-clamp method (Fig. 4). Cells were depolarized from $-70$ mV to 0 mV, and $I_{Ca}$ density was measured over a period of 12 min. In the whole cell configuration, $I_{Ca}$ deteriorated to 50% of its original value within ~2 min, and to 20% after 12 min, regardless of the level of intracellular $Ca^{2+}$ buffering (Fig. 4). In the perforated-patch configuration, $I_{Ca}$ remained relatively stable and was only reduced by 10% over the entire 12-min period. Therefore, the perforated-patch, voltage-clamp technique was used in all subsequent experiments when measuring $I_{Ca}$. Interestingly, when using 5 mM EGTA or 5 mM BAPTA, the $I_{Ca}$ density ($3.8 \pm 1.0$ pA/pF and $4.2 \pm 0.3$ pA/pF, respectively) was similar to perforated-patch $I_{Ca}$ density ($3.2 \pm 0.5$ pA/pF). However, when using 25 $\mu$M EGTA, $I_{Ca}$ density was considerably lower ($1.54 \pm 0.36$ pA/pF). This suggests the buffering capacity of turtle myocytes is greater than 25 $\mu$M EGTA.

$I_{Ca}$ density, kinetics and voltage relations. The current-voltage relationship for turtle ventricular myocytes is shown in Fig. 5. $I_{Ca}$ activated at approximately $-40$ mV, peaked at 0 mV, and reversed at 60 mV. At peak $I_{Ca}$ density ($-3.2 \pm 0.5$ pA/pF), the time constant for fast inactivation (ri) and slow inactivation (rs) was 28.7 ± 1.5 ms and 169.2 ± 8.6 ms, respectively ($n = 15$). Because of this relatively slow inactivation time, charge transfer, and therefore total $Ca^{2+}$ influx through L-type $Ca^{2+}$ channels, was particularly high in turtle myocytes (Table 2).

Steady-state activation and inactivation of $I_{Ca}$. Activation of $I_{Ca}$ began positive at $-40$ mV and was half maximal ($V_h$) at $-3.9 \pm 2.3$ mV, while inactivation of $I_{Ca}$, or channel availability, began decreasing positive at $-40$ mV and was half complete at $-22.5 \pm 1.3$ mV (Fig. 6). The slopes of activation...
and inactivation \( (k) \) were 6.4 ± 0.7 and 4.4 ± 0.3, respectively. At voltages positive to 10 mV, channel inactivation was attenuated, probably due to a reduced driving force and consequently less \( Ca^{2+} \)-dependent inactivation. As a result of overlap between activation and inactivation curves, a window current was evident between −40 and 0 mV. The window current was maximal at approximately −18 mV, where it contributed 5% of maximal conductance (Fig. 6, inset).

\( I_{Ca} \) recovery from inactivation. The recovery of \( I_{Ca} \) from inactivation at −70 mV following 1-s pulses to 0 mV is shown in Fig. 7. The number of recovered channels increased as the duration between the prepulse and the test pulse was lengthened. The time constant of recovery from inactivation \( (\tau) \) was 157.3 ± 18.6 ms, thus at physiologically relevant frequencies of contraction (0.2–1 Hz), incomplete restitution of turtle L-type \( Ca^{2+} \) channels is unlikely to occur.

NCX Current Properties

Characterization of \( I_{NCX} \). The capacity of the NCX to transfer charge is highly dependent on the intracellular [\( Na^{+} \)]. In mammalian cardiac myocytes, resting intracellular \( Na^{+} \) concentration \( ([Na^{+}]_{i}) \) varies between species with a range of 4–15 mM (7) Because in vivo \( [Na^{+}]_{i} \), is not known for turtle myocytes, we investigated the capacity of the NCX at a low (7 mM) and a high (14 mM) amount of \( Na^{+} \) in the patch pipette. Membrane current was measured in the presence of 50 \( \mu \)M nifedipine to block \( I_{Ca} \), and a combination of 10 \( \mu \)M ryanodine and 2 \( \mu \)M thapsigargin was used to inhibit possible SR \( Ca^{2+} \) release and reuptake. Under these conditions, a square-wave voltage pulse from −70 mV to 0 mV for 500 ms gave rise to a maintained outward current, which could be blocked with 10 mM NiCl\(_{2}\), confirming the presence of a \( Ni^{2+} \)-sensitive \( Na^{+}/Ca^{2+} \) exchange current \( (I_{NCX}) \). The \( Ni^{2+} \)-sensitive currents elicited with 14 mM and 7 mM [\( Na^{+} \)], are shown in Fig. 8. \( I_{NCX} \) was integrated to give a measure of charge transfer so that total \( Ca^{2+} \) influx through the NCX could be calculated. At 14 and 7 mM, intracellular \( Na^{+} \), NCX charge transfer was 0.24 ± 0.1 and 0.12 ± 0.1, respectively, giving a total \( Ca^{2+} \) influx through the NCX at 0 mV of 58.5 ± 7.7 \( \mu \)mol/l and 26.7 ± 3.2 \( \mu \)mol/l, respectively (\( n = 9 \) and 7, Fig. 8). Total \( Ca^{2+} \) influx through the NCX was significantly greater with 14 mM than with 7 mM [\( Na^{+} \)], \( (P = 0.004) \).

NCX voltage sensitivity. The voltage-dependence of the \( Na^{+}/Ca^{2+} \) exchange current was measured using a voltage ramp protocol in the presence of 50 \( \mu \)M nifedipine (Fig. 9, inset). \( I_{NCX} \) was identified as the \( Ni^{2+} \)-sensitive current. The measured reversal potential of \( I_{NCX} \) (ENCX) was ~30 mV, regardless of the intracellular \( Na^{+} \) concentration (Fig. 9). The calculated ENCX under the present experimental conditions is 28.29 mV for 14 mM [\( Na^{+} \)], and 80.68 mV for 7 mM [\( Na^{+} \)]. Thus, although the experimentally derived ENCX at 14 mM

![Figure 5](image-url)  
**Fig. 5.** L-type \( Ca^{2+} \) channel current-voltage relationships in turtle ventricular myocytes. \( I_{Ca} \) was normalized to myocyte capacitance (pF) and expressed as current density (pA pF\(^{-1}\)). Values are expressed as means ± SE (\( n = 15 \)).

![Figure 6](image-url)  
**Fig. 6.** Steady-state activation and inactivation of \( I_{Ca} \) in turtle ventricular cardiac myocytes. A: voltage protocol used to measure steady-state activation and inactivation. B: representative current recording from a turtle ventricular myocyte (capacitance = 78 pF) subjected to the voltage protocol given in A; C: mean steady-state inactivation and activation profiles. Values are means ± SE (\( n = 10 \)). Inactivation is measured by depolarizing from −70 mV to a test potential for 1 s and then testing the remaining available \( I_{Ca} \) at 0 mV. Activation is calculated by dividing peak current by apparent driving force \( (\text{applied membrane potential (Em)} - \text{reversal potential (Erev)} \) according to Ohm’s law. 

<table>
<thead>
<tr>
<th>Capacitance, pF</th>
<th>( I_{Ca} ) Density, pA/pF</th>
<th>Charge Transfer, pC</th>
<th>Charge Density, pA/pF</th>
<th>( \Delta [Ca^{2+}] ) + Non-M, ( \mu )mol/l</th>
</tr>
</thead>
<tbody>
<tr>
<td>42.4 ± 1.9</td>
<td>3.2 ± 0.5</td>
<td>−11.3 ± 1.7</td>
<td>−0.27 ± 0.04</td>
<td>64.1 ± 9.3</td>
</tr>
</tbody>
</table>

Data are expressed as means ± SE, \( n = 15 \). \( \Delta [Ca^{2+}] \) + Non-M, Change in total cellular \([Ca^{2+}]\) expressed as a function of nonmitochondrial cell volume.
[Na\textsuperscript{+}]_i agrees well with the calculated value, it is expected that decreasing [Na\textsuperscript{+}]_i will increase ENCX. A possible explanation for this discrepancy is that diastolic Ca\textsuperscript{2+} levels may also change when altering [Na\textsuperscript{+}]_i, which will also affect the measured ENCX. At both 7 mM and 14 mM intracellular [Na\textsuperscript{+}], the outward INCX current showed a steep increase with increasing voltage (outward rectification) (Fig. 9), while the inward current peaked at approximately -10 to -30 mV and then decreased at more negative voltages. I\textsubscript{NCX} was significantly larger with 14 mM than with 7 mM [Na\textsuperscript{+}]_i, at almost all membrane voltages. Importantly, in the absence of the L-type Ca\textsuperscript{2+} channel and SR Ca\textsuperscript{2+} release, I\textsubscript{NCX} was able to trigger contraction independently. This was the case with both concentrations of intracellular Na\textsuperscript{+}; however, at 14 mM [Na\textsuperscript{+}], a greater degree of myocyte shortening occurred compared with 7 mM [Na\textsuperscript{+}]_i (data not shown).

**DISCUSSION**

In the hearts of most ectothermic vertebrates, the SR seems to contribute little to contraction and relaxation (see Ref. 45). The cardiac muscle of many ectothermic species, including the turtle, are ryanodine insensitive and exhibit a postrest decay of force, suggesting SR independence, at least under normal physiological conditions (10, 12, 17, 41). In the present study, we tested SR involvement in cellular Ca\textsuperscript{2+} flux directly, and we show that inhibition of SR function with a combination of ryanodine and thapsigargin had little effect on [Ca\textsuperscript{2+}]_i in turtle ventricular myocytes, supporting earlier findings on isolated muscle preparations (12). Thus, in the absence of a functional SR, the turtle heart will depend strongly on transsarcolemmal Ca\textsuperscript{2+} cycling for both the contraction and relaxation of the myocyte.

Fig. 7. Recovery of I_{Ca} from inactivation in turtle ventricular myocytes. A: voltage protocol used to measure recovery from inactivation. B: representative trace from a turtle ventricular myocyte (capacitance = 78.0) subjected to the displayed voltage protocol. C: relationship between current recovery and interpulse duration for ventricular myocytes. Values are expressed as means ± SE, n = 10.

Fig. 8. Measurement of the Na\textsuperscript{+}/Ca\textsuperscript{2+} exchange current (I\textsubscript{NCX}) using square wave voltage steps in turtle ventricular myocytes. A: voltage waveform protocol was applied in the absence and then the presence of NiCl\textsubscript{2} (10 mM), a blocker of the NCX. B: I\textsubscript{NCX} was identified as the Ni\textsuperscript{2+}-sensitive current at each intracellular concentrations of Na\textsuperscript{+}: 14 and 7 mM.

Fig. 9. Measurement of the I\textsubscript{NCX} using ramp protocols (inset) in turtle ventricular myocytes. I\textsubscript{NCX} was identified as the Ni\textsuperscript{2+}-sensitive current during the hyperpolarizing phase of the ramp pulse and was measured at two different intracellular concentrations of Na\textsuperscript{+}: 7 mM Na\textsuperscript{+} (n = 5) and 14 mM Na\textsuperscript{+} (n = 5). Inset shows voltage ramp protocol. Values are means ± SE. *Significant difference between data at 7 mM Na\textsuperscript{+} and 14 mM Na\textsuperscript{+} (one-way repeated-measures ANOVA; P < 0.05).
The importance of extracellular Ca\textsuperscript{2+} cycling in turtles, and the apparent lack of SR involvement has a clear ultrastructural basis. The morphological design of turtle myocytes reveals a system primed for transsarcolemmal Ca\textsuperscript{2+} flux. Turtle myocytes are spindle-shaped (long and thin) and lack T-tubules, which is typical of ectothermic vertebrates (45). The surface area-to-volume ratio (18.3) is high compared with mammalian ventricular myocytes (rabbit, 4.6; rat, 6–8) (2, 30), but similar to other ectotherms (trout, 18.2; crucian carp, 19.2; bluefin tuna, 14–17; mackerel, 18–22) (32, 42). This large surface area-to-volume ratio will enhance the efficacy of sarcosomal Ca\textsuperscript{2+} transport by reducing the diffusional distance that Ca\textsuperscript{2+} has to travel to the myofilaments. Moreover, the myofilaments of ectothermic myocytes are subsarcolemmal (42), further promoting the close association of the sarcolemmal membrane and the contractile elements of the cell.

In most ectotherms, the majority of extracellular Ca\textsuperscript{2+} enters the cell through L-type Ca\textsuperscript{2+} channels (18, 34, 42). The central role of these channels can be demonstrated by an almost complete inhibition of force production by L-type Ca\textsuperscript{2+} channel blockers in fish (1). Our study demonstrates the turtle is no exception to this trend and relies strongly on this form of Ca\textsuperscript{2+} influx for contraction. The density and kinetics of turtle I\textsubscript{Ca} are typical of other ectothermic species (20, 32, 42, 43). Steady-state activation and inactivation curves and the time taken for recovery from inactivation also coincide with those found in fish (32, 34, 42, 43). A sizeable window current exists at room temperature, which probably contributes to the prolongation of the action potential, and may be important in Ca\textsuperscript{2+} cycling at lower body temperatures (see Ref. 34).

Interestingly, although the density and kinetics of tonic I\textsubscript{Ca}, with the exception of I\textsubscript{Ca} time course of inactivation (see below), may be similar to other ectothermic species, the pharmacology suggests the heart may predominantly contain L-type Ca\textsuperscript{2+} channels composed of a different subunit than most vertebrate cardiac L-type channels. The L-type Ca\textsuperscript{2+} channel is made up of a number of subunits, but the a\textsubscript{1L}-subunit is the main functional unit of the channel, responsible for pore formation and voltage sensitivity (35). In cardiac muscle, it exists as two main isoforms: a\textsubscript{1C} and a\textsubscript{1D} (8). The a\textsubscript{1C} subunit is most commonly found in the mammalian myocardium and is characteristically sensitive to dihydropyridines (24). Conversely, the a\textsubscript{1D} is found abundantly in neurons or pacemaker regions of the heart and requires a higher dose of dihydropyridines (~10–20-fold higher) to completely inhibit I\textsubscript{Ca} (4, 22, 38, 46). Thus the pharmacology of the turtle L-type Ca\textsuperscript{2+} current may suggest the heart contains a large complement of a\textsubscript{1D} subunits in their L-type Ca\textsuperscript{2+} channels. It is not clear what the functional significance of these channels may be in turtle ventricular myocytes; however, L-type Ca\textsuperscript{2+} channels containing the a\textsubscript{1D} subunit have been linked with conduction of the action potential in the mammalian heart (23, 37, 47).

Our results indicate the turtle L-type Ca\textsuperscript{2+} channel is the dominant source of extracellular Ca\textsuperscript{2+}, with a total Ca\textsuperscript{2+} influx of 64.1 ± 9.3 of nonmitochondrial (n = 15) space. This value is 5 times that found in adult mammalian ventricular myocytes (5, 29) and approximately double that of certain fish ventricular cells (34, 42, 43). This difference is probably due to the relatively slow time course of inactivation of turtle I\textsubscript{Ca} and may be a direct result of less Ca\textsuperscript{2+}-induced inactivation of the L-type Ca\textsuperscript{2+} channel and the lack of a functional SR. However, it must be noted that in similar experiments with fish, measurements were made in the whole cell configuration with intracellular buffering, which can increase the amplitude of I\textsubscript{Ca} and therefore the rate of Ca\textsuperscript{2+}-dependent inactivation of the channel. Nevertheless, when compared with mammalian studies using a perforated-patch configuration, our results indicate the turtle L-type Ca\textsuperscript{2+} channel may contribute an enormous amount of Ca\textsuperscript{2+} for contraction, probably enough to support myocyte contraction independently of other cellular cycling mechanisms.

The large amount of Ca\textsuperscript{2+} that enters the cell during contraction via the L-type channels has to be removed to allow relaxation, and if the SR is not involved in the relaxation process, then another mechanism must be in place. In mammals, the SR and the NCX compete for the removal of Ca\textsuperscript{2+} during relaxation (6). Thus, in the absence of a functional SR, it is expected that the NCX will now act as the primary Ca\textsuperscript{2+} efflux pathway in turtle ventricular myocytes. In some species the NCX also contributes to contraction by entering "reverse mode" (Ca\textsuperscript{2+} in, Na\textsuperscript{+} out) during the upsweep of the action potential. In the hearts of neonatal mammals, which are similar to turtle cardiac myocytes in both structure and function, the NCX accounts for 65–75% of total Ca\textsuperscript{2+} influx (27, 28), and among ectotherms, the NCX alone can activate contraction in the crucian carp or rainbow trout, with up to 50% of Ca\textsuperscript{2+} entry mediated through this reverse NCX activity (19, 44). Thus, in ectothermic vertebrates, and also mammalian neonates, the NCX may provide a substantial proportion of the activator Ca\textsuperscript{2+} necessary for contraction.

Measurement of NCX activity is complicated by the lack of information regarding the intracellular concentration of Na\textsuperscript{+} in turtle ventricular myocytes. In fish, varying [Na\textsuperscript{+}], greatly influences the activity of the NCX, with a reduction from 16 to 10 mM [Na\textsuperscript{+}], leading to a 58% reduction in Ca\textsuperscript{2+} entry through the NCX (19). Thus we have measured the capacity of the NCX at two different concentrations of [Na\textsuperscript{+}]: 14 and 7 mM. Depolarizing voltage steps to 0 mV in the presence of L-type channel and SR blockade led to a nickel-sensitive outward current corresponding to a total Ca\textsuperscript{2+} influx of 58.5 ± 7.7 μmol/l (n = 9) and 26.7 ± 3.2 μmol/l (n = 7) at 14 and 7 mM [Na\textsuperscript{+}], respectively. Importantly, at either concentration of [Na\textsuperscript{+}], NCX Ca\textsuperscript{2+} entry was sufficient to support myocyte contraction independently. If we speculate that [Na\textsuperscript{+}], in turtle ventricular myocytes is somewhere between 7 and 14 mM and we combine these figures with Ca\textsuperscript{2+} entry via L-type channels, we could expect a total Ca\textsuperscript{2+} influx at 0 mV via both mechanisms to amount to ~100 μmol/l. Of this, ~35% of activator Ca\textsuperscript{2+} originates from the NCX. This percentage is slightly higher than that seen in the crucian carp (~25%) when NCX activity was measured using similar voltage protocols and intracellular constituents. It is important to note that in the present study, L-type Ca\textsuperscript{2+} channels were inhibited with nifedipine, while measuring NCX activity. Previous studies measuring NCX activity in fish have suggested the presence of an intact Ca\textsuperscript{2+} current via L-type channels will increase forward mode NCX activity, due to the concentration-dependent nature of the exchanger and the reduced driving force for Ca\textsuperscript{2+} entry (19, 44). Thus our study may have overestimated the amount of Ca\textsuperscript{2+} entry via the NCX, although Hove-Madsen et al. (19) found that preserving I\textsubscript{Ca} during depolarizing voltage steps had little effect on NCX Ca\textsuperscript{2+} entry at 0 mV. In any case, it seems...
clear that the NCX is capable of contributing activator Ca\(^{2+}\) for contraction in turtle myocytes, and importantly, it is able to support myocyte contraction independent of the SR and L-type Ca\(^{2+}\) channels.

**Perspectives**

This is the first study to address cellular Ca\(^{2+}\) cycling in any reptilian species. Our results show that turtles, similar to most fish, rely predominantly on the L-type Ca\(^{2+}\) channel for delivering extracellular Ca\(^{2+}\) for contraction under routine environmental conditions. We also show that the NCX is a powerful route for both Ca\(^{2+}\) entry and Ca\(^{2+}\) removal. Freshwater turtles are renowned for their ability to tolerate a wide range of environmental challenges, and some species hibernate in aquatic habitats for up to 6 mo, subjecting themselves to large temperature fluctuations, long periods of hypoxia and anoxia, and consequential acidosis (15, 40). The mammalian myocardium is particularly sensitive to environmental change (6, 9, 11), which suggests the turtle heart may have physiological specializations in E-C coupling, which allow them to cope with their changing environment. The present study provides the groundwork for cellular studies on ion regulation of contractility in reptilian cardiomyocytes. Thus future studies should be aimed at investigating how these Ca\(^{2+}\) flux pathways are affected by temperature, hypoxia, and acidosis to provide mechanistic insight into the stress tolerance of the turtle heart.

**ACKNOWLEDGMENTS**

We especially thank Prof. David Eisner and Dr Andrew Trafford for their helpful advice and technical expertise.

**GRANTS**

This study was funded by The BBSRC, The Welcome Trust, The Company of Biologists, and The Anglo-Danish Society.

**REFERENCES**


