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Caveolin-3 regulates compartmentation of cardiomyocyte beta2adrenergic receptor-mediated cAMP signaling

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Abstract

The purpose of this study was to investigate whether caveolin-3 (Cav3) regulates localization of β_2 -adrenergic receptor (β_2AR) and its cAMP signaling in healthy or failing cardiomyocytes. We co-expressed wildtype Cav3 or its dominant-negative mutant (Cav3DN) together with the Förster resonance energy transfer (FRET)-based cAMP sensor Epac2-camps in adult rat ventricular myocytes (ARVMs). FRET and scanning ion conductance microscopy were used to locally stimulate β_2AR and to measure cytosolic cAMP. Cav3 overexpression increased the number of caveolae and decreased the magnitude of β_2AR -cAMP signal. Conversely, Cav3DN expression resulted in an increased β_2AR -cAMP response without altering the whole-cell L-type calcium current. Following local stimulation of Cav3DN-expressing ARVMs, β_2AR response could only be generated in T-tubules. However, the normally compartmentalized β_2AR -cAMP signal became diffuse, similar to the situation observed in heart failure. Finally, overexpression of Cav3 plays a crucial role for the localization of β_2AR and compartmentation of β_2AR -cAMP signaling to the T-tubules.

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tubules of healthy ARVMs, and overexpression of Cav3 in failing myocytes can partially restore the disrupted localization of these receptors.

Keywords

beta-adrenergic receptors; cardiomyocytes; T-tubules; FRET; SICM

1. INTRODUCTION

The beta-adrenergic receptors (β AR) [1] are the main G-protein coupled receptors which mediate the functional effects of catecholamines in the heart. There are two main β AR subtypes (β_1 and β_2 AR) which are expressed in human and rodent cardiomyocytes [1–3]. Selective acute or chronic stimulation of β_1 or β_2 AR elicits different cellular responses with respect to contractility, calcium cycling, hypertrophy and apoptosis. Whilst chronic β_1 AR stimulation usually promotes hypertrophy and apoptosis, β_2 AR elicits rather protective effects unless overexpressed at very high levels [4–7].

Highly localized activation of signaling pathways within different subcellular microdomains may be the key reason for these differences. β_1AR has been shown to act exclusively by increasing cAMP levels and stimulating CAMKII via the stimulatory G-proteins (G_s), while β_2AR can also activate inhibitory G-proteins (G_i), thereby inhibiting cAMP production, activating anti-apoptotic pathways and altering target protein phosphorylation through actions on phosphatases [5, 8, 9]. Previously, we used Förster resonance energy transfer (FRET)-based imaging to investigate subtype-specific cAMP signaling by β_1AR and β_2AR . Using a transgenically expressed FRET sensor in mouse cardiomyocytes, we demonstrated that β_1 -ARmediated cAMP signals diffused over long distances throughout the cell, whereas β_2AR -receptor signals were locally confined [10].

Such differences in spatial cAMP dynamics might result from differential localization of both receptor subtypes in caveolae micro-domains, whilst β_1AR has been found in both caveolar and non-caveolar membrane fractions of neonatal and adult cardiomyocytes; β_2AR was shown to exclusively localize in caveolae [11–15]. Caveolae are invaginations of the plasma membrane enriched in cholesterol, glycophospholipids, and glycosylphosphatidylinositol-anchored proteins [16–20]. Caveolin, the main protein component of caveolae, recruits components of various signaling pathways, including G_i proteins [21], endothelial nitric oxide synthase [22, 23] and several protein kinases.

Neonatal cardiomyocytes lack transverse tubules (T-tubules), but have an increased caveolae density. Caveolae may be developmental precursors of T-tubules and share some of their functions [24]. T-tubular development in striated muscle depends on cholesterol and caveolin-3 (Cav3), the principal protein component of cardiac caveolae [17]. Several studies using transgenic mice overexpressing Cav3 revealed increased number of sarcolemmal muscle cell caveolae [25]. Overexpression of Cav3 in Duchene muscular dystrophy skeletal myofibers increased caveolae number [26]. In contrast, Cav3 knockout mice showed complete loss of cardiomyocyte caveolae, with associated T-tubular disorganization and dramatic cardiomyopathy [27, 28]. Experiments in adult rat ventricular myocytes (ARVMs)

indicated that cholesterol depletion disrupted β_2AR coupling to inhibitory G-proteins, and acute chemical caveolae disruption using methyl- β -cyclodextrin (M β CD) led to an increase in β_2AR mediated cAMP and cell contractility [29]. This agrees with experimental evidence placing the β_2AR in a functional relationship with the L-type Ca²⁺ channel, both of which are predominantly localized in the T-tubules and caveolae [30, 31].

However, no studies have addressed subtype-specific β AR signaling and its regulation by caveolae in failing cardiomyocytes. Recently, we and others have shown that during the development of heart failure (HF) after myocardial infarction, a dramatic remodelling of intracellular T-tubular network occurs which is characterized by its disorganisation and dilation at earlier stages [32], as well as loss of cell surface T-tubular openings in severe chronic HF [33]. Importantly, the latter condition leads to a redistribution of β_2 AR from T-tubules to detubulated membrane areas. This triggers a loss of β_2 AR-cAMP signal confinement, the diffusion of which is normally restricted to this microdomain in healthy cells, as revealed by a novel scanning ion conductance microscopy (SICM)/FRET-based imaging technique [34].

In this study, we tested the hypothesis that Cav3 regulates T-tubular localization of β_2AR and its signalling to cAMP which are altered in HF. Using SICM/FRET and adenovirusmediated expression of Cav3 or the dominant-negative mutant of Cav3 (Cav3DN), we found that Cav3 selectively modulates the spatial compartmentation of β_2AR -cAMP responses in the T-tubular compartment which is altered in HF. Cav3 overexpression in failing cardiomyocytes reversed the pathological redistribution of β_2AR -cAMP signalling.

2. MATERIALS AND METHODS

2.1 Heart Failure Model and Cell isolation

All animal surgical procedures and perioperative management were carried out in accordance with the Guide for the Care and Use of Laboratory Animals published by the U.S. National Institutes of Health under assurance number A5634-01. All animal surgical procedures and perioperative management conformed to the UK Animals (Scientific Procedures) Act 1986. Adult male Sprague-Dawley rats (250-300 g) underwent proximal coronary ligation to induce myocardial infarction as described [33]. Anaesthesia was induced by administration of 5% isoflurane for induction and reduced to 2% isoflurane once intubated and ventilated. It was ensured that pain reflexes were absent prior to an incision being made by testing pedal and palpebral reflexes. Preoperatively, buprenorphine was administered subcutaneously at a dose of 0.05 mg/kg to ensure adequate analgesia in addition to anaesthesia. Further doses of buprenorphine were administered as required postoperatively if there was any sign of distress. In addition, enrofloxacin (5 mg/kg) and 0.9 % saline (10 ml/kg) were administered pre-operatively. Cardiac failure was assessed via biometric and echocardiographic means. Heart weight corrected to tibia length provided a measure of hypertrophy. Echocardiography was performed under anaesthesia (2 % isoflurane) in the week of sacrifice. B-Mode echocardiographic images were acquired in the parasternal long axis at 70 Hz using a Visualsonics Vevo 770. This chronic HF model associated with massive hypertrophy and dilation has been previously well characterized by

histological and functional analysis [33]. Isolated failing cardiomyocytes have been shown to have lower $\beta_1 AR$ and unchanged $\beta_2 AR$ densities [34].

Hearts were harvested for cell isolation 16 weeks post myocardial infarction, when a chronic HF phenotype has developed. Rats were sacrificed by cervical dislocation following brief exposure to 5% isoflurane until righting reflex was lost. Myocytes from healthy or failing hearts were isolated by the Langendorff perfusion method [34], plated on laminin coated coverslips and infected for 48h with Epac2-camps, Cav3 or Cav3DN adenoviral vectors [35]. For control experiments shown in Figures 3 and 4, cells were isolated from agematched sham-operated control animals.

2.2 Electron Microscopy

Cardiomyocytes were fixed with 2.5 % glutaraldehyde for 2 to 4h and then centrifuged at 500g for 5 min and the pellet was left overnight. The pellet was washed three times in cacodylate buffer and fixed in 1% osmium-tetroxide, followed by a 5–10 min washing with pure water. A small amount (25 to 50 μ l) of liquid 2% agar at 45°C was added to the pellet. Drops were left to solidify on polythene, providing agar blocks with evenly distributed cells. The blocks were dehydrated through a series of graded alcohols, propylene oxide, and embedded in araldite. For low power examination by light microscopy before EM examination, 1 μ m thick sections were cut and stained with 1% toluidine blue in 1% borax. For transmission electron microscopy, ultra-thin sections were stained with uranyl acetate and lead citrate. The ultrastructural features of cardiac myocytes, especially the membrane area were examined.

2.3 Western blot analysis

ARVMs were transfected with the control (LacZ), Cav3 or Cav3DN adenoviruses (all at MOI 500). 48h later cell were washed once and homogenized in the lysis buffer containing: 300mM sucrose, 150mM NaCl, 1mM EGTA, 2mM CaCl₂, 1% Triton-100, 10 mM HEPES, pH=7.4. 5 µg protein samples were separated on a 15 % SDS-polyacrylamide gel and blotted onto nitrocellulose membrane (Millipore Corp., Bedford, MA, USA). Membranes were blocked for 1h at room temperature with 5 % non-fat milk in PBS containing 0.05 % Tween-20 and incubated with primary monoclonal Cav3 antibodies (1:5000, Santa Cruz Biotech, USA) overnight at 4°C. After washing, the blots were probed with a 1:5000 dilution of horseradish peroxidase (HRP)-conjugated anti-mouse IgG (Sigma, St Louis, MO, USA) and visualized using the ECL kit (Amersham Biosciences).

2.4 Radioligand binding studies

Radioligand binding studies were performed as previously described [36]. Briefly, isolated cell membranes were incubated for 1h at 30°C with 60–100 pM ¹²⁵Icyanopindolol (¹²⁵I-CYP) (PerkinElmer Life Sciences, Dreieich, Germany) and increasing concentrations of ICI118, 551. Competition binding curves were fitted and analyzed with the Prism software (GraphPad, San Diego, CA).

2.5 FRET imaging of cAMP in living cardiac myocytes

FRET was performed in cells infected for 48h with Epac2-camps adenovirus [37]. Cells were washed once and measured at room temperature in the buffer containing 144mM NaCl, 5.4mM KCl, 1mM MgCl₂, 2mM CaCl₂, and 10mM HEPES, pH=7.3. The imaging system was build around the Nikon TE2000 microscope equipped with halogen lamp illuminator pillar, EX436/20 excitation filter combined with DM455 dichroic mirror. Cell fluorescence was split into YFP and CFP channels using the DualView (Optical Insights, equipped with 535/40 and 480/30 emission filters) and monitored by the ORCA-ER CCD camera (Hamamatsu Photonics, Welwyn Garden City, UK). Cell images were analyzed using SimplePCI software (Hamamatsu). FRET ratios were corrected for the bleed through of CFP into the YFP channel and analyzed using the Origin software (OriginLab Corporation, Northhampton, MA).

2.6 SICM and SICM/FRET experiments

SICM/FRET measurements were performed exactly as previously described [34]. These included scanning of the living cell surface, local ligand application into single T-tubules via the scanning pipette and sub-cellular analysis of the induced cAMP signal by FRET microscopy. For high-resolution SICM, resistance of the pipette tip was around 200M Ω . To calculate the Z-groove index, we measured the maximum length of Z-groove sobserved on single SICM images and divided this value by the total estimated Z-groove length as previously described [33].

2.7 Electrophysiological recordings from adult cardiac myocytes

Macroscopic currents were recorded using the whole-cell patch clamp technique with the external recording solution containing 1mM CaCl₂, 0.5mM MgCl₂, 5mM HEPES, 140mM choline chloride, 5mM CsCl, 5.5mM glucose, pH 7.4 with CsOH, ~305mOsm, and the internal pipette solution containing 130mM Cs-methanesulphonate, 11mM EGTA, 10mM HEPES, 2mM MgCl₂, 5mM Mg-ATP, 0.3mM Na-GTP, pH 7.2 with CsOH, ~290mOsm/kg.

2.8 Statistical analysis

Data were analyzed using Origin Pro 8.6 software (OriginLab Corporation, Northhampton, MA) and presented as means \pm SE from the indicated number of independent experiments, animals or cells isolated from several rats per condition, as indicated in the figure legends. Differences were tested using one-way ANOVA with Bonferroni post-test and considered significant at p<0.05.

2.9 Computer simulations

The β -adrenergic signaling formulation of Heijman et al. [39] was modified to represent one dimensional spatial distribution along the long axis of a simulated ventricular myocyte. A brief summary of aspects of the Heijman et al. model that are not directly germane to the simulations shown here can be found in the Supplementary Text. Crucial aspects are listed here together with modifications for the present study. The model includes distinct caveolar, extra-caveolar, and cytosolic signaling domains (cav, ecav, and cyt, respectively) and separate representations for β_1 and β_2 ARs and their associated G-proteins. Here, we

distinguish T-tubular from non-T-tubular, or "crest" sarcolemma (named based on topological appearance in SICM scans). In both membrane types, β_2ARs reside primarily in cav domains (85%, with 15% in ecav in T-tubule; 100% cav in crest). The cyt domain is without β_2ARs .

The 100 µm, 38e-6 µL (volume) cell was divided into 120 evenly spaced, equal volume segments. Segments alternated between T-tubule and crest type: 6 of each within 10 µm, as in SICM scans. Diffusion of cAMP and other molecules (i.e. PKA, phosphodiesterases, and PKI) between ecav and cyt domains in adjacent segments was at a rate of 1.0e-7 µL/s. The specific choice of 1.0e-7 µL/s was similar to the inter-domain diffusion rates in the original formulation by Heijman et al. and was supported by the empirical observation that either a ten-fold increase (to 1.0e-6 µL/s) or a ten-fold decrease (1.0e-8 µL/s) prevented cAMP concentration gradients from forming in disease cases where a diffusion gradient was expected (e.g Cav3DN, heart failure).

Cav domains are discrete and isolated from one another. Therefore, there was no direct inter-segmental diffusion of cAMP emanating from cav domains. However, indirect inter-segmental diffusion between adjacent T-tubule and crest segments could occur via ecav and cyt domains, which were represented as connected and contiguous. That is, ecav was connected to same-segment cav and cyt domains and to the neighboring segment ecav domain; cyt was connected to same-segment cav and ecav domains and to the neighboring segment cyt domain.

3. RESULTS

3.1 Caveolin-3 regulates the number of caveolae in cardiomyocytes

To analyze the effect of the functional Cav3 protein on the amount of caveolae in adult cardiomyocytes, we manipulated the function of endogenous Cav3 in two ways. First, we overexpressed the c-myc-tagged human Cav3 via the previously described adenoviral vector which has been shown to protect NRCM from hypertrophy [35]. Transduction of ARVMs for 48 h with this Cav3 adenovirus resulted in a 3-fold increase caveolae number, as revealed by electron microscopy of longitudinal cardiomyocyte sections (Figure 1A, B). We quantified the degree of Cav3 overexpression in ARVMs by Western blot analysis with a monoclonal Cav3 antibody, which detects both endogenous and the c-myc tagged Cav3 bands, and observed a 3.8 ± 0.1 fold increase (mean \pm SE, n=3) in the amount of Cav3. In living Cav3 overexpressing ARVMs, cell-surface caveolae could be detected by highresolution SICM (Figure 1D). Second, we used an adenovirus which encodes an untagged mutant of Cav3 lacking 3 amino acids (threonine, phenylalanine and threonine) in the caveolin scaffolding domain and behaves in a dominant-negative fashion (Cav3DN) [40], causing limb-girdle muscular dystrophy in humans [35]. Cav3DN could also be detected in Cav3 western blot, at a 2.9 ± 0.4 fold (mean \pm SE, n=3) higher level than the endogenous Cav3 (Figure 1C). In contrast to wildtype Cav3, expression of the Cav3DN mutant led to a moderate but significant decrease in caveolae number (see Figure 1A, B).

3.2 Caveolin-3 regulates βAR-cAMP signaling in a subtype-specific manner

We next investigated whether Cav3 might have an effect on β AR subtype-specific cAMP signaling. To test whether Cav3 overexpression had an effect on the total and relative numbers of β_1 and $\beta_2 ARs$, we prepared cell membranes from LacZ or Cav3 transfected ARVMs and used them in total radioligand binding and competition displacement experiments. There was no significant difference in total amounts of βAR receptors (~0.5 fmol/µg membrane protein) and in the relative fractions of β_1 and β_2 ARs as measured by radioligand displacement with ICI 118551 (Supplementary Figure 1A). To elucidate whether functional activity of β_1 or β_2 AR is affected by Cav3 overexpression, we performed FRETbased imaging of cAMP levels in living ARVMs. 48 hours after co-transfection with adenoviruses for the Epac2-camps FRET biosensor and Cav3, we monitored intracellular cAMP levels upon selective stimulation of *ARVM* with either β_1 or β_2 AR ligands (Figure 2). We stimulated entire cardiomyocytes (bath application) with saturating concentrations of the β AR agonist isoproterenol (ISO, 100 nM) in presence of selective antagonists of either β_2 AR (ICI118551, 50 nM) or β_1 AR (CGP20712A, 100 nM). Subsequently, the adenylyl cyclase activator forskolin, 10 µM, was applied to determine the maximal cAMP response. In control cardiomyocytes, cAMP signals from β_2AR were smaller than β_1AR responses, consistent with our previous results in adult mouse and rat cardiomyocytes [10, 34]. Selective stimulation of β_1 AR led to similar amounts of cAMP produced in case of control or Cav3 expressing cells (Figure 2A, B). In sharp contrast, the cAMP signals upon selective β_2 AR stimulation were ~2-fold smaller when Cav3 was overexpressed, without any change in the FRET-responses to the direct adenylyl cyclase activation (Figure 2C, D).

3.3 Cav3DN increases the levels and intracellular diffusion of β2AR-cAMP

We hypothesized that endogenous Cav3 restricts local cAMP gradients generated by $\beta_2 AR$ stimulation to the T-tubular compartment. To test this, we studied whether the expression of the dominant negative Cav3 mutant affects the spatio-temporal characteristics of the β_2 ARcAMP responses. In Cav3DN expressing cells, selective whole-cell β_2 AR stimulation led to significantly higher total levels of intracellular cAMP (~2-fold increase in the FRET signal amplitude) (Figure 3A). To study whether this might have any effect on $\beta_2 AR$ localization and the sub-cellular cAMP dynamics, we performed SICM/FRET experiments in Cav3DN expressing cells. Previously, using this technique we showed that in healthy ARVMs, functional β_2 ARs are restricted to T-tubules, *while* in failing cardiomyocytes or in cells after chemical cholesterol depletion using M β CD, β_2 AR was redistributed from the T-tubules to non-tubular membrane areas where this receptor induced far-reaching cAMP signals [34]. Here we took advantage of Cav3DN as a more specific tool to address the role of Cav3 in β₂AR localization and signaling. In contrast to our previous MβCD results, Cav3DN expressing cells did not show any $\beta_2 AR$ redistribution (see Figure 3B). However, the spatial distribution of the B2AR-cAMP signals after local stimulation of single T-tubules was dramatically changed. In contrast to control cells where these signals are stringently localized (Figure 3C), Cav3DN expression led to far-reaching cAMP signals (Figure 3D) similar to those which can be observed in failing cardiomyocytes isolated from rats 16 weeks after myocardial infarction (Figure 3E). These results suggest that Cav3 plays a crucial role in the confinement of β_2 AR-cAMP signaling to the T-tubular compartment.

Similar to the previously published results with M β CD [41], the density of the whole-cell L-type Ca²⁺ channel currents were not affected by Cav3DN (Supplementary Figure 2), suggesting that the basal activity of these channels is not altered by the disruption of the T-tubular cAMP micro-domain.

3.4 Overexpression of Cav3 partially restores the disrupted $\beta_2 AR$ localization in heart failure

Finally, we asked whether Cav3 overexpression might improve the abnormal localization of β_2AR and β_2AR -cAMP signals found in failing cardiomyocytes [34]. We isolated cells from the same rat chronic HF model used in this former study (Supplementary Figure 3), and overexpressed Cav3 together with Epac2-camps for 48 h to analyse the membrane structure and receptor localization by SICM/FRET. Cav3 overexpression for this short period of time did not result in any significant increase in the membrane curvature or the appearance of the T-tubular openings, as judged based on overall membrane topography and on the Z-groove index (Figure 4A, B). However, cells analyzed by SICM/FRET showed redistribution of β_2AR back into the remaining T-tubules (Figure 4C) in comparison to heart failure cells without Cav3 overexpression. This suggests that Cav3 is important for T-tubular localization of β_2AR , and its overexpression can restore the disrupted β_2AR distribution in the failing cardiomyocytes. However, Cav3 overexpression also reduces the amplitude of β_2AR responses as demonstrated by the bath application experiments described above. (Figure 4D-**representative trace**).

3.5 Mechanisms by which Cav3 regulates localization of the β2ARs response

We developed a theoretical framework so that a hypothetical explanation for Cav3 interactions with β_2 ARs under the various conditions considered here and in our previous related work [34] could be proposed. The conditions considered include control, Cav3 overexpression (Cav3), expression of dominant negative mutant Cav3 (Cav3DN), application of M β CD, heart failure, and failure with Cav3 overexpression (failure+Cav3). The framework is illustrated in Figure 5.

The working components of the model, which associate with β_2ARs , include Cav3, Cav3DN, and fully formed caveolae; β_2ARs can also be unassociated (lone). Pairings of β_2AR with caveolae or Cav3 were considered to be discrete "cav" signaling domains, where cAMP was local due to spatial isolation (see Methods section 2.9) and where relatively high phosphodiesterase (PDE) activity partially muted signals (see Supplementary Text). Lone β_2ARs or β_2ARs paired with dysfunctional Cav3DN constituted ecav signaling domains. cAMP generated by ecav β_2ARs was readily diffusible because ecav elements were contiguous (not discrete like cav, c.f. Methods 2.9) and because PDE activity was low enough to permit cAMP accumulation. Ecav lacks PDE type 3, which is found in cav, and ecav has a volume that is twice that of cav, causing relative dilution of PDE activity (see Supplementary Text). The cytosol was devoid of β_2ARs . Passive diffusion of cAMP in cyt occurred, but signals were greatly diminished by the fact that cyt volume was roughly tenfold larger than sub-membrane volume (cav+ecav).

Our electron micrographs showed fully formed, flask-shaped caveolae in crest regions, but they were *not observed in* T-tubules. Recently, Wong et al. *presented a similar finding on the basis of a highly novel fluorescent microscopy technique coupled with electron microscopy (their* Supplementary Figure S4 [38]). *Cav3 protein is apparently abundant in T-tubules but it is not clear whether fully stabilized caveolae domains can be formed in this region* (e.g. [38]). Thus, to represent the control case, β_2 ARs were placed in caveolae in the crest and were Cav3-associated in T-tubules (exception: 15% of T-tubular β_2 ARs were lone; Figure 5, top middle).

Cav3 overexpression (Figure 5, top right) paired the few lone T-tubular β_2ARs with Cav3 and added nonfunctional Cav3 and caveolae lacking β_2ARs . We hypothesized that Cav3DN (Figure 5, top left) affected T-tubular β_2AR :Cav3 pairs, replacing 75% of native Cav3 with mutant Cav3DN, and changing their designation from cav to ecav in the model. Importantly, we hypothesized that β_2AR :caveolae pairs at the crest were immune to Cav3DN replacement. This concept is supported further below. We hypothesized that formation of β_2AR :caveolae and β_2AR :Cav3 complexes requires cholesterol. Therefore, M β CD was represented by strictly lone β_2ARs in crest and T-tubules (Figure 5, bottom left). Failure was represented by Cav3 loss (affecting 75% of β_2AR :Cav3 pairs in intact T-tubules) and also loss of T-tubules (half of them, as indicated by Z-groove index reduction shown here and previously [34]). Lost T-tubules purged their contents to the crest, dissociating Cav3 from 90% of β_2ARs in the process (Figure 5, bottom middle). Finally, overexpressing Cav3 in heart failure was represented as pairing lone β_2ARs with Cav3 (1000% and 300% increase in β_2ARs :Cav3 pairings in crest and T-tubules, respectively, Figure 5, bottom right).

The central assumption of this theoretical framework is summarized as follows (supported below). β_2AR associations with caveolae, found in the crest, are stable in the face of Cav3DN expression, while associations with Cav3 protein(s) in the T-tubules are not. This assumption can alternatively be viewed as causing limited pools of Cav3 to prioritize the crest over T-tubule when it comes to the localization of cAMP by forming stronger cAMP compartments. Other assumptions (e.g. the % changes above), though not incidental, were of quantitative but not qualitative consequence and needed to be made in order to realize a computational model to test the validity of the framework in Figure 5.

Figure 6 presents simulation results. Plotted are sub-membrane cAMP concentration landscapes ([cAMP] in μ M) computed in space and time along a longitudinal line scan (100 microns) following β_2 AR stimulation at either a distal T-tubular site (panel A) or at a distal crest site (panel B). We chose to measure [cAMP] at the membrane where the majority of important signaling targets reside (rationale outlined further in Supplementary Text and Supplementary Figure 4). Pipette placement over a T-tubule stimulates many β_2 ARs, all along the entire depth of the structure, while stimulation at crest sites affects only the few β_2 ARs immediately beneath the pipette (e.g. pipette cartoon in Figure 5, bottom middle). Simulations accounted for this with a 1000-fold T-tubule to crest β_2 AR capture ratio (results were not different using a lower estimate of 100-fold).

T-tubular stimulation resulted in sharp local spikes in [cAMP] at the stimulation site in all cases. However, diffuse cAMP (i.e. smooth concentration gradient from the stimulus site)

occurred only in Cav3DN, failure, and M β CD cases, but not in cases of control, Cav3 overexpression, or heart failure with Cav3 overexpression. This matches experiments shown here and in our previous work: Cav3DN – Figure 3D, heart failure – Figure 3E, M β CD – Supplementary Figure S8C of Nikolaev et al. [34], control – Figure 3C, and failure+Cav3 – Figure 4D. The cAMP signal was reduced by Cav3 overexpression (integrated total cAMP was ~70% of control). This is in agreement with Figure 2D. Cav3DN simulations showed an increase in overall cAMP relative to control (73% increase in integrated total cAMP). Although the amplitude is not different, more cAMP response is present due to propagation throughout the cell. This is in agreement with Figure 3A,

Simulation results showed that a response to crest stimulation was only registered for the M β CD and heart failure cases. This is in accord with experiments (no response: control – Figure 2G of Nikolaev et al. [34], Cav3DN – Figure 3B, and failure+Cav3 – Figure 4C. response: M β CD – Supplementary Figure S8B of Nikolaev et al. [34], and failure – Figure 4C). Importantly, when we relaxed the central assumption that Cav3DN does not disrupt β_2 AR associations with caveolae in the crest, there was a robust crest response in Cav3DN simulations, in violation of Figure 3B experimental results. Without this central assumption, the Cav3DN case was nearly identical to the heart failure case. Though crest caveolae were not degraded in failure, T-tubule loss caused redistribution of newly lone β_2 ARs to the crest. This distinguished the failure response from the Cav3DN response.

Simulations predicted baseline cAMP elevation in pathological states (i.e. Cav3DN, failure and M β CD). This was caused by the greater proportion of β_2 ARs in the ecav domains, where PDE activity was relatively low. FRET experiments cannot support or refute this prediction.

4. DISCUSSION

The β_2AR has been shown to associate with caveolin and caveolae in neonatal and adult ventricular cardiomyocytes. However, functional consequences of the receptor-caveolin interactions remained poorly defined, especially in failing cardiomyocytes. In this work, we analyzed the role of Cav3 in β_2AR localization and in β_2AR -cAMP compartmentation in healthy and failing ARVMs and showed that β_2AR localization and its compartmentalized signaling are directly regulated by Cav3.

Overexpression of Cav3 in healthy ARVMs led to an increase in caveolae number (see Figure 1) and to a subtype-specific decrease of β_2AR -cAMP production which was not associated with altered receptor densities or with changes in the β_1/β_2AR expression ratio (see Figure 2, Supplementary Figure 1). This suggests that Cav3 restricts cAMP accumulation or its diffusion from β_2AR into the cytosol. Conversely, adenoviral gene transfer of the dominant-negative mutant Cav3DN led to an increase of the total β_2AR -cAMP levels (see Figure 3A). Interestingly, Cav3 had no effect on the amplitude of the β_1AR -stimulated cAMP response. Recently, it was demonstrated that β_1AR -mediated cAMP production in the cytosolic compartment of ARVM assessed by the Epac2-camps sensor was unchanged after cholesterol depletion by M β CD. However, it was increased in the compartment associated with caveolae and the type II cAMP-dependent protein kinase,

suggesting that there is a population of $\beta_1 AR$ specifically coupled to caveolae which cannot be functionally detected by our cytosolic cAMP sensor Epac2-camps [13].

Previously, Calaghan and White demonstrated that β_1 AR-dependent modulation of cell shortening and Ca^{2+} transient was not potentiated by caveolar disruption, while $\beta_2 AR$ stimulation significantly increased phospholamban phosphorylation, cell shortening and Ca^{2+} transients after M β CD treatment [29]. Therefore, β_2AR association with caveolae is required for the localized signaling and relative dampening of the positive inotropic and lusitropic response mediated by this receptor. Using a specific Cav3 inhibitory peptide, the same group showed that selective disruption of Cav3 results in increased β_2 AR-stimulated phospholamban phosphorylation. This was not only due to an increase in cAMP levels but also due to enhanced inhibitory modulation of phosphatases at the sarcoplasmic reticulum which can be explained by changes in receptor-cAMP compartmentation [42]. However, the role of Cav3 in β_2 AR localization has not been studied before due to the lack of appropriate techniques. To directly visualize the functional interactions of $\beta_2 AR$ with the sub-cellular pools of cAMP, we used the recently developed SICM/FRET method which can correlate the localization of the functional receptor with the cAMP microdomains. In our previous study, we used M β CD to disrupt cardiomyocyte caveolae and found β_2 AR responses to be present both in the T-tubular and in non-tubular areas of the cell surface. Stimulating $\beta_2 AR$ in either T-tubules or crests of MBCD-treated cardiomyocytes generated far-reaching cAMP signals. This contrasted with the normal cell in which $\beta_2 AR$ signal is exclusively present in T-tubules and is stringently confined [34]. Here we disrupted caveolae more precisely by expressing a dominant-negative Cav3DN mutant. We found that the β_2AR signal does not migrate from T-tubules to the non-tubular cell surface areas, suggesting unaltered receptor localization. However, upon stimulation of the $\beta_2 AR$ in the T-tubules, the resulting cAMP signal becomes diffuse, similar to its dynamics in failing cardiomyocytes (see Figure 3). This strongly suggests that Cav3 may be responsible for the compartmentation of the β_2 ARcAMP response to the T-tubules of healthy cardiomyocytes.

Since the β_2AR signaling and its regulation by caveolae in failing cardiomyocytes has not been systematically analyzed before, we studied how Cav3 overexpression might affect the altered β_2AR localization and cAMP dynamics in HF. It is widely accepted that β_2AR plays a cardio-protective role in the healthy heart [43]. Whilst transgenic overexpression of Cav3 in mouse cardiomyocytes induces endogenous cardiac protection from ischemia/reperfusion injury via increased activation of survival kinases, this effect is absent in Cav3 knockout mice [44]. The previously reported down-regulation of Cav3 in various animal HF models [45–48], and the adverse cardiac phenotype and T-tubular abnormalities in Cav3 knockout mice [27, 28] support the idea of Cav3 overexpression as a beneficial treatment option for HF. We tested this possibility in cells isolated from the chronic rat HF model where we overexpressed Cav3 for 48h (see Figure 4). This short treatment did not lead to any measurable improvement of cell surface morphology, but it was associated with redistribution of β_2AR 's functional effects back into the T-tubules, suggesting that Cav3 might potentially restore the β_2AR localization and its confined signaling to cAMP.

That the computational model was able to reproduce essential experimental observations is support for our hypothesis and its underlying assumptions. That is, β_2AR associations with

Cav3 molecules in T-tubules are more readily disrupted than are β_2AR associations with fully formed caveolae. The exact reasons for this should be the subject of future work. We speculate that binding among the 14–16 Cav3 molecules in the formation of a caveolae [17] is cooperative, and that this could be the stabilizing factor. The mechanism by which caveolae or Cav3 association can confine cAMP response resulting from β_2AR stimulation is likely to be indirect. We speculate that individual β_2AR :Cav3 and β_2AR :caveolae are diffusively separated from the general milieu. Moreover, we speculate that relatively high phosphodiesterase activity in the cav domain (included in the Heijman et al. model, see Supplementary Text for more detail), can mute cAMP levels and thereby limit its reach.

Implications of this study extend beyond Cav3 mutations and heart failure. A recent publication showed that a diet high in palmitate disrupted Cav3 in ventricular myocytes [49]. High dietary palmitate may lead to cellular behavior similar to Cav3DN case, where local cAMP responses to sympathetic β_2 AR stimulation were lost. This may have consequences for individuals' cardiovascular physiology. *Finally, although the rat cardiomyocyte is the model presented in this paper the authors are aware of the limitations of this species with regard to the necessity of culture. The t-tubular network is observed to degrade during the course of the culturing process. Equally, adenovirus appears to be the only method of gene delivery robust enough for use with this cell type. The physical and molecular changes caused by adenoviral entry and t-tubular disruption as well as the effects of unloading in culture may reduce the relevance of data from studies using this cell to failing cells in vivo.*

Future studies should analyze β_2AR localization and cAMP signaling by SICM/FRET in cardiomyocyte-specific Cav3 transgenic and knockout mice where the differences in membrane morphology might be even more pronounced due to constitutive expression or deletion of Cav3. *Transgenic mice are in existence, which constitutively express FRET sensors similar to the one employed by this study* [34]. *Crossing these mice with Cav3 transgenics may provide an avenue to studying the role of Cav3 in modulating* β_2AR *function in freshly isolated cells.* Finally, measurements in failing human myocytes with Cav3 re-expression could be performed when technically possible, *as this would also require culturing.* These studies might pave the way towards a new gene therapy option for HF patients.

5. CONCLUSIONS

In healthy adult rat cardiomyocytes, Cav3 regulates β_2 AR-cAMP signaling, playing a crucial role for the compartmentation of cAMP signals to the T-tubular compartment. Disruption of Cav3 function with a dominant-negative mutant leads to far-reaching β_2 AR-cAMP signals similar to those observed in heart failure. In failing cardiomyocytes, Cav3 overexpression can partially restore the disrupted localization of these receptors.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

β-AR	β -adrenergic receptor
ARVM	adult rat ventricular myocytes
Cav3	caveolin-3
Cav3DN	caveolin-3 dominant negative mutant
FRET	fluorescence resonance energy transfer
HF	heart failure
ΜβCD	methyl-β-cyclodextrin
SICM	scanning ion conductance microscopy

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HIGHLIGHTS

- Caveolin-3 (Cav-3) controls cardiomyocyte β₂AR signaling *via effects on cAMP responses*
- Cav-3 is necessary for proper compartmentation of β_2 AR-cAMP responses
- Loss of Cav-3 function disrupts β_2AR -cAMP's spatial compartmentation specifically at the T-tubules
- Overexpression of Cav-3 in failing myocytes can restore T-tubular $\beta_2 AR$ localization and spatial compartmentation



Figure 1. Cav3 overexpression increases caveolae number, while the expression of the dominant-negative Cav3 mutant slightly decreases it

A) Representative electron microscopy image of longitudinal cardiomyocyte sections showing caveolae (marked with black arrows) in control cells (infected with LacZ adenovirus) as well as in cells overexpressing wildtype Cav3 or Cav3DN mutant (all at MOI 500).

B) Quantification of the number of caveolae per μ m cross-section of the membrane from the electron microscopy images. Data are means \pm SE, n=4–5 per condition. *, ** - Differences are significant at P<0.05 or 0.01, respectively.

C) Western blots analysis of the Cav3 overexpression 48 h after transduction with the control (LacZ), Cav3 or Cav3DN adenoviruses (all at MOI 500).

D) High resolution SICM image of an ARVM overexpressing Cav3 (MOI 500) which shows clearly visible caveolae.



Figure 2. FRET-based cAMP measurements upon selective stimulation of β_1 or β_2AR in cardiomyocytes overexpressing Cav3

A) Cav3 overexpression does not affect β_1AR responses. ARVMs were infected for 48 h with Epac2-camps cAMP sensor adenovirus in combination with LacZ or the Cav3 virus (all at MOI 500). cAMP levels are shown as % change of the YFP/CFP ratio. A decrease in this ratio corresponds to an increase in intracellular cAMP. Cells were first treated with 100 nM isoproterenol (ISO) plus 50 nM of the β_2AR blocker ICI118551 (ICI) and subsequently fully stimulated with 10µM forskolin. Quantification of the FRET responses from β_1AR is presented in **B** (means ± SE, n=5–14cells).

C) Cav3 overexpression decreases cAMP production after β_2AR stimulation. Cells were first treated with 100nM ISO plus 100nM of the β_1AR blocker CGP20712A (CGP) and subsequently fully stimulated with 10µM forskolin. Quantification of the FRET responses from β_2AR is shown in **D** (means ± SE, n=7–12 cells). * - Differences between the control and Cav3 groups are statistically significant at *P*<0.05.



Figure 3. Cav3 modulates cAMP levels and its spatial distribution after β_2AR stimulation A) Expression of the dominant negative Cav3DN construct results in an increase in intracellular cAMP stimulated by β_2AR . Cells were transduced with either LacZ (control) or Cav3DN together with the Epac2-camps sensor adenovirus (all at MOI 500) and stimulated as described in Figure 2C and D. Data are means \pm SE (n=12–13 cells). * - Difference from control is significant at *P*<0.05.

B) SICM/FRET analysis of β_2AR distribution in Cav3DN expressing cells. β_2AR was locally stimulated with the scanning pipette in either single T-tubules or in cell crests located between the Z-lines. The experiment was performed exactly as previously described [34]. β_2AR -cAMP signals can be detected only upon local receptor stimulation in the T-tubules, suggesting that β_2AR localization is unaltered in Cav3DN expressing cardiomyocytes. Data are presented as means±SE (n=5–8 cells). ** - Difference is significant at *P*<0.01. C) Local β_2AR stimulation in single T-tubules (white arrow denotes the position of the SICM pipette) of control (LacZ expressing) cardiomyocytes results in a highly localized cAMP signal which is measurable only in the subcellular region adjacent to the pipette. Data

in the graph show YFP/CFP ratios in different color-labeled regions across the cell cytosol. Representative experiment, n=>6 cells

D) Local β_2AR stimulation in single T-tubules of Cav3DN expressing cardiomyocytes results in a far-reaching cAMP gradient which diffuses across the entire cell. Data are presented as in C. Representative experiment, n=>8 cells.

E) Local T-tubular β_2AR stimulation in failing cardiomyocytes leads to a similar farreaching cAMP gradient. Representative experiment, n=10 cells.



Figure 4. Cav3 overexpression in failing cardiomyocytes partially restores the T-tubular localization of $\beta_2 AR$

A) Cav3 overexpression for 48 h (MOI 500) does not lead to a significant improvement of the cell surface morphology. Representative SICM image of a failing cardiomyocyte overexpressing Cav3 (n=20 cells).

B) Quantification of the Z-groove index in these cells compared to healthy (Control) and failing (HF) cardiomyocytes without Cav3 overexpression (LacZ transfected, MOI 500). Shown are means \pm SE (n=12–16 cells). ** - Difference is significant at *P*<0.01. NS - no significant difference.

C) SICM/FRET analysis of β_2AR localization in Cav3 expressing cells, isolated from rats 16 weeks post-MI. FRET responses are significantly higher when agonist is applied to the t-tubules compared to the cellular crests. Graphs show Mean ± SE (n=15–19 cells) *P<0.05. This suggests some restoration of the normal healthy character of β_2AR has been achieved by Cav3 overexpression; this is in contrast to cells derived from the same animal without Cav3 overexpression.

E) A representative trace of the $\beta_2 AR$ response from a heart failure cell with Cav3 overexpressed.



Figure 5.

Illustration of the relationships represented in the computational model. Model components included $\beta_2 AR$ (red diamonds), Cav3 protein (black diamonds), mutant Cav3 protein (Cav3DN, yellow shaded diamonds), and fully formed caveolae (black crescents). cAMP emanating from discrete cav domains remained local, while that from ecav domains diffused freely. The cyt domain was passive, lacking β_2 ARs. In control (top middle), nearly all β_2ARs were in cav domains. In the crest all β_2ARs were caveolae associated. In T-tubules, 85% of β_2 ARs were cav3 associated (lone β_2 ARs were rare, 15%). Cav3 overexpression (top right) associated lone β_2 ARs with Cav3 in the T-tubules. Cav3DN (top left) replaced 75% of β_2 AR:Cav3 associations with β_2 AR:Cav3DN. M β CD (bottom left) removed all caveolae and Cav3, leaving only lone β_2 ARs. Heart failure (bottom middle) involved disassociation of β_2 AR:Cav3 pairs (75%) and T-tubule loss — represented by relocation of previously Ttubular β_2 AR:Cav3 pairs and lone β_2 ARs to the crest. Overexpression of Cav3 in the case of failure (bottom right) restored cav functionality to all lone β_2 ARs at the crest, and to 75% of lone β_2 ARs in remaining T-tubules. As depicted above the heart failure case, pipette stimulation at a T-tubule site captured many β_2 ARs down the entire length of a T-tubule; in contrast, at the flat crest surface, far fewer $\beta_2 ARs$ could be captured beneath the pipette and stimulated (1000-fold fewer than at a T-tubule).



Figure 6.

Simulations based on the relationships proposed in Figure 5 qualitatively reproduced cAMP experimental measurements. Shown are surface maps of cAMP concentration at the membrane ([cAMP], from 0.5 to 4 µM) along the cell length (100 microns) during the time just prior to and minutes following $\beta_2 AR$ stimulation. Stimulation was at distal T-tubule (panel A, top) or distal crest sites (panel B, bottom) — indicated in space and time by black rectangles labeled "B2AR stim.". The protocol was designed to mimic experiments shown in Figures 3C, D, E and 4D. Results for control, Cav3 overexpression, Cav3DN, M\betaCD, heart failure, and heart failure with Cav3 overexpression are shown. Cases are arranged as in Figure 5. A) Following Ttubular stimulation, cAMP was robustly generated at the stimulus site in all cases. However, it did not diffuse in control, Cav3, and failure+Cav3 cases (flat [cAMP] landscapes, "no diffusion" straight arrow). Pathological diffusion was exhibited by Cav3DN, MBCD, and heart failure (contoured [cAMP] landscapes, "diffusion" curved arrow). B) Crest stimulation did not always generate cAMP: e.g. the non-disease cases of control and Cav3, but also Cav3DN and failure+Cav3 cases. cAMP was generated following crest stimulation for M β CD and failure cases, and it diffused along the cell. Color scales are the same for all cases. Z-axes scales are the same for each case.