Antisense Oligonucleotide-Mediated Silencing of Mitochondrial Fusion and Fission Factors Modulates Mitochondrial Dynamics and Rescues Mitochondrial Dysfunction

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Mitochondria are highly dynamic organelles that produce ATP and maintain metabolic, catabolic, and redox homeostasis. Mitochondria owe this dynamic nature to their constant fission and fusion—processes that are regulated, in part, by fusion factors (MFN1 and MFN2) and fission factors (DRP1, FIS1, MFF, MIEF1, MIEF2) located on the outer mitochondrial membrane. While mitochondrial fusion and fission are known to influence mitochondrial morphology and function, a key question is whether rebalancing mitochondrial morphology can ameliorate mitochondrial dysfunction in the context of mitochondrial pathology. In this study, we used antisense oligonucleotides (ASOs) to systematically evaluate the effects of fusion and fission factors in vitro. Free uptake by cells of fusion or fission factor ASOs caused robust decreases in target gene expression and altered a variety of mitochondrial parameters, including mitochondrial size and respiration, which were dose dependent. In Mfn1 knockout mouse embryonic fibroblasts (MEFs) and MFN2-R94Q (Charcot-Marie-Tooth Type 2 Disease-associated mutation) MEFs, two cellular models of mitochondrial dysfunction, we found that ASO-mediated silencing of only Drp1 restored mitochondrial morphology and enhanced mitochondrial respiration. Together, these data demonstrate in vitro proof-of-concept for rebalancing mitochondrial morphology to rescue function using ASOs and suggest that ASO-mediated modulation of mitochondrial dynamics may be a viable therapeutic approach to restore mitochondrial homeostasis in diseases driven by mitochondrial dysfunction.

Keywords: antisense, oligonucleotides, mitochondria, mitochondrial dynamics

Introduction

Mitochondria are appreciated for their primary function of energy production to sustain life. While mitochondria have their own genome that encodes 13 mitochondrial proteins, the majority of the mitochondrial proteome is derived from >1,000 nuclear-encoded genes that are synthesized in the cytosol and imported into mitochondria. Among these nuclear-encoded genes are mitochondrial fission and fusion factors that endow these endosymbiotic organelles with their characteristically dynamic nature [1].

Mitochondria exist as a complex interconnected network in the cytoplasm, and their ability to dynamically reshape their morphology is an essential component of cellular homeostasis and adaptive physiology. Mitofusin 1 (MFN1) and mitofusin 2 (MFN2) are GTPases that mediate the outer membrane fusion of neighboring mitochondria, while optic atrophy 1 (OPA1) mediates inner membrane fusion. Reduction of mitochondrial fusion factors results in mitochondrial network fragmentation and has been reported in various contexts [2].

During fission events, the GTPase dynamin-related protein 1 (DRP1) is recruited to the outer mitochondrial membrane where it binds to outer membrane-anchored adaptor proteins (mitochondrial fission 1, FIS1; mitochondrial fission factor, MFF; mitochondrial elongation factor 1, MIEF1 or MID51; mitochondrial elongation factor 2, MIEF2 or MID49) to mediate membrane constriction [2]. FIS1 was originally described as a DRP1 adaptor protein in yeast. However, recent evidence suggests that mammalian FIS1 inhibits the GTPase activity of MFN1, MFN2, and OPA1, thereby preventing fusion rather than activating fission [3]. The function of MFF is clear as a DRP1 adaptor in mammalian cells [4,5], while MIEF1 and MIEF2 have more complex mechanisms of...
Fission and fusion are processes critical for maintaining mitochondrial health as mutations in the genes that encode fission and fusion factors cause mitochondrial dysfunction and are associated with various human pathologies. Certain neuropathies, for example, are caused by mutations in Opa1 (autosomal dominant optic atrophy, ADOA) and Mfn2 (Charcot-Marie-Tooth disease type 2A, CMT2A), where patient cells have clear mitochondrial morphology imbalance and dysfunction. Abnormal expression of fission/fusion factors and aberrant mitochondrial morphology have also been widely implicated in neurodegenerative, cardiovascular, and metabolic diseases [12].

It is clear that mitochondrial morphology and function are inextricably linked as the structure of the mitochondrial network often dictates a cell’s ability to meet energy demands and selectively degrade damaged mitochondria through mitophagy [13]. Thus, in diseases caused by an imbalance of mitochondrial network morphology, a potential therapeutic approach could be to restore normal morphology and function by targeting the opposing fission or fusion pathway to ameliorate disease.

Altering the levels of mitochondrial fission or fusion factors to rebalance mitochondrial morphology has been explored over the past decade. For example, in vitro studies in COS7 cells showed that silencing Drp1, Miffl/Mift51, or Miffl/Mirt49 prevented carbonyl cyanide m-chlorophenyl hydrazone (CCCP)-induced mitochondrial fragmentation [6]. In HeLa cells, it was shown that OPA1 siRNA-induced fragmentation could be rescued by either Drp1 or Mff siRNA, while another study demonstrated that either Mfn1 or Mfn2 siRNA reversed mitochondrial enlargement in Drp1 knockout (KO) cells [4,14]. In mouse embryonic fibroblasts (MEFs), normal mitochondrial morphology was restored in Cofilin KO MEFs using Drp1 siRNA [15].

Another study used the small molecule lefunomide to rebalance mitochondrial morphology that has been explored over the past decade. For example, in vitro studies in COS7 cells showed that silencing Drp1, Miffl/Mift51, or Miffl/Mirt49 prevented carbonyl cyanide m-chlorophenyl hydrazone (CCCP)-induced mitochondrial fragmentation [6]. In HeLa cells, it was shown that OPA1 siRNA-induced fragmentation could be rescued by either Drp1 or Mff siRNA, while another study demonstrated that either Mfn1 or Mfn2 siRNA reversed mitochondrial enlargement in Drp1 knockout (KO) cells [4,14]. In mouse embryonic fibroblasts (MEFs), normal mitochondrial morphology was restored in Cofilin KO MEFs using Drp1 siRNA [15].

Antisense oligonucleotides (ASOs) provide a unique opportunity to target mitochondrial fission or fusion to restore the balance of mitochondrial morphology and function that has been disrupted in disease. ASOs affect target gene expression through various mechanisms, including RNase H1-mediated RNA degradation and splicing modulation [17,18], and are routinely used as both tools in research and as therapeutics to affect cellular function and disease [19,20]. In fact, several ASOs have been approved by the FDA to treat a broad range of diseases, and many others are currently being evaluated in the clinic for additional indications [21–26]. The wide use of ASOs in research and the clinic supports the pursuit of antisense therapeutics targeting mitochondrial dynamics.

In this study, we utilize ASOs to potently silence gene expression of each outer mitochondrial membrane fusion and fission factor and systematically evaluate their effects on mitochondrial dynamics and function. We demonstrate that these ASOs modulate mitochondrial morphology, along with changes in mitochondrial respiration and mitophagy. Importantly, we show that ASOs targeting Drp1 can rescue mitochondrial morphology and respiration by reducing mitochondrial fission in cells that exhibit excess mitochondrial fragmentation and dysfunction, including an in vitro disease model of CMT2A. This study shows for the first time that ASOs targeting mitochondrial fission factor Drp1 can bring balance to mitochondrial network morphology and restore respiration in cellular models of mitochondrial dysfunction.

### Materials and Methods

#### Synthesis of ASOs

All ASOs are 16 nucleotides in length and chemically modified with a phosphorothioate backbone and 2’-O-2′-aminoethoxyethyl (cEt) in the three nucleotides at each end (Supplementary Fig. S1). ASOs were synthesized at Ionis Pharmaceuticals, as previously described [27]. In brief, ASOs were synthesized at 40 μmol scale using UnyLinker™ solid support. cEt bridged nucleic acid (BNA) phosphorodiamidites were synthesized using reported procedures [27]. A measure of 0.1 M solution of cEt BNA in 50% toluene in acetonitrile and standard sulfur transfer, detritylation, and capping reagents were used for ASO synthesis. For each of the modified analogs fourfold excess of modified nucleoside 3’-phosphoramidite was delivered with a 12-min coupling time.

Postsynthetically, all oligonucleotides were treated with 1:1 triethylamine:acetonitrile to remove cyanoethyl protecting groups from the backbone phosphorothioate linkages. Subsequently, solid support bearing ASOs were treated with aqueous NH4OH (28–30 wt%) and heated at 55°C for 18 h. The solid support was then filtered and washed with water. Filtrate and washings were pooled together and purified by high-performance liquid chromatography (HPLC) on a strong anion exchange column (GE Healthcare Life Sciences SOURCE 30Q) using a linear gradient (buffer A: 100 mM NH4OAc in water containing 30% acetonitrile, buffer B: 1.5 M NaBr in buffer A, flow 4 or 10 mL min−1, λ 260 nm). Fractions containing full length ASO were pooled together, concentrated, and desalted by HPLC on a reverse phase column. Purity and mass of oligonucleotides were determined using ion-pair liquid chromatography—mass spectrometry analysis. ASOs were formulated in PBS, sterile filtered, and added to cell culture media. For simultaneous administration of two ASOs, 5 μM of each ASO were used.

#### Cell culture and reagents

All cell cultures were maintained at 37°C with 5% CO2, mouse hepatocellular SV40 large T-antigen carcinoma (MHT) cell line (described previously, [28]), WT MEFs (ATCC), Mfn1 KO MEFs (ATCC), Mfn2 KO MEFs (ATCC), Mfn2-R94Q MEFs, and Lenti-X-293T (Clontech) cell lines were cultured in DMEM with 10% FBS. All cell lines tested negative for mycoplasma contamination (IDEXX BioAnalytics). Lentiviral particles were produced using the Lenti-X-293T cells and Lenti-X Single Shots system (Clontech) and used to transduce MHT and MEF cell lines. Cells with stable expression of vectors were selected with 1 μg/mL puromycin.
MFN2-R94Q lentiviral vectors were created by performing site-directed mutagenesis (QuikChange II Kit; Agilent Technologies) on a mouse \textit{Mfn2} tagged TrueORF clone (Origene) and then cloned into lentiviral gene expression vector pLenti-C-Myc-DDK-P2A-Puro (Origene).

Basal mitophagy was quantified using the Mito-QC mitophagy reporter [29]. Briefly, MHT cells were transduced with pLVX-mCherry-GFP-FIS1 (MRC PPU Reagents No. DU55501) lentiviral particles, and stably expressing clones were selected with 1 \textmu g/mL puromycin.

**Western blot analysis**

Whole cell lysates were prepared from cultured cells using RIPA buffer (Thermo Fisher Scientific) and cComplete U-LTRA Protease Inhibitor Cocktail tablets (Millipore-Sigma). All lysates were sonicated then clarified by centrifugation at 10,000 rpm for 10 min at 4°C. Total protein in supernatants was quantified using the DC Protein Assay (Bio-Rad). Thirty micrograms of protein from each sample was denatured with 4 \times NuPAGE LDS Sample Buffer (Thermo Fisher Scientific) supplemented with 8% \textbeta-mercaptoethanol, separated on NuPAGE 4%–12% Bis-Tris SDS-PAGE gels with 1 \times MES running buffer (Thermo Fisher Scientific), and transferred to nitrocellulose membranes (Bio-Rad). Membranes were blocked with 5% nonfat milk (Bio-Rad) in 1 \times PBS supplemented with 0.02% Tween-20 (PBST) at room temperature blocked with 5% nonfat milk (Bio-Rad) in 1 \times PBS and then fixed in 4% paraformaldehyde diluted in culture media for 15 min at 37°C and stained with Hoechst 33342 solution (Thermo Fisher Scientific) diluted 1:1,000 in 1 \times PBS to detect nuclei. Confocal imaging was conducted on fixed cells, and 30-step Z-stacks were acquired (Leica SP8, 63X oil objective). Images were processed and analyzed using the Imaris software package (Bitplane). Three-dimensional (3D) models of the MitoTracker signal were constructed using the Imaris Surfaces Module, and mitochondrial length was quantified using the Imaris Filament Tracer module.

For mitophagy quantification, cells stably expressing the lentiviral mCherry-GFP-FIS1 mitophagy reporter vector were cultured in chambered coverslip slides (Ibidi), treated with ASOs, and then stained with 200 nM MitoTracker Red CMX-Ros (Thermo Fisher Scientific) for 30 min at 37°C. Cells were then fixed in 4% paraformaldehyde diluted in culture media for 15 min at 37°C and stained with Hoechst 33342 solution (Thermo Fisher Scientific) diluted 1:1,000 in 1 \times PBS to detect nuclei. Confocal imaging was conducted on fixed cells, and 30-step Z-stacks were acquired (Leica SP8, 63X oil objective). Images were processed and analyzed using the Imaris software package (Bitplane). Three-dimensional (3D) models of the MitoTracker signal were constructed using the Imaris Surfaces Module, and mitochondrial length was quantified using the Imaris Filament Tracer module.

**Immunofluorescent microscopy and image analysis**

For mitochondrial morphology studies, cells were cultured in chambered coverslips (Ibidi), treated with ASOs, and then stained with 200 nM MitoTracker Red CMX-Ros (Thermo Fisher Scientific) for 30 min at 37°C. Cells were then fixed in 4% paraformaldehyde diluted in culture media for 15 min at 37°C and stained with Hoechst 33342 solution (Thermo Fisher Scientific) diluted 1:1,000 in 1 \times PBS to detect nuclei. Confocal imaging was conducted on fixed cells, and 30-step Z-stacks were acquired (Leica SP8, 63X oil objective). Images were processed and analyzed using the Imaris software package (Bitplane). Three-dimensional (3D) models of the MitoTracker signal were constructed using the Imaris Surfaces Module, and mitochondrial length was quantified using the Imaris Filament Tracer module.

**Mitochondrial DNA analysis**

Total DNA was isolated from cells using a DNeasy 96 Kit (Qiagen). For each biological replicate, quantitative PCR was performed as described above, excluding the SuperScript Mix reagent in the RT-qPCR Kit and using the following thermocycling protocol (50°C for 2 min, 95°C for 10 min, then 40 cycles of 95°C for 15 s and 60°C for 1 min). Relative mitochondrial DNA (mt-DNA) amount was determined using a mt-DNA-specific primer probe set (\textit{mt-Nd2}) and normalized by a reference gene of nuclear origin (\textit{Gusb}). Primer and probe information can be found in the Supplementary Data.
before performing the assay. Cells were stimulated sequentially with one injection of 1 μM oligomycin, 2 μM trifluromethoxy carbonyl cyanide phenylhydrazone (FCCP), and 0.5 μM rotenone/antimycin A. Mix, wait, and measure times were 3 min each. Three measurements were made following each injection. Oxygen consumption rates (OCRs) were normalized by relative cell number using CyQUANT following each injection. Oxygen consumption rates (OCRs) were measured with a Spark microplate reader (Tecan). Data were analyzed with Wave software (Agilent) and exported to Prism (GraphPad).

**Flow cytometry analysis**

Cells were cultured and treated with ASOs as described and then stained with 20 nM MitoTracker Green (Thermo Fisher Scientific) or 50 nM tetracyethylrhodamine ethyl ester (TMRE) (Millipore-Sigma) for 30 min at 37°C. Trypsinized cells were resuspended in PBS supplemented with 1% FBS and analyzed using the Accuri C6 instrument (BD Biosciences). Cell debris was excluded, and live cells were gated based on forward scatter versus side scatter dot plots. Live cells were plotted on histograms, and mean fluorescent intensity was determined using FlowJo software. Unstained cells were used as negative controls, and CCCP treated cells were used as a positive control for TMRE staining (data not shown).

**Statistical analyses**

Data are reported as mean or median ± SEM. Statistical analyses were performed with an unpaired t-test, one-way ANOVA, or two-way ANOVA with multiple comparisons using Prism (GraphPad). Appropriate statistical tests and sample sizes are indicated in figure legends.

**Results**

**ASOs targeting mitochondrial fusion and fission factors exhibit potent gene silencing and alter mitochondrial morphology**

To determine whether ASOs can be used to modulate mitochondrial dynamics, we administered a panel of ASOs targeting mitochondrial fusion and fission factors (Mfn1, Mfn2, Drp1, Fis1, Mff, Mief1, Mief2) to MHT (mouse hepatocellular carcinoma) cells in culture [28]. MHT cells freely internalize ASOs in media, and after 48 h of treatment, we found that all ASOs potently reduce their target mRNA in a dose-responsive manner, with IC_{50} values ranging from 5.6 to 160 nM (Fig. 1A, B). ASO treatment also resulted in corresponding decreases in target protein levels (Fig. 1C). Treatment with a nontargeting control ASO did not alter the expression of any target we measured, and while some modest compensatory changes in protein expression of other fusion/fission factors were observed following target ASO treatment, factor levels remained generally unchanged across all conditions (Fig. 1C and Supplementary Fig. S2A–C).

Given that decreasing the levels of mitochondrial fission and fusion factors has been widely reported to alter mitochondrial network structure, we next examined how ASO treatments change mitochondrial morphology *in vitro*. MHT cells treated with fusion or fission ASOs at 5 μM for 48 h exhibited dramatic alterations in mitochondrial size when visualized with MitoTracker (Fig. 2A). Specifically, ASOs targeting fusion factors Mfn1 and Mfn2 decreased mean mitochondrial length, while the opposite was observed with ASOs targeting fission factors Drp1, Fis1, Mff, Mief1, and Mief2 (Fig. 2B). ASO-dependent changes in mitochondrial length distribution were also observed, with fusion factor ASOs increasing the proportion of small mitochondria (<1 μm) and fission factor ASOs increasing the proportion of large mitochondria (>10 μm; Fig. 2C). Interestingly, ASO-dependent changes in mitochondrial size were prevented when MHT cells were treated with two opposing ASOs simultaneously. Coadministration of one fission factor ASO together with one fusion factor ASO led to a normalization of mitochondrial sizes (Fig. 2B, C, and Supplementary Fig. S2D). These data show that modulation of mitochondrial morphology is possible through ASO-mediated gene silencing and that it occurs rapidly within 48 h.

**ASOs targeting mitochondrial fusion and fission factors alter mitochondrial respiration**

Since changes in mitochondrial morphology have been linked to changes in function, we next sought to determine if ASO-mediated silencing of mitochondrial fusion and fission factors alters mitochondrial functional parameters. Previous reports have shown a direct association between mitochondrial fragmentation and decreased mitochondrial respiratory function, and fragmentation is often a prerequisite to apoptosis [30,31]. Changes in mitochondrial morphology have also been linked to alterations in mitochondrial membrane potential, a necessary component of oxidative phosphorylation (OXPHOS)-dependent ATP production [32]. Based on these studies, we hypothesized that ASO-mediated elongation of mitochondrial networks would lead to an increase in respiration capacity, while ASO-mediated fragmentation would cause a decrease in respiration.

To measure mitochondrial respiration, we treated MHT cells with 5 μM ASOs for 48 h and performed oxygen consumption analysis using the Seahorse XFe24 Analyzer [33]. We found that basal oxygen consumption is largely unchanged upon fusion/fission factor ASO administration, although Mff, Mief1, and Mief2 ASOs did cause a slight decrease (Fig. 3A, B, E, F). Importantly, we found that maximal respiration and spare respiratory capacity were significantly affected by most ASO treatments (Fig. 3C, D, G, H). Surprisingly, however, only the Drp1 ASO enhanced maximal oxygen consumption and spare respiratory capacity, suggesting that not all forms of mitochondrial network elongation affect respiration in the same way and that additional functions of the fission factors may play a role.

Mfn1, Mief1, and Mief2 ASOs caused a decrease in oxygen consumption parameters, suggesting that mitochondrial dysfunction can occur in the context of mitochondrial fragmentation, as well as elongation. Mfn2 ASO did not change oxygen consumption parameters, despite causing mitochondrial fragmentation, suggesting that MFN2 is dispensable for mitochondrial function in MHT cells. Together, these data show that mitochondrial morphology does not always correlate with mitochondrial function and that one strategy to enhance function is by Drp1 ASO-mediated mitochondrial elongation (Fig. 3I).
ASOs targeting mitochondrial fusion and fission factors exhibit potent gene silencing in vitro. Dose-responsive activity of ASOs targeting mitochondrial fusion factors (A) and fission factors (B) measured using RT-qPCR of mRNA from MHT cells following 48 h of ASO treatment. Open circles represent Control ASO, and filled circles represent indicated ASO. Gene expression levels reported are mean (±SEM, n = 3) relative to cells treated with ASO vehicle (% of PBS). IC₅₀ values for each ASO are 6.4 nM (Mfn1), 12.8 nM (Mfn2), 5.6 nM (Drp1), 14.2 nM (Mff), 11.9 nM (Fis1), 65.7 nM (Mief1), and 160 nM (Mief2) and were determined by performing nonlinear regressions with least squares fit in Prism (GraphPad). (C) Western blots of whole cell lysates from MHT cells treated with 5 μM of indicated ASO for 48 h. Primary antibodies used to detect protein levels are indicated. Beta-actin (ACTB) served as a loading control. Western blot quantification is shown in Supplementary Fig. 2C. ASOs, antisense oligonucleotides; MHT, mouse hepatocellular SV40 large T-antigen carcinoma cell line; RT-qPCR, reverse transcription quantitative real-time PCR.

**FIG. 1.** ASOs targeting mitochondrial fission and fusion factors exhibit potent gene silencing in vitro. Dose-responsive activity of ASOs targeting mitochondrial fusion factors (A) and fission factors (B) measured using RT-qPCR of mRNA from MHT cells following 48 h of ASO treatment. Open circles represent Control ASO, and filled circles represent indicated ASO. Gene expression levels reported are mean (±SEM, n = 3) relative to cells treated with ASO vehicle (% of PBS). IC₅₀ values for each ASO are 6.4 nM (Mfn1), 12.8 nM (Mfn2), 5.6 nM (Drp1), 14.2 nM (Mff), 11.9 nM (Fis1), 65.7 nM (Mief1), and 160 nM (Mief2) and were determined by performing nonlinear regressions with least squares fit in Prism (GraphPad). (C) Western blots of whole cell lysates from MHT cells treated with 5 μM of indicated ASO for 48 h. Primary antibodies used to detect protein levels are indicated. Beta-actin (ACTB) served as a loading control. Western blot quantification is shown in Supplementary Fig. 2C. ASOs, antisense oligonucleotides; MHT, mouse hepatocellular SV40 large T-antigen carcinoma cell line; RT-qPCR, reverse transcription quantitative real-time PCR.

**ASOs targeting mitochondrial fission and fusion factors alter total mitochondrial content**

To determine if ASO-dependent changes in mitochondrial function were caused by changes in total mitochondrial content or membrane potential, we used flow cytometry to quantify total mitochondrial mass (MitoTracker Green) and mitochondrial membrane potential (TMRE) in MHT cells treated with 5 μM ASOs for 48 h. Interestingly, of all the fission factor ASOs, only the Drp1 ASO caused a significant increase in total mitochondrial mass (Fig. 4A), despite all causing mitochondrial network elongation (Fig. 2). In contrast, Mief1 and Mief2 ASOs caused a decrease in mitochondrial mass (Fig. 4A) despite a clear increase in mitochondrial length (Fig. 2), in agreement with their detrimental effects on mitochondrial respiration (Fig. 3E).

When we evaluated fusion factor ASOs, we found that Mfn1 ASO caused a decrease in total mitochondrial mass, while it remained unchanged with Mfn2 ASO treatment (Fig. 4A), despite both causing mitochondrial fragmentation.
This observation is in agreement with the disparate effects of Mfn1 and Mfn2 ASO treatment on mitochondrial respiration (Fig. 3A). Drp1 ASO also caused an increase in membrane potential, while Mfn1 ASO caused a decrease (Fig. 4B), although these changes are likely due to corresponding changes in total mitochondrial mass (Fig. 4A).

mt-DNA content, another readout for total mitochondrial mass, was largely unchanged across ASO treatments, except for Mfn1 ASO which caused a decrease (Fig. 4C). We found similarly, by western blot of mitochondrial markers, that proteins from the various mitochondrial compartments (outer mitochondrial membrane, OMM; inner mitochondrial membrane, IMM; matrix) are largely unchanged for most ASOs (Fig. 4D and Supplementary Fig. S3). However, we found that Drp1 ASO increased, while Mfn1 ASO decreased, levels of the OXPHOS components Complex II (CII) and Complex III (CIII), consistent with their effects on other readouts of total mitochondrial content and respiration (Figs. 3A and 4A–C). Overall, these data suggest that ASO-mediated changes in mitochondrial morphology may be

FIG. 2. ASOs targeting mitochondrial fission and fusion factors modulate mitochondrial morphology. (A) Representative confocal microscopy images of MHT cells treated with 5 µM of indicated ASOs for 48 h and stained with MitoTracker-Red-CMX-Ros to visualize mitochondria. Scale bars are 5 µm. (B) Average mitochondrial length (n = 6-9 images per group, one-way ANOVA with Dunnett’s multiple comparison test, *P < 0.05, **P < 0.005, ***P < 0.0005). Data are box plots from one of two separate experiments, with the center lines representing medians. Box limits indicate the 25th and 75th percentiles. Whiskers extend to the 5th and 95th percentiles. (C) Distribution of mitochondrial lengths as a percentage of total mitochondria quantified from all images in each group.

(Fig. 2). This observation is in agreement with the disparate effects of Mfn1 and Mfn2 ASO treatment on mitochondrial respiration (Fig. 3A). Drp1 ASO also caused an increase in membrane potential, while Mfn1 ASO caused a decrease (Fig. 4B), although these changes are likely due to corresponding changes in total mitochondrial mass (Fig. 4A).
FIG. 3. ASOs targeting mitochondrial fission and fusion factors alter mitochondrial respiration. Mitochondrial respiration measured by OCR over time in MHT cells treated with 5 μM of indicated ASOs for 48 h prior (A, E). Assay compounds were added at indicated timepoints (Oligomycin, FCCP, Rot/AA). Basal respiration (B, F), maximal respiration (C, G), and spare capacity (D, H) calculated from experiments in (A, E), respectively. (I) Summary of data shown in A–H. All data shown are from individual experiments representative of at least two separate experiments and reported as mean ± SEM, n = 4 (A–E), one-way ANOVA with Dunnett’s multiple comparison test, *P < 0.05, **P < 0.005, ***P < 0.0005. FCCP, trifluoromethoxy carbonyl cyanide phenylhydrazone. OCR, oxygen consumption rate; Rot/AA, rotenone/antimycin A.

FIG. 4. ASOs targeting mitochondrial fission and fusion factors modulate mitochondrial content and basal mitophagy. Total mitochondrial mass (A) and mitochondrial membrane potential (B) quantified by flow cytometry of MHT cells treated with 5 μM of indicated ASOs for 48 h, then stained with the indicated fluorescent dyes (MitoTracker-Green, TMRE). (C) Mitochondrial DNA content of MHT cells treated with indicated ASOs quantified as the ratio of mitochondrial DNA gene (mt-Nd2) to nuclear DNA gene (Gusb) and normalized to cells treated with ASO vehicle (% of PBS). (D) Western blots of whole cell lysates of MHT cells treated with 5 μM of indicated ASOs for 48 h. Primary antibodies used to detect protein levels are indicated. (E) Representative confocal microscopy images and corresponding 3D reconstructions of MHT cells stably expressing the mitophagy reporter Mito-QC and treated with 5 μM of indicated ASOs for 48 h before imaging. Scale bars are 5 μm for the upper panels and 1 μm for the lower panels. (F) Mitophagy was quantified by calculating the Mitophagy Index, defined as the ratio of the total volume of mitophagy events to the total volume of mitochondria per cell (see Materials and Methods section for details). Data are box plots from one of two separate experiments, with the center lines representing medians. Box limits indicate the 25th and 75th percentiles. Whiskers extend to the 5th and 95th percentiles (n = 6–9 images per group, one-way ANOVA with Dunnett’s multiple comparison test, *P < 0.05). All other data are shown as mean ± SEM, n = 3, one-way ANOVA with Dunnett’s multiple comparison test, *P < 0.05, **P < 0.005, ***P < 0.0005. CI, CII, CIII, CV, OXPHOS complex subunits; 3D, three-dimensional; IMM, inner mitochondrial membrane; MFI, mean fluorescence intensity; OMM, outer mitochondrial membrane; OXPHOS, oxidative phosphorylation; TMRE, tetramethylrhodamine ethyl ester.
linked to changes in function by their effects on mitochondrial content and mitochondrial membrane potential. However, it remains unclear why ASO-mediated mitochondrial elongation results in divergent effects on mitochondrial mass and function. Moreover, these data underscore the importance of utilizing multiple methods to determine total mitochondrial content in cells.

ASOs targeting mitochondrial fusion and fission factors alter basal mitophagy

Previous studies have shown that mitochondrial dynamics plays a role in modulating levels of mitophagy, which has effects on cellular mitochondrial content [34,35]. Mitophagy is the major intracellular pathway of mitochondrial removal and is
known to influence mitochondrial functions [36–38]. During mitophagy, adaptor proteins P62/SQSTM and OPTN accumulate on the outer membrane of damaged mitochondria and recruit autophagosomes by their interactions with LC3B [39]. LC3B is an autophagosome membrane component whose lipidosed form (LC3B-II) is a common readout for autophagy pathway activation [40]. Given our results suggesting differential effects of mitochondrial fusion and fission factor ASOs on mitochondrial function and total mitochondrial content, we next sought to determine if basal mitophagy is altered by these ASOs.

To investigate ASO-mediated effects on mitophagy, we focused on Drp1 and Mfn1 ASOs due to their clear effects on mitochondrial morphology, function, and content in MHT cells (Figs. 2 and 3). First, we evaluated the effects of Drp1 and Mfn1 ASOs on autophagy and mitophagy by western blot analysis and found that 5 μM treatment of MHT cells for 48 h did not cause obvious changes in LC3B, P62, and OPTN levels (Supplementary Fig. S4). These results suggest that global changes in autophagy and mitophagy are not being induced by ASO-mediated silencing of Drp1 or Mfn1. However, Drp1 ASO treatment did cause a slight increase in P62, suggesting that there could be an accumulation of mitophagy intermediates indicative of a delay in mitophagy, as previously reported [41,42].

To monitor basal mitophagy events in a more sensitive and quantitative manner, we used MHT cells stably expressing the lentiviral mitophagy reporter, Mito-QC [29,43]. Mito-QC is an outer mitochondrial membrane-targeted, pH-sensitive, tandem fluorescent reporter (mCherry-GFP). In the low pH environment of lysosomes, GFP fluorescence is quenched, and the remaining mCherry-only signals represent mitophagic events (mitochondrial fragments within lysosomes). Three-dimensional modeling of confocal images provides a quantitative readout of basal mitophagy levels. Using MHT cells stably expressing Mito-QC, we found that basal mitophagy is altered by ASO-mediated reduction of DRP1 and MFN1 (5 μM, free uptake for 48 h). Drp1 ASO treatment decreased, while Mfn1 ASO treatment increased, the level of basal mitophagy (Fig. 4E, F) suggesting that Drp1 and Mfn1 ASO treatment may be affecting mitochondrial content and, in turn, the function by modulating basal mitophagy.

Together, these results show that ASO-dependent modulation of mitochondrial morphology influences basal mitophagy in the absence of overt mitochondrial damage or stress. In addition, these results provide a mechanism that links the ASO-mediated changes in mitochondrial content with mitochondrial function, suggesting that effects on total mitochondrial content are likely due to modulation of the pathway that orchestrates their removal and degradation. Moreover, these results support the notion that mitochondrial fission is a necessary step in mitophagy, as previously reported [34], and that enabling excess mitochondrial fusion (by inhibiting fission) is sufficient to restrict basal mitophagic events. Conversely, enabling excess mitochondrial fission (by inhibiting fusion) is sufficient to enhance basal mitophagy, suggesting that mitochondrial size is an important factor in autophagosome engulfment during mitophagy.

Changes in mitochondrial morphology and respiration are ASO-dose dependent

To evaluate the dose-responsive effects of Drp1 and Mfn1 ASOs on mitochondrial morphology, we treated MHT cells with Drp1 and Mfn1 ASOs in separate dose–response experiments, then analyzed target gene expression, and quantified mitochondrial morphology. As observed previously, Drp1 and Mfn1 ASO treatments resulted in dose responsive reductions in Drp1 and Mfn1 expression (Fig. 5A, C). Similarly, we found that both Drp1 and Mfn1 ASOs also modulated mitochondrial length in a dose-dependent manner (Fig. 5A, C). Interestingly, we found significant differences in the sensitivity of MHT cells to ASO-mediated reductions in Drp1 compared to Mfn1. We observed that at least 50%–80% reduction of Drp1 expression is necessary to observe an increase in mitochondrial length, while a minimum of only 25%–40% reduction of Mfn1 expression is needed to observe a decrease in mitochondrial length (Fig. 5A, C).

We found similar ASO dose-dependent effects on mitochondrial function. In three-point ASO dose–response experiments, a dose-dependent increase in maximal respiration (OCR) was observed for Drp1 ASO, while the opposite was observed for Mfn1 ASO (Fig. 5B, D). Reflecting the ASO dose-dependent effects on mitochondrial morphology, enhancement of maximal respiration required more than ~80% reduction of Drp1, while a decrease in maximal respiration was observed at all Mfn1 ASO concentrations tested (>80% Mfn1 reduction). All together, these data suggest that for Drp1 and Mfn1 ASOs the effects on mitochondrial morphology and function are likely linked and can be modulated incrementally in response to ASO dose. Furthermore, these data showcase the enhanced functional sensitivity of cells to mitochondrial fragmentation—as opposed to mitochondrial elongation—shedding light on the dose requirements needed to effectively modulate mitochondrial dynamics in cells with ASOs.

ASO-mediated silencing of Drp1, but not other fission factors, rescues mitochondrial fragmentation and dysfunction in Mfn1 KO and MFN2-R94Q MEFs

To test whether ASO-mediated modulation of mitochondrial dynamics can rescue mitochondrial dysfunction in cells with perturbed mitochondrial morphology, we utilized two Mfn KO MEF cell lines as in vitro model systems of excess mitochondrial fragmentation: (1) Mfn1 KO MEFs and (2) Mfn2 KO MEFs stably expressing exogenous Mfn2 harboring a CMT2A mutation R94Q [44], referred to from now on as MFN2-R94Q. Both cell lines exhibit mitochondrial fragmentation and dysfunction compared to WT MEFs (Supplementary Fig. S5A), as reported previously [30,45,46]. MFN2-R94Q expression was confirmed in lentivirally transduced Mfn2 KO MEFs at the protein and mRNA level (Supplementary Fig. S5B). We sought to correct the mitochondrial dysfunction in these cells with treatment of fission factor ASOs in an attempt to restore mitochondrial homeostasis. We hypothesized that despite the negative effects on respiration of some fission factor ASOs in unperturbed cells (Fig. 3E–I), treatment of fission factor ASOs in a disease context may provide a beneficial effect.

To measure the effects of ASOs on mitochondrial respiration in MEFs, we performed oxygen consumption analysis using the Seahorse XF24 Analyzer. Administration of fission factor ASOs to MEF cell lines by free uptake at 5 μM for 48 h caused significant changes in mitochondrial function (Fig. 6A, B). In Mfn1 KO MEFs, Drp1 ASO enhanced basal OCR, maximal OCR, and spare respiratory capacity, while
ASOs targeting other fission factors were either detrimental to or did not change respiration (Fig. 6A and Supplementary Fig. S5C, F). Similarly, Drp1 ASO consistently enhanced basal OCR, maximal OCR, and spare respiratory capacity in MFN2-R94Q MEFs (Fig. 6B and Supplementary Fig. S5C, G–I). However, unlike in Mfn1 KO MEFs, Fis1 ASO enhanced basal and maximal OCR, but did not spare respiratory capacity (Supplementary Fig. S5G–I). Although Mief1 and Mief2 ASOs decreased spare capacity, other ASOs targeting fission factors did not greatly change mitochondrial function in MFN2-R94Q MEFs (Fig. 6B and Supplementary Fig. S5G–I). Paradoxically, Mief2 ASO caused an increase in basal OCR, in the absence of an increase in other OCR measures (Supplementary Fig. S5G–I). Overall, only Drp1 ASO enhanced the various OCR measures in both cellular models of mitochondrial dysfunction (Fig. 6C).

Since Drp1 ASO was the only fission factor ASO that consistently enhanced mitochondrial function in Mfn1 KO and MFN2-R94Q MEFs, we next sought to evaluate the effects of Drp1 ASO treatment on mitochondrial morphology in these cells. In accordance with enhanced mitochondrial function, Drp1 ASO caused an increase in mitochondrial length in both Mfn1 KO and MFN2-R94Q MEFs (Fig. 6D, E). Interestingly, Drp1 ASO failed to restore mitochondrial morphology in Mfn1/Mfn2 double KO MEFs, suggesting that the presence of either MFN1 or MFN2 is necessary for Drp1 ASO-mediated rescue.

Together, these results provide a novel proof-of-concept for ASO-mediated rescue of mitochondrial morphology and function in contexts of mitochondrial dysfunction due to an imbalance in mitochondrial dynamics. Furthermore, the data suggest that while Drp1 ASO consistently rescued
FIG. 6. ASO-mediated silencing of Drp1 rescues mitochondrial dysfunction in Mfn1 KO and MFN2-R94Q MEFs. Mitochondrial respiration measured by OCR over time in Mfn1 KO MEFs (A) and MFN2-R94Q MEFs (B) treated with the 5 μM of indicated ASOs 48 h prior. Assay compounds were added at indicated timepoints. Data shown are from individual experiments representative of at least two separate experiments and reported as mean ± SEM, n = 4. See Supplementary Fig. 5 for basal OCR, maximal OCR, and spare respiratory capacity measures. (C) Table summarizing the effects of indicated ASOs on mitochondrial respiration in Mfn1 KO and MFN2-R94Q MEFs. (D) Representative confocal microscopy images of Mfn1 KO, MFN2-R94Q, and Mfn1/2 KO MEFs treated with 5 μM of indicated ASOs for 48 h and stained with MitoTracker-CMX-Ros to visualize mitochondria. Scale bars are 5 μm. (E) Average mitochondrial length (n = 4 images per group, two-way ANOVA with Sidak’s multiple comparisons test, *P < 0.05, ***P < 0.0005). Data are box plots from one of two separate experiments, with the center lines representing medians. Box limits indicate the 25th and 75th percentiles. Whiskers extend to the 5th and 95th percentiles. KO, knockout; MEFs, mouse embryonic fibroblasts.
mitochondrial dysfunction, the functional effects of targeting Mff, Fis1, Mief1, and Mief2 are likely context dependent. Notably, Drp1 ASO was able to overcome the dominant negative nature of the R94Q mutation in MFN2 suggesting that other dominant negative mutations affecting mitochondrial fusion may be amenable to this approach.

Discussion

The dynamic regulation of mitochondrial shape is essential for the various roles mitochondria play in the cell. Metabolic adaptation, stress response, apoptosis, and even oogenesis and fertilization require mitochondrial dynamism through the action of mitochondrial fission and fusion factors [47–49]. With the growing number of diseases that are associated with mitochondrial dysfunction [50], it is critical to discover novel therapeutics that aim to restore mitochondrial homeostasis.

This study reports, for the first time, the use of ASOs to systematically evaluate the effects of reducing fission and fusion factors on both mitochondrial morphology and mitochondrial respiration in unperturbed cells in a dose-dependent manner. Furthermore, we demonstrate that a Drp1-targeting ASO, but not other fission factor ASOs, can rescue mitochondrial dysfunction in two cellular disease models, including one containing a disease-relevant dominant-negative mutation in MFN2 (R94Q). Altogether, this study is a proof-of-concept for the beneficial rebalancing of mitochondrial morphology to restore mitochondrial function and paves the way for further drug discovery for various mitochondrial genetic diseases for which there is significant unmet medical need.

Targeting mitochondrial fission pharmacologically has been previously reported using peptides and small molecules in various disease models with some success. The peptide P110 inhibits DRP1’s association with FIS1 and caused increased mitochondrial size and membrane potential, in agreement with the present study [51]. Importantly, P110 prevented neurotoxicity in multiple mouse models and prevented Escherichia coli-mediated mitochondrial fragmentation associated with gut permeability in Crohn’s disease [52–54]. Mdivi-1, a small molecule inhibitor of DRP1’s GTPase activity, has also been used to prevent disease-associated mitochondrial fragmentation; however, more recently the specificity and mechanism of action of Mdivi-1 have come into question [55].

In contrast, small-molecule peptide mimics have been used to restore mitochondrial fusion activity in disease models in vitro through allosteric activation of MFN1 and MFN2 [56]. While these efforts represent significant advancements in the field, more studies are needed to fully characterize the specificity and off-target effects of these compounds before their advancement into the clinic. ASOs, on the other hand, present a more targeted approach to modulating mitochondrial fission and fusion and possess a clinically favorable profile [57,58].

In this study, we show how reducing DRP1 with ASOs enhances mitochondrial function in both unperturbed cells and models of mitochondrial dysfunction. We attribute this effect to the reduction of mitochondrial removal through the mitophagy pathway, as similarly described in previous studies [34,48,59]. However, the functional effects of reducing DRP1 in cells are mixed in the literature. For instance, one study reported reduced mitochondrial respiration in cardiomyocytes lacking Drp1 [42], while another reported enhanced respiration in Drp1 KO MEFs [60]. These discrepancies support the notion that the effect of DRP1 reduction and mitophagy is cell-type specific. While long-term developmental reduction of DRP1 is likely detrimental to unperturbed cells due to reductions in mitophagy, we show here that a transient reduction in DRP1 using ASOs may be a safer approach to restoring mitochondrial homeostasis.

It has not escaped our notice that there are discrepancies in the mitochondrial morphology-function relationship outside of our focus on DRP1 and MFN1. We found that ASO-mediated reduction of MFN2 did not alter mitochondrial function while MFN1 did, suggesting that MHT cells don’t require MFN2 to sustain proper respiration in the face of mitochondrial fragmentation. Differential functions of MFN1 and MFN2 have been reported previously suggesting a potential reason for this finding [61]. In addition, the increase in mitochondrial length that we reported with ASOs targeting Mff, Fis1, Mief1, and Mief2 and the resulting lack of mitochondrial mass and respiration enhancement is counterintuitive.

It is possible that DRP1’s extramitochondrial effects may be influencing respiration in this context. For instance, DRP1 is a known regulator of peroxisome fission [62,63], and we observed elongated peroxisomes in MHT cells treated with Drp1 ASO (data not shown). Furthermore, peroxisomes perform beta-oxidation of fatty acids, and we found that MHT cells treated with Drp1 ASO have an enhanced dependence on fatty acids as fuel for mitochondrial respiration (data not shown). This shift in fuel preference, in combination with increased mitochondrial content, may be contributing to the enhancement in respiration observed with Drp1 ASO.

In contrast, it is possible that additional known or unknown functions of MFF, FIS1, MIEF1, and MIEF2, other than their roles in mitochondrial fission, may counteract the beneficial effects of suppressed fission on mitochondrial respiration. These are interesting hypotheses that are beyond the scope of this study and may be the focus of future studies. Nevertheless, our systematic evaluation of ASOs targeting each outer mitochondrial membrane fission factor is the first study to show the effects each fission factor has on mitochondrial morphology and function in unperturbed cells, as well as effects on respiration in an in vitro disease model of CMT2A.

The use of ASOs provides the benefit of being able to fine-tune mRNA expression level in a dose-dependent manner, which we show leads to concomitant changes in mitochondrial morphology and function. While previous studies have shown that a partial reduction in Drp1 expression results in significant changes in mitochondrial morphology [64], our study is the first to our knowledge to explore this notion further in multidose–response experiments. These findings have important implications for ASO drug development since knowing how much target reduction is required to observe a functional effect (eg, mitochondrial respiration) can relieve potential off-target liabilities in off-target tissues. Overall, this study provides novel proof-of-concept for the use of ASOs in restoring the balance of mitochondrial homeostasis in the context of disease and paves the way for future ASO drug development for the increasing number of diseases associated with mitochondrial dysfunction.
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Supplementary Material

Supplementary Data
Supplementary Figure S1
Supplementary Figure S2
Supplementary Figure S3
Supplementary Figure S4
Supplementary Figure S5

References

1. Chan DC. (2020). Mitochondrial dynamics and its involvement in disease. Annu Rev Pathol Mech Dis 15:235–259.
2. Sebastián D, M Palacín and A Zorzano. (2017). Mitochondrial dynamics: coupling mitochondrial fitness with healthy aging. Trends Mol Med 23:201–215.
3. Yu R, S-B Jin, U Lendahl, M Nistér and J Zhao. (2019). Human Fis1 regulates mitochondrial dynamics through inhibition of the fusion machinery. EMBO J 38:e99748.
4. Otera H, C Wang, MM Cleland, K Setoguchi, S Yokota, RJ Youle and K Mihara. (2011). Mff is an essential factor for mitochondrial recruitment of Drp1 during mitochondrial fission in mammalian cells. J Cell Biol 191:1141–1158.
5. Liu R and DC Chan. (2015). The mitochondrial fission receptor Mff selectively recruits oligomerized Drp1. Mol Biol Cell 26:4466–4477.
6. Palmer CS, LD Osellame, D Laine, OS Koutsopoulos, AE Frazier and MT Ryan. (2011). MiD49 and MiD51, new components of the mitochondrial fission machinery. EMBO Rep 12:565–573.
7. Yu R, T Liu, S-B Jin, C Ning, U Lendahl, M Nistér and J Zhao. (2017). MIEF1/2 function as adaptors to recruit Drp1 to mitochondria and regulate the association of Drp1 with Mff. Sci Rep 7:1–16.
8. Otera H, N Miyata, O Kuge and K Mihara. (2016). Drp1-dependent mitochondrial fission via MiD49/51 is essential for apoptotic cristae remodeling. J Cell Biol 212:531–544.
9. Osellame LD, AP Singh, DA Stroud, CS Palmer, D Stojanovski, R Ramachandran and MT Ryan. (2016). Cooperative and independent roles of the Drp1 adaptors Mff, MiD49 and MiD51 in mitochondrial fission. J. Cell Sci 129:2170–2181.
10. Palmer CS, KD Elgass, RG Parton, LD Osellame, D Stojanovski and MT Ryan. (2013). Adaptor proteins MiD49 and MiD51 can act independently of Mff and Fis1 in Drp1 recruitment and are specific for mitochondrial fission. J Biol Chem 288:27584–27593.
11. Kalia R, RY-R Wang, A Yusuf, PV Thomas, DA Agard, JM Shaw and A Frost. (2018). Structural basis of mitochondrial receptor binding and constriction by DRP1. Nature 558:401–405.
12. Archer SL. (2013). Mitochondrial dynamics—mitochondrial fission and fusion in human diseases. N Engl J Med 369:2236–2251.
13. Mattie S, M Krols and HM McBride. (2019). The enigma of an interconnected mitochondrial reticulum: new insights into mitochondrial fusion. Curr Opin Cell Biol 59:159–166.
14. Ota A, T Ishihara and N Ishihara. (2020). Mitochondrial nucleoid morphology and respiratory function are altered in Drp1-deficient HeLa cells. J Biochem (Tokyo) 167:287–294.
15. Rehlkau K, L Hoffmann, CB Gurniak, M Ott, W Witke, L Scorrano, C Culmsee and MB Rust. (2017). Cofilin1-dependent actin dynamics control DRP1-mediated mitochondrial fission. Cell Death Dis 8:e3063.
16. Miret-Casals L, D Sebastián, J Brea, EM Rico-Leo, M Palacín, PM Fernández-Salgueiro, MI Loza, F Albercio and A Zorzano. (2018). Identification of new activators of mitochondrial fusion reveals a link between mitochondrial morphology and pyrimidine metabolism. Cell Chem Biol 25:268–278.e4.
17. Crooke ST, JL Witztum, CF Bennett and BF Baker. (2018). RNA-targeted therapeutics. Cell Metab 27:714–739.
18. Levin AA. (2019). Treating disease at the RNA level with oligonucleotides. N Engl J Med 380:57–70.
19. Aghajan M, SL Booten, M Althage, CE Hart, A Ericsson, I Maxvall, J Ochaba, A Menschik-Lundin, J Hartleib, et al. (2019). Antisense oligonucleotide treatment ameliorates FN-γ-induced proteinuria in APOL1-transgenic mice. JCI Insight 4:e126124.
20. Ochaba J, AF Powers, KA Tremble, S Greenlee, NM Post, JE Matson, AR MacLeod, S Guo and M Aghajan. (2019). A novel and translational role for autophagy in antisense oligonucleotide trafficking and activity. Nucleic Acids Res 47:11284–11303.
21. Raal FJ, RD Santos, DJ Blom, AD Marais, M-J Chang, WC Cromwell, RH Lachmann, D Gaudet, JL Tan, et al. (2010). Mipomersen, an apolipoprotein B synthesis inhibitor, for lowering of LDL cholesterol concentrations in patients with homozygous familial hypercholesterolaemia: a randomised, double-blind, placebo-controlled trial. Lancet Lond Engl 375:998–1006.
22. Mercuri E, BT Darras, CA Chiriboga, JW Day, C Campbell, AM Connolly, ST Iannaccone, J Kirschner, NL Kutz, et al. (2018). Nusinersen versus sham control in later-onset spinal muscular atrophy. N Engl J Med 378:625–635.
23. Ackermann EJ, S Guo, MD Benson, S Booten, S Freier, SG Hughes, T-W Kim, T Jesse Kwoh, J Matson, et al. (2016). Suppressing transthyretin production in mice, monkeys and humans using 2nd-Generation antisense oligonucleotides. Amyloid Int J Exp Clin Investig 23:148–157.
24. Cohn DM, NJ Viney, LM Fijen, E Schneider, VJ Alexander, S Xia, GE Kaeser, C Nanavati, BF Baker, et al. (2020). Antisense inhibition of prekallikrein to control hereditary angioedema. N Engl J Med. [Epub ahead of print]: DOI: 10.1056/NEJMoa1915035.
25. Miller T, M Cudkowicz, PJ Shaw, PM Andersen, N Atassi, RC Bucelli, A Genge, J Glass, S Ladha, et al. (2020). Phase 1–2 trial of antisense oligonucleotide tofersen for SOD1...
ALS, N Engl J Med, [Epub ahead of print]; DOI: 10.1056/NEJMoa2003715.

26. Witztum JL, D Gaudet, SD Freedman, VJ Alexander, A Digienio, KR Williams, Q Yang, SG Hughes, RS Geary, et al. (2019). Volanesorsen and triglyceride levels in familial chylomicronemia syndrome. N Engl J Med 381:531–542.

27. Seth PP, G Vasquez, CA Allerson, A Berdeja, H Gaus, GA Kinberger, TP Prakash, MT Migawa, B Bhat and EE Swayne. (2010). Synthesis and biophysical evaluation of 2′,4′-constrained 2′-methoxyethyl and 2′,4′-constrained 2′-O-ethyl nucleic acid analogues. J Org Chem 75:1569–1581.

28. Koller E, TM Vincent, A Chappell, S De, M Manoharan and CF Bennett. (2011). Mechanisms of single-stranded phosphorothioate modified antisense oligonucleotide accumulation in hepatocytes. Nucleic Acids Res 39:4795–4807.

29. McWilliams TG, AR Prescott, GFG Allen, J Tamjdar, MJ Munson, C Thomson, MMK Muqit and IG Ganley. (2016). mito-QC illuminates mitophagy and mitochondrial architecture in vivo. J. Cell Biol 214:333–345.

30. Chen H, A Chomyn and DC Chan. (2005). Disruption of fusion results in mitochondrial heterogeneity and dysfunction. J Biol Chem 280:26185–26192.

31. Frank S, B Gaume, ES Bergmann-Leitner, WW Leitner, EG Robert, F Catez, CL Smith and RJ Youle. (2001). The role of dynamin-related protein 1, a mediator of mitochondrial fission, in apoptosis. Dev Cell 1:515–525.

32. Jones E, N Gaytan, I Garcia, A Herrera, M Ramos, D Agarwala, M Rana, W Innis-Whitehouse, E Schuenzel and R Gilkerson. (2017). A threshold of transmembrane potential is required for mitochondrial dynamic balance mediated by DRP1 and OMA1. Cell Mol Life Sci 74:1347–1363.

33. Divakaruni AS, A Paradyse, DA Ferrick, AN Murphy and M Jastroch. (2014). Analysis and interpretation of microplate-based oxygen consumption and pH data. In: Methods in Enzymology. Murphy AN, Chan DC, eds. vol. 547. Elsevier, USA, pp 309–354.

34. Twigg G, A Elorza, AJA Molina, H Mohamed, JD Wikstrom, G Walzer, L Stiles, SE Haigh, S Katz, et al. (2008). Fission and selective fusion govern mitochondrial segregation and elimination by autophagy. EMBO J 27:433–446.

35. Shirihai OS, M Song and GW Dorn. (2015). How mitochondrial dynamism orchestrates mitophagy. Circ. Res 116:1835–1849.

36. Palikaras K, E Lionaki and N Tavernarakis. (2018). Mechanisms of mitophagy in cellular homeostasis, physiology and pathology. Nat Cell Biol 20:1013–1022.

37. Pickles S, P Vigié and RJ Youle. (2018). Mitophagy and quality control mechanisms in mitochondrial maintenance. Curr Biol 28:R170–R185.

38. Youle RJ and DP Narendra. (2011). Mechanisms of mitophagy. Nat Rev Mol Cell Biol 12:9–14.

39. Gustafsson ÅB and GW Dorn. (2019). Evolving and expanding the roles of mitophagy as a homeostatic and pathogenic process. Physiol Rev 99:853–892.

40. Mizushima N and T Yoshimori. (2007). How to interpret LC3 immunoblotting. Autophagy 3:542–545.

41. Yamada T, D Murata, Y Adachi, K Itoh, S Kameoka, A Igarashi, T Kato, Y Araki, RL Huganir, et al. (2018). Mitochondrial stasis reveals p62-mediated ubiquitination in Parkin-independent mitophagy and mitigates nonalcoholic fatty liver disease. Cell Metab 28:588–604.e5.

42. Kageyama Y, M Hoshijima, K Seo, D Bedjia, P Sysa-Shah, SA Andrabi, W Chen, A Höke, VL Dawson, et al. (2014). Parkin-independent mitophagy requires Drp1 and maintains the integrity of mammalian heart and brain. EMBO J 33:2798–2813.

43. Marcassa E, A Kalinos, J Jardine, EV Rusilowicz-Jones, A Martinez, S Kuehl, M Islinger, MJ Clague and S Urbé. (2018). Dual role of USP30 in controlling basal pexophagy and mitophagy. EMBO Rep 19:e45595.

44. Züchner S, IV Mersiyanova, M Muglia, N Bissar-Tadmouri, J Rochelle, EL Dadali, M Zappa, E Nèlis, A Patitucci, et al. (2004). Mutations in the mitochondrial GTFase mitofusin 2 cause Charcot-Marie-Tooth neuropathy type 2A. Nat Genet 36:449–451.

45. Detmer SA and DC Chan. (2007). Complementation between mouse Mfn1 and Mfn2 protects mitochondrial fusion defects caused by CMT2A disease mutations. J Cell Biol 176:405–414.

46. Chen H, SA Detmer, AJ Ewald, EE Griffin, SE Fraser and DC Chan. (2003). Mitofusins Mfn1 and Mfn2 coordinately regulate mitochondrial fusion and are essential for embryonic development. J Cell Biol 160:189–200.

47. Liesa M and OS Shirihai. (2013). Mitochondrial dynamics in the regulation of nutrient utilization and energy expenditure. Cell Metab 17:491–506.

48. Rambold AS, B Kostelecky, N Elia and J Lippincott-Schwartz. (2011). Tubular network formation protects mitochondria from autophagosomal degradation during nutrient starvation. Proc Natl Acad Sci U S A 108:10190–10195.

49. Mishra P and DC Chan. (2014). Mitochondrial dynamics and inheritance during cell division, development and disease. Nat Rev Mol Cell Biol 15:634–646.

50. Chan DC. (2020). Mitochondrial dynamics and its involvement in disease. Annu Rev Pathol Mech Dis 15:235–259.

51. Qi X, N Qvit, Y-C Su and D Mochly-Rosen. (2013). A novel Drp1 inhibitor diminishes aberrant mitochondrial fission and neurotoxicity. J Cell Sci 126:789–802.

52. Haileselassie B, AU Joshi, PS Minhas, R Mukherjee, KI Andreasson and D Mochly-Rosen. (2020). Mitochondrial dysfunction mediated through dynamin-related protein 1 (Drp1) propagates impairment in blood brain barrier in septic encephalopathy. J Neuroinflammation 17:36.

53. Mancini NL, L Goudie, W Xu, R Soubany, S Rajeev, A Wang, N Esquerre, A Al Rajabi, TS Jayme, et al. (2020). Perturbed mitochondrial dynamics is a novel feature of colitis that can be targeted to lessen disease. Cell Mol Gastroenterol Hepatol 10:287–307.

54. Mancini NL, S Rajeev, TS Jayme, A Wang, ÁV Keita, ML Workentine, S Hamed, JD Söderholm, F Lopes, et al. (2020). Crohn’s disease pathobiont adherent-invasive e coli disrupts epithelial mitochondrial networks with implications for gut permeability. Cell Mol Gastroenterol Hepatol 11:551–571.

55. Smith G and G Gallo. (2017). To mdivi-1 or not to mdivi-1: is that the question? Dev Neurobiol 77:1260–1268.

56. Rocha AG, A Franco, AM Krezel, JM Runsey, JM Ahlbert, WC Knight, N Biris, E Zacharioudakis, JW Janetka, et al. (2018). MFN2 agonists reverse mitochondrial defects in preclinical models of Charcot-Marie-Tooth disease type 2A. Science 360:336–341.

57. Shen X and DR Corey. (2018). Chemistry, mechanism and clinical status of antisense oligonucleotides and duplex RNAs. Nucleic Acids Res 46:1584–1600.
58. Roberts TC, R Langer and MJA Wood. (2020). Advances in oligonucleotide drug delivery. Nat Rev Drug Discov 19: 673–694.
59. Gomes LC, G Di Benedetto and L Scorrano. (2011). During autophagy mitochondria elongate, are spared from degradation and sustain cell viability. Nat Cell Biol 13:589–598.
60. Serasinghe MN, SY Weider, TT Renault, R Elkholi, JJ Asciolla, JL Yao, O Jabado, K Hoehn, Y Kageyama, et al. (2015). Mitochondrial division is requisite to RAS-induced transformation and targeted by oncogenic MAPK pathway inhibitors. Mol Cell 57:521–536.
61. Ishihara N, Y Eura and K Mihara. (2004). Mitofusin 1 and 2 play distinct roles in mitochondrial fusion reactions via GTPase activity. J Cell Sci 117:6535–6546.
62. Koch A, M Thiemann, M Grabenbauer, Y Yoon, MA McNiven and M Schrader. (2003). Dynamin-like protein 1 is involved in peroxisomal fission. J Biol Chem 278:8597–8605.
63. Kamerkar SC, F Kraus, AJ Sharpe, TJ Pucadyil and MT Ryan. (2018). Dynamin-related protein 1 has membrane constricting and severing abilities sufficient for mitochondrial and peroxisomal fission. Nat Commun 9: 5239.
64. Fonseca TB, Á Sánchez-Guerrero, I Milosevic and N Raimundo. (2019). Mitochondrial fission requires DRP1 but not dynamins. Nature 570:E34–E42.

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