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Resting Tension Participates in the Modulation of Active Tension in Isolated Guinea Pig Ventricular Myocytes

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We studied active and passive properties of intact isolated guinea-pig ventricular myocytes in auxotonic conditions. Cells were attached using carbon fibres. The passive properties of the myocytes, in the presence of the stretch-activated channel blocker streptomycin sulphate, could be separated into two groups: stiff cells (stiffness slope \(= 2.88 \pm 0.93 \text{nN/}\mu \text{m}^3 \), \(n=63 \text{ cells} \)) and compliant cells (stiffness slope \(= 0.91 \pm 0.35 \text{nN/}\mu \text{m}^3 \), \(n=52 \text{ cells} \)). The study and the localization of the different kind of cells indicated that endocardium is mainly constituted of stiff cells (80%) while the epicardium contained more compliant cells (60%). When a longitudinal strain was applied to compliant cells, an increase in resting tension, diastolic sarcomere length and active tension were observed. On the other hand, in stiff cells, it induced an increase in resting tension and active tension with little change of diastolic sarcomere length. In both kinds of cells, strain had no effect on \(\text{Ca}^{2+} \) transient amplitude and shape. Plotting active tension v diastolic sarcomere length also clearly showed two separated populations of cells, corresponding to stiff and compliant cells. The results of the two groups of cells when plotting active tension v resting tension could not be distinguished. We conclude that resting tension is an important factor in the modulation of active tension by stretch in addition to interfilament lattice spacing or sarcomere length.

Key Words: Cardiac myocytes; Excitation–contraction coupling; Stretch; Frank–Starling law; Length–tension relationship.

Introduction

Stretching a striated muscle normally results in an increase of the active tension developed during a twitch. It is known that the positive inotropism of stretch, in skeletal muscle, is mainly due to geometrical rearrangements of contractile proteins. Even, if this participates to the cardiac length–tension curve, it is admitted that the steep positive inotropism of stretch is a consequence of an increase of the \(\text{Ca}^{2+} \) sensitivity of the contractile machinery (Hibberd and Jewell, 1982; Babu et al., 1988).

The mechanisms by which stretch modulates the \(\text{Ca}^{2+} \) sensitivity of troponin C (TnC) are not understood. It is established that force developed during cross-bridge interactions may modulate \(\text{Ca}^{2+} \) binding to TnC in rigor bridges (Hofmann and Fuchs, 1987) or during calcium activation (Allen and Kentish, 1988) independently of cell length. With stretch, cell parameters like cell length, cell width and height (and consequently interfilament spacing) and resting tension may change. Some authors have suggested that cell length may modulate active tension through modifications of the
interfilament spacing (Hibberd and Jewell, 1982; DeBeer et al., 1988). The interfilament spacing varies inversely with cell length because cells have a constant volume, i.e. as the cell length increased the interfilament spacing decreases. Recent works on skinned muscle cardiac fibres have demonstrated that interfilament spacing could be an important modulator of the force of contraction (McDonald and Moss, 1995; Wang and Fuchs, 1995; Fuchs and Wang, 1996). The idea is that this lattice spacing may have an ideal size below and above which the actomyosin complex would form less efficiently.

In 1994, Gannier et al. showed that streptomycin, a stretch activated channel blocker, could inhibit the large increase in resting calcium induced by stretch of guinea-pig ventricular myocytes. This work revealed two distinct populations of ventricular myocytes which were distinguished by their passive stiffness. In the present study, we confirm the existence of the two populations of cells. We compare the effects of stretch on passive tension, active tension and sarcomere length (SL) on these two kinds of cells. We find that resting tension, in addition to sarcomere length or interfilament spacing per se, is important in the modulation of active tension during a cell stretch.

Materials and Methods

Cells isolation

Guinea pigs were killed by cervical dislocation following stunning, and hearts were excised. Ventricular myocytes were enzymatically isolated as previously described (Le Guennec et al., 1990). Following isolation, cells were placed in the experimental chamber of a Nikon diaphot inverted microscope and superfused with a physiological saline solution (PSS) of the following composition (in mM): 140 NaCl, 5.4 KCl, 1 MgCl₂, 1.8 CaCl₂, 11 glucose, 0.33 NaH₂PO₄, 0.04 streptomycin sulphate, 10 HEPES; pH adjusted to 7.3 with NaOH. Streptomycin sulphate was added to the Tyrode solution to prevent the large increase in resting calcium induced by stretch (White et al., 1993; Gannier et al., 1994, 1996), and thus to unmask the two populations of cells. In some experiments (see Results), cells from the apex, base, endocardium or epicardium of left and right ventricles were obtained by cutting the digested heart with fine scissors.

Measurement of [Ca²⁺].

Myocytes were loaded with the fluorescent Ca²⁺ indicator, Indo-1 (AM form, Sigma, France), as described previously (Gannier et al., 1994). The ratio of emitted fluorescence at 405 (F405) and 480 nm (F480) was used to calculate [Ca²⁺], according to the equation:

\[ [Ca^{2+}] = K_c \beta (R - R_{min})/(R_{max} - R) \]

where \( R_{min} \) and \( R_{max} \) are the fluorescence ratios F405/F480 obtained in the absence of Ca²⁺ and at saturating [Ca²⁺] respectively (see later). \( \beta \) is the ratio of the 480 nm signal in the absence and at saturating [Ca²⁺]. A value of 380 nm was assumed for the \( K_d \) (the dissociation constant) at the temperature at which experiments were performed (26–27°C; Bers et al., 1989). The calibration of Indo-1 was performed in cells as already described by Gannier et al. (1994). Due to uncertainties inherent in the calibration procedure of the AM form of the dye (O’Neill et al., 1990), two scales (ratio and concentration) were always given.

A shutter (UniBlitz D122 shutter driver, USA) was placed between the Xenon lamp and the objective of the microscope. The shutter was opened for 1.5 s every 4 s in order to reduce photobleaching of the Ca²⁺-indicator. Cells were field stimulated by platinum electrodes every 4 s (White et al., 1993).

Stretching cells and the measurement of sarcomere length

In order to attach and stretch cardiac cells, we used carbon fibres (Le Guennec et al., 1990; White et al., 1995). We attached these fibres close to the ends, and perpendicular to the long axis of the cells. A stiff fibre, with a compliance of 0.3 μm/µN was attached to one end of the cell and a more supple fibre with a compliance of approximately 50 μm/µN was attached to the other end of the cell. After attachment, the cell under study was lifted to avoid friction between the cell and the coverglass. The compliance of each carbon fibre was determined using a force transducer (Ackers AE801) (Le Guennec et al., 1990). During a twitch, the supple fibre moved and active tension was calculated from the displacement. For each cell, we first measured the maximum shortening during a twitch under isotonic conditions to be sure that, after attachment to carbon fibres, auxotonic contractions are not limited by restoring forces (see White et al., 1995). Cells were stretched by manually displacing the stiff fibre along the longitudinal axis of the cell.
induced changes in cell and SL of varying importance depending on the stiffness of the individual cell. For each cell, we measured its width. Force was normalized to the cross-section area assuming a cell height of 16 μm (Le Guenne et al., 1990) since it is a quite conserved value (see Fabiato, 1981). When we compared the width of the different groups, we did not find any differences. This allows us to make the assumption that the morphology of the cross section of the cells is the same in the different groups. Cells were positioned with a remotely controlled rotatable experimental chamber (Kohl et al., 1994). Sarcomere length was determined along the cell by using a fast Fourier transform algorithm on an acquired video image of the cell (Gannier et al., 1993). The sampling frequency of video image acquisition (50 Hz) is fast enough to have a precise idea of the peak tension during a twitch. Resting tension was determined, assuming no tension at slack sarcomere length.

Statistical analysis

To compare mean results obtained from the two populations of cells, the non-parametric Mann–Whitney U test and unpaired Student t-test were used (both tests gave the same results). For correlation, we used the Bravais Pearson test. For all tests, the level of significance was set at 1%. Results are expressed as mean ± standard deviation (n = number of cells studied).

Results

Description of two groups of cells on the basis of their passive properties

A first batch of 43 cells was used to study the effects of stretch on passive and active tension. Then the passive properties of a second batch of 84 cells were studied to detect any localization of the two kind of cells (see below). For each cell, we measured the SL at slack length and after stretching to different lengths. Resting tension was also measured in these conditions assuming no resting tension at slack length. The results are shown in Figure 1 and revealed that two kinds of cells can be observed: (1) cells in which stretch induced cell and sarcomere lengthening was associated with an increase in resting tension. These cells are termed compliant cells; (2) cells in which stretch induced a large increase in resting tension associated with less cell and sarcomere lengthening. We term these cells stiff or rigid cells. A stiffness slope of 1.7 nN/μm² was set as the threshold between stiff and compliant cells (Gannier et al., 1994). There were 21 stiff cells and 22 compliant cells obtained from the whole ventricles. As reported by Gannier et al. (1994), the resting calcium and, above all, the slack SL (17 ± 19 nm, 1830 ± 30 nm and 24 ± 47 nm, 1850 ± 40 nm for stiff and compliant cells, respectively) were not significantly different in the two cell groups, excluding a high stiffness resulting from calcium overload of some cells. Also, the calcium transients evoked by field stimulation of the cells were not different in the two cell groups: the calcium transient amplitude (227 ± 67 nm and 194 ± 59 nm for stiff and compliant cells, respectively) and shape (calcium transient duration measured at 50% relaxation 780 ± 240 ms and 775 ± 168 ms for stiff and compliant cells, respectively) were not significantly different. Moreover, these transients were insensitive to stretch in the two cell populations (Fig. 2) as already described (White et al., 1995). It is worth noting that both kinds of cells were found for the same cell isolation. We never observed just one kind of cell per isolation procedure. To test if there is a spatial localization of the different kind of cells in the heart, we measured the resting tension, at different lengths, of cells obtained from the left or right ventricle, from the

![Figure 1](image-url) Distinction of different populations of cells based upon their passive stiffness. Resting tension is plotted $v$ sarcomere length after different stretches for all studied cells ($n = 43$). Compliant cells (closed circles, $n = 22$) are distinguishable in this representation from stiff cells (open circles, $n = 21$) by the slope of the length–tension curve. Two significant correlations were found with slopes of $2.3 ± 0.2$ and $0.74 ± 0.05$ nN/μm² for stiff and compliant cells, respectively. Lines represent the linear regression for each group of cells. Correlation coefficients are 0.82 and 0.87 for stiff and compliant cells, respectively.
Table 1 Repartition of compliant and stiff cells in different parts of the heart. The percentage of cells, classified as compliant and stiff, found in the part of the heart given in the first column is given in the second and third columns, respectively. Bracketed numbers are numbers of cells found in the given category out of the total of studied cells. A stiffness slope of 1.7 nN/μm³ was determined to distinguish both kind of cells. The cells given in this table do not include the 43 cells of the rest of the study.

<table>
<thead>
<tr>
<th>Part</th>
<th>Compliant</th>
<th>Stiff</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left</td>
<td>40% (17/42)</td>
<td>60% (25/42)</td>
</tr>
<tr>
<td>Right</td>
<td>40% (17/43)</td>
<td>60% (26/43)</td>
</tr>
<tr>
<td>Apex</td>
<td>44% (8/18)</td>
<td>56% (10/18)</td>
</tr>
<tr>
<td>Base</td>
<td>33% (9/27)</td>
<td>67% (18/27)</td>
</tr>
<tr>
<td>Endocardial</td>
<td>20% (3/16)</td>
<td>80% (13/16)</td>
</tr>
<tr>
<td>Epicardial</td>
<td>60% (14/24)</td>
<td>40% (10/24)</td>
</tr>
</tbody>
</table>

Apex or base of the ventricles, from the endocardium or epicardium of both ventricles. The results are summarized in Table 1. It is clear on this table that there is a tendency for stiff cells to be localized in the endocardium and for compliant cells in the epicardium (see Discussion).

Active properties of the two groups of cells with stretch

Stretching compliant cells resulted in an increase of resting tension, diastolic SL and active tension (Fig. 3). Care was taken to avoid restoring forces since cell contractions were in auxotonic conditions. For stiff cells, the behaviour was quite different, since, although a stretch induced an increase in resting tension and active tension, it was not associated with a measurable increase in diastolic SL (Fig. 4). Care was taken to ensure that cells were not attached to the coverglass so that our observations were not the consequence of lifting the cell from the bottom of the chamber. Thus, it is clear that stretch may induce a positive inotropism without any change in SL.

In order to determine if there are any factors in common in the modulation of active force by stretch for the two groups of cells, we correlated active tension with different parameters. For each cell, we plotted active tension v diastolic SL [Fig. 5(a)]. If we correlate pooled data from stiff and compliant cells, a non-significant correlation is obtained ($r = 0.37$). Correlating active tension and diastolic sarcomere length only gave significant results when the data were separated into stiff and compliant cells. The same results were obtained when plotting active tension v sarcomere shortening (not shown)

Figure 2 Calcium transients evoked by field stimulation at 0.25 Hz, obtained from a compliant cell (a) and a stiff cell (b). The transients were obtained at slack length and after two stretches inducing an increase of active tension of 135 and 172% for the stiff cell and of 150 and 176% in compliant cell. In both cases, there was no obvious modification of the calcium transient which almost perfectly superimpose in this figure.
Figure 3  Contractile properties of a compliant cell at different levels of stretch. (a) Evolution of tension (upper trace) and of sarcomere length (lower trace) before and after two stretches (S1 and S2). The cell was electrically stimulated at 0.25 Hz. (b) Examples of individual contractions after subtraction of the resting tension on an expanded time scale before and after the two stretches (L1 and L2).
Figure 4  Contractile properties of a stiff cell at different levels of stretch. Labels and experimental protocol have the same meaning as in Figure 3.
Resting tension modulates active tension—surprisingly and interestingly, in stiff cells, we observed that it may induce an increase in active tension without a measurable increase in SL.

Concerning the two populations of cells, we can argue that they are physiological and not artefacts: both kinds of cell were observed in the same cell isolation; the resting calcium and the slack SL are also identical. Concerning the calcium, it is known that the absolute quantification is difficult. These difficulties were also strengthened by the low stimulation frequency (0.25 Hz) inducing a lower resting calcium level which was more difficult to measure. This is the reason why we also measured slack SL. This parameter is very sensitive to changes in intracellular calcium, pH, ATP, etc. occurring during isolation procedures. Thus, it is a very sensitive parameter to the physiological integrity of the cells.

When we observed the distribution of both kinds of cell among the ventricles, we observed that right and left ventricles are constituted of 60% stiff cells and 40% compliant cells. The same proportions were broadly observed in the apex and the base of the ventricle. In the endocardium, we found mainly stiff cells (80%) while the epicardium is the only part where we found mainly compliant cells (60%). However, it must be underlined that the epicardium is very thin, which made it very difficult to study. In our conditions, we probably studied a mixture of cells which contained more epicardial cells than in the whole ventricle, but not exclusively. If so, then there is a tendency for compliant cells to be localized in the epicardium. In fact, it is likely that from the endocardium to epicardium there is a progressive decrease of the ratio stiff/compliant cells. During blood filling, the endocardium is less stretched than is the epicardium (MacKenna et al., 1996). Thus, it is not surprising that endocardial cells are mainly stiff and epicardial cells are mainly compliant to follow changes in ventricular shape during diastolic filling. To support our hypothesis, Kang et al. (1996) found that, in bovine hearts, the endocardium is stiffer than is the epicardium in the low strain range. Even if in their case the collagen may play an important role, the difference of stiffness in the heart may have a partial intracellular origin. It must be noted that two such populations have not been found in previous studies on cell stiffness. Some hypothesis may be proposed: (1) species differences. The majority of studies have been performed on rat cells while we worked on guinea pig cells. (2) Cell preparation differences. Studies on cell stiffness have been mainly performed on skinned cells. It is possible that the treatment to skin the cells changes the elastic properties of these cells.

Figure 5  Effects of stretch on the active tension of stiff and compliant cells. (a) Plot of active tension vs diastolic sarcomere length (SL). The stiff cells (open circles) develop an active tension superior to that of compliant cells (closed circles) for a similar diastolic SL. Correlating SL and active tension for the total data was not significant, while significant correlations for SL and active tension were found for the stiff cells ($r = 0.73$, slope $= 2.5 \pm 0.3 \text{nN/\mu m}^2$) and compliant cells ($r = 0.75$, slope $= 1.0 \pm 0.1 \text{nN/\mu m}^2$) separately. (b) Active tension is plotted vs resting tension for all cells ($n = 43$). In this case, combined or separated correlations gave the same values ($r = 0.8$). The correlations for two groups of data were not significantly different from that of combined data.

Discussion

In this study, we observed two important things: first, there are two kinds of cell which can be distinguished on the basis of their stiffness; second, putting a strain on compliant cells induced an increase in resting tension, diastolic SL and active tension—surprisingly and interestingly, in stiff cells, we observed that it may induce an increase in active tension without a measurable increase in SL.
To explain the difference in stiffness between the two populations of cells, at least two main hypotheses may be proposed: (1) differences in the constitution or proportion of intracellular proteins involved in the resting tension. Indeed, it is now established that titin, a giant protein link between Z and M lines of the sarcomeres, is mainly responsible for the high resting tension observed in cardiac preparations (Granzier and Irving, 1995). We can predict that different isoforms of this protein exist. (2) It is established that weak cross-bridges exist at rest (Solott et al., 1996). Some differences in the number of weak cross-bridges may be present at rest.

Concerning the active properties, there is evidence that cell length may modulate active force in cardiac cells and in skeletal muscle by changing the interfilament spacing (Cecchi and Bagni, 1994; McDonald and Moss, 1995; Wang and Fuchs, 1995; Fuchs and Wang, 1996). Our experimental results cannot be explained on this basis, since the height of the cells did not change very much after stretch because cell length did not. Something else is needed to explain our observations. The auxotonic conditions cannot explain the inotropism of stretch when diastolic SL did not change. Whether cells are in auxotonic or isometric conditions, if diastolic SL and calcium transient are unchanged, the peak of active tension cannot change since it is the lengthening of diastolic SL which is responsible for the inotropism. Our results can be accounted for if resting tension is implied in the modulation of active tension, since, presumably, it is the only measured parameter which changed. In particular, the absence of changes of SL in some stiff cells excludes it as the determinant factor in the relationship between resting and active tension. Indeed, we found that when plotting active tension vs passive tension, the two cell groups could not be distinguished. The auxotonic conditions render the quantification of the relation between force and SL impossible.

To explain the relationship between active and passive tension, we can work on the two hypotheses we proposed above on the origin of the difference of stiffness between the two groups of cells:

1. Titin, the protein mainly involved in the resting tension, may interact with thin filaments. This interaction may modulate the velocity of actin sliding on myosin in vitro (Li et al., 1995; Kellermayer and Granzier, 1996) and in turn contraction. Titin may be the link between the importance of resting tension and changes in TnC affinity for Ca$^{2+}$. Since titin is different in skeletal and cardiac muscle, this may explain why the affinity to Ca$^{2+}$ of cardiac TnC varies with length in cardiac muscle (Babu et al., 1988) while it does not vary in skeletal muscle (Moss et al., 1991).

2. It is known that the force developed during a twitch is modulated by the cross-bridge interactions and indicates a co-operativity in the cross-bridge formation. If we assume that more weak cross-bridges are present in stiff cells and are responsible for the difference in stiffness, we can imagine that the force applied to these bridges may increase the co-operativity and consequently the force developed during a twitch. Our preliminary experiments using 2,3-Butanedione monoxime, an actin-myosin interaction inhibitor, seem to indicate that resting tension of stiff cells does not involve weak cross-bridges.

In conclusion, in this study we confirm the presence of two populations of cells in the guinea-pig ventricles. Stiff cells seem to be mainly localized in the endocardium and compliant cells in the epicardium. We show that stretching intact isolated cardiac cells induces an increase in active force. Even if a relationship exists between SL (and thus interfilament spacing) and active force, it seems that the primary effect of stretch is to increase resting tension which in turn modulates this positive inotropism. This is clear in experiments on stiff cells where stretch is able to increase the force during a twitch without measurable diastolic sarcomere lengthening.

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