Multiscale modeling of twitch contractions in cardiac trabeculae

Srboljub M. Mijailovich, Momcilo Prodanovic, Corrado Poggesi, Michael A. Geeves, and Michael Regnier

Understanding the dynamics of a cardiac muscle twitch contraction is complex because it requires a detailed understanding of the kinetic processes of the Ca^{2+} transient, thin-filament activation, and the myosin–actin cross-bridge chemomechanical cycle. Each of these steps has been well defined individually, but understanding how all three of the processes operate in combination is a far more complex problem. Computational modeling has the potential to provide detailed insight into each of these processes, how the dynamics of each process affect the complexity of contractile behavior, and how perturbations such as mutations in sarcomere proteins affect the complex interactions of all of these processes. The mechanisms involved in relaxation of tension during a cardiac twitch have been particularly difficult to discern due to nonhomogeneous sarcomere lengthening during relaxation. Here we use the multiscale MUSICO platform to model rat trabecular twitches. Validation of computational models is dependent on being able to simulate different experimental datasets, but there has been a paucity of data that can provide all of the required parameters in a single experiment, such as simultaneous measurements of force, intracellular Ca^{2+} transients, and sarcomere length dynamics. In this study, we used data from different studies collected under similar experimental conditions to provide information for all the required parameters. Our simulations established that twitches either in an isometric sarcomere or in fixed-length, multiple-sarcomere trabeculae replicate the experimental observations if models incorporate a length–tension relationship for the nonlinear series elasticity of muscle preparations and a scheme for thick-filament regulation. The thick-filament regulation assumes an off state in which myosin heads are parked onto the thick-filament backbone and are unable to interact with actin, a state analogous to the super-relaxed state. Including these two mechanisms provided simulations that accurately predict twitch contractions over a range of different conditions.

Introduction

In cardiac muscle, transient increases in the cytosolic calcium concentration trigger twitchlike contractions during the systolic phase of heart function (Gordon et al., 2000). Ca^{2+} binding to cardiac troponin (cTn) results in movement of tropomyosin (Tpm) over the F-actin surface of the thin filament. This exposes myosin-binding sites on actin that allow for myosin–actin cross-bridge formation, cyclic interactions of myosin heads with actin, force generation, and contraction. After reaching a peak, intracellular Ca^{2+} concentration ([Ca^{2+}]) is rapidly removed from the cytosol, lowering [Ca^{2+}], enhancing additional dissociation of Ca^{2+} from cTn, and allowing Tpm to return to its blocking position. The increasing number of Tpms in the blocked position reduces overall binding flux, reduces the number of bound cross-bridges, and results in muscle relaxation. Understanding the dynamics of this process in a single twitch contraction is complicated because it requires a detailed understanding of the kinetics of each step of the process: the Ca^{2+} transient, the dynamics of the troponin (Tn)–Tpm switch, and the myosin cross-bridge chemomechanical cycle. Each of these steps is well defined individually, but how all three processes work in combination is not understood. Here we build on our previous work on modeling the different elements of the contraction cycle to define a novel multiscale model of a cardiac muscle twitch contraction.

Computational models have the potential to interconnect multiple types of experiments and protocols and can be a useful tool for generating and testing hypotheses regarding the mechanisms of contractile function in healthy hearts and contractile dysfunction in diseased hearts. Early models defined structural cooperative effects in sarcomeres, such as end-to-end
interactions between adjacent Tpm–Tn regulatory units, between cross-bridges bound to actin and the thin-filament regulatory units, and between neighboring bound cross-bridges (Campbell et al., 2001; Rice et al., 2008; Tanner et al., 2012). Several of these models are derived from the original formulation by Razumova et al. (1999, 2000), where the muscle tension is proportional to the product of fractional occupancy of the myosin bound states and the mean cross-bridge distortion of each state population. The approximation of strain-dependent distributions by mean distortions is mathematically effective because the model can be defined by a system of ordinary differential equations. The newer generations of spatially explicit models, coupled with a cross-bridge cycle and thin-filament regulation by Ca²⁺, show significantly improved relationships between molecular interactions and muscle fiber behavior (Chan et al., 2000; Tanner et al., 2012; Mijailovich et al., 2016; Mijailovich et al., 2019). The key feature of these models is the ability to predict mechanical response to Ca²⁺ transients reflecting cardiac muscle behavior in a living heart. The dynamic relationships in a contraction are defined by the coupling of thin-filament regulation by Ca²⁺ and the strain-dependent kinetics of myosin binding to actin. However, just as with experiments studying force generation and cross-bridge cycling, this type of model has been limited primarily to simulating force generation under steady-state Ca²⁺ conditions (Chase et al., 2004; Kataoka et al., 2007; Tanner et al., 2007; Tanner et al., 2008; Tanner et al., 2012). Modeling twitch contractions has been approached with different levels of complexity, ranging from models based on solution of ordinary differential equations (Campbell et al., 2001; Niederer et al., 2006; Rice et al., 2008; Land et al., 2017), to a spatially detailed cardiac sarcomere model (Washio et al., 2012), to a continuous time Markov (regulatory) chain coupled with a generalized Huxley 57 model (Regazzoni et al., 2020), to more detailed models based on solution of partial differential equations (Chung et al., 2017; Campbell et al., 2018).

Moreover, the latest models include mechanisms that could alter cardiac muscle contractility by structural changes in thick filaments. Possible mechanisms could include “mechanosensing” in thick filaments (Linari et al., 2015; Irving, 2017) and interfilament mechanical signaling by myosin-binding protein C (MyBP-C; Kampourakis et al., 2016; Irving, 2017; Brunello et al., 2020). Following these ideas, Campbell et al. (2018) hypothesized that the transition rate from the parked state (PS) to the myosin on state, which permits binding to actin, is dependent on force. They successfully fitted the length-dependent activation data, where the twitch contractions only include fitting of the observed tension traces, but they did not take into account the observed significant shortening and lengthening at the sarcomere level known to occur in trabeculae held at fixed length (Janssen and de Tombe, 1997; Ferrantini et al., 2014; Caremani et al., 2016). In addition, these fits were achieved in the model using simplified cross-bridge cycle kinetics and sarcomere geometry. The simulations of twitch contractions, in which transition rate from PS state to M.D.Pi (myosin head with nucleotides ADP and phosphate, Pi) are force dependent, using a more detailed model (muscle simulation code, MUSICO), however, showed that the relaxation of tension is slow at higher levels of tension even when the [Ca²⁺] falls to low values. Our preliminary results suggest that the transition rate from PS state should be defined by the relationship of some process other than a linear dependence on tension because the PS should act as a sink for myosin heads during relaxation and a source for additional myosin head recruitment during contractions (Prodanovic et al., 2020).

Taken together, all of these models can more or less successfully replicate observed tension traces in response to Ca²⁺ transients. However, they all include a large number of simplifications, so that changes in molecular interactions within sarcomeres, such as by mutations of contractile and regulatory proteins, may not be transparent across the multiple length scales. Thus, these models may not be able to translate the changes at the molecular level to the observed changes in cardiac muscle behavior in associated diseases.

Furthermore, a major challenge in validating computational models by simulating experimental data has been the variability in conditions used by investigators. For example, some experimenters have recorded tension and sarcomere length (SL) transients but not the Ca²⁺ transient (Caremani et al., 2016), or they have recorded Ca²⁺ and tension transients at different temperatures without monitoring changes in the SL during the twitch in fixed-length trabeculae (Janssen et al., 2002). The only studies that we are aware of that reported [Ca²⁺] tension, and SL simultaneously were reported by Janssen and de Tombe (1997) and Ferrantini et al. (2014). Studies that do not include SL control do not account for the series elastic (SE) compliance, i.e., for significant changes in SLs during contractions of intact muscle preparations held at a fixed length. Attempts to take this SE compliance into account include the study by Kataoka et al. (2007), where the series compliance of trabeculae was exclusively attributed to the compliance of thin and thick filaments but did not account for SL changes caused by SE compliance of trabeculae. This series compliance was considered by Rice et al. (2008), who demonstrated a profound effect on the tension response during twitch contractions. However, Rice et al. used a model that showed only the sensitivity of the responses to a large range of simulated compliances, with little reference to experimental data. To more precisely simulate the steady-state and dynamic properties of striated muscle contraction and relaxation, all compliances and SL changes must be considered because cross-bridge kinetics is sensitive to the changing load as the sarcomeres shorten or lengthen.

In this study, we report a model that more precisely defines the multiple sarcomere parameters that vary during contraction and relaxation, using the methodology implemented in the MUSICO computational platform (Mijailovich et al., 2016; Mijailovich et al., 2019). This model allows simulation of the tension developed during a twitch contraction, as well as the kinetics of contraction and relaxation. Previous models have successfully simulated the kinetics but have accomplished this using a normalized force value. We initially simulated the twitch data of Janssen and de Tombe, 1997; Caremani et al., 2016 without normalizing tension using a five-state cross-bridge model. This resulted in an elevation of resting force and prolonged relaxation that were addressed by addition of two
features: (1) SE compliance to account for shortening and lengthening of sarcomeres during a twitch in a trabecula kept at fixed length and (2) expansion of the cross-bridge cycle to include a myosin “off state” that acts as a source for myosin recruitment during activation of contraction and as a sink for taking myosin out of the cycling pool during relaxation (analogous to the super-relaxed [SRX] state of the thick filament; Limari et al., 2015; McNamara et al., 2015; Irving, 2017). Addition of this state to the cross-bi”dge model accelerated relaxation kinetics and reduced resting tension, producing simulated twitches that matched the experimental twitch data of Janssen and de Tombe, 1997; Caremani et al., 2016. Additional validation of the model was performed by simulations of a twitch for multiple datasets, including varying temperature, and the force-Ca²⁺ relationship. Thus, the model can predict both twitch transients and the Ca²⁺ sensitivity of force. Preliminary reports of this work have been published previously in the forms of abstracts and proceedings (Prodanovic et al., 2020; Mijailovich et al. 2018. Biophys. J. 114:Abstract 500A. https://doi.org/10.1016/j.bpj.2017.11.2737; Mijailovich et al. 2019. Biophys. J. 116:Abstract 116A. https://doi.org/10.1016/j.bpj.2018.11.654).

Materials and methods
Here we outline the essential definitions and relationships developed in previous studies, including (1) the multiscale structural organization of intact muscle (Mijailovich et al., 2016) and (2) Ca²⁺-dependent kinetics that regulate Tn-Tpm interaction with F-actin in cardiac muscle (Geeves et al., 2011; Mijailovich et al., 2012; Mijailovich et al., 2019; Prodanovic et al., 2020). We also describe new features implemented in MUSICO to model transient contractions in cardiac trabeculae, namely (3) an expanded cross-bridge cycle that contains five essential states and further expansion to six states, (4) intrinsic series compliance that permits SL changes during twitch in fixed-length trabeculae, and (5) thick-filament activation.

Multi-scale model of trabeculae
Intact ventricular trabeculae from the rat heart are composed of parallel cardiac muscle cells, or myocytes, embedded in an extracellular matrix that includes the collagen structure laterally connecting the muscle fibers. Myocytes occupy ~70% of a trabecular volume, and the cardiac cells have a diameter ranging between 10 and 20 µm and a length between 80 and 100 µm. The myocyte contractile substructure consists of myofibrils aligned in parallel, where each myofibril is composed of 40–50 sarcomeres in a series. The sarcomeres are composed of interdigitated thick and thin filaments, where each half thick filament consists of ~150 myosin molecules projecting on each side of the M-line and each thin filament contains from 360 to 400 actin monomers emerging from Z-discs on opposite sides of a sarcomere. The thick filaments are associated with the auxiliary protein titin, connecting it via the thick-filament backbone with the Z-disc and the M-line, and MyBP-C connecting thick and thin filaments. The thin filaments contain the regulatory proteins Tpm and Tn essential for Ca²⁺ regulation of contraction and other auxiliary proteins regulating thin-filament length and other functions.

Trabecular geometry and elasticity
The contractile characteristics of cardiac muscles are typically derived from isometric and isotonic twitch contractions in vivo preparations. However, twitchs of trabeculae that are isometric at the muscle level are not isometric at the level of the sarcomere (Janssen and de Tombe, 1997; Caremani et al., 2016). During twitches, when force rises, the sarcomeres shorten; during relaxation, they lengthen. The origin of the elasticity in trabeculae is complex and may include muscle cell alignment and connective tissue serial elasticity. The simplest method is to include a trabecula SE element in series with the contractile element (CE) consisting of multisarcomere muscle fibers (Fig. 1).

Spatially explicit 3-D model of a sarcomere
In the 3-D sarcomere, an overlapping hexagonal lattice of thick and thin filaments has a well-defined interfilament spacing where the mean spacing is measured by x-ray diffraction (Higuchi et al., 1995; Millman, 1998). Explicit formulation of the sarcomere lattice is defined by the discrete position of two-headed myosin molecules and their binding sites on actin filaments. For simplicity, we consider that only one myosin head per myosin molecule is active at any one time and is denoted as a cross-bridge. Overall, the 3-D sarcomere structure is viewed as an array of thin and thick filaments connected by cross-bridges and other elastic elements (e.g., titin and MyBP-C) in a lattice network, and all of these elements can be represented as linear springs and elastic beams (Daniel et al., 1998; Mijailovich et al., 2016). The instantaneous equilibrium between actin and myosin filaments interconnected by cross-bridges is defined by a stiffness matrix, which includes the elasticity of thick and thin filaments, attached cross-bridges and titin, and load containing a vector of all external forces and internal forces generated by the action of cross-bridges. The loading conditions are defined at the boundaries of the muscle system, and they follow the experimental protocol of the relevant experiment. Typically, these include changes of length or imposed tension at the muscle end(s). The stiffness matrix is constantly changing as actomyosin bonds are created or disrupted, and active forces are also changing during conformational changes in attached myosins.

In this study, we used experimental conditions such as isometric (half) sarcomeres and fixed-length trabeculae. The latter case includes variable loading (tension) on multisarcomere (muscle fiber) structure due to serial elasticity in fixed-length trabeculae. Regarding the MUSICO simulations of trabeculae, it is possible to prescribe at the end of trabeculae arbitrary changes of length, such as in previous studies (Janssen and de Tombe, 1997; Caremani et al., 2016) where trabecular length was changed in order to keep SL constant or approximately constant or tension was applied on the trabecular ends. This kind of loading is shown in Fig. S3. The solutions of the equations defining the mechanical system of sarcomeres in series are obtained by standard finite element procedures for nonlinear systems using an iterative procedure (Bathe and Mijailovich, 1988; Mijailovich et al., 1993; Bathe, 1996).
Strain-dependent kinetics of cross-bridge cycle in the 3-D sarcomere lattice

The discrete nature of the myosin-binding sites in two helically arranged strands in each actin filament and the regular arrangement of myosin crowns along the myosin filaments require that myosin heads in a 3-D sarcomere lattice move not only axially but also radially and azimuthally to reach binding sites on actin monomers with the correct orientation (Squire, 1992). The relative spatial positions between myosin heads and binding sites on the thin filament are obtained from the current spatial positions of the sites on deformed thin and thick filaments (Mijailovich et al., 2016; Mijailovich et al., 2019), as shown in Fig. S1. The cause of deformation in the elastic filaments originates from local forces imposed by bound cross-bridges and other elastic elements (e.g., titin), but, in turn, the forces in cross-bridges depend on state and the relative positions between actin and myosin sites. The strain-dependent rate of myosin binding to actin in the 3-D sarcomere lattice is modulated by the axial strain component, \( x \), as used in all cross-bridge kinetic models, and by radial and azimuthal spatial arrangement relative to binding sites on the actin filaments (Fig. S1), displayed by interfilament spacings, \( d_{\alpha-M} = 2 d_{\alpha}/3 \), and azimuthal angles \( \alpha \) and \( \beta \) (Mijailovich et al., 2016, 2019).

Minimal cross-bridge cycle including five essential states

Simple kinetic models include only three states (e.g., Daniel et al., 1998; Duke, 1999; Mijailovich et al., 2016; Mijailovich et al., 2019), where multiple states are lumped into a single state. A cross-bridge model with additional states is desirable to relate the underlying well-defined molecular kinetics to the simulations of, for example, twitches in cardiac trabecula and allow simulation of tension levels in units of the observations. The additional states provide closer proximity to measurements of state transition rates from assays in solution and motility assays and thus provide experimentally measured model parameters at the molecular scale rather than parameters estimated from fits of data from higher length scale experiments. Hence, we expanded the above models into a five-state cross-bridge model that introduces a separate strain-dependent ADP release to distinguish it from ATP binding associated with cross-bridge detachment and the hydrolysis step that separates M.T (myosin head with ATP) and M.D.Pi states (Fig. 2 A). The former defines the maximum cycling rate of the cross-bridge cycle as a function of load. The hydrolysis step defines the minimum lifetime of the detached cross-bridge. Separation of M.T and M.D.Pi is convenient for introduction of the concepts associated with thick-filament elasticity.
that the rate of a reaction is limited by the energy barrier presented by the point of highest Gibbs energy on the reaction path. The general form of the state transition rates in models with a power stroke is formulated by Hill (1974) as the ratio of forward to backward rates that must satisfy Gibbs’s thermodynamic identity. The rates are defined in terms of the Gibbs energies of the initial and final states, including the elastic strain energy derived from the cross-bridge tension (Hill, 1974; Wood and Mann, 1981).

The state transition rates between states in the cross-bridge cycle are derived from the free energy of the cross-bridge states in terms of the axial strain component \( x \). The complete set of the state transition rate constants is described in Appendix A. Moreover, in the 3-D sarcomere lattice, the strain-dependent rate of myosin binding to actin is modulated, in addition to the axial strain component \( x \), by the spatial position of the actin filaments and the azimuthal departure of a myosin-binding site on an actin filament from the plane passing through actin longitudinal axes (Fig. S1). This effect is implemented in MUSICO via weight factors \( C_\alpha \) and \( C_\beta \) for the azimuthal departures and, in addition, includes the normalization factor, \( f_\alpha \), that takes into account the number of actin sites that a cross-bridge can reach and bind (for further details, see Mijailovich et al., 2016; Mijailovich et al., 2019).

### Thin-filament regulation by calcium

Muscle contraction and relaxation are regulated by the Ca\(^{2+}\)-dependent azimuthal movements of Tpm–Tn complexes over the surface of the actin filament. Structurally, Tpm covers seven monomers on one strand of the actin double helix and is associated with one Tn molecular complex. The Tn complex consists of troponin T, troponin C, and troponin I, denoted as TnT, TnC, and TnI, respectively. The affinity of myosin for the actin filament is controlled by the azimuthal position of Tpm, where Tpm molecules are aligned along each strand of the actin double-stranded helix. Most regulatory models assume that each Tpm molecule moves between three discrete orientations: blocked, closed, and myosin bound, originally called the “open state” (McKillop and Geeves, 1993; Lehman et al., 2000; Pirani et al., 2005; Poole et al., 2006). These Tpm positions are defined by TnI bound to actin, a free chain interacting with actin, and a chain displaced by strongly bound myosin heads, respectively. Calcium regulation of these processes is defined by calcium binding or dissociation from cardiac TnC and change in the affinity of TnI to actin surface induced by a conformational change in the Tn molecule.

### Calcium-dependent kinetics of Tpm–Tn complex with actin in cardiac muscle

Ca\(^{2+}\) regulation of the interaction of the Tpm–Tn complex with actin is defined by open and closed allosteric states of TnC with one Ca\(^{2+}\) ion bound in cardiac muscle or with two Ca\(^{2+}\) ions in skeletal muscle (McKay et al., 2000). Ca\(^{2+}\) binding to TnC exposes a hydrophobic region in the N-domain of TnC, providing a binding site for the segment of TnI, strengthening the interaction of TnC with TnI, and weakening the affinity of TnI to actin. A minimal description of the allosteric mechanism in cardiac muscle contraction and relaxation is provided by the following scheme (Fig. 2).

---

**Figure 2. Model of the cross-bridge cycle.** (A) Five-state model of the actomyosin cycle includes biochemical states consistent with observed structural states: a detached state 1, M.D.Pi; weakly bound state 2, A.M.D.Pi; strongly bound post–power stroke state 3, A.M.D; rigor-like state 4, A.M; and detached state 5, M.T. The strain-dependent state transition rates are also associated with conformational changes defining the structural conformations of myosin in each state (i.e., power stroke, \( d_p \), associated with Pi release, and stroke, \( S \), associated with the ADP release, ATP binding and cross-bridge detachment, and hydrolysis and reverse stroke). (B) Addition of a state representing interaction of myosin heads with the thick filament backbone, the so-called “parked state”, PS, denoted as state 6, into a five-state model from A. The PS is a partial M.D.Pi state with structural conformation associated with thick-filament backbone, reducing the population of the M.D.Pi state capable of binding to actin and therefore fluxes from M.D.Pi state to A.M.D.Pi state (myosin binding to actin) or the reverse hydrolysis to M.T state. The transition rate from PS to M.D.Pi, \( k_{PS} \), is assumed to be strongly dependent on \([Ca^{2+}]\), and the transition rate from M.D.Pi state to PS, \( k_{PS} \), is independent on \([Ca^{2+}]\). (C) Kinetic scheme of calcium binding to TnC and interaction of TnI with actin in cardiac muscle. Calcium binding to TnC, with the equilibrium rate \( K_{Ca} = k_{Ca}(Ca^{2+})/k_{Ca} \), forms a Ca\(^{2+}\)-Tn.C complex and reduces affinity of TnI to actin. The detachment rate of TnI from actin is defined by the equilibrium rate \( \lambda \) forming the CaTnC.TnI state. In CaTnC state, Tpm is free to move, mostly azimuthally, permitting myosin binding and force generation. The dissociation from A.TnI to TnC.TnI state without bound Ca\(^{2+}\) is slow, attenuated by \( e_x \ll 1 \), and the calcium binding to TnI into the CaTnC.TnI complex is accelerated by a factor \( 1/e_x \).
muscle includes four states (Fig. 2C), comprising two TnC closed states, where TnC has no or one bound Ca$^{2+}$ to N-terminal site II, denoted as TnC and CaTnC, and similarly two TnC open states. In the presence of Ca$^{2+}$, the open state is favored over the closed state, and TnI binds preferentially to the open TnC, forming CaTnC.TnI state. In the absence of Ca$^{2+}$, the closed form of TnC is favored, and TnI binds preferentially to actin, forming A.TnI state. The equilibrium rate constant of Ca$^{2+}$ binding to TnC closed states, $K_{Ca} = \frac{k_{Ca}}{\epsilon_{o}} \cdot [Ca^{2+}]$, is effectively defined via the forward constant, $k_{Ca} = \epsilon_{o} \cdot [Ca^{2+}]$, where $k_{Ca}$ linearly depends on $[Ca^{2+}]$ and the Ca$^{2+}$-independent dissociation constant, $k_{Ca}$. The dissociation constant of Tnl from closed A.TnI to CaTnC.TnI is defined by a first-order affinity $\lambda$ when one Ca$^{2+}$ is bound. With no bound Ca$^{2+}$, the transition from A.TnI to TnC.TnI shows slow dissociation of Tnl from actin with an attenuated rate, $\epsilon_{o}$, where allosteric fraction $\epsilon_{o}$ is $\ll 1$. On the other hand, Ca$^{2+}$ binding to TnC in a TnC.TnI complex is accelerated to $k_{Ca}/\epsilon_{o}$, keeping the population of the A+TnC.TnI state low.

**Thin-filament regulation of myosin interactions with actin**

The kinetics of interactions between the Tpm–Tn complexes with an actin filament are, at present, best described by the long-range cooperative continuous flexible chain (CFC) model (Geeves et al., 2011; Mijailovich et al., 2012) that includes structural constraints between Tpm–Tn regulatory units. The model is built on structural evidence that neighboring Tpms overlap and that one end of TnT is bound to a specific site on Tpm, whereas its N terminus overlaps with the adjacent Tpm, forming linked Tpm–Tpm regions. The interconnected neighboring Tpm–Tn units form the appearance of a CFC on each strand of an actin (Lorenz et al., 1993; Vibert et al., 1997) rather than a set of the independent Tpm–Tn units.

For modeling thin-filament regulation as a CFC, we follow a Monte Carlo approach with spatially explicit myosin binding to regulated actin filaments in solution (Geeves et al., 2011; Mijailovich et al., 2012) that we later expanded to the model of thin filaments in a 3-D sarcomere lattice (Mijailovich et al., 2019). The coupled algorithm for thin-filament regulation and the actomyosin cycle consists of three main steps: (1) calculation of the state transitions between actin–Tpm–Tn states as a function of $[Ca^{2+}]$, (2) calculation of mean CFC angular positions and their azimuthal angular fluctuations along the actin filament, and (3) calculation of state transitions between actin–myosin states. These processes are interrelated and regulated by the $[Ca^{2+}]$ where CFC angular position is modulated by either TnI bound to actin or dissociated from actin after calcium binding to TnC. The release of TnI from the actin surface allows Tpms angular movement, exposing myosin-binding sites on actin and thus permitting myosin binding and force generation by cross-bridges. The CFC moves azimuthally at a high frequency, and the mean CFC angular position and the thermally induced azimuthal fluctuations are defined by the instantaneous spatial positions of the pinning sites, at which the TnI and myosin are bound to actin (Geeves et al., 2011; Mijailovich et al., 2012; Mijailovich et al., 2019). General definitions, calculations of CFC angular position and its variance, and the coupling between the calcium-regulated position of the CFC and myosin cycle states are described in Appendix B.

**Thick-filament regulation by calcium**

There is now substantial evidence for a myosin off state in the relaxed structure of the thick filament in which the two myosin heads self-associate and pack down onto the thick filament to form what is referred to here as the “PS.” This state is denoted as a disordered relaxed state (DRX) associated with the thick-filament backbone (Anderson et al., 2018; Brunello et al., 2020) or an ordered SRX state (McNamara et al., 2016). The kinetics of the equilibrium between the PS and the on state of myosin on the thick filament is necessary to define how the filament is activated and relaxed and how the transition between these thick-filament states is modified by mechanosensing in thick filaments (Linari et al., 2015; Irving, 2017), interfilament mechanical signaling by cardiac MyBP-C (McNamara et al., 2016; Brunello et al., 2020), and phosphorylation of the regulatory light chain (RLC) and cardiac MyBP-C (Kampourakis et al., 2016) and, perhaps, titin.

In the absence of precise information concerning the molecular details of each of these prospective mechanisms, we began by implementing a minimal version of the PS. We treated each myosin head as an independent force generator that can isomerize from the M.D.Pi state to the PS. For reducing high levels of tension at low [Ca$^{2+}$] (Prodanovic et al., 2020), we modeled the PS to be activated by Ca$^{2+}$ association with a component of the thick filament so that the activation follows the Ca$^{2+}$ transient. This idea is supported by the observation of Huxley et al. (1994) that, during fast Ca$^{2+}$ activation, myosin heads move radially away from the thick-filament backbone much faster than they attach to the thin filament. Thus, this process could be attributed to a [Ca$^{2+}$]-dependent transition from the PS to M.D.Pi state that can reduce resting tension at low levels of Ca$^{2+}$ and accelerate tension relaxation when [Ca$^{2+}$] falls. Alternative hypotheses are considered in the Discussion section.

The full implementation of a cross-bridge model including a PS in the MUSICO platform requires knowledge of the equilibrium between the PS and the myosin on state on the thick filament (Fig. 2B). In the absence of strong experimental evidence regarding the mechanism defining the transition from or into the PS, for simplicity, we consider the rate constant $k_{PS}$ that could potentially depend on [Ca$^{2+}$]. We define the rate $k_{PS}(Ca)$ in the form of a Hill curve:

\[ k_{PS} = k_{PS}(Ca^{2+}) = \frac{k_{PS}}{k_{PS}^{MAX} + \frac{[Ca^{2+}]^{b}}{[Ca^{2+}]^{b}}} \]

where $k_{PS}$ is the baseline rate, $k_{PS}^{MAX}$ is the rate at high calcium (e.g., at pCa = 4.6), $b$ is the Hill coefficient of the rate sigmoidal $k_{PS}$ rise, and $[Ca^{2+}]_{50}$ is $[Ca^{2+}]$ when $k_{PS}$ is equal to $k_{PS}^{MAX}/2$. The reverse rate, $k_{PS}$, is assumed to be constant.

**Monte Carlo simulations of rate-dependent stochastic processes**

In Monte Carlo simulations, we used the standard Metropolis algorithm, where a kinetic transition in a time step $\Delta t$ occurs...
when a random number in \((0, 1)\) lies in the range \((0, k\Delta t)\), where \(k\) is state transition rate constant. \(\Delta t\) must be much smaller than the inverse of the fastest rate constant of the system, \(k_{\text{max}}\) and, in practice, \(k_{\text{max}}\Delta t < 0.001\) was sufficient to achieve satisfactory statistics and avoid interference between multiple transitions within a single subsystem and the negligibly small interference between the subsystems.

The coupling between thin-filament regulatory processes and the actomyosin cycle requires two sets of Monte Carlo random number drawings within each time step. The first set of random number drawings defines transitions of TnI–actin states from the calcium kinetics scheme in Fig. 2C, depending on \([\text{Ca}^{2+}]\). The position of bound TnI to actin after updating and bound myosins from the previous time step set the chain configuration for obtaining the mean CFC angular position along the actin filament strand and its azimuthal variation. The second set of random number drawings defines the changes in actomyosin states regulated by the CFC, setting the configuration for the calculation of the mechanical equilibrium with external forces and constraints.

For each TnI or cross-bridge, we use one Monte Carlo random number drawing to define whether the TnI or cross-bridge remains in its current state or will change its state into one of the other possible states within the current time step \(\Delta t\). The change of state is defined by the probability, in the range from 0 to 1, that is divided into probability bins, \(P_{ij}\), in a specified order, including the set of probability bins associated with a TnI or cross-bridge changing state and a bin associated with the probability of remaining in the current state. Depending which bin the drawn random number falls into, the fate of a particular TnI or cross-bridge is defined and set for the following time step. The calculations of the transition probabilities from the transition rates between TnI–actin states and between actomyosin states in the 3-D sarcomere lattice are as described previously (Mijailovich et al., 2012; Mijailovich et al., 2016; Mijailovich et al., 2019). The coupling between cross-bridge cycling and calcium regulation of the thin filament is described in Appendix C.

**Stochastic process, model size, and myofibril edge effects**

Due to stochastic transitions in the cross-bridge cycle, the force per myosin filament can vary between the filaments by ±15% at any instant of time and also by about the same amount in the same filament over time. In contrast, the variation in average force per myosin filament is small; for example, if the number of myosin filaments matches the number of filaments in an average myofibril, the fluctuations in isometric force are minimal (<0.1%) and reflect observed tension variations.

In the parameter exploration phase, we limited the stochastic simulations to a half-sarcomere with 200 myosin and 400 actin filaments that is approximately one-third of the number of filaments in a cross-section of a typical myofibril. This number of filaments is sufficient for stable simulations and statistical averaging without requiring running the simulation multiple times. In the final simulations, we increased the number of myosin filaments per half-sarcomere to 500 in order to get realistically smooth tension responses.

**Procedure for estimation of model parameters**

The MUSICO platform is formulated on three principal pillars: the 3-D explicit (multi)sarcomere geometry, a strain-dependent cross-bridge cycle, and calcium-dependent thin- and thick-filament regulation. This model provides the precision necessary to quantify the effects of kinetics and the structural changes, associated with mutations in contractile and regulatory proteins, on muscle function, but it requires a large number of model parameters to properly define muscle fiber structure, cross-bridge kinetics, and thin-filament regulation by Ca²⁺. We contend that conventional fitting of the data is not ideal, because the published experimental data show that large variations and fitting many free parameters can result in unreasonable estimates. Thus, we used established structural data and best estimates deduced from experimentally established kinetic data to define as many parameters as possible (Table 1).

In this study, we then varied only a few parameters depending on the particular sets of data that model prediction had to match. In most of the cases, we adjusted (1) the myosin–actin binding rate, \(k_{\text{bi}}\), to account for changes of interfilament spacing, high speed of sarcomere shortening, and possible myosin activation by stretching thick filaments via titin at longer SLs; (2) the power stroke Gibbs free energy change, \(\Delta G_{\text{stroke}}\), and the ADP release rate, \(k_{\text{ADP}}\), because these parameters are load dependent and the degree of shortening and sarcomere disorder is difficult to define, particularly during relaxation; (3) the kinetics of \(\text{Ca}^{2+}\)-binding, \(\dot{k}_{\text{Ca}}\), and dissociation, \(k_{\text{Diss}}\), from TnC that reciprocally defines interaction of TnI with actin; and (4) the \([\text{Ca}^{2+}]\) for the half-maximal rate from PS, \([\text{Ca}^{2+}]_{\text{PS}}\), a key parameter defining the state transition rate from PS, \(k_{\text{PS}}([\text{Ca}^{2+}]_{\text{PS}})\), due to large changes in the cytosol from intact cells to the bathing solution in demembranated muscles. The parameters used in simulations are described in detail in Appendix D, and the complete set of parameters used in simulations of trabecular twitch responses to calcium transients is shown in Table 1. In a few simulations, different parameters were used to accommodate large differences in experimental conditions, such as a large change in interfilament spacing, \(d_0\), due to demembranation of trabeculae, or to achieve good fits of the relaxation phase of the twitch contractions that must account for inhomogeneous lengthening between sarcomeres.

A sensitivity analysis is typically needed when fitted parameters have a large range of possible values (Mijailovich et al., 2010; Mijailovich et al., 2012; Ujfalusi et al., 2018). In this study, we also performed a sensitivity analysis of the parameters \(k_{\text{bi}}\), \(k_{\text{ADP}}\), \(k_{\text{PS}}\), \(\Delta G_{\text{stroke}}\), \(\dot{k}_{\text{Ca}}\), and \([\text{Ca}^{2+}]_{\text{PS}}\) and b in \(k_{\text{PS}}([\text{Ca}^{2+}]_{\text{PS}})\), which were necessary to be estimated (usually no more than three, depending on specific case) to match all experimental data. The sensitivity analysis of these and other important parameters is shown in the supplemental text (see bottom of the PDF and Figs. S8–S13 and S16–S23). We used the sensitivity analysis of a set of free parameters to estimate parameters for the next simulation by methodology described by Mijailovich et al. (2010). From those estimates, the changes in parameters were filtered, using our experience, in order to avoid imposing changes in parameters that marginally affect the predicted response and therefore move the next model prediction more quickly toward the
observation. Using this approach, we achieved good fits with a minimal number of long simulation runs.

The adjusted parameters that differ from data in Table 1 are shown in comparative tables and in the figure legends associated

### Table 1. MUSICO parameters for the simulations of twitch contractions in intact rat trabeculae at fixed length and at temperature 27.2°C in Fig. 5 (Caremani et al., 2016) and at 27.5°C in Fig. 6 (Janssen et al., 2002)

| Description                                                                 | Parameter | Isometric trabeculae | Notes                                      |
|------------------------------------------------------------------------------|-----------|-----------------------|--------------------------------------------|
| Cross-bridge cycle                                                          |           |                       |                                            |
| Myosin–actin binding rate                                                   | $k_{ca}$  | 226 s$^{-1}$          |                                            |
| Myosin–actin detachment rate$^a$                                             | $k_a$     | 46 s$^{-1}$           |                                            |
| Myosin stroke cap rate                                                       | $k_{cap}^+$ | 1,000 s$^{-1}$   | Mijailovich et al., 2016, 2019             |
| Myosin reverse stroke cap rate                                               | $k_{cap}^-$ | 100 s$^{-1}$    | Mijailovich et al., 2016, 2019             |
| Power stroke Gibbs energy change                                            | $\Delta G_{stroke}$ | $-13 k_BT$       | Mijailovich et al., 2016, 2019             |
| Working stroke                                                               | $d$       | 10.5 nm               | Duke, 1999; Mijailovich et al., 2016, 2019 |
| Second working stroke                                                        | $\delta$  | 1 nm                  | Mijailovich et al., 2016, 2019             |
| ADP release rate$^a$                                                         | $k_{D}$   | 60 s$^{-1}$           |                                            |
| ATP binding and myosin detachment$^a$                                       | $k_T^+$   | $10^6$ s$^{-1}$      |                                            |
| Hydrolysis forward rate$^a$                                                  | $k_H^+$   | $10^6$ s$^{-1}$      |                                            |
| Hydrolysis backward rate$^a$                                                 | $k_H^-$   | $10^5$ s$^{-1}$      |                                            |
| Cross-bridge stiffness                                                       | $\kappa$  | 1.3 pN/nm             | Duke, 1999; Mijailovich et al., 2016, 2019 |
| $k_BT$ at 27.5°C                                                             | $k_BT$    | 4.147 pN/nm           |                                            |
| Calcium kinetics                                                             |           |                       |                                            |
| Calcium binding to TnC equilibrium rate                                       | $\tilde{K}_{Ca}$ | 1 µM$^{-1}$   | Smith and Geeves, 2003; Geeves et al., 2011 |
| Calcium binding rate to TnC                                                  | $K_{Ca}$  | 75.4 µM$^{-1}$s$^{-1}$ | Smith and Geeves, 2003; Geeves et al., 2011 |
| Calcium dissociation rate from TnC                                           | $k_{Ca}$  | 75.4 s$^{-1}$        | Kreutziger et al., 2011; Wang et al., 2012, 2013 |
| TnI–actin equilibrium rate constant at high Ca$^{2+}$                       | $\lambda$ | 10                   |                                            |
| TnI–actin backward rate constant                                             | $\lambda$ | 375 s$^{-1}$         |                                            |
| Allosteric TnC–TnI–actin–Ca$^{2+}$ parameter                                 | $\epsilon_o$ | 0.01           | Smith and Geeves, 2003; Geeves et al., 2011 |
| CFC                                                                          |           |                       |                                            |
| Tpm pinning angle                                                            | $\phi$   | $-25^\circ$           | Poole et al., 2006                        |
| Myosin Tm angular displacements                                              | $\phi_+$  | $10^\circ$           | Poole et al., 2006                        |
| Angular SD of free CFC                                                       | $\sigma_0$ | 29.7$^\circ$   | Pirani et al., 2005; Mijailovich et al., 2012 |
| Persistence length of Tm–Tn confined chain                                   | $1/\xi$  | 50 nm                 | Mijailovich et al., 2012                   |
| PS                                                                           |           |                       |                                            |
| Transition rate to PS                                                        | $k_{PS}$  | 200 s$^{-1}$          | Assumed                                   |
| Baseline rate                                                                | $k_{PS}^0$ | 5 s$^{-1}$          | Assumed                                   |
| Amplitude                                                                   | $k_{PS}^{max}$ | 400 s$^{-1}$ | Assumed                                   |
| Calcium Hill function slope                                                 | $b$       | 5                     | Assumed                                   |
| Half-activation point of the Hill function [Ca$^{2+}$]$^{50}$                |           | 1 µM                  | Assumed                                   |
| Sarcomere                                                                    |           |                       |                                            |
| Length of sarcomere                                                          | $SL$      | 2.2 µm                | Robinson and Winegrad, 1977, 1979          |
| Length of actin filament                                                     | $L_s$     | 11 µm                 | Robinson and Winegrad, 1977, 1979          |
| Interfilament spacing                                                       | $d_{ij}$  | 33.83 nm              | Irving and Maughan, 2000                   |
| Thin-filament elastic modulus                                               | $AE_a$    | 65 nN                 | Huxley et al., 1994; Kojima et al., 1994   |
| Thick-filament elastic modulus                                              | $AE_{cm}$ | 132 nN                | Huxley et al. 1994                        |

$^a$Based on mouse and human α-myosin values in (Deacon et al., 2012a, 2012b) with corrections for temperature; ionic strength is as documented in (Mijailovich et al., 2017).
with the specific simulations where the adjustments were necessary.

The MUSICO software environment and simulation details
The MUSICO software was developed as a C++ object-oriented application that includes the LAPACK linear algebra package and the deal.II finite element library. In parts of MUSICO, where parallelization provides a significant reduction in computational time, OpenML is implemented. Typical run times for these simulations depend on the number of actin and myosin filaments. The simulation of 200 myosin filaments over 1 s with a time step of 1 µs requires ~10 h on the AEG IS04-KG grid site, consisting of six nodes, each equipped with two AMD Opteron 6276 16-core processors and 96 GB RAM, totaling 192 processors.

Online supplemental material
The supplemental text includes (i) the spatial positions between myosin heads and binding sites on the thin filament in 3-D sarcomere lattice; (ii) the correction of Caremani et al. (2016) observed tension under isometric half-sarcomere length due to the loss of trabecular length control during the sarcomere-lengthening phase of a twitch; (iii) how the accelerated relaxation can be achieved by variation of parameters in the five-state model; (iv) measurements of passive SE compliance during twitch in isometric trabeculae; (v) ATPase consumption during twitch contractions in fixed-length trabeculae; (vi) sensitivity analysis of twitch tension to model parameters in fixed-length trabeculae; (vii) estimation of the active tension increase due to decrease of titin passive tension during twitch contractions in fixed-length trabeculae where large changes in SL are observed (Caremani et al., 2016); and (viii) sensitivity analysis of twitch tension transients to model parameters in the absence of a PS in isometric half-sarcomere. The following supplemental figures are available online in a ZIP package. Fig. S1 shows the interaction between myosin heads and actin filaments in three dimensions defined by the triple-helical arrangement of myosin molecules along the myosin filament and the double-helical arrangement of monomers (myosin-binding sites) along the actin filaments. Fig. S2 shows the effect of the five-state cross-bridge model rate $k_C^D$ on relaxation of twitch tension in isometric half-sarcomeres (IsoSarc). Fig. S3 shows the SL controlled isometric tension of a half-sarcomere. Fig. S4 shows the significance of the PS in twitch relaxation phase. Fig. S5 shows the comparative tension–displacement loops obtained from Janssen and de Tombe, 1997; Caremani et al., 2016. Fig. S6 shows a comparison of ATP consumption rate and ATPase during twitch contractions of isometric half-sarcomere and fixed-length trabeculae. Fig. S7 shows that ATPase is strongly correlated to strations of isometric half-sarcomere and fixed-length son of ATP consumption rate and ATPase during twitch con-

Fig. S8 shows the changes in twitch tension from Fig. 5 B by ±20% variation in baseline rate from the PS. Fig. S14 shows active tension corrected for the decrease in titin force due to shortening of a half-sarcomere. Fig. S15 shows the effect of allosteric TnC–Tnl–actin–Ca$^{2+}$ interaction parameter, $e_0$, on resting tension at observed baseline calcium level and when [Ca$^{2+}$] was reduced to 0.05 µM. Fig. S16 shows changes in twitch tension predicted by the five-state model for change in $k_C^D$ for up to ±50% from those used for Fig. 3, denoted as original $k_C^D$. Fig. S17 shows changes in twitch tension predicted by the five-state model for change in $k_0^D$ or up to ±50% from those used in Fig. 3, denoted as original $k_0^D$. Fig. S18 shows changes in twitch tension predicted by the five-state model for change in $k_{ii}$ for up to ±50% from those used in Fig. 3. Fig. S19 shows changes in twitch tension predicted by the five-state model for change in $k_d$ for up to 24-fold from those used in Fig. 3. Fig. S20 shows changes in twitch tension at isometric half-sarcomere obtained from simulations by the five-state model. Fig. S21 shows changes in twitch tension relaxation rates with increase of $k_C^D$ predicted by the five-state model. Fig. S22 shows the best fit to the observations with the five-state model requiring the change in power stroke rate $k_{p1}$. Fig. S23 shows the effect of changes in baseline calcium level on resting and twitch tension. Fig. S24 shows a comparison of tension–pCa relationships between the cross-bridge cycle models with or without PS (i.e., the six-state and five-state models).

Results
Isometric fixed end twitch contractions in cardiac muscle result from Ca$^{2+}$ transients that lead to transient tension responses and SL changes. In intact trabeculae, this involves the combined effects of multiple underlying mechanisms acting together. Thus, to test the effects of two novel features of our new model, namely thick filament activation during contraction and the series elasticity observed in cardiac muscle, we first address them separately. To establish the kinetic parameters of the Ca$^{2+}$ activation scheme and the cross-bridge cycle, we first simulated twitches in a single isometric half-sarcomere. Then, using these parameters, we simulated twitches of a fixed-length trabecula using a multisarcomere system in series that contains an elastic component to predict changes in the SLs.

Why a PS is necessary for twitch contractions of an isometric sarcomere
We tested the ability of the five-state cross-bridge model (Fig. 2 A), coupled to a Ca$^{2+}$ regulatory scheme (Fig. 2 C), to simulate twitch contractions in an isometric half-sarcomere using MUSICO. There are only a few published trabecular twitch experiments where SL was kept approximately constant (Janssen and de Tombe, 1997; Caremani et al., 2016). The experiments of Janssen and de Tombe recorded Ca$^{2+}$ and tension transients at 22°C, but the data have an unusual flat top to the tension transient, which is not typical of the transients recorded in the literature and is hard to describe in simulations (see Fig. S4; and Niederer et al., 2006). Caremani et al. recorded tension but not
Ca\(^{2+}\) at 27.2°C. To provide the Ca\(^{2+}\) transient for the Caremani et al. experiment, we adapted the calcium transient of Janssen et al. (2002) collected at similar temperature (i.e., 27.5°C) during twitch contractions in fixed-length trabecula experiments. The transients typically achieve a higher peak [Ca\(^{2+}\)] in fixed-length trabeculae, where internal shortening can occur, than in isometric sarcomere twitches where feedback length is controlled (Janssen and de Tombe, 1997). We rescaled the adapted Ca\(^{2+}\) transient for the Caremani et al. trabecula to the isometric half-sarcomere using the ratio of peak Ca\(^{2+}\) recorded in both protocols by Janssen and de Tombe, 1997 at the lower temperature (for details, see Appendix D).

In the isometric half-sarcomere twitches at 27.2°C observed by Caremani et al. (2016), the length of half-sarcomeres was well controlled during the rising phase of tension (Fig. S3). However, it was far more difficult to control tension during relaxation, where inhomogeneity in the sarcomeres can occur, leading to differential shortening and lengthening of sarcomeres. We considered whether a loss of SL control could induce a discrepancy in the recorded tension. Specifically, the Caremani et al. tension transients showed some lengthening at the sarcomere level during the relaxation phase (Fig. S3). This lengthening increased tension in a passive parallel elastic (PE) component that is not present if the half-sarcomere is kept isometric. Consequently, the adjusted tension for the effect of PE (dark gray line in Fig. 3) had faster relaxation and lower resting tension than the observed tension in the Caremani et al. experiment.

The obtained prediction of the twitches using our five-state model shows a good match to the peak tension, but tension rise was slightly slower, and there was a poor overall fit to the relaxation phase (dark green line in Fig. 3), with the fall in tension being too slow, compared with the observed tension transient (dark gray line in Fig. 3). The parameters used in the simulations of the isometric half-sarcomere are listed in Table 1, except parameters \(k_{A1}^C, k_{A1}^T,\) and \(k_{D1},\) which are presented in the text for the five-state model and in Table 2 for the six-state cross-bridge cycle. For comparison, mean force per myosin filament and corresponding (observed) muscle tension (Exp.) are shown on the vertical axes. For consistency in all plots, resting tension is subtracted; thus, the responses to calcium transient display net change in active tension.
We further explored the possibility of various combinations of parameters (see Figs. S16, S17, S18, S19, S20, S21, S22, and S23) so that model predictions with the five-state model match the observed tension responses. The best match of MUSICO five-state model predictions with experimental observations was achieved by varying values of \( k_{A_r} \), \( k_{i_0} \), and shifting forward power stroke rate \( k_{23}(x) \) for \( \Delta x_e \) 0.5 nm (Fig. S22). The shift (see inset in Fig. S22) effectively increased the number of cross-bridges that can complete the power stroke and go through a full ATPase cycle. The latest step increased the cross-bridge detachment and rate of tension relaxation, enabling the model predictions to match the observations (Fig. S22). However, this was only achieved by a reduced level of baseline [Ca\(^{2+}\)], as used in Fig. 3 for simulations with the five-state model. Moreover, the simulations with the observed Ca\(^{2+}\) transients showed the high resting tension and slightly higher tension peak, and they slowed the rate of tension relaxation (Fig. S23). We further explored other values of parameters, but we were not able to reduce the resting tension. The reason is shown in Fig. S24, where the tension-pCa relationship, obtained using the parameters that achieved very good fit to the observation (displayed in Fig. S22), shows increased Ca\(^{2+}\) sensitivity at low [Ca\(^{2+}\)]. Importantly the tension at baseline [Ca\(^{2+}\)] shown in Fig. S24 is approximately the same as the resting tension in Fig. S23 at the same [Ca\(^{2+}\)].

An alternative way to reduce the number of available M.D.Pi cross-bridges, accelerate relaxation, and lower resting tension is to incorporate a regulated thick-filament relaxed state (see Fig. 2 B). Such a relaxed state of the thick filament, called the “PS,” has been widely discussed in the literature in recent years, and the background to this is introduced in the Materials and methods section (Linari et al., 2015; McNamara et al., 2015; Irving, 2017).

There are several ways in which a relaxed thick filament could be regulated and implemented in our model, as outlined in the Materials and methods section. The PS could be regulated via strain in the thick filament, via phosphorylation of the RLC and MyBP-C, or Ca\(^{2+}\)-dependent association/dissociation from the thick filament backbone. To implement any of these in the spatially explicit 3-D model, we would need to know details of myosin head packing onto the thick filament, the degree of cooperativity in the transition between the packed and active states, and any necessary differences between, for example, the C and D zones of the thick filament, as suggested by Brunello et al. (2020). In the absence of direct experimental evidence for a detailed thick-filament activation mechanism, and for simplicity, we made the PS Ca\(^{2+}\) sensitive with a Ca\(^{2+}\) affinity of 1 µM. This allowed thick-filament regulation to mirror that of the thin filament (see Eq. 1 in the Materials and methods section for a definition of Ca\(^{2+}\)-dependent state transition rates). The simulations using the six-state model (red line in Fig. 3) substantially improved matching of the observed experimental resting tension and tension transients, and, importantly, the relaxation time course was close to observations. The parameters used in our simulations of isometric sarcomere experiments that were different from those listed in Table 1 are shown in Table 2.

We also did attempt to fit the transient of Janssen and de Tombe (see Fig. S4). We could generate a reasonable description of tension rise and peak tension; however, the relaxation phase was inconsistent with the observed tension trace by Janssen and de Tombe. This could be consistent with a problem controlling the SL during relaxation similar to that of Caremani et al. The parameters used in these simulations were adjusted for the 5.2°C-lower temperature than that used by Caremani et al., and they are listed in Table 2.

Importantly, simulations with the cross-bridge model with the PS showed significantly reduced resting tension and had little effect on the rate of tension rise or peak tension, but they accelerated the relaxation phase and resulted in a near-perfect fit to the adjusted tension transient (Fig. 3, red line). Thus, this six-state model was able to effectively simulate relaxation, which has been a major challenge in modeling data where tension is not normalized, because detaching myosin heads entering the PS was not available to reattach and contribute to the maintenance of tension.

**Tension and SL responses to Ca\(^{2+}\)-evoked twitch transients in fixed-end trabeculae**

As noted above, sarcomeres are not isometric during an observed “fixed-end” trabecular twitch, as shown by Janssen and de Tombe, 1997; Caremani et al., 2016, who simultaneously measured the tension and half-sarcomere displacement during the twitch. The sarcomere-level shortening and lengthening due to trabecular compliance affects the time course and magnitude of the observed tension (Caremani et al., 2016). The observed tension-displacement loop shows higher tensions during sarcomere shortening and lower tensions during sarcomere lengthening (Fig. 4 A, orange circles). Accordingly, sarcomeres shortened during tension rise and lengthened during relaxation. The next step in development of the model was to incorporate a compliance originating from the series elasticity of cardiac muscle structural components (Fig. 1).

For simplicity, we derived a nonlinear serial spring-like elasticity (SE) from the average loop tension–displacement relationship (gray line in Fig. 4 A). The difference of tension (orange circles) to the mean (gray line) in Figs. 4 A showed an approximately symmetric shape along the path of the loop (Fig. 4 B), reflecting the viscoelastic effects due to shortening and lengthening velocities. A similar relationship was derived from the Janssen and de Tombe experiments (see Fig. S5), where the magnitudes of the peak tensions and displacements show about the same values, but viscoelastic effects were a little different, more so during relaxation.

Caremani et al. did not observe [Ca\(^{2+}\)] transients for the fixed-length trabecula at 27.2°C. Therefore, as for the isometric sarcomere twitches above, we used the time course of the [Ca\(^{2+}\)] transient from Janssen et al. (2002) at 27.5°C, but with a reduced peak from 1.51 to 1.20 µM to match the lower tension at the lower temperature. The corrected [Ca\(^{2+}\)] trace, proportionally scaled down from Janssen et al., is shown as a pink line in Fig. 5 B.

The tension and [Ca\(^{2+}\)] transients shown on Fig. 5, A and B, reflect the change in temperature between the two conditions. In all simulations, we used the six-state model with the same parameters as for the isometric sarcomere data in Fig. 3, and the nonlinear trabecular SE stiffness (Fig. 4 A and Fig. S5).
MUSICO simulations did not produce a good match to the observed transients (not shown). To produce a match to the tension transient, three parameters needed to be adjusted: the forward and reverse rates for myosin attachment to actin $k_{A}^{o}$, $k_{A}^{-}$, and the power stroke free energy change, $\Delta G_{\text{stroke}}$. Their values used are listed in Table 2. The simulations, shown in Fig. 5, A and B (green line), describe the tension transients (red line) quite well in both the rising and relaxing tension phases. However, during tension relaxation in both experiments, the predicted half-sarcomere lengthening was slower than experimentally observed.

A principal cause of the inability to fit both tension and displacement could be the oversimplified nonlinear stiffness used for the SE component, which neglects the hysteresis in the observed tension–displacement data caused by the viscoelastic effect shown in Fig. 4 A and Fig. S5. However, the model predictions were able to describe tension and the size of the SL changes in fixed-length trabeculae (Fig. 5). Because measurements of tension are more precise and more reliable than measurements of displacements, in the following analyses and comparisons, we focus on the tension transients.

The adjustments required to simulate the fixed-length trabecula versus isometric sarcomere tension transients were similar at both temperatures used (Table 2). The values of $k_{A}^{o}$ and $k_{A}^{-}$ increased approximately twofold in each case (range, 1.8–2.2-fold), whereas $k_{i0}$ remained unaltered and $\Delta G_{\text{stroke}}$ increased by ~15%. This adjustment can be accounted for to a significant extent by the shortening/lengthening of sarcomeres that occurs during the twitch. This change in structure affects the number of accessible actin-binding sites for myosin through changing the alignment of the thick and thin filaments and alterations to the strain-dependent rate constants ($k_{A}, k_{A}^{-}, k_{A}^{o}$). Note that because both $k_{A}$ and $k_{i0}$ change by the same amount, there is no change in the equilibrium constant of M.D.Pi binding to actin. For a sensitivity analysis of the adjusted parameters, see Figs. S8, S9, S10, S11, S12, and S13.

It is interesting to note that lengthening and shortening also result in significant changes in the cross-bridge cycling rates (ATPase rates; Figs. S6 and S7), as expected from the Fenn effect (Fenn, 1924). Length effects on Tn Ca$^{2+}$ release/uptake rates have also been reported (Kataoka et al., 2007). Moreover, it is worth noting that correcting for the 5.2°C temperature change between the two datasets was identical for the isometric sarcomere and the fixed-end twitches. This is consistent with the temperature effect being due to the temperature sensitivity of the underlying molecular events (actin binding and ADP release).

To test our model further, we took a third independent fixed-length rat trabecular twitch dataset from Janssen et al. (2002) collected at three different temperatures (see Fig. 6). These data were collected under conditions similar to the earlier measurements and indicate a slight increase in peak tension, as well as marked acceleration of both tension rise and relaxation with a temperature increase from 22.5°C to 30°C (Fig. 6, red lines). Note that these transients are not identical to those of Fig. 5 because the [Ca$^{2+}$] transient has a slightly higher peak and hence also a higher peak tension. This provides a reasonable test of the precision of our model. For simulations of tension transients at the three temperatures (Fig. 6, green lines), only three parameters needed to be adjusted (Table 2). Our simulations used the identical set of parameters, including the serial elastic element parameters, used for the 22°C (Janssen and de Tombe, 1997) and 27.2°C (Caremani et al., 2016) datasets (Table 1 and Table 2). The only exception is a slightly increased rate of $k_{A}^{o}$ from 140 to 149 s$^{-1}$ for an increase in temperature of 0.5°C (from 22 to 22.5°C). The simulations at temperatures paralleling the previous two experiments (Fig. 6, green lines) show an excellent match to tension traces for the Janssen et al. (2002) experiments (Fig. 6 A, red lines). Also, the predicted tension-[Ca$^{2+}$] loops (solid lines) matched the observations (circle symbols) at all three temperatures (Fig. 6 B). Notably, further increasing temperature to 30°C also produced a modest increase in peak tension and in the rates of tension rise and relaxation. The simulations required a further increase of parameters $k_{A}^{o}$, $k_{A}^{-}$, and $k_{i0}$ by factors of 1.13, 1.4, and 1.5, respectively, for the 2.5°C temperature increase (from 27.5 to 30°C). This is in line with those required for the ~5°C temperature change from 22 to 22.7°C. Thus, this independent dataset with higher Ca$^{2+}$ transients and higher peak tensions required no special adjustments from experiments used in Figs. 3 and 5.

| XB cycle | Parameter | Isometric HS | Fixed-length trabeculae |
|----------|-----------|-------------|--------------------------|
|          |           | 22°C$^a$ | 27.2°C$^b$ | 30°C$^c$ | 22°C$^d$ | 27.2°C$^e$ | 30°C$^f$ |
| Myosin–actin binding rate | $k_{A}^{o}$ | 72 | 125 | 140 | 226 | 256 |
| Myosin–actin detachment rate | $k_{A}^{-}$ | 10 | 21 | 22 | 46 | 66 |
| Power stroke Gibbs energy change | $\Delta G_{\text{stroke}}$ | -15.3 | -15.3 | -13 | -13 | -13 |
| ADP release rate | $k_{i0}$ | 40 | 60 | 40 | 60 | 92 |

Displayed are only parameters that are different from those shown in Table 1. Rates $k_{A}^{o}$, $k_{A}^{-}$, and $k_{i0}$ are per second; $\Delta G_{\text{stroke}}$ is in units of kBT. HS, half-sarcomere; XB, cross-bridge.

$^a$Janssen and de Tombe (1997).

$^b$Caremani et al. (2016).

$^c$Janssen et al. (2002) at slightly different temperatures of 22.5°C and 27.5°C; the rate constants used for the SE component, which neglects the hysteresis in the observed tension–displacement data caused by the viscoelastic effect shown in Fig. 4 A and Fig. S5. However, the model predictions were able to describe tension and the size of the SL changes in fixed-length trabeculae (Fig. 5). Because measurements of tension are more precise and more reliable than measurements of displacements, in the following analyses and comparisons, we focus on the tension transients.

The adjustments required to simulate the fixed-length trabecula versus isometric sarcomere tension transients were similar at both temperatures used (Table 2). The values of $k_{A}^{o}$ and $k_{A}^{-}$ increased approximately twofold in each case (range, 1.8–2.2-fold), whereas $k_{i0}$ remained unaltered and $\Delta G_{\text{stroke}}$ increased by ~15%. This adjustment can be accounted for to a significant extent by the shortening/lengthening of sarcomeres that occurs during the twitch. This change in structure affects the number of accessible actin-binding sites for myosin through changing the alignment of the thick and thin filaments and alterations to the strain-dependent rate constants ($k_{A}, k_{A}^{-}, k_{A}^{o}$). Note that because both $k_{A}$ and $k_{i0}$ change by the same amount, there is no change in the equilibrium constant of M.D. Pi binding to actin. For a sensitivity analysis of the adjusted parameters, see Figs. S8, S9, S10, S11, S12, and S13.

It is interesting to note that lengthening and shortening also result in significant changes in the cross-bridge cycling rates (ATPase rates; Figs. S6 and S7), as expected from the Fenn effect (Fenn, 1924). Length effects on Tn Ca$^{2+}$ release/uptake rates have also been reported (Kataoka et al., 2007). Moreover, it is worth noting that correcting for the 5.2°C temperature change between the two datasets was identical for the isometric sarcomere and the fixed-end twitches. This is consistent with the temperature effect being due to the temperature sensitivity of the underlying molecular events (actin binding and ADP release).

To test our model further, we took a third independent fixed-length rat trabecular twitch dataset from Janssen et al. (2002) collected at three different temperatures (see Fig. 6). These data were collected under conditions similar to the earlier measurements and indicate a slight increase in peak tension, as well as marked acceleration of both tension rise and relaxation with a temperature increase from 22.5°C to 30°C (Fig. 6, red lines). Note that these transients are not identical to those of Fig. 5 because the [Ca$^{2+}$] transient has a slightly higher peak and hence also a higher peak tension. This provides a reasonable test of the precision of our model. For simulations of tension transients at the three temperatures (Fig. 6, green lines), only three parameters needed to be adjusted (Table 2). Our simulations used the identical set of parameters, including the serial elastic element parameters, used for the 22°C (Janssen and de Tombe, 1997) and 27.2°C (Caremani et al., 2016) datasets (Table 1 and Table 2). The only exception is a slightly increased rate of $k_{A}^{o}$ from 140 to 149 s$^{-1}$ for an increase in temperature of 0.5°C (from 22 to 22.5°C). The simulations at temperatures paralleling the previous two experiments (Fig. 6, green lines) show an excellent match to tension traces for the Janssen et al. (2002) experiments (Fig. 6 A, red lines). Also, the predicted tension-[Ca$^{2+}$] loops (solid lines) matched the observations (circle symbols) at all three temperatures (Fig. 6 B). Notably, further increasing temperature to 30°C also produced a modest increase in peak tension and in the rates of tension rise and relaxation. The simulations required a further increase of parameters $k_{A}^{o}$, $k_{A}^{-}$, and $k_{i0}$ by factors of 1.13, 1.4, and 1.5, respectively, for the 2.5°C temperature increase (from 27.5 to 30°C). This is in line with those required for the ~5°C temperature change from 22 to 22.7°C. Thus, this independent dataset with higher Ca$^{2+}$ transients and higher peak tensions required no special adjustments from experiments used in Figs. 3 and 5.
Predictions of tension–pCa relationship in intact muscles with parameters from twitch contraction simulations

A major challenge in simulating cardiac muscle contraction is to predict the [Ca\textsuperscript{2+}]\textsubscript{so} and Hill coefficient in the force–pCa relationship of intact muscle. We took the parameters from Table 1, and the cross-bridge kinetics at 22.5°C listed in Table 2, used to generate fits of the twitch data of Janssen et al. (2002) at 22.5°C (Fig. 6 A), to simulate values of the force–pCa relationship (Fig. 7) for fixed-length trabeculae. The simulations with the six-state model (Fig. 7, red line) matched very well the force–pCa data of Janssen et al. (2002) (Fig. 7, black filled squares), including the pCa\textsubscript{50} and Hill coefficient (n\textsubscript{H}) values. The simulations also fit the intact muscle tension–pCa data from Gao et al. (1994) (Fig. 7, gray triangles), which were almost identical to those of Janssen et al. (2002). It is notable that the simulations, with no change in values from those used in Figs. 5 and 6, generated not only the correct pCa\textsubscript{50} but also the observed high value of the Hill coefficient. Models that rely solely on Ca\textsuperscript{2+} regulation of the thin filament via CFC (i.e., intrinsic cooperativity) do not typically produce values of the Hill coefficient above 3 in cardiac muscle (Mijailovich et al., 2019). The higher Hill coefficient, shown in Fig. 7, is a consequence of the cooperativity assumed in the regulation of the thick filament, in addition to thin-filament mechanisms. It is important to note that other models based on cooperativity factors can achieve higher cooperativity factors (Campbell et al., 2001; Rice et al., 2008), but these are parametric models, and these kind of models cannot be compared with structurally based models such as CFC (Geeves et al., 2011; Mijailovich et al., 2012; Mijailovich et al., 2019).

Discussion

The MUSICO platform, which includes explicit 3-D sarcomere geometry, several multistate cross-bridge cycles, and Ca\textsuperscript{2+} regulation of thin-filament activation by a CFC (Mijailovich et al., 2016; Mijailovich et al., 2019), has now been further developed to allow simulations of twitch contraction in cardiac trabeculae. The key new implemented features reported here include a cross-bridge cycle with a PS that enables regulation of thick-filament activation and an SE component observed in twitch experiments in fixed-end trabeculae. In this study, we showed that both features are necessary for our model to simulate twitches and predict the tension–pCa relationship from twitches.

Our simulations show that we can describe three independent datasets for fixed-end trabecula twitches (Janssen and de Tombe, 1997; Janssen et al., 2002; Caremani et al., 2016) using the same parameters after corrections for the different temperatures. The corrections required to the three parameters (k\textsubscript{A}, k\textsubscript{D}, and k\textsubscript{P}) listed in Table 2 were modest and in line with expectations of the temperature dependence of protein reactions. Similarly, we have simulated the Caremani et al. (2016) data in which SL was held constant during the rising phase of tension at 27.2°C and the corrected tension during twitch relaxation phase for the distortions of the PE elements caused by imperfect sarcomere control during relaxation. The corrections between the isometric sarcomere and the fixed-end twitches were twofold decreases in the rates of attachment and detachment of myosin to actin, which may reflect changes in accessibility to actin sites during lengthening/shortening.

In addition to the sensitivity analysis in the supplemental material, two other features came out of the model that were not part of the initial objective. The same parameters used to describe the twitches in Figs. 5 and 6 were able to generate a force–pCa relationship observed in two previous studies (Gao et al., 1994; Janssen et al., 2002). Similarly, the same sets of parameters were able to predict the change in the ATPase/cross-bridge cycling rate during the rising and falling phases of the tension transient. The cycling rates, as expected, change due to the shortening and lengthening of the sarcomeres (Fenn, 1924).
Why a PS may be necessary for cardiac muscle and normal ventricular function

Simulations using a minimal five-state cross-bridge model, coupled with thin-filament activation by Ca\(^{2+}\) in intact muscle, presented three apparently unrelated problems: slow tension relaxation during the twitch (Fig. 3), high levels of resting tension, and unrealistic values for the myosin (cross-bridge) detachment rate, \(k_{oA}\), to bring the tension relaxation rate toward the observations (Fig. S2). High resting tension was also predicted in simulations of demembranated cardiac muscle fibers and at low Ca\(^{2+}\) (Prodanovic et al., 2020). These observations suggest a missing component in the underlying mechanisms of activation and relaxation. This conclusion is supported by recent studies indicating that, in addition to thin-filament activation, thick-filament activation may be an important process (Linari et al., 2015; McNamara et al., 2015; Irving, 2017; Reconditi et al., 2017). This is especially true when considering length-dependent activation (Ait-Mou et al., 2016; Caremani et al., 2019) and the contractile abnormalities that occur in cardiac myopathies associated with mutations in thick-filament proteins (Hooijman et al., 2011; Spudich, 2015; Nag et al., 2017; Anderson et al., 2018). The thick filament may contain one or more “off” states where myosins are not available to bind to the thin filament. For example, both structural (Zoghbi et al., 2008) and biochemical energetic studies suggest an ordered state, the so-called SRX state proposed by Cooke (McNamara et al., 2015), and a disordered state still associated with the thick-filament backbone (Anderson et al., 2018; Brunello et al., 2020) that has extremely low probability for myosin heads to bind to actin. One possibility for the transition from a myosin PS to an “on” (i.e., M.D.Pi) state could be triggered by thick-filament strain (Linari et al., 2015; Irving, 2017). This kind of mechanism has
been implemented in a model of cardiac contraction where the rate of transition from the myosin PS state to M.D.Pi state is proportional to force (Campbell et al., 2018). This model provides reasonable predictions of length-dependent activation and twitch contractions in cardiac muscle, but the model used simplified cross-bridge cycle kinetics and sarcomere geometry and did not take into account the observed significant SL changes (Janssen and de Tombe, 1997; Ferrantini et al., 2014; Caremani et al., 2016). In addition, these good fits are achieved by an empirically defined flux of thin-filament regulatory units to an on state tightly coupled to the Ca2+ transient and inclusion of a cooperativity factor that amplifies the number of on states during calcium rise and attenuating during the fall of [Ca2+] levels. The cooperativity factor was necessary to close units during tension relaxation because the thick filament is strongly activated at high tension, and relaxation was prolonged without this factor. The concept of thick-filament activation rates coupled to force should be further investigated to provide possible mechanisms. Moreover, a recent paper (Ma et al., 2018a) suggested that thick filament–based regulation is likely to be more complex than the previous studies implied and that other mechanisms of thick-filament activation could be operating during twitch contractions, as suggested, for example, in our preliminary studies (Prodanovic et al., 2020) and justified here.

In our model, the PS acts as a Ca2+-dependent buffer for myosin cross-bridges, increasing the available myosin heads on Ca2+ activation and reducing the population of M.D.Pi state during relaxation. This was achieved in the model by the transition rate out of the PS, $k_{PS}(\text{Ca}^{2+})$, having a slow rate at low [Ca2+] and a fast rate at high [Ca2+] concentrations (Eq. 1). Such a model may be supported by x-ray diffraction observations that myosin heads rapidly move radially away from the thick-filament backbone during fast Ca2+ activation of frog skeletal muscle (Huxley et al., 1994), much faster than the increase in the number of attached cross-bridges and subsequent development of tension. These unbound myosin heads in the proximity of actin filaments increase the pool of cross-bridges available to bind to the thin filament. A Ca2+-dependent transition rate from PS is one of multiple alternative possibilities for activation of the thick filament that needs to be further investigated experimentally before it is fully confirmed.

Our implementation of the PS illustrates how the presence of this PS can slow the rate of tension rise, reduce resting tension, and accelerate relaxation by limiting the number of myosin heads available to interact with actin. It follows, therefore, that if the occupancy of the PS can be manipulated by phosphorylation events (e.g., RLC, MyBP-C), small molecules (e.g., mavacamten, deoxyadenosine ADP), or mutations in myosin or MyBP-C, then alterations in all three twitch parameters can be expected. Indeed, several recent studies have suggested that many hypertrophic cardiomyopathy–linked mutations in myosin or MyBP-C cause hypercontraction by reducing the occupancy of the SRX state, allowing more myosin heads to participate in a contraction, and that this can be reversed by molecules such as mavacamten. The SRX and DRX states observed in experiments is equivalent to our PS in the model presented here. Our model may therefore be able to predict the results of mutations and small molecules on the cardiac myocyte twitch. Effects on twitch relaxation times similar to those anticipated in the absence of PS here were reported by Toepfer et al. (2020) for cardiac induced pluripotent stem cells expressing hypertrophic cardiomyopathy–linked mutations in myosin that were reversed by mavacamten.

Series elasticity of trabeculae
Fixed-length contractions of cardiac muscle can result in relatively large changes in SLs during twitch contractions (~7%; Janssen and de Tombe, 1997; Caremani et al., 2016), indicating significant compliance in intact muscle preparations. Current simulations of the cardiac twitch contraction typically use models of the half-sarcomere (though, on rare occasions, multiple sarcomeres in series) where the thick- and thin-filament

![Figure 7. Increase in calcium sensitivity in intact muscle normalized tension–pCa relationships deduced from the twitch simulations versus those observed in intact and demembranated muscles. MUSICO simulations of the tension–pCa relationships matched the observations in intact muscles by Gao et al. (1994) and Janssen et al. (2002). The muscle tension is normalized by maximum tension at fully activated muscle (T/Tmax). The parameters for MUSICO simulations are the same as in twitch simulations at 22.5°C shown in Fig. 6. However, the sensitivity in intact muscles was significantly higher than in demembranated (Demem.) muscles. The experiments performed on the same muscle before and after demembranation showed a somewhat smaller increase in the muscle sensitivity (gray arrow) than between Kreutziger et al. (2011) demembranated muscle (red arrow). Interestingly, the intact muscle tension–pCa relationships show only small differences between the Gao et al. (1994) and Janssen et al. (2002) observations. Change in [Mg2+] from 1.2 mM (gray dashed line) to the physiological value of 0.5 mM increased calcium sensitivity (green dotted line) toward that observed in intact trabeculae (green arrows; Gao et al., 1994).]
compliances account for only a small fraction of the observed compliance in the cardiomyocytes or tissue. The first attempt to model the SE component was reported by Parmley and Sonnenblick (1967), who proposed schemes of CE in series and in parallel with the elastic elements (SE and PE). This kind of model has been used in simulations of twitch contractions in trabeculae; however, prediction of a highly simplified model only showed the effect of the variation of SE and PE elements but not in milieu or experimentally observed data (Rice et al., 2008). In a more detailed model (Campbell et al., 2018), SE is apparently incorporated, but the changes in SLs reported are small and largely underestimate the observed values. It is likely that they considered only compliance of thick and thin filaments and not compliance of trabeculae. Our results presented here and elsewhere (Prodanovic et al., 2020) suggest that analysis of subtle differences in cardiac muscle dynamic contractions associated with, for example, mutations in sarcomere proteins requires a thorough description of the series elasticity observed in cardiac muscle in order to be most accurate and informative.

A simplified definition of the series elasticity, derived from the observations of Caremani et al. (2016), was used in our simulations and provided reasonable but not exact fits of tension transients and sarcomere displacements during twitch contractions of fixed-end trabeculae (Fig. 5). Specifically, it was difficult to achieve good fits for both tension and displacements, particularly during relaxation. The reasons for this may come from both experimental and model limitations. On the experimental side, measurements of SLs are obtained by different techniques, often with different outcomes. For example, Caremani et al. (2016) used a striation follower to monitor SL in the middle portion of the trabeculae, whereas Ferrantini et al. (2014) used laser diffraction. In the first case, the change in a half-sarcomere length, denoted as displacement, represents an average of half-sarcomere length changes in the middle portion of trabeculae, where the SLs are more uniform. In the laser diffraction case, the half-sarcomere length is determined as an average of all sarcomeres reflected from the laser beam. The use of different measurement techniques resulted in large variations in recorded displacement traces, such as between observations of Caremani et al. (2016) and Ferrantini et al. (2014). In addition, either of these techniques cannot fully control the inhomogeneity within these SLs during tension relaxation, particularly in the chaotic phase of relaxation (Brunello et al., 2009).

The use of simplified elastic characteristics in the model instead of the observed hysteretic loop (Fig. 3) is in part responsible for imperfect matches of displacements in simultaneous traces compared with the observations of sarcomere lengthening during tension relaxation (Fig. 5, A and B). The exact tension–length relationship (orange circles in Fig. 4 A) can be implemented in MUSICO, but we prefer the simplified model (thick gray line in Fig. 4 A). This is because the observed force loop depends on the magnitude and rate of developed tension and needs to be recorded separately for each experiment, information that is usually not available. Thus, we believe that fitting the tension traces is a more general and more robust approach than fitting displacements.

The effect of changes in calcium sensitivity on twitch contractions

The majority of studies on cardiac muscle contraction have focused on force generation and the Ca²⁺ sensitivity of steady-state force that can be relatively easily assessed as a function of [Ca²⁺] in the activation solutions. Accordingly, an abundance of experimental data in the form of force–pCa relationships provides evidence for functional changes in cardiac muscle associated with cardiomyopathies and their responses to treatment by prospective drugs. However, these data cannot be directly translated to physiological transient contractions in the living heart. Our simulations predicted the tension–pCa data of Janssen et al. (2002) and Gao et al. (1994) (Fig. 7) with the same model parameters as for the twitches at the same temperature. This is a promising step toward the possibility of interrelating the Ca²⁺ sensitivity from steady-state measurements to tension responses to Ca²⁺ transients. We believe that the reverse process is possible: to predict twitches with parameters estimated from tension–pCa relationships of cardiac muscles under similar experimental conditions. However, the problem is that most force–pCa data originate from the measurements in demembranated cardiac muscle.

The experimentally observed difference in the Ca²⁺ sensitivity of tension between intact and demembranated cardiac muscle is well recognized, but there are only a few studies that have quantitatively assessed the magnitude of these differences (Gao et al., 1994). The [Ca²⁺] used to determine the force–pCa relationship observed in demembranated muscle is well controlled, but interfilament spacing and experimental conditions such as SL, pH, phosphate, and ionic strength can be significantly different from those in intact muscle cells. To the best of our knowledge, only Gao et al. (1994) have measured the force–pCa relationship for both conditions in the same muscle (Fig. 7). The sensitivity in demembranated muscles was significantly lower than in intact muscles, and it is strongly dependent on experimental conditions (Fig. 7). The differences in experimental conditions between these preparations affecting the Ca²⁺ sensitivity, including pCa50 and cooperativity, nH, are shown in Table 3. Bringing the experimental conditions in demembranated muscles close to those observed in intact muscle reduces the differences in both pCa50 and nH between the preparations. The strongest effect is reduction of free Mg²⁺ to the physiological value of 0.5 mM that reduces the difference between intact and demembranated muscle to only ~0.2 pCa units. Interestingly, this difference in sensitivity can be attributed to the effect of the large increase in interfilament spacing, from δ₀ = 33.83 nm in intact muscle to 41.66 nm in demembranated muscle (both at SL = 2.2 μm). This increase in sensitivity is similar to that observed experimentally when the lattice of demembranated muscles is compressed back to intact muscle values with dextran (Cazorla et al., 2001; Martyn et al., 2004). Our simulations of these experiments predicted a change of 0.15–0.25 pCa units (Mijailovich et al., 2019) that can bring the sensitivity values of demembranated muscle at 0.5 mM free Mg²⁺ to the values observed in intact muscle. This is a promising result, but it is based on a single experiment, and to fully develop methodology to translate the force–pCa data from demembranated muscle to
Our current model includes the Ca\textsuperscript{2+} transient, mechanisms of cooperativity between thick- and thin-filament activation processes, series elasticity of cardiac tissue, and the force-generating cross-bridge cycle. This was sufficient to provide an accurate description of a twitch contraction of rat cardiac muscle, enabling simulations of tension and SL changes in fixed-length trabeculae as a response to intracellular Ca\textsuperscript{2+} transients. To accomplish this, two new features needed to be introduced: an SE element and a process for thick-filament myosin activation and deactivation. An elastic component that is adjustable during tension rise and fall is essential to account for the in-ternal sarcomere shortening that occurs in fixed-end “isometric” cardiac muscle contractions. Using thin-filament regulation the same kind of experiments from different laboratories with slight or no obvious changes in experimental conditions. There are also differences in preparations (e.g., intact versus demembranated trabeculae) and in modes of contraction, such as transient twitch responses and steady-state contractions usually displayed as force–pCa relationships. Here, we minimized the number of fitting parameters by using experimentally determined values whenever possible. Examples include SL and lattice spacing and other well-established experiments such as ATPase measurements and motility assays. The remaining parameters are estimated as best guesses based on experience and experimentation. However, some parameters can inherit hidden deviations due to specific conditions of the experiments that could be quite different in intact or demembranated muscles. For example, demembranated muscles show a large increase in sarcomere lattice spacing that is associated with decreases in myosin-binding rate (k\textsubscript{m}f\textsubscript{0}) and Ca\textsuperscript{2+} sensitivity compared with intact muscle. Experiments often have their own large variation, with differences in magnitude and shape of tension responses during twitches that may reflect seemingly minor differences in experimental conditions. A second large hurdle is the incompleteness of most datasets, where simultaneously measurements of tension and changes in SL during twitch contractions of trabeculae with fixed lengths and Ca\textsuperscript{2+} transients are not available. Thus, with these multiple uncertainties on the experimental side, it is exceedingly difficult to fit all experiments with the same set of input parameters. The strength of the detailed modeling approach presented here is the ability to distinguish the contributions of each effect individually to the overall recorded responses where multiple effects operate in aggregate. For example, the contribution of changes in interfilament spacing between demembranated and intact trabeculae to the Ca\textsuperscript{2+} sensitivity of tension development can be estimated from MUSICO simulations (Mijailovich et al., 2019). The estimate of such an increase in sensitivity of demembranated muscle due to an increase in lattice spacing is only a small fraction of the total increase in Ca\textsuperscript{2+} sensitivity. The remaining differences need to be attributed to other mechanisms when they are known.

Conclusions

Developing an accurate model of a twitch contraction is a serious challenge that requires multiple theoretical concepts to be integrated into simulations to account for diverse experiments. Our current model includes the Ca\textsuperscript{2+} transient, mechanisms of cooperativity between thick- and thin-filament activation processes, series elasticity of cardiac tissue, and the force-generating cross-bridge cycle. This was sufficient to provide an accurate description of a twitch contraction of rat cardiac muscle, enabling simulations of tension and SL changes in fixed-length trabeculae as a response to intracellular Ca\textsuperscript{2+} transients. To accomplish this, two new features needed to be introduced: an SE element and a process for thick-filament Ca\textsuperscript{2+} transients.

The shortening during a twitch starting from SL = 2.2 µm decreases force in titin from ∼16.5 pN per myosin filament to ∼0 at SL = 1.9 µm; thus, the active tension has slightly higher peak magnitude than reported in the literature, where initial (passive) tension is simply subtracted from total tension measured (Fig. S14).

Table 3. Hill coefficient and pCa\textsubscript{50} in intact and demembranated rat trabeculae

| Source                        | Temp. (°C) | SL (µm) | Free Mg\textsuperscript{2+} (mM) | n\textsubscript{H} | pCa\textsubscript{50} |
|-------------------------------|------------|---------|---------------------------------|------------------|----------------------|
| Intact                        |            |         |                                 |                  |                      |
| Gao et al., 1994 (Fig. 8)     | 22.0       | 2.2     | 4.87                            | 6.21             |                      |
| Janssen et al., 2002          | 22.5       | 2.2     | 4.8                             | 6.16             |                      |
| Yue et al., 1986 (Fig. 2)     |            |         | 5.59                            | 6.34             |                      |
| MUSICO                       | 22.5       | 2.2     | 5.1                             | 6.2              |                      |
| Demembranated                 |            |         |                                 |                  |                      |
| Gao et al., 1994 (Fig. 5)     | 20.0–22.0  | 2.2–2.3 | 1.2                             | 2.44             | 5.5                  |
| Gao et al., 1994 (Fig. 5)     | 20.0–22.0  | 2.2–2.3 | 1.2                             | 2.84             | 5.61                 |
| Gao et al., 1994 (Fig. 8)     | 20.0–22.0  | 2.2–2.3 | 0.5                             | 3.75             | 6.03                 |
| Dobesh et al., 2002          | 15         | 2.25    | 1.0                             | 6.9              | 5.48                 |
| Kreutziger et al., 2011       | 15         | 2.29    | 5.74                            | 5.37             |                      |

The observed values are complemented with MUSICO predictions, shown in Fig. 7, for intact muscles. Displayed in parantheses are the figure numbers in the cited references from which data were taken.

twitches, it is necessary to confirm it by stronger experimental evidence. This is additional evidence of how strongly coupled experimental data and computational simulations could translate valuable data to physiologically relevant cases.

Active tension and role of titin in length-dependent calcium sensitivity

Tension in cardiac muscle is measured as the aggregate of active and passive tension. The active tension reflects forces generated by bound cross-bridges and the passive tension as the collective contribution of titin and collagen (Cazorla et al., 2001). The passive tension is ∼0 at slack length, but at longer SLs the tension progressively increases nonlinearly (Mijailovich et al., 2019). During twitch contractions of the isometric trabeculae, a significant shortening of sarcomeres, up to 80 nm per half-sarcomere, can be observed and is due to series elasticity (Fig. 6). The shortening during a twitch starting from SL = 2.2 µm decreases force in titin from ∼16.5 pN per myosin filament to ∼0 at SL = 1.9 µm; thus, the active tension has slightly higher peak magnitude than reported in the literature, where initial (passive) tension is simply subtracted from total tension measured (Fig. S14).

Potential model limitations: variation of the model parameters, uncertainties of experimental evidence, and parameter estimation methodology

Fitting multiple sets of data presents numerous challenges. The model is imperfect (as are all models), and multiple sets of parameters can fit the desired set of data. On the other hand, there is a need to reconcile the large variability in the data obtained in

Mijailovich et al.

Computational model of cardiac twitch
alone was not sufficient to match the observed relaxation patterns. The main reason for simulations having slow relaxation was the large population of myosin heads in the M.D.Pi state. These heads can rebind to actin filament and therefore impede the relaxation process. Inclusion of a PS (either DRX or SRX state), where myosin heads interact with the thick-filament backbone, provided a mechanism to remove heads from the cycling cross-bridge pool. Our simulations also account for the effect of change in sarcomere interfilament spacing and differences in local experimental conditions (e.g., ionic strength, SL, and temperature). Thus, our simulations demonstrate the need for spatially explicit models to understand the multiple interacting processes that occur within cardiac muscle during twitch contractions. Furthermore, in future studies, the MUSICO platform developed here may provide a powerful tool to explore the effects of point mutations in sarcomeric proteins on twitch transient responses, the effect of the degree of penetration of the mutated protein(s), and the potential influence of small molecules on rebalancing altered twitch kinetics.

Appendix A

Strain-dependent rates in the actomyosin cycle

The 6-state actomyosin cycle is defined by 11 state transition rates (Fig. 2), of which five are strain dependent. For myosin binding to actin, the strain-dependent rate in quadratic form is derived from a Langevin type of equation balancing thermal fluctuations of the detached myosin molecule, elastic restoring, inertial, and viscous drag forces (Kramers, 1940; Papoulis, 1991; Hunt et al., 1994; Daniel et al., 1998). Most of the cross-bridge kinetics models used the following the equilibrium relation:

\[ K_{23}(x) = \frac{k_{12}(x)}{k_{23}(x)} = e^{-\frac{\Delta G_{bind}}{k_B T} + \frac{x^2}{2k_B T}}, \]  

\[ (A1) \]

where \( \Delta G_{bind} < 0 \) is the reduction in free energy due to myosin binding in units of \( k_B T \), \( k_B \) is the Boltzmann constant, and \( T \) is absolute temperature in degrees Kelvin. At zero strain (\( x = 0 \)), the equilibrium binding rate represents the ratio of unstrained binding and detaching rates, \( k_{12}/k_{23} = k_{A}^e/k_{A}^d = e^{-\Delta G_{bind}} \). For simplicity, the detachment rate, \( k_{A}^d \), is usually kept constant, and the attachment rate has quadratic dependence on (strain) \( x \). This approach has worked well for steady-state solutions where there is sufficient time to achieve the equilibrium cross-bridge distributions except during lengthening. During transient changes, typical for trabecular twitch contractions, sarcomeres shorten during force rise due to serial elasticity and then lengthen during force relaxation. The above approach can satisfactorily replicate transient shortening but shows, however, prolonged transition relaxation during transient lengthening. To speed up relaxation, detachment rates during lengthening should progressively increase with strain (for \( x > 0 \)). Hill explored different distributions of attachment and detachment rates that are consistent with an equilibrium rate constant \( K_{23} \) (Hill, 1974). Following his approach, we used modified attachment and detachment rates:

\[ k_{12}(x) = k_{A}(x) = k_{A}^e e^{-\frac{x^2}{2f_{da}}} k_B T \]  

\[ (A2a) \]

and

\[ k_{23}(x) = k_{A}(x) = k_{A}^d e^{-\frac{x^2}{2f_{da}}} k_B T \]  

\[ (A2b) \]

where, for \( x > 0 \), factor \( f_{da} \) increases both attachment and detachment rates but, for larger values of \( x \), increase in attachment rates becomes smaller, whereas detachment rate progressively increases. For \( x \leq 0 \), we set \( f_{da} = 1 \), leading to \( k_{23}(x) = k_A(x) = k_{A}^d \) (i.e., the formulation in standard form we used previously; Mijailovich et al., 2016; Mijailovich et al., 2019).

The transitions between the two attached states, A.M.D.Pi and A.M.D, are rapid, and this transition includes inorganic phosphate (Pi) release accompanied by a large (negative) change in chemical free energy \( \Delta G_{stroke} \), in units of \( k_B T \) and the displacement of the lever arm carrying out the power stroke, \( d \):

\[ K_{23}(x) = \frac{k_{23}(x)}{k_{32}(x)} = e^{\frac{\Delta G_{stroke} + \frac{x^2}{2k_B T}}{2k_B T}}. \]  

\[ (A3) \]

Here, \( \Delta G_{stroke} \) defines the equilibrium rate in an unstrained system, such as in experiments in solution, and factor \( \kappa d^2/2k_B T \) is free energy increase by stretch due to a power stroke from arbitrary \( x \) at state 2. The latter term shows a constant shift to higher energy at any \( x \), whereas decay of \( K_{23}(x) \) is defined by the \( k_{32}/k_B T \) term. The forward and backward power stroke rates, \( k_{23}(x) \) and \( k_{32}(x) \), can be obtained in the simplified form by assuming that the backward rate of \( k_{32} \) is independent of \( x \) but, in this formulation, forward rate \( k_{23}(x) \) reaches extremely large values with a decrease of \( x \) even to moderate negative values. Taking into account bending of the lever arm, \( k_{23}(x) \) rates are attenuated, reaching more moderate values with decreases of \( x \) (Smith et al., 2008).

To keep this pattern, the exponential rise of the forward rates and backward rates, for convenience, are capped to the assigned maximums of forward and backward constants (Smith and Mijailovich, 2008; Mijailovich et al., 2016; Mijailovich et al., 2019). For example, for assigned values of \( k_{23}^{cap} \) and \( k_{23}^{cap} \) (i.e., \( k_{23}^{cap} \), \( k_{23}^{cap} \)), and cross-bridge stiffness \( \kappa \), the forward rate including capping is defined as

\[ k_{23}(x) = k_{23}^{cap} \]  

\[ for \ x \leq x_0 \]

\[ k_{23}(x) = k_{23}^{cap} e^{-\frac{d(x-x_0)}{k_B T}} \]  

\[ for \ x > x_0 \]  

\[ (A4) \]

where \( x_0 = -0.5d - \frac{\Delta G_{stroke} + \ln(k_{23}^{cap}/k_{23}^{cap}) \kappa d}{(\kappa d)} \) is the cross-bridge strain where large exponential rates are capped. When the cap value is reached, the reverse rate \( k_{32}(x) \) changed to decay exponentially to satisfy the equilibrium constant, \( K_{23}(x) \) (Eq. A3).

For the strain-dependent ADP release (i.e., A.M.ADP → A.M), the forward rate is defined as a function of the rate of ADP release when the elastic element is relaxed, \( k_{34} \), and of the displacement that the lever arm must move to open the nucleotide pocket, \( \delta \).

\[ k_{34}(x) = k_{34} \]  

\[ = k_{34} e^{\frac{x\delta d}{2(k_B T)}} \]  

\[ (A5) \]

The backward rate, \( k_{34}(x) \), depends on ADP concentration and for small ADP concentrations can be assumed to be \( k_{34}(x) \). For ATP binding and cross-bridge detachment rates (i.e., transition between A.M. and M.T states), we assume for
now that $k_{45} = \text{constant}$, although it depends on ATP concentration, and $k_{\text{diss}} = 0$ due to a large increase in $\Delta O_{\text{ATP}}$.

The state transition rates between detached states M.T and M.D.Pi, $k_{51} = k_{15}$ and $k_{15} = k_{51}$, are independent of strain. The forward rate, $k_{15}$, involves a conformational change in the head following the opening of the 50-kD cleft leading to the recovery stroke and hydrolysis of ATP transitioning into the M.D.Pi state.

The transition of the M.D.Pi state into the FS is defined by the rate constant $k_{52}$ that could potentially depend on $[Ca^{2+}]$, myosin light chain, and MyBP-C phosphorylation. For simplicity, we first consider only the effect of $[Ca^{2+}]$ by setting, for example, $k_{52} = \text{constant}$ and $k_{52}(Ca^{2+})$ in the form of a Hill curve described by Eq. 1.

### Appendix B

**Thin-filament regulation by calcium**

**CFC angular position and its variance**

The angular displacements of the Tpm–Tn chain are defined by the function of thermally excited chain configurations defined by the Feynman path integral (Feynman and Hibbs, 1965; Smith, 2001), where the current configuration of bound TnIs and myosins at defined positions, $s$, along F-actin, the path of minimum energy represents the mean chain angle $\bar{\phi}(s)$ and the SD $\sigma_\phi(s)$ of chain angles arising from thermal excitation (Smith, 2001; Smith et al., 2003; Geeves et al., 2011; Mijailovich et al., 2012).

If there are no constraints, the mean configuration of the CFC is in the closed state sitting on the bottom of the confined potential at the angle $\phi = 0$, and the SD $\sigma_\phi$ has a value of about $\sigma_\phi = (k_4 T/8 k_{\text{CFC}} \xi) ^{1/2}$, where $k_{\text{CFC}} = k_{\text{M.T}} R^2$, $R$ is the radius at which Tpm sits on the actin filament, and $1/\xi = (4 k_{\text{CFC}}/\alpha_{\text{DP}})^{1/4}$ is the confined persistence length of the CFC (Smith et al., 2003). When the CFC is constrained, the chain is pinned at the positions where TnIs are bound to actin at an angle $\phi = 0$, forming a local blocked state, whereas the chain is assumed to be conditionally constrained at positions where myosins are strongly bound to actin, preventing the CFC from reaching angles $\phi < \phi_c$, forming a local open state. At positions where myosins are weakly bound, the chain is conditionally constrained, preventing the CFC from reaching angles $\phi < \phi_c$. The pinning and conditional constraints guarantee that the blocked, open, and closed states correspond to the three orientations seen in cryo-EM images defined by angular displacements $\phi$, $\phi_0$, and $\phi_c$. (Pirani et al., 2005; Poole et al., 2006).

### Appendix C

**The coupling between cross-bridge cycling and thin-filament regulation by calcium**

**Modulation of TnI–actin transitions by bound myosins**

In the absence of $Ca^{2+}$, most TnIs are bound to actin, but in the presence of $Ca^{2+}$, calcium binds TnC in a concentration-dependent manner. For simplicity, we assume that $k_{-1} = \lambda$, has a fixed value, whereas the binding rate of TnI to actin is a weighted function of the rate of TnI binding to actin from the unweighted closed (or open) state rate, $k_{1} = \lambda$. (Mijailovich et al., 2012). The transition probability of attachment of TnI to actin is $P_{1} = C_{1}^{\text{CFC}} \Delta \lambda$, where $C_{1}^{\text{CFC}}$ is a weight factor and the transition probability for the detachment of TnI from actin is simply $P_{-1} = \lambda \cdot \Delta \lambda$. The first Monte Carlo drawing is performed over all TnCs along all CFCs, where each actin filament has two CFC strands. If any change of TnI state is drawn, the TnI state is updated for calculation of the CFC configuration in the current time step.

**Modulation of myosin–actin transitions by the CFC**

The probabilities of changing state in the five-state (and six-state) cross-bridge model are constructed so that each state can transition to two or three neighboring states. The transition from the detached state M.D.Pi, associated with axial strain-dependent rates, includes attachment probability, $P_{12}$, adjusted for binding to multiple binding sites on actin. For the cross-bridges in the detached state, the attachment probability is shared between all reachable actin states defined as $P_{12} = [\sum_{x=1}^{x_{\text{max}}} P_{12}(\phi_x)]$, where $P_{12}(\phi_x) = \frac{k_{12}}{C_{0}} \left( \phi_x - \phi_0 \right) \cdot C_x \left( \phi_x - \phi_0 \right) \cdot C_{0} \left( \phi_0 \right) + C_{0}^{\text{CFC}}(\phi_x, \sigma_\phi) \Delta \lambda$ and $k_{12} = k_{o} f_{x, 12}$. Because $P_{12}$ is the sum of the probabilities of attaching myosin heads to each reachable sites $l = 1$ to $l_{\text{max}}$, the equivalent axial strain–dependent binding rate $k_{12}$ is set to provide the same flux as the probabilistic binding rate $k_{12}$.

The scaling factor, $f_{x, 12}$, decreases the magnitude of $k_{12}$ at each $x$ by an average number of reachable binding sites to myosin heads (Mijailovich et al., 2016). The weight factors $C_x$ and $C_{0}$ are associated with the azimuthal position of actin filaments in the sarcomere lattice relative to the myosin head, angle $\alpha$, and azimuthal angle of actin site $\beta$ (see Fig. S1 and Mijailovich et al., 2016). The weight factors $C_{0}^{\text{CFC}}(\phi_x, \sigma_\phi)$ are proportional to the fraction of time step $\Delta \lambda$ when these transitions are possible (i.e., to the weighted probabilities.
that the CFC is at positions $\phi_4^i \geq \phi_s$ that modulates weak myosin binding, denoted as subscript $M-\text{wb}$ (Mijailovich et al., 2012).

The transitions from the weakly to the strongly attached states (2 and 3) are defined by the probabilities $P_{23} = \frac{C_{\text{CFC}}^{M-\text{wb}} k_{23}(x)}{k_{23}(x) \Delta t}$ for the prestroke state 2 (A-M-D-P) and $P_{32} = k_{32}(x) \Delta t$ for the poststroke state 3 (A-M-D). Calculation of these probabilities is almost identical to that reported by Mijailovich et al. (2016), except that $P_{23}$ includes the weight factor $C_{\text{CFC}}^{M-\text{wb}}$ that modulates weak to strong myosin binding, denoted as subscript $M-\text{wb}$. The factor $C_{\text{CFC}}^{M-\text{wb}}$ is proportional to the fraction of time step $\Delta t$ when the isomerization is possible (i.e., $\phi \geq \phi_s$; Mijailovich et al., 2012).

Appendix D
Model parameters
Sarcomere geometry
The 3-D sarcomere lattice in vertebrate striated muscle is composed of myosin filaments, each surrounded with six actin filaments and each actin filament with three myosin filaments. Each half-length myosin filament is associated with six titin molecules (Mijailovich et al., 2019). Actin filaments can vary in length, depending on species, muscle type, and presence or absence of nebulin in skeletal muscle. The actin filaments have a monomer spacing of 2.7359 nm, and the half-period of one strand is 35.56 nm under relaxed conditions (Huxley et al., 1994; Wakabayashi et al., 1994; Prodanovic et al., 2016). The length of a myosin filament is $\sim 1.58$ µm, having 50 crowns (i.e., 150 myosin molecules per half-thick filament), with a crown spacing of 14.3 nm (Luther et al., 2008). This number of crowns and amount of spacing provides maximum overlap with a thin filament of $\sim 0.7$ µm. The actin radius is $r_a = 3.5$ nm, and the myosin radius is $r_m = 7.9$ nm (Mijailovich et al., 2016). The lattice interfilament spacing depends on SL, the specificities of the muscle type, and experimental conditions. The values of the spacing $d_{10}$ are specified for each set of simulations.

Interfilament lattice spacing
The interfilament spacings in intact trabeculae are much smaller than in demembranated trabeculae having values of $\sim 33.83$ nm at SL 2.2 µm. All intact trabecula fixed-length simulations used this value. In contrast, the interfilament spacings of demembranated cardiac trabecula muscle were obtained at SL 2.25 µm, having $d_{10}$ values of $\sim 41.36$ nm (Irving et al., 2000; Cazorla et al., 2001) and used in simulations of force-pCa relationships observed by Kreutziger et al. (2011). The interfilament spacing is an important factor in modulation of myosin binding to actin and possibly other rates in the cross-bridge cycle (Williams et al., 2013; Mijailovich et al., 2019).

Myofilament elasticity
Actin and myosin filaments are extensible with filament moduli (elastic modulus times cross-sectional area) derived from x-ray diffraction or direct measurement: for actin, $K_a = 65$ nN; for myosin, $K_m = 132$ nN (Huxley et al., 1994; Kojima et al., 1994). We used the modulus, $AE$, rather than stiffness, $AE/L$, because the reported stiffness values depend on the filament length, $L$, and cross-sectional area $A$ is not well defined for myosin and actin filaments. The only nonlinear element included in calculations is titin, whose nonlinear elastic characteristics were described previously (Mijailovich et al., 2019). In filaments, the deformation in contracting muscle is small: $<1\%$ (Huxley et al., 1994; Wakabayashi et al., 1994; Ma et al., 2018b). The elasticity of cross-bridges originates from multiple sources and can be nonlinear (Kaya and Higuchi, 2010) but, for simplicity, here we used a linear elastic model, as is usually done in sliding filament models.

Cross-bridge model parameters
To achieve the observed rate of twitch relaxation, we used a six-state cycle that includes a PS (Fig. 2, A and B). Following the approach of Duke (Duke, 1999; Mijailovich et al., 2016; Mijailovich et al., 2019), the state transition rate constants are defined by specifically designed experiments, estimated from indirect observations, or deduced from multiple sources. The simulations used a large number of the same parameters, except a few, that can strongly vary with the experimental conditions. For clarity, we show the complete set of cross-bridge model parameters, where variable ones are taken from simulations of trabecula twitch experiments at 27.5°C (see Table 1). When different parameter values are used for simulations of the experiments performed under different conditions, such as at what temperature, SL, and interfilament lattice spacings and whether the muscle is demembranated or intact, they are shown in the main text or in figure legends.

The cross-bridge cycle parameters for myosin binding to and detachment from actin (Eq. A2) are $k_{dA}^p = 226$ s$^{-1}$, $k_A^p = 46$ s$^{-1}$, and $f_{\text{rel}} = 1.004$; for the power stroke, the equilibrium constant (Eq. A3) is defined by $\Delta G_{\text{stroke}} = -13 k_BT$, the power stroke $d = 10.5$ nm; for ADP release, $k_{dO}^p = 60$ s$^{-1}$, and the second power stroke $\delta = 1.0$ nm. Because of the exponential forms in the expressions for the state transition rates, Eq. A3 and Eq. A5 can become very large and can generate numerical problems; they are capped to $k_{dA}^p = 1000$ s$^{-1}$, $k_{\text{cap}}^p = 100$ s$^{-1}$, and $k_{\text{cap}}^A = 10^4$ s$^{-1}$. When the cap value is reached, the reverse rates are changed to decay exponentially to satisfy the equilibrium constant, $K_b(x)$.

The remaining rates of the six-state cycle (Fig. 2) are defined as follows: ATP-binding and myosin detach rate, $k_{\text{bind}} = 10^6$ s$^{-1}$; hydrolysis forward rate $k_{sH} = 100$ s$^{-1}$ and backward rate $k_{sH} = 10$ s$^{-1}$; the transition rate to PS $k_{\text{PS}} = 200$ s$^{-1}$ and the calcium-dependent transition rate from PS (Eq. 1), where base line rate is $k_{sPS}^p = 5$ s$^{-1}$, the rate at high $[\text{Ca}^{2+}]$, $k_{sPS}^p = 400$ s$^{-1}$, Hill coefficient of the rate sigmoidal $[\text{Ca}^{2+}]$ rise $b = 5$ and calcium concentration, when $k_{\text{PS}} = \text{equal to } k_{sPS}^p/2$, is $[\text{Ca}^{2+}] = 1$ µM. In all simulations, cross-bridge stiffness is taken to be $k = 1.3$ pN/nm and the value for $k_BT = 4.147$ pN nm at 27.5°C or calculated at the temperature of the experiment.

CFC model parameters
For calcium-binding kinetics to TnC and interaction of TnI with actin (Fig. 2 C), the equilibrium rate constant of calcium binding to TnC along TnC closed states, $K_C = \bar{K}_{\text{Ca}} \cdot [\text{Ca}^{2+}]$, is effectively defined via forward constant $k_{\text{Ca}} = \bar{k}_{\text{Ca}} \cdot [\text{Ca}^{2+}]$, where
the number of thick filaments per unit of cross-sectional area, and calcium-independent dissociation constant $k_{Ca} = 75.4 \times 10^7 M^{-1} s^{-1}$ and calcium-independent dissociation constant $k_{Ca} = 75.4 \times 10^7 M^{-1} s^{-1}$ and $\text{TnT-actin interaction equilibrium rate constant } \lambda = 10$, and $\text{TnT attachment rate } k_T = \lambda = 375 s^{-1}$, and cooperativity coefficient $\sigma_c = 0.01$.

For the CFC model, we used the same parameters reported previously (Mijailovich et al., 2012): a Tpm pinning angle, $\phi = -28^\circ$; myosin-imposed Tpm angular displacement, $\phi = 10^\circ$; the persistence length of Tpm–Tn confined chain $1/\xi = (4k_{CFC}/\alpha_p)^{1/4} = 50$ nm; and angular SD $\sigma_\phi = 29.7^\circ$ (Pirani et al., 2005). The strength of the chain confining potential is weaker than used for skeletal muscle (Mijailovich et al., 2012), having a value for $\alpha_p$ of 0.149 pN.

**Calcium transients: measurements, conversions, and derivations when they are not available**

Used in twitch simulations are [Ca$^{2+}$]$^-$ transients reported by Janssen and de Tombe, 1997 using the calibration equation $[Ca^{2+}]$ 

$$[Ca^{2+}] = K_0 (R - R_{min}) / (R_{max} - R)$$

(Backx and Ter Keurs, 1993; Gao et al., 1994; Backx et al., 1995), where $K_0 = 2.95$, $R_{max} = 10.21$, and $R_{min} = 0.47$ (Fig. 5 A and Fig. S4), or deduced from these measurements when calcium transients are not reported (Figs. 3 and 5 B). In the latter case, we adapted the Janssen et al. (2002) Ca$^{2+}$ transient recorded at 27.5°C in the fixed-length trabecula to the Caremani et al. (2016) experiment at similar temperature (27.2°C) by reducing the peak concentration from 1.51 to 1.20 µM to match lower observed maximum twitch tension (lower by 27.3%) in the Caremani et al. (2016) experiment.

Because the calcium transients typically achieve a higher peak [Ca$^{2+}$] in fixed-length muscles than in isometric sarcomere twitches (Janssen and de Tombe, 1997), we rescaled the adapted calcium transient for the Caremani et al. fixed-length trabecula experiment (from the peak of 1.2 µM to the isometric half-sarcomere using the ratio of the peaks of [Ca$^{2+}$] recorded by Janssen and de Tombe at 22°C).

**Appendix E**

**Conversion between observed tension and isometric force per myosin filament**

Due to the stochastic process of myosin interacting with actin, the forces in the myofilaments fluctuate in time and each filament experiences somewhat different force. For comparison with observed isometric tensions, we include in all plots both the tension in kilopascals and the average force per myosin filament, $F$, in piconewtons. The scales are related by a factor that takes into account how many myosin filaments there are per unit of the fiber cross-sectional area (Linari et al., 1998; Mijailovich et al., 2016). Because the total number of thick filaments in a myofibril or muscle fiber does not change with experimental conditions, for the estimation of the number of thick filaments per unit of cross-sectional area, we used the lattice spacing, $d_{l,0}$, at slack length (i.e., at the same length where the muscle cross-sectional area used in the tension calculations is measured).

The lattice spacing in demembranated rat cardiac trabeculae cells at slack SL (1.9 µm) is $d_{l,0} = 43.5$ nm (Irving et al., 2000) and the number of thick filaments per µm² myofibril is ~458. Taking into account the fraction of the cross-sectional area in cardiac cells occupied by myofibrils as 68% (Cazorla et al., 2000), this reduces the number of myosin filaments over the muscle cell cross-section to ~311 per µm². In this case, the scaling factor provides that $T_c = 40$ kPa corresponds to $F_c = 128.5$ pN per myosin filament or 64.25 pN per actin filament.

In intact trabeculae, the lattice spacing is smaller and at 2.2 µm, where the cross-sectional area was measured, is $d_{l,0} = 33.83$ nm (Irving et al., 2000), and the number of thick filaments per µm² myofibril is ~757. Taking into account the fraction of the cross-sectional area in rat trabeculae occupied by myofibrils to be ~61%, as observed in previous studies (Schaper et al., 1985; Barth et al., 1992), reduces the number of myosin filaments over the trabecula cross-section to ~462 per µm². In this case, the scaling factor provides that $T_c = 40$ kPa corresponds to $F_c = 86.7$ pN per myosin filament or 43.35 pN per actin filament.

**Acknowledgments**

Henk L. Granzier served as editor.

We gratefully acknowledge the help of Prof. Thomas C. Irving with editing of the final version of the manuscript.

This project is supported by National Institutes of Health grants R01 HL128368 and RM1 GM131981 (to M. Regnier) and by the European Union’s Horizon 2020 research and innovation program under grant agreement 777204 (to S.M. Mijailovich, M. Regnier, M.A. Geeves, and C. Poggesi). This article reflects only the authors’ view. The European Commission is not responsible for any use that may be made of the information the article contains.

The authors declare no competing financial interests.

Author contributions: S.M. Mijailovich conceptualized the study, introduced new additions into MUSICO platform, including formulation of thin filaments calcium regulation, implementation of updated cross-bridge cycle, series elastic element in the model of trabecula contraction, new structural elements and changes in sarcomere geometry, designed numerical procedures, analyzed the data from simulations, wrote the first draft of the manuscript, and supervised each step of the work. M. Prodanovic assisted in designing numerical procedures and monitored implementation of changes computer code, performed all simulations and data analyses, contributed in graphical design, and assisted in the writing of the manuscript. C. Poggesi provided interpretation of experimental data and contributed in the comparative analysis of simulations and experimental data. M.A. Geeves contributed to development of the model of cross-bridge cycle and thin filament regulation, to formulating the key points in the Discussion, and in writing the final draft of the manuscript. M. Regnier contributed by providing experimental data, formulating the elucidating important points in the Discussion, and assisted in writing the final draft of the manuscript.
Chung, C.S., C.W. Hoopes, and K.S. Campbell. 2017. Myocardial relaxation is accelerated by fast stretch, not reduced afterload. J. Mol. Cell. Cardiol. 103:65–73. https://doi.org/10.1016/j.yjmcc.2017.01.004

Daniel, T.L., A.C. Trimble, and P.B. Chase. 1998. Compliant realignment of binding sites in muscle: transient behavior and mechanical tuning. Biophys. J. 74:1611–1621. https://doi.org/10.1016/S0006-3495(98)7875-5

Deacon, J.C., M.J. Bloemink, H. Rezavandi, M.A. Geeves, and L.A. Leinwand. 2012a. Erratum to: Identification of functional differences between recombinant human α and β cardiac myosin motors. Cell. Mol. Life Sci. 69:4239–4235. https://doi.org/10.1007/s00018-012-1111-5

Deacon, J.C., M.J. Bloemink, H. Rezavandi, M.A. Geeves, and L.A. Leinwand. 2012b. Identification of functional differences between recombinant human α and β cardiac myosin motors. Cell. Mol. Life Sci. 69:2261–2277. https://doi.org/10.1007/s00018-012-0927-3

Dobesh, D.P., J.P. Konhilas, and P.P. de Tombe. 2002. Cooperative activation in cardiac muscle: impact of sarcomere length. Am J Physiol Heart Circ Physiol. 282:H1055–1062.

Duke, T.A. 1999. Molecular model of muscle contraction. Proc. Natl. Acad. Sci. USA. 96:2770–2775. https://doi.org/10.1073/pnas.96.6.2770

Eyiring, H. 1935. The activated complex in chemical reactions. J. Chem. Phys. 3:107–115. https://doi.org/10.1063/1.1749604

Fenn, W.O. 1924. The relation between the work performed and the energy liberated in muscular contraction. J. Physiol. 58:373–395. https://doi.org/10.1113/jphysiol.1924.sp000241

Ferrantini, C., R. Coppini, L. Sacconi, B. Tosi, M.L. Zhang, G.L. Wang, E. de Vries, E. Hopenbrouwers, F. Pavone, E. Cerbai, et al. 2014. Impact of detubulation on force and kinetics of cardiac muscle contraction. J. Gen. Physiol. 143:783–797. https://doi.org/10.1085/jgp.201311125

Feynman, R.P., and A.R. Hibbs. 1965. Quantum Mechanics and Path Integrals. McGraw-Hill, New York.

Gao, W.D., P.H. Backx, M.A. Backx, and E. Marban. 1994. Myofilament Ca2+ sensitivity in intact versus skinned rat ventricular muscle. Circ. Res. 74:408–415. https://doi.org/10.1161/01.RES.74.3.408

Gao, W.D., N.G. Perez, and E. Marban. 1998. Calcium cycling and contractile activation in intact mouse cardiac muscle. J. Physiol. 507:175–184. https://doi.org/10.1113/1.14469759.1998.115877b

Geeves, M., H. Griffiths, S. Mijailovich, and D. Smith. 2011. Cooperative [Ca2+]–dependent regulation of the rate of myosin binding to actin: solution data and the tropomyosin chain model. Biophys. J. 100:2679–2687. https://doi.org/10.1016/j.bpj.2011.04.020

Glassstone, S., J. Laidler, and K. Eyiring. 1941. The Theory of Rate Processes. McGraw Hill, New York.

Gordon, A.M., E. Homsher, and M. Regnier. 2000. Regulation of contraction in striated muscle. Physiol. Rev. 80:853–924. https://doi.org/10.1152/physrev.2000.80.3.853

Higuchi, H., T. Yanagida, and Y.E. Goldman. 1995. Compliance of thin filaments in skinned fibers of rabbit skeletal muscle. Biophys. J. 69:1000–1010. https://doi.org/10.1016/S0006-3495(95)79957-1

Hill, T.L. 1974. Theoretical formalism for the sliding filament model of contraction of striated muscle. Part I. Prog. Biophys. Mol. Biol. 28:267–340. https://doi.org/10.1016/0032-0941(74)90020-0

Hooijman, P., M.A. Stewart, and R. Cooke. 2011. A new state of cardiac myosin with very slow ATP turnover: a potential cardioprotective mechanism in the heart. Biophys J. 100:1969–1976. https://doi.org/10.1016/j.bpj.2011.02.061

Hunt, A.J., F. Gittes, and J. Howard. 1994. The force exerted by a single kinesin molecule against a viscous load. Proc. Natl. Acad. Sci. USA. 91:6776–6781. https://doi.org/10.1073/pnas.91.15.6776

Huxley, H.E., A. Stewart, H. Sosa, and T. Irving. 1994. X-ray diffraction measurements of the extensibility of actin and myosin filaments in contracting muscle. Biophys. J. 67:2421–2421. https://doi.org/10.1016/0006-3495(94)70278-3

Irving, M. 2017. Regulation of contraction by the thick filaments in skeletal muscle. Biophys. J. 113:2579–2594. https://doi.org/10.1016/j.bpj.2017.09.037

Irving, T.C., and D.W. Maughan. 2000. In vivo x-ray diffraction of indirect flight muscle from Dro sophila melanogaster. Biophys. J. 78:2531–2558. https://doi.org/10.1016/S0006-3495(00)76796-8

Irving, T.C., J. Konhilas, D. Perry, R. Fischetti, and P.P. de Tombe. 2000. Myofilament lattice spacing as a function of sarcomere length in isoleted rat myocardium. Am. J. Physiol. Heart Circ. Physiol. 279:H2568–H2573. https://doi.org/10.1152/ajpheart.2000.279.5.H2568

Janssen, P.M., and P.P. de Tombe. 1997. Uncontrolled sarcomere shortening increases intracellular Ca2+ transient in rat cardiac trabeculae. Am. J. Physiol. 272:H1892–1897. https://doi.org/10.1152/ajpheart.1997.272.2.H1892

Janssen, P.M., L.B. Stull, and E. Marban. 2002. Myofilament properties comprise the rate-limiting step for cardiac relaxation at body temperature in the rat. J. Gen. Physiol. 119:383–405. https://doi.org/10.1085/jgp.201812604

Mijailovich et al. Computational model of cardiac twitch
McNamara, J.W., A. Li, N.J. Smith, S. Lal, R.M. Graham, K.B. Koolker, S.J. van Dijk, C.G.D. Remedios, S.P. Harris, and R. Cooke. 2016. Ablation of cardiac myosin binding proteins disrupts the super-relaxed state of myosin in murine cardiomyocytes. J. Mol. Cell. Cardiol. 94:55–71. https://doi.org/10.1016/j.yjmcc.2016.03.009

Mijailovich, S.M., D. Stamenoğlu, and J.J. Frederking. 1993. Toward a kinetic theory of connector tissue micromechanics. J Appl Physiol (1985) 74: 665–681. https://doi.org/10.1152/jappl.1993.74.2.665

Mijailovich, S.M., J.J. Frederking, and J.P. Butler. 1996. On the theory of muscle contraction: filament extensibility and the development of isometric force and stiffness. Biophys. J. 71:1475–1484.

Mijailovich, S.M., X. Li, J.C. del Alamo, R.H. Griffiths, V. Kecman, and M.A. Geeves. 2010. Resolution and uniqueness of estimated parameters of a model of thin filament regulation in solution. Comput. Biol. Chem. 34: 19–33. https://doi.org/10.1016/j.compbiolchem.2009.11.002

Mijailovich, S.M., O. Kayser-Herold, X. Li, H. Griffiths, and M.A. Geeves. 2012. Cooperative regulation of myosin-S1 binding to actin filaments by a continuous flexible Tn-Tnch chain. Eur. Biophys. J. 41:1015–1032. https://doi.org/10.1007/s00249-012-0859-8

Mijailovich, S.M., O. Kayser-Herold, B. Stojanovic, D. Nedic, T.C. Irving, and M.A. Geeves. 2016. Three-dimensional stochastic model of actin-myosin binding in the sarcomere lattice. J. Gen. Physiol. 148:459–488. https://doi.org/10.1085/jgp.201611608

Mijailovich, S.M., D. Nedic, M. Svicevic, B. Stojanovic, J. Walklate, Z. Ufalsa, and M.A. Geeves. 2017. Modeling the actin-myosin ATPase cross-bridge cycle for skeletal and cardiac muscle myosin isoforms. Biophys. J. 112: 984–996. https://doi.org/10.1016/j.bpj.2017.01.021

Mijailovich, S.M., M. Stojanovic, S.M. Mijailovich, M.A. Geeves, T.C. Irving, and H.L. Granzier. 2019. Nebulin and titin modulate cross-bridge cycling and length-dependent calcium sensitivity. J. Gen. Physiol. 151:680–704. https://doi.org/10.1085/jgp.201812165

Millman, B.M. 1998. The filament lattice of striated muscle. Physiol. Rev. 78: 359–391. https://doi.org/10.1152/physrev.1998.78.2.359

Nag, S., D.V. Trivedi, S.S. Sarkar, A.S. Adhikari, M.S. Sushita, S. Sutton, K.M. Ruppel, and J.A. Spudich. 2017. The myosin mesa and the basis of hypercontractility caused by hypertrophic cardiomyopathy mutations. Nat. Struct. Mol. Biol. 24:525–533. https://doi.org/10.1038/nsmb.3408

Niederer, S.A., P.J. Hunter, and N.P. Smith. 2006. A quantitative analysis of cardiac myocyte relaxation: a simulation study. Biophys. J. 90:1697–1722. https://doi.org/10.1529/biophysj.105.069534

Papoulis, A. 1991. Probability, Random Variables, and Stochastic Processes. McGraw-Hill, New York.

Parmley, W.W., and E.H. Sonnenblick. 1967. Series elasticity in heart muscle. Its relation to contractile element velocity and proposed muscle models. Circ. Res. 20:112–123. https://doi.org/10.1161/01.RES.20.1.112

Pirani, K., C. Xu, V. Hatch, R.L. Moss. 2005. Single particle analysis of relaxed and activated muscle thin filaments. J. Mol. Biol. 346:761–772. https://doi.org/10.1016/j.jmb.2004.12.013

Poole, K.J., M. Lorenz, G. Evans, G. Rosenbaum, A. Pirani, R. Craig, L.S. Tobacman, and J.A. Spudich. 2017. The myosin mesa and the basis of hysteresis in cardiac myosin binding in the sarcomere lattice. J. Mol. Biol. 384:60–72. https://doi.org/10.1016/j.jmb.2008.09.013

Ma, W., H. Gong, and T. Irving. 2018a. Myosin head configurations in resting and contracting murine skeletal muscle. Int. J. Mol. Sci. 19:2643. https://doi.org/10.3390/ijms19092643

Ma, W., H. Gong, B. Kiss, E.J. Lee, H. Granzier, and T. Irving. 2018b. Thick-filament extensibility in intact skeletal muscle. Biophys. J. 115: 1580–1588. https://doi.org/10.1016/j.bpj.2018.08.038

Martyn, D.A., B.B. Adhikari, M. Regnier, J. Gu, S. Xu, and L.C. Yu. 2004. Response of equatorial x-ray reflections and stiffness to altered sarcomere development in human cardiomyocytes. J. Mol. Cell. Cardiol. 39:168–183. https://doi.org/10.1016/j.yjmcc.2004.09.015

Mijailovich et al.

Computational model of cardiac twitch

Journal of General Physiology

23 of 24

https://doi.org/10.1085/jgp.202012604

Downloaded from http://rupress.org/jgp/article-pdf/153/3/e202012604/1409072/jgp_202012604.pdf by guest on 22 January 2022
Regazzoni, F., L. Dedè, and A. Quarteroni. 2020. Biophysically detailed computational models of a multicellular cardiac active mechanics. PLOS Comput. Biol. 16:e1008294. https://doi.org/10.1371/journal.pcbi.1008294

Rice, J.J., F. Wang, D.M. Bers, and P.P. de Tombe. 2008. Approximate model of cooperative activation and crossbridge cycling in cardiac muscle using ordinary differential equations. Biophys. J. 95:2368–2390. https://doi.org/10.1529/biophysj.107.119487

Robinson, T.F., and S. Winegrad. 1977. Variation of thin filament length in heart muscles. Nature. 267:74–75. https://doi.org/10.1038/267074a0

Robinson, T.F., and S. Winegrad. 1979. The measurement and dynamic implications of thin filament lengths in heart muscle. J. Physiol. 286:607–619. https://doi.org/10.1113/jphysiol.1979.sp012640

Schaper, J., E. Meiser, and G. Stämmier. 1985. Ultrastructural morphometric analysis of myocardium from dogs, rats, hamsters, mice, and from human hearts. Circ. Res. 56:377–391. https://doi.org/10.1161/01.RES.56.3.377

Smith, D. 2001. Path integral theory of an axially-confined worm-like chain. J. Phys. Math. Gen. 34:4507–4523. https://doi.org/10.1088/0305-4470/34/21/307

Smith, D.A., and M.A. Geeves. 2003. Cooperative regulation of myosin-actin interactions by a continuous flexible chain I: actin-tropomyosin-troponin and regulation by calcium. Biophys. J. 84:3168–3180. https://doi.org/10.1016/S0006-3495(03)70041-1

Smith, D.A., and S.M. Mijailovich. 2008. Toward a unified theory of muscle contraction. II: predictions with the mean-field approximation. Ann. Biomed. Eng. 36:1359–1371. https://doi.org/10.1007/s00439-008-9514-z

Smith, D.A., R. Mayrum, and M.A. Geeves. 2003. Cooperative regulation of myosin-actin interactions by a continuous flexible chain I: actin-tropomyosin systems. Biophys. J. 84:3155–3167. https://doi.org/10.1016/S0006-3495(03)70040-X

Smith, D.A., M.A. Geeves, J. Sleep, and S.M. Mijailovich. 2008. Towards a unified theory of muscle contraction. I: foundations. Ann. Biomed. Eng. 36:1624–1640. https://doi.org/10.1007/s00439-008-9536-6

Spudich, J.A. 2015. The myosin mesa and a possible unifying hypothesis for the molecular basis of human hypertrophic cardiomyopathy. Biochem. Soc. Trans. 43:64–72. https://doi.org/10.1042/BST20140324

Squire, J.M. 1992. Muscle filament lattices and stretch-activation: the matching-mismatch model reassessed. J. Muscle Res. Cell Motil. 13:183–189. https://doi.org/10.1007/BF01874155

Tanner, B.C., Y. Yue, D.T. Marban, and W.G. Wier. 1986. Relationship between force and intracellular [Ca2+] in tetanized mammalian heart muscle. J Gen Physiol. 87:223–242.

Zoghbi, M.E., J.L. Woodhead, R.L. Moss, and R. Craig. 2008. Three-dimensional structure of vertebrate cardiac muscle myosin filaments. Proc. Natl. Acad. Sci. USA. 105:2386–2390. https://doi.org/10.1073/pnas.0708912105
Supplemental material

All supplemental figures are available online in a ZIP package.

Methods

The relative spatial positions of myosin heads and binding sites on the thin filament in the 3-D sarcomere lattice
The 3-D sarcomere is composed of thick and thin filaments arranged in an overlapping hexagonal lattice where myosin filaments extend from the central M-band toward Z-lines and actin filaments extending from Z-lines forming a hexagonal arrangement in cross-section. The actin monomers are helically arranged in a double-stranded helical structure, and orientationally favorable myosin-binding sites on actin filament are associated with a few myosin heads. Each myosin molecule is attached to the backbone of myosin filament via the S2 rod and has two heads at the free end, but only one head per dimer is shown (Fig. S1). The pairs of myosin heads form a triple-helix along the myosin filament. The myosin heads are arranged in layers, and each layer forms a “crown” with three pairs of heads. The crowns are axially separated by 14.3 nm and rotated by 40°, forming different angular arrangements with the actin filaments, but only those that might interact with the actin filament are shown. In the axial direction, each pair of heads and multiple binding sites (target zones) on surrounding actin filaments form a large number of arrangements defined by the relative axial distances, \( x_{\text{ml}} \), between the unstrained position of the myosin head or cross-bridge \( m \) and the nearest actin-binding site \( l \). In the radial direction, in the azimuthal plane, up to three myosins can attach to each actin filament, forming azimuthal angles \( \alpha \) and \( \beta \) (Fig. S1 B). The hexagonal sarcomere lattice with a 2:1 actin-to-myosin filament ratio shows the spatial arrangement of crowns where the three (pairs of) heads in crown 1 interact with one actin filament displayed by azimuthal angles \( \alpha = 0 \) and \( \beta \) (Fig. S1 C), and the heads in crowns 2 and 3 are rotated for 40° and two times 40°, respectively, having different azimuthal spatial arrangements relative to binding sites on the actin filaments displayed by azimuthal angles \( \alpha \neq 0 \) and \( \beta \) (Fig. S1 D). The angles \( \alpha \) and \( \beta \) define weight factors \( C_{\alpha} \) and \( C_{\beta} \) of myosin binding in the 3-D sarcomere lattice (Mijailovich et al., 2016).

Can accelerated relaxation be achieved by a five-state model?
We have explored the possibility of speeding up the tension relaxation rate with a five-state model by varying the rate constants \( k_{oA}^{+} \) and \( k_{oA}^{-} \). The superscript \( o \) denotes rates when the elastic element is relaxed and is omitted in the relevant figures for simplicity of notation. In Fig. S2, the predicted tensions with the five-state model are shown, where the parameters used in the simulations are taken from the isometric half sarcomere simulations with the five-state model displayed in Fig. 3. The predicted tension shows significantly slower relaxation (dark green line) as compared to the Caremani et al. (2016) experiment (dark gray line). This slow relaxation rate is caused by an increasing number of cross-bridges in the A.M.D.Pi state during lengthening that cannot be detached through ADP release and the ATP-binding part of cross-bridge cycle, but only through a large increase in the detachment rate \( k_{oA}^{-} \). An increase of \( k_{oA}^{-} \) by 17-fold significantly increased the tension relaxation rate (green line), but it still is not sufficient to reach the observations (black line). This large increase in \( k_{oA}^{-} \) is far above physiological range and is significantly larger than rate \( k_{oA}^{+} \).

Correction for loss of trabecula length control during sarcomere-lengthening phase of twitch
Caremani et al. (2016) attempted to keep the SL fixed by controlling the length of trabeculae (gray line in Fig. S3 A). However, this was only partially achieved by increasing the length of trabeculae during the rise of twitch tension, while, when the tension reached value of the twitch peak, length control failed to control SL (Fig. S3 A) and sarcomeres started to lengthen (gray line in Fig. S3 C). Consequently, the observed tension was greater during relaxation, and relaxation was significant slower (gray line in Fig. S3 B) than if the SL were kept constant.

Because the observed half-sarcomere length increases, mainly during relaxation of the twitch, the imposed lengthening increased tension in the SE component and extension of titin, which would not be present if a half-sarcomere length were kept fixed. After the correction, the resulting tension was lower than observed (pink line versus gray in Fig. S3 B). For reference, the reduction in tension due to the SE component only (red line in Fig. S3 B) is also shown. These corrections also preserve the trabecula length change (i.e., motor position in pink line; Fig. S3 A), where the motor reaches the position held at the beginning of the experiment when muscle was fully relaxed.

The twitch simulations with the five- and six-state cross-bridge cycles in the isometric half-sarcomere
For comparison between the five- and six-state cross-bridge cycles, we also attempted to fit the twitch transient of Janssen and de Tombe (1997) in fixed-length half-sarcomeres at 22°C (see Fig. S4). The simulations with the five-state model matched the peak tension but showed slightly slower tension rise and a poor overall fit to the relaxation phase (dark green line in Fig. 3), with the fall in tension being too slow.

Mijailovich et al.
Computational model of cardiac twitch

Journal of General Physiology
https://doi.org/10.1085/jgp.202012604
Note that the five-state model also predicted a high level of resting tension, which we corrected by lowering baseline calcium (pink dashed line in Fig. S4) to generate this fit. The simulations using the six-state model (red line in Fig. S4) substantially increased the rate of tension relaxation, but fit of the Janssen and de Tombe data was poor during the relaxation. For comparison of the net active tension change, we subtracted the resting tension from all traces. The parameters used in the simulations with the five-state model are listed in Table 1, except $k_{o_d}$, $k_{o_A}$, and $k_{o_D}$, having values of 29, 10, and 160 s$^{-1}$, respectively, and for the six-state model in Table 2. Interestingly, the increase in temperature from 22°C to 27.2°C increases the rate $k_{o_A}$ by ~70% in the simulations with the six-state cycle that is almost the same increase in $k_{o_A}$ obtained by the five-state cycle (i.e., from 29 s$^{-1}$ at 22°C to 50 s$^{-1}$ at 27.2°C).

**Measurements of passive SE compliance during a twitch in isometric trabeculae**

Large changes in SL during twitch contractions in fixed-length cardiac muscle indicate significant compliance in the intact muscle preparations. Simultaneous measurements of tension and changes in a half-sarcomere length can define the compliance of intact trabecula or other cardiac muscle. We compare tension length loops observed by Janssen and de Tombe (1997) and Caremani et al. (2016) in order to extract characteristics of the nonlinear series compliance. Comparison of the observed loops is shown in Fig. S5. Because of the uncertainties in the descending part of Janssen and de Tombe’s loop (red symbols and line), we have derived average tensions over the range of the Caremani et al. loop (gray line) and used these values, for simplicity, in all MUSICO simulations.

**ATPase consumption during twitch contractions in fixed-length trabeculae**

During contraction of fixed-length trabeculae, the sarcomeres shorten during the tension rise and lengthen during the tension fall. These large changes in length are associated with a large increase in the rate of ATP consumption, up to 20-fold compared with that of a half-sarcomere under truly isometric conditions (Fig. S6). The ATPase in isometric half-sarcomeres, with rigid filaments at peak twitch tension, is ~1 s$^{-1}$, whereas with extensible filaments, it is ~1.15 s$^{-1}$, where the differences can be explained by the unloading of cross-bridges due to local sliding between the filaments, leading to detachment of a significant fraction of previously attached cross-bridges and their reattachment (Mijailovich et al., 1996). In fixed-length trabeculae, realignment of myocytes and other series elasticities involves SL changes, and this results in slower force development, lower tension peak (blue solid line in Fig. S6), and also detachment and reattachment of cross-bridges, thereby resulting in a large increase in ATPase (blue symbols). The ATPase peak correlates with maximum shortening velocity in the early phase twitch tension development and then monotonically decays as shortening velocity decreases. Similar behavior is observed in simulations of force development in fixed-end trabeculae. In both cases, the ATPase normalized to shortening velocity per SL shows stable values, increasing at slower shortening velocities (Fig. S7).

**Sensitivity analysis of twitch tension to model parameters in fixed-length trabeculae**

The estimated free model parameters for twitchs in fixed-length trabeculae showed (in several cases) a large range of values depending on experimental conditions and model constraints. In most cases, the chosen parameter was varied by ±20%, though in a few cases the variations were larger to emphasize the significance of the parameter change. Parameters used for twitch simulations of the Caremani et al. (2016) data by the six-state cross-bridge cycle were varied, and the predicted tension responses are compared with the experimental responses shown in Fig. 5 B. The changes in the key cross-bridge parameters include rates of myosin binding to actin, $k_{o_d}$ (Fig. S8), $k_{o_A}$ (Fig. S9), and ADP release, $k_{o_D}$ (Fig. S10).

The variation in parameters defining the transition rate from the PS to the M.D.Pi state (defined by Eq. 1) includes the Hill coefficient of the sigmoidal rate of rise, $b$, [Ca$^{2+}$] at the half $k_{PS}$ rise, [Ca$^{2+}$]$_{SO}$, and the baseline rate from the PS, $k_{o_P}$ (Figs. S11, S12, and S13).

**Estimate of active force during twitch contractions in isometric trabeculae**

To estimate the effect of changes in half-sarcomere length (up to 80 nm) on tension during twitch contractions in fixed-length trabeculae (Caremani et al., 2016), simulations showed the effect of an SE component on tension due to titin (Fig. S14). Typically, the active tension is estimated from the observed tension transient when the resting tension is subtracted. However, resting tension at SL <2.2  ㎛ originates primarily from titin elasticity, and the tension in titin relaxes as SL decreases (green line). The force per myosin filament at a SL of 2.2  ㎛ at 22.5°C (Janssen et al., 2002) has a passive tension contribution from titin of ~16.5 pN per myosin filament or muscle tension of ~7.5 kPa. When this force is subtracted from the predicted twitch force (red line), the observed active tension matches well (black line) with the data of Janssen et al. (2002). However, due to the large degree of internal shortening of sarcomeres in isometrically held cardiac muscle (displacements; red line), the titin force falls to ~4 pN, so active force is higher by about the same amount as decrease in titin force (pink line). Under these conditions, maximum active tension (pink line) is ~9% higher than reported (black line).

**Sensitivity analysis of twitch tension to model parameters in the absence of a PS**

We show in Fig. 3 that the five-state model, without a PS, displays high resting tension and much slower tension relaxation than is observed. In order to separate the effects of altering cross-bridge cycle parameters with or without a PS from the effects of SL changes, we performed simulations on an isometric half-sarcomere following the experiments of Caremani et al. (2016) and Janssen...
and de Tombe (1997). Because resting tension was high at the observed baseline \([\text{Ca}^{2+}]\) of 0.143 \(\mu\text{M}\), we reduced \([\text{Ca}^{2+}]\) to 0.05 \(\mu\text{M}\) in the simulations using the five-state cross-bridge model. This reduction of baseline \([\text{Ca}^{2+}]\) decreased resting tension from 57 kPa to a more reasonable value of 20 kPa (Fig. S15).

The parameters \(K_{\text{Ca}} = R_{\text{Ca}} \cdot [\text{Ca}]\) and \(\lambda\) (Fig. 2 C) define the kinetics of opening regulatory units and enabling tension generation. At low calcium levels, \(K_{\text{Ca}}\) is attenuated, but the regulatory units can still be open through pathways independent of \([\text{Ca}^{2+}]\) (Fig. 2 C). The transition rate between the blocked state A.TnI + TnC and the open state A + TnC.TnI (that permits myosin binding) is defined by the equilibrium rate \(e_o \cdot \lambda\), where \(e_o\) is established from experiments in solution having values from 0.005 to 0.033 (Geeves et al., 2011). Consequently, a significant resting tension can be developed at observed \([\text{Ca}^{2+}]\) levels, as shown in Fig. S15, for \(e_o = 0.01\). Decreasing \(e_o\) by three orders of magnitude (i.e., to \(10^{-9}\)) significantly reduces opening of regulatory units and decreased the resting tension by \(\sim 10\%\) at the observed calcium levels (0.143 \(\mu\text{M}\)) and by \(\sim 25\%\) at reduced baseline calcium level (0.05 \(\mu\text{M}\)). These data show that reducing \(e_o\) to much lower values than observed cannot sufficiently reduce resting tension (Fig. S15) to match the observations; thus, other mechanisms should be explored and implemented.

As described in the main text, the key feature for keeping resting tension low at observed baseline \([\text{Ca}^{2+}]\) and allowing fast relaxation of tension can be achieved by maintaining a low population of M.D.Pi at low levels of \([\text{Ca}^{2+}]\), such as by including the PS as introduced in this paper. However, there are other models that can match the observations, but they are based on less-detailed cross-bridge cycles and regulatory schemes, simplified sarcomere geometry, and a less-constrained scheme for the estimation of the model parameters. Inspired by the accomplishments with these models, we explored the possibility of achieving better fits with the five-state model with systematic parameter estimation.

The five-state model simulations showed relatively high resting tension (Prodanovic et al., 2020) and slow relaxation (Figs. 3 and S2). The main reason for these responses is the relatively high population of myosin in heads in the M.D.Pi state at low concentrations of \([\text{Ca}^{2+}]\), resulting in a high binding flux, and also fast rebinding of detached cross-bridges, maintaining tension for a prolonged period of time after reaching the peak tension. From the cross-bridge cycle scheme in Fig. 2 A, we explored possibilities to keep the population of M.D.Pi state low at low \([\text{Ca}^{2+}]\) levels and therefore populations of force-generating states without involving a PS. We explored the sensitivity of the twitch tension on critical cross-bridge rates \(k_{A}^{+}\), \(k_{D}^{-}\), \(k_{H}^{-}\), and \(k_{A}^{-}\). These rates can potentially maintain a population of cross-bridge states at the levels that can produce observed tension during relaxation phase at low \([\text{Ca}^{2+}]\). Figs. S16, S17, S18, and S19 show tension transients including changes of the peak and resting tensions in response to up to \(\pm 50\%\) change in each parameter value from the corresponding value used in the simulations shown in Fig. 3 (dark green line), denoted in these figures as original (i.e., orig.) values. In all simulations, the resting tension is subtracted to match the observations, and a reduced level of baseline \([\text{Ca}^{2+}]\) = 0.05 \(\mu\text{M}\) is used.

The increase in binding rate \(k_{D}^{-}\) increases the rate of tension rise, peak tension, and resting tension, but the rate of relaxation at all \(k_{D}^{-}\) is at least twofold slower than observed (Fig. S16). The relaxation rate is estimated from the time required for tension to decrease from the peak tension to 50% of the value, denoted as TP50. With decreasing of rate \(k_{D}^{-}\), TP50 also decreases, but even at \(k_{D}^{-}\) reduced to 50% of the reference value, TP50 is at least two times slower than the observed TP50 along with much lower tension peak (Fig. S16).

The increase in the rate of ADP release, \(k_{D}^{+}\), reduces the population of force-producing cross-bridge states and therefore reduces the overall tension response. Consequently, an increase in \(k_{D}^{+}\) decreases the rate of tension rise, peak tension, and resting tension, but the rate of relaxation is about the same for all \(k_{D}^{+}\) values and is about twofold slower than that observed (Fig. S17).

A decrease in the rate of ATP hydrolysis, \(k_{H}\), can possibly reduce the population of M.D.Pi state and therefore reduce the rebinding flux of available and recently detached cross-bridges. However, decreasing \(k_{H}\) from 150% to 50% of the original value used in Fig. 3 only modestly decreases peak tension and resting tension, whereas the rate of relaxation is about the same in all simulations (Fig. S18).

Another possible way to attenuate the binding flux due to the high population of myosin heads in the M.D.Pi state is to increase the rate of detachment from A.M.D.Pi state, \(k_{A}^{-}\) (i.e., to reduce the affinity of myosin to actin). Increasing \(k_{A}^{-}\) shows a significant decrease in both the peak and resting tension and in the rate of tension rise (Fig. S19). There are two distinctive regions in these changes: (1) the peak tensions modestly decrease for \(k_{A}^{-}\) increases up to twofold, and resting tension sharply decreases for \(k_{A}^{-}\) increases up to fourfold; and (2) for larger increases in \(k_{A}^{-}\), the peak tension falls rapidly, and resting tension reaches very low levels. Similarly, the rate of tension relaxation increases for \(k_{A}^{-}\) increases up to fourfold and then decreases. This limit in relaxation rate increase could be overcome by changes in other cross-bridge cycle rates, so the predicted rate could possibly match the observed relaxation rate. One promising combination of parameters showed a significant increase in rate of tension relaxation (Fig. S2), but it still fell short of reaching the observed rate.

Following this analysis, we further explored how changes in the combination of parameters can increase the tension relaxation rate. Starting with parameters for the best fit with the six-state model (Fig. 3 and Table 2), after excluding the PS, the simulation shows, as expected, much higher tension than observed and a slow tension relaxation rate (black line in Fig. S20). Increase of \(k_{D}^{-}\) by 2.5-fold matched the peak tension and rate of tension rise, but the relaxation was too slow (dark green line in Figs. S2 and S20). We accelerated the relaxation rate by \(\sim 2.4\)-fold by increasing of \(k_{A}^{-}\) 17-fold (green line in Figs. S2 and S20), but it still did not match the observations. Further increasing \(k_{A}^{-}\) showed a decrease in the peak tension and a slight decrease in the rate of the tension relaxation.
We further tested the possibility of matching both the peak tension and the rate of tension relaxation by decreasing $k_{oD}^{-A}$ by 20\% (i.e., to 120 s$^{-1}$), which allowed a further increase in $k_{oD}^{-A}$ to 420 s$^{-1}$ in order to reach the observed tension peak, but the rate tension relaxation changed only marginally compared with the best match to the observations (shown as the green line in Fig. S20). It is interesting that a further decrease in $k_{oD}^{-A}$ not only decreased the peak tension but also modestly decreased the rate of tension relaxation (Fig. S21).

In order to reach the observations, we also changed forward power stroke rate. Shifting the forward rate $k_{23}(x)$ for $\Delta x_o \sim 0.5$ nm via increase of $k_{23}^{\text{cap}}$ (Eq. A4) increased the relaxation rate in tension, enabling matching the rate of relaxation with the observations (Fig. S22). In these simulations, we kept the power stroke equilibrium rate, defined by $\Delta G_{\text{stroke}} = 13 k_B T$, and cross-bridge stiffness, $\kappa = 1.3$ pN/nm, and we kept the reverse power stroke rate the same (inset in Fig. 22). Changing $k_{23}(x)$ increased the number of post-power stroke (i.e., A.M.D) cross-bridges during tension rise and increased the rate of detachment for these cross-bridges during relaxation.

A good match of the experimental data using the five-state model would suggest that PS may not be necessary. However, in these simulations, we used a calcium transient with reduced baseline [Ca$^{2+}$] levels (pink dashed line in Figs. 3, S16–S23) compared with the observed (dark pink solid line in Fig. S23). The simulations with the observed calcium transient and with the same parameters as in Fig. S22 showed high resting tension ($\sim 18.5$ kPa), a slightly higher tension peak ($\sim 6.5\%$), and $\sim 25\%$ slower relaxation rate (Fig. S23). Further changes in $k_{oD}^{-A}$, $k_{oD}$, and $k_{23}(x)$ were not able to bring the model predictions to match the observations.

The effect of baseline Ca$^{2+}$ levels can be seen in tension–pCa relationships. Fig. S24 compares predictions of tension–pCa relationships using the cross-bridge cycle with the PS (the six-state model) and without a PS (the five-state model). The five-state model simulations show increased tension at low calcium levels compared with the observations and the predictions with the six-state model. Importantly, the tension at the observed baseline [Ca$^{2+}$] = 0.143 $\mu$M is at about the same value as the resting tension in Fig. S23.

Taken together, the cross-bridge cycle without a PS (the five-state model) cannot simultaneously match the observed resting tension, peak tension, and tension relaxation rate during twitch contraction with the observed [Ca$^{2+}$] transients. The reason for this inability of the five-state model is illustrated in Fig. S24, where the tension–pCa relationship shows a high level of tension at low levels of [Ca$^{2+}$]. This is strongly supported by the similar magnitude of tension at the observed baseline Ca$^{2+}$ level in twitch contractions (Fig. S23) and in the tension–pCa relationship at the same [Ca$^{2+}$] level (Fig. S24). This analysis suggests that some other mechanisms are necessary to resolve the problem of matching the model predictions to the observations, such as by introducing a PS, as proposed in this study.
Interaction between myosin heads and actin filaments in 3-D is defined by the triple-helical arrangement of myosin molecules along the myosin filament and the double-helical arrangement of monomers (myosin-binding sites) along the actin filaments. The 3-D geometry of myosin head binding domains and binding sites on actin in a sarcomere requires both longitudinal position matching and angular matching in the azimuthal plane. (A) Each myosin head $m$ can move from its undeformed position $X_m^0$ along actin filament due to thermal agitation and can reach a few neighboring binding sites on actin. The myosin-binding domain is shown as a yellow oval at the tip of the myosin head, and binding sites on actin are shown as bright blue circles. The range of axial movement is shown as a pale red bar. The relative axial position of a myosin head (cross-bridge) $X_m^0(t)$ and adjacent actin sites $X_i^{a,l}(t)$, where superscripts $n + l$ and $l$ denote the index of an adjacent site on actin, and $l = 0, 1, 2, … (L_a - 1)$ is the index of accessible sites on actin in neighborhood of $m$, respectively. The maximum number of adjacent sites on actin reachable by a myosin head $m$ is denoted as $L_a$. To bind the site $+l$, the cross-bridge, including S2 and a myosin head, needs to stretch or compress axially for displacement $x_{lm}$. (B) In the 3-D sarcomere lattice, the actin and myosin filaments are separated by spacing $d_{A-M}$, and sites on the actin filament (strand) are at an azimuthal angle $\beta$. In addition, a cross-bridge needs to turn from its equilibrium position by an angle $\alpha$ to reach an actin filament that is not aligned with its equilibrium angular position. For precise calculations of the angles, it is necessary to know the myosin equilibrium angular positions $\theta_m$, angular position of site on actin filament $\theta_a$, and diameters of myosin and actin filaments $2r_m$ and $2r_a$, respectively. The angular range of movement is denoted as a pale red arc around the actin filament. The azimuthal weight factors $C_\beta$ and $C_\alpha$ of myosin binding in 3-D sarcomere lattice are defined as in C and D. (C) When myosin heads in crown 1 are directly aligned with three actin filaments, then $C_\alpha = 1$ and $C_\beta$ weights the azimuthal departure of a myosin-binding site on the actin filament from the plane passing through myosin and actin longitudinal axes, where the angle $\beta$ is a function of the axial departure from perfect matching, $\xi$, which resembles the preference for myosin heads to bind to favorably oriented sites on the actin filament. (D) When myosin heads are not directly aligned with the surrounding actin filaments, such as with crowns 2 and 3, the weight factor $C_\alpha$ takes into account the departure by the angle $\alpha$ from perfect alignment between the heads on the crown and the reachable actin filaments. Figure S1 is adapted from Mijailovich et al. (2016).
Figure S2. **Effect of the five-state cross-bridge model rate $k_{oA}$ on relaxation of twitch tension in isometric half-sarcomeres (IoSarc).** The simulation, with parameters obtained from fits of isometric half-sarcomere tension at 27.2°C (Fig. 3), shows slow relaxation (dark green line). An additional increase of $k_{oA}$ by $\sim$17-fold significantly increases the rate of relaxation (green line and arrows), but it is still insufficient to reach the observation (dark gray line). The superscript o is omitted from the figure display for clarity.
Figure S3. SL controlled isometric tension of a half-sarcomere. (A) Observed change in length of trabeculae per half-sarcomere (gray line) and corrected length of trabeculae to truly keep a half-sarcomere length isometric (pink line), which shows a decrease in trabecula length (i.e., motor position) toward the initial length of relaxed muscle. (B) Correction of the tension transient during the twitch due the change in SL (gray line in C). The observed tension (gray line) and the tension after correcting for the PE passive tension due to change of half-sarcomere length (red line) and also after additional correction for titin (passive) forces (pink line). (C) Change in half-sarcomere length (gray line) shows the effect of losing control of HS length that apparently should be constant (pink line). The tension of the trabecula length was fixed (black line in B) is shown for reference. The direction and color of the arrows signify changes in length and tension.
Figure S4. **Significance of the PS in twitch relaxation phase.** The simulations of a half-sarcomere isometric twitch with the five-state model showed slow relaxation (dark green line) compared with adjusted observations of Janssen and de Tombe, 1997 (gray line), even when the baseline [Ca$^{2+}$] was reduced (pink dashed line). The simulations also showed high muscle resting tension at observed baseline [Ca$^{2+}$] (not shown), similar to that shown by Prodanovic et al. (2020) at comparable low [Ca$^{2+}$]. In contrast, the simulations with the six-state model, which includes the PS, significantly increased the speed of relaxation (indicated by arrow), but the model predictions state (red line) followed the poorly observed unusual flat top to the tension transient reported by Janssen and de Tombe, 1997. However, the model predictions approximately matched the simulation of tension reported by Niederer et al. (2006) (dark gray dotted line), suggesting possible inconsistency of the observation with other similar data. In the simulations, we used the same model parameters as shown in Table 1, except $k_1^+$, $k_2^+$, and $k_3^+$, which are presented in the text here for the five-state model and in Table 2 for the six-state cross-bridge cycle. For comparison, mean force per myosin filament and corresponding (observed) muscle tension are shown on the vertical axes. For consistency, resting tension is subtracted in all plots; thus, the responses to calcium transient display net change in active tension.
Figure S5. Comparative tension–displacement loops obtained from Janssen and de Tombe and Caremani et al. (red and orange symbols and lines, respectively). Related to Janssen and de Tombe, 1997; Caremani et al., 2016. Both loops indicated viscoelastic behavior showing higher tension during shortening and lower tension during relaxation accompanied with lengthening. In both cases, the peak force per myofilament and maximum shortening were about the same, but Janssen and de Tombe’s loop is much wider and more asymmetric about the mean value than that of Caremani et al. (see also Fig. 4 B). The Janssen and de Tombe loop (red line) also shows a large drop in tension after reaching maximum shortening and, after the drop in tension, relaxes slowly during lengthening, having a higher value of tension than the initial value at displacement <4 nm. This deviation of the descending part of the loop shows some uncertainty in the measurements compared with that of Caremani et al. Thus, for simplicity, we derived a nonlinear serial springlike elasticity from Caremani et al.’s average tension–displacement relationship (thick gray line) that shows more consistent behavior at displacement <40 nm and a nicely rounded loop at the tip of shortening. Because the conversion factor between force per myofilament and tension depends on the length at which the trabecula cross-section was measured, the corresponding tensions are shown in red letters for Janssen and de Tombe’s loop and in orange letters for Caremani et al.’s loop.
During twitch in an isometric half-sarcomere, the ATPase rate increases up to 15% due to relative movement between extensible filaments compared with the ATPase with rigid filaments (Mijailovich et al., 1996). Therefore, ATPase shows only a minor increase if the extensibility of thick and thin filaments is taken into account. In contrast, the ATPase in fixed-length trabeculae shows more than an order of magnitude change during tension rise, having a peak value $\sim 21 \text{s}^{-1}$. This large increase is associated with a large degree of sarcomere shortening (80 nm) that requires multiple detachments and reattachments of the cross-bridges. The ATPase is calculated as the rate of cross-bridge detachment over the number of cross-bridges participating in an active cross-bridge cycle; that is, the cross-bridge in PS is excluded.

Figure S6. Comparison of ATP consumption rate, ATPase, during twitch contractions of isometric half-sarcomere and fixed-length trabeculae. During twitch in an isometric half-sarcomere, the ATPase rate increases up to 15% due to relative movement between extensible filaments compared with the ATPase with rigid filaments (Mijailovich et al., 1996). Therefore, ATPase shows only a minor increase if the extensibility of thick and thin filaments is taken into account. In contrast, the ATPase in fixed-length trabeculae shows more than an order of magnitude change during tension rise, having a peak value $\sim 21 \text{s}^{-1}$. This large increase is associated with a large degree of sarcomere shortening (80 nm) that requires multiple detachments and reattachments of the cross-bridges. The ATPase is calculated as the rate of cross-bridge detachment over the number of cross-bridges participating in an active cross-bridge cycle; that is, the cross-bridge in PS is excluded.
Figure S7. ATPase is strongly correlated to sarcomere-shortening velocities in fixed-length trabeculae during force development (in SL per second). The nondimensional ratio of the ATPase and sarcomere-shortening velocity slowly increases from ∼25 at 1 SL/s to ∼50 at 0.25 SL/s. The progressive increase of the normalized ATPase suggests that there is a steady-state component of ATPase that becomes more evident at low shortening velocities <0.25 SL/s. Interestingly, about the same behavior is observed for the simulations of twitches and force development in fixed-length trabeculae. The shortening velocities rise sharply up to 3 SL/s and quickly drop to ∼1 SL/s and further decrease as the rate of force rise decreases.
Figure S8. Changes in twitch tension from Fig. 5B (black line) by ±20% variation in binding rate, $k_{oA}$. Increase in $k_{oA}$ increases the peak of the twitch tension and shows delay in relaxation. The superscript $o$ is omitted from the figure display for clarity.
Figure S9. Changes in twitch tension from Fig. S B (black line) by ±20% variation in detachment rate, $k_{-A}$. Increase in $k_{-A}$ increases the rate in tension relaxation. The superscript o is omitted from the figure display for clarity.
Figure S10. Changes in twitch tension from Fig. 5 B (black line) by ±20% variation in ADP release rate, $k_{o}^{+\text{D}}$. Increase in $k_{o}^{+\text{D}}$ only slightly increases the peak of the twitch tension and the relaxation starts at earlier times. The superscript o is omitted from the figure display for clarity.
Figure S11. Changes in twitch tension from Fig. 5 B (black line) by ±40% variation in Hill coefficient of the rate sigmoidal rise, $b$. Increase in $b$ decreases the peak of the twitch tension, and the relaxation starts earlier. These changes are much smaller for values of $b > 5$. 
Figure S12. Changes in twitch tension from Fig. 5 B (black line) by ±20% variation in \([\text{Ca}^{2+}]\) at the half of \(k_{\text{PS}}\) rise, \([\text{Ca}^{2+}]_{50}\). Increase in \([\text{Ca}^{2+}]_{50}\) decreases the peak of the twitch tension, and the relaxation starts earlier.
Figure S13. Changes in twitch tension from Fig. 5B in the main text (black line) by ±40% variation in baseline rate from the PS, $k_{PS}^0$. Increases in $k_{PS}^0$ increase the resting muscle tone and the peak of the twitch tension, and the relaxation is slower and slightly delayed.
Figure S14. **Active tension corrected for the decrease in titin force due to shortening of a half-sarcomere.** The observed tension is usually corrected for resting tension associated with the force in titin and other parallel connective tissue. The observed tension (black line) after the subtraction of the resting tension is usually reported as active tension. However, the resting tension in titin and other passive components decreases the tension due to shortening of the half-sarcomere denoted as displacements. The decrease in the (titin) force (green line) during twitch effectively increases active force (red to pink line) by the same amount. Force per myosin filament from MUSICO simulations is shown in parallel with the observed tension.
Figure S15. Effect of allosteric TnC–Tnl–actin–Ca$^{2+}$ interaction parameter, $\varepsilon_o$, on resting tension at observed baseline calcium level (0.143 µM) and when [Ca$^{2+}$] is reduced to 0.05 µM. MUSICO simulations using the five-state cross-bridge cycle predicted a slight decrease in resting tension at both baseline [Ca$^{2+}$] for decrease by three orders of magnitude in $\varepsilon_o$, but significant resting tension remains even at extremely low $\varepsilon_o$ values. Parameters used in simulations are the same as used in the five-state model simulations shown in Fig. 3.
Figure S16. Changes in twitch tension predicted by the five-state model for change in $k_{+A}$ up to ±50% from those used for Fig. 3 (dark green line), denoted as orig. $k_{+A}$. The main effects of changes in $k_{+A}$ on the tension responses are displayed in changes in the peak tension, resting tension, and the rate of tension rise, but the rate of relaxation only marginally changes in all cases and is much slower than the observed. The superscript o is omitted from the figure axis label for clarity.

Figure S17. Changes in twitch tension predicted by the five-state model for change in $k_{+D}$ up to ±50% from those used in Fig. 3 (dark green line), denoted as orig. $k_{+D}$. The main effects of changes in $k_{+D}$ on the tension responses are displayed in changes in the peak tension, resting tension, and the rate of tension rise, but the rate of relaxation changes only marginally in all cases and is about twofold slower than that observed. The superscript o is omitted from the figure axis label for clarity.
Figure S18. Changes in twitch tension predicted by the five-state model for change in $k_{+H}$ up to $\pm 50\%$ from those used in Fig. 3 (dark green line), denoted as orig. $k_{+H}$. The main effect of changes in $k_{+H}$ on the tension responses are displayed in modest changes in the peak tension, resting tension, the rate of tension rise, and the rate of relaxation. For all values of $k_{+H}$, the rate of relaxation is about twofold slower than that observed.

Figure S19. Changes in twitch tension predicted by the five-state model for change in $k_{-A}$ up to 24-fold from those used in Fig. 3 (dark green line), denoted as orig. $k_{-A}$. The main effect of changes in $k_{-A}$ on the tension responses are displayed as changes in the peak tension, resting tension, and the rate of tension rise. There are two distinctive regions in the peak and resting tensions: (1) the peak tensions modestly decrease for $k_{-A}$ increase up to twofold and resting tension decreases for $k_{-A}$ increase up to fourfold; and (2) for larger increases in $k_{-A}$, the peak tension rapidly falls, and resting tension reaches very low levels. Similarly, the rate of tension relaxation increases for $k_{-A}$ increase up to fourfold and then decreases. The superscript o is omitted from the figure axis label for clarity.
Figure S20. Changes in twitch tension at isometric half-sarcomere obtained from simulations with the five-state model. The tension transient predicted by the cross-bridge model without the PS, using parameters estimated by the best fit with the six-state model (Fig. 3 and Table 2 at 27.2°C) shows much higher peak tension and slow relaxation (black line). Increase of $k_{+A}$ by 2.5-fold significantly decreased peak tension (dark green line) and, in addition, increase of $k_{-A}$ for $\sim$17-fold brings the peak tension to the observed level, significantly increasing the rate of the relaxation (green line), but it is still not sufficient to reach the observed rate (gray line). Further decreasing $k_{-A}$ decreased the peak tension and slowed the relaxation rate. The superscript $o$ is omitted from the figure display for clarity.
Figure S21. Changes in twitch tension relaxation rates with increase predicted by the five-state model. The tension relaxation rate first increases to approach the observed values for $k_{-A}^o$, up to $\sim 350$ s$^{-1}$ (green arrow), but with further increases of $k_{-A}^o$, the relaxation rate starts to decrease (red arrow). At a lower value of $k_{+D}^o$ (gray triangle), the relaxation rate is slightly higher but still short of reaching the observed rate value. The superscript o is omitted from the figure display for clarity.
Figure S22. The best fit to the observations with the five-state model requires the change in power stroke rate $k_{+\Pi}$ (inset) in addition to changing $k_{o}^{-}$ and $k_{o}^{+}$. Effectively, a shift of $k_{\Pi}(x)$ for $\Delta x_{o} \approx 0.5$ increases the number of cross-bridges that can complete an ATPase cycle and enables faster relaxation. This change is achieved by an increase in $k_{cap}^{o}$ while keeping the equilibrium rate $K_{Pi}$ and the reverse power stroke rate unchanged (see Eq. A4). The superscript $o$ is omitted from the figure display for clarity.
Figure S23. **Effect of changes in baseline calcium level on resting and twitch tension.** In all simulations with cross-bridge cycle without PS (i.e., five-state model), the calcium transient (dashed pink line) with baseline $[\text{Ca}^{2+}]$ levels was reduced from observed values of 0.143 µM to 0.05 µM. MUSICO simulations with the observed $[\text{Ca}^{2+}]$ transient (dark pink line) with the same model parameters as shown in Fig. S22 predicted significantly elevated resting tension, slightly higher peak tension, and slower tension relaxation rate (dark green line). After returning to low $[\text{Ca}^{2+}]$ levels (i.e., at time >0.4 s), tension remains about the same as the resting tension. For reference, the twitch tension response to $[\text{Ca}^{2+}]$ transients with reduced baseline $[\text{Ca}^{2+}]$ level is shown as a green line.
Figure S24. Comparison of tension–pCa relationships between the cross-bridge cycle models with or without PS (i.e., the six- and the five-state models). The simulations of tension–pCa relationships in fixed-length intact trabeculae with the six-state model (red line) agree well with the experiments of (Gao et al., 1998; Janssen et al., 2002) at 22.5°C (symbols). The parameters used in these simulations are the same as those used in Fig. 7. The simulations of tension–pCa relationships in isometric half-sarcomeres with the five-state model (green line), using the parameters from the simulations that matched the observations (in Fig. S22), showed significantly higher sensitivity of tension at low levels of [Ca\(^{2+}\)] (pCa > 6.6) and lower cooperativity (i.e., Hill coefficient) than observed. In contrast, the simulations with the six-state model (dashed cyan line) using the same parameters as those for simulations of twitch contractions of isometric half-sarcomeres at 27.2°C (Fig. 3) matched the observations and showed lower levels of tension for pCa > 6.6. Notably, the five-state model predicted about the same resting tension at [Ca\(^{2+}\)] 0.143 μM (cyan-green arrow) as at the same baseline [Ca\(^{2+}\)] shown in Fig. S23.