The anti-viral dynamin family member MxB participates in mitochondrial integrity

Hong Cao¹², E.W. Krueger², Jing Chen², Kristina Drizyte-Miller³, Mary E. Schulz² & Mark A. McNiven¹²

The membrane deforming dynamin family members MxA and MxB are large GTPases that convey resistance to a variety of infectious viruses. During viral infection, Mx proteins are known to show markedly increased expression via an interferon-responsive promoter to associate with nuclear pores. In this study we report that MxB is an inner mitochondrial membrane GTPase that plays an important role in the morphology and function of this organelle. Expression of mutant MxB or siRNA knockdown of MxB leads to fragmented mitochondria with disrupted inner membranes that are unable to maintain a proton gradient, while expelling their nucleoid-based genome into the cytoplasm. These findings implicate a dynamin family member in mitochondrial-based changes frequently observed during an interferon-based, anti-viral response.
Mitochondria are dynamic organelles that undergo fission and fusion events in response to changes in metabolic needs and important cellular processes such as cell division and apoptosis. Central to these alterations in mitochondrial form are the dynamin family of large GTPases, which are known to bind and remodel membranes. Distinct from the conventional dynamins (Dyn1, 2, 3) that are known to support membrane scission at distinct cellular sites such as the plasma membrane, the Golgi apparatus, as well as endosomes and autolysosomes, the dynamin-related proteins have been implicated in the dynamics of both mitochondria and peroxisomes. Although highly conserved within the N-terminal GTPase domains, these dynamin-related proteins are only modestly conserved in the middle domain and generally lack pleckstrin binding and proline-rich domains found in the conventional family members. The mitofusins (MFN1/MFN2) are believed to support the outer mitochondrial membrane fusion and optic atrophy 1 (OPA1) has been implicated in the fusion of the inner membrane. Reciprocally, dynamin-related protein 1 (DRP1) has been shown to constrict the outer mitochondrial membrane with the final scission event perhaps driven by conventional Dyn2.

In addition to the mitochondria-associated dynamins are the myxovirus resistance GTPases (MX1/2 human and MxA/B mouse), which are <40% similar to the conventional dynamins but over 60% identical in sequence to each other. These related proteins are expressed in cells upon exposure to type 1 interferon (IFN) and convey a cellular innate immune response to a variety of different pathogenic viruses and influenza, hepatitis B (MxA), and HIV-1 (MxB). We and others have found that MxA can assemble into polymers that act to deform membranes and appear to reside on components of the smooth endoplasmic reticulum (ER). MxB also assembles into polymeric structures and was originally found to associate with nuclear pores where it was predicted to restrict the access of a viral genome to the host transcription machinery. MxB has been described to bind to the HIV-1 genome, while impairing its nuclear localization and possibly the fusion of the inner mitochondrial membrane. The long form of MxB associates with both organelles and has been shown to be expressed in two forms, one of which contains an N-terminal sequence required for nuclear targeting. The distribution of the long and short MxB forms have been examined in significant detail by others. In our hands we find that the short form has modest affinity for the nucleus or mitochondria regardless of the tag used in comparison with the long form that associates with both organelles.

### Results

**MxB associates with mitochondria in cultured cells.** As the related family member MxA has been shown to convey a partial resistance to the hepatocyte-centric HBV, while associating with membranous structures in hepatocytes, our goal is to compare the distribution and expression of the highly related family member MxB in hepatocyte cell lines and in the whole liver. Using several different antibodies and tagged constructs to MxB, we observe the punctate nuclear staining previously observed by others. In addition to this localization, we observe a fusiform and linear staining pattern in the hepatoma cell line Hep3B and HeLa cells, using a previously published MxB antibody and exogenously expressed green fluorescent protein (GFP)/Myc-tagged constructs (Fig. 1a–d and Supplementary Fig. 1a–c). Co-staining these cells with a variety of different organelle marker antibodies reveals that the MxB labeling represents mitochondria as confirmed by antibodies to cytochrome c oxidase 4 (Fig. 1a–d and Supplementary Fig. 1a–c). This localization is consistent when staining with three distinct MxB antibodies or expressing either GFP or Myc-tagged MxB, whereas the characteristics of labeling using these different probes varies from fine to large puncta coating the mitochondria.

Western blot (WB) analysis of various human tissues for endogenous MxB shows that MxB appears enriched in the liver, lymph node, testis, tonsil, and pig liver. Of note is the fact that the human lymph node and tonsil tissues possess two bands, whereas the liver, testis, and pig liver indicate a single band (Fig. 1e). Accordingly, appreciable levels of endogenous MxB are observed in several human hepatoma cell lines (Hep3B, HepG2, Huh7) and primary hepatocytes isolated from human and pig livers (Fig. 1f). The liver. Cells appear to have modest MxB levels that become more apparent upon longer exposures (Supplementary Fig. 4). As reported previously, MxB expression is markedly increased in isolated cells treated with 1000 IU ml⁻¹ of IFNα (Fig. 1g).

MxB has been shown to be expressed in two forms, one of which contains an N-terminal sequence required for nuclear targeting. The distribution of the long and short MxB forms have been examined in significant detail by others. In our hands we find that the short form has modest affinity for the nucleus or mitochondria regardless of the tag used in comparison with the long form that associates with both organelles (Supplementary Fig. 1d–j).

As our initial morphological observations (Fig. 1 and Supplementary Fig. 1) show MxB associating with mitochondria as either small or large puncta, or more diffuse in nature, we think it is important to analyze live cells expressing MxB-mCherry to test for changes in MxB localization over time. One example of this dynamic distribution is depicted in Fig. 2a, showing a single Hep3B cell expressing MxB-mCherry viewed over a 16 h time period. Dramatic and unexpected changes in MxB localization are observed, as the tagged protein appears to cycle between diffuse and punctate forms during the extended viewing period. Subsequently, cells co-transfected with Mito-GFP as a mitochondrial marker test whether the coalescing MxB puncta might associate with this organelle. Indeed, as shown in the images from movies of two distinct live cells (Fig. 2b, c), MxB-mCherry appears diffuse and barely detectable at early viewing times but subsequently coalesces into numerous large puncta that associate with the tips of undulating mitochondria (inserts). As observed in Fig. 2a, these puncta appear to dissolve over time, leaving the MxB in a diffuse state.

**MxB manipulations disrupt mitochondrial structure and function.** Based on the morphological observations described above, we next test whether manipulation of MxB levels might alter mitochondrial morphology. Hep3B cells transfected with either GFP-tagged or untagged wild-type (WT) or GTPase-defective mutant K131A MxB for 24 h are followed by staining with the CoxIV marker antibody. Representative images displayed in Fig. 3a, b show that mutant MxB-transfected cells possess fragmented mitochondria and significantly reduced CoxIV staining compared with adjacent untransfected cells. Importantly, in many transfected cells, mitochondrial distribution, size, and shape appear abnormal. These altered phenotypes include fragmented, twisted, or clustered morphologies (Fig. 3c and Supplementary Fig. 3a). It is particularly surprising that 25–30% of the cells examined have markedly attenuated CoxIV staining (Fig. 3d). To define these changes at a higher resolution, Hep3B cells transfected to express WT or K131A mutant MxB for 2 days are then fixed and embedded for electron microscopy (EM). Although untransfected cells display normal mitochondrial morphologies, many Hep3B cells expressing either of the...
constructs possess mitochondria with a greatly reduced number of cristae and/or numerous vacuoles or blisters (Fig. 3e–g). Similar results are seen in transfected HeLa cells, which lose almost all of their mitochondrial cristae, leaving hollow, elongated, organelle shells (Supplementary Fig. 2a–c). To better understand the effects of the mutant K131A form, we compare the distribution of this mutant with the WT form in Hep3B cells that are transfected with GFP-tagged constructs then fixed and viewed by fluorescence microscopy. Quantification of mitochondrial and nuclear fluorescence using ImageJ software shows a near-equal preference of the WT form for both organelles, while the K131A mutant exhibits increased association with mitochondria (Supplementary Fig. 2d). As a biochemical comparison, Hep3B cells expressing either the WT or K131A form subjected to subcellular fractionation to obtain nuclear, mitochondrial enriched, and cytosol fractions, are blotted for corresponding organelle markers and MxB (Supplementary Fig. 2e). By this approach, a near-equal distribution of both forms is observed.

To further define the role of MxB in maintaining mitochondrial morphology, Hep3B cells are subjected to MxB knockdown using both small interfering RNA (siRNA) and short hairpin RNA (shRNA) approaches. Following 5 days of treatment, cells are fixed and stained for CoxIV (Fig. 3h–k) or prepared for EM

Fig. 1 MxB is localized to mitochondria in cultured cells. Fluorescence images of Hep3B cells or HeLa cells, stained with a polyclonal antibody (IJ-GP) to MxB (a,c), or expressing exogenous human MxB-GFP (b,d). All cells were stained for the inner mitochondrial membrane marker CoxIV. MxB is visualized as small or large puncta that align closely with mitochondrial membranes. Boxed regions provide a higher magnification for color overlays showing the alignment of MxB (green) with mitochondria (red) in both cell types. Scale bars, 10 μm. Western blot analysis of different human tissues (Br = brain, Lng = lung, Liv = liver, LN = lymph node, SP = spleen, Tst = testis, Trns = tonsil) for endogenous MxB that appears enriched in the liver, lymph node, testis, tonsil, and pig liver. Western blotting comparing MxB expression in HeLa cells, three different hepatocyte cell lines, and a human monocyte cell line (THP-1), as well as primary human and pig hepatocytes. Expression levels are exceptionally high in primary hepatocytes, the HepG2 and Huh7 cell lines (H3B = Hep3B, HG2 = HepG2, Hu7 = Huh7, THP = THP-1, Hu = human hepatocytes). Western blotting of four distinct cell lines comparing endogenous MxB expression under control conditions vs. treatment with 1,000 IU/ml of IFN-α-2A. HeLa, Hep3B and HepG2 cells induced MxB expression by IFN treatment, but not Huh7 cells.
As observed in cells expressing exogenous WT or mutant (K131A) MxB protein, these knockdown cells display altered mitochondrial morphologies including fragmented, twisted, or clustered shapes. A comprehensive table showing the effects of MxB knockdown, or overexpression, on mitochondrial shape is provided in Supplementary Fig. 3a. In contrast to the cells overexpressing tagged MxB proteins, the knockdown cells did not exhibit reduced staining for CoxIV or Tom20. EM of siRNA-treated Hep3B cells, however, did reveal highly altered mitochondria with deformed shapes, large vacuoles, and vesiculated cristae (Fig. 3m–o). Based on our findings from the expression of either WT or mutant MxB proteins, and two distinct knockdown strategies, we conclude that this dynamin family member plays an important role in the maintenance of mitochondrial shape and integrity.

A central property of the Mx proteins is a substantially increased expression induced by IFN. Although MxB, but not MxA, is constitutively expressed in some cell and tissue types examined (Fig. 1e–g), the expression levels of both Mx proteins is markedly increased in cells treated with IFN (Fig. 1g, Supplementary Fig. 3j). We found that Hep3B cells treated with IFN for 48 h possess fragmented mitochondria in comparison with control cells (Supplementary Fig. 3b–d, h) and slightly increased levels of MxB on the nuclear envelope vs. mitochondria by immunostaining (Supplementary Fig. 3e–g,i). As a biochemical comparison, Hep3B cells treated as above with IFN for 48 h prior to lysis and differential centrifugation provides a rudimentary separation of nuclei from the mitochondria and cytosol. These fractions are run on SDS-polyacrylamide gel electrophoresis (PAGE) and probed by WB with antibodies to MxB and MxA.

![Fig. 2 MxB distribution in living cells is dynamic. a Time-lapse imaging of Hep3B cells expressing mCherry-MxB and viewed over a 16 h time period. Initially, MxB is diffuse in distribution, although condenses into several fine puncta by 5 h that subsequently dissolve but reform into much larger structures by 9 h. Surprisingly, these large aggregates also dissolve or disassemble several hours later (see Supplementary Movie 1). b, c Time-lapse imaging of Hep3B cells co-expressing mCherry-MxB (red) and Mito-GFP (white), and viewed over a 16 h time period to test for dynamic associations of MxB with mitochondria. Low-magnification images of cells show the tagged MxB changing from a diffuse to punctate, to diffuse distribution in close proximity of mitochondria. Higher magnification viewing (inserts) reveals that many of these formed puncta associate along or at the tips of the dynamic mitochondria (arrows) (see Supplementary Movies 2 and 3). Scale bars, 10 μm.

![Fig. 3m–o.](image)
Supplementary Fig. 3j). This blotting is consistent with the cell staining and suggests that although there is a large increase in MxB associated with both organelles, a slight preference is observed with the nuclear fraction. As a control comparison the IFN-induced MxA protein is observed exclusively in the cytosolic fraction.

As light microscopy imaging of MxB in cultured cells (Fig. 1) suggests an intra-mitochondrial localization consistent with the structural defects we observe by EM (Fig. 3), we utilize biochemical approaches to further define MxB localization. To this end, standard subcellular fractionation methods using pig liver as a tissue source are used. As displayed in Fig. 4a, a substantial enrichment of CoxIV is observed through sequential gradient centrifugation steps when comparing liver homogenate with a highly enriched mitochondrial fraction (Pure-Mito). Importantly, MxB protein levels are markedly increased in the “pure” mitochondrial fraction compared with all other isolated fractions, including the nuclear pellet fraction in which MxB is expected to reside. We next subject this “pure” mitochondrial preparation to a protease protection assay.
used often to define the topology of proteins associated with membranous organelles\textsuperscript{40}. A freshly isolated, enriched, mitochondrial fraction is treated with different concentrations of trypsin for 20 min at 4 °C then solubilized in sample buffer prior to SDS-PAGE and WB analysis for MxB and the mitochondrial proteins OPA1, Tom20, or CoxIV (Fig. 4b). Each of these four proteins display distinct sensitivities to the protease with the outer mitochondrial transmembrane protein Tom20 showing the most degradation followed by OPA1. MxB appears to be largely protected from the protease treatment while the inner membrane protein CoxIV remains intact, perhaps due to a tight association with the 12 other members of the respiratory Complex IV.

To support the morphological and biochemical observations described above, we attempt to define the localization of MxB at the ultrastructural level using a recently developed APEX2 approach for EM, in which cultured cells express the protein of interest tagged with a 28 kDa ascorbate peroxidase (APEX2)\textsuperscript{41}. The chimeric protein maintains its activity within cells despite aldehyde fixation and allows the focused deposition of an electron-dense reaction product where the protein is localized. This application has been used to provide accurate localization for a series of known cellular proteins including several to different parts of the mitochondria including the outer mitochondrial membrane, the intermembrane space, and the matrix\textsuperscript{41}. Accordingly, Hep3B cells expressing an MxB-APEX2 construct...
are then processed and viewed by EM. As shown in Fig. 4c, mock-transfected cells that are subsequently treated the same as MxB-APLEX2-transfected cells, possess normal mitochondria of low contrast due to the lack of post staining. In contrast, cells expressing MxB WT-APLEX2 exhibit dense staining of nuclear pores (Fig. 4d, arrowheads), consistent with the findings of others along with darkly stained outer and inner mitochondrial membranes (arrows). Many of these mitochondria also appear to have reduced number of cristae. Cells expressing the mutant MxB K131A-APLEX2 display mitochondria filled with considerably darker reaction product in the matrix (Fig. 4f, g), which is also observed in cells viewed by fluorescence microscopy (Fig. 4h, i).

As the biochemical and morphological localizations of MxB are consistent with the observed changes in mitochondrial cristae in MxB-altered cells, it is important to test whether mitochondria within these cells had normal or impaired function. To this end, Hep3B cells are transfected to express vector, WT, or mutant MxB protein (Fig. 5a–c) or, alternatively, are treated with either non-targeted siRNA or siRNA to MxB for 5 days (Fig. 5d, e) prior to incubation with 50 nM Rho123 dye for 30 min as a probe for mitochondrial membrane potential. The capacity of the mitochondria to actively sequester this dye is widely used to measure the bioenergetics of this organelle. Importantly, all cells in which MxB levels are manipulated show substantially less Rho123 fluorescence than did control cells (Fig. 5f), suggesting that cells with morphologically altered mitochondria due to MxB manipulation have reduced cristae and are functionally compromised.

**Discussion**

In this study, we report an unexpected subcellular distribution and function for the dynamin family member MxB at the mitochondria. Although MxB is well described to exhibit an IFN-inducible expression, we find it is also expressed at basal levels in several untreated tissues including human liver, lymph node, testis, and tonsil, as well as pig liver (Fig. 1e).

Further, MxB is expressed in a variety of hepatocyte cells including primary hepatocytes (Fig. 1f) and this expression is markedly increased upon IFN treatment (Fig. 1g). This basal level of expression may suggest that MxB performs housekeeping functions in cells in addition to its anti-viral action. Although HeLa cells do not show an MxB band in the blot provided, this can be resolved upon higher protein loading and/or longer exposure of the blot. In addition, alterations of MxB function in these cells lead to disrupted mitochondria (Fig. 1 and Supplementary Fig. 2). Further, despite our continued efforts, we have been unable to isolate viable MxB-knockout clones using CRISPR/Cas9 in HeLa or Hep3B cells, suggesting a critical function of this protein. The higher molecular weight band of MxB in human liver (Fig. 1e), Hep3B, HepG2, and human primary hepatocytes (Fig. 1f) showed prominent expression. Interestingly, when treating cells with IFN, the short form of MxB is induced, indicating that it may exist constitutively with the long form of MxB in the liver/hepatocytes. However, this lower molecular weight form of MxB needs to be confirmed to indeed be a truncated form of MxB.

The widely reported distribution of MxB at the cyttoplasmic face of nuclear pores is consistent with this protein containing an N-terminal nuclear localization sequence (NLS), making it readily available to prevent the nuclear uptake and chromosome integration of lentiviruses such as HIV-1 into the host genome. In addition to the reported localization at nuclear pores, numerous other studies have described a cytoplasmic distribution for MxB, which we believe may have represented the mitochondria. From the data described above, the localization of MxB at the mitochondria is observed consistently in several different cell types, while using multiple MxB antibodies or tagged constructs (Fig. 1a–d and Supplementary Fig. 1a, b). Although all of these reagents indicate a nuclear and mitochondrial distribution for MxB, distinctions in the types of mitochondrial staining were observed, which varied from evenly distributed to small or large puncta. Digital recording of live Hep3B cells expressing MxB-mCherry demonstrated that a single cell may display all of these distributions (Fig. 2). It was particularly surprising that such large aggregates of MxB can form and disassociate, as static images of these structures suggested the formation of insoluble or denatured aggregates of a tagged protein. The dissolution of these aggregates suggests otherwise and may provide insights into the function of this antiviral dynamin protein. Whether this cycling process leads to an association of MxB with the surface or interior of the mitochondria needs to be determined. We do observe that the mitochondrial distribution is significantly increased in cells expressing an MxB K131A mutant (Fig. 4h, i), a finding that provided us with the initial motivation for pursuing the mitochondrial association.

**Loss of MxB expells mitochondrial genome into the cytoplasm.**

From the combined observations detailed above showing altered mitochondrial function, form, and disrupted cristae upon manipulations of MxB levels, we predict that these mitochondria might also exhibit alterations in their genome stability. Indeed, the mammalian dynamin family member OPA1 is also named MGM1 in yeast (for mitochondria genome maintenance), as the loss of this gene leads to disorganized cristae and loss of mitochondrial DNA. The mitochondrial genome of 37 genes is packaged into spherical, punctate structures known as nucleoids, which can be observed using various DNA or protein antibodies. To test the effects of MxB on the morphology and distribution of mitochondrial nucleoids, Hep3B cells are treated with siRNA or shRNA to MxB, as well as non-targeted siRNA or vector-alone shRNA (TRC2) as a control for 5 days prior to fixation and staining with antibodies to mitochondrial DNA (mtDNA) and CoxIV (Fig. 5g–k). OPA1 levels also reduced via siRNA are used as a comparison (Fig. 5i). In control cells, each mitochondria contains one to ten nucleoids (Fig. 5g, j). In striking contrast, the mitochondria of MxB knockout cells are markedly fragmented (Fig. 5h, k) with fewer nucleoids and many of the nucleoids appear to reside within the matrix. Morphological quantification of these cells shows a three to four fold increase in cytoplasmic distribution of mitochondrial nucleoids in MxB-knockdown cells vs. control cells (Fig. 5i). This loss of mtDNA is substantially less pronounced in cells subjected to a knockdown of the inner mitochondrial dynamin OPA1 (Fig. 5i, l). To support these morphological observations, we utilize quantitative PCR (qPCR) methods to measure total mtDNA as well as cytosolic mtDNA content from Hep3B cells subjected to siRNA knockdown of MxB and/or OPA1 (Fig. 5m, n). Interestingly, total DNA levels are nearly doubled in the knockdown cells compared with control cells. Consistent with the morphological measurements, cytoplasmic mtDNA levels in OPA1-knockdown cells are modestly increased compared with control cells, whereas these levels are markedly increased (three to four fold) in MxB-knockdown cells. To test whether the MxB-induced loss of mtDNA is dependent upon OPA1 function, Hep3B cells treated with siRNAs to both OPA1 and MxB are subjected to morphometry and qPCR as described above. When both of these dynamin family members are reduced, the loss of mtDNA into the cytoplasm trends toward control levels, suggesting that OPA1 may play a role in the genomic instability induced by a reduction of the MxB protein.

**Discussion**

In this study, we report an unexpected subcellular distribution and function for the dynamin family member MxB at the mitochondria. Although MxB is well described to exhibit an IFN-inducible expression, we find it is also expressed at basal levels in several untreated tissues including human liver, lymph node, testis, and tonsil, as well as pig liver (Fig. 1e). Further, MxB is expressed in a variety of hepatocyte cells including primary hepatocytes (Fig. 1f) and this expression is markedly increased upon IFN treatment (Fig. 1g). This basal level of expression may suggest that MxB performs housekeeping functions in cells in addition to its anti-viral action. Although HeLa cells do not show an MxB band in the blot provided, this can be resolved upon higher protein loading and/or longer exposure of the blot. In addition, alterations of MxB function in these cells lead to disrupted mitochondria (Fig. 1 and Supplementary Fig. 2). Further, despite our continued efforts, we have been unable to isolate viable MxB-knockout clones using CRISPR/Cas9 in HeLa or Hep3B cells, suggesting a critical function of this protein. The higher molecular weight band of MxB in human liver (Fig. 1e), Hep3B, HepG2, and human primary hepatocytes (Fig. 1f) showed prominent expression. Interestingly, when treating cells with IFN, the short form of MxB is induced, indicating that it may exist constitutively with the long form of MxB in the liver/hepatocytes. However, this lower molecular weight form of MxB needs to be confirmed to indeed be a truncated form of MxB.

The widely reported distribution of MxB at the cyttoplasmic face of nuclear pores is consistent with this protein containing an N-terminal nuclear localization sequence (NLS), making it readily available to prevent the nuclear uptake and chromosome integration of lentiviruses such as HIV-1 into the host genome. In addition to the reported localization at nuclear pores, numerous other studies have described a cytoplasmic distribution for MxB, which we believe may have represented the mitochondria. From the data described above, the localization of MxB at the mitochondria is observed consistently in several different cell types, while using multiple MxB antibodies or tagged constructs (Fig. 1a–d and Supplementary Fig. 1a, b). Although all of these reagents indicate a nuclear and mitochondrial distribution for MxB, distinctions in the types of mitochondrial staining were observed, which varied from evenly distributed to small or large puncta. Digital recording of live Hep3B cells expressing MxB-mCherry demonstrated that a single cell may display all of these distributions (Fig. 2). It was particularly surprising that such large aggregates of MxB can form and disassociate, as static images of these structures suggested the formation of insoluble or denatured aggregates of a tagged protein. The dissolution of these aggregates suggests otherwise and may provide insights into the function of this antiviral dynamin protein. Whether this cycling process leads to an association of MxB with the surface or interior of the mitochondria needs to be determined. We do observe that the mitochondrial distribution is significantly increased in cells expressing an MxB K131A mutant (Fig. 4h, i), a finding that provided us with the initial motivation for pursuing the mitochondrial association.
In addition to the observed fluorescence localization of MxB to the mitochondria, substantial amounts of MxB were found in isolated organelle fractions from pig liver that coincided with mitochondrial enrichment as assessed by CoxIV levels (Fig. 4a). Equally supportive are the findings from the organelle protein protection assays, suggesting that, as observed for OPA1 (Fig. 4b)\textsuperscript{40,48}, MxB is protected from mild trypsin exposure of 0.2 mg ml\(^{-1}\) and it may reside within the inner mitochondrial membrane. In contrast to OPA1, MxB remains resistant to proteolysis at higher trypsin concentrations, as is the case for CoxIV, suggesting MxB may indeed reside on the inner membrane or in the matrix within a protective complex. The mitochondrial localization of WT MxB is less clear at the ultrastructural level using the APEX2-EM method, as both the inner and outer mitochondrial
membranes appeared darker than observed in control untransfected cells (Fig. 4c–e and Supplementary Fig. 4). It is unclear, however, if this indicates that MxB resides on both the inner and outer membranes or the WT protein has a more modest, and perhaps transient, interaction with the inner membrane. A pronounced mitochondrial staining was observed in cells expressing an APEX2-tagged MxB K131A mutant protein. In these cells, the reaction product appeared to fill the matrix in its entirety with the most intense staining coating the cristae membranes while leaving, what we interpret to be, the intermembrane space unstained. This enhanced association of the GTPase mutant was also observed in transfected cells viewed by fluorescence microscopy (Fig. 4h, i). It will be important to define how MxB is targeted to the mitochondria as opposed to the nuclear envelope and how these localizations are exclusive to MxB compared with MxA. Defining this targeting process will further strengthen the concept of MxB as a bona fide mitochondrial dynamin family member, while providing new insights into its function.

GTPase defective mutants in the conventional dynamins are generally believed to bind GTP, at perhaps a reduced affinity, but remain in a bound state as hydrolysis is defective49–51. GTP-bound dynamin is thus more likely to favor polymer assembly while associating with membranes in cells52, a state also induced by the GTP analog GTPγS as observed using in vitro assays53,54. Thus, the analogous GTPase mutation in MxB could explain the marked increase in mitochondrial D-loop primer set (m, n, n = 3 independent experiments). m, n Graphs depicting quantitative PCR of either total cellular mtDNA (m) or cytosolic (n) mtDNA. Mean fold change in mtDNA copy number between NTsi, MxB, OPA1, and MxB/OPA1 siRNA-treated cells was measured by qPCR using a mitochondrial D-loop primer set (m, n, n = 3 independent experiments). Data were analyzed using a two-tailed t-test, error bars represent SD. m NTsi vs. MxBsi, OPA1si, MxBsi /OPA1si, TRC2 vs. shRNA#44: all P < 0.0001, OPA1si vs. MxBsi/OPA1si: P = 0.0007; NS, not significant (P = 0.1827). m NTsi vs. MxBsi: P = 0.000082, OPA1si: P = 0.00069, MxB/OPA1si: P = 0.00023, nNTsi vs. MxBsi: P = 0.0027, OPA1si: N5 (P = 0.2334), MxB/OPA1si P = 0.00305. Scale bars, 10 μm.

As additional insights into the functions of the Mx proteins expand, it is likely that newly identified intracellular localizations will also increase. Conventional dynamins have been reported to participate in a wide variety of key cellular processes beyond the liberation of caveolae or clathrin-coated pits from the plasma membrane. Some of these functions include budding of nascent secretory vesicles from the trans-Golgi network, vesiculation of autophagosomes and lysosomes, as well as centriole separation and midbody scission during mitosis5,6,50. MxB has been reported to possess a NLS at the N-terminus and is likely to be expressed in the mitochondria along with a profound loss of mitochondrial function. This indicates that MxB resides on both the inner and outer membranes.

The stability of the mitochondrial genome appears to be directly related to the integrity of mitochondrial cristae, as has been suggested for the yeast protein Mgm1, named after its involvement with mitochondrial genome maintenance60. A putative role for MxB in cristae fusion or genome tethering based on its localization and effects on mitochondrial structure is certainly less developed than what is now known about the yeast Mgm1 and mammalian OPA1 proteins after many years of
investigations\textsuperscript{16,61,62}. Future detailed studies on the role of MxB on the inner mitochondrial membrane dynamics will be important, although challenging, as defining the functions of putative inner mitochondrial membrane proteins are particularly difficult. It is well documented that shape, as well as crista number and structure, are often altered in response to disease, energy burden or metabolic state of the host cells\textsuperscript{63,64}. Indeed, since the discovery of yeast DNMT\textsuperscript{15} and the mammalian homolog DRP1/ DLP1\textsuperscript{66} almost two decades ago, its precise role in mitochondrial fission remains somewhat elusive, particularly with the observation that conventional Dyn2 is recruited to the constriction site and may provide the final scission event\textsuperscript{38}.

From our perspective, what makes the findings of this current study so exciting is the concept of a dynaminc family member that is markedly induced in response to viral infection also participates in mitochondrial form, function, and integrity. Further, the two known viruses which MxB attenuates are known to utilize or modify mitochondria during infection. HIV-1 infection is well described to cause alterations in mitochondrial function that engages the DNA damage response (DDR) and induces mitochondrial-centric apoptosis\textsuperscript{67,68}. In addition, herpes virus, a recently identified MxB target\textsuperscript{38}, was originally observed to have an intimate association with the host’s cell mitochondria over 60 years ago\textsuperscript{69}. It is now well known that infection by this virus results in a rapid elimination of mtDNA from the host cells\textsuperscript{70,71}. Thus, it is especially attractive to postulate that MxB could be inducing genomic changes in the host’s cell mitochondria as part of the innate immune response. It is now clear that the inappropriate discharge of mtDNA into the cytoplasm is recognized by the host cell as a “proapoptotic infection event” that engages the DNA sensor cGAS to promote STING-dependent signaling\textsuperscript{45,72,73}. It will be particularly interesting to test whether MxB is responsible for the observed mitochondrial release of mtDNA during infection by herpes virus and other pathogens.

\textbf{Methods}

\textbf{Antibody and reagents.} Rabbits were immunized with the keyhole limpet hemocyanin-conjugated human MxB N-terminal peptide, 5′-KPWPYRRSOSFSSRKYLKKEMNSFQQPO′-3′, and the crude antiserum was affinity purified using an agarose column conjugated with the appropriate high-performance liquid chromatography-purified synthetic peptide and a low pH elution buffer, according to the manufacturer’s directions (Pierce Chemical Co., Rockford, IL). The purified anti-MxB antibody (immunofluorescence staining (IF) 5 μg ml\textsuperscript{-1}; WB 1:1000) was diazoylated (molecular weight cut-off 12,000 Da), concentrated and applied to 0.04% NaN\textsubscript{3} in 100 mM KCl, 0.9 mM CaCl\textsubscript{2}, 0.5 mM MgCl\textsubscript{2} containing 0.04% NaN\textsubscript{3}. The purified antibody (immunofluorescence staining (IF) 5 μg ml\textsuperscript{-1}; WB 1:1000) was added\textsuperscript{74}. The other anti-MxB antibodies [Guinea Pig (IF 1:200; WB 1:1000) and Rabbit (IF 5 μg ml\textsuperscript{-1}; WB 1:500)] were a generous gift from Dr Ilkka Julkunen\textsuperscript{70} and Dr Chen Liang\textsuperscript{72}, the anti-MX2 rabbit polyclonal antibody (NBPI-81018) (IF 1:100; WB 1:1000) was from Novus (Centennial, CO). The CovX (E1) (IF 1:250; WB:1:1000) rabbit mAb and (AD11-B3-E8) (IF 1:100; WB:1:1000) mouse mAb; Tom20 (D18TAN); (WB:1:1000) rabbit mAb, OPA1 (D6UN); (WB:1:1000) rabbit mAb; GM130 (D6B1) XP (WB:1:1000) rabbit mAb; and GAPDH (D16H11) XP (WB:1:1000) rabbit mAb were from Cell Signaling (Danvers, MA). The OPA1 mouse mAb; GM130 (D6B1) XP (WB:1:1000) rabbit mAb; and GAPDH (D16H11) XP (WB:1:1000) rabbit mAb were from Cell Signaling (Danvers, MA). The OPA1 mouse mAb (WB:500) was from BD Biosciences (San Jose, CA). The anti-Actin (WB 1:2000) was from Santa Cruz Biotechnology, CA.

\textbf{Tissue, cell culture, and transfection.} Human tissue was from the Mayo Clinic Biobank for Gastrointestinal Health Research (Institutional Review Board (IRB) 622-0401). Consistent from previous studies because the IRB determined the frozen tissue did not constitute research involving human subjects as defined under 45 CFR 46.102. Primary pig hepatocytes isolated and cultured from female, Large Domestic Cross-bred White Pigs, 2–3 months old\textsuperscript{80}, were a kind gift from Dr Scott Nyberg, Mayo Clinic. All animals received humane care and the procedures were performed with the approval of the Institutional Care and Use Committee at Mayo Clinic and are in accordance with guidelines set forth by the National Institutes of Health. Human hepatocytes were from Biovit (Westbury, NY). HeLa cells, an adenocarcinoma from human cervix (ATCC CCL-2), and Huh-7, a hepatocellular carcinoma from human liver\textsuperscript{30}, were grown in Dulbecco’s modified Eagle medium, plus 10% fetal bovine serum, 100 μg ml\textsuperscript{-1} penicillin, and 100 μg ml\textsuperscript{-1} streptomycin ( Gibco, Waltham, MA). Hep3B and HepG2 cells, both hepatocellular carcinoma from human liver (ATCC HB-8064) and (ATCC HB-8065), respectively, were maintained in Minimum Eagle’s medium with 10% fetal bovine serum, 100 μ/mL streptomycin, and 100 μg/ml penicillin (Gibco, Waltham, MA). THP-1 cells are monocyte from acute leukemia (ATCC TIB-202). All cells are cultured in 5% CO\textsubscript{2} and 95% air at 37 °C incubator. Cells were cultured in T-75 flasks (Fisher Scientific, Pittsburgh, PA). Cells were transfected using Lipofectamine 2000 (Invitrogen, Carlsbad, CA) according to the manufacturer’s protocol.

\textbf{Immunofluorescence and electron microscopy.} For IF microscopy, cells on coverslips were rinsed with D-PBS and fixed for 20 min with 2.5% formaldehyde in PBS buffer (0.1 M Pipes pH 6.95, 3 mM MgSO\textsubscript{4}, 1 mM EGTA). After rinsing cells were permeabilized with 0.1% Triton X-100 for 2 min, rinsed in D-PBS, and incubated in blocking buffer (5% normal goat serum and 5% glycerol in D-PBS) for 1 h at 37 °C. Cells were incubated in primary antibodies diluted in blocking buffer and rinsed repeatedly in D-PBS before incubating in the appropriate fluorescently labeled secondary antibodies diluted in blocking buffer. Cells were then washed extensively with D-PBS, rinsed with distilled water, and mounted on a glass slide with mounting reagent Prolong gold (Invitrogen, Carlsbad, CA). Fluorescence micrographs were acquired using an AxioObserver D.1 epi-fluorescence microscope (Carl Zeiss, Thornwood, NY) equipped with a 100 W mercury lamp using a x63, 1.4 NA objective lens, an Orca II digital camera (Hamamatsu, Japan), T9000, and AxioVision software (Carl Zeiss, Microscopy LLC, Thornwood, NY). Images were processed using Adobe Photoshop (Adobe Systems Incorporated, San Jose, CA).

For standard transmission EM, cells on carbon-coated coverslips were rinsed in 37 °C Hank’s buffered salt solution and fixed with 37 °C primary fixative (100 mM cacodylate, 10 mM MnCl\textsubscript{2}, 4, 6, 10 mM sucrose) followed by postfixation (100 mM cacodylate pH 7.4, 200 mM sucrose) and mounted on a glass slide with mounting reagent (100 mM cacodylate pH 7.4, 200 mM sucrose) for 1 h at room temperature, rinsed three times with washing buffer (100 mM cacodylate pH 7.4, 200 mM sucrose) then fixed in the secondary fixative (50 mM cacodylate pH 7.4, 100 mM sucrose, 1% Os\textsubscript{4}O\textsubscript{4}) for 1 h at room temperature, and rinsed three times in water then fixed in 1% uranyl acetate in water for 1 h at room temperature. Samples were then dehydrated in a graded ethanol series, embedded in Qutel 651 (Ted Pella, Redding, CA), and polymerized in a 65 °C oven overnight. After removal from the oven, the coverslip was removed from the bottom of the sample, the block trimmed down to a trapezoid 1 mm wide at the base, 100 mm thin sections were cut and viewed on a Joel 1200 transmission electron microscope (Leo Ltd, Tokyo, Japan).

For cells expressing the APEX2 tag, after primary fixation for 30 min, an electron-dense reaction product was generated by incubating the samples with 1 mg ml\textsuperscript{-1} DAB-HCI (3,3′-diaminobenzidine tetrahydrochloride; Electron microscopy).
Mitochondrial puriﬁed mitochondria (mito) fractions were isolated from pig liver using a modiﬁed protocol at 4 °C. Approximately 10 g of pig liver was resuspended in 50 ml of IB-1 buffer (225 mM mannitol, 75 mM sucrose, 0.5% BSA, and 30 mM Tris-HCl pH 7.4) and homogenized 10 or more strokes at 1500 r.p.m. Homogenate was then centrifuged at 10,000 × g for 10 min. The mitochondrial pellet was resuspended in 2.25 ml of MRB (250 mM mannitol, 5 mM HEPES, and 0.5 mM EGTA). To isolate pure mito fraction, 1.8 ml of crude mito fraction was layered on Percoll gradient followed by centrifugation at 95,000 × g for 10 min. The mitochondrial pellet was resuspended in a small volume of MRB and used for mitochondria protease protection assay40. Briefly, 100 μg of freshly isolated pure mito fraction was treated with 0.2% or 1 μg/ml lysotracker in 100 μl of mitochondria digestion buffer (10 mM sucrose, 0.1 mM EGTA, and 10 mM Tris-HCl pH 7.4) for 20 min on ice. The reaction was terminated by boiling the samples in Laemmli buﬀer + β-mercaptoethanol for 5 min.

Quantitative methods and statistical analysis. For optical measurements of mitochondrial DNA, ﬂuorescence microscopy of Hep3B cells stained for CoxIV and mtDNA were analyzed with ImageJ software and the degree of overlap was calculated using the Just Another Colocalization Plugin77. Both nuclei and the whole cell were treated with carbonyl cyanide m-chlorophenylhydrazone (CCCP) to depolarize the mitochondria and their ﬂuorescence intensity set to zero to calibrate the graphs.

Mitochondrial proteolipid and protease protection assay. Enriched mitochondrial (mito) fractions were isolated from pig liver using a modiﬁed protocol at 4 °C. Approximately 10 g of pig liver was resuspended in 50 ml of IB-1 buffer (225 mM mannitol, 75 mM sucrose, 0.5% BSA, and 30 mM Tris-HCl pH 7.4) and centrifuged twice at 10,000 × g for 10 min. After the ﬁnal centrifugation, crude mito pellet was resuspended in 2.25 ml of MRB (250 mM mannitol, 5 mM HEPES, and 0.5 mM EGTA). To isolate pure mito fraction, 1.8 ml of crude mito fraction was layered on Percoll gradient followed by centrifugation at 95,000 × g for 30 min. A dense band containing pure mito at the bottom of the tube was collected using a glass Pasteur pipette and diluted 10× with MRB followed by two centrifugations at 6300 × g for 10 min. The ﬁnal pellet was resuspended in a small volume of MRB and used for mitochondria protease protection assay40. Briefly, 100 μg of freshly isolated pure mito fraction was treated with 0.2% or 1 μg/ml lysotracker in 100 μl of mitochondria digestion buffer (10 mM sucrose, 0.1 mM EGTA, and 10 mM Tris-HCl pH 7.4) for 20 min on ice. The reaction was terminated by boiling the samples in Laemmli buﬀer + β-mercaptoethanol for 5 min.

Rigor and reproducibility. The type of statistical test, n-values, and P-values are all listed in the ﬁgure legends or in the ﬁgures. All microscopy data were quantiﬁed using ImageJ or Fiji, and ﬁgures were assembled using Adobe Photoshop and are representative of at least three independent experiments with similar results. All experiments were performed at least three times, except for Fig. 4a-b, which was fractionated one time and blotted three times, Supplementary Fig. 2a–c, which was done one time and Supplementary Movie 1 is representative of 24 cells over 6 experiments and Supplementary Movies 2 and 3 are representative of 33 cells over 5 experiments.

Live-cell imaging. Hep3B cells were transfected with MxB-mCherry and mito-GFP in a six-well plate. After transfection, cells were re-plated into 35 mm glass-bottomed imaging dishes (Cell E&G, San Diego, CA) in MEM media with 10% fetal bovine serum, 100 U ml−1 penicillin, and 100 μg ml−1 streptomycin ( Gibco, Waltham, MA). The 35 mm imaging dish was placed in a stage top incubator at 37 °C and 5% CO2 on a Zeiss AxioObserver microscope (Carl Zeiss, Thornwood, NY) equipped with a Colibri 7 LED light source, a Zeiss Axioscam 702 digital camera (Carl Zeiss, Thornwood, NY) and images were captured every 5 min for 16–24 h with Zen software (Carl Zeiss, Thornwood, NY).
23. Liu, Z. et al. The interferon-inducible MxB protein inhibits HIV-1 infection. *Cell Host Microbe* **14**, 398–410 (2013).

24. Buffone, C., Schulte, B., Opp, S. & Díaz-Griffero, F. Contribution of MxB oligomerization to HIV-1 capsid binding and restriction. *J. Virol.* **89**, 3285–3294 (2015).

25. Accola, M. A., Huang, B., Al Masri, A. & McNiven, M. A. The antiviral dynamin family member, MxA, tubulates lipids and localizes to the smooth endoplasmic reticulum. *J. Biol. Chem.* **280**, 21879–21885 (2005).

26. von der Maalsburg, A., Abutbul-Ionita, I., Haller, O., Kochs, G. & Danino, D. stalk domain of the dynamin-like MxA GTPase protein mediates membrane binding and liposome tubulation via the unstructured L4 loop. *J. Biol. Chem.* **286**, 37858–37865 (2011).

27. Stertz, S. et al. Interferon-Induced, antiviral human MxA protein localizes to a distinct subcompartiment of the smooth endoplasmic reticulum. *J. Interferon Cytokine Res.* **26**, 650–660 (2006).

28. Álvarez, F. J. D. et al. CryoEM structure of MxB reveals a novel oligomerization interface critical for HIV restriction. *Sci. Adv.* **3**, e1701264 (2017).

29. Haller, O. Dynamins are forever: MxB inhibits HIV-1. *Cell Host Microbe* **14**, 352–365 (2013).

30. Fricke, T. et al. MxB binds to the HIV-1 core and prevents the uncoating process of HIV-1. *Retrovirology* **11**, 68 (2014).

31. Schulte, B. et al. Restriction of HIV-1 requires the N-terminal region of MxB as a capsid-binding motif but not as a nuclear localization signal. *J. Virol.* **89**, 8599–8610 (2015).

32. Yi, D. R. et al. Human MxB inhibits the replication of hepatitis C virus. *J. Virol.* **93**, https://doi.org/10.1128/JVI.01285-18 (2019).

33. Li, N. et al. MxA inhibits hepatitis B virus replication by interaction with hepatitis B core antigen. *Hepatology* **56**, 803–811 (2012).

34. Melén, K. et al. Human Mx protein, an interferon-a-inducible GTPase, contains a nuclear targeting signal and is localized in the heterochromatin region beneath the nuclear envelope. *J. Biol. Chem.* **271**, 23478–23486 (1996).

35. Melén, K. & Julkunen, I. Nuclear cotransplant mechanism of cytoplasmic human MxB protein. *J. Biol. Chem.* **272**, 32353–32359 (1997).

36. King, M. C., Raposo, G. & Lemmon, M. A. Inhibition of nuclear import and cell-cycle progression by mutated forms of the dynamin-like GTPase MxB. *Proc. Natl Acad. Sci. USA* **101**, 8957–8962 (2004).

37. Yoon, Y., Krueger, E. W., Oswald, B. J. & McNiven, M. A. The mitochondrial replication protein Mfs1 regulates mitochondrial fusion in mammalian cells through an interaction with the dynamin-like protein DLP1. *Mol. Cell. Biol.* **23**, 5409–5420 (2003).

38. Lee, Y. H., Tan, H. T. & Chung, M. C. Subcellular fractionation methods and strategies for proteomics. *Proteomics* **10**, 3935–3956 (2010).

39. Wieckowski, M. R., Giorgi, C., Lebiedzinska, M., Duszyński, I. & Pinton, P. Isolation of mitochondria-associated membranes and mitochondria from animal tissues and cells. *Nat. Protoc.* **4**, 1582 (2009).

40. Wei, Y., Chang, W.-C., Sumpter, R., Mishra, P. & Levine, B. Prohibitin 2 is an inner mitochondrial membrane mitochondrial protein. *J. Biol. Chem.* **271**, 23478–23486 (1996).

41. Lam, S. S. et al. Directed evolution of APEX2 for electron microscopy and strategies for proteomics. *Protein Cell* **16**, 224–238.e210 (2015).

42. Verhelst, J., Huliansky, P. & Saels, X. Mx proteins: antiviral gatekeepers that restrain the uninvited. *Microbiol. Mol. Biol. Rev.* **77**, 551–566 (2013).

43. Elachouri, G. et al. OPA1 links human mitochondrial genome maintenance to mitochondrial fusion. *Cell. Mol. Life Sci.* **75**, 389–399 (2018).

44. Takei, K. et al. Generation of coated intermediates of clathrin-mediated endocytosis on protein-free liposomes. *Cell* **91**, 131–141 (1998).

45. Takei, K., Slepnev, V. I., Rocke, V. & De Camilli, P. Functional partnership between amphiphysin and dynamin in clathrin-mediated endocytosis. *Nat. Cell Biol.* **1**, 33–39 (1999).

46. Nigg, P. E. & Pavlovic, J. Oligomerization and GTP-binding requirements of MxK for viral target recognition and antiviral activity against influenza A virus. *J. Biol. Chem.* **270**, 21879–21886 (2005).

47. Haller, O., Gao, S., von der Maalsburg, A., Daumke, O. & Kochs, G. Dynamin-like MxA GTPase: structural insights into oligomerization and implications for antiviral activity. *J. Biol. Chem.* **285**, 28419–28424 (2010).

48. Mitchell, P. S., Young, J. M., Emerman, M. & Malki, H. S. Evolutionary analyses suggest a function of Mx immunity proteins beyond lentinivirus restriction. *PLoS Pathog.* **11**, e1005304 (2015).

49. Crameri, M. et al. MxB is an interferon-induced restriction factor of human herpesviruses. *Nat. Commun.* **9**, 1980 (2018).

50. Yu-Wai-Man, P. et al. OPA1 mutations cause cytochrome c oxidation deficiency due to loss of wild-type mtDNA molecules. *Hum. Mol. Genet.* **19**, 3043–3052 (2010).

51. Guan, K., Farh, L., Marshall, T. K. & Deschenes, R. J. Normal mitochondrial structure and genome maintenance in yeast requires the dynamin-like protein of the Mgm1 gene. *Curr. Genet.* **24**, 141–148 (1993).

52. Meersen, S. et al. Mitochondrial inner-membrane fusion and cristae maintenance requires the dynamin-related GTPase Mgm1. *Cell* **127**, 383–395 (2006).

53. Ackermann, W. W. & Kurtz, H. The relation of herpes virus to host cell membrane fusion. *Annu. Rev. Cell Biol.* **1**, 131–157 (1995).

54. Accola, M. A., Huang, B., Al Masri, A. & McNiven, M. A. A novel dynamin-like protein associates with cytoplasmic vesicles and tubules of the endoplasmic reticulum in mammalian cells. *J. Cell Biol.* **170**, 779–793 (1998).

55. Badley, A. D., Roussinier, T., Lumin, J. & Kroemer, G. Mitochondrian-mediated apoptosis in HIV−1 infection. *Trends Pharmacol. Sci.* **24**, 298–303 (2005).

56. Arnaud, D., Petit, F., Lelièvre, J. D. & Estaurant, J. Mitochondria in HIV−1-induced apoptosis. *Biochem. Biophys. Res. Commun.* **304**, 561–574 (2003).

57. Ackermann, W. W. & Kurtz, H. The relation of herpes virus to host cell membrane fusion. *Annu. Rev. Cell Dev. Biol.* **16**, 311–16 (2000).

58. West, A. P. & Shadel, G. S. Mitochondrial DNA in innate immune responses and inflammatory pathology. *Nat. Rev. Immunol.* **17**, 363–375 (2017).

59. Hickey, R. D. et al. Curative ex vivo liver-directed gene therapy in a pig model of hereditary tyrosinemia type I. *Sci. Transl. Med.* **8**, 349ra39 (2016).

60. Nakabayashi, H., Takeka, K., Miyano, K., Yamane, T. & Sato, J. Growth of human hepatoma cell lines with differentiated functions in chemically defined medium. *Cancer Res.* **42**, 3858–3863 (1982).

61. Schneider, C. A., Rasband, W. S. & Eliceiri, K. W. NIH Image to ImageJ: 25 years of image analysis. *Nat. Methods* **9**, 671–675 (2012).

62. Bolte, S. & Cordelieres, F. P. A guided tour into subcellular colocalization analysis in light microscopy. *Microsc. Microanal.* **22**, 213–232 (2006).

Acknowledgements
We thank the members of the McNiven laboratory for helpful discussions, and special thanks to R.J. Schulte for helping with cell fractionation and to H.M. Salmonowicz for qPCR of mtDNA quantification technical support. This study was supported by NIH R01DK044650 and R01AA020735 and the Mayo Clinic Center for Cell Signaling in Gastroenterology (NIHDK 5P30DK084567).

Author contributions
M.A.M. wrote the manuscript. H.C. designed and performed the molecular biology, cell culture work, siRNA/shRNA knockdown, morphological analysis, and analyzed experimental data. E.W.K. performed cell culture work, time-lapse microscopy, electron microscopy, Rho 123 assay, and Fig. S1 quantification. J.C. performed all biochemistry work and analyzed experimental data. K.D.M. performed mitochondrial purification and protease protection assays. M.E.S. conducted IF and WB work, and did critical reading of the manuscript. All authors reviewed the results and approved the final version of the manuscript.
Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41467-020-14727-w.

Correspondence and requests for materials should be addressed to M.A.M.

Peer review information Nature Communications thanks Luca Scorrano and the other, anonymous, reviewers for their contribution to the peer review of this work.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.