Tropomyosin flexural rigidity and single Ca$^{2+}$ regulatory unit dynamics: implications for cooperative regulation of cardiac muscle contraction and cardiomyocyte hypertrophy

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Striated muscle contraction is regulated by dynamic and cooperative interactions among Ca$^{2+}$, troponin, and tropomyosin on the thin filament. While Ca$^{2+}$ regulation has been extensively studied, little is known about the dynamics of individual regulatory units and structural changes of individual tropomyosin molecules in relation to their mechanical properties, and how these factors are altered by cardiomyopathy mutations in the Ca$^{2+}$ regulatory proteins. In this hypothesis paper, we explore how various experimental and analytical approaches could broaden our understanding of the cooperative regulation of cardiac contraction in health and disease.

Keywords: tropomyosin, thin filament, calcium activation, persistence length, cooperativity, heart, sarcomere, cardiomyopathy

Cardiac muscle contraction is regulated by a Ca$^{2+}$-dependent switch mechanism. Each basic contractile unit, or sarcomere, consists of a staggered array of thin and thick filaments. Contraction takes place when myosin heads in thick filaments engage in the cross-bridge cycle to generate isometric force or to slide adjacent thin filaments toward a sarcomere’s center, thereby shortening the sarcomere. Ca$^{2+}$ regulation of striated muscle contraction is achieved primarily by the orchestrated action of troponin (Tn) and tropomyosin (Tm) on thin filaments. Tn is a three-subunit complex consisting of TnC, TnI, and TnT, where TnC is the Ca$^{2+}$ sensor, TnI is the inhibitory subunit that holds the other two Tn subunits in a ternary complex and also binds actin, and TnT anchors the Tn complex to Tm. Tm is a dimeric, α-helical coiled-coil protein that binds along both grooves of an actin filament’s helix (O’Brien et al., 1971; Milligan et al., 1990; Perry, 2001). αTm is the predominant Tm isoform in adult human hearts. The molecular contour length ($L_c$) of αTm is about 40 nm (Smillie, 1996; Perry, 2001). All native Tm molecules polymerize into a long strand through head-to-tail overlap that involves about nine amino acids from the carboxy- and the amino-termini of adjacent molecules (McLachlan and Stewart, 1975). Systolic contraction is initiated by elevation of cytoplasmic Ca$^{2+}$ that binds to Tn, which undergoes a conformational change that induces azimuthal movement of Tm on the thin filament to uncover myosin binding sites, thereby permitting cross-bridge cycling and concomitant ATP hydrolysis.

As demonstrated by various biochemical (Lehrer et al., 1997; Maytum et al., 1999) and physiological (Metzger, 1995; Iwamoto, 1998; Dobesh et al., 2002) studies, cooperativity is a central feature of striated muscle activation and contraction. In the following discussion, unless otherwise specified, we focus on the Ca$^{2+}$-induced cooperative activation of cardiac thin filaments in which Tm plays a central role through the end-to-end overlap between adjacent molecules.

**THIN FILAMENT ACTIVATION: TWO STATE VERSUS THREE STATE MODEL**

Several models have been proposed depicting the mechanism by which thin filaments are activated. Hill’s model assumes two states that correspond to the thin filament being in an “on” or “off” conformation (Hill et al., 1980). The entire regulatory unit switches between states where myosin can bind either weakly or strongly. This model is appealing because it is based on well-established structural and thermodynamic details, but aspects of it are not entirely consistent with subsequent observations of thin filament structural data which show Ca$^{2+}$ binding causes a major movement of Tm (Vibert et al., 1997).

A widely accepted model for striated muscle regulation based on kinetic studies was proposed by McKillop and Geeves (1993); it describes thin filament regulatory units in equilibrium between three states, termed Blocked, Closed, and Open. In each state, Tm...
Cooperativity is an essential feature of striated muscle activation, especially in cardiac muscle because the heart functions in a highly coordinated manner. In addition to local cooperativity within a regulatory unit implied above, three models have been proposed for longer range cooperativity between regulatory units along a thin filament. An initial model was proposed by Hill et al. (1980). Hill suggested that each Tm–Tn and associated seven actin monomers act as a single unit that changes state individually, but the state of one unit affects the probability of activation of the adjacent, nearest-neighbor units; this can result in positive or negative cooperativity between nearest-neighbor units. In a later model of Geeves and Lehrer (1994), the concept of cooperative unit size n was introduced which represents the number of adjacent actin monomers that become available for myosin binding when a single regulatory unit turns on. This model incorporated Tm as a continuous, flexible filament allowing signal propagation along the surface of thin filaments. A third model was developed by Tobacman and Butters (2000) and involves actin–actin cooperativity which allows long range propagation of actin structural changes along a thin filament.

All three models have been shown to apply for a particular set of in vitro data, but currently none can fully explain the range of experimental data available (Boussouf and Geeves, 2007). This may be due, in part, to the complexity of the myofilament lattice in striated muscle. Each thin filament is comprised of several hundred molecules, and can interact with a similarly large number (tens to hundreds) of calcium ions and myosin molecules, with binding sites distributed along its length. Thus not only are there several different types of cooperative mechanisms in striated muscle, there are an even larger number of possible cooperative interactions (Gordon et al., 2000) that could affect muscle function. In Table 1, we summarize various cooperative mechanisms that may be involved in striated muscle regulation: Ca2+ binding to one regulatory unit may induce Ca2+ binding to an adjacent unit; formation of strong actomyosin crossbridges in one regulatory unit may induce Ca2+ binding to and/or crossbridge formation in the same and/or an adjacent unit; displacement of one tropomyosin (e.g., by Ca2+ binding to troponin and/or crossbridge formation) leads to the spread of activation which allows further crossbridge binding not only within that regulatory unit but also, through end-to-end contacts between adjacent tropomyosin molecules, in adjacent units; alternatively, it was proposed that activation might spread along the thin filament through actin monomers. All of these mechanisms might involve tropomyosin directly or indirectly. For instance, binding of Ca2+ to a regulatory unit leads to displacement of tropomyosin which affects the neighboring subunit, possibly through the Tm end-to-end overlap, and may induce Ca2+ binding to the neighboring Tn. Tm does in fact play a central role in cooperativity and is involved in the different mechanisms proposed above; it is indispensable for a coordinated activation of the muscle. There is an additional mechanism specific to skeletal muscle regulation, i.e., Ca2+ binding to one trigger site in the N-lobe of skeletal troponin C may induce Ca2+ binding to the second trigger site in the N-lobe of that troponin molecule; this only applies to skeletal muscle because the N-lobe of skeletal troponin C has two physiologically active EF-hand Ca2+-binding sites, while cardiac troponin C only has one. Furthermore, the mechanical properties – particularly myofilament compliance – of the myofilament assemblies could modulate both the actual cooperative interactions and the apparent cooperativity of force generation in the sarcomere (Chase et al., 2004; Kataoka et al., 2007).

A quantitative model described by Hill treats Tm as a rigid rod having end-to-end overlap with adjacent molecules of Tm (Hill et al., 1980). If only a single regulatory unit was activated, however, this simplification would mean that the Tm strand would be broken as one Tm molecule changes its structural state. It was later demonstrated experimentally that Tm is a semi-flexible molecule and Tn increases communication between neighboring structural units (Geeves and Lehrer, 1994). It was further shown that skeletal Tm induces cooperative binding of S1-ADP to actin.

Table 1 | Cooperative mechanisms in muscle regulation where tropomyosin might play a central role.

| Type of cooperativity | Mechanism | Tropomyosin central role |
|-----------------------|-----------|--------------------------|
| Calcium               | Binding to one subunit induces Ca2+ binding to the next | Possibly |
| Crossbridge formation | Formation in one regulatory unit induces Ca2+ binding to the same or next unit | Possibly |
| Tm activation         | Formation in one regulatory unit induces crossbridge formation in same or next unit | Yes |
| Actin activation      | Spread of activation through tropomyosin | Yes |
|                       | Spread of activation through actin | No |
in solution (Hill et al., 1980), and cooperative activation of actomyosin subfragment 1 (acto-S1) solution ATPase activity which is manifested by a sigmoidal ATPase versus (S1) relationship in the presence of skeletal Tm (Lehrer and Morris, 1982). Using a variety of complementary assays at different levels of biological organization, cardiac Tm confers statistically similar degrees of apparent cooperativity when compared with skeletal Tm (Clemmens et al., 2005; Boussouf et al., 2007; Jagatheesan et al., 2010); interestingly, these studies demonstrate that additional aspects of Ca$^{2+}$-activated actomyosin function – such as the maximum isometric force – may be influenced by which isoform of Tm is present, and also that the isoform of troponin is important. While this work is in accord with the idea that both Ca$^{2+}$ and crossbridges cooperatively activate the thin filament regardless of Tm (and Tn) isoform, the experiments do not address their relative importance for cardiac versus skeletal muscle, as dissected by Gillis et al. (2007) and Regnier et al. (2002). Regardless of whether thin filament activation depends more on Ca$^{2+}$ or crossbridges (Table 1), these data indicate that cooperative spread of activation from one regulatory unit to the next depends critically on both the presence of, and the molecular composition of tropomyosin.

**DOES Ca$^{2+}$ ACTIVE THE THIN FILAMENT AS A SINGLE UNIT?**

The primary structural regulatory unit (SRU) responsible for the Ca$^{2+}$ switch, consisting of one Tm molecule, one Tn complex, and seven actin monomers, is approximately the length of one single Tm molecule. The functional regulatory unit (FRU) is defined by the length of the thin filament activated by Ca$^{2+}$ binding to a single Tn. In case of negative cooperativity the FRU would be shorter than the SRU, whereas for positive cooperativity, the FRU would be longer than the SRU. It was suggested from isometric force measurements that Ca$^{2+}$ is measured prior to, and $k_{TR}$ is measured following mechanical maneuvers that consist of a rapid length release which unloads the muscle preparation, and a sudden re-stretch to the original length (Brenner, 1988; Sweeney and Stull, 1990). At saturating Ca$^{2+}$ levels, $P$ is primarily determined by the number of strong crossbridges whereas $k_{TR}$ reports the rates of crossbridge transitions between weak, non-force states and strongly bound, force generating states (Brenner and Eisenberg, 1986). When Ca$^{2+}$ concentration is varied over the activating range, the relationship between $k_{TR}$ and $P$ is such that at low $P$ ($P < 50\% P_{max}$), $k_{TR}$ is slow and unchanging. When the level of Ca$^{2+}$ activation is increased and $P$ approaches $P_{max}$, $k_{TR}$ increases 10- to 15-fold in skeletal muscle preparations (Metzger and Moss, 1992; Chase et al., 1994; Regnier et al., 1996, 1998).

To explain the activation dependence of $k_{TR}$, a simple model was evaluated (Landesberg and Sideman, 1994; Hancock et al., 1997; Regnier et al., 1999). A version of the model with four states (Figure 1) was necessary to describe the relationship between $k_{TR}$ and $P$ in skeletal muscle, while only three states (no state 4 in Figure 1) were required to describe the less-steep relationship in cardiac muscle (Hancock et al., 1997). In addition to the variable [Ca$^{2+}$], the skeletal and cardiac versions of the model have two pairs of kinetic rate parameters: $f$ and $g$ which reflect the processes associated with strong crossbridge formation and dissociation, respectively, and $k_{on}$ ($k_{on}^f$) and $k_{off}$ ($k_{off}^g$) which reflect the processes associated with thin filament regulatory unit activation by Ca$^{2+}$ and deactivation, respectively. The four-state version of the model for skeletal muscle has additional parameters, $f'$ and $g'$: $g'$ reflects strong cross-bridge dissociation from regulatory units that have lost Ca$^{2+}$; the rate parameter $f'$ reflects the low probability of strong crossbridge formation at regulatory units in the blocked state, and is included for completeness. There is no inherent cooperative mechanism taken into consideration in the model, whether at the level of Ca$^{2+}$ regulation or crossbridge formation. While this does not allow simulation of the steep dependence of $P$ on Ca$^{2+}$ concentration, the model can readily predict the thin filament activation dependence of $P$ and $k_{TR}$ under a variety of conditions. Modeling and experiments, taken together, indicate that $k_{TR}$ at submaximal Ca$^{2+}$ activation typically reflects the kinetics of individual thin filament regulatory units without requiring cooperative interactions between regulatory units, unlike steady-state tension where cooperativity plays a central role and simulations are not possible without it (Regnier et al., 1999; Moreno-Gonzalez et al., 2007). This shows that under physiologically relevant conditions, $k_{TR}$ is regulated by the dynamics of thin filament activation at submaximal Ca$^{2+}$ levels and Ca$^{2+}$ controls the rate limiting step in tension development, whereas at maximal Ca$^{2+}$ the increase of $k_{TR}$ with force is governed by the
FIGURE 1 | Schematic diagram of four-state model. On the left hand side are states where myosin forms weak, non-force-producing crossbridges. On the right hand side are force generating states where myosin binds strongly to actin. The upper states represent Ca$^{2+}$ free states whereas the lower states represent Ca$^{2+}$ bound. Ca$^{2+}$ binds to TnC and activates regulatory units with rate constants $k_{on}$ (left) and $k_{on}'$ (right), while $k_{off}$ (left) and $k_{off}'$ (right) reflect the kinetics of processes associated with regulatory units returning to the blocked state. Strong crossbridge formation and dissociation from thin filaments are governed by rate constants $f$ and $g$, respectively. $g'$ describes the rate of crossbridge detachment from regulatory units without Ca$^{2+}$, as may occur during relaxation of skeletal muscle; the rate of strong crossbridge formation ($f'$) is practically non-existent in this condition. State 4 (upper right) is needed to model P−$k_{TR}$ relationships in skeletal muscle, but can be omitted when modeling cardiac muscle (Hancock et al., 1997).

rate of crossbridge cycling (Chase et al., 1994; Regnier et al., 1996, 1998).

While the experiments described above provide information about the dynamics of individual regulatory units within the sarcomere, all biochemical and skinned-fiber experiments performed thus far to characterize cooperativity involved ensemble measurements. There is no direct measurement of cooperativity between individual neighboring SRUs, which is necessary to determine whether a thin filament activates as a single unit or activation involves numerous cooperative units.

Hypothesizing that we are able to incorporate a small number of reporters, each at an individual regulatory subunit along an individual thin filament, we could study the dynamics of activation of single regulatory units, as well as cooperative interactions between regulatory subunits. The reporter changes state when the regulatory unit turns on or off, reflecting the state of that regulatory unit (Figure 2). Statistical analysis of the signal from a single reporter would yield the dynamics of activation of the associated regulatory unit. We could also examine the cooperativity along the thin filament as a function of distance between two reporters. As shown in Figure 2A, two regulatory units separated by a short distance could show a highly correlated signal. Reporter signals from two regulatory units that are far apart, however, might not be correlated (Figure 2B). Introduction of such a novel technique would allow us to study cooperativity directly, and test the hypothesis of long range cooperativity along a thin filament.

FAMILIAL HYPERTROPHIC CARDIOMYOPATHY ALTERS COOPERATIVE Ca$^{2+}$-ACTIVATION OF CARDIAC THIN FILAMENTS

Independent of whether a thin filament activates as a single unit, familial hypertrophic cardiomyopathy (FHC)-related mutations have been found to alter Ca$^{2+}$-sensitivity using in vitro functional assays. FHC is an inherited disease characterized by thickening of the myocardium. The disease affects an estimated 0.2% of the population and may be relatively benign, or may lead to heart failure or sudden cardiac death (Maron, 1997; Maron et al., 1998). A number of FHC-related mutations have been found in human cardiac α-tropomyosin (αTm), along with many other mutations primarily in cardiac cytoskeletal proteins (Bing et al., 2000; Fatkin and Graham, 2002; Roberts, 2002; Towbin and Bowles, 2002; Takeda, 2003; Wolska and Wieczorek, 2003). FHC-related αTm mutants have been linked to decreased thermal stability (Hilario et al., 2004; Kremneva et al., 2004; Wang et al., 2011) and a lower binding affinity for actin (Bing et al., 1997; Kremneva et al., 2004) compared to WT. In vitro studies with mutant αTm using myofibrillar ATPase activity, motility assays, and isometric force generation show significantly enhanced Ca$^{2+}$-sensitivity and/or reduced cooperativity (Bing et al., 2000; Chang et al., 2005; Bai et al., 2011; Mathur et al., 2011; Wang et al., 2011). Cardiomyopathy mutations in αTm also affect skeletal muscle physiology (Bottinelli et al., 1998) because both α- and β-tropomyosins (WT and, in patients mutant αTm) are expressed in skeletal muscles, while cardiac muscle contains predominantly αTm; because the major pathology is in the heart,
however, the consensus has been to look at the cardiac muscle and its proteins when studying the effect of FHC mutations on muscle contraction. Although the mechanistic relationship between the mutations and these observations is not yet clearly established (Tardiff, 2011), reduced rigidity of human cardiac αTm due to the mutations might be expected to affect the regulatory function of the molecule, leading to observed changes in Ca²⁺-sensitivity and/or cooperativity.

**IS TROPOMYOSIN A SEMI-FLEXIBLE MOLECULE?**

The flexural rigidity of a semi-flexible linear molecule can be characterized by its persistence length ($L_p$) which is the length over which the molecule loses directional correlation. $L_p$ of tropomyosin from chicken and turkey gizzard smooth muscle, rabbit skeletal muscle, or rabbit and bovine cardiac muscle has been estimated to be 55–170 nm by various techniques at different temperatures (Swenson and Stellwagen, 1989; Phillips and Chacko, 1996; Li et al., 2010; Sousa et al., 2010), consistent with measurements obtained for other α-helical coiled-coil proteins (Hvidt et al., 1982; Howard and Spudich, 1996). Among these studies, Li et al. (2010) suggested that previous experimental estimates of $L_p$ (i.e., apparent $L_p$), including that obtained for bovine cardiac tropomyosin in the same study, measured a combined effect of the inherent curved molecular structure and the true mechanical flexibility of tropomyosin. The apparent $L_p$ is related reciprocally to the $L_p$ due to true mechanical flexibility (i.e., dynamic $L_p$) and the $L_p$ due to inherent molecular curvature (i.e., intrinsic $L_p$; Eq. 1):

$$\frac{1}{L_{p\text{ apparent}}} = \frac{1}{L_{p\text{ intrinsic}}} + \frac{1}{L_{p\text{ dynamic}}}.$$  

The two effects were decoupled in a molecular dynamics simulation, which determined the dynamic $L_p$ of Tm to be on the order of 500 nm, or ~12 times the length of a single Tm molecule. We note that an $L_p$ of ~500 nm would imply that Tm behaves as a rigid body over the span of ~12 SRU and thus the length of a FRU would be equivalent to approximately half the length of a thin filament within a sarcomere. This implication is consistent with the hypothesis that a thin filament activates as a single unit (Figure 3A, lower left). In contrast, however, previous studies showed the length of a FRU is no more than 12–14 actin monomers, which is approximately equivalent to the length of two SRUs, or more simply two Tm molecules (Regnier et al., 2002; Figure 3A, lower right). It is particularly noteworthy here that, assuming the intrinsic $L_p$ of 135 nm determined by Li et al. (2010), Eq. 1 suggests the correction to apparent $L_p$ due to intrinsic $L_p$ diminishes drastically and
non-linearly as apparent \( L_p \) approaches \( L_c \) (Figure 3); in case of a rigid Tm with apparent \( L_p > 2.5 \, L_c \), the dynamic \( L_p \) is above 400 nm, or \( \sim 10 \) SRU, consistent with the hypothesis that a thin filament activates as a single unit (Brandt et al., 1980; Fraser and Marston, 1995); on the other hand, in case of a semi-flexible Tm with apparent \( L_p \sim L_c \), dynamic \( L_p \) of the molecule is in the order of 60 nm, or \( \sim 1.5 \, L_c \), consistent with the hypothesis that the length of a FRU approximately equals 1–2 SRUs (Regnier et al., 2002; Gillis et al., 2007). It remains to be determined whether differences in experimentally determined values of apparent \( L_p \) over this crucial range (Figure 4) reflect differences inherent in the proteins or in experimental methodologies. Due to the non-linearity evident in Figure 4, a slight reduction in the experimentally measurable apparent \( L_p \), depending on its exact value, may potentially imply a profound change in the true mechanical flexibility of Tm, as represented by its dynamic \( L_p \).

REDUCTION IN TROPOMYOSIN RIGIDITY DECREASES MECHANICAL CORRELATION ALONG THE MOLECULE

Transmission of a mechanical perturbation along the length of a Tm molecule can be modeled as an exponential decay according to the cosine correlation function (CCF; Howard, 2001). Since a Tm molecule follows the helical structure of a thin filament along its length, and each end of a Tm molecule is associated with a distinct Tn complex (at the head-to-tail overlap regions with neighboring molecules of Tm), the middle of a Tm molecule is expected to be the least perturbed region during activation at high \( \text{Ca}^{2+} \) levels (i.e., when the Tn complexes at both ends of a Tm molecule have \( \text{Ca}^{2+} \) bound). Assuming that configurations sampled by Tm on the thin filament are mainly determined by its mechanical properties, as illustrated in Figure 3, CCF can predict the propagation of activation. Figure 5A depicts the variation of perturbation signal transmitted from the ends to the middle of a Tm molecule at decreasing flexural rigidity, with initial \( L_p = 65 \) or 170 nm (as the lower and upper limit in crystallographic and solution studies of skeletal Tm). The effect of reduced signal transmission is noticeable in both cases, but is more significant for a more flexible Tm (\( L_p = 65 \) nm). On the other hand, since cytoplasmic \( \text{Ca}^{2+} \) does not normally achieve levels that fully saturate thin filament during systolic activation of cardiac muscle, only some but not all SRUs will have \( \text{Ca}^{2+} \) bound to the Tn complexes at both ends. Thus we also have to consider the case where \( \text{Ca}^{2+} \) binds to the Tn complex at only one end of a Tm molecule. In that situation, the opposite, \( \text{Ca}^{2+} \)-free end would be the least perturbed region, and Figure 5B shows the variation of transmitted signal at the \( \text{Ca}^{2+} \)-free end when Tm rigidity is altered. Compared to the case where \( \text{Ca}^{2+} \) binds to the Tn complexes at both ends of a Tm molecule, activation signal transmitted to the least perturbed region is significantly decreased at any given reduction in rigidity of Tm. Readers should note especially the case of a more flexible Tm (initial \( L_p = 65 \) nm), where signal transmission is nearly halved over the considered range of decrease in rigidity. It is clear that in most cases during systolic activation, when \( \text{Ca}^{2+} \) may be bound to the Tn complex only at one end of a Tm molecule, the transmission of mechanical perturbation is significantly impeded by reduction in rigidity of Tm, especially when the initial \( L_p \) is comparable to \( L_c \). It is evident from this simple model that a drop in rigidity of a Tm molecule (e.g., associated with a disease-related mutation)

![Figure 4](image128x145to468x381.png)

**FIGURE 4** Correction due to intrinsic curvature of tropomyosin is less significant when apparent \( L_p \) approaches the contour length. According to Eq. 1, the percentage correction to apparent \( L_c \) due to intrinsic \( L_c \) (dashed line, right axis) is non-linear and is much smaller at values of apparent \( L_p \leq L_c \) than at longer values. Dynamic \( L_c \) (solid line, left axis), measurement of the true mechanical flexibility obtained after the correction, is also close to apparent \( L_c \) when apparent \( L_c \leq L_c \). For reference, the linear relation obtained for molecules with no intrinsic bend is also shown (dot-dashed line, left axis). Also for reference, \( L_c (\sim 40 \text{ nm}) \) for Tm molecules is highlighted by the vertical, gray dashed line.
can lead to a significant difference in mechanical correlation along its length, the extent of which depends on both the initial flexural rigidity of the molecule and where along the length this effect is measured. This effect is likely to have important functional consequences which are discussed further below. It can also underlie mechanical tuning at the molecular level through evolution, where sensitivity of Tm as a key component of the regulatory switch for thin filament activation may have been optimized within different physiological situations.

**IMPLICATIONS OF REDUCED CORRELATION ON COOPERATIVE THIN FILAMENT REGULATION**

We expect that a reduction in mechanical correlation would correspond to reduced cooperativity in transmission of activation along thin filaments, and could have major impacts on thin filament regulation.

First, decreased correlation along the length of a Tm molecule could influence ordered assembly of thin filaments by reducing the overall affinity of αTm strands for F-actin. It is therefore easier for αTm to move away from myosin binding sites during transition from the “blocked” to “closed” state during Ca2⁺-induced activation. Reduced mechanical correlation also implies a smaller turning moment and thus a lesser extent of conformational change of Tn is required to initiate azimuthal movement of αTm on the actin filament. Taken together, these can affect the functional Ca2⁺ sensitivity (i.e., pCa50) of regulation. Secondly, reduced correlation also means the mechanical turning moment due to Tn conformational change does not propagate as effectively along the length of Tm (Figure 5B, right). This may in turn affect the number of myosin binding sites uncovered and thus the force generated by each SRU at a given Ca2⁺ level. The exact functional implication and significance of these effects depend heavily on the innate flexibility of the wildtype αTm, which should be considered in three regimes: very flexible (Lp ≪ Lc), very rigid (Lp ≫ Lc), and semi-flexible (Lp ∼ Lc).

In the case of αTm that is already very flexible (Lp ≪ Lc), little or no intrinsic cooperative activation would exist between SRUs through end-to-end overlap. A further increase in flexibility (reduction in Lp) would result in even less mechanical correlation along the molecule, such that activation within each SRU would be reduced at all levels of Ca2⁺. In other words, an increase in flexibility of a highly flexible αTm leads to predicted reductions in both Ca2⁺ sensitivity and maximum force. The condition Lp ≪ Lc, however, is outside the range of existing measurements on α-helical coiled-coil proteins (see Is Tropomyosin a Semi-Flexible Molecule?), and thus this regime is not considered in detail.

Conversely, if αTm is inherently very rigid (Lp ≫ Lc), the reduction in mechanical correlation due to increased flexibility may be relatively insignificant (Figures 5A,B, dotted-dash line), such that the probability of uncovering any number of myosin binding sites does not change at all levels of Ca2⁺. Therefore, in this case, an increase in flexibility leads to the prediction of no or minimal change in Ca2⁺ sensitivity, cooperativity, and maximum force.

Lastly, for a semi-flexible αTm (Lp ∼ Lc), an increase in flexibility will result in a significant loss in correlation (Figure 5B, solid line), while the activation signal would still be able to propagate along a limited span of the molecule that is longer than an actin monomer. In this case where cooperativity is reduced, it would be mechanically more favorable for a Ca2⁺ bound Tn...
to induce sufficient movement of αTm to uncover at least some myosin binding sites, leading to increased functional Ca2⁺ sensitivity. As a result, the thin filament can be activated at a lower level of Ca2⁺, such as in early and late stages of systole or, in extreme cases that might be associated with cardiac disease, diastole. At saturating Ca2⁺, however, the reduced correlation implies that Ca2⁺ dependent azimuthal displacement at each end of Tm may not be fully transmitted to the middle of the molecule (Figure 5A, solid line). If \( L_p \) is near the low end of this regime and the increase in flexibility is sufficiently large, then some myosin binding sites would be more likely to remain blocked at high Ca2⁺ levels. On the other hand, if \( L_p \) is near the high end of this regime, there could be a tolerance in the system such that cardiac sarcomeres might maintain maximum force at systolic Ca2⁺ level despite a moderate increase in flexibility of αTm. Taken together, increased flexibility of a semi-flexible αTm enhances the probability to uncover at least some myosin binding sites at low Ca2⁺, but may slightly reduce the probability to uncover all myosin binding sites at high Ca2⁺.

**POSSIBLE LINK TO CARDIAC HYPERTROPHY**

We reason by considering the mechanical correlation along the tropomyosin molecule that reduced Tm rigidity will contribute to higher Ca2⁺-sensitivity in cooperative thin filament regulation. This implies that human cardiac thin filaments harboring less rigid αTm mutants will undergo a prolonged systolic activation and perhaps diastolic dysfunction. Cardiac thin filaments will become activated earlier during systole, remain activated longer during the relaxation phase, and in extreme cases could possibly stay partially activated during diastole (Ho et al., 2009; Bai et al., 2011; Campbell and McCulloch, 2011). These effects would be expected to markedly impact the overall mechanics of the heart and cardiac output. Therefore, we hypothesize that at least some FHC-related mutations of human cardiac Tm exert their major influence on sarcomere mechanics through altered flexural rigidity of the molecule. This may in turn lead to functional changes in cooperative Ca2⁺ induced activation of cardiac thin filaments, such as previously observed in *in vitro* studies (Bing et al., 2000; Chang et al., 2005; Boussouf et al., 2007; Bai et al., 2011; Mathur et al., 2011; Wang et al., 2011). Techniques such as that illustrated in Figure 2 (described above) can directly decipher at the molecular level the effects FHC-related mutations impose on the dynamics of activation and cooperative interactions of individual and multiple regulatory units, respectively. While it is likely that additional factors will need to be considered to fully understand the complex phenotype observed in patients with these inherited cardiomyopathies, we fully anticipate that measurements of Tm’s flexural rigidity will provide an improved molecular understanding (Figures 3–5) of an important mechanistic link between FHC-related mutations of the molecule and hypertrophy of the heart due to increased workload.

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