FARS2 deficiency in *Drosophila* reveals the developmental delay and seizure manifested by aberrant mitochondrial tRNA metabolism

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ABSTRACT

Mutations in genes encoding mitochondrial aminoacyl-tRNA synthetases are linked to diverse diseases. However, the precise mechanisms by which these mutations affect mitochondrial function and disease development are not fully understood. Here, we develop a *Drosophila* model to study the function of dFARS2, the *Drosophila* homologue of the mitochondrial phenylalanyl–tRNA synthetase, and further characterize human disease-associated FARS2 variants. Inactivation of dFARS2 in *Drosophila* leads to developmental delay and seizure. Biochemical studies reveal that dFARS2 is required for mitochondrial tRNA aminoacylation, mitochondrial protein stability, and assembly and enzyme activities of OXPHOS complexes. Interestingly, by modeling *FARS2* mutations associated with human disease in *Drosophila*, we provide evidence that expression of two human FARS2 variants, p.G309S and p.D142Y, induces seizure behaviors and locomotion defects, respectively. Together, our results not only show the relationship between dysfunction of mitochondrial aminoacylation system and pathologies, but also illustrate the application of *Drosophila* model for functional analysis of human disease-causing variants.

INTRODUCTION

The aberrant aminoacylation in mitochondrial tRNAs have been linked to a wide spectrum of clinical presentations (1–6). Aminoacylation, the attachment of an amino acid to a tRNA, is catalyzed by mitochondrial aminoacyl-tRNA synthetases (mt-aaRSs) such as mitochondrial phenylalanyl–tRNA synthetase (FARS2) (7–9). In animal, 22 mitochondrial tRNAs are encoded by its own genome (mtDNA) (10,11), while all 19 mt-aaRSs are encoded by the nuclear genome, synthesized in cytosol and subsequently imported into mitochondria (12–14). Clinical presentations linked to aminoacylation defects caused by these mt-aaRS defects usually display a marked bias for the central nervous system, including the encephalopathy, leukodystrophy, Perrault syndrome, sensorineural deafness and visual impairment (15–22). The clinical spectrum of *FARS2* deficiency ranges from the infantile-onset phenotype, characterized by epileptic mitochondrial encephalopathy, to the later-onset phenotype, characterized by spastic paraplegia, less severe neurologic manifestations, and longer survival (23–30). In particular, patients carrying the homozygous p.Y144C and p.G309S or compound heterozygous p.G309S with p.R153G mutations in the *FARS2* developed infantile-onset epileptic mitochondrial encephalopathy with seizure, developmental delay and truncal hypotonia from the birth to age of six month (23,25,27,29). The primary defects in these *FARS2* mutations were the deficient aminoacylation of tRNAPhe (26,28). The aberrant tRNA^Phe^ metabolism impaired mitochondrial translation and subsequent deficiencies of ox-
ative phosphorylation (25, 28). However, the pathophysiology of FARS2 deficiency is still poorly understood due to the lack of animal disease model.

*Drosophila* is a powerful model organism for understanding pathophysiology of human diseases (31). To investigate whether *FARS2* deficiency caused the epileptic mitochondrial encephalopathy and spastic paraplegia *in vivo*, we produced the *Drosophila dFARS2* knockout mutants by genome editing using the CRISPR/Cas9 system and *dFARS2* knockdown model using RNAi approach. These *dFARS2* knockout and knockdown fly models were characterized for the phenotypes including developmental delay and seizure. Biochemical analysis further revealed that *dFARS2* deficiency led to defects in mitochondrial tRNA<sup>phe</sup> metabolism, translation, assembly and activity of oxidative phosphorylation system (OXPHOS) complexes. Finally, we produced humanized fly models by introducing human wild type FARS2, p.G309S and p.D142Y pathogenic variants into *Drosophila dFARS2* mutants and analyzed the pathologic consequence of human disease-causing FARS2 p.G309S and p.D142Y mutations.

**MATERIALS AND METHODS**

*Drosophila* stocks

All flies were reared on standard corn medium at 25°C. The following fly stocks were used: *UAS-dFARS2-RNAi* (Bloomington *Drosophila* Stock Center, BDSC64869), *Daughterless-Gal4* (Da-Gal4), *elav-Gal4*, y<sup>1</sup> M{vas-int.Dm}ZH-2A w<sup>®</sup>; M{3xP3-RFP.attP}ZH-86Fb (Bloomington *Drosophila* Stock Center, BDSC24749) and y<sup>1</sup> M{vas-Cas9}ZH-2A w<sup>118</sup>/FM7c (Bloomington *Drosophila* Stock Center, BDSC51323).

Generation of *Drosophila* dFARS2 mutants and transgenic flies expressing dFARS2 or human FARS2

*Drosophila* dFARS2<sup> ΔKO </sup>mutant allele was generated through the CRISPR-Cas9 system (32–34). A guide RNA sequence against *dFARS2* gene was designed by a CRISPR guide RNA analysis tool (http://tools.flycrispr.molbio.wisc.edu/targetFinder/). The DNA template for transcription of single-stranded guide RNA (sgRNA) was synthesized by PCR amplification according to the manufacturer’s protocol with the forward primer 5′-TAATACGACTCACTATAGGGTACACTTGTTGTATTTGAAGTG-3′ and reverse primer 5′-CCGGATCGGGGTACATGCTTCTGACGCTTC-3′. The PCR products were sequenced to assess the mutation in the target region. To generate the pAttB-dFARS2 construct, a 2356 bp genomic DNA fragment containing the *Drosophila FARS2* gene (CG13348) was amplified with the corresponding primers (Supplementary Table S1), and then subcloned into the pAttB vector. To generate the pUAST-FARS2-AttB construct, a human FARS2 CDS fragment (1356 bp) was amplified with the corresponding primers (Supplementary Table S1), and then subcloned into the pUAST-AttB vector. Mutation of c.G925A (p.G309S) and c.G424T (p.D142Y) were introduced through PCR based site-directed mutagenesis to give rise to pUAST-FARS2-p.G309S-AttB or pUAST-FARS2-p.D142Y-AttB construct. Primers used are listed in Supplementary Table S1. All constructs were sequenced and inserted at the 86FB landing site (BDSC24749) on the third chromosome using *phi-C31* integrase-mediated site-specific transgenesis.

**S2 cell culture and transfection**

*Drosophila* S2 cells were cultured in Schneider’s medium (Gibco, 21720–001) with 10% FBS (Gibco, 10099-141) at 25°C. The full length *dFARS2* cDNA fragment was amplified by PCR using *FARS2* gene-specific primers (Forward: 5′-CCGGATCGGGGTACATGCTTCTGACGCTTC-3′ and Reverse: 5′-CCGGATCGGGGTACATGCTTCTGACGCTTC-3′), and subcloned into a pAC5.1-Flag vector to generate the pAC5.1-dFARS2-Flag construct. S2 cells were transfected with pAC5.1-dFARS2-Flag plasmid using Effectene Transfection Reagent (QIAGEN, 301425).

Mitochondrial tRNA analysis

Total RNA was extracted with TRIzol (Invitrogen, 15596018) from the second instar larvae and adult fly heads. The tRNA Northern blot analysis was performed as detailed elsewhere (22,35,36). Digoxigenin (DIG) labeled Probe sequences for mitochondrial tRNA<sup>phe</sup>, tRNA<sup>lys</sup>, tRNA<sup>thr</sup>, tRNA<sup>leu</sup>(CUN), tRNA<sup>asp</sup>, tRNA<sup>his</sup>, tRNA<sup>tyr</sup>, tRNA<sup>ile</sup>s and 5S rRNA were provided in Supplementary Table S2. The hybridization and quantification of density in each band were performed as detailed previously (37).

The aminoacylation assays were carried out as detailed elsewhere (37,38). To further distinguish non-aminoacylated tRNA from aminoacylated tRNA, samples of tRNAs were decacylated with the treatment of 0.2 M NaOH after heating for 10 min at 60°C (pH 9) and then run in parallel (37,38). Oligodeoxynucleotide probes for mitochondrial tRNA<sup>phe</sup>, tRNA<sup>lys</sup>, tRNA<sup>thr</sup> were described as above. The hybridization and quantification were carried out as described previously (37,38).

**qRT-PCR and mtDNA copy number analysis**

For qRT-PCR, total RNA was extracted from the second instar larvae with TRIzol (Invitrogen). Reverse transcription was performed using HiScript II 1st Strand cDNA Synthesis Kit (Vazyme, R211-02). For the measurement of mtRNA copy number, total genomic DNA was isolated from the second instar larvae using MiniBEST Universal Genomic DNA Extraction Kit (TAKARA, 9765) according to the manufacturer’s protocol. Quantitative PCR analysis was carried out using an ABI 7900HT Fast Real-Time
PCR System with a ChamQ SYBR qPCR Master Mix (Vazyme, Q311-02). The primers used were provided in the Supplementary Table S3.

**Isolation of mitochondria**

Mitochondria were isolated from second instar larvae according to the method described by previously (39). Approximately 2 ml of second instar larvae were homogenized in 5 ml ice-cold mitochondria isolation buffer [225 mM d-mannitol, 75 mM sucrose, 30 mM Tris–HCl, pH 7.4] containing the PhosSTOP™ phosphatase inhibitor (Roche, 4906845001) and cOmplete™ protease inhibitor cocktail (Roche, 4693132001). The homogenate was centrifuged twice at 600 × g for 10 min at 4°C. After each centrifugation, the supernatant was transferred to a new tube and centrifuged at 7000 × g for 10 min at 4°C. The resulting pellet was resuspended in 1 ml ice-cold mitochondria isolation buffer. The resuspended pellet was homogenized again and then centrifuged at 9000 × g for 10 min at 4°C. The final pellet containing mitochondria was stored at −80°C.

**Western blot analysis**

Second instar larvae or adult fly heads were collected and homogenized in RIPA lysis buffer [50 mM Tris–HCl, 150 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.5% SDS] containing the PhosSTOP™ phosphatase inhibitor (Roche, 4906845001) and cOmplete™ protease inhibitor cocktail (Roche, 4693132001) by grinding at 4°C for 30 min. The protein concentration was measured with BCA assay Kit (Generay, gk5012) according to the manufacturer’s instructions. Samples were electrophoresed on a 10% SDS-PAGE and transferred to a polyvinylidene fluoride (PVDF) membrane. The membrane was blocked with 5% non-fat powdered milk dissolved in 1× TBST [20 mM Tris–HCl, 500 mM NaCl, 0.1% Tween 20, pH 7.4] for 1 h at room temperature. After incubation with the primary antibodies overnight at 4°C, the membrane was probed with the secondary antibodies for 1 h at RT. The primary antibodies used include anti-NDUFS3 (1:1000; Abcam, ab147711), anti-UQCRFS1 (1:1000; Abcam, ab14746), anti-CO3 (1:1000; Abcam, ab110259), anti-NDUFS1 (1:1000; Proteintech, 12444-1-AP), anti-ATPSA (1:2000; Abcam, ab14748), anti-SHDA (1:1000; Abcam, ab209986), anti-TFAM (1:100; BOSTER, ba2827) and anti-Porin (1:2000; Abcam, ab14734). Anti-dFARS2 antibody was raised in rabbit against the C-terminal 14 amino acids ([C]VDKFHKPKTGKSSV of the *Drosophila* FARS2 (1:1000; GenScript). Horseradish peroxidase conjugated Goat anti-mouse IgG (GenScript, A00160) and goat anti-rabbit IgG (GenScript, A00098) were used as secondary antibodies. Proteins were detected using the Chemi Lucent ECL detection reagents (Millipore, WBKLS0500). The blot was scanned using a gel imaging system and the image was analyzed by ImageJ software.

**Biochemical assays of respiratory chain complexes and in-gel activity assays**

BN-PAGE was performed on mitochondrial protein extracted from second instar larvae as detailed elsewhere (40–42). Briefly, mitochondrial proteins were isolated with solution buffer A [50 mM NaCl, 50 mM imidazole, 2 mM 6-amino-caproic acid, 1 mM EDTA, 2% Triton X-100, pH 7.4] containing the PhosSTOP™ phosphatase inhibitor (Roche, 4906845001) and cOmplete™ protease inhibitor cocktail (Roche, 4693132001). 50 μg of mitochondrial protein was mixed with BN-PAGE loading buffer [final concentration: 0.5% (w/v) Coomassie Blue G-250, 50 mM 6-amino-caproic acid, 20% glycerol, pH 7.0] and separated on a 3~11% BN-PAGE gel. Cathode buffer [50 mM Tricine, 7.5 mM imidazole, 0.02% Coomassie Blue G-250, pH 7.0] and anode buffer [25 mM imidazole, pH 7.0] were used for electrophoresis. The samples in the gel were soaked in 5% Coomassie Blue G-250 for Coomassie blue staining or transferred to a PVDF membrane. The membrane was probed with primary and secondary antibodies as described above.

For in-gel activity assays, the BN-PAGE gels were incubated in ice-cold ddH₂O at 4°C for 30 min, then transferred to complex specific solutions at room temperature for 5 h. The Complex I solution was as follows: 100 mM Tris-HCl pH 7.4, 1 mg/ml Nitrotetrazolium blue chloride (NBT) and 0.1 mg/ml NADH. The Complex II solution was as follows: 5 mM Tris–HCl pH 7.4, 2.5 mg/ml NBT, 0.2 mM phenazine methasulphate and 20 mM sodium succinate (43).

**Measurement of mitochondrial respiratory complex activity**

Mitochondria were isolated as described above. Enzymatic activities of single respiratory chain complexes were assayed as previously described (44–46). Briefly, complex I (NADH ubiquinone oxidoreductase) activity was determined by following the oxidation of NADH with ubiquinone as the electron acceptor. The activity of Complex II was examined through the artificial electron acceptor DCPIP. Complex III (ubiquinone cytochrome c oxidoreductase) activity was measured as the reduction of cytochrome c (III) using D-ubiquinol-2 as the electron donor. The activity of complex IV (cytochrome c oxidase) was monitored by following the oxidation of cytochrome c (II). The activity of complex V was explored through the NADH oxidation via conversion of phosphoenolpyruvate to lactate by two step reactions. Citrate synthase (CS) activity was determined by catalyzing the reaction of oxaloacetic acid and acetyl-CoA with 5,5′-dithio-bis-nitrobenzoic acid as the chromogenic agent. All mitochondrial respiratory complex activities were expressed relative to citrate synthase (CS) activity.

**Immunofluorescence staining**

For MitoTracker staining, S2 cells were incubated for 40 min in 100 nM MitoTracker Red (Cell Signaling Technology, #9082), and then fixed with 1× PBS containing 4% paraformaldehyde (PFA) for 20 min at RT. Next, S2 cells were washed three times with 1× PBS containing 0.3% Triton-X (PBT) for 20 min at RT. The samples were blocked with 3% BSA in PBT for 1 h, and then incubated with primary antibodies overnight at 4°C. After three washes with 1× PBT, the samples were incubated with secondary antibodies for 1 h at RT. DAPI (1:1000; Invitrogen, D1306) was added for the last 10 min. The samples were washed three
more times with 1× PBT and mounted in Vectashield. The primary antibodies used was rabbit anti-flag (1:1000; Sigma-Aldrich, F2555). The secondary antibodies used was goat anti-rabbit Alexa Fluor 568 (1:1000; Thermo Fisher Scientific, A-11036). Images were obtained with an Olympus FV1000 confocal microscope and processed using ImageJ and Photoshop.

**Bang-sensitive behavioral testing**

Bang-sensitive behavioral testing was performed as described previously (47,48). Flies were collected under CO2 at 3 days after eclosion and kept at 10 flies/vial for up to 10 days. Flies were then transferred to an empty vial 1 h before behavioral analysis. For testing, vials were mechanically stimulated by placement in a vortex mixer (Vortex-Genie 2) at the maximum speed for 20 s. The number of flies that displayed the bang-sensitive phenotype (seizure and paralysis) after the vortex were recorded. These flies were also used to analyze the recovery time. The time for each bang-sensitive fly to right itself after vortexing immediately was recorded. All flies were tested only once and the results of the same genotype were combined.

**Transmission electron microscope**

Fly brains were dissected and fixed with 2.5% glutaraldehyde in 1× PBS overnight and then washed three times with 1× PBS for 15 min at each step. Samples were post fixed with 1% OsO4 in 1× PBS for 1 h and washed three times with 1× PBS for 15 min at each step. The fixed samples were dehydrated by a graded series of ethanol (30%, 50%, 70%, 80%, 90% and 95%) for about 15 min at each step and absolute ethanol for 20 min, and then transferred to absolute acetone for 20 min. Samples were then infiltrated with and embedded in Spurr resin. Thin sections were cut using ultramicrotome (Leica, EM UC7) and stained with uranyl acetate and alkaline lead citrate. Afterwards, the sections were examined using a Hitachi Model H-7650 transmission electron microscope.

**Survival rate analysis**

For survival rates from the first instar larvae to adult, 30 first instar larvae were collected and cultured in vials with the standard medium at 25°C, and the number of pupa and eclosed adults was counted every 24 h.

**Climbing assay**

Flies were collected as described above. For the assay, groups of 10 female flies were placed in empty vials and tapped to the bottom of the vial, and the number of flies that climbed 8 cm above the bottom within 10 s was counted.

**Statistical analysis**

Data shown in the figures are reported as mean ± SD. Statistical analysis was performed using the unpaired, two-tailed Student’s t-test contained in the Microsoft-Excel program. Significance levels are indicated in the figure legends.

**RESULTS**

*Drosophila* FARS2 is a highly conserved mitochondrial protein

*Drosophila* FARS2 (CG13348, hereafter referred to as dFARS2) encodes a 453 amino acid protein with the typical mitochondrial target sequence with 27 residues at the N-terminus, predicted by MitoProt program (49). Alignment of dFARS2 with its homologs of other organisms, including *Homo sapiens*, *Mus musculus*, *Danio rerio*, *Drosophila melanogaster* and *Saccharomyces cerevisiae* revealed an extensive conservation of protein sequence (Supplementary Figure S1). In particular, dFARS2 shares an overall amino acid identity of 50% and 68% similarity with human FARS2, respectively (Supplementary Figure S1).

To test whether dFARS2 is localized to mitochondria, we transfected a dFARS2 construct tagged with a C-terminal FLAG into *Drosophila* S2 cells and examined subcellular localization by immunofluorescence analysis. Using an antibody against FLAG and Mito Tracker probes, a dye to label the mitochondrial network, we observed a complete overlap of both fluorescence signal in the transfected cells (Figure 1A). These data demonstrate the mitochondrial localization of dFARS2.

**Generation of *Drosophila* dFARS2 deficiency**

To explore the pathological consequences of dFARS2 mutation, we first generated a *Drosophila* model with the loss-of-function dFARS2 mutants through the CRISPR/Cas9 system (Figure 1B) (32,50). One mutant allele, dFARS2KO, was obtained. This allele harbored a 1 bp deletion followed by an 11 bp insertion in exon 1 of dFARS2 (Homozygous and heterozygous dFARS2KO mutant flies were described as KO and het.KO respectively, while wild type flies were described as WT). In fact, this deletion caused a frame shift at codon 87 and a new stop codon at codon 101 downstream of the translation start (Figure 1C). This allele was confirmed by Sanger sequencing, PCR and Western blot analysis (Figure 1D–F). We also generated fly models with the knockdown of dFARS2 by using a UAS-dFARS2 RNAi line in combination with the GAL4/UAS binary expression system (dFARS2 knockdown flies were described as RNAi and control flies were described as Ctrl). The knockdown of dFARS2 by RNAi with a ubiquitously expressed *Daughterness-Gal4* (*Da-Gal4*) or a pan-neuronal Gal4 (*elav-Gal4*) was further confirmed by the reduced level of dFARS2 protein in whole body (Figure 1G) or fly heads (Figure 1H) using western blot analysis.

**dFARS2 deficiency causes developmental delay**

Mutations in FARS2 are associated with several pathological states in humans, notably developmental delay and seizure (29). However, the cause-and-effect relationship has not been established in a highly tractable animal model. We took advantage of *Drosophila* dFARS2 deficiency models generated in this study to examine the pathological consequence.

The homozygous dFARS2KO mutants were embryonic viable, and the newly hatched larvae were able to grow and
Figure 1. Generation of Drosophila models with dFARS2 deficiency. (A) Mitochondrial localization of dFARS2 in S2 cells. Cells were stained for Mito-Tracker (Red) and anti-Flag antibody (Green). DAPI was used to label DNA (Blue). Scale bar represents 10 μm. (B) Genome editing of dFARS2 using the CRISPR/Cas9 system. sgRNA sequence is indicated in blue and PAM sequence is indicated in orange. (C) Schematic representation of the dFARS2 wild type (WT) and dFARS2 mutant (KO) proteins. MTS indicates mitochondrial targeting sequence. (D) Partial Sanger sequencing chromatograms of the sgRNA targeting regions showing for WT and dFARS2KO. A hyphen indicates a deletion and a red letter indicates an insertion. (E) PCR analysis of the sgRNA targeting region at the dFARS2 locus. (F–H) Western blot analysis of dFARS2 in protein extracts from WT and dFARS2KO second-instar larvae (F), the control and Da-Gal4 driven dFARS2 knockdown second-instar larvae (G), the control and elav-Gal4 driven dFARS2 knockdown fly heads (H).

dFARS2 deficiency causes seizure phenotype

Seizure in the Drosophila model shares many characteristics similar to those of seizures in humans (47,50,51). Bang sensitivity assay has been widely used to measure seizure susceptibility in Drosophila (47,48). In this assay, the time required for the flies to right themselves and move after a mechanical stress was recorded to determine seizure susceptibility (47,48). The number of flies displaying bang-sensitive phenotype was also counted. In response to a strong mechanical stress, wild type flies recovered quickly while bang-
Figure 2. dFARS2 deficiency leads to the developmental delay. (A) Images showing wild type (WT) and dFARS2 knockout (KO) larvae, from 1 to 5 days after egg laying (AEL). Rescue denotes the dFARS2KO carrying the genomic rescue transgene for dFARS2. Scale bars: 1 mm. (B) Graph showing the percentage of survivors for WT, dFARS2KO and rescue larvae from 1 to 5 days AEL. n = 5. (C) Graph showing the relative size of WT, dFARS2KO and rescue larvae at 5 days AEL. n = 10. (D) Images showing control and Da-Gal4 driven dFARS2 knockdown animals at various developmental stages. Scale bars: 1 mm. (E) Graph showing the percentage of survivors for control and Da-Gal4 driven dFARS2 knockdown larvae. n = 5. (F) Graph showing the relative size of control and Da-Gal4 driven dFARS2 knockdown larvae at 5 days AEL. n = 10. (G) Images showing control and elav-Gal4 driven dFARS2 knockdown animals at various developmental stages, from 1 to 15 days AEL. Scale bars: 1 mm. (H) Graph showing pupariation curves for control and elav-Gal4 driven dFARS2 knockdown larvae. n = 3. (I) Graph showing eclosion curves for control and elav-Gal4 driven dFARS2 knockdown pupae. n = 4. (J) Graph showing the survival rate of control and elav-Gal4 driven dFARS2 knockdown adult flies. n = 3. Data are presented as means ± SD. ****P < 0.0001. NS, not significant.
sensitive mutants showed a characteristic seizure pattern (47, 48). Ten-day-old elav-Gal4 driven dFARS2 knockdown and control flies were tested for bang sensitivity to assess their susceptibility to seizure. The dFARS2 knockdown female and male flies were bang sensitive and took a long time to recover from a 20-s vortex, whereas the control flies were able to right themselves immediately or within a few seconds (Figure 3A–C, supplementary video 1). Due to the larval lethality of the homozygous dFARS2KO mutants, the heterozygous dFARS2KO mutants were examined and no seizure behaviors were observed (Supplementary Figure S3C, D). Furthermore, the brain of dFARS2 knockdown adult flies appeared to be smaller than in the controls (Figure 3D). These data suggest that dFARS2 deficiency induces seizure phenotype in Drosophila.

Next, we examined the ultrastructure of mitochondria in the central brain. Thin sections of the central brain regions of 10-day-old elav-Gal4 driven dFARS2 knockdown and age-matched control flies were examined using transmission electron microscopy. Mitochondria in the controls were densely packed with intact cristae. In contrast, dFARS2 knockdown flies displayed loosely packed mitochondria with disorganized cristae (Figure 3E and F). The cristae became round and gave mitochondria a honeycomb-like appearance (Figure 3E and F). The mitochondria morphological defects in the dFARS2 knockdown flies suggest that the seizure phenotypes could be derived from mitochondrial dysfunction.

**Reduced aminoacylation of mitochondrial tRNA^Phe**

As dFARS2 is a mitochondrial phenylalanyl-tRNA synthetase, we next performed a series of biochemical assays to further confirm the validity of our dFARS2 deficiency models.

To evaluate whether the dFARS2 deficiency affected the mitochondrial tRNA aminoacylation, we measured the level of aminoacylation using Northern Blot. In order to do this, total RNA was isolated from the larvae or adult fly heads and separated on an acid urea polyacrylamide gel which allows separation of uncharged tRNA from slow migrating aminoacylated tRNA. The presence of an mt-tRNA was assayed by hybridization to the complementary oligonucleotide probes. As a control for deacylated tRNA, an aliquot of each sample was deacylated under alkaline conditions (DA) and separated on the gel along with the untreated sample (Amino). The level of aminoacylated mt-tRNA^Phe was reduced in dFARS2KO mutants compared to those in wild type larvae (Figure 4A). However, the aminoacylation levels of two other mitochondrial tRNA, mt-tRNA^Lys and mt-tRNA^Thr, were similar in both wild type and dFARS2KO mutant larvae (Figure 4A). These results were also confirmed in dFARS2 knockout larvae (Figure 4B). Da-Gal4 or elav-Gal4 driven dFARS2 RNAi caused a reduction of aminoacylated mt-tRNA^Phe in third instar larvae or fly heads, as compared to the control (Figure 4B).

**Increased steady-state levels of mitochondrial tRNAs**

To examine the effect of dFARS2 deficiency on the steady-state level of mitochondrial tRNAs, total RNAs were isolated and analyzed by Northern blot. Loss of dFARS2 resulted in obvious increases in the steady-state levels of eight mitochondrial tRNAs (tRNA^Phe, tRNA^Lys, tRNA^Thr, tRNA^Leu(CUN), tRNA^Asp, tRNA^His, tRNA^Tyr, tRNA^Leu) analyzed (Figure 4C). For comparison, the levels of mitochondrial tRNAs were quantified by normalizing to the reference 5S rRNA. The mean values of tRNA^Phe, tRNA^Lys, tRNA^Thr, tRNA^Leu(CUN), tRNA^Asp, tRNA^His, tRNA^Tyr, tRNA^Leu in dFARS2KO mutant larvae were 200.9%, 166.3%, 236.6%, 194.0%, 145.1%, 137.4%, 187.9% and 193.5%, compared to those in control larvae (Figure 4E). Similarly, the mean value of tRNA^Phe in dFARS2KO, tRNA^Thr, tRNA^Leu(CUN), tRNA^Asp, tRNA^His, tRNA^Tyr, tRNA^Leu in Dau-Gal4 driven dFARS2 RNAi larvae were 251.3%, 210.1%, 185%, 200.2%, 161%, 135%, 146.1% and 170.8%, compared to those in control larvae (Figure 4D and F). Moreover, an increased level of mitochondrial tRNA^Phe (153.5%) was also observed in elav-Gal4 driven dFARS2 knockdown fly heads, compared to that in control fly heads (Figure 4D and F). The increase in the steady-state levels of mitochondrial tRNA in our dFARS2 deficiency models suggests a potential compensatory response of mitochondrial defects.

We next investigated the effect of dFARS2 deficiency on mitochondrial transcription. The protein level of mitochondrial transcription factor TFAM was slightly elevated in both dFARS2KO mutant and RNAi larvae (Supplementary Figure S4A). In addition, the levels of mtDNA-encoded transcripts, including ND4, CYTB, ATP6 and 16S rRNA were increased in both dFARS2KO mutant and RNAi larvae, suggesting an increase of overall mitochondrial transcription (Supplementary Figure S4B and C). Furthermore, we performed qPCR to examine the level of mitochondrial DNA. Both dFARS2KO mutant and RNAi knockdown larvae exhibited increased mtDNA levels, as shown for ND4, CYTB, ATP6 and 16S rRNA genomic DNA fragments (Supplementary Figure S4D and E). These results are consistent with the notion that there is a compensatory response in dFARS2 deficiency.

**Reductions in the levels of the subunits of OXPHOS complexes**

Having shown that dFARS2 is required for mitochondrial tRNA^Phe aminoacylation, we proceeded to test whether loss of dFARS2 has an effect on the levels of OXPHOS components. Western blot analysis showed that the levels of the mtDNA-encoded complex IV subunit (CO3) and the nuclear gene encoded complex III subunit (UQCRFS1), complex I subunits (NDUFS1 and NDUFS3), complex V (ATP5A) and were reduced in dFARS2KO mutant as well as dFARS2 RNAi larvae (Figure 5A). However, the level of complex II subunit (SDHA) was not altered in both dFARS2KO mutant and RNAi larvae (Figure 5A). The levels of NDUFS1, NDUFS3, SDHA, UQCRFS1, CO3 and ATP5A were 40.6%, 19.9%, 105%, 30.2%, 38.7% and 71.4% in the dFARS2KO larvae, 31.1%, 23%, 98%, 29%, 25.8% and 86.6% in the dFARS2 knockdown larvae, relative to the mean values measured in the control larvae, respectively (Figure 5B).
Defective assembly of OXPHOS complexes

We then examined the assembly of the individual OXPHOS complexes. Mitochondria were isolated from the second instar larvae and analyzed by blue native polyacrylamide gel electrophoresis (BN-PAGE) and Western blot. The levels of assembled OXPHOS complexes I and V were obviously decreased in both dFARS2KO mutant and dFARS2 RNAi larvae (Figure 6A). Consistent with the BN-PAGE results, Western blot analyses revealed that depletion of dFARS2 caused substantial reduction in OXPHOS complexes I, III, IV and V components, but not in complexes II components (Figure 6B). The levels of complex I, II, III, IV and V were 11.7%, 103.3%, 5.4%, 17.6% and 36.9% in the dFARS2KO larvae, 10.8%, 95%, 3.8%, 21.8% and 42% in the dFARS2 knockdown larvae, relative to the mean values measured in the control larvae, respectively (Figure 6C).

Altered activities of OXPHOS complexes

OXPHOS assembly defects can affect respiratory chain activity. Thus, we tested the effect of dFARS2 depletion on the oxidative phosphorylation using established biochemical assays to measure the activities of individual respiratory complexes. Complex I activity was measured as the rate of NADH oxidation in the presence of ubiquinone. The nucleus-encoded Complex II activity was determined using dichlorophenolindophenol (DCPIP) as an artificial electron acceptor and succinate as the substrate. Complex III activity was measured by the rate of cytochrome c reduction using dibuquinol-2 as an electron donor. Complex IV activity was quantified by the rate of cytochrome c oxidation. Complex V activity was determined as the rate of NADH oxidation using phosphoenolpyruvate as an electron acceptor. Our results showed that the enzyme activities of OXPHOS complexes I, III, IV and V were reduced in dFARS2KO mutant and dFARS2 RNAi larvae, whereas the complex II activity was unaffected (Figure 7A and B). For comparison, the levels of mitochondrial complex activity were normalized to the average content of citrate synthase (CS). The mean values of Complex I, II, III, IV and V activities in dFARS2KO mutant were 25.6%, 101.8%, 40.4%, 68.8% and 65.7%, relative to the mean values measured in control larvae, respectively (Figure 7A). Similarly, the mean values of complexes I, II, III, IV and V activities were 45.0%, 92.6%, 32.4%, 41.5% and 49.5% in dFARS2 RNAi larvae, compared to those in control larvae, respectively (Figure 7B).
Figure 4. The effects of dFARS2 deficiency on the aminoacylation and steady-state levels of tRNAs. (A, B) In vivo aminoacylation of mitochondrial tRNA assays. Fifty or one hundred μg of total RNAs purified from various larvae (A) the whole body of dFARS2 KO and WT second instar larvae; (B) the whole body of Da-Gal4 driven dFARS2 knockdown second instar larvae or the head of elav-Gal4 driven dFARS2 knockdown and control flies) under acid conditions were electrophoresed through an acid (pH 5.2) 9% polyacrylamide-7 M urea gel, electroblotted, and hybridized with DIG-labeled oligonucleotide probes specific for the tRNA^Phe, tRNA^Lys and tRNA^Thr, respectively. The charged (upper band) and uncharged (lower band) forms of different tRNAs were separated by the gel system. Samples were deacylated (DA) by heating for 10 min at 60°C at pH 9, electrophoresed, and hybridized with DIG-labeled oligonucleotide probes as described above. (C, D) Northern blot analysis of eight tRNAs. Ten μg of total RNA from various larvae were electrophoresed through a denaturing polyacrylamide gel, electroblotted, and hybridized with DIG-labeled oligonucleotide probes specific for tRNA^Phe, tRNA^Lys, tRNA^Thr, tRNA^Leu(CUN), tRNA^Asp, tRNA^His, tRNA^Tyr, tRNA^Ile and 5SrRNA, respectively. (E, F) Quantification of relative tRNA levels. The content of each tRNA was normalized to that of 5S rRNA. Calculations were based on three independent experiments. Error bars indicate two standard deviations (SD) of the means. * P < 0.05, ** P < 0.01, *** P < 0.001, **** P < 0.0001. NS, not significant.

Figure 5. Western blotting analysis of mitochondrial proteins. (A) Ten micrograms of total proteins from various larvae were electrophoresed through a denaturing polyacrylamide gel, electroblotted and hybridized with antibodies for 6 subunits of OXPHOS (mtDNA-encoded CO3, complex IV; nuclear-encoded UQRCFS1, complex III; NDUF53 and NDUF51, complex I; ATP5A, complex V; SDHA, complex II), and Porin as a loading control, respectively. (B) Quantification of relative mitochondrial protein levels. Average content of each polypeptide was normalized to the average content of Porin in each genotype. n = 3. Calculations were based on three independent experiments. Graph details and symbols are explained in the legend to Figure 4.
Figure 6. Defective assembly of OXPHOS complexes. (A) BN-PAGE analysis of mitochondrial respiratory chain complexes in mitochondrial protein extracts from larvae with different genotypes. Left: WT and dFARS2KO larvae; Right: control and Da-Gal4 driven dFARS2 knockdown second instar larvae. Positions of specific complexes are indicated. (B) Western blot analysis of mitochondrial proteins after BN-PAGE. Mitochondria extracted from various larvae were solubilized with 20% Triton X-100, electroblotted and hybridized with antibodies specific for subunits of OXPHOS, respectively, and with Porin as a loading control. Asterisk indicates unknown complexes. (C) Quantification of the levels of complexes I, II, III, IV and V in various larvae. The calculations were based on three independent determinations. Graph details and symbols are explained in the legend to Figure 4.

Characterization of two human FARS2 pathogenic variants in the dFARS2 mutant flies

To further utilize the Drosophila dFARS2 knockout mutant model and determine the functional consequence of disease variants identified in human patients, we performed rescue experiments through expression of either wild type or mutated human FARS2 into dFARS2KO mutant flies. We chose two human FARS2 variants, p.G309S and p.D142Y, for this analysis. p.G309S has been linked with the early-onset epileptic mitochondrial encephalopathy, whereas p.D142Y is associated with the less severe spastic paraplegia (27,29). We first generated transgenic flies with UAS-human wild type FARS2 and FARS2 bearing p.G309S or p.D142Y variant. To ensure the same expression levels, the constructs were subcloned into the same vector and inte-
grated into the same genomic landing site in the *Drosophila* genome. When ubiquitously expressed under the control of Da-Gal4, wild type and two mutated versions of human FARS2 were able to partially rescue the larval growth defects and lead to the eclosion of adult flies (Figure 8A). We noticed that while there was a slightly delay during pupal developmental for *dFARS2*KO mutants expressing human wild type FARS2, the developmental delay was much severer in *dFARS2*KO mutants carrying p.G309S or p.D142Y variant (Figure 8B). In order to provide further evidence for the disease-causing nature of the variants, we also tested for their seizure susceptibility in those surviving adult flies carrying wild type or mutated FARS2 transgenes. *dFARS2*KO mutant flies expressing wild type FARS2 showed a low level of bang sensitivity comparable to wild type controls (Figure 8C and D, supplementary video 2). Notably, there was an induced seizure phenotype in *dFARS2*KO mutants upon the expression of FARS2 p.G309S variant (Figure 8C and D, supplementary video 2). In contrast, there was no obvious seizure phenotype in *dFARS2*KO mutants carrying p.D142Y variant of human FARS2 (Figure 8C and D, supplementary video 2). Given that p.D142Y variant of human FARS2 was reported in patients with spastic paraplegia, we then performed the climbing assay to test whether these flies display locomotion defects. Flies were placed into an empty vial and tapped down to the bottom of the vial, and the number of flies able to pass an 8-cm mark successfully within 10 s was recorded to generate a climbing index. Indeed, *dFARS2*KO mutant flies carrying p.D142Y variant of human FARS2 showed a reduced climbing ability compared to controls (Figure 8E). A slightly reduced climbing ability was also observed for *dFARS2*KO mutant flies with wild type FARS2 or p.G309S variant (Figure 8E). These results suggest that our fly models could separate the disease phenotypes associated with two distinct mutations. Moreover, the efficiency of aminoacylated mt-tRNA^Phe^ in the *dFARS2*KO mutant larvae carrying wild type human FARS2 was comparable with that in wild type larvae (Figure 8F and G). Notably, *dFARS2*KO mutant larvae carrying p.G309S mutation reduced about 15%, while the levels of aminoacylation in *dFARS2*KO mutants carrying p.D142Y mutation was comparable with that in wild type larvae (Figure 8F and G). The steady levels of mitochondrial tRNA^Phe^ and tRNA^12s^ were not changed among these larvae (Figure 8H). The in-gel activity analysis revealed that introducing p.G309S or p.D142Y variant into *dFARS2*KO mutant background caused a slight reduction of Complex I activity, but did not affect Complex II activity (Figure 8I). These data provide evidences that human FARS2 p.G309S mutation causes developmental delay and seizure and p.D142Y mutation is associated with spastic paraplegia.

**DISCUSSION**

In the present study, we have generated *Drosophila* *dFARS2* knockout and knockdown models and analyzed disease related phenotypes. We found that *dFARS2* mutations or ubiquitously knocked down of *dFARS2* result in developmental delay and lethality at the second instar larval stages. Similar phenotypes have been reported in *Drosophila SARS2* and *MARS2* mutants (52,53). Furthermore, the lethality is also observed in mouse *DARS2* knockout (54). These findings in model organisms are consistent with the absence of patients reported so far with an allele combination of complete loss of function of mitochondrial aminoacyl-tRNA synthase, as such mutations might be incompatible with life.

Neurological disorder is one major phenotype in patients with various mitochondrial aminoacyl-tRNA synthetase mutations (55,56). However, a proper animal model for studying these neuronal defects was not available. We specifically knocked down *dFARS2* in the nervous system and observed that *dFARS2* reduction leads to a developmental delay and induces seizure. These results clearly suggest that mitochondrial tRNA aminoacylation and protein translation are crucial for development as well as the maintenance of neuronal functions. The mitochondria of *dFARS2* knockdown fly brains display abnormal morphology, which indicates their dysfunctions. The changes in mitochondrial morphology have previously been reported in flies depleted of SARS2 or SARS2-related gene (53,57). The phenotypes in these *dFARS2* knockdown flies mirror the defects reported in patients with FARS2 mutations, indicating the potential use of these models in dissecting the molecular mechanisms of FARS2 deficiency (29). Interestingly, previous reported *MARS2* knockout and *SARS2* knockdown fly models have shown some features related to neurological defects (52,53). However, the seizure behaviors of *MARS2* and *SARS2* models were not further examined (52,53). It might be interesting to see whether seizure induction is a more general phenotype associated with *ARS2* mutations in *Drosophila* in the future.

We have shown that mitochondrial tRNA^Phe^ aminoacylation activity was severely reduced in *dFARS2* mutant and knockdown larvae, indicating *dFARS2* is essential for aminoacylation of tRNA^Phe^ . Synthesis of cognate mitochondrial aminoacyl tRNAs by specific ARS2s is important for the fidelity in mitochondrial protein translation (8,29). In the absence of *dFARS2*, decreased charged tRNA^Phe^ can impair mitochondrial protein synthesis and reduce the efficiency of protein translation. Our western blot analysis revealed that the levels of both mitochondria and nuclear encoded OXPHOS components for complex I, III, IV and V were decreased in *dFARS2* deficiency larvae, indicating reduced protein synthesis and/or protein stability. Moreover, the ability of these proteins to assemble into the OXPHOS complexes was compromised in *dFARS2* deficiency, as demonstrated by the Blue native gel assay. The defective assembly of OXPHOS complexes further indicates that balanced production of nuclear and mitochondrial OXPHOS polypeptides is important for respiratory complex assembly. *dFARS2* deficiency specifically reduces the activity of OXPHOS complexes that contain mtDNA encoded subunits. However, the entirely nuclear encoded complex II activity was not affected. These differential effects on the OXPHOS complexes have been observed for mutations in several other mitochondrial genes. Interestingly, we found that the levels of a subset of mitochondrial mRNAs and tRNAs were increased in *dFARS2* mutants. Moreover, the mitochondrial DNA levels were also upregulated. This could be explained by a compensatory response to mitochondrial dysfunction (58,59). However, the increase in the steady-state levels of these mt-tRNAs was not able to over-
Figure 8. Functional analysis of two human FARS2 variants in dFARS2 mutants. (A) Images showing the larvae with different genotypes at 5 days AEL (left) and adult female flies at 15 days AEL (right). Scale bars represent 1 mm. (B) Graph showing pupariation curves for control, dFARS2KO larvae expressing human wild type FARS2, FARS2 p.G309S mutation or FARS2 p.D142Y mutation. n = 4. (C) Graph showing the percentage of female flies (control, KO + FARS2, KO + FARS2G309S or KO + FARS2D142Y) with BS paralytic phenotypes. n = 4. (D) Graph showing the recovery time of flies as described in (C) after BS paralysis. n = 7 for control; n = 8 for KO + FARS2; n = 17 for KO + FARS2G309S; n = 9 for KO + FARS2D142Y. (E) Graph showing the climbing index of flies as described in (C). n = 5. (F) In vivo aminoacylation of tRNA^Phe assay in control and dFARS2KO third instar larvae expressing human wild type FARS2 (KO + FARS2), FARS2 p.G309S mutation (KO + FARS2G309S) or FARS2 p.D142Y mutation (KO + FARS2D142Y). (G) Quantification of aminoacylation level of tRNA^Phe among larvae with different genotypes. (H) Northern blot analysis of tRNA^Phe and tRNA^Lys in various third instar larvae. (I) In-gel activity analysis from various third instar larvae. The left panel shows in gel activity of complex I, and the right panel shows in-gel activity of complex II. The calculations were based on 3–4 independent experiments. Graph details and symbols are explained in the legend to Figure 4.

come the loss of dFARS2, as the OXPHOS complexes were not assembled properly and resulted in compromised mitochondrial respiration. Together, these data demonstrate the biochemical characteristics observed in dFARS2 deficiency Drosophila larvae largely resemble the phenotypes in human cell lines carrying the mutations of FARS2 gene.

Moreover, an important finding of our study is that expression of human disease-causing variants in Drosophila dFARS2 mutants can partially recapitulate some features of the disease. Expression of human wild type FARS2 and FARS2 carrying the p.G309S or p.D142Y variant in the dFARS2 mutants could rescue the viability of the mutants. Compared with human wild type FARS2, expression of human FARS2 with p.G309S or p.D142Y variant in dFARS2 mutants leads to a severer developmental delay during the pupal stage. Furthermore, the adult dFARS2 mutant flies carrying human FARS2 with p.G309S variant display a seizure defect. In contrast, the adult dFARS2 mutant flies...
carrying human FARS2 with p.D142Y variant display a strong locomotion defect. These data together provide experimental evidence for the pathogenicity of the p.G309S and p.D142Y variants and also indicate the phenotypic difference between these two variants. Our results highlights the importance of the Drosophila in vivo model to verify the pathogenic effects of FARS2 variants with respect to the human disease.

In summary, our Drosophila dFARS2 knockout and knockdown models accurately recapitulate many phenotypic features of human disease caused by FARS2 mutations. These models will be valuable for identifying the underlying pathomechanism of ARS2 diseases and screening the suppressors of ARS2 mutations as well as drugs with a therapeutic potential.

DATA AVAILABILITY
The authors declare that [the/all other] data supporting the findings of this study are available within the article [and its supplementary information files].

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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