Suppression of Mitochondrial Function by Oxidatively Truncated Phospholipids Is Reversible, Aided by Bid, and Suppressed by Bcl-XL*

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Oxidatively truncated phospholipids are present in atherosclerotic lesions, apoptotic cells, and oxidized low density lipoproteins. Some of these lipids rapidly enter cells to induce apoptosis by the intrinsic pathway, but how such lipids initiate this process is unknown. We show the truncated phospholipid hexadecyl azelaoyl glycerophosphocholine (Az-LPAF), derived from the fragmentation of abundant sn-2 linoleoyl residues, depolarized mitochondria of intact cells. Az-LPAF also depolarized isolated mitochondria and allowed NADH loss, but did not directly interfere with complex I function. Cyclosporin A blockade of the mitochondrial permeability transition pore partially prevented the loss of electrochemical potential. Depolarization of isolated mitochondria by the truncated phospholipid was readily reversed by the addition of albumin that sequestered this lipid. Ectopic expression of the anti-apoptotic protein Bcl-XL in HL-60 cells reduced apoptosis by the truncated phospholipid by protecting their mitochondria. Mitochondria isolated from these cells were also protected from Az-LPAF-induced depolarization. Conversely mitochondria isolated from Bid−/− animals that lack this pro-apoptotic Bcl-2 family member were resistant to Az-LPAF depolarization. Addition of recombining full-length Bid, which has phospholipid transfer activity, restored this sensitivity. Thus, phospholipid oxidation products physically interact with mitochondria to continually depolarize this organelle without permanent harm, and Bcl-2 family members modulate this interaction with full-length Bid acting as a co-factor for pro-apoptotic, oxidatively truncated phospholipids.

Vascular cells are exposed to oxidizing radicals during normal metabolism, but especially so during physiologic and pathologic inflammatory processes. The double bonds of polyunsaturated fatty acyl residues are particularly prone to attack by radicals because the C-H bond situated between two double bonds is relatively weak, allowing a more facile abstraction of hydrogen to produce a radical. Because polyunsaturated fatty acyl residues are abundant and are generally esterified in the sn-2 position of the glycerol backbone, common products of oxidative attack on cells and circulating lipoproteins are phospholipids that have been peroxidized at their sn-2 position. These peroxide radicals abstract hydrogen to form hydroperoxy phospholipids, may be reduced to the corresponding alcohol, rearrange (2, 3), or fragment to generate a host of oxidatively truncated phospholipids (4–7).

The shortened sn-2 residue of truncated phospholipids, which may also contain a newly introduced polar oxygen function, does not intercalate into the membrane well and is energetically favored to protrude into the aqueous phase, a conformation that disorders phospholipid packing into a bilayer (8–10). Oxidatively truncated phospholipids are more water soluble than their phospholipid precursors and readily associate with plasma albumin (11), plasma membranes (12), and even traffic into cells to lysosomes (12) or mitochondria (13) depending on the structure of the truncated phospholipid.

Phospholipid oxidation products can be cytotoxic (14, 15), and at least some of these are toxic because they initiate the apoptotic process of regulated cell death (13). The manner by which oxidatively truncated phospholipids alter cell viability has been ascribed to solubilization of the plasma membrane (14), addition of mitochondrial proteins (17), temporary physical distortion of the plasma membrane (18), or activation of acid sphingomyelinase activity that alters plasma membrane microdomains by generating ceramide (15, 19). We found that a common oxidatively truncated phospholipid, containing a 9-carbon azelaoyl fragment derived from fragmentation of sn-2 linoleoyl residues, induces apoptosis by the intrinsic caspase cascade with loss of mitochondrial function and not, apparently, from damage of the plasma membrane (13).

Members of the Bcl-2 family modulate mitochondria-dependent apoptosis either by promoting apoptosis (Bid, Bad, and Bax) or obstructing this event (Bcl-2 and Bcl-XL). Aggregation of Bax on the mitochondrial outer membrane forms ion conducting pores and Bcl-XL associates with mitochondrial outer membranes to suppress this Bax activity (20). In contrast, Bid promotes apoptosis after cleavage to truncated Bid, a regulatory event catalyzed by activated caspase 8 (21). Bid, alone among Bcl-2 family members, displays homology to plant lipid transfer proteins and both truncated and full-length Bid will incorporate fluorescent phospholipids, and not the cognate fluorescent fatty acid, into mitochondrial membranes (22).

We determined whether mitochondrial integrity or function were directly affected by oxidatively truncated phospholipids, and then whether Bcl-2 family members alter these effects as they do in other, established apoptotic signaling pathways. We
Oxidized Phospholipids Depolarize Mitochondria

find that truncated phospholipids accumulated from the extracellular environment depolarize intracellular mitochondria, that these bilayer challenged phospholipids reversibly interact with mitochondria to continually reduce their transmembrane potential, and that Bcl-2 family members modulate this interaction.

EXPERIMENTAL PROCEDURES

Materials

JC-1, annexin V, propidium iodide, and the mitoprobe transition pore assay kit were from Invitrogen. Recombinant murine Bid was from ProteinX Lab (San Diego, CA). Tetramethylrhodamine methyl ester (TMRM),2 CCCP (carbonyl cyanide m-chlorophenyl hydrazone), NADH, coenzyme Q10, rotenone, phosphate-buffered saline (PBS), cyclosporin A, bovine serum albumin (BSA), and other reagents were from Sigma. The Cleveland Clinic medium core prepared RPMI 1640 serum albumin (BSA), and other reagents were from Sigma. The Cleveland Clinic medium core prepared RPMI 1640 medium. Fetal bovine serum was from Fisher Scientific (Pittsburgh, PA). [acetyl-3H]PAF was supplied by PerkinElmer Life Sciences. Az-LPAF and butyroyl-PAF (C4PAF) were from Cayman Chemical (Ann Arbor, MI). Lyso-PAF, carbamoyl-PAF, PAF, and BEL were from BioMol Research Laboratories (Plymouth Meeting, PA). Palmitoyl glutaroyl phosphatidylcholine, palmitoyl oxoaleryl phosphatidylcholine, and lysophosphatidic acid were from Avanti Polar Lipids, Inc. (Alabaster, AL).

Cell Culture

HL-60 cells were cultured in RPMI 1640 supplemented with streptomycin (100 units/ml), penicillin (100 μg/ml), and 10% fetal bovine serum. The cells were maintained in log phase growth at 37 °C in a humidified atmosphere of 5% CO2.

Mitochondria Preparation

Livers from adult Sprague-Dawley rats, or C57 BL6 wild type or Bid−/− mice (generously provided by Dr. Xiao-Ming Yin, University of Pittsburgh) in a protocol approved by the Cleveland Clinic IACUC were excised and minced on ice in EB medium (200 mM D-mannitol, 70 mM sucrose, 20 mM Hepes, pH 7.4, 0.5 g/liter of defatted BSA, and 1 mM EGTA). Homogenates were centrifuged at 1,000 g for 5 min twice before these supernatants were centrifuged at 9,500 × g for 10 min. The resulting pellets were resuspended in the same volume of EB medium without BSA and centrifuged at 9,500 × g for 10 min. The final pellet was resuspended in EB medium to 35 mg of protein/ml. HL-60 cells (3 × 10⁸) stably transfected with Bcl-XL or its empty vector were washed twice in PBS and once in EB medium before the cells were suspended in 10 ml of EB medium containing BSA (2 mg/ml), and then mechanically homogenized and mitochondria isolated as above. Cellular organelles were prepared from liver homogenates pre-cleared by centrifugation (1,000 × g) and then fractionated by centrifugation through a sucrose density gradient. Sucrose concentrations of the recovered fractions were determined by refractometry.

Phospholipid Hydrolysis

Density purified subcellular fractions were recovered and assayed for PAF hydrolysis with or without a 30-min preincubation with the iPLA2 inhibitor BEL at 10 μM to achieve complete inhibition. Hydrolysis of [acetyl-3H]PAF was quantitated by separation and recovery of [3H]acetate over C8-silica cartridges as described (23). Hydrolysis of Az-LPAF used freezing and thawing to lyse mitochondria (40 mg). This was diluted into Hanks’ balanced salt solution containing 400 nM Az-LPAF and incubated in triplicate at 37 °C for the stated times before the reaction was stopped with methanol and the lipid recovered by extraction (24). [3H]PAF was used as an internal standard. Quantitation of samples or defined amounts of Az-LPAF used liquid chromatography/electrospray ionization/tandem mass spectrometry with a Quattro Ultima triple quadrupole mass spectrometer (Micromass, Wythenshawe, UK) with collision-induced dissociation by argon gas. Analyses were performed using electrospray ionization in the positive-ion mode with multiple reaction monitoring. The multiple reaction monitoring transitions used to detect the choline phospholipids were the mass to charge ratio (m/z) for the molecular ion [M + H]+ and the m/z 184 phosphocholine daughter ion.

Complex I Activity Measurement

Mitochondria were frozen and thawed before these lysed mitochondria (1 mg) were added to 1 ml of a reaction mixture containing 15 mM Hepes, pH 7.3, 3.5 mM phosphate, 50 mM KCl, 80 μM coenzyme-Q10, and the stated amount of Az-LPAF. The reaction mixture was equilibrated at room temperature (3 min) prior to the addition of 0.2 mM NADH to initiate the reaction. NADH oxidation was monitored at 340 nm at 37 °C over time.

Mitochondrial Membrane Potential

Rat liver mitochondria (0.1 mg/ml) were mixed with 0.5 μM TMRM, a positively charged fluorescent dye accumulated by energized mitochondria, and the fluorescence intensity of the organelle was determined using stirred, thermostatically controlled cuvettes. Fluorescence was determined using excitation wavelengths of 546 and 573 nm and an emission wavelength of 590 nm. The ratio between these signals reflects mitochondrial membrane potential. Changes in mitochondrial NADH levels were determined by incubating mitochondria (0.4 mg/ml) at room temperature in medium composed of 250 mM sucrose, 10 mM Hepes (pH 7.4), 25 mM EGTA, and 2 mM phosphate at 24 °C as fluorescence was determined by excitation at 340 nm with emission monitored at 460 nm. Fluorescence measurements used an LPS-200B fluorometer (Photon Technologies International, Lawrenceville, NJ) operated by FeliX32 software with slit widths for both emission and excitation set at 2.5 nm.

Flow Cytometry

HL-60 cells were washed and resuspended in serum-free RPMI 1640 and incubated with the stated amount of Az-LPAF for 4 h. One aliquot of washed cells was irradiated by 400 J/m²

2 The abbreviations used are: TMRM, tetramethylrhodamine methyl ester; BEL, bromeno lactone; CCCP, carbonyl cyanide 3-chlorophenylhydrazone; PAF, platelet-activating factor; iPLA2, calcium-independent phospholipase A2; PBS, phosphate-buffered saline; BSA, bovine serum albumin; Az-LPAF, alkyl azelaoyl glycerophosphocholine (azelaoyl lysoPAF).
(measured using a IL1700 radiometer and a SED240 UVB detector; International Light, Newburyport, MA) for 5 min, whereas a third aliquot remained unstimulated prior to the 4-h culture.

**Apoptosis Quantification**—To measure surface phosphatidylserine and the ability to exclude a nuclear dye, HL-60 cells were washed with PBS and stained with Annexin V-Alex 488 (1/100 dilution) and 1 μg/ml propidium iodide in binding buffer (140 mM NaCl, 10 mM Hepes, pH 7.4, and 2.5 mM CaCl₂) for 15 min at room temperature. At the end of this incubation the samples were diluted 4-fold with binding buffer and fluorescence was assessed by two-color flow cytometry.

**Assessment of Mitochondrial Potential**—Mitochondrial membrane potential was detected by incubating the cells with the potentiometric dye JC-1 (10 μg/ml) for 15 min, before the cells were washed twice and resuspended for two-color flow cytometry.

**RESULTS**

The Truncated Phospholipid Az-LPAF Initiates Apoptosis and Depolarizes Intracellular Mitochondria—Oxidatively modified phospholipids affect cell function through G-protein coupled and nuclear hormone receptors, but also act in currently undefined, receptor-independent ways. For example, the truncated phospholipid Az-LPAF stimulates PAF receptor signaling (25), but low micromolar concentrations of this oxidatively truncated phospholipid rapidly depolarized HL-60 cell mitochondria (Fig. 1A). Because these cells do not express the PAF receptor (26), mitochondria are one non-receptor dependent target of the oxidized phospholipids. The electrochemical potential across the mitochondrial inner membrane in this experiment was monitored by JC-1 fluorescence. This cationic dye is accumulated by polarized mitochondria where its high concentration promotes aggregation, changing its green fluorescence to a reddish orange. Two-color flow cytometry of HL-60 cells showed that as little as 1.25 μM Az-LPAF doubled the number of cells with dysfunctional mitochondria. The number of these cells increased as the concentration was increased to 5 μM Az-LPAF, rivaling that of the positive control, UVB irradiation.

HL-60 cells exposed to these same concentrations of Az-LPAF also lost the phospholipid asymmetry of their plasma membrane, an early event in apoptotic cell death. The data show (Fig. 1B) that over half of the cells exposed to 5 μM Az-LPAF displayed phosphatidylserine on their surface, which was comparable with the 40% of cells that expressed phosphatidylserine on their surface after UVB irradiation. Few cells exposed to either Az-LPAF or UVB at this time had progressed to the point where they were unable to exclude propidium iodide from their nucleus.

Az-LPAF Depolarizes Isolated Mitochondria—We determined whether Az-LPAF physically affected mitochondrial function using isolated rat liver mitochondria. Isolated mitochondria were provided glutamate and malate to oxidize (shown by the first arrow, Fig. 2A), which increased the transmembrane potential detected by increased TMRM fluorescence. The addition of 2.5 μM Az-LPAF at the time shown by the second arrow produced a rapid, but partial, loss of mitochondrial potential. There was an immediate and extensive loss of fluorescence when this truncated phospholipid was added at a final concentration of 5 μM. Complete depolarization of the isolated mitochondria was achieved by addition of the protonophore CCCP (third arrow, Fig. 2A). This agent had no effect on mitochondria previously exposed to 5 μM Az-LPAF, so this concentration of the truncated phospholipid had completely depolarized the organelle.

Az-LPAF, although the Most Effective Truncated Phospholipid, Is Not the Sole Lipid to Depolarize Mitochondria—We first determined whether the sn-1 ether bond of Az-LPAF was required for mitochondrial depolarization. We found that Az-LPAF was about twice as effective as its diacyl homolog 1-palmitoyl-2-azelaoyl-sn-glycero-3-phosphocholine (95 ± 2% depolarization by 5 μM Az-LPAF versus 40 ± 15% for diacyl palmitoyl azelaoyl glycerophosphocholine). We next tested related phospholipids and lysophospholipids, and many of these also caused a rapid loss of the energy gradient across intact mitochondria (Fig. 2B). Phospholipids with very short sn-2 residues (e.g. those with the 2-carbon acetyl residue of PAF and carbamoyl-PAF, and the 4-carbon long residues of butyroyl-LPAF) reduced the transmembrane potential by about one-third, which was similar to the effect of lyso-PAF that lacks any sn-2 residue. Phospholipids containing sn-2 residues 5 carbon atoms long (e.g. palmitoyl glutaroyl and palmitoyl oxovalleryl phosphatidylcholines) that, like Az-LPAF, possess ω-oxy functions on the fragmented sn-2 residue, were also no more than one-third as effective as Az-LPAF. Apparently, the combination of an ω-acidic function in conjunction with a medium length sn-2 acyl chain is particularly devastating to mitochondrial function. Only the strongly acidic phospholipid lysophosphatidic acid approached Az-LPAF effectiveness in depolarizing mitochondria.

Mitochondria Are Not Irretrievably Damaged by Az-LPAF—Amphipathic truncated phospholipids bearing a carboxyl function are detergents that disrupt membrane structure (14, 27), and they physically bind to cytochrome c (28). We therefore determined whether Az-LPAF irreversibly damaged mitochondrial function, as would be the case after solubilization, by depleting mitochondria of this lipid by sequestering it with excess albumin. Addition of Az-LPAF to mitochondria oxidizing glutamate and malate resulted, as before, in an immediate loss of transmembrane potential (Fig. 3A). Addition of BSA to the Az-LPAF de-energized mitochondria produced an immediate recovery of most, although not all, of the lost potential energy. Mass spectrometry (not shown) confirmed that an albumin wash removed 80 to 90% of the Az-LPAF associated with mitochondria. These data show that Az-LPAF must be continuously present to fully depolarize mitochondria.

We determined whether a transmembrane potential gradient, perhaps aiding internalization of the lipid, was essential for depolarization by Az-LPAF by treating mitochondria with Az-LPAF in the absence of oxidizable substrates. We found (Fig. 3B) that the resting transmembrane potential was rapidly and fully depolarized by CCCP and by about half with Az-LPAF. Albumin alone did not affect organelle function, but the introduction of albumin induced a small increase in transmembrane potential of mitochondria exposed to the lipids Az-LAPF or

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We determined whether the permeability transition pore participated in the loss of barrier function using isolated mitochondria which recovered intact electron transport lipoprotein complexes (29), to find all macromolecular complexes were still present after exposure to Az-LPAF (not shown). Staining these gels with the oxidizable substrate nitro blue tetracium dye indicated that complex I function was normal after Az-LPAF exposure (data not shown).

**Mitochondria Metabolize Az-LPAF**—Mitochondria contain iPLA₂ that protects against staurosporine-induced apoptosis (30). Mitochondria also express the calcium-dependent phospholipase A₂γ that hydrolyzes phosphatidylcholine and then can also reacylate the resulting lyso-phosphatidylcholine (31). We determined whether mitochondria also displayed PAF acetylhydrolase-like activity, that is, the ability to hydrolyze PAF in the absence of Ca²⁺. We found that the fraction in a sucrose density gradient containing intact mitochondria (fraction V) effectively hydrolyzed PAF, and that this activity was displayed by the other subcellular fractions (Fig. 4A). The mechanism-based iPLA₂ inhibitor BEL (32) reduced PAF hydrolysis in fraction V by half, indicating that mitochondrial iPLA₂ was able to hydrolyze short-chain phospholipids. Plasma (33) and type II (34) PAF acetylhydrolases additionally will hydrolyze oxidized phospholipids such as Az-LPAF. The ability to hydrolyze this truncated phospholipid was also present in mitochondria. Thus, isolated mitochondria completely catabolized Az-LPAF by 10 min (Fig. 4B), with an increase in lyso-PAF (Fig. 4C). Lyso-PAF was not stable in isolated mitochondria and its concentration fell over the subsequent 10 min.

**Az-LPAF Opens the Mitochondrial Permeability Transition Pore**—One route to loss of membrane potential, with subsequent irreversible effects, is through the opening of the mitochondrial permeability transition pore that allows molecules below ~2000 daltons to move across the inner membrane. To determine whether Az-LPAF opened this channel, we loaded HL-60 cells with the fluorescent dye calcein, which distributes between mitochondria and cytoplasm, and treated them with either cobalt chloride or buffer. The cobalt cation quenches calcein fluorescence, but cannot do so when an intact permeability barrier excludes the ion from the mitochondrial matrix. We found that, first, this dye was retained by cells exposed to Az-LPAF, which shows that the plasma membrane barrier function was uncompromised by the truncated phospholipid. Second, we found that the internalized dye content was the same between control and Az-LPAF-treated cells (Fig. 5A, left). We also found that the cobalt ion had no effect on the fluorescence of control cells, but that fluorescence in the majority of cells exposed to Az-LPAF was sharply reduced by cobalt exposure (Fig. 5A, right). This means the cobalt ion had access to the inner mitochondrial compartment, which in turn implies that the mitochondrial permeability barrier had been breached.

We determined whether the permeability transition pore participated in the loss of barrier function using isolated mitochondria...
treated with varied concentrations of cyclosporin A to block the cyclophilin D component of the pore. This inhibitor produced a partial, concentration-dependent increase (Fig. 5B) in mitochondrial potential in mitochondria that would have been completely depolarized by 5 μM Az-LPAF. Cyclosporin A also reduced mitochondrial swelling secondary to collapse of the electrical potential, but again was only partially effective in this (not shown).

Opening of the permeability transition pore allows NADH to leach from mitochondria, compromising their function. To test for the effect of Az-LPAF on mitochondrial NADH content we treated mitochondria with cyclosporin A, or not, and then with 5 μM Az-LPAF. We found (Fig. 5C, left, first arrow) that there was a time-dependent loss of NADH fluorescence, and that this change was less severe when the permeability transition pore was blocked by cyclosporin A. We then added CCCP (second arrow) to stimulate maximal electron transport and this, as expected, pulled electrons from NADH causing a rapid loss of fluorescence at 340 nm as non-fluorescent NAD⁺ accumulated. Finally, we added rotenone (third arrow) to block the flow of electrons from complex I to complex II. In this case, NADH produced by glutamate and malate oxidation cannot be reduced by the electron transport chain. Thus, rotenone addition allowed a full recovery of NADH fluorescence when the permeability transition pore was blocked by cyclosporin A, but allowed only a partial recovery of NADH fluorescence in mitochondria exposed to Az-LPAF when the permeability transition pore was not blocked.

Az-LPAF-induced Mitochondrial Dysfunction Is Regulated by Bcl-2 Family Members—Bcl-XL is a member of the Bcl-2 family that physically interacts with the mitochondrial outer membrane and suppresses apoptosis (35). Overexpressing Bcl-XL in HL-60 cells conferred a significant protection from Az-LPAF-induced apoptosis as shown by the reduced abundance of cell surface phosphatidylserine (Fig. 6A). Bcl-XL overexpression also reduced the loss of mitochondrial membrane potential in cells exposed to Az-LPAF, although Bcl-XL overexpression itself modestly decreased mitochondrial function (Fig. 6B).

The protection afforded by overexpressing Bcl-XL was a direct effect on mitochondria because the protective effect remained with mitochondria isolated from the overexpressing
Exposing mitochondria isolated from control HL-60 cells to Az-LPAF in 1.25 mM increments produced a progressive loss of transmembrane potential with each addition of Az-LPAF (Fig. 6C). This loss of transmembrane potential was greatly reduced when the mitochondria were obtained from Bcl-XL overexpressing cells. For example, when mitochondria from control cells were nearly completely depolarized by 3.75 mM Az-LPAF, mitochondria from Bcl-XL expressing cells still retained about 50% of their charge.

Bid is a pro-apoptotic member of the Bcl-2 family that uniquely possesses phospholipid transferase activity. Accordingly, we found that recombinant murine Bid (rBid) sensitized the mitochondria isolated from wild type animals to the effects of small amounts of Az-LPAF (Fig. 7A). Bid itself had no effect on mitochondrial potential, nor did 1 mM Az-LPAF, but the combination of the two resulted in a steady loss of the accumulated charge.

Conversely, a lack of Bid suppressed mitochondrial sensitivity to Az-LPAF. Mitochondria isolated from the livers of Bid knock-out animals behaved like mitochondria from control mice when given oxidizable substrates, but were depolarized at a much slower rate than their normal counterparts when exposed to Az-LPAF (Fig. 7B). Ultimately, when mitochondria from wild type animals had been completely depolarized by Az-LPAF, mitochondria from Bid null animals still retained half their original membrane potential. Mitochondria from Bid knock-out animals remained sensitive to Bid, however, because the addition of recombinant Bid sensitized these organelles to low concentrations of Az-LPAF (Fig. 7C). Here, the combination of Az-LPAF and recombinant Bid initiated a time-dependent loss of membrane potential by mitochondria isolated from Bid-/- animals that did not occur when just Az-LPAF was present.

**DISCUSSION**

The truncated phospholipid Az-LPAF rapidly enters cells, and a significant portion of this internalized phospholipid associates with mitochondria (13). Here, we find that mitochondria exposed to this oxidatively truncated phospholipid rapidly lose cells. Exposing mitochondria isolated from control HL-60 cells to Az-LPAF in 1.25 mM increments produced a progressive loss of transmembrane potential with each addition of Az-LPAF.
their transmembrane potential at least partially through opening of the permeability transition pore. As with other apoptotic stimuli, Bcl-XL protected mitochondria from Az-LPAF, whereas Bid augmented this mitochondrial dysfunction. What we did not find in this study was that truncated phospholipids “ruined” the membrane structure (36). The critical site of bilayer perturbation by the various truncated phospholipids was at the mitochondrial membrane, and not at the plasma membrane (14) with (15) or without (18) the participation of acid sphingomyelinase that hydrolyzes sphingomyelin to non-bilayer forming ceramide. We conclude this because cells exposed to Az-LPAF retained cytoplasmic fluorescent dyes, and did not allow entry of extracellular Ca$^{2+}$ (not shown). Although the barrier to NADH or cobalt cations presented by mitochondrial membranes was functionally compromised by Az-LPAF, the mitochondrial membrane organization was not impaired to the extent that the electron transport chain was compromised. Moreover, washing the organelle with albumin to remove this phospholipid allowed the mitochondria to rapidly recover most of their membrane potential. This shows that mitochondrial depolarization primarily was an ongoing process that required the continual presence of Az-LPAF to reduce transmembrane potential. The small amount of non-reversible change induced by Az-LPAF might result from the release of a portion of matrix NADH.

Model systems show that truncated phospholipids like Az-LPAF greatly perturb the membrane bilayer structure because the ionizable carboxylate function orients the residue toward the solvent (8–10) rather than into the hydrophobic phase. This extension of the sn-2 residue into the solvent disrupts phospholipid packing, distorts the bilayer, causes phase separation of normal and truncated phospholipids (8, 9, 37), and changes membrane surface potential (8). Phospholipid oxidation products that disrupt membrane structure and function in these ways are physiologically relevant because these types of damaged phospholipids are present in mitochondria isolated from the livers of animals oxidatively stressed by CCl$_4$ intoxication (36), and are generated by mitochondria themselves during state IV respiration (38).
Az-LPAF was the most effective depolarizing phospholipid we examined, and, interestingly, was more effective than glutaroyl phosphatidylcholine derived from oxidative fragmentation of sn-2 arachidonoyl residues. The sn-2 glutaroyl fragment, like the azelaoyl fragment derived from oxidative cleavage of the more abundant linoleoyl residues, contains an ionizable carboxyl function at the ω end of the shortened sn-2 fatty acyl fragment. The glutaroyl fragment is shorter than the azelaoyl fragment by 4 methylene molecules, and the decreased hydrophobicity of the glutaroyl residue had a less severe effect on bilayer organization than the longer azelaoyl fragment (39). Still, glutaroyl phosphatidylcholine will induce apoptotic cell death (40) and, like Az-LPAF, is internalized (12, 41). However unlike Az-LPAF that was primarily found in mitochondria (13), fluorescent glutaroyl phosphatidylcholine preferentially accumulates in lysosomes (12). Whether this disparate localization reflects differences in the length of the sn-2 fragment or structural differences, e.g. the sn-1 ether bond of Az-LPAF or the presence of a bulky fluorescent group of the glutaroyl phosphatidylcholine probe, remains to be determined.

Disruption of phospholipid packing by Az-LPAF intercalation into mitochondrial membranes was not the sole agent of mitochondrial dysfunction because opening of the mitochondrial permeability transition pore had a role in extending the loss of the transmembrane potential. Thus we found that addition of cyclosporin A, which occludes the pore through its interaction with cyclophilin D (42), protected a portion of the transmembrane potential from the effect of Az-LPAF. Free fatty acids, which like Az-LPAF contain a full negative charge on a carboxylate function, depolarize mitochondria, and do so in part by opening the permeability transition pore in a cyclosporin-dependent way (43).

In fact, the charge on these lipids may not be the only, or even the major, element in opening the pore because ceramide also depolarizes mitochondria in a fashion that is reduced by about half by the presence of cyclosporin A (44). This suggests the common element of these lipids is their ability to physically intercalate into mitochondrial membranes and change the highly ordered nature of these membranes. The central effect of membrane organization on pore opening is reinforced by the observation that the membrane intercalating peptide mastoparan also opens a cyclosporin-sensitive permeability transition pore (45). Indeed gangliosides GD3 and GM3, which normally reside in cholesterol- and sphingomyelin-rich microdomains at serine.

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**FIGURE 6.** The anti-apoptotic protein Bcl-XL protects cells and isolated mitochondria from Az-LPAF-induced depolarization and apoptosis. A, Bcl-XL suppresses Az-LPAF-induced surface expression of phosphatidylserine. HL-60 cells stably transfected with human Bcl-XL or empty vector were treated with 5 μM Az-LPAF or buffer for 6 h as described in the legend to Fig. 1. The cells were then stained with Alexa 488-conjugated annexin V to detect externalized phosphatidylserine and propidium iodide before cellular fluorescence was quantified by flow cytometry also as described in the legend to Fig. 1 (n = 3).

B, Bcl-XL suppressed Az-LPAF-induced depolarization of mitochondria within HL-60 cells. HL-60 cells stably transfected with human Bcl-XL or empty vector were incubated with 5 μM Az-LPAF for 4 h, and then loaded with the dye JC-1 for 30 min before monomeric fluorescence in FL1 (x axis) and mitochondria internalized dye was assessed in FL2 (y axis) (n = 3).

C, Bcl-XL protects isolated mitochondria from Az-LPAF-induced depolarization. Mitochondria isolated from HL-60 cells expressing Bcl-XL or empty vector were labeled with TMRM as described in the legend to Fig. 2, and given glutamate and malate at the times shown by the first arrow. Az-LPAF was then added in 1.25 μM increments at the times shown by subsequent arrows, and then the mitochondria were completely depolarized by the addition of CCCP (n = 3).
Mitochondria have the ability to protect themselves and limit their exposure to oxidatively truncated phospholipids because they were able to hydrolyze short-chain phospholipids PAF and Az-LPAF. PAF acetylhidrolases specifically hydrolyze short-chain phospholipids (33, 34). Whether iPLA₂ also can hydrolyze these lipids has not been tested, but specific inhibition of this enzyme showed that almost half of the PAF hydrolyase activity of mitochondria could be ascribed to this enzyme. iPLA₂ reduces peroxide-initiated cell death (47) and protects mitochondria against oxidative damage (30). This suggests turnover of oxidatively damaged phospholipids, including truncated phospholipids, and preserves mitochondrial function and cell viability.

The permeability transition pore (48) and mitochondrial transmembrane potential are regulated by Bcl-2 family members (49–52), and we found that Bcl-X₅ and the BH3-only protein Bid exerted opposing effects on the sensitivity of mitochondria to Az-LPAF. We observed that neither intact Bid nor low concentrations of Az-LPAF affected mitochondrial transmembrane potential, but their combination did so. Because Bid interacts with microsomal lipid microdomains (46), it is well positioned to promote the intercalation of Az-LPAF into mitochondrial membranes. It is notable that Bid, at least after proteolytic cleavage to truncated Bid, induces cytochrome c escape in a way that is not dependent on the permeability transition pore (53). Potentially, this unidentified mechanism also participates in the pore-independent effect of Az-LPAF.

Bid, alone among Bcl-2 family members, displays weak homology to plant lipid transfer proteins (54), and in fact functions as a lipid transporter. Bid enhances the mobility of acidic phospholipids (55, 56) and lysophosphatidic acid (22), but not unesterified fatty acids. A mutated Bid that does not bind lysophospholipids also is not pro-apoptotic (22). Lipid transfer activity is present whether or not Bid has been cleaved to its truncated form (22), and we found that full-length Bid promoted mitochondrial dysfunction by Az-LPAF. This then means that sensitivity to oxidatively truncated phospholipids is not regulated by Bid cleavage. Instead, it is the accumulation of oxidatively truncated phospholipids, e.g. during CCl₄ intoxication (36), that determines whether mitochondria depolarize, swell, and release cytochrome c to complete the apoptosome under conditions that promote formation of phospholipid oxidation products.

These studies focused on levels of truncated phospholipid sufficient to evoke large changes in mitochondrial function associated with apoptotic events. Our data, however, also show that the effect of these lipids is progressive and that lower levels of exposure still induce noticeable decreases in mitochondrial polarization. Truncated phospholipids, and lysophosphatidic acid, are uncoupling agents that are more effective in this than other lipid uncoupling agents such as fatty acids, hydroperoxy fatty acids, or their Co-A thioesters (57, 58). Mitochondrial uncoupling reduces ATP generation that can compromise cellular function, but also reduces superoxide production and so may be protective during some pathologic challenges (59).
Oxidatively truncated phospholipids are specifically catabolized by PAF acetylhydrolase, either within or outside cells, to soluble fatty acid fragments and lysophospholipids (60, 61). This is a critical protective mechanism against exogenous oxidized phospholipids, tert-butyl hydroperoxide oxidative stress, or Ca\(^{2+}\) overloading of neuronal cells because overexpression of PAF acetylhydrolase protects cells from all these apoptotic insults (13, 16, 34, 62, 63). This means that for these oxidative insults it is the intact oxidized phospholipid that is responsible for the apoptotic event. Conversely, neither the lysophospholipid nor the fragmented acyl chain products of the phospholipase reaction, or even proteinaceous components such as Bid, are sufficient to initiate the apoptotic cascade in these circumstances. We now understand that the target of these pro-apoptotic phospholipids are the mitochondria, and that Bid and Bcl-X\(_L\) are co-factors that act in opposing directions to alter mitochondrial sensitivity to oxidatively truncated phospholipids.

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REFERENCES

1. Buettner, G. R. (1993) Arch. Biochem. Biophys. 300, 535–543
2. Morrow, J. D., Hill, K. E., Burk, R. F., Nammour, T. M., Badr, K. F., and Roberts, L. J. (1990) Proc. Natl. Acad. Sci. U.S.A. 87, 9383–9387
3. Salomon, R. G., and Miller, D. B. (1985) Adv. Prostaglandin Thromboxane Leukot. Res. 15, 323–326
4. Tomkunova, A., Toujima, M., Yoshioka, Y., and Fukuzawa, K. (1996) Lipids 31, 1251–1258
5. Reis, A., Domingues, P., Ferrer-Correia, A. J., and Domingues, M. R. (2004) Rapid Commun. Mass Spectrom. 18, 2849–2858
6. Podrez, E. A., Poliakov, E., Sheng, Z., Deng, Y., Sun, M., Finton, P. J., Shan, L., Gugiu, B., Fox, P. L., Hoff, H. F., Salomon, R. G., and Hazen, S. L. (2002) J. Biol. Chem. 277, 38503–38516
7. Chen, X., Zhang, W., Laird, J., Hazen, S. L., and Salomon, R. G. (2008) J. Lipid Res. 49, 832–846
8. Sabatini, K., Mattila, J. P., Megli, F. M., and Kinnunen, P. K. (2006) Biochim. Biophys. Acta 1778, 2041–2050
9. Brooks, P. S., Pinner, A., Ramachandran, A., Coward, L., Barnes, S., Kim, H., and Darley-Usmar, V. M. (2002) Proteomics 2, 969–977
10. Zelczev, K., Zhao, C., Zhang, X. H., Song, K., and Ma, Z. A. (2006) J. Biol. Chem. 281, 22275–22288
11. Yamashita, A., Tanaka, K., Kamata, R., Kumazawa, T., Suzuki, N., Koga, H., Waku, K., and Sugura, T. (2009) Biochim. Biophys. Acta. in press
12. Hazen, S. L., Zupan, L. A., Weiss, R. H., Getman, D. P., and Gross, R. W. (1991) J. Biol. Chem. 266, 7227–7232
13. Stremler, K. E., Stafforini, D. M., Prescott, S. M., and McIntyre, T. M. (1991) J. Biol. Chem. 266, 11095–11103
14. Kono, N., Inoue, T., Yoshiida, Y., Sato, H., Matsuue, T., Itabe, H., Niki, E., Aoki, J., and Arai, H. (2008) J. Biol. Chem. 283, 1628–1636
15. Kroemer, G. (1999) Biochem. Soc. Symp. 66, 1–15
16. Megli, F. M., and Sabatini, K. (2004) FEBS Lett. 573, 68–72
17. Megli, F. M., Russo, L., and Sabatini, K. (2005) FEBS Lett. 579, 4577–4584
18. Megli, F. M., and Sabatini, K. (2003) FEBS Lett. 550, 185–189
19. Megli, F. M., and Russo, L. (2008) Biochim. Biophys. Acta 1778, 143–152
20. Fruhwirth, G. O., Moutzui, A., Loidl, A., Inoglic, E., and Hermetter, A. (2006) Biochim. Biophys. Acta 1761, 1060–1069
21. Rhodes, S., Gruri, R., Brameshuber, M., Hermetter, A., and Schütz, G. J. (2009) J. Biol. Chem. 284, 2258–2265
22. Waldmeir, P. C., Zimmermann, K., Qian, T., Tintelnot-Blomley, M., and Lemasters, J. J. (2003) Curr. Med. Chem. 10, 1485–1506
23. Wieckowski, M. R., Birczka, D., and Wojtczak, L. (2000) FEBS Lett. 484, 61–64
24. Novgorodov, S. A., Szule, Z. M., Lutberto, C., Jones, A. J., Bielawski, I., Bielawska, A., Hannun, Y. A., and Obeid, L. M. (2005) J. Biol. Chem. 280, 16096–16105
25. Pfeffer, D. R., Gudz, T. I., Novgorodov, S. A., and Erdahl, W. L. (1995) J. Biol. Chem. 270, 4923–4932
26. Garofalo, T., Giammarioli, A. M., Missa, R., Tinari, A., Manganelli, V., Gambardella, L., Pavan, A., Malorni, W., and Sorice, M. (2005) Cell Death Differ. 12, 1378–1389
27. Kinsey, G. R., Blum, J. L., Covington, M. D., Cummings, B. S., McHowat, J., and Schnellmann, R. G. (2008) J. Lipid Res. 49, 1477–1487
28. Vander Heiden, M. G., and Thompson, C. B. (1997) Nat. Cell Biol. 1, E209–E216
29. Shimizu, S., Eguchi, Y., Kamiike, W., Funahashi, Y., Mignon, A., Aramone, V., Matsuda, H., and Tsujimoto, Y. (1998) Proc. Natl. Acad. Sci. U.S.A. 95, 1455–1459
30. Matsuysama, S., Lloips, J., Deveraux, Q. L., Tsien, R. Y., and Reed, J. C. (2000) Nat. Cell Biol. 2, 318–325
31. Shimizu, S., Shinohara, Y., and Tsujimoto, Y. (2000) Oncogene 19, 4309–4318
32. Vander Heiden, M. G., Chandel, N. S., Williamson, E. K., Schumacker, P. T., and Thompson, C. B. (1997) Cell 91, 627–637
33. Kim, T. H., Zhao, Y., Barber, M. I., Kuhlarsky, D. K., and Yin, X. M. (2000) J. Biol. Chem. 275, 39474–39481
34. Degl Esposti, M. (2002) Biochim. Biophys. Acta 1553, 331–340
35. Esposti, M. D., Erler, J. T., Hickman, J. A., and Dive, C. (2001) Mol. Cell Biol. 21, 7268–7276
56. Zhai, D., Miao, Q., Xin, X., and Yang, F. (2001) Eur. J. Biochem. 268, 48–55
57. Korge, P., Honda, H. M., and Weiss, J. N. (2003) Am. J. Physiol. Heart Circ. Physiol. 285, H259–H269
58. Jaburek, M., Miyamoto, S., Di Mascio, P., Garlid, K. D., and Jezek, P. (2004) J. Biol. Chem. 279, 53097–53102
59. Echtay, K. S. (2007) Free Radic. Biol. Med. 43, 1351–1371
60. Stafforini, D. M. (2009) Cardiovasc. Drugs Ther. 23, 73–83
61. McIntyre, T. M., Prescott, S. M., and Stafforini, D. M. (2009) J. Lipid Res. 50, (Suppl.) S255–S259
62. Matsuzawa, A., Hattori, K., Aoki, J., Arai, H., and Inoue, K. (1997) J. Biol. Chem. 272, 32315–32320
63. Hirashima, Y., Ueno, H., Karasawa, K., Yokoyama, K., Setaka, M., and Takaku, A. (2000) Brain Res. 885, 128–132