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UPR\textsuperscript{mt} scales mitochondrial network expansion with protein synthesis via mitochondrial import

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Abstract

As organisms develop, individual cells generate mitochondria to fulfill physiologic requirements. However, it remains unknown how mitochondrial network expansion is scaled to cell growth and impacted by environmental cues. The mitochondrial unfolded protein response (UPR\textsuperscript{mt}) is a signaling pathway mediated by the transcription factor ATFS-1 which harbors a mitochondrial targeting sequence (MTS)\textsuperscript{1}. Here, we demonstrate that ATFS-1 mediates an adaptable mitochondrial expansion program that is active throughout normal development. Developmental mitochondrial network expansion required the relatively inefficient MTS\textsuperscript{2} in ATFS-1, which allowed the transcription factor to be responsive to parameters that impact protein import capacity of the entire mitochondrial network. Increasing the strength of the ATFS-1 MTS impaired UPR\textsuperscript{mt} activity throughout development due to increased accumulation within mitochondria. The insulin-like signaling-TORC1\textsuperscript{3} and AMPK pathways affected UPR\textsuperscript{mt} activation\textsuperscript{4,5} in a manner that correlated with protein synthesis. Manipulation to increase protein synthesis caused UPR\textsuperscript{mt} activation. Alternatively, S6 kinase inhibition had the opposite effect due to increased mitochondrial accumulation of ATFS-1.

However, ATFS-1 with a dysfunctional MTS\textsuperscript{6} constitutively increased UPR\textsuperscript{mt} activity independent of TORC1 function. Lastly, expression of a single protein with a strong MTS, was sufficient to expand the muscle cell mitochondrial network in an ATFS-1-dependent manner. We propose that mitochondrial network expansion during development is an emergent property of the synthesis of highly expressed mitochondrial proteins that exclude ATFS-1 from mitochondrial import, causing UPR\textsuperscript{mt} activation. Mitochondrial network expansion is attenuated once ATFS-1 can be imported.
Main

The UPR\textsuperscript{mt} is a mitochondrial-to-nuclear signal transduction pathway regulated by the transcription factor ATFS-1 that is required for development and longevity during mitochondrial dysfunction\textsuperscript{1,7,8}. Because ATFS-1 harbors a MTS and a nuclear localization sequence (NLS), its transcription activity is regulated by subcellular localization. If ATFS-1 is imported into mitochondria, it is degraded by the protease LONP-1\textsuperscript{1} (Fig. 1a). However, if a percentage of ATFS-1 fails to be imported into mitochondria, it traffics to the nucleus to activate a transcriptional response that includes mitochondrial chaperones\textsuperscript{9,10}. Perturbations to OXPHOS or mitochondrial proteostasis activate the UPR\textsuperscript{mt} as both processes are required for mitochondrial protein import\textsuperscript{11}.

A role for ATFS-1 in mitochondrial network maintenance and expansion

We previously found that OXPHOS dysfunction due to deleterious mtDNA heteroplasmy caused an atfs-1-dependent expansion of the mitochondrial network that was observed only when mitophagy was impaired\textsuperscript{12}. Similarly, OXPHOS dysfunction caused by mutations in the ubiquinone biogenesis gene \textit{clk-1}, induced the UPR\textsuperscript{mt} and lead to an increase in mtDNA (Extended Data Fig. 1a) suggesting a role for the UPR\textsuperscript{mt} in mitochondrial biogenesis or network expansion.

\textit{atfs-1(et18)} worms constitutively activate the UPR\textsuperscript{mt} due to an amino acid substitution in the MTS which impairs import into mitochondria even in the absence of mitochondrial stress\textsuperscript{6}. Impressively, \textit{atfs-1(et18)} worms harbored more mtDNAs relative to wildtype worms (Fig. 1b), suggesting that UPR\textsuperscript{mt} activation is sufficient to expand the mitochondrial network. Conversely, worms lacking the entire \textit{atfs-1} open reading frame (\textit{atfs-1(null)})\textsuperscript{13} had reduced mtDNAs (Fig. 1c). Moreover, TMRE staining indicated that \textit{atfs-1(null)} or
atfs-1 RNAi treated worms harbor fewer functional mitochondria relative to wildtype worms, in intestinal cells (Fig. 1d-f) suggesting the UPR\textsuperscript{mt} is actively involved in the maintenance and expansion of the mitochondrial network during development. Importantly, both atfs-1\textit{(null)} and atfs-1\textit{(et18)} caused developmental delays and impaired respiration (Fig. 1g-i). The reduction in respiratory capacity in the atfs-1\textit{(et18)} strain was consistent with reduced TMRE staining in intestinal cells (Extended Data Fig. 1b-c). Thus, in the absence of the UPR\textsuperscript{mt}, mtDNAs and functional mitochondrial are reduced, while continuous UPR\textsuperscript{mt} activation results in a partial expansion of the mitochondrial network that yields dysfunctional mitochondria.

ATFS-1 mediates a mitochondrial expansion program during development

To elucidate the role of ATFS-1 in mitochondrial expansion and homeostasis, we compared transcriptional profiles of wildtype, atfs-1\textit{(null)} and atfs-1\textit{(et18)} worms during development in the absence of mitochondrial stress. Remarkably, atfs-1\textit{(et18)} worms induced mitochondrial genes including the proteostasis components associated with the UPR\textsuperscript{mt} (Fig. 2a-d, Extended data Figure 2a, Supplementary table 1). Furthermore, over 50 genes required for mitochondrial ribosome function were upregulated, as were genes required for mtDNA replication, and cardiolipin biosynthesis pathway genes required for mitochondrial inner membrane synthesis. Lastly, genes required for both mitochondrial protein import and OXPHOS complex assembly were also upregulated.

Conversely, most of the mitochondrial genes induced in the atfs-1\textit{(et18)} strain were downregulated in the atfs-1\textit{(null)} worms compared to wildtype worms (Fig. 2a, 2e-g, Supplementary table 2-3), consistent with less mtDNA and TMRE staining in intestinal cells.
cells (Fig 1). Interestingly, the OXPHOS protein NDUFS3 was decreased in the atfs-1(null), while unaffected in atfs-1(et18) worms (Fig. 2h, Extended Data Fig. 2b). To exclude potential effects of mitochondrial degradation via mitophagy, these studies were performed in a strain lacking pdr-1 (Parkin)\(^{14}\). Interestingly, multiple metabolic components including those of the TCA cycle and OXPHOS were repressed in atfs-1(et18) relative to wildtype worms (Supplementary table 4), consistent with UPR\(^{\text{mt}}\) limiting expression of highly expressed mitochondrial proteins\(^{10}\) and the reduced TMRE staining (Extended data Fig. 1c). Lastly, while many mitochondrial mRNAs were expressed at lower levels in atfs-1(null) worms, mtDNA replication and mitophagy components were upregulated suggesting an alternative stress response(s) is induced in the absence of atfs-1 (Fig. 2a).

Because of the alterations in mtDNA levels, TMRE, and transcription of mitochondrial components, we visualized mitochondria via transmission electron microscopy. Impressively, mitochondria in atfs-1(null) worms were smaller and appeared defective in both intestine and muscle cells, along with pervasive muscle cell aberrations (Fig. 2i-l, Extended data Fig. 2c). In contrast, mitochondria in atfs-1(et18) were elongated, particularly visible in the intestine (Fig. 2i-l, Extended data Fig. 2c). Combined, our results suggest that ATFS-1 regulates a mitochondrial expansion program.

**A weak MTS regulates ATFS-1 and mitochondrial network expansion**

We next sought to determine how ATFS-1 is regulated, or excluded from mitochondria, during development. Because ATFS-1 harbors a MTS along with a NLS, we have proposed that the UPR\(^{\text{mt}}\) is regulated by protein import capacity of the entire mitochondrial
network\(^1\). ATFS-1 is predicted to have a relatively weak, or inefficient, MTS compared to other mitochondrial-targeted proteins such as mitochondrial chaperones and OXPHOS components\(^{15,16}\) (Fig. 3a). To compare the MTS strength of the OXPHOS protein ATP synthase subunit 9 (Su9) to ATFS-1, the amino-terminus of each was fused to GFP and expressed in HEK293T cells. As expected, both GFP-fusion proteins accumulated within mitochondria, but unlike Su9\(^{(1-69)}::\text{GFP}\), ATFS-1\(^{(1-100)}::\text{GFP}\) fluorescence also accumulated within the cytosol, but to a lesser extent than that of ATFS-1\(^{\text{et18}(1-100)}::\text{GFP}\) (Fig. 3b). Additionally, import of ATFS-1\(^{(1-100)}::\text{GFP}\) was limited compared to Su9\(^{(1-69)}::\text{GFP}\) in an \textit{in vitro} import assay (Fig. 3d) consistent with ATFS-1 harboring a weak MTS.

We hypothesized that the inefficient MTS allows ATFS-1 import and UPR\(^{\text{mt}}\) activation to be sensitized to conditions that impact mitochondrial import capacity including mitochondrial stress, total mitochondria, and potentially the flux of other proteins into mitochondria. Thus, we sought to generate a worm strain expressing ATFS-1 with a stronger, or more efficient, MTS. Amino acid substitutions of T10 and D24 to arginine are predicted to increase MTS strength (Fig. 3a, 3c). Similar to Su9\(^{(1-69)}::\text{GFP}\), ATFS-1\(^{\text{R/R}(1-100)}::\text{GFP}\) only accumulated within mitochondria and not in the cytosol in HEK293T cells (Fig. 3b). Furthermore, more ATFS-1\(^{\text{R/R}(1-100)}::\text{GFP}\) accumulated within mitochondria than ATFS-1\(^{(1-100)}::\text{GFP}\) in an \textit{in vitro} import assay, consistent with increased MTS strength (Fig. 3d).

Via CRISPR-Cas9, mutations were introduced at the endogenous \textit{atfs-1} locus to generate ATFS-1\(^{\text{R/R}}\). We first examined accumulation of ATFS-1\(^{\text{R/R}}\) within mitochondria during normal development by raising worms on \textit{lonp-1}(RNAi), which impairs ATFS-1
degradation within the matrix\textsuperscript{1}. Strikingly, more ATFS-1\textsuperscript{R/R} accumulated within mitochondria compared to wildtype ATFS-1 or ATFS-1\textsuperscript{et18} during normal development (Fig. 3e). And, ATFS-1\textsuperscript{R/R} worms expressed less hsp-6\textsubscript{pr}::gfp relative to wildtype or atfs-1\textsuperscript{et18} worms during normal development (Fig. 3f, Extended Data Fig. 3a) and had reduced expression of hsp-6 and timm-23 mRNAs (Extended Data Fig. 3b-c). ATFS-1\textsuperscript{R/R} also impaired UPR\textsuperscript{mt} activation caused by ethidium bromide (EtBr) exposure (Fig. 3g, Extended Data Fig. 3d). However, timm-23(RNAi) which impairs a component required for import of most proteins harboring amino-terminal MTSs\textsuperscript{11}, caused UPR\textsuperscript{mt} activation in both ATFS-1\textsuperscript{R/R} and wildtype worms (Fig. 3g, Extended Data Fig. 3d) indicating ATFS-1\textsuperscript{R/R} is a functional transcription factor likely impaired due to increased mitochondrial accumulation. Similar to worms lacking atfs-1, worms expressing ATFS-1\textsuperscript{R/R} developed slower (Fig. 3h) and exhibited a perturbed and fragmented mitochondrial network in both intestine and muscle cells along with a reduction in mtDNA (Fig. 3i-j, Extended Data Fig. 3e-g).

Combined, these data suggest that ATFS-1 regulates a transcriptional program to expand mitochondrial biomass that is active throughout development and reliant on an inefficient MTS that confers sensitivity to conditions that impact mitochondrial import capacity. These findings suggest that during development a percentage of ATFS-1 cannot be imported into mitochondria of growing cells resulting in modest UPR\textsuperscript{mt} activation and mitochondrial network expansion (Fig. 3k).

**Interplay between protein synthesis, mitochondrial import, and ATFS-1**
Previous screens for components required for UPR\textsuperscript{mt} activation identified multiple regulators of growth-related protein synthesis including the insulin-like receptor \textit{daf-2}, \textit{rheb-1}, \textit{mTOR} (\textit{let-363}), and \textit{rsks-1} (S6 kinase).\textsuperscript{4,5} TORC1 regulates protein synthesis rates in response to diverse inputs including growth signals and cellular energetics.\textsuperscript{3} Insulin-like signaling-TORC1 promotes protein synthesis by phosphorylating RSKS-1, which in turn, phosphorylates a ribosomal subunit\textsuperscript{3} (Extended Data Fig. 4a). Alternatively, the 5' AMP-activated protein kinase (AMPK) limits TORC1 activity and protein synthesis when ATP levels are low.\textsuperscript{17,18}

As expected, DAF-2 inhibition impaired induction of \textit{hsp-6}\textsuperscript{pr}::\textit{gfp} (Extended Data Fig. 4b),\textsuperscript{19} but \textit{hsp-6} and \textit{atfs-1} mRNAs were also reduced during normal development and during mitochondrial stress (Extended Data Fig. 4c-d). Inhibition of TORC1 components \textit{rheb-1}, \textit{raga-1}, \textit{mTOR}, and \textit{rsks-1} also reduced \textit{hsp-6}\textsuperscript{pr}::\textit{gfp} (Extended Data Fig. 4e-g), as well as \textit{hsp-6} and \textit{atfs-1} mRNA levels, as did starvation (Fig. 4a-b, Extended Data Fig. 4h-j). Conversely, inhibition of AMPK, which increases TORC1 activity, resulted in increased \textit{hsp-6} mRNA in an \textit{atfs-1} and \textit{mTOR}-dependent manner (Fig. 4c-d) while AMPK activation\textsuperscript{20} reduced \textit{hsp-6} transcripts (Fig. 4e). Combined, our results indicate that TORC1 and RSKS-1 are required for UPR\textsuperscript{mt} during development and suggest that increased protein synthesis during development stimulates UPR\textsuperscript{mt} activity.

In mammals, TORC1 promotes protein synthesis by phosphorylating S6 kinase and 4EBP, which requires a TOR signaling (TOS) motif in each protein.\textsuperscript{21} Interestingly, ATFS-1 also harbors a canonical TOS motif (-FEMDI-) (Fig. 3C), which we mutated at the endogenous locus to yield -AEMDI- (\textit{atfs-1}(-TOS)). UPR\textsuperscript{mt} activation was attenuated in \textit{atfs-1}(-TOS) worms relative to wildtype worms upon EtBr exposure (Extended Data Fig.
4k). Importantly, the TOS motif was also required for the increased \( hsp-6_{pr}::gfp \) (Extended Data Fig. 4l) and mtDNA (Fig. 1b) in \( atfs-1(et18) \) worms suggesting the TOS motif promotes nuclear function of ATFS-1, similar to the TOS motif found in the transcription factor HIF-1α\(^{22} \). Interestingly, \( hsp-6_{pr}::gfp \) induction caused by EtBr or complex III-deficiency \( (isp-1(qm150))^{23} \), was impaired further in \( rsks-1(ok1255);atfs-1(\Delta TOS) \) worms relative to either \( atfs-1(\Delta TOS) \) or \( rsks-1(ok1255) \) worms (Extended Data Fig. 4m-o), suggesting that RSKS-1 promotes UPR\(^{mt} \) activation independent of the TOS motif.

We next examined the effect of \( rsks-1 \) inhibition on expression and trafficking of ATFS-1. One possibility is that \( rsks-1 \) inhibition simply reduces synthesis of ATFS-1 limiting its nuclear transcription activity. Thus, we examined the mitochondrial accumulation of ATFS-1, by inhibiting LONP-\(^{1} \). Interestingly, more ATFS-1 accumulated within mitochondria in \( rsks-1(ok1255) \) worms relative to wildtype worms (Fig. 4f). Thus, rather than reduced ATFS-1 expression, UPR\(^{mt} \) impairment in worms lacking RSKS-1 is due to mitochondrial accumulation of ATFS-1, similar to ATFS-1\(^{R/R} \) (Fig. 3e), which prevents trafficking to nuclei. Consistent with \( rsks-1(ok1255) \) impairing UPR\(^{mt} \) activation by increasing mitochondrial import capacity, \( rsks-1(\text{RNAi}) \) did not reduce \( hsp-6_{pr}::gfp \) in \( atfs-1(et18) \) worms with an impaired MTS (Fig. 4g) or when treated with \( timm-23(\text{RNAi}) \) (Extended Data Fig. 5a). Moreover, activation of ATFS-1 persisted during starvation in \( atfs-1(et18) \) worms and \( timm-23(\text{RNAi}) \) treated worms (Extended Data Fig. 5b-c).

Because RSKS-1 is required for protein synthesis during cell growth, we hypothesized that the high rate import of proteins into mitochondria may cause UPR\(^{mt} \) activation and mitochondrial network expansion during normal development. As a test of this model, we sought to determine the impact of overexpressing a single protein with a
relatively strong MTS on the mitochondrial network. The mitochondrial network was examined in muscle cells of worms expressing GFP or mtGFP via the strong myo-3 promoter\textsuperscript{24}. Importantly, mtGFP harbors a relatively strong MTS (aa 1-24) from the enzyme aspartate aminotransferase (AST) (Extended Data Fig. 5d), and the myo-3 promoter is expressed throughout development\textsuperscript{25}(Extended Data Fig. 5e). Relative to GFP, mtGFP expression increased accumulation of functional mitochondria as determined by TMRE staining (Fig. 4h-i), mtDNA number (Fig 4j), and hsp-6, timm-17 and timm-23 mRNAs (Fig. 4k-l, Extended Data Fig. 5f) despite mtGFP being expressed at a lower level than GFP (Extended Data Fig. 5g). Expansion of TMRE staining by mtGFP was impaired by atfs-1(RNAi) (Fig. 4m-n), atfs-1(null) (Fig. 4o-p) and in ATFS-1\textsuperscript{R/R} worms (Fig. 4q), however mtGFP transcription was not affected by the atfs-1(null) allele (Extended Data Fig. 5h).

To determine if the perturbed mitochondrial network was due to the inability of ATFS-1 to traffic to the nucleus and activate the UPR\textsuperscript{mt}, we used CRISPR-Cas9 to generate impaired NLS in ATFS-1 (R426A). ATFS-1\textsuperscript{(\textsuperscript{\text{\text{\text{\text{\text{\text{ANLS}}})}}))} accumulated within mitochondria similar to wildtype ATFS-1 (Extended Data Fig. 5i), but failed to induce hsp-6\textsubscript{pr::gfp} or endogenous hsp-6 transcripts during mitochondrial stress elicited by knockdown of the mitochondrial protease SPG-7 (Extended Data Fig. 5j-k). Similar to ATFS-1\textsuperscript{R/R}, or atfs-1(null) worms, ATFS-1\textsuperscript{(\textsuperscript{\text{\text{\text{\text{\text{\text{ANLS}}})}}})} resulted in fragmented mitochondrial morphology in muscle cells indicating that the nuclear activity of ATFS-1 is essential for its function (Extended Data Fig. 5l). Combined, these finding indicate that expression of a single protein with a strong MTS is sufficient to expand the mitochondrial network in an atfs-1-dependent manner.
Discussion

In summary, we have found that ATFS-1 regulates a mitochondrial expansion program that is active throughout normal development. Developmental mitochondrial expansion required the inefficient MTS of ATFS-1 and TORC1 activity suggesting an interplay between protein synthesis, mitochondrial protein import capacity, and nuclear activity of ATFS-1. Consistent with these findings, OXPHOS transcripts are among the most highly expressed mRNAs in worms (Extended Data Fig. 6a). And, C. elegans ribosome profiling data indicates that OXPHOS proteins are translated primarily during the early stages of worm development (L1,L2), and are reduced or absent by the L4 stage (Extended Data Fig. 6b). Interestingly, the ATFS-1 ribosome profile mirrors the OXPHOS profiles early in development and is also diminished at L4 (Extended Data Fig. 6c), consistent with the observation that the UPR\textsuperscript{mt} can only be activated by stress prior to the L4 stage\textsuperscript{26}. Intriguingly, it was recently reported that mitochondrial metabolic proteins are prone to stalling within mitochondrial import channels under basal conditions in growing cells\textsuperscript{27,28}, suggesting import or intra-mitochondrial protein processing can be overwhelmed during normal cell growth. We propose a model where the high levels of mitochondrial protein synthesis that occurs during development drives mitochondrial network expansion by excluding a percentage of ATFS-1 from mitochondrial import. And, network expansion continues until import is sufficient to import ATFS-1 and terminate the UPR\textsuperscript{mt}. These findings are conceptually similar to the endoplasmic reticulum expansion that occurs in response to increased protein flux via the UPR\textsuperscript{ER}, which is regulated by IRE1 and XBP1\textsuperscript{29}.
We propose that as a function of the mitochondrial import flux or mitochondrial protein processing, ATFS-1 scales mitochondrial network expansion with cell growth.
Methods

Worms, plasmids and staining

The reporter strain hsp-6::gfp for visualizing UPR^{mt}, the myo-3::gfp and the myo-3::mtgfp for visualization of mitochondrial mass and atfs-1(null) worms have been previously described^{13,30,31}. The MTS in the myo-3::mtgfp is the first 24 amino acids from the enzyme aspartate aminotransferase from *Coturnix japonica* (1-MALLQSRLLLSAPRRAATARRSS-24) fused to GFP. The atfs-1(et18) strain was a gift from Marc Pilon. N2(wildtype), isp-1(qm150), rsks-1(ok1255) and daf-2(e1370), were obtained from the Caenorhabditis Genetics Center (Minneapolis, MN).

The atfs-1^{R/R}(cmh16), the atfs-1(ΔTOS)(cmh17) and the atfs-1(ΔNLS)(cmh18) strains were generated via CRISPR-Cas9 in hsp-6::gfp worms as described^{13}. The atfs-1(ΔTOS) was generated in both the wildtype worms as well as in the atfs-1(et18) strain.

The crRNAs (IDT) were co-injected with purified Cas9 protein, tracrRNA (Dharmacon), repair templates (IDT) and the pRF4::rol-6(su1006) plasmid as described^{32,33}. The crRNAs and repair templates used in this study are listed in Supplementary Table 5. The pRF4::rol-6 (su1006) plasmid was a gift from Craig Mello^{34}. The ATFS-1^{1-100}:GFP expressing plasmid was previously described^{1}. The ATFS-1^{1-100(R4C)}:GFP and the ATFS-1^{1-100(T10R, D24R)}:GFP were generated by introducing mutations to yield the described amino acid substitutions in the ATFS-1^{1-100}:GFP expressing plasmid. The subunit 9 of the F0-ATPase (SU9)^{1-69}:GFP PQCXIP expression plasmid was a gift from Xuejun Jiang.

Worms were raised HT115 strain of *E. coli* and RNAi performed as described^{35}. Ethidium bromide (EtBr) and TMRE experiments were performed by synchronizing and raising worms on plates previously soaked with M9 buffer containing EtBr or 2µM TMRE.
Worms were analyzed at the L4 larvae stage except for EtBr treated worms that led to developmental arrest. EtBr treated worms were analyzed at the same time as the control.

**Protein analysis and antibodies**

Synchronized worms were raised on plates with control(RNAi) or *lonp-1*(RNAi) to the L4 stage prior to harvesting. Whole worm lysate preparation was previously described\(^3^0\). Antibodies against α-tubulin were purchased from Calbiochem (CP06), GFP and for NDUFS3 from Abcam (ab6556 and ab14711 respectively). Antibodies for ATFS-1 were previously described\(^1\). Immunoblots were visualized using ChemiDoc XRS+ system (Bio-Rad). All western blot experiments were performed at least three times.

**mtDNA quantification**

mtDNA quantification was performed using a qPCR-based method similar to previously described assays\(^3^6\). 20–30 worms were collected in 30 µl of lysis buffer (50 mM KCl, 10 mM Tris-HCl (pH 8.3), 2.5 mM MgCl\(_2\), 0.45% NP-40, 0.45% Tween 20, 0.01% gelatin, with freshly added 200 µg/ml proteinase K) and frozen at −80°C for 20 minutes prior to lysis at 65°C for 80 minutes. Relative quantification was used for determining the fold changes in mtDNA between samples. 1 µl of lysate was used in each triplicate qPCR reaction. qPCR was performed using the Thermo-Scientific SyBr Green Maxima Mix and the MyiQ2 Two-Color Real-Time PCR Detection System (Bio-Rad Laboratories). Primers that specifically amplify mtDNA are listed in Supplementary table 5. Primers that amplify a non-coding region near the nuclear-encoded ges-1 gene were used as a control. mtDNA was harvested from synchronized worms at the L4 stage. All qPCR results have
been repeated at least 3 times and performed in triplicates. A Student’s t-test was employed to determine the level of statistical significance.

**RNA isolation and qRT-PCR**

RNA isolation and qRT-PCR analysis were previously described\(^\text{12}\). Worms were synchronized by bleaching, raised on HT115 *E. coli* and harvested at the L4 stage. Total RNA was extracted from frozen worm pellets using RNA STAT (Tel-Test) and 500 ng RNA was used for cDNA synthesis with qScript™ cDNA SuperMix (QuantaBio). qPCR was performed using iQ™ SYBR® Green Supermix (Bio-Rad Laboratories). qPCR primers are listed in Supplementary Table 5. All qPCR results were repeated at least 3 times and performed in triplicates. A Student’s t-test was employed to determine the level of statistical significance.

**Oxygen Consumption**

Oxygen consumption was measured using a Seahorse XFe96 Analyzer at 25°C similar to that described previously\(^\text{37}\). In brief, L4 worms were transferred onto empty plates and allowed to completely digest the remaining bacteria for 1 hour, after which 10 worms were transferred into each well of a 96-well microplate containing 180 µl M9 buffer. Basal respiration was measured for a total of 30 minutes, in 6 minute intervals that included a 2 minute mix, a 2 minute time delay and a 2 minute measurement. To measure respiratory capacity, 15 µM FCCP was injected, the OCR (oxygen consumption rate) reading was allowed to stabilize for 6 minutes then measured for five consecutive intervals.
Mitochondrial respiration was blocked by adding 40mM Sodium azide. Each measurement was considered one technical replicate.

**Cultured cells and imaging**
HEK293T cells were transfected with 0.5 µg of the expression plasmids: SU91-69::GFP with ATFS-11-100::GFP, ATFS-11-100(R/R)::GFP and ATFS-11-100(et18)::GFP via Lipofectamine. The cells were imaged sixteen hours post transfection.

**RNA-sequencing and differential expression analysis**
cDNA libraries were constructed with standard Illumina P5 and P7 adapter sequences. The cDNA libraries were run on an Illumina HiSeq2000 instrument with single-read 50-bp (SR50). RNA reads were then aligned to WBcel235/ce11 reference genome and differential gene expression analysis was performed with edgeR. Differences in gene expression between atfs-1(et18) and atfs-1(null) compared to wildtype are listed in Supplementary Tables 6 and 7 respectively.

**Analysis of worm development**
Worms were synchronized via bleaching and allowed to develop on HT115 bacteria plates for 3 days at 20°C. Developmental stage was quantified as a percentage of the total number of animals. Each experiment was performed three times. For the comparison of wildtype and atfs-1(null) worms; N=162 (wildtype), and 282 (atfs-1(null)). For the comparison of wildtype to atfs-1(R/R) worms; N=158 (wildtype) and N=256 (atfs-1(R/R)).
Statistics

All experiments were performed at least three times yielding similar results and comprised of biological replicates. The sample size and statistical tests were chosen based on previous studies with similar methodologies and the data met the assumptions for each statistical test performed. No statistical method was used in deciding sample sizes. No blinded experiments were performed, and randomization was not used. For all figures, the mean ± standard deviation (s.d.) is represented unless otherwise noted.

Microscopy

C. elegans were imaged using either a Zeiss AxioCam 506 mono camera mounted on a Zeiss Axio Imager Z2 microscope or a Zeiss AxioCam MRc camera mounted on a Zeiss SteREO Discovery.V12 stereoscope. Images with high magnification (63×) were obtained using the Zeiss ApoTome.2. Exposure times were the same in each experiment. Cell cultures were imaged with the Zeiss LSM800 microscope. All images are representatives of more than three images. Quantification of fluorescent intensity as well as creating binary skeleton-like structures were done with the Fiji software.

Gene set enrichment analysis

The OXPHOS gene set was downloaded from WormBase Ontology Browser. mRNA abundance was measured and ranked by reads per kilobase per million reads (RPKM) from RNA-seq data. Pre-ranked gene set enrichment analysis was performed with GSEA3.0 software with 'classical' scoring.
Transmission Electron Microscopy

L4 larvae were transferred to 2.5% glutaraldehyde in 0.1 M Sodium Cacodylate buffer pH 7.2 for 10 min. The tail and head of each worm were dissected out and the main body was transferred to fresh 2.5% glutaraldehyde in 0.1 M Sodium Cacodylate buffer and kept at 4°C overnight. Samples were processed and analyzed at the University of Massachusetts Medical School Electron Microscopy core facility according to standard procedures. Briefly, the samples were rinsed three times in the same fixation buffer and post-fixed with 1% osmium tetroxide for 1h at room temperature. Samples were then washed three times with ddH₂O for 10 minutes and then dehydrated through a graded ethanol series of 20% increments, before two changes in 100% ethanol. Samples were then infiltrated first with two changes of 100% Propylene Oxide and then with a 50%/50% propylene oxide/SPI-Pon 812 resin mixture. The following day, five changes of fresh 100% SPI-Pon 812 resin were performed before the samples were polymerized at 68°C in flat pre-filled embedding molds. The samples were then reoriented, and thin sections (approx. 70nm) were placed on copper support grids and contrasted with Lead citrate and Uranyl acetate. Sections were examined using a CM10 TEM with 100Kv accelerating voltage, and images were captured using a Gatan TEM CCD camera.

Ribosome profiling data analysis

Ribosome profiling sequencing data was downloaded from the NCBI Sequence Read Archive (SRA) (http://www.ncbi.nlm.nih.gov/sra/) under accession number SRA055804. Data was analyzed as previously described. Data analysis was done with the help of Unix-based software tools. First, the quality of raw sequencing reads was determined by
FastQC\textsuperscript{42}. Reads were then filtered according to quality via FASTQ for a mean PHRED quality score above 30\textsuperscript{43}. Filtered reads were mapped to the \textit{C. elegans} reference genome (Wormbase WS275) using BWA (version 0.7.5) and SAM files were converted into BAM files by SAMtools (version 0.1.19). Coverage data for specific genes (including 5'UTR, exons and 3'UTR) were calculated by SAMtools and coverage data for each gene was plotted using R\textsuperscript{44}.

\textbf{Mitochondria isolation and in vitro protein import}

Cells (budding yeast W303) were grown to logarithmic phase in YPD (1\% yeast extract, 2\% peptone, 2\% glucose), collected by centrifugation and washed once with water. Cells were then resuspended in 0.1 M Tris pH 9.4, 10 mM DTT and incubated for 20 min at 30°C. Cell walls were disturbed by incubation in 1.2M sorbitol, 20mM K2HPO4 pH 7.4, 1\% zymolyase for 1 h at 30°C. Dounce homogenization was used to lyse the cells in 0.6M sorbitol, 10mM Tris pH 7.4, 1mM EDTA, fatty acid free 0.2\% BSA and 1mM PMSF. Mitochondria were then isolated by differential centrifugation as described previously\textsuperscript{45} and resuspended in SEM buffer (0.25M sucrose, 10mM MOPS KOH pH 7.2 and 1mM EDTA).

The coupled Transcription/Translation system (T7 Quick for PCR DNA, Promega) was used to express ATFS-1 from a PCR template. Precursor proteins (ATFS-1\textsuperscript{1-100}::GFP, ATFS-1\textsuperscript{1-100(R/R)}::GFP and Su9\textsuperscript{1-69}::GFP) were synthesized in reticulocyte lysate in the presence of [35S]methionine (T7 Quick for PCR DNA, Promega). Import into isolated mitochondria was performed in import buffer (3 \% (w/v) BSA, 250 mM sucrose, 80 mM KCl, 5 mM methionine, 5 mM MgCl2, 2 mM KH2PO4, 10 mM MOPS-KOH, pH
7.2, 4 mM NADH, 2 mM ATP, 5 mM creatine phosphate, 0.1 mg/ml creatine kinase) at 25°C. The import reaction was stopped on ice or by addition of AVO (8 μM antimycin A, 20 μM oligomycin, 1 μM valinomycin). To dissipate Δψ, AVO was added before the import experiment. Samples were treated with 25 μg/ml proteinase K for 15 min on ice, following by treatment with 2 mM PMSF for 5 min on ice. Mitochondrial were washed twice with SEM buffer and analyzed by electrophoresis on SDS-PAGE.

Data availability

The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih.gov/geo (accession no. GSE114951). Data also available from the corresponding author upon reasonable request.
References


Acknowledgements We thank W. Mair, M. Pilon and the *Caenorhabditis* Genetics Center for providing *C. elegans* strains (funded by NIH Office of Research Infrastructure Programs (P40 OD010440), and the UMass Medical School Core facilities for deep sequencing and electron microscopy. This work was supported by HHMI, the Mallinckrodt Foundation, and National Institutes of Health grants (R01AG040061 and R01AG047182) to C.M.H. and (SI0OD021580) to L.S. The authors are solely responsible for the content.


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Data deposition The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih.gov/geo (accession no. GSE114951).
Fig 1. ATFS-1 regulates mitochondrial network expansion.

a. Schematic of ATFS-1 regulation.

b. Quantification of mtDNA in wildtype, atfs-1(et18) and atfs-1(et18;ΔTOS) as determined by qPCR. N=4. Error bars mean +/- s.d, *p<0.05 (Student’s t-test).

c. Quantification of mtDNA in wildtype and atfs-1(null). N=4. Error bars mean +/- s.d, *p<0.05 (Student’s t-test).

d. Quantification of TMRE intensity in wildtype and atfs-1(null) worms. N=21(wildtype), N=33 (atfs-1(null)). Error bars mean +/- s.d, ***p<0.001, (Student’s t-test).

e. TMRE staining of wildtype worms raised on control or atfs-1(RNAi) and atfs-1(null) worms. Skeleton-like binary backbone is presented (bottom). Scale bar 10µm.

f. Quantification of TMRE intensity in wildtype and atfs-1(RNAi) worms. N=25. Error bars mean +/- s.d, ****p<0.0001 (Student’s t-test).

g. Developmental stages of 3 day old wildtype, atfs-1(et18) or atfs-1(null) worms. N=546 (wildtype), N=597 atfs-1(et18) and 627 (atfs-1(null)).

h-i. Oxygen consumption rates (OCR) in wildtype, atfs-1(et18) and atfs-1(null). Basal respiration (h), maximal respiration (i). N=9. Error bars mean +/- s.d, **p<0.01,***p<0.001,****p<0.0001 (Student’s t-test).
Figure 2

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Fig 2. ATFS-1 mediates a mitochondrial network expansion program.

a. Number of differentially regulated genes induced in atfs-1(et18) worms relative to wildtype (WT) or reduce in atfs-1(null) worms relative to wildtype. Green-upregulated, red-downregulated and the number of overlapping genes between atfs-1(et18) and atfs-1(null) worms are listed in the last column.

b-g. Transcript levels of translation elongation factor mitochondrial 1 (tsfm-1) and G elongation factor mitochondrial 1 (gfm-1) (b,e), translocase of inner mitochondrial membrane 17B.1 (timm-17B.1) and translocase of inner mitochondrial membrane 23 (timm-23) (c,f), heat shock protein 6 (hsp-6) and NADH:ubiquinone oxidoreductase complex assembly factor 3 (nuaf-3) (d,g) as determined by qRT-PCR in wildtype and atfs-1(et18) (b-d) or in wildtype and atfs-1(null) (e-g). N=3 except for gfm-1(e) and nuaf-3(g) N=4. Error bars mean +/- s.d, *p<0.05, **p<0.01 ***p<0.001, ****p<0.001 (Student’s t-test).

d. SDS-Page immunoblots of lysates from pdr-1(tm598), atfs-1(null);pdr-1(tm598) and atfs-1(et18);pdr-1(tm598) worms. NDUFS3 is a component of the NADH:Ubiquinone Oxidoreductase complex I and tubulin (Tub) was used as a loading control.

i-k. Transmission electron microscopy of body wall muscle of wildtype (i), atfs-1(null) (j) and atfs-1(et18) worms (k). Scale bar 1 µm.

l. Transmission electron microscopy of intestinal cells from wildtype, atfs-1(null) and atfs-1(et18) worms. Mitochondria are highlighted in the middle panel. Scale bar 1 µm (left) and 200nm (right).
**Figure 3**

(a) **ATFS-1** expression levels in different cell lines, showing high levels in control conditions.

(b) Western blot analysis of **ATFS-1** and other mitochondrial proteins under normal and mutated conditions.

(c) GFP expression in cells with and without **ATFS-1** expression.

(d) Time course analysis of **ATFS-1** expression over a period of 0 to 100 minutes, showing increased expression levels.

(e) Protein levels of **ATFS-1** and other proteins in WT and mutant conditions, with arrows indicating the direction of the protein mobility.

(f) Western blot analysis of **ATFS-1** and other mitochondrial proteins under different conditions.

(g) GFP expression in cells with and without **ATFS-1** expression.

(h) **ATFS-1** expression levels in different developmental stages, with a percentage scale.

(i) TMRE staining in WT and mutant conditions, with a color gradient indicating the intensity of the staining.

(j) Fold change in mtDNA expression levels in different conditions, with a line graph showing the trend.

(k) Schematic representation of the **ATFS-1** pathway, illustrating the role of **ATFS-1** in mitochondrial function.

If a percentage of **ATFS-1** cannot be imported into mitochondria, it traffics to the nucleus to activate the **UPR** until **ATFS-1** can be imported into mitochondria.
Fig 3. **UPR\textsuperscript{mt} requires the weak MTS in ATFS-1.**

a. Mitochondrial targeting sequence probability prediction using MitoFates. OXPHOS proteins (red), mitochondrial chaperones (blue), ATFS-1 (black).

b. HEK293T cells expressing ATFS-1\textsuperscript{1-100}::GFP, ATFS-1\textsuperscript{1-100(R/R)}::GFP, ATFS-1\textsuperscript{1-100(et18)}::GFP or SU9\textsuperscript{1-69}::GFP. Scale bar 10 µm.

c. ATFS-1 schematic highlighting ATFS-1\textsuperscript{R/R} amino acid substitutions.

d. In organelle import of radiolabeled ATFS-1\textsuperscript{1-100}::GFP, ATFS-1\textsuperscript{1-100(R/R)}::GFP and Su9\textsuperscript{1-69}::GFP into isolated mitochondria. After the indicated time points, mitochondria were washed and analyzed by SDS-PAGE electrophoresis. ATFS-1, Su9, mature (m) are marked.

e. SDS-Page immunoblots of lysates from wildtype, atfs-1\textsuperscript{R/R} and atfs-1(et18) worms raised on control or lonp-1(RNAi). ATFS-1 is marked (●) and tubulin (Tub) was used as a loading control.

f. *hsp-6\textsubscript{Pr}::gfp* in wildtype, atfs-1(et18) and atfs-1\textsuperscript{R/R} worms. Scale bar 0.1 mm.

g. *hsp-6\textsubscript{Pr}::gfp* in wildtype and atfs-1\textsuperscript{R/R} worms raised on 50µg/ml EtBr, control or *timm-23*(RNAi). Scale bar 0.1 mm.

h. Developmental stages of 3 day old wildtype or atfs-1\textsuperscript{R/R} worms. N=158 (wildtype) and N=256 (atfs-1\textsuperscript{R/R}).

i. TMRE staining of wildtype and atfs-1\textsuperscript{R/R} worms. Skeleton-like binary backbone is presented (right). Scale bar 10 µm.

j. Quantification of mtDNA in wildtype and atfs-1\textsuperscript{R/R} worms as determined by qPCR. N=3. Error bars mean +/- s.d, *p<0.05 (Student’s t-test).

k. Proposed model for ATFS-1 mediated mitochondria expansion.
Figure 4

(a) [Graph showing fold increase in RFU for hsp-6 and atfs-1.]

(b) [Graph showing fold increase in RFU for rsks-1.]

(c) [Image showing control and atfs-1 (RNAi) conditions, with GFP and TMREMerge.]  

(d) [Graph showing TMRE intensity (AU) for hsp-6.]  

(e) [Graph showing fold increase in RFU for atfs-1 (null) and atfs-1 (RNAi).]

(f) [Image showing ATFS-1 and Tub with WT and rsks-1 (ok1255) conditions.]  

(g) [Image showing GFP, TMREMerge, and mGFP mtGFP conditions.]  

(h) [Image showing control and atfs-1 (RNAi) conditions, with GFP and TMREMerge.]  

(i) [Graph showing TMRE intensity (AU) for atfs-1 (RNAi).]  

(j) [Graph showing fold change in mtDNA for GFP and mGFP.]  

(k) [Graph showing fold increase in RFU for timm-23 and hsp-6.]  

(l) [Graph showing fold increase in RFU for atfs-1 (null) and atfs-1 (RNAi).]  

(m) [Image showing control and atfs-1 (RNAi) conditions, with GFP and TMREMerge.]  

(n) [Graph showing TMRE intensity (AU) for atfs-1 (RNAi).]  

(o) [Image showing myo-3p::mtgfp and atfs-1 (RNAi) conditions, with GFP and TMREMerge.]  

(p) [Graph showing TMRE intensity (AU) for atfs-1 (RNAi).]  

(q) [Image showing myo-3p::mtgfp and atfs-1 (null) conditions, with GFP and TMREMerge.]
Fig 4. TORC1-mediated protein synthesis promotes ATFS-1 activation.

a-b. Transcript levels of heat shock protein-6 (hsp-6) (a) and of activated transcription factor stress-1 (atfs-1) (b) as determined by qRT-PCR in wildtype and rsks-1(ok1255) worms. N=4. Error bars mean +/- s.d, *p<0.05, ***p<0.001 (Student’s t-test).

c. hsp-6pr::gfp and aak-2(rr48);hsp-6pr::gfp worms raised on control, let-363 or atfs-1 (RNAi). Scale bar 0.1 mm.

d-e. Transcript levels of heat shock protein-6 (hsp-6) as determined by qRT-PCR in wildtype and aak-2(rr48) worms N=3 (d) and in wildtype, agd383 and agd383;atfs-1(et18) strains N=5 (e). Error bars mean +/- s.d, *p<0.05, **p<0.01 (Student’s t-test).

f. SDS-Page immunoblots of wildtype and rsks-1(ok1255) worms, raised on control or lonp-1(RNAi). Tubulin (Tub) was used as a loading control.

g. atfs-1(et18);hsp-6pr::gfp worms raised on control or rsks-1(RNAi). Scale bar 0.1mm.

h. TMRE staining (red) of worms expressing myo-3pr::gfp or myo-3pr::mtgfp. Scale bar 10 µm.

i. Quantification of TMRE intensity in muscle cells of worms expressing myo-3pr::gfp or myo-3pr::mtgfp. N=34 (myo-3pr:: gfp), N=24 (myo-3pr:: mtgfp). Error bars mean +/- s.d, *p<0.05 (Student’s t-test).

j. Quantification of mtDNA in myo-3pr:: gfp and myo-3pr:: mtgfp worms as determined by qPCR. N=6. Error bars mean +/- s.d, *p<0.05 (Student’s t-test).

k-l. Transcript levels of translocase of inner mitochondrial membrane-23 (timm-23) (k) and of heat shock protein-6 (hsp-6) (l) as determined by qRT-PCR in myo-3pr::gfp and in myo-3pr::mtgfp worms. N=4. Error bars mean +/- s.d, *p<0.05, **p<0.01 (Student’s t-test).
m. TMRE staining of worms expressing myo-3pr::mtgfp raised on control or atfs-1(RNAi). Scale bar 10 µm.

n. Quantification of TMRE intensity in muscle cells of worms expressing myo-3pr::mtgfp raised on control or atfs-1(RNAi). N=18 (control), N=13 (atfs-1(RNAi)). Error bars mean +/- s.d, ***p<0.001 (Student’s t-test).

o. TMRE staining of wildtype and atfs-1(null) worms expressing myo-3pr::mtgfp. Scale bar 10 µm.

p. Quantification of TMRE intensity in muscle cells of wildtype, and atfs-1(null) worms. N=15 (wildtype), N=24 (atfs-1(null)). Error bars mean +/- s.d, **p<0.01 (Student’s t-test).

q. TMRE staining of wildtype and atfs-1R/R worms expressing myo-3pr::mtgfp. Scale bar 10 µm.