Decreasing pdzd8-mediated mitochondrial-ER contacts in neurons improves fitness by increasing mitophagy

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Abstract

The complex cellular architecture of neurons combined with their longevity makes maintaining a healthy mitochondrial network particularly important and challenging. One of the many roles of mitochondrial-ER contact sites (MERCs) is to mediate mitochondrial quality control through regulating mitochondrial turnover. Pdzd8 is a newly discovered MERC protein, the organismal functions of which have not yet been explored. Here we identify and provide the first functional characterization of the Drosophila melanogaster ortholog of Pdzd8. We find that reducing pdzd8-mediated MERCs in neurons slows age-associated decline in locomotor activity and increases lifespan in Drosophila. The protective effects of pdzd8 knockdown in neurons correlate with an increase in mitophagy, suggesting that increased mitochondrial turnover may support healthy aging of neurons. In contrast, increasing MERCs by expressing a constitutive, synthetic ER-mitochondria tether disrupts mitochondrial transport and synapse formation, accelerates age-related decline in locomotion and reduces lifespan. We also show that depletion of pdzd8 rescues the locomotor defects characterizing an Alzheimer’s disease (AD) fly model over-expressing Amyloidβ1-42 (Aβ42) and prolongs the survival of flies fed with mitochondrial toxins. Together, our results provide the first in vivo evidence that MERCs mediated by the tethering protein pdzd8 play a critical role in the regulation of mitochondrial quality control and neuronal homeostasis.

Keywords
Mitochondria, ER (Endoplasmic reticulum), organelle contact site, mitoQC, Amyloid beta, mitophagy, lifespan, Drosophila melanogaster, aging, neurodegeneration, healthspan, MERCs, Alzheimer’s disease.

Introduction

Since the vast majority of neurons are postmitotic, maintaining functional neurons throughout an organism’s lifetime requires tight regulation of organelle functions and stress responses. Mitochondria and the endoplasmic reticulum (ER) extend throughout neuronal processes including axons and dendrites, and both are vital and interdependent contributors to neuronal health (Wu et al, 2017). Mitochondria-ER contacts (MERCs) are controlled by a variety of contact site proteins and contribute to a range of functions required for proper development and maintenance of postmitotic neurons, including regulation of calcium homeostasis, lipid biogenesis, organelle reshaping and dynamics, and metabolic signalling (Giacomello & Pellegrini, 2016; Paillusson et al, 2016). Dysregulation of MERCs is particularly damaging to neurons as they are particularly susceptible to calcium overload, oxidative and ER stresses and to altered mitochondrial function, localization and transport (Lee et al, 2018a; Misgeld & Schwarz, 2017).

As MERCs are modulated by a number of different protein complexes, the detrimental effects of dysregulation of MERCs are varied due to the diversity of these contact site functions (Martino Adami et al, 2019). The critical function of MERCs in regulating cellular responses to damage and stress is underscored by the finding that many human patient cellular models and animal models of age-related neurodegenerative diseases have been shown to have disrupted MERCs. Both reduced MERCs (De Vos & Hafezparast; Sepulveda-Falla et al, 2014) and increased MERCs (Area-Gomez et al, 2012; Gómez-Suaga et al, 2019; Zampese et al, 2011) have been implicated in neurodegenerative diseases. Consequently, there is still little consensus on how altered MERCs contribute to neurodegeneration, even within a single disease model.
(Erpapazoglou et al., 2017). The various functions of MERCs make it likely that multiple mechanisms might be involved and, with an ever-expanding toolkit, we can now better define the molecular identities and specific functions of ER-mitochondria tethering complexes and begin to unify many of the seemingly conflicting discoveries in this rapidly growing field (Csordas et al., 2018).

Pdzd8 is one of the most recently discovered proteins that mediates mammalian MERCs (Hirabayashi et al., 2017) and is a paralog of Mmm1 (Wideman et al., 2018), a component of the fungal-specific ER mitochondria encounter structure (ERMES) and first MERC complex identified (Kornmann et al., 2009). We used Drosophila melanogaster to study the consequences of neuronal depletion of pdzd8 both at the cellular and at the organismal level. Importantly, we also describe how the phenotypes associated with neuron-specific depletion of pdzd8 change with age and may contribute to healthy aging. We show the MERCs mediated by the pdzd8 tethering protein regulate mitochondrial turnover through mitophagy and that reducing these contacts prolonged locomotor activity and lifespan in Drosophila melanogaster.

Results

Characterization of fly pdzd8

The Drosophila melanogaster gene CG10362 encodes an uncharacterized protein in the PDZK8 family (Lee & Hong, 2006). CG10362 has a similar predicted domain structure to mammalian Pdzd8 (Figure 1A). Expression of CG10362 in flies is low but is most highly expressed in the central nervous system (FlyAtlas 2; Figure S1A) (Leader et al., 2018) and is enriched in neurons over glia (Figure S1B) (Davie et al., 2018). This specificity in expression in Drosophila provided an excellent opportunity to explore the neuronal functions of this newly discovered MERC protein and also to investigate the functional relevance of MERCs in adult and aging neurons. We propose that CG10362 encodes the fly ortholog of mammalian Pdzd8 and, based on the data presented in this paper, we propose to name it pdzd8.

To characterize the function of pdzd8 in flies, we used the UAS-GAL4 system to manipulate its expression (Brand & Perrimon, 1993). Ubiquitous expression of an RNAi construct targeting pdzd8 strongly reduces its mRNA levels in larvae (Figure S1C). To establish that MERCs were decreased in neurons expressing pdzd8-RNAi, we analysed adult fly brains by transmission electron microscopy (TEM) and manually identified contacts between ER and mitochondria (Figure 1B). Accordingly, the proportion of mitochondria in the soma of adult fly neurons in contact with ER is reduced in flies expressing pdzd8-RNAi (Figure 1C and S1F).

To confirm that pdzd8-RNAi reduces MERCs in axons as well as soma, we adapted the recently developed MERC quantification tool, a split-GFP-based contact site sensor (SPLICS) (Cieri et al., 2017) to create a SPLICS transgenic reporter line. The SPLICS construct targets β-strands 1-10 of GFP to the mitochondrial outer membrane and β-strand 11 to the ER membrane, and where these membranes are in close proximity fluorescent puncta are produced by reconstitution of the split-GFP (Figure S2A). To validate this tool in Drosophila, we expressed SPLICS in motor neurons (Figure S2B) and compared the number of puncta in control axons and those expressing a well-characterized artificial ER-mitochondrial tether developed by Csordas et al. (Basso et
al., 2018; Csordas et al., 2006). The tether construct induces formation of ~5 nm MERCs through transmembrane domains that anchor it in both the mitochondrial outer membrane and the ER (Figure S2C). The density of SPLICS puncta in the axons expressing the tether was four times higher than controls (Figure S2D), indicating the SPLICS reporter was able to detect the increased MERCs resulting from synthetic tether expression in neurons in vivo.

Using this SPLICS construct we also detected a significant decrease in the density of SPLICS puncta in central larval axons (bundles projecting to segments A7 and A8) expressing the pdzd8-RNAi compared to LacZ-RNAi controls (Figure 1D-E). While pdzd8 expression is low beyond the nervous system, ubiquitous knockdown of pdzd8 also reduces the extent of MERCs measured by fluorescence colocalization of ER and mitochondrial signals by super-resolution microscopy (structured illumination microscopy, SIM) in larval epidermal cells (Figure 1F-G and S1E), corroborating our results using TEM and SPLICS analysis.

Consistent with the first reported function of Pdzd8 at MERCs in mouse neurons (Hirabayashi et al., 2017), we observed a reduced number of MERCs (Figure 1), but no obvious changes in mitochondrial or ER morphology in fly larval or adult neurons upon pdzd8 knockdown (See Figures 1B, 1D, 1F, S1D-E, 4A-B, 4F-I, 5A and 5C). Together, these data show that CG10362 encodes the Drosophila homolog of Pdzd8 and functions like mammalian Pdzd8 to regulate MERCs. We next sought to investigate the consequences of loss of pdzd8 from neurons on organismal phenotypes.

**Reduced mitochondria-ER contacts are protective in aging neurons**

Knockdown of pdzd8 in fly neurons produces viable adults. There was no impact on the locomotor performance of pan-neuronal pdzd8-RNAi knockdown in young flies assessed in a climbing assay (Figure 2A). Surprisingly, we found that loss of pdzd8 dramatically slowed the age-associated decline in locomotor activity assessed by the climbing assay (Figure 2A). Importantly, this effect was reproduced by motor neuron-specific pdzd8 knockdown (Figure S3A). Strikingly, the increase in locomotor activity was accompanied by a significant increase in lifespan (Figure 2C).

In contrast, increasing MERCs by expression of a synthetic mitochondria-ER tether in all neurons resulted in a climbing defect in young flies and a significant acceleration of the age-related decline in climbing (Figure 2B), consistent with previous reports (Basso et al., 2018). This climbing defect was exacerbated with age (Figure 2B) and associated with a substantially reduced lifespan (Figure 2D). Consistent with these results, increased expression of pdzd8 also resulted in decreased climbing ability with age (Figure S3B). Notably, this effect was suppressed by co-expression of the pdzd8-RNAi, further validating the specificity of this transgene (Figure S3B). Therefore, decreasing pdzd8-mediated MERCs in neurons prolonged lifespan and protected against locomotor decline with age, while increased MERCs in neurons were found to speed up the age-related decrease in locomotor activity and decreased lifespan.

**Loss of neuronal pdzd8 promotes survival in the presence of mitochondrial toxins**

To investigate how reduction of pdzd8-mediated MERCs improved fitness i.e. prevented age-related decline in locomotor activity and increased lifespan, we assessed whether reducing pdzd8 expression in neurons may protect from additional stresses during aging. We assessed lifespan in flies aged on food with limited nutrients (5 %
sucrose, 1 % agar), and found that in contrast to flies aged on a rich diet (Figure 2C), neuronal expression of pdzd8-RNAi no longer extended the lifespan in comparison to controls (Figure 3A). When an additional oxidative stress was introduced by adding hydrogen peroxide to the food, the flies expressing pdzd8-RNAi died faster than controls (Figure 3B). Therefore, in the presence of general stresses, pdzd8-RNAi is also not protective.

Due to the function of pdzd8 at MERCs, we examined whether the protective effects caused by the pdzd8 depletion in neurons were associated with mitochondrial damage. To address this, we fed the flies mitochondrial toxins: Rotenone, a complex I inhibitor or antimycin A, a complex III inhibitor, both block the electron transport chain and result in dysfunctional mitochondria. Reducing pdzd8 levels in neurons significantly prolonged the survival of flies fed with mitochondrial toxins rotenone (Figure 3C) or antimycin A (Figure 3D) compared to control flies. As improved mitochondrial function could also contribute to the protective effects of loss of pdzd8 in neurons we measured ATP levels in young or aged fly heads expressing pdzd8-RNAi but found no significant differences (Figure S3D). These results indicated that neuronal loss of pdzd8 protects flies from damage induced by mitochondrial toxins.

**Modulating MERCs causes axonal transport and NMJ defects**

While mitochondrial motility is important for neuronal health, it remains an open question whether decline of mitochondrial transport in neurons contributes to aging (reviewed in (Mattioli and Vagnoni 2019)). We first tested the hypothesis that decreased ER mitochondrial tethering contributes to the protective effect of pdzd8 downregulation in aging through changes in mitochondrial motility. We examined the distribution and morphology of mitochondria in axons of CCAP efferent neurons (Park et al, 2003). We found no significant change in mitochondrial length or density in larval axons when comparing control to pdzd8-RNAi or synthetic ER-mitochondria tether-expressing neurons (Figure 4A-C). However, increasing tethering dramatically decreased mitochondrial motility (Figure 4D-E). In contrast, knockdown of pdzd8 to reduce MERCs had no effect on axonal transport in larvae (Figure 4D-E).

To better understand the effects of altered MERCs in neurons, we analysed the morphology of mitochondria located in 1s boutons of larval neuromuscular junctions (NMJs) on muscle 4. Knockdown of pdzd8 led to smaller NMJs and a significant reduction of mitochondrial volume, but overall no change in mitochondrial density compared to control flies (Figure 4F-G), showing that mitochondria distribute normally in these smaller NMJs. Increased tethering, however, resulted in severely deformed NMJs (Figure S3C) and made NMJs type 1s and 1b synaptic boutons indistinguishable, making it impossible to quantify mitochondrial density in 1s boutons. Together, these results show that increasing tethering has dramatic and detrimental effects early in development but reduced tethering through pdzd8-RNAi expression has more limited effects during these early stages of neuronal development.

**Reducing pdzd8 expression increases mitophagy in aged neurons**

We hypothesized that the reduced sensitivity to mitochondrial toxins may be due to improved mitochondrial quality control mechanisms. Thus, we analysed the levels of mitophagy, the clearance of damaged mitochondria by autophagy, in these neurons using the mitoQC mitophagy reporter (Allen et al, 2013; Lee et al, 2018b). This reporter is a pH-sensitive mCherry-GFP fusion targeted to the mitochondrial outer membrane
which provides a read out of mitophagy in the form of mCherry-only signal where the
acidic lysosomal environment has quenched the GFP. Mitophagy was detected in the
soma of nSyb-GAL4-expressing neurons in both larval and adult brains (Figure 5A and
C). There was no significant difference in mitophagy levels in larval neurons expressing
pdzd8-RNAi or control (Figure 5A-B). Quantification of mitoQC mCherry puncta did
not indicate an age-dependent increase in mitophagy in the brains of control adults,
however mitophagy was significantly increased in the brains of 20-day-old pdzd8-
RNAi animals compared to controls (Fig 5C-D).

The majority of mitochondria in neurons are found in the neurites, so to examine
mitophagy in axons of aged flies, we analysed mitoQC signal in axons of the adult fly
wing in situ (Vagnoni & Bullock, 2016) (Figure 5E). Here, in contrast to the adult brain
cell bodies, we did observe an age dependent increase in mitophagy in axons of control
flies (Figure 5F). Consistent with our previous results, pdzd8 knockdown further
increased mitophagy in axons of aged flies (Figure 5E-F). Together, these results show
that loss of pdzd8 promotes the turnover of mitochondria in aging neurons.

Reduced MERCs is protective in a fly model of Alzheimer’s disease

Our results indicate that the loss of pdzd8 is protective against mitochondrial insults,
prevents the age-related decrease in locomotion and increases lifespan. Since
mitochondrial dysfunction is a common feature of many neurodegenerative diseases,
and altered MERCs has been documented in some, we next sought to explore the
neuroprotective potential of pdzd8 depletion in an age-related neurodegenerative
disease model. To test this, we turned to an Alzheimer’s disease fly model where the
expression of pathogenic Aβ42 has been shown to cause neural dysfunction, due in part
to oxidative stress (Rival et al, 2009).

Since increased MERCs have also been associated with Alzheimer’s disease (Area-
Gomez et al, 2012; Del Prete et al, 2017), we first used the SPLICS reporter to
determine the number of MERCs in larval axons. Consistent with other cellular and
organismal models, flies expressing the Aβ42 showed an increase in SPLICS puncta,
indicating MERCs are increased in axons of this model of Alzheimer’s disease (Figure
6A-B). Flies expressing Aβ42 in neurons exhibit a significant climbing defect in young
flies that worsens rapidly with age (Figure 6C)(Crowther et al, 2005). However, pan-
neuronal pdzd8 knockdown in combination with Aβ42 was sufficient to substantially
ameliorate the decline in locomotor activity observed in young and 10-day-old Aβ42
flies (Figure 6C). Thus, reducing pdzd8-mediated MERCs is protective in this
progressive neurodegenerative disease model with increased MERCs.

Discussion

Here we have identified and characterized the putative Drosophila homolog of the
newly discovered mammalian MERC tethering protein Pdzd8 (Hirabayashi et al,
2017). The sequence divergence between Pdzd8 and its yeast paralog Mmm1 and the
additional domains present in Pdzd8 (Wong & Levine, 2017), made the relationship
between these paralogs difficult to identify (Wideman et al, 2018). While the conserved
predicted domain structure strongly suggests Drosophila CG10362 encodes the fly
homolog of mouse Pdzd8, the low, 21 % overall sequence identity of these proteins
raises the interesting possibility of evolution of more species-specific functional
specialisation.
Using RNAi, we characterized the effects of depletion of pdzd8 in Drosophila with a focus on neurons where this protein is most highly expressed. Knockdown of pdzd8 reduces contacts between the ER and mitochondria in epidermal cells measured using super resolution microscopy of ER and mitochondria labeled with fluorescent reporters, in motor neurons, monitored by the contact site reporter SPLICS, and in the soma of adult neurons using transmission electron microscopy. These data suggest that pdzd8, like its mammalian ortholog, functions as a tether between ER and mitochondria. The only other neuronally expressed tethering protein that has been characterized in flies is the Drosophila ortholog of Mfn2, Marf (Hwa et al., 2002). However, the analysis of Marf is complicated by its additional roles in mitochondrial and ER morphology (Debattisti et al., 2014; El Fissi et al., 2018; Sandoval et al., 2014). While mammalian Ptdz8 is expressed in a range of tissues (Hirabayashi et al., 2017), in flies, pdzd8 mRNA expression is low outside the nervous system. Knockdown of pdzd8 and expression of a synthetic tether therefore provided a unique opportunity to simply and selectively examine the function of MERCs in Drosophila neurons.

We found increased ER-mitochondrial tethering in neurons strongly impairs climbing ability and reduces lifespan of flies. Such detrimental effects are consistent with previous reports using this line (Basso et al., 2018). Other similar manipulations have also been shown to result in dopaminergic neuron loss (Lee et al., 2018c) and detrimental effects on sleep in ventral lateral neurons (Valadas et al., 2018). Here we also show highly abnormal NMJ development, with increased MERCs and smaller, but otherwise structurally intact NMJs upon pdzd8 knockdown. The function of the yeast paralog Mmml and of its Synaptotagmin-like mitochondrial lipid-binding protein (SMP) domains, suggests that disruptions in lipid transfer due to less pdzd8 might contribute to these developmental defects (Jeong et al., 2017; Kawano et al., 2018; Shirane et al., 2020), however whether lipid biogenesis defects could contribute to a protective effect of pdzd8-RNAi in neurons remains an open question.

Transport of mitochondria is intimately linked to the health of neurons, at least in the peripheral nervous system (De Vos et al., 2008; Habauer, 2017; Misgeld & Schwarz, 2017). Although there is some evidence that increased MERCs may be directly associated with decreased mitochondrial motility (Krols et al., 2018), this has not been shown in neurons which are particularly sensitive to mitochondrial transport imbalance (Maday et al., 2014). Our data suggests axonal mitochondrial transport defect contributes to the detrimental effects of increased tethering and adds to the evidence that efficient mitochondrial transport is essential for healthy aging neurons, as seen in many models of neurodegenerative motor disorders (Baldwin et al., 2016).

In contrast to the detrimental effects of increased tethering, reducing MERCs by knockdown of pdzd8 in Drosophila neurons dramatically delayed age-associated decline in locomotor activity and significantly extended median lifespan compared to control animals. Since decline in mitochondrial transport is proposed to contribute to neuronal aging (Vagnoni & Bullock, 2018), we hypothesized that reducing tethering in the aging flies might be protective by allowing sustained mitochondrial motility (Mattedi & Vagnoni, 2019). However, we detected no change in the percentage of motile mitochondria in larval neurons with reduced pdzd8 expression. Since knockdown of pdzd8 also prolonged the survival of flies fed mitochondrial toxins, this suggests that the protective effects of reducing MERCs might instead be result from...
more efficient clearance of the damaged mitochondria accumulating with age or more acutely by feeding flies mitochondrial toxins.

To explore the clearance of damaged mitochondria, we examined the age dependence of mitophagy using the mitoQC reporter. Clearance of damaged cellular components by autophagy is critical for the maintenance of healthy neurons with age (Stavoe & Holzbaur, 2019). More specifically, the clearance of damaged mitochondria via mitophagy is also thought to be a key factor in healthy aging of neurons (Ma et al., 2018; Pickrell & Youle, 2015; Whitworth & Pallanck, 2017). It remains unclear, however whether, increased or decreased mitophagy in neurons is protective during aging (Montava-Garriga & Ganley, 2020) and if specific neuronal populations are more susceptible to mitochondrial turnover. Studies examining mitophagy in aging brains have produced inconsistent results, with differences observed between brain regions and cell types (Cornelissen et al., 2018; Lee et al., 2018b; Sun et al., 2015). Consistent with our previous result in dopaminergic neurons (Lee et al., 2018b), we also did not detect any age-related change in mitophagy levels in the cell bodies in the brains of control flies.

There is accumulating evidence showing that axonal mitophagy occurs in cultured mammalian neurons (Ashrafi et al., 2014; Zheng et al., 2019), but other results suggest axonal maintenance in flies may not require mitophagy (Cao et al., 2017). There is evidence of mitophagy in vivo in neurites in young mice (McWilliams et al., 2018; McWilliams et al., 2019), but mitoQC has not previously been used to detect mitophagy in neurites during aging (Cornelissen et al., 2018). Using mitoQC, we detected a significant amount of mitophagy in wing axons of control flies which increased with age. Our results suggest mitophagy in axon but not cell bodies increases with age, but future experiments should specifically examine variations in mitophagy with age across brain regions, cell types and even other subcellular compartments such as dendrites.

Here we provide the first in vivo evidence that mitophagy may be regulated by pdzd8-mediated MERCs. Mitophagy levels did not change in young flies with less pdzd8, but when aged, these flies displayed significant increases in mitophagy in both soma and axons. There is also accumulating evidence that defects in mitophagy contribute to the early stages of Alzheimer’s disease (AD) (Kerr et al., 2017; Lee et al., 2019) and boosting mitophagy has been shown to be protective in worm and mouse AD models (Fang et al., 2019). Here we show increased MERCs in an AD fly model consistent with the altered MERCs identified in both patients and cell models of AD (Area-Gomez et al., 2012). Expression of pdzd8-RNAi in the AD flies significantly slowed their age-associated decline in climbing, consistent with reduced MERCs allowing more efficient mitophagy and removal of damaged mitochondria.

Mitochondrial dysfunction and associated metabolic changes are considered a hallmark of aging (Lopez-Otin et al., 2013). Boosting mitophagy has been suggested to increase lifespan and protect animals from mitochondrial toxins in both D. melanogaster (Aparicio et al., 2019; Rana et al., 2017) and C. elegans (Palikaras et al., 2015). Early work on starvation induced autophagosome biogenesis showed MERCs could contribute to the formation of both the autophagosome (Hailey et al., 2010; Hamasaki et al., 2013) and the mitophagosome (Yang & Yang, 2013). As postmitotic, long-lived cells with complex morphologies, neurons require more careful regulation of
mitophagy than other cell types (Evans & Holzbaur, 2019), but the mechanisms that regulate this process in neurons are still poorly understood (Evans & Holzbaur, 2020).

MERCs mediated by pdzd8 may limit the rate of mitophagy, analogous to the protective role that mitochondrial fusion is thought to play during starvation-induced autophagy (Gomes et al., 2011; Rambold et al., 2011; Rana et al., 2017; Twig et al., 2008). Recently it was reported that increasing several different types of MERC proteins can slow toxin-induced mitophagy in non-neuronal cultured cells (McLelland & Fon, 2018; McLelland et al., 2018). There is also some recent evidence that MERCs might be directly involved in regulation of mitophagy in neurons (Puri et al., 2019). Pdzd8 has also been found to mediate contacts between the ER and lysosomes (Guillen-Samander et al., 2019) and three way contacts between the ER, mitochondria and late endosomes (Elbaz-Alon et al., 2020; Shirane et al., 2020). This may indicate that Pdzd8, like other MERC proteins, can play multiple roles at different organelle contacts, or that Pdzd8 is functional at three-way contacts between the ER, lysosomes and mitochondria (Wong et al., 2019). Pdzd8 function at any of these contact sites could potentially alter mitophagy and future work should explore the mechanisms behind our observed increases in mitophagy.

While our data suggests that increased mitophagy contributes to the protective effects of the loss of pdzd8-mediated MERCs, we have not ruled out contributions from the other specialized functional and signalling roles of MERCs in metabolic regulation, lipid biogenesis and calcium handling (Lee et al., 2018a; Rowland & Voeltz, 2012). The increased mitophagy we detected in aged neurons expressing pdzd8-RNAi correlates with improved survival and aged locomotor activity, suggesting that reducing the extent of pdzd8-mediated MERCs may facilitate more efficient mitophagy and removal of damaged mitochondria from aging neurons. We propose that reducing pdzd8-mediated MERCs may be protective in aging neurons by allowing more efficient turnover of mitochondria as damage accumulates with age. As regulators of mitophagy, manipulating MERCs may provide an avenue for enhanced mitochondrial quality control to help promote healthy aging of neurons.

**Methods**

**Husbandry**

Flies were raised under standard conditions at 25°C on food containing agar, cornmeal, molasses, malt extract, soya powder, propionic acid, nipagin and yeast in a 12 h:12 h light:dark cycle.

**Genetics**

*Drosophila* lines were obtained as indicated in Table 1, or generated as described below. All mutant lines used in this study were backcrossed to an isogenic w^1118^ strain (RRID:BDSC 6326), for 4-6 generations before use. For all integration events, multiple independent lines were initially isolated, verified by PCR and assessed for consistent effects before selecting a single line of each integration site for further study. Wherever possible inert UAS lines such as UAS-LacZ-RNAi and UAS-mitoCherry are used as dilution controls to ensure equal numbers of UAS constructs in control and experimental conditions. Unless otherwise stated male flies were used in all experiments.
**New lines**

**SPLICS**

The SPLICSs construct with an 8-10 nm range from Tito Cali & Marisa Brini (Cieri et al., 2017), was amplified from pSYC-SPLICSs-P2A using (TAAGCAAGCGGCGCTGATTTAGGTGACACTATAG) and T7 forward primer (TAATACGACTCACTATAGGG) and cloned into pUAST-AttB between NotI and XbaI sites. Flies were injected by BestGene to insert into attP16 (II) and attP2 (8622, III) and the attP16 site gave better signal and so was used in this work. The number of puncta produced in the axon bundles driven by nSyb-GAL4 varied with a consistently more puncta than in the central axons bundles in the peripheral bundles (Fig S2B).

**pdzd8-HA**

pdzd8-HA was synthesized by Genewiz based on the cDNA Genbank sequence LD34222 (AY118553.1) (Sayers et al., 2020), and cloned into pUAST.attB between EcoRI and Xbal. The University of Cambridge Department of Genetics Fly Facility generated lines by injection of this construct into the attP40 landing site.

**Climbing**

The startle induced negative geotaxis (climbing/locomotor) assay was performed as described previously (Andreazza et al., 2019). Briefly, a maximum of 23 males were placed into the first tube of a countercurrent apparatus, tapped to the bottom, and given 10 s to climb 10 cm. This procedure was repeated five times (five tubes), and the number of flies that has remained into each tube counted and the climbing performance expressed as a climbing index (Greene et al., 2003). The same flies were aged and assayed again on the indicated days post-eclosion.

**Lifespan**

For lifespan experiments groups of approximately 20-25 males were collected with minimal time under anesthesia (CO2), placed into separate vials with food and maintained at 25°C. Flies were transferred into fresh vials every 2-3 days, and the number of dead flies were recorded. Percent survival was calculated using https://flies.shinyapps.io/Rflies/. To assess lifespan in a diet with restricted nutrients flies were raised in standard conditions then transferred to tubes containing food made from 5 % sucrose and 1 % agar and flipped every 2-3 days. Lifespans in the presence of mitochondrial toxins and hydrogen peroxide were also performed on food made from 5 % sucrose and 1 % agar cooled to less than 50°C before adding toxin at 1:1000. Rotenone (Sigma R8875) was dissolved in DMSO (1 mM final concentration) and antimycin A (Sigma A8674) (4 μg/mL final concentration) dissolved in 70 % ethanol. Flies in toxin assays were starved for 5 h before being placed on food containing toxins. Flies in rotenone assays were monitored twice a day and flipped every two days. Flies in antimycin A assays were monitored three times a day and flipped every two days.

**Fluorescence Microscopy**

Imaging of larval axons was performed as described in (Wang & Schwarz, 2009) with the following variations: wandering third instar larvae were pinned at each end dorsal side up to a reusable Sylgard (Sigma 761028) coated slide using pins (Fine Science Tools FST26002-10) cut to ~5 mm and bent at 90°. The larvae were cut along the dorsal midline using micro-dissection scissors. Internal organs were removed with forceps without disturbing the ventral ganglion and motor neurons. Larvae were then covered in dissection solution (Godena et al., 2014). The cuticle was then pulled back with four
additional pins. The anterior pin was adjusted to ensure axons are taut and as flat as possible for optimal image quality.

Movies were taken using a Nikon E800 microscope with a 60x water immersion lens (NA 1.0 Nikon Fluor WD 2.0) and an LED light source driven by Micromanager 1.4.22 Freeware (Edelstein et al., 2014). A CMOS camera (01-OPTIMOS-F-M-16-C) was used to record 100 frames at a rate of 1 frame per 5 s (8 min 20 s total). Axons were imaged within 200 μm of the ventral ganglion in the proximal portion of the axons and no longer than 1 h after dissection. Movies were converted into kymographs using Fiji (Schindelin et al., 2012) and mitochondrial motility quantified manually with the experimenter blinded to the condition.

For SPLICS imaging in axon bundles at least three ROI 50 μm x 12 μm were quantified per animal and averages for each larva were plotted. For SPLICS quantification puncta intensity varied considerably so blinded manual counting was used.

To image NMJs larvae were dissected as described above and fixed for 20 min in 4 % formaldehyde in PBS. After blocking for 1 h in 1 % BSA/0.3 % Triton X-100/PBS solution, anti-HRP was added at 1:500 and samples agitated gently overnight at 4°C. After three washes in 0.3 % Triton X-100/PBS at room temperature, samples were incubated with Alexa Fluo 594 at 1:500 for 1 h in 1% BSA/0.3% Triton X-100/PBS solution. Samples were then washed in 3x in PBS before being mounted in Prolong Diamond. NMJs were imaged on a Nikon Eclipse TiE inverted microscope with appropriate lasers using an Andor Dragonfly 500 confocal spinning disk system, using an iXon Ultra 888 EMCCD camera (Andor), coupled with Fusion software (Andor) using a 60x NA 1.49 objective. NMJs on muscle 4 from segments A3 and A4 (NMJs on these segments are the same size (Nijhof et al., 2016)) were captured in Z stacks with 0.3 μm step size and analysed using Imaris (x64 9.2.0) to determine NMJ volume, mitochondrial volume and mitochondrial number.

For MitoQC imaging, samples were fixed for 30 min in 4 % formaldehyde (16 % VWR 100503) diluted in pH 7.0 PBS. Adult brains were mounted in Prolong Diamond Antifade Mountant (Thermofisher P36961) using spacers and imaged on a Carl Zeiss LSM880 confocal laser-scanning system on an Axio Observer Z1 microscope (Carl Zeiss), coupled with ZEN software (Carl Zeiss) using a 100x Plan-APOCHROMAT /1.4 oil DIC objective. Images are shown in false colour with magenta puncta representing mCherry signal indicating where the reporter is in an acidic environment of a lysosome and the GFP has been quenched (Allen et al., 2013).

Imaging of wings was performed as described in Vagnoni & Bullock (Vagnoni & Bullock, 2016). Briefly, flies were anaesthetized with CO₂, and immobilised with their wings outstretched on a cover glass with a fine layer of Halocarbon oil (VWR). A second cover glass was then added on top of the fly to stabilize the sample. Live imaging in the wing nerves was performed using a Nikon spinning disk system essentially as described previously (Morotz et al., 2019). The mitoQC puncta were annotated with Fiji using the Cell Counter plugin and quantified with the experimenter blinded to the genotype.

For imaging of the larval epidermal cells, the larvae were dissected as described above but the nervous system was also removed before fixation. The samples were washed in
PBS and the muscles were then removed (Tenenbaum & Gavis, 2016). The dissected
filets were mounted in Prolong Diamond Antifade Mountant using No. 1.5H High
Precision Deckglaser cover slips and placed under a weight for 24 h. N-Actin
Structured Illumination Microscopy) imaging was performed on a Nikon Ti Eclipse
with an Andor DU-897 X-5835 camera and SR Apo TIRF 100x (NA1.5) objective run
using NIS-Elements 4.60. Images were analysed in Fiji (Schindelin et al, 2012) using
the Coloc2 plugin.

Transmission Electron Microscopy
Transmission electron microscopy (TEM) was performed at Cambridge Advanced
Imaging Center (CAIC). Brains of 2 day old adult flies were fixed in 2 %
gluteraldehyde/ 2 % formaldehyde in 0.1 M sodium cacodylate buffer pH 7.4
containing 2 mM CaCl2 and 0.1 % Tween20 (based on method described in (Celardo
et al, 2016)), overnight at 4°C. Samples were then washed 5x with 0.1 M sodium
cacodylate buffer and then treated with osmium for 2 days at 4°C (1 % OsO4, 1.5 %
potassium ferrocyanide, 0.1 M sodium cacodylate buffer pH 7.4). Samples were then
washed 5x in distilled water and treated with 0.1 % aqueous thiocarbohydrazide for 20
min in the dark at room temperature. Samples were washed another 5x in distilled water
then treated with osmium a second time for 1 h at room (2 % OsO4 in distilled water).
Samples were then washed another 5x in distilled water before being treated with uranyl
acetate bulk stain for 3 days at 4°C (2 % uranyl acetate in 0.05 M maleate buffer pH
5.5). After a final 5x wash in distilled water, samples were dehydrated in 50/70/95/100
% ethanol, 3x in each for at least 5 min each. Dehydration was completed by two further
treatments with 100 % dry ethanol, 2x in 100 % dry acetone and 3x in dry acetonitrile
for at least 5 min each. Quetol resin mix (12 g Quetol 651, 15.7 g NSA, 5.7 g MNA,
0.5 g benzylidimethylamine) made with an equal volume of 100 % dry acetonitrile and
samples placed in this mix for 2 h at room temperature. Samples were then incubated
in pure Quetol resin mix for 5 days, exchanging the samples to fresh resin mix each
day. After 5 days the brains were embedded in coffin moulds and cured at 60°C for at
least 48 h. Ultrathin sections were cut on a Leica Ultracut E at 70 nm thickness. Sections
were mounted on 400 mesh bare copper grids and viewed in a FEI Tecnai G20 electron
microscope run at 200 keV using a 20 µm objective aperture. Quantification of the
percentage of clearly identifiable mitochondria in contact with ER was performed
manually as described in (Celardo et al, 2016) and the experimenter was blinded to the
genotype.

qPCR
Five female wandering third instar larvae per sample were washed briefly in 1xPBS,
placed in RNAse free tubes and frozen on dry ice. Larvae were homogenized in Trizol
and RNA isolated by phenol:chloroform extraction and isopropanol precipitation.
DNAse treatment using Invitrogen TURBO DNA-free rigorous procedure was
performed before measuring RNA concentration with a Qubit® RNA HS Assay Kit
(Molecular Probes, Life Technologies). Reverse transcription reactions used 1.32 µg of
RNA using SuperScript III Reverse Transcriptase (Invitrogen) with Oligo(dT)23VN as
per manufacturer’s instructions. The resulting cDNA was used for qPCRs using
PowerUp SybrGreen (Applied Biosystems A25742). Primers for pdzd8 were PDZD8-
F TTCTGTGTGGCTTCTCCTGG, PDZD8-R TTGAGGAACTGCAGACTGATC
designed using RealTime qPCR Assay Entry (idtdna.com), aTub84B (Fwd:
TGGGCCCCGTCTGGACCACAA, Rev: TCGCCGTACCAGGACTCCAT), vkg (Fwd:
CGAGGATGTTACCCAGAGATC, Rev: TGCGTCCCTTGATTCCTTTG), COX8
(Fwd: CAGAGCCGTTCGCCAGTC, Rev: CTTGTCGCCCCTTGATGCT), and Rpl32
(Fwd: AAACGCGGTTCTGCATGAG, Rev: GCCGCTTCAAGGGACAGTATCTG)
were used as housekeeping genes with their values combined to compare knockdown
to the geometric mean (Taylor et al, 2019).

ATP
ATP levels were measured in two- and twenty-day-old fly heads with 40 flies per
genotype and three biological replicates. The ATP levels were measured as described
in Tufi, Gleeson et al. (Tufi et al, 2019) with minor modifications. Briefly, heads were
homogenized in 6 M guanidine-Tris/EDTA extraction buffer and subjected to rapid
freezing in liquid nitrogen. Luminescence produced from homogenates mixed with the
CellTiter-Glo Luminescent Cell Viability Assay (Promega) was measured with a
SpectraMax Gemini XPS luminometer (Molecular Devices) and normalized to total
protein, quantified using the Pierce BCA method (Thermo Fisher Scientific).

Quantification & Statistical analysis
Statistical analyses were performed using GraphPad Prism 8 software. Data are
reported as mean ± 95 % CI unless otherwise stated in figure legends. Climbing was
assessed using a Kruskal-Wallis non-parametric test with Dunn’s post-hoc correction
for multiple comparisons. Lifespans were compared using log-rank Mantel-Cox tests.
Number of flies and p-values are reported in the figure legends.

Mitochondrial transport was analysed using ordinary one-way ANOVA and Holm-
Sidak’s multiple comparison. ATP measurements were analysed by two-tailed t-test.
Values are not significantly different to controls unless otherwise stated.

SCope (http://scope.aertslab.org/) was used to visualize transcriptome data from the
unfiltered adult fly brain dataset (Davie et al, 2018).

Table S1: Fly Lines

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<th>Label</th>
<th>Line</th>
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<th>Reference</th>
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<td>(Park et al, 2003)</td>
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### Table S2: Genotypes in figures

<p>| Figure 1 (B, C) | LacZ-RNAi/+; nSyb/+ pdzd8-RNAi/nSyb |
| Figure 1 (D, E) | SPLICS/LacZ-RNAi; nSyb/+  SPLICS/+; pdzd8-RNAi/nSyb |
| Figure 1 (F, G) | ER-Tom, mitoGFP/ LacZ-RNAi; da/+ ER-tom, mitoGFP/+; da/pdzd8-RNAi |
| Figure 2 (A, C) | LacZ-RNAi/+; nSyb/+ pdzd8-RNAi/nSyb |
| Figure 2 (B, D) | LacZ-RNAi/+; nSyb/+ tether/nSyb |
| Figure 3 (A-D)  | LacZ-RNAi/+; nSyb/+ pdzd8-RNAi/nSyb |
| Figure 4 (A-E)  | CCAP/+; mitoGFP/+ CCAP/+; pdzd8-RNAi/mitoGFP |
| Figure 5 (A-F)  | OK371, mitoGFP/LacZ-RNAi OK371, mitoGFP/+; pdzd8-RNAi/+ |
| Figure 6 (A-B)  | LacZ-RNAi/SPLICS; nSyb/+ Aβ/SPLICS; nSyb/+ |
| Figure 6 (C)    | LacZ-RNAi/+; nSyb/+ LacZ-RNAi/Aβ; nSyb/+ LacZ-RNAi/Aβ; pdzd8-RNAi/nSyb |
| Figure S1 (C)   | Act/LacZ-RNAi (females) Act/+; pdzd8-RNAi/+ (females) |
| Figure S1 (D)   | ER-Tom, mitoGFP/LacZ-RNAi, nSyb/+ ER-Tom, mitoGFP/+; nSyb/pdzd8-RNAi |
| Figure S1 (E)   | ER-Tom, mitoGFP/ LacZ-RNAi; da/+ ER-tom, mitoGFP/+; da/pdzd8-RNAi |
| Figure S2 (B)   | SPLICSs/+; nSyb-GAL4/+ |
| Figure S2 (D)   | SPLICS/LacZ-RNAi; nSyb/+ SPLICS/+; tether/nSyb |</p>
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<th>Figure S3 (A)</th>
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**Figure Legends**

**Figure 1.**

**Expression of pdzd8-RNAi reduces mitochondria-ER contacts.** (A) Domain organisation of Drosophila pdzd8 (CG10362) compared to mouse Pdzd8 showing percentage identities of conserved domains based on Clustal Omega alignments. Overall percentage identity of the amino acid sequences is 21 %. SMP (Synaptotagmin-like mitochondrial lipid-binding proteins) 33 % identical, PDZ (PSD95/DLG/ZO-1) 36 % identical, C1 (C1 protein kinase C conserved region 1 also known as Zn finger phorbol-ester/DAG-type signature) 47 % identical; TM: predicted transmembrane domain, CC: coil-coil domain. (B) Electron microscopy images of 2-day-old adult brains showing representative images of ER, mitochondria and MERCs in soma from nSyb> LacZ-RNAi and nSyb> pdzd8-RNAi flies. Scale bar 500 nm. Mitochondria without identifiable ER contacts marked with magenta *, mitochondria forming ER contact marked with yellow * with yellow arrow indicating contact location, organelles that did not contain clear cristae are marked with a cyan * and were excluded from the analysis (C) Percentage of mitochondria in contact with the ER from controls and pan-neuronal nSyb> pdzd8-RNAi flies quantified from EM images of 2-day-old adult brains. N = 3 brains per genotype. (D) SPLICS puncta indicating MERCs in axon bundles of larval neurons from controls and nSyb> pdzd8-RNAi flies. Quantified puncta highlighted with V. (E) Quantification of SPLICS puncta in C. n = 11 animals per genotype, p=0.0198, unpaired t-test with Welch’s Correction, scale bar 5 μm. (F) Representative binarized SIM images of ER (green) and mitochondria (purple) in larval epidermal cells from controls and da> pdzd8-RNAi flies. Scale bar 500 nm. (G) Quantification of colocalization of ER and mitochondria using Mander’s Correlations compared using an unpaired t-test with Welch’s Correction. p=0.012.

**Figure 2.**

**Lifespan and locomotor activity changes in aged flies with pan-neuronal (nSyb) driven alterations in tethering.** (A, B) Locomotor activity of flies was assessed during aging by negative geotaxis climbing assays on the indicated days. n>50 flies per genotype. Flies expressing (A) pdzd8-RNAi or (B) synthetic tether were compared to LacZ-RNAi controls. (C, D) Lifespans in standard growth conditions and food. (C) Flies expressing pdzd8-RNAi were compared to LacZ-RNAi controls. n = 97, 108 per genotype, median survival 44 vs 52 days, p<0.0001. (D) Flies expressing the synthetic tether were compared to LacZ-RNAi controls. n = 74, 85 per genotype, median survival 52 vs 33 days, p<0.0001.

**Figure 3.**

**Knockdown of pdzd8 protects flies against mitochondrial toxins.** Flies expressing pan-neuronal nSyb> pdzd8-RNAi were compared to nSyb> LacZ-RNAi controls when aged on a restricted diet of food containing 1 % agar with 5 % sucrose. (A) Lifespan

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with dietary restriction alone. N = 62 vs 64, median survival 20 days vs 19 days, difference ns. (B) Lifespan with addition of 5% hydrogen peroxide. Median survival: 63 h vs 74 h, n = 67, 74, p<0.0002. (C) Lifespan with addition of 1 mM rotenone. Median survival: 11 days vs 20 days, n = 66, 57, p<0.0001. (D) Lifespan with addition of 5 µg/mL antimycin A. Median survival: 74 h vs 68 h, n = 72, 68, p=0.0002.

**Figure 4.**
Knockdown of pdzd8 in larval neurons causes minor defects while increasing MERCs is detrimental to in axonal mitochondria size and motility. (A) Representative images of mitochondrial morphology and distribution in larval axons. Mitochondria were detected using CCAP>mitoGFP in controls and pdzd8-RNAi expressing larvae. Scale bar 5 µm. (B,C) Mitochondrial length (B), and mitochondrial density (C) in the larval axons shown in A were analysed using ordinary one-way ANOVA and Holm-Sidak’s multiple comparisons. n = 10, 10, 13 animals, data points represent different axons, all differences ns. (D) Representative kymographs showing motility of CCAP>mitoGFP signal in controls and pdzd8-RNAi expressing larvae. Stationary mitochondria appear as vertical lines, moving mitochondria form diagonal lines in anterograde or retrograde directions, t = 500s (E) Quantification of mitochondrial transport shown in D, analysed using ordinary one-way ANOVA and Holm-Sidak’s multiple comparisons, n = 14-25, p<0.0001. (F) Representative images of NMJs and mitochondria of controls and pdzd8-RNAi labelled using OK371>mitoGFP. magenta = mitoGFP, green = anti-HRP (neuronal membrane), scale bar 10 µm. (G-I) Quantifications of NMJ volume, p=0.0036, (G), mitochondrial volume, p=0.002 (H) and mitochondrial density (I) were compared using an unpaired t-test with Welch’s Correction.

**Figure 5.**
Pan-neuronal pdzd8-RNAi increases mitophagy during aging. (A) Representative images of MitoQC signal in wandering L3 larval ventral ganglia. magenta = mCherry, green = GFP, images show a single plane of a Z-stack, scale bar 2 µm. (B) Quantification of MitoQC puncta shown in (A), n = 9, differences ns. (C) Representative images of MitoQC signal in adult brains in two- and ten-day old flies, magenta = mCherry, green = GFP, scale bar 5 µm, image shows a single plane of a Z-stack. (D) Quantification of MitoQC signal in adult brains and compared using an unpaired t-test with Welch’s correction (n = 7-9, p=0.0072) (E) The Representative images of MitoQC signal in 14 day-old fly wings. Only mCherry signal (magenta) is shown for clarity, white outlines edges of wing nerves, scale bar 5 µm. (F) Quantification of MitoQC signal in aged fly wings at 2, 14 and 30 days post eclosion using a one-way ANOVA with Holm-Sidak’s multiple comparisons. n (2 days) = 33, 26 (14 days) = 24, 31, (30 days) = 32, 12.

**Figure 6.**
Reducing pdzd8-mediated MERCs rescues the locomotor defects in an Alzheimer’s disease Drosophila model. (A) CCAP>SPLICS signal in larval axons expressing Aβ42 compared to controls. Quantified puncta highlighted with V. (B) Quantification of SPLICS signal in (A) using an unpaired t-test with Welch’s correction, n = 6, p=0.008. (C) Aged climbing assay of flies expressing nSyb>Aβ42, n>75 flies were aged and climbed on the indicated days. p <0.001, 0.002, <0.001, <0.001.
Figure S1.
Expression and tissue specificity of pdzd8. (A) Tissue specific expression of pdzd8 in 7 day old adult males from (Leader et al, 2018) FPKM: fragments per kilobase of exon model per million reads mapped providing a normalized estimation of gene expression based on RNA-seq data. (B) SCope transcriptome data from the unfiltered adult fly brain dataset. (C) Relative abundance of pdzd8 transcript in controls compared to Tub>pdzd8-RNAi normalised to the relative to geometric mean of housekeeping genes αTub84B, vkg, COX8 and Rpl32; p=0.0134. (D) Representative SIM images of ER (green) and mitochondria (purple) in L3 larval neurons, scale bar 5 µm. (E) Representative SIM images of ER and mitochondria in larval epidermal cells, scale bar 5 µm. (F) Close up inlays (F'-F''') of example MERCs in Figure 1E identified in nSyb>LacZ-RNAi brains imaged using electron microscopy. Scale bar 500 nm.

Figure S2.
SPLICS and tether constructs used in this study. (A) Cartoon of SPLICS targeting and mode of action (Created with BioRender.com). (B) Density of SPLICS puncta in axons is different in different axon bundles, scale bar 5 µm. (C) Cartoon of the synthetic tether construct targeting and mode of action (Created with BioRender.com). (D) nSyb>Tether construct targeting and mode of action in larval axons, scale bar 5 µm, n = 9 per genotype.

Figure S3.
Phenotypic characterization of altered tethering in motor neurons. (A) Motor neuron-specific aged climbing assay showing OK371>pdzd8-RNAi compared to LacZ-RNAi controls. (B) pdzd8-RNAi rescues aged climbing defect resulting from pdzd8 overexpression. (C) nSyb>tether expression resulted in severely deformed NMJs on muscle 4 and made it impossible to distinguish NMJs type 1s and 1b synaptic boutons (therefore mitochondrial density could not be quantified). (D) ATP levels in fly heads analysed showing mean ± standard deviation, n = 3, 40 flies per replicate, compared using a two-tailed t-test, all differences ns.

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Conflict of Interests
The authors declare that they have no conflict of interest.

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Author Contributions
VLH, LM-F, SA, JP, FM and AV designed and/or performed experiments, and analysed the data. VLH wrote the manuscript with input from all authors. AJW, AV and FP supervised the work.

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Figure 1. Expression of *pdzd8*-RNAi reduces mitochondria-ER contacts.

A

B

LacZ-RNAi  pdzd8-RNAi

*C* Mitochondrion without ER contact  *M* Mitochondrion with ER contact  *U* Unassigned organelle  *Y* MERC

C

% mitochondria in contact with the ER

0  20  40  60

LacZ-RNAi  pdzd8-RNAi  201  195

D

E

SPLICS puncta per 50 µm

0  5  10  15  20

LacZ-RNAi  pdzd8-RNAi

F

G

Manders' Correlation

0.0  0.2  0.4  0.6  0.8

LacZ-RNAi  pdzd8-RNAi
Figure 2. Lifespan and locomotor activity changes in aged flies with pan-neuronal (nSyb) driven alterations in tethering.
Figure 3. Knockdown of *pdzd8* protects flies against mitochondrial toxins.

**A** Starvation

**B** Hydrogen peroxide

**C** Rotenone

**D** Antimycin A

*ns* = not significant

**Statistical significance**

- ***p* = 0.0002
- **** *p* < 0.0001
- *** *p* = 0.0002
Figure 4. Knockdown of *pdzd8* in larval neurons causes minor defects while increasing MERCs is detrimental to axonal mitochondria size and motility.
Figure 5. Pan-neuronal pdz8-RNAi increases mitophagy during aging.

A) mitoQC:  
- LacZ-RNAi
- pdz8-RNAi

B) Box plots showing puncta per cell for LacZ-RNAi and pdz8-RNAi.

C) Images showing 2 days and 20 days for LacZ-RNAi and pdz8-RNAi.

D) Box plots showing puncta per ROI (400 µm²) for LacZ-RNAi and PDZD8-RNAi at 2 days and 20 days.

E) Images showing LacZ-RNAi and pdz8-RNAi at different time points.

F) Bar charts showing puncta per 50 µm for LacZ-RNAi and pdz8-RNAi at 2 days, 14 days, and 30 days.
Figure 6. Reducing pdzd8-mediated MERCs rescues the locomotor defects in an Alzheimer’s disease *Drosophila* model.
Figure S1. Expression and tissue specificity of pdzd8.

A. FPKM values for different tissues.

B. Image showing expression patterns with different colors indicating different tissues.

C. Bar graph showing relative pdzd8 mRNA levels with error bars.

D. Confocal images comparing LacZ-RNAi and pdzd8-RNAi treatments.

E. Magnified view of LacZ-RNAi and pdzd8-RNAi treatments.

F. Transmission electron microscopy images showing mitochondrial and ER contacts.
Figure S2. SPLICS and tether constructs used in this study.

A

SPLICS

~8-10 nm

mitochondrial targeting sequence
ER membrane anchor
GFP β strands 1-10
GFP β strand 11

B

Peripheral axon bundles projecting to other more proximal segments
Central axons

C

synthetic tether

~5 nm

mitochondrial targeting sequence
ER membrane anchor

D

control
tether

E

Puncta per 50 µm

p < 0.0001

control tether
Figure S3. Phenotypic characterization of altered tethering in motor neurons.

A

![Climbing index graph](image)

B

![Climbing index graph](image)

C

![Immunofluorescence images](image)

D

![ATP normalized graph](image)