Transgenic sensors reveal compartment-specific effects of aggregation-prone proteins on subcellular proteostasis during aging

Graphical abstract

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In brief

Curley et al. develop transgenic sensors of subcellular proteostasis in Drosophila based on misfolding-prone Fluc^{DM} protein variants that localize to the cytoplasm, nucleus, and mitochondria. They demonstrate that these tools can be used to monitor how cellular stress, interorgan signaling, and aggregation-prone proteins impact subcellular proteostasis during aging in Drosophila.

Highlights

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- **e** Establishes transgenic sensors of subcellular proteostasis in Drosophila
- Demonstrates their use to monitor proteostasis in different cell compartments
- Examines how inter-organ signaling impacts subcellular proteostasis
- \bullet Shows that pathogenic tau^{V337M} differentially remodels proteome solubility

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Transgenic sensors reveal compartment-specific effects of aggregation-prone proteins on subcellular proteostasis during aging

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MOTIVATION Eukaryotic cells have evolved complex functions via their compartmentalization into organelles and subcellular microenvironments. Such specialization of the cellular space enabled optimal function but also resulted in compartment-specific challenges to proteostasis. However, due to the paucity of tools, how subcellular proteostasis is regulated in physiological and pathological conditions remains largely uncharted. In this study, we generated *Drosophila* strains that ubiquitously express misfolding-prone Fluc^{DM} protein variants that have been engineered to localize to the nucleus and mitochondria, along with an untargeted variant that localizes primarily to the cytoplasm. By validating these tools in several contexts, we demonstrate that compartment-targeted Fluc^{DM} variants can be used to either perturb or to monitor subcellular proteostasis, depending on the experimental design.

SUMMARY

Loss of proteostasis is a hallmark of aging that underlies many age-related diseases. Different cell compartments experience distinctive challenges in maintaining protein quality control, but how aging regulates subcellular proteostasis remains underexplored. Here, by targeting the misfolding-prone Fluc^{DM} luciferase to the cytoplasm, mitochondria, and nucleus, we established transgenic sensors to examine subcellular proteostasis in Drosophila. Analysis of detergent-insoluble and -soluble levels of compartment-targeted Fluc^{DM} variants indicates that thermal stress, cold shock, and pro-longevity inter-organ signaling differentially affect subcellular proteostasis during aging. Moreover, aggregation-prone proteins that cause different neurodegenerative diseases induce a diverse range of outcomes on Fluc^{DM} insolubility, suggesting that subcellular proteostasis is impaired in a disease-specific manner. Further analyses with Fluc^{DM} and mass spectrometry indicate that pathogenic tau^{V337M} produces an unexpectedly complex regulation of solubility for different Fluc^{DM} variants and protein subsets. Altogether, compartment-targeted Fluc^{DM} sensors pinpoint a diverse modulation of subcellular proteostasis by aging regulators.

INTRODUCTION

Age-related diseases arise from interconnected, degenerative events that occur at multiple levels in organs, tissues, cells, and subcellular compartments. $1-5$ Although the decline of proteostasis is a defining feature of many age-related diseases, $6-8$ little is known about how protein quality control is regulated in different cellular organelles and compartments during aging. $1-3$ Because each cellular compartment/organelle is characterized by its folding environment and proteolytic capacities, disease

processes and environmental stressors that challenge protein quality control may impact each organelle differently.¹

The previous development of tools in *C. elegans* and cell culture has provided remarkable insight into this subject. For example, redox reporters targeted to the endoplasmic reticulum (ER) and the cytoplasm indicate that the redox state of these cell compartments is profoundly remodeled during aging in *C. elegans*. Specifically, the ER is oxidizing in young animals but shifts toward reducing conditions during aging, whereas the cytosol becomes more oxidizing.^{[9](#page-16-3)} Another study used a general (i.e., cytoplasmic)

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Figure 1. Generation of compartment-targeted, misfolding-prone Fluc^{DM} variants to perturb and sense subcellular proteostasis in **Drosophila**

(A) Generation of transgenic organelle-targeted sensors of protein quality control based on a misfolding-prone mutant firefly luciferase (Fluc^{DM}) fused to EGFP. The mitochondrial targeting sequence from the human mitochondrial COX VIII protein was utilized to generate the mito-Fluc^{DM} variant, whereas a standard nuclear targeting sequence was used to generate the NLS-Fluc^{DM}. General (untargeted) sensors and reporters for mitochondria and the nucleus were site integrated and are expressed ubiquitously (downstream of a tubulin promoter) and at similar levels, as indicated by qRT-PCR with 3 batches of flies and the mean ± SD (no significant changes, one-way ANOVA).

(B) Immunostaining and confocal microscopy of enterocytes indicate that Fluc^{DM} variants exhibit the expected specificity in subcellular localization. General (untargeted) Fluc^{DM} is detected in the cytoplasm (but also in the nucleus and plasma membrane), mito-Fluc^{DM} is detected in ATP5A-stained mitochondria, and NLS-Fluc^{DM} is found in the nucleus. Scale bars represent 20 and 15 μ m, as indicated.

(C) Immunostaining and confocal microscopy of brain cells from the antennal lobe indicates a similar localization. The untargeted Fluc^{DM} is detected in the cytoplasm, mito-Fluc^{DM} is detected in ATP5A-positive mitochondria, and NLS-Fluc^{DM} is found in the nucleus. Scale bars represent 10 and 5 um.

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sensor of proteostasis based on a structurally destabilized firefly luciferase mutant (Fluc^{DM}) and found that aging leads to an increase in sensor insolubility in *C. elegans*. [10](#page-16-4) Organelle-targeted sensors can also be used to identify organelle-specific components of proteostasis. For example, by using model substrates targeted to the nucleus and the cytoplasm, studies in yeast have unveiled that proteostasis in these compartments is regulated by distinct sets of proteins and that the ubiquilin Dsk2 is required to clear nuclearly misfolded proteins.^{[11](#page-16-5)} Moreover, compartmentspecific sensors of proteostasis may also be used to monitor inter-organelle crosstalk, which is a component of age-related processes. $1,2,12-14$ $1,2,12-14$ $1,2,12-14$ In addition to working as sensors, previous studies have found that misfolding-prone proteins^{15,[16](#page-17-1)} and organelle-targeted reporters such as mito-GFP¹⁷ can impact the specific compartment where they are targeted and induce adaptive and maladaptive stress responses. $18-21$ On this basis, depending on the experimental design, organelle-targeted reporter proteins may be utilized both as sensors of proteostasis or as probes to perturb proteostasis in a specific subcellular compartment.

Here, we expanded the toolkit to study subcellular proteostasis by generating transgenic *Drosophila* strains that express misfolding-prone firefly luciferase Fluc^{DM} variants that localize to the mitochondria, the nucleus, and the cytoplasm. Fluc^{DM} was chosen as the starting point for generating sensors of subcellular proteostasis because of the previous studies demonstrating its utility in other model organisms $10,22$ $10,22$ and because it is an exogenous protein that may be less likely to perturb cellular functions compared to the transgenic expression of an endogenous misfolding-prone protein. By comparing the impact of experimental versus control interventions in the presence of each Fluc^{DM}, we find that $Fluc^{DM}$ variants can be utilized as sensors of proteostasis. By monitoring the levels of detergent-soluble and -insoluble $Fluc^{DM}$, we find that subcellular proteostasis is differentially regulated by environmental stress, endocrine signaling, and aggregation-prone proteins (such as microtubule-associated protein tau [MAPT]) during aging in *Drosophila*. Moreover, we find that these tools can be utilized to perturb proteostasis in a compartment-specific manner when a Fluc^{DM} variant is compared to an isogenic control with no Fluc^{DM} expression. Altogether, these studies indicate that organelle-targeted misfolding-prone proteins may find several experimental applications and that such tools may provide insight into how subcellular proteostasis is differentially cross-regulated in distinct compartments by age-related diseases and anti-aging interventions.

RESULTS

Establishment of transgenic, compartment-targeted, misfolding-prone Fluc^{DM} proteins to sense and perturb subcellular proteostasis in Drosophila

 $Fluc^{DM}$ is a misfolding-prone firefly luciferase protein variant that has been used as a reporter of cytoplasmic proteostasis in cell culture and in *C. elegans*[10](#page-16-4) and, more recently, *in vivo* in

mice.^{[22](#page-17-4)} Detergent-soluble Fluc^{DM} is functional, whereas sequestration of Fluc^{DM} into detergent-insoluble fractions (which typically correspond to protein aggregates) indicates the occurrence of misfolding and the consequent loss of solubility. In addition to increased misfolding, retention of Fluc^{DM} into detergent-insoluble fractions can also indicate deficits in the degradation of misfolded proteins. In line with this model, aging increases $Fluc^{DM}$ insolubility in *C. elegans*, [10](#page-16-4) confirming that cytoplasmic proteo-stasis declines with aging.^{[6,](#page-16-1)[7](#page-16-7),[10,](#page-16-4)[23](#page-17-5)}

Based on the previously characterized misfolding-prone lucif-erase,^{[10](#page-16-4)} we generated compartment-targeted Fluc^{DM} variants tagged with enhanced green fluorescent protein (EGFP) for *in vivo* use in *Drosophila* [\(Figure 1](#page-2-0)A). To generate a Fluc^{DM} variant targeted to mitochondria (mito-Fluc^{DM}-EGFP), the mitochondrial import sequence from the human mitochondrial COX VIII protein (cytochrome c oxidase subunit 8, a component of the respiratory chain located in the inner mitochondrial membrane) $24,25$ $24,25$ was fused to the N terminus of Fluc^{DM}-EGFP. A standard nuclear tar-geting sequence (PAAKRVKLD)^{[26](#page-17-8)} was fused to the Fluc^{DM} N ter- $\frac{3}{2}$ minus to generate a Fluc^{DM}-EGFP variant with a nuclear localization signal (NLS-Fluc^{DM}-EGFP). Lastly, transgenic flies were also generated for untargeted Fluc^{DM}-EGFP ([Figure 1](#page-2-0)A), which has been found previously to display diffuse cytoplasmic localization in *C. elegans* and cell culture.^{[10](#page-16-4)}

To ensure expression in the physiological range, all Fluc^{DM} variants were engineered to be expressed downstream of a ubiquitous tubulin promoter and to be site integrated at the same position into the genome via the phiC31 integrase system, 27 leading to similar expression levels, as assessed by qRT-PCR [\(Figure 1A](#page-2-0)).

Next, immunostaining and confocal microscopy were utilized to determine the subcellular localization of Fluc^{DM} protein variants. Considering that the tubulin promoter drives the expression of Fluc^{DM} transgenes ubiquitously, Fluc^{DM} intracellular localization was monitored in enterocytes, which are a convenient system for these analyses given their large cell size, and in brain cells because of the importance of proteostasis for neurodegeneration.^{[7](#page-16-7)} Confocal microscopy indicates that Fluc^{DM} variants exhibit distinct subcellular localization [\(Figures 1B](#page-2-0) and 1C): general (untargeted) Fluc^{DM} is detected in the cytoplasm, nucleus, and plasma membrane of enterocytes ([Figure 1](#page-2-0)B) but only in the cytoplasm of brain cells [\(Figure 1](#page-2-0)C); NLS-Fluc^{DM} is detected in the nucleus of both en-terocytes and brain cells ([Figures 1B](#page-2-0) and 1C); and mito-Fluc^{DM} is detected in mitochondria, defined by co-staining for the mitochon-drial marker ATP5a (ATP synthase F1 subunit alpha) [\(Figures 1](#page-2-0)B and 1C). However, not all mito-Fluc^{DM} is detected in mitochondria, presumably because mito- $Fluc^{DM}$ is synthesized in the cytosol, and its import into mitochondria is suboptimal. Altogether, we established transgenic *Drosophila* strains that express general (i.e., cytosolic) and compartment-targeted Fluc^{DM} variants.

It has been found previously that heat shock results in the acc-umulation of cytoplasmic aggregates of Fluc^{DM} in cell culture.^{[10](#page-16-4)} On this basis, we tested whether the intracellular localization of

⁽D) Immunostaining of enterocytes from heat-shocked and control flies identifies Fluc^{DM}-GFP aggregates that accumulate in the cytoplasm in response to thermal stress compared to non-heat-shocked controls. Similar heat-induced cytoplasmic aggregates are also found in heat-shocked NLS-Fluc^{DM}-EGFP and mito-Fluc^{DM}-EGFP cells. In the case of mito-Fluc^{DM}-EGFP, these aggregates are recognizable because they produce larger puncta than the staining that corresponds to mito-Fluc^{DM}-EGFP-positive mitochondria (B). Scale bar, 10 μ m.

compartment-targeted Fluc^{DM} variants is remodeled by thermal stress. To this purpose, we examined the subcellular localization of the EGFP-tagged Fluc^{DM} variants by immunostaining the enterocytes of heat-shocked flies and controls with anti-GFP antibodies. In agreement with previous findings in cell culture, 10 the diffuse localization of the general Fluc^{DM}-EGFP sensor shifted to a spotty pattern upon heat shock, indicative of the sequestration of a fraction of the Fluc^{DM}-EGFP into cytoplasmic aggregates ([Figure 1](#page-2-0)D). Similar cytoplasmic aggregates were also found in the heat-shocked NLS-Fluc^{DM}-EGFP and mito-Fluc^{DM}-EGFP cells [\(Figure 1](#page-2-0)D). In the case of mito-Fluc^{DM}-EGFP, these aggregates were recognized because they were bigger than the speckles that correspond to mitochondria with mito-Fluc^{DM}-EGFP [\(Figure 1D](#page-2-0)), which largely co-localizes with the mitochon-drial marker ATP5a ([Figures 1B](#page-2-0) and 1C). Altogether, these findings indicate that thermal stress induces the sequestration of a fraction of the $Fluc^{DM}$ variants into cytoplasmic aggregates ([Figure 1D](#page-2-0)).

Organism-wide perturbation of subcellular proteostasis with compartment-targeted Fluc^{DM} variants extends lifespan and delays neuromuscular aging

Protein misfolding restricted to a specific cell compartment has been found previously to induce adaptive and maladaptive responses locally and even systemically.^{[16](#page-17-1)[,18–21](#page-17-3)} For example, the expression of a metastable sarcomeric protein in *C. elegans* skeletal muscle induces a local and systemic stress response characterized by transcriptional induction of the chaperone *Hsp90*. [15](#page-17-0) Moreover, it was recently appreciated that expression of mito-GFP, a GFP with a mitochondrial import sequence that is utilized to identify mitochondria by microscopy, leads to perturbation of mitochondrial proteostasis and induc-tion of the mitochondrial unfolded protein response.^{[17](#page-17-2)} On this basis, with an appropriate experimental design, Fluc^{DM} variants may be utilized to perturb proteostasis in a subcellular compartment and determine the corresponding stress responses induced by such compartment-restricted disruption of proteostasis compared to controls with no Fluc^{DM}.

To determine the overarching results of perturbing proteostasis in distinct subcellular compartments at the organism level, we examined whether Fluc^{DM} variants regulate lifespan and neuromuscular function during aging compared to isogenic controls with no Fluc^{DM} expression. Survival analyses indicate that Fluc^{DM} variants extend lifespan, although this occurs differently for each compartment-targeted Fluc^{DM}: at 18°C and 25°C, the lifespan extension is more pronounced for the untargeted (i.e., cytoplasmic) Fluc^{DM} and for mito-Fluc^{DM} compared to NLS-Fluc^{DM} ([Figures 2A](#page-5-0) and 2B). However, all Fluc^{DM} variants similarly extend lifespan at 29° C (Figure $2C$), a temperature that corresponds to mild chronic thermal stress for *Drosophila*. Interestingly, 2 copies of the mito-Fluc^{DM} and NLS-Fluc^{DM} transgenes were not more effective in extending lifespan than single copies, suggesting that an optimal adaptive response is already reached with 1 copy and that additional copies of the transgene may not further induce the response or even be detrimental [\(Figures 2A](#page-5-0)-2C).

We next examined negative geotaxis; i.e., the startle-induced escape response from gravity.^{28–30} This is an innate behavior that depends on the function of the brain and skeletal muscle

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and that senesces with aging in *Drosophila*. [28–30](#page-17-10) These studies indicate that the percentage of flies able to climb progressively decreases with aging in control flies but less so in isogenic flies that express untargeted Fluc^{DM}, mito-Fluc^{DM}, and NLS-Fluc^{DM} at 18°C, 25°C, and 29°C ([Figures 2D](#page-5-0)–2F). Altogether, these findings indicate that organism-wide expression of a misfolding-prone protein targeted to distinct cell compartments induces, to varying degrees, a hormetic response that extends lifespan and preserves neuromuscular function during aging in *Drosophila*.

It has been shown previously that metastable proteins can trigger an adaptive response that partially preserves or even re-stores protein quality control.^{[18–21](#page-17-3)} On this basis, we next tested whether Fluc^{DM} variants regulate proteostasis in a Huntington's disease model based on the expression of GFP-tagged pathogenic huntingtin in the retina and in which Htt-polyQ72-GFP aggregates progressively form with aging.^{31–34} Compared with control mCherry and with no transgene expression, all Fluc^{DM} variants significantly reduced Htt-polyQ72-GFP aggregates [\(Figure 2G](#page-5-0)), suggesting that the stress response induced by Fluc^{DM} improves proteostasis compared to control flies with no $Fluc^{DM}$ expression.

In summary, these studies indicate that compartmenttargeted Fluc^{DM} variants can be utilized to perturb subcellular proteostasis and to identify the consequent hormetic stress responses that are induced compared to isogenic controls with no Fluc^{DM} expression.

Transcriptional stress responses induced by organelletargeted perturbation of proteostasis with Fluc^{DM} variants

Having established that Fluc^{DM} variants induce a non-canonical stress response at the organismal and cellular levels [\(Figure 2](#page-5-0)). we next sought to define the underlying transcriptional responses. To this purpose, RNA sequencing (RNA-seq) was done from Fluc^{DM} variants and isogenic control flies reared at 18°C, 25°C, and 29°C [\(Table S1\)](#page-16-8).

Heatmaps of the genes with the highest *Z* scores indicate that Fluc^{DM} flies display substantial gene expression changes compared to isogenic controls with no Fluc^{DM} [\(Figure 3A](#page-6-0)). Analysis of the Gene Ontology (GO) terms that are modulated by Fluc^{DM} include several categories involved in proteostasis and aging, such as proteases, lipid metabolism, and innate immunity [\(Figure 3B](#page-6-0)). Examples of differentially regulated genes include the mitochondrial heat shock protein 22 (Hsp22), 35 which is among the most highly induced genes in all $Fluc^{DM}$ strains compared to isogenic controls: while *Hsp22* expression normally rises with heat from 18°C to 29°C ([Figure 3C](#page-6-0)), Fluc^{DM} further significantly increases its levels both at 25°C and 29°C. Because previous studies have found a key role for Hsp22 in lifespan regu-lation,^{[35](#page-17-12)} these findings suggest that Hsp22 is part of the Fluc^{DM}induced stress response that extends lifespan and improves neuromuscular function ([Figure 2](#page-5-0)). Consistent with this model, NLS-Fluc^{DM}, which improved climbing and lifespan less than Fluc^{DM} and mito-Fluc^{DM} [\(Figure 2\)](#page-5-0), induced *Hsp22* expression less than the other $Fluc^{DM}$ variants ([Figure 3](#page-6-0)C). Additional Fluc^{DM}-regulated genes include the APP (amyloid precursor protein)-cleaving beta-secretase *Bace* (which is downregulated by FlucDM) and the ubiquitin ligase *CG32581/sordd2* (suppression of retinal degeneration disease 1 upon overexpression 2), which

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Figure 2. Use of compartment-targeted Fluc^{DM} variants to perturb subcellular proteostasis

Fluc^{DM} are misfolding-prone proteins and, therefore, they may challenge proteostasis in the subcellular compartment to which they are targeted. On this basis. Fluc^{DM} variants can be used as tools to induce a moderate perturbation of subcellular proteostasis when compared to an isogenic control with no Fluc^{DM} expression.

 $(A-C)$ The untargeted (cytoplasmic) Fluc^{DM}, the mitochondrially targeted mito-Fluc^{DM}, and the nucleus-targeted NLS-Fluc^{DM} extend lifespan ($p < 0.001$, log -rank test, with *n* indicated) when compared to isogenic controls with no Fluc^{DM}. Distinct Fluc^{DM} variants have effects of different magnitudes; the untargeted Fluc^{DM} is more effective in extending lifespan at 18°C (A) and at 25°C (B) compared to NLS-Fluc^{DM}, whereas similar lifespan extension is seen at 29°C (C) for all Fluc^{DM} variants. These findings suggest that moderate perturbation of subcellular proteostasis by Fluc^{DM} variants induces a stress response that extends lifespan. A single (orange) or 2 copies (gray) of the Fluc^{DM} transgenes similarly extend lifespan compared to the isogenic controls with no Fluc^{DM} (blue).

(D–F) Negative geotaxis assays indicate that expression of misfolding-prone Fluc^{DM} proteins targeted to distinct subcellular compartments reduces agerelated neuromuscular dysfunction during aging compared to isogenic controls that do not express Fluc^{DM}. A similar protection is found at 18°C (D), 25°C (E), and 29 C (F). These findings suggest that moderate perturbation of subcellular proteostasis by compartment-targeted Fluc^{DM} variants induces an adaptive stress response that improves neuromuscular function; *p* < 0.001 (log -rank test) with *n* indicated (A–C).

(G) Aggregates of GFP-tagged pathogenic huntingtin-polyQ can be seen in the retina of *GMR>Htt-polyQ72-GFP* flies at 30 days of age, but the total area of such aggregates is higher in controls (mCherry and no transgene, +) compared to flies that express $Fluc^{DM}$ variants. This suggests that moderate stress induced by $Fluc^{DM}$ can induce a hormetic stress response that improves proteostasis. The *n* (biological replicates) and the mean ± SD are indicated, with **p* < 0.05 and ***p* < 0.01 (one-way ANOVA).

is homologous to human RNF185 (ring finger protein 185) and is highly induced by all $Fluc^{DM}$ variants [\(Figure 3](#page-6-0)C).

Further analysis of the transcriptional responses induced by mito-Fluc^{DM} and NLS-Fluc^{DM} compared to the untargeted Fluc^{DM} revealed several gene categories that are differentially regulated by distinct Fluc^{DM} proteins: lipid metabolism and antimicrobial peptides were commonly induced by mito-Fluc^{DM} and NLS-Fluc^{DM} compared to the untargeted Fluc^{DM}, whereas components of the ER and gustatory receptors were enriched categories among the genes regulated by mito-Fluc^{DM} and by NLS-FlucDM, respectively ([Figure 3](#page-6-0)D). An example of gene that is differentially regulated by distinct Fluc^{DM} variants is the triacylglycerol lipase *CG11598*, which is homologous to human LIPN (lipase family member N) and related lipases: *CG11598* is more highly expressed in NLS-Fluc^{DM} versus mito-Fluc^{DM} and even more so compared to the untargeted Fluc^{DM} [\(Figure 3](#page-6-0)E). Altogether, transcriptional profiling identifies differentially expressed genes that are induced consistently or distinctly by Fluc^{DM} variants compared to isogenic controls with no Fluc^{DM} expression.

Compartment-targeted Fluc^{DM} sensors indicate that heat and cold shock distinctly impact subcellular proteostasis

In addition to perturbing proteostasis in subcellular compart-ments ([Figures 2](#page-5-0) and [3](#page-6-0)), Fluc^{DM} variants are primarily utilized as sensors of proteostasis, as found previously for a general (i.e., cytoplasmic) FlucDM in *C. elegans*[10](#page-16-4) and for compartment-targeted Fluc^{DM} in mice.²² To test whether this is also possible in

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Figure 3. Compartment-targeted Fluc^{DM} variants induce a transcriptional stress response compared to isogenic controls with no Fluc^{DM} (A) Heatmap of the genes that are most differentially regulated indicate that Fluc^{DM} variants induce transcriptional adaptive responses that differentiate them from isogenic controls with no FlucDM expression. The average *Z* scores are color coded; upregulated genes are shown in red and downregulated genes in blue. (B) Several gene categories are commonly modulated by compartment-targeted Fluc^{DM} variants, including genes that encode for proteins with monooxygenase activity, proteases, and proteins involved in lipid catabolism.

(C) Differentially modulated genes such as *Hsp22*, *Bace*, and *CG32581/sordd2* may provide a mechanistic explanation for how stress responses induced by perturbation of subcellular proteostasis are protective during aging [\(Figure 2\)](#page-5-0).

(D) Apart from transcriptional stress responses that are commonly induced by all Fluc^{DM} variants (B and C), there are also gene categories that are differentially regulated by mito-Fluc^{DM} and by NLS-Fluc^{DM} versus the untargeted Fluc^{DM}.

(E) An example of a gene that is differentially modulated by distinct Fluc^{DM} variants is the triacylglycerol lipase CG11598, which is not regulated by NLS-Fluc^{DM}, is downregulated by mito-Fluc^{DM}, and even more downregulated by the untargeted Fluc^{DM} compared to isogenic controls with no Fluc^{DM} expression.

In (C) and (E), $n = 3$ (biological replicates) with the mean \pm SD indicated; for each genotype, the mRNA levels (transcripts per million [TPM]) at 18°C (gray), 25°C (yellow), and 29°C (orange) are shown. For the genes in (C) and (E), the color-coded *p* value summary (one-way ANOVA) is reported for the comparison of Fluc^{DM} variants to the control at each temperature. Additional statistical cross-comparisons of each Fluc^{DM} to the control at each temperature are reported in the source data file [\(Table S4\)](#page-16-8).

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Drosophila, two environmental stressors (cold and heat shock) were utilized to determine how they impact subcellular proteostasis *in vivo*. While thermal stress (heat shock) is a well-known challenge to proteostasis because of heat-induced protein unfolding,^{36–38} cold shock has been conversely found to improve proteostasis and delay aging.^{39–43} However, the impact of cold and heat shock on subcellular proteostasis remains largely unexplored. To address this question, Fluc^{DM} strains were exposed to heat stress (36 $^{\circ}$ C for 3 h) or cold shock (4 $^{\circ}$ C for 2 h) and compared to isogenic controls kept at the standard temperature of 25° C. Subsequently, the solubility of the untargeted (i.e., cytoplasmic) Fluc^{DM}, the mitochondrial mito-Fluc^{DM}, and the nuclear NLS-Fluc^{DM} was analyzed via western blot of detergent-soluble and -insoluble fractions. As expected based on its capacity to cause protein unfolding, $36,37$ $36,37$ thermal stress induced profound changes in Fluc^{DM} solubility that were, however, of different magnitudes in distinct subcellular compartments. In general, there was a decrease in the soluble levels of all Fluc^{DM} variants and a corresponding increase in their detergent-insoluble levels; these heat-induced changes in protein insolubility were the highest for the untargeted (i.e., cytoplasmic) Fluc^{DM}, intermediate for the nuclear NLS-Fluc^{DM}, and minimal for the mitochondrial mito-Fluc^{DM} ([Figure 4](#page-7-0)A). Altogether, these findings suggest that certain cellular compartments (e.g., mitochondria) may be relatively resilient to heat-induced protein unfolding compared to the nucleus and the cytoplasm.

Cold shock impacts the cell differently than thermal stress.³⁹⁻⁴³ Consistent with this scenario, western blots indicated that there were no changes in the soluble levels of any Fluc^{DM} variants, whereas the detergent-insoluble levels were conversely regulated depending on the compartment [\(Figure 4B](#page-7-0)): detergent-insoluble levels of the (cytoplasmic) Fluc^{DM} were reduced by cold shock but trended toward increasing for NLS-Fluc^{DM}, whereas minimal changes were found for mito-Fluc^{DM} ([Figure 4](#page-7-0)B). Altogether, the folding status of compartment-targeted Fluc^{DM} variants indicates that proteostasis is affected differentially by heat and cold shock in distinct subcellular compartments.

The solubility of compartment-targeted Fluc^{DM} proteins is correspondingly regulated by cytoplasmic and mitochondrial chaperones

Proteostasis in distinct subcellular compartments critically relies on dedicated proteolytic systems and chaperones, such as the

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mitochondrial chaperone Hsp22 and the cytoplasmic chaperone Hsp70.^{[35,](#page-17-12)[44–46](#page-17-16)} On this basis, we tested whether modulation of Hsp22 and Hsp70 impacts the folding status of Fluc^{DM} variants. Knockdown of Hsp70 resulted in an \sim 70% decline in Hsp70 mRNA levels [\(Figure S1](#page-16-8)A) and in a significant decline in the detergent-soluble levels of the untargeted (i.e., cytoplasmic) Fluc^{DM} [\(Figure S1](#page-16-8)B), whereas there were no changes in the detergentsoluble and -insoluble levels of mito-Fluc^{DM} and NLS-Fluc^{DM} [\(Figures S1](#page-16-8)C and S1D). Moderate $(\sim 3$ -fold) overexpression of the mitochondrial chaperone *Hsp22* [\(Figure S1](#page-16-8)A) did not significantly affect the detergent-soluble and -insoluble levels of Fluc^{DM} and NLS- Fluc^{DM} ([Figures S2A](#page-16-8) and S2C) but significantly reduced the levels of detergent-insoluble mito-Fluc^{DM} [\(Fig](#page-16-8)[ure S2](#page-16-8)B), consistent with the known role of Hsp22 in preserving mitochondrial proteostasis.^{[35](#page-17-12)} Altogether, these findings indicate that chaperones with specific subcellular localizations correspondingly impact the folding status of the Fluc^{DM} sensors that are targeted to the same subcellular compartment.

Previous studies in *C. elegans* have found that aging leads to an increase in Fluc DM sensor insolubility.^{[10](#page-16-4)} On this basis, we examined whether the detergent-soluble and -insoluble levels of Fluc^{DM} variants are modulated by aging in extracts from whole flies, heads (enriched for tissues of the central nervous system), and thoraces (enriched for skeletal muscle). In whole flies, there was a significant increase in the detergent-insoluble levels of the untargeted (i.e., cytoplasmic) Fluc^{DM} sensor with aging, whereas mito-Fluc^{DM}, and NLS-Fluc^{DM} were not modulated ([Figure S3](#page-16-8)). Analysis of extracts from heads and thoraces indicates that there are minimal or no changes in the insoluble levels of Fluc^{DM}, mito-Fluc^{DM}, and NLS-Fluc^{DM} during aging ([Figure S3](#page-16-8)).

Similar results were also found by monitoring the folding status of the firefly luciferase Fluc^{DM}-GFP sensors with luciferase assays: the signal for Fluc^{DM} does not significantly change with aging (10, 30, and 60 days) in fly thoraces. Moreover, the luciferase signal (normalized by total protein content) was higher for the general Fluc^{DM} compared to NLS-Fluc^{DM} and mito-Fluc^{DM}, suggesting that there could be detection limits that can impair the use of luciferase assays for monitoring the folding status of Fluc^{DM} variants [\(Figure S4\)](#page-16-8).

Altogether, these findings indicate that the folding status of Fluc^{DM} variants is relatively well preserved in the cytoplasm, nucleus, and mitochondria of aging cells in *Drosophila* in the absence of environmental shocks or genetic perturbations.

Figure 4. Use of compartment-targeted Fluc^{DM} as sensors of subcellular proteostasis

Fluc^{DM} are misfolding-prone proteins and, therefore, can be used to determine whether an experimental intervention modulates proteostasis compared to a control intervention. Specifically, Fluc^{DM} protein misfolding leads to its detergent insolubility, which indicates defects in protein quality control.

(A) Testing compartment-targeted Fluc^{DM} with thermal stress (red) versus controls (gray). Western blots of detergent-soluble and -insoluble fractions from whole flies with anti-GFP antibodies detect the EGFP-tagged Fluc^{DM} variants targeted to the mitochondria (mito-Fluc^{DM}) and the nucleus (NLS-Fluc^{DM}) and the untargeted Fluc^{DM} that localizes primarily to the cytoplasm. Ponceau staining and ß-actin are shown as normalization controls, and the graphs refer to the GFP levels normalized by β-actin. Heat shock significantly decreases the detergent-soluble levels of all compartment-targeted Fluc^{DM}. This corresponds to an increase in the detergent-insoluble levels of the untargeted (i.e., cytoplasmic) Fluc^{DM}, NLS-Fluc^{DM}, and, to a lower extent, mito-Fluc^{DM}. Altogether, these findings indicate that thermal stress compromises proteostasis across the cell but more prominently in the cytoplasm and nucleus compared to the mitochondria. $n = 3$ (biological replicates) with the mean \pm SD indicated; γp < 0.05 and $\pi\gamma p$ < 0.001 (unpaired two-tailed t test).

(B) Testing compartment-targeted Fluc^{DM} with cold shock (blue) versus controls (gray). Shown are western blots of detergent-soluble and -insoluble fractions from whole flies with anti-GFP and anti-ß-actin antibodies. Cold shock reduces the detergent-insoluble levels of the untargeted (i.e., cytoplasmic) Fluc^{DM}, whereas there are no changes in the detergent-soluble levels and in the detergent-insoluble levels of mito-Fluc^{DM} and NLS-Fluc^{DM}. Altogether, these findings indicate that cold shock improves cytoplasmic proteostasis but does not impact protein quality control in the nucleus and mitochondria. *n* = 3 (biological replicates) with the mean \pm SD indicated; ** p < 0.01 (unpaired two-tailed t test).

Modulation of subcellular proteostasis by endocrine signaling during aging

Aging is characterized by changes in multiple organ systems that collectively decrease the ability of the organism to maintain homeostasis. [6](#page-16-1),[47–49](#page-17-17) Previous studies have shown that inter-organ endocrine signaling regulates systemic aging and lifespan.[50–59](#page-18-0) Skeletal muscle has emerged as an important tissue in the systemic regulation of aging because of its capacity to secrete a variety of signaling factors (myokines) that contribute to inter-organ signaling in response to exercise, stress, and nutrient sensing.^{[60–68](#page-18-1)} On this basis, we tested the function of the stress-induced amylase Amyrel, which regulates protein quality control in the central nervous system during aging via endocrine maltose/SLC45 signaling, ^{[68,](#page-18-2)[69](#page-18-3)} and the adipokinetic hormone (Akh), a glucagon-like hormone that antagonizes insu-lin signaling and delays aging.^{[70–72](#page-18-4)} In these studies, detergentsoluble and -insoluble fractions from whole flies were examined to determine the systemic outcome of muscle-expressed Akh. A ~4-fold overexpression of *Akh* in muscle [\(Figure S1A](#page-16-8)) did not impact the detergent-soluble and -insoluble levels of $Fluc^{DM}$ and mito-Fluc^{DM} [\(Figures 5](#page-10-0)A and 5B), apart from a significant reduction in the soluble levels of Fluc^{DM} in old age ([Figure 5A](#page-10-0)). However, overexpression of *Akh* in muscle significantly reduced the detergent-insoluble levels of NLS-Fluc^{DM} across all ages ([Figure 5C](#page-10-0)), indicating that muscle-derived Akh improves nuclear proteostasis. Altogether, these findings indicate that signaling by the glucagon-like hormone Akh has a compartment-specific effect on subcellular proteostasis during aging.

We next tested whether another endocrine regulator of aging, Amyrel/maltose signaling,^{[69](#page-18-3)} similarly impacts subcellular proteostasis. In these studies, detergent-soluble and -insoluble fractions from heads and thoraces were examined to determine the outcome of muscle-expressed Amyrel on the central nervous system (head extracts) or skeletal muscle (thoraces). Western blots of detergent-soluble and -insoluble fractions indicate that muscle-specific induction of Amyrel reduces the insoluble levels of the untargeted (i.e., cytoplasmic) $Fluc^{DM}$ in heads ([Figure 5D](#page-10-0)), whereas mito-Fluc^{DM} and NLS-Fluc^{DM} are not affected ([Figures 5E](#page-10-0) and 5F). These findings, therefore, indicate that Amyrel promotes proteostasis in the cytoplasm but not in the mitochondria and nuclei, which is consistent with the previous finding that Amyrel induces the expression of the cytoplasmic chaperone *Hsp23*. [69](#page-18-3) Apart from the preservation of cytoplasmic proteostasis in head extracts (i.e., central nervous system), there was no effect of muscle-induced Amyrel on muscle proteostasis, as assessed from the analysis of detergent-soluble and insoluble levels of Fluc^{DM}, mito-Fluc^{DM}, and NLS -Fluc^{DM} in thoracic fractions [\(Figure S5\)](#page-16-8). This is consistent with previous findings that have shown that overexpression of the maltose-producing enzyme Amyrel in skeletal muscle regulates protein quality control in the brain and retina but not in skeletal muscle during aging,^{[69](#page-18-3)} presumably because of the high expression of SLC45 maltose transporters in the brain versus the skeletal muscle.^{[69](#page-18-3)} Altogether, these findings indicate that anti-aging endocrine interventions may act by selectively improving proteostasis in specific subcellular compartments.

Divergent modulation of subcellular proteostasis by distinct aggregation-prone proteins

Aggregation-prone proteins are the culprits of several agerelated diseases. $23,73-77$ $23,73-77$ Although they are primarily localized in a specific cell compartment, aggregation-prone proteins perturb the function of many organelles.^{1-3[,14](#page-16-9)[,78](#page-18-6)[,79](#page-18-7)} However, it remains largely unknown whether such changes in cell function arise from corresponding changes in the protein quality control of subcellular compartments. To address this question, we monitored how the solubility of Fluc^{DM}, mito-Fluc^{DM}, and NLS-Fluc^{DM} is modulated by aggregation-prone proteins in skeletal muscle, an abundant tissue in which proteostasis is compromised with aging. $8,32,33,63$ $8,32,33,63$ $8,32,33,63$ $8,32,33,63$ In particular, we utilized flies that express a protein with 49(GGGGCC) repeats that model those found in human C9orf72 and that are associated with the etiology of amyotrophic lateral sclerosis and frontotemporal dementia.^{[80](#page-18-9)} Such a protein with 49(GGGGCC) repeats forms toxic cytoplasmic and nuclear protein aggregates^{[81](#page-18-10)} compared to a control non-toxic pro-tein with 8(GGGGCC) repeats^{[80](#page-18-9)} and to mCherry. Western blots of detergent-soluble and -insoluble fractions indicate that 49(GGGGCC) increases the insoluble levels of all $Fluc^{DM}$ reporters (Fluc^{DM}, mito-Fluc^{DM}, and NLS-Fluc^{DM)} compared to the non-toxic controls, 8(GGGGCC)-containing proteins, and/ or mCherry [\(Figures 6A](#page-11-0)–6C). Altogether, these findings indicate that a toxic protein with 49(GGGGCC) repeats compromises proteostasis widely across the cell in the cytoplasm, mitochondria, and the nucleus ([Figures 6A](#page-11-0)–6C).

In a second set of studies, we expressed a C-terminal fragment of the human Machado-Joseph disease/spinocerebellar ataxia type 3 protein with a 78-repeat polyglutamine tract expansion (hATXN3.tr-Q78), 82 which forms toxic aggregates that are primarily located in the nucleus.⁸³⁻⁸⁵ As a control, hATXN3.tr-Q78 was compared to a non-toxic hATXN3 with only 27 repeats (hATXN3.tr-Q27) 82 and to mCherry. These analyses indicate that hATXN3.tr-Q78 has no substantial effect on the solubility of the untargeted (i.e., cytoplasmic) $Fluc^{DM}$ reporter ([Figure 6D](#page-11-0)). There was minimal impact of hATXN3.tr-Q78 on the solubility of mito-Fluc^{DM} and NLS-Fluc^{DM}, and these effects were inconsistent when comparing hATXN3.tr-Q78 to distinct controls, hATXN3.tr-Q27 and mCherry ([Figures 6E](#page-11-0) and 6F). Altogether, pathogenic ataxin-3 seems to minimally impact the folding status of Fluc^{DM} sensors located in distinct cell compartments, suggesting that pathogenic ataxin-3 may have a relatively limited impact on subcellular proteostasis.

In summary, the findings indicate that distinct pathogenic proteins have strikingly different impacts on subcellular protein quality control: while a toxic protein with 49(GGGGCC) repeats widely disrupts proteostasis in the cytoplasm, nucleus, and mitochondria, pathogenic ataxin-3 has minimal influence on the folding status of $Fluc^{DM}$ proteins targeted to these compartments.

Compartment-specific effects of pathogenic tau on subcellar proteostasis during aging in Drosophila

Aggregation of MAPT deranges proteostasis, and it is a cause of Alzheimer's disease and related dementias.^{[86–88](#page-19-0)} While tau is a cytoplasmic protein, many other cell compartments, such as the nucleus and mitochondria, are perturbed by pathogenic

Figure 5. Organelle-targeted sensors of proteostasis indicate that distinct endocrine signaling factors differentially impact subcellular proteostasis

(A–C) Western blot analyses of detergent-soluble and insoluble fractions from control whole flies (*Mhc>+*, gray) and isogenic flies with muscle-specific overexpression of Akh (*Mhc>Akh*, blue), the *Drosophila* functional homolog of glucagon and related peptides. Western blotting with anti-GFP antibodies indicates the levels of Fluc^{DM}-EGFP sensors, whereas Ponceau staining and β -actin are used as normalization controls. Akh differentially impacts the detergent-insoluble levels of Fluc^{DM}-EGFP variants targeted to distinct cell compartments during aging. Akh reduces the detergent-insoluble levels of nuclearly localized NLS-Fluc^{DM} at all ages (C), whereas it does not substantially impact the detergent-insoluble levels of cytoplasmic Fluc^{DM} (A) and mitochondrial mito-Fluc^{DM} (B). The ages analyzed for each genotype are 10, 30, and 60 days. *n* = 3 (biological replicates) with the mean \pm SD indicated; $\gamma p < 0.05$, $*^{*}p$ < 0.01, $*^{*}p$ < 0.001 (unpaired two-tailed t test).

(D–F) Western blot analysis of detergent-soluble and insoluble fractions from head extracts (enriched for the CNS) from control flies (*Mhc>+*, gray) and isogenic flies with muscle-specific overexpression of Amyrel (*Mhc>Amyrel*, green), which preserves proteostasis via maltose/SLC45 signaling. Amyrel reduces the detergent-insoluble levels of the untargeted (i.e., cytoplasmic) Fluc^{DM} (D) but not of mito-Fluc^{DM} (E) or NLS- $Fluc^{DM}$ (F). Altogether, these findings indicate that musclederived Amyrel improves cytoplasmic proteostasis in the CNS. The ages analyzed for each genotype are 10, 30, and 45 days. $n = 3$ (biological replicates) with the mean \pm SD indicated; p < 0.05, α *y* < 0.01 (unpaired two-tailed t test).

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Figure 6. Distinct aggregation-prone proteins differentially impact subcellular proteostasis

(A–C) Analyses of detergent-soluble and insoluble fractions from skeletal muscle from 10-day-old flies that express a toxic, aggregation-prone protein with 49(GGGGCC) repeats (orange) compared to non-toxic controls, 8(GGGGCC)-containing proteins (gray), and/or mCherry (white). Western blots with anti-GFP antibodies detect the levels of Fluc^{DM}-EGFP, whereas Ponceau staining and α -tubulin are used as normalization controls. The toxic 49(GGGGCC) protein increases the detergent-insoluble levels of Fluc^{DM} variants that localize to the cytoplasm (A), mitochondria (B), and the nucleus (C), indicating that aggregationprone proteins with 49(GGGGCC) repeats generally disrupt proteostasis across multiple cell compartments. *n* = 3 (biological replicates) with the mean ± SD indicated; **p* < 0.05, ***p* < 0.01, ****p* < 0.001 (one-way ANOVA).

(D–F) Western blots of detergent-soluble and -insoluble fractions from skeletal muscles of flies that express pathogenic ataxin-3 with poly-glutamine tract expansion (hATXN3.tr-Q78; red) versus a non-pathogenic ataxin-3 (hATXN3.tr-Q27, gray) and mCherry controls (white). There is no modulation of detergentsoluble and -insoluble Fluc^{DM} levels (D), whereas the effects of hATXN3.tr-Q78 on mito-Fluc^{DM} (E) and NLS-Fluc^{DM} (F) levels are inconsistent when compared to the hATXN3.tr-Q27 versus the mCherry control. $n = 3$ (biological replicates) with the mean \pm SD indicated; *p < 0.05, **p < 0.01 (one-way ANOVA). Altogether, these findings indicate that distinct aggregation-prone proteins have strikingly different impacts on subcellular proteostasis.

tau.^{[7,](#page-16-7)[89–93](#page-19-1)} For example, tau deranges nucleocytoplasmic transport 94 and nuclear architecture and chromatin organization, $95-97$ and tau nuclear aggregates alter the composition and organiza-tion of nuclear speckles.^{[98](#page-19-4)} However, it remains largely unknown whether the effects of cytosolic tau on the nucleus and organelles derive from corresponding changes in the protein quality control of these cell compartments.

To address this question, we utilized knockin fly models^{[99,](#page-19-5)[100](#page-19-6)} in which the endogenous *Drosophila* tau was substituted by pathogenic human tau^{V337}, which carries a mutation that causes

frontotemporal dementia in humans.¹⁰¹⁻¹⁰³ Previous studies have shown that these knockin *Drosophila* models develop progressive signs of tauopathy during aging when heterozygous, including neurodegeneration and behavioral deficits in memory, locomotion, and sleep.^{[99](#page-19-5)[,100](#page-19-6)} On this basis, we tested whether tau^{V337M} impacts the detergent-soluble and -insoluble levels of Fluc^{DM}, mito-Fluc^{DM}, and NLS-Fluc^{DM} compared to isogenic controls. For these studies, flies with ubiquitous expression of tau^{V337M} and of Fluc^{DM} variants were examined at 10, 30, and 60 days, which correspond to young, intermediate, and old

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Figure 7. Pathogenic tau^{v337M} impacts subcellular proteostasis during aging in a contrasting manner

Western blots of detergent-soluble and -insoluble fractions from control flies (gray) and isogenic knockin flies that express human tau^{V337M} (green), a mutant tau that causes frontotemporal dementia in humans. Western blotting with anti-GFP antibodies indicates the levels of Fluc^{DM}-EGFP sensors, whereas Ponceau staining and β -actin are used as normalization controls.

(A–C) Analysis of detergent-soluble and -insoluble fractions from head extracts (enriched for tissues of the CNS) indicates that heterozygous tau^{V337M} mutations increase the detergent-insoluble levels of the cytoplasmic Fluc^{DM} reporter during aging (at 30 and 60 days) but not at a young age (10 days) compared to isogenic controls. There is little effect on the solubility of mito-Fluc^{DM} and NLS-Fluc^{DM}, indicating that tau^{V337M} reduces cytoplasmic proteostasis but has no substantial effect on mitochondrial and nuclear proteostasis in the CNS.

(D–F) Western blots of detergent-soluble and insoluble fractions from skeletal muscle (thoracic fractions) of flies with heterozygous tau^{V337M} mutations. There is no significant effect of pathogenic tau on Fluc^{DM} (D) and mito-Fluc^{DM} (E) levels, suggesting that the skeletal muscle may be resistant to tau^{V337M}-induced defects in cytoplasmic proteostasis (D) compared to the CNS (A). However, heterozygous tau^{Y337M} mutations significantly reduce the detergent-insoluble levels of nuclearly localized NLS-Fluc^{DM} (F), suggesting that pathogenic tau^{V337M} may derange protein import into the nuclei of skeletal muscle.

In (A)–(F), the ages analyzed for each genotype are 10, 30, and 60 days. *n* = 3 (biological replicates) with the mean ± SD indicated; **p* < 0.05, ***p* < 0.01 (unpaired two-tailed t test).

(G-I) Ultra-deep-coverage TMT mass spectrometry indicates that pathogenic tau^{y337M} divergently regulates the solubility of distinct protein sets. These proteomics analyses (18-plex TMT) are based on detergent-soluble and -insoluble fractions from isogenic knockin flies that express human tau^{WT} and pathogenic human tau $\frac{V337M}{N}$ at 10 and 60 days of age.

ages, and fly heads (enriched for the central nervous system) and thoraces (enriched for skeletal muscles) were analyzed separately.

Western blots of detergent-soluble and -insoluble fractions from fly heads revealed that, as expected, tau^{V337M} worsens cytosolic proteostasis, as indicated by a significant increase in the detergent-insoluble levels of the untargeted $(\sim$ cytosolic) Fluc^{DM}, whereas its soluble levels were not affected ([Figure 7](#page-12-0)A). Interestingly, these effects were significant at day 30 and day 60 but not at day 10 ([Figure 7](#page-12-0)A), consistent with the previous finding that the development of tauopathy is progressive and age dependent in heterozygous tau^{v337M} flies.^{[99,](#page-19-5)[100](#page-19-6)} There were minimal effects of tau^{V337M} on mitochondrial and nuclear proteostasis [\(Figures 7B](#page-12-0) and 7C), apart from a decrease in the insoluble mito-Fluc^{DM} levels at a young age ([Figure 7](#page-12-0)B). Altogether, these analyses indicate that Fluc^{DM} insolubility is promoted by tau^{V337M} primarily in the cytoplasm and in an age-dependent manner.

Although pathogenic tau has been reported to impact skeletal muscle,^{[104–106](#page-19-8)} our analyses revealed minimal effects of pathogenic tauV337M on cytosolic and mitochondrial proteostasis in this tissue, as assessed by monitoring the detergent-soluble and insoluble levels of the untargeted Fluc^{DM} ([Figure 7](#page-12-0)D) and mito-Fluc^{DM} ([Figure 7E](#page-12-0)) in muscle. Therefore, these findings suggest that skeletal muscle may be relatively resistant to derangement of cytoplasmic proteostasis by tau^{V337M} compared to the central nervous system. There was, however, an unexpected decline in the insoluble levels of NLS-Fluc^{DM} in response to tau^{V337M}, which may indicate derangement of nuclear protein import, whereas the soluble levels were unaffected ([Figure 7F](#page-12-0)). Together, these findings indicate that pathogenic tau influences protein solubility in the nuclei of skeletal muscle cells ([Figure 7F](#page-12-0)). Because such a reduction in insoluble NLS-Fluc^{DM} levels was not seen in fly head extracts ([Figure 7C](#page-12-0)), which are enriched for the central nervous system, these findings suggest that the impact of pathogenic tau on subcellular proteostasis varies depending on the tissue and cell type.

Ultra-deep-coverage mass spectrometry indicates that pathogenic tau regulates, in opposite manners, the solubility of distinct protein sets during aging in **Drosophila**

To complement these analyses and determine how pathogenic tau impacts the solubility of the entire proteome, we next utilized ultra-deep-coverage tandem mass tag (TMT) mass spectrometry to analyze detergent-soluble and insoluble fractions. Also, in these studies, we utilized knockin tau models in which the endogenous *Drosophila* tau was replaced by wild-type human tau (tau^{WT}) or by pathogenic human tau^{v337M [99,100](#page-19-5)} On this basis, an 18-plex TMT analysis was done from whole flies with homozygous tau V337M and control tau WT expression at day 60, which

corresponds to an old age in which behavioral deficits are man-ifest in tau^{V337M} versus tau^{WT} fly strains.^{[99](#page-19-5),[100](#page-19-6)} In addition, homozygous young (10-day-old) tau^{V337M} flies were examined to determine the age-related changes in protein solubility that occur with aging. This TMT-based profiling indicates that the solubility of several proteins is reshaped by tau^{V337M} compared to control, age-matched, isogenic tau WT flies at 60 days of age [\(Figures 7](#page-12-0)G–7I; [Table S2](#page-16-8)).

Because many proteins are expressed at the threshold of their solubility,^{[77](#page-18-13)[,107–114](#page-19-9)} it has been hypothesized previously that aggregation-prone proteins generally tip the balance toward insolubility for other proteins. However, our TMT-based analyses indicate that pathogenic tau^{V337M} has complex effects on protein solubility: while some proteins become more insoluble, others display increased solubility upon expression of tau^{V337M} versus tau^{WT} [\(Figures 7](#page-12-0)G–7I, [Figures S6A](#page-16-8), and S6B).

We next examined the proteins with a significant difference $(p < 0.05)$ in protein solubility (insoluble/soluble levels) in tau $\frac{V337M}{V}$ versus tau^{WT} flies to determine whether any specific subcellular compartment is impacted. While several mitochondrial proteins were among the proteins with solubility regulated by tau^{V337M} versus tau^{WT} (with primarily increased insolubility in response to tau^{V337M}), the magnitude of these changes, although significant $(p < 0.05)$ was overall low [\(Figure 7](#page-12-0)G). Of the 3 significant proteins with nuclear localization, only one (nucleolar GTP-binding protein 1, CG8801) had increased insolubility in tau^{V337M} versus tau^{WT} [\(Figure 7](#page-12-0)G).

Further analysis of over-represented protein categories [\(Table S3](#page-16-8)) indicates that transmembrane proteins (e.g., tetraspanins), proteins with disordered domains, and proteins that are components of cell projections (e.g., axons) are enriched among proteins with tau^{V337M}-induced protein insolubility ($p < 0.05$ and $\Delta > 0.5$ for insoluble/soluble levels in tau^{V337M} versus tau^{WT}). Curated analyses further indicate the presence of several proteins involved in synaptic transmission and brain function (e.g., the kainate-type ionotropic glutamate receptor subunit 1D, the potassium:chloride symporter Kazachoc, axotactin, syntaxin-1A, adenylate cyclase, and synaptosomal-associated protein 25) and proteins enriched in microvilli and rhabdomeres and, hence, required for photoreceptor function (e.g., prominin, chaoptin, and opsin Rh1) [\(Figures 7H](#page-12-0) and [S6](#page-16-8)C). Together, these findings suggest that tau $\frac{V337M}{M}$ may impact synaptic transmis-sion^{[88,](#page-19-10)[115–118](#page-19-11)} and light detection by photoreceptors (as reported for human patients and mouse models of tauopathy $119-123$) by perturbing the solubility of specific components of the synapse and of the light-sensing apparatus. Altogether, organism-wide profiling of proteome solubility indicates that tau^{V337M} particularly impacts the solubility of proteins involved in the function of the central nervous system. Conversely, proteins with tauV337M-induced upregulation of solubility (*p* < 0.05 and $\Delta < -0.5$ for insoluble/soluble levels in tau V337M versus tau $^{WT})$

⁽G) The difference in the insoluble/soluble protein levels in tau^{V337M} versus tau^{WT} at approximately day 60. There are 432 proteins of 4,012 quantified proteins that display significantly ($p < 0.05$) increased ($n = 140$; $n = 20$ with log2R > 1) and decreased ($n = 292$, $n = 43$ with log2R <-1) insolubility in tau^{v337M} versus tau^{WT} at day 60. These include mitochondrial (pink) and nuclear (green) proteins.

⁽H and I) Examples of proteins with increased (H) and decreased (I) insolubility in response to tau^{V337M} in old age compared to tau^{WT} at the same age (black line of comparison) and compared to tau^{V337M} at a young age (green line). $n = 3$ (biological replicates) with the mean \pm SD indicated; *p < 0.05, $^{**}p$ < 0.01, *** p < 0.001 (one-way ANOVA).

are enriched in peptidases/proteases, glycoproteins, and proteins with disordered regions. Further refined analyses indicate that several secreted factors (e.g., Ance-3, Swim, teq, and fon) and the chaperone crystallin are among the proteins with improved solubility in tau^{V337M} versus control tau^{WT} flies ([Figures](#page-12-0) [7](#page-12-0)I and [S6C](#page-16-8)).

In summary, these proteomics analyses identify remarkably diverse impacts of pathogenic tau^{V337M} on protein solubility and also highlight subcellular compartments (cell projections and plasma membrane) that are particularly affected.

DISCUSSION

Eukaryotic cells have evolved complex functions via their compartmentalization into organelles and subcellular microenvi-ronments.^{[124](#page-20-1)} Such specialization of the cellular space has enabled optimal organelle function but also resulted in compart-ment-specific challenges to proteostasis.^{[9](#page-16-3)[,23](#page-17-5),[125](#page-20-2)} Consequently, organelle-specific stress responses and quality control mechanisms are in place to ensure proteostasis within each compart-ment.^{[2,](#page-16-2)[18,](#page-17-3)[19](#page-17-20)[,126](#page-20-3)} Although the loss of proteostasis is a key hallmark of aging, 47 it remains largely unknown how subcellular proteostasis is impacted by environmental interventions and endocrine signaling factors that regulate aging. Moreover, while aggregation-prone proteins are the culprit of many age-rela-ted diseases.^{[7](#page-16-7)} it remains undetermined whether they perturb proteostasis across the cell or only in specific subcellular compartments.

In this study, we have generated *Drosophila* strains that ubiquitously express misfolding-prone Fluc^{DM} protein variants that were engineered to selectively localize to the nucleus and mitochondria, along with an untargeted variant that localizes primar-ily to the cytoplasm ([Figure 1](#page-2-0)). By using these tools, we demonstrate that Fluc^{DM} protein variants can be utilized to perturb proteostasis in a compartment-targeted manner in the cytoplasm, nucleus, and mitochondria when compared to isogenic flies with no Fluc^{DM} expression. Analysis of the organism-wide consequences of these ubiquitous subcellular perturbations indicates that mild stress induced by expression of compartment-targeted Fluc^{DM} variants extends lifespan, impedes ageassociated neuromuscular aging, and promotes proteostasis ([Figure 2](#page-5-0)). At the molecular level, there are both common and variant-specific transcriptional responses that are induced by Fluc^{DM} ([Figure 3](#page-6-0)). Apart from a strong induction of the mitochondrial small heat shock protein Hsp22 by all $Fluc^{DM}$ variants, chaperones were not an overrepresented category among the differentially regulated genes [\(Figure 3](#page-6-0)B), suggesting that the transcriptional stress response induced by $Fluc^{DM}$ variants is different from the heat shock response (HSR).^{[127,](#page-20-4)[128](#page-20-5)} Likewise, apart from Hsp22, there was no upregulation of Hsp60 ([Table S1;](#page-16-8) [Figure S7\)](#page-16-8), a mitochondrial chaperone that is stereotypically induced by the mitochondrial unfolded protein response (UPR^{mt}).^{20[,129](#page-20-6)} Altogether, these findings indicate that perturbation of subcellular proteostasis by $Fluc^{DM}$ does not simply induce the HSR or the UPR mt [\(Figure 3\)](#page-6-0). Previous studies in cultured</sup> cells engineered to express nuclear and cytoplasmic proteins with destabilizing domains similarly found that the transcriptional stress response induced by these misfolding-prone proteins is

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distinct from the HSR and the UPR.^{[16](#page-17-1)} Altogether, these findings suggest that transcriptional responses different from the HSR and the UPR can be induced by misfolding-prone proteins targeted to specific subcellular compartments. Interestingly, Hsp22 levels have been identified previously as an important predictor of survival in *Drosophila* during aging,^{[35](#page-17-12)[,45](#page-17-22)} indicating a possible role of this chaperone as part of the stress response and lifespan extension induced by Fluc^{DM} [\(Figure 2\)](#page-5-0). However, apart from Hsp22, chaperones do not appear to be a major category of genes transcriptionally induced by $Fluc^{DM}$ [\(Fig](#page-16-8)[ure S7\)](#page-16-8). Other gene categories were rather induced by Fluc^{DM}, including lipases and proteases, which may, respectively, help maintain metabolic homeostasis and protein quality control in cooperation with the ubiquitin-proteasome system and autophagy.^{[130](#page-20-7)}

In addition to utilizing $Fluc^{DM}$ variants as tools to perturb subcellular proteostasis [\(Figures 2](#page-5-0) and [3](#page-6-0)), we found that they can be used as reporters of proteostasis. Specifically, by measuring the detergent-soluble and insoluble levels of each Fluc^{DM}, the impact of genetic and environmental interventions on subcellular proteostasis can be determined compared to controls. In partic-.
ular, the detergent insolubility of the untargeted Fluc^{DM}, mito-Fluc^{DM}, and NLS-Fluc^{DM} sensors increases to different degrees upon heat shock [\(Figure 4A](#page-7-0)). Moreover, the beneficial effects of cold shock may derive from an improvement in cytoplasmic proteostasis, as indicated by a reduction in the detergent-insoluble levels of the untargeted Fluc^{DM}, whereas mito-Fluc^{DM} and NLS-Fluc^{DM} were not affected ([Figure 4](#page-7-0)B). Different endocrine signals that regulate aging (the glucagon-like hormone Akh^{70-72} and Amyrel/maltose signaling 69) also elicited remarkably distinct outcomes on subcellular proteostasis: while Akh improved nuclear proteostasis (by reducing the detergent-insoluble levels of NLS-Fluc^{DM}), Amyrel reduced the insoluble levels of the untargeted, largely cytoplasmic Fluc^{DM}. Because we induced Akh and Amyrel specifically in skeletal muscle and assessed the outcome systemically and/or in head extracts (which are enriched for the central nervous system), these findings suggest that endocrine signaling mediated by different muscle-secreted factors (myokines) may have remarkably distinct impacts on subcellular proteostasis. Physiologically, this may occur in response to muscle contraction because Amyrel is induced by cell stress,^{[69](#page-18-3)} which is a component of exercise,^{[131–135](#page-20-8)} and because glucagon-like peptide-1 (GLP-1, possibly related to Akh) is expressed by skeletal muscles in response to exercise in mice. 136 Altogether, these findings indicate that compartment-targeted Fluc^{DM} can be utilized as a sensor to determine whether environmental and genetic interventions regulate subcellular proteostasis.

We also examined how distinct aggregation-prone proteins impact the solubility of Fluc^{DM} sensors during aging. Aggregation-prone proteins can interact with and disrupt the folding status of other native proteins and impede proteasome activity.^{[7](#page-16-7)[,23](#page-17-5),[74,](#page-18-14)[76,](#page-18-15)[125](#page-20-2),[137](#page-20-10)[,138](#page-20-11)} On this basis, the toxicity of aggregation-prone proteins may derive from a widespread loss of protein quality control.^{[7](#page-16-7),[23,](#page-17-5)[74](#page-18-14)[,76](#page-18-15)[,125,](#page-20-2)[137](#page-20-10),[138](#page-20-11)} To test this hypothesis, here, we estimated the impact of different aggregation-prone proteins on the detergent-insoluble levels of compartment-targeted Fluc^{DM} reporters of proteostasis [\(Figures 6](#page-11-0) and [7](#page-12-0)). Our analyses

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indicate a range of diverse outcomes. A toxic protein with 49(GGGGCC) repeats, associated with amyotrophic lateral sclerosis and frontotemporal dementia, 80 widely impaired proteostasis across cell compartments (cytoplasm, mitochondria, and nucleus). Conversely, pathogenic ataxin-3 (hATXN3.tr-Q78) associated with Machado-Joseph disease/spinocerebellar ataxia^{[82](#page-18-11)} had minimal effects on the detergent-insoluble levels of all compartment-targeted Fluc^{DM} reporters [\(Figure 6](#page-11-0)).

In addition to 49(GGGGCC) and hATXN3.tr-Q78, we examined how a mutant form of MAPT^{[90](#page-19-12),[139](#page-20-12)} impacts Fluc^{DM} solubility. Normally, tau is predominantly (>90%) attached to microtubules, but its mutation and/or hyperphosphorylation promote its aggregation and accumulation into cytoplasmic neurofibrillary tangles in Alzheimer's disease and related dementias.^{[7,](#page-16-7)[90,](#page-19-12)[101](#page-19-7),[139](#page-20-12)[,140](#page-20-13)} Recent evidence indicates that cytoplasmic, soluble oligomers of tau, generated during tangle formation, are the most toxic tau species that cause neurodegeneration.^{[7](#page-16-7)[,89–91](#page-19-1)} Although the decline of proteostasis is a defining feature of tauopathies, $7,89-91$ $7,89-91$ it remains largely undetermined whether cytoplasmic mutant tau regulates protein quality control in other cell compartments and organelles apart from the cytosol. By utilizing untargeted (\sim cytoplasmic) Fluc^{DM}, mito-Fluc^{DM}, and NLS-Fluc^{DM}, our analyses now indicate that pathogenic tau^{V337M} impairs proteostasis in the cytoplasm but not in the nucleus and mitochondria ([Figures 7A](#page-12-0)–7C) and that this occurs in the central nervous system but not in the skel-etal muscle ([Figures 7](#page-12-0)D–7F). Therefore, although tau^{V337M} is known to derange the function of many cell compartments (e.g., mitochondria and the nucleus), $90,139$ $90,139$ this does not seem to occur via a general perturbation of proteostasis in these compartments (as inferred from the detergent-insoluble levels of mito-Fluc^{DM} and NLS-Fluc^{DM}) but only in the cytoplasm.

In summary, our analyses indicate that the impact of aggregation-prone proteins on subcellular proteostasis is widely diverse: there is an overt decline in cytoplasmic, nuclear, and mitochondrial proteostasis in response to 49(GGGGCC) expression ([Fig](#page-11-0)[ure 6](#page-11-0)), whereas tau V337M impedes only cytoplasmic protein quality control [\(Figure 7\)](#page-12-0), and hATXN3.tr-Q78 has minimal impact ([Fig](#page-11-0)[ure 6](#page-11-0)). Altogether, these findings pinpoint that the toxicity of aggregation-prone proteins is not necessarily coupled to widespread disruption of proteostasis across cell compartments.

By analyzing the effects of pathogenic tau^{V337M} on subcellular proteostasis in head extracts (consisting mostly of tissues from the central nervous system, CNS) versus thoraces (consisting of skeletal muscle), we found that tau^{V337M} impedes cytoplasmic proteostasis in the CNS but not in muscle ([Figure 7](#page-12-0)), consistent with the fact that tauopathies have primary manifestations in the brain and less so in peripheral tissues.^{86–88} Interestingly, tau^{V337M} expression reduced the detergent-insoluble levels of NLS-Fluc^{DM} in muscle fractions [\(Figures 7](#page-12-0)D–7F), suggesting that pathogenic tau^{V337M} may derange nuclear protein import in muscle cells.

To further explore how tau^{V337M} impacts the solubility of the proteome, we utilized TMT mass spectrometry to profile the detergent-soluble and -insoluble levels of >4,000 proteins in whole flies that expressed endogenous levels of human tau V337M compared to tau^{WT}. These analyses indicate that pathogenic tau^{V337M} does not simply skew the proteome toward insolubility but, rather, remodels protein solubility in a balanced manner: \sim 3.4% and \sim 7.2% of detected proteins displayed significantly

increased and decreased insolubility, respectively, in tau^{V337M} versus tau^{WT} ([Figures 7](#page-12-0)G–7I). While proteins with tau^{V337M}-modified solubility encompass several cell compartments, including the nucleus and mitochondria, there is an enrichment for transmembrane proteins and components of cell projections (e.g., axons). Functional categories of such proteins include synaptic function and light perception, suggesting that the solubility of proteins necessary for key functions of the CNS is prominently impacted by tau V337M. These studies therefore indicate a complex remodeling of proteome solubility by pathogenic tau.

Altogether, transgenic *Drosophila* strains that express compartment-targeted Fluc^{DM} sensors provide tools to survey subcellular proteostasis and guide subsequent analyses. Depending on the experimental design, compartment-targeted $Fluc^{DM}$ variants can be utilized as reporters or as tools to perturb subcellular proteostasis. On this basis, transgenic Fluc^{DM} strains may be utilized to determine how protein quality control is regulated across cell compartments during the progression of agerelated diseases and by interventions that delay aging.

Limitations of the study

This study examined how Fluc^{DM}-based sensors of subcellular proteostasis can be utilized to either monitor or perturb subcellular proteostasis (depending on the experimental design) in the context of aging and age-associated conditions. To perturb subcellular proteostasis, Fluc^{DM} variants targeted to different subcellular compartments are compared to isogenic controls with no Fluc^{DM} to detect responses that are triggered by compartment-targeted Fluc^{DM}. However, we did not systematically examine the proteomic alterations resulting from $Fluc^{DM}$ and how these compare to other established proteostatic challenges occurring during normal aging or experimental interventions. In a second application, Fluc^{DM} variants can be utilized to monitor the impact of an experimental intervention (genetic, pharmacologic, etc.) on subcellular proteostasis by comparing such interventions to controls that also express the same Fluc^{DM}. For this application, the measurement of luciferase activity does not appear to be a convenient readout for the mitochondrial and nuclear luciferases (mito-Fluc^{DM}-EGFP and NLS-Fluc^{DM}-EGFP) because of the low levels detected, which are presumably due to the low luciferase activity and/or low stability of these Fluc^{DM} variants. Because mitochondrial and nuclear proteins are translated in the cytoplasm and then imported into mitochondria and nuclei, the insolubility of mito-Fluc^{DM}-GFP and NLS-Fluc^{DM}-GFP may also reflect defects in protein import into these cellular compartments. Another limitation is that the Fluc^{DM} sensors indicate the general status of proteostasis but not necessarily that of individual proteins or specific protein categories. Our mass spectrometry data indeed indicate that pathogenic tauV337M differentially remodels the solubility of distinct protein subsets [\(Figure 7\)](#page-12-0).

RESOURCE AVAILABILITY

Lead contact

Requests for further information, resources, and reagents should be directed to and will be fulfilled by the lead contact, Fabio Demontis [\(Fabio.Demontis@](mailto:Fabio.Demontis@stjude.org) [stjude.org](mailto:Fabio.Demontis@stjude.org)).

Materials availability

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There are no restrictions on the availability of the data and tools generated by this study.

Data and code availability

- The data supporting the findings of this study are available within the paper [\(Figures 1,](#page-2-0) [2](#page-5-0), [3,](#page-6-0) [4](#page-7-0), [5,](#page-10-0) [6](#page-11-0), [7](#page-12-0) and [S1–S7](#page-16-8) and [Tables S1–S5\)](#page-16-8). The TMT mass spectrometry proteomics data have been deposited in the ProteomeXchange Consortium via the PRIDE partner repository and are accessible with the dataset identifier PRIDE: PXD045325. The RNA-seq data have been deposited in the Gene Expression Omnibus (GEO) with the identifier GEO: GSE243224.
- \bullet This paper does not report original code.
- Any other information required to reanalyze the data reported in this paper is available from the [lead contact](#page-15-0) upon request.

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AUTHOR CONTRIBUTIONS

M.R. generated the Fluc^{DM} transgenic sensors. M.C. did most of the experiments with the Fluc^{DM} sensors together with M.R., C.-L.C., A.S., Z.C., and M.R.-M. V.P. did the TMT mass spectrometry and corresponding data analyses. Y.-D.W. analyzed the RNA-seq data. J.P. supervised the mass spectrometry studies. F.D. supervised the project and wrote the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

STAR+METHODS

Detailed methods are provided in the online version of this paper and include the following:

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. [QUANTIFICATION AND STATISTICAL ANALYSIS](#page-26-0)

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.crmeth.2024.100875) [crmeth.2024.100875](https://doi.org/10.1016/j.crmeth.2024.100875).

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STAR+METHODS

KEY RESOURCES TABLE

(*Continued on next page*)

Article

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Drosophila husbandry

Flies were kept (~30 flies/tube) at 25°C, 60% humidity, and a 12h/12h light-dark cycle in tubes containing cornmeal/soy flour/yeast fly food. The fly food was changed regularly every 2-3 days. Flies were aged to 10, 30, and 60 days at 25°C (unless otherwise indicated). Similar procedures as previously reported were utilized for measuring survival during aging and for negative geotaxis.^{[141–144](#page-20-14)} All experiments were done with male flies.

METHOD DETAILS

Drosophila stocks

The following fly stocks were utilized in this study: *Mhc-Gal4*, [141–144](#page-20-14) *GMR-Gal4*, [33](#page-17-19) *UAS-mCherry* (Bloomington stock center #35787), *UAS-8(GGGGCC)* repeats (non-toxic control; #84724), *UAS-49(GGGGCC)* repeats (toxic poly-GA proteins, as found in C9orf72 associated with ALS and FTD; #84727),⁸⁰ UAS-hATXN3.tr-Q27 (non-toxic control; #8149), UAS-hATXN3.tr-Q78 (expresses a toxic C-terminal fragment of the human Machado-Joseph Disease/Spinocerebellar Ataxia 3 protein with a 78 repeat polyglutamine tract; #8150), *UAS-Htt-Q72-GFP*, [31–33](#page-17-11) and knock-in tau models in which the endogenous *Drosophila* tau has been replaced by wild-type

human tau (tauWT) and by pathogenic human tauV337M. [99,100](#page-19-5) These additional fly stocks were also utilized: *UAS-mCherryRNAi* (#35785), *UAS-Hsp70RNAi* (#33948), and *UAS-Hsp22* (#20055). FlucDM stocks (generated as indicated below), *UAS-Akh* (#27343), and *UAS-Amyrel*[69](#page-18-3) were backcrossed against the *w¹¹¹⁸* background and compared to their respective litter-mate isogenic controls with no transgene expression.

Cloning and establishment of compartment-targeted Fluc^{DM} transgenic sensors in Drosophila

A *pCaSpeR5-attB-tubulin promoter* vector was generated by PCR-mediated amplification and cloning of a *tubulin* promoter and *attB* sequences by using *Kpn*I with *EcoR*I-HF, and *EcoR*I-HF, respectively. To express a general, untargeted version of a misfolding-prone luciferase (Fluc^{DM}),^{[10](#page-16-4)} Fluc^{DM} fused with EGFP was PCR-amplified from the *pCI-Fluc^{DM}-EGFP* plasmid¹⁰ and cloned into the *pCaSpeR5-attB-tubulin promoter* vector with *Kpn*I-HF and *Xba*I. A similar procedure was used to clone a mitochondriallytargeted version of Fluc^{DM}-EGFP (mito-Fluc^{DM}), which consists of a mitochondrial target sequence from human COX VIII (MRLRLTVFCGVLRAIMSVLTPLLLRGLTGSARRLPVPRAK, previously utilized for establishing the mito-GFP *Drosophila* stock, Bloo-mington #8442),^{24,[25](#page-17-7)} a linker (RSS), and Fluc^{DM}-EGFP. For establishing Fluc^{DM} targeted to the nucleus (NLS-Fluc^{DM}), the canonical PAAKRVKLD nuclear localization signal²⁶ was added by PCR to the N-terminus of Fluc^{DM}-EGFP. These plasmids were injected into *w1118;attP40* embryos and *tubulin-FlucDM-EGFP* transgenic *Drosophila* stocks were established by phiC31-mediated site integration.¹⁴⁵

Immunostaining and laser scanning confocal microscopy of Drosophila tissues

Drosophila brains and guts from Fluc^{DM} transgenic stocks, established as described above, were immunostained according to stan-dard procedures.^{[32,](#page-17-18)[144](#page-20-16),[146](#page-20-17)} Specifically, brains and guts were dissected and immediately fixed with 4% paraformaldehyde and 0.2% Triton X-100 for 30 min, washed, and then incubated with 1:200 chicken anti-GFP (Aves #GFP-1010) primary antibodies, followed by AlexaFluor488-conjugated anti-chicken secondary antibodies (ThermoFisher #A11039) together with DAPI (ThermoFisher #D1306) and AlexaFluor635-conjugated phalloidin (Life Technologies #A22284). Co-staining with mouse anti-ATP5a (abcam #14748) was done to assess the degree of mito-Fluc^{DM} localization to mitochondria, identified by the mitochondrial marker ATP5a. The samples were mounted on glass slides with SlowFade Gold antifade reagent (Invitrogen #S36937) and #1.5 coverslips and imaged by using a Zeiss LSM780 confocal microscope. Enterocytes of the midgut and brain cells of the antennal lobes are shown in the representative images in [Figures 1](#page-2-0)B–1D.

RNA sequencing and hierarchical clustering

Samples for RNA sequencing were prepared with TRIzol (Ambion #15596018) from whole flies and the RNA was extracted by iso-propanol precipitation from the aqueous phase.^{[147–149](#page-20-18)} RNA sequencing libraries for each sample were prepared from 1 µg total RNA by using the Illumina TruSeq RNA Sample Prep v2 Kit per the manufacturer's instructions, and sequencing was completed on the Illumina NovaSeq 6000. The 100-bp paired-end reads were trimmed, filtered against quality (Phred-like Q20 or greater) and length (50-bp or longer), and aligned to the Drosophila melanogaster reference genome (BDGP6/dm6) by using CLC Genomics Workbench v12.0.1 (Qiagen). For gene expression comparisons, we obtained the TPM (transcripts per million) counts from the CLC RNA-Seq Analysis tool. The differential gene expression analysis was performed by applying the non-parametric ANOVA using Kruskal-Wallis and Dunn's tests on log-transformed TPM, implemented in the Partek Genomics Suite v7.0 software (Partek Inc.). Hierarchical clustering of RNA-seq data into heatmaps was done by using the UPGMA clustering method (unweighted pair group method with arithmetic mean) with similarity measure of correlation, implemented in Spotfire v7.5.0 software (TIBCO). The RNA-seq data dis-cussed in this publication are available in [Table S1](#page-16-8) and have been deposited in the NCBI's Gene Expression Omnibus (GEO) with accession number GEO: GSE243224. GO terms analysis was done with DAVID.^{[150](#page-21-0)}

qRT-PCR

For qRT-PCR, cDNAs were reverse transcribed with the iScript cDNA synthesis kit (Bio-Rad #1708840) from 1 μ g total RNA. qRT-PCR was performed by using the IQ Sybr Green supermix (Bio-Rad #170–8885). *Tub84B* was utilized as normalization. The following qRT-PCR oligos were used:

Hsp22: 5′-CTTACCGATGTTTTGGCGCA-3′ and 5′-TCGTGGAAGAAGGCGTGAAA-3' *Hsp70Bb: 5′-*AAATCGGATGGAGAGTTGGC-3′ and 5′-TGTAGGCGGGTTTTTGTTT-3' A*kh*: 5'-AAGCACCGCGAGTAGATAGC-3' and 5'-TGTGTGTGCGTGCTAGACAT-3' Tub84B: 5'-GCTGTTCCACCCCGAGCAGCTGATC-3' and 5'-GGCGAACTCCAGCTTGGACTTCTTGC-3' Fluc *(firefly luciferase): 5'*-CCCTGGTTCCTGGAACAATTGC-3' and 5'-AAGAATTGAAGAGAGTTTTCACTGC-3'

Analysis of aggregates of pathogenic huntingtin

Pathogenic huntingtin-polyQ72-GFP protein aggregates were imaged as previously done^{[33](#page-17-19)} by using a ZEISS SteREO Discovery.V12 epifluorescence microscope. Subsequently, the images were analyzed using Cell Profiler 3.0.0 ([cellprofiler.org\)](http://cellprofiler.org) to quantify the total area of protein aggregates (Huntingtin-polyQ72-GFP speckles) normalized by the retinal tissue area. This analysis was done with male flies after aging them at 25° C for 30 days.

Heat shock and cold shock

For the analysis of Fluc^{DM} solubility upon thermal stress, 14-days-old wild-type flies were heat shocked at 36°C for 3 h. For cold shock, 14-days-old flies were housed at 4° C for 2 h. Only flies that were alive after the heat and cold shocks were utilized for subsequent analyses.

Western blots of detergent-soluble and insoluble protein fractions

Western blots of detergent-soluble and insoluble fractions were obtained substantially as described previously.^{[142](#page-20-19),[151](#page-21-1)} Specifically, whole bodies, heads, or thoraces were homogenized in 60 µL ice-cold Triton X-100 buffer (1% Triton X-100 in PBS containing protease inhibitors and phosphatase inhibitors) for 5 min at the highest speed. Homogenates were centrifuged at 14,000 rpm at 4° C for 10 min and the supernatant was collected (Triton X-100 soluble fraction). The remaining pellet was washed in 400 μ L Triton X-100 buffer and centrifuged twice at 14,000 rpm for 5 min at 4° C. The pellet was then resuspended at room temperature in 60 µL RIPA buffer containing 8M urea and 5% SDS, centrifuged at 14,000 rpm at 4°C for 10 min, and the supernatant collected (Triton X-100 insoluble fraction). 8 µL of soluble/insoluble protein extracts were boiled with sample buffer containing DTT and used for SDS-PAGE. Detergent-soluble and insoluble fractions were analyzed on 4–20% SDS-PAGE with anti-GFP antibodies (1:500, dissolved in TBST; Cell Signaling Technologies, #2956). Ponceau S staining (ThermoFisher #A40000279) and anti-b-actin and/or anti-a-tubulin antibodies (Cell Signaling Technologies, #8457 and #2125) were used as loading controls. The western blots with head samples were typically normalized by β -actin whereas the blots with thoracic samples were normalized by Ponceau staining or α -tubulin.

Luciferase assays

For luciferase assays, 10 thoraces/replicate were homogenized in 170 µL ice-cold homogenization buffer (250 mM sucrose, 50 mM Tris-HCl, 5 mM MgCl₂, and protease inhibitors) in a NextAdvance bullet blender at 4° C. Subsequently, 90 µL of the homogenate was transferred to a new tube and gently mixed with an equal volume of Dual-Glo luciferase reagent (Promega).^{[141](#page-20-14)[,142,](#page-20-19)[152](#page-21-2)} After a 10-min incubation, the luminescence was read with a Tecan Infinite 200 Pro. Luciferase readings were normalized by the protein content, as estimated with the Pierce BCA protein assay kit (Thermo Scientific).

Preparation of detergent-soluble and insoluble fractions, protein digestion, and peptide isobaric labeling by tandem mass tags

For TMT of *Drosophila* samples, 25 male flies per sample (*n* = 3 biological replicates per condition) were collected at day 10 and day \sim 60 from eclosion and homogenized in 200 µL NP40 cell lysis buffer (Invitrogen #FNN0021) with protease and phosphatase inhibitors and 0.5-mm zirconium beads in a NextAdvance bullet blender.^{[153,](#page-21-3)[154](#page-21-4)} After homogenization, the samples were centrifuged at maximum speed for 10 min, and the supernatant was collected (corresponding to the detergent-soluble fraction, typically consisting of >100 µg). The remaining pellet was washed 3x in PBS and homogenized in insoluble lysis buffer containing 8M urea, PBS, and 1% SDS (with protease and phosphatase inhibitors). The resulting supernatant was collected (corresponding to the detergent-insoluble fraction, typically consisting of >10 µg). All the samples were then prepared for subsequent analyses by adding SDS-blue loading buffer and DTT (Cell Signaling Technologies #7722), and by heating them for 5 min at 95°C. The protein concentration of the lysates was determined by Coomassie-stained short gels using bovine serum albumin (BSA) as standard.^{[155](#page-21-5)} Specifically, the samples were run on 7.5% SDS-page gels at 60V for 10–15 min, or until all the protein was through the initial lane. BSA was used as a standard at 0.2, 1, and 5 mg. The gels were washed in distilled water and stained with Coomassie blue for 1–4 h with gentle shaking at room temperature, shielded from light. The gels were then de-stained with distilled water overnight at 4°C with gentle shaking. The bands were excised from the gels, cut into several pieces, collected into 1.5 mL-eppendorf tubes on ice, and stored at -80° C.

In preparation for TMT, 1^{56} the short gel bands were washed twice with 50% acetonitrile and dried. The dried gel bands were then incubated with trypsin at an enzyme-to-substrate ratio of 1:10 (w/w) for overnight digestion. Following the overnight digestion, the peptide solution from the short gel bands was extracted, and the peptides were reduced by adding 1 mM DTT for 30 min at room temperature followed by alkylation with 10 mM iodoacetamide (IAA) for 30 min in the dark at room temperature. The unreacted IAA was quenched with 30 mM DTT for 30 min. Finally, the digestion was terminated and acidified by adding trifluoroacetic acid (TFA) to 1%, desalted using C18 cartridges (Harvard Apparatus), and dried by speed vac. The peptide mixture was resuspended in 50 mM HEPES (pH 8.5) and labeled with 18-plex Tandem Mass Tag (TMT) reagents (ThermoScientific) following the manufacturer's recommendations.

Two-dimensional HPLC and mass spectrometry

The TMT-labeled samples were mixed equally, desalted, and fractionated on an offline HPLC (Agilent 1220) using basic pH reversephase liquid chromatography (pH 8.0, XBridge C18 column, 4.6 mm \times 25 cm, 3.5 μ m particle size, Waters). The fractions were dried and resuspended in 5% formic acid and analyzed by acidic pH reverse phase LC-MS/MS analysis. The peptide samples were loaded on a nanoscale capillary reverse phase C18 column (New objective, 75 μ m ID \times ~25 cm, 1.9 μ m C18 resin from Dr. Maisch GmbH) by an HPLC system (Thermo Ultimate 3000) and eluted by a 60-min gradient. The eluted peptides were ionized by electrospray ionization and detected by an inline Orbitrap Fusion mass spectrometer (ThermoScientific). The mass spectrometer is operated in datadependent mode with a survey scan in Orbitrap (60,000 resolution, 1×10^6 AGC target and 50 m maximal ion time) and MS/MS high-resolution scans (60,000 resolution, 2 \times 10⁵ AGC target, 120 m maximal ion time, 32 HCD normalized collision energy, 1 *m/z* isolation window, and 15 s dynamic exclusion). 157

MS data analysis

The MS/MS raw files were processed by the tag-based hybrid search engine JUMP.^{[158](#page-21-8)} The raw data were searched against the UniProt *Drosophila* database concatenated with a reversed decoy database for evaluating false discovery rates. Searches were performed using a 15-ppm mass tolerance for both precursor and product ions, fully tryptic restriction with two maximal missed cleavages, three maximal modification sites, and the assignment of *a*, *b*, and *y* ions. TMT tags on Lys and N-termini (+304.20715 Da) were used for static modifications and Met oxidation (+15.99492 Da) was considered as a dynamic modification. Matched MS/MS spectra were filtered by mass accuracy and matching scores to reduce the protein false discovery rate to \sim 1%. Proteins were quantified by summing reporter ion intensities across all matched PSMs using the JUMP software suite.^{[159](#page-21-9)} Categories enriched in protein sets were identified with DAVID.^{[150](#page-21-0)}

The TMT mass spectrometry proteomics data are reported in [Table S2](#page-16-8) and have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PRIDE: PXD045325.

QUANTIFICATION AND STATISTICAL ANALYSIS

Data organization, scientific graphing, and statistical analyses were done with Microsoft Excel (version 14.7.3) and GraphPad Prism (version 8). The unpaired two-tailed Student's t test was used to compare the means of two independent groups to each other. Oneway ANOVA was used for multiple comparisons of more than two groups of normally distributed data. Survival and climbing data were analyzed with the log-rank test. The n for each experiment can be found in the figures and represents independently generated samples. Bar graphs represent the mean \pm SD or the mean \pm SEM, as specified in the figure legend. A significant result was defined as *p* < 0.05. Throughout the figures, asterisks indicate the significance of *p* values: **p* < 0.05, ***p* < 0.01, and ****p* < 0.001.