Gain-of-function mutations in the ALS8 causative gene VAPB have detrimental effects on neurons and muscles

Citation for published version:
Sanhueza, M, Zechini, L, Gillespie, T & Pennetta, G 2014, 'Gain-of-function mutations in the ALS8 causative gene VAPB have detrimental effects on neurons and muscles' Biology Open, vol 3, no. 1, pp. 59-71. DOI: 10.1242/bio.20137070

Digital Object Identifier (DOI):
10.1242/bio.20137070

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Biology Open

Publisher Rights Statement:
© 2013. Published by The Company of Biologists Ltd
This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution and reproduction in any medium provided that the original work is properly attributed.

General rights
Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.
Gain-of-function mutations in the ALS8 causative gene VAPB have detrimental effects on neurons and muscles

Mario Sanhueza*, Luigi Zechini*, Trudy Gillespie and Giuseppa Pennetta‡

Centre for Integrative Physiology, Euan MacDonald Centre for Motor Neuron Disease Research, School of Biomedical Sciences, University of Edinburgh, Edinburgh EH8 9XD, UK

*These authors contributed equally to this work
‡Author for correspondence (g.pennetta@ed.ac.uk)

Biology Open 3, 59–71
doi: 10.1242/bio.20137070
Received 4th November 2013
Accepted 26th November 2013

Summary

Amyotrophic Lateral Sclerosis (ALS) is a motor neuron degenerative disease characterized by a progressive, and ultimately fatal, muscle paralysis. The human VAMP-Associated Protein B (hVAPB) is the causative gene of ALS type 8. Previous studies have shown that a loss-of-function mechanism is responsible for VAPB-induced ALS. Recently, a novel mutation in hVAPB (V234I) has been identified but its pathogenic potential has not been assessed. We found that neuronal expression of the V234I mutant allele in Drosophila (DVAP-V260I) induces defects in synaptic structure and microtubule architecture that are opposite to those associated with DVAP mutants and transgenic expression of other ALS-linked alleles. Expression of DVAP-V260I also induces aggregate formation, reduced viability, wing postural defects, abnormal locomotion behavior, nuclear abnormalities, neurodegeneration and upregulation of the heat-shock-mediated stress response. Similar, albeit milder, phenotypes are associated with the overexpression of the wild-type protein. These data show that overexpressing the wild-type DVAP is sufficient to induce the disease and that DVAP-V260I is a pathogenic allele with increased wild-type activity. We propose that a combination of gain- and loss-of-function mechanisms is responsible for VAPB-induced ALS.

© 2013. Published by The Company of Biologists Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution and reproduction in any medium provided that the original work is properly attributed.

Key words: ALS, Drosophila, Genetics, Neurodegeneration, VAPB

Introduction

Amyotrophic Lateral Sclerosis (ALS) is a progressive neurodegenerative disorder characterized by the degeneration of motor neurons leading to muscle atrophy, spasticity and, eventually, death. ALS is traditionally classified into two categories: familial ALS (FALS) and sporadic ALS (SALS) (Pasinelli and Brown, 2006). Missense mutations (P56S and T46I) at the N-terminal region of the human VAMP-associated protein B gene (hVAPB) have been identified in both patients with FALS and SALS (Nishimura et al., 2004; Funke et al., 2010; Millecamps et al., 2010; Landers et al., 2008; Chen et al., 2010). VAP proteins contain an N-terminal domain, which is highly homologous to the nematode major sperm protein (MSP), a central domain that forms a coiled-coil structure and a C-terminal transmembrane domain (Nishimura et al., 1999; Kaiser et al., 2005).

Members of the highly conserved VAP family have been involved in a number of seemingly unrelated functions including regulation of ER morphology, lipid transfer, vesicular trafficking and dendritic morphogenesis (Soussan et al., 1999; Skehel et al., 2000; Amarilio et al., 2005; Teuling et al., 2007; Lev et al., 2008; Peretti et al., 2008; Kuippers et al., 2013a). DIVAP, the Drosophila orthologue of hVAPB, controls synaptic structure, microtubule cytoskeleton architecture and composition of post-synaptic glutamate receptors (Pennetta et al., 2002; Chai et al., 2008). In C. elegans and Drosophila, the MSP domain is cleaved, secreted and acts as a ligand for Robo and Lar-like receptors to control muscle mitochondria morphology, localization and function (Tsuda et al., 2008; Han et al., 2012). Previous studies have implicated ALS mutant alleles in an abnormal unfolded protein response and in the disruption of the anterograde axonal transport of mitochondria (Chen et al., 2010; Kanekura et al., 2006; Langou et al., 2010; Suzuki et al., 2009; Gkogkas et al., 2008; Mörotz et al., 2012). Transgenic expression in Drosophila of either DVAP-P58S or DVAP-T48I, two ALS-linked alleles, mirrors major hallmarks of the human disease including neurodegeneration, aggregate formation, locomotion defects and chaperone upregulation (Chai et al., 2008; Chai et al., 2010). Several lines of evidence support the notion that P56S mutation is a loss-of-function mutation, possibly via a dominant negative effect. P56S mutant protein forms aggregates in which the wild-type protein is recruited (Teuling et al., 2007). This together with the analysis of mutant phenotypes associated with transgenic expression of the P56S protein in several disease models, confirms that P56S is a loss-of-function mutation (Teuling et al., 2007; Chen et al., 2010; Chai et al., 2008;
Moreover, we show that nuclear abnormalities represent a highly conserved transmembrane domain with an Isoleucine (V234I in humans and V260I in Drosophila) (van Blitterswijk et al., 2012). Surprisingly, we found that nuclear abnormalities at the position 234 of the highly conserved transmembrane domain in one ALS patient who has also a pathogenic repeat expansion in C9ORF72 (chromosome 9 open reading frame 72), another ALS causative gene. The mutation within the hVAPB gene replaces the Valine at the position 234 of the highly conserved transmembrane domain with an Isoleucine (V234I in humans and V260I in Drosophila) (van Blitterswijk et al., 2012). Moreover, no information is available about the dominant or recessive inheritance of the mutation in humans and theoretical predictions based on bioinformatics approaches give contradictory results as to whether V234I mutation has a damaging effect on the function of the protein (van Blitterswijk et al., 2012; Ingre et al., 2013). To directly assess the pathogenic effect of this mutation and to unveil potential new mechanisms of disease pathogenesis, we carried out a series of functional studies in a well-established Drosophila model of ALS.

Here we report that transgenic expression of DVAP-V260I recapitulates major hallmarks of the human disease including aggregate formation, reduced viability, neuromuscular defects, abnormal locomotion behavior, neurodegeneration and upregulation of the heat-shock-mediated stress response. Moreover, we show that nuclear abnormalities represent a novel aspect of ALS pathogenesis as expression of DVAP-V260I either in neurons or muscles induces disruption in nuclear architecture, position and shape. Surprisingly, we found that transgenic expression of DVAP-V260I at the Drosophila larval neuromuscular junction (NMJ) induces an increase in the number of synaptic boutons and a decrease in their size. This phenotype is highly reminiscent of the phenotype associated with the neuronal overexpression of DVAP-WT protein and opposite to that of DVAP loss-of-function mutations and transgenic expression of the DVAP-P58S allele in neurons (Pennetta et al., 2002; Forrest et al., 2013). In addition, overexpression of DVAP-WT protein either in neurons or muscles induces phenotypes similar, albeit milder, than those associated with V260I expression. Altogether these data lead to the fundamental conclusion that DVAP-V260I is a pathogenic allele with an increased wild-type activity and that a combination of loss- and gain-of-function mechanisms are responsible for VAP-induced ALS. In conclusion, on the basis of the data reported here and those on previously identified mutations, we propose that VAPB levels or/and activity must be tightly regulated to keep neurons and muscles healthy as slight disturbances in one direction or the other may induce cell dysfunction and death.

Results
Presynaptic expression of either DVAP-V260I or DVAP-WT transgenes leads to an overproduction of small synaptic boutons

The V234I mutation in the hVAPB gene recently identified in one ALS patient but not in a large number of healthy controls, is located within the conserved and functionally important transmembrane domain of VAP proteins (van Blitterswijk et al., 2012) (supplementary material Fig. S1). Conversely, the previously identified mutations, P56S and T46I, are localized in a highly conserved stretch of amino acids within the MSP domain (supplementary material Fig. S1C).

Presynaptic expression of the ALS-linked allele DVAP-P58S induces a NMJ phenotype characterized by a decrease in the number of boutons and an increase in their size (Chai et al., 2008; Ratnaparkhi et al., 2008). These data prompted us to examine the effect of DVAP-V260I expression on bouton formation and synaptic structure. We generated several transgenic lines expressing DVAP-V260I using the bipartite UAS/GAL4 system and the panneural elav-Gal4 as a driver (Lin and Goodman, 1994). To examine basic synaptic morphology, NMJs were stained with anti-HRP antibodies that label and allow for visualization of the entire presynaptic membrane. Changes in NMJ structure were assessed by counting the number of synaptic boutons at muscles 12 and 13 of the abdominal segment A3. This analysis revealed that transgenic expression of DVAP-V260I has clear consequences on the expansion of boutons as compared to controls. We were surprised to observe that, in this case, there is an increase in the total number of synaptic boutons when compared to controls (514 ± 3 in elav;DVAP-V260I versus 272 ± 2 in controls; P < 0.001) (Fig. 1A,C). The size of boutons, however, is dramatically reduced.

We have previously shown that an increase in the number of boutons with a concomitant decrease in their size is associated with the presynaptic overexpression of DVAP-WT protein and that this phenotype is highly dependent on the dosage of the DVAP (Pennetta et al., 2002). To further confirm and extend this analysis, we generated a number of additional transgenic lines expressing different levels of the DVAP-WT protein and found that they all exhibited similar synaptic phenotypes. We selected a weak line (DVAP-WT1) in which the number of boutons on the muscles 12 and 13 of the segment A3 is 1.3-fold higher than in controls and a strong DVAP-WT2 line that exhibits a 1.7-fold increase in the number of synaptic boutons (Fig. 1A,C; P < 0.001 in both cases).

We also noticed that in all transgenic lines the overall morphology of the synapse has changed. In control NMJs, synaptic boutons within a branch, resemble a string of beads with boutons connected to one another by a short neuritic process (Fig. 1B). In contrast, in larvae overexpressing either DVAP-V260I or DVAP-WT transgenes, many small boutons appear to bud off from a central bouton or from neuronal processes connecting two boutons (Fig. 1B, arrows). Clusters of smaller boutons surrounding central larger boutons are especially common in DVAP-V260I overexpressors (Fig. 1B,D). Boutons of similar size and morphology have been described in other overgrowth synaptic phenotypes and they have been called “satellite” boutons (Torroja et al., 1999; Franco et al., 2004; Kamimura et al., 2013). To correct for variations in total bouton number, we quantified the amount of “satellite” boutons for each genotype as a percentage of the mean total bouton number. We found that presynaptic expression of DVAP-V260I using the neuronal elav-Gal4 driver, induces a 2.6-fold increase in the percentage of “satellite” boutons compared to controls (Fig. 1B,D; P < 0.001). Neuronal expression of either DVAP-WT or DVAP-WT2 transgenes using the same Gal4 driver, leads to 1.7-fold and 2.2-fold increase in satellite bouton number, respectively (Fig. 1B,D; P < 0.001 in both cases). In summary, expression of either DVAP-WT or DVAP-V260I transgenes induces a dramatic change on the overall morphology of the
synapse due, not only, to an increase in the total number of boutons but also to an elevated number of “satellite” boutons. It was surprising to find an overgrowth synaptic phenotype associated with DVAP-V260I as transgenic expression of the previously identified DVAP-P58S allele in neurons leads to an opposite phenotype characterized by a decrease in the number of boutons and an increase in their size (Pennetta et al., 2002; Chai et al., 2008). The DVAP-P58S-induced phenotype is similar to that associated with DVAP loss-of-function mutations (Pennetta et al., 2002) leading to the hypothesis that DVAP-P58S is a loss-of-function allele (Pennetta et al., 2002; Ratnaparkhi et al., 2008; Chai et al., 2008). Accordingly, we found that in DVAP-P58S transgenics the endogenous wild-type protein accumulates in the aggregates distributed along the nerves and is depleted from its normal synaptic localization (supplementary material Fig. S2E, arrow). In particular, quantification of DVAP-positive staining at NMJs of DVAP-P58S expressing neurons results in nearly 45% decrease in the DVAP immunoreactivity compared to controls (supplementary material Fig. S2A,E,F; P<0.001). We then assessed the distribution and the local expression levels of DVAP at NMJs of both the DVAP-V260I and the two DVAP-WT lines. In agreement with the observed synaptic phenotypes, we found that elaGal4-driven expression of every transgene results in a rather homogenous distribution of DVAP at the synapse and quantification of DVAP-positive immunoreactivity revealed that in every case the amount of total protein is higher than in controls (supplementary material Fig. S2). Specifically, the increase in DVAP signal is 2.2-fold higher than in controls for the DVAP-WT2 line while it is 1.7-fold and 1.8-fold over the endogenous levels for DVAP-V260I and DVAP-WT1, respectively (supplementary material Fig. S2, P<0.001 in all cases). It is worth to note that although DVAP-WT1 and DVAP-V260I lines express comparable amounts of their respective transgenes, expression of the DVAP-WT1 transgene leads to synaptic phenotypes and indeed to a number of other phenotypes (see below) that are milder than those associated with the transgenic expression of the mutant allele (compare supplementary material Fig. S2 and Fig. 1). Furthermore, phenotypes induced by the expression of DVAP-V260I in eight tested lines are consistently more severe than those observed in a comparable number of lines expressing the DVAP-WT transgene (data not shown). Taken together, these results indicate that an increase in the levels of DVAP wild-type protein is sufficient to induce an overgrowth synaptic phenotype and that the DVAP-V260I mutant allele acts as a hypermorphic allele as it has an increased ability to promote bouton formation at the NMJ.

Presynaptic transgenic expression of DVAP-WT and DVAP-V260I leads to bouton expansion by affecting microtubule cytoskeleton architecture

Formation of synaptic boutons at axon terminals is achieved via microtubules invading and promoting bouton budding from the plasma membrane (Roos et al., 2000). Within the Drosophila NMJ, excess of microtubule loop formation is associated with overgrowth phenotypes, particularly those with an increased number of “satellite” boutons (Franco et al., 2004; Kamimura et al., 2013). Thus, we asked whether the increase in bouton number in NMJs expressing either DVAP-WT or DVAP-V260I transgenes is related to alterations in microtubule cytoskeleton architecture. To address this question, we carried out immunostainings against the Drosophila MAP1B homologue Futsch, a marker of neuronal microtubule structures (Hummel et al., 2000; Roos et al., 2000). DVAP-WT1, DVAP-WT2 and DVAP-V260I transgenes were each expressed in neurons using the elaGal4 driver and the number of Futsch-positive microtubule loops was quantified for every genotype. Presynaptic overexpression of DVAP-V260I shows a drastic increase in the number of Futsch-positive loops as
compared to controls (38±0.4% versus 11±0.4% in controls, P<0.001) (Fig. 2A,B,E). A significant increase in the proportion of boutons containing a microtubule loop-like structure was also observed following the presynaptic expression of DVAP-WT transgenes (26±0.5% for DVAP-WT1 and 31±0.5% for DVAP-WT2). However, in both cases, the change is less than that associated with DVAP-V260I overexpressing mutants (Fig. 2C,D,E). Of note, an increase in number of boutons with more disorganized microtubule structure represented by splayed/punctuate Futsch-positive immunostaining was reported in larvae expressing DVAP-P58S and in DVAP loss-of-function mutants (Pennetta et al., 2002; Forrest et al., 2013). As expected, these NMJs exhibit an opposite synaptic phenotype characterized by a decrease in number of boutons and an increase in their size (Pennetta et al., 2002; Forrest et al., 2013). Thus, expression of either DVAP-V2601 or DVAP-WT transgenes in neurons, affects the organization of microtubule architecture, albeit at a different degree, with consequences on synaptic bouton formation and division.

Targeted expression of either DVAP-V2601 or DVAP-WT transgenes induces aggregate formation and nuclear abnormalities

DVAP is a ubiquitous protein and pathological alterations in muscles during ALS have been considered to result from toxicity of the mutant protein in motor neurons rather than a direct effect of the pathogenic protein on striated muscles. In order to explore whether the muscle is a direct target of DVAP-V2601 and DVAP-WT toxicity, the expression of either the wild-type or mutant allele was selectively targeted to striated muscles, by using the muscle-specific driver BG57-Gal4 (Budnik et al., 1996).

Abnormal aggregates containing disease proteins are a common hallmark of several neurodegenerative disorders. We therefore stained dissected NMJs with DVAP antibodies to assess the localization of DVAP proteins in controls and in muscles expressing either the DVAP-WT or the DVAP-V2601 transgenes. We found that while in controls DVAP is expressed throughout the muscle, in DVAP-V2601 muscles numerous and punctuate, DVAP-immunopositive inclusions are present (Fig. 3A,B, Fig. 5A,B, staining in red). Interestingly, similar inclusions are also found in both the weak (DVAP-WT1) and strong (DVAP-WT2) transgenic lines overexpressing the DVAP-WT protein (Fig. 3C,D, Fig. 5C,D, staining in red). These data indicate that overexpression of the DVAP-WT protein is sufficient to induce the formation of aggregates and that DVAP-V2601 is a pathogenic allele.

Nuclear defects have been correlated with ageing as well as with a number of pathological manifestations in humans including Parkinson’s disease (Worman et al., 2010; Liu et al., 2012). To determine whether nuclear architecture or/and positioning are affected in our ALS fly model, we visualized nuclei in muscles overexpressing either DVAP-V2601 or DVAP-WT transgenes using a nuclear marker and an anti-lamin antibody. We found that in controls, nuclei are evenly spaced along the muscle fibre while in muscles expressing either the DVAP-V2601 or the DVAP-WT transgenes, nuclei are closely associated and exhibit a tendency to form clusters (Fig. 3). To better evaluate the relative position of nuclei within a muscle, a “nearest-neighbor” analysis was conducted for each genotype (see Materials and Methods for details). We found that, compared to controls, the average shortest distance between nuclei is severely reduced in muscles overexpressing the DVAP-V2601 transgene (P<0.001) while the overexpression of the DVAP-WT transgenes has a significant but milder effect even in the case of the strongest DVAP-WT2 line (Fig. 3E; P<0.01 for DVAP-WT1 and P<0.001 for DVAP-WT2).

We also noticed a drastic deterioration in nuclear architecture. In particular, overexpression of either DVAP-WT or DVAP-V2601 transgenes results in deformed nuclei with an elongated structure. To quantify this phenotype, we measured the width to length ratio of nuclei within muscles per each genotype. As the nuclei in control muscles have a distinct round shape, the width to length ratio is close to 1. Any deviation from this value indicates a loss of circularity and therefore a change in shape. We found that nuclei within muscles overexpressing the DVAP-V2601 transgene are less round compared to nuclei in control muscles (Fig. 3A,B,F; P<0.001) and a similar, albeit milder effect, was observed in muscles overexpressing the DVAP-WT transgenes (Fig. 3A,C,D,F; P<0.01 for DVAP-WT1 and P<0.001 for DVAP-WT2). We also found that nuclei in muscles expressing the DVAP-V2601 transgene display a marked enlarged nuclear volume compared to controls (Fig. 3G; P<0.001) as do the nuclei

Fig. 2. Expression of either DVAP-V2601 or DVAP-WT transgenes in neurons affects synaptic microtubule cytoskeleton. (A) Representative images of branches of NMJs of third instar elav-Gal4/+ control larvae, (B) elav;DVAP-V2601, (C) elav;DVAP-WT1 and (D) elav;DVAP-WT2 larval NMJs labeled with antibodies against Futsch to show microtubule loops. Arrows in every panel indicate examples of Futsch loops. Scale bars: 10 μm. (E) Quantitative assessment of Futsch-positive loops at A2 and A3 muscle 4 NMJs for each indicated genotype. The highest increase in the percentage of boutons exhibiting looped Futsch staining (relative to the total number of boutons for each NMJ) was observed when DVAP-V2601 was expressed presynaptically. Asterisks denote statistical significance compared to controls (***P<0.001).
of muscles expressing the $DVAP-WT$ transgenes, although to a lesser extent (Fig. 3G; $P < 0.001$ for the $DVAP-WT^1$ and $P < 0.01$ for $DVAP-WT^2$). Intriguingly, a fraction of DVAP protein that is predominantly a cytoplasmic protein was found to localize to the nucleus in $DVAP-V260I$ overexpressing muscles while in both $DVAP-WT$ lines this aberrant DVAP localization is nearly absent...
(Fig. 3H–K). A nuclear localization for DVAP has never been described before but previous studies have shown that translocation from the cytoplasm to the nucleus of α-synuclein associates with increased toxicity and neurodegeneration in Parkinson’s disease (Kontopoulos et al., 2006). Moreover, an increased number of small and condensed pyknotic nuclei was predominantly observed in muscles overexpressing the DVAP-V260I transgene while such nuclei are significantly less in the weak DVAP-WT1 line (Fig. 3, white arrows).

Finally, we noticed that larger and malformed nuclei are also observed in neurons of larval brains expressing either the DVAP-WT or DVAP-V260I transgenes and again the phenotype appears to be more severe when the mutant transgene is expressed (Fig. 4). In addition, expression of DVAP-V260I in neurons also induces accumulation of prominent inclusions that appear to be very large and intensely immunoreactive to DVAP antibodies (Fig. 4A,B). In contrast, in neurons of the weakest DVAP line (DVAP-WT1), small and isolated foci accumulating DVAP were observed. Although not directly quantified, the number of these foci appear to be increased in neurons of the strongest DVAP-WT line (Fig. 4C,D). Interestingly, no small pyknotic nuclei were observed in these neurons that present an aberrant nuclear architecture.

Taken together, these data demonstrate that transgenic expression of either DVAP-V260I or DVAP-WT in neurons as well as in muscles elicits the formation of aggregates and a severe disruption in the architecture, size and positioning of nuclei.

Induction of the heat-shock response in DVAP-WT and DVAP-V260I overexpressors

In many neurodegenerative diseases including ALS, aggregates are thought to be formed from the accumulation of misfolded proteins that trigger the upregulation of chaperone proteins (Vabulas et al., 2010). We assessed whether this was also the case for larvae expressing either DVAP-V260I or DVAP-WT transgenes in their muscles. In controls, Hsp70 is barely detectable and results in faint and diffuse staining throughout the cytoplasm (Fig. 5A). In transgenic animals expressing the DVAP-V260I construct specifically in the striated muscles, there is an increase in the expression levels of Hsp70 that appears to concentrate in puncta mainly localized inside the nucleus (Fig. 5B). In transgenic muscles expressing the DVAP-WT transgenes, an increase in the accumulation of puncta immunoreactive to the Hsp70 was observed. These puncta are partially localized into the nucleus in muscles expressing the strongest wild-type transgene DVAP-WT2 (Fig. 5D). Conversely, in muscles expressing the weak DVAP-WT1 transgene, puncta are still present in the cytoplasm, although the majority of them preferentially accumulate in the perinuclear region (Fig. 5C). The phenomenon of the Hsp70 relocating from the cytoplasm to the nucleus during a stress response is poorly understood; however, it is clear that this relocation is directly linked and is part of the cellular response to stress (Vabulas et al., 2010). Taken together, these data indicate that in DVAP-V260I as well as in DVAP-WT overexpressors, a stress response mediated by heat-shock proteins...
has been induced and that the intensity of this response directly correlates with the toxicity of the overexpressed protein.

Expression of DVAP-V260I and DVAP-WT in adult flies induces reduced viability, compromised locomotion behavior and neuromuscular defects.

We showed that expression of DVAP-V260I and DVAP-WT transgenes in larval neurons and muscles causes a number of phenotypes closely mirroring the human pathology. We then sought to test whether the same transgenes have any effect on the adult neuromuscular junctions. We first expressed the DVAP-V260I or the DVAP-WT transgenes in the adult nervous system by using the panneural driver elav-Gal4. Flies expressing either the DVAP-V260I transgene or any of the two wild-type transgenic constructs, are able to pupate and adults eclose from pupae. However, the rate of eclosion is different depending on the overexpressed transgene (Fig. 6A). Flies expressing the DVAP-WT transgene have a small but statistically significant decrease in the rate of eclosion (Fig. 6A; P<0.05) while for flies expressing the strongest wild-type allele, DVAP-WT², the rate of eclosion is 75% (Fig. 6A; P<0.001). A strong reduction in the number of flies eclosed as viable adults is observed when the expression of the DVAP-V260I was targeted in the adult neurons. In this case only 35% of pupae eclose to give an adult fly (Fig. 6A). However, all flies, irrespective of their genotypes, display distinctive postural and locomotion defects as viable adults. While in controls, wings run dorsal and parallel to the body, surviving mutants exhibit different wing posture phenotypes including droopy and held-up wings (Fig. 6B–D). Additionally, surviving adult flies are severely uncoordinated and within a few days following eclosion, get stuck to the food and die. When the expression of either DVAP-WT or DVAP-V260I constructs is targeted to all muscles by using the MHC-Gal4 driver, the progeny exhibits the same wing posture phenotypes as for the neuronal expression of the same transgenes but their eclosion ratio is similar for every transgene and closer to that of controls (supplementary material Fig. S3). Similar adult phenotypes have been described in a number of fly models of human neurodegenerative diseases including Fragile X syndrome and Parkinson’s disease and have been attributed to neuromuscular dysfunction (Zhang et al., 2001; Clark et al., 2006).

Targeted expression of either DVAP-WT or DVAP-V260I transgenes in the adult Drosophila eye causes neuronal degeneration.

Neuronal degeneration in Drosophila models of human neurodegenerative disorders including ALS is usually assessed in the adult fly eye. To determine whether overexpression of DVAP-WT protein and transgenic expression of DVAP-V260I causes neurodegeneration, we used the ey-Gal4 driver to specifically target the expression of these transgenes in the adult fly eye (Halder et al., 1998; Hauck et al., 1999). Adult fly eyes expressing either DVAP-WT or DVAP-V260I transgenes display a range of structural abnormalities. As compared with
controls, eyes expressing either the *DVAP-WT* or *DVAP-V260I* transgenes are reduced in size and show a marked roughness over the entire surface (Fig. 7A). A striking aspect of the eye phenotype induced by the overexpression of either *DVAP-WT* or the *DVAP-V260I* transgenes, is the high degree of heterogeneity in the severity of the phenotype within the same line and even within the same fly as, in many cases, one of the eyes exhibits a more extensive degeneration than the eye on the opposite side. We therefore quantified the phenotype in both control and overexpressor lines by measuring the eye size in a comparable number of flies for every genotype. Overall, the eye size of *DVAP-V260I* transgenic flies is less than 50% of the control size while the measured reduction in eye size for *DVAP-WT*\(^1\) and *DVAP-WT*\(^2\) was 30% and 20%, respectively (Fig. 7C; P<0.001 in all cases).

Histopathological analysis of paraffin embedded sections of transgenic eyes expressing either *DVAP-WT* or *DVAP-V260I* confirmed the degeneration of cells beneath the external surface of the eye (Fig. 7B). In particular, photoreceptor morphology is disrupted and several vacuoles are visible. Vacuolization has

---

**Fig. 6.** *DVAP-V260I* and *DVAP-WT* overexpressors display reduced viability and wing postural defects. (A) Eclosion rate of flies of the designated genotype. (B) elav-Gal4/+ control flies with a dorsal wing posture. (C) Held-up wings and (D) droopy wing flies. See text for comments. Asterisks denote statistical significance. **P<0.001, *P<0.05.**

**Fig. 7.** Expression of either *DVAP-V260I* or *DVAP-WT* transgenes in adult *Drosophila* eyes induce neurodegeneration. (A) Stereomicroscope images of ey-Gal4/+ control eyes and eyes of the indicated genotypes. (B) Frontal sections of control and transgenic eyes of the indicated genotypes stained with H&E. White arrows point to vacuoles in photoreceptors while the black arrowheads indicate areas of extensive tissue degeneration. (C) Quantification of the eye surface area of every genotype. Scale bars: 50 μm. Asterisks denote statistical significance (**P<0.001).**
been observed in a number of fly models of neurodegenerative diseases and the appearance of vacuoles has been correlated to neurodegeneration (Chan and Bonini, 2000). Of note, the strongest phenotype for both the reduction in the eye size and the degeneration of the internal eye is that associated with the expression of the DVAP-V260I transgene (Fig. 7). The internal tissues of transgenic eyes expressing the DVAP-WT protein are disrupted in a similar way, albeit to a lesser extent, especially when the weakest line DVAP-WT1 is considered (Fig. 7B). In conclusion, these data indicate that increasing the level of expression of the DVAP-WT protein is sufficient to trigger neurodegeneration, although transgenic expression of the mutant DVAP-V260I allele appears to be more efficient in inducing neuronal cell death.

Discussion

Recently, a novel mutation in the gene hVAPB has been identified in one ALS patient who is also known to have a pathogenic repeat expansion in the C9ORF72 gene. The missense mutation replaces the Valine with an Isoleucine at codon 234, a highly conserved residue within the C-terminal transmembrane domain of VAP proteins (van Blitterswijk et al., 2012). In Drosophila, the transmembrane domain of DVAP binds the phosphoinositide phosphatase Sac1 to control phosphoinositide levels whose upregulation is responsible for neurodegeneration and a number of synaptic phenotypes including altered synaptic morphology and mislocalization of several post-synaptic markers (Forrest et al., 2013). Moreover, in mammals the transmembrane-mediated interaction of VAPB with the ER–Golgi recycling protein YIF1A controls dendritic remodeling in the central nervous system (Kuijpers et al., 2013a). It is therefore plausible that mutations targeting this important functional domain may have a pathogenic effect.

Here we report that transgenic expression of DVAP-V260I in Drosophila, induces mutant phenotypes that closely mirror major hallmarks of ALS such as structural remodeling at NMJs, formation of aggregates, abnormal locomotion behavior, reduced viability, neuromuscular defects and upregulation of the heat-shock-mediated stress response. In addition, muscles and neurons expressing the DVAP-V260I transgene exhibit striking abnormalities in nuclear positioning, shape and size. Interestingly, nuclear abnormalities have never been linked to ALS pathogenesis.

Preceding identified disease mutations in hVAPB are loss-of-function mutations, possibly by a dominant negative mechanism (Ratnaparkhi et al., 2008; Forrest et al., 2013; Kuijpers et al., 2013a). We were therefore surprised to find that transgenic expression of DVAP-V260I construct consistently produces similar but more severe phenotypes than those associated with the overexpression of the wild-type protein. Collectively, our results show that a gain-of-function mechanism is responsible for VAPB-induced ALS and that the V234I is a pathogenic allele with an increased wild-type activity. In conclusion, we propose that both gain- and loss-of-function effects play an important role in driving the damage, which ultimately leads to cellular downfall and clinical expression of the disease.

Nuclear abnormalities in ALS pathogenesis

Mispositioning of nuclei has been associated with decreased cell survival in vertebrate central nervous system (Tsujikawa et al., 2007). In striated muscles, myonuclei appear as rounded structures evenly spaced along muscle fibers separated from the bulk of the cytoplasm, which contains densely arranged myofibrils. The mechanisms establishing and maintaining this highly ordered distribution of nuclei and its relevance to muscle function is not known but several human myopathies are caused by mutations in genes such as Nesprins that control the correct position and distribution of nuclei in muscles (Puckelwartz et al., 2010). Furthermore, Nesprin-1 knockout mice exhibit 50% lethality and those surviving display progressive muscle wasting and abnormal gait (Puckelwartz et al., 2009). Nuclear architecture defects have been shown to correlate with aging as well as with a number of human diseases, including Parkinson’s disease (Liu et al., 2012).

To study the role that nuclear position and structure might have in ALS, we targeted the expression of DVAP-WT and DVAP-V260I in neurons and muscles. Our morphometric analysis revealed that overexpression of the DVAP-V260I results in deformed nuclei with an elongated structure and a marked enlarged nuclear volume. We also found that in DVAP-V260I transgenic muscles the average distance between nuclei is remarkably reduced and sometimes, nuclei are found to aggregate into clusters. A similar deterioration in nuclear architecture and positioning was observed in muscles overexpressing DVAP-WT transgenes but, overall, the defects are milder than those associated with the expression of DVAP-V260I even when the strongest DVAP-WT2 transgene is expressed. In summary, nuclear abnormalities are more prominent in transgenic conditions associated with a more aggressive disease phenotype.

Finally, small and condensed pyknotic nuclei were observed in muscles expressing DVAP-V260I and to a lesser extent in those overexpressing the DVAP-WT transgene. Although the presence of pyknotic nuclei is a hallmark of apoptosis, we do not believe that the observed nuclear abnormalities are a mere consequence of apoptotic cell death. No condensed, pyknotic nuclei are present in neurons expressing any DVAP-WT transgenes or DVAP-V260I protein at the time when nuclear abnormalities are already visible. These results suggest that defects in nuclear architecture and distribution may precede the appearance of apoptotic hallmarks. Indeed recent studies suggest that alterations in nuclear structure and positioning represent an irreversible trigger of apoptosis rather than a consequence of this process (Liu et al., 2012).

Taken together, these studies identify the nucleus as a novel cellular organelle involved in ALS pathogenesis and open new avenues for the potential development of therapies targeting this fundamental cellular structure.

Expression of either DVAP-V260I or DVAP-WT transgenes recapitulates major ALS-related hallmarks

The data discussed in the previous paragraph suggest that our DVAP-V260I fly model may function as a platform for the discovery and study of novel and, at the moment, elusive aspects of ALS pathogenesis. Our data also suggest that this model recapitulates many of the phenotypes known to be associated with ALS and indeed with a number of other neurodegenerative diseases.

In Drosophila, DVAP interacts with the adhesion molecule Dscam to affect synaptic connectivity and its homologue in mammals, VAPB, is required for intracellular membrane trafficking and normal dendritic growth (Yang et al., 2012;
Kuijpers et al., 2013a). In addition, loss-of-function mutations in the SMN (Survival motor neuron) gene, the causative gene of the motor neuron disease spinal muscular atrophy (SMA), induces defects in axons and synaptic structure (Torres-Benito et al., 2012). Similar results have been reported for the ALS-linked gene VCP (Valosin Containing Protein) as mutations in this gene lead to impaired synaptogenesis and spinogenesis (Wang et al., 2011). To study the effect of DVAP-WT and mutant DVAP-V260I transgenes on synaptic remodeling and structure, we focused on the Drosophila larval NMJ. Morphological analysis shows that at NMJs overexpressing either DVAP-WT or DVAP-V260I transgenes, boutons are smaller, more numerous and tend to form dense clusters especially at the ends of terminal branches. Overgrowth synaptic phenotypes have been associated with alterations in microtubule networks and specifically with an increase in the percentage of boutons containing microtubule loop-like structures (Torroja et al., 1999). Indeed we found that at NMJs expressing DVAP-V260I there is a specific increase in this category of boutons. Similar, but less severe, phenotypes are associated with the transgenic expression of the DVAP-WT protein. In contrast, expression of the ALS-linked allele DVAP-P58S in neurons and loss-of-function mutations in DVAP induce a reduction in number of boutons and an increase in their size. A shift in microtubule structures towards less organized morphologies, which is often associated with undergrowth synaptic phenotypes, was also observed in this transgenic condition (Pennetta et al., 2002). In agreement with these data, we show that there is a local increase in DVAP levels at DVAP-V260I expressing NMJs and a decrease in synapses expressing DVAP-P58S. In summary, we propose that disruption of synaptic remodeling and structure may represent a fundamental and common aspect of disease pathogenesis in ALS and that the DVAP-V260I allele, in contrast to the DVAP-P58S allele, exhibits an increased wild-type activity.

A common feature of neurodegenerative diseases is the presence of misfolded protein aggregates in affected regions of the nervous system. Many neurodegenerative proteins behave like prion proteins (Polymenidou and Cleveland, 2012). In vivo and in vitro studies have shown that these proteins exhibit a high tendency to self-aggregate because of their intrinsic ability to undergo a series of kinetically unfavorable conformational changes to produce misfolded proteins. Increasing the concentration of the native protein favors the formation of aggregates and disease-linked mutations often appear to confer to the protein an increased ability to generate inclusions (Eisele, 2013). Further studies are necessary to assess whether DVAP is a prion-like protein. However, as expected for a prion-like protein, DVAP-WT overexpression is sufficient to induce the formation of aggregates in a dosage-dependent manner and transgenic expression of the mutant DVAP-V260I protein effectively leads to the accumulation of large protein inclusions even at relatively low doses.

In our Drosophila model, muscles expressing the DVAP-V260I transgene, which associates with the most aggressive disease phenotype, exhibit a relocalization of DVAP protein into the nucleus. DVAP remains predominantly localized to the cytoplasm in controls and in all the other transgenic conditions exhibiting a milder disease phenotype. The mechanism by which DVAP enters the nucleus is unclear; however, this relocalization closely resembles the relocalization from the cytoplasm to the nucleus reported for α-synuclein, a protein involved in Parkinson’s disease (PD). Normal and PD-linked α-synucleins promote toxicity by relocating to the nucleus where they bind to histones and inhibit their acetylation. Administration of histone deacetylase (HDAC) inhibitors restores the histone acetylation state and rescues α-synuclein-mediated toxicity (Kontopoulos et al., 2006). Remarkably, administration of HDAC inhibitors to ALS mouse models successfully decreases cell toxicity (Yoo and Ko, 2011).

Implications for ALS pathogenesis

Loss or depletion of VAPB activity in zebrafish, worms and mice induce a relatively mild phenotype possibly because of a functional redundancy between closely related VAP proteins in these models (Kabashi et al., 2013; Han et al., 2012). In Drosophila, only one VAP gene is present and therefore its inactivation leads to the appearance of a more aggressive mutant phenotype (Pennetta et al., 2002; Forrest et al., 2013). Mice expressing the VAPB-P56S transgene have been generated and they exhibit no overt neurodegenerative phenotype (Aliaga et al., 2013; Qiu et al., 2013; Tudor et al., 2010; Kuijpers et al., 2013b). However, a recent report describes degeneration of a subset of motor neurons in mice expressing higher levels of the VAPB-P56S transgene (Aliaga et al., 2013). In addition, neurodegeneration has been associated with the expression of the same transgene in cell culture systems (Langou et al., 2010; Suzuki and Matsuoka, 2011; Teuling et al., 2007). Taken together these considerations further underscore the importance of VAPB dosage in inducing ALS-like phenotypes, including neurodegeneration. It is not uncommon that genetic models fail to reproduce the full range of clinical manifestations associated with a human disease and for this non-human primate models should represent a better option. Genetic models, however, remain invaluable tools to dissect mechanistic aspects of disease pathogenesis.

Studies in several model systems showed that the P56S mutation induces ALS by a loss-of-function mechanism, possibly due to a dominant negative effect (Ratnaparkhi et al., 2008; Tsuda et al., 2008; Suzuki et al., 2009; Forrest et al., 2013; Kuijpers et al., 2013a). Furthermore, a reduction in VAP protein levels have been reported in sporadic ALS patients, SOD1 mutant mice as well in induced pluripotent stem cells derived from ALS patients (Teuling et al., 2007; Anagnostou et al., 2010; Mitne-Neto et al., 2011). The dominant negative effect of P56S mutation is further supported by the observation that the mutant protein accumulates into intracellular inclusions, which also sequester the wild-type protein (Teuling et al., 2007; Chen et al., 2010). Here we uncover a novel mechanism for the disease pathogenesis and we show that increasing the levels and/or activity of DVAP may be detrimental as well. The notion that increased hVAPB levels are sufficient to induce neurodegeneration would be greatly supported by finding that duplications or mutations in the non-coding regions of the hVAPB gene are associated with ALS in humans. For hVAPB, however, such mutations have yet to be identified. These and other studies indicate that, in many cases, a strict dichotomy between loss-of-function versus gain-of-function mechanisms do not respond to a biological reality as both mechanisms seem to be at play (Robberecht and Philips, 2013). In support of this hypothesis, variations in the dosage of the SMN gene have been shown to induce two closely related motor neuron diseases as loss-of-function mutations in SMN cause Spinal Muscular
Atrophy while duplications of the same gene have been linked to ALS (Bürghlen et al., 1995; Blauw et al., 2012).

Finally, our data are compatible with a model in which, in DVAP-V260I-induced ALS, gain- and loss-of-function mechanisms may coexist within the same neuron depending on the subcellular compartment. At the presynaptic compartment, a local increase in DVAP levels induces a remodeling of synaptic morphology and microtubule architecture compatible with a gain-of-function mechanism. Conversely, in neuronal cell bodies the inherent ability of wild-type hVAPB to be included into aggregates that may also function as a sink for other important proteins, suggests that DVAP could be pathogenic by a possible loss-of-function mechanism. The complexity of such a scenario pinpoints to novel opportunities for possible therapies but at the same time makes the task of finding an effective therapeutic strategy particularly challenging.

Materials and Methods

Fly strains and husbandry
Flies were reared in standard cornmeal–molasses medium and maintained at room temperature unless otherwise specified. To drive the expression of DVAP transgenes in neurons, larval muscles and adult eyes we used the elav-Gal4 line (Lin and Goodman, 1994), the BG57-Gal4 line (Budnik et al., 1996) and the ey-Gal4 line (Halder et al., 1998; Hauck et al., 1999), respectively. We used the ey-Gal4 instead of the GMR-Gal4 driver because GMR-Gal4 has been shown to induce eye neurodegeneration on its own (Rezavály et al., 2007). Transgenic expression in adult muscles was achieved by using the MHC-Gal4 line.

Generation of DVAP-WT and DVAP-V260I transgenic lines
Site-directed mutagenesis on DVAP cDNA was carried out using the Quick Change Site Directed Mutagenesis Kit (Agilent, Colorado Springs, CO, USA) following manufacturer’s instructions. Gal4-responsive transgenic lines were constructed cloning DVAP-WT and DVAP-V260I cDNAs in the pUAST vector (Brand and Perrimon, 1993) using the cloning strategy previously described (Pennetta et al., 2002). Constructs were injected into embryonic germ cells and transgenic lines were established following standard procedures (Genetic Services, Inc., Sudbury, MA, USA). Given the strong dosage-dependent effect on mutant proteins, suggests that DVAP could be pathogenic by a possible gain-of-function mechanism. Conversely, in neuronal cell bodies the inherent ability of wild-type hVAPB to be included into aggregates that may also function as a sink for other important proteins, suggests that DVAP could be pathogenic by a possible loss-of-function mechanism. The complexity of such a scenario pinpoints to novel opportunities for possible therapies but at the same time makes the task of finding an effective therapeutic strategy particularly challenging.

Statistical analysis
Data throughout the paper are presented as mean ± s.e.m. A minimum of 10 larvae were analyzed per genotype. At least 20 flies per genotype were analyzed in scoring the adult eye phenotype.

Analysis and quantification of the Drosophila adult eye phenotype
For the eye morphology, two-dimensional images were obtained using a Nikon D5100 DSLR camera attached to a SZX9 Nikon stereomicroscope (Nikon, Kingston upon Thames, Surrey, UK). To quantify the eye surface area images were analyzed with ImageJ software. For histology, flies were raised at 25°C and collected immediately after eclosion. Heads were severed and place in fresh 4% paraformaldehyde in 0.1 M sodium phosphate buffer (pH 7.4) for 24 hours at 4°C. Heads were then washed and placed in 70% ethanol and processed into paraffin using standard histological procedures. 10 μm serial sections were obtained and rehydrated with PBS. Sections were stained with H&E.

Adult phenotypes
Eclosion rates were determined by counting numbers of empty versus full (dead) pupae on the sides of vials in which flies of different genotypes had been allowed to lay for comparable time periods. Food was covered with a thick layer of yeast to prevent flies from desiccating before eclosion. Flies exhibiting an aberrant posture in their wings were collected and stored at –70°C for analysis. The percentage of flies with an aberrant posture was calculated for each genotype. Flies were scored blind using a dissecting microscope to assign phenotype to each fly.

Acknowledgements
We thank the Developmental Studies Hybridoma Bank for monoclonal antibodies; Bing Zhang and Hermann Haberle for fly stocks; Paul Fisher for the polyclonal lamin antibody.

Funding
This work was supported by the Wellcome Trust [grant number: Pennetta8920] and by the Motor Neuron Disease Association [grant number: Pennetta6231].
Author Contributions
L.Z., M.S. and G.P. performed the experiments; M.S., T.G. and G.P. analyzed data; G.P. conceived and designed the experiments, and wrote the paper.

Competing Interests
The authors have no competing interests to declare.

References
Aliaga, L., Lai, C., Yu, J.-H., Chen, S., Shim, H., Sun, L., Xie, C., Yang, W. J., Lin, X., O’Donovan, M. J. et al. (2013). Amyotrophic lateral sclerosis-related PVP56 mutation differentially affects the function and survival of corticospinal and spinal motor neurons. Hum. Mol. Genet. 22, 4293-4305.

Amarillo, R., Ramachandran, S., Sabanay, H. and Lev, S. (2005). Differential regulation of endoplasmic reticulum structure through VAP-Nir protein interaction. J. Biol. Chem. 280, 5934-5944.

Anagnostou, G., Akbar, M. T., Paul, P., Angelinette, C., Steiner, T. J. and de Bellerocche, J. (2010). Vesicle associated membrane protein B (VAPB) is decreased in ALS spinal cord. Neurobiol. Aging 31, 969-985.

Bischof, J., Maeda, R. K., Hediger, M., Karch, F. and Basler, K. (2007). An open transgenesis system for Drosophila using germ-line specific phiC31 integrases. Proc. Natl. Acad. Sci. USA 104, 3132-3137.

Blaauw, P., Bame, F., Fister, V., van Vught, P. W., van Rheenen, W., Verheul, M., Cuppen, E., Veldink, J. H. and van den Berg, I. H. (2012). SMN1 gene splice variants are associated with sporadic ALS. Neurology 78, 776-780.

Brand, A. H. and Perrimon, N. (1993). Targeted gene expression as a means of altering cell fates and generating dominant phenotypes. Development 118, 401-415.

Budnik, V., Kob, Y. H., Guan, B., Hartmann, B., Hough, C., Woods, D. and Gorczyca, M. (1996). Regulation of synapse structure and function by the Drosophila tumor suppressor gene dig. Neuron 17, 627-640.

B¨urglen, L., Spiegel, R., Ignatiu, I., Cobben, J. M., Landrieu, P., Lefebvre, S., Munnich, A. and Melki, J. (1995). SMN gene deletion in variant of infantile spinal muscular atrophy. Lancet 346, 316-317.

Chai, A., Withers, J., Kob, Y. H., Parry, K., Bao, H., Zhang, B., Budnik, V. and Pennetta, G. (2008). VAPB, the causative gene of a heterogeneous group of motor neuron diseases in humans, is functionally interchangeable with its Drosophila tumor suppressor domain. EMBO J. 27, 1979-1988.

Chen, H. Y. and Bonini, N. M. (2000). Drosophila models of human neurodegenerative disease. Cell Death Differ. 7, 1075-1080.

Chen, H. J., Anagnostou, G., Chai, A., Withers, J., Morris, A., Adhikaree, J., Pennetta, G. and de Bellerocche, J. S. (2010). Characterization of the properties of a novel mutation in VAPB in familial amyotrophic lateral sclerosis. J. Biol. Chem. 285, 40266-40271.

Clark, I. E., Dodson, M. W., Jiang, C., Cao, J. H., Huh, J. R., Seol, J. H., Yoo, S. J., Chan, H. Y. and Bonini, N. M. (2007). An Futsch/22C10 is a MAP1B-like protein required for dendritic and axonal growth cone guidance receptors. Neuron 56, 564-569.

Da Fonseca, A., Freitas, C., Ferreira, R., Fonseca, J., Ribeiro, M. and Brandao, A. H. (2013). SOD1, VAP, TARDBP and mutations in familial amyotrophic lateral sclerosis: genotype-phenotype correlations. J. Med. Genet. 47, 554-560.

D’Ambrosio, G., M¨uller, B., Lasch, K., Brinckmann, F., Mann, M., Brinkmann, B. D., Hiesinger, P. R., Meinertzhagen, I. A. and Bellen, H. J. (2010). Characterization of the properties of a novel mutation in VAPB/ALS8 major calcium homeostasis to disrupt axonal transport of mitochondria. Hum. Mol. Genet. 20, 3642-3652.

Disruption of nesprin-1 produces an Emery Dreifuss muscular dystrophy-like disease in mice. J. Cell Sci. 125, 2199-2203.

Doherty, P., Dahan, N., Hirschberg, K. and Lev, S. (2008). VAPB protein family from cellular functions to motor neuron disease. Trends Cell Biol. 18, 282-290.

Doherty, P., Dahan, N., Hirschberg, K. and Lev, S. (2008). VAPB protein family from cellular functions to motor neuron disease. Trends Cell Biol. 18, 282-290.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.

Doherty, P. R., Dahan, N., Hirschberg, K. and Lev, S. (2008). Prion-like spread of protein aggregates in neurodegeneration. Neuron 56, 776-780.
Rezával, C., Werbajh, S. and Ceriani, M. F. (2007). Neuronal death in Drosophila triggered by GAL4 accumulation. *Eur. J. Neurosci.* 25, 683-694.

Robberecht, W. and Philips, T. (2013). The changing scene of amyotrophic lateral sclerosis. *Nat. Rev. Neurosci.* 14, 248-264.

Roos, J., Hummel, T., Ng, N., Klämbt, C. and Davis, G. W. (2000). Drosophila Futsch regulates synaptic microtubule organization and is necessary for synaptic growth. *Neuron* 26, 371-382.

Skehel, P. A., Fabian-Fine, R. and Kandel, E. R. (2000). Mouse VAP33 is associated with the endoplasmic reticulum and microtubules. *Proc. Natl. Acad. Sci. USA* 97, 1101-1106.

Sousan, L., Burakov, D., Daniels, M. P., Toister-Achituv, M., Porat, A., Yarden, Y. and Elazar, Z. (1999). ERG30, a VAP-33-related protein, functions in protein transport mediated by COPI vesicles. *J. Cell Biol.* 146, 301-312.

Suzuki, H. and Matsuoka, M. (2011). Amyotrophic lateral sclerosis-linked mutant VAPB is cleaved, secreted, and acts as a ligand for Eph receptors. *Cell* 133, 963-977.

Tsuji, W., Omori, Y., Biyanwila, J. and Malicki, J. (2007). Mechanism of positioning the cell nucleus in vertebrate photoreceptors. *Proc. Natl. Acad. Sci. USA* 104, 14819-14824.

Tudor, E. L., Galtrey, C. M., Perkinton, M. S., Lau, K. F., De Vos, K. J., Mitchell, J. C., Ackerley, S., Horta, G., Vámos, E., Leigh, P. N. et al. (2010). Amyotrophic lateral sclerosis mutant vesicle-associated membrane protein-associated protein-B transgenic mice develop TAR-DNA-binding protein-43 pathology. *Neuroscience* 167, 774-785.

Valiunas, R. M., Raychaudhuri, S., Hayer-Hartl, M. and Hartl, F. U. (2010). Protein folding in the cytoplasm and the heat shock response. *Cold Spring Harb. Perspect. Biol.* 2, a004390-a004390.

van Blitterswijk, M., van Es, M. A., Koppers, M., van Rheenen, W., Medic, J., Schel涵as, H. J., van der Kooi, A. J., de Visser, M., Veldink, J. H. and van den Berg, L. H. (2012). VAPB and C9orf72 mutations in 1 familial amyotrophic lateral sclerosis patient. *Neurobiol. of Aging* 33, 2950e1-2950e4.

Wang, H. F., Shih, Y. T., Chen, C. Y., Chao, Y. C., Chao, H. W., Lee, M. J. and Hsueh, Y. P. (2011). Valosin-containing protein and neurofibrillar interact to regulate dendritic spine density. *J. Clin. Invest.* 121, 4820-4837.

Worman, H. J., Ostlund, C. and Wang, Y. (2010). Diseases of the nuclear envelope. *Cold Spring Harb. Perspect. Biol.* 2, a000760.

Yang, Z., Huh, S. U., Drennan, J. M., Kathuria, H., Martinez, J. S., Tsuda, H., Hall, M. C. and Clements, J. C. (2012). Drosophila Vap-33 is required for axonal localization of Dscam isoforms. *J. Neurosci.* 32, 17241-17250.

Yoo, Y. E. and Ko, C. P. (2011). Treatment with trichostatin A initiated after disease onset delays disease progression and increases survival in a mouse model of amyotrophic lateral sclerosis. *Exp Neurol.* 231, 147-159.

Zhang, Y. Q., Bailey, A. M., Matthies, H. J., Renden, R. B., Smith, M. A., Speese, S. D., Rubin, G. M. and Broadie, K. (2001). Drosophila fragile X-related gene regulates the MAP1B homolog Futsch to control synaptic structure and function. *Cell* 107, 591-603.
Fig. S1. Identification of the V234I/V260I mutation on VAP proteins. (A) Predicted functional domains in hVAPB: a transmembrane domain (blue) at the C-terminus, a coil-coiled domain (brown) in the middle and a domain (green) at the N-terminus showing a significant homology to the nematode major sperm protein (MSP). The Val234 residue (indicated by an arrow) is located within the transmembrane domain and it has been replaced by an Isoleucine. (B) High degree of conservation of the transmembrane domain in VAP proteins from different species. (C) Sequences of human VAPB (hVAPB) protein and its Drosophila orthologue DVAP were aligned by using the ClustalW version 1.82 alignment program available from EMBL–EBI. Asterisk indicates an identity match; colon indicates conservation between amino acids with strongly similar properties (scoring 0.5 in the Gonnet PAM 250 matrix) and period indicates a conserved substitution between amino acids with weakly similar properties (<0.5 in the Gonnet PAM 250 matrix). Red boxes highlight the amino acid residues changed by the ALS8 causing mutations identified so far. The Drosophila protein (DVAP) exhibits the same functional domains as the hVAPB.

Fig. S2. Synaptic levels of DVAP in different genetic contexts. (A) Control elav/+ NMJs and (B) elav:DVAP-V260I, (C) elav:DVAP-WT1, (D) elav:DVAP-WT2, (E) elav:DVAP-P58S NMJs stained with antibodies specific for DVAP and for anti-HRP. Scale bars: 10 µm. (F) Quantification of synaptic DVAP intensity for the reported genotypes. DVAP fluorescence intensity is presented as a relative ratio of fluorescence intensity values of DVAP against those of controls. Note that in DVAP-P58S transgenic line aggregates are evident in the terminal part of the nerve (arrow in panel E) and the endogenous protein at the synapse has decreased to about 66±4% of the control value. Conversely, compared to controls DVAP levels at the synapse are significantly increased in all the other transgenic lines (P<0.001 in all cases). However, upregulation of DVAP levels in neurons expressing DVAP-V260I is comparable to that induced by the DVAP-WT1 transgene (P<0.05) and significantly lower than that associated with the strongest DVAP-WT2 overexpressing line DVAP-WT (P<0.001). The difference in DVAP levels between DVAP-WT1 and DVAP-WT2 lines was statistically significant (P<0.001).
Fig. S3. Overexpression of *DVAP-WT* and transgenic expression of *DVAP-V260I* in muscles does not affect viability. Eclosion rate of flies of the indicated genotypes. *MHC-Gal4/+* flies were used as controls. *P* > 0.05.