Discovery of bactericides as an acute mitochondrial membrane damage inducer

Ryan Houston, Yusuke Sekine, Mads B Larsen, Kei Murakami, Steven J. Mullett, Stacy G. Wendell, Derek P. Narendra, Bill B. Chen, and Shiori Sekine

ABSTRACT Mitochondria evolved from endosymbiotic bacteria to become essential organelles of eukaryotic cells. The unique lipid composition and structure of mitochondrial membranes are critical for the proper functioning of mitochondria. However, stress responses that help maintain the mitochondrial membrane integrity are not well understood. One reason for this lack of insight is the absence of efficient tools to specifically damage mitochondrial membranes. Here, through a compound screen, we found that two bis-biguanide compounds, chlorhexidine and alexidine, modified the activity of the inner mitochondrial membrane (IMM)-resident protease OMA1 by altering the integrity of the IMM. These compounds are well-known bactericides whose mechanism of action has centered on their damage-inducing activity on bacterial membranes. We found alexidine binds to the IMM likely through the electrostatic interaction driven by the membrane potential as well as an affinity for anionic phospholipids. Electron microscopic analysis revealed that alexidine severely perturbated the cristae structure. Notably, alexidine evoked a specific transcriptional/proteostasis signature that was not induced by other typical mitochondrial stressors, highlighting the unique property of alexidine as a novel mitochondrial membrane stressor. Our findings provide a chemical-biological tool that should enable the delineation of mitochondrial stress-signaling pathways required to maintain the mitochondrial membrane homeostasis.

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INTRODUCTION
Mitochondria are essential and multifunctional organelles of the cell that are involved in energy production, metabolic processes, and cellular signaling. Evolutionally, mitochondria evolved from α-proteobacteria that invaded into host eukaryotic cells. Mitochondria are surrounded by two membranes: the outer mitochondrial membrane (OMM) and the inner mitochondrial membrane (IMM). Both membranes have a characteristic phospholipid composition and structure. Reflecting the endosymbiotic origin of mitochondria, the IMM shares similarity with the lipid composition of bacterial membranes including high levels of cardiolipin (CL) (Horvath and Daum, 2013; Tamura et al., 2020; Tatsuta and Langer, 2017). CL is a nonbilayer forming phospholipid that destabilizes the lipid order in bilayers and induces high membrane curvature. Together with phosphatidylethanolamine (PE), which has similar biophysical properties as CL, these nonbilayer lipids make up approximately ~50% of the phospholipids in the IMM and are more abundant in this membrane than in any other cellular membranes (Horvath and Daum, 2013; Tatsuta and Langer, 2017; Tamura et al., 2020). CL and PE contribute to the characteristic highly folded structure of the IMM as represented by cristae, the concave membrane structure of the IMM (Cogliati et al., 2016). Among the biological membranes in the cells, the IMM is known to possess the highest protein density, allowing various essential bioenergetic reactions to occur. The activities of the IMM-embedded enzymes, including the OXPHOS proteins, rely on the defined lipid composition and structures of the IMM (Cogliati et al., 2016). Therefore, the integrity of the IMM must be carefully monitored and maintained in the face of internal or external insults.

Various mitochondrial stress responses that maintain healthy mitochondrial network have been discovered (Youle, 2019). These include mitophagy, an autophagic degradation of damaged mitochondria (Youle and Narendra, 2011), as well as the mitochondrial unfolded protein response (mtUPR) (Anderson and Haynes, 2020), that up-regulates a specific transcriptional program to relieve mitochondrial proteotoxic stress. Importantly, the existence of small molecules that can mimic a distinct type of mitochondrial damage has significantly contributed to the discovery and understanding of these crucial mitochondrial stress-signaling pathways. For example, the mitochondrial protonsponge, carbonyl cyanide m-chlorophenyl hydrazone (CCCP) and the combination of OXPHOS inhibitors, antimycin and oligomycin, have been widely used for the mechanistic analysis of PTEN-induced kinase 1 (PINK1)/Parkin-mediated mitophagy (Narendra et al., 2008; Matsuda et al., 2010; Lazarou et al., 2015). The treatment of the cells with these compounds results in mitochondrial membrane potential loss, one of the hallmarks of OXPHOS dysfunction, triggering mitophagy. In mammalian cells, the mtUPR can be induced by CDDO, an inhibitor of the matrix-resident protease LONP, or GTTP1, a mitochondrial HSP90 inhibitor (Munch and Harper, 2016). Moreover, mitochondrial proteotoxic stress can be induced by actinonin (Richter et al., 2015; Burman et al., 2017), an inducer of mito-ribosome stalling that results in the blockade of mitochondrial protein translation. A series of mitochondrial import blockers, so-called MitoBloCKs, can impair mitochondrial protein import pathways (Hasson et al., 2010; Dabir et al., 2013). Recent omics analyses of the mammalian cells that were treated with these mitochondrial stressors revealed that many of these stressors commonly activate the integrated stress response (ISR) (Quiros et al., 2017). The ISR induces the expression of particular cytoprotective genes through the activation of the transcription factor (ATF4), suggesting the existence of an intimate mitochondria-nuclear communication to activate the proper stress response following specific mitochondrial stressors. Recent studies using cell genetic screens for genes involved in the mitochondrial stressor-induced ISR revealed that mitochondrial proteolysis plays a critical role in activating the ISR (Fessler et al., 2020; Guo et al., 2020). These examples clearly highlight the importance of chemical compounds that can induce a specific mitochondrial stress in identifying and analyzing mitochondrial stress-signaling pathways.

Recent identification of sets of lipid synthesis enzymes and lipid transfer proteins significantly advanced our understanding of lipid metabolism within mitochondria (Tatsuta and Langer, 2017; Tamura et al., 2020). However, it is still not known how mitochondrial membrane homeostasis is preserved under conditions that disturb mitochondrial membrane integrity. This is partly due to a lack of established compounds that can specifically perturb the phospholipid environment of mitochondrial membranes.

Here, through our unbiased small-compound screen that targeted the IMM-integral protease OMA1, we found that two small compounds, chlorhexidine and alexidine, acutely disrupted the integrity of mitochondrial membranes and thereby secondarily alter OMA1 activity. Interestingly, these compounds are known as bactericides that have damage-inducing activities on bacterial membranes. Our biochemical analyses revealed that alexidine had an affinity to mitochondrial membranes and particularly damage the cristae membranes in the IMM. Moreover, we found that the alexidine treatment induced transcriptional and proteostatic signatures that were not observed with other typical mitochondrial stressors. Our discovery therefore provides a unique chemical-biological tool that can acutely and selectively perturb membrane homeostasis in the IMM.

RESULTS
Compound screen of OMA1 inhibitors identifies bactericides
In healthy mitochondria which maintain mitochondrial membrane potential, PINK1 is cleaved by the IMM-resident protease PARL just after the mitochondrial import of PINK1 (Jin et al., 2010) (Supplemental Figure S1A, left panel). The cleaved product of PINK1 is retro-translocated into the cytosol for proteasomal degradation (Yamano and Youle, 2013). In contrast, mitochondrial depolarization induces the mitochondrial import arrest of PINK1, which results in the accumulation of the full-length form of PINK1 and its kinase activation on the OMM of damaged mitochondria (Matsuda et al., 2010; Narendra et al., 2010; Okatsu et al., 2012) (Supplemental Figure S1A, middle panel). Activated PINK1 promotes the autophagic elimination of damaged mitochondria, so-called mitophagy, cooperatively working with the cytosolic E3 ligase Parkin (Youle and Narendra, 2011). Mutations in PINK1 or Parkin cause recessive early onset Parkinson’s disease (PD), suggesting a protective role of mitophagy in PD pathogenesis (Youle and Narendra, 2011). We have previously reported that several PD-related PINK1 mutants are insensitive to the mitochondrial stress-dependent import arrest and fail to accumulate in the OMM (Sekine et al., 2019, Sekine, 2020). While these PINK1 mutants are cleaved by PARL in a similar way to PINK1 wild type (WT) in healthy mitochondria, the missorted PINK1 mutants in depolarized mitochondria are instead cleaved by another IMM proteases and induces high membrane curvature. Together with phosphatidylethanolamine (PE), which has similar biophysical properties as CL, these nonbilayer lipids make up approximately ~50% of the phospholipids in the IMM and are more abundant in this membrane than in any other cellular membranes (Horvath and Daum, 2013; Tatsuta and Langer, 2017; Tamura et al., 2020). CL and PE contribute to the characteristic highly folded structure of the IMM as represented by cristae, the concave membrane structure of the IMM (Cogliati et al., 2016). Among the biological membranes in the cells, the IMM is known to possess the highest protein density, allowing various essential bioenergetic reactions to occur. The activities of the IMM-embedded enzymes, including the OXPHOS proteins, rely on the defined lipid composition and structures of the IMM (Cogliati et al., 2016). Therefore, the integrity of the IMM must be carefully monitored and maintained in the face of internal or external insults.

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FIGURE 1: Identification of bactericides as a stabilizer of PD-related PINK1 (C125G) mutant. (A) PINK1 KO HeLa cells stably expressed PINK1 (C125G)-EYFP were transfected with control or OMA1 siRNA. After 72 h, cells were treated with 20 μM CCCP or 10 μM MG132 for 4 h. The lysate was analyzed by SDS-PAGE. Note that OMA1 is known to be degraded on CCCP through autocatalytic activation (see also Supplemental Figure S1B). (B) PINK1 KO HeLa cells stably expressed PINK1 (C125G)-EYFP were seeded on 384 well plate and treated with each compound (5 μM, FDA-approved compounds). After 18 h, cells were treated with 20 μM CCCP for 4 h. EYFP fluorescence intensity of each well was measured by the high content image analyzer. (C, D) The chemical structure of a hit compound, chlorhexidine (C), and a similar compound, alexidine (D). Guanidium groups that have delocalized positive charges are highlighted in blue. (E, F) PINK1 KO HeLa cells stably expressed PINK1 (C125G)-EYFP were treated with the indicated drugs for the indicated time period, and the lysate was analyzed by SDS–PAGE. (G, H) HeLa cells stably expressed PINK1 (C125G)-EYFP were transfected with control or OMA1 siRNA. After 72 h, cells were treated with 20 μM CCCP or 5 μM alexidine for 8 h and subjected to ICC. High-magnification images in the indicated regions (G) are shown in H. TOM20 was utilized as a mitochondrial marker. Scale bars; 25 μm (G) and 5 μm (H).
and 6), indicating constitutive proteasomal degradation of PINK1 (C125G)-EYFP after PARL-mediated cleavage in healthy mitochondria. Importantly, PINK1 (C125G)-EYFP could not accumulate in response to CCCP in control siRNA-treated cells (Figure 1A, lane 2), while the full-length form of PINK1 (C125G)-EYFP did accumulate when OMA1 expression was suppressed by an OMA1-specific siRNA (Figure 1A, lane 5). This observation was confirmed by an independent OMA1 siRNA that targets different regions of OMA1 mRNA (Supplemental Figure S1B). Thus, in depolarized mitochondria, PINK1 (C125G)-EYFP is degraded through OMA1.

We realized this assay system might allow for the discovery of OMA1 inhibitors since the level of PINK1 (C125G)-EYFP under depolarized conditions was dependent on OMA1 activity. We, therefore, next measured the intensity of the YFP signal of PINK1 (C125G)-EYFP after CCCP treatment using a high-content image analyzer. The Z'-factor (a factor that can evaluate the effectiveness of a high-throughput screening) calculated by the YFP signal intensity derived from negative control siRNA versus OMA1 siRNA yielded a significantly high value (Z'-factor = 0.74) (Supplemental Figure S1C), indicating the potential robustness of our screening system (Z' < 0.0, not a suitable assay; 0.0 < Z' < 0.5, a marginal assay; 0.5 < Z' < 1.0, an excellent assay; Z' = 1.0, an ideal assay) (Zhang et al., 1999). The initial screen was performed with an FDA-approved library consisting of approximately 1100 compounds (Supplemental Table S1). The top three hit compounds that induced the accumulation of PINK1 (C125G)-EYFP were proteasome inhibitors (Figure 1B and Supplemental Table S1), confirming that the degradation of PINK1 (C125G)-EYFP occurs through proteasome. Among several other hit compounds, we focused on chlorhexidine and alexidine hydrochloride, which were independently identified within top 10 hits (Figure 1B and Supplemental Table S1). Chlorhexidine is a bis-biguanide compound (Figure 1C) that is clinically used as a bactericide, particularly in hand washing and oral care products (McDonnell and Russell, 1999; Cieplik et al., 2019). There is a similar bis-biguanide compound called alexidine (Figure 1D) (McDonnell and Russell, 1999). These compounds share structural similarities in that they contain symmetrical biguanide units tethered by a long alkyl chain. Strikingly, the treatment with chlorhexidine or alexidine, but not CCCP, significantly promoted PINK1 (C125G) accumulation in a dose- and time-dependent manner (Figure 1, E and F). While both chlorhexidine and alexidine lowered the mitochondrial membrane potential similar to CCCP (Supplemental Figure S1D), these observations suggest a membrane depolarization-independent mechanism for the PINK1 (C125G) accumulation by these bactericides. Alexidine was chosen for further study, as it showed a stronger stabilization activity for PINK1 (C125G) than did chlorhexidine. Alexidine, but not CCCP, induced the accumulation of PINK1 (C125G) on mitochondria (Figure 1, G and H). These results suggest that the identified bactericides somehow prevented OMA1-mediated PINK1 (C125G) degradation.

Alexidine demonstrates a substrate-dependent inhibition of OMA1-mediated proteolysis

OMA1 is a stress-responsive protease whose proteolytic activity is enhanced in response to mitochondrial damage including CCCP-induced mitochondrial depolarization (Ehres et al., 2009; Head et al., 2009). OMA1 activation is achieved by its self-cleavage that eventually leads to the complete degradation of OMA1 (Baker et al., 2014; Zhang et al., 2014) (Supplemental Figure S1B). Thus, the degradation of OMA1 indicates its activation. Although chlorhexidine and alexidine appeared to inhibit the OMA1-mediated degradation of PINK1 (C125G) (Figure 1, E and F, PINK1 pan-
FIGURE 2: Alexidine shows the substrate-dependent inhibition on OMA1-mediated proteolysis. (A, B) PARL KO HeLa cells were pretreated with alexidine for the indicated time period (A), or at the indicated concentration for 30 min (B), and after, treated with 20 μM CCCP for 2 h. Note that in PARL KO cells, the CCCP-dependent PGAM5 cleavage is mediated by OMA1. The lysate was analyzed by SDS–PAGE. (C) WT HeLa cells were pretreated with 5 μM alexidine for 1 h, and after, treated with 150 μM actinonin for 2 h. The lysate was analyzed by SDS–PAGE. (D) HeLa cells transiently expressed with DEL1-HA was pretreated with 5 μM alexidine for 1 h, and after, treated with 20 μM CCCP for 4 h. The lysate was analyzed by SDS–PAGE. (E) Alexidine showed the inhibitory effect on the OMA1-mediated proteolysis in a substrate-dependent manner. (F–H) The indicated HeLa cells were transfected with control or PHB2 siRNA. After 72 h, cells were harvested, and the lysate was analyzed by SDS–PAGE; 10 μM doxycycline was added for last 24 h to induce DEL1-HA in H. *Nonspecific bands. (I) PHB complex differentially regulates the OMA1-mediated proteolysis in a substrate-dependent manner.
deletion (Figure 2F, OPA1 panel, lanes 7–9). PGAM5 cleavage was also induced by PhB2 KD in WT cells (Figure 2F, PGAM5 panel, lanes 1–3). However, unlike OPA1, this was not prevented by OMA1 deletion (Figure 2F, PGAM5 panel, lanes 7–9). Rather, consistent with a recent report (Yan et al., 2020), PARL deletion prevented the PhB2 KD-induced PGAM5 cleavage (Figure 2F, PGAM5 panel, lanes 4–6). We found PhB2 deletion also showed different effects on other OMA1 substrates. PhB2 KD had only slight or no obvious effects on the degradation of CHCHD2 and CHCHD10 (Figure 2G), while it clearly induced DELE1 cleavage (Figure 2H). These results suggest that PhB2 deletion induces the OMA1-mediated proteolysis in a substrate-dependent manner (Figure 2I), which was apparently similar to the effects of alexidine (Figure 2E).

**Alexidine has an affinity for the IMM**

We next tried to identify a target of alexidine to further address the underlying molecular mechanism of the observed substrate-specific action of alexidine on OMA1-mediated proteolysis. In addition to antimicrobial properties, chlorhexidine and alexidine are also reported as inhibitors of PTPMT1 (Doughty-Shenton et al., 2010), a mitochondrial matrix-localized phosphatase that dephosphorylates phosphatidylglycerol-phosphate, an essential intermediate in CL biosynthesis (Xiao et al., 2011; Zhang et al., 2011). However, KD of PTPMT1 in our PINK1 (C125G)-EYFP stable HeLa cells did not promote PINK1 (C125G) stabilization (Supplemental Figure S2A, lane 5), suggesting that alexidine appears to have a different target in this context.

The proposed mechanism of action of chlorhexidine as a bactericidal agent is based on its interaction with bacterial membrane damage-inducing ability through its interaction with phospholipids (McDonnell and Russell, 1999; Cieplik et al., 2019). Alexidine also has a similar activity on the bacterial membranes (McDonnell and Russell, 1999). Guanidinium groups of these compounds possess delocalized positive charges at physiological pH (Langmaier et al., 2016) (Figure 1, C and D). The delocalized positive charges have higher lipophilicity compared with groups that have localized charges, which is considered to confer the efficient binding ability of chlorhexidine and alexidine to phospholipids, together with their long alkyl chain between two symmetric guanidinium groups (McDonnell and Russell, 1999; Cieplik et al., 2019). These observations led us to examine the effect of alexidine on the phospholipids in the IMM. We first tested a phospholipid dye 10-N-Nonyl acridine orange (NAO) staining with or without alexidine treatment. NAO is a lipophilic and positively charged molecule that is often utilized to monitor anionic phospholipids in bacterial membranes (Lin and Weibel, 2016). In eukaryotic cells, NAO selectively accumulates in the IMM (Wol et al., 2019; Kondadi et al., 2020). Although NAO is originally developed as a CL dye, PTPMT1 deficiency did not alter the mitochondrial NAO staining in HeLa cells (Supplemental Figure S2B), suggesting that NAO has a broader affinity for mitochondrial phospholipid species. We found that treatment with alexidine but not CCCP dramatically reduced mitochondrial NAO staining (Figure 3A). NAO staining was rapidly lost after alexidine treatment (Figure 3B). In contrast, the fluorescent intensity of Su9-mCherry (matrix marker) was not affected, indicating that mitochondria themselves were still present (Figure 3B). These results indicate that alexidine has effects on the IMM phospholipids. To directly evaluate the binding affinity of alexidine to phospholipids, we examined the effect of alexidine on in vitro binding between NAO and anionic phospholipid species coated on microplate wells (Nomura et al., 2000; Rodriguez et al., 2008). Preincubation with alexidine was found to reduce the fluorescent intensity derived from NAO bound to anionic phospholipids in a dose-dependent manner (Figure 3C). Consistent with the NAO staining in cells (Figure 3A), CCCP did not reduce the NAO intensity in this assay (Figure 3C). A lipidomic analysis of mitochondria isolated from mock- or alexidine-treated cells did not show any decrease in mitochondrial phospholipid species (Supplemental Figure S3), excluding the possibility that alexidine decreased the amount of mitochondrial membrane lipids. Altogether, these results suggest that alexidine has an affinity to anionic phospholipids and competes with NAO to bind to these IMM phospholipids.

Intriguingly, the aforementioned chemical property of guanidinium groups is also often utilized to target drugs to mitochondria, because it is known to preferentially accumulate in the IMM which has an electrochemical gradient (Sibrian-Vazquez et al., 2008; Battagotch et al., 2018). Therefore, we tested whether mitochondrial depolarization prevents the action of alexidine on the IMM. CCCP pretreatment partially attenuated the alexidine-induced reduction of NAO staining (Figure 3, D and E). These results suggest that the mitochondrial membrane potential can be a primary driving force for the mitochondrial targeting of alexidine, and the high lipophilicity of alexidine promotes the accumulation of this compound in the hydrophobic lipid environment of the IMM.

**Alexidine induces an acute perturbation of IMM integrity**

Since alexidine appeared to interact with the IMM phospholipids, we investigated the effects of alexidine on IMM structure and on IMM-shaping proteins. The IMM is structurally subdivided into two domains: the inner boundary membrane (IBM), where the IMM is in close proximity with the OMM, and the cristae, baglike structures where the IMM invaginates into the matrix (Cogliati et al., 2016; Rampelt et al., 2017). These two IMM domains are connected by narrow, necklike structures called cristae junctions (CJs) (Cogliati et al., 2016; Rampelt et al., 2017) (Figure 4A). To examine IMM structure, we performed an electron microscopic (EM) analysis of mitochondria with or without alexidine treatment. The EM images clearly revealed that the IMM structure was severely disrupted after alexidine treatment (Figure 4A, right panels). The alteration of the cristae membrane was the most striking feature. The cristae membrane was

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**FIGURE 3:** Alexidine has an affinity for the IMM. (A and B) WT HeLa cells or HeLa cells stably expressed Su9-mCherry (a matrix marker) were treated with 5 μM alexidine or 20 μM CCCP for the indicated time period. After the drug-treatment, cells were washed with PBS for twice and stained with NAO. NAO staining was analyzed by live-cell imaging. Scale bars; 25 μm (A) and 10 μm (B). (C) NAO fluorescence intensity (Ex 485 nm /Em 535 nm) in individual wells of a microtiter plate which were coated with or without the indicated phospholipid species was measured by microtiter plate reader. Alexidine or CCCP was added at the indicated concentration for 30 min before the NAO staining. Data are shown as mean ± SD (n = 3 or n = 6 per condition). **p < 0.01, ***p < 0.001, and ****p < 0.0001 (one-way ANOVA followed by Turkey’s multiple comparison). (D) WT HeLa cells were treated with 5 μM alexidine, 20 μM CCCP, or the combination of these drugs for the indicated time period. After the drug-treatment, cells were washed with PBS twice and stained with NAO. NAO staining was analyzed by live-cell imaging. Scale bars; 50 μm. (E) FACS analysis of NAO fluorescence intensity in D.
**A**

Mock, CCCP, and Alexidine treatments are shown in various images. The images depict cellular structures with a focus on the mitochondrial outer membrane (OMM) and inner mitochondrial membrane (IMM).

**B**

Panel B displays images of OMA1 (E328Q)-EYFP stable OMA1 KO with mock, Alexidine (4 hour), and CCCP (4 hour) treatments. The images show the localization of OMA1 in comparison to PHB2 and COX IV proteins.

**C**

Panel C is similar to panel B but focuses on OMA1 (E328Q)-EYFP stable OMA1 KO with Alexidine (4 hour) and CCCP (4 hour) treatments.

**D**

Panels D show images of MIC60 and TIM50 proteins under mock, Alexidine (2 hour), and CCCP (2 hour) conditions. The images are merged to highlight the colocalization of these proteins.

**E**

A graph showing Pearson's R value for different treatments including Mock, Alexidine, and CCCP is presented, with statistical significance indicated by asterisks.
pinched off from the IBM and often appeared bunched up in an
onionlike ball in the matrix (Figure 4A, right panels, and Supplemental
Figure S4). Mitochondria were swollen and the matrix content
seemed to be diluted. However, the OMM and the IBM did not ex-
hibit apparent morphological changes and remained in place, sug-
gesting that alexidine may particularly influence the cristae mem-
branes and CJs. In contrast, CCCP treatment only displayed a mild
disturbance in the IMM structure (Figure 4A, middle panels, and
Supplemental Figure S4).

We next examined the effects of alexidine on proteins that help
shape the IMM. Because we observed substrate-specific effects of
alexidine on OMA1-mediated proteolysis (Figure 2E), and PHB2 de-
dletion showed similar effects (Figure 2I), we examined the localization
of OMA1 and PHB2. To prevent the stress-dependent autocatalytic
degradation of OMA1, we stably expressed an OMA1 protease activity-
dead mutant, OMA1(E328Q) (Sekine et al., 2019), in OMA1 KO
cells. The localization of OMA1 and PHB2 mostly overlapped under
steady-state and CCCP-treated conditions (Figure 4B, upper and
lower panels). However, under alexidine-treated conditions, the local-
ization of these proteins diverged, with part of the OMA1 pool now
segregated from the PHB2-positive IMM (Figure 4B, middle panels).

When the IMM was stained with Cox IV, a subunit of the cytochrome
oxidase complex, the localization of OMA1 overlapped with the
Cox IV-positive IMM even under alexidine-treated conditions (Figure
4C, middle panels), indicating that OMA1 still exists in the IMM.

The MICOS complex is located at CJs where it stabilizes
membrane curvature and forms contact sites (CSs) between the
OMM and the IMM (Rampelt et al., 2017) (Figure 4A). It is
reported that the MICOS complex genetically interacts with the CL
synthesis pathway (Hopkins et al., 2011; Friedman et al., 2015),
and that some components (Mic60 and Mic27) of the MICOS
complex directly bind to CL in vitro (Weber et al., 2013; Michaud
et al., 2016). Among the seven components of the metazoan MI-
COS complex (Rampelt et al., 2017), we examined MIC60 local-
ization before and after the alexidine treatment. Under the resolu-
tion of conventional confocal microscopy, MIC60 shows a
uniform distribution along the mitochondrial string at steady-
state conditions (Figure 4D, upper panels, and Supplemental
Figure S5A). However, after alexidine treatment, MIC60 was lo-
calized in a restricted region of each fragmented mitochondrion
and showed an intense, punctalike localization within the IMM
(Figure 4, D and E, and Supplemental Figure S5, A–C). In contrast
to the alexidine-treated cells, MIC60 was uniformly distributed in
each fragmented mitochondrion after CCCP treatment (Figure 4,
D and E, and Supplemental Figure S5, A–C), indicating that the
intense MIC60 puncta formation is specifically induced by alexi-
dine. Despite the distribution changes of PHB2 and MIC60, their
high weight molecular complex formation was not affected by
alexidine treatment (Supplemental Figure S5, D and E). Taken

Alexidine evokes a unique transcriptional/proteostasis
signature
From the observations above, we hypothesized that alexidine could
be used as an acute mitochondrial membrane damage inducer.
Therefore, we decided to characterize the cellular response elicited
by the alexidine-induced mitochondrial membrane perturbation.
For this purpose, we performed TMT-based quantitative proteomics
(Figure 5A, and Supplemental Table S2) and RNA-sequencing analy-
sis (Figure 5C, and Supplemental Table S3). As described so far,
alexidine induced mitochondrial alterations that were distinct from
CCCP treatment. To identify the proteins or mRNAs whose expres-
sion was specifically changed in response to the alexidine-induced
mitochondrial membrane stress, we compared three different con-
ditions: mock, alexidine, and CCCP treatment. Gene Ontology
analysis for proteins that were significantly changed in the alexidine-
treated cells showed a significant enrichment of mitochondria-re-
lated proteins (Figure 5B), suggesting that alexidine preferentially
affects mitochondria among several other organelles. Many proteins
or mRNAs were commonly up-regulated or down-regulated in both
the alexidine- and CCCP-treated cells (for example, ISR-targeted
genes were up-regulated in both alexidine- and CCCP-treated cells
in RNA-seq) (Figure 5, A and C), which may be attributed to the
observation that alexidine also induces mitochondrial depolariza-
tion at almost the same level as CCCP (Supplemental Figure S1D).
Notably, the expression of some proteins was specifically altered in
the alexidine-treated cells. Twenty-seven proteins were specifically
identified as down-regulated proteins in alexidine-treated cells (fold
change < 0.8, t test q value < 0.05), (Figure 5D and Supplemental
Table S2). Among these, 13 proteins were mitochondrial proteins
(Figure 5E). These include OXPHOS proteins, proteins which are in-
volved in Coenzyme Q biosynthesis, and PTPMT1. As up-regulated
proteins (fold change > 1.5, t test q value < 0.05), only four nonmito-
chondrial proteins were specifically up-regulated in response to
alexidine (Figure 5F and Supplemental Table S2). These include
metallothioneins (MTs) and heme oxygenase 1 (HMOX1), an essen-
tial enzyme in heme catabolism (Igarashi and Sun, 2006) (Figure 5G).
We next sought to confirm the alexidine-specific expression changes
obtained from the proteomics analysis. IB analysis confirmed that
HMOX1 was up-regulated while COA7, COX17, and PTPMT1 were
down-regulated on alexidine treatment (Figure 5H). Strikingly, these
changes were only observed by alexidine treatment and not by
other well-known mitochondrial stressors such as CCCP, rotenone
(a Complex I inhibitor), actinonin, or CDDO (Figure 5H). We con-
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Supplemental Figure S6C), indicating the existence of a mitochondrial-nuclear communication in response to alexidine treatment. In contrast, the mRNA levels of the alexidine-specific down-regulated mitochondrial proteins including COA7 and COX17 did not significantly change after alexidine treatment, indicating that alexidine likely induces the posttranscriptional degradation of these proteins (Figure 5E and Supplemental Figure S6C). LONP KD prevented the alexidine-induced down-regulation of PTPMT1 (Supplemental Figure S6D), suggesting that at least PTPMT1 is degraded within the mitochondria. Consistent with the observation that PTPMT1 deficiency affected neither PINK1 (C125G) accumulation nor the NAO staining unlike alexidine (Supplemental Figure S2), it did not show the induction of HMox1 or the degradation of COA7 (Supplemental Figure S6E).

Collectively, our results suggest that alexidine can evoke unique mitochondrial responses that are not induced by other typical mitochondrial stressors, presumably through the perturbation of the IMM integrity (Figure 5I).

DISCUSSION

In this study, we found two bactericides, chlorhexidine and alexidine, as small molecules that can induce the acute perturbation of mitochondrial membrane integrity. EM analysis of mitochondria after alexidine treatment showed a strikingly altered cristae membrane structure, while keeping the OMM and the IBM largely in place (Figure 4A). This suggests that alexidine can specifically damage the cristae among several distinct membrane compartments within mitochondria (Figure 5I). It has been noted that cristae membranes are the membranes where OXPHOS proteins are concentrated (Cigliati et al., 2016). Our TMT-based quantitative proteomics analysis indicated that many OXPHOS proteins were down-regulated after alexidine treatment (Figure 5E), which also supports the specific action of alexidine on the cristae membrane. Interestingly, several recent studies identified alexidine (and chlorhexidine) as an agent that can alter cellular metabolism. This metabolic shift ultimately resulted in various effects on cells: the anti-invasive and metastatic activity on tumor cells (Kenny et al., 2015; Commander et al., 2020), the maintenance of the quiescent status of stem cells (Liu et al., 2015), the enhanced glucose utilization in vivo (Nath et al., 2015), or the transcription factor TFEB nuclear translocation through AMPK activation (Wang et al., 2017). Some of these reports found that the alexidine treatment reduced the oxygen consumption (Liu et al., 2015) and preferentially shifted the energy source from OXPHOS to glycolysis (Commander et al., 2020). Because PTPMT1 was reported as a metazoan target of alexidine (Doughty-Shenton et al., 2010), some studies indicated above speculated that the observed metabolic effects may result from PTPMT1 inactivation. However, the cristae membrane-disrupting activity of alexidine, which we identified in this study, must now be considered as a basis for the acute effects of alexidine on the cellular metabolism.

It still remains elusive how alexidine specifically affects the cristae membrane. As predicted from the chemical properties of guanidine groups of alexidine (Sibrian-Vazquez et al., 2008; Bhattogtokh et al., 2018), our NAO staining assay indicated that mitochondrial membrane potential can be a driving force for the mitochondrial targeting of alexidine (Figure 3D and E). Because of the high abundance of OXPHOS proteins in the cristae (Cigliati et al., 2016), the cristae membranes have higher membrane potential than the IBM (Wolf et al., 2019). This unique feature of the cristae may explain the specific effect of alexidine on this membrane compartment within the IMM. Also, it has been suggested that the high curvature of the cristae is created by high amounts of nonbilayer lipids such as CL and PE (Horvath and Daum, 2013; Cigliati et al., 2016). We demonstrated that alexidine has a reasonable affinity for CL as it is able to compete with NAO (Figure 3C). Therefore, this property of alexidine may also contribute to the accumulation of alexidine in the cristae membrane.

As a result of alexidine treatment, we observed a robust induction of HMox1, a heme-degrading enzyme, and several MTs, metal chelators (Figure 5, F–H). The direct link between the alexidine-mediated mitochondrial membrane damage and the induction of HMox1 and MTs is not known. Early studies indicate that HMox1 and MTs are simultaneously induced by heme addition to the culture media (Smith, 2000). Subsequent studies have demonstrated that HMox1 induction was mediated by nuclear factor erythroid 2-related factor 2, a transcription factor involved in the antioxidant response (Igarashi and Sun, 2006). Mitochondria are known as a site for heme biosynthesis (Xu et al., 2013). Also, the mitochondrial matrix has a pool of heavy metal copper (Xu et al., 2013). Together with Fe-S clusters that are synthesized in mitochondria, heme and copper are utilized as important cofactors for various enzymes including OXPHOS proteins. Due to their harmful radical-formation activity, the export of newly synthesized heme across the mitochondrial membranes is tightly regulated by a membrane-embedded heme exporter, while copper chaperones ensure the safe delivery of...
copper to target proteins (Xu et al., 2013). It is possible that the alexidine-mediated IMM perturbation disturbed this regulation and resulted in the heme and copper leakage from mitochondria. Ultimately, it might lead to the induction of HMOX1 and MTs as a preventive strategy. Complex IV utilizes a heme-copper center to reduce oxygen (Xu et al., 2013). We observed that the alexidine treatment strongly degraded two Complex IV assembly factors, COA7 and COX17 (Xu et al., 2013; Kozjak-Pavlovic et al., 2014; Mohanraj et al., 2019) (Figure S5H). The exact role of COA7 in the assembly of Complex IV was not known, but very recently, its heme-binding/chaperoning activity was proposed (Formosa et al., 2021). COX17 is well known as a copper chaperone that delivers copper to Complex IV (Xu et al., 2013). Therefore, in addition to the direct leakage of heme and copper from the mitochondria, it is also possible that heme and copper released from degraded OXPHOS proteins activate the transcription of HMOX1 and MTs. In either case, the induction of HMOX1 and MTs can be used as a sensitive marker of the mitochondrial membrane damage.

In addition to these transcriptional/proteostasis alterations, alexidine remodeled the IMM-resident membrane proteins including PHB2, OMA1, and MIC60 (Figure 4, B and D). The single particle EM analysis of PHB complex suggested that it forms a ringlike structure in the IMM (Tatsuta et al., 2005). It is predicted that the ringlike PHB complex can exert a partitionlike function in the IMM, where it can define the lateral distribution of specific lipids, including CL and PE, or proteins such as IMM-resident proteases including OMA1 (Osman et al., 2009b). As alexidine showed an affinity to CL (Figure 3C), alexidine might be able to accumulate in the PHB complex-organized CL/PE-enriched domain of the IMM. Previous reports revealed that PHB deletion can activate OMA1 proteolytic activity and induce subsequent OPA1 cleavage without mitochondrial membrane depolarization (Merkwirth et al., 2008; Merkwirth et al., 2012; Korwitz et al., 2016), indicating that PHB complex may hold OMA1 in the inactive state presumably through restricting the protease to specific IMM microdomains. We observed the segregation of OMA1 from PHB2-positive IMM after alexidine treatment (Figure 4B). These observations may indicate that alexidine causes OMA1 to dissociate from the PHB complex-organized microdomain and that once OMA1 is released from PHB complex-mediated inhibition, it is proteolytically active. However, OMA1 did not cleave or degrade certain substrates such as PINK1 (C125G), PGAMS, CHCHD2, and CHCHD10 on alexidine treatment (Figure 2E). PHB2 deletion provided similar results (Figure 2I). These observations may imply the existence of an additional layer of regulation that allows OMA1 to access some substrates.

MIC60, one of the important components of the MICOS complex (Rampelt et al., 2017), showed a punctalike localization within the IMM after alexidine treatment (Figure 4, D and E, and Supplemental Figure S5, A–Q). Among several components of the MICOS complex, it has been suggested that Mic60 can self-assemble and form puncta within the IMM when all other MICOS components are absent in yeast (Friedman et al., 2015). Moreover, the recent analysis showed the de novo formation of CJs by drug-controlled expression of MIC60 in reconstituted MIC60 KO cells (Stephan et al., 2020), establishing a critical role for MIC60 in CJ formation. Upstream determinants of MIC60 localization are not fully understood. However, recent studies in yeast have revealed that the aforementioned Mic60 puncta formed in the absence of other components of the MICOS complex are often observed in proximity to ER-mitochondria CJs, where the ERMES complex exists (Tirrell et al., 2020). The ERMES complex physically tethers the ER and mitochondria in yeast and creates membrane CJs to allow efficient lipid transfer between the ER and the mitochondria (Tamura et al., 2020). Together with the fact that MICOS and ERMES genetically interact with each other (Hoppins et al., 2011), cooperative functions of MICOS and ERMES have been suggested in mitochondrial membrane architecture (ERMIONE) (van der Laan et al., 2012). Mitochondrial lipid homeostasis depends on both interorganelle (mainly from the ER) and intraorganelle (between the OMM and the IMM) lipid trafficking (Tatsuta and Langer, 2017; Tamura et al., 2020). Therefore, the coordinated regulation of ERMES-MICOS localization may be functionally linked to allow for efficient lipid trafficking across the membranes. Although it has not been examined whether the MICOS complex is involved in interorganelle lipid trafficking, it is known that the MICOS complex is involved in intramitochondrial lipid metabolism (Aaltonen et al., 2016; Michaud et al., 2016). Our EM analysis revealed that the cristae structure was severely damaged following alexidine treatment (Figure 4A and Supplemental Figure S4). In such a case, it is expected that mitochondrial lipid demand is significantly increased in order to help restore the highly folded cristae structure. It is tempting to speculate that the alexidine-induced MIC60 puncta formation plays a role in this process as part of a mitochondrial membrane stress response. However, we cannot deny the possibility that the distribution change of MIC60 is a consequence of the cristae disruption. It may be also of note that several reports indicate a link between the MICOS complex and the COA7 or COX17 (robustly degraded proteins after the alexidine treatment). For example, in yeast, it was shown that Cox17 physically interacts with Mic60 and modulates the MICOS complex formation (Chojnacka et al., 2015). Other studies suggest that the sustained KD of MIC60 (or its interactor, SAM50) resulted in the degradation of COA7 in mammalian cells (Kozjak-Pavlovic et al., 2014). COA7 was also identified as the possible interactor of MIC10, another important component of MiCOS complex (Alkhaja et al., 2012). These previous observations may explain the reason why we observed the relatively specific and robust degradation of COA7 and COX17 among over 100 components of OXPHOS system.

In conclusion, we discovered alexidine and chlorhexidine as small molecules that enable us to acutely and preferentially perturb the mitochondrial membrane architecture in the IMM (Figure 5I). Our findings therefore offer a useful chemical-biological tool for delineating mitochondrial membrane stress responses.

**MATERIALS AND METHODS**

**Cell culture, transfection, and treatments**

HEK293T and HeLa cells were cultured in DMEM (Life Technologies) supplemented with 10% fetal bovine serum (WWR Life Science), 10 mM HEPES (Life Technologies), 1 mM sodium pyruvate (Life Technologies), nonessential amino acids (Life Technologies), and GlutaMAX (Life Technologies). For RNA interference, 20 nM Stealth siRNAs (Stealth RNAi Negative Control Med GC Duplex #2, 12935112; LONP, HSS113887) (Thermo Fisher Scientific) or 5 nM Silencer select siRNAs (Silencer Select Negative Control #1 siRNA, 4390843; OMA1, s41776; LONP, s17903; PTPM1, s229947) (Thermo Fisher Scientific) were transfected using Lipofectamine RNAi max transfection reagent (Thermo Fisher Scientific) at the same time as cell seeding. For drug treatment experiments, cells were incubated in medium containing one or more of the following compounds: CCCP (Cayman Chemical), MG132 (Sigma), chlorhexidine (Cayman Chemical), alexidine (Cayman Chemical), rotenone (Cayman Chemical), actinonin (Cayman Chemical), CDDO (Cayman Chemical). For examination of mitochondrial membrane potential or mitochondrial ROS production, 20 nM TMRM (Thermo Fisher Scientific) or 50 nM MitoSOX (Thermo Fisher Scientific), respectively, was directly added...
to cell culture media and incubated for 15 min. For NAO staining, cells were washed twice with phosphate-buffered saline (PBS) and incubated with 100 nM NAO (Thermo Fisher Scientific) for 15 min. Cells were washed and replaced with normal medium followed by live-cell imaging using a 63x/1.4 NA oil immersion objective on a Leica SP8 LIGHTNING confocal microscope (Leica) or FACS analysis using Attune NxT Acoustic Focusing Cytometer (Thermo Fisher Scientific).

**Plasmids**

Site-directed mutagenesis of pLenti-CMV-Neo-PINK1 (C125G)-EYFP or pLVX-puro-OMA1 (E328Q)-EYFP was performed by PCR amplification (CloneAmp HiFi PCR Premix, Takara or Q5 High-Fidelity DNA Polymerase system, NEB) of PINK1 or OMA1 encoding plasmid using appropriate primers followed by Gibson assembly (In-Fusion HD Cloning system, Clontech) into the Sall-Xhol of the pLenti-CMV-Neo vector, or into the EcoRI site of pLVX-puro vector (Clontech). pLVX-puro-DELE1-HA and pLVX-puro-Su9-mCherry was created by PCR amplification and subcloning into the EcoRI site of pLVX-puro vector. pRetroX-Tight-puro-DELE1-HA was created by PCR amplification and subcloning into the BamHI/NotI site of pRetroX-Tight-puro vector (Clontech). All constructs were confirmed by DNA sequencing.

**Generation of stable cell lines**

To generate stably transfected cell lines, lentiviruses (for plasmids within pLenti-CMV-neo or pLVX-puro vectors) and retroviruses (for plasmids within pRetroX-Tight-puro vector) were packaged in HEK293T cells. HeLa cells were transduced with viruses with 10 μg/ml polybrene (Sigma) then optimized for protein expression via an antibiotics selection. PINK1 KO HeLa cells stably expressed PINK1 (WT)-EYFP or PINK1 (C125G)-EYFP were monocloned, and clone #21 or clone #23, respectively, was used in this study. For generating the Tet-on DELE1-HA stable cell line, Retro-X-Tet-on Inducible Expression System (Clontech) was used according to the manufacturer's instruction.

**Generation of KO cell lines**

PTPMT1 KO HeLa cell lines were generated using lentCRISPRv2 system (Sanjana et al., 2014; Shalem et al., 2014). Briefly, after the infection of lentiviruses that express hSpCas9 and PTPMT1 sgRNA (5′-TGGCGGCTCAAGCTCCGCAA-3′), infected cells were selected via the treatment with 500 μg/ml hygromycin (Sigma) for 24 h. The selected cells were cultured for more than 2 wk at least to ensure the efficient gene deletion and used in experiments. The following HeLa KO cell lines were kindly provided by Richard J. Youle (NIH, NINDS): PINK1 KO (Nezich et al., 2015), OMA1 KO (Sekeine et al., 2019), and PARL KO (Sekeine et al., 2019).

**Immunoblotting (IB)**

For SDS–PAGE, cells were lysed with 1× NuPAGE LDS sample buffer (Thermo Fisher Scientific) supplemented with 100 mM dithiothreitol (DTT) (Sigma) and boiled with shaking for 15 min. Approximately, 50 μg of protein per sample was separated on 7.5, 10, or 4–20% gradient Mini-PROTEAN TGX Precast Gel (Bio-Rad) or Criterion TGX Gels (Bio-Rad) and then transferred to a nitrocellulose membrane (Bio-Rad) or PVDF membrane (Bio-Rad). The membrane was blocked with Odyssey Blocking Buffer (LI-COR) and incubated with the indicated primary antibodies at 4°C overnight. After washing with PBS-T (PBS + 0.05% Tween-20), the membrane was incubated with HRP-conjugated secondary antibodies (Thermo Fisher Scientific) and washed again with PBS-T. Detection was performed with iBright CL1000 Imaging System (Thermo Fisher Scientific). For Phos-tag SDS–PAGE, cells were lysed with 1% Triton buffer [1% Triton X-100, 150 mM NaCl, 50 mM Tris-HCl, pH 7.4, 1 mM EDTA, phosphatase inhibitors (PhosSTOP, Sigma), and protease inhibitors (Complete, Sigma)]. After centrifugation, the lysate that contains 10 μg of protein per sample was mixed with 2× Laemmli sample buffer (Bio-Rad) supplemented with 2 M 2-mercaptoethanol (Bio-Rad) and boiled for 3 min. Samples were separated on 7.5% Mini-Gel (TGX FastCast Acrylamide Solutions, Bio-Rad) containing 50 μM Phos-tag AAL-107 (Wako) and 10 mM MnCl2 (Sigma) according to the manufacturer’s instructions. For the elimination of the manganese ion from the gel, the gel was soaked with a transfer buffer containing 1 mM EDTA for 10 min, washed with a transfer buffer without EDTA for 10 min, and then transferred to a PVDF membrane (Bio-Rad).

**Immunocytochemistry (ICC)**

Cells were seeded into Lab-Tek Chambered Coverglass with 4 wells (Thermo Fisher Scientific). Cells were rinsed in PBS and fixed with PBS containing 4% formaldehyde for 15 min at room temperature. Cells were permeabilized with PBS containing 0.1% Triton X-100 for 10 min at room temperature. Blocking was performed using PBS containing 2% bovine serum albumin (BSA) for 30–60 min at room temperature. For immunostaining, cells were incubated with primary or secondary antibodies (Alexa Fluor, Thermo Fisher Scientific) diluted in PBS containing 2% BSA overnight at 4°C or about 2 h at room temperature, respectively. Cells were imaged using a 63x/1.4 NA oil immersion objective on Leica SP8 LIGHTNING confocal microscope (Leica).

**Antibodies**

The following antibodies were used in IB or ICC: PINK1 (Cell Signaling, 6946S), OMA1 (Santa Cruz, sc-515788), PGAMS (Thermo Fisher Scientific, PA5-57894), CHCHD2 (Proteitech, 66302-1-lg), and CHCHD10 (Sigma, HPA003440), OPA1 (BD Biosciences, 612607), HSP90 α/β (Santa Cruz, sc-7947 or sc-13119), HSP60 (Santa Cruz, sc-13115), Tim50 (Santa Cruz, sc-39678), Tom20 (Santa Cruz, sc-17764), PHB2 (Proteitech, 66424-1-lg) for ICC, PHB2 (Proteitech, 12295-1-1g) for IB, Cox IV (Thermo Fisher Scientific, PA5-19471), Mic60 (Proteitech, 10179-1-AP), HMOX1 (GeneTex, GTX101147), PTPMT1 (Santa Cruz, sc-390901), COX17 (Proteitech, 11464-1-AP), COA7 (Proteitech, 25361-1-AP), LONP (Novus Biologicals, NB181734), GFP (Novus Biologicals, NB600-597 or NB-600-308), and HA (Cell Signaling, 3724).

**FDA-approved compound library screening**

The FDA-approved compound library (Selleck, 100 nl per drug) was stamped to black 384-well plates with glass bottoms using CyBio Well vario (Analytik Jena). PINK1 KO HeLa cells stably expressed PINK1 (C125G)-EYFP (clone #23) were then added to give density of 4000 cells per well and a final drug concentration of 5 μM. After 18 h of treatment, CCCP was added to a final concentration of 20 μM and incubated for 4 h followed by fixation in 4% paraformaldehyde and counterstaining with Hoechst 33342. Fluorescence was detected using an ImageXpress Micro XLS (Molecular devices) high-content imager and the cellular EYFP fluorescence signal was calculated using CellProfiler software (McQuin et al., 2018).

**In vitro NAO assay**

Binding of NAO to each anionic phospholipid was studied in lipid monolayers (Nomura et al., 2000; Rodriguez et al., 2008), with slight modifications; 96-well microtiter plates (Corning, 3915) were coated with 50 μl of 20 μM each anionic phospholipid
in ethanol and evaporated at 37°C for 5 h. An increasing concentration of allexidine (final; 0–100 μM) or CCCP (final; 10 μM) in 50 μl PBS containing 2% BSA was added and incubated at 37°C for 30 min. Then, NAO (final; 100 nM) in 50 μl PBS containing 2% BSA was added and incubated at 37°C for 30 min protected from light. After the incubation, each well of the plates was washed with 150 μl PBS for five times. Finally, NAO fluorescence intensity was measured by SpectraMax i3x Multi-Mode Detection Platform (Molecular Devices) using Ex 485 nm/Em 535 nm. The following anionic phospholipids purchased from Avanti Polar Lipids were used: Heart CA (840012P), Egg PA (840101P), Egg PG (841138P), Liver PI (840042P), and Brain PS (840032P).

Transmission EM
HeLa cells were fixed with 4% glutaraldehyde (Electron Microscopy Services) in EM buffer (0.1 N sodium cacodylate at pH 7.4 with 2 mM calcium chloride) for 30 min at room temperature and then at 4°C for at least 24 h. Samples were washed with buffer and treated with 1% osmium tetroxide in 0.1 N cacodylate buffer at pH 7.4 for 1 h on ice, washed, and en bloc stained with 0.25–1% uranyl acetate in 0.1 N acetate buffer at pH 5.0 overnight at 4°C, dehydrated with a series of graded ethanol, and finally embedded in epoxy resins. Ultrathin sections (70 nm) were stained with lead citrate and imaged with a JEOL 1200 EXII Transmission electron microscope.

RNA isolation and RT PCR
Total RNAs were isolated using RNeasy Mini Kit (Qiagen) and reverse-transcribed to cDNA using iScript cDNA Synthesis Kit (Bio-Rad) according to the manufacturer's instructions. RT PCR was performed using SYBR Green Master Mix (Bio-Rad) and QuantStudio 3 RT-PCR system (Thermo Fisher Scientific). All expression levels were normalized to that of RPS18 mRNA. The following RT-PCR primers were used: Heart CA (840012P), Egg PA (840101P), Egg PG (841138P), Liver PI (840042P), and Brain PS (840032P).

RNA-seq
RNA-seq was performed by Illumina TruSeq stranded mRNA kit was used to prepare libraries according to the manufacturer's instructions. Resulting libraries were sequenced on NovaSeq 6000 using paired-end 100 base pair sequencing at the depth of 40-M reads per sample.

Transmission EM
HeLa cells were fixed with 4% glutaraldehyde (Electron Microscopy Services) in EM buffer (0.1 N sodium cacodylate at pH 7.4 with 2 mM calcium chloride) for 30 min at room temperature and then at 4°C for at least 24 h. Samples were washed with buffer and treated with 1% osmium tetroxide in 0.1 N cacodylate buffer at pH 7.4 for 1 h on ice, washed, and en bloc stained with 0.25–1% uranyl acetate in 0.1 N acetate buffer at pH 5.0 overnight at 4°C, dehydrated with a series of graded ethanol, and finally embedded in epoxy resins. Ultrathin sections (70 nm) were stained with lead citrate and imaged with a JEOL 1200 EXII Transmission electron microscope.

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RNA-seq
RNA-seq was performed by Illumina TruSeq stranded mRNA kit was used to prepare libraries according to the manufacturer's instructions. Resulting libraries were sequenced on NovaSeq 6000 using paired-end 100 base pair sequencing at the depth of 40-M reads per sample.

TMT-based quantitative proteomics
TMT-based quantitative proteomics was performed by MS Bioworks.

Sample preparation. Cells were lysed in 8 M urea, 50 mM Tris-HCl (pH 8.0), 1x complete protease inhibitor (Roche), and 1x PhosStop (Roche) with a sonic probe (3 x 30 s at 80% amplitude) with subsequent mixing at room temperature for 1 h at 1000 rpm on a Thermomixer. Lysates were quantified by Qubit fluorometry (Life Technologies); 50 μg of each sample was digested overnight with trypsin. Briefly, samples were reduced for 1 h at room temperature in 12 mM DTT followed by alkylation for 1 h at room temperature in 15 mM iodoacetamide. Trypsin was added to an enzyme:substrate ratio of 1:20. Each sample was acidified in formic acid and subjected to SPE on an Empore SD C18 plate. Each sample was lyophilized and reconstituted in 140 mM HEPES (pH 8.0), 30% acetonitrile for TMT labeling. Peptides were labeled using TMT 10-plex (Thermo Fisher Scientific) according to the manufacturer's instructions. Briefly, 40 μl of acetonitrile was added to each TMT tag tube and mixed aggressively. Tags were incubated at room temperature for 15 min; 30 μl of label was added to each peptide sample and mixed aggressively. Samples were incubated in an Eppendorf Thermomixer at 300 rpm at 25°C for 1.5 h. Reactions were terminated with the addition of 8 μl of fresh 5% hydroxylamine solution and 15 min incubation at room temperature. Each labeled sample was pooled, frozen, and lyophilized and subjected to SPE on a High-Density 3M Empore SDB-XC column. The eluent was lyophilized. Peptides were fractionated using high pH reverse-phase chromatography on an Agilent 1100 HPLC system using a Waters XBridge C18 column (2.1 mm ID × 150 mm length, 3.5-μm particle size) at 300 μl/min. The following gradient was employed: 0.5% B initial conditions, 0.5–3.0% B from 0 to 1 min, 3–25% B from 1 to 36 min, 25–45% B from 36 to 44 min, 45–90% B from 44 to 47 min, 90% B from 47 to 49 min, and 90–0.5% buffer B from 49 to 50 min (buffer A: 100% H2O, 10 mM NH4OH; buffer B: 100% CH3CN, 10 mM NH4OH). Every 12th well was combined to create 12 pools. Each pool was lyophilized. Mass spectrometry: Peptides (10% per pool) were analyzed by nano LC/MS/MS with a Waters NanoAcuity HPLC system interfaced to a ThermoFisher Fusion Lumos mass spectrometer. Peptides were loaded on a trapping column and eluted over a 75-μm analytical column at 350 nl/min; both columns were packed with Luna C18 resin (Phenomenex). Each high pH RP pool was separated over a 2-h gradient (24 h instrument time total). The mass spectrometer was operated in data-dependent mode, with MS and MS/MS performed in the Orbitrap at 60,000 FWHM resolution and 50,000 FWHM resolution, respectively. A 3-s cycle time was employed for all steps. Data analysis: Data were analyzed using MaxQuant v1.6.2.3 (Max Planck) and searched against the combined forward and reverse Swissprot Homo sapiens protein database. The database was appended with common background proteins. Search parameters were precursor mass tolerance 7 ppm, product ion mass tolerance 20 ppm, 2 missed cleavages allowed, fully tryptic peptides only, fixed modification of carbamidomethyl cysteine, variable modifications of oxidized methionine and protein N-terminal acetylation. Data were filtered 1% protein and peptide level false discovery rate sand requiring at least one unique peptide per protein. Reporter ion intensities were exported for further analysis.

Untargeted high-resolution LC-HRMS lipidomic analysis
Sample preparation. Metabolic quenching, lysis, and lipid extraction was performed by adding 500 μl ice-cold PBS. Crude mitochondrial samples were homogenized using MP Bio Matrix A tubes
at 60 Hz for 1 min; 400 μl uncleared supernatant was transferred to a clean glass tube containing 10 μl LipidSplash deuterated lipid internal standards (Avanti Polar Lipids, Alabaster, AL) and subjected to a Folch extraction. Samples were rested on ice for 10 min before phase separation via centrifugation at 2500 × g for 15 min; 700 μl of organic phase was dried to completed under nitrogen gas and re-suspended in 1:1 acetone/trifluoroisopropanol, and 3 μl of sample was subjected to online LC-MS analysis.

**LC-HRMS method.** Analyses were performed by untargeted LC-HRMS. Briefly, samples were injected via a Thermo Vanquish UHPLC and separated over a reversed-phase Thermo Accucore C-18 column (2.1 × 100 mm, 5 μm particle size) maintained at 55°C. For the 30 min LC gradient, the mobile phase consisted of the following: solvent A (50:50 H2O:ACN 10 mM ammonium acetate/0.1% acetic acid) and solvent B (90:10 IPA:ACN 10 mM ammonium acetate/0.1% acetic acid). Initial loading condition is 30% B. The gradient was the following: over 2 min, increase to 43% B, continue increasing to 55% B over 0.1 min, continue increasing to 65% B over 10 min, continue increasing to 85% B over 6 min, and finally increasing to 100% B over 2 min. Hold at 100% for 5 min, followed by equilibration at 30% B for 5 min. The Thermo IDX tridix mass spectrometer was operated in both positive and negative ESI mode. A data-dependent MS2 method was used for scanning in Full MS mode from 200 to 1500 m/z at 120,000 resolution with an AGC target of 5e4 for triggering ms2 fragmentation using stepped HCD collision energies at 20, 40, and 60% in the orbitrap at 15,000 resolution. Source ionization settings were 3.5 and 2.4 kV spray voltage, respectively, for positive and negative mode. Source gas parameters were 35 sheath gas, 5 auxiliary gas at 300°C, and 1 sweep gas. Calibration was performed prior to analysis using the Pierce FlexMix Ion Calibration Solutions (Thermo Fisher Scientific). Internal standard peak areas were then extracted manually using Quan Browser (Thermo Fisher Xcalibur ver. 2.7), normalized to weight and internal standard peak area, then graphed using GraphPad PRISM (ver. 9.0). Untargeted differential comparisons were performed using LipidSearch 4.2 (Thermo Fisher) to generate a ranked list of significant lipid compounds at the class and species-specific levels.

**Statistical analysis**
Statistical significances were determined using Prism software (GraphPad Software) as indicated in the figure legends.

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