Phosphorylation of cMyBP-C Affects Contractile Mechanisms in a Site-specific Manner

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ABSTRACT Cardiac myosin binding protein-C (cMyBP-C) is a cardiac-specific, thick-filament regulatory protein that is differentially phosphorylated at Ser²⁷³, Ser²⁸², and Ser³⁰² by various kinases and modulates contraction. In this study, phosphorylation-site-specific effects of cMyBP-C on myocardial contractility and cross-bridge kinetics were studied by sinusoidal analysis in papillary and trabecular muscle fibers isolated from t/t (cMyBP-C-null) mice and in their counterparts in which cMyBP-C contains the ADA (Ala²⁷³-Asp²⁸²-Ala³⁰²), DAD (Asp²⁷³-Ala²⁸²-Asp³⁰²), and SAS (Ser²⁷³-Ala²⁸²-Ser³⁰²) mutations; the results were compared to those from mice expressing the wild-type (WT) transgene on the t/t background. Under standard activating conditions, DAD fibers showed significant decreases in tension (~50%), stiffness, the fast apparent rate constant $2\pi c$, and its magnitude C, as well as its magnitude H, but an increase in the medium rate constant $2\pi b$, with respect to WT. The t/t fibers showed a smaller drop in stiffness and a significant decrease in $2\pi c$ that can be explained by isoform shift of myosin heavy chain. In the pCa-tension study using the 8 mM phosphate (Pi) solution, there was hardly any difference in Ca²⁺ sensitivity (pCa₅₀) and cooperativity ($n_{\rm H}$) between the mutant and WT samples. However, in the solutions without Pi, DAD showed increased $n_{\rm H}$ and slightly decreased pCa₅₀. We infer from these observations that the nonphosphorylatable residue 282 combined with phosphomimetic residues Asp²⁷³ and/or Asp³⁰² (in DAD) is detrimental to cardiomyocytes by lowering isometric tension and altering cross-bridge kinetics with decreased $2\pi c$ and increased $2\pi b$. In contrast, a single change of residue 282 to nonphosphorylatable Ala (SAS), or to phosphomimetic Asps together with the changes of residues 273 and 302 to nonphosphorylatable Ala (ADA) causes minute changes in fiber mechanics.

INTRODUCTION

Cardiac myosin binding protein-C (cMyBP-C) is a 140 kDa thick-filament-associated protein (1) that plays both structural and functional roles in sarcomeres (2,3). Mutations in the cMyBP-C gene (MYBPC3) cause >40% of familial hypertrophic cardiomyopathy (FHC) (4), the leading cause of sudden cardiac death in young adults. cMyBP-C differs from the skeletal isoforms in that it contains multiple phosphorylation sites. These are phosphorylated in a site-specific way by at least six kinases: protein kinase A (PKA) (5,6), protein kinase C (PKC) (5), protein kinase D (PKD) (7), Ca²⁺/calmodulin-dependent protein kinases II (CaMKII) (6), ribosomal s6 kinase (8), and glycogen synthase kinase 3β (GSK3 β) (9). cMyBP-C is extensively phosphorylated under basal conditions (10), but its phosphorylation level decreases under pathological conditions (10-15). Despite extensive research on cMyBP-C, the physiological role of its phosphorylation is not well understood.

Three phosphorylation sites identified in mice and their homologous sites in humans are: Ser²⁷³, Ser²⁸², and Ser³⁰² (6,14,16). These residues are located in the M domain between domains C1 and C2. It was found that PKA phosphorylates all three sites, PKC phosphorylates Ser²⁷³ and Ser³⁰², and CaMKII phosphorylates only Ser²⁸² (17). Being

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a substrate of PKA and PKC (5,6), cMyBP-C is thought to be important in regulating myocardial contraction through a β -adrenergic signaling mechanism (18). Mutation of the three Ser residues to phosphorylation-deficient Ala results in impaired cardiac function, reduced contractile reserve, and cardiac hypertrophy in transgenic mice (19,20). In contrast, mutation of Ser to the phosphor-mimetic Asp does not impair cardiac function and results in a cardioprotective effect (21).

The phosphorylation sites in cMyBP-C are functionally unequal and hierarchically regulated (6). That is, the three phosphorylation sites do not act independently, but cooperatively (22). Among these three sites, Ser^{282} appears to play a central role, as evidenced by the inactivation of cMyBP-C phosphorylation at other sites when Ser²⁸² is mutated to Ala (6,17). It may be the consequence of a conformational change that makes other sites inaccessible to kinases (6). In a previous investigation, we studied the importance of Ser²⁸² phosphorylation on cardiac morphology and function, using the transgenic mouse model SAS (Ser²⁷²-Ala²⁸²-Ser³⁰²), in which Ser^{282} was mutated to Ala (17). At the same time, two additional mouse models, DAD (Asp²⁷³-Ala²⁸²-Asp³⁰², in which Ser²⁷³ and Ser³⁰² were mutated to the phosphorylation-mimetic Asp), and ADA (Ala²⁷³-Asp²⁸²-Ala³⁰²), were studied (17). It was found that the hearts of DAD mice, but not those of their ADA or SAS counterparts, showed significant hypertrophy, fibrosis, and myocyte disarray, mimicking the t/t (null) phenotype (17). Based on these observations, we



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hypothesized that DAD diminishes cardiac contractility and cross-bridge kinetics. The objective of this study was to determine the consequences of phosphorylation at different cMyBP-C sites for myocardial contractility and cross-bridge kinetics, using the ADA, DAD, and SAS mutant mice. The wild-type (WT) cMyBP-C transgene was introduced on the cMyBP-C null (t/t) background, and this cDNA was subjected to site-directed mutagenesis to generate ADA, DAD, and SAS transgenes. The results from each mouse model were compared with those from mice expressing the WT transgene. Our results indicate that the cross-bridge kinetics are significantly altered in mice carrying the DAD mutation, and that this results in a reduction of isometric tension.

MATERIALS AND METHODS

Animal models

The transgenic mouse models used in this study were reported in a previous publication (17). The t/t mouse is a hypertrophic cardiomyopathy mutant in which modifications to the carboxyl end of the cMyBP-C protein cause it to be absent from sarcomeres (23). The strain referred to as WT is a transgenic line in which the WT cMyBP-C gene was introduced into the t/t background. Three mutant transgenes were also created, in which Ser residues of cMyBP-C at positions 273, 282, and 302 were either unchanged or mutated to Ala or Asp to produce a phosphorylation-deficient or phosphorylation-mimetic effect, respectively. In the mutant names, Ala is coded as A, Asp is coded as D, and Ser is coded as S, in the standard way, and each mutant is identified by the three-letter code: ADA (expressing Ala²⁷³-Asp²⁸²-Ala³⁰²), DAD (Asp²⁷³-Ala²⁸²-Asp³⁰²), and SAS (Ser²⁷³-Ala²⁸²-Ser³⁰²). Information on the number of mice used, including their age and gender, is listed in Table 1. These animal experiments were in accordance with the guidelines of the Institutional Animal Care and Use Committees at Loyola University in Chicago and The University of Iowa, as well as those listed in the Guide for the Use and Care of Laboratory Animals published by the National Institutes of Health.

Muscle preparations

Each mouse was euthanized through intraperitoneal injection of Na pentobarbital at 50 mg/kg, and the heart was removed. The whole heart was

TABLE 1 Gender and age for the adult mice used in this study

Mouse model	Gender	Age (months)
WT	Male	5.8
WT	Female	4.5
WT	Male	2.3
t/t	Male	2.7
t/t	Male	5.1
t/t	Male	5.1
t/t	Male	2.4
t/t	Male	2.4
ADA	Male	5.0
ADA	Male	8.0
ADA	Male	8.0
DAD	Male	7.4
DAD	Male	5.5
SAS	Male	10.4
SAS	Female	4.4
SAS	Male	2.9

perfused using the Langendorff system with perfusion buffer at 37°C for 5 min. The hearts were then moved sequentially to Na skinning solution for 15–30 min, K skinning solution for 30 min, and storage solution on ice overnight. On the next day, the storage solution was replaced once. The compositions of the solutions used have previously been reported (24). The hearts were then shipped on ice to Iowa City, where they were stored at -20° C until used for biophysical studies. The experiments were performed two days to four weeks after shipment. There were hardly any changes in the performance, with respect to either isometric tension or rate constants, during this period. In the cases of preparations for which a change in one of these parameters was observed, we disposed of the data rather than proceeding with further analysis. With these criteria, many of the preparations were usable beyond four weeks.

The method used for the mechanical measurements has been described previously (25); the compositions of the solutions are found in our earlier report (24). In brief, papillary and trabecular muscle fibers from both ventricles ~1 mm in length and 100 μ m in diameter were dissected and mounted on the experimental apparatus by attaching their ends to two stainless steel wire hooks using nail polish. One hook was connected to a length driver, and the other to a tension transducer. Fibers were soaked for 3–5 min in 600 μ L relaxing solution, followed by further skinning in the relaxing solution containing 1% Triton X-100 for 20 min. Fibers were then washed with the relaxing solution and stretched to remove the slack. This procedure generally resulted in a sarcomere length of 2.1–2.2 μ m, as judged by optical diffraction using a He-Ne laser (wavelength: 0.6328 μ m) (25). Fiber length (L_0) and cross-sectional area were recorded at this time. All the solutions were applied twice (600 μ l each) to ensure that significant amounts of the old solution did not remain.

Solutions

Each fiber was tested in the standard activating solution, followed by the pCatension and pCa-stiffness study. The standard activating solution contained (in mM) 6 K2CaEGTA, 6.1 Na2H2ATP, 15 Na2CP (creatine phosphate), 6.6 MgAc₂ (Ac, acetate), 8 K_{1.5}H_{1.5}Pi (phosphate), 13 NaAc, 83 KAc, 10 MOPS, and 80 U/mL CK (creatine kinase); the ionic compositions of this solution were pCa $(-\log_{10}[Ca^{2+}])$ 4.55, Mg²⁺ 1 mM, and MgATP²⁻ 5 mM, pH 7.00, and ionic strength (IS) was 200 mM (adjusted by KAc). The pCa-tension and pCa-stiffness relations were studied in two sets of solutions, 8Pi and 0Pi. The 8Pi solution is our standard activating solution, in which the total Ca concentration was changed from 0.94 mM to 6 mM to produce pCa levels in the range 7.00-4.55, respectively. The OPi solution did not contain added Pi, and contained (in mM) 7 K2CaEGTA, 2 Na2H2ATP, 15 Na2CP, 3 MgAc2, 81 KAc, 10 MOPS, and 80 U/ml CK; the ionic compositions of this solution series were Mg^{2+} 0.78–0.89 mM and $MgATP^{2-}$ 1.62 mM, pH 7.00, with IS 150 mM (Table 1 of Bubb et al. (26)). pCa was adjusted to 7.00-4.40 by changing the total Ca concentration across the range of 1.1-7.0 mM, respectively. The OPi solution has been used by many other investigators (27-31), whereas we have been using the 8Pi solution (24,33-35).

Sinusoidal analysis

The sinusoidal analysis is complementary to step analysis, in which length changes are limited to a fraction of 1% of the total length; the details of this analysis method are described in Wang et al. (24). In brief, when tension reached a plateau, fiber length was oscillated in sine waves of varying frequencies in the range 0.5–140 Hz at a small amplitude (0.125%, corresponding to 1.3 nm at the half-sarcomere level). Both length (strain) and tension (stress) time courses were digitized simultaneously using two 16-bit A/D converters without filtration. The complex modulus Y(f) was defined as the ratio of the stress change to the strain change expressed in the frequency domain. Y(f) consists of two components: the viscous modulus (imaginary part of Y(f)) and the elastic modulus (real part of Y(f)). For cardiac muscle, Y(f) was fitted to Eq. 1 to resolve into two exponential processes, B and C (33):

Process B Process C

$$Y(f) = H - \frac{Bfi}{b+fi} + \frac{Cfi}{c+fi}$$
(1)

Processes B and C reflect the active interaction of myosin cross-bridges with the thin filament, which leads to the transduction of chemomechanical energy, because they are absent in the rigor condition (33). Process B corresponds to phase 3 of step analysis, which is a medium frequency-exponential delay during which the muscle generates oscillatory work. Process C corresponds to phase 2, which is a high-frequency exponential advance during which the muscle absorbs work. In fast-twitch skeletal muscle fibers, an additional exponential process A was observed (33-35). In cardiac muscles, process A is observed only at temperatures $\geq 30^{\circ}$ C (see Fig. 3, C, F, I, L and O in Lu et al. (36)). The study described here was performed at 20°C to result in two exponential processes, as shown by Eq. 1. The rate constants of delayed tension $2\pi b$, and fast tension recovery $2\pi c$, are the apparent rate constants of processes B and C, and B and C are their respective magnitudes (amplitudes); H is the O(zero) frequency stiffness. These were deduced by fitting Y(f) to Eq. 1. The complex modulus Y(f) extrapolated to infinite (∞) frequency is: $Y_{\infty} = H - B + C$. Y_{∞} corresponds to phase 1 of the step analysis, and is loosely called stiffness in muscle mechanics literature.

pCa-tension and pCa-stiffness studies

Tension increased when the pCa was changed sequentially from high to low, as indicated in Fig. 1. The pCa-tension relationship was individually fitted to the Hill equation:

Tension =
$$\frac{T_{\text{max}}}{1 + \left(\frac{\text{Ca}_{50}}{[\text{Ca}^{2+}]}\right)^{n_H}}$$
 (2)

where T_{max} is the maximum tension developed at saturating [Ca²⁺], Ca_{50} is the Ca²⁺ concentration at half-saturation (apparent Ca²⁺ dissociation constant), and n_{H} is the Hill factor (describes cooperativity among contractile proteins, including troponin, tropomyosin, myosin, and cMyBP-C) based on tension. pCa₅₀ = $-\log_{10}Ca_{50}$ indicates Ca²⁺ sensitivity. The pCastiffness relationship was fitted to a similar equation:

Stiffness =
$$\frac{S_{\text{max}}}{1 + \left(\frac{Ca_{50S}}{[Ca^{2+}]}\right)^{n_{HS}}}$$
 (3)



FIGURE 1 Slow time courses of length (*A*) and tension (*B*), demonstrating the experimental protocol. This example is from an ADA fiber preparation (0.9 mm in length and 87 μ m in diameter). The amplitude of sinusoidal oscillation was kept at 0.125% L₀ in all frequencies, but because the signal was filtered with a second-order low-pass filter (cutoff at 10 Hz), the amplitude looks attenuated at higher frequencies (to the left). R, relaxing solution; A, standard activating solution. *2 indicates that the solution was changed twice.

where S_{max} is the maximum stiffness developed at saturating [Ca²⁺], Ca_{50S} is the Ca²⁺ concentration at half-saturation, and n_{HS} is the Hill factor (which describes cooperativity determined based on stiffness). pCa_{50S} = $-\log_{10}Ca_{50S}$ indicates Ca²⁺ sensitivity based on measurement of stiffness.

Identification of myosin heavy chain isoforms

To determine myosin heavy chain (MHC) isoform compositions, α - and β -isoforms were separated on a 6.25% acrylamide Hoefer gel (99:1 acrylamide:bis-acrylamide) by SDS-PAGE. Gels were fixed in 10% acetic acid followed by overnight staining with Sypro-Ruby. Gels were imaged and the percentage of myosin isoforms was determined using Image J software.

Statistics

In each experiment, >10 fibers were dissected from multiple hearts from each mouse model (as indicated in Table 1) and studied. All data are expressed as the mean \pm SE. One-way ANOVA was performed to determine the significance of the difference among test groups. Post hoc tests (least significant difference) that allow pairwise multiple comparisons were applied to locate the difference among groups. The MHC expression level measured from SDS-PAGE was compared with Bonferroni post-hoc test (see Fig. 8). A significant difference is defined as $p \le 0.05$, and highly significant as $p \le 0.01$.

RESULTS

All the values of biophysical parameters reported in this section are listed in Table S1.

Tension and stiffness during maximal Ca²⁺ activation

The cardiac fiber was soaked first in the relaxing solution and then in the standard activating solution. When the steady state in tension was achieved, the computer was triggered to perform sinusoidal analysis and to collect the fast-time-course data for length and tension signals as reported (37). The slow time courses were traced by a pen recorder, and typical records are shown in Fig. 1 *A* (length) and Fig. 1 *B* (tension). The fiber was subsequently relaxed. After this initial test, the fiber was activated in a series of solutions containing increasing amounts of Ca^{2+} .

The above experiments were repeated for 11–38 fibers from each mouse model and the results were averaged. The complex modulus data were fitted to Eq. 1 to extract two exponential processes. Their parameters were used to calculate the elastic modulus Y_{∞} , which is called stiffness. Isometric tension and stiffness are plotted for five groups of mice at pCa 4.55 in the 8Pi and 200IS solution (Fig. 2 *A*) and at pCa 4.40 in the 0Pi and 150IS solution (Fig. 2 *B*). These results demonstrate that tension and stiffness were significantly lower (by ~48%) in the DAD group than in the WT group (p < 0.01) in both solutions. It was also noted that stiffness was significantly lower in the t/t group in the 8Pi solution (Fig. 2 *A*), and in the ADA group in the 0Pi solution (Fig. 2 *B*), than it was in the WT mice.



FIGURE 2 Isometric tension (T_{max} in Eq. 2; *left ordinate*) and stiffness (S_{max} in Eq. 3; *right ordinate*) in 8Pi solution (A) and 0Pi solution (B) at saturating [Ca²⁺]. The 8Pi solution contained 8 mM Pi and its IS was 200 mM, whereas the 0Pi solution contained no added Pi and its IS was 150 mM. **p < 0.01 and *p < 0.05 compared to WT. The number of experiments (n) in A is 31 for WT, 38 for t/t, 32 for ADA, 31 for DAD, and 27 for SAS; in B, n = 14 for WT, t/t, ADA, and DAD, and n = 11 for SAS.

Tension was, as anticipated, 50–60% greater in the 0Pi solution (Fig. 2 *B*) than in the 8Pi solution (Fig. 2 *A*). This is due to the differences in both [Pi] and IS, as an increase in either can decrease isometric tension in skeletal (38,39) and cardiac (40,41) muscle fibers. Similarly, stiffness was 20–30% greater in the 0Pi solution than in the 8Pi solution.

Cross-bridge kinetics

During the tension plateau of the standard activation, sinusoidal analysis was performed as described in Materials and Methods, and the complex modulus data, Y(f), were collected and plotted as discrete points in Fig. 3. These data are equivalents of tension transients in response to step changes in length. Fig. 3 includes plots of the elastic modulus versus frequency (Fig. 3 *A*), the viscous modulus versus the elastic modulus (Nyquist plots) (Fig. 3 *C*). Each Nyquist plot shows two contiguous semicircles. Hence, the complex modulus data can be resolved into two exponential processes, B and C, as shown in Eq. 1. Fig. 3 reveals that although the WT, t/t, ADA, and SAS groups had fairly

similar complex modulus values, those of the DAD group were considerably smaller, as evidenced by the considerably smaller diameter of the semicircle in the Nyquist plot at frequencies >20 Hz (Fig. 3 *C*).

The complex modulus data were fitted to Eq. 1, and the best-fit data are shown in Fig. 3 as continuous curves. Fig. 3 A shows that the elastic modulus remained small in the low-frequency range (<7 Hz), reached its minimum value at ~7 Hz (this frequency has been referred to as $f_{\rm min}$ by some investigators (42)), and increased significantly in the high-frequency range (>7 Hz). Fig. 3 B indicates that the viscous modulus had small negative values in the lowfrequency range (<5 Hz), reached a shallow minimum at ~2 Hz, increased steeply in the range 5-25 Hz, reached its maximum value at ~30 Hz, and decreased again in the high-frequency range (50-140 Hz). The frequency at which the viscous modulus is at its minimum approximates the characteristic frequency, b, and that at which it is at its maximum approximates the characteristic frequency, c. In Fig. 3 B, it is evident that the frequency associated with the maximum viscous modulus is shifted to the left in the t/t group, indicating that $2\pi c$ is slower.



FIGURE 3 Averaged complex modulus data in the 8Pi solution. Data were plotted as elastic modulus (real Y(f)) versus frequency (f) (A), viscous modulus (imaginary Y(f)) versus frequency (B), and viscous modulus versus elastic modulus (Nyquist plot) (C). n = 31 for WT, 38 for t/t, 32 for ADA, 31 for DAD, and 27 for SAS. Solid lines represent Eq. 1 with best-fit parameters. In C, frequency increases in the clockwise direction; its value can be found in A and B.

The fitting of the data in Fig. 3 to Eq. 1 yielded two apparent rate constants, $2\pi b$ and $2\pi c$, and their respective magnitudes, B and C, as well as magnitude H, all of which are plotted in Fig. 4 for the 8Pi solution at pCa 4.55, and in Fig. 5 for the 0Pi solution at pCa 4.40. Fig. 4 A (*left ordinate*) reveals that the rate constant $2\pi b$ for the DAD and SAS groups was significantly larger than that for the WT mice, and $2\pi b$ of the t/t and ADA groups was similar to that of the WT group. The rate constant $2\pi c$ (Fig. 4 A, right ordinate) for the t/t and DAD groups was significantly less than that for the WT, but $2\pi c$ for the ADA and SAS groups was similar to that for the WT. The different directions of the two rate constants are not surprising, because each rate constant reflects a different event in the cross-bridge cycle (43,44). Fig. 4 B (left ordinate) indicates that magnitude B for the t/t and SAS groups was larger than that for the WT group (p < 0.05), and that B of the ADA and DAD groups did not differ significantly from that of WT. Magnitudes C and H were significantly smaller in the DAD group than in the WT group (p < 0.05), but were similar across the other groups (Fig. 4 B). In general, magnitude B was about half of magnitude C but twice magnitude H in the 8Pi solution.

The apparent rate constants $2\pi b$ and $2\pi c$ deduced from the OPi solution (Fig. 5 A) were $\sim 1/5$ and 1/3, respectively, of those deduced from the 8Pi solution (Fig. 4 A). Similarly, magnitude B was ~1/10, and magnitude C ~1/2, of the corresponding values in the 8Pi solution. Thus, the resolution of processes B and C was much less in the OPi solution than in the 8Pi solution. Because of this, the relative error bars plotted in Fig. 5 are larger than those in Fig. 4. Due to the small magnitude and large scatter of the data, the only statistical difference in the rate constants measured among the five groups was a decreased $2\pi c$ in the t/t group. The magnitudes B and C did not differ significantly among all groups, but were somewhat lower in the DAD group. Magnitudes H were significantly smaller in the DAD group than in the WT group (p < 0.05). In all groups, magnitude H was three times larger in the OPi solution than in the 8Pi solution.

pCa-tension and pCa-stiffness studies

pCa-tension and pCa-stiffness results are plotted in Fig. 6 (at 8Pi and 200IS) and Fig. 7 (at 0Pi and 150IS), as discrete points after proper normalization. The data for each preparation were fitted to Eq. 2 (tension) or Eq. 3 (stiffness), and the fitted parameters were averaged. The best-fit curves are shown by solid lines in Figs. 6 and 7. The averaged pCa₅₀ and $n_{\rm H}$ values are presented in Figs. 6, *B* and *D*, and 7, *B* and *D*.

In the 8Pi solution (Fig. 6 *B*, *left ordinate*), pCa₅₀ and $n_{\rm H}$ (Fig. 6 *B*, *right ordinate*) were centered at ~5.56 and ~6.0, respectively, based on tension, and neither differed significantly between the mutant and WT groups. Almost identical results were obtained based on stiffness: pCa_{50S} was centered at ~5.56, $n_{\rm HS}$ was centered at ~6.0, and the values from the five groups were not significantly different (Fig. 6 *D*).

In the OPi solution (Fig. 7, B and D), pCa_{50} was larger and $n_{\rm H}$ was smaller compared to their values in the 8Pi solution. pCa₅₀ and pCa_{50S} centered at ~5.68, and $n_{\rm H}$ and $n_{\rm HS}$ centered at ~2.7. Thus, it can be said that cardiac muscle fibers are more sensitive to Ca^{2+} in the OPi solution than in the 8Pi solution (by 0.12 pCa unit), but cooperativity in the OPi solution is about half that in the 8Pi solution. These results are consistent with our recent study on myosin regulatory light chain mutations in mice (24). In addition, values among the five groups varied more extensively in the OPi solution, and differed significantly in the OPi versus the 8Pi solution. The cooperativity was significantly larger in the DAD group than in the WT group with respect to both tension (p < 0.05) and stiffness (p < 0.01). Ca²⁺ sensitivity was less in the DAD group than in the WT group with respect to both tension (p = 0.054) and stiffness (p < 0.05).

Cross-bridge kinetics during partial activation

As indicated in Fig. 1, sinusoidal analysis was also performed during the pCa study with 8Pi series solutions.



FIGURE 4 Parameters of exponential processes in the 8Pi solution (pCa 4.55). (A) Apparent rate constants $2\pi b$ (*left ordinate*) and $2\pi c$ (*right ordinate*). (B) Magnitudes of processes B (*left ordinate*) and C (*right ordinate*). (C) Magnitude H. Error bars represent the mean \pm SE of 26–38 experiments carried out in each group (for numbers of mice, see Table 1). n = 31 for WT, 38 for t/t, 32 for ADA, 31 for DAD, and 27 for SAS. **p < 0.01 and *p < 0.05 compared to WT.

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FIGURE 5 Parameters of exponential processes in the 0Pi solution (pCa 4.40). (*A*) Apparent rate constants $2\pi b$ (*left ordinate*) and $2\pi c$ (*right ordinate*). (*B*) Magnitudes of processes B (*left ordinate*) and C (*right ordinate*). (*C*) Magnitude *H*. n = 14 for WT, t/t, ADA, and DAD, and n = 11 for SAS. **p < 0.01 and *p < 0.05 compared to WT.

The rate constants $2\pi b$ and $2\pi c$ are plotted in Fig. 8. At lower levels of activation (pCa ≥ 6.0), the signal-to-noise ratio was too small to resolve the cross-bridge kinetics. Between pCa 5.8 and 5.4, $2\pi b$ increased steeply and $2\pi c$ decreased steeply. Between pCa 5.4 and 4.5, $2\pi b$ decreased slightly, and $2\pi c$ remained approximately the same. At all pCa points, the order of $2\pi b$ and $2\pi c$ values among all mouse groups remained the same as that shown in Fig. 4 *A*, with larger $2\pi b$ in the DAD and SAS groups, and smaller $2\pi c$ in the t/t and DAD groups.

Myosin α - and β -isoform expression

Because it has been known that MHC isoforms affect the apparent rate constant $2\pi c$ (45) and the shifts in isoform distribution have been observed in some transgenic mice (46), we assessed the MHC isoform contents as described in

Materials and Methods. A representative gel is shown in Fig. 9 *A*, and the analysis of the averaged data from three gels is shown in Fig. 9 *B*. This analysis demonstrated that WT mouse hearts contained 95.2 \pm 0.2% α -MHC (and 4.8% β -MHC), that ADA contained 86 \pm 6% α -MHC, and that DAD contained 84 \pm 2% α -MHC; the values for the latter two groups were not significantly different from that of WT. In contrast, α -MHC was reduced to 57 \pm 4% in t/t mice and to 74 \pm 3% in SAS mice, and β -MHC increased significantly to 43 \pm 4%, and 26 \pm 3%, respectively.

Results excluding female mice

There may be a gender difference. Since we mixed results from 14 male mice and two female mice (Table 1), we constructed plots for the males only (Figs. S2–S7)







FIGURE 7 Effect of pCa in the 0Pi solution in five groups of mice. Nomenclatures and conventions used are as in Fig. 6. n = 14 for WT, t/t, ADA, and DAD, and n = 11 for SAS. ** $p \le 0.01$, *0.01 < $p \le 0.05$, and (*)0.05 < $p \le 0.1$ compared to WT.

to examine whether this makes a difference. The figure numbers, panel designations, and plotting conventions are the same as those in the main text. When the data were limited only to males, there were no substantial changes in the results, but the error bars became larger because of the smaller population. The exceptions were the larger active tension (Figs. S2 A) and magnitudes B, C, and H (Fig. S4) in the SAS group, in which the increase in tension and magnitudes C and H became significant ($p \le 0.05$), and B became more significant ($p \le 0.01$), compared to WT.

DISCUSSION

The purpose of this study was to characterize the biophysical functions of cardiac muscle fibers that carry ADA, DAD, SAS, and null (t/t) forms of cMyBP-C, and to thereby advance our understanding of the role of cMyBP-C phosphorylation. Tension and stiffness, together with the apparent rate constants ($2\pi b$ and $2\pi c$) and the magnitudes (*B* and *C*) of two exponential processes were studied. Rate constants $2\pi b$ and $2\pi c$ are closely associated with the elementary steps of the cross-bridge cycle (25), including the binding of ATP and the release of Pi and ADP. These parameters can be used to characterize contractile properties and cross-bridge functions. pCa studies have been performed to establish the Ca²⁺ sensitivity and cooperativity.

In this report, two sets of solutions were used. The first is termed 8Pi solution and contains 5 mM MgATP²⁻, 8 mM Pi, and 200 mM IS, close to the composition in the cytosol of contracting cardiomyocytes, which is reported to be



FIGURE 8 Apparent rate constants $2\pi b$ (*A*) and $2\pi c$ (*B*) plotted against pCa in the 8Pi solution. Values at pCa ≥ 6.0 are not included, because they were seriously contaminated with noise. n = 17 for WT, 13 for t/t, 14 for ADA, 18 for DAD, and 12 for SAS.



FIGURE 9 Expression of myosin α - and β -isoforms in each mouse group. (*A*) A representative experiment. Isoforms detected using specific antibodies after Sypro stained SDS-PAGE analysis. The molecular mass of α -MHC is 223.565 kDa, and that of β -MHC is 222.849 kDa. (*B*) Fractions of α - and β -MHC, as averaged from three gels (n = 3), similar to that represented in *A*, for WT and mutant cMyBP-C. Error bars are drawn for each fraction. * $p \le 0.05$.

6–9 mM [Pi] (47), 4–6 mM [MgATP^{2–}] (47), and 215 mM IS (48). The [ATP] and [Pi] data were obtained using the chemical method in working hearts with adequate oxygenation (47). Given that the physiological ionic strength of cardiomyocytes has not been reported, the [IS] is based on resting skeletal muscle fibers (48). The second solution is termed 0Pi solution, and it has been used by many other investigators to characterize pCa-tension relationships (27–30,49,50). This solution does not have any added Pi, [MgATP^{2–}] is 1.6 mM, and IS is 150 mM. Because of the contaminating Pi (mostly via ATP and phosphocreatine) and the liberation of Pi from ATP hydrolysis, the actual [Pi] in the 0Pi solution is in the range of 0.15–0.8 mM (51–53).

The tension values in Fig. 2 (8Pi solution) do not differ from those previously reported in cardiac fibers (54) or from those discussed in our previous work (24). The [Pi] significantly influenced muscle contraction in many aspects. The increased [Pi] caused less active tension and stiffness (Fig. 2 A versus Fig. 2 B), larger rate constants and magnitudes B and C (Fig. 4, A and B, versus Fig. 5, A and B), but less magnitude H (Fig. 4 C versus Fig. 5 C); $n_{\rm H}$ became larger and pCa₅₀ became smaller (Fig. 6, B and D, versus Fig. 7, B and D) regardless of mouse genotype in this study. The differences between the mouse groups in either tension or stiffness under OPi and 8Pi solutions were similar (Fig. 2). Most of the significant changes in the cross-bridge kinetics occurred in the 8Pi solution (Fig. 4). The trend was similar in the OPi solution (Fig. 5), although this was not a significant difference owing to the smaller magnitudes B and C, which made the rate-constant data more scattered. In contrast, with the pCa-tension study, the genotype difference was more clear in the 0Pi solution (Fig. 7, B and D) than in the 8Pi solution (Fig. 6, B and D). The kinetics of the cross-bridge cycle appear to be modified through the thin filament activation mechanism; therefore, in partial activation (pCa > 5.4), $2\pi b$ became slower and $2\pi c$ became faster compared to the full activation (pCa \leq 5.0) (Fig. 8). Rate constant $2\pi c$ became significantly larger than $2\pi b$ (up to 20 times). These changes are similar to those found in skeletal muscle fibers (55). The ~75% increase in $2\pi b$ and the ~25% decrease in $2\pi c$ in DAD compared to WT (Fig. 4 A) demonstrate the profound effect of DAD on the elementary steps of the cross-bridge cycle.

Electron microscopy studies (56,57) suggest that cMyBP-C is present only in the central 53% of the thick filaments (9 stripes out of the total 17: C-zone), and absent in the remaining 47% (8 stripes out of 17) (57). Therefore, cross-bridges can be considered as a mixture of two kinds-one-half (53%: C-zone) with cMyBP-C, and the other half (47%) without cMyBP-C-arranged in parallel. The role of cMyBP-C in modulating cardiac contractility appears to be localized in the C-zone as shown by in vitro motility assay (58). Our results demonstrate that isometric tension and stiffness are significantly reduced (by ~48%) in the DAD fibers compared to the WT fibers (Fig. 2). Assuming that the cross-bridges outside of the C zone (~47%) function normally, the cross-bridges containing cMyBP-C DAD protein (~53%) must be almost inactive, or their function is extremely diminished to $\sim 5\%$ (= 53% - 48%). However, it is also true that the cross-bridges that form with the DAD protein must be partially effective, as evidenced by the altered kinetics of the rate constants $2\pi b$ and $2\pi c$ in the 8Pi solution (Fig. 4 A). Therefore, it is possible that the DAD protein influences the cross-bridges beyond the C-zone.

In mouse cardiac muscles, PKA phosphorylates all three sites (Ser²⁷³, Ser²⁸², and Ser³⁰²), whereas PKC phosphorylates only two sites (Ser²⁷³ and Ser³⁰²) (17). The significantly compromised tension-generating capacity and the reduced number of cross-bridges in DAD mice, but not in ADA and SAS observed in this study (Fig. 2), lead us to propose that PKC-mediated phosphorylation of cMyBP-C at Ser²⁷³ and Ser³⁰² either serves as a compensatory mechanism that protects cardiomyocytes from overexertion of the already dysfunctional heart, or results in maladaptive contraction that ultimately compromises cardiac function. Our biophysical data from DAD mice are consistent with the cardiac hypertrophy, myocyte disarray, and interstitial fibrosis previously observed (17). A recent study investigated three additional cMyBP-C transgenic lines in mice: DAA, AAD, and SDS, together with AAA, ADA, and t/t (22). It was found that DAA and AAD mice were significantly affected, showing more severe cardiac pathology than t/t mice, with left ventricle dilation, interstitial fibrosis, irregular cardiac rhythm, and sudden cardiac death (22). Also, the sarcomere structure was seriously damaged in DAD fibers (17). When these data and our results are combined, we can conclude that a nonphosphorylatable Ala²⁸² with phosphomimetic Asp²⁷³ and/or Asp³⁰² (in AAD, DAA, and DAD) are detrimental to cardiac structure and function. We further conclude that the combination of the phosphorylation pattern is critical in the function of cMyBP-C.

An increase in active tension was previously found after partial (60-70%) extraction of cMyBP-C (59); some investigators have reported accelerated cross-bridge kinetics and increased power output in t/t compared to WT mice (60-63). We are not certain about the reasons for these discrepancies, but several differences in the experimental conditions may have contributed. First, differences in the methods used and the parameters investigated may account for the differences. In previous studies, the skinned fibers were produced by homogenization of the thawed frozen heart (60,61,63), but we generated skinned fibers from fresh hearts. It is also worth noting that the accelerated cross-bridge kinetics reported in previous studies were found to be significant in partial activation (50% or 25% of maximal force) in the absence of added Pi and in the 180 mM IS solution (60-63), whereas our standard solution contained 8 mM Pi and 200 mM IS. In addition, previous studies measured the rate constants of stretch-induced force decay (k_{rel}) and delayed force development (k_{df}) in stretch activation (60,61), and the rate constant of force redevelopment (k_{tr}) through large release-restretch experiments (62,63). We measured the rate constants of delayed tension, $2\pi b$, and of fast tension recovery, $2\pi c$, through sinusoidal analysis under near-isometric conditions.

Compared to the DAD group, the ADA and SAS groups exhibited little change from the WT. In contrast to DAD, 100% phosphomimetic Asp²⁸² within the context of nonphosphorylation at positions 273 and 302 (ADA) appears to have little effect other than a decreased stiffness in the OPi solution (Fig. 2 *B*). Because Ser^{282} plays an important role in the phosphorylation of Ser^{273} and Ser^{302} (6,17), it is reasonable to postulate that in the SAS mutation (with phosphorylation-deficient Ala²⁸²), the phosphorylation of Ser²⁷³ and Ser³⁰² is significantly reduced. The smaller changes in SAS compared to DAD suggest that when residue 282 cannot be phosphorylated, 100% phosphomimetic Asp²⁷³ and Asp³⁰² (DAD) may be detrimental for cardiac function. The SAS actually exhibited somewhat opposite effects to DAD in active tension, and in magnitudes B and H; these changes became significant when the data of one female SAS mouse was excluded (Figs. S2 and S4). This does not mean, however, that these parameters are lower in female SAS mice, because there were not adequate data in females.

Several studies found that a 13–19% increase in β -MHC expression (and a concomitant decrease in α -MHC expression) occurs in t/t mouse hearts (10,46,61). It has also been observed that a 100% shift from α - to β -MHC reduces $2\pi c$ by ~75% without changing the isometric tension (45). Thus, our observations on $2\pi c$ may be confounded by two factors: a mutation in cMyBP-C and an MHC isoform shift. For this reason, we studied the MHC isoform content (Fig. 9). We found that our conclusions on DAD and ADA are not significantly affected by the isoform shift, because

the β -MHC contents of DAD and ADA were small and did not differ significantly from that of WT (Fig. 9 B). In the case of t/t, there was a significant (43 \pm 4%) decrease of the α -MHC content (Fig. 9 B), which could account for much of the observed decrease in $2\pi c$ (42% in Fig. 4 A and 33% in Fig. 5 A), because $75\% \times 43\% = 32\%$. The absence of cMyBP-C causes a problem in cardiac muscles, as demonstrated by disorganized and damaged sarcomere structures in t/t mice (17) and disordered thick filaments with varying myosin heads extending from the backbone of the thick filament, as well as an ~30% reduction in maximal force after partial (29 \pm 3%) extraction of cMyBP-C (64). The reduced $2\pi c$ in t/t mouse hearts (Fig. 4 A) is consistent with the diminished left ventricle function based on in vivo measurements (61,65,66), depressed cross-bridge kinetics (67), slower shortening velocity, and reduced power output (68,69). All of these lines of evidence support the hypothesis that the absence of cMyBP-C results in impaired myocardial contractility.

In the case of SAS, there was a significant $(26 \pm 3\%)$ decrease in α -MHC contents (Fig. 9 B), which predicts a 20% (75% \times 26%) decrease in $2\pi c$. Because the actual decrease in $2\pi c$ was ~8% (Figs. 4 A and 5 A), the effect of mutation in SAS may be to increase this rate constant by ~12% (20% - 8%), and the two antagonizing effects may compensate each other. Although there are no reports on the effects of MHC isoforms on $2\pi b$, it is possible that the isoform content does not affect this parameter, given that little change was observed in $2\pi b$ of t/t in two different solutions (Figs. 4 A and 5 A). The increase in $2\pi b$ in DAD must be due to this mutant form of cMyBP-C, because there is no significant isoform shift (Fig. 9 B). The involvement of isoform shifting does not modify our conclusions on tension, stiffness, or magnitude parameters, because tension is not sensitive to the shifting (45), and parameters Y_{∞} , B, C, and H are usually scaled with tension.

CONCLUSION

This study investigated cardiac fiber contractility and cross-bridge kinetics in transgenic mouse models carrying cMyBP-C mutations that affect three different phosphorylation sites in its M-domain. Our investigation revealed that a lack of cMyBP-C diminishes the rate constant of fast tension recovery, $2\pi c$, owing to the isoform shifting of MHC. Phosphomimetic Asp²⁷³ and Asp³⁰² with a nonphosphorylatable Ala substituted for residue 282 (DAD) significantly diminishes active tension and altered cross-bridge kinetics by increasing the rate constant of delayed tension $(2\pi b)$, and decreasing the rate constant of fast tension recovery $(2\pi c)$. However, neither the single substitution of a nonphosphorylatable residue 282 (SAS), nor the combination of nonphosphorylatable residues at 273 and 302 and the presence of a 100% phosphomimetic Asp²⁸² (ADA), causes significant changes in fiber tension and cross-bridge kinetics. In conclusion, cMyBP-C influences cardiac contractility and cross-bridge kinetics in a manner specific to the phosphorylation sites, and the nonphosphorylatable residue 282 combined with phosphomimetic residues 273 and/or 302 is detrimental to cardiomyocytes.

SUPPORTING MATERIAL

Five figures and one table are available at http://www.biophysj.org/ biophysj/supplemental/S0006-3495(14)00127-1.

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Phosphorylation of cMyBP-C affects contractile mechanisms in a site-specific manner

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Due to the possibility of a gender difference in these experiments, we eliminated the data from females (one from WT and one from the SAS group). Data was averaged only in males and plotted in this section. The figure numbers (S2, S4, etc.) correspond to those in the main text (Figs. 2, 4, etc.). Figure numbers S1 and S3 are absent.



FIGURE S2 Isometric tension (T_{max} in Eq. 2: left ordinate) and stiffness (S_{max} in Eq. 3: right ordinate) in the 8Pi solution (in *A*) and the 0Pi solution (in *B*). The 8Pi solution contained 8 mM Pi and its ionic strength (IS) was 200 mM, whereas the 0Pi solution contained no Pi and its IS was 150 mM. **p < 0.01 and *p < 0.05 compared to WT. The number of experiments (*n*) is in (*A*), 23 for WT, 28 for t/t, 32 for ADA, 31 for DAD, and 14 for SAS; In (*B*), n = 9 for WT, 14 for t/t, ADA, and DAD, and n = 5 for SAS.



FIGURE S4 **Parameters of exponential processes in the 8Pi solution.** (*A*) The apparent rate constants $2\pi b$ (left ordinate) and $2\pi c$ (right ordinate). (*B*) the magnitudes of processes B (left ordinate) and C (right ordinate). (*C*) magnitude *H*. Error bars represent SE of the averages of 14-38 experiments carried out in each group (for the number of mice, see Table 1). n = 23 for WT, 38 for t/t, 32 for ADA, 31 for DAD, and 14 for SAS. **p < 0.01 and *p < 0.05 compared to WT.



FIGURE S5 **Parameters of exponential processes in the 0Pi solution.** (*A*) The apparent rate constants $2\pi b$ (left ordinate) and $2\pi c$ (right ordinate). (*B*) the magnitudes of processes B (left ordinate) and C (right ordinate). (*C*) magnitude *H*. n = 9 for WT, 14 for t/t, ADA, and DAD, and n = 5 for SAS. **p < 0.01 and *p < 0.05 compared to WT.



FIGURE S6 The effect of $[Ca^{2+}]$ in the 8Pi solution in five groups of mice. (*B*) pCa₅₀ (left ordinate) and n_H (right ordinate), as deduced by fitting the individual data of pCa-tension relationship to Eq. 2, and averaged. (*D*) pCa_{50S} (left ordinate) and n_{HS} (right ordinate), as deduced by fitting the data of pCa-stiffness relationship to Eq. 3, and averaged. In (*B*) and (*D*), no significant difference was found in any of the parameters measured. n = 9 for WT, 13 for t/t, 14 for ADA, 18 for DAD, and 5 for SAS.



FIGURE S7 The effect of $[Ca^{2+}]$ in the 0Pi solution in five groups of mice. The same nomenclature and conventions are used as in Fig. S6. n = 9 for WT, 14 for t/t, ADA, and DAD, and n = 5 for SAS. ** $p \le 0.01$, *0.01 , and (*) <math>0.05 compared to WT.

Study	Parameter	WT	t/t	ADA	DAD	SAS
	$2\pi b (s^{-1})$	40.0 ± 1.4	36.0 ± 2.6	43.7 ± 3.2	$67.5 \pm 2.5 **$	$55.6 \pm 6.2 $ **
	$2\pi c (s^{-1})$	198 ± 10	$115 \pm 5^{**}$	197 ± 8	$146 \pm 10^{**}$	183 ± 16
8Pi,	Magnitude B (kPa)	221 ± 17	$329 \pm 42*$	221 ± 30	255 ± 39	$365 \pm 64*$
Standard	Magnitude C (kPa)	760 ± 68	711 ± 68	661 ± 59	$486 \pm 79*$	956 ± 111
activation	Magnitude H (kPa)	116 ± 10	111 ± 10	93 ± 5	$84 \pm 11*$	137 ± 12
	Tension (kPa)	15.4 ± 1.2	13.6 ± 1.0	14.4 ± 1.4	8.2 ± 1.1 **	18.1 ± 2.2
	Stiffness (kPa)	655 ± 60	$489 \pm 38*$	534 ± 50	$335 \pm 63 **$	729 ± 93
	$2\pi b (s^{-1})$	8.3 ± 1.4	8.5 ± 1.4	6.5 ± 1.0	11.8 ± 1.6	10.4 ± 1.3
	$2\pi c (s^{-1})$	70.5 ± 5.6	$37.9 \pm 4.2 **$	64.5 ± 4.8	59.4 ± 4.7	64.6 ± 6.6
0Pi.	Magnitude B (kPa)	30.1 ± 16.5	42.8 ± 17.3	33.2 ± 11.9	5.5 ± 3.7	49.7 ± 38.0
Standard	Magnitude C (kPa)	384 ± 63	349 ± 68	267 ± 37	218 ± 98	378 ± 78
activation	Magnitude H (kPa)	373 ± 48	420 ± 36	312 ± 27	211 ± 25*	429 ± 66
	Tension (kPa)	24.3 ± 2.5	20.0 ± 1.6	19.7 ± 1.7	12.4 ± 2.2 **	24.0 ± 3.3
	Stiffness (kPa)	761 ± 101	645 ± 64	$558 \pm 52*$	$423 \pm 115*$	757 ± 109
8Pi, pCa-	pCa_{50} (Ca ²⁺ sensitivity)	5.562 ± 0.011	5.555 ± 0.010	5.586 ± 0.012	5.548 ± 0.012	5.548 ± 0.015
tension	$n_{\rm H}$ (cooperativity)	5.7 ± 0.3	5.1 ± 0.3	6.3 ± 0.3	6.2 ± 0.3	5.9 ± 0.4
8Pi, pCa-	pCa_{50S} (Ca^{2+} sensitivity)	5.565 ± 0.012	5.566 ± 0.009	5.595 ± 0.014	5.565 ± 0.012	5.563 ± 0.015
stiffness	$n_{\rm HS}$ (cooperativity)	6.2 ± 0.3	5.4 ± 0.3	6.7 ± 0.3	6.8 ± 0.3	5.9 ± 0.5
0Pi. pCa-	pCa_{50}	5.726 ± 0.030	5.768 ± 0.017	5.712 ± 0.038	$5.640 \pm 0.030^{(*)}$	5.714 ± 0.023
tension	$n_{\rm H}$	2.3 ± 0.2	$2.8 \pm 0.2^{(*)}$	2.6 ± 0.2	3.1 ± 0.3*	2.7 ± 0.2
0Pi. pCa-	pCa ₅₀₈	5.760 ± 0.041	5.786 ± 0.016	5.715 ± 0.037	$5.659 \pm 0.023^{*}$	5.728 ± 0.022
stiffness	n _{HS}	2.4 ± 0.2	2.7 ± 0.2	2.8 ± 0.3	3.5 ± 0.4**	2.6 ± 0.2

TABLE S1. The biophysical parameters (**p < 0.01 and *p < 0.05 compared to WT) including two female mice.

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