Neurobiology of Disease

Loss of Activity-Induced Mitochondrial ATP Production Underlies the Synaptic Defects in a *Drosophila* Model of ALS

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Mutations in the gene encoding vesicle-associated membrane protein B (VAPB) cause a familial form of amyotrophic lateral sclerosis (ALS). Expression of an ALS-related variant of *vapb* (*vapb*^{P58S}) in *Drosophila* motor neurons results in morphologic changes at the larval neuromuscular junction (NMJ) characterized by the appearance of fewer, but larger, presynaptic boutons. Although diminished microtubule stability is known to underlie these morphologic changes, a mechanism for the loss of presynaptic microtubules has been lacking. By studying flies of both sexes, we demonstrate the suppression of *vapb*^{P58S}-induced changes in NMJ morphology by either a loss of endoplasmic reticulum (ER) Ca²⁺ release channels or the inhibition Ca²⁺/calmodulin (CaM)-activated kinase II (CaMKII). These data suggest that decreased stability of presynaptic microtubules at *vapb*^{P58S} NMJs results from hyperactivation of CaMKII because of elevated cytosolic [Ca²⁺]. We attribute the Ca²⁺ dyshomeostasis to delayed extrusion of cytosolic Ca²⁺. Suggesting that this defect in Ca²⁺ extrusion arose from an insufficient response to the bioenergetic demand of neural activity, depolarization-induced mitochondrial ATP production was diminished in *vapb*^{P58S} neurons. These findings point to bioenergetic dysfunction as a potential cause for the synaptic defects in *vapb*^{P58S}-expressing motor neurons.

Key words: ALS; Drosophila neurobiology; ER calcium channels; mitochondrial ATP production; neurodegeneration; VAPB

Significance Statement

Whether the synchrony between the rates of ATP production and demand is lost in degenerating neurons remains poorly understood. We report that expression of a gene equivalent to an amyotrophic lateral sclerosis (ALS)-causing variant of vesicle-associated membrane protein B (VAPB) in fly neurons decouples mitochondrial ATP production from neuronal activity. Consequently, levels of ATP in mutant neurons are unable to keep up with the bioenergetic burden of neuronal activity. Reduced rate of Ca^{2+} extrusion, which could result from insufficient energy to power Ca^{2+} ATPases, results in the accumulation of residual Ca^{2+} in mutant neurons and leads to alterations in synaptic vesicle (SV) release and synapse development. These findings suggest that synaptic defects in a model of ALS arise from the loss of activity-induced ATP production.

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Introduction

Amyotrophic lateral sclerosis (ALS) is an untreatable neurodegenerative disease characterized by the progressive loss of motor function leading to paralysis and death by respiratory failure (Taylor et al., 2016). Mutations in many genes have been implicated in the onset of the familial forms of the disease, which result in one or more of the following, disrupted nucleocytoplasmic transport, proteostatic imbalance, alterations in RNA metabolism, genome instability, mitochondrial dysfunction, aberrant Ca²⁺ homeostasis, neuronal hyperexcitability, and neuroinflammation (Ling et al., 2013; Selfridge et al., 2013; Taylor et al., 2016; Lin et al., 2017; Frere and Slutsky, 2018). In this study, we sought to examine the mechanisms of neuronal dysfunction in flies expressing a missense variant of vapb, which is equivalent to the human variant that causes a familial form of ALS (ALS8) in humans (Nishimura et al., 2004, 2005; Kanekura et al., 2006; Marques et al., 2006; Landers et al., 2008). We chose vesicle-associated membrane protein B (VAPB) because of the importance of this protein to neuronal viability. In addition to the missense variant that functions via a dominant-negative mechanism (Ratnaparkhi et al., 2008), spinal cords from patients with sporadic cases of ALS exhibit decreased VAPB expression (Anagnostou et al., 2010; Mitne-Neto et al., 2011). Defective VAPB function is also observed in Parkinson's disease (Kun-Rodrigues et al., 2015; Paillusson et al., 2017; Boczonadi et al., 2018).

At the molecular level, VAPB is a single-pass endoplasmic reticulum (ER) membrane protein that has been proposed to regulate mTOR signaling, autophagy, lysosomal acidification, proteasomal degradation/ER quality control, and formation of interorganellar contacts that link the ER to endosomes, peroxisomes, Golgi, and mitochondria (Peretti et al., 2008; De Vos et al., 2012; Deivasigamani et al., 2014; Moustaqim-Barrette et al., 2014; Roulin et al., 2014; Dong et al., 2016; Stoica et al., 2016; Hua et al., 2017; Zhao et al., 2018; Chaplot et al., 2019; Mao et al., 2019; Şentürk et al., 2019). Via its role in the formation of ERmitochondria contacts, VAPB orchestrates the transfer of Ca²⁺ from ER into the mitochondrial matrix, and thereby, regulates ATP synthesis (De Vos et al., 2012; Stoica et al., 2016; Gomez-Suaga et al., 2017; Smith et al., 2017; Xu et al., 2020; Wong et al., 2021).

Expression of the ALS-related variants of vapb in Drosophila neurons has been shown to result in morphologic alterations in both axons and dendrites (Chai et al., 2008; Ratnaparkhi et al., 2008; Kamemura et al., 2021). Although it is known that defects in the development of the axon termini stem from diminished stability of presynaptic microtubules (Chai et al., 2008; Ratnaparkhi et al., 2008), exactly how mutant VAPB disrupts the microtubule cytoskeleton has remained unknown. In order to uncover the underlying molecular mechanism, we sought to examine genetic interactions between the vapb variant and genes encoding ER Ca²⁺ channels whose absence elicits phenotypes that ostensibly resemble those evoked by mutant vapb (Wong et al., 2014). These studies revealed that the phenotypes arising from the expression of mutant vapb are a consequence of presynaptic Ca²⁺ dyshomeostasis, which could stem from the inability of the neurons to synthesize the ATP molecules needed for Ca²⁺ extrusion. Our findings agree with the insights gleaned from in silico modeling of ALS neurons (Le Masson et al., 2014), and provide a possible explanation for the presynaptic defects associated with the expression of the ALS-causing variant of vapb in Drosophila motor neurons.

Materials and Methods

Drosophila husbandry

Flies were reared at 21°C on standard fly food (11 of food contained: 95 g of agar, 275 g of Brewer's yeast, 520 g of cornmeal, 110 g of sugar, 45 g of propionic acid, and 36 g of Tegosept) unless otherwise stated. The following fly lines were obtained from Bloomington *Drosophila* Stock Center: *ok371-GAL4* and *d42-GAL4* (Parkes et al., 1998; Mahr and Aberle, 2006), *RyR*¹⁶ (Sullivan et al., 2000), *UAS-Dcr1* (Dietzl et al., 2007), *UAS-CKII-I.Ala* (*UAS-CKII*^{ala}; Joiner and Griffith, 1997), *UAS-mCherry-mito-OMM* (Vagnoni and Bullock, 2016), and *UAS-itpr*^{RNAi} (*TRiP.JF01957*; Wong et al., 2021). Other strains used in the study were: *UAS-vapb*^{WT} and *UAS-vapb*^{P58S} (Tsuda et al., 2008), *UAS-GCaMP5G-tdTomato* (Wong et al., 2021), *Canton-S*, *UAS-PercevalHR* (Wong et al., 2021), *iav*¹ (also called *iav*^{hypoB-1}; Wong et al., 2014), and *UAS-iav* (Wong et al., 2014). *Canton-S* flies were used as the wild-type controls in the various crosses.

Analysis of RNAi-mediated gene knock-down

Flies expressing the relevant RNAi transgenes or *UAS-Luc* under the control of heat shock–inducible promoter (hs-GAL4) and hs-GAL4/+ controls were heat shocked on three consecutive days by placing them in a 37°C water bath for 1-h each time. The day after the third heat shock, RNA was extracted from whole-fly extracts using RNeasy mini kit (QIAGEN) by following the manufacturer's instructions. Using the high-capacity cDNA reverse transcription kit (Applied Biosystems), 1 μ g of total RNA was reverse-transcribed. Real-time qPCR was performed using PowerUP SYBR Green Master Mix (Applied Biosystems) by following the manufacturer's instructions. The primers used were as follows:

RpL32 forward: TACAGGCCCAAGATCGTGAA RpL32 reverse: TCTCCTTGCGCTTCTTGGA vapb (Vap33) forward: TGAAGTGCGTTTTCGAGATGC vapb (Vap33) reverse: CTGAGCTAGTATTGGCACCCG

Neuromuscular junction (NMJ) immunohistochemistry and confocal microscopy

Dissection and immunostaining of NMJ was performed as described previously (Wong et al., 2014, 2015). Briefly, wandering third instar larvae were filleted in ice-cold PBS to remove all visceral organs except the brain and nerves. For microtubule staining, larvae were dissected in HL-3 (70 mm NaCl, 5 mm KCl, 1 mm CaCl₂, 20 mm MgCl₂, 10 mm NaHCO₃, 115 mm sucrose, 5 mm trehalose, and 5 mm HEPES; pH 7.2), before treated with HL-3 containing 50 μ M nocodazole or 0.1% DMSO for 30 min at room temperature. The fillets were fixed in 4% paraformaldehyde in PBS for 30 min. The fixed fillets were washed with 0.1% Triton X-100 in PBS before incubation with primary antibodies overnight at 4° C. Primary antibody dilutions were 1:100 mouse anti-discs large (DLG), 1:50 mouse anti-Futsch, and 1:200 mouse anti- α -tubulin. The monoclonal antibodies against DLG (4F3), Futsch (22C10), and α -tubulin (12G10) were obtained from the Developmental Studies Hybridoma Bank (DSHB) developed under the auspices of the NICHD and maintained by The University of Iowa, Department of Biology, Iowa City, IA 52 242. The samples were then washed and probed with 1:200 FITC-conjugated anti-HRP (Jackson ImmunoResearch) and Alexa Fluor 568-conjugated anti-mouse secondary antibodies (ThermoFisher) at room temperature for 1.5 h, and then mounted on glass slides with Vectashield (Vector Labs). Confocal images were obtained using a Nikon A1 Confocal Laser Microscope System (Nikon). For NMJ bouton counting, a $60 \times$ oil objective was used to focus on the NMJs on abdominal segment 3.

To count the number of microtubule loops within the type 1b NMJ boutons in larvae stained with anti- α -tubulin antibody, we applied an HRP mask to remove the α -tubulin signal outside the boutons. Briefly, confocal images were opened in the Fiji image processing package. We first separated the merged Z-stacks into the anti-HRP and anti-tubulin channels. Next, we applied the "Make Binary" function to the anti-HRP channel using default settings, and having checked "Calculate threshold for each image" and "Black background (of binary masks)." We then generated a maximum intensity Z-projection of the binary image, and inverted the color to generate the mask. Using the "Image Calculator"

dialogue window, we subtracted the mask from the anti- α -tubulin stack to remove the anti- α -tubulin signal outside the boutons.

To analyze mitochondria within NMJ boutons, we first separated merged Z-stack images into the mCherry-mito-OMM and anti-HRP signals. We generated an HRP mask exactly as described above to remove any mCherry signal outside the NMJs. To determine the number, volume, and surface area of the mitochondria at the NMJ, we used the "3D Object Counter v2.0" dialogue box with "Threshold" set at 50 and minimum "Size filter" set to 1. We then determined the aspect ratio (volume/ surface area ratio) for each entity identified by the software. Operating under the assumption that the aspect ratio of each mitochondrion should be distinct, akin to a fingerprint, we filtered out all the counted particles whose aspect ratio was duplicated in the dataset for that NMJ. The remaining particles were counted in the analyses.

Larval crawling assay

Larval crawling assay was performed as reported previously (Kashima et al., 2017). Briefly, a 100 mm Petri dish half-filled with 2% agar was placed on an LED drawing pad in a dark room. A black paper strip was wrapped around the circumference of the Petri dish to form a 20-mmtall wall. Late third-instar larvae were collected from fly food-containing vials, briefly rinsed with water, and placed in a 35-mm Petri dish. A paint brush was used to transfer one larva to the center of the agar plate. A digital camera connected to a laptop computer was mounted at ~80 cm above the Petri dish to capture time-lapse images at 2-s intervals (0.5 frame per second). Image recording continued until the larva reached the perimeter of the dish or for a maximum of 5 min. We used each larva in the assay only once. For analysis, we superimposed the time-lapse images of every 10 s (five frames) using ImageJ (NIH). A translucent ruler with millimeter marks on the LED pad was used as calibration scale. Position of each larva was tracked on the superimposed image to measure the total crawling distance in millimeters. Mean velocity was calculated by dividing the distance by the duration of the time. At least six larvae were recorded and analyzed for each genotype.

NMJ electrophysiology

Wandering third instar larvae were dissected in ice-cold HL-3 (70 mm NaCl, 5 mm KCl, 20 mm MgCl₂.6H₂O, 10 mm NaHCO₃, 115 mm sucrose, 5 mm trehalose, and 5 mm HEPES; pH 7.2) and rinsed with HL-3 containing 0.5 mm Ca²⁺. Recordings were made from body-wall muscle 6 (abdominal segment 3) with sharp electrodes filled with a 2:1 mixture of 2 M potassium acetate and 2 M potassium chloride. Data were collected from samples with resting membrane potential below -60 mV. Excitatory postsynaptic junctional potentials (EJPs) were evoked by directly stimulating the A3 hemisegmental nerve through a glass capillary electrode (internal diameter, $\sim 10 \,\mu\text{m}$) at 0.2 Hz. Stimulus pulses were generated by pClamp 10 software (Molecular Devices Inc). The applied currents were 6 \pm 3 μ A with fixed stimulus duration at 0.3 ms. A total of 20-30 evoked EJPs were recorded for each muscle for analysis. Miniature EJP (mEJP) events were collected for 5 min. Both EJPs and mEJPs were amplified with an Axoclamp 900A amplifier (Molecular Devices) under bridge mode, filtered at 10 kHz and digitized at 10 kHz (EJPs) and 40 kHz (mEJPs) with pClamp 10. Experiments were performed at 21°C. EJPs and paired-pulse stimulation were analyzed with pClamp 8.0 software (Molecular Devices). The mEJPs were analyzed using the Mini Analysis Program (Synaptosoft Inc.). The EJPs paired-pulse amplitudes were corrected by nonlinear summation (Feeney et al., 1998). Paired-pulse ratio was calculated as the ratio of second to first peak. The quantal content of evoked release was calculated from individual muscles by the ratio of the average EJP amplitude over the average mEJP amplitude. For high-frequency stimulation, nerves were stimulated for 10 min at 10 Hz. By fitting the decay portions of each highfrequency trace to first order exponential function, we obtained the rate constant of decays. Rundowns were extracted from the rate-constants of decay.

Dissociation of Drosophila neurons

We dissociated primary motor neurons from *Drosophila* as described previously (Wong et al., 2021). Briefly, the exterior of wandering third

instar larvae was sterilized by brief submersion in ethanol, and then washed with sterilized H₂O before dissection in filtered Schneider's medium (S0146; Sigma-Aldrich) containing 10% fetal bovine serum (FBS), antibiotic/antimycotic solution (A5955; Sigma-Aldrich) and 50 μg/ml of insulin (I6634; Sigma-Aldrich). Brains dissected from these larvae were washed in separate wells containing filtered Schneider's medium before being transferred to a filtered HL-3 solution (70 mm NaCl, 5 mm KCl, 1 mм CaCl₂, 20 mм MgCl₂, 10 mм NaHCO₃, 115 mм sucrose, 5 mм trehalose, and 5 mm HEPES) supplemented with 0.423 mm L-cysteine (Calbiochem) and 5 U/ml papain (Worthington; note, after L-cysteine addition but before papain addition, the pH of the solution was recalibrated to 7.4). The brains were then enzymatically digested in the papain solution for 20 min before transfer to a 1.5-ml tube containing 1 ml of filtered Schneider's medium. Cells were centrifuged at 100 G for 1 min before decantation of Schneider's medium. The solution was replaced with 1 ml of fresh filtered Schneider's medium. This process was repeated twice before neurons were dissociated by pipetting repeatedly until the solution was homogeneous. The solution with dissociated neurons was then placed on 35-mm glass bottom dishes (D35-10-0-N; Cellvis) that had been coated with concanavalin A (C2010; Sigma-Aldrich). Cells were cultured in Schneider's medium supplemented with 10% FBS, antibiotic/antimycotic solution (A5955; Sigma-Aldrich) and 50 μg/ml of insulin (I6634; Sigma-Aldrich) at room temperature in a humidified container at room temperature. After each day in culture, cells were washed with PBS to remove any yeast contamination or debris remaining from dissociation.

Live-cell imaging of fly primary neurons

Live imaging of dissociated neurons was performed as described previously (Wong et al., 2021). Briefly, culture media on plates of dissociated neurons was first replaced with HL-3 (70 mm NaCl, 5 mm KCl, 1 mm CaCl₂, 20 mm MgCl₂, 10 mm NaHCO₃, 115 mm sucrose, 5 mm trehalose, and 5 mm HEPES; pH 7.2, room temperature). For measurements of cytosolic Ca²⁺, GCaMP5G and tdTomato were sequentially excited at 488 and 561 nm, respectively, by an A1 laser confocal microscope with a 40× objective (Nikon). Emission signals at 525 and 595 nm were recorded. Backgrounds were measured from a cell-free region of interest (ROI). Baselines were established for 1 min before addition of muscarine (1 mm). Cytosolic Ca²⁺ transients were evoked by store-depletion with the SERCA inhibitor, thapsigargin (TG), as described. Amplitudes of the GCaMP5G/tdTomato ratio represented cytosolic free [Ca²⁺]. More than 50 cells from a minimum of three independently conducted experiments for each condition were used for the calculations.

PercevalHR signals were recorded by measuring the ratio of fluorescence emissions at 525 nm sequentially excited at 487.5 and 407.8 nm. An A1R laser confocal microscope with 40× objective (Nikon) was used for measurement. Background emission signals were measured from a cell-free ROI. Baselines were established for 2 min before the bath was replaced with high [K $^+$] (52 mm) HL-3. Oligomycin A (oligoA; 10 μ m) were added as needed and signals were recorded. Amplitudes of the emission ratio represented the cytosolic [ATP]/[ADP] ratio (Tantama et al., 2013; Wong et al., 2021). Data were quantified as change in the PercevalHR ratio over the baseline that was set to the value at the time point immediately before the addition of the drug. We determined the cumulative change in the ratio using a custom R code and the area under the curve (AUC) per unit time. Cells from a minimum of three to four independently conducted experiments for each condition were used for the calculations.

Analysis of fly lifespan

Newly eclosed adult flies of both sexes were collected and transferred to vials containing standard fly food (\leq 15 flies per vial). Flies were kept at room temperature (\sim 21°C) and transferred to new vials twice per week. Dead flies at the bottom of the old vials were counted after each transfer until all the animals in a cohort died.

Experimental design and statistical analyses

Excel (Microsoft), and Prism 8 (GraphPad) were used for statistical analyses and curve-fitting. We used either parametric or nonparametric tests

of statistical significance on the basis of whether the data were normally distributed. We assessed the normality of data using the Shapiro-Wilk test. Descriptions of the data distribution are provided in the tables. Multiple comparisons were made by ANOVA or Kruskal–Wallis tests. Custom R code used for quantifying [Ca²+] and PercevalHR data. Statistical significance was defined as a p < 0.05. P-values were shown on the figures as asterisks: P0.005, P0.01, P0.001, P0.001. Lifespan (Kaplan–Meier) curves were generated using Prism 8. We used the log-rank (Mantel–Cox) test to determine P-values. Other test-specific values are provided in the tables.

Results

Expression of an ALS-related variant of *vapb* results in defective presynaptic bouton development

Either the deletion of *Drosophila vapb* (also called *Vap33*) or the expression of a transgene equivalent to an ALS-causing variant (vapb^{P58S}) disrupts presynaptic microtubules at the larval NMJ resulting in the appearance of fewer, but morphologically larger, boutons (Pennetta et al., 2002; Chai et al., 2008; Ratnaparkhi et al., 2008). We confirmed that in comparison to the GAL4 and *UAS* controls, ectopic expression of *vapb*^{P58S}, but not wild-type vapb (vapb WT), in motor neurons (using VGlut ok371-GAL4, herein referred to as ok371-GAL4) led to a significant reduction in bouton number (Fig. 1A-E; Brand and Perrimon, 1993; Mahr and Aberle, 2006; Tsuda et al., 2008). NMJ boutons in neurons expressing *vapb*^{P58S} also exhibited significant increases in bouton area (Fig. 1A,B,F). Although animals expressing vapbWT in motor neurons exhibited a slight, yet significant, increase in bouton area (Fig. 1D,F) relative to the UAS controls, bouton numbers in these animals were not significantly altered (Fig. 1E). Animals expressing $vapb^{P58S}$, however, exhibited significantly fewer and larger boutons in relation to the GAL4/ *UAS* controls and those expressing $vapb^{WT}$ (Fig. 1*E,F*). These data agree with prior findings regarding the effects of VAPBP58S on Drosophila larval NMJ synapse morphology (Chai et al., 2008; Ratnaparkhi et al., 2008).

${ m VAPB}^{ m P58S}$ -induced defects in presynaptic bouton development are ameliorated by lowering the expression of genes encoding ER ${ m Ca}^{2+}$ release channels

We previously showed that decreased abundance of ER Ca²⁺ release channels, Inactive (Iav), ryanodine receptor (RyR), and inositol trisphosphate receptor (IP₃R), result in NMJs that exhibit fewer boutons that are larger in size (Fig. 2A; Wong et al., 2014). The ostensible similarities between the morphology of NMJ boutons in iav^{I} mutants (also called $iav^{hypoB-I}$, strong hypomorphs of iav; Homyk and Sheppard, 1977; Wong et al., 2014) and animals expressing *vapb*^{P58S} implies a common underlying mechanism. Therefore, we speculated that the coincidence of iav^{I} and $vapb^{P58S}$ would enhance the respective phenotypes leading to an even greater reduction in bouton number (Fig. 2A). To our surprise, expression of vapb^{P58S} in the iav¹ background restored, rather than worsened, both bouton number and morphology to control levels (Fig. 2B,C,J, K). Whereas bouton numbers and area elicited by $vapb^{\overline{P}58S}$ were not significantly different from those in *iav*¹, these parameters were restored to wild-type levels on the coincidence of vapb^{P58S} and iav¹ (Fig. 2J,K). In contrast, overexpression of vapbWT did nothing to alter the paucity of boutons in iav¹ (Fig. 2D,E,L). Therefore, decreased abundance of Iav rescued the synaptic growth phenotype stemming from the expression of $vapb^{P585}$.

Next, we asked whether the rescue of *vapb*^{P58S}-induced NMJ phenotypes was specific to the loss of Iav, or whether similar suppression would occur in response to lowered abundance of other

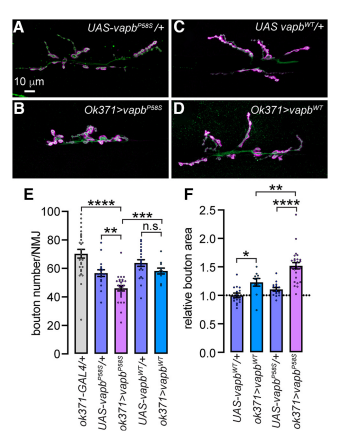


Figure 1. Expression of $vapb^{PS85}$ in Drosophila motor neurons led to significant changes in presynaptic bouton development at the larval NMJ. A-D, Representative confocal images of larval NMJs dissected from animals of the indicated genotypes stained with antibodies against HRP (green) and DLG (magenta). Scale bar shown in A on the top left applies to all panels. Please note that in all figures, UAS-transgene/+ refers to the presence of the noted UAS-transgene construct without a GAL4 driver, and driver-transgene refers to the UAS-transgene, whose expression was driven using the driver-GAL4. E, F, Bar graphs showing quantification of the larval NMJ bouton numbers (F) and relative bouton area (F) in animals of the indicated genotypes. Values represent mean \pm SEM; *p < 0.05, **p < 0.01, ***p < 0.001, ***p < 0.001; n.s., not significant, t tests with Bonferroni correction. Dots represent values from distinct NMJs. Refer to Table 1 for statistical information.

ER Ca²⁺ channels. Simultaneous knock-down of the gene encoding the fly IP₃R (itpr) using the d42-GAL4 motor neuron driver and a validated RNAi line (*UAS-itpr*^{RNAi}) ameliorated the effect of vapb^{P58S} expression on bouton number (Fig. 2G,M; Venkatesh and Hasan, 1997; Parkes et al., 1998; Wong et al., 2014, 2021). The d42-GAL4/+ and UAS-vapb^{P58S}/+ control animals did not exhibit a decrease in bouton numbers (Fig. 2M). Absence of comparable suppression in animals coexpressing vapb^{P58S} and a neutral UAS transgene (UAS-Dcr1; Fig. 2F,M) demonstrates that the suppression brought about by coexpression of *itpr*^{RNAi} was not because of GAL4 dilution stemming from the presence of a second UAS transgene. Concomitant absence of one copy of RyR ($RyR^{16}/+$), which mimics the iav1 NMJ growth phenotype, also prevented vapb^{P58S}-induced alterations in bouton numbers (Fig. 2H,I,N; Sullivan et al., 2000; Wong et al., 2014). Taken together, these data demonstrate that VAPB^{P58S}-induced defects in presynaptic bouton development are ameliorated by cell autonomous reduction in the expression of genes encoding ER Ca²⁺ channels.

CaMKII overactivation underlies alterations in presynaptic development in motor neurons expressing $vapb^{P58S}$

The aforementioned findings are consistent with the notion of VAPB^{P58S} inducing an increase in cytosolic [Ca²⁺] such that the

Table 1. Statistical information for the data shown in Figure 1

Figure 1 <i>E</i>			
Genotype	Normally distributed?	Values	Number of animals
ok371-GAL4/+ UAS-vapb ^{P585} /+ ok371>vapb ^{P585} UAS-vapb ^{WT} /+ ok371>vapb ^{WT}	Yes Yes Yes Yes Yes	70.3 ± 3.1 56.7 ± 2.4 46.1 ± 1.9 63.8 ± 2.3 58.2 ± 2.0	14 (n = 28 NMJs) 9 (n = 17 NMJs) 13 (n = 26 NMJs) 11 (n = 21 NMJs) 6 (n = 11 NMJs)
Comparison between	Test used	<i>P</i> -value	Test statistics
ok371-GAL4/+ ok371>vapb ^{PS85} UAS-vapb ^{PS85} /+ ok371>vapb ^{PS85} ok371>vapb ^{PS85} ok371>vapb ^{WT} UAS-vapb ^{WT} /+ ok371>vapb ^{WT} Figure 1F	t test followed by Bonferroni correction	0.0039 0.0024 0.0006 0.24	t = 3.519, df = 33.74 t = 3.767, df = 28.97 t = 4.379, df = 26.47 t = 1.807, df = 28.68
Genotype	Normally distributed?	Values	Number of animals
UAS-vapb ^{WT} /+ ok371>vapb ^{WT} UAS-vapb ^{P585} /+ ok371>vapb ^{P585}	Yes Yes Yes No	$\begin{array}{c} 1 \pm 0.04 \\ 1.23 \pm 0.07 \\ 1.11 \pm 0.03 \\ 1.52 \pm 0.06 \end{array}$	8 (n = 15 NMJs) 5 (n = 10 NMJs) 8 (n = 16 NMJs) 13 (n = 26 NMJs)
Comparison between	Test used	<i>P</i> -value	Test statistics
$UAS-vapb^{WT}/+ ok371>vapb^{WT}$ $ok371>vapb^{WT} ok371>vapb^{P585}$ $UAS-vapb^{P585}/+ ok371>vapb^{P585}$	t test followed by Bonferroni correction Mann–Whitney followed by Bonferroni correction Mann–Whitney followed by Bonferroni correction	0.02 0.0096 <0.0001	t = 2.901, df = 15.24 U = 27 U = 27

attenuation of ER Ca²⁺ release restores homeostasis, and thus, mitigates the bouton development phenotypes. Our data also imply that either an increase or decrease in presynaptic [Ca²⁺] results in morphologically identical bouton phenotypes. Indeed, we previously showed that either the absence or overexpression of *iav* results in comparable reduction in the number of NMJ boutons (Wong et al., 2014). How might this bell-shaped relationship between presynaptic [Ca²⁺] and NMJ bouton development come about?

We posit that the key to understanding these outcomes is the stability of presynaptic microtubules. We and others have shown that destabilization of presynaptic microtubules, which manifests as loss of characteristic intrabouton microtubule loops, results in an increase bouton size, and a concomitant decrease in bouton number (Roos et al., 2000; Pennetta et al., 2002; Viquez et al., 2006; Wong et al., 2014). Indeed, the number of presynaptic microtubule loops at NMJs of animals expressing vapb P58\$ in motor neurons was significantly lower than that in animals expressing vapb^{WT} (Fig. 3A,B). Treatment of NMJs with the microtubule depolymerizing agent, nocodazole, led to a reduction in the number of microtubule loops at the NMJs of animals expressing $vapb^{WT}$, whereas we observed no further decrease in the number of microtubule loops in nocodazole-treated NMJs in animals expressing $vapb^{P58S}$ (Fig. 3A,B). The number of microtubule loops in nocodazole-treated *vapb*^{WT} NMJs was statistically indistinguishable from those in nocodazole-treated or -untreated vapb^{P58§} NMJs (Fig. 3B). These data are consistent with microtubule depolymerization underlying the fewer microtubule loops within the *vapb*^{P58S} NMJs.

We showed previously that reduction in presynaptic resting [Ca²⁺] and the attendant attenuation of the Ca²⁺/calmodulin (CaM) responsive phosphatase, calcineurin (CanA1), destabilize presynaptic microtubules (Wong et al., 2014). In the event of a Ca²⁺-responsive kinase having the same target as CanA1, increased activity of that kinase could also result in hyperphosphorylation of

the target. If so, one might expect that either an increase or decrease in cytosolic [Ca²⁺] would result in higher fractions of that target being phosphorylated. Since activation of the Ca²⁺/CaM-dependent protein kinase II (CaMKII or CKII) by high [Ca²⁺] destabilizes microtubules by phosphorylating microtubule-associated proteins (MAPs; Baratier et al., 2006; McVicker et al., 2015; Oka et al., 2017), we asked whether elevated ER Ca²⁺ release compels the observed changes in NMJ development because of unrestrained CaMKII activity. Mirroring the decrease in the number of presynaptic microtubule loops, NMJs in animals expressing vapb^{P58S} also exhibited fewer loops of the fly MAP1b homolog, Futsch (Hummel et al., 2000; Roos et al., 2000; Fig. 3C,D). Coexpression of the CaMKII inhibitory peptide, CaMKII^{ala} (CKII^{ala}; Joiner and Griffith, 1997), with *vapb*^{P58S} led to recovery of the numbers of Futsch loops (Fig. 3C,D) and boutons (Fig. 3E). In iav¹ synapses, which exhibit lower resting [Ca²⁺] (Wong et al., 2014), expression of CKII^{ala} did not restore the bouton number (Fig. 3E). Suggesting a specific rescue of the $vapb^{P58S}$ phenotype rather than a general increase in bouton number, overexpression of CKII^{ala} without vapb^{P58S} was not sufficient to elevate the number of presynaptic boutons (Fig. 3E). Indicating the sufficiency of CaMKII downstream of elevated [Ca²⁺], in iav-overexpressing neurons, which exhibit higher presynaptic [Ca²⁺] (Wong et al., 2014), bouton number was restored to wild-type level on coexpression of CKII^{ala} (Fig. 3F). Taken together, these data suggest a biphasic dependency of presynaptic microtubule stability and bouton number on cytosolic [Ca²⁺], and that vapb^{P58S}-induced alterations in presynaptic development stem from aberrant CaMKII activation (Fig. 3*G*).

Next, we asked whether $vapb^{P58S}$ -induced phenotypes other than NMJ development also stem from elevated presynaptic [Ca²⁺] and CaMKII activation. Since $vapb^{P58S}$ is equivalent to the ALS-causing variant of human VAPB (Nishimura et al., 2004, 2005; Kanekura et al., 2006; Marques et al., 2006; Landers et al., 2008), we examined locomotor activity in larvae expressing

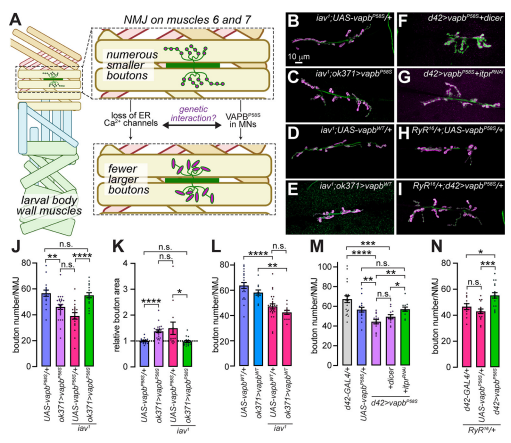


Figure 2. VAPB^{P-S8S}-induced defects in presynaptic bouton development are ameliorated on concomitant loss of ER Ca²⁺ release channels. **A**, Model showing that the development of control NMJs on muscles 6 and 7 of the larval body wall muscles leads to the appearance of numerous small boutons. Animals either expressing $vapb^{PS8S}$ in motor neurons or harboring mutations that diminish ER Ca²⁺ release are characterized by the appearance of fewer, but morphologically larger, boutons. Ostensible similarities in bouton development phenotypes raises the question of genetic interactions between these conditions. Image was created with BioRender. **B–I**, Representative confocal images of larval NMJs dissected from animals of the indicated genotypes stained with antibodies against HRP (green) and DLG (magenta). Scale bar shown in **B** applies to all panels. **J–N**, Bar graphs showing quantification of the larval NMJ bouton numbers (**J**, **I–N**) and relative bouton area (**K**) in animals of the indicated genotypes. Values represent mean \pm SEM; *p < 0.05, **p < 0.01, ****p < 0.001, ****p < 0.0001; n.s., not significant, paired t tests with Bonferroni correction. Dots represent values from distinct NMJs. Quantification of bouton numbers in Figure 1 and this figure were from the same experiment. Refer to Table 2 for statistical information.

this transgene in motor neurons. To this end, we placed wandering third instar larvae in an open field and recorded their movement toward the periphery of the field (Fig. 4A). In comparison to larvae expressing $vapb^{WT}$, those expressing $vapb^{P58S}$ exhibited a significant reduction in velocity (Fig. 4A,B). Upon coexpression of either $itpr^{RNAi}$ or $CKII^{ala}$, the velocities of $vapb^{P58S}$ -expressing larvae were no longer significantly different from the animals expressing $vapb^{WT}$ with $itpr^{RNAi}$ or $CKII^{ala}$, respectively (Fig. 4B). Expression of $vapb^{P58S}$ in motor neurons shortens Drosophila

Expression of $vapb^{P58S}$ in motor neurons shortens Drosophila lifespan via overactivation of $PLC\beta$ – IP_3R signaling, and either the deletion of the gene encoding a fly $PLC\beta$ (norpA; Bloomquist et al., 1988) or the concomitant knock-down of itpr strongly suppresses the effects of $VAPB^{P58S}$ on animal longevity (Wong et al., 2021). In contrast to the ameliorative effects on the NMJ bouton phenotype, neither iav^I nor the $RyR^{I6}/+$ mutations influenced the lifespan of animals expressing $vapb^{P58S}$ (Wong et al., 2021). Therefore, the effects of $vapb^{P58S}$ on NMJ bouton development and adult lifespan occur via distinct mechanisms. In agreement, coexpression of $CKII^{ala}$, which mitigated the NMJ growth and larval velocity phenotypes in animals expressing $vapb^{P58S}$, did not influence the effect of mutant VAPB on adult lifespan (Fig. 4C). Collectively, these data show that although larval phenotypes in $vapb^{P58S}$ -expressing animals involves CaMKII, the same is not the case for adult longevity.

Expression of $vapb^{P58S}$ results in diminished extrusion of cytosolic Ca^{2+}

Lower presynaptic resting [Ca²⁺] in the absence of Iav results in reduced probability of synaptic vesicle (SV) release (Wong et al., 2014). NMJs in *iav*¹ animals, therefore, exhibit reduced amplitudes of evoked EJPs and paired-pulse facilitation in response to a second stimulus applied after a delay of 50 ms (Wong et al., 2014). Conversely, overexpression of iav led to an increase in presynaptic resting [Ca²⁺], higher EJP amplitudes, and pairedpulse depression indicating elevated SV release probability (Wong et al., 2014). These data point to a dose-dependent effect of ER Ca²⁺ release on presynaptic resting [Ca²⁺] and SV release probability. Given that the NMJ bouton phenotypes in animals overexpressing either *vapb*^{P58S} or *iav* were suppressed by CKII^{ala}, we asked whether VAPB^{P58S} also elevates SV release probability because of an increase in resting [Ca²⁺]. If so, expression of vapb^{P58S} would result in increased EJP amplitude and pairedpulse depression. However, EJP amplitudes in *vapb*^{P58S} neurons were not significantly different from those in $\hat{vap}b^{WT}$ neurons (Fig. 5A,B), which is in agreement with the report that ectopic expression of the ALS-causing human transgene (hVAPP56S) had no effect on the amplitude of evoked potentials at the Drosophila larval NMJ (Chai et al., 2008). Furthermore, relative to neurons expressing vapb^{WT}, those expressing vapb^{P58S} showed paired-

Table 2. Statistical information for the data shown in Figure 2

Genotype	Normally distributed?	Values	Number of animals
UAS-vapb ^{P58S} /+	Yes	56.7 ± 2.4	9 (<i>n</i> = 17 NMJs)
ok371>vapb ^{P585}	Yes	46.1 ± 1.9	13 ($n = 26 \text{ NMJs}$)
iav ¹ ;UAS-vapb ^{P585} /+	Yes	39.2 ± 2.6	9 ($n = 18 \text{ NMJs}$)
iav ¹ ;ok371>vapb ^{P585}	Yes	55.1 ± 2	9 ($n = 17 \text{ NMJs}$)
Comparison between	Test used	<i>P</i> -value	Test statistics
$UAS-vapb^{P58S}/+ ok371>vapb^{P58S}$	t test followed by Bonferroni correction	0.0024	t = 3.767, df = 28.97
UAS - $vapb^{P58S}$ /+ iav^{1} ; $ok371$ > $vapb^{P58S}$	t test followed by Bonferroni correction	1	t = 0.5174, df = 30.85
ok371>vapb ^{P585} iav ¹ ;UAS-vapb ^{P585} /+	t test followed by Bonferroni correction	0.12	t = 2.183, df = 33.57
iav ¹ ;UAS-vapb ^{P58S} /+ iav ¹ ;ok371>vapb ^{P58S}	t test followed by Bonferroni correction	< 0.0001	t = 4.966, df = 31.30
Figure 2K			
Genotype	Normally distributed?	Values	Number of animals
UAS-vapb ^{P585} /+	Yes	1 ± 0.03	8 (n = 15 NMJs)
ok371>vapb ^{P585}	No	1.37 ± 0.05	5 ($n = 10 \text{ NMJs}$)
iav ¹ ;UAS-vapb ^{P58S} /+	No	1.49 ± 0.24	7 ($n = 13 \text{ NMJs}$)
iav ¹ ;ok371>vapb ^{P585}	No	0.98 ± 0.04	9 (<i>n</i> = 17 NMJs)
Comparison between	Test used	<i>P</i> -value	Test statistics
UAS - $vapb^{P585}$ /+ $ok371$ > $vapb^{P585}$	Mann-Whitney followed by Bonferroni correcti		<i>U</i> = 27
UAS - $vapb^{P585}$ /+ iav^1 ; $ok371$ > $vapb^{P585}$	Mann–Whitney followed by Bonferroni correcti		U = 99
ok371>vapb ^{P585} iav ¹ ;UAS-vapb ^{P585} /+	Mann-Whitney followed by Bonferroni correcti		<i>U</i> = 124
iav^1 ;UAS-vapb ^{P585} /+ iav^1 ;ok371>vapb ^{P585}	Mann—Whitney followed by Bonferroni correcti	ion 0.018	U = 46
Figure 2L			
Genotype	Normally distributed?	Values	Number of animals
UAS-vapb ^{WT} /+	Yes	63.8 ± 2.3	11 ($n = 21 \text{ NMJs}$)
ok371>vapb ^{WT}	Yes	58.2 ± 2.0	6 ($n = 11 \text{ NMJs}$)
iav ¹ ;UAS-vapb ^{WT} /+	Yes	42.5 ± 1.6	6 ($n = 12 \text{ NMJs}$)
iav ¹ ;ok371>vapb ^{WT} Comparison between	Yes Test used	47.2 \pm 2.0 <i>P</i> -value	13 (n = 25 NMJs) Test statistics
UAS-vapb ^{WT} /+ iav ¹ ;UAS-vapb ^{WT} /+ ok371>vapb ^{WT} iav ¹ ;ok371>vapb ^{WT}	t test followed by Bonferroni correction t test followed by Bonferroni correction	<0.0001 0.0018	t = 7.535, df = 30.86 t = 3.853, df = 27.60
ok371>vapb lav ,0k371>vapb ok371>vapb lav ,0k371>vapb lav ,0k3	t test followed by Bonferroni correction	0.0018	t = 3.860, df = 33.96
Figure 2M	t test followed by bomerfolli correction	0.21	t = 1.000, ut = 33.90
Genotype	Normally distributed?	Values	Number of animals
d42-GAL4/+	Yes	67.3 ± 3.5	10 (n = 20 NMJs)
UAS-vapb ^{P585} /+	Yes	56.7 ± 2.4	8 (n = 15 NMJs)
d42>vapb ^{P585}	Yes	44.5 ± 2.2	7 (n = 14 NMJs)
d42>vapb ^{P58S} +dicer	Yes	49.5 ± 2.1	6 ($n = 12 \text{ NMJs}$)
d42>vapb ^{P58S} +itpr ^{RNAi}	Yes	57.2 ± 2.1	6 (<i>n</i> = 12 NMJs)
Comparison between	Test used	<i>P</i> -value	Test statistics
$d42$ -GAL4/+ $d42$ >vapb P585	t test followed by Bonferroni correction	< 0.0001	t = 5.507, df = 30.15
d42-GAL4/+ d42>vapb ^{P58S} +dicer	t test followed by Bonferroni correction	0.0004	t = 4.368, df = 28.63
$UAS-vapb^{P585}/+ d42>vapb^{P585}$	t test followed by Bonferroni correction	0.0032	t = 3.767, df = 28.97
$UAS-vapb^{P585} + d42 > vapb^{P585} + itpr^{RNAi}$	t test followed by Bonferroni correction	1	t = 0.1462, df = 26.92
d42>vapb ^{P585} d42>vapb ^{P585} + dicer d42>vapb ^{P585} d42>vapb ^{P585} + itpr ^{RNAi}	t test followed by Bonferroni correction	0.44	t = 1.647, df = 23.98 t = 1.860, df = 33.96
d42>vapb +1tpr d42>vapb ^{P585} +dicer d42>vapb ^{P585} +itpr ^{RNAi}	t test followed by Bonferroni correction t test followed by Bonferroni correction	0.0012 0.04	t = 1.800, df = 33.90 t = 2.606, df = 22.00
Figure 2N	t test followed by boilleffolli correction	0.04	t — 2.000, ut — 22.00
Genotype	Normally distributed?	Values	Number of animals
RyR ¹⁶ /+;UAS-vapb ^{P585} /+	·		
RyR ¹⁶ /+;d42-GAL4/+	Yes Yes	43.1 ± 1.8 46.6 ± 2.3	8 (n = 16 NMJs) 6 (n = 11 NMJs)
$RyR^{16}/+;d42>vapb^{P585}$	Yes	$\frac{40.0 \pm 2.3}{55.3 \pm 2.1}$	9 ($n = 11 \text{ NMJs}$)
Comparison between	Test used	<i>P</i> -value	Test statistics
$RyR^{16}/+;UAS-vapb^{P58S}/+RyR^{16}/+;d42-GAL4/+$	t test followed by Bonferroni correction	0.67	t = 1.246, df = 20.68
$RyR^{16}/+;d42-GAL4/+RyR^{16}/+;d42>vapb^{P585}$	t test followed by Bonferroni correction	0.0003	t = 4.430, df = 32
$RyR^{16}/+;UAS-vapb^{P58S}/+RyR^{16}/+;d42>vapb^{P58S}$	t test followed by Bonferroni correction	0.027	t = 2.813, df = 23.92

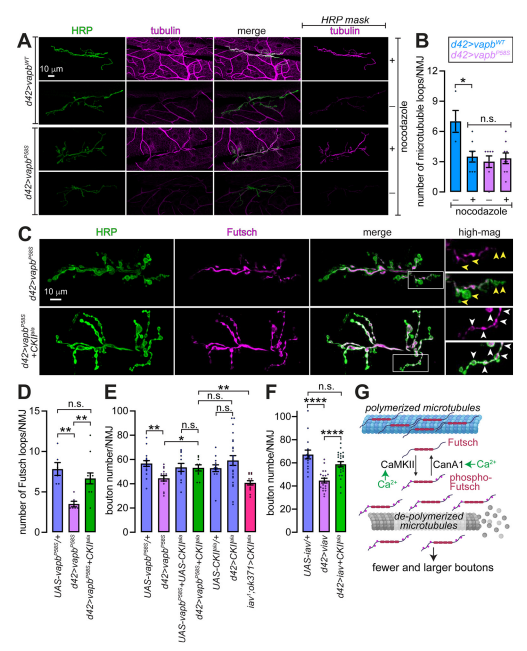


Figure 3. CaMKII underlies the alterations in presynaptic bouton development in motor neurons expressing $vapb^{PS85}$. A, Representative confocal images of larval NMJs dissected from animals of the genotypes indicated on the left, stained with antibodies against HRP (green) and tubulin (magenta) as indicated on the top. Images derived by application of an HRP mask to the antitubulin images are shown. Whether or not the larvae were treated with nocodazole is indicated with + or - on the right. Scale bar shown in top left image applies to all the images in the panel. B, Bar graph showing quantification of the number of microtubule loops within each NMJ. Values represent mean \pm SEM; *p < 0.05; t tests; and n.s., not significant, Kruskal–Wallis. Dots represent values from distinct NMJs. C, Representative confocal images of NMJs expressing $vapb^{PS85}$ either with or without $CKII^{pla}$ as indicated. Samples were stained with antibodies against HRP (green) and Futsch (magenta). White arrowheads point to individual boutons and highlight recovery of Futsch loops in neurons expressing both $vapb^{PS85}$ and $CKII^{pla}$. Higher magnification images (high-mag) represent the regions shown in the box overlaid on the merged images. Scale bar shown on the top left applies to all panels. D-F, Bar graphs showing quantification of the number of Futsch loops per NMJ (D) and larval NMJ bouton numbers (E, F) in animals of the indicated genotypes. Values represent mean \pm SEM; *p < 0.05, **p < 0.05, **

pulse facilitation, rather than paired-pulse depression (Fig. 5A,C). Amplitudes of mini-EJPs (mEJPs) and quantal content were also virtually identical in $vapb^{WT}$ -expressing and $vapb^{P58S}$ -expressing NMJs (Fig. 5D,E). Taken together, these data argue against elevations in presynaptic resting

 $[Ca^{2+}]$ and SV release probability in $vapb^{P58S}$ -expressing motor neurons.

What then could explain the hallmarks of elevated [Ca²⁺] in *vapb*^{P58S}-expressing motor neurons? Elevations in cytosolic [Ca²⁺] could stem either from greater Ca²⁺ influx or from

Table 3. Statistical information for the data shown in Figure 3

Figure 3B			
Genotype	Normally distributed?	Values	Number of animals
$d42>vapb^{WT}$ $d42>vapb^{WT} + \text{nocodazole}$ $d42>vapb^{P585}$ $d42>vapb^{P585} + \text{nocodazole}$	Yes Yes No Yes	7.0 ± 1.1 3.5 ± 0.5 3.0 ± 0.6 3.3 ± 0.5	2 (n = 4 NMJs) 4 (n = 8 NMJs) 4 (n = 7 NMJs) 5 (n = 10 NMJs)
Comparison between	Test	t used <i>P</i> -value	Test statistics
$\frac{d42>vapb^{WT}}{d42>vapb^{WT}} + \text{nocodazole} \\ d42>vapb^{WT} + \text{nocodazole} \\ d42>vapb^{PS85} \\ d42>vapb^{PS85} $	t te b^{P585} + nocodazole Krus	st 0.03 skal–Wallis 0.9379	t = 2.904, df = 4.533 Kruskal-Wallis statistic = 0.1282
Figure 3D			
Genotype	Normally distributed?	Values	Number of animals
UAS-vapb ^{PS8S} /+ d42>vapb ^{PS8S} d42>vapb ^{PS8S} +CKIf ^{ala}	Yes No Yes	7.8 ± 0.8 3.5 ± 0.3 6.7 ± 0.7	3 (n = 6 NMJs) 3 (n = 6 NMJs) 6 (n = 6 NMJs)
Comparison between	Test used	P-valu	e Test statistics
UAS-vapb ^{P585} /+ d42>vapb ^{P585} d42>vapb ^{P585} d42>vapb ^{P585} +CKII ^{ala} UAS-vapb ^{P585} /+ d42>vapb ^{P585} +CKII ^{ala}	Mann–Whitney followed by Bonfe Mann–Whitney followed by Bonfe t test followed by Bonferroni corre	erroni correction 0.007	U = 0 U = 5 t = 1.119, df = 12.06
Figure 3 <i>E</i>			
Genotype	Normally distributed?	Values	Number of animals
$\begin{array}{l} \textit{UAS-vapb}^{\textit{PS8S}}/+\\ \textit{d42}>\textit{vapb}^{\textit{PS8S}}\\ \textit{UAS-vapb}^{\textit{PS8S}}+\textit{UAS-CKII}^{\textit{ala}}\\ \textit{d42}>\textit{vapb}^{\textit{PS8S}}+\textit{CKII}^{\textit{ala}}\\ \textit{d42}>\textit{vapb}^{\textit{PS8S}}+\textit{CKII}^{\textit{ala}}\\ \textit{UAS-CKII}^{\textit{ala}}/+\\ \textit{d42}>\textit{CKII}^{\textit{ala}}\\ \textit{iav}^{1};\textit{ok371}>\textit{CKII}^{\textit{ala}} \end{array}$	Yes Yes Yes Yes No Yes	56.7 ± 2.4 44.5 ± 2.2 53.5 ± 3.3 53.1 ± 2.5 52.9 ± 2.8 59.2 ± 4.0 40.8 ± 1.8	8 (n = 15 NMJs) 7 (n = 14 NMJs) 6 (n = 12 NMJs) 5 (n = 10 NMJs) 7 (n = 14 NMJs) 12 (n = 24 NMJs) 7 (n = 14 NMJs)
Comparison between	Test used	<i>P</i> -value	Test statistics
UAS-vapb ^{P585} /+ d42>vapb ^{P585} d42>vapb ^{P585} d42>vapb ^{P585} + CKII ^{ala} UAS-vapb ^{P585} + UAS-CKII ^{ala} d42>vapb ^{P585} + CKII ^{ala} d42>vapb ^{P585} + CKII ^{ala} d42>CKII ^{ala} d42>vapb ^{P585} + CKII ^{ala} iav ¹ ; ok371 > CKII ^{ala}	 t test followed by Bonfer 	roni correction 0.03 roni correction 1 roni correction 0.6	t = 3.767, df = 28.97 t = 2.555, df = 19.88 t = 0.09618, df = 19.53 t = 1.272, df = 32.00 t = 3.977, df = 17.07
Figure 3F			
Genotype	Normally distributed?	Values	Test statistics
UAS-iav/+ d42>iav d42>iav+CKII ^{ala}	Yes Yes Yes	67.3 ± 3.8 44.7 ± 2.1 58.8 ± 2.2	8 (n = 15 NMJs) 10 (n = 19 NMJs) 12 (n = 23 NMJs)
Comparison between	Test used	<i>P</i> -value	Test statistics
UAS-iav/+ $d42$ >iav $d42$ >iav $d42$ >iav+ $CKII^{ala}$ UAS -iav/+ $d42$ >iav+ $CKII^{ala}$	 t test followed by Bonferroni correction t test followed by Bonferroni correction t test followed by Bonferroni correction 	< 0.0001	t = 5.214, df = 21.89 t = 4.611, df = 40.00 t = 1.907, df = 23.78

diminished Ca²⁺ extrusion from the cytosol. Since the former is ruled out by the aforementioned data, we asked whether the effects of VAPB^{P58S} stem from diminished Ca²⁺ extrusion (Fig. 5F). We reasoned that diminished Ca²⁺ extrusion would promote the accumulation of residual Ca²⁺, boost SV release in response to rapidly delivered stimuli, and thereby, explain the paired-pulse facilitation in *vapb*^{P58S}-expressing neurons (Fig. 5A, C). To examine the rates of Ca²⁺ extrusion, we performed livecell imaging of larval motor neurons expressing GCaMP5G-tdTomato (Daniels et al., 2014; Wong et al., 2014, 2021). To ensure that our assessment of Ca²⁺ extrusion focused on transfer

to the extracellular medium rather than uptake into the ER, we applied the SERCA inhibitor, TG (Lytton et al., 1991; Wong et al., 2021). Incidentally, TG-treatment evokes cytosolic Ca^{2+} transients because of leak of ER stores into the cytosol (Wong et al., 2021), and the kinetics of the return of these transients to baseline reflect the rate of Ca^{2+} extrusion. We found that the decay of TG-evoked GCaMP5G-tdTomato elevations followed a first order exponential function (Fig. 5G). Half-life of this decay was significantly higher in neurons expressing $\operatorname{vapb}^{P58S}$ compared with those expressing vapb^{WT} (Fig. 5G). These data argue in favor of diminished Ca^{2+} extrusion in neurons expressing $\operatorname{vapb}^{P58S}$.

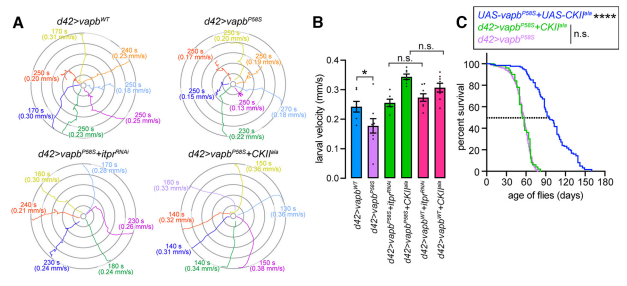


Figure 4. $vapb^{0'585}$ -induced alterations in the velocity of larval peristalsis and adult longevity stem from overlapping and distinct underlying mechanisms. **A**, Images showing the trajectory of larval crawling in an open field. Each colored line represents the path taken by distinct individual larvae. For each larva, duration for which video capture and mean velocity of crawling are noted. Genotypes of the larvae are indicated. **B**, Bar graph showing quantification of crawling velocity of larvae of the indicated genotypes. Values represent mean \pm SEM; *p < 0.05; n.s., not significant, t tests. Dots represent values from distinct larvae. **C**, Lifespan of flies of the indicated genotypes; *****p < 0.0001; n.s., not significant, log-rank tests. Refer to Table 4 for statistical information.

Table 4. Statistical information for the data shown in Figure 4

Figure 4B				
Genotype	Normally distributed?	Values	Values	
$d42 > vapb^{WT}$	Yes	0.24 ±	± 0.02	7
$d42>vapb^{P585}$	Yes	0.18 =	± 0.02	9
$d42 > vapb^{P58S} + itpr^{RNAi}$	Yes	0.23 =	± 0.01	6
$d42 > vapb^{P58S} + CKII^{ala}$	Yes	0.34 =	± 0.01	7
$d42 > vapb^{WT} + itpr^{RNAi}$	Yes	0.27 =	± 0.01	8
$d42 > vapb^{WT} + CKII^{ala}$	Yes	0.30 =	± 0.01	8
Comparison between	Test used	ı	P-value	Test statistics
$d42 > vapb^{WT} d42 > vapb^{P58S}$	t test	(0.04	t = 2.154, df = 13.69
$d42 > vapb^{P58S} + itpr^{RNAi} d42 > vapb^{WT} + itpr^{RNAi}$	t test	(0.3	t = 1.024, df = 11.78
$d42 > vapb^{P58S} + CKII^{ala} d42 > vapb^{WT} + CKII^{ala}$	t test	(0.06	t = 2.114, df = 11.35
Figure 4C				
Comparison between (number of animals)		Test used	<i>P</i> -value	Test statistics
$\frac{UAS-vapb^{P585}+UAS-CKII^{ala}\ (125)\ d42>vapb^{P585}+CKII^{ala}\ (50)\ d42>vapb^{P585}\ (350)}{d42>vapb^{P585}+CKII^{ala}\ (50)\ d42>vapb^{P585}\ (350)}$		log-rank long-rank	<0.0001 0.5	$\chi^2 = 283.9$, df = 2 $\chi^2 = 0.4$, df = 1

Rates of ATP production are unable to keep up with demand in $vapb^{P58S}$ -expressing motor neurons

Electrochemical homeostasis in healthy neurons is maintained by pumps powered by mitochondrially-derived ATP (Fig. 6A). While SERCA and PMCA restore depolarization-induced changes in cytosolic [Ca²⁺] to resting levels, the Na⁺/K⁺ ATPase restores the membrane potential (Fergestad et al., 2006; Chouhan et al., 2012; Ivannikov and Macleod, 2013; Le Masson et al., 2014). The purported coupling of ATP production to neuronal activity, therefore, ensures the availability of ATP for maintenance of Ca²⁺ homeostasis and for membrane repolarization (Fig. 6A, left; Le Masson et al., 2014; Rangaraju et al., 2014; Wong et al., 2021). In ALS neurons, defects in mitochondrial function are predicted to result in a paucity of ATP, which in turn, attenuates the activity of Ca²⁺ and Na⁺/K⁺ ATPases (Fig. 6A, right; Le

Masson et al., 2014). In silico modeling has revealed that the outcome of mitochondrial dysfunction and diminished pump activity is a "deadly loop" comprised of a rapidly burgeoning demand for ATP that remains unmet, and ${\rm Ca}^{2^+}$ dyshomeostasis (Le Masson et al., 2014). Since our findings of diminished ${\rm Ca}^{2^+}$ extrusion in neurons expressing $vapb^{P58S}$ (Fig. 5G) aligns with the notion of ${\rm Ca}^{2^+}$ dyshomeostasis, we asked whether the expression of $vapb^{P58S}$ compromises mitochondrial ATP production, which in principle, could preclude a suitable response to depolarization and ${\rm [Ca}^{2^+}]$ elevation.

Given that mitochondrial Ca²⁺ uptake, needed for energizing those organelles and stimulating TCA dehydrogenases, is disrupted in *vapb*^{P58S}-expressing neurons, it stood to reason that "on-demand" ATP production would be attenuated in these neurons (Duchen, 1992; McCormack and Denton, 1993; Dumollard

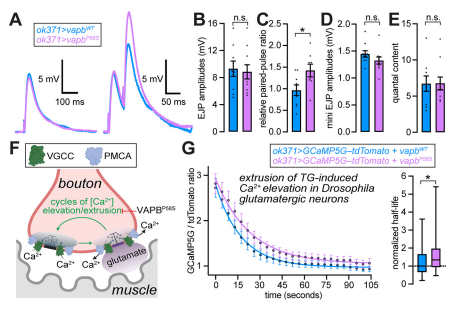


Figure 5. Expression of vapb^{P585} results in delayed extrusion of cytosolic Ca²⁺. A, Representative EJP (left) and pairedpulse EJP (right) traces recorded from larval NMJs isolated from animals expressing vapb^{WT} (blue trace) or vapb^{PS8S} (magenta trace) in motor neurons using ok731-GAL4. B, Bar graph showing quantification of the EJP amplitudes from the data shown in **A**. Values represent mean \pm SEM; n.s., not significant, paired t tests. Dots represent values from distinct NMJs. **C**, Bar graph showing quantification of the paired-pulse ratio (fractional change in the amplitude of the second EJP to that of the first EJP when the two stimulatory pulses were applied 50 ms apart) of the data shown in A. Values represent mean \pm SEM; *p < 0.05, t test. Dots represent values from distinct NMJs. **D**, Bar graph showing quantification of the mini EJP amplitudes. Values represent mean \pm SEM; n.s., not significant, paired t tests. Dots represent values from distinct NMJs. E, Bar graph showing quantification of quantal content (ratio of amplitudes of EJP and mini EJP). Values represent mean \pm SEM; n.s., not significant, paired t tests. Dots represent values from distinct NMJs. F, Model showing that the rates of $[Ca^{2+}]$ elevation and extrusion are necessary for maintaining the fidelity of synaptic transmission. Paired-pulse facilitation without a change in the amplitude of the first pulse, as seen in \mathbf{A} , could be explained by VAPB^{P58S} decreasing the rates of Ga^{2+} extrusion. Image was created with BioRender. G. Left, Traces showing the decay of GCaMP5G/tdTomato ratio after TG-induced cytosolic [Ga²⁺] elevations in motor neurons dissociated from animals of the indicated genotypes. Values were fit to a first order exponential function. Values represent mean \pm SEM of traces from >50 neurons of each genotype. Right, Box plots showing quantification of half-lives of decay of the signals in individual neurons of the indicated genotypes; *p < 0.05, Mann-Whitney test. Refer to Table 5 for statistical information.

et al., 2004; Cárdenas et al., 2010; Ding et al., 2018; Wong et al., 2021). To test this idea, we monitored the cytosolic [ATP]/ [ADP] ratio in larval motor neurons using the genetically-encoded sensor, PercevalHR (Tantama et al., 2013; Wong et al., 2021). In neurons coexpressing $vapb^{WT}$, depolarization by the application of high [K $^+$] (52 mm) led to a transient decrease in the PercevalHR ratio (i.e., [ATP]/[ADP] ratio), whereas the ratio remained unchanged in neurons not challenged with high [K $^+$] (Fig. 6B). In $vapb^{P58S}$ -expressing neurons, however, high [K $^+$] led to a more sustained decline in the PercevalHR ratio (Fig. 6B, see below for quantification of cumulative decline). Since prior reports have indicated that $vapb^{P58S}$ exerts its

Since prior reports have indicated that $vapb^{P58S}$ exerts its effects in fly neurons via a dominant-negative mode of action (Ratnaparkhi et al., 2008), we asked whether a reduction in vapb expression would result in a similar defect in sustaining the [ATP]/[ADP] ratio on depolarization. We obtained a transgenic line for vapb knock-down, $vapb^{RNAi}$, which significantly reduced vapb mRNA levels (Fig. 6C). Upon coexpression with PercevalHR in motor neurons, $vapb^{RNAi}$ led to a sustained decrease in the [ATP]/[ADP] ratio in response to depolarization with high [K⁺] (Fig. 6D). Comparison of the cumulative change in the PercevalHR ratio per unit time revealed that neurons expressing $vapb^{P58S}$ or $vapb^{RNAi}$ exhibited a significantly greater cumulative decrease in PercevalHR ratio in response to high [K⁺] than did neurons expressing $vapb^{WT}$ (Fig. 6E). In contrast, the cumulative

changes in vapb^{P58S} or vapb^{RNAi} were not statistically significant (Fig. 6E). These data indicate that neurons expressing vapb^{P58S} or $vapb^{RNAi}$ were less capable of responding to the ATP burden of depolarization than were neurons with expressing $vapb^{WT}$. Recovery of the PercevalHR ratio during a 30-s window after application of high [K⁺] (Fig. 6B,D, green lines), was significantly greater in neurons overexpressing vapb^{WT} than they were in neurons expressing vapb^{P58S} or vapb^{RNAi} (Fig. 6F). These data show that the rate at which the [ATP]/ [ADP] ratio recovers after depolarization is diminished in neurons expressing vapb^{P58S} or *vapb*^{RNAi}. Finally, we asked whether knock-down of vapb also delays the extrusion of cytosolic Ca^{2+} as we had observed in neurons expressing $vapb^{P58S}$. Relative to neurons not expressing the RNAi transgene, those expressing $vapb^{RNAi}$ exhibited delayed recovery from TG-induced [Ca²⁺] elevation (Fig. 6G). Together, these data indicate that either the knock-down of vapb or the expression of vapb^{P58S} in Drosophila motor neurons compromised the neurons' response to the bioenergetic burden of depolarization, and delayed the rates at which they extrude Ca2+. These conclusions agree with the predictions made by in silico modeling of ALS neurons (Fig. 6A; Le Masson et al.,

Next, we examined the role of OXPHOS in the observed bioenergetic alterations. We found that application of the ATP synthase inhibitor, oligoA, led to the expected decrease of the [ATP]/[ADP] ratio in neurons expressing $vapb^{WT}$, $vapb^{P58S}$, or

vapb^{RNAi}, regardless of whether the neurons were exposed to normal or high [K⁺] (Fig. 7A,B). The cumulative decreases in ratio after inhibition of OXPHOS were comparable in the vapb WT and vapb^{P58S} neurons (Fig. 7A,B). These data imply that on cessation of ATP production, the [ATP]/[ADP] ratio comes to rest at similar values in neurons expressing either $vapb^{WT}$ or $vapb^{P58S}$. These data argue that neuronal [ADP] is not significantly altered in neurons of the two genotypes. In agreement, the cumulative decrease in [ATP]/[ADP] ratio was reduced in depolarized neurons because the burden of depolarization had lowered the [ATP]/[ADP] ratio, even before the application of oligoA. Taken together, our data indicate that the greater cumulative decline in the [ATP]/[ADP] ratio in depolarized vapb^{P58S} neurons, as evident from Figure 6E, reflects the diminished rates of ATP synthesis rather than changes in substrate (i.e., ADP) availability. In neurons expressing vapb NAil, however, oligoA-induced reduction in the [ATP]/[ADP] ratio was significantly higher than that in the other two genotypes, both at normal and high [K⁺] (Fig. 7A,B). These data argue in favor of higher [ADP] in those neurons, which is likely a compensatory response to *vapb* knock-down.

Loss of synaptic transmission during high-frequency stimulation in *vapb*^{P58S}-expressing motor neurons

Many *vapb*^{P58S} phenotypes we describe in this study, including elevated [Ca²⁺] and limited [ATP], are also observed in *drp1*-

Table 5. Statistical information for the data shown in Figure 5

Figure 5B						
Genotype	Normall	ormally distributed? Values		ies	Number of animals	
d42>vapb ^{WT} d42>vapb ^{P58S}	Yes Yes		9.3 ± 1.1 8.9 ± 1.0		5 (n = 10 NMJs) 5 (n = 10 NMJs)	
Comparison betwee	n	Test used	<i>P</i> -value		Test statistics	
$d42>vapb^{WT}d42>$	vapb ^{P58S}	t test	0.77		t = 0.2961, df = 16.68	
Figure 5C						
Genotype	Normally	distributed?	Value	S	Number of anima	
$d42 > vapb^{WT}$ $d42 > vapb^{P58S}$	Yes Yes		•		5 (n = 10 NMJs) 5 (n = 10 NMJs)	
Comparison betwee	n	Test used	P-v	ralue	Test statistics	
$\overline{d42}>vapb^{WT}d42>$	vapb ^{P58S}	t test	0.0	3	t = 2.388, df = 14.1	
Figure 5D						
Genotype	Normally	distributed?	Value	S	Number of anima	
d42>vapb ^{WT} d42>vapb ^{P58S}	Yes Yes		1.44 ± 0.06 1.32 ± 0.07		5 (n = 10 NMJs) 5 (n = 10 NMJs)	
Comparison between		Test used	<i>P</i> -value		Test statistics	
$\overline{d42}>vapb^{WT}d42>$	vapb ^{P58S}	t test	0.1	9	t = 1.372, df = 17.8	
Figure 5E						
Genotype	Normally	y distributed?	Value	es	Number of anima	
d42>vapb ^{WT} d42>vapb ^{P58S}	Yes No		6.73 ± 1.1 6.78 ± 0.9		5 (n = 10 NMJs) 5 (n = 10 NMJs)	
Comparison betwee	n	Test used		<i>P</i> -value	Test statisti	
$d42 > vapb^{WT} d42 > vapb^{$	vapb ^{P58S}	Mann-Whit	ney	0.9	U = 43	
Figure 5G						
Genotype		Normally distrib	uted?	Values		
$d42 > vapb^{WT}$ $d42 > vapb^{P58S}$ Comparison between $d42 > vapb^{WT}$ $d42 > vapb^{WT}$		No No Test used Mann—Whitney		median = median = P-value 0.04		

deficient animals (Verstreken et al., 2005). Since the paucity of ATP in drp1 mutant NMJs precludes the maintenance of synaptic transmission during high-frequency stimulation (Verstreken et al., 2005), we asked whether the shortage of ATP in $vapb^{P58S}$ motor neurons would result in similar rundown of SV release. Although stimulation of the vapb^{P58S} NMJs at 10 Hz led to EJP amplitudes that initially plateaued at values that were \sim 35% higher than baseline, those values transitioned to periods of sustained decay after ~5 min of high-frequency stimulation (Fig. 7C, left, traces with filled circles, green arrow indicates onset of decay). EJP amplitudes in 100% of the traces recorded from vapb^{P58S} animals stimulated at 10 Hz exhibited sustained periods of decay after initial, transient elevations. We reasoned that superimposed on the liminal increase in EJP amplitudes were the rundowns of SV release, which become apparent after ~5 min of high-frequency stimulation. We extracted the rundown components from the traces using the rate-constants of the decay (Fig. 7C, left, traces with filled squares). Over the 10 min duration of the high-frequency stimulation, calculated rundowns in vapb^{P58S} NMJs dropped significantly lower than baseline (Fig. 7C, left). The decay phase of high-frequency traces recorded from $vapb^{WT}$ NMJs was relatively less-pronounced (Fig. 7*C*, right), with only 40% of the traces exhibiting sustained periods of declining EJP values. Calculated rundown in $vapb^{WT}$ NMJs was not significantly lower than baseline (Fig. 7*C*, right). Therefore, high-frequency stimulation leads to a greater loss of synaptic transmission in $vapb^{P58S}$ -expressing motor neurons. In agreement, calculated rundowns in $vapb^{P58S}$ were predicted to plateau at 3.82 mV [95% confidence interval (CI), 3.69–3.94, p<0.0001], whereas rundowns were predicted to plateau at 11.96 mV (95% CI, 11.81–12.11, p<0.0001) in $vapb^{WT}$.

Bioenergetic dysfunction and attendant changes in SV cycling at the drp1 mutant NMJs stem from the absence of local mitochondria because of defects in mitochondrial trafficking to the axon termini (Verstreken et al., 2005). In contrast, coexpression of the mitochondrial marker, mCherry-mito-OMM (Vagnoni and Bullock, 2016), with the vapb variants belied putative defects in the availability of mitochondria at the NMJ (Fig. 7D,E). Mitochondria in the *vapb*^{P58S}-expressing neurons did, however, exhibit subtle morphologic changes, slight yet significant increases in the relative volume and surface-area of each mitochondrion at the termini (Fig. 7F,G). Mitochondrial aspect ratio (i.e., the volume/surface area ratio of each mitochondrion) was \sim 2% lower in $vapb^{P58S}$ NMJs (Fig. 7H), which argues against meaningful changes in fission-fusion dynamics. Collectively, our data argue against changes in mitochondrial trafficking or dynamics as the underlying cause for rundown of synaptic transmission in *vapb*^{P58S} NMJs.

Discussion

Our findings explain the characteristic changes to NMJ morphology observed in animals that harbor vapb deletions or express an ALS-causing variant of the gene (*vapb*^{P58S}; Pennetta et al., 2002; Chai et al., 2008; Ratnaparkhi et al., 2008). While it was known that the reduction in NMJ bouton number and increase in bouton size stem from diminished stability of presynaptic microtubules (Pennetta et al., 2002; Chai et al., 2008; Ratnaparkhi et al., 2008), the molecular underpinnings of these cytoskeletal changes had remained unclear. Our finding that nocodazole decreased the number of presynaptic microtubule loops in neurons expressing $vapb^{WT}$, but not in neurons expressing vapb^{P58S}, argues in favor of microtubule-depolymerization being constitutively elevated in the latter. The additional finding that the morphologic defects in *vapb*^{P58S} NMJs were attenuated by inhibition of CaMKII implicates aberrant CaMKII activation in the observed NMJ phenotypes. These conclusions agree with prior reports of CaMKII regulating microtubule stability by phosphorylating microtubule-associated proteins (Baratier et al., 2006; McVicker et al., 2015; Oka et al., 2017).

We speculate that the MAP1b homolog, Futsch (Hummel et al., 2000; Roos et al., 2000), is the relevant CaMKII target in this context. Previous studies have shown that hyperphosphorylation of Futsch, which provokes microtubule destabilization, results in the loss of Futsch-loops within NMJ boutons (Gögel et al., 2006; Viquez et al., 2006; Wong et al., 2014). The potential involvement of Futsch also agrees with prior reports of reduced Futsch abundance at the NMJ in a fly model of TDP-43-induced proteinopathy, primarily because of diminished translation of *futsch* mRNA (Coyne et al., 2014). Therefore, microtubule destabilization with the underlying involvement of Futsch could be a common occurrence in *Drosophila* models of ALS (Pennetta et al., 2002; Ratnaparkhi et al., 2008; Coyne et al., 2014). Interestingly,

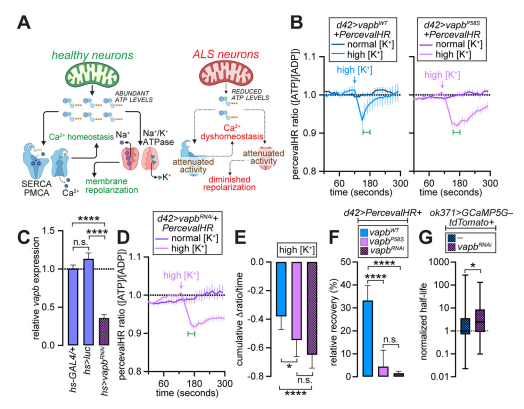


Figure 6. Response to ATP burden of depolarization in motor neurons depends on the presence of functional vapb. A, Model adapted from Le Masson et al. (2014) showing the role for neuronal ATP in mediating ionic homeostasis. In healthy neurons, ATP derived from mitochondrial oxidative phosphorylation (0XPHOS) powers Ca^{2+} ATPases, such as SERCA and PMCA, to maintain Ca^{2+} homeostasis, and the Na^+/K^+ ATPase, which is needed for setting the resting membrane potential and for repolarization of the membrane potential after bouts of depolarization. In ALS neurons, a decrease in mitochondrial ATP production would attenuate the activities of Ca^{2+} and Na^+/K^+ -ATPases. This would be predicted to set in motion a self-reinforcing "deadly loop" comprised of continuously increasing ATP consumption coupled with diminished ATP availability, Ca^{2+} dyshomeostasis, and the eventual loss of membrane potential. Image was created with BioRender. B, D, Representative traces showing normalized PercevalHR ratios in *Drosophila* motor neurons coexpressing the indicated transgenes. Values represent mean \pm SEM; *****p0 neurons of each genotype. Arrows indicate treatments. C0, Bar graph showing relative vapb expression in animals of the indicated genotypes. Values were normalized to the mean in hs-GAL4/+, and represent mean \pm SEM; *****p0 0.0001; n.s., not significant, t1 tests with Bonferroni correction. E1, Bar graph showing the cumulative change in the PercevalHR ratio per unit time in neurons dissociated from animals of the indicated genotypes. Values represent median t20% Cls; **t2000, ***t30, ***t40, **t40, **t50, ***t50, **t50, ***t50, **t50, **t50,

expression of a gain-of-function variant of *vapb* results in the opposite phenotype, increased number of synaptic boutons and Futsch loops (Sanhueza et al., 2014). Given the likelihood that VAPB^{P58S} acts via a dominant-negative mode of action (Ratnaparkhi et al., 2008), there appears to be a direct correlation between the functionality and dosage of VAPB, Futsch-dependent microtubule stability, and bouton number at the larval NMJ (Pennetta et al., 2002; Sanhueza et al., 2014).

Ectopic expression of an overactive TRPV4 channel variant that elevates intracellular [Ca²⁺] and induces neuropathology via CaMKII hyperactivation (Woolums et al., 2020) sets a precedent for the notion that aberrant CaMKII activation in motor neurons implies higher cytosolic [Ca²⁺]. Blocking the sources of presynaptic Ca²⁺ elevation, then, would be expected to restore Ca²⁺ homeostasis, and thereby, ameliorate the consequences of CaMKII hyperactivation. Indeed, reduced expression of ER Ca²⁺ release channels in *Drosophila* neurons counteracts the phenotypes stemming from activating mutations in a voltage-gated Ca²⁺ channel (Brusich et al., 2018). We found that lowering the abundance of any one of the three presynaptic ER Ca²⁺ release channels, Iav, RyR, and IP₃R, restored bouton morphology and number in neurons

expressing *vapb* ^{P58S}. These data also point to the existence of a biphasic, bell-shaped relationship between presynaptic [Ca²⁺] and NMJ morphology, whereby either a supraphysiological increase or decrease in [Ca²⁺] elicits morphologically indistinguishable changes at the NMJ.

Do the effects of ER Ca²⁺ channel knock-down or CKII^{ala} expression on NMJ development in larvae expressing vapb^{P58S} correlate with functional alterations in locomotion? To answer this question, we first asked whether expression of vapb^{P58S} in motor neurons leads to alterations in larval locomotion. We found that crawling velocity in an open field was significantly reduced in larvae expressing vapb^{P58S} relative to those expressing vapb^{WT}. Concomitant knockdown of the gene encoding IP₃R or expression of CKII^{ala} prevented this decrease in the velocity of peristalsis in vapb^{P58S}-expressing larvae. Expression of vapb^{P58S} in motor neurons also leads to premature lethality in adult animals (Wong et al., 2021). In contrast to its restorative effects on NMJ development and larval locomotion, CKII^{ala} had no effect on the abbreviated lifespan observed in adult flies expressing vapb^{P58S} in motor neurons. Therefore, the devel-

opmental and longevity phenotypes stem from partially

Table 6. Statistical information for the data shown in Figure 6

Figure 6C					
Genotype	Normally dist	ributed?	Values		
hs-GAL4/+ hs>luc hs>vapb ^{P58S}	Yes Yes Yes		1.0 ± (1.1 ± (0.4 ± (0.08	
Comparison between	Test used		<i>P</i> -value		Test statistics
hs-GAL4/+ hs>vapb ^{P585}	t test followed by	r Bonferroni correction r Bonferroni correction r Bonferroni correction	0.6 <0.0001 <0.0001		t = 1.414, df = 8.165 t = 9.808, df = 9.999 t = 8.539, df = 8.209
Figure 6E					
Genotype	Normally di	stributed?	Values (median, lo	wer Cl, upper Cl)	
$d42 > PercevalHR + vapb^{WT}$ $d42 > PercevalHR + vapb^{P585}$ $d42 > PercevalHR + vapb^{RNAi}$	No No No		(-0.38, -0.47, - (-0.55, -0.66, - (-0.65, -0.74, -	-0.42)	
Comparison between		Test used		<i>P</i> -value	Test statistics
$\begin{array}{l} d42 > \textit{PercevalHR} + \textit{vapb}^{\textit{WT}} \ d42 > \textit{PercevalHR} + \textit{vapb}^{\textit{PS85}} \\ d42 > \textit{PercevalHR} + \textit{vapb}^{\textit{PS85}} \ d42 > \textit{PercevalHR} + \textit{vapb}^{\textit{RNAi}} \\ d42 > \textit{PercevalHR} + \textit{vapb}^{\textit{WT}} \ d42 > \textit{PercevalHR} + \textit{vapb}^{\textit{RNAi}} \end{array}$	Ai	Mann—Whitney followed by Bor Mann—Whitney followed by Bor Mann—Whitney followed by Bor	ferroni correction	0.01 <0.0001 0.45	U = 44192 U = 39586 U = 35319
Figure 6F					
Genotype	Normally di	stributed?	Values (median, lo	wer Cl, upper Cl)	
$d42 > PercevalHR + vapb^{WT}$ $d42 > PercevalHR + vapb^{P585}$ $d42 > PercevalHR + vapb^{RNAi}$	No No No		(33.24, 26.88, 39.7 (4.47, 0.33, 11.59) (1.55, 1.06, 2.38)		
Comparison between		Test used		<i>P</i> -value	Test statistics
$\frac{d42}{PercevalHR} + vapb^{WT} \frac{d42}{PercevalHR} + vapb^{PS85} \frac{d42}{PercevalHR} + vapb^{PS85} \frac{d42}{PercevalHR} + vapb^{RNAi} $	Ai	Mann—Whitney followed by Bor Mann—Whitney followed by Bor Mann—Whitney followed by Bor	ferroni correction	<0.0001 0.21 <0.0001	U = 15458 U = 24263 U = 13935
Figure 6G					
Genotype	Norr	nally distributed?	Values (mediar	ı, lower CI, upper CI)	
ok731>GCaMP5G-tdTomato ok731>GCaMP5G-tdTomato+vapb ^{RNAi}	No No		Median = 1 Median = 2.5		
Comparison between		Test used		<i>P</i> -value	Test statistics
ok731>GCaMP5G-tdTomato ok731>GCaMP5G-tdTomato+vapb ^{RNAi}		Mann-Whiti	ney 0.01		<i>U</i> = 1410

overlapping, yet distinct, molecular pathways. While the NMJ and larval motor phenotypes were suppressed by concomitant reductions in the abundance of Iav and RyR, or by the expression of the CaMKII inhibitory peptide, none of these manipulations influenced the effects of VAPB^{P58S} on adult lifespan. Only with the knock-down of the gene encoding IP₃Rs do we observe suppression of NMJ development phenotypes, larval locomotion, and adult lethality, though the former two likely involve CMKII attenuation while the latter, as we showed previously (Wong et al., 2021), stems from mitigation of endolysosomal [Ca²⁺] overload.

Given that the NMJ phenotypes in neurons expressing $vapb^{P58S}$ were suppressed by the loss of ER Ca^{2+} channels, which are needed for maintaining presynaptic resting $[Ca^{2+}]$ (Wong et al., 2014), we asked whether VAPB^{P58S} was triggering an increase in resting $[Ca^{2+}]$. Higher presynaptic resting $[Ca^{2+}]$ augments the probability of SV release resulting in increased amplitudes of evoked postsynaptic potentials that is accompanied by paired-pulse depression (Wong et al., 2014). In agreement with prior reports (Chai et al., 2008), we found that expression of $vapb^{P58S}$

induced neither a change in the amplitude of evoked potentials nor the appearance of paired-pulse depression. Rather, $vapb^{P58S}$ NMJs exhibited paired-pulse facilitation, which in combination with the absence of changes in the amplitude of evoked potentials, was consistent with poststimulus accumulation of residual ${\rm Ca^{2+}}$ because of delayed extrusion of presynaptic ${\rm Ca^{2+}}$. Indeed, direct examination of cytosolic ${\rm [Ca^{2+}]}$ using GCaMP5G-tdTomato revealed significantly delayed ${\rm Ca^{2+}}$ extrusion in $vapb^{P58S}$ -expressing motor neurons. These data indicate that the aberrant activation of CaMKII in $vapb^{P58S}$ -expressing neurons arose from the accumulation of residual ${\rm Ca^{2+}}$ because of delayed extrusion.

What could explain the defects in Ca^{2+} extrusion in neurons expressing *vapb*^{P58S}?

The plasma membrane-resident Ca²⁺ ATPase, PMCA, has been shown to play a major role in the extrusion of presynaptic Ca²⁺ in *Drosophila* motor neurons (Ivannikov and Macleod, 2013; Rossano et al., 2013). Given the bioenergetic burden of powering PMCA, we reasoned that a shortage of ATP could underlie the Ca²⁺ dyshomeostasis observed in *vapb*^{P58S}-expressing neurons.

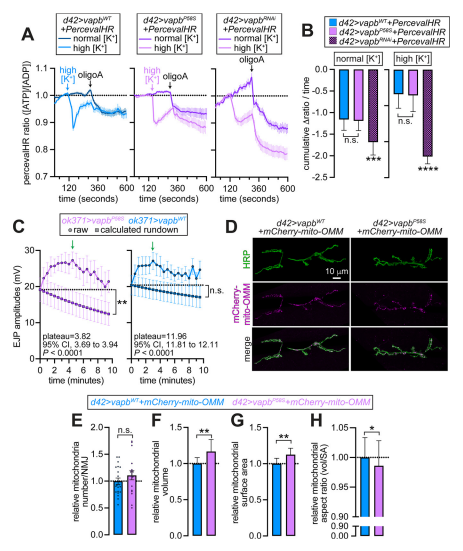


Figure 7. oligoA-induced changes in the [ATP]/[ADP] ratio, and effects of high-frequency stimulation in neurons expressing variants. A, Traces showing PercevalHR ratio after application of oligoA in neurons dissociated from animals of the indicated genotypes. Values represent mean \pm SEM of traces from >20 neurons of each genotype. Arrows indicate treatments. B, Bar graph showing the cumulative change in the PercevalHR ratio per unit time in neurons dissociated from animals of the indicated genotypes. Values represent median \pm 95% Cls; ***p < 0.001, ****p < 0.0001; n.s., not significant, Kruskal–Wallis and Mann-Whitney tests. C, Traces showing EJP amplitudes recorded from NMJs of animals of the indicated genotypes stimulated at 10 Hz for 10 min. Traces connecting filled circles represent raw values that were determined experimentally, and the traces connecting filled-squares represent calculated rundowns fit to first order exponential decay. Green arrows indicate the onset of decay. All values represent mean \pm SEM of EJP values recorded from five NMJ preparations of each genotype; **p < 0.01; n.s., not significant, paired t tests. **D**, Representative confocal images of larval NMJs dissected from animals of the genotypes indicated on the top and stained with antibodies against HRP (green). Mitochondria were labeled by mCherry-mito-OMM (magenta). Scale bar shown in top left image applies to all the images in the panel. *E-H*, Bar graphs showing quantification of the parameters of mitochondrial morphology in animals of the indicated genotypes. Values were normalized to the control mean (\pmb{F}) or to the control median (\pmb{F} - \pmb{H}). Values represent mean \pm SEM (\pmb{F}) or median \pm 95% CIs (\pmb{F} - \pmb{H}). In \pmb{E} , n.s., not significant, t test, and circles represent values from distinct NMJs. In F-H, *p < 0.05, **p < 0.01; Mann—Whitney tests. Refer to Table 7 for statistical information.

In support of this model, mutated VAPB has been shown to disrupt interorganellar transfer of Ca^{2+} from the ER to mitochondria in both mammalian cells and $\operatorname{Drosophila}$ neurons (De Vos et al., 2012; Stoica et al., 2016; Gomez-Suaga et al., 2017; Smith et al., 2017; Xu et al., 2020; Wong et al., 2021). Given the role of matrix $[\operatorname{Ca}^{2+}]$ in ATP production (Duchen, 1992; McCormack and Denton, 1993; Dumollard et al., 2004; Cárdenas et al., 2010; Ding et al., 2018), we asked whether expression of $\operatorname{vapb}^{P58S}$ compromised mitochondrial ATP production. By examining the $[\operatorname{ATP}]/[\operatorname{ADP}]$ ratio in live neurons, we found that

the bioenergetic response to depolarization was relatively stunted in vapb^{P58S}expressing neurons. The cumulative decrease in [ATP]/[ADP] ratio in depolarized *vapb*^{P58S}-expressing neurons was greater than that in neurons expressing $vapb^{WT}$. Eventual recovery of the [ATP]/[ADP] ratio in depolarized neurons was faster if they overexpressed $vapb^{WT}$, which points to the sufficiency of vapb in sculpting the kinetics of neurons' response to the bioenergetic burden of depolarization. Consistent with VAPB^{P58S*} having a dominant negative mode of action (Ratnaparkhi et al., 2008), neurons expressing an RNAi line against vapb also exhibit larger cumulative decrease in the [ATP]/[ADP] ratio, and delayed extrusion of cytosolic Ca^{2+} .

Upon cessation of OXPHOS by application of the ATP synthase inhibitor, oligoA, we found that cumulative decreases in the [ATP]/[ADP] ratio were comparable in neurons expression either vapb variant. Strictly speaking, the levels at which the cytosolic [ATP]/[ADP] ratio settles following application of oligoA, are functions of two parameters: (1) levels of a pool of ATP that is either not available for hydrolysis or is generated from a source other than OXPHOS; and (2) free [ADP]. Absence of evidence supporting the presence of nonhydrolyzable depots of ATP in the cytosol of neuronal cell bodies, however, argues that the cumulative decrease in the [ATP]/[ADP] reflects the abundance of ADP available for ATP synthesis. By extension, the sluggish response to bioenergetic demands of depolarization in *vapb*^{P58S} neurons does not stem from diminished substrate availability. Rather, the rates of ATP synthesis from ADP are likely lower in those neurons, although elevated rates of ATP hydrolysis could also be contributing to the phenotype. Indeed, in silico modeling has predicted that mitochondrial dysfunction in ALS neurons is sufficient to induce a toxic self-reinforcing loop of increased ATP consumption and a progressive decrease in the availability of ATP (Le Masson et al., 2014). In neurons expressing the RNAi line against vapb,

oligoA-induced cumulative decrease in [ATP]/[ADP] ratio was significantly greater than in the other genotypes. These data argue in favor of basally higher net [ADP] on *vapb* knock-down, which could be a compensatory response to sustained defects in ATP production. The fact that these neurons were still unable to meet the bioenergetic demands of depolarization speaks to profound defects in activity-dependent ATP synthesis in neurons with lower *vapb* expression. The apparent absence of compensation in neurons expressing *vapb*^{P588} could imply a role for *vapb*

Table 7. Statistical information for the data shown in Figure 7

Figure 7B					
Genotype	Normally distributed?		Values (median, lower CI, upper CI)		
d42>PercevalHR+vapb ^{WT} norm K d42>PercevalHR+vapb ^{PS8S} norm K d42>PercevalHR+vapb ^{RNAI} norm K d42>PercevalHR+vapb ^{WT} high K d42>PercevalHR+vapb ^{PS8S} high K	No No No No No		(-1.16, -1.41, -1.01) (-1.19, -1.42, -0.78) (-2.08, -1.98, -1.48) (-0.34, -0.54, -0.23) (-0.36, -0.58, -0.28)		
$d42 > PercevalHR + vapb^{RNAi}$ high K	No		(-1.21, -1.31, -0.99)		
Comparison between	Test used		<i>P</i> -value	Test statistics	
d42>PercevalHR+vapb ^{WT} norm K d42>PercevalHR+vapb ^{PS8S} norm K d42>PercevalHR+vapb ^{WT} norm K d42>PercevalHR+vapb ^{PS8S} norm K	Mann—Whitney followed by Bonfo Kruskal—Wallis	erroni correction	1 0.0004	U = 2637Kruskal—Wallis statistic = 15.58	
d42>PercevalHR+vapb ^{RNAi} norm K d42>PercevalHR+vapb ^{WT} high K d42>PercevalHR+vapb ^{PS85} high K	Mann—Whitney followed by Bonfo	erroni correction	0.9	<i>U</i> = 1706	
d42>PercevalHR+vapb ^{WT} norm K d42>PercevalHR+vapb ^{PS8S} norm K d42>PercevalHR+vapb ^{RNAi} norm K	Kruskal—Wallis		<0.0001	Kruskal—Wallis statistic = 65.55	
Figure 7C					
Comparison between	Test used		<i>P</i> -value	Test statistics	
ok371>vapb ^{WT} ok371>vapb ^{P585}	paired <i>t</i> test paired <i>t</i> test		0.18 0.001	t = 1.632, df = 4 t = 8.347, df = 4	
Figure 7E					
Genotype	Normally distributed	?	Values	Number of animals	
d42>mCherry-mito-OMM+vapb ^{WT} d42>mCherry-mito-OMM+vapb ^{P585}	Yes Yes		1.0 ± 0.06 1.1 ± 0.08	11 (n = 21 NMJs) 9 (n = 18 NMJs)	
Comparison between		Test used	<i>P</i> -value	Test statistics	
$d42>m$ Cherry-mito-OMM $+$ vapb WT $d42>m$ Ch	erry-mito-OMM+vapb ^{P58S}	t test	0.3	t = 1.058, df = 30.65	
Figure 7F Genotype	Normally distributed?	Values ((median, lower CI, upper CI)	Number of animals	
$d42>m$ Cherry-mito- 0 MM $+vapb$ WT $d42>m$ Cherry-mito- 0 MM $+vapb$ P585	No No	(1.0, 0.92, 1.1) (1.2, 1.1, 1.3)		11 $(n = 21 \text{ NMJs})$ 9 $(n = 18 \text{ NMJs})$	
Comparison between		Test used	<i>P</i> -value	Test statistics	
$d42>mCherry-mito-OMM+vapb^{WT}d42>$	erry-mito-OMM+vapb ^{P58S}	Mann-Whitney	0.0025	<i>U</i> = 2,042,530	
Figure 7G					
Genotype	Normally distributed?	Values (median, lower CI, upper CI)		Number of animals	
$d42>m$ Cherry-mito- $OMM+vapb^{WT}$ $d42>m$ Cherry-mito- $OMM+vapb^{P58S}$	No No	(1.0, 0.94, 1.1) (1.12, 1.1, 1.2)		11 (n = 21 NMJs) 9 (n = 18 NMJs)	
Comparison between		Test used	<i>P</i> -value	Test statistics	
$d42>m$ Cherry-mito-OMM $+$ vapb WT $d42>m$ Ch	erry-mito-OMM+vapb ^{P58S}	Mann-Whitney	0.0071	<i>U</i> = 2,055,518	
Figure 7H					
Genotype	Normally distributed?	Values (median, lower CI, upper CI)		Number of animals	
d42>mCherry-mito-OMM+vapb ^{WT} d42>mCherry-mito-OMM+vapb ^{P585}	No No	(1.0, 0.97, 1.03) (0.98, 0.92, 1.03)		11 (n = 21 NMJs) 9 (n = 18 NMJs)	
Comparison between		Test used	<i>P</i> -value	Test statistics	
$d42>mCherry-mito-OMM+vapb^{WT}d42>$	erry-mito-OMM+vapb ^{P58S}	Mann-Whitney	0.01	<i>U</i> = 2,060,657	

dosage in initiating the compensatory response. Nevertheless, the paucity of [ATP] in $vapb^{P58S}$ -expressing neurons during periods of depolarization (i.e., neuronal activity), is consistent with diminished ${\rm Ca}^{2^+}$ extrusion, which in-turn could provoke the

hyperactivation of CaMKII and attendant changes in NMJ morphology.

Maintenance of synaptic transmission at *Drosophila* larval NMJs during periods of high-frequency stimulation depends on

a steady supply of ATP. A local shortage of ATP, for instance, because of an absence of presynaptic mitochondria in drp1 mutants, results in rundown of SV release during high-frequency stimulation (Verstreken et al., 2005). These rundowns occur because of the inability of the synapses to meet the energy demands of SV cycling, and diminished recruitment of the reserve pool of SVs that are otherwise mobilized to replace the rapid depletion of the readily-releasable pool of vesicles (Verstreken et al., 2005). In either the $vapb^{\overline{W}T}$ -expressing or vapb^{P58S}-expressing NMJs, high frequency stimulation led to initial elevation of synaptic transmission. Superimposed over these liminal elevations were rundowns that became obvious after ~5 min of stimulation, and were significantly greater in vapb^{P58S}-expressing neurons. These data agree with enhanced rundown of SV release in motor neurons expressing vapb^{P58S}. The magnitude of this phenotype, however, was smaller than that observed in the drp1 mutants (Verstreken et al., 2005). This discrepancy is explained by the absence of presynaptic mitochondria in drp1 mutant neurons (Verstreken et al., 2005), while we observed no significant changes in the number of presynaptic mitochondria in vapb^{P58S}. The relative severity of rundown at drp1 mutant NMJs is likely because of constitutively lower [ATP] in those mutants (Verstreken et al., 2005), whereas shortage of ATP in $vapb^{P58S}$ -expressing neurons is contingent on depolarization.

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