The Conformation of Myosin Heads in Relaxed Skeletal Muscle: Implications for Myosin-Based Regulation

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ABSTRACT In isolated thick filaments from many types of muscle, the two head domains of each myosin molecule are folded back against the filament backbone in a conformation called the interacting heads motif (IHM) in which actin interaction is inhibited. This conformation is present in resting skeletal muscle, but it is not known how exit from the IHM state is achieved during muscle activation. Here, we investigated this by measuring the in situ conformation of the light chain domain of the myosin heads in relaxed demembranated fibers from rabbit psoas muscle using fluorescence polarization from bifunctional rhodamine probes at four sites on the C-terminal lobe of the myosin regulatory light chain (RLC). The order parameter ($<P^2>$) describing probe orientation with respect to the filament axis had a roughly sigmoidal dependence on temperature in relaxing conditions, with a half-maximal change at ~19°C. Either lattice compression by 5% dextran T500 or addition of 25 μM blebbistatin decreased the transition temperature to ~14°C. Maximum entropy analysis revealed three preferred orientations of the myosin RLC region at 25°C and above, two with its long axis roughly parallel to the filament axis and one roughly perpendicular. The parallel orientations are similar to those of the so-called blocked and free heads in the IHM and are stabilized by either lattice compression or blebbistatin. In relaxed skeletal muscle at near-physiological temperature and myofilament lattice spacing, the majority of the myosin heads have their light chain domains in IHM-like conformations, with a minority in a distinct conformation with their RLC regions roughly perpendicular to the filament axis. None of these three orientation populations were present during active contraction. These results are consistent with a regulatory transition of the thick filament in skeletal muscle associated with a conformational equilibrium of the myosin heads.

INTRODUCTION

Contraction of striated muscle is initiated by calcium-induced changes in the structure of the actin-containing thin filament. Calcium ions released in the cytoplasm following electrical stimulation bind to troponin, triggering the movement of tropomyosin around the filament and uncovering the actin binding sites for the motor or head domains of myosin emerging from the overlapping thick filament (1,2). In skeletal muscle these structural changes in the thin filament are faster than the binding of myosin heads to actin, suggesting that the rate of force generation in physiological conditions may be controlled by the kinetics of structural changes in the thick filament (3,4). Thick filament-based regulatory mechanisms are best characterized in smooth muscle (5), where they replace the troponin/tropomyosin system as the primary regulatory system, and they have received much less attention in skeletal muscle.

Early x-ray diffraction studies of skeletal muscle suggested that the myosin heads in resting muscle are arranged in a quasihelical array near the surface of the thick filaments, and that this helical arrangement is lost on activation (6).

Subsequent electron microscopy (EM) studies of isolated thick filaments from invertebrate skeletal muscle identified an asymmetric arrangement of the two heads of each myosin molecule folded back against the myosin tails in the filament backbone (7,8). This conformation, called the J motif or interacting heads motif (IHM), inhibits the ATPase activity of intrinsically regulated myosin filaments (i.e., in muscles lacking the troponin/tropomyosin system) and is associated with the relaxed or OFF state of smooth muscle myosin (9). The IHM has also been observed in the C-zone of cardiac thick filaments, the region containing myosin-binding protein C (MyBP-C) (10,11), and in isolated molecules of vertebrate muscle myosin (12). No high-resolution structures are available for isolated thick filaments from vertebrate skeletal muscle, but x-ray interference studies are consistent with the presence of the IHM in resting intact muscle (3). Functionally, such a structurally inhibited state of myosin in resting skeletal muscle could explain the slow ATP turnover observed in relaxed demembranated fibers from mammalian muscle, which has been called a super-relaxed state (SRX) (13,14). However, the interaction between this potential regulatory structural switch in the thick filament and the well-known regulatory switch in the thin filaments remains obscure.

To better understand the role of myosin head conformations in the concept of thick filament regulation summarized above, we measured the orientation of the regulatory light
chain (RLC) regions of the myosin heads in the native sarcomeres of demembranated fibers from mammalian muscle. The structural relationship between the thick and thin filaments is preserved in this preparation, and RLC orientation can be measured using fluorescence polarization from bifunctional sulforhodamine (BSR) probes at four different sites on the C-terminal lobe of the RLC. X-ray diffraction studies on this preparation showed that the quasihelical organization of myosin heads on the thick filament surface in relaxing conditions depends strongly on temperature (15–18). We therefore used this protocol as a model system to investigate the transition between conformations of the myosin heads with different degrees of helical order, overcoming limitations of a previous fluorescence polarization study by using a skeletal muscle isoform of RLC rather than a smooth muscle isoform (19), and a milder exchange protocol to introduce labeled RLCs into muscle fibers. To further constrain the interpretation, we also investigated how the relationship between myosin head conformation and temperature depends on the lateral spacing between the thick and thin filaments, and on inhibition of myosin by blebbistatin (20).

**MATERIALS AND METHODS**

**Muscle fiber preparation**

–6 mm long segments of demembranated fibers were dissected from rabbit psoas muscle as previously described (21). Fibers were mounted in the experimental trough between a force transducer and a loudspeaker motor, and the extremities were fixed using 5% glutaraldehyde in rigor solution and glued to comeres of demembranated fibers from mammalian muscle.

**Preparation of BSR-RLCs**

Four double-cysteine mutants of chicken skeletal RLC (D95C/V103C (E helix), E131C/A138C (G helix), K151C/T158C (H helix), and T122C/K134C (linking F and G helices)) were obtained by site-directed mutagenesis, expressed in *Escherichia coli*, and purified as previously described (19,22) (Fig. S1 in the Supporting Material). The two native cysteines at position 125 and 154 were replaced by alanine to avoid nonspecific labeling. Each pair of introduced cysteines was cross-linked with BSR (B10621, Invitrogen, Waltham, MA) to give 1:1 BSR-RLC conjugates that were purified by reverse-phase high-performance liquid chromatography to >95% homogeneity. Specificity and stoichiometry of BSR labeling were determined by reverse-phase high-performance liquid chromatography and mass spectrometry.

**Exchange of BSR-RLC mutants into muscle fibers**

Muscle fibers were first transferred to rigor solution at –1°C, and then RLC-extracting solution (20 mM EDTA, 50 mM KPr, 10 mM potassium phosphate buffer, pH 7.1) at 1°C for 30 min, and finally to RLC-extracting solution containing 0.5 mg/ml (~25 μM) BSR-RLC at 19°C for 30 min. Fibers were washed in relaxing solution and then incubated at 10°C for 20 min in relaxing solution containing 0.5 mg/ml of rabbit skeletal muscle troponin complex (Tn, Life Diagnostics, West Chester, PA) and for a further 20 min in relaxing solution containing 0.5 mg/ml of wild-type chicken skeletal muscle troponin C (TnC), expressed in E. coli, and purified as previously described (23), to replace Tn/TnC lost during the RLC exchange. Before and after the exchange protocol the fiber was activated twice at 11°C by temperature jump (21). A ramp-release (4% of the initial fiber length (L0) complete in 3 ms) and a step-release (0.2% L0 complete in 0.12 ms) after the force redevelopment following unloaded shortening were applied at the plateau of the isometric contraction to check the mechanical performance of the RLC-exchanged fiber (Fig. S2). The concentration of BSR-RLC in the fiber (C_{BSR-RLC}), calculated from fluorescence intensity (4), was 39 ± 10 μM (mean ± SD, n = 6 fibers) and therefore the exchanged fraction was 26% assuming that the concentration of native RLC in demembranated fibers from rabbit psoas muscle is 150 μM (24). Maximum isometric force was 224 ± 23 kPa at 11°C and force recovery after RLC exchange was 95 ± 4% (mean ± SD, n = 12 fibers). Samples for confocal microscopy (Fig. S3) were prepared by stretching the relaxed fiber to a sarcomere length of 3.0 μm, and then fixing in 4% formaldehyde at room temperature for 5 min. The fiber was then incubated in antmyosin antibody (A4.1025, kindly provided by Prof. Mathias Gautel) for 1 h at room temperature and in secondary antibody labeled with Alexa Fluor 488 for 1 h at room temperature, and finally transferred to a microscope slide in a mounting medium. Images were acquired using a Zeiss LSM 510 Meta (Zeiss, Jena, Germany).

**Experimental protocol and data analysis**

The temperature of the experimental trough was increased in steps of 3°C in the range 2.5–33.0°C, and the polarized fluorescence intensities from the four BSR-RLCs were recorded in relaxing conditions at each temperature using a custom-built setup (4) and used to calculate the second- and fourth-rank order parameters of the orientation distribution of the BSR dipole, (P2) and (P4), respectively (25). The order parameter (P2) quantifying rapid probe motion was calculated from the polarized intensities recorded with X- and Y- illumination (in line with and perpendicular to the measured fluorescence emission, respectively (4)). In relaxing conditions (P2) was 0.867 ± 0.004 for the E helix probe, 0.849 ± 0.018 for the G

**TABLE 1 Composition of the Physiological Solutions**

<table>
<thead>
<tr>
<th>Solution</th>
<th>Imidazole</th>
<th>EGTA</th>
<th>Ca-EGTA</th>
<th>HDTA</th>
<th>KPr</th>
<th>MgAc2</th>
<th>Na2ATP</th>
<th>Na2CP</th>
<th>pCa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relaxing</td>
<td>25</td>
<td>10</td>
<td>–</td>
<td>–</td>
<td>11.8</td>
<td>6.89</td>
<td>5.56</td>
<td>20</td>
<td>9</td>
</tr>
<tr>
<td>Preactivating</td>
<td>25</td>
<td>0.1</td>
<td>–</td>
<td>9.9</td>
<td>11.7</td>
<td>6.48</td>
<td>5.56</td>
<td>20</td>
<td>9</td>
</tr>
<tr>
<td>Activating</td>
<td>25</td>
<td>–</td>
<td>10</td>
<td>–</td>
<td>11.7</td>
<td>6.39</td>
<td>5.65</td>
<td>20</td>
<td>4.7</td>
</tr>
<tr>
<td>Rigor</td>
<td>25</td>
<td>10</td>
<td>–</td>
<td>–</td>
<td>100.3</td>
<td>1.50</td>
<td>–</td>
<td>–</td>
<td>9</td>
</tr>
</tbody>
</table>

All the concentrations are in mM. EGTA, HDTA, 1,6 diaminohexane-N,N,N’,N’-tetraacetic acid; KPr, potassium propionate; MgAc2, magnesium acetate; ATP, Na2CP, phosphocreatine disodium salt; pCa = –log[Ca2+]. Mg2+-ATP concentration is 5 mM and free Mg2+ is 1 mM.

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helix probe, 0.760 ± 0.009 for H helix probe, and 0.827 ± 0.017 for the FG probe (mean ± S.D., n = 3 fibers), indicating that the amplitude of such motion is small. \( \langle P_{30} \rangle \) was not sensitive to temperature in the range studied here. The pH of the solutions was lower at higher temperature, but order parameters measured at 25°C in a relaxing solution with pH set to 7.1 at 11°C were not significantly different from those with pH set to 7.1 at 25°C, indicating that the probe orientation in relaxing solution is not sensitive to temperature-induced changes in pH. The effect of dextran on the temperature dependence of \( \langle P_{30} \rangle \) and \( \langle P_{4} \rangle \) in relaxing conditions was investigated in three independent experiments for each RLC probe \((n = 12 \text{ fibers})\) (see Fig. 2). The effect of blebbistatin in the presence and in the absence of dextran on the relaxed values of \( \langle P_{30} \rangle \) and \( \langle P_{4} \rangle \) was tested in one experiment for each mutant \((n = 4)\). The errors shown for relaxing conditions (see Fig. 2) are pooled standard deviations for all the data for each probe in the presence and absence of dextran; the original data for each temperature and condition without error pooling are shown in Fig. S4.

To describe the in situ RLC-orientation, we defined a reference frame (the EG frame) using the crystallographic structure of nucleotide-free myosin S1 from chicken skeletal muscle (see Fig. 1 (27)). The orientation of the RLC domain is described by \( \beta \), the angle between the E helix (see Fig. 1, magenta) and the thick-filament axis, and \( \gamma \), the twist angle around the E helix. \( \gamma = 0 \) when the G helix (see Fig. 1, orange) is in the plane defined by the E helix and the thick-filament axis, and \( \gamma > 0 \) for a counterclockwise rotation of the lobe viewed from the + end of the E helix. The orientation of each BSR probe in this frame was described by two angles calculated from the coordinates of the \( \beta \) carbons of the two attachment points, assuming that the dipole axis is parallel to the line joining those points. \( \theta \) is the angle between the dipole and the E helix, and \( \phi \) is the angle between the plane defined by the E helix and the dipole and that defined by the E and G helices (Table S1; Fig. S1). These angular coordinates and the measured order parameters for the four probes were used in a maximum entropy (ME) algorithm (28, 29) to calculate the smoothest distribution of RLC-orientations consistent with the data. The two-dimensional contour map denoting the probability of each \((\beta, \gamma)\) orientation (with \( 0^\circ < \beta < 180^\circ \) and \( 0^\circ < \gamma < 360^\circ \) ) was projected onto the surface of a sphere using Origin software (OriginLab, Northampton, MA), in which \( \beta \) is the polar angle (latitude) and \( \gamma \) the azimuthal angle (longitude). This representation has the advantage of avoiding distortion of the distribution introduced in a Cartesian representation and of showing only one of the two equivalent solutions \((\beta, \gamma)\) and \((180^\circ - \beta, 180^\circ + \gamma)\) associated with the dipole ambiguity and the antiparallel arrangement of myosin motors in the two halves of the thick filament (30). The methods used for integration of specific regions of the ME maps are discussed in the Supporting Material.

RESULTS

Temperature dependence of the order parameters of RLC probes in relaxed muscle fibers

Mutants of chicken skeletal RLC, with native cysteines replaced by alanine and new cysteine pairs introduced in its C-lobe at positions 95–103 (E helix), 131–138 (G helix), 151–158 (H helix), or 122–134 (in the F and G helices, respectively), were labeled by cross-linking cysteine pairs with BSR (Fig. 1; Fig. S1; Movie S1). The E helix probe was designed with its dipole axis approximately parallel to the myosin lever arm to be sensitive to lever arm tilting. Each BSR-RLC was exchanged into permeabilized fibers from rabbit psoas muscle using a mild exchange protocol to better preserve the native structure and function of the thick filaments (see Materials and Methods section). This protocol led to replacement of ~26% of the native RLC by BSR-RLC, which was uniformly distributed along the A-band of the sarcomere and across the fiber width (Fig. S3). Moreover, the function of myosin motors was not affected by RLC exchange: isometric force recovery after exchange was >95%, and both the rate of force redevelopment following unloaded shortening and the force transient elicited by a 0.2% step release in fiber length were similar to those before exchange (Fig. S2).

The polarized fluorescence from the four RLC probes was measured in the temperature range 2.5–33.0°C in relaxing conditions at maximum myofilament overlap (sarcomere length 2.40 µm) and used to calculate the second- and fourth-rank order parameters of the orientation distribution of each BSR-RLC in the fiber, \( \langle P_{30} \rangle \) and \( \langle P_{4} \rangle \), respectively. \( \langle P_{2} \rangle \) is a measure of how parallel the probe dipole is to the filament axis, and varies from +1 for parallel to −1
for perpendicular. \( \langle P_4 \rangle \) is a higher-order harmonic term, giving higher resolution angular information (25).

Increasing the temperature in relaxing solution induced reproducible changes in \( \langle P_2 \rangle \) for each probe (Fig. 2, solid line and solid circles), with smaller changes in \( \langle P_4 \rangle \) (Fig. S4). The temperature dependence of \( \langle P_2 \rangle \) for each probe had a roughly sigmoidal shape with a transition temperature (the temperature at which the change in \( \langle P_2 \rangle \) is half maximal) of \( \sim 19^\circ \text{C} \). The largest change in orientation was observed for the E helix, with \( \langle P_2 \rangle \) becoming more positive at high temperature, indicating that the E helix became more parallel to the filament axis (at high temperature, similar to the effect of osmotic compression by 5% dextran). The effects of blebbistatin and dextran were additive; in the presence of both 25 \( \mu \text{M} \) blebbistatin and 5% dextran T500 the transition temperature was reduced by \( \sim 10^\circ \text{C} \) (Fig. 2, dot-dashed line and open diamonds).

**RLC-orientation distribution in relaxed muscle fibers**

The orientation of RLC with respect to the filament or fiber axis was described by two angular coordinates \( (\beta, \gamma) \) using a reference coordinate frame defined by the E and G helices (magenta and orange helices, respectively, in Fig. 1). \( \beta \) is the angle between the E helix and the filament axis, whereas \( \gamma \) is the rotation of the RLC around the E helix using the G helix to define \( \gamma = 0 \) (Figs. 1 and S6). The in situ orientation distribution of the RLC in this EG frame was calculated using a ME algorithm (28), a model-free method that determines the smoothest probability distribution of orientations that reproduces the experimental \( \langle P_2 \rangle \) and \( \langle P_4 \rangle \) values for each probe (Figs. 2 and S4), given the angular coordinates of the probe dipoles in the EG frame (Table S1). These probability distributions were plotted on a sphere (Fig. 3), where \( \beta \) is the latitude and \( \gamma \) is the longitude (\( \beta = 90^\circ \) at the equator and \( \gamma = 0^\circ \) at the meridian), as contour maps with hotter colors denoting more likely orientations.

![FIGURE 2 Temperature dependence of the \( \langle P_2 \rangle \) order parameter for the RLC probes. Order parameter \( \langle P_2 \rangle \) (mean \( \pm \) pooled SD) of E (A), G (B), H (C), and FG (D) probes in the temperature range 2.5–33.0°C in standard relaxing solution (solid circles and solid line, \( n = 3 \) fibers), in the presence of 5% dextran T-500 (open circles and dashed line, \( n = 3 \) fibers) or 25 \( \mu \text{M} \) blebbistatin (solid diamonds and dotted line, \( n = 1 \) fiber), and in the presence of both 5% dextran and 25 \( \mu \text{M} \) blebbistatin (open diamonds and dashed/dotted line, \( n = 1 \) fiber). \( \langle P_2 \rangle \) values during active contraction and in rigor at 11.0°C and 24.8°C are shown as stars (\( n = 5 \) fibers) and squares (\( n = 3 \) fibers), respectively (mean \( \pm \) SD).](https://www.biophysicsjournal.org/content/109/4/783.full.pdf)
The in situ distribution of RLC orientations at 2.5°C in standard relaxing solution (in the absence of dextran and blebbistatin) showed a major equatorial population at $\beta = 90^\circ$, $\gamma = -90^\circ$ with $\sim 20^\circ$ angular dispersion in $\beta$ and $\gamma$ (RX1, Fig. 3 A). At 11°C two minor populations appeared, one close to the pole (RX2, $\beta = 165^\circ$, $\gamma = -70^\circ$) and one close to the equator ($\beta = 85^\circ$, $\gamma = -15^\circ$) (Fig. 3 B). RX2 increased progressively at higher temperatures with a consequent depletion of RX1, and became dominant above 19°C (Fig. 3, C–E; Movie S2). As the temperature was increased above 19°C, the minor equatorial population with $\gamma$ near 0° became more parallel to the filament axis ($\beta = 130^\circ$, $\gamma = 0^\circ$, Fig. 3 D) and more intense (Fig. 3, D and E); this orientation is denoted RX3. Addition of 5% dextran at 11°C increased the intensity of RX2 and decreased that of RX1, so that the two peaks were roughly equal in intensity in these conditions (Fig. 3 G). Addition of 25 μM blebbistatin at 11°C increased the intensity of RX2 and RX3 (Fig. 3 H), so that the overall orientation distribution was similar to that in normal relaxing solution at 25°C (Fig. 3 D).

The hypothesis that the RX2 and RX3 peaks observed at higher temperatures in situ are similar to the conformation of the myosin heads in isolated thick filaments from tarantula muscle inferred from EM studies (3DTP (8), Fig. 3 F) was tested by calculating the ($\beta$, $\gamma$) orientation of the two RLCs in that structure. The orientations of the RLCs of the free and blocked heads of the IHM, shown as yellow ($\beta = 158^\circ$, $\gamma = -60^\circ$) and red ($\beta = 131^\circ$, $\gamma = 0^\circ$) spheres, respectively, in the spherical plots of Fig. 3, are similar to the orientation distribution of the two polar lobes (Table S2, Fig. S6). The RLC orientation of the free head (yellow) is close to RX2 ($\beta = 165^\circ$, $\gamma = -70^\circ$), whereas that of the blocked head is close to RX3 ($\beta = 130^\circ$, $\gamma = 0^\circ$), becoming more prominent at higher temperatures (Fig. 3 E).

The fraction of myosin heads with their RLC regions in either the RX2 or RX3 orientation ($F_{RX(2+3)}$) was estimated by integrating over the corresponding orientation region (see Fig. S8 for details) and normalizing by the total intensity at each temperature (Supporting Material). $F_{RX(2+3)}$ had a roughly sigmoidal dependence on temperature (Fig. 4, solid circles), which is similar to that of $\langle P_2 \rangle$ for the E helix probe (Fig. 2). At 2.5°C $\sim 35\%$ of the light chain domains are in the RX2 or RX3 orientation. $F_{RX(2+3)}$ increased with temperature and reached a maximum of $\sim 70\%$ at 25°C. At each temperature below 25°C the addition of dextran and/or blebbistatin induced an increase in $F_{RX(2+3)}$ (Fig. 4, diamonds and open circles), but did not significantly increase the limiting value of $F_{RX(2+3)}$ at 25°C and above.

**Orientation distribution of the RLC region during active isometric contraction and in rigor**

Fibers were calcium activated at 11.0°C or 24.8°C at fixed length using a temperature-jump protocol (21) in the absence of dextran. Active isometric force was 46 ± 4% (mean ± SE, n = 17) higher during activation at 24.8°C than at 11.0°C. In general, the order parameters during active contraction were distinct from those during relaxation (Figs. 2 and S4, stars). In the case of $\langle P_2 \rangle$ for the E helix probe the difference was small at 11.0°C, but about $-0.2$ units at 24.8°C, indicating that the E helix, and therefore the long axis of the lever arm, becomes significantly less parallel to the filament axis on activation at this temperature.

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**FIGURE 3** Orientation distributions of the RLC region of myosin. ME contour maps of the probability distribution of RLC orientations in skeletal muscle fibers shown as spherical plots, in which $\beta$ is the latitude and $\gamma$ is the longitude; $\beta = 90^\circ$ at the equator and $\gamma = 0^\circ$ at the meridian (black lines). (A–E) Standard relaxing solution in the temperature range 2.5–33.0°C (F) IHM structure (3DTP (8)) with the RLCs of the blocked and free heads in red and yellow, respectively, (G) relaxing solution with 5% dextran at 11°C (H) relaxing solution with 5% dextran + 25 μM blebbistatin at 11°C, (I) active contraction, and (J) rigor at 24.8°C. The RLC orientations for the blocked ($\beta = 131^\circ$, $\gamma = 0^\circ$), and free head ($\beta = 158^\circ$, $\gamma = -60^\circ$) in the EG frame are shown as red and yellow spheres, respectively, in the ME plots. The errors in the ME maps are analyzed in Fig. S9.
The orientation distribution of the RLC region of the myosin heads during active isometric contraction at 11.0°C showed a broad peak centered on (\(\beta = 110^\circ\), \(\gamma = -55^\circ\)) with \(\pm 30^\circ\) dispersion in \(\beta\) and \(\gamma\), respectively (Fig. S7). The corresponding orientation distribution at 24.8°C was also broad, and centered on (\(\beta = 115^\circ\), \(\gamma = -50^\circ\)) (Figs. 3 I and S7). The RX1, RX2, and RX3 orientations observed in relaxing conditions at 11.0°C and 24.8°C (Fig. 3, B and D) disappeared on activation.

In rigor conditions the order parameters of the E, FG, and H probes decreased from their relaxed values, and were insensitive to a temperature increase from 11.0°C to 24.8°C (Fig. 2, squares). \(\langle P_2 \rangle\) for the E helix decreased by \(-0.2\) units at 11.0°C and by \(-0.5\) units at 24.8°C, indicating that the E helix axis and the myosin lever arm becomes much more perpendicular to the filament axis in rigor. The ME orientation distribution of the RLC region in rigor (Fig. 3 J) consisted of a single broad peak centered on (\(\beta = 90^\circ\), \(\gamma = -80^\circ\)) with a larger dispersion in \(\gamma\). The center of the rigor peak is close to the RX1 orientation seen in relaxing conditions.

**DISCUSSION**

**Conformational equilibrium of myosin heads in relaxed muscle fibers**

We used fluorescence polarization from four bifunctional probes on the C-lobe of the RLC to investigate the in situ conformation of the myosin heads in mammalian skeletal muscle. First, we characterized the conformational changes in the RLC region of the myosin heads during the temperature-induced structural transition in the thick filament in relaxing conditions (15–18). A temperature increase from 2.5°C to 33.0°C induced changes in the order parameters describing the orientation of each probe with respect to the filament or fiber axis (Fig. 2). The largest change was observed for the E helix probe dipole, which is almost parallel to the myosin lever arm: \(\langle P_2 \rangle\) of the E helix increased by \(-0.4\) units as the temperature was increased, suggesting a large tilting of the probe, and therefore of the lever arm, toward an orientation more parallel to the filament. This is the first study, to our knowledge, in which such large temperature-induced changes of RLC orientation have been observed in relaxing conditions. A previous electron paramagnetic resonance spectroscopy study using RLC probes revealed temperature-dependent changes in the orientation and mobility of the probes in relaxing conditions, but did not provide a detailed description of the changes in RLC orientation (33). A previous fluorescence polarization study using bifunctional probes labeling the C-lobe of a smooth muscle isoform of RLC in skeletal muscle fibers showed very small effects of temperature in relaxing conditions (19). Here, we overcame some limitations of that study by using a skeletal muscle isoform of RLC rather than a smooth muscle isoform, and a milder RLC exchange protocol that replaces only \(-25\%\) of the native RLC. The mechanical properties of exchanged fibers were unaltered by this RLC exchange protocol (Fig. S2) and the large signals from the probes associated with temperature changes in relaxing conditions (Fig. 2) suggest that structure of the thick filaments was also preserved.

The results presented here reveal a temperature-dependent conformational equilibrium between the three main RLC orientations (RX1, RX2, and RX3) in relaxed muscle fibers. At 2.5°C, the most prominent RLC orientation has the E helix (and the myosin lever arm) roughly perpendicular to the filament axis (RX1, \(\beta = 90^\circ\); Fig. 3), similar to the orientation observed for actin-bound heads in rigor (Fig. 3 J). At 25°C, in contrast, the prominent orientations have the E helix much more parallel to the filament axis (RX2, \(\beta = 170^\circ\) and RX3, \(\beta = 130^\circ\); Fig. 3). The RX2 and RX3 orientations are similar to those of the free and blocked heads, respectively, in the IHM observed in isolated thick filaments from invertebrate muscle (8). These results indicate that the majority of the myosin heads in relaxed muscle fibers at near-physiological temperature have their RLC regions in conformations similar to those in the IHM. The limitations of the technique presented here and the ME analysis, and of fitting crystal structures into EM electron densities, means that these conformations are not necessarily identical. Moreover, the results presented here indicate a higher fraction of RX2 (free-like) than RX3 (blocked-like) orientations at the higher temperatures, whereas these two fractions would by definition be equal.

![FIGURE 4 Temperature dependence of the fraction of myosin heads in the RX2 and RX3 orientations in relaxed muscle fibers.](image-url)
for the IHM. Although ME analysis cannot give a precise measure of the relative probabilities of two closely-spaced orientations (28), the relatively large difference between RX2 and RX3 intensities suggests the presence in relaxed muscle fibers of some myosin molecules in a novel conformation, with one parallel (RX2- or free-like) and one perpendicular (RX1-like) light chain domain.

The sigmoidal temperature dependence of the fraction of parallel (RX2 or RX3) orientations (Fig. 4) is similar to that of the intensity of the first myosin layer line observed in x-ray studies (16). Moreover, the temperature-dependent increase in the fraction of RX2 or RX3 orientations is similar to that in the fraction of helically ordered heads estimated from the change in myosin layer-line intensities (17,34,35). Although the latter depend on other parameters of myosin head orientation, including their radial and azimuthal position and the conformation of the catalytic domain of the heads that cannot be detected by dipole probes on the RLC, there is a close correlation between a helically ordered conformation of the myosin heads and a parallel orientation of their light chain domains. Both of these sets of structural parameters are also correlated at least qualitatively with the formation of the IHM at higher temperature, which in turn is correlated with the increase in the fraction of heads in the so-called SRX state characterized by its slow ATP-turnover. The SRX state is estimated to be occupied by ~75% of the heads at physiological temperature in demembranated fibers from rabbit muscle (13,14), similar to the fraction of light chain domains in parallel (RX2 or RX3) orientations estimated here for similar conditions. Thus, the previous concept of a correlation between the IHM and SRX states is extended here by a direct correlation with the orientation of the RLC region of the myosin heads.

However, the results presented here also confirm previous evidence from relaxed muscle fibers at close to physiological temperature that ~1/3 of the myosin heads have their light chain regions in a more perpendicular conformation that is not consistent with the IHM. Modeling of x-ray data for these conditions suggested the coexistence of at least two myosin head orientations, corresponding to two radial positions with respect to the filament backbone (17), that are likely to be associated with different regions of the thick filament (36). EM-based three-dimensional reconstructions of the C-zone of vertebrate cardiac thick filaments showed two different conformations in the three axial layers of heads of each 43-nm repeat (10). Thick filaments from knockout mice lacking MyBP-C exhibit a more disordered structure (10), suggesting that some heads outside the C-zone may not form the IHM. Our results show that the RX1 myosin heads have a defined orientation of their light chain domains perpendicular to the filament axis, although these heads appear to be helically disordered in x-ray and EM studies (13,17,37). This distinction is likely to be related to the lack of sensitivity of the technique presented here to radial and azimuthal parameters, and to effects of changes in the conformation of the catalytic domains of the heads on the x-ray and EM data, as noted previously. The comparative advantage of the technique presented here is that it isolates the (β,γ) orientation of the RLC region of the myosin heads from those other parameters, and consequently shows that the helically disordered heads have RLC regions with ordered (β,γ) orientations. Our results show that the previous concept of the temperature-dependent change in myosin head conformation as a disorder-to-order transition (15–18) does not apply to the RLC region, which instead exhibits a temperature-dependent conformational equilibrium. The functional implications of this distinction are discussed below.

Factors that influence the conformational equilibrium of the RLC region of myosin

It has been shown that the formation of helical order in thick filaments requires the closed structure of the nucleotide-binding pocket of the myosin head (35,38). Therefore, the temperature dependence of the helical order has been explained on the basis that the open conformation is preferred at lower temperature, and the closed conformation at higher temperature (35). The open-closed equilibrium can also be influenced by ligands (32,35): in particular, the myosin inhibitor blebbistatin favors the closed conformation (20) and stabilizes the IHM in isolated myosin molecules (12), the helical order of the thick filament (32,39), and the SRX state of myosin (37). Our results show that blebbistatin also stabilizes the parallel orientations of RLC in relaxed muscle fibers by lowering the transition temperature of its conformational equilibrium without altering the maximum fraction of parallel RLC orientations at high temperature (Fig. 4).

We found that the filament lattice spacing also influences RLC orientation under relaxing conditions. The recovery of the filament lattice spacing typical of muscle cells with an intact surface membrane by adding 5% dextran T500 to relaxing solution (31) stabilized the parallel orientations of the RLC region in the relaxed demembranated fiber. The effect was similar to that of blebbistatin (Fig. 4). Moreover, the effects of blebbistatin and dextran on RLC orientation were additive, suggesting that both interventions alter the conformational equilibrium without altering the low- or high-temperature end states. In the presence of 5% dextran the center-to-center distance of thick and thin filaments decreases from 28 to 23 nm (31). Assuming a radius of 3.5 and 7.5 nm for thin and thick filaments, respectively, the distance between the filament surfaces decreased from 17 to 12 nm in the compressed lattice. Thus, either steric or charge effects may favor the parallel RLC orientation states if, as expected, these are associated with a lower mean radius of the myosin heads with respect to the thick filament axis. Alternatively, the effect of dextran on the
off structure of the thick filament might involve MyBP-C. The N-terminal region of MyBP-C is thought to bind to the thin filament, and therefore to be roughly perpendicular to the thick filament axis (40,41). These MyBP-C links have been shown to stabilize the resting structure of the thick filament by preserving its resting periodicity and the axial perturbations of the myosin layers in each 43-nm repeat (42). Therefore, the recovery of the physiological thick-thin filament distance by osmotic compression could promote the parallel RLC orientations by allowing the formation of MyBP-C links.

The experiments presented in this work were performed with unphosphorylated BSR-RLCs and the protocol used to prepare muscle samples yielded fibers in which the native RLC is unphosphorylated (13,43). However, under physiological conditions the conformational equilibrium of myosin could also be modulated by the phosphorylation level of RLC. It has been shown that under relaxing conditions RLC-phosphorylation can disorder the helical array of heads (13,44) and weaken the interactions between blocked and free heads (8). Therefore, RLC-phosphorylation is expected to disrupt the parallel orientation and shift the equilibrium toward the perpendicular orientation. This might be expected to bring the catalytic domains of the heads closer to the actin filaments and might therefore lead to changes in the kinetics of activation following electrical stimulation, as suggested by the finding that RLC-phosphorylation potentiates the twitch force in skeletal muscle (45).

Implications for myosin-based regulation of mammalian skeletal muscle

The results presented above indicate that under relaxing conditions at near-physiological temperature and lattice spacing, and at low RLC-phosphorylation levels, ~70% of the myosin heads have parallel RLC (RX2 and RX3) orientations, similar to those in the IHM/SRX state, with the myosins outside the C-zone in a conformation that inhibits the myosin ATPase and the interaction with the overlapping thin filament. The remaining 30% of the heads have an RLC orientation (RX1) perpendicular to the filament axis and inconsistent with the IHM state. The 70% fraction of IHM-like heads appears to be an upper limit, because the addition of a stabilizing agent such as blebbistatin does not further increase either the fraction of heads with a parallel RLC region, as shown here, or the helical order (32).

Because the IHM seems to be confined to the C-zone of the thick filaments in vertebrate muscle (10,11), it is likely that the perpendicular RX1 orientation is confined to the remainder of the filament. The C-zone occupies ~1/3 of the filament, so the simplest interpretation of the distribution of RX1, RX2, and RX3 states observed here (Figs. 3 and 4), would be that all myosins in the C-zone in resting muscle are in the IHM state, but that the remaining two-thirds of the myosins outside the C-zone have, on average, one head with a parallel (RX2-like) RLC orientation and the other with a perpendicular (RX1-like) orientation.

The RX1, RX2, and RX3 orientations disappear on activation and are replaced by a broad orientation distribution lying between them (Figs. 3 I and S7). This transition is likely to be related to the complete loss of helical order of the heads on activation as measured by x-ray diffraction (3,6,42) and electron paramagnetic resonance (46). The redistribution of RLC orientations on activation of skeletal muscle does not occur on activation of cardiac muscle (26). This difference may be related to differences in the docking of the RLC onto the thick filament surface associated with the different isoforms of myosin and myosin binding protein C and with the different regulatory mechanisms and functional requirements in the two muscle types.

The signaling pathway that triggers the change in RLC orientation and the loss of helical order of the heads on activation of vertebrate skeletal muscle is unknown. In intrinsically regulated myosins this signal is provided by RLC phosphorylation, which weakens the interactions stabilizing the IHM and can be considered to switch myosin on. However, in skeletal muscle the structure of the thick filament becomes indistinguishable from the fully active state a few tens of milliseconds after the start of electrical stimulation (3). Because the skeletal isoform of myosin light chain kinase operates on a slower time scale (47), it cannot account for the fast activation of myosin. Thick filaments must therefore be switched on by a signaling pathway that rapidly transmits the on state of the thin filament to the thick filament. MyBP-C is a possible component of this signaling pathway, given its role in the stabilization of the resting structure of the thick filament and in the sensitization to calcium of the thin filament (48). MyBP-C links could sense the structural changes in the thin filament initiated by calcium binding to the regulatory proteins and trigger the exit of the heads from the IHM in the C-zone of the thick filament. Cooperative propagation of this structural change along the thick filament would be required to switch on all the myosin heads.

Although MyBP-C links may be involved in the regulation of the structural changes associated with myosin activation, our results suggest the possibility of an additional component involving the minor population of non-IHM heads in the resting thick filament. These heads might be more available for actin binding and could potentially act as sensors of the activation level of the thin filament (14). When the thin filament is switched on, allowing strong actin-myosin interactions, these perpendicular RX1 heads might contribute to disrupting the conformation of the IHM heads. Further work will be required to clarify the role of these different populations of myosin heads, of MyBP-C, and perhaps of titin, in thick filament regulation in vertebrate skeletal muscle.
CONCLUSIONS

In relaxed demembranated muscle fibers from mammalian skeletal muscle the RLC region of the myosin heads are in a temperature-dependent orientational equilibrium. A temperature increase from 2.5 to 33.0°C induces a shift of the conformational equilibrium from more perpendicular to more parallel orientations with respect to the thick filament axis. The latter are similar to the RLC orientations in the free and blocked heads of the IHM, and are stabilized by both blebbistatin and myofilament lattice compression. Close to physiological temperature and lattice spacing the heads with parallel RLC regions coexist with a significant fraction of myosin heads with a perpendicular orientation. During active contraction all these defined RLC orientations are replaced by a broad intermediate distribution. Modulation of the conformational equilibrium of the RLC region of the myosin heads may be a component of the physiological signaling pathway of muscle activation.

SUPPORTING MATERIAL

Supporting Discussion, nine figures, two tables, and two movies are available at http://www.biophysj.org/biophysj supplemental/S0006-3495(15)00621-9.

AUTHOR CONTRIBUTION

L.F. and M.I. designed research, Z.H. contributed analytic tools, L.F. performed research and analyzed data, L.F. and M.I. wrote the article.

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