Genetically Encoded Biosensors Reveal PKA Hyperphosphorylation on the Myofilaments in Rabbit Heart Failure

Federica Barbagallo,* Bing Xu,* Gopireddy R. Reddy, Tony West, Qingtong Wang, Qin Fu, Minghui Li, Qian Shi, Kenneth S. Ginsburg, William Ferrier, Andrea M. Isidori, Fabio Naro, Hemal H. Patel, Julie Bossuyt, Donald Bers, Yang K. Xiang

Rationale: In heart failure, myofilament proteins display abnormal phosphorylation, which contributes to contractile dysfunction. The mechanisms underlying the dysregulation of protein phosphorylation on myofilaments is not clear.

Objective: This study aims to understand the mechanisms underlying altered phosphorylation of myofilament proteins in heart failure.

Methods and Results: We generate a novel genetically encoded protein kinase A (PKA) biosensor anchored onto the myofilaments in rabbit cardiac myocytes to examine PKA activity at the myofilaments in responses to adrenergic stimulation. We show that PKA activity is shifted from the sarcolemma to the myofilaments in hypertrophic failing rabbit myocytes. In particular, the increased PKA activity on the myofilaments is because of an enhanced β2 adrenergic receptor signal selectively directed to the myofilaments together with a reduced phosphodiesterase activity associated with the myofibrils. Mechanistically, the enhanced PKA activity on the myofilaments is associated with downregulation of caveolin-3 in the hypertrophic failing rabbit myocytes. Reintroduction of caveolin-3 in the failing myocytes is able to normalize the distribution of β2 adrenergic receptor signal by preventing PKA signal access to the myofilaments and to restore contractile response to adrenergic stimulation.

Conclusions: In hypertrophic rabbit myocytes, selectively enhanced β2 adrenergic receptor signaling toward the myofilaments contributes to elevated PKA activity and PKA phosphorylation of myofilament proteins. Reintroduction of caveolin-3 is able to confine β2 adrenergic receptor signaling and restore myocyte contractility in response to β adrenergic stimulation. (Circ Res. 2016;119:931-943. DOI: 10.1161/CIRCRESAHA.116.308964.)

Key Words: adrenergic receptor ■ heart failure ■ myofibrils ■ phosphorylation ■ protein kinase A

In animal hearts, elevated sympathetic activity stimulates the β-adrenergic signal to promote inotropy and lusitropy via the concerted protein kinase A (PKA) phosphorylation of substrates at multiple subcellular locations. PKA is spatially and temporally regulated through a set of A-kinase anchoring protein complexes.1 As a result, the precise PKA phosphorylation of myofilament proteins, such as cardiac troponin I (TnI) and myosin-binding protein C, and Ca2+-handling proteins, such as phospholamban (PLB) and ryanodine receptor on the sarcoplasmic reticulum (SR), is necessary to co-ordinate positive inotropic and lusitropic cardiac effects.2,3

Chronic adrenergic signaling also promotes structural and functional remodeling in the myocardium, which contributes to cardiac hypertrophy and eventually heart failure in a variety of clinical conditions. A desensitized cardiac β-adrenergic receptor (βAR) signaling pathway,4 a hallmark of the failing heart, is associated with a reduced number of β1AR- but not β2AR-binding sites at the plasma membrane (PM). As a consequence, diminished βAR signaling may lead to lower PKA-mediated protein phosphorylation after adrenergic stimulation.5 The disturbed βAR signaling and reduced PKA phosphorylation result in blunted contractile response after administration of βAR agonists.6 Interestingly, recent

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Nonstandard Abbreviations and Acronyms

<table>
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<tr>
<th>Acronym</th>
<th>Description</th>
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<tr>
<td>AKAR</td>
<td>A-kinase activity reporter</td>
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<tr>
<td>βAR</td>
<td>β adrenergic receptor</td>
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<tr>
<td>CS3D</td>
<td>scaffolding domain of caveolin-3</td>
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<tr>
<td>FRET</td>
<td>fluorescence resonant energy transfer</td>
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<tr>
<td>HF</td>
<td>heart failure</td>
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<tr>
<td>ISO</td>
<td>isoproterenol</td>
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<tr>
<td>PDE</td>
<td>phosphodiesterase</td>
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<tr>
<td>PKA</td>
<td>protein kinase A</td>
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<tr>
<td>PLB</td>
<td>phospholamban</td>
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<tr>
<td>PM</td>
<td>plasma membrane</td>
</tr>
<tr>
<td>SR</td>
<td>sarcoplasmic reticulum</td>
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<tr>
<td>TnI</td>
<td>troponin I</td>
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Progress suggests that βAR can continuously signal both at the cell surface and after catecholamine-induced endocytosis. These βAR signals may play a role in adaptive cardiac hypertrophic remodeling and promote compensatory contractility in the heart. Until now, it remains poorly understood how remodeling of βAR signaling affects PKA activity in different subcellular locations for PKA-dependent phosphorylation of substrates in failing myocytes.

Recent efforts have been put forward to understand β-adrenergic signaling at the sarcolemma and the SR because of their essential roles in calcium handling and arrhythmia in cardiac diseases. By using genetically encoded biosensors, studies show that adrenergic-induced PKA activity at the SR can be selectively inhibited by chronic insulin and the proinflammatory cytokines show that adrenergic-induced PKA activity at the SR can be selectively inhibited by chronic insulin.

Statistical Analysis

Statistical analysis was performed using GraphPad Prism 6 software (La Jolla, CA). Expanded methods can be found online (Online Data Supplement).

Results

Novel Biosensors Reveal Highly Localized βAR-Induced PKA Activity That Does not Reach the Myofilaments in Healthy Rabbit Cardiac Myocytes

To analyze the dynamics of PKA activity on the myofilaments, we took a novel strategy to anchor FRET-based A-kinase activity reporter 3 (AKAR3) onto the myofilaments by linking AKAR3 to the C terminus of troponin T (myofilament-AKAR3; Figure 1A). Myofilament-AKAR3 was used together with 2 other targeted PKA biosensors developed previously: PM-AKAR3 is linked to a PM-targeting sequence by combined aortic insufficiency and stenosis as previously described. These data indicate that the molecular pathways are preserved on the expression of biosensors, and the targeted probes are suitable for analysis of PKA activity in the microdomains of adult cardiac myocytes.

Fluorescence Resonance Energy Transfer Measurements

Images were acquired using a Leica DMi300B inverted fluorescence microscope (Leica Biosystems, Buffalo Grove, IL) with a 40× oil-immersion objective lens and a charge-coupled device camera controlled by Metafluor software (Molecular Devices, Sunnyvale, CA). Fluorescence images for fluorescence resonance energy transfer (FRET) analysis were obtained as described previously.

Statistical Analysis

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Methods

Experimental Animals, Cardiac Myocyte Isolation, and Culture

The animal care and experimental protocols followed US National Institutes of Health guidelines and were approved by the Institutional Animal Care and Use Committees of the University of California at Davis. Heart failure (HF) was induced in New Zealand White rabbits by combined aortic insufficiency and stenosis as previously described.

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We further characterized the AKAR3 biosensors in response to a set of incremental doses of β-adrenergic agonist isoproterenol in adult rabbit myocytes. Stimulation of myocytes with isoproterenol induces dose-dependent increases in FRET ratio of the PM-, SR-, and myofilament-anchored AKAR3 biosensors, and the maximal responses are higher at the PM than those at the SR and myofilament (Figure 1D; Online Figure 1). The responses of PM-AKAR3 are also more sensitive to isoproterenol stimulation than those of SR-AKAR3 and myofilament-AKAR3 (Figure 1D and 1E; Online Figure 1). These data are consistent with previous studies showing that PKA phosphorylation of substrates at the PM is β-adrenergic signal and restores contractile response to adrenergic stimulation.
Figure 1. Generation of cardiac myofilament-targeted fluorescence resonant energy transfer (FRET) biosensors. A, Schematic representation of subcellular localized protein kinase A (PKA) A-kinase activity reporter 3 (AKAR3) biosensors. Yellow (YFP) and cyan (CFP) fluorescent proteins flank a PKA substrate and a forkhead-associated (FHA) domain that recognizes the phosphorylated PKA substrate. AKAR3 is linked to a Kras-derived sequence for plasma membrane (PM-AKAR3) localization, to a phospholamban (PLB)-derived sequence for sarcoplasmic reticulum (SR-AKAR3) localization, and to troponin T for myofilament (MF-AKAR3) localization. B, Representative confocal images of biosensor (green) expressed in young rabbit cardiac myocytes. Cells are immunostained with subcellular specific markers (red) for the SR (ryanodine receptor [RyR]), the PM (wheat germ agglutinin [WGA]), and (Continued)
more sensitive to lower doses of isoproterenol stimulation than those at the intracellular compartments. The isoproterenol-induced maximal responses are not because of saturation of individual biosensors because cotreatment of myocytes with forskolin and IBMX (3-isobutyl-1-methylxanthine) induces higher increases in FRET ratios of biosensors than those induced by isoproterenol, respectively (Figure 1F). Notably, the isoproterenol-induced maximal responses in these localized biosensors are equivalent when normalized by those induced by cotreatment of forskolin and IBMX (Figure 1G), suggesting that these biosensors are functionally comparable in response to adrenergic stimulation in rabbit myocytes.

To analyze the alteration of local PKA dynamics at the myofilaments in failing heart, we used cardiac myocytes isolated from rabbits with hypertrophic HF induced by combined aortic insufficiency and stenosis. Myocytes from age-matched rabbits with hypertrophic HF induced by combined aortic insufficiency and stenosis. Myocytes from age-matched SHAM myocytes, stimulation of the βAR with a saturated dose of isoproterenol (100 nmol/L) leads to an increase of PKA activity, which is completely blocked by βAR selective antagonist CGP20712A but not by βAR selective antagonist ICI118551 (Figure 2A). Accordingly, stimulation of SHAM myocytes with isoproterenol significantly promotes PKA phosphorylation of TnI at serine 23/24 in response to isoproterenol (ISO) stimulation (1 or 100 nmol/L) in young rabbit myocytes infected with MF-AKAR3 probes. Young rabbit myocytes expressing PM-AKAR3, SR-AKAR3, and MF-AKAR3 are stimulated with a set of incremental doses of ISO. Time courses show AKAR3 FRET responses after stimulation with ISO. Normalized isoproterenol-induced, dose-response curves of AKAR3 biosensors (EC50 PM-AKAR3 at 2.16×10⁻⁹ mol/L, SR-AKAR3 at 2.57×10⁻⁹ mol/L, and MF-AKAR3 at 3.89×10⁻⁹ mol/L). F. Maximal increases in AKAR3 FRET ratio after stimulation with ISO (1 μmol/L) or after cotreatment with forskolin (10 μmol/L) and IBMX (3-isobutyl-1-methylxanthine; 100 μmol/L). G. Normalized maximal FRET responses of individual AKAR3 biosensors against the increases induced by cotreatment with forskolin and IBMX, respectively. SERCA indicates sarco/endoplasmic reticulum Ca²⁺-ATPase.

Redistribution of Phosphodiesterase Activity Contributes to Elevated PKA Activity on the Myofilaments in Failing Heart

Recent studies have also revealed essential roles of alteration in phosphodiesterase (PDE) expression in remodeling of adrenergic signal in HF myocytes. We next studied the involvement of various cAMP PDEs on the modification of local PKA activity in HF myocytes. In SHAM myocytes, inhibition of PDE3 with cilostamide induces a strong increase in PKA activity at the myofilaments and the SR but not at the PM (Figure 5A through 5C). In comparison, inhibition of PDE4 with rolipram induces a strong response in PKA activity at the PM and, to a lesser extent, at the myofilaments and the SR (Figure 5D through 5F). Consistent with these observations, PDE3 displays colocalization with the myofilament-binding protein actin, whereas both PDE4D and PDE4B display colocalization with the PM marker caveolin-3 (Online Figure II). In HF myocytes, the myofilament-associated PDE3 activity is decreased (Figure 5A and 5G), whereas the PM-associated PDE3 activity is increased relative to SHAM controls (Figure 5C and 5I), suggesting a shifted distribution of PDE3 activity from the myofilaments to the sarcolemma. In comparison, the PM-associated PDE4 activity in HF myocytes is decreased relative to SHAM controls (Figure 5F and 5I). Meanwhile, inhibition of PDE4 induces a small increase in PKA activity at the SR in SHAM myocytes, which is enhanced in HF cells (Figure 5E and 5H). However, the SR-associated PDE3 activity is not significantly altered in HF cells (Figure 5B and 5H). In addition, inhibition
of PDE2 with erythro-9-(2-hydroxy-3-nonyl)adenine induces a small increase in PKA activity at the PM and myofilaments in SHAM myocytes; the erythro-9-(2-hydroxy-3-nonyl)adenine--induced response is increased at the PM in HF cells, whereas the erythro-9-(2-hydroxy-3-nonyl)adenine--induced response is reduced at the myofilaments in HF cells (Online Figure III). Although the PDE2 activity at the SR is minimal in SHAM myocytes, it is increased in HF cells (Online Figure III).

Together, in healthy rabbit myocytes, the PKA activity at the PM is under predominant control of PDE4, whereas PDE3 hydrolyzes preferentially a pool of cAMP in the vicinity of the myofilaments (Figure 5). In HF rabbit myocytes, the presence of PDE4 activity at the PM is reduced, whereas the activity of PDE3 and PDE2 become more prominent. In contrast, the activity of PDE3 and PDE2 is decreased at the myofilaments (Figure 5; Online Figure III). These data...
indicate dysregulation of these hydrolytic enzymes in HF myocytes. Accordingly, Western blotting analysis shows a reduction of PDE3 and an upregulation of PDE4 expression in left ventricles after HF (Figure 6A). Moreover, PDE4 is shifted to internal compartments, whereas PDE3 is enriched at the sarcolemma region in HF myocytes relative to SHAM controls (Online Figure IV). The membrane-enriched PDE3 does not colocalize with β2AR in HF myocytes (Online Figure IV). Therefore, in HF myocytes, PDEs may not be effectively coupled to adrenergic receptors. Indeed, although inhibition of PDE4 and PDE3 potentiates adrenergic stimulation–induced PKA phosphorylation of TnI in SHAM myocytes, the effects of these PDE inhibitors are diminished in the HF cells (Online Figure V).
Here, we reveal an 80% reduction of protein expression of ion channels at the PM. Previous studies have demonstrated that the expression of caveolin-3 is decreased in different mouse and human cardiac diseases including heart failure and hypertrophy. Loss of caveolin-3 is associated with redistribution of β2AR signal on the PM in failing myocytes.

In a separate set of experiments, we applied PKA inhibitors in the FRET measurement to assess the baseline subcellular PKA activity in HF and SHAM myocytes (Online Figure VI). In SHAM myocytes, we detect a significant reduction of PKA activity by H89 at the myofilaments and the SR but minimal change in PKA activity at the PM (Online Figure VI). In HF myocytes, the H89-sensitive PKA activity at the PM and the SR does not change (Online Figure VI). However, the H89-sensitive PKA activity at the myofilaments is increased in HF cells, indicating an elevation of baseline PKA activity associated with myofilaments in HF myocytes (Online Figure VI).

Reintroduction of Caveolin-3 Restores Distribution of βAR-Induced PKA Signal, PKA Phosphorylation, and Contractility in Failing Myocytes

β2AR is enriched in the caveolae of cardiac myocytes, which confines the receptor action to the local PM domains. Disruption of caveolae leads to enhanced β2AR stimulation of ion channels at the PM. Previous studies have demonstrated that the expression of caveolin-3 is decreased in different mouse and human cardiac diseases including heart failure and hypertrophy. Loss of caveolin-3 is associated with redistribution of β2AR signal on the PM in failing myocytes. Here, we reveal an 80% reduction of protein expression of caveolin-3 in the failing rabbit myocytes. This reduction is associated with an increase of PKA phosphorylation of TnI (Figure 6A). Caveolin-3 has a scaffolding domain (C3SD) that is critical for the structure and function of caveolae. A membrane-permeable peptide corresponding to C3SD modulates β2AR regulation of L-type calcium channel. We applied the membrane-permeable C3SD peptide to healthy rabbit myocytes to examine whether inhibition of caveolin-3 promotes β2AR signal to the myofilaments. Treatment with C3SD significantly enhances β2AR-induced PKA activity at the myofilaments but not at the SR (Figure 6B and 6C). In contrast, a scrambled peptide does not enhance β2AR-induced PKA activity at the myofilament. Consistent with the FRET-based measurement of PKA activity, C3SD significantly promotes β2AR-induced PKA phosphorylation of TnI at the myofilaments (Figure 6D). These data suggest that caveolin-3 plays an essential role in preventing the β2AR from signaling to the myofilaments; therefore, loss of caveolin-3 in HF myocytes may permit β2AR to signal to the myofilaments to increase PKA activity and substrate phosphorylation.

We sought to restore the expression of caveolin-3 in hypertrophic failing myocytes and to evaluate βAR signaling and PKA activity on the myofilaments. Reintroduction of caveolin-3 partially restores the distribution of PDEs (Online Figure VI).
Figure IV) and their activity in HF myocytes, including an increase in PDE4 activity at the PM and increases in PDE3 and PDE2 activity at the myofilaments (Online Figures III and VII). Moreover, reintroduction of caveolin-3 partially normalizes the distribution of β2AR (Online Figure VIII) and PKA activity at the subcellular compartments, including increases in PKA activity at the PM and decreases in PKA activity at the SR and myofilaments after stimulation with isoproterenol (Figure 7A through 7C). Although the β2AR-induced PKA activity is increased at the PM, the β2AR-induced PKA activity is decreased throughout the myocytes. Notably, after reintroduction of caveolin-3, the β2AR-induced PKA activity does not reach the myofilaments; instead, it is confined to the PM and the SR (Figure 7A through 7C). Accordingly, the phosphorylation of TnI under β2AR stimulation is significantly diminished after overexpression of caveolin-3 (Figure 7D). Finally, overexpression of caveolin-3 restores β-adrenergic–induced contractile shortening response in hypertrophic failing rabbit myocytes (Figure 7E).

**Discussion**

The cardiac PKA activity is fine-tuned to ensure co-ordinated phosphorylation of substrates in different compartments for excitation–contraction coupling at baseline and after adrenergic stimulation. In this study, we have revealed a structural–functional remodeling in hypertrophic failing rabbit myocytes that links the redistribution of βAR signal to enhanced PKA phosphorylation of myofilament proteins (Figure 8). Although the β2AR-induced signal is restricted along the sarcolemma and the SR in healthy myocytes, an augmented β2AR signal associated with loss of caveolin-3 is selectively directed to the myofilaments in hypertrophic failing cells. This augmented β2AR signal, together with reduced PDE activity associated with myofibrils, promotes strong PKA activity and PKA phosphorylation of the myofilament protein TnI in hypertrophic failing myocytes. Moreover, reintroduction of caveolin-3 is able to normalize the distribution of adrenergic signaling and restores contractile response to adrenergic stimulation in hypertrophic failing rabbit myocytes.

A tonic PKA activity is maintained by adenylyl cyclase–dependent cAMP production and PDE-dependent cAMP hydrolysis.19,20 Our results show that PDE4 is the major enzyme that controls local PKA activity along the sarcolemma, whereas PDE3 mediates primarily cAMP hydrolysis on the myofilaments (Figure 5). In agreement, both PDE4D and PDE4B display overlap with the membrane marker caveolin-3 (Online Figure II), consistent with a recent report showing that both enzymes have a binding motif to caveolin-3;27 loss of caveolin-3 may lead to loss of PDE4 activity at the sarcolemma. In contrast, PDE3 displays a colocalization with myofibrils (Online Figure II), supporting its primary role in controlling local cAMP and PKA activity there (Figure 5). Meanwhile, we observe a high tonic PKA activity on the myofilaments, which is consistent with a relatively high PKA phosphorylation of myosin-binding protein C and TnI in healthy cardiac tissues.28

Although the PKA activity is usually depressed in the late stage of human heart failure,29,30 there are reports...
showing preserved or elevated PKA phosphorylation on TnI and PLB. It is not completely understood how the cellular PKA activity in myocytes evolves during the course of heart failure development. Here, we observe a redistribution of PKA activity in hypertrophic failing myocytes, including an increased PKA activity at the myofilaments. In agreement, a recent report also shows that PKA phosphorylation of myofilament proteins is elevated in the spontaneous hypertensive rat model. It is therefore possible that PKA activity is redistributed as part of the remodeling of βAR signaling during the compensatory stage before a general depression of PKA activity at the late stage of heart failure. In a classic view, βARs are desensitized in HF due in part to the reduced β1AR density at the cell surface, rendering the heart unable to respond to catecholamine stimulation. However, recent progress suggests that βARs can continuously signal at the cell surface and after catecholamine-induced endocytosis. Studies show that βARs are accumulated at the endosome under chronic elevation of sympathetic activity in HF. These βARs may signal from endosomes, offering a potential mechanism on redistribution of PKA activity from the PM to the myofilaments in failing myocytes (Figures 2 and 4). Further analysis reveals

Figure 6. Inhibition of caveolin-3 promotes adrenergic signaling on the myofilaments in heart failure (HF) cardiac myocytes. A, Immunoblots show expression of caveolin-3, PDE3A, PDE4, troponin I (TnI), phospholamban (PLB), and phospholemman (PLM), as well as pTnI at serine 23/24, pPLB at serine 16, and pPLM at serine 68 in SHAM and HF heart lysates. Bar graphs represent mean±SEM. N=3; *P<0.05, **P<0.01, and ***P<0.001 by Student t test. B and C, Healthy young rabbit myocytes expressing SR-A-kinase activity reporter 3 (AKAR3) or MF-AKAR3 are stimulated with 100 nmol/L isoproterenol (ISO) and 300 nmol/L CGP20712A after incubation with 1 μmol/L C3SD or scrambled peptide for 1 h. Time courses of changes in AKAR3 fluorescence resonant energy transfer (FRET) ratio are shown, and the maximal increases in FRET ratio are plotted in bar graph. ***P<0.001 by 1-way ANOVA followed by post hoc Bonferroni test. D, Healthy young rabbit myocytes are stimulated with 100 nmol/L ISO in the presence of 1 μmol/L C3SD or scrambled peptide. PKA phosphorylation of troponin I at serine 23/24 is detected in Western blots and quantified in bar graph. N=3; **P<0.01 by 1-way ANOVA followed by post hoc Bonferroni test.
that the diminished β2AR signaling contributes to lower PKA activity at the sarcolemma (Figure 4), which is consistent with previous observations that β2AR density is downregulated in this rabbit HF model. In comparison, β1AR density is not reduced; rather an augmented β1AR signal may be preferentially directed to the myofilaments. Moreover, the shifted PKA activity can be further exacerbated by relocation of PDE3 and PDE2 activity away from the myofilaments to the sarcolemma (Figure 5; Online Figure III). Together, these observations suggest that the βAR signal undergoes relocation intracellularly during the development of heart failure. Further studies will likely yield novel information on βAR-PKA signal remodeling in subcellular compartments in models of cardiac disease, which may offer insight into cardiac hypertrophic remodeling and compensatory contractility in the heart.

Detubulation and loss of caveolae are two prominent features during cardiac structural remodeling; these alterations can in turn promote signal remodeling in cardiac myocytes. Although the β2AR is confined in the tubular region in healthy myocytes, chemical disruption of caveolae with a cholesterol-depleting agent, methyl-β-cyclodextrin, results in >60% increase in β2AR-dependent PKA phosphorylation of PLB and TnI, as well as lusitropic responses in adult myocytes. In addition, a membrane-permeable peptide representing the C3SD leads to reduction of β2AR-dependent PKA phosphorylation of L-type calcium channel at the PM. Detubulation and loss of caveolae in hypertrophic failing myocytes isolated from a mouse model yield broad distribution of β2AR signaling along the PM crest. The loss of close proximity of β2AR to caveolin-3–associated PDE4 can also permit a much broader reach of receptor signaling outside of the T-tubular region in failing cells. In this study, we have observed an amplified β2AR signal associated with loss of caveolin-3 in failing rabbit myocytes, which preferentially propagates toward the myofilaments (Figures 2 and 6). These data, for the first time, link the redistribution of β2AR to amplified PKA signaling in subcellular compartments in models of cardiac disease, which may offer insight into cardiac hypertrophic remodeling and compensatory contractility in the heart.
PKA phosphorylation of TnI (Figure 8). Moreover, reintroduction of caveolin-3 could normalize the functional relationship between βAR and the L-type Ca^{2+} channel at both T-tubules and caveolae.16,21 As a result, the caveolin-3–expressing myocytes become more responsive to adrenergic stimulation to enhance contractility (Figure 7). Together, these data support the notion that caveolin-3 is essential in regulation of compartmented cardiac βAR signaling.25,40
Together, this study reveals that structural remodeling leads to βAR signal remodeling in failing myocytes, which yields a decrease in PKA activity at the PM and an increase in β2AR-PKA activity and phosphorylation of substrates on the myofilaments in hypertrophic rabbit hearts. Our study suggests caveolin-3 as an essential regulator of subcellular adrenergic signal at the myofilaments in myocytes.

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Disclosures
None.

References
What Is Known?
- Abnormal protein kinase A phosphorylation of substrates on myofilaments directly contributes to inotropic and lusitropic dysfunction in heart failure (HF).
- Adrenergic signaling is altered in HF myocytes, including downregulation of β1 adrenergic receptor (β1AR)-induced signals and an increase in the β2AR-induced signal.

What New Information Does This Article Contribute?
- In HF myocytes, β2AR-induced signaling is selectively downregulated at the sarcolemma, whereas β1AR-induced signaling gains access to myofilament.
- Reintroduction of caveolin-3 in HF myocytes prevents β2ARs from sending signal to myofilaments for substrate phosphorylation.
- Reintroduction of caveolin-3 normalizes βAR-induced contractile responses in HF myocytes.

Novelty and Significance
These studies reveal that βAR signaling undergoes remodeling in HF myocytes, which shifts the protein kinase A activity from the cell surface to the intracellular compartments including the sarcoplasmic reticulum and myofilaments. In particular, β2AR-induced signals gain access to the myofilament, which contributes to abnormal protein kinase A phosphorylation of troponin I and contractile dysfunction. Preclinical studies to assess the effects of inhibition of β2AR signal on HF are indicated.
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Online expanded Methods

Development of PKA FRET-based biosensor and adenovirus construction
The regular PKA activity biosensor AKAR3 was linked to the C-terminus of cardiac troponin T (provided by Dr. Jianpin Jin, Wayne State University, OH) to generate a myofilament-anchored AKAR3, termed MF-AKAR3. The recombinant construct in pcDNA3.1 (Invitrogen, CA, USA) was verified by DNA sequencing. MF-AKAR3 cDNA was subsequently subcloned into a shuttle vector pAdTrack-CMV to generate recombinant adenovirus according to the manufacturer’s instructions (Quantum Biotechnologies, Republic of South Africa). The PM-AKAR3 and SR-AKAR3 have been reported previously.2, 3

Experimental animals, cardiac myocyte isolation and culture
The animal care and experimental protocols followed US National Institutes of Health guidelines and were approved by the Institutional Animal Care and Use Committees of the University of California at Davis. HF was induced in New Zealand White rabbits by combined aortic insufficiency and stenosis as previously described.4 HF progression was monitored by echocardiography and myocytes were isolated 7.1±1.9 months after aortic constriction when left ventricle end-systolic dimension exceeded 1.4 cm. Briefly, rabbits were sacrificed under general anesthesia (induction with propofol 2 mg/kg followed by 2-5% isoflurane in 100% oxygen). After thoracotomy, the heart was quickly excised and rinsed in cold nominally Ca-free MEM. The right atrium was removed and the aorta opened to visualize the left coronary ostia which was then cannulated using a 4F catheter.5 Perfusion of the left ventricle and left atrium was established before removal of the right ventricle free wall and application of a purse-string suture to secure the catheter in place. The remainder of the isolation procedure was then essentially as previously described.6

For myocyte cultures, cells were plated on natural mouse laminin (Life Technologies, Grand Island, NY) coated dishes in culture medium PC-1 (Lonza Walkersville, MD, USA) supplemented with 1% Penicillin-Streptomycin, PH 7.4, filtered) and infected, when specified, with PM-AKAR3, SR-AKAR3, or MF-AKAR3 biosensor together with GFP-caveolin-3 or caveolin-3 for at least 28 hours.

Immunostaining
Confocal microscopy was performed using Zeiss LSM700 microscope (Zeiss, Pleasanton, CA) equipped with a Plan-Apochromat 63X oil-immersion objective. For co-localization experiments, cells were fixed for 15 min with PFA 4%, rinsed 3 times with PBS and permeabilized with blocking solution (0.5% NP40, 2% goat serum) for 15 min. Cells were then washed and stained 2 hours with anti-RyR antibody (obtained from the DSHB, the University of Iowa, IA), anti-caveolin-3 (BD, NJ, USA #610420), anti-PDE3 (a gift from Yan Chen, University of Rochester) anti-PDE4B (a gift from Marco Conti, University of California at San Francisco) antibodies, anti-PDE4D (Ab14613) (Abcam, Cambridge, UK) followed by 1 hour of the secondary anti-mouse or anti-rabbit Alexa Fluor® 488 antibodies (A-21063 and A-11055, respectively) (Thermo Fisher,
Western blotting
Left ventricular extracts or isolated cardiac myocytes were prepared in lysis buffer (25mM Tris HCl PH 7.6, 150 mM NaCl, 1% IGEPAL, 1% Sodium deoxycholate, 0.1% SDS, 1mM EDTA) with protease and phosphatase inhibitors (Na3F 100mM, Na2VO4 1mM, glycerol 1mM, NaPO2 2.5 mM, leupeptin 10 µg/ml, PMSF 1mM, aprotinin 10 µg/ml), and protein concentration was measured by BCA assay (Pierce, IL). Lysates were resolved by SDS-PAGE before being transferred onto a PVDF membrane (Merck Millipore ltd. MA, USA) and incubated overnight with primary antibodies followed by IRDye 680CW or 800CW secondary antibodies and acquired with an Odyssey scanner (LI-COR Biosciences Lincoln, NE, USA). The primary antibodies used for Western blotting were as follows: anti-β2AR (sc-570) (SCBT, TX, USA), anti-PDE3A (1098-1115, NIH), anti-PDE4D (Ab14613) (Abcam, Cambridge, UK) anti-phospho-PLB (Ser16, Badrilla), anti-PLB (MA3-922) (Affinity Bioreagent, CO, USA), anti-phospho-troponin I (Ser23/24, #4004) and anti-troponin I (#4002) (Cell Signaling, MA, USA). Signal intensity was quantitated by Image Studio software version 2.1 (LI-COR Biosciences Lincoln, NE, USA).

Fluorescence resonance energy transfer (FRET) measurements
Images were acquired using a Leica DMI3000B inverted fluorescence microscope (Leica Biosystems, Buffalo Grove, IL) with a 40X oil-emersion objective lens and a charge-coupled device camera controlled by Metafluor software (Molecular Devices, Sunnyvale, CA). Fluorescence images were recorded by exciting the donor fluorophore at 430-455 nm and measuring emission fluorescence with two filters (475DF40 for cyan and 535DF25 for yellow). Images were subjected to background subtraction, and were acquired every 20 seconds with exposure time of 200 ms. The donor/acceptor FRET ratio was calculated and normalized to the ratio value of baseline. The binding of cAMP to AKAR3 increases CFP/YFP FRET ratio. 1

Adult cardiomyocyte contractility
Fresh isolated adult cardiomyocytes were infected with GFP or GFP-caveolin-3 adenovirus for 18 hours. Cells were then placed in a dish with 3 ml beating buffer (NaCl 120 mM, KCl 5.4 mM, NaH2PO41.2mM, MgSO41.2 mM, HEPES 20mM, Glucose 5.5 mM, CaCl21mM, PH 7.1, filtered). Cells were paced at 1 Hz with voltage of 30 V using a SD9 stimulator (Grass Technology, Warwick, RI). Contractile shortening of myocytes were recorded on an inverted microscope (Zeiss AX10) at 20X magnification using Metamorph software (Molecular Devices, Sunnyvale, CA) with the following settings: 40 frames per second for 5 seconds, at an interval of 1 minute for a duration of 10 minutes. Isoproterenol (100 nM) was added into the dish after two movies were acquired. The percentage of myocyte fractional shortening (FS) was calculated as (maximal cell length – minimal cell length)/ maximal cell length * 100%. Analysis was performed using Metamorph.

Statistical analysis
All data were expressed as mean ± SEM. Statistical analysis was performed using GraphPad
Prism 6 software (La Jolla, CA). The sample size for each group was shown in the figure legends. The cells were from at least three sets of independent experiments. Differences between two groups were evaluated by 2-tailed Student’s t-test; comparisons of multiple groups were performed using one-way ANOVA followed by post hoc Bonferroni test. p < 0.05 was defined as statistically significant.

Reference

Supplemental Figure legend

Supplemental Figure I. Dose-dependent increases in AKAR3 FRET biosensors after stimulation of βAR in young adult rabbit myocytes. Rabbit myocytes expressing PM-AKAR3, SR-AKAR3, and MF-AKAR3 are stimulated with increasing doses of ISO. The maximal increases in different doses of ISO are plotted as dose response curves for individual biosensors, respectively.

Supplemental Figure II. Distribution of PDEs in SHAM and HF rabbit cardiac myocytes
Immunostaining shows distribution between PDE4B or PDE4D (red) and caveolin-3 (green), and between PDE3A (green) and the myofilaments (phalloidin, red) in healthy, HF, or HF myocytes with caveolin-3 expression.

Supplemental Figure III. Redistribution of PDE2 activity in HF rabbit cardiac myocytes
Quantification of the maximal changes in FRET ratio in SHAM and HF myocytes with or without expression of caveolin-3. Cells expressing PM-AKAR3, SR-AKAR3 or MF-AKAR are stimulated with the PDE2 selective inhibitor erythro-9-(2-hydroxy-3-nonyl)adenine (EHNA, 1 μmol/L). * p < 0.05 and ** p < 0.01 by one-way ANOVA followed by post hoc Bonferroni test.

Supplemental Figure IV. Distribution of PDEs in SHAM and HF rabbit cardiac myocytes
A and B) Immunostaining shows distribution between PDE4D (red) and caveolin-3 (green), and between PDE3A (green) and the myofilaments (phalloidin, red) in SHAM, HF, or HF myocytes with caveolin-3 expression. C) HF rabbit myocytes expressing flag-β2AR are fixed for immunostaining to show distribution of between β2AR and PDE3A or PDE4D.

Supplemental Figure V. PDE regulates PKA-mediated phosphorylation of myofilaments in SHAM and HF rabbit cardiac myocytes
Immunoblots show PKA phosphorylation of TnI in SHAM or HF cells after stimulation with ISO (100 nmol/L, 5 min) in the presence of PDE2 inhibitor
Supplemental Figure VI. Detection of baseline PKA activity at the subcellular compartments in SHAM and HF rabbit myocytes. Data shows quantification of the maximal changes in FRET ratio in SHAM and HF rabbit myocytes. A) Young adult myocytes expressing MF-AKAR3 are stimulated with ISO (100 nmol/L) in the presence of PKA inhibitor myr-PKI (10 μmol/L), KT5720 (10 μmol/L) or H89 (10 μmol/L). *** p < 0.001 by one-way ANOVA followed by post hoc Bonferroni test. B) SHAM and HF rabbit myocytes expressing PM-AKAR3, SR-AKAR3 or MF-AKAR are treated with PKA inhibitor H89 (10 μmol/L). * p < 0.05 by student’s t-test.

Supplemental Figure VII. Caveolin 3 modulates subcellular distribution of PDE3 and PDE4 activity in HF rabbit cardiac myocytes. Data shows quantification of the maximal changes in FRET ratio in SHAM myocytes and HF myocytes with or without expression of caveolin-3. Cells expressing PM-AKAR3 or MF-AKAR are treated with PDE3 selective inhibitor cilostamide (Cilo, 1 μmol/L) or PDE4 selective inhibitor Rolipram (Roli, 10 μmol/L). * p < 0.05 by student’s t-test.

Supplemental Figure VIII. Localization of β2AR and caveolin-3 in HF rabbit cardiac myocytes. HF rabbit myocytes expressing flag-β2AR alone (lower panels) or together with caveolin-3 (upper panels) are fixed for immunostaining to show distribution of β2AR and caveolin-3.
Supplemental Figure IV

A

SHAM

HF

HF + Cav3

PDE4D

Cav3

Merge
Supplemental Figure IV Continued

B

SHAM

HF

HF + Cav3

PDE3A
Phalloidin
Merge

C

HF

b2AR

PDE3A
Merge

HF

b2AR

PDE4D
Merge
Supplemental Figure V

Supplemental Figure VI

A

B

Baseline PKA activity