Pseudomonas aeruginosa homoserine lactone triggers apoptosis and Bak/Bax-independent release of mitochondrial cytochrome C in fibroblasts

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Summary

Pseudomonas aeruginosa use N-(3-oxododecanoyl)-homoserine lactone (C12) as a quorum-sensing molecule to regulate gene expression in the bacteria. It is expected that in patients with chronic infections with P. aeruginosa, especially as biofilms, local [C12] will be high and, since C12 is lipid soluble, diffuse from the airways into the epithelium and underlying fibroblasts, capillary endothelia and white blood cells. Previous work showed that C12 has multiple effects in human host cells, including activation of apoptosis. The present work tested the involvement of Bak and Bax in C12-triggered apoptosis in mouse embryo fibroblasts (MEF) by comparing MEF isolated from embryos of wild-type (WT) and Bax−/−/Bak−/− (DKO) mice. In WT MEF C12 rapidly triggered (minutes to 2 h): activation of caspases 3/7 and 8, depolarization of mitochondrial membrane potential (ΔΨmito), release of cytochrome C from mitochondria into the cytosol, blebbing of plasma membranes, shrinkage/condensation of cells and nuclei and, subsequently, cell killing. A DKO MEF line that was relatively unaffected by the Bak/Bax-dependent proapoptotic stimulants staurosporine and etoposide responded to C12 similarly to WT MEF: activation of caspase 3/7, depolarization of ΔΨmito and release of cytochrome C and cell death. Re-expression of Bak or Bax in DKO MEF did not alter the WT-like responses to C12 in DKO MEF. These data showed that C12 triggers novel, rapid proapoptotic Bak/Bax-independent responses that include events commonly associated with activation of both the intrinsic pathway (depolarization of ΔΨmito and release of cytochrome C from mitochondria into the cytosol) and the extrinsic pathway (activation of caspase 8). Unlike the proapoptotic agonists staurosporine and etoposide that release cytochrome C from mitochondria, C12’s effects do not require participation of either Bak or Bax.

Introduction

Pseudomonas aeruginosa are opportunistic bacteria that accumulate and form biofilms in the lungs of cystic fibrosis (CF) patients. P. aeruginosa uses N-(3-oxododecanoyl)-homoserine lactone (C12) as a quorum-sensing molecule to control bacterial gene expression, including production of biofilms and virulence factors (Schuster and Greenberg, 2006; Rumbaugh, 2007; Irie and Parsek, 2008). C12 elicits proapoptotic effects in host cells that are likely to be relevant biologically since it is expected that in patients exposed in CF lungs, including epithelial cells, leucocytes, fibroblasts and endothelial cells (Smith et al., 2001; 2002; Tatada et al., 2003; Li et al., 2004; 2009; Kravchenko et al., 2006; 2008; Jacobi et al., 2009; Schwarzer et al., 2012). Inflammatory responses may vary depending on whether C12 was added in the absence (DiMango et al., 1995; Smith et al., 2001; 2002; Jacobi et al., 2008) or presence (Kravchenko et al., 2008) of other agonists.

The goal of the present study was to test whether Bak and Bax were involved in mediating apoptotic responses of mouse embryo fibroblasts (MEF) to C12. We used MEF for these studies for two reasons. First, fibroblasts are relevant biologically since it is expected that in patients with chronic infections with P. aeruginosa biofilms local [C12] will be high and, since C12 is lipid soluble, it will diffuse from the airways across and into the epithelium and underlying fibroblasts. Second, MEF are advantageous experimentally since MEF from wild-type (WT) and Bak−/−/Bax−/− (DKO) mice are readily available, and complete knockout of these genes in airway epithelia using siRNA or similar method is extremely difficult to achieve.
Bak/Bax are commonly involved in apoptosis, particularly in the release of cytochrome C from mitochondria to the cytosol to trigger downstream activation of caspases 3/7 and 9 (Degenhardt et al., 2002; Mikhailov et al., 2003; Oakes et al., 2003; Scorrano et al., 2003; Wang et al., 2011). Schwarzer et al. (2012) previously showed that C12 rapidly triggered events commonly associated with activation of apoptosis (membrane blebbing, activation of caspases 3/7, 8 and 9; depolarization of mitochondria and release of cytochrome C). Recent experiments (Valentine et al., 2013a) that were reported while work for this article was in progress showed that C12 caused ZVAD-blocked cleavage and increased activity of caspase 3/7 and killing in MEF over 4–8 h. These experiments showed that these C12-triggered proapoptotic events were totally dependent on IRE1α and XBP-1 in MEF. The present experiments performed direct tests of the roles of Bak and Bax in C12-triggered apoptosis in MEF. We measured/assayed cellular and nuclear morphology, activities of caspases 3/7 and 8, cell killing and three different aspects of mitochondrial function/morphology in MEF from WT and DKO mice and also in DKO MEF that had been transduced to re-express Bak and/or Bax. The results indicated that C12 triggers multiple proapoptotic events in MEF, including release of cytochrome C from mitochondria into the cytosol, and Bak or Bax were not required for any of these effects. As far as we can tell, C12 is a unique bacterial product in that it is a membrane-permeant small molecule that triggers cytochrome C release from mitochondria without the involvement of Bak or Bax.

Results

Effects of C12 on caspase 3/7 and cell morphology in WT MEF

We tested for C12-induced apoptosis in WT MEF by performing assays of caspases, membrane blebbing and cellular and nuclear shrinkage. In dose–response experiments conducted after 1 h exposure, C12 increased caspase 3/7 activity at 10 and 50 μM in WT MEF, but not at 1 μM (Fig. 1). Control, untreated cells showed typical, flattened morphology with oval nuclei and BCECF distributed throughout the cytosol and nuclei. 50 μM C12 caused WT cells and nuclei to shrink/condense, and plasma membrane had multiple blebs containing BCECF, indicating continuity with the cytosol. These effects were very prominent by 2 h of treatment (Fig. 2).

Effects of C12 on caspases 3/7 and 8 and cell killing in WT, DKO and Bak or Bax-corrected MEF

The roles of Bak and Bak in apoptotic responses were tested using MEF from WT and DKO mice (Wang et al., 2011). DKO MEF were also transduced with retrovirus expressing Bak or Bax as a further test of the role of Bak and Bax in the MEF responses to C12. We first tested for Bak and Bax expression in the cell lines. Western blots showed that WT MEF and the GFP-transduced MEF (used as control for transductions) and Bak- and/or Bax-corrected DKO MEF expressed Bak and Bax as predicted (Fig. 3). C12 caused activation of caspases 8 and 3/7 after 2 h in both WT and DKO MEF (Fig. 4). Further, re-expression of Bak or Bax in DKO MEF did not change caspase 3/7 responses to C12 (Fig. 5). Staurosporine, a commonly used proapoptotic agonist, was also tested to assure that the DKO and Bak/Bax-corrected DKO MEF were behaving as predicted (Oakes et al., 2003). Staurosporine had no effect on caspase 3/7 in DKO MEF, but did activate caspase 3/7 activity in DKO + Bak and DKO + Bax MEF. We also tested whether the anti-apoptotic Bcl-2 affected C12-induced apoptosis. WT MEF were transduced with Bcl-2 using a retroviral vector (Supplementary Data Fig. S1A). However, C12-induced activation of caspase 3/7 occurred similarly in WT MEF and in WT MEF expressing either GFP (as control for transduction) or BCI-2 (Supplementary Data Fig. S2B).

Measurements of cell death (uptake of propidium iodide, PI) after 24 h treatment with C12 were also used to test the roles of Bax/Bak. PI responses to the common proapoptotic agonist etoposide were used to assure WT and DKO and Bak/Bax-corrected DKO MEF were behaving as predicted. Preliminary dose–response experiments showed that 50 μM C12 caused roughly 20% killing, and 100 and 150 μM C12 caused equivalent 70–75% killing of WT MEF (Zhao and Li, data not shown). We used 100 μM HSL triggers Bax/Bak-independent apoptosis

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C12 for cell killing experiments. Consistent with results measuring caspase 3/7 activity, C12 caused PI uptake (killing) of both WT and DKO MEF, and re-expression of Bak and/or Bax in DKO MEF did not change killing responses (Fig. 6A). Etoposide caused predictable cell killing in WT MEF and in DKO MEF that expressed Bak or Bax, but much less in DKO and DKO + GFP MEF (Fig. 6B). C12 also caused roughly equivalent, dose-dependent killing of WT and Bax−/−/Bak−/− cells in another cell line, NIH3T3 MEF (Fig. S2A and B). Overall, these data showed that C12 caused activation of caspase 3/7 (Fig. 5) and cell killing (Fig. 6) in both WT and DKO MEF, and C12-triggered responses in DKO MEF’s did not increase in DKO MEF in which Bak/Bax had been re-expressed.

Fig. 2. C12 causes cell and nuclear shrinkage in WT MEF. WT MEF were observed under DIC (A, C) or fluorescence (B, D) optics at 2 h. Fluorescence images of blue Hoechst-stained nuclei and green BCECF-stained cytosol were overlain using Adobe Photoshop. Control, untreated cells showed typical, flattened morphology with oval nuclei and BCECF distributed throughout the cytosol and nuclei. Compared to control conditions, C12-treated WT cells and nuclei were obviously shrunken, and plasma membrane had multiple blebs (arrows) containing BCECF. Images typical of > 15 each in three different coverglasses each.

Fig. 3. Expression of Bax and Bak in WT and DKO MEF. Western blots to identify Bak and Bax were performed on WT and DKO MEF transduced with GFP (served as control for transductions), Bak or Bax. Results typical of two similar.

Fig. 4. Effects of C12 on caspases 3/7 and 8 in WT and DKO MEF. WT and DKO MEF were exposed to 50 μM C12 for 2 h, then prepared for assays of caspase 8 (A) and caspase 3/7 (B). C12 activated caspases 3/7 and 8 in WT and in DKO MEF. Average ± SD, n = 3 experiments each. *P < 0.05 for comparisons between C12-treated versus control.
Testing roles of Bax and Bak in C12-induced depolarization of $\Delta\psi_{\text{mito}}$ and release of cytochrome C from mitochondria in MEF

Imaging of the voltage-sensitive dye JC-1 was used to test for Bak/Bax-dependent effects of C12 in mitochondria of MEF. JC1 accumulated in WT MEF in punctate patterns throughout the cytosol consistent with mitochondrial localization. Depolarization of $\Delta\psi_{\text{mito}}$ was assessed from the increase in fluorescence as JC1 was released from mitochondria into the cytosol and nucleus. In WT MEF C12 caused a slow increase of JC1 fluorescence over the course of 15 min (Fig. 7A), similar to effects in airway epithelial cells (Schwarzer et al., 2012). Complete depolarization of $\Delta\psi_{\text{mito}}$ was triggered by FCCP (10 $\mu$M) treatment at the end of each experiment (Fig. 7A). C12 also caused depolarization of $\Delta\psi_{\text{mito}}$ in DKO MEF (Fig. 7B) that was very similar to that in WT MEF (Fig. 7A). Averages from these time-course experiments are summarized in Fig. 8: C12 caused 70% depolarization of $\Delta\psi_{\text{mito}}$ (i.e. steady-state values in response to C12 compared to effects of FCCP 100%) in WT and DKO MEF.

We performed both immunofluorescence and ELISA assays to test whether C12’s proapoptotic effects included Bak/Bax-dependent release of cytochrome C from mitochondria into the cytosol (Goldstein et al., 2005). Typical images of control and C12-treated MEF are shown in Fig. 9. Under control conditions WT MEF exhibited characteristic cytochrome C staining throughout the cytosol and typical oval nuclei with smooth boundaries that excluded cytochrome C. In control conditions DKO MEF exhibited less distinct and more punctate staining of cytochrome C-labelled mitochondria, consistent with the mitochondrial morphology described in previous work (Karbowski et al., 2006). Cytochrome C was largely

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excluded from the nuclei of both WT and DKO MEF in control conditions. In WT and DKO MEF C12 caused cytochrome C staining to become diffuse in both the cytosol and nuclei (see arrows). Nuclei of C12-treated WT and DKO MEF were often shrunken and fragmented (arrowheads). These results indicated cytochrome C had been released from mitochondria into the cytosol and had diffused into the nuclei of WT and DKO MEF.

ELISA assays for cytochrome C were performed on cytosolic and mitochondrial fractions from WT and DKO MEF exposed to 50 μM C12 for 1 h. The same fractions were also analysed for the presence of mitochondria by performing Western blots with antibodies to the mitochondrial marker VDAC (voltage-dependent anion channel; Huang et al., 2013) and actin, as a loading control. C12 increased cytosol/mitochondria ratio of cytochrome C in WT and DKO MEF (Fig. 11A). Western blots showed that VDAC was present in the mitochondrial fractions but not in the cytosolic fractions of all cells under all conditions (Fig. 11B). Actin staining indicated that equal amounts of protein were loaded to each lane. Thus, Western blots were consistent with the idea that the C12-induced increase in cytosol/mitochondria cytochrome C ratio in WT and DKO MEF resulted from release of cytochrome C into the cytosol and not from an artefact resulting from ineffective cell fractionation or homogenization of mitochondria. Immunomicroscopy and ELISA data were therefore consistent in showing that C12 caused cytochrome C release from mitochondria into the cytosol in WT and DKO cells.

Discussion

An important conclusion from this study was that C12 rapidly triggered multiple events associated with apoptosis.
in MEF. C12 caused apoptosis-characteristic activation of caspases 3/7 and 8, release of cytochrome C from mitochondria into the cytosol, shrinkage of cells and nuclei and fragmentation of nuclei and blebbing of plasma membranes. A striking aspect of these effects of C12 was the rapidity of the proapoptotic responses: C12-induced depolarization of mitochondrial membrane potential began within 5 min and was complete within 20 min; release of cytochrome C from mitochondria into the cytosol and activation of executioner caspase 3/7 occurred within 60 min. In contrast, many other proapoptotic agonists require hours or even days for apoptosis to become manifest (Goldstein et al., 2000; 2005). C12 appears also to trigger similar apoptosis in other cell types, including mast cells, macrophages, neutrophils, airway epithelial cells and mammary epithelial cells (Tateda et al., 2003; Li et al., 2004; Kravchenko et al., 2006; Shiner et al., 2006; Schwarzer et al., 2012).

C12 induced killing of WT and DKO MEF after 24 h. Although PI uptake could have resulted from either necrosis or apoptosis, PI uptake also occurred in response to the proapoptotic agonist etoposide, a result consistent with the idea that C12-induced PI uptake may have similarly resulted from apoptosis. While C12-induced cell killing varied between 1.5- and 8-fold over control levels (Fig. 6), caspase activations varied between 8- and 25-fold activation over controls. This apparent quantitative discrepancy may result from differences in the methods. Caspase measurements were performed after 2 h on one set of cells while cell killing was performed after 24 h on cells that had different passage number and cell density, both of which could affect responses. Further tests of the quantitative relationship between C12-induced caspase activation and cell killing will require performing experiments over similar time-courses on cells in which growth patterns were con-
Despite this quantitative inconsistency in killing versus caspase assays, the data were clear in showing that C12 induced very rapid activation of proapoptotic responses.

A second major conclusion was that C12's proapoptotic effects occurred without the involvement of Bak or Bax. WT and DKO MEF exhibited equivalent depolarization of mitochondrial membrane potential and release of cytochrome C into the cytosol. C12 also induced equivalent activation of executioner caspase 3/7 and cell killing in WT and DKO MEF. Further, expressing Bak or Bax in DKO MEF did not increase C12's ability to activate caspase 3/7 or to kill MEF. In contrast, effects of staurosporine and etoposide, known activators of the intrinsic pathway to apoptosis (Degenhardt et al., 2002; Mikhailov et al., 2003) were dependent on Bak or Bax. Thus, staurosporine activated caspase 3/7 in WT and DKO MEF expressing Bak or Bax, but not in DKO MEF. And etoposide caused cell killing in WT and DKO MEF expressing Bak or Bax, but not in DKO MEF. Experiments on WT and DKO NIH3T3 cells (Supplementary Data Fig. S2) confirmed dose-dependent cell killing by C12 in WT and DKO MEF used for all the other experiments (Fig. 6A).

These results showing no involvement of proapoptotic Bak/Bax in C12-triggered apoptosis were consistent with the finding that overexpression of anti-apoptotic Bcl-2 in WT MEF also did not alter C12's ability to activate caspase 3/7. A previous study showed that C12-triggered depolarization of $\Delta \psi_{\text{mito}}$ and activation of caspases 3 and 8 and cell killing was reduced in Jurkat cells that overexpressed Bcl-2 (Tateda et al., 2003), consistent with the idea that Bcl-2 and related proteins including Bak and Bax were intimately involved in C12-triggered apoptosis. Different cell lines or methods or even genes in apparently matched cell lines may explain such apparent discrepancies. Recent work (Valentine et al., 2013a) showed that C12-induced apoptosis in MEF required IRE1$\alpha$ and XBP1, also consistent with the idea that something besides Bak and Bax was mediating C12-induced apoptosis.

The most striking and important results of this study were that C12 caused cytochrome C release from mitochondria into the cytosol equally well in WT and DKO MEF. It is generally thought that following activation by upstream BH3-only proteins and inactivation of Bcl-2 and Bcl-xl, Bak and Bax oligomerize in the outer membrane of mitochondria and facilitate release of cytochrome C into the cytosol (Bender and Martinou, 2013). C12's ability to release cytochrome C from mitochondria into the cytosol without the involvement of Bak and Bax is, as far as we can tell, novel for bacteria-derived products. The triterpenoid plant derivative betulnic acid causes similar release of cytochrome C from mitochondria and activation of apoptosis, an effect that appears to be mediated by the opening of the mitochondrial permeability transition pore or PTP (Mullauer et al., 2009; 2010). The role of the PTP in C12-induced apoptosis should be tested.

Although it is clear that C12's proapoptotic effects are Bak/Bax-independent, the specific molecules or pathways triggered by C12 remain unknown. C12 clearly affects mitochondria and downstream caspase 3/7, so it seems likely that C12 activates molecule(s) in the intrinsic pathway to apoptosis in MEF. In addition, C12 appears also to activate the extrinsic pathway (via caspase 8 activation), implicating a role for C12 activating a receptor like the TNF receptor or TNFR-like receptor (Speeckaert et al., 2012; de Wilt et al., 2013). Previous studies showed that C12 triggers events often associated with receptor...
activation: changes in activities of Stat 3 (Li et al., 2004), PPARγ (Jahoor et al., 2008), Ca2+ and cAMP signalling (Schwarzer et al., 2010; 2012), Ca2+-triggered phosphorylation (Vikström et al., 2009; 2010) and activation of IRE1α, splicing of XBP1 mRNA and production of XBP1s protein (Valentine et al., 2013a). However, the roles of and connections among these potential receptors, pathways and molecules to activation of the wide array of C12-induced proapoptotic responses remain unknown. A recent study showed that C12’s effects to inhibit NF-κB activation were blocked by several triazolo[4,3-a]quinolones with nM affinities, indicating that C12 elicits at least some of its effects selectively (Valentine et al., 2013b). Further tests of these compounds on C12’s activation of proapoptotic and inflammatory responses may provide insights into whether C12’s multiple effects result from one or multiple activation steps. Although the molecular mechanisms remain to be determined, it seems possible that C12, with its high lipid solubility, activates a pattern associated molecular pattern (PAMP) receptor or damage-associated molecular pattern receptor (DAMP) (Kumar et al., 2011) in the cytosol. In lungs or other tissues infected with P. aeruginosa biofilms, where local C12 > 10 μM, epithelia, dendritic cells, white cells, fibroblasts, neutrophils, smooth muscle and endothelial cells, may all exhibit increased proapoptotic signalling. Such effects may contribute to the persistence of P. aeruginosa infections in CF lungs.

Experimental procedures

Reagents

Unless otherwise specified, reagents and chemicals were obtained from Sigma. C12 was dissolved in DMSO and frozen in separate vials and then thawed for single experiments. Preliminary experiments showed that C12 lost potency with repeated thaw–freeze–thaw cycles.

Generation and culture of WT and DKO and Bax/Bak-corrected MEF

Uncloned population of wild-type and Bak−/−Bax−/− (DKO) MEF immortalized by SV40 large T antigen expression have been described previously (Wang et al., 2011). DKO MEF were resistant to the apoptosis stimuli etoposide. Bax/Bak-corrections of DKO MEF were performed. The cDNA of murine Bak or murine Bax was subcloned into the retroviral expression vector pBABE-IRES-EGFP with the marker protein enhanced green fluorescent protein (EGFP) expressing from an internal ribosomal entry site (IRES). To generate retrovirus, the packaging cell line human embryonic kidney (HEK)-293T was transfected with the plasmid carrying the respective gene of interest or the empty vector along with the retroviral helper plasmids pUVMC and pMDG2.0 using the jetPRIME® transfection reagent (Polyplus-transfection; New York, NY). Medium containing retrovirus was collected 48–72 h following transfection and supplemented with 10 μg ml−1 polybrene (Sigma; St. Louis, MO) to enhance infection efficiency. Bak, Bax or the control GFP alone was expressed in DKO MEF by retroviral infection. Infected cells were again subcloned by limited dilution to identify cells that expressed high levels of GFP.

Wild-type and Bak−/−Bax−/− murine fibroblasts immortalized by the NIH3T3 spontaneous immortalization method were a gift from Dr Wei-xing Zong (Zong et al., 2003). All cells were cultured in DMEM/high glucose medium (Mediatech; Manassas, VA) with 10% (v/v) fetal bovine serum (Gemi; West Sacramento, CA). 100 μl ml−1 penicillin and 100 μg ml−1 streptomycin (Mediatech). Cells were grown in a humidified 95/5% air/CO2 incubator at 37°C. The cells were passaged at 1:5–1:10 dilutions and the remaining cell suspension was seeded directly onto a 96-well, 24-well or 12-well tissue culture plate (BD Falcon, Bedford, MA) or onto coverglasses for imaging.

Western blot analysis of Bax and Bak

Whole-cell lysates with equal number of cells (5 × 10⁴) were loaded on a 4–12% Bis-Tris gel (Bio-Rad; Hercules, CA) and transferred onto PVDF membrane (Millipore; Billerica, MA). The membrane was incubated with appropriate primary or secondary antibodies either at 4°C overnight or at room temperature for 3 h in blotting buffer (1 × PBS + 0.2% tween-20) with 10% (w/v) non-fat dry milk (Bio-Rad). Protein levels were detected using the enhanced chemiluminescent detection system (Pierce; Rockford, IL). Antibodies used for Western blot analysis were anti-β-actin mAb (Sigma), anti-Bak pAb (Enzo; Farmingdale, NY), and anti-Bax pAb (Santa Cruz; Santa Cruz, CA).

Assay for caspases 3/7 and 8

Caspase 3/7 and caspase 8 activities were measured by cell-based homogeneous luminescent assays (Caspase-Glo, Promega, Madison, WI), in which a specific substrate that contains the tetrapeptide DEVD (specific for caspase 3/7) and LETD (specific for caspase 8) was cleaved by the activated caspases from the cells to release aminoluciferin reacting with the luciferase and resulting in the production of light. WT and DKO MEF were plated on a clear-bottom, white 96-well plate in 100 μl media per well for 4–5 days until they were confluent. During the experiment, cells were treated with drugs in the 37°C incubator for 60–120 min, depending on the experiment, or were left untreated as controls. On the same plate, some wells without cells but 100 μl of the same media served as blanks. After treatment, 100 μl reagent was added to each well with cells (treated or controls) and their media, or blank (media only). The plate was incubated at room temperature for 1 h on a shaker, and the end-point luminescence was measured in a plate-reading luminometer (LmaxI 384, Molecular Devices, Sunnyvale, CA). Data were background (blank) subtracted and averaged.

Cell viability assay

The indicated cell lines were plated (2 × 10⁴) in each well of a 48-well tissue culture plate and grown for 24 h. The different cell lines were treated with various concentrations of C12 or with 10 μM etoposide for a further 24 h, then incubated with Trypsin-
were measured at the end of experiments by treating cells with FCCP. Data have been reported as relative ratios. Maximum JC-1 ratios were corrected for background (region without cells). Images were corrected for background (region without cells).

City, CA) controlled both light exposure and collection of data. Measurements of Δψ_mito were made by diluting stock solutions into Ringer's solution at the temperature, and then washed three times with Ringer's solution to remove the extra dye. Dye-loaded cells were mounted onto a chamber on the stage of either a wide field imaging microscope or a spinning disk confocal imaging microscope. Cells were maintained at room temperature during the experiment. Treatments were made by diluting stock solutions into Ringer's solution at the concentrations stated in the text. Fluorescence imaging measurements of Δψ_mito were performed using equipment and methods that have been reported previously (Fu et al., 2007; Hybiske et al., 2007; Schwarzer et al., 2008). Briefly, a Nikon Diaphot inverted microscope was used with a 40× Neofluar objective (1.4 NA). A CCD camera collected emission images (em: > 510 nm for JC-1) during excitation [490 ± 5 nm for JC-1] using a filter wheel (Lambda-10, Sutter Instruments, Novato, CA). Axon Imaging Workbench 5.0 (Axon Instruments, Foster City, CA) controlled both light exposure and collection of data. Images were corrected for background (region without cells). Data have been reported as relative ratios. Maximum JC1 ratios were measured at the end of experiments by treating cells with 10 μM FCCP to completely depolarize Δψ_mito. Minimum JC1 fluorescence was defined as the starting value, and maximal depolarization was defined as fluorescence measured in presence of FCCP.

**Cytochrome C, nuclear and cellular morphology in fixed and stained MEF**

MEF grown on coverslips were left untreated or incubated with C12 (50 μM) for times mentioned in the text; cells were then rinsed with PBS, fixed for 30 min with 2% paraformaldehyde in PBS at 4°C, rinsed with PBS and permeabilized with 0.2% Triton X-100 in PBS for 15 min. After blocking with 2% BSA-PBS for 60 min, cells were incubated for 1 h at room temperature with an anti-cytochrome C (Invitrogen, Grand Island, NY) followed by thorough rinses with PBS (±0.05% Tween-20). Finally, cells were incubated for 1 h with an Alexa Fluor 546 anti-rabbit – or Alexa Fluor 488 anti-mouse secondary antibodies (Invitrogen, Grand Island, NY) followed by 10 min incubation with 1 μM Hoechst 33342. Fluorescence images were obtained using an inverted confocal microscope with a 63× objective (Zeiss 710). Quantification of resulting cytochrome C distribution was performed on these images using NIH Image. Using Adobe Photoshop, we outlined images of nuclei in blue images. These outlines were then overlaid on the cytochrome C (green images). Equal-sized areas of the outlined nuclei and cytosol (area outside the nuclei) were corrected for background (area without cells) and the nuclear/cytosol ratio was calculated for each cell: (nucleus – background)/(cytosol – background). At least 10 cells in three fields from each slide were measured and averaged for quantification of nuclear/cytosol ratio for cytochrome C.

**ELISA assays of cytosolic and mitochondrial cytochrome C**

ELISA assays of cytosolic cytochrome C were performed to test whether C12 altered cytosol/mitochondrial distribution of WT and DKO MEF, which were grown on plastic plates, rinsed with Ringer’s and exposed for 1 h to Ringer’s alone or Ringer’s + 50 μM C12. Cells were then homogenized and processed using differential centrifugation to isolate cytosolic and mitochondrial fractions as previously described (Frezza et al., 2007). Protein content of fractions was measured by Bradford and equal amounts of protein per well were assayed by Quantikine ELISA (R&D Systems, Minneapolis, MN) to quantify rat/mouse cytochrome C content. Aliquots of the same fractions were also analysed in Western blots using anti-VDAC (voltage-dependent anion channel, Huang et al., 2013) and anti-actin antibodies (Cell Signaling Technology, Danvers, MA, and MP Biomedicals, Solon, OH) to determine whether the cytosolic fractions had contamination from mitochondria that occurred during preparation of the cells and to assure equivalent protein loading into the lanes of the gel.

**Statistics**

Quantitative data were presented as averages ± SD, where n = number of different biological samples or experiments. Statistical comparisons between C12 treatment and control (untreated) were determined using t-tests for paired or unpaired samples as appropriate; P < 0.05 was considered statistically significant.

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Conflict of interest
The authors have no conflicts of interest.

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Supporting information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

**Fig. S1.** Bcl-2 expression in WT MEF and (lack of) effect on C12-induced activation of caspase 3/7. WT MEF grown on coverglasses (A) or in plastic dishes (B) were infected with retroviruses expressing GFP or Bcl-2 + GFP. The retroviral vector DNA plasmids (MSCV-PIG) expressing GFP or Bcl2-IRES-GFP were obtained from Dr A. Winoto (Univ. California, Berkeley). The Bcl2 vector has been described in Bassik et al. (2004). These plasmids were used to transfect Phoenix cells and produce retroviruses based on the system developed by Dr G. Nolan (Stanford Univ., http://www.stanford.edu/group/nolan/retroviral_systems/phx.html). Briefly, Phoenix cells were plated and grown overnight, then transfected with retroviral vector and Nolan helper vectors. The culture media containing retroviruses was collected 24–48 h post-transfection. The collected viral-containing media was filtered with 0.45 µm low protein-binding filter and kept frozen in small aliquots for future infection. WT MEF were plated and grown overnight to 50% confluence. Then the cells were infected by adding the retrovirus-containing media mixed with fresh media at 1:1 ratio. The expression of GFP and Bcl2-IRES-GFP was visualized by GFP fluorescence 24–48 h after the infection. This Bcl-2 construct elicits efficient reduction of apoptosis in Jurkat cells (Bassik et al., 2004). A. Retroviruses expressing GFP and Bcl-2 + GFP expressed in nearly 100% of cells on the coverglasses.

B. Treatment of MEF with 50 µM C12 caused equivalent increases in caspase 3/7 activity in WT, GFP- and Bcl-2-infected MEF. WT and DKO murine fibroblasts immortalized by the NIH3T3 spontaneous immortalization method (Zong et al., 2003) were used for measurements of PI uptake (cell death) or Western blots to test for Bak/Bax expression.

**Fig. S2.** C12-induced killing of WT and DKO NIH3T3 MEF. WT and DKO murine fibroblasts immortalized by the NIH3T3 spontaneous immortalization method (Zong et al., 2003) were used for measurements of PI uptake (cell death) or Western blots to test for Bak/Bax expression.

A. WT and DKO NIH3T3 cells were incubated with 10–50 µM C12 for 24 h then processed for PI uptake. There were no differences in cell killing between the two cell lines.

B. Western blots for Bak and Bax of WT and DKO NIH3T3 cells confirmed Bak/Bax expression in WT but not DKO cells. Data are averages ± SD, n = 3 experiments in each case.