Genome-wide RNAi screen for regulators of UPR\textsuperscript{mt} in \textit{Caenorhabditis elegans} mutants with defects in mitochondrial fusion

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UPR<sup>mt</sup> screen in fzo-1(<i>tm133</i>) mutants (37 characters)

**Keywords:** mitochondrial unfolded protein response, mitochondrial dynamics, fzo-1, IP<sub>3</sub> signaling, Mitoguardin

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ABSTRACT

Mitochondrial dynamics play an important role in mitochondrial quality control and the adaptation of metabolic activity in response to environmental changes. The disruption of mitochondrial dynamics has detrimental consequences for mitochondrial and cellular homeostasis and leads to the activation of the mitochondrial unfolded protein response (UPR\textsuperscript{mt}), a quality control mechanism that adjusts cellular metabolism and restores homeostasis. To identify genes involved in the induction of UPR\textsuperscript{mt} in response to a block in mitochondrial fusion, we performed a genome-wide RNAi screen in Caenorhabditis elegans mutants lacking the gene \textit{fzo-1}, which encodes the ortholog of mammalian Mitofusin, and identified 299 suppressors and 86 enhancers. Approximately 90\% of these 385 genes are conserved in humans, and one third of the conserved genes have been implicated in human disease. Furthermore, many have roles in developmental processes, which suggests that mitochondrial function and the response to stress are defined during development and maintained throughout life. Our dataset primarily contains mitochondrial enhancers and non-mitochondrial suppressors of UPR\textsuperscript{mt}, indicating that the maintenance of mitochondrial homeostasis has evolved as a critical cellular function, which, when disrupted, can be compensated for by many different cellular processes. Analysis of the subsets ‘non-mitochondrial enhancers’ and ‘mitochondrial suppressors’ suggests that organelar contact sites, especially between the ER and mitochondria, are of importance for mitochondrial homeostasis. In addition, we identified several genes involved in IP\textsubscript{3} signaling that modulate UPR\textsuperscript{mt} in \textit{fzo-1} mutants and found a potential link between pre-mRNA splicing and UPR\textsuperscript{mt} activation.

(244/250 words)
INTRODUCTION

Mitochondria are important for cellular adenosine triphosphate (ATP) production, iron-sulfur-cluster biogenesis, lipid metabolism and apoptosis, and therefore, mitochondrial homeostasis is tightly regulated by several quality control mechanisms (Tatsuta and Langer, 2008; Kornmann, 2014). Moreover, mitochondria are required to respond to environmental challenges, which are often accompanied by alterations in energy demand (Youle and van der Bliek, 2012). Mitochondrial dynamics controls mitochondrial shape and distribution, thus playing a central role in both mitochondrial homeostasis and the adjustment to changing energy demands (Yaffe, 1999; van der Bliek et al., 2013). Dynamics of mitochondrial membranes is controlled by large guanosine triphosphate-binding proteins (GTPases) of the dynamin-like family, which are conserved from yeast to humans (Hales and Fuller, 1997; Otsuga et al., 1998; Smirnova et al., 1998; Bleazard et al., 1999; Labrousse et al., 1999; Shepard and Yaffe, 1999; Chen et al., 2003; Santel et al., 2003; Ichishita et al., 2008; Kanazawa et al., 2008). In the nematode Caenorhabditis elegans, fusion of the outer and inner mitochondrial membrane (OMM and IMM) is facilitated by FZO-1MFN1,2 (Ichishita et al., 2008) and EAT-3OPA1 (Kanazawa et al., 2008), respectively. Conversely, fission of the OMM and IMM is carried out by DRP-1DRP1 (Labrousse et al., 1999), whose ortholog in Saccharomyces cerevisiae (Dnm1p) has been shown to form constricting spirals around mitochondria (Ingerman et al., 2005). The disruption of mitochondrial dynamics has detrimental consequences for mitochondrial and ultimately cellular homeostasis and is associated with several human diseases. Thus, mitochondrial homeostasis is controlled by several additional protective quality control mechanisms, including the mitochondrial unfolded protein response (UPRmt) and mitophagy (Chen and Chan, 2004; Youle and van der Bliek, 2012; van der Bliek et al., 2013; Kornmann, 2014). How these quality control mechanisms are coordinated with mitochondrial dynamics is not fully understood. Recently,
disruption of mitochondrial dynamics has been shown to induce UPR$^{mt}$ (Kim and Sieburth, 2018; Zhang et al., 2018; Rolland et al., 2019; Haeussler et al., 2020). UPR$^{mt}$ has been studied extensively in the past decade using genome-wide RNAi screens in *C. elegans* (Haynes et al., 2007; Runkel et al., 2013; Bennett et al., 2014; Liu et al., 2014; Rolland et al., 2019). Upon mitochondrial stress and the concomitant decrease in mitochondrial membrane potential, the master regulator of UPR$^{mt}$, ‘activating transcription factor associated with stress 1’ (ATFS-1$^{ATF4,5}$), instead of being imported into mitochondria, translocates from the cytosol to the nucleus, where it activates a broad transcriptional program (Haynes et al., 2010; Nargund et al., 2012; Rolland et al., 2019). UPR$^{mt}$ activation leads to the expression of a large set of cytoprotective genes including genes encoding chaperones (e.g. hsp-6$^{mtHSP70}$ and hsp-60$^{HSP1}$, whose transcription is commonly used to monitor UPR$^{mt}$ activation (Yoneda et al., 2004)) or proteases, and has been shown to promote mitochondrial biogenesis and coordinate cellular metabolism (Nargund et al., 2012; Rauthan et al., 2013; Liu et al., 2014; Ranji et al., 2014; Nargund et al., 2015; Oks et al., 2018; Haeussler et al., 2020). (All genes that are specifically up- or downregulated upon induction of UPR$^{mt}$ are referred to as UPR$^{mt}$ effectors.) Moreover, UPR$^{mt}$ has been shown to act in a cell non-autonomous way, and once activated in a certain tissue can result in a systemic response (Durieux et al., 2011; Shao et al., 2016; Kim and Sieburth, 2018; Zhang et al., 2018; Kim and Sieburth, 2020).

In this study, we performed a genome-wide RNAi screen to identify regulators of UPR$^{mt}$ in *fzo-1(tm1133)* mutants and identified 299 suppressors and 86 enhancers. We analyzed this dataset using bioinformatic tools, such as GO enrichment analysis, gene network analysis and analysis of transcription factor binding sites in promotors of candidate genes. Furthermore, we determined the specificities of the candidates identified with respect to their ability to modulate UPR$^{mt}$ using secondary screens. Finally, we identified the *C. elegans* ortholog of the mammalian genes *Miga1* and *Miga2*, which have been implicated in mitochondrial fusion,
and demonstrate that the loss of the *C. elegans* ortholog leads to mitochondrial fragmentation and the induction of UPR\textsuperscript{mt}. 
METHODS

General C. elegans methods and strains

C. elegans strains were cultured as previously described (Brenner, 1974). Bristol N2 was used as the wild-type strain. All experiments were carried out at 20°C and all strains were maintained at 20°C. The following alleles and transgenes were used: LGI: spg-7(ad2249) (Zubovych et al., 2010); LGII: fzo-1(tm1133) (National BioResource Project); eat-3(ad426) (Kanazawa et al., 2008); LGIV: drp-1(tm1108) (National BioResource Project); bcSi9 (P_hsp-6::gfp::unc-54 3’UTR) (Haeussler et al., 2020); LGV: miga-1(tm3621) (National BioResource Project). Additionally, the following multi-copy integrated transgenes were used: zcIs9 (P_hsp-6::gfp::unc-54 3’UTR), zcIs13 (P_hsp-6::gfp::unc-54 3’UTR) (Yoneda et al., 2004); bcIs78 (P_myo-3::gfp*) (Rolland et al., 2013).

RNA-mediated interference

RNAi by feeding was performed using the updated ‘Ahringer’ RNAi library (Kamath and Ahringer, 2003), which covers around ~87% of the currently annotated C. elegans protein coding genes. For the primary and secondary screens with the multi-copy zcIs13 transgene in the fzo-1(tm1133), drp-1(tm1108), eat-3(ad426) or spg-7(ad2249) background, RNAi clones were cultured overnight in 100 µL of LB containing carbenicillin (100 µg/mL) in a 96 well plate format at 37°C and 200 rpm. 10 µL of each RNAi culture was used to seed one well of a 24 well RNAi plate containing 0.25% Lactose (w/v) as described previously (Rolland et al., 2019). The plates were incubated at 20°C in the dark. 24 hours later, 3 L4 larvae of all strains carrying the fzo-1(tm1133) and spg-7(ad2249) allele, and 2 L4 larvae of drp-1(tm1108) were transferred to each well of the RNAi plates. The F1 generation was scored by eye for fluorescence intensity of the P_hsp-6_mHSP70gfp reporter after 4-12 days and compared to worms of the respective genotype on the negative control sorb-1(RNAi).
For double-RNAi experiments (Figure S1), RNAi clones were cultured as described above but experiments were conducted in three independent experiments using RNAi plates containing 6 mM IPTG. \( rps-1(\text{RNAi}) \) was diluted 1:1 with either empty vector RNAi (L4440) or \( kgb-I(\text{RNAi}) \).

**Screening procedure and sequencing of RNAi-clones**

For the primary screen, all RNAi clones of the library were tested once. Bacterial RNAi clones that enhanced or suppressed the \( P_{hsp-6 \text{mHSP70}}\text{gfp} \) reporter were picked from the wells and inoculated in 100 μL of LB containing carbenicillin (100 μg/mL) in a 96 well plate format and cultured overnight at 37°C and 200 rpm. Glycerol stocks of these overnight cultures were prepared the following day by adding 100 μL of LB containing 30% glycerol and frozen at -80°C. After all RNAi clones of the library were tested, the 657 identified candidates were re-tested at least three times in duplicates for verification of the observed phenotype. The RNAi clones that reproduced the suppression or enhancement phenotype at least three out of six times were considered as verified candidates.

The 385 verified RNAi clones were sequenced. For this, colony PCRs were performed directly from the glycerol stocks using the primers \( L4440F \) and \( L4440R \). To remove excessive primers and nucleotides, PCR products were treated with ExoSAP-IT™ (Applied Biosystems, Cat.no. 78200.200.UL) according to manufacturer’s protocol. After PCR clean-up, samples were sent for sequencing using \( L4440F \) primer.

\[
\begin{align*}
L4440 F &\ 5’-\text{TGGATAACCGTATTACCGCC-3’} \\
L4440 R &\ 5’-\text{GTTTTCCAGTCACGACGTT-3’}
\end{align*}
\]

According to our sequencing results, seven of the RNAi clones covered two genes. These are indicated in column B (‘Sequence’) in Table S1. These RNAi clones were assigned to the GO
group of the gene, which was predominantly covered by our sequencing result and all subsequent analysis were carried out using this gene.

Subsequently, the verified and sequenced clones were rescreeened in technical duplicates in three independent experiments in the secondary screens in *drp-1(tm1108), eat-3(ad426)* and *spg-7(ad2249)* mutant backgrounds.

**Identification of human orthologs**

Human orthologs and OMIM data (Amberger *et al.*, 2018) were extracted from wormbase.org using https://intermine.wormbase.org (Harris *et al.*, 2019). Human orthologs were then manually verified using ‘alliancegenome.org’ (The Alliance of Genome Resources, 2019), ‘orthodb.org’ (Kriventseva *et al.*, 2018), ‘ensembl.org’ (Hunt *et al.*, 2018) and ‘uniprot.org’ (Consortium, 2018).

**Prediction of mitochondrial localization and mitochondrial targeting sequences**

First, https://intermine.wormbase.org (Harris *et al.*, 2019) was used to identify all candidate genes, which are related to any mitochondrial processes/pathways. To that end, we extracted all 698 genes currently associated with at least one of the 404 GO-terms containing ‘mitochond’ and checked how many of our 385 candidate genes are among them. Additionally, we used the online platform ‘MitoProt’ (https://ihg.gsf.de/ihg/mitoprot.html) (Claros and Vincens, 1996) for computational prediction of mitochondrial targeting sequences. Proteins for which the value of a mitochondrial targeting sequence was ≥0.5 in this analysis were predicted to be mitochondrial.

**Gene ontology enrichment analysis using DAVID**
In search of enriched gene ontology terms, we used the DAVID tool (version 6.8 (Huang et al., 2008, 2009)) and ran the list of candidates against all genes of the C. elegans genome as a background list. Using an EASE score from the modified fisher-exact test, the clustering algorithm groups genes based on their association in GO categories and assigns a significance value to the group (Huang et al., 2007). The clustered groups were then plotted using modified functions from the GO plot package (R version 1.0.2 (Walter et al., 2015)).

**Transcription factor enrichment analysis**

We searched for enriched transcription factors using the tool g:Profiler (a tool for functional enrichment analysis using over-representation (Raudvere et al., 2019)). The two input lists (suppressors and enhancers of fzo-1(tm1133)-induced UPR\textsuperscript{mt}) with WBGene-IDs of the identified candidate genes were used to search in the Transfac database (annotations: TRANSFAC Release 2019.1 classes: v2 (Knüppel et al., 1994; Matys et al., 2006)).

**Construction of gene networks of FZO-1 and MFN1/2, and the UPR\textsuperscript{mt}**

The C. elegans interactomes were compiled for FZO-1 or all 16 genes that are currently associated with the GO-term ‘mitochondrial unfolded protein response’ (GO:0034514) from scientific literature (Durinck et al., 2009; Simonis et al., 2009) and databases such as mentha (Calderone et al., 2013), BioGRID3.5 (Oughtred et al., 2018), IntAct (Orchard et al., 2014) and STRING (Szklarczyk et al., 2018) (STRING was only used to build the FZOome). The human orthologs of those genes were identified and were searched as well. Whenever possible, the interaction partners were converted back to C. elegans genes using biomaRt (Durinck et al., 2009) and available scientific literature (Shaye and Greenwald, 2011; Kim et al., 2018). The complete list of interactions was uploaded to cytoscape (v.3.7.2 (Shannon et
and a network was calculated, highlighting both enhancers and suppressors from the screening results.

**Image acquisition, processing and analysis**

For double-RNAi experiments (Figure S1), 10-20 *fzo-1(tm1133)* mutants carrying *bcSi9* (*P*$_{hsp-6\,\text{mtHSP70}}$*gfp*) were immobilized with M9 buffer containing 10 mM levamisole on 2% agarose pads and imaged using a Nikon SMZ18 dissecting microscope and Nikon-Elements software. For each mutant in Figure S2, 10-20 animals were immobilized with M9 buffer containing 150 mM sodium azide on 2% agarose pads and imaged using a Leica GFP dissecting microscope (M205 FA) and Leica Application Suite software (3.2.0.9652).

For the analysis of mitochondrial morphology a strain carrying *bcIs78* (*P*$_{\text{myo-3}:\text{gfp}^{mt}}$) was imaged using a Zeiss Axioskop 2 with a 63x objective and MetaMorph software (Molecular Devices). Subsequently, mitochondrial morphology was assessed using the deep learning algorithm MitoSegNet (Fischer *et al.*, 2020).
RESULTS & DISCUSSION

Genome-wide RNAi screen for suppressors and enhancers of fzo-1(tm1133)-induced UPR^{mt} identifies highly conserved set of genes with relevance to human health

The disruption of mitochondrial dynamics in C. elegans induces the mitochondrial unfolded protein response (UPR^{mt}) (Kim and Sieburth, 2018; Zhang et al., 2018; Rolland et al., 2019; Haeussler et al., 2020). To identify genes affecting mitochondrial homeostasis in animals with defects in mitochondrial dynamics, we used a loss-of-function mutation of fzo-1^{MFNI.2}, tm1133, (National BioResource Project) to induce the UPR^{mt} reporter P_{hsp-6 mtHSP70}gfp (zcIs13) and screened the C. elegans genome for modifiers. To that end, we used RNA-mediated interference (RNAi) and targeted ~87% of the currently annotated protein coding genes (Kamath and Ahringer, 2003) (Figure 1A). The moderate induction of the P_{hsp-6 mtHSP70}gfp reporter in the fzo-1(tm1133) background allowed the identification of both suppressors and enhancers of the response. Using a protocol in which the F1 generation is scored for P_{hsp-6 mtHSP70}gfp expression levels in the fourth larval stage of development (L4), we initially identified 657 candidate genes of which 385 reproduced. Of the 385 candidates identified, 299 act as suppressors upon knock-down and 86 as enhancers (Figure 1B and Table S1). In addition, upon knock-down, many candidates result in synthetic slow growth and/or reduced fertility (indicated in the ‘Overview’ sheet in Table S1). In order to assess whether the 86 identified enhancers are specific to the fzo-1(tm1133) background or if their depletion induces UPR^{mt} also in the absence of mitochondrial stress, we knocked them down in a wild-type background and tested for induction of the P_{hsp-6 mtHSP70}gfp reporter. All except three genes (copd-1^{ARCN1}, F25H9.6^{PPCDC}, metl-17^{METTL17}) induce P_{hsp-6 mtHSP70}gfp expression when knocked-down in wild-type animals, suggesting that the induction of UPR^{mt} by depletion of these candidates is independent of the loss of fzo-1. (Candidates that encode mitochondrial
proteins and that induce UPR\textsuperscript{mt} in a wild-type background upon knock-down were included in a recent publication, which reported the systematic identification of mitochondrial inducers of UPR\textsuperscript{mt} (Rolland\textit{et al.}, 2019).

Among the 299 suppressors, only 25 (8\%) have previously been found to suppress UPR\textsuperscript{mt} induced by other means upon knock-down (Haynes\textit{et al.}, 2007; Runkel\textit{et al.}, 2013; Liu\textit{et al.}, 2014). Similarly, among the 86 enhancers, only 15 (17\%) have previously been shown to induce UPR\textsuperscript{mt} upon knock-down (indicated ‘Previously identified’ in the ‘Overview’ sheet of Table S1). This may be due to different genetic backgrounds and to differences in RNAi-protocols. Moreover, false negatives in RNAi screens have been estimated to vary between 10\% and 30\%, even if the same protocol is used by the same laboratory (Simmer\textit{et al.}, 2003).

Using ‘alliancegenome.org’ (The Alliance of Genome Resources, 2019), ‘orthodb.org’ (Kriventseva\textit{et al.}, 2018), ‘ensembl.org’ (Hunt\textit{et al.}, 2018), ‘uniprot.org’ (Consortium, 2018) and ‘wormbase.org’ (Harris\textit{et al.}, 2019) databases, we found that approximately 90\% of the suppressors and enhancers (348) have at least one ortholog in humans (indicated ‘Human ortholog’ in the ‘Overview’ sheet of Table S1). For comparison, the overall conservation of genes from \textit{C. elegans} to humans is only about 41\% (Shaye and Greenwald, 2011; Kim\textit{et al.}, 2018). Moreover, we found that the orthologs of 36\% (126) of the conserved candidates have previously been associated with human disease and are listed in the ‘Online Mendelian Inheritance in Man’ database (Amberger\textit{et al.}, 2018) (indicated ‘OMIM’ in the ‘Overview’ and ‘OMIM’ sheet of Table S1). In summary, we identified a set of predominantly conserved genes, many of them relevant to human health, which when knocked-down affect mitochondrial homeostasis in mutants with defects in mitochondrial fusion.
Genes with functions in development, receptor-mediated endocytosis and metabolism modulate UPR\textsuperscript{mt} signaling

In order to obtain an overview of the type of processes that affect fzo-1(tm1133)-induced UPR\textsuperscript{mt}, we analyzed the gene ontology (GO) terms of all 385 candidates, sorted them into ‘functional groups’ (Figure 1B) and performed a clustered gene enrichment analysis using DAVID (Huang \textit{et al.}, 2008, 2009) (Figure 2 and Table S2). (Thirty-one suppressors and enhancers could not be assigned to functional groups since these genes are uncharacterized in \textit{C. elegans} and/or lack orthologs in humans. For this reason, they were assigned to the functional group ‘uncharacterized’ (Figure 1B)).

In the clustered gene enrichment analysis, we found that the majority of both suppressors and enhancers are associated with at least one of the following GO-terms: ‘nematode larval development’, ‘embryo development ending in birth or egg hatching’ or ‘reproduction’ (Table S2). It has been shown that reducing the functions of some genes encoding components of the ETC (e.g. \textit{cox-5B(RNAi)}) in specific tissues and at specific times during development can lead to both systemic activation of UPR\textsuperscript{mt} and longevity (Dillin \textit{et al.}, 2002; Rea \textit{et al.}, 2007; Durieux \textit{et al.}, 2011). This indicates that the activity levels of mitochondria in an individual animal are ‘set’ at a specific developmental stage and, once set, are maintained throughout development and adult life. Our results demonstrate that disrupting development compromises this process, thereby affecting an animal’s ability to cope with mitochondrial stress and to respond to UPR\textsuperscript{mt} activation, which is expected to indirectly affect processes such as its lifespan. In support of this notion, we found that approximately 20% of the suppressors carry the GO-term ‘determination of adult lifespan’ (Table S2).

Among the suppressors, the GO-term ‘receptor-mediated endocytosis’ is enriched (Figure 2A and Table S2). It contains many genes with roles in vesicular trafficking and vesicle budding. Genes required for vesicular trafficking have been shown to affect mitochondrial morphology...
and homeostasis when inactivated, and it has been proposed that this is the result of altered contact sites between organelles and altered lipid transfer into mitochondria (Altmann and Westermann, 2005). Furthermore, we recently demonstrated that approximately half of the candidates in this GO-category are negative regulators of autophagy. Upon knock-down, these genes suppress *fzo-1(tm1133*)-induced UPR$^{\text{mt}}$ most probably by inducing autophagy thereby causing changes in lipid metabolism (Haeussler et al., 2020). Moreover, many cellular signaling pathways originate at the plasma membrane and, thus, are dependent on endocytosis (Sorkin and von Zastrow, 2009; Di Fiore and von Zastrow, 2014). Therefore, we speculate that depletion of the genes associated with the GO-term ‘receptor mediated endocytosis’ may either cause changes in lipid metabolism thereby suppressing UPR$^{\text{mt}}$ or disrupt cell non-autonomous UPR$^{\text{mt}}$ signaling.

The functional group ‘ribosome biogenesis’ contains 78 (26%) of the suppressors (Figure 1B) and includes both small- and large ribosomal subunits, as well as proteins with roles in the maturation or transport of ribosomal subunits and rRNAs. Accordingly, in all three GO-domains (Biological Process, Cellular Compartment, Molecular Function), we found that several GO-terms related to the ribosome were significantly enriched (Figure 2A and Table S2). (The GO-term ‘apoptotic process’ also contains many ribosomal subunits leading to its enrichment in our analysis.)

Moreover, we assigned a substantial part of the suppressors to the groups ‘RNA processing’ (38), ‘transcription’ (35) and ‘translation’ (27) (Figure 1B). Hence, we found five GO-terms related to translation-, two to transcription- and one to RNA-related processes to be enriched in a statistically significant manner in the GO enrichment analysis (Figure 2A and Table S2). These results raise the question whether knock-down of the candidates involved in cytosolic translation specifically suppresses UPR$^{\text{mt}}$ or simply reduces the expression of the P$_{\text{hsp-6mtHSP70gfp}}$ reporter. A previous study also identified many genes related to ribosome...
biogenesis and cytosolic translation in a screen for suppressors of paraquat-induced UPR\textsuperscript{mt} (Runkel \textit{et al.}, 2013). Runkel and colleagues reported reduced levels of two other reporters (P\textsubscript{hsp-16.2 CRYAB}\textsubscript{gfp}, P\textsubscript{hsp-4 HSPA5}\textsubscript{gfp}) upon attenuation of cytosolic translation by \textit{rpl-36}(RNAi). In contrast, they showed that P\textsubscript{gst-4 HPGDS}\textsubscript{gfp} was slightly hyperactivated (Runkel \textit{et al.}, 2013), as previously shown for this reporter upon knock-down of several other genes related to cytosolic translation (Melo and Ruvkun, 2012). We recently showed that knock-down of the cytosolic tRNA synthetase \textit{hars-1}\textsuperscript{HARS1}, which we found to suppress P\textsubscript{hsp-6} \textit{mHSP70}\textsubscript{gfp} expression in \textit{fzo-1}(\textit{tm1133}) and which presumably also compromises cytosolic translation, results in reduced expression of a control reporter that is unrelated to other stress responses, P\textsubscript{ges-1 \textit{GES2}}\textsubscript{gfp} (Haeussler \textit{et al.}, 2020). Taken together, we cannot exclude the possibility that the knock-down of candidates related to the functional groups of transcription, RNA processing, ribosome biogenesis and translation may, at least to some extent, interfere with reporter expression \textit{per se}. In addition, Runkel and colleagues showed that depletion of \textit{kgb-1}\textsuperscript{MAPK10}, a JNK-like MAP-kinase mediating cellular surveillance-activated detoxification and defenses (cSADDs) in \textit{C. elegans} (Melo and Ruvkun, 2012), derepresses UPR\textsuperscript{mt} induced by paraquat upon attenuation of cytosolic translation (Runkel \textit{et al.}, 2013). Therefore, we tested whether knock-down of \textit{kgb-1} also relieves the induction of UPR\textsuperscript{mt} upon knock-down of \textit{rps-1} and found that P\textsubscript{hsp-6} \textit{mHSP70}\textsubscript{gfp} expression was partially restored under these conditions (Figure S1). Thus, attenuation of cytosolic translation may activate cSADDs through \textit{kgb-1}\textsuperscript{MAPK10}, thereby preventing UPR\textsuperscript{mt} induction in \textit{fzo-1}(\textit{tm1133}) mutants.

Among the enhancers, we assigned most candidates to the functional groups ‘metabolism’ and ‘mitochondrial ribosome biogenesis’ as well as ‘cellular trafficking’, ‘mitochondrial translation’ and ‘ETC assembly’ (Figure 1B). Accordingly, GO analysis of the enhancers shows that the cellular compartments ‘mitochrondrion’, ‘mitochondrial small ribosomal subunit’, ‘mitochondrial large ribosomal subunit’, ‘mitochondrial inner membrane’,
‘mitochondrial matrix’ and ‘ribosome’ are enriched (Figure 2B and Table S2). In addition, the biological processes ‘translation’ (which also includes ‘mitochondrial translation’), ‘tricarboxylic acid cycle’ and ‘receptor-mediated endocytosis’ are enriched as is the molecular function ‘structural constituent of ribosome’ (Figure 2B and Table S2). Among the enhancers carrying the GO-term ‘receptor-mediated endocytosis’, we identified many subunits of the mitochondrial ribosome and genes required for mitochondrial translation, which are most likely misannotated and therefore led to enrichment of this GO-term. In summary, we showed that disrupting mitochondrial translation and metabolism induces UPR\textsuperscript{mt} in \textit{fzo-1(tm1133)}. Disruption of these processes has also previously been shown to induce UPR\textsuperscript{mt} in wild type (Durieux \textit{et al.}, 2011; Houtkooper \textit{et al.}, 2013). Therefore, we conclude that reducing mitochondrial function induces UPR\textsuperscript{mt} independently of the genetic background.

In summary, the GO enrichment analysis revealed that depletion of the majority of candidates in our dataset may modulate UPR\textsuperscript{mt} due to their role in development. Furthermore, we propose that the suppressors with roles in endocytosis modulate UPR\textsuperscript{mt} signaling indirectly and speculate that cellular signaling and/or alterations in organelar contact sites may influence mitochondrial metabolism and hence, UPR\textsuperscript{mt} signaling. Finally, we find disruption of mitochondrial metabolism and translation to robustly enhance UPR\textsuperscript{mt} signaling in \textit{fzo-1(tm1133)}.

\textbf{Mitochondrial fitness balances cellular homeostasis}

Next, we determined which fraction of the identified enhancers and suppressors encode proteins that have a mitochondrial function or localize to mitochondria. We extracted all 698 genes that are associated with at least one of the 404 GO-terms containing ‘mitochond’ using the ‘WormMine’ database (https://intermine.wormbase.org) (Harris \textit{et al.}, 2019), and then determined how many of our candidate genes are associated with any of these GO-terms.
Using this approach, we identified 11 suppressors and 59 enhancers that encode proteins that localize to mitochondria or play a role in mitochondrial metabolism and dynamics, respectively (indicated ‘GO mitochond’ in ‘Overview’ and ‘Mitochondrial’ sheet of Table S1). Next, we used the online platform ‘MitoProt’ (https://ihg.gsf.de/ihg/mitoprot.html) (Claros and Vincens, 1996) for computational prediction of mitochondrial targeting sequences and identified an additional 5 suppressors and 14 enhancers that are predicted to localize to mitochondria (cut-off value ≥ 0.5) (indicated ‘MitoProt prediction’ in ‘Mitochondrial’ sheet of Table S1). Third, by literature searches, we found that the orthologs of 3 enhancers localize to mitochondria (Shafqat et al., 2003; Spaan et al., 2005; Cambier et al., 2012). In summary, 76 out of 86 (88%) enhancers and 16 out of 299 (5%) suppressors encode proteins that have a mitochondrial function. This suggests that only few processes exist outside of mitochondria that can perturb mitochondrial homeostasis when compromised. Conversely, many processes and mechanisms exist outside of mitochondria that can compensate for mitochondrial dysfunction, thereby ensuring mitochondrial and consequently cellular homeostasis.

Among the 10 ‘non-mitochondrial’ enhancers of UPR\textsuperscript{mt} are three genes (\textit{F29B9.8, Y61A9LA.11, C25H3.10}) with yet unknown functions, which lack orthologs in other systems. \textit{ORC-1\textsuperscript{ORC1}} is a component of the origin recognition complex and plays a role in DNA replication (Gavin et al., 1995; Ohta et al., 2003; Tatsumi et al., 2003). The disruption of DNA replication or cell cycle progression has previously not been reported to lead to UPR\textsuperscript{mt} induction. We speculate that disruption of DNA replication leads to developmental defects and therefore induces UPR\textsuperscript{mt}. \textit{F25H9.6\textsuperscript{PPCDC}} is the \textit{C. elegans} ortholog of phosphopantothenoylcysteine decarboxylase, an enzyme required for biosynthesis of coenzyme A (CoA) (Daugherty et al., 2002). Thus, knock-down of \textit{F25H9.6\textsuperscript{PPCDC}} may interfere with critical biosynthetic and metabolic pathways (including the TCA cycle) and therefore enhance UPR\textsuperscript{mt}. \textit{NHR-209\textsuperscript{HNF4A,G}} is orthologous to Hepatocyte Nuclear Factor 4α.
(HNF4A) and belongs to the family of nuclear hormone receptors, a class of cofactor and ligand-inducible transcription factors (TFs) that regulate various cellular processes, including metabolism, development and homeostasis (Aranda and Pascual, 2001; Bolotin et al., 2010).

Interestingly, long-chain fatty acids are ligands of HNF4A and, depending on their chain length and degree of saturation, activate or repress the transcriptional activity of HNF4A (Hertz et al., 1998; Dhe-Paganon et al., 2002; Wisely et al., 2002; Duda et al., 2004). Furthermore, HNF4α activity has been shown to be required for β-oxidation of fatty acids both in mice and Drosophila melanogaster (Palanker et al., 2009; Chen et al., 2020). Thus, NHR-209HNF4A,G may have a similar role in C. elegans and act as a metabolic sensor, which when deactivated, enhances UPR^mt in fzo-1(tm1133). Moreover, we identified cpa-3^{CPNE5,8,9}, an ortholog of mammalian copine family members, a class of calcium dependent phospholipid binding proteins with roles in intracellular signaling and membrane trafficking (Creutz et al., 1998; Tomsig et al., 2003; Tomsig et al., 2004; Ramsey et al., 2008). Previously, another gene of the copine family, gem-4^{CPNE8}, has been shown to be upregulated upon UPR^mt activation (Nargund et al., 2012). Therefore, we speculate that signaling via copine family members may be important for UPR^mt regulation. Another non-mitochondrial enhancer, copd-1^{ARCNI}, encodes a protein orthologous to the delta subunit of coatamer in S. cerevisiae and humans (RET2 and ARCN1, respectively), which is involved in the formation of coat protein complex I (COPI) vesicles. COPI vesicles play a central role in the secretory pathway and are required for the retrieval of lipids and proteins from the Golgi apparatus and the subsequent retrograde transport of these lipids and proteins to the ER (Lee et al., 2004; Beck et al., 2009). Furthermore, the trafficking to their final destination of most non-mitochondrial and non-peroxisomal transmembrane proteins, as well as proteins required for the release of neurotransmitters, such as SNARE proteins, is dependent on COPI-mediated transport (Beck et al., 2009). Thus, disruption of the secretory pathway affects many intra- and intercellular signaling pathways, including the Ras and TOR signaling pathways, as well as signaling via
G-protein-coupled receptors (GPCRs) and receptor tyrosine kinases (Farhan and Rabouille, 2011). Moreover, disruption of the retrograde transport system has been shown to lead to erroneous secretion of ER resident proteins (e.g. ER chaperones) and, consequently, to the activation of UPR in the ER (UPRER) (Aguilera-Romero et al., 2008; Izumi et al., 2016). Therefore, we speculate that the enhancement of UPRmt induction in fzo-1(tm1133) animals upon copd-1(RNAi) may be due to alterations in one of the above-mentioned signaling pathways. This notion is supported by the finding that phospholipase C (PLC-1PLCE1), a GPCR associated enzyme, is among the non-mitochondrial enhancers, as well as srh-40 (serpentine receptor class H), which is predicted to encode a GPCR. Taken together, we identified many genes among the ‘non-mitochondrial’ enhancers, which regulate intra- and intercellular signaling cascades, and we speculate that these may play a role in signaling of UPRmt, both in a cell autonomous and cell non-autonomous fashion. In addition, we identified ‘non-mitochondrial’ enhancers that directly regulate metabolic homeostasis and, thus, enhance UPRmt in fzo-1(tm1133) mutants.

Among the 16 identified ‘mitochondrial suppressors’ of UPRmt are candidates, such as TFG-1TFG and GBF-1GBF1, that encode proteins that have been shown to associate with mitochondria but also other organelles. GBF-1GBF1 is a guanine nucleotide exchange factor (GEF) for the small GTPase ARF-1.2ARF1, which in yeast recruits ARF-1.2ARF1,3 to ER-mitochondria contact sites (Ackema et al., 2014). Depletion of GBF-1GBF1 leads to altered ARF-1.2ARF1,3 localization and changes in mitochondrial morphology both in yeast and C. elegans and this appears to be independent of their roles in endosomal transport (Ackema et al., 2014). Ackema and colleagues observed an increase in mitochondrial connectivity upon GBF-1GBF1 depletion, similar to that observed upon knock-down of miro-1MIRO1 and vdac-1VDAC, both of which encode proteins that also localize to ER-mitochondria contact sites. However, the alterations in mitochondrial morphology of FZO-1MFN1,2 depleted animals were
shown to be epistatic to the changes in mitochondrial morphology observed upon gbf-1(RNAi) and arf-1.2(RNAi). Therefore, the suppression of UPR\textsuperscript{mt} observed in fzo-1(tm1133) animals upon gbf-1(RNAi) may not be due to a rescue of the mitochondrial morphology defect but rather be the consequence of changes in ER-mitochondria contact sites. This highlights the importance of organelar contact sites for the maintenance of mitochondrial and consequently cellular homeostasis. Furthermore, we identified TFG-1\textsuperscript{TFG}, a component of the secretory pathway via COPII vesicles (Witte \textit{et al.}, 2011), as a suppressor of fzo-1(tm1133)-induced UPR\textsuperscript{mt}. COPII vesicles transport newly synthesized proteins and lipids from specialized ER zones, so called ER exit sites (ERES), to the Golgi apparatus (Budnik and Stephens, 2009; Kurokawa and Nakano, 2018). Similar to what we propose for copd-1(RNAi) (see above), we speculate that disruption of the secretory pathway may lead to alterations in cellular signaling, ER-mitochondria contact sites and, depending on the context, either to suppression or enhancement of UPR\textsuperscript{mt}. Taken together, we demonstrate that the perturbation of primarily mitochondrial processes leads to the enhancement of UPR\textsuperscript{mt}. However, the identification of non-mitochondrial enhancers demonstrates that disruption of processes taking place outside of mitochondria can also compromise mitochondrial function and activate or enhance UPR\textsuperscript{mt}. Alterations in cellular signaling pathways and/or organelar contact sites may play a role in this respect. Moreover, we find that the majority of suppressors of fzo-1(tm1133)-induced UPR\textsuperscript{mt} are non-mitochondrial, suggesting that many cellular pathways outside of mitochondria exist that can compensate for mitochondrial stress and, hence, ensure mitochondrial homeostasis. In line with this notion, we identified a few ‘mitochondrial suppressors’, most of which are involved in the maintenance of contacts to other organelles, especially the ER.
Defects in mitochondrial fusion and fission are suppressed and enhanced by the same pathways

In order to define the specificity of the 299 suppressors and 86 enhancers, we carried out secondary screens. To identify general modifiers of UPR\textsuperscript{mt}, we rescreened the candidates in the background of spg-7(ad2249), which induces UPR\textsuperscript{mt} (Figure S2). spg-7\textsuperscript{AFG3L2} encodes a mitochondrial matrix AAA-protease, which induces UPR\textsuperscript{mt} when depleted and which is commonly used as a positive control for UPR\textsuperscript{mt} activation (Yoneda et al., 2004; Haynes et al., 2007; Haynes et al., 2010). To identify genes in our dataset that specifically modify UPR\textsuperscript{mt} induced by defects in mitochondrial membrane fusion, we rescreened all candidates in the eat-3(ad426) background, in which IMM fusion is blocked. Finally, to identify genes that may modulate UPR\textsuperscript{mt} induced by defects in mitochondrial dynamics, we rescreened all candidates in the drp-1(tm1108) background, in which mitochondrial fission is blocked. In the drp-1(tm1108) background, of the 385 candidates, 291 suppress and 59 enhance. In the eat-3(ad426) background, 242 suppress and, 49 enhance. Finally, in the spg-7(ad2249) background, 181 suppress and 54 enhance (Table S1). (Of note, there is an inverse correlation between the level of P\textsubscript{hsp-6} \textsubscript{mt}HSP70\textsubscript{gfp} expression in the above-mentioned mutant background and the number of candidates that reproduce. Hence, the level of reporter expression may correlate with the number of false negatives in a given dataset of the secondary screens, for both suppressors and enhancers.) Since more suppressors reproduced in drp-1(tm1108) and eat-3(ad426) compared to spg-7(ad2249), we conclude that defects in mitochondrial dynamics, to some extent, are suppressed or enhanced by the same pathways. Moreover, the suppressors of fzo-1(tm1133)-induced UPR\textsuperscript{mt} that were sorted into the functional groups ‘ribosome biogenesis’, ‘RNA processing’ and ‘translation’, reproduced comparably well in all secondary screens. Thus, attenuation of cytosolic translation may either be a general mechanism to suppress UPR\textsuperscript{mt} or, as discussed above, interfere with reporter expression.
Among the enhancers, genes that sorted into the functional groups ‘ETC assembly factors’, ‘mitochondrial ribosome biogenesis’ and ‘mitochondrial translation’ showed the highest overlap among the secondary screens (Table S1), which demonstrates that disruption of mitochondrial translation robustly enhances UPR\textsuperscript{mt}, independent of genetic background.

While we did not identify any suppressors that act exclusively in the fzo-1(tm1133) background, we found six enhancers (slc-25A26\textsuperscript{SLC25A26}, frh-1\textsuperscript{FXN}, sdha-1\textsuperscript{SDHA}, sucg-1\textsuperscript{SUCLG2}, metl-17\textsuperscript{METTL17}, K03B4.1) that did not reproduce in any of the secondary screens. Among these, metl-17\textsuperscript{METTL17}, a methyltransferase required for mitochondrial ribosome assembly and mitochondrial translation in mice (Shi \textit{et al.}, 2019), also did not induce UPR\textsuperscript{mt} expression in wild type and, thus, specifically enhances fzo-1(tm1133)-induced UPR\textsuperscript{mt}.

Twelve candidates that suppressed UPR\textsuperscript{mt} in the primary screen using fzo-1(tm1133), enhanced UPR\textsuperscript{mt} in one or more of the secondary screens. Conversely, ten enhancers of fzo-1(tm1133)-induced UPR\textsuperscript{mt} suppress UPR\textsuperscript{mt} in at least one of the mutants in the secondary screens (listed in the ‘Opposing UPR\textsuperscript{mt} phenotypes’ sheet in Table S1). For example, knockdown of icd-1\textsuperscript{βNAC} suppresses \textit{phsp-6} \textit{mHSP70gfp} in all mitochondrial dynamics-related backgrounds, but enhances \textit{spg-7(ad22449)}-induced UPR\textsuperscript{mt}. Knock-down of \textit{icd-1}\textsuperscript{βNAC} in \textit{C. elegans} has been reported to induce UPR\textsuperscript{ER} in wild-type embryos (Arsenovic \textit{et al.}, 2012). Furthermore, \textit{icd-1}\textsuperscript{βNAC} has been described as a cytosolic stress sensor, which in the absence of stress associates with ribosomes to promote cytosolic translation, and which acts as a chaperone in the cytosol upon heat stress (Kirstein-Miles \textit{et al.}, 2013). We recently showed that \textit{icd-1}\textsuperscript{βNAC} is a negative regulator of autophagy and that increased autophagic flux fuels mitochondria with certain triacylglycerols, thereby suppressing UPR\textsuperscript{mt} in fzo-1(tm1133) and drp-1(tm1108) mutants (Haeussler \textit{et al.}, 2020). Thus, blocking mitochondrial dynamics may reduce the flux of lipids into mitochondria, which can be compensated for by the induction of autophagy and we speculate that this mechanism may also apply to \textit{eat-3(ad426)} mutants.
Conversely, we speculate that defects in mitochondrial homeostasis induced by a point mutation in spg-7, may exert stress to the cytosol and that this is normally compensated for by factors, such as icd-1\textsuperscript{BNAC}. Knocking-down icd-1\textsuperscript{BNAC} may therefore increase cytosolic stress, which in turn enhances UPR\textsuperscript{mt} in spg-7(ad2249) mutants. Taking the candidates into account that have opposing UPR\textsuperscript{mt} phenotypes in the secondary screens, 95\% of the suppressors and 66\% of the enhancers reproduce in drp-1(tm1108), while 79\% of the suppressors and 57\% of the enhancers reproduce in eat-3(ad426). We found the lowest overlap of candidate genes in spg-7(ad2249) mutants, with 59\% of the suppressors and 60\% of the enhancers reproducing in this background. Taken together, the results of the secondary screens show that there are candidates that, when depleted, act to influence UPR\textsuperscript{mt} signaling in general whereas others are specific to a certain type of UPR\textsuperscript{mt} induction, such as the disruption of mitochondrial dynamics.

Transcription factor enrichment analysis identifies factors with roles in development, metabolism and oxidative stress response

Next, we identified TF binding sites in the promoters of our candidates using ChIP-seq datasets from the modENCODE project (Celniker et al., 2009) in order to test for enrichment of TFs that bind to these sites. To that end, we used g:Profiler, a tool for functional enrichment analysis using over-representation (Raudvere et al., 2019), which utilizes TRANSFAC resources (Knüppel et al., 1994; Matys et al., 2006). Using this approach, we found 15 TFs to be enriched in a statistically significant manner (Figure 3 and Table S3). Ten of these TFs only bind promotor regions of suppressors (7) or enhancers (3) (‘suppressor- or enhancer specific’). The remaining five TFs bind to promotor regions of both suppressors and enhancers (‘shared’). The ‘shared’ TFs have previously been implicated in cell fate determination or developmental timing. Five out of seven ‘suppressor specific’ TFs have been
shown to exclusively control developmental processes. The remaining two ‘suppressor-specific’ TFs are \( \text{ELT-3}^{\text{GATA3,4}} \) and \( \text{HLH-11}^{\text{TFAP4}} \), which have been shown to play a role in development, ageing and the response to oxidative stress (Gilleard et al., 1999; Budovskaya et al., 2008; Hu et al., 2017) and to act as a dietary sensor that regulates metabolic gene expression, respectively (Soo-Ung et al., 2009; Watson et al., 2013).

Three TFs (SKN-1\(^{\text{NFE2,NFE2L1,2,3}}\), HLH-29 and VAB-7\(^{\text{EVX2}}\)) were identified to be ‘enhancer-specific’ (Figure 3 and Table S3). VAB-7\(^{\text{EVX2}}\) and HLH-29 are both required for certain aspects of development (Ahringer, 1996; Esmaeili et al., 2002; Pocock et al., 2004; Neves and Priess, 2005; McMiller et al., 2007; Grove et al., 2009) and HLH-29 has additional roles in fatty acid metabolism and energy homeostasis (McMiller et al., 2007; Quach et al., 2013). Furthermore, HLH-29 and SKN-1\(^{\text{NFE2,NFE2L1,2,3}}\) are regulators of the oxidative stress response (An and Blackwell, 2003; An et al., 2005; Inoue et al., 2005; Quach et al., 2013) and SKN-1\(^{\text{NFE2,NFE2L1,2,3}}\) has previously been implicated in the UPR\(^{\text{mt}}\) pathway in \textit{C. elegans} (Nargund et al., 2012; Nargund et al., 2015; Wu et al., 2018). In summary, we identified several TFs that bind to promoters of our candidate genes, which have previously been implicated in oxidative stress response, cellular metabolism and development in \textit{C. elegans}. Interestingly, \textit{fzo-1(tm1133)} mutants have previously been shown to be slightly sensitive to oxidative stress and have increased levels of carbonylated proteins, a measure for oxidative damage (Yasuda et al., 2011). Moreover, in \textit{isp-1(qm150)} and \textit{clk-1(qm30)} mutants, both of which have increased levels of reactive oxygen species (ROS) (Van Raamsdonk et al., 2010; Yang and Hekimi, 2010; Dues et al., 2017), UPR\(^{\text{mt}}\) activation has been shown to lead to ATFS-1\(^{\text{ATF4,5}}\)-dependent expression of genes required for detoxification of reactive oxygen species (Wu et al., 2018). This induction is orchestrated by ATFS-1\(^{\text{ATF4,5}}\) but may, to some extent, additionally be facilitated through activation of ELT-3\(^{\text{GATA3,4}}\) and HLH-29, as it has previously been shown for SKN-1\(^{\text{NFE2,NFE2L1,2,3}}\) (Nargund et al., 2012; Nargund et al., 2015;
The identification of many TFs controlling developmental processes is in agreement with our finding that GO-terms related to developmental processes are enriched among our dataset. This again highlights that the activity levels of critical cellular processes and responses in somatic tissues appear to be set during development. Finally, we previously found that the induction of autophagy suppresses UPR\textsuperscript{mt} in fzo-1(tm1133) mutants most likely through increased metabolic activity (Haeussler \textit{et al.}, 2020). In our analysis, we identified two TFs, which regulate energy homeostasis and metabolic gene expression. This supports the notion that UPR\textsuperscript{mt} in fzo-1(tm1133) mutants acts to compensate for metabolic defects. In summary, we identified several TFs with roles in development, oxidative stress response and metabolism that previously have not been connected to UPR\textsuperscript{mt} signaling. These TFs may be specific to UPR\textsuperscript{mt} in fzo-1(tm1133) but some may generally be involved in UPR\textsuperscript{mt} signaling.

**Interactome of UPR\textsuperscript{mt} reveals potential new regulators**

In order to determine whether any of the suppressors or enhancers that we identified have previously been shown to interact with fzo-1\textsuperscript{MFN1,2} or its mammalian orthologs \textit{MFN1} or \textit{MFN2}, we built a gene network containing all known interactions of fzo-1\textsuperscript{MFN1,2} and its mammalian orthologs \textit{MFN1} and \textit{MFN2}. Using the interaction databases ‘string-db.org’, ‘IntAct’, ‘BioGRID3.5’, ‘Genemania’, ‘CCSB’ and ‘mentha’ (Warde-Farley \textit{et al.}, 2010; Calderone \textit{et al.}, 2013; Orchard \textit{et al.}, 2014; Rolland \textit{et al.}, 2014; Oughtred \textit{et al.}, 2018; Szklarczyk \textit{et al.}, 2018), we included genetic and physical interactions (but not predicted interactions or co-expression data) and uploaded them to the cytoscape software (Shannon \textit{et al.}, 2003) to calculate a complete interaction network. The resulting network contains 38 genes and 67 interactions (Figure S3). None of the 10 interactors of fzo-1\textsuperscript{MFN1,2} in \textit{C. elegans} was identified in our screen (turquoise dots in Figure S3). Next, we manually annotated the \textit{C. elegans} orthologs of 24 interactors of \textit{Mfn1} or \textit{Mfn2} in mammals (except FAF2, MAVS,
TCHP, SLC25A38 for which we did not find any orthologs in C. elegans, indicated in dark blue in Figure S3) but again did not find any overlap between the gene network and our screen dataset (orange dots in Figure S3). In summary, in our screen for modifiers of fzo-1(tm1133)-induced UPR\textsuperscript{mt}, we did not find any previously known interactors of fzo-1\textsuperscript{MFN1,2}. These could either have been missed in the RNAi screen, be essential in the fzo-1(tm1133) background or not have a function in mitochondrial homeostasis and, hence, UPR\textsuperscript{mt} signaling.

Similar to the approach described above, we used the 16 C. elegans genes currently associated with the GO-term ‘mitochondrial unfolded protein response’ (GO:0034514) (referred to as ‘input genes’), identified their human orthologs and included known physical and genetic interactors from the interaction databases ‘BioGRID3.5’, ‘IntAct’ and ‘mentha’ (Calderone et al., 2013; Orchard et al., 2014; Oughtred et al., 2018) to calculate an interaction network containing 2603 genes and 4655 interactions (Figures S4, Figure S5, Figure S6). In this ‘UPR\textsuperscript{mt}ome’, we identified 129 genes (including the 16 ‘input genes’), 36 of which are enhancers and 77 of which are suppressors of fzo-1(tm1133)-induced UPR\textsuperscript{mt}, with a total of 213 interactions (Figure 4 and Table S4). For the ‘input gene’ atfs-1\textsuperscript{ATF4,5}, we found five interactors (gtf-2\textsuperscript{GTF2F2}, lin-54\textsuperscript{LIN54}, rps-6\textsuperscript{RPS6}, spr-2\textsuperscript{SET}, tbp-1\textsuperscript{TBP}) that suppress fzo-1(tm1133)-induced UPR\textsuperscript{mt} and the gene products of four of these localize to the nucleus (Sopta et al., 1989; Lichtsteiner and Tjian, 1993; Wen et al., 2000; Thomas et al., 2003; Harrison et al., 2006; Tabuchi et al., 2011). These could potentially facilitate or directly be involved in the transcription of UPR\textsuperscript{mt} effectors upon activation of the UPR\textsuperscript{mt} response.

Moreover, for the ‘input gene’ ubl-5\textsuperscript{UBL5}, we found four interactors that overlap with our dataset of suppressors, three of which are splicing factors (pqbp-1.2\textsuperscript{PQBP1}, sfa-1\textsuperscript{SF1}, snr-3\textsuperscript{SNRPD1}) (Thomas et al., 1988; Krämer, 1992; Arning et al., 1996; Imafuku et al., 1998; Kambach et al., 1999; Mazroui et al., 1999; Waragai et al., 1999). Of note, HUB1, the ortholog of UBL-5\textsuperscript{UBL5} in Saccharomyces pombe, has been shown to interact with
components of the spliceosome. Furthermore, the loss of HUB1 results in reduced splicing efficiency of a variety of mRNAs (Wilkinson et al., 2004). However, in C. elegans, ubl-5(RNAi) has previously been reported to not cause splicing defects (Haynes et al., 2007). Thus, the identification of the splicing factor genes pqbp-1.2PQP1, sfa-1SF1, snr-5SNRPD1 in our dataset presents an interesting potential link between UPRmt activation and pre-mRNA splicing via UBL-5UBL5. In addition, we identified taf-4TAF4, which encodes an associated factor of transcription factor TFIID, to interact with the ‘input gene’ sphk-1SHPK1,2 and to suppress fzo-1(tm1133)-induced UPRmt upon knock-down. taf-4TAF4 has previously been shown to be required for life span extension in isp-1(qm150), clk-1(qm30) and tpk-1(qm162) mutants, (Walker et al., 2001; Walker et al., 2004; Khan et al., 2013). Finally, we identified many genes interacting with the ‘input gene’ bar-1JUP,CTN1, which has previously been shown to be involved in cell non-autonomous propagation of UPRmt signaling (Zhang et al., 2018). Among these interactors is phospholipase C (plc-1PLCE), which enhances fzo-1(tm1133)-induced UPRmt and plays a central role in the inositol triphosphate (IP3) signaling pathway (Clandinin et al., 1998; Kariya et al., 2004). In summary, we identified several genes in our dataset using gene network analysis that have previously not been identified to play a role in UPRmt signaling in C. elegans. The genes with roles in pre-mRNA splicing and IP3 signaling may be particularly interesting in this respect. Furthermore, we propose that these genes may directly influence UPRmt signaling through interactions with known players of the UPRmt pathway.

**Interactome analysis reveals involvement of IP3 signaling pathway in UPRmt regulation in fzo-1(tm1133)**

In our gene network analysis, we identified plc-1PLCE, which encodes phospholipase C, as an interactor of bar-1β-catenin (Byrne et al., 2007). Interestingly, we and others found several genes
that play a role in inositol triphosphate (IP₃) signaling (Figure 5) (Liu et al., 2014). The IP₃ pathway is well known for its role in the regulation of intracellular calcium levels and transmits signals from the extracellular space via GPCRs and second messengers to the ER (Berridge, 2009). Thus, this signaling pathway may have a role in cell non-autonomous propagation of UPR^mt. We identified the enzyme CDGS-1^{CDS1}, which is essential for the production of phosphatidylinositol (PI) (Wu et al., 1995; Vance, 1998), and EFR-3^{EFR3B}, which targets PI-4-kinase (PI4K) to the plasma membrane (Nakatsu et al., 2012). Furthermore, we identified the sole type I PIP kinase in C. elegans, PPK-1^{PPSK1A} (Weinkove et al., 2008), which phosphorylates PI4P to form PI(4,5)P₂ (Ishihara et al., 1996; Loijens and Anderson, 1996). PLC-1^{PLCE} is activated via GPCR and hydrolyzes PI(4,5)P₂ to generate the second messengers DAG and IP₃, known regulators of several signal transduction pathways (Clandinin et al., 1998; Kariya et al., 2004). One mechanism that is dependent on IP₃-signaling is the release of calcium from the ER (Clandinin et al., 1998; Kariya et al., 2004; Kovacevic et al., 2013). Interestingly, the IP₃ receptor at the ER, ITR-1^{ITPR1}, has previously also been identified as a suppressor of antimycin-induced UPR^mt (Liu et al., 2014). Thus, it is tempting to speculate that altering IP₃ signaling influences cellular calcium signaling in fzo-1(tm1133), thereby affecting mitochondrial homeostasis and consequently UPR^mt signaling. Moreover, we propose that the effect on UPR^mt signaling may be indirect since we previously showed that knock-down of mitochondrial genes controlling calcium homeostasis does not induce UPR^mt in wild type (Rolland et al., 2019). Furthermore, we propose that fzo-(tm1133) mutants may be more prone to changes in IP₃ signaling and, consequently, calcium signaling since these mutants may have altered ER-mitochondria contact sites, as shown in tissue culture cells lacking the mammalian ortholog MFN2 (de Brito and Scorrano, 2008; Cosson et al., 2012; Filadi et al., 2015, 2016; Leal et al., 2016; Naon et al., 2016; Basso et al., 2018).
*miga-1(tm3621)* mutants show mitochondrial fragmentation and induce UPR$^{mt}$

One of the enhancers we identified is *K01D12.6*, which is conserved from *C. elegans* to humans. The *D. melanogaster* ortholog of this gene has previously been identified in a screen for genes, which when knocked-down induce photoreceptor cell neurodegeneration. Furthermore, it was shown to be required for the maintenance of mitochondrial morphology and hence, named ‘*Mitoguardin*’ (Zhang *et al.*, 2016). Moreover, the two orthologs of this gene in mammals (*MIGA1, MIGA2*) were found to regulate mitochondrial fusion and to be critical for mitochondrial function in human tissue culture cells and in mice (Liu *et al.*, 2016; Zhang *et al.*, 2016; Liu *et al.*, 2017). Therefore, we named *K01D12.6 ‘mitoguardin homolog-1 (miga-1)’*. We verified UPR$^{mt}$ induction using the $P_{hsp-60}$*HSPD1*Gfp (*zcs9*) reporter in the *miga-1(tm3621)* mutant background (Figure 6A). On average, the induction of $P_{hsp-60}$*HSPD1*Gfp is higher in *miga-1(tm3621)* animals compared to *fzo-1(tm1133)* animals. Moreover, we tested the effects of *miga-1(tm3621)* on steady-state mitochondrial morphology, which, in *C. elegans*, is carried out using a mitochondrial matrix-targeted GFP under a promoter that expresses the transgene in body wall muscle cells ($P_{myo-3}$*MYH*Gfp$^{mt}$) (Labrousse *et al.*, 1999; Ichishita *et al.*, 2008; Rolland *et al.*, 2013). While wild-type worms show a tubular network of mitochondria, *miga-1(tm3621)* mutants have a ‘fragmented mitochondria’ phenotype, which is less severe than that caused by the loss of *fzo-1* (Figure 6B). In addition, we analyzed mitochondrial morphology using the MitoSegNet algorithm (Fischer *et al.*, 2020) and confirmed the ‘fragmented mitochondria’ phenotype of *miga-1(tm3621)* mutants. Specifically, for most of the shape descriptors analyzed, *miga-1(tm3621)* mutants were statistically different from wild type but distinct from *fzo-1(tm1133)* mutants (Figure 6C). In summary and in line with previous observations in other organisms, we see drastic changes in mitochondrial morphology in *miga-1(tm3621)* mutants, which are accompanied by the induction of UPR$^{mt}$. 
Data Availability

Strains are available upon request. Figure S1 contains data about involvement of the cSADDS response in suppression of UPR\textsuperscript{mt} upon attenuation of cytosolic translation. Figure S2 shows different mutants inducing the UPR\textsuperscript{mt} reporter. Figure S3 shows the FZOome. Figure S4 contains a subset of the UPR\textsuperscript{mt}ome coming from direct interactions in C. elegans. Figure S5 depicts a subset of the UPR\textsuperscript{mt}ome coming from interactions of human orthologs. Figure S6 shows the complete UPR\textsuperscript{mt}ome. Table S1 contains all suppressors and enhancers of fzo-1(tm1133)-induced UPR\textsuperscript{mt} identified in a genome-wide RNAi screen in C. elegans. Table S2 contains the gene ontology enrichment analysis of suppressors and enhancers of fzo-1(tm1133)-induced UPR\textsuperscript{mt}. Table S3 contains transcription factor enrichment analysis of suppressors and enhancers of fzo-1(tm1133)-induced UPR\textsuperscript{mt}. Table S4 contains the results of the interactome analysis (UPR\textsuperscript{mt}ome). The authors affirm that all data necessary for confirming the conclusions of the article are present within the article, figures, and tables. The supplemental material is available at figshare: https://doi.org/10.25387/g3.14262425.

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**CONFLICT OF INTEREST**

The authors have declared that no competing interests exist.

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FIGURE LEGENDS

Figure 1: Overview of genome-wide RNAi screen for suppressors and enhancers of fzo-1(tm1133)-induced UPR$^{mt}$. (A) Schematic overview of the RNAi screening procedure using the RNAi feeding library (Kamath and Ahringer, 2003) in fzo-1(tm1133) mutants that express the UPR$^{mt}$ reporter $P_{hsp-6\text{mtHSP70}gf}p$ (zcIs13). The moderate induction of the reporter in the fzo-1(tm1133) background allowed screening for both suppressors and enhancers of the response. (B) The screen resulted in identification of 299 suppressors and 86 enhancers of fzo-1(tm1133)-induced UPR$^{mt}$, which were sorted into categories that we defined according to their function. ETC: electron transport chain.

Figure 2: Gene ontology enrichment analysis of suppressors and enhancers of fzo-1(tm1133)-induced UPR$^{mt}$ using DAVID. (A) Results of the clustered gene ontology enrichment analysis of suppressors of fzo-1(tm1133)-induced UPR$^{mt}$ using DAVID (Huang et al., 2008, 2009). (B) Results of the clustered gene ontology enrichment analysis of enhancers of fzo-1(tm1133)-induced UPR$^{mt}$ using DAVID. (A) & (B) Statistically significant ($P>0.05$) enriched GO-terms, except the nematode specific GO-terms, of fzo-1(tm1133)-induced UPR$^{mt}$ are depicted. Circle size correlates with the number of genes associated with a specific GO-term.

Figure 3: Enrichment analysis of transcription factors binding to promoters of candidate genes that suppress or enhance fzo-1(tm1133)-induced UPR$^{mt}$. (A) Transcription factor (TF) binding sites were identified using the modENCODE database (Celniker et al., 2009) and enrichment analysis was performed separately for suppressors and enhancers of fzo-1(tm1133)-induced UPR$^{mt}$ using g:profiler (Knüppel et al., 1994; Raudvere
et al., 2019). TFs that are statistically enriched among the candidate genes are shown. References: [1] (Grove et al., 2009), [2] (Hallam et al., 2000), [3] (Horn et al., 2014), [4] (Huang et al., 2014), [5] (Armakola and Ruvkun, 2019), [6] (Ceol and Horvitz, 2001), [7] (Garbe et al., 2004), [8] (Chi and Reinke, 2006), [9] (Miller et al., 2016), [10] (Baugh et al., 2005), [11] (Maduro et al., 2005), [12] (Lei et al., 2009), [13] (Schwarz et al., 2012), [14] (Gillear et al., 1999), [15] (Budovskaya et al., 2008), [16] (Hu et al., 2017), [17] (Soo-Ung et al., 2009), [18] (Watson et al., 2013), [19] (An and Blackwell, 2003), [20] (An et al., 2005), [21] (Inoue et al., 2005), [22] (Nargund et al., 2012), [23] (Nargund et al., 2015), [24] (Kim and Sieburth, 2018), [25] (Wu et al., 2018) [26] (Neves and Priess, 2005), [27] (McMiller et al., 2007), [28] (Quach et al., 2013), [29] (Ahringer, 1996), [30] (Esmaeili et al., 2002), [31] (Pocock et al., 2004), [32] (Jacquemin et al., 2003), [33] (Furuno et al., 2008), [34] (Klimova et al., 2015), [35] (Ambros and Horvitz, 1984), [36] (Chang et al., 2003), [37] (Uchida et al., 2003), [38] (Etchberger et al., 2007), [39] (Rahe and Hobert, 2019), [40] (Huang et al., 1995), [41] (Wilanowski et al., 2002), [42] (Venkatesan et al., 2003), [43] (Pradel et al., 2007), [44] (Kim et al., 2015). (B) Graphical representation of enriched TFs and the cellular processes they control. ‘Suppressor specific’ TFs are indicated in blue, ‘enhancer specific’ TFs in orange and ‘shared’ TFs in green. The number of candidate genes controlled by a certain group of TFs is indicated in each circle below the functional group name.

**Figure 4: Analysis of a gene network – the UPR\textsuperscript{mt}ome.** Interactors of all genes that are currently associated with the GO-term ‘mitochondrial unfolded protein response’ and of their human orthologs were identified to build the complete UPR\textsuperscript{mt}ome using ‘IntAct’, ‘BioGRID3.5’ and ‘mentha’ databases (Calderone et al., 2013; Orchard et al., 2014; Oughtred et al., 2018). 129 genes are depicted, which overlapped between the complete UPR\textsuperscript{mt}ome and the candidate list of our screen in *fzo-1(tm1133)* mutants. Turquoise circles: ‘input genes’
currently associated with GO-term ‘mitochondrial unfolded protein response’, red arrowheads: suppressors of fzo-1(tm1133)-induced UPR\textsuperscript{mt} that overlap with the complete UPR\textsuperscript{mt}ome, green triangles: enhancers of fzo-1(tm1133)-induced UPR\textsuperscript{mt} that overlap with the complete UPR\textsuperscript{mt}ome. Interactions of two genes that were identified for \textit{C. elegans} genes are indicated with green lines, interactions that were identified in human orthologs are indicated with blue lines.

**Figure 5: Candidate genes with roles in IP\textsubscript{3} signaling.** We identified four genes in our dataset that either play a direct role in the IP\textsubscript{3} signaling pathway or are crucial for the synthesis of phosphatidylinositol-4,5-biphosphate (PI(4,5)P\textsubscript{2}). The IP\textsubscript{3} receptor has previously been identified (Liu \textit{et al.}, 2014). Suppressors are shown in yellow boxes, enhancers in green boxes. PA phosphatidic acid, CDP-DAG cytidine biphosphate-diacylglycerol, PI phosphatidylinositol, PI(4)P phosphatidylinositol-4-phosphate, IP\textsubscript{3} inositol triphosphate, ER endoplasmic reticulum, GPCR G-protein coupled receptor.

**Figure 6: \textit{miga-1(tm3621)} mutants induce UPR\textsuperscript{mt} and have altered mitochondrial morphology.** (A) Fluorescence images of L4 larvae expressing P\textsubscript{hs3-60}HSPD1\textsuperscript{gfp} (zcIs9) in wild type (+/+), \textit{miga-1(tm3621)} or \textit{fzo-1(tm1133)} mutants. Scale bar: 200 μm (B) Fluorescence images of L4 larvae expressing mitochondrial targeted \textit{gfp} (P\textsubscript{myo-3}gfp\textsuperscript{mt}) in wild type (+/+), \textit{miga-1(tm3621)} or \textit{fzo-1(tm1133)} mutants. Representative images are shown. Scale bar: 10 μm. (C) Fluorescence images of L4 larvae expressing mitochondrial targeted \textit{gfp} (P\textsubscript{myo-3}gfp\textsuperscript{mt}) in wild type (+/+), \textit{miga-1(tm3621)} or \textit{fzo-1(tm1133)} mutants were quantified using the MitoSegNet algorithm (Fischer \textit{et al.}, 2020). ns: not significant, *\textit{P}<0.05, **\textit{P}<0.01, ***\textit{P}<0.001, ****\textit{P}<0.0001 using Kruskal-Wallis test with Dunn’s post hoc test for multiple comparison among all three genotypes, \textit{n} ≥ 15. px: pixel.
### A

| Gene name | Human Or奇妙物 | Sequence Logo | P-value | Reference |
|-----------|----------------|--------------|---------|-----------|
| hth-2:ord-1 | NEUROD1,4,8 | ![Sequence Logo](image1) | 3.26E-13 | [1],[2] |
| hthp-1 | PRDM1 | ![Sequence Logo](image2) | 4.82E-08 | [3],[4],[5] |
| ehh-1 | EPHA4, EPHA5 | ![Sequence Logo](image3) | 4.54E-08 | [6],[7],[8],[9] |
| psr-1 | C22X1 | ![Sequence Logo](image4) | 0.000402 | [10],[11],[12] |
| hth-2:hth-10 | TCP12, ASCL4 | ![Sequence Logo](image5) | 0.001189 | [1],[13] |
| enh-3 | GABA3.4 | ![Sequence Logo](image6) | 0.020084 | [14],[15],[16] |
| hth-11 | TFAP4 | ![Sequence Logo](image7) | 0.032103 | [1],[17],[18] |

### B

#### number of candidate genes controlled

![Diagram](image8)

- Development & Cell Fate: 163
- Metabolic: 11
- Oxidative stress: 32
- Aging: 23
