Adrenergic signaling regulates mitochondrial Ca2+ uptake through Pyk2-dependent tyrosine phosphorylation of the mitochondrial Ca2+ uniporter.

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Adrenergic Signaling Regulates Mitochondrial Ca\textsuperscript{2+} Uptake Through Pyk2-Dependent Tyrosine Phosphorylation of the Mitochondrial Ca\textsuperscript{2+} Uniporter

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Abstract

Aims: Mitochondrial Ca\textsuperscript{2+} homeostasis is crucial for balancing cell survival and death. The recent discovery of the molecular identity of the mitochondrial Ca\textsuperscript{2+} uniporter pore (MCU) opens new possibilities for applying genetic approaches to study mitochondrial Ca\textsuperscript{2+} regulation in various cell types, including cardiac myocytes. Basal tyrosine phosphorylation of MCU was reported from mass spectroscopy of human and mouse tissues, but the signaling pathways that regulate mitochondrial Ca\textsuperscript{2+} entry through posttranslational modifications of MCU are completely unknown. Therefore, we investigated \(\alpha_1\)-adrenergic-mediated signal transduction of MCU posttranslational modification and function in cardiac cells. Results: \(\alpha_1\)-adrenoceptor (\(\alpha_1\)-AR) signaling translocated activated proline-rich tyrosine kinase 2 (Pyk2) from the cytosol to mitochondrial matrix and accelerates mitochondrial Ca\textsuperscript{2+} uptake via Pyk2-dependent MCU phosphorylation and tetrameric MCU channel pore formation. Moreover, we found that \(\alpha_1\)-AR stimulation increases reactive oxygen species production at mitochondria, mitochondrial permeability transition pore activity, and initiates apoptotic signaling via Pyk2-dependent MCU activation and mitochondrial Ca\textsuperscript{2+} overload. Innovation: Our data indicate that inhibition of \(\alpha_1\)-AR-Pyk2-MCU signaling represents a potential novel therapeutic target to limit or prevent mitochondrial Ca\textsuperscript{2+} overload, oxidative stress, mitochondrial injury, and myocardial death during pathophysiological conditions, where chronic adrenergic stimulation is present. Conclusion: The \(\alpha_1\)-AR-Pyk2-dependent tyrosine phosphorylation of the MCU regulates mitochondrial Ca\textsuperscript{2+} entry and apoptosis in cardiac cells. Antioxid. Redox Signal. 21, 863–879.

Introduction

Mitochondrial Ca\textsuperscript{2+} homeostasis determines numerous cell functions, including energy metabolism, reactive oxygen species (ROS) generation, spatiotemporal dynamics of Ca\textsuperscript{2+} signaling, cell growth/development, and death (19, 24, 56, 57). The mitochondrial Ca\textsuperscript{2+} uniporter (mtCU), which is inhibited by lanthanides and ruthenium red (26), is the primary mechanism for mitochondrial Ca\textsuperscript{2+} influx in all cell types (8). Although functionally characterized several decades ago (18, 56), the complete molecular identity of the mtCU has yet to be fully elucidated. However, groundbreaking studies recently uncovered the molecular identity of the mtCU pore (MCU), the coiled-coil domain-containing protein 109A (\textit{CCDC109A}) (3, 14), as well as the regulatory proteins (56). These findings open up exciting new opportunities for using genetic approaches to elucidate mechanisms that regulate mitochondrial Ca\textsuperscript{2+} uptake in various cell types, including cardiac myocytes (17). For example, basal tyrosine phosphorylation of \textit{CCDC109A} was
Innovation

This report is the first to show the regulation of mitochondrial Ca^{2+} uptake, reactive oxygen species generation, and cell death signaling via mitochondrial Ca^{2+} uniporter pore (MCU) posttranslational modification. Our data provide significant and broad implications for understanding the regulation of MCU in cell signaling across all cell types, including cardiomyocytes. In addition, the results from this study suggest that inhibition of MCU tyrosine phosphorylation represents a potential, novel therapeutic target to prevent mitochondrial Ca^{2+} overload, oxidative stress, and mitochondrial/cell injury.

reported from mass spectroscopy analyses of human and mouse samples (see online database PhosphoSitePlus) (33). However, specific signaling pathways that control mitochondrial Ca^{2+} entry through posttranslational modifications of MCU are completely unknown.

In cardiac cells, adrenoceptor (AR) stimulation, either through β- or α_{1}-ARs, is a main determinant of physiological and pathophysiological cell signaling, predominantly through serine/threonine kinases [e.g., protein kinase A and Ca^{2+}/calmodulin-dependent protein kinase II (CaMKII)](55). In addition, we reported that protein tyrosine kinases (PTKs) are also activated during α_{1}-AR stimulation (α_{1}-ARS) in cardiac myocytes (54), and another report showed that a Ca^{2+}-dependent PTK named proline-rich tyrosine kinase 2 (Pyk2) is activated in response to α_{1}-ARS (27). Pyk2 is expressed abundantly in cancers (42, 42), and a recent study using human tissue showed that Pyk2 is also highly expressed in the heart and significantly activated during heart failure (40). Moreover, several reports indicate that Pyk2 is not only localized in the cytosol, but also in mitochondria (1, 23). Therefore, we hypothesize that α_{1}-AR-Pyk2 signaling regulates mitochondrial Ca^{2+} entry through MCU tyrosine phosphorylation in cardiac cells.

In this study, we report that Pyk2 activated by α_{1}-AR signaling directly phosphorylates MCU, which enhances mitochondrial Ca^{2+} uptake by promoting MCU channel oligomerization and formation of tetrameric channels. Moreover, we demonstrate that α_{1}-ARS and Pyk2 activation initiates ROS production in mitochondria (mitochondrial ROS: mROS), the activity of mitochondrial permeability transition pore (mPTP), apoptotic signaling, and cardiomyocyte death. This study is the first to show the modulation of MCU by adrenergic signaling and the regulation of ROS generation and cell death signaling via a posttranslational modification of MCU in cardiac cell.

Results

α_{1}-AR stimulation accelerates mitochondrial Ca^{2+} uptake

To explore whether adrenergic signaling regulates mitochondrial Ca^{2+} uptake in cardiac cells, we monitored changes in the mitochondrial matrix Ca^{2+} concentration ([Ca^{2+}]_{mt}) by expressing Mitycam, a mitochondria-targeted Ca^{2+}-sensitive inverse pericam (39, 53), in intact cardiac H9c2 myoblasts (Fig. 1). In this cell, Mitycam localized exclusively in mitochondria by cotransfecting with mitochondrial matrix-targeted RFP (53) (mt-RFP) (Fig. 1A, B). We observed the peak amplitude of the changes in Mitycam fluorescence (decrease in Mitycam fluorescence) to evaluate the magnitude of mitochondrial Ca^{2+} uptake (39, 53) (see also online Materials and Methods section). The changes in the cytosolic Ca^{2+} concentration ([Ca^{2+}]_{c}) were also monitored by Fura-red (53) (Fig. 1 and Supplementary Fig. S1; Supplementary Data are available online at www.liebertpub.com/ars). Mitycam responded to elevations in [Ca^{2+}]_{mt} in response to [Ca^{2+}]_{c} elevations induced by an inhibitor of the sarco/endoplasmic Ca^{2+}-ATPase (SERCA), thapsigargin (TG, 3 μM) (41) (Fig. 1D and Supplementary Fig. S2). The increase in [Ca^{2+}]_{mt} observed in response to an elevation in cytosolic Ca^{2+} induced by TG was almost abolished by expression of a dominant-negative pore-forming subunit of MCU (MCUB) (63), confirming the MCU dependence of this mitochondrial Ca^{2+} uptake (Fig. 1D–G). In addition, a significant increase in [Ca^{2+}]_{c} was detected in cells over-expressing MCUB due to reduced mitochondrial buffering.

FIG. 1. α_{1}-adrenoceptor (α_{1}-AR) stimulation accelerates mitochondrial Ca^{2+} uptake in H9c2 cells. (A) Expression of mitochondria-targeted Ca^{2+}-sensitive probe, Mitycam. Mitochondrial location was determined using mt-RFP. A cell stably transfected with only mt-RFP is also shown in the top row to confirm almost no background fluorescence in the GFP channel. Mitycam localized to mitochondria. Scale bars, 20 μm. (B) Color scatter plotting (right) and frequency scatter plotting (left) obtained from representative pictures in (A). (C) Time course of Mitycam fluorescence intensity during 3 μM thapsigargin (TG) treatment. Scale bar, 10 μm. (D) Left: Representative time courses of [Ca^{2+}]_{mt} (top) and [Ca^{2+}]_{c} (bottom) in cells stimulated by TG in control cells (CTR: nontransfected cells) (black), MCU (red), and MCUB (blue)-overexpressed cells. Arrow indicates TG application. [Ca^{2+}]_{mt} and [Ca^{2+}]_{c} were measured using Mitycam and Fura-Red, respectively. Mitycam fluorescence (F) was converted into 1 – (ΔF/ΔF_{o}), which shows the changes in [Ca^{2+}]_{mt}, where ΔF_{o} stands for initial fluorescence levels. Averaged data are shown in Supplementary Fig. S2. Right: summary data of left panels. [Ca^{2+}]_{mt} was measured at 60 s after stimulation, For [Ca^{2+}]_{c} peak [Ca^{2+}]_{c}, within 120 s was measured. *p < 0.05, compared to CTR. (E) Effect of Phe pretreatment on [Ca^{2+}]_{mt} and [Ca^{2+}]_{c}. Representative time course of [Ca^{2+}]_{mt} (top) and [Ca^{2+}]_{c} (bottom) after TG stimulation (arrow) in cells with (red) or without (black) pretreatment with Phe (100 μM, 15 min). [Ca^{2+}]_{mt} and [Ca^{2+}]_{c} were simultaneously measured using Mitycam and Fura-red, respectively, in a single cell (see Supplementary Materials and Methods section). Summary data are shown in Supplementary Fig. S5. (F) Left: Averaged time course of [Ca^{2+}]_{mt} after TG stimulation (arrow) in cells with (red) or without (black) pretreatment with Phe (100 μM, 15 min). [Ca^{2+}]_{mt} in cells over-expressing MCU-DN (blue) or MCUB (sky blue) is also shown for comparison. Right: Averaged time course of [Ca^{2+}]_{mt} after TG stimulation (arrow) in MCU-transfected cells pretreated with (red) or without (black) Phe. (G) Summary data of (F) at 60 s after stimulation. [Ca^{2+}]_{mt} in cell pretreatment with Phe and prazosin (Praz, 1 μM) (purple) is also shown (see also Supplementary Fig. S5). *p < 0.05, compared to CTR without Phe treatment. **p < 0.05, N.S., not significant. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
On the other hand, the increase in $[\text{Ca}^{2+}]_{\text{mt}}$ in response to TG was enhanced and a significant reduction in the $[\text{Ca}^{2+}]_c$ in response to TG was observed in MCU overexpressing cells (Fig. 1D–G and Supplementary Fig. S2). This greater mitochondrial response was not secondary to alterations of the cytosolic response, but a rather significant reduction in the $[\text{Ca}^{2+}]_c$ transient was observed, due to the increased $\text{Ca}^{2+}$ clearance by mitochondria, consistent with previous reports (3, 14, 17).

To test whether $\alpha_1$-ARS regulates mitochondrial $\text{Ca}^{2+}$ uptake, changes in $[\text{Ca}^{2+}]_{\text{mt}}$ and $[\text{Ca}^{2+}]_c$ induced by TG were simultaneously measured in cells pretreated for 15 min with the $\alpha_1$-AR agonist phenylephrine (Phe, 100 $\mu$M). In H9c2 cells, not like other Gq-protein-coupled receptors (GqPCR) expressed in this cell line (41, 53, 59), acute Phe treatment did not induce global cytosolic $\text{Ca}^{2+}$ elevation indicating a lack of $\text{Ca}^{2+}$ release from the sarco/endoplasmic reticulum (SR/ER) through inositol 1,4,5-trisphosphate (IP$_3$) receptor (Supplementary Fig. S1), possibly due to the lower endogenous expression levels of $\alpha_1$-AR and subsequent IP$_3$ generation upon stimulation (Supplementary Fig. S3) in this cell line. Phe was able to activate downstream signaling equivalent to other GqPCR agonists (Supplementary Fig. S4). TG induced a higher $[\text{Ca}^{2+}]_{\text{mt}}$ increase in Phe-pretreated cells compared with untreated cells (Fig. 1E, F). A significant reduction in the $[\text{Ca}^{2+}]_c$ transient was also observed, due to the increased $\text{Ca}^{2+}$ clearance by mitochondria (Fig. 1E and Supplementary Fig. S5). The increase in $[\text{Ca}^{2+}]_{\text{mt}}$ observed in Phe-pretreated cells was almost abolished in the presence of the $\alpha_1$-AR antagonist prazosin (1 $\mu$M), confirming that this
effect is mediated through α1-ARS (Fig. 1G and Supplementary Fig. S5). Interestingly, peak [Ca^{2+}]_mt in Phe-treated cells reached similar levels as that observed in MCU overexpressing cells without Phe pretreatment (Fig. 1F, G).

We also observed the concentration-dependent effect of Phe on [Ca^{2+}]_mt elevation. Half maximal effective concentration of Phe was \( \leq 2\) μM (Supplementary Fig. S5). In addition, to test whether β-AR stimulation, which is the major AR isoform in cardiomyocytes, regulates mitochondrial Ca^{2+} uptake, TG-induced [Ca^{2+}]_mt changes were measured in cells pretreated for 15 min with the β-AR agonist isoproterenol (Iso, 1 μM). In Iso-pretreated cells, the increase in [Ca^{2+}]_mt was similar to that in nontreated cells, indicating that this effect is specific to α1-AR signaling (Supplementary Fig. S5). In cells expressing MCUB (63) or a dominant-negative MCU mutant (MCU-DN) (14), TG-induced [Ca^{2+}]_mt uptake was significantly reduced and Phe pretreatment did not augment this uptake, demonstrating that Phe stimulation enhances MCU function (Fig. 1F–G). On the other hand, in MCU overexpressing cells, TG-induced [Ca^{2+}]_mt uptake reached higher levels following Phe pretreatment (Fig. 1F, G). We also confirmed that in native cultured rat neonatal cardiomyocytes (NCMs), α1-AR signaling (pretreatment of Phe or a physiological agonist, norepinephrine) can accelerate mitochondrial Ca^{2+} uptake (Supplementary Fig. S6). Thus, α1-ARS upregulates mitochondrial Ca^{2+} uptake by augmenting MCU function.

**α1-AR stimulation triggers activated Pyk2 translocation from cytosol to mitochondria and MCU tyrosine phosphorylation**

The next question is how α1-ARS can activate MCU function. Our hypothesis is that activated Pyk2 by α1-ARS phosphorylates MCU and accelerates mitochondrial Ca^{2+} uptake. To begin with, we first established a HEK293T cell line stably overexpressing MCU (HEK293T-MCU-Flag cells) to circumvent potential issues related to low endogenous expression of MCU in cultured cells (3, 14) (Fig. 2). We found that a small amount of endogenous Pyk2 was present in mitochondria-enriched fractions and α1-ARS increased the amount of mitochondrial Pyk2 (named mPyk2) (Fig. 2A). We quantitatively analyzed the ratio of cytosolic and mitochondrial Pyk2 and found that Pyk2 translocated from the cytosol to mitochondria upon Phe stimulation (Fig. 2A). In addition, the amount of activated Pyk2 (autophosphorylated Pyk2) was significantly increased in mitochondria-enriched fraction after α1-ARS, possibly due to the significant activated Pyk2 translocation from the cytosol to mitochondria (Fig. 2A, B). GPCR signaling, including α1-AR, increases ROS (20) and Pyk2 is known to be activated by ROS (22). Consistent with this mechanism, pretreatment with an antioxidant, N-acetylcysteine or a mitochondria-targeted antioxidant, mito-TEMPO [a specific scavenger of mitochondrial superoxide (mSO)] (73) significantly inhibited α1-AR-mediated activated Pyk2 translocation from cytosol to mitochondria (Fig. 2A, B and Supplementary Fig. S7). We next used the anti-phosphotyrosine (p-Tyr) antibody and observed tyrosine phosphorylation of MCU in mitochondria-enriched fraction after Phe stimulation. The intensity of a p-Tyr band identical to the molecular weight of MCU was significantly increased after Phe stimulation, indicating that tyrosine residue(s) in MCU might be phosphorylated after mPyk2 activation (Fig. 2B).

To investigate whether Pyk2 could directly phosphorylate MCU, we determined the submitochondrial localization of Pyk2 using Proteinase K (PK) digestion with a series of detergent concentrations to differentially permeabilize the outer mitochondrial membranes (OMM) and inner mitochondrial membranes (inner mitochondrial membranes).

**FIG. 2. α1-ARS stimulation induces activated Pyk2 translocation from cytosol in mitochondria and tyrosine phosphorylation of MCU in HEK293T cells.** (A) **Right:** Pyk2 activation in HEK293T cells stably overexpressing MCU-Flag after 100 μM Phe (15 min). HEK293T cells stably overexpressing MCU-Flag were treated by 100 μM Phe for 15 min and whole cell lysates (WCL) were prepared. Proteins were separated by SDS-PAGE (50 μg/well). Activated Pyk2 was detected using the anti-phospho-Tyr402-Pyk2 (p-Pyk2) antibody (see Supplementary Materials and Methods section). Summary data are shown at the bottom. *p < 0.05. **Left:** Pyk2 translocation into mitochondria in HEK293T cells stably overexpressing MCU-Flag after 100 μM Phe (15 min). HEK293T cells stably overexpressing MCU-Flag were treated by 100 μM Phe for 15 min in the presence of absence of N-acetylcysteine (NAC) and mitochondria-enriched fraction (Mito) and cytosolic fraction (including ER/SR) were prepared. Proteins were separated by SDS-PAGE (12.5 μg/well for each fractionated protein). Summary data are shown at the bottom. *p < 0.05. (B) Pyk2 activation and MCU phosphorylation in mitochondrial fractionation (50 μg/well). To increase the signals from P-Pyk2 and Pyk2, 50 μg protein of mitochondria-enriched fraction was loaded in each well. MCU phosphorylation was detected by anti-p-Tyr antibody. MCU phosphorylation was estimated by the p-Tyr bands identical to the molecular weight of MCU. *p < 0.05. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
membranes (IMM) (11). Proteins with known localization are immunoblotted and labeled according to their topology (Supplementary Fig. S8). Similar to the matrix protein cyclophilin D (Cyp-D), MCU is resistant to PK proteolysis even under high digitonin concentrations (up to 0.1%), consistent with the idea that MCU termini face to the matrix side. All these proteins, including Cyp-D and MCU, were substrates of PK as evidenced by significant digestion in the presence of Triton X-100. In addition, Pyk2 was also digested only with high digitonin, suggesting that endogenous mPyk2 primarily exists in the matrix and/or is bound to IMM from the matrix side at resting conditions. Interestingly, higher concentrations of digitonin were required to digest Pyk2 after Phe stimulation, suggesting that the amount of matrix mPyk2 increases after Phe stimulation (Supplementary Fig. S8).

Next, we confirmed that Pyk2 could specifically and directly bind to MCU and phosphorylate MCU by in vitro binding (72) and kinase assays (37) using purified MCU and Pyk2 proteins (Fig. 3A, B). The MCU-Pyk2 interaction and Pyk2-dependent phosphorylation of MCU were also observed in situ by immunoprecipitation (IP) in whole cell lysates (WCL) (Fig. 3C, D) or mitochondrial-enriched protein fractions (Fig. 3E). The MCU-Pyk2 interaction was enhanced after Phe stimulation concomitant with MCU phosphorylation (Fig. 3D, E). The MCU phosphorylation by Phe was inhibited by pretreatment with Mito-TEMPO, confirming that this mechanism is mROS dependent (Supplementary Fig. S7). Moreover, these effects were abolished by expression of a kinase-dead Pyk2 mutant (Pyk2-KD) (Supplementary Fig. S9) or transfection of siRNA targeted to Pyk2 (siRNA-Pyk2) (Fig. 3F, G).

Together, these results demonstrate that mPyk2 is activated by z1-AR-dependent and mROS-dependent signaling, which leads to mPyk2 binding to MCU and direct phosphorylation of MCU in the matrix.

Next, we tested whether this z1-AR-Pyk2-MCU signaling cascade also exists in H9c2 cells. Pyk2 activation, Pyk2 translocation into mitochondria, MCU-Pyk2 interaction, and Pyk2-dependent MCU phosphorylation were detected in this cell line (Fig. 4A–D), similar to those observed in HEK293T cells. We also observed Pyk2 translocation into mitochondria using confocal microscopy under Phe stimulation by cotransfection of GFP-Pyk2 and MCU-DsRed (Fig. 4E). Pyk2 localized in both the cytosol and mitochondria at baseline. After Phe stimulation, a more punctate pattern of GFP-Pyk2 localized in both the cytosol and mitochondria at baseline. Interestingly, higher concentrations of digitonin were required to digest Pyk2 after Phe stimulation, suggesting that the amount of matrix mPyk2 increases after Phe stimulation (Supplementary Fig. S8).

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Finally, we tested whether Pyk2 activation during z1-AR enhances mitochondrial Ca2+ uptake in H9c2 cells using the protocol shown in Figure 1. Pyk2 is known as a Ca2+- and ROS-dependent kinase, but 10 min of TG treatment did not activate Pyk2 in this cell line, confirming that our protocol for triggering mitochondrial Ca2+ accumulation itself did not significantly activate Pyk2 through increased [Ca2+]i (Supplementary Fig. S12). The increase in [Ca2+]i observed in Phe-pretreated cells was abolished in the presence of PF-431396, indicating that this effect is mediated by Pyk2 or FAK activity (Fig. 7A, right). The treatment with PF-431396 itself did not change the peak [Ca2+]i (Fig. 7A, left). Next, we transfected the plasmid containing siRNA-Pyk2 (67) into H9c2 cells (Fig. 7B and Supplementary Fig. S13). The increase in [Ca2+]i observed in Phe-pretreated cells was abolished in the cells transfected by siRNA-Pyk2, indicating
that Pyk2 is necessary for this effect (Fig. 7B). Collectively, these results show that α₁-AR-Pyk2-dependent MCU tyrosine phosphorylation promotes MCU oligomerization, leading to the facilitation of mitochondrial Ca²⁺ uptake through MCU.

α₁-AR-Pyk2 signaling triggers MCU-dependent ROS generation and proapoptotic protein release

Excessive mitochondrial Ca²⁺ uptake increases ROS production from mitochondria in cardiomyocytes (31). This suggests that the interaction between Pyk2 and MCU is crucial for regulating mitochondrial Ca²⁺ homeostasis and signaling pathways that may contribute to the development of cardiac dysfunction in various cardiac diseases.
addition, mitochondrial Ca\(^{2+}\) overload results in mitochondrial membrane potential (\(\Delta V_{\text{m}}\)) depolarization, ROS overproduction, and release of the proapoptotic proteins into the cytosol, ultimately resulting in cell injury and death. Our data clearly showed that \(\alpha_1\)-AR-Pyk2 signaling strongly elevates \([Ca^{2+}]_{\text{mt}}\) (Figs. 1–7) and that this increase is maintained at higher levels after \(\alpha_1\)-ARS. This sustained increase in mitochondrial Ca\(^{2+}\) could significantly enhance mROS production and apoptotic signals. Therefore, we tested whether \(\alpha_1\)-ARS increases the mitochondrial levels of ROS in H9c2 cells using the mSO indicator MitoSOX-Red (53) (Fig. 8).

Basal MitoSOX-Red intensity was significantly decreased by Mito-TEMPO pretreatment, confirming that MitoSOX-Red is sensitive to mSO (Fig. 8B). We found that Phe treatment significantly increased mSO levels and this effect was abolished by the pretreatment of Mito-TEMPO (Fig. 8A, B). Next, cells were pretreated by a potent FAK/Pyk2 inhibitor PF-431396. Pretreatment of PF-431396 dramatically blocked Phe-induced ROS increase (Fig. 8B). Moreover, we used the cells transfected with \(\text{siRNA-Pyk2}\) (Supplementary Fig. S13). Basal MitoSOX-Red intensity in cells transfected by \(\text{siRNA-Pyk2}\) was significantly lower than that in cells treated with control vector, indicating that Pyk2 is involved in the maintenance of basal mSO levels (Fig. 8B). Pyk2 knockdown significantly blocked the Phe-induced increase in MitoSOX-Red intensity, confirming that the increase in mSO levels by...
Phe was mediated through $\alpha_1$-AR-Pyk2 signaling. In addition, in cells overexpressing MCUB, MitoSOX-Red intensity in basal conditions significantly decreased compared with those in control cells and the Phe-induced increase in MitoSOX-Red intensity was completely abolished (Fig. 8B). These results indicate that Phe stimulates mSO levels via $\alpha_1$-AR-Pyk2-MCU signaling.

Because high levels of mROS trigger the release of mitochondrial intermembrane space proteins to the cytosol to initiate apoptosis through increasing the OMM permeability.

**FIG. 5.** $\alpha_1$-AR stimulation activates Pyk2 and phosphorylates MCU in native cardiomyocytes. (A) Neonatal cardiomyocytes (NCMs) were treated by 100 $\mu$M Phe (in the presence of 1 $\mu$M propranolol) for 15 min with or without Pyk2 inhibitor (PF) and WCL were prepared. *$p<0.05$ compared to the cells without stimulation (black). (B) Detection of MCU tyrosine phosphorylation after IP by anti-MCU antibody using WCL. MCU phosphorylation was detected by anti-P-Tyr antibody. $p<0.05$ compared to the cells without stimulation (black). (C) Top: Pyk2 translocation and MCU phosphorylation after Phe stimulation in NCMs. Bottom: Summary data. *$p<0.05$ compared to the cells without stimulation (black). (D) Top: Isolated adult cardiomyocytes (ACMs) were treated by 100 $\mu$M Phe (in the presence of 1 $\mu$M propranolol) for 30 min with or without $\alpha_1$-AR antagonist prazosin (Praz) or Pyk2 inhibitor (PF) and WCL were prepared. Bottom: Summary data. *$p<0.05$. (E) Detection of MCU tyrosine phosphorylation after IP by anti-MCU antibody using WCL. $p<0.05$ compared to the cells without stimulation (black). To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars

**FIG. 6.** $\alpha_1$-AR stimulation enhances the oligomerization of MCU via Pyk2 activation. (A) Representative Native PAGE of WCL from HEK293T cells stably overexpressing MCU-Flag treated by Phe. Membranes were blotted with anti-Flag antibody. MCU monomer bands in Native PAGE and SDS PAGE are shown with an arrow. MCU monomer bands in SDS PAGE are shown as loading control. A higher band ~140 kDa (smear distribution shown as a red square bracket), which is compatible with a tetramer, indicating that MCU monomers were oligomerized in higher order complexes in situ. (B) Mito-TEMPO pretreatment inhibits Phe-induced MCU oligomerization. MCU oligomerization (within a red square bracket) was quantified by normalizing with MCU monomer in SDS-PAGE (bottom). *$p<0.05$. N.S., not significant. (C) Pyk2 knockdown inhibits Phe-induced MCU oligomerization (see also Fig. 3F, G). MCU oligomerization (within a red square bracket) was quantified by normalizing with MCU monomer in SDS-PAGE (bottom). *$p<0.05$. N.S., not significant. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
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FIG. 7. \(\alpha_1\)-AR-Pyk2 signaling accelerates mitochondrial Ca\(^{2+}\) uptake. (A) Effect of Phe pretreatment on [Ca\(^{2+}\)]\(_{\text{mt}}\) and [Ca\(^{2+}\)]\(_{\text{c}}\) in the presence or absence of FAK/Pyk2 inhibitor (PF). [Ca\(^{2+}\)]\(_{\text{mt}}\) (right) and [Ca\(^{2+}\)]\(_{\text{c}}\) (left) after TG stimulation in cells with or without pretreatment of Phe and PF are shown. PF was applied before Phe stimulation (2 h before Phe application). [Ca\(^{2+}\)]\(_{\text{mt}}\) and [Ca\(^{2+}\)]\(_{\text{c}}\) were measured using Mitycam and Fura-red, respectively (see Supplementary Materials and Methods section). *p<0.05. (B) Effect of Phe pretreatment on [Ca\(^{2+}\)]\(_{\text{mt}}\) and [Ca\(^{2+}\)]\(_{\text{c}}\) in Pyk2-knockdown H9c2 cells. Knockdown efficiency is shown in Supplementary Fig. S13. [Ca\(^{2+}\)]\(_{\text{mt}}\) (right) and [Ca\(^{2+}\)]\(_{\text{c}}\) (left) after TG stimulation in cells with or without transfection of a plasmid containing siRNA-Pyk2 are shown. *p<0.05. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars

(e.g., by mPTP opening) (75), we determined whether \(\alpha_1\)-ARS promotes proapoptotic protein release. We used a fluorescence-tagged mitochondrial intermembrane space protein, GFP-tagged second mitochondrial-derived activator of caspase (Smac), to follow the time-dependent release of proapoptotic proteins during \(\alpha_1\)-ARS (Fig. 8C–E) (49). We analyzed changes in pixel intensity standard deviation (punctate/diffuse index) as an index of Smac-GFP release from mitochondria (49). The mitochondrial uncoupler carbonyl cyanide 4-(trifluoromethoxy)phenylhydrazone (FCCP) was used as a positive control to depolarize \(\Delta W_{\text{m}}\) to trigger mPTP opening (15). After FCCP treatment, Smac-GFP release into the cytosol (monitored by the decrease in punctate/diffuse index) was within 5 min (Fig. 8C, D). Over 15–20 min of continuous stimulation, high-dose Phe releases a small amount of Smac-GFP into the cytosol (Fig. 8C–E). Phe-induced Smac-GFP release was completely blocked by either the \(\alpha_1\)-AR antagonist prazosin, overexpression of MCU-DN, or addition of PF-431396 (2, 65) (Fig. 8D, E). In addition, the Phe-induced proapoptotic protein release was confirmed by detecting cytochrome C in enriched cytosolic and mitochondrial fractions (Fig. 8F). Furthermore, significant activation of caspase 3 was observed following prolonged Phe stimulation, consistent with the idea that \(\alpha_1\)-ARS initiated destructive apoptotic cascades following proapoptotic protein release (Fig. 8G). The results from Figure 8 indicate that Phe stimulation induces activation of apoptotic signaling cascades through the \(\alpha_1\)-AR-Pyk2-MCU-mROS signaling pathway (Fig. 9A, B).

Furthermore, we examined whether \(\alpha_1\)-AR-Pyk2-mROS signaling initiates destructive apoptotic cascades in ACMs. Significant caspase 3 activation was observed after 30 min of Phe stimulation, consistent with the results observed in H9c2 cells (Fig. 9C), but 30 min of Phe stimulation did not increase cardiomyocyte death (data not shown). Next, we observed the ACM viability after Phe stimulation up to 24 h of treatment by cell morphology (counting the number of rod-shaped cells) under a light microscope (Fig. 9D and Supplementary Fig. S14). We found that over 6 h of Phe stimulation significantly increased ACM death. Conversely, pretreatment with prazosin or PF-431396 inhibited the increase of mSOF induced by Phe, indicating that the Phe-induced mSOF generation is mediated through the \(\alpha_1\)-AR-Pyk2 signaling pathway (Fig. 9A, B).

We next determined whether this cell death signaling through the \(\alpha_1\)-AR-Pyk2-MCU-mROS cascade occurs in ACMs. We used mitochondria-targeted circularly permuted YFP (mt-YFP) to monitor mSO before and after Phe stimulation. This biosensor was recently reported from our groups, which can detect a stochastic and transient superoxide burst from either single or restricted clusters of interconnected mitochondria or isolated mitochondria (termed as a “mitochondrial superoxide flash” (mSOF)) (73). The mSOF generation is a result of a small increase in constitutive ROS production in mitochondria, which transiently opens a large channel of mPTP to evoke transient \(\Delta W_{\text{m}}\) depolarization, and subsequently stimulates the ETC to produce a burst in O\(_2^+\) production. Using this biosensor, we found that Phe significantly increased the frequency of mSOF in ACMs (Fig. 9A, B). Pretreatment with prazosin or PF-431396 inhibited the increase of mSOF induced by Phe, indicating that the Phe-induced mSOF generation is mediated by the \(\alpha_1\)-AR-Pyk2 signaling pathway (Fig. 9A, B).

\(\alpha_1\)-AR-Pyk2 signaling triggers mSOF generation and cell death in native cardiomyocytes

We next determined whether this cell death signaling through the \(\alpha_1\)-AR-Pyk2-MCU-mROS cascade occurs in ACMs. We used mitochondria-targeted circularly permuted YFP (mt-YFP) to monitor mSO before and after Phe stimulation. This biosensor was recently reported from our groups, which can detect a stochastic and transient superoxide burst from either single or restricted clusters of interconnected mitochondria or isolated mitochondria (termed as a “mitochondrial superoxide flash” (mSOF)) (73). The mSOF generation is a result of a small increase in constitutive ROS production in mitochondria, which transiently opens a large channel of mPTP to evoke transient \(\Delta W_{\text{m}}\) depolarization, and subsequently stimulates the ETC to produce a burst in O\(_2^+\) production. Using this biosensor, we found that Phe significantly increased the frequency of mSOF in ACMs (Fig. 9A, B). Pretreatment with prazosin or PF-431396 inhibited the increase of mSOF induced by Phe, indicating that the Phe-induced mSOF generation is mediated through the \(\alpha_1\)-AR-Pyk2 signaling pathway (Fig. 9A, B).

Furthermore, we examined whether \(\alpha_1\)-AR-Pyk2-mROS signaling initiates destructive apoptotic cascades in ACMs. Significant caspase 3 activation was observed after 30 min of Phe stimulation, consistent with the results observed in H9c2 cells (Fig. 9C), but 30 min of Phe stimulation did not increase cardiomyocyte death (data not shown). Next, we observed the ACM viability after Phe stimulation up to 24 h of treatment by cell morphology (counting the number of rod-shaped cells) under a light microscope (Fig. 9D and Supplementary Fig. S14). We found that over 6 h of Phe stimulation significantly increased ACM death. Conversely, pretreatment with prazosin or PF-431396 inhibited the increase of cell death induced by Phe. We also found that ACMs with abnormal morphology after Phe treatment had a higher activity of caspase-3/7, confirming that this Phe-induced cell death is mediated via the apoptotic cascade (Fig. 9E, F). The pretreatment of prazosin, PF-431396, or mito-TEMPO attenuated cell apoptosis induced by 6 h of Phe treatment. These data indicate that the \(\alpha_1\)-AR-Pyk2 signaling pathway induces cell death in ACMs possibly through an increase in mSO.
In this study, we identified a signaling pathway that regulates mitochondrial Ca\(^{2+}\) uptake, ROS production, and apoptotic signaling through a posttranslational modification of MCU (Fig. 10). Specifically, we found that Pyk2 activated downstream of \(\alpha_1\)-ARS, translocates into the mitochondria matrix, where it directly phosphorylates MCU tyrosine residue(s), which leads to an increase in the number of functional tetrameric MCU channels by promoting MCU oligomerization.
Moreover, this \( \alpha_1 \)-AR-Pyk2-MCU signaling cascade enhances mitochondrial \( Ca^{2+} \) uptake, mSO generation, and proapoptotic protein release, which initiates the apoptotic cascade and finally leads to cardiomyocyte death (Fig. 9). These results provide new insights into the molecular basis of adrenergic modulation of mitochondrial \( Ca^{2+} \) handling and apoptosis. Furthermore, the results of this study unveil a new pathway that could be targeted for the development of novel drugs to protect against the pathophysiological conditions, where increased \( \alpha_1 \)-ARS (36) and mitochondrial injury (25) are known to coexist.

**Molecular mechanism underlying enhanced mitochondrial \( Ca^{2+} \) uptake by \( \alpha_1 \)-AR-Pyk2-dependent phosphorylation of MCU**

We showed here that Pyk2 activation during \( \alpha_1 \)-ARS increases mitochondrial \( Ca^{2+} \) uptake through tyrosine phosphorylation of MCU, which enhances MCU oligomerization. This evidence strongly supports the hypothesis that Pyk2-mediated MCU oligomerization is directly linked to an increase in mitochondrial \( Ca^{2+} \) uptake following \( \alpha_1 \)-ARS. We also confirmed the MCU topology at IMM and detailed mPyk2 submitochondrial localization: (i) MCU termini are located at the matrix side as reported (9, 11, 46), (ii) mPyk2 is predominantly localized in the matrix, and (iii) Phe stimulation induces Pyk2 translocation from the cytosol to matrix. This observation suggests that Pyk2 can interact with MCU from the matrix side and directly phosphorylate the channel on either the N- or C-termini, which are known to be important for channel function (12). Recent broad screening of Pyk2-specific substrates using phosphoproteomics to identify consensus substrate motifs for Pyk2 has not yet been established (6). Human MCU contains 15 tyrosine residues; 5 in the N-terminus and 6 in the C-terminus, which are conserved across all eukaryotic species. No tyrosine residues are located in the pore-forming region (14). However, only three of these tyrosine residues (Y158 at N-terminus, Y289, and Y317 at C-terminus) are predicted to be potential PTK phosphorylation sites using the NetPhos 2.0 phosphorylation prediction program (5) and only one of these sites (Y289) is predicted to be a potential FAK/Pyk2-specific phosphorylation site using the Group-based Prediction System (74). Interestingly, basal

**FIG. 9.** \( \alpha_1 \)-AR-Pyk2 signaling accelerates mSO generation and initiates cell death in adult cardiomyocytes. (A) Representative images of mSO flashes (mSOF) observed in ACMs after Phe stimulation (10 min, 100 \( \mu \)M with 1 \( \mu \)M propranolol) by using mitochondrial-targeted cpYFP (see Supplementary Materials and Methods section). (B) Summary data of mSOF frequency. Pretreatment of prazosin (Praz) (30 min, 1 \( \mu \)M) or PF-431396 (PF) (2 h, 10 \( \mu \)M) efficiently inhibits the increase of mSOF induced by Phe. mSOF were collected from 18 cells isolated from three rats. *p<0.05. Scale bar, 10 \( \mu \)m. (C) Top: Caspase 3 activation by \( \alpha_1 \)-AR signaling. Cells were stimulated by Phe for 30 min and WCL were prepared to detect procaspase 3 and cleaved caspase 3. Caspase 3 activity was evaluated by the cleaved caspase 3/procaspase 3 ratio. Bottom: Summary data. (D) Top: Six-hour treatment with Phe increased ACM death. Cell variability was monitored by cardiomyocyte morphology. Pretreatment of Praz or PF efficiently inhibits the increase in cell death induced by Phe. Bottom: Summary data. *p<0.05. (E) Phe induces apoptosis in primary cultured cardiomyocytes. The pretreatment of Praz (30 min, 1 \( \mu \)M), PF (2 h, 10 \( \mu \)M), or Mito-TEMPO (1 h, 1 \( \mu \)M) attenuated cell apoptosis induced by 6-h Phe treatment. Apoptosis was detected by confocal (exitation/emission:488 nm/525 nm) in live cells after staining with a fluorogenic substrate for activated caspase-3/7 (CellEvent\textsuperscript{TM} Caspase-3/7 Green Detection Reagent) (see also Supplementary Materials and Methods section). Cell morphology was also observed under brightfield. As a positive control, cells were treated by staurosporine (Stauro, 1 \( \mu \)M, 6 h). Images obtained from cells without staining with the detection reagent were also shown as a negative control. Scale bars, 200 \( \mu \)m. (F) Summary data of (E). Images were randomly collected from 11–14 image fields from two rats. *p<0.05. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
Y289 phosphorylation was reported following mass spectrometry of human tissue (PhosphoSite) (32, 33). Future studies are needed to definitively determine Pyk2-specific phosphorylation site(s) in MCU and to test the functional importance of these sites using nonphosphorylation and phosphorylation mimetic MCU mutants. In addition, since the basal tyrosine phosphorylation of MCU was not completely abolished by Mito-TEMPO or siRNA-Pyk2 application (Supplementary Fig. S7), suggesting that not only Pyk2 but also other PTK in the mitochondrial matrix (e.g., Src) might regulate the basal tyrosine phosphorylation level of MCU in situ (28). Two potential CaMKII phosphorylation motifs in the N-terminus of MCU were identified using a nonphosphomutagenesis approach and mutation of these sites abolishes CaMKII-dependent activation of MCU current in mitoplast patch-clamp experiments (38). In this study, we also tested the effect of β-AR signaling on [Ca^{2+}]_{int}, which can activate a variety of serine/threonine kinase, including CaMKII, but this signaling did not significantly alter TG-induced [Ca^{2+}]_{int} elevation in H9c2 cells (Supplementary Fig. S5C). The amount of mitochondrial Ca^{2+} uptake is generally regulated by the magnitude of [Ca^{2+}]_{c} elevation in all cell types (56). Therefore, it is a reasonable idea that β-AR signaling has a more prominent effect on the amount of mitochondrial Ca^{2+} uptake due to its strong modulation of the cytosolic Ca^{2+} transient (CaT) size compared to α1-AR signaling in ACMs. Our observation of α1-AR-Pyk2-dependent MCU phosphorylation and its channel activation may serve as a supportive or additional mechanism for accelerating mitochondrial Ca^{2+} in response to the enhanced CaT under the physiological adrenergic stimulation. In addition, α1-AR might play more significant roles when β-AR signaling is downregulated, such as during heart failure. Future study will address the role of α1-AR-Pyk2-MCU signaling during beat-to-beat CaT and/or in the presence of main adrenergic signaling, β-AR stimulation in ACMs.

Our results also do not rule out the possibility that α1-AR-Pyk2-dependent tyrosine phosphorylation of MCU alters regulation of MCU by its auxiliary proteins (e.g., MICU1) (11), which may change its Ca^{2+} sensitivity and enhance the mitochondrial Ca^{2+} uptake rate. Indeed, Phe is capable of increasing both mROS production and OMM permeability in the absence of TG-induced global Ca^{2+} elevation (Figs. 8 and 9). These data suggest that there might be another mechanism for the enhancement of the mitochondrial Ca^{2+} clearance by α1-AR-Pyk2 signaling in addition to the MCU oligomerization. In cardiac cells, α1-AR increases the resting (diastolic) [Ca^{2+}]_{c} levels (34, 44, 61) and/or spontaneous Ca^{2+} release (45) although its effect to peak [Ca^{2+}]_{c} is relatively small. Therefore, after 15 min of Phe stimulation, α1-AR-Pyk2 signaling may change the Ca^{2+} sensitivity of MCU through channel tyrosine phosphorylation and mitochondria might accumulate Ca^{2+} more efficiently under the enhancement of spontaneous Ca^{2+} releases from ER/SR and/or the increase of resting [Ca^{2+}]_{c} levels by Phe. Our data using native PAGE strongly support the idea that the MCU channel is a tetramer (see also (63)) and that the tetrameric assembly is augmented by Pyk2-mediated tyrosine phosphorylation (18) (Fig. 6), although Baughman et al. reported a larger complex (3). The difference between these two studies may be due to the distinct solubilization methods used, such that the larger complex may also include additional mtCU auxiliary proteins, such as MICU1 (11). Further studies are needed to clarify whether tyrosine phosphorylation of MCU modulates the interaction of the channel pore with its regulatory proteins and changes its channel property such as Ca^{2+} sensitivity.

**Role of MCU in ROS generation and proapoptotic protein release**

In this study, we found that α1-AR-Pyk2-MCU signaling induced mROS generation (especially mSOF), proapoptotic protein release through mitochondrial Ca^{2+} overload, and

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**FIG. 10.** Working model: Cardiac α1-AR-Pyk2-dependent MCU phosphorylation initiates mitochondrial Ca^{2+} entry, mSOF generation, and cell death. To see this illustration in color, the reader is referred to the web version of this article at www.liebertpub.com/ars
ultimately, initiating a classic apoptotic signaling cascade and cell death. In addition, \( \alpha_1 \)-AR-Pyk2-MCU-mediated mROS generation and mitochondrial release of proapoptotic proteins were prevented by either a potent FAK/Pyk2 inhibitor or expression of MCU-DN/MCUB. These findings raise the distinct possibility of targeting the Pyk2 activity as a novel therapeutic strategy to prevent mitochondrial Ca\(^{2+} \) overload, ROS generation, and cell injury (see also next section).

The alterations in mitochondrial Ca\(^{2+} \) homeostasis are frequently observed under cardiac pathological conditions (15, 16, 25), where AR signaling also exists (36). Recently, \( G_{\alpha_i} \)PCR stimulation, including via \( \alpha_1 \)-AR, was shown to increase ROS in cardiomyocytes that activate ROS-sensitive downstream signaling processes (e.g., Pyk2 and CaMKII) (21, 27). However, the detailed mechanism by which \( \alpha_1 \)-ARS induces oxidative stress is unclear. In this study, we confirmed that \( \alpha_1 \)-ARS increases ROS levels, as previously reported in native cardiomyocytes (70). Moreover, we showed that the main source of ROS generation during Phe stimulation is the mitochondria and Pyk2-MCU signaling is critical for mSO generation. We also demonstrated that mROS increases the Pyk2 activity and mitochondrial localization (Supplementary Fig. S9). These data also indicate that persistent \( \alpha_1 \)-ARS activates a vicious cycle of ROS-induced ROS generation (75) as a result of ROS-sensitive activation of Pyk2 (Fig. 10).

By being the primary source of cellular ROS production, mitochondria play a crucial role for ROS signaling both in physiological and pathophysiological conditions (51, 66, 71). With sustained elevations in \([Ca^{2+}]_{\text{intr}}\), mROS generation and \( \Delta\Psi_{\text{mit}} \) depolarization trigger the mPTP opening that leads to apoptosis (7, 35). Thus, \([Ca^{2+}]_{\text{intr}}\) is commonly regarded as an important determinant in cell sensitivity to apoptotic stimuli in various cell types (64). Sites of intimate ER-mitochondria contact enable efficient local Ca\(^{2+} \) transmission between the two organelles, which serves as a potential regulatory site for apoptotic signaling (60), possibly via \([Ca^{2+}]_{\text{intr}}\)-ROS-mPTP pathways (see Figs. 8–10). We reported that the mitochondrial voltage-dependent anion channel 1 (VDAC1) selectively interacts with the IP\(_3\) receptor in the ER to direct the apoptotic Ca\(^{2+} \) release signals into mitochondria (13). Finally, we demonstrate here the direct involvement of Ca\(^{2+} \) influx through MCU in the regulation of mROS generation, apoptotic signaling, and cell death in native cardiomyocytes (Fig. 9).

In this study, a supraphysiologically concentration of an \( \alpha_1 \)-AR agonist Phe (100 \( \mu \)M) was used for activating the cell death cascade to mimic the pathophysiological conditions (e.g., chronic AR stimulation or a neurohumoral injury state). However, what is the physiological relevance of \( \alpha_1 \)-ARS-Pyk2-MCU signaling? We also tested the concentration-dependent effect of Phe on \([Ca^{2+}]_{\text{intr}}\) elevation and found that Phe had a significant effect for activating mitochondrial Ca\(^{2+} \) uptake through MCU in the range of the subphysiological concentration (EC\(_{50}\) = 2 \( \mu \)M) (Supplementary Fig. S5). We also showed that a significant reduction in the \([Ca^{2+}]_{\text{c}}\) transient was also observed, due to the increased Ca\(^{2+} \) clearance by mitochondria after \( \alpha_1 \)-ARS in H9C2 cells (Fig. 1E and Supplementary Fig. S5). These results are consistent with our previous observation that \( \alpha_1 \)-ARS decreases the Ca\(^{2+} \) transient by inhibiting (i) Ca\(^{2+} \) influx into the cell and (ii) by enhancing the Ca\(^{2+} \) clearance from the cytosol through PTK activity (53). Recently, we proposed that MCU may serve as an important mechanism for cytosolic Ca\(^{2+} \) buffering when \([Ca^{2+}]_{\text{c}}\) increases at the ER/SR contact sites (17, 54). Therefore, physiological \( \alpha_1 \)-ARS may regulate/enhance the Ca\(^{2+} \) uptake by mitochondria, which may promote mitochondrial function and/or more efficient buffering of cytosolic Ca\(^{2+} \). However, it is still controversial whether mitochondrial Ca\(^{2+} \) uptake contributes to the kinetics of beat-to-beat Ca\(^{2+}\)T formation in ACMs although mitochondria occupy 35% of cytosolic space and are well known to uptake Ca\(^{2+} \) (62). Shannon et al. estimated that the mitochondria contribute to only ~1% of the total Ca\(^{2+} \) removal from the cytosol during Ca\(^{2+}\)T; SERCA and sarcoplasmic Na\(^{+}\)/Ca\(^{2+}\) exchanger (NCX) are responsible for almost all Ca\(^{2+}\) removal from the cytosol (69). Further studies are required for quantitatively evaluating the contribution of the \( \alpha_1 \)-AR-Pyk2-MCU signaling pathway for enhancing the mitochondrial Ca\(^{2+}\) buffering capacity during beat-to-beat Ca\(^{2+}\)T under AR stimulation.

**\( \alpha_1 \)-AR-Pyk2 signaling as a novel therapeutic target for preventing mitochondrial Ca\(^{2+} \) overload, ROS generation, and cardiac injury**

Pyk2 is abundantly found in various types of cancers and is well established as a key regulator of cancer proliferation, migration, and invasion (42, 42). Therefore, Pyk2 (or FAK) is a potent therapeutic target for cancer treatment, and currently, several FAK/Pyk2 inhibitors are already being evaluated in clinical trials (68). In cardiomyocytes, Pyk2 regulates the transcription factors that lead to cardiac remodeling, including hypertrophy (4, 29, 30). Therefore, Pyk2 is well studied as a component of nuclear signal transduction in cancer biology and cardiac remodeling. However, essentially, no information is available related to the effect of Pyk2 on mitochondrial functions. Interestingly, Pyk2 contains a predicted (using Mitoprot (10)) N-terminus mitochondria-targeting sequence (MSGVSEPLSRVKLGTLRRP), and a few reports have shown Pyk2 to be localized in mitochondria (1, 23). However, the functional role of mPyk2 has not been established. In this study, we confirmed that Pyk2 exists in the mitochondrial matrix in cardiac cells and found that \( \alpha_1 \)-ARS leads to activated Pyk2 translocation from the cytosol to mitochondrial matrix. Moreover, we showed that mPyk2 regulates MCU and initiates proapoptotic protein release from mitochondria. Although potent FAK/Pyk2 inhibitors (e.g., PF-431396 and PF-562271) are widely used in the field of cancer research, there are no reports regarding the effects of these drugs in heart disease and mitochondrial injury. Our results suggest that the small-molecule Pyk2/FAK inhibitors or Pyk2 knockdown, may prevent mitochondrial Ca\(^{2+} \) overload, mROS generation, proapoptotic protein release, and cell death in the cardiomyocytes during sustained \( \alpha_1 \)-ARS or in the specific pathophysiological conditions where \([Ca^{2+}]_{\text{c}}\) overload and ROS overproduction coexist (e.g., ischemic–reperfusion).

Recently, Pyk2 was shown to be activated in human no-ischemic heart failure (40). However, in heart failure, it has been well reported that there is a significant elevation of cytosolic Na\(^{+}\)/Ca\(^{2+}\) concentration \([Na^{+}]_{\text{c}}\), which enhanced cytosolic Ca\(^{2+}\) extrusion from the cell during diastole through NCX and reduced the \([Ca^{2+}]_{\text{c}}\) transient (52). In
addition, the [Na\(^{2+}\)]\(_e\) elevation and mitochondrial NCX activity in heart failure dictate mROS generation through a decrease in the mitochondrial Ca\(^{2+}\) uptake (50). Indeed, several studies showed that mitochondrial Ca\(^{2+}\) uptake is reduced in failing cardiomyocytes (43, 47). These reports demonstrated that elevated [Na\(^{2+}\)]\(_e\) and mitochondrial NCX activity also critically contribute to decreased [Ca\(^{2+}\)]\(_{\text{int}}\), especially during heart failure. Since we did not observe the detailed relationship between the Pyk2-dependent MCU regulation and [Na\(^{2+}\)]\(_e\), or resting [Ca\(^{2+}\)]\(_e\), in this study, it is still not clear whether activated Pyk2 during heart failure has a significant impact on the MCU function.

**Limitation of this study**

In this study, most of the experiments were performed in cultured cell lines. While important experiments validate using native cardiomyocytes, we still need to take into account that our finding cannot be directly applicable to the in vivo situation. Animal models such as the neurohumoral injury model using the z\(_1\)-AR agonist will be indispensable for exploring the role of the cardiac z\(_1\)-AR-Pyk2-MCU signaling pathway in physiological and pathophysiological conditions in vivo as well as validating the Pyk2 inhibition as a possible novel therapeutic strategy for preventing mitochondrial injury and cell death.

**Conclusion**

This study is the first to investigate the regulation of MCU by cardiac adrenergic signaling. Elucidation of these regulatory mechanisms may lead to the design of novel targets for the pharmacological management of heart failure, where chronic adrenergic stimulation, mitochondrial injury, oxidative stress, and myocardial death are present.

**Materials and Methods**

An expanded fully described Materials and Methods section is available in Supplementary Data.

**Cell cultures**

H9c2 cells and HEK293T cells were maintained, transfected with plasmids, and used for experiments (37, 53, 54). HEK293T-MCU-Flag cells were generated by transfecting with pcDNA3.1(+) -MCU-Flag and maintained in the presence of G-418 (53). NCMs and ACMs were isolated from Sprague-Dawley rats, plated, cultured, and used for experiments (37). NCMS were transfected with plasmids and ACMs were infected with adenovirus (17, 37). All animal experiments were performed in accordance with the Guideline on Animal Experimentation of Thomas Jefferson University (TJU). The study protocol was approved by the Animal Care Committee of TJU. The investigation conformed to the Guideline for the Care and Use of Laboratory Animals published by the US National Institutes of Health (NIH).

**Mitochondrial protein isolation, Western immunoblot analysis, in vitro kinase, and binding assay**

Whole cell lysates and mitochondria-enriched and cytosolic proteins were prepared (53). For electrophoresis under nondissociating conditions, mitochondrial proteins or whole cell lysates were separated by 10% polyacrylamide gel (Native-PAGE) (63). IP (37), in vitro binding (72), and kinase assays (37) were performed as described previously.

**Live cell imaging in confocal microscopy and image analysis**

The [Ca\(^{2+}\)]\(_{\text{int}}\) measurement using mitochondria-targeted Ca\(^{2+}\) biosensor Mitycam (39, 53), [Ca\(^{2+}\)]\(_e\) measurement using Fura-Red (53), FRET measurement (63), mSO measurement by MitoSOX-Red (53), visualizing the release of mitochondrial intermembrane space proteins (49), measurement of mSOF by mT-cyFP (73), detection of caspases 3/7 activity (48) were performed in live cells using the laser scanning confocal microscope (Olympus, Tokyo, Japan). There were no significant changes in the initial Mitycam fluorescence levels between the different experimental settings used in this study (see Supplementary Fig. S15). All images were analyzed with ImageJ software (NIH).

**Data and statistical analysis**

All results are shown as mean standard error. Unpaired Student’s t-test was performed for two data sets. For multiple comparisons, one-way ANOVA followed by the post hoc Tukey test was performed. Statistical significance was set as a p value of <0.05.

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**Author Disclosure Statement**

No competing financial interests exist.

**References**

ADRENERGIC REGULATION OF MITOCHONDRIAL Ca2+ UNIPORTER


ADRENERGIC REGULATION OF MITOCHONDRIAL Ca\textsuperscript{2+} UNIPORTER


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**Abbreviations Used**

- $\alpha_1$-AR = $\alpha_1$-adrenoceptor
- $\alpha_1$-ARS = $\alpha_1$-adrenoceptor stimulation
- $\Delta W_m$ = mitochondrial membrane potential
- ACM = isolated adult cardiomyocytes
- $[Ca^{2+}]_c$ = cytosolic Ca$^{2+}$ concentration
- $[Ca^{2+}]_{mt}$ = mitochondrial matrix Ca$^{2+}$ concentration
- CaMKII = Ca$^{2+}$/calmodulin-dependent protein kinase II
- CCDC109A = coiled-coil domain-containing protein 109A
- Cyp-D = cyclophilin D
- DN = dominant negative
- ER = endoplasmic reticulum
- FAK = focal adhesion kinase
- FCCP = carbonyl cyanide 4-(trifluoromethoxy) phenylhydrazone
- FRET = Förster resonance energy transfer
- GqPCR = G$\text{q}$-protein coupling receptor
- HEK293T-MCU-Flag cells = HEK293T cells stably overexpressing MCU-Flag
- IMM = inner mitochondrial membranes
- IP = immunoprecipitation
- KD = kinase dead
- MCU = mitochondrial Ca$^{2+}$ uniporter pore
- mPTP = mitochondrial permeability transition pore
- mPyk2 = mitochondrial Pyk2
- mROS = ROS production in mitochondria
- mSO = mitochondrial superoxide
- mSOF = mitochondrial superoxide flashes
- mitochondrial Ca$^{2+}$ uniporter
- mt-RFP = mitochondria-targeted RFP
- NCM = neonatal cardiomyocyte
- OMM = outer mitochondrial membranes
- Phe = phenylephrine
- PK = Proteinase K
- P-Tyr = phosphotyrosine
- Pyk2 = proline-rich tyrosine kinase 2
- ROS = reactive oxygen species
- SERCA = sarcoendoplasmic reticulum/endoplasmic reticulum Ca$^{2+}$-ATPase
- Smac = second mitochondrial-derived activator of caspases
- SR = sarcoplasmic reticulum
- TG = thapsigargin
- VDAC = voltage-dependent anion channel
- WCL = whole cell lysates