Congestive heart failure typically arises from cardiac myocyte necrosis/apoptosis, associated with the pathological opening of the mitochondrial permeability transition pore (mPTP). mPTP opening decreases the mitochondrial membrane potential leading to the activation of Ca2+-independent phospholipase A2γ (iPLA2γ) and the production of downstream toxic metabolites. However, the array of enzymatic mediators and the exact chemical mechanisms responsible for modulating myocardial mPTP opening remain unclear. Herein, we demonstrate that human heart failure activates specific myocardial mitochondrial phospholipases that increase Ca2+-dependent production of toxic hydroxyeicosatetraenoic acids (HETEs) and attenuate the activity of phospholipases that promote the synthesis of protective epoxyeicosatrienoic acids (EETs). Mechanistically, HETEs activated the Ca2+-induced opening of the mPTP in failing human myocardium, and the highly selective pharmacological blockade of either iPLA2γ or lipoxigenases attenuated mPTP opening in failing hearts. In contrast, pharmacological inhibition of cytochrome P450 epoxygenases opened the myocardial mPTP in human heart mitochondria. Remarkably, the major mitochondrial phospholipase responsible for Ca2+-activated release of arachidonic acid (AA) in mitochondria from non-failing hearts was calcium-dependent phospholipase A2ζ (cPLA2ζ) identified by sequential column chromatographies and activity-based protein profiling. In contrast, iPLA2γ predominated in failing human myocardium. Stable isotope kinetics revealed that in non-failing human hearts, cPLA2ζ metabolically channels arachidonic acid into EETs, whereas in failing hearts, increased iPLA2γ activity channels AA into toxic HETEs. These results mechanistically identify the sequelae of pathological remodeling of human mitochondrial phospholipases in failing myocardium. This remodeling metabolically channels AA into toxic HETEs promoting mPTP opening, which induces necrosis/apoptosis leading to further progression of heart failure.

Congestive heart failure is a public health crisis of epidemic proportions that typically results from cardiac myocyte necrosis/apoptosis leading to progressive hemodynamic compromise (1–3). In animal models, it is generally believed that cardiac myocyte cell death and heart failure are intimately associated with the pathological opening of the mitochondrial permeability transition pore (mPTP) (3–5). Opening of this pore dissipates the mitochondrial membrane potential and facilitates the production of toxic metabolites, the activation of membrane potential-sensitive calcium-independent phospholipase A2γ (iPLA2γ) (6), and the release/activation of proteins mediating cell death-signaling programs (e.g. cytochrome c). The resultant dropout of cardiac myocytes during heart failure by either apoptosis or necrosis leads to the progression of heart failure.

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This article was selected as one of our Editors’ Picks.

This article contains Figs. S1–S5.

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7 The abbreviations used are: mPTP, mitochondrial permeability transition pore; AA, arachidonic acid; HETE, hydroxyeicosatetraenoic acid; EET, epoxyeicosatrienoic acid; PGE2, prostaglandin E2; TXB2, thromboxane B2; MAFP, methyl arachidonyl fluorophosphonate; iPLA2γ, calcium-independent phospholipase A2γ; cPLA2ζ, calcium-dependent phospholipase A2ζ; LOX, lipoxigenase; CYP450, cytochrome P450; HRAM, high resolution accurate mass; ESI, electrospray ionization; BEL, bromoenol lactone; bis-Tris, 2-[bis(2-hydroxyethyl)aminio]-2-(hydroxymethyl)propan-1,3-diol; AACC0F, arachidonoyl trifluoromethyl ketone; Ibu, ibuprofen; Baic, baicalein; MSPPOH, N-(methylsulfonoyl)-2-(2-propyloxy)-benzenehexanamide; LPC, lysophosphatidylcholine; CyPd, cyclophilin D; [14C]PAPC, palmitoyl-2-arachidonoyl-sn-glycero-3-phosphocholine; SR, sarcoplasmic reticulum; AMPP, N-(4-aminomethylphenyl)pyridinium; Pyr, pyridoline; VAD, ventricular assist device.
Ca²⁺-activated phospholipases A₂ in human heart mitochondria

failure, further hemodynamic compromise, and ultimately death (7, 8). However, the precise chemical mechanisms, enzymatic mediators, and molecular events responsible for modulating the amount and duration of mPTP opening in both animal and human myocardium remain largely undefined.

Previous studies have demonstrated that human myocardium contains multiple different phospholipase activities encoded by distinct genes (9–11). Recently, we demonstrated that genetic knock-out of the active site of iPLA₂γ (PNPLA8) results in the attenuation of mPTP opening and that iPLA₂γ is largely responsible for the calcium-dependent production of oxidized fatty acid metabolites (e.g. eicosanoids, docosanoids, and oxidized linoleic acid metabolites) emanating from the mitochondrial compartment in murine hepatic and myocardial tissues (12–14). Moreover, loss of mitochondrial transmembrane potential evoked by either inhibition of mitochondrial electron transport chain complexes or prolonged calcium transients results in the further activation of iPLA₂γ (6). Collectively, this sequence of events results in a feed-forward loop promoting the autoamplification of mitochondrial lipid second messenger generation. Mitochondria contain, or are associated with, multiple different types of phospholipases in humans. However, their molecular identity, their contributions to production of signaling lipids, and the effects of the resultant downstream metabolites in non-failing versus failing human myocardium are unknown.

During heart failure in animal models, increases in mitochondrial calcium concentration and changes in calcium dynamics are present resulting from alterations in sarcoplasmic reticulum (SR)-mitochondrial subcellular architecture, pathological SR calcium coupling, and reduced Ca²⁺ sequestration (15, 16). Because phospholipase-catalyzed arachidonic acid (AA) release is the rate-limiting step in the generation of eicosanoid-derived signaling moieties, knowledge of the types of mitochondrial phospholipases, Ca²⁺-mediated changes in their activities, and metabolic fates of their downstream metabolites are essential in the mechanistic understanding of the pathology resulting from the diverse array of toxic oxidized lipids produced during human heart failure.

Recent studies have demonstrated that specific eicosanoids may confer either protective or deleterious effects on myocardial function (17–19). For example, increased levels of 14,15-epoxyeicosatrienoic acid (14,15-EET) in murine myocardium produced by either overexpression of the cytochrome P450 enzyme CYP2J3 or by genetic knock-out of soluble epoxide hydrolase resulted in decreased mPTP opening probability, improved preservation of mitochondrial membrane potential (ΔΨₘ), and attenuated damage after ischemia/reperfusion (17, 20, 21). The diverse effects of prostaglandin E₂ (PGE₂), a cyclooxygenase-2 product, on cardiac function have been documented, which include the reduction of cardiac ischemia/reperfusion injury and a contribution to myocardial hypertrophy via prostaglandin E₂ receptor signaling (22, 23). In contrast, 12-HETE and 20-HETE induce increases in mitochondrial calcium concentration and myocardial infarct size during ischemia/reperfusion in mice (24–26).

Previously, we identified the prominent production of eicosanoids (including HETEs, EETs, and prostaglandins) in calcium-activated murine heart tissue and mitochondria that were markedly reduced by genetic ablation of iPLA₂γ (13, 27). Accordingly, we hypothesized that abrupt activation of mitochondrial phospholipases by calcium uptake into the mitochondria in conjunction with the loss of membrane potential would result in the pathological activation of iPLA₂γ leading to the generation of a complex array of eicosanoid metabolites and fatty acids that contribute to the opening of the mPTP and progression of human heart failure. Here, we demonstrate the dramatic increase in calcium-activated HETE production at the expense of cardioprotective EETs in failing human hearts in comparison with non-failing human myocardium. These changes collectively result in the increased sensitivity of mitochondria to calcium-induced mPTP opening and further depolarization leading to cardiac myocyte damage. Furthermore, we present primary evidence of metabolic channeling of AA into discrete eicosanoid products through the remodeling of mitochondrial phospholipase activities from cPLA₂ζ in non-failing human myocardium to iPLA₂γ during heart failure. Collectively, this study mechanistically identifies the pathological remodeling of cardiac phospholipases and their downstream products resulting from altered metabolic channeling as a principal mechanism increasing lipid-mediated mPTP opening leading to mitochondrial dysfunction and the progression of human heart failure.

Results
Remodeling of human mitochondrial eicosanoid production in non-failing versus failing human myocardium

To assess whether alterations in eicosanoid molecular species are present in failing versus non-failing human heart mitochondria, we incubated mitochondria from non-failing or failing human myocardium in the absence or presence of calcium ion as described under “Experimental procedures.” Reaction products were identified and quantified using charge-switch derivatization with LC-MS/MS (14, 28) and selected reaction monitoring with HRAM (high resolution accurate mass) mass spectrometry of product ions as reported previously (14). Intriguingly, mitochondria (and their tethered/associated membranes) prepared from non-failing human myocardium exhibited a substantial increase in the initial rate of production of a wide range of eicosanoid molecular species after calcium challenge, including cytoprotective EETs (Fig. 1, A–D). In sharp contrast, similar calcium challenge of mitochondria from failing human myocardium produced robust increases in multiple lipoxygenase metabolites, including 5-, 8-, 11-, 12-, and 15-HETEs. Critically, in failing myocardium, calcium stimulation did not result in significant increases in salutary 14,15-EET production (Fig. 1, A and B), although the enzymes to catalyze the oxidation of released AA to 14,15-EET were clearly present in failing myocardium (see below). Moreover, large Ca²⁺-dependent increases in proinflammatory PGE₂ were manifest in conjunction with decreases in cardioprotective prostacyclins (29–31) (measured as their stable end products, 6-keto-PGF₁α) in mitochondria from failing hearts.
Ca\(^{2+}\)-activated phospholipases A\(_2\) in human heart mitochondria

Failing human myocardium is more susceptible to calcium-induced opening of mPTP in comparison with non-failing human myocardium

Despite the importance of the mPTP channel in mediating cardiac myocyte cell death programs, to the best of our knowledge, no previous investigations have explored alterations in cardiac myocyte cell death programs, to the best of our knowledge. Accordingly, we incubated isolated human heart mitochondria with calcium ion and measured mitochondrial swelling indicative of mPTP opening. Although high-resolution respirometry revealed that oxidative phosphorylation of mitochondria isolated from non-failing and failing hearts was not significantly different (Fig. S1), calcium challenge of mitochondria dramatically increased swelling of mitochondria from failing hearts in comparison with non-failing hearts (Fig. 2A). Inclusion of inorganic phosphate (Pi), a potent inducer of mPTP opening in animal models, only moderately increased swelling of human myocardial mitochondria without affecting the relative sensitivity of non-failing versus failing heart mitochondria to calcium-induced mPTP opening (Fig. 2B). The expression levels of known transition pore regulators (32, 33) (e.g. adenine nucleotide translocase (ANT), voltage-dependent anion channel (VDAC), and cyclophilin D (CypD)) were not different in non-failing versus failing hearts as assessed by Western blot analyses (Fig. 2C). Thus, the observed susceptibility of failing heart mitochondria to calcium-induced swelling was not due to alterations in the content of pore-regulating proteins, but rather to their maladaptive regulation by different modulators in non-failing versus failing human hearts.

HETEs activate the calcium-induced mitochondrial permeability transition pore opening

To determine whether the dramatic increase in HETE production was mechanistically responsible for the increase in sensitivity to mitochondrial swelling, mitochondria from failing human myocardium were treated with specific inhibitors of either cyclooxygenases (COX), lipoxygenases (LOX), or cytochrome P450 monoxygenases (CYP450). Remarkably, inhibition of LOX activity with baicalein (Baic) completely abolished calcium-activated mPTP opening of failing heart mitochondria, whereas CYP450 inhibition with MSPPOH dramatically exacerbated swelling (Fig. 3A). Because MSPPOH-enhanced

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**Figure 1.** Mass spectrometric analysis and quantitation of eicosanoids generated by non-failing and failing human heart mitochondria stimulated by calcium. Myocardial mitochondria were isolated from non-failing (control) and failing hearts by differential centrifugation as described under “Experimental procedures” and then sonicated in HEPES buffer (10 mM HEPES (pH 7.4) containing 10% glycerol and 1 mM DTT). Mitochondrial homogenates were incubated immediately isolated by solid-phase extraction, derivatized with AMPP, and analyzed by HRAM mass spectrometry. The resultant production of HETEs (A), EETs (B), and Prostaglandins (PGs) (C) is shown with the average ± S.E. (n = 6 for non-failing control hearts and n = 6 for failing hearts). *, p < 0.05, and **, p < 0.005 for control (non-failing) versus failing hearts. †, p < 0.005, and ‡, p < 0.005 for control (non-failing) versus failing hearts. *p < 0.05, and **p < 0.005 for control (non-failing) versus failing hearts.

**Figure 2.** Inclusion of inorganic phosphate (Pi), a potent inducer of mPTP opening in animal models, only moderately increased swelling of human myocardial mitochondria without affecting the relative sensitivity of non-failing versus failing heart mitochondria to calcium-induced mPTP opening (Fig. 2B). The expression levels of known transition pore regulators (32, 33) (e.g. adenine nucleotide translocase (ANT), voltage-dependent anion channel (VDAC), and cyclophilin D (CypD)) were not different in non-failing versus failing hearts as assessed by Western blot analyses (Fig. 2C). Thus, the observed susceptibility of failing heart mitochondria to calcium-induced swelling was not due to alterations in the content of pore-regulating proteins, but rather to their maladaptive regulation by different modulators in non-failing versus failing human hearts.
mitochondrial swelling appears inhibitable with cyclosporine A, inhibition of CYP450 likely facilitate cyclophilin D-dependent mPTP opening (Fig. 3B). Inhibition of COX enzymes with ibuprofen (Ibu) did not affect human failing heart mitochondrial swelling. Similarly, mPTP opening of non-failing heart mitochondria was dramatically promoted by preincubation with MSPPOH, although the induction time for mitochondrial swelling was significantly longer than that of failing heart (Fig. 3C). These distinct effects resulting from selective inhibition of discrete oxygenases demonstrate the prominent role of signal-ling eicosanoids in the enhancement or attenuation of mPTP opening in failing human myocardium. Mitochondrial swelling data with oxygenase-specific inhibitors are consistent with the HRAM mass spectrometric analyses of mitochondria dem-onstrating higher HETE levels in mitochondria from failing hearts accompanied by decreased calcium-stimulated EET produc-
tion in comparison with non-failing hearts (Fig. 1). Importantly, exogenous provision of selected HETEs reversed the protection from mPTP opening conveyed by baicalein treatment (Fig. 3D). Although multiple HETEs were promoters of mitochondrial swelling (Fig. 3D), the lipoxigenase products 12-HETE and 15-HETE are likely of greater pathophysiological significance because they are prominent in human myocardium, and Ca\(^{2+}\) stimulation generated substantially greater amounts of these eicosanoids in failing human heart mitochondria (Fig. 1). This notion is further supported by the observation of the inability of CYP450 hydroxylase(s) to generate effective amounts of 20-HETE after calcium stimulation in either non-failing or failing human myocardial mitochondria (Fig. 1).

**Oxidation of exogenously added deuterated arachidonic acid versus endogenously produced arachidonic acid in mitochondria from non-failing and failing myocardum**

To determine whether alterations in the enzymic activities of AA oxidases in non-failing versus failing heart mitochondria were responsible for the differences in the profiles of eicosanoids products, we compared their rate of production using deuterated isotopes of arachidonic acid examined under initial rate conditions. Mitochondrial sonicates from non-failing or failing human hearts were incubated with exogenous deuterated arachidonic acid (d\(_6\)-AA) in the absence or presence of Ca\(^{2+}\). The results demonstrated minimal differences in deuterated oxylipin production using d\(_6\)-AA in the presence of EGTA or Ca\(^{2+}\) treatments in either non-failing and failing myocardial mitochondria (Fig. 4). Moreover, these results demonstrate that AA availability to the downstream oxidative enzymes was rate-determining in the synthesis of the measured eicosanoids because if exogenously added d\(_6\)-AA was present, no substantive differences in deuterated eicosanoid production in non-failing versus failing hearts occurred. In stark contrast, non-labeled HETE (H−) production from endogenous AA in failing heart mitochondria, even in the presence of exogenous d\(_6\)-AA, was markedly higher than that in non-failing heart mitochondria (Fig. 4A). The production of virtually all non-labeled salu-
tary EETs in failing human heart mitochondria was drastically reduced in comparison with non-failing heart mitochondria. However, substantive amounts of d\(_6\)-labeled EETs (especially 14,15-EET) were readily produced when exogenous d\(_6\)-AA was present. These results demonstrate that the enzymes for EET production were present in failing myocardium, but the endogenously generated AA upon calcium challenge could not be metabolically channeled to EET production (Fig. 4B). Cyclooxygenase-mediated prostaglandin E\(_2\) (PGE\(_2\)) production using endogenous AA was Ca\(^{2+}\)-dependent, and the rate of production of both thromboxane and prostacyclin was dramatically reduced in mitochondria from failing hearts (Fig. 4C). Mitochondrial preparations from failing human hearts possess less capacity for the production of prostacyclins (e.g. 6-keto-PGF\(_{1\alpha}\)), even after the addition of exogenous substrate, demonstra-
ning a deficiency of prostacyclin synthase activity and/or failure of metabolic channeling of the released AA to reach its thromboxane synthase target (Fig. 4C). Collectively, these results demonstrate the importance of calcium for the produc-
tion of eicosanoid products, the tight metabolic regulation of
mitochondrial phospholipases in non-failing hearts, and the differential metabolic channeling of released AA in non-failing versus failing human myocardium.

**Mitochondrial-associated phospholipases A₂ in non-failing versus failing human myocardium**

The remarkable differences in the rates of eicosanoid synthesis in non-failing versus failing hearts led us to further investigate the Ca²⁺ dependence and sensitivity of the mitochondrial phospholipase activity responsible for the release of the AA precursor for downstream oxygenation. Incubation of non-failing heart mitochondria with Ca²⁺ resulted in the dramatic and selective release of AA with only modest increases in other saturated and unsaturated fatty acids (Fig. 5 and Fig. S2). Concomitant with AA release, only lysophosphatidylcholines (LPCs) containing 16:0-diacyl or plasmenyl 16:0- (P16:0-) vinyl ether linkages were significantly increased upon calcium challenge. These results suggest that a highly active Ca²⁺-stimulated phospholipase preferentially hydrolyzes arachidonoylated phosphatidylcholine and plasmenylcholine (PC) molecular species present in the non-failing human heart (Fig. 5). In contrast, mitochondria from failing hearts demonstrated a markedly reduced capacity to release AA in response to calcium challenge with substantially less AA selectivity and attenuated LPC production (Fig. 5 and Fig. S2).

Next, we sought to identify the major heart mitochondrial PLA₂ enzyme(s) responsible for the release of AA from human mitochondria from non-failing and failing human myocardium. The results demonstrated that ~80% of Ca²⁺-activated AA release and LPC production containing either 16:0 fatty acyl or plasmenyl residue in non-failing heart mitochondria were inhibited by a non-selective serine hydrolase inhibitor, methyl

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**Figure 3. Differential effects of eicosanoid molecular species on mPTP opening.** A, isolated mitochondria from failing hearts (n = 6) were preincubated for 10 min with 10 μM baic, 20 μM ibu, or 40 μM MSPPOH for selective inhibition of lipoygenase, cyclooxygenase, or cytochrome P450 enzymes, respectively. After preincubation, mitochondrial swelling was initiated by addition of Ca²⁺. **p < 10⁻⁵** and ¶, p < 10⁻⁶ when compared with Ca²⁺ alone (Ctl). $\delta$, intact mitochondria from failing hearts were preincubated with 1 μM cyclosporine A (CsA) with or without 40 μM MSPPOH for 10 min at 25 °C. Mitochondrial swelling was monitored by addition of Ca²⁺ (left panel). The final ΔA₅₄₀ end-point measurements at 30 min are presented (right panel, n = 4 – 8). C, isolated mitochondria from non-failing control (n = 5) and failing hearts (n = 7) were preincubated with or without 40 μM MSPPOH for 10 min at 25 °C for inhibition of cytochrome P450(α). Mitochondrial swelling was monitored after addition of Ca²⁺ (left panel). The final ΔA₅₄₀ end point measurements at 20 and 30 min are presented in the middle and right panels, respectively (n = 4 – 8). $\delta$, p < 0.005. N.S., not significant. D, isolated mitochondria from failing hearts (n = 4) were preincubated for 10 min with 10 μM baic in the absence or presence of either 5-, 12-, 15-, or 20-HETE (4 μM final concentrations). Mitochondrial swelling was initiated by addition of Ca²⁺. Representative tracings for mitochondrial swelling and the absorbance decreases at 540 nm after 30 min of incubation are presented. *, p < 0.05, and **, p < 0.0005.
Ca\(^{2+}\)-activated phospholipases \(A_2\) in human heart mitochondria

![Figure 4. Deuterated (d\(_5\)) and non-labeled (H\(_n\)) eicosanoid production in the presence of exogenous d\(_6\)-arachidonic acid in non-failing control and failing human heart mitochondria. Isolated mitochondria (1 mg) were placed in 10 mM HEPES buffer (pH 7.4) containing 10% glycerol and 1 mM DTT, and homogenized by sonication on ice. Mitochondrial phospholipase activity was activated by addition of either 2 mM EGTA or 0.2 mM free Ca\(^{2+}\) in the presence of 8 nM of d\(_6\)-AA at 35 °C. After 20 min of incubation, the reaction was terminated by addition of methanol to 20% (v/v) final concentration at 4 °C. Eicosanoids were immediately extracted by solid phase extraction with internal standards, derivatized with AMPP, and analyzed by high-resolution mass accuracy mass spectrometry. The quantities of deuterated (d\(_5\)) and non-labeled (H\(_n\)) eicosanoids, including HETEs (A), EETs (B), and prostaglandins (C), were determined by calculations based on either internal standards, derivatized with AMPP, and analyzed by high-resolution mass accuracy mass spectrometry.

Identification of the predominant calcium-activated phospholipase activity in non-failing human mitochondria as cPLA\(_{2x}\) by sequential column chromatographies followed by activity-based protein profiling

In aggregate, these results suggested the presence of a previously unidentified phospholipase activity in human myocardial mitochondria that is calcium-sensitive and AA-selective and is present in either greater abundance or possesses a higher specific activity in non-failing relative to failing myocardium. These intriguing observations led us on a campaign to identify this unknown mitochondrial PL\(A_2\) activity in failing human hearts. Our strategy employed protein extraction at high pH for solubilization and sequential column chromatographies, including DEAE, chromatofocusing (Mono P), and Mono Q FPLC (Fig. 7A, Fig. 5A, and Table 1) followed by activity-based protein profiling using a serine hydrolase inhibitor as probe. The highly purified active fractions collected from the final Mono Q column were loaded onto an SDS-polyacrylamide gel, silver-stained, and either trypsinized in situ or utilized for activity-based protein profiling with desthiobiotin-fluorophosphonate (Fig. 7B). Tryptosynthesis of the active protein from the Mono Q column and analysis of the resultant peptides by ESI-nano LC-MS/MS identified multiple signature fragmentation peptides of cPLA\(_{2x}\) (i.e. \(526\)YGAYVPTELFGSELFMR\(^{533}\) and \(552\)ICYLQGMWGSATLSDEIFKpTAGpSGLSFLEW-pYR\(^{537}\)) (Fig. 8). Furthermore, the peptides from trypsinolysis of SDS-polyacrylamide gel (red dotted box in Fig. 7) were also

![Figure 5A. Control/EGTA](control EGTA), Control/Ca\(^{2+}\), Failing/EGTA, Failing/Ca\(^{2+}\)

![Figure 5B. Control/EGTA](control EGTA), Control/Ca\(^{2+}\), Failing/EGTA, Failing/Ca\(^{2+}\)

![Figure 5C. Control/EGTA](control EGTA), Control/Ca\(^{2+}\), Failing/EGTA, Failing/Ca\(^{2+}\)
identified as cPLA₂ζ fragmentation (i.e. 199GDGpTAPREEYGSRQLQLAVPGAYEK215 and 1MLWALWPRWLADK13) (Fig. S5). These results clearly indicated that cPLA₂ζ was the unknown mitochondrial phospholipase in non-failing myocardium responsible for the production of the majority of arachidonate release (Fig. 8). Although the protein that co-chromatographed with the PLA₂ activity was not readily detectable by silver staining due to its high specific activity and low abundance, the power of activity-based protein profiling with des-thiobiotin labeling followed by biotin immunoblotting confirmed the identity of cPLA₂ζ as the major PLA₂ in non-failing human myocardial mitochondria.

Discussion

Myocardial mitochondria undergo marked changes in size, lipid content, lipid second messenger generation, and bioenergetics in animal models after prolonged opening of the mPTP by various pathological stimuli (e.g. ischemia/reperfusion). These changes result from maladaptive calcium dynamics, the dissipation of membrane potential (ΔΨm), and the increase utilization of ketone bodies that feed into the tricarboxylic acid cycle at succinate likely increasing the generation of reactive oxygen species (7, 34–36). However, previous reports as well as results from this study did not demonstrate alterations in mitochondrial oxygen consumption, respiratory coupling, or different responses to various respiratory agonists and antagonists in mitochondria from failing human hearts (37, 38). Accordingly, the conclusions from those studies suggested that alterations in mitochondrial function were not responsible for the progression of human heart failure. However, no previous work has investigated alterations in human mitochondrial lipid content, lipid second messenger generation, or the identity and changes in the enzymatic mediators responsible for maladaptive lipid signaling upon pathological stimuli (e.g. calcium overload and decreased membrane potential) manifest in heart failure. The present results demonstrate the following: 1) the profound remodeling of Ca²⁺-dependent mitochondrial phospholipase activities during heart failure from cPLA₂ζ to iPLA₂γ; 2) the dramatic alterations in lipid second messenger production in failing hearts resulting in the generation of toxic HETEs and decreases in salutary EETs; 3) the differential metabolic channeling of AA released by cPLA₂ζ in comparison with iPLA₂γ in non-failing versus failing myocardial mitochondria; and 4) the ability of HETEs to promote Ca²⁺-induced mPTP opening further activating iPLA₂γ and its production of toxic HETEs in an autoamplification process (Fig. 9). Collectively, these results are the first to identify human mitochondrial Ca²⁺-activated phospholipases as mediators of alterations in the metabolic channeling of AA to discrete oxidized metabolites during heart failure and the bidirectional roles of oxidized arachidonic acid in either the protection or damage to human mitochondria challenged by calcium.

Alterations in the amplitude and duration of cardiac transients in failing myocardium/myocytes have been found to be elevated during diastole, decreased during systole, and prolonged during diastolic relaxation in animal models (39, 40). The delay in the relaxation phase has been correlated with deficient calcium uptake due to decreases in sarcoplasmic reticulum Ca²⁺-ATPase (SERCA2a) expression and activity as well as decreased phosphorylation of phospholamban during heart failure (39, 40). The resultant alterations in calcium uptake and delayed sequestration during heart failure likely predispose heart mitochondria to mPTP opening through Ca²⁺ activation of mitochondrial phospholipase(s) and subsequent production of cardiotoxic lipid second messengers. Consistent with this notion, our recent findings have demonstrated that murine myocardial mitochondrial iPLA₂γ is activated by calcium and that iPLA₂γ loss of function in cardiac myocytes markedly attenuates Ca²⁺-induced mPTP opening, generation of proinflammatory oxidized lipid metabolites (e.g. eicosanoids, docosanoids, and oxidized linoleic acid metabolites), and membrane potential dissipation resulting in reduction of infarct size during cardiac I/R in mice (13, 27).
Ca\(^{2+}\)-activated phospholipases A\(_2\) in human heart mitochondria

Figure 6. Pharmacological inhibition of PLA\(_2\) activities differentiates non-failing myocardial mitochondrial PLA\(_{2}\)-s from those present in failing human heart. Isolated mitochondria from human non-failing control (n = 5) (A and B) and failing hearts (n = 8) (C) were placed in 10 mM HEPES buffer (pH 7.4) containing 10% glycerol and 1 mM DTT and homogenized by sonication on ice. Mitochondrial homogenates were preincubated with 10 \(\mu\)M (S)-BEL, 10 \(\mu\)M (R)-BEL, 2 \(\mu\)M Pyr, or 10 \(\mu\)M MAFP for 10 min or with 25 \(\mu\)M AACCOCF\(_3\) for 30 min at 25 °C. Mitochondrial phospholipase reactions were initiated by addition of 2 mM EGTA or 0.6 mM Ca\(^{2+}\) at 35 °C. After 10 min of incubation, the reactions were terminated by addition of chloroform/methanol (1:1, v/v). The lipid products including AA, 16:0-LPC, and 16:0 plasmenyl phosphatidylcholine were extracted by a modified Bligh-Dyer method in the presence of lipid internal standards and quantified by mass spectrometry. 

The cardioprotective effects of CYP450 products and the toxic effects of certain HETEs produced by both CYP450 and LOX enzymes have been studied in animal models of heart failure. For example, EETs and a soluble epoxide hydrolase inhibitor have been demonstrated to reduce infract size during I/R and improve cardiac function by preventing cardiac fibrosis (20, 41, 42). In contrast, cardiac overexpression of 12/15-LOX was reported to result in systolic dysfunction, inflammation, cardiac fibrosis, and macrophage infiltration (43, 44).

The results of this study identify dramatic increases in both the rate of production and steady-state amounts of toxic HETEs in mitochondria from failing human myocardium. As far as we are aware, this study is the first to demonstrate the prominent roles of HETEs in regulating mPTP opening in either experimental animals or in human myocardium after calcium challenge. Our results obtained by utilizing selective pharmacological inhibitors suggest the intriguing possibility that the sensitivity of failing myocardium to stress and its predisposition to necrosis/apoptosis is increased by the up-regulation of Ca\(^{2+}\)-activated phospholipase-initiated HETE production at the expense of salutary EETs.

A completely unanticipated finding in this study is the remodeling of mitochondrial phospholipases during heart failure. It was originally thought that AA precursor release for subsequent eicosanoid production in human hearts would be detrimental to cardiac function on multiple levels, including lethal arrhythmias, inflammation, and hemodynamic performance. However, the results of initial rate analyses using stable isotopes in conjunction with HRAM mass spectrometry indicate that the generation of endogenous AA by mitochondrial phospholipase(s) and the resultant downstream eicosanoid metabolites in non-failing hearts are remarkably different from those in human failing hearts. Accordingly, it became important to identify the salutary mitochondrial phospholipase(s) in non-failing hearts. Through sequential chromatographies in conjunction with activity-based protein profiling and high mass accuracy proteomics (45, 46), we quite unexpectedly identified the predominant phospholipase activity associated with non-failing human heart mitochondria as cPLA\(_{2}\). In stark contrast, the predominant mitochondrial phospholipase activity in failing hearts was catalyzed by iPLA\(_{2}\gamma\) generating toxic HETEs which, in combination with a prominent decrease in protective EETs, likely results in alterations in surface charge and potential in the mitochondrial membrane (27, 47). Moreover, previous studies by our group reported the regiospecific PLA\(_{2}\) activity of iPLA\(_{2}\gamma\) with phospholipids containing polyunsaturated fatty acids at the sn-2 position. This results in palmitate release in the plane of the inner membrane. Palmitate has previously been shown to be a potent activator of mPTP opening that, when generated in the plane of the inner membrane, could rapidly diffuse laterally to amplify mPTP opening (11, 27, 48). Thus, a single enzyme, iPLA\(_{2}\gamma\), has dual regulating pathways for opening of the mPTP, each of which are dependent upon the generation of lipid-signaling molecules (e.g. palmitate and oxidized PUFAs). Furthermore, these results identify inhibition of iPLA\(_{2}\gamma\) as a high value pharmacological target to prevent the progression of heart failure by attenuating cardiac myocyte dropout from necrosis and apoptosis.

A second unexpected outcome of this study is the demonstration that AA generated in human myocardial mitochondria is not due to cPLA\(_{2}\alpha\), which is considered to be the canonical pathway for AA generation as reported in numerous other studies. Instead, AA release in mitochondria from non-failing human myocardium is largely the result of cPLA\(_{2}\zeta\) that is under tight control to release AA that is channeled into the production of cardiac protective metabolites. These findings are particularly relevant considering that mitochondria represent 30% of myocardial volume, and human mitochondrial phospholipids contain large amounts of esterified AA, thereby emphasizing the important roles of mitochondrial phospholipases in human heart function during health and disease. Although cPLA\(_{2}\zeta\) from mouse has been cloned (49, 50), its primary struc-
**Ca^{2+}**-activated phospholipases A_2 in human heart mitochondria

To detect the putative serine hydrolase(s) responsible for the observed PL_{A2} activity, mitochondria isolated from non-failing human myocardium were resuspended in highly basic buffer (20 mM Na$_2$CO$_3$ (pH 11.4)), and the soluble fraction was collected by ultracentrifugation at 100,000 x g. Next, the mitochondrial extract was applied to sequential DEAE and chromatofocusing Mono P columns as described under “Experimental procedures” (Fig. S4). Fractions with phospholipase A$_2$ activity (identified using [14C]PAPC as substrate) were collected at each chromatographic step for high-resolution separation utilizing a Mono Q FPLC column. The elution profile of the identified PL_{A2} activity following Mono Q chromatography as monitored by UV absorbance (280 nm) and release of [14C]arachidonic acid from [14C]PAPC substrate. Active fractions from the Mono Q FPLC column were resolved by SDS-PAGE and visualized by silver staining (B) or were incubated with desthiobiotin-fluorophosphonate for labeling of serine hydrolases prior to separation by SDS-PAGE and subsequent Western blot analysis to visualize resultant biotinylated proteins (C). A protein band was identified in which the intensity of labeling by desthiobiotin-fluorophosphonate correlated well with the observed PL_{A2} catalytic activity in each fraction (fractions 25–27 in the red dotted box in B). Hsp70, 70-kDa heat shock protein; MCAD, medium-chain acyl-CoA dehydrogenase; SCAD, short-chain acyl-CoA dehydrogenase.

**Table 1**

Purification table for the predominant human mitochondrial phospholipase A$_2$

Myocardial mitochondria (200 mg) were isolated from human non-failing human heart, resuspended in 20 mM Na$_2$CO$_3$ buffer (pH 11.4, containing 10% glycerol, 2 mM EGTA, and 2 mM EDTA), and homogenized by brief sonication. Phospholipase A$_2$ activity in the supernatant after ultracentrifugation was purified by multiple chromatographic steps, including DEAE, Mono P chromatofocusing, and Mono Q FPLC through following the release of [14C]AA from [14C]PAPC substrate. It should be noted that the initial yield of crude homogenates assigned as 100% is certainly underestimated due to substantial isotope dilution of the radioactive substrate by endogenous lipids present in the mitochondrial homogenate. The amount of protein in the active fractions from each chromatographic step was determined by a Bradford protein assay with BSA as standard. The details for each protein purification step are described under “Experimental procedures.”

| Mitochondria (200 mg) | Total protein mg | Total activity mmol/min | Specific activity mmol/min/mg | Purification Yield fold | % |
|-----------------------|------------------|-------------------------|-------------------------------|-------------------------|---|
| Crude homogenates     | 200              | 0.94                    | 0.0047                        | 1                       | 100 |
| pH 11.4 extraction    | 62               | 0.53                    | 0.0085                        | 1.8                     | 53  |
| DEAE                  | 6.8              | 2.9                     | 0.43                          | 91                      | 306 |
| Mono P                | 0.5              | 1.3                     | 2.7                           | 510                     | 138 |
| Mono Q                | 0.018            | 1.6                     | 87                            | 18,596                  | 170 |

In contrast, the human heart cultured is substantially different (only 73% homology to the human isoform) and there have not been any studies published to the best of our knowledge on human cPLA$_{2\gamma}$. Critically, failing myocardium has a deficiency in generating or directing the cPLA$_{2\gamma}$-mediated release of arachidonic acid to the synthesis of beneficial eicosanoid metabolites. Furthermore, these results show that the necessary oxygenase enzymes for conversion of AA to salutary eicosanoids products are present in failing human myocardium, but that they cannot be accessed by AA release from phospholipases active in failing human hearts. We note that cPLA$_{2\gamma}$ has not been previously reported in human myocardium nor has its utility in the production of salutary eicosanoids been recognized.

Collectively, this study demonstrates the profound remodeling of mitochondrial phospholipase activities and metabolic channeling of arachidonic acid in non-failing versus failing human myocardium resulting in dramatic alterations in eicosanoid second messenger generation and metabolite profiles in failing hearts. Accordingly, myocardial phospholipases are prime targets for pharmacological interventions because they likely initiate a series of cellular events that collectively conspire to accelerate the progression of human heart failure. Considering the activation of iPLA$_{2\gamma}$ mediated by calcium or membrane potential and its resultant production of toxic HETEs, which open the mPTP, future studies should target the prevention of the activation of human myocardial iPLA$_{2\gamma}$ and/or the generation of its detrimental downstream toxic products.
Ca\(^{2+}\)-activated phospholipases A\(_2\) in human heart mitochondria

Figure 8. Identification of the purified human myocardial mitochondrial PLA\(_2\) as cPLA\(_2\)\(_{\gamma}\) by mass spectrometric proteomic analysis. Active fractions from the Mono Q FPLC column (Fig. 7) were trypsinized, and the resultant peptides were analyzed by mass spectrometry (NanoLC-MS/MS) utilizing a protein sequence database program (SEQUEST). An excellent correlation between observed and predicted b and y ions for the tryptic peptides corresponding to residues 526–543 (A) and 553–587 (B) of human cPLA\(_2\)\(_{\gamma}\) was obtained. Interestingly, multiple phosphorylated residues were identified in the 553–587 peptide. p, phosphorylation; #, oxidation.

Figure 9. Schematic comparison of phospholipases responsible for Ca\(^{2+}\)-activated arachidonic acid release and its differential downstream eicosanoid metabolites in non-failing versus failing human myocardium mitochondria. The predominant PLA\(_2\) activity in non-failing human heart mitochondria is from cPLA\(_2\)\(_{\gamma}\) which releases AA that is channeled to cardioprotective EET production. In stark contrast, iPLA\(_2\)\(_{\gamma}\) in failing heart mitochondria is a major PLA\(_2\) activity responsible for the dramatic increase in toxic HETEs upon calcium challenge facilitating mPTP opening. Deuterated eicosanoid metabolite profiles generated by the oxidation of exogenous d\(_9\)-AA did not differ significantly in non-failing versus failing myocardial mitochondria suggesting the presence of differential metabolic channelling of endogenously released AA mediated by pathological remodeling of mitochondrial PLA\(_2\) isoforms. P450, cytochrome P450.

**Experimental procedures**

**Reagents**

Ibuprofen, baicalein, MSPPOH, AACOCF\(_3\), BEL, MAFP, 5-HETE, 12-HETE, 15-HETE, 20-HETE, arachidonic acid, d\(_8\)-arachidonic acid, TXB\(_2\)-d\(_4\), PGE\(_2\)-d\(_4\), and 12-HETE-d\(_8\) were purchased from Cayman Chemical. N-((2S,4R)-4-(Biphenyl-2-ylmethyl-isobutyl-amino)-1-[2-(2,4-difluorobenzoyl)-benzoyle]-pyrroli din-2-ylmethyl)-3-[4-(2,4-dioxothiazolidin-5-ylidenemethyl)-phenyl]acrylamide (cPLA\(_2\)\(_{\gamma}\) inhibitor, a pyrroline derivative) was obtained from EMD Millipore. Des-thiobiotin-fluorophosphonate was purchased from Thermo Fisher Scientific. Lipid internal standards, including 17:0-LPC and d\(_8\)-16:0-fatty acid, were purchased from Avanti Polar Lipids, Inc., and Cambridge Isotope Laboratories, Inc. A rabbit polyclonal antibody directed against human iPLA\(_2\)\(_{\gamma}\) was generated in our laboratory as described previously (12). Antibodies against biotin, VDAC, ANT, and CypD were obtained from Santa Cruz Biotechnology. Unless otherwise indicated, all reagents for chromatographic protein purification, proteomic analysis, and mitochondrial studies were obtained from Sigma or Thermo Fisher Scientific.

**Collection of human non-failing and failing heart tissue**

All human tissue was collected with the written informed consent received from participants as a part of an Institutional Review Board (IRB) approved protocol at Washington University in St. Louis and the Human Research Committee at Harvard Medical School, Boston, MA.
University School of Medicine. Clinical variables from a variety of fields were collected on each heart failure patient by the clinical heart failure and heart transplant service as a part of a separate IRB-approved protocol. Human heart tissue was obtained from Mid-America Transplant (St. Louis, MO) and the Washington University Translational Cardiovascular Tissue Core. Tissue from the left ventricular apex of failing human hearts was collected at the time of ventricular assist device (VAD) implantation. Tissue from the left ventricle and left ventricular apex of non-failing human hearts was obtained from organ donors having a heart deemed unsuitable for transplantation due to donor age, epicardial (non-occlusive) coronary vascular disease, or high risk behavioral profile. All tissue was trimmed of excess fat and washed in cold phosphate-buffered saline and then placed in ice-cold mitochondrial isolation buffer or freeze-clamped at the temperature of liquid nitrogen for later mass spectral and metabolic analyses.

Isolation of human heart mitochondria

Human heart tissue from non-failing and VAD (failing) donors was collected in Mitochondrial Isolation Buffer (MIB: 0.21 M mannitol, 70 mM sucrose, 0.1 mM potassium-EDTA, 1 mM EGTA, 10 mM Tris-HCl, 0.5% fatty acid-free BSA (pH 7.4)) at 4 °C. Fat and fibrous tissues were removed prior to extensive washing of the myocardial sample with isolation buffer to remove residual blood. After cutting into small pieces and chopping briefly with a razor blade on ice (4 °C ambient temperature) in mitochondrial isolation buffer, the heart tissue was homogenized using 12–15 passes with a Teflon homogenizer at a rotational speed of 120 rpm. Next, the homogenate was centrifuged for 5 min at 850 × g, and the supernatant was collected and centrifuged at 7200 × g for 10 min. The mitochondrial pellet was resuspended in MIB without BSA and EGTA and then centrifuged again for 7200 × g for 10 min. The resultant pellet was resuspended in either mitochondrial respiration buffer, mitochondrial swelling buffer, or HEPES buffer (for the determination of PLA2 activity).

Mitochondrial swelling assay

Isolated human myocardial mitochondria were placed in Mitochondrial Swelling Buffer (3 mM HEPES (pH 7.4), containing 0.23 M mannitol, 0.07 M sucrose, 5 mM succinate, and 2.5 μM rotenone in the presence or absence of 1 mM KH2PO4) at 4 °C. For experiments examining the effects of PLA2 inhibition or arachidonic acid oxygenase inhibition, mitochondria were pretreated with the following inhibitors: 10 μM (R)- or (S)-BEL, (E)-6-(bromomethylene)-3-(1-naphthylenyl)-2H-tetrahydropyran-2-one, 2 μM pyrrolidine cPLA2α inhibitor, 10 μM baicalein (5,6,7-trihydroxyflavone), 40 μM MSPPOH (N-(methylsulfonyl)-2-(2-propynoloxy)-benzenehexanamide), 20 μM ibuprofen (2-(4-isobutylphenyl)propanoic acid), or dimethyl sulfoxide (DMSO) vehicle alone (0.5% v/v) for 10 min at 23 °C. Mitochondrial swelling was initiated by addition of 70 μM CaCl2 (final concentration) or 10 μM EGTA as control. Decreases in the absorbance at 540 nm indicative of mitochondrial swelling were measured at 15-s intervals using a Spectra-Max M5e microplate reader (Molecular Devices, Sunnyvale, CA).

High-resolution mitochondrial respirometry

Human heart tissue collected from control (non-failing) and VAD (failing) donors was placed in ice-cold Mitochondrial Isolation Buffer (MIB: 0.21 M mannitol, 0.070 M sucrose, 0.1 mM K-EDTA, 10 mM Tris-HCl, 1 mM EGTA, 0.5% BSA (pH 7.4)) in a Petri dish on ice. Heart tissue was immediately diced into small millimeter-sized pieces with a razor blade, transferred to a 15-ml Potter-Elvehjem Teflon-pestle glass-mortar tissue grinder with 7 ml of MIB. The tissue was homogenized using a motor-driven pestle operated at 120 rpm. The homogenate was then centrifuged in 10 ml of MIB for 7 min at 850 × g. The supernatant was carefully collected and centrifuged at 10,000 × g for 10 min. The final pellet was resuspended in MIB with no BSA. High-resolution respirometry was performed using a 2-ml chamber OROBOROS® Oxygraph 2K (Innsbruck, Austria). Respiratory measurements were performed at 30 °C utilizing 50 μg of mitochondria in MiRO5 buffer. The rate of oxygen consumption was calculated as a time derivative of oxygen concentration using the DatLab Analysis software (OROBOROS®, Innsbruck, Austria). Respiration was started by the addition of either pyruvate (5 mM)/malate (5 mM), palmitoylcarinitine (20 μM)/malate, glutamate (10 mM)/malate, or pyruvate (5 mM)/glutamate (10 mM)/malate (State 2) followed by sequential addition of ADP (1.25 mM) (State 3), succinate (5 mM) (State 3 Max), rotenone (0.5 μM), oligomycin (2.5 μM) (State 4), N,N,N',N'-tetramethyl-p-phenylenediamine (5 μM) with ascorbate (0.5 μM), and antimycin A (1 μM). All values reflect the subtraction of residual oxygen consumption after addition of antimycin A. The quantity of mitochondrial protein was determined using a Pierce™ BCA protein assay kit according to the manufacturer’s instructions.

Determination of phospholipase activity in mitochondrial homogenates

Isolated human myocardial mitochondria placed in HEPES buffer (10 mM HEPES (pH 7.4) containing 1 mM DTT and 10% glycerol) were sonicated for 20 s with a Branson sonicator using 1-s pulses at 30% power. The protein contents of the sonicated mitochondria were then determined using a Bradford protein assay (Bio-Rad) with bovine serum albumin as standard. For experiments examining the effects of PLA2 inhibition, mitochondria were preincubated with the inhibitors (10 μM (R)-BEL, 10 μM (S)-BEL, 25 μM arachidonyl trifluoromethyl ketone (AACOCF3), 10 μM methyl arachidonoyl fluorophosphonate (MAFP), or 2 μM pyrrolidine) or DMSO vehicle alone (1% v/v, final) for 15 min at 23 °C. Mitochondrial phospholipase activity was initiated by adding either 2 mM EGTA or CaCl2 at the indicated concentrations at 35 °C for 10 min, and then reactions were terminated by sequential addition of 2 ml of chloroform/methanol (1:1, v/v), internal standards (d14-16:0 free fatty acid and 17:0-LPC), and 700 μl of 10 mM LiCl. After vigorous vortexing, samples were centrifuged to promote phase separation, and the chloroform layer was collected and dried under nitrogen stream. The remaining lipids were then re-extracted with 3
ml of chloroform/methanol/water (1:1:1, v/v/v), and those lipids present in the chloroform layer were dried under nitrogen stream. Identification and quantitation of extracted free fatty acid and LPC molecular species were performed in the negative and positive ion modes, respectively, utilizing a TSQ Quantum Ultra mass spectrometer (Thermo Fisher Scientific, San Jose, CA) equipped with an automated nanospray apparatus (Advion Bioscience, Ithaca, NY) as described previously (11). In some cases, fatty acids were also analyzed in positive ion mode after chemical derivatization with AMPP under the same reaction conditions as eicosanoid AMPP derivatization (see below).

Identification and quantitation of eicosanoids by LC-MS/MS

Mitochondrial homogenates in HEPES buffer at the indicated concentrations were incubated in the presence of either 2 mM EGTA or 0.2 mM free Ca²⁺ for 20 min at 35 °C. The reaction was stopped by addition of methanol (20% v/v final concentration) at 4 °C and diluted with water after addition of internal standards (250 pg each of TXB₂-d₄, PGF₂α-d₄, and 12-HETE-d₄). The solution was immediately applied to a Strata-X solid-phase extraction cartridge preconditioned with 1 ml of methanol followed by 1 ml of 10% methanol. The cartridge was then washed with 1 ml of 5% methanol twice, and the remaining solvent was flushed out with N₂ at a pressure of 5 p.s.i. Bound eicosanoids were eluted with 1 ml of methanol containing 0.1% glacial acetic acid. All cartridge steps were carried out using a vacuum manifold attached to a house vacuum line. After evaporation of the organic solvent with a SpeedVac, isolated eicosanoids were derivatized with N-(4-aminomethylphenyl) pyridinium (AMPP), as described previously (28). Briefly, eicosanoids were dissolved in 12.5 μl of ice-cold acetonitrile/N,N-dimethylformamide (4:1, v/v) followed by addition of 12.5 μl of ice-cold 640 mM (3-(dimethylamino)propyl)ethyl carbodiimide hydrochloride in HPLC grade water. Eicosanoid derivatization was initiated by adding 25 μl of 5 mM N-hydroxysulfonofluorophosphate and 15 mM AMPP and heating at 60 °C for 30 min. The derivatized eicosanoids were separated by RP-HPLC utilizing a C18 reverse-phase column (Ascentis Express, 2.7-μm particles, 150 × 2.1 mm) at 23 °C using a linear mobile phase gradient with solvent A (0.1% glacial acetic acid in water) and solvent B (0.1% glacial acetic acid in acetonitrile) at a flow rate of 0.2 ml/min. The solvent gradient was programmed as follows: 0.0–5.0 min, 25% B; 5.0–7.0 min, 25–35% B; 7.0–20.0 min, 35–60% B; 20.0–20.1 min, 60–100% B; 20.1–24.0 min, 100% B; 24.0–25.0 min, 100% B to 25% B and 10-min isocratic hold at 25% B (14). The separated derivatized eicosanoids were analyzed after infusion into a hybrid tandem mass spectrometer (LTQ-Orbitrap, Thermo Fisher Scientific) via selected reaction monitoring in the positive ion mode with sheath, auxiliary, and sweep gas flows (arbitrary units) of 30, 5, and 1, respectively. The capillary temperature was set to 275 °C, and the electrospray voltage was 4.1 kV. Capillary voltage and the tube lens voltage were set to 2 and 100 V, respectively. Instrument control and data acquisition were performed using Thermo Xcalibur Version 2.1 software. For stable isotope labeling experiments utilizing d₈-arachidonic acid (d₈-AA), it is critical to first remove residual oxidized d₈-AA from the original stock by reverse-phase HPLC. In addition, mitochondrial phospholipase activity assay described above was performed in the presence of 8 nmol of d₈-AA, which was purified by HPLC to remove residual oxidized AA, during the 20-min reaction time. Quantitation of eicosanoids was performed by using either internal standards, including 250 pg each of TXB₂-d₄, PGF₂α-d₄, or calibration of an external standard such as 250 pg of d₈-12-HETE.

Purification of human heart mitochondrial phospholipase A₂

Mitochondria isolated from non-failing human hearts were placed in 20 mM Na₂CO₃ buffer ([pH 11.4] containing 10% glycerol, 2 mM EGTA, and 2 mM EDTA) and sonicated for 20 s with 1-s pulses at 30% power. Insoluble membrane material was removed by ultracentrifugation at 100,000 × g for 1 h, and the soluble fraction was diluted with 20 mM Tris-Cl (pH 7.4), loaded to a 1-ml HiTrap-DEAE (GE Healthcare) column equilibrated with the same buffer and eluted with a stepwise NaCl gradient (0–0.5 M NaCl). Active fractions were identified by a PL₂ assay utilizing 1-α-1-palmitoyl-2-arachidonoyl[arachidonoyl-1-14C]-sn-glycero-3-phosphocholine ([14C]PAPC) (PerkinElmer Life Sciences)/1-(12-octadecenyl)-2-oleyl-sn-glycero-3-phosphoethanolamine (7:5) as substrate. The released [14C]AA was extracted into chloroform using the Bligh-Dyer method and isolated by TLC, and its quantity was determined by liquid scintillation counting. After desalting active fractions using a gel filtration column (Bio-Rad) with Mono P equilibration buffer (20 mM bis-Tris-Cl (pH 6.3) containing 2% glycerol), the active PL₂ was loaded onto a Mono P chromatofocusing column (GE Healthcare) and resolved with a linear gradient from pH 6.0 to 4.0 by utilizing Polybuffer 74 (GE Healthcare). Mono P active fractions determined by radioactive PL₂ assay were diluted with Mono Q equilibration buffer (20 mM Tris-Cl (pH 7.4) containing 10% glycerol), loaded onto Mono Q column (GE Healthcare), and resolved with a linear NaCl gradient (0–0.4 M). Active Mono Q fractions were determined using [14C]PAPC as substrate. The amount of protein in active fractions from each of the chromatographic steps was determined using a Bradford Protein Assay (Bio-Rad) with BSA as standard. For activity-based protein profiling, the fractions from the Mono Q column were incubated with 0.5 μM des-thiobiotin-fluorophosphonate for 30 min in the presence of 1 mM Ca²⁺. Reactions were terminated by addition of SDS-PAGE sample loading buffer. Following electrophoresis, immunoblot analysis was performed to identify biotinylated target proteins.

In-gel digestion of proteins for identification by mass spectrometric analyses

Protein bands visualized with silver staining were excised from the gel, cut into small pieces (∼1 mm³), and destained twice by incubating with a freshly prepared 1:1 (v/v) solution (300 μl) of 30 mM potassium ferricyanide and 100 mM sodium thiosulfate for 8 min. The solution was discarded, and the gel pieces were washed four times with 1 ml of water. After incubation with 100% acetonitrile for 15 min at 23 °C, the solution was discarded. The gel pieces were dried utilizing a
SpeedVac, rehydrated in 25 mM ammonium bicarbonate (pH 7.8) containing 12.5 μg/ml trypsin, and then incubated on ice for 45 min. Excess liquid was removed, and 25 mM ammonium bicarbonate (pH 7.8) was added to cover the gel pieces. Following overnight incubation at 37 °C, the supernatant was removed and transferred to a siliconized microcentrifuge tube on ice. Next, 30 μl of 25 mM ammonium bicarbonate (pH 7.8) was added and incubated for 20 min at 23 °C. The supernatant was removed and combined with the supernatant from the previous step. Finally, 30 μl of 50% acetonitrile containing 5% formic acid was added to the gel pieces, which were then incubated for 20 min at 23 °C. The resultant supernatant was combined with the supernatants from the previous steps. This extraction step was repeated once, and the peptide extracts were combined and evaporated to near dryness using a SpeedVac. The peptide extract was reconstituted in 5% acetonitrile containing 0.1% formic acid for analysis by ESI-NanoLC-MS/MS.

**ESI-NanoLC-MS/MS analysis**

Trypsinized peptide samples were loaded onto a PepMap100 C18 precolumn (300 μm × 1 cm, Dionex, Sunnyvale, CA) by a Surveyor autosampler (Thermo Fisher Scientific, San Jose, CA). The precolumn was then washed with 0.1% formic acid for 5 min. Trypsinized peptides from the precolumn were eluted with a gradient from 95% of mobile phase A (0.1% formic acid in water) and 5% of mobile phase B (0.1% formic acid in acetonitrile) to 10% of A and 90% of B onto a reverse-phase C18 analytical PepMap100 Nano-LC column (75 μm × 15 cm, Dionex, Sunnyvale, CA) at a flow rate of 200 nL/min for 150 min. The eluted peptides were injected into an LTQ-Orbitrap mass spectrometer (Thermo Fisher Scientific, San Jose, CA) at a flow rate of 200 nl/min for 150 min. The eluted peptides were injected into an LTQ-Orbitrap mass spectrometer (Thermo Fisher Scientific, San Jose, CA) by a Surveyor autosampler (Thermo Fisher Scientific, San Jose, CA) operated in a data-dependent acquisition mode. Each full scan analysis was set at 30 ms with an activation parameter q = 0.25, and precursor ions were isolated within a mass window range of 2 Th. The acquired data were searched against a customized protein sequence database using the SEQUEST algorithm. All positive identifications were individually confirmed.

**Statistical analyses**

A two-tailed Student t test was routinely performed to determine the significance of differences between two groups. A p value of <0.05 was considered significant. All data were reported as average ± S.E.

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