A new myofilament contraction model with ATP consumption for ventricular cell model

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Received: 20 May 2017 / Accepted: 14 July 2017 / Published online: 2 August 2017 © The Physiological Society of Japan and Springer Japan KK 2017

Abstract A new contraction model of cardiac muscle was developed by combining previously described biochemical and biophysical models. The biochemical component of the new contraction model represents events in the presence of Ca2+-crossbridge attachment and power stroke following inorganic phosphate release, detachment evoked by the replacement of ADP by ATP, ATP hydrolysis, and recovery stroke. The biophysical component focuses on Ca2+ activation and force ($F_b$) development assuming an equivalent crossbridge. The new model faithfully incorporates the major characteristics of the biochemical and biophysical models, such as $F_b$ activation by transient Ca2+ ($[\text{Ca}^{2+}]$–$F_b$), $[\text{Ca}^{2+}]$–ATP hydrolysis relations, sarcomere length–$F_b$, and $F_b$ recovery after jumps in length under the isometric mode and upon sarcomere shortening after a rapid release of mechanical load under the isotonic mode together with the load–velocity relationship. ATP consumption was obtained for all responses. When incorporated in a ventricular cell model, the contraction model was found to share approximately 60% of the total ATP usage in the cell model.

Keywords Myofilament model · Mechano-energetics · Actomyosin–ATPase · Crossbridge kinetics · Troponin system

Introduction

Activation of muscle contraction results in a cycle of crossbridge (CB) attachment–detachment between the two major sliding filaments, actin and myosin. During this cycle, the CB converts the biochemical energy to contractile mechanical force by hydrolyzing ATP catalyzed by actomyosin (AM) ATPase. Since this is the major component of ATP consumption in the cardiac muscle [1], a detailed mathematical model of CB dynamics and associated ATPase activity is a prerequisite for analyzing the energetics of cardiac muscle contraction under physiological conditions.

A variety of whole-cell models have been developed to analyze mechanisms underlying the physiological regulation of cardiac contraction. Recently we developed a human ventricular cell model that includes both a detailed calcium ion (Ca2+)-induced Ca2+ release model and a contraction model in addition to membrane excitation supported by the regulation of intracellular ion concentrations [2, 3]. However, to our knowledge, no cell model has been reported to date which calculates ATP consumption during electrical excitation and contraction of the muscle based on the detailed reaction cascade of ATP hydrolysis by the S1 segment of myosin.

Recent studies have clarified many details of the biochemical and structural states of the CB cycle, and 11 predominant reaction states of AM ATPase have been identified in skeletal muscle. Lombardi and Piazzesi [4], Månsson [5], and Månsson et al. [6] reduced the scheme to four to six essential lumped states. These myofilament models are based on the Huxley hypothesis of independent CB behavior and calculate the state transition of CB based on the hypothetical Gibb’s free energy profile as a function of relative distance between myosin head and the nearest...
in the Månsson model, were simplified in reference to the original rate constants, which were theoretically elaborated state transition of the myofilament reactions in situ. The direct measure the rates of CB movement as well as the or isotonic contraction modes, since such experiments responses to square pulse application during the isometric tension evoked by applying step changes in the isometric tension evoked by the membrane excitation, as well as the events of the cardiac muscle described to date in bio-

The objective of the study reported here is to develop a new hybrid model which combines the reaction steps of ATP hydrolysis by the S1 segment (‘Månsson model’) with the biophysical contraction models that describe the Ca$^{2+}$-mediated activation of the myofilaments as well as the CB kinetics using mean-field approximation [8–14] instead of spatial explicit approaches [15, 16]. The Negroni and Lascano model (‘NL model’ [13, 14]) assumes a relatively simple equivalent CB (eCB), the distortion of which is represented with a linear spring and its sliding along the actin filament calculated by assuming a movable viscosity head. These approximations seem to be roughly in common with the Rice model [12]. The NL model has also been used to develop various types of ventricular models. Thus, we refer to the NL model in the present study because of its use in developing the hybrid model. The ‘NL model’ successfully reconstructs most of the key mechanical events of the cardiac muscle described to date in biological studies, such as the time course of developing the tension evoked by the membrane excitation, as well as the tensions evoked by applying step changes in the isometric and isotonic contraction modes at various [Ca$^{2+}$]. The model also successfully reconstructs the relationships of [Ca$^{2+}$] and force, of force and half-sarcomere length, and of force and velocity of shortening [17, 18].

The development of such a hybrid model may critically depend on reconstruction of the rapid time course of responses to square pulse application during the isometric or isotonic contraction modes, since such experiments directly measure the rates of CB movement as well as the state transition of the myofilament reactions in situ. The original rate constants, which were theoretically elaborated in the Månsson model, were simplified in reference to the kinetics of eCB. Thus, in the present study, we readjusted the rate of state transitions to obtain a new contraction model. When the hybrid contraction model was incorporated into the human ventricular model [3], ATP consumption by AM ATPase was approximately 60% of the total usage of the cell model. ATP consumption proportional to the pressure–volume trajectory area was obtained when the myofilament model was used in the ‘Laplace heart’ in the multi-scale hemodynamic model of cardiovascular system [19].

Materials and methods

The model of CB dynamics in the NL model is characterized by the introduction of the eCB, which largely facilitates the calculation of the developed force of contraction in a continuous manner. Even though the simplification was achieved without truly developing a specific algorithm for averaging the behavior of all CBs, the magnitude as well as the time-course of the reconstructed $F_h$ agrees well with the experimental findings [17]. This eCB assumption was adopted in the present model as in the original study. Details on kinetic equations in the NL model are described in the Electronic Supplementary Materials (ESM), with the exception of those described in the following subsections. For details on the rate constants defined in the Månsson model, the reader is referred to the original publications [4–6].

Model reaction scheme

The lumped four-state biochemical model developed by Månsson [5] well represents the four essential steps of ATP hydrolysis. In the new state–transition scheme shown in Fig. 1, these four states are represented by symbols of $cAM_{DPw}$, $cAM_{D}$, $cAM_{T}$, and $cM_{DP}$ ($cM_{T}$), respectively, where A, M, T, D, P, and c refer to actin, myosin, ATP, ADP, inorganic phosphate (P), and Ca$^{2+}$, respectively, and the lowercase letters w and s indicate weakly and strongly bound conformations of CB, respectively. The correspondence of each state with the original $A_{0}$, $A_{1}$, $A_{2}$ and $A_{3}$ states defined in the Månsson model is indicated below each state in the scheme shown in Fig. 1. A schematic composition of each state is indicated in the right panel with supplemental troponin–Ca$^{2+}$ binding. Note that calmodulin is aligned on the thin filament, and the Ca$^{2+}$-binding sites are occupied in all $cM_{DP}$, $cAM_{DPw}$, $cAM_{D}$, $cAM_{T}$, and $cM_{DP}$ ($cM_{T}$), in contrast to the three unoccupied states, $AM_{D}$, $AM_{T}$, $M_{DP}$ ($M_{T}$) aligned on the left relaxation pathway and associated with the same rate constants, $Z_{d}$ and $Z_{h}$, respectively, for simplicity. The angle of the lever arm in connection to the S1 segment shows the configuration of the myosin lever arm, i.e., the power stroke at $cAM_{D}$ and the recovery stroke at $cM_{DP}$ and $M_{DP}$. Thus, the Ca$^{2+}$-binding steps of $Q_{1}–Q_{2}$ and $Q_{10}–Q_{11}$ as well as the CB attachment steps $Q_{3}–Q_{4}$ are adopted from the NL model, while the other steps are all based on the Månsson model.
Rate constants for these four Ca\(^{2+}\)-bound states of ATP hydrolysis were adjusted in the present study to approximate the overall rate of ATP hydrolysis in situ at physiological temperature \([1,20]\), with reference to the theoretical ones determined as a function of \(\Delta G\) of each conformation in the original Månsen model. The NL model is also well adapted to the kinetics of the ventricular myocardium at 37\(\degree\) C. The time courses of the Ca\(^{2+}\) transition states followed by CB association and dissociation are quite similar to those observed in experiments \([21]\); namely, at systolic free intracellular Ca\(^{2+}\) (\([\text{Ca}^{2+}]_i\)) of approximately 0.5–1 \(\mu\)M at physiological temperature, the increase in \([\text{Ca}^{2+}]_i\) and the Ca\(^{2+}\)-induced troponin-switch, which corresponds to the \(\text{MDP}\rightarrow\text{cMDP}\) step, takes places within approximately 20 ms \([22,23]\).

**Rate function for state transitions in the NL model**

In the original NL model \([14]\), the state transition from \(\text{MDP}\) to \(\text{cMDP}\) represents lumped steps of the Ca\(^{2+}\) binding to troponin and the subsequent conformation change in the tropomyosin complex. Therefore, in the new NL model this step was conserved as in the original model. In the original model, a hypothetical troponin system (TS) was assumed to consist of three sequential troponin–tropomyosin regulatory units, assuming a nearest neighbor influence in the Ca\(^{2+}\) binding reaction \([10,15]\). However, the resultant Hill coefficient was <4 and still lower than the experimental value of 4–7 in the overall Ca\(^{2+}\)–F\(_b\) relationship in the experiments \([24]\). Thus, we readjusted the number of Ca\(^{2+}\) (\(n\text{Ca}\)) bound to the TS only for a phenomenological description. We found that an assumption of simultaneous binding of five Ca\(^{2+}\) (\(n\text{Ca} = 5\)) gave an overall Hill coefficient of 6–7 in the experimental Ca\(^{2+}\)–F\(_b\) relation (Fig. 2). Thus, the binding rate of Ca\(^{2+}\) to the TS is calculated as,

\[
Q_1 = Y_b \cdot [\text{MDP}] \cdot [\text{Ca}^{2+}]^{n\text{Ca}}
\]

\[
Q_{11} = Z_t \cdot [\text{MDP}] \cdot [\text{Ca}^{2+}]^{n\text{Ca}}
\]

The concentrations of all states in the scheme of state transition (Fig. 1) are given in terms of TS. Thus, it was assumed that a single TS controls simultaneously a number
The rate functions and equations for calculating state transitions are described in the ESM Tables S2 and S3. Equations and parameters to determine $F_b$ are the same as those in the original NL model, with the exception of a slight modification of $B_w$, as listed in ESM Table S1.

Concentration of the myosin S1 segment and magnitude of contraction

In the cell models published to date [26–34], variable troponin concentrations of approximately 70 μM were assumed. For the $[mS1]$, a molar ratio of actin:troponin concentration of 7:1 [35] and a ratio of myosin S1 segment:actin of 1:4.1 [36] were reported in cardiac muscle. These findings give a ratio of myosin S1 segment:troponin of 1.71, or 119.7 μM [mS1] in the previous cell models. This [mS1] is well within the range of direct biochemical measurements of the myosin head in ventricular tissue in various species, which are 200 (in pig [37]), 157 (in guinea pig [38]), and 144 (in rabbit [39]) μmol/kg wet weight. Here, we tentatively adopt the [mS1] of 200 μM.

In the NL model [14] the converting factors (stiffness of elastic CB structure) of $A_p$ and $A_w$ were used to calculate the magnitude of developed tension for a unitary cross-section area (in mm$^2$) of cardiac muscle for the powered and weak CB concentrations (in μM), respectively. In the present study, CB force ($F_b$) is given by the sum of weak CB force ($F_{bw}$) and powered CB force ($F_{bp}$), as the following equation (in mN/mm$^2$).

$$F_b = F_{bw} + F_{bp}$$

$$F_{bw} = nCa \cdot A_w \cdot [cAM_{DPw}] \cdot h_w$$

$$F_{bp} = nCa \cdot A_p \cdot ([cAM_{Ds}] + [AM_{Ds}]) \cdot h_p$$

Here, the $h_p$ and $h_w$ represent elongation of the elastic component of the strong-bound CB states ($cAM_{Ds}$ and $AM_{Ds}$) and weak-bound CB state ($cAM_{DPw}$), respectively. Note that variable CB elongations within the whole population of CB within a cell are represented by a single eCB having an ‘average’ CB elongation, $h_p$ and $h_w$, for the powered and weak eCBs, respectively in the NL model. The rate of change in eCB elongation and the amount of Ca$^{2+}$ bound to the TS were calculated in the same way as in the original NL model. The force of the parallel elastic element ($F_p$) is given by Eq. 14.

$$F_p = K_e \cdot (halfSL - hSL_0)^5 + L_e \cdot (halfSL - hSL_0)$$

where $hSL_0$ is the slack length of the parallel elastic element. The rate of change in the eCB elongation ($h_w$ and $h_p$) was calculated essentially in the same way as in the original NL model. The parameter values were adopted from Negroni et al. [25], as described in ESM Table S1.
The ATP consumption rate is calculated from two state transitions from $cAM_{Ds}$ to $cAM_T$ ($Q_2$) and from $AM_{Ds}$ to $AM_T$ ($Q_{12}$).

**Time-integration of ordinary differential equations**

The numerical integration was performed by Euler’s method with a time step of 0.01 or 0.005 ms. The Ca$^{2+}$ transient was generated by the empirical equations given in the NL model [14]. The model parameters and equations are described in the ESM Tables S1 to S5.

The following dimensions were applied; micromole (μM) for concentrations, milliseconds (ms) for time, milliNewton per millimeter squared (mN/mm$^2$) for force of contraction, and micrometer (μm) for length. All codes of the simulation program were prepared using Microsoft Visual Basic on Visual Studio Community 2013 (Microsoft Corp., Redmond, WA).

**Results**

**ATP hydrolysis is activated by [Ca$^{2+}$] in the new hybrid contraction model**

The AM ATPase is activated indirectly by Ca$^{2+}$, and repetitive cycles of hydrolysis are maintained in the presence of [Ca$^{2+}$]. The rate of ATP hydrolysis with accompanying developed tension $F_b$ was examined by applying various [Ca$^{2+}$] to the hybrid model. As the magnitude of $F_b$ is dependent on halfSL, the relationship between [Ca$^{2+}$] and the rate of ATP hydrolysis was examined at different halfSL (Fig. 2a, b). Although the underlying state transitions following Ca$^{2+}$ binding, including the A$_v$–A$_3$ steps are quite complicated (Fig. 1), it would appear that these [Ca$^{2+}$]–$F_b$ or [Ca$^{2+}$]–$v_{ATPase}$ relationships conform well to a saturation kinetic mechanism (Eq. 15).

$$v = \frac{v_{max}}{1 + \left(\frac{K_{0.5}}{[Ca^{2+}]_i}\right)^n_{H}} = \{F_b, v_{ATPase}\}$$

$$v_{max} = \{F_{b, max}, v_{ATPase, max}\}$$

Figure 2c and d indeed show approximately linear relationships at every [Ca$^{2+}$] with nearly constant value of $n_{H}$. The slope of the relationship was determined as a mean $n_{H}$ over an interval of −0.3 to 0.3 on the abscissa, which increased as 5.348, 5.787, 6.272, and 6.494 at halfSL of 0.8, 0.9, 1.0, and 1.1 μm, respectively. Simultaneously, the $K_{0.5}$ decreased as 1.271, 1.105, 0.980, and 0.923 μM in both the [Ca$^{2+}$]–$F_b$ and [Ca$^{2+}$]–$v_{ATPase}$ relationships. The $V_{max,ATPase}$ increased as 163.03, 245.87, 279.16, and 289.66 μM/ms and the $V_{max,Fb}$ as 737.3, 1112.0, 1262.6, and 1310.1 mN/mm$^2$ at 0.8, 0.9, 1.0, and 1.1 μm halfSL, respectively. Comparison of the relative amplitudes between these values revealed that $V_{max,ATPase}$ is quite proportional to $V_{max,Fb}$. The scope of the relationship became slightly shallower with increasing [Ca$^{2+}$], mostly due to the limited number of TS. Note that cooperativity was assumed only for Ca$^{2+}$ binding to troponin. These results are all quite consistent with the experimental findings [37, 41].

**Contraction and ATPase activities evoked by the idealized Ca$^{2+}$ transient**

The proportionality between $F_b$ and $v_{ATPase}$ in Fig. 2 is expected in the present hybrid model. Namely, both $F_b$ and $v_{ATPase}$ are largely determined by [cAM$_{Ds}$] and [AM$_{Ds}$] in both Eqs. 13 and 17. The first term in Eq. 12, which represents the contribution of weakly bound CB, is a minor component of the total $F_b$ during the steady presence of Ca$^{2+}$ because the $h_{w}$ quickly relaxes to a negligibly small $h_{w} = 0.0001$ μm. It may be noted that the denominator in Eq. 17 is constant since [ATP] and [ADP] are both given constants during the time course in Fig. 2.

$$v_{ATPase} = nCa \cdot (Q_7 + Q_{12})$$

$$= nCa \cdot \frac{Z_d \cdot ([cAM_{Ds}] + [AM_{Ds}])}{ATP} \cdot \left(1 + \frac{[ADP]}{K_{ADP}}\right)$$

ATPase activity was examined during the usually developed tension evoked by the transient Ca$^{2+}$ at a halfSL = 1.05 μm (Fig. 3). The model was activated by the standard transient Ca$^{2+}$ given by ESM Eqs. S14 and S15. The component of AM$_T$ was not included in the calculation of $F_b$, since this state in the new hybrid model represents a sum of (AM$_T$ + MT$_T$) in the Månssohn A$_2$ and A$_3$ states. In Fig. 3c, the time course of $v_{ATPase}$ is nearly similar to that of $F_b$ in Fig. 3b. During a single stimulus interval of 800 ms, the amount of ATP used was 3969 μM, which gives an average of ATPase turnover rate of 19.84/twitch (=3969/800 = 200 μM). The ATP consumption rate via transition step $Q_{12}$ during the removal of Ca$^{2+}$ was much smaller if compared with that in the step $Q_7$. 

\[ \text{Fig. 2c and d indeed show approximately linear relationships at every [Ca}^{2+}] 	ext{with nearly constant value of } n_{H}. \text{The slope of the relationship was determined as a mean } n_{H} \text{ over an interval of } -0.3 \text{ to } 0.3 \text{ on the abscissa, which increased as 5.348, 5.787, 6.272, and 6.494 at halfSL of 0.8, 0.9, 1.0, and 1.1 μm, respectively. Simultaneously, the } K_{0.5} \text{ decreased as 1.271, 1.105, 0.980, and 0.923 μM in both the [Ca}^{2+}]–F_b \text{ and [Ca}^{2+}]–v_{ATPase} \text{ relationships. The } V_{max,ATPase} \text{ increased as 163.03, 245.87, 279.16, and 289.66 μM/ms and the } V_{max,Fb} \text{ as 737.3, 1112.0, 1262.6, and 1310.1 mN/mm}^2 \text{ at 0.8, 0.9, 1.0, and 1.1 μm halfSL, respectively. Comparison of the relative amplitudes between these values revealed that } V_{max,ATPase} \text{ is quite proportional to } V_{max,Fb}. \text{The slope of the relationship became slightly shallower with increasing [Ca}^{2+}], \text{mostly due to the limited number of TS. Note that cooperativity was assumed only for Ca}^{2+} \text{ binding to troponin. These results are all quite consistent with the experimental findings [37, 41].} \]
Isotonic contraction

Compared to the isometric contraction, the isotonic contraction was characterized by the delayed peak in the developed tension $F_b$ and the ATP flux. This delayed peak was caused by the shortening of $h_p$ and thereby also of halfSL (Fig. 4e, f), in contrast to the isometric contraction. This decrease in $h_p$ largely inverted $h_w$ at the onset of contraction, followed by gradual relaxation toward the stable length of $h_w$ (0.0001 μm) during contraction. Thus, the CB detachment rate $g$, given as a function of $h_w$ deviation from $h_w^r$ (Eq. 5), was largely increased (Fig. 4g). On the other hand, the attachment rate $f$ was nearly constant, though increased only slightly. Through these changes in CB kinetics, the development of cAM$_{Ds}$ was largely delayed and its magnitude was smaller, when compared with the isometric contraction. Thereby, the peak tension and the peak of v_ATPase were delayed according to the time course of the state cAM$_{Ds}$. It should be noted that AM$_{Ds}$ is not visible due to overlapping cM$_{DP}$ (cM$_T$) (Fig. 4).

Taken together, Figs. 3 and 4 indicate that the time course of $F_b$ is quite similar to that in the original NL model [14]. Thus, it is evident that the addition of the ATP hydrolysis cycle to the NL model did not affect the general time course of contraction obtained in the new hybrid model.

The length clamp experiment in the hybrid model

The measurements of relaxation time course in $F_b$ evoked by applying a step change in halfSL during halfSL-controlled conditions provide essential parameters for the kinetics of the state transitions of the eCB conformations. Thus, we tested our new model by applying the length step shown in Fig. 5a and examined the $F_b$ (Fig. 5b) and ATP hydrolysis (Fig. 5c) at a constant $[Ca^{2+}]$ of 0.647 μM. At the onset of the length step, $h_p$ decreased to approximately 0 and then rapidly recovered within the subsequent 5 ms due to the rapid intrinsic rate of $dh_p/dt$ (Fig. 5e). The negative deflection of $h_w$ at the pulse onset caused rapid detachment of the eCB by approximately 80% through the
The transient increase in \( g \) according to Eq. 5. During the following slow relaxation phase of approximately 900-ms interval, the \( F_b \) recovered due to the re-equilibration of state transitions (Fig. 5d) as in the original NL model.

The time course of \( v_{\text{ATPase}} \) change (Fig. 5c) was parallel to the time course of \([cAM_D]_w\) and \([cAM_D]_b\), as shown in Fig. 5d. This finding indicates that ATP consumption should be transiently decreased and at the onset of the shortening length pulses. Thereafter, ATPase activity will gradually recover during the \( F_4 \) phase in actual experiments.

**The rapid-release experiment in the isotonic contraction mode**

The rapid-release experiment revealed the three phases in the time course of \( \text{halfSL} \) shortening, namely, the initial jump (phase 1), the rapid hyperbolic shortening (phase 2) within the initial 50 ms, and the subsequent slow almost linear shortening (phase 3), as observed typically in skeletal muscle (Fig. 6b, e). These simulation results are quite comparable to the relaxation time course demonstrated in the original NL model [14]. Similar time courses have been obtained in experiments using cardiac muscle preparation, but obviously at a low time resolution due to the intrinsic complexity of the trabeculae in the cardiac tissue [18].

The force clamp experiment is quite straightforward, namely the product of \( h_p \) multiplied by the number of eCB is proportional to the applied load (\( F_{\text{ext}} \)). The quick release instantaneously decreased \( h_p \) (upper line, Fig. 6d) or negatively deflected \( h_w \) (lower line, Fig. 6d). The deviation in \( h_w \) from \( h_w \) decreased the number of attached eCBs by accelerating the rate of detachment (\( g \)) of eCBs represented by Eq. 5, as in the length jump simulation. This decrease in the number of eCB increased \( h_p \) to balance the applied \( F_{\text{ext}} \). Through these two opposite influences of decreasing and increasing \( h_p \), the magnitude of \( h_p \) relaxed approximately to a new steady level during phases 1–2. The fraction of \( cAM_D_w \) reached a new equilibrium level through an almost simple sigmoidal time course. Therefore, the rate of ATP hydrolysis was almost linearly related to the level of test external load during the phase 3, as shown in Fig. 6g, which differs from the length clamp experiment.

It should be noted that the magnitude of \( h_p \) remained nearly equal to 6 nm during the phase 4 shortening. This means that the shortening of \( \text{halfSL} \) is attributable exclusively to the movement of eCB attachment point along the actin fiber. The rate of shortening in phase 4 was nearly an exponential function of the mechanical load in Fig. 6f, which is in rough agreement with experimental findings (Fig. 6e). On the other hand, the rate of ATP usage increased in proportion to the magnitude of \( F_{\text{ext}} \) from almost zero in the absence of \( F_{\text{ext}} \) (Fig. 6g). In summary, the turnover rate of the AM ATPase largely varied depending on both the magnitude of \( F_{\text{ext}} \) and \([Ca^{2+}]_c\) in the cytosol (\([Ca^{2+}]_c\)).

**Integration of the new contraction model into the ventricular whole cell model**

Parameters of the new hybrid model of contraction were also tuned by integrating the model in the human ventricular cell model (HuVEC model [3]). This was done because whole cell ATP consumption by the S1 segment has been well established by macroscopic measurements rather than by the study of dissected preparations in vitro. It should be noted that the estimation of ATP consumption by ionic pump activities in the whole cell cardiac cell models is now well established by the refined magnitude of ionic...
Ca$^{2+}$ content in the SR. Under these conditions, ATP consumption by the contraction was 58.6% of the sum of ATP consumption by the myofilaments, SR Ca$^{2+}$-ATPase (SERCA), Na/K, and plasma membrane Ca$^{2+}$ ATPase (PMCA) per cycle. This value is quite consistent with the approximately 60% consumption by the AM–ATPase in the beating blood perfused heart [1]. Other components, including SERCA, Na/K-ATPase, and PMCA, used 36, 4.5, and 0.6% of the total ATP consumption, respectively, under the physiological condition. A slightly different result was reported by Schramm et al. [45], who attributed 76% of the whole ATP consumption to AM–ATPase and, 15 and 9% to the SERCA and Na/K-ATPase, respectively.

Applicability of the present hybrid model to estimate the myocardial ATP usage in the simple blood circulation model

The NL model has been used to examine the hemodynamics in the multi-scale model of the cardiovascular system using ‘Laplace’s left ventricular hemispherical model’ [19, 30, 46]. We replaced the hybrid model for the original NL model in the hemodynamic model of Utaki et al. [46] and estimated ATP consumption during the pressure–volume (PV) trajectory. As expected from the faithful reproduction of the developed tension described above (Figs. 2, 3), the PV trajectory obtained by the new system model seems to be quite consistent with the results of previous studies (Fig. 8). The pressure–volume area (PVA) of the left ventricle, which correlates well with myocardial oxygen consumption per beat [1], changed (enlarged) with increasing preload scale (reviewed by Utaki et al. [46]) (standard $K_{rp}$: 0.8, 1.0, 1.2, 1.4, 1.6, and 1.8 in Fig. 8). ATP usage in the hybrid contraction model was simultaneously plotted in the same figure, but at different scale (Fig. 8a, shown in gray). During the ejection period, the ATP hydrolysis rate decreased because of the decrease in the number of powered CBs. This is different from the gradual increase in ventricular pressure due to the decrease in the radius of the hemisphere Laplace heart. The relationship between PVA (mmHg × ml) and the corresponding integration of ATP usage ($\mu$M/ms) was examined in Fig. 8b. The linear relationship between PVA and $v_{ATPase}$ definitely indicates the relevance of using the hybrid model in calculating ATP consumption.

Discussion

Brief summary

A new cardiac contraction model was developed by combining the biochemical model elaborated by Månsson [5]...
with the biophysical NL model. The new model is the successor to the dynamic CB models developed by Lombardi and Piazzesi [4], Piazzesi et al. [47], and Edman et al. [48], all of which are based on tetanus contraction in skeletal muscle. In contrast, the newly proposed NL model of the TS reconstructs \( F_b \) in the cardiac myocytes over a wide variety of physiological experimental findings. In the new hybrid model, the scheme of state transition among the \( A_0 \)–\( A_3 \) states in the Månsson model was used as it is, but the rate constants for individual state transitions were simplified. In the original Månsson model, the rate was calculated by assuming a Gibbs free energy profile as a function of the distance (\( x \)) between the CB head and the nearest binding site on actin fiber. In the hybrid model, a constant rate was used for individual state transitions of \( eCB \) by referring to the rate change based on the Gibbs free energy profile. The new model inherited well the major characteristics of both types of the two models, such as the concentration–response relation of \([Ca^{2+}]_c\)–\( F_b \) or \([Ca^{2+}]_c\)–\( v_{\text{ATPase}} \) (Fig. 2), the time course of the developed tension activated by the intracellular transient \( Ca^{2+} \) (Figs. 3, 4), the turnover rate of the AM ATPase, the \( F_b \) responses to jumps in the \( \text{halfSL} \) (Fig. 5), and \( F_{\text{ext}} \) (Fig. 6) under the isometric and isotonic contraction modes, respectively, and the load–velocity relation (Fig. 6). When the new hybrid model was incorporated into a ventricular cell model, ATP consumption by contraction was approximately 60% of the whole cell ATP usage at a cycle length of 0.8 s (Fig. 7). When the new model was implemented in the hemodynamic blood circulation model, ATP consumption was proportional to the PVA of the hemisphere Laplace heart (Fig. 8).

**Simultaneous reconstruction of ATP consumption and the developed tension**

The minimum requirement of any mathematical model of cardiac contracting fibers should be the capture of the following four essential steps of the CB cycle tightly coupled to the accompanying ATP hydrolysis:

**Step 1:** The binding of ATP in place of ADP to the catalytic domain of the S1 segment rapidly
detaches the myosin head from the actin binding site, resulting in relaxation of the rigor state [49–52].

Step 2: ATP is hydrolyzed in association with a structural change of a swing of the myosin lever arm (a recovery stroke), leaving products of ATP hydrolysis, ADP, and Pi bound to the active site of myosin head [50, 53, 54].

Step 3: Ca$^{2+}$ binding to troponin dislocates the tropomyosin complex during the time transient Ca$^{2+}$ is within the cell [55] and thereby allows the myosin head (carrying ADP and Pi) to bind with actin filament, forming a weakly bound state of CB [51, 56].

Step 4: The attachment of the myosin head to the actin binding site causes an approximately 100-fold increase of the rate of Pi release [6]. Dissociation of Pi from the S1 segment is tightly coupled with the power stroke of the CB, resulting in the sliding motion between the thin and thick filaments to stretch the elastic elements in the CB [57–59], which is responsible for the developed tension $F_b$.

Steps 1, 2, and 4 are precisely described in the biochemical models (Månsson model), while the NL model simulated well step 3 based on biophysical experimental findings, such as the time course of $F_b$ evoked by the transient Ca$^{2+}$, the response of the shortening of the halfSL evoked by the rapid release protocol in the isotonic contraction, and the time course of $F_b$ recovery evoked by the shortening step of the halfSL in the isometric contraction mode. The hyperbolic increase in shortening velocity with decreasing external load was reconstructed based on the time course of halfSL shortening reconstructed by the numerical integration in the model, by applying the quick release protocol according to the experimental findings. In the Månsson model, this relationship was theoretically predicted indirectly from the transition states. The present model clearly proposes that ATP usage is minimum in the absence of external load and increases in proportion to the magnitude of $F_{ext}$ over the physiological range (Fig. 6g).

We have shown that the new hybrid model successfully calculated the rate of ATP hydrolysis simultaneously with the accompanying development of $F_b$. Thus, the presented hybrid model largely facilitated development of new whole cell models for analyzing cardiac energetics, development of the force of contraction, as well as the ATP dependency of the muscle contraction, as shown in Fig. 7. The merit of using the NL model may be largely due to the introduction of the eCB, which represents the average behavior of the whole population of CBs within a cell. This kind of spatial average model of the whole population of CBs has been used in a variety of simplified models (for references see Introduction). In the NL model, the behavior of the eCB was thoroughly fitted to the classical experimental findings ([60–62] and models [8, 63, 64]). The time-dependent change of eCB elongation, $h_b$ or $h_w$, is continuous as described in ESM Eqs. S3 or S4, and the rate g of detachment of CB is given as a function of $h_w$ (2008 NL model [14]) or $h_p$ (1996 NL model [13]). This g provides the basis for explaining the halfSL dependence of the [Ca$^{2+}$–$F_b$] relationship [17] (Fig. 2). The relationship between the $F_{ext}$ and $dh/dt$ is also explained by the eCB kinetics, the rate of quick recovery of $F_b$ at the onset of the halfSL jump, and the time course of the shortening of halfSL evoked by the release of $F_{ext}$.

In the original model of Lombardi and Piazzesi [4], the fractions of CBs in the $A_1$, $A_2$, and $A_3$ states were calculated by applying the numerical integration method with a discrete interval of ($\Delta x$) of <0.5 nm at each x-point. It was also assumed that the elongation of CB was increased by an interval of the neighboring actin binding sites for each state transition of $A_1$–$A_2$, and $A_2$–$A_3$. When calculating the force–velocity relationship, the smooth continuous change in the velocity as well as the force of CBs were obtained theoretically by averaging for whole population of CB states without reconstructing the experimental response to quick release. Using a linear stiffness of the CB elongation, the force of contraction (for example, in units of mN/mm$^2$) was calculated from the sum of $A_2$ (weakly bound) + $A_3$ (strongly bound) over the range of $x$.

Negroni and Lascano elegantly simplified the calculation of developed tension and the CB kinetics by assuming the eCB might roughly represent the average CB elongation $h_b$ or $h_w$. Consequently, the unitary developed tension ($f_i$) is calculated with a stiffness ($a$) of the elastic structure ($f_i = a \times h$), and the movement rate of the CB head along the actin fiber is calculated as $dh/dt$. The $dh/dt$ is defined for both $h_p$ and $h_w$ as:

$$\frac{dh}{dt} = -B \cdot (h - h_c) \quad (18)$$

It should be noted that Eq. 18 is equivalent to ESM Eq. S3 provided that halfSL is constant during the time interval under consideration, as in the $L_1$ phase in the isometric contraction (Fig. 6). Thus, $dh/dt$ is used to calculate the movement of the CB head when attached. In the detached CB head, $h$ relaxes quickly to the resting elongation ($h_r$). Note that the $x_0$ giving the energy minima in the detached $A_3$ state of the Månsson model [5] is about 7–8 nm distant from the corresponding position of energy minima in the $A_1$ and $A_2$ state, which is close to $h_c$ (=6 nm) in the NL model, although Piazzesi et al. [47] and Edman et al. [48] assumed slightly different $x_0$ of energy minima. In the Månsson model the binding rate is described as a function...
of the velocity of the CB head movement along the actin fiber [5]; this assumption may correspond to the $h$-dependent detaching rate of $g$ in the NL model.

**Turnover rate of ATP hydrolysis in the new hybrid model**

The sequential steps of $Q_7^{-}Q_9$ and $Q_{12^{-}}Q_{13}$ in the new hybrid model were adopted from the Månsson model, in which the weakly bound CB was separated from the powered CB according to experimental findings [65–70] and then combined with the $Q_6^{-}Q_4$ and $Q_{10^{-}}Q_{11}$ steps of the 2008 version of the NL model. This modification did not largely modify the time course of developed tension when compared to the NL model because the newly implemented cycle of ATP hydrolysis is much faster than that in the entire state transition cycle in the original NL model. In the new hybrid model, ATP binding to the catalytic domain is also assumed in the unbound Ca$^{2+}$ state, $AM_{DS}$, which appears during the relaxing phase (shown in Fig. 1). This assumption is justified by the recent finding that ATP hydrolysis occurs in the reconstructed myosin fiber in the absence of both Ca$^{2+}$ and actin [54]. In our simulation (Fig. 3), the contribution of $AM_{DS}$, however, was only 1.5% of the total ATP consumption.

The average ATP turnover rate of the new contraction model was 19.85/s (twich in Fig. 3). Namely, quite consistent rates of 2.7–3.3/s [71–73] have been reported in a rat model of chemically skinned trabeculae in the presence of a saturating concentration of Ca$^{2+}$ at room temperature (20–21 °C). If these rates are corrected for the physiological temperature using the $Q_{10}$ of approximately 3.5 obtained by de Tombe and Stienen [74], an experimental turnover rate of 20–24/s is expected, which is slightly larger than the present simulation results. However, it should be noted that the rate of ATP consumption depends on several factors, such as the difference between isotonic and the isometric contraction modes (Figs. 3, 4), the $[Ca^{2+}]_{cyt}$ (Fig. 2), the external load (Fig. 5), and the halfSL (Figs. 5, 6). It should also be noted that the measurements of the turnover rate as well as $Q_{10}$ showed variations. A much larger ATP turnover rate (approx. 10/s per myosin head at 24 °C) has been reported in skinned rat trabeculae [75] and, in contrast, a lower value has been described in a pig cardiac preparation (approx. 0.5/s at 24 °C [38]) and in a rat preparation (approx. 3/s at 20 °C) using different experimental protocols [41, 71–73]. The temperature coefficient $Q_{10}$ of ATP hydrolysis is also variable (Siemankowski et al. [76], $Q_{10}$ = approx. 5; Burchfield and Rall [77], $Q_{10}$ = approx. 4; de Tombe and Stienen [74], $Q_{10}$ = approx. 3.5). Species-specific differences in the ATP hydrolysis rate might be attributed to differences in the composition of isoforms of the S1 myosin isoform [78].

The variability of the rate of ATP consumption, as described above, indicates the difficulty in comparing the turnover rate of ATPase activity in simulation and experiments due to the high variability of the recording conditions and data possibly not being completely described in experiments. Thus, any comparison of the relative weight of ATP consumption within a given cell model between the contraction and the ion pumps may be relevant in testing AM ATPase activity. Although additional fine-tuning of several parameters was required in both the contraction model and the cell model, the whole cell model presented here reconstructed well the experimental measurement of approximately 60% consumption by the contraction with reference to the sum of contraction and ion pumps (Fig. 7).

**The stiffness of single CB in ventricular whole cell**

In the present study, the CB concentration of ([CB] = 200 μM) was assumed by referring to corresponding values in conventional cardiac cell models and also to the direct biochemical measurements of CB (see Materials and methods). In the study reported here, we have examined if the [CB] assumed in our model is consistent with the stiffness of single CB of $\varepsilon = 0.5–3$ pN/nm which has been assumed to date in the Månsson model for calculating the $F_b$.

The value of $\varepsilon$ is calculated by the following equation using the force of contraction ($f_b$) and the elongation of the single CB ($h$).

$$\varepsilon = \frac{f_b}{h} = \frac{F_b}{f_b \cdot n \cdot h}$$  \hspace{1cm} (19)

A representative value set was $F_b = 20$ mN/mm$^2$, the fraction of activated CBs $f_b = 0.05$, $h = 6$ nm, and $half\text{SL} = 1$ μm. $n$ is the number of CBs within a volume of muscle fiber given as a product of $[half\text{SL} \times \text{unit cross-section area } (A) = 1 \text{ mm}^2]$ and is determined as:

$$n = N_A \cdot [CB] \cdot half\text{SL} \cdot A$$  \hspace{1cm} (20)

where, $N_A$ is the Avogadro’s number. The $\varepsilon$ thus obtained was 0.55 pN/nm, which is well within the range of 0.5–3 pN/nm assumed by Månsson [5] and Månsson et al. [6].

**Limitation of the new hybrid model**

Taken together, the simulation results of the hybrid model presented here are in good agreement to experimental data published to date. However, the limitation of this new contraction mode is obvious, since the ability to generate
realistic response does not prove that the underlying biological mechanisms are correctly represented.

Author Contributions YM and AN designed the study and developed the mathematical model. YM analyzed the experimental simulations and drafted the manuscript. AN and AA interpreted the data, discussed the results, and revised the manuscript. All authors have approved the final version of the submitted manuscript.

Compliance with ethical standards

Funding The authors received no external funding for this research.

Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval This article does not contain any studies with human participants or animals performed by any of the authors.

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